SUPERSEDED GUIDANCE - NEWER VERSION AVAILABLE

Guidance on the Biocidal Products Regulation

Volume II Efficacy - Assessment and Evaluation (Parts B+C)

Version 1.0
February 2017
LEGAL NOTICE

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Guidance on the BPR: Volume II Efficacy - Assessment and Evaluation (Parts B+C)

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PREFACE

The Guidance on the Biocidal Products Regulation (BPR) is to be applied to applications for active substance approval and product authorisation as submitted from 1 September 2013, the date of application (DoA) of the Biocidal Product Regulation (the BPR).

This document describes the BPR obligations and how to fulfil them.

The scientific guidance provides technical scientific advice on how to fulfil the information requirements set by the BPR (Part A), how to perform the risk assessment and the exposure assessment for the evaluation of the human health and environmental aspects and how to assess and evaluate the efficacy to establish the benefit arising from the use of biocidal products and that it is sufficiently effective (Parts B & C).

In addition to the BPR guidance, the Biocidal Products Directive (BPD) guidance and other related documents are still considered applicable for new submissions under the BPR in the areas where the BPR guidance is under preparation. Furthermore these documents are still valid in relation to the applications for active substance approval or applications for product authorisation under the BPD that may still be under evaluation. Also the Commission has addressed some of the obligations in further detail in the Biocides competent authorities meetings documents which applicants are advised to consult. Please see ECHA Biocides Guidance website for links to these documents: [https://echa.europa.eu/guidance-documents/guidance-on-biocides-legislation].

Applicability of Guidance


¹ Link available under Working Procedures (right column) [https://echa.europa.eu/about-us/who-we-are/biocidal-products-committee]
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NOTES to the reader:

In this document text cited from the Biocidal Products Regulation (EU) No 528/2012 is indicated in green boxes.

This symbol highlights text to be noted.

Section 5.4.5 PT5 Drinking water disinfectants: A preliminary draft text for PT 5 is included in this section: the section is currently under review within the ECHA “Disinfectants Project”. (see section for full details of note).

Section 5.6 and sub-sections for PT10, PT11, PT12, PT15, PT16, PT17, PT19 (non-arthropods) and PT20: please refer to the General sections 1-3 of this guidance and the TNsG.
# List of Abbreviations

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<td>AS</td>
<td>Active substance</td>
</tr>
<tr>
<td>BP</td>
<td>Biocidal product</td>
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<td>BPD</td>
<td>Biocidal Products Directive 98/8/EC</td>
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<tr>
<td>BS</td>
<td>British standard</td>
</tr>
</tbody>
</table>
| CA/CAs eCA   | Competent Authority/Competent Authorities  
|              | - Evaluating CA (eCA) is the Competent Authority that evaluates the application for an active substance approval or an application for a Union authorisation.  
<p>|              | - Receiving CA is the Competent Authority that receives an application for a National Authorisation. |
| CAR          | Competent Authority Report, (also known as the assessment report). |
| CEN          | Comité Européen de Normalisation; European Committee for Standardisation <a href="http://www.cen.eu/">http://www.cen.eu/</a> |
| CFU          | Colony forming units |
| CIP          | Cleaning-in-Place |
| CT           | Concentration x Time |
| CV           | Critical value |
| DIN          | Deutsches Institut fuer Normung; German national organisation for standardisation <a href="http://www.din.de/">http://www.din.de/</a> |
| DVG          | Deutsche Veterinaermedizinische Gesellschaft; German Veterinary Medical Society <a href="http://www.dvg.net/">http://www.dvg.net/</a> |
| EN           | European Standard |
| EPPO         | European and Mediterranean Plant Protection Organization <a href="http://www.eppo.org">www.eppo.org</a> |</p>
<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td>ESL</td>
<td>Estimated service life</td>
</tr>
<tr>
<td>EU</td>
<td>European Union + Norway, Iceland and Lichtenstein</td>
</tr>
<tr>
<td></td>
<td>Please note the BPR applies to the European Economic Area (EEA) and thus all references to the EU in the text should be understood as EEA (EU + Norway, Iceland and Lichtenstein)</td>
</tr>
<tr>
<td>GLP</td>
<td>Good laboratory practice</td>
</tr>
<tr>
<td>ISO</td>
<td>International Organization for Standardisation <a href="http://www.iso.org/">http://www.iso.org/</a></td>
</tr>
<tr>
<td>KD</td>
<td>Knock down</td>
</tr>
<tr>
<td>KD$_{50}$</td>
<td>Knock down for 50% of the group of tested animals</td>
</tr>
<tr>
<td>KT$_{50}$</td>
<td>Knock down time for 50% of the group of tested animals</td>
</tr>
<tr>
<td>LD$_{50}$</td>
<td>Lethal dose for 50% of the group of tested animals</td>
</tr>
<tr>
<td>MAD</td>
<td>Mutual acceptance of data</td>
</tr>
<tr>
<td>prEN</td>
<td>Draft European Standard</td>
</tr>
<tr>
<td>PAR</td>
<td>Provisional assessment report</td>
</tr>
<tr>
<td>PEG</td>
<td>Partner expert group</td>
</tr>
<tr>
<td>PT</td>
<td>Product-type</td>
</tr>
<tr>
<td>SPC</td>
<td>Summary of Product Characteristics</td>
</tr>
<tr>
<td>TC</td>
<td>Technical Committee</td>
</tr>
<tr>
<td>TM</td>
<td>Technical Meeting</td>
</tr>
<tr>
<td>TNsG</td>
<td>Technical Notes for Guidance</td>
</tr>
<tr>
<td>TVC</td>
<td>Total viable count</td>
</tr>
<tr>
<td>UC</td>
<td>Use Class</td>
</tr>
<tr>
<td>US-EPA</td>
<td>United States Environmental Protection Agency <a href="http://www.epa.gov/">http://www.epa.gov/</a></td>
</tr>
<tr>
<td>VAH</td>
<td>Verbund fuer Angewandte Hygiene; Association for Applied Hygiene <a href="http://www.vah-online.de/">http://www.vah-online.de/</a></td>
</tr>
<tr>
<td>VOC</td>
<td>Volatile organic compound</td>
</tr>
</tbody>
</table>
# Glossary of Terms

<table>
<thead>
<tr>
<th>Standard term</th>
<th>Explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Activity against enveloped viruses (see also Virucidal activity and Limited spectrum virucidal activity)</td>
<td>A claim for hygienic hand and skin disinfectants with activity against enveloped viruses only.</td>
</tr>
<tr>
<td>Algaecide</td>
<td>A product or active substance used to control (inhibit the growth) or kill algae.</td>
</tr>
<tr>
<td>Algaecidal activity</td>
<td>The capability of a product or active substance to produce a reduction in the number of viable algae cells under defined conditions.</td>
</tr>
<tr>
<td>Antimicrobial product</td>
<td>A product which prevents the growth of/reduces the number of/mitigates the growth of micro-organisms</td>
</tr>
<tr>
<td>Bactericide</td>
<td>A product or active substance which irreversibly inactivates vegetative bacteria under defined conditions</td>
</tr>
<tr>
<td>Bactericidal activity</td>
<td>The capability of a product or active substance to produce a reduction in the number of viable bacterial cells of relevant test-organisms under defined conditions</td>
</tr>
<tr>
<td>Bacteriostatic activity</td>
<td>Capability of a product or active substance to inhibit the growth of bacteria under defined conditions</td>
</tr>
<tr>
<td>Biocidal product/Biocide</td>
<td>BPR Article 3(1)(a): — any substance or mixture, in the form in which it is supplied to the user, consisting of, containing or generating one or more active substances, with the intention of destroying, deterring, rendering harmless, preventing the action of, or otherwise exerting a controlling effect on, any harmful organism by any means other than mere physical or mechanical action, — any substance or mixture, generated from substances or mixtures which do not themselves fall under the first indent, to be used with the intention of destroying, deterring, rendering harmless, preventing the action of, or otherwise exerting a controlling effect on, any harmful organism by any means other than mere physical or mechanical action. A treated article that has a primary biocidal function shall be considered a biocidal product.</td>
</tr>
<tr>
<td>Biofilm</td>
<td>An accumulation of microbial cells immobilised on a substratum and embedded in an organic polymer matrix of microbial origin</td>
</tr>
<tr>
<td>Biostatic product</td>
<td>A product which inhibits the growth of micro-organisms under defined conditions</td>
</tr>
<tr>
<td>Curative effect on biofilm</td>
<td>The biocide is added after the biofilm is formed and acts on biofilm stability, facilitating the biocide interaction with cells – it may or may not act as detergent and detach the biofilm from the surface</td>
</tr>
<tr>
<td>Standard term</td>
<td>Explanation</td>
</tr>
<tr>
<td>---------------------------------------------------</td>
<td>-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Disinfectant within PT 2, 3, 4 and 5</td>
<td>A disinfectant is a product that reduces the number of micro-organisms in or on an inanimate matrix- achieved by the irreversible action of a product, to a level judged to be appropriate for a defined purpose</td>
</tr>
<tr>
<td>Disinfection within PT 2, 3, 4 and 5</td>
<td>Disinfection is the reduction of the number of micro-organisms in or on an inanimate matrix- achieved by the irreversible action of a product, to a level judged to be appropriate for a defined purpose</td>
</tr>
<tr>
<td>Skin disinfection within PT1</td>
<td>Skin disinfection is the reduction of the number of micro-organisms on skin, achieved by the irreversible action of a product, to a level judged to be appropriate for a defined purpose</td>
</tr>
<tr>
<td>Efficacy</td>
<td>The ability of a product or active substance to produce an effect as described in the label claims made for it, when used under actual use conditions.</td>
</tr>
<tr>
<td>Flow condition (for biofilm)</td>
<td>Biofilm is formed on supports of different nature placed along a tube or a chamber where the medium (inoculated and/or fresh) is circulated in a closed (reservoir-pump-tubing) or open (reservoir-pump-tubing-outlet) system</td>
</tr>
<tr>
<td>Fungicide</td>
<td>A product or active substance which irreversibly inactivates fungi (vegetative mycelia, budding yeasts and/or their spores) under defined conditions</td>
</tr>
<tr>
<td>Fungicidal Activity</td>
<td>The capability of a product or active substance to produce a reduction in the number of viable vegetative yeast cells and mould spores of relevant test organisms under defined conditions</td>
</tr>
<tr>
<td>Fungistatic activity</td>
<td>The capability of a product or active substance to inhibit the growth of fungi under defined conditions</td>
</tr>
<tr>
<td>Hygienic hand disinfectants</td>
<td>A hygienic hand disinfectant is a hygienic handrub disinfectant or a hygienic handwash disinfectant</td>
</tr>
<tr>
<td>Hygienic handrub disinfectant</td>
<td>Product used for post-contamination treatment that involves rubbing hands, without the addition of water, which is directed against transiently contaminating microorganisms to prevent their transmission regardless of the resident skin flora</td>
</tr>
<tr>
<td>Hygienic handwash disinfectant</td>
<td>Product used for post-contamination treatment that involves washing hands with water, which is directed against transiently contaminating microorganisms to prevent their transmission regardless of the resident skin flora</td>
</tr>
<tr>
<td>Limited spectrum virucidal activity</td>
<td>Limited spectrum virucidal activity is a claim for hygienic hand and skin disinfectants using Adenovirus and Murine Norovirus as test organisms, thus including activity against the test viruses and all enveloped viruses (see Appendix 5).</td>
</tr>
<tr>
<td>Log reduction / log₁₀ reduction / lg reduction</td>
<td>Reduction presented in a logarithmic scale. Example 1: when a disinfection reduces 10⁸ bacteria to 10² bacteria, this is a lg reduction of 6. Example 2: when a disinfection reduces 5.10⁷ fungal spores to 8.10³ fungal spores this is a lg reduction of 3.79.</td>
</tr>
<tr>
<td>Standard term</td>
<td>Explanation</td>
</tr>
<tr>
<td>---------------</td>
<td>-------------</td>
</tr>
<tr>
<td>Microbes/micro-organisms</td>
<td>bacteria (including vegetative cells bacterial spores and mycobacteria) fungi (including yeasts, moulds and fungal spores) algae, viruses (including bacteriophages), protozoa (including cysts and other permanent states), etc.</td>
</tr>
<tr>
<td>Mycobactericide</td>
<td>A product or active substance which irreversibly inactivates mycobacteria under defined conditions</td>
</tr>
<tr>
<td>Mycobactericidal activity</td>
<td>The capability of a product or active substance to produce a reduction in the number of viable mycobacterial cells of relevant test organisms under defined conditions</td>
</tr>
<tr>
<td>Neutraliser</td>
<td>A chemical agent or formulation which suppresses the residual activity of an disinfectant within a test but does not inhibit or inactivate micro-organisms</td>
</tr>
<tr>
<td>Performance standard</td>
<td>Regulatory or scientific standard for biocides that is either quantitative or qualitative (that may also be specified in the test method) by which a decision is taken on the acceptability of a claim.</td>
</tr>
<tr>
<td>Preventive effect on biofilm</td>
<td>The biocide is present before the biofilm is formed and may act both on cell viability and/or on cell adhesion/biofilm maturation</td>
</tr>
<tr>
<td>Product type (PT)</td>
<td>Product types (PT) are defined in BPR annex V</td>
</tr>
<tr>
<td>Sporicide</td>
<td>A product or active substance which inactivates dormant bacterial spores under defined conditions</td>
</tr>
<tr>
<td>Sporicidal activity</td>
<td>The capability of a product or active substance to produce a reduction in the number of viable bacterial spores of relevant test organisms under defined conditions</td>
</tr>
<tr>
<td>Sporistatic activity</td>
<td>The capability of a product to inhibit the germination of dormant bacterial spores under defined conditions</td>
</tr>
<tr>
<td>Static condition (for biofilm)</td>
<td>Biofilm is formed on supports such as microplates without agitation after an incubation time that depends on the micro-organism considered</td>
</tr>
<tr>
<td>Surgical hand disinfectants</td>
<td>A surgical hand disinfectant is a surgical handrub disinfectant or a surgical handwash disinfectant</td>
</tr>
<tr>
<td>Surgical handrub disinfectant</td>
<td>Product used for preoperative treatment that involves rubbing hands, without the addition of water, which is directed against the flora of microorganisms on hands to prevent the transmission of microorganisms into the surgical wound</td>
</tr>
<tr>
<td>Surgical handwash disinfectant</td>
<td>Product used for preoperative treatment that involves washing hands with water, which is directed against the flora of microorganisms on hands to prevent the transmission of microorganisms into the surgical wound</td>
</tr>
<tr>
<td>Treated article</td>
<td>A treated article is any substance, mixture or article which has been treated with, or intentionally incorporates, one or more biocidal products</td>
</tr>
<tr>
<td>Tuberculocide</td>
<td>A product or active substance which irreversibly inactivates Mycobacterium tuberculosis under defined conditions</td>
</tr>
<tr>
<td>Standard term</td>
<td>Explanation</td>
</tr>
<tr>
<td>-------------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Tuberculocidal activity</td>
<td>The capability of a product or active substance to irreversibly inactivate <em>Mycobacterium tuberculosis</em>, demonstrated by the capability to produce a reduction in the number of viable cells of the test organism <em>Mycobacterium terrae</em> under defined conditions</td>
</tr>
<tr>
<td>Virucide</td>
<td>A product or active substance which irreversibly inactivates viruses under defined conditions</td>
</tr>
</tbody>
</table>
| Virucidal activity (see also Limited spectrum virucidal activity + Activity against enveloped viruses) | The capability of a product or active substance to produce a reduction in the number of infectious virus particles of relevant test organisms under defined conditions  
“Full spectrum” virucidal activity is a claim for biocidal products using relevant test organisms and thus showing activity against the enveloped and non-enveloped viruses. |
| Yeasticide                    | A product or active substance which irreversibly inactivates yeast under defined conditions                                                                                                                |
| Yeasticidal activity          | The capability of a product or active substance to produce a reduction in the number of viable vegetative yeast cells of relevant test organisms under defined conditions                                                 |
1. General Introduction

Evaluation and Assessment

The process of evaluation of active substance applications is given in Article 8 (BPR) and the common principles for the evaluation of dossiers for biocidal products (including the representative biocidal product in the context of the active substance approval) is given in Annex VI (BPR).

The evaluating or receiving CA uses the data submitted in support of an application for active substance approval or authorisation of a biocidal product to make a risk assessment based on the proposed use of the (representative) biocidal product. The general principles of assessment are given in Annex VI (BPR) and the evaluation is carried out according to these general principles. The evaluating body will base its conclusions on the outcome of the evaluation and decide whether or not the biocidal (representative) product complies with the criteria for authorisation set down in Article 19(1)(b) and/or whether the active substance may be approved.

Efficacy data are a fundamental component in the regulatory management and decision making process for biocidal products. Efficacy data are required to establish the benefit arising from the use of biocidal products and must be balanced against the risks their use poses to man and the environment.

Authorisation of a biocidal product will only be granted according to Art. 19 (1) b of the BPR if that product is shown to be sufficiently effective.

Even for the requirement to limit the use to the minimum necessary and the general requirement of sustainable use of biocidal products (Art. 17 and 18 BPR), it is crucial that the biocide in questions delivers the expected effect.

The information and data required relevant to the effectiveness of the active substance(s) to be employed in biocidal products are outlined in Annex II, BPR, title 1 No. 6 and 7 and title 2 No 5 and 6. For biocidal products the data required are set out in Annex III, Title 1 No 6 and 7, and title 2, No 6 and 7.

These general sections at the beginning of this guidance, (namely sections 1, 2 and 3), provide a general overview for the efficacy evaluation; the more specific requirements for each Product Type (PT), which must be met and should be followed in the first instance, are described in the later sections.

2. Claims

2.1 Introduction

The evaluation of the efficacy of a biocidal product is carried out in order to determine whether the claims made for the activity of the active substance (within the product) or the product itself, are supported by suitable efficacy data. A claim is the precondition and base for efficacy testing.

Claims should comprise of the description of the problem and the way it is suggested to be solved by the biocidal treatment. Claims include information given in an active substance dossier, information on the label of a product, information provided on a web-site or in product-associated leaflets. All claims should be consistent.
Claims can range from simple to complex, depending on the activity and benefits the applicant wishes to claim as resulting from the use of the active substance/biocidal product. This should include as a minimum the following information:

- The purpose of the claim (e.g. prevent destruction of material by insect infestations, disinfect surface);
- The function of the product (e.g. insecticide, wood preservative, disinfectant, etc.);
- The (group of) target organisms which will be controlled;
- In-use concentration;
- Use conditions and area of use;
- The effect which will result from using the product on the target organisms (e.g. kill, control, repel, prevent, etc.);
- Any products, organisms or objects to be protected.

Some examples are available in the different claim matrices and PT specific guidance sections (see later sections).

However this basic information can be supplemented by additional claims which further describe the effects of the active substance/product where appropriate, such as:

- How fast the effect is produced;
- The duration of the effect (residuality) or lifespan;
- The types of surface on which the product can be used (e.g. hard porous and non-porous surfaces, softwood).

For products used to treat articles, additional information should be provided:

- Durability of the effect in relation to the expected life-span of the treated article;
- Resilience towards ageing, weathering or other use conditions as for instance washing;
- Where relevant, leaching/migration data for different materials or different use conditions.

All claims made should be supported by data or a suitably robust scientifically based reasoned case.

### 2.2 Label claims and directions for use

The directions for use and the claims made for the biocidal product are included in a summary of biocidal product characteristics (SPC) in accordance with Article 22(2) (BPR).

A label claim is information which is provided to the user which describes the biocidal effects that will result from using a biocidal product under its normal conditions of use (e.g. when it is used at the recommended dose/application rate, by the recommended application method(s) and in the appropriate areas, etc.). The product label can only include claims that are in line with the authorised uses, as given in the SPC.

Label claims should be as specific as possible, or if more general claims (such as “fast acting”) are made, then they should be further clarified on the label where possible (e.g. “fast acting – acts within 5 minutes”). If no clarification is provided, the evaluating Competent Authority should ask the applicant to specify the claim. A judgement as to what a normal user would reasonably expect from the claim should be made. Evaluation should be made according to this claim and the directions for use should be taken into account.
An application for a product authorisation must include a draft SPC and additionally should include a copy of the draft product label containing the claims made for the product.

Applications for product families should include the entire range of the claims proposed for the products within the family.

### 3. General considerations for the development and reporting of efficacy data

#### 3.1 Efficacy

Efficacy is defined as the ability of a product to fulfil the claims made for it when used according to the directions for use on the proposed product label (as given in the SPC): Is the product actually sufficiently effective against the claimed organisms under the conditions specified? The applicant must provide sufficient information to clearly specify the field of use of the product. In addition, studies must be provided to demonstrate that the product, when used in accordance with the use instructions (concentration, application method, etc.), is sufficiently effective.

#### 3.1.1 Efficacy tests

The applicant must submit studies which clearly demonstrate the efficacy of the active substance/product.

We distinguish various types of studies:

- Screening tests
- Laboratory studies
- Simulation tests in laboratory
- Field tests

**Screening tests** are usually not related to practical/field conditions and are often not implemented with the complete product but only with the active substance. Such tests are therefore primarily useful for providing supplementary information, for example to demonstrate that the concentration used is optimal.

**Laboratory studies** are performed to validate the efficacy in a laboratory according to criteria defined. These tests permit to validate for example a level of mortality during a given time, a knock down (KD) effect and if need be the palatability of the product.

**Simulation tests** are more linked to practical/field conditions and can, in some cases, be sufficient for demonstrating the efficacy. Simulation tests can include factors like ageing, weathering, UV, washing, etc. Example: For disinfecting products aimed at controlling bacteria on hard surfaces, it is sufficient to carry out a suspension test and a surface test in accordance with the relevant EN standards.

**Field tests** provide a good indication of how the product works in practice/under field conditions, to evaluate how the efficacy can be affected by a variety of factors (the weather, population density, natural fluctuation of the population over time etc.). The experimental setup is important in these tests. The results of the tests should be compared to the results achieved with a control object which has not been treated or with the situation prior to treatment: however, in some cases it is not possible to include a control sample in field tests.
Screening tests, laboratory studies and simulation tests must always include an untreated control without active substance (i.e. a negative control); it is preferred that this is the formulated product without active substance. However, providing it can be justified, this can be, a control with only the solvent, e.g. water. There are few exceptions to this rule, such as the EN disinfection test, and all exceptions should be justified by the methodology.

Tests should preferably be carried out in accordance with standard protocols, e.g. CEN, ISO, OECD, ASTM, etc. If standard protocols are not available or are not suitable for the field of use concerned, other methods may also be used on condition that the studies concerned have a sound scientific basis. Preferably, available standard methods should be modified to meet the actual application in such cases. Ideally, tests are carried out in accordance with Good Laboratory Practice (GLP) or similar quality assurance systems (ISO), although this is not mandatory for efficacy tests.

3.1.2 Test report

Some standard tests (e.g. EN tests) contain examples of appropriate reports, which should be used as a template. In all other cases the test report must contain the following elements:

- introduction
- materials and methods (e.g. tested product composition, conditions of the test temperature, humidity,)
- tested organisms
- results and raw data
- conclusion/discussion based on criteria defined in guidance

The introduction must indicate the goal of the test. When a standard test is used the name and/or number of the test should be stated. The section on materials and methods must provide a complete description of the test method. If an internationally recognised standard method is used, it is sufficient to provide a brief description of the test. The product used and the concentration of the active substance must be specified. If the name of the product tested is not the same as the product for which the application is being submitted (e.g. a name used outside the EU or an internal company code for the product), the complete composition of the product tested must be provided in a separate document. The test organisms used must correspond to the organisms against which the product is intended to be used, or they must be adequate representatives. For example, if a product is intended for use against bacteria in hospitals, it is not possible to test the product on all possible species of bacteria. Instead, four standard species of bacteria are usually tested. The conditions under which the negative control tests were carried out must also be described (e.g. treated with product not containing the active substance, not treated, or treated with water for example).

The materials and methods should be described well. In case of standard test protocols all the deviations should be indicated and justified.

The section on the results of the test must provide quantitative data. It is not sufficient to present only tables or figures in which the results have been processed. The raw data must also be included. In case of repetitions performed in the test, the results should also be subjected to a statistical analysis, when appropriate. At the end of the report, a conclusion must be presented. Sometimes, it is necessary to discuss and/or present further arguments for the conclusion. For field tests in particular, the results obtained in repeated tests may
differ. If an explanation is provided for such differences in results, a test may possibly still be approved.

Example: In test 1, the product was “washed away by rainfall” and was therefore not effective, but tests 2 and 3 do demonstrate the efficacy. In such case the tests can be accepted and a remark will be made on the SPC that the product should not be used when rain is expected within x hours, because this will influence the efficacy negatively.

When applying for authorisation all the efficacy tests should be summarised in the PAR. The PAR format includes a table. This table should be filled out in a way that it gives an overview of all the efficacy results. When the test is not a standard test a short description of the method should be included. The test column “test system/concentration applied/ exposure time” should include all the relevant information on the test, the test parameter (e.g. contact time, temperature, replicates) in way that it can be compared to the intended use. The results should be specified (e.g. x% mortality, log reduction >x) and not just “test passed”. In some cases it might be easier to summarise the results in the text instead of the table (e.g. field trials).

Below the table the tests should be discussed and an explanation should be given on how the test results demonstrated the efficacy of the product for the different uses under use conditions.

3.2 Resistance

The topic of resistance is discussed in the general part of the TNsG on Product Evaluation (Section 6). Information on resistance should be given for active substances and biocidal products. Additionally, in support of the review for each active substance, information on resistance is given in the Competent Authority Report (CAR) of this active substance.

Resistance will be assessed on the basis of expert judgement. This section of the guidance will be updated in the future in the light of experience gained in evaluation of resistance.

4. Active substance approval

4.1 Introduction

According to Article 4 of the BPR, an active substance must be approved if at least one biocidal product containing that active substance may be expected to meet the criteria laid down in point (b) of article 19(1), and more particularly for the context of this guidance the paragraph (i), which says “the biocidal product is sufficiently effective”.

During the review of an active substance at the active substance approval stage, both the efficacy of the active substance and of the representative biocidal product are assessed in a relevant matrix. At this approval stage, it is the activity of the active substance which must be demonstrated, both in its own right and when formulated into a biocidal product.

Although a biocidal product containing the active substance is evaluated at the active substance approval stage, this part of the BPR process is concerned primarily with the efficacy of the active substance itself. The purpose of this section is to provide guidance for applicants and competent authorities on the principles for evaluation of efficacy at the active substance approval stage, and to help determine whether the information provided in an application for approval of an active substance is sufficient for inclusion of the substance in the Union list. For guidance on data requirement see Volume II Part A of ECHA’s guidance under the BPR.
4.2 General principles

4.2.1 Intended use

When making an application for approval of an active substance, the applicant must clearly describe the uses for which the active substance is intended. This information is required to allow a proper evaluation of the efficacy to be carried out, and must include, for every product type separately:

- The purpose of the claim (e.g. prevent destruction of material by insect infestations, decrease risk of infection by bacterial contamination);
- The function of the active substance (e.g. bactericide, fungicide, rodenticide, insecticide);
- The (group of) target organism(s) to be controlled;
- The effects on representative target organism(s) (e.g. attracting, killing, inhibiting);
- Any products, organisms or objects to be protected.
- The likely concentration at which the active substance will be used in products and, where appropriate, in treated articles. This likely concentration should be demonstrated to be effective according to the requirements described in section 4.2.2.1.

In the application, the applicant may choose to provide information on all of the intended target organisms at the active substance approval stage, or a representative selection. However, in order for approval of the active substance to be granted, efficacy must be demonstrated for at least one main target organism (or group of target organisms e.g. bacteria). Use against additional target organisms may be applied for at the product authorisation stage.

For active substances used in treated articles, see section 4.5 and sub-sections 4.5.2 and 4.5.3.

4.2.2 Evaluation of efficacy

Efficacy of an active substance has to be demonstrated both in part A of the CAR (related to the intrinsic efficacy of the active substance) and in part B (where the active substance is incorporated in a formulated product). Evaluation of each part is described below.

4.2.2.1 Active substance efficacy (part A):

As the testing of an active substance is normally carried out using the technical active substance, or a simple dilution of the active substance in water or an appropriate matrix (so that the testing is carried out in the absence of other substances which may affect the efficacy), an extensive data package and evaluation is not required at this stage.

However, efficacy studies should be submitted on the active substance, and these data should be capable of demonstrating the innate activity of the active substance against representatives of the proposed target organisms at the concentration relevant for the risk assessment. For that purpose, innate activity of an active substance could be defined as the capacity of an active substance to provide a sufficient effect on one or several relevant target organisms, for the use considered.

The following minimum requirements should be fulfilled to demonstrate innate activity:
• For main group 1 (disinfectants: PT1, 2, 3, 4 and 5), innate activity is at least a “cidal” activity demonstrated in a suspension test and has to be demonstrated against one or more representative target organism(s) for the activity claimed (e.g. bactericide, yeasticide), preferably according to the CEN norms (phase 1 tests and phase 2 step 1 tests). Test organism(s) should be that or those specified in the respective norm. Phase 1 tests are sufficient for the active substance if a phase 2 step 1 test is available for the representative product. When only specific biostatic activity (e.g. bacteriostatic, fungistatic) is claimed, an appropriate method should be used.

• For main group 2 (preservatives: PT6, 7, 8, 9, 10, 11, 12 and 13), innate activity is generally a static activity demonstrated in challenge tests on several and relevant target organisms, in the relevant matrix. However, if curative effects are claimed, cidal activity is requested. To demonstrate efficacy against one target organism only could also be acceptable in the case of a strictly defined use relevant for the PT (e.g. the control of Legionella in cooling water in PT11). For PT8, CEN norms are available to support efficacy testing and give indications on representative target organisms to be tested. Growth in the untreated control is essential to show the validity of the test. If the claim is only for a curative effect, it is sufficient to show that the decline in the microbial population in the treated samples is statistically significantly more than in the untreated control samples.

• For main group 3 (pest control: PT14, 15, 16, 17, 18, 19 and 20), innate activity can be demonstrated for one target organism only (for instance, control of mice or control of bedbugs).

• For main group 4 (other biocidal products: PT21 and PT22), innate activity is generally supported on a group of organisms (algae, animals, bacteria) and examples of appropriate target organisms are available in the Efficacy guidance for PT21 and PT22.

When minimum requirements are not met this should be justified.

Generally, efficacy data are generated from laboratory tests, performed by the applicant. Nevertheless efficacy data from literature could also be acceptable if the application rate, target organisms, area of use and the identity of the active substance is described and are relevant. If cited literature is used to support a preserving effect it must also show that untreated test specimens supported growth. When curative effects are claimed the cited literature must demonstrate the efficacy of the active substance according to the requirements per PT. The use of cited literature should be agreed between the applicant and the evaluation CA (eCA) on a case by case basis.

The level of efficacy demonstrated at this stage of the process need not be high, as an active substance in a simple solution may not be as effective as when it is used in a fully formulated product. For that reason an active substance should still be considered suitable for approval if the levels of efficacy demonstrated fulfil the minimum requirements above. In the case where the levels of efficacy of the active substance alone are lower than expected, efficacy tests performed with the representative product has to show a sufficient/basic efficacy, according to the requirements above. If both are insufficient, approval for the Union list should not be proposed.

If no efficacy tests with the active substance itself are available, but only tests with a formulation, a justification has to be given by the applicant regarding the possible influence of co-formulants on the efficacy. If the co-formulants used potentially have biocidal activity,
it is essential to demonstrate that the efficacy is due to the active substance and not to the co-formulants, e.g. a control should be performed with all co-formulants but without the active substance.

4.2.2.2 Product efficacy (part B):

Although approval for the Union list is primarily concerned with the active substance, efficacy data is also required for a representative product. Ideally efficacy data on an existing biocidal product should be submitted. If this is not possible data on a dummy product could be acceptable in order to demonstrate that the active substance is capable of producing an effect on the target organism and in a relevant matrix according to the proposed use, when included in a formulated product.

However, a detailed evaluation of the effectiveness of the product (including an evaluation of the proposed label claims) is not in all cases required at the active substance approval stage. This may for example be the case where no marketed product is available.

Nevertheless, the level of efficacy (e.g. the kind of activity “biocidal” or “biostatic”) have to be consistent with the uses claimed and fulfil the minimum requirements mentioned in the active substance part (part A).

4.2.3 Overall evaluation for active substance approval

It is concluded that efficacy data are required on the active substance, to demonstrate on the one hand the innate activity of the substance (either the technical grade active substance or a dilution in water or a solvent) and on the other hand the efficacy of the representative product against one or more of the proposed target organisms. Efficacy should be demonstrated in accordance with the use(s) considered in the risk assessment. If for some justified reasons, the results of the biocidal product do not completely fulfil the requirements described above, this could still be acceptable as long as the results of the active substance are sufficient to demonstrate efficacy. The other way around, if the results of the active substance do not fulfil the requirements described above acceptable data of the biocidal product may be sufficient as long as it can be excluded that the co-formulants contribute to the efficacy of the product.

Where the levels of efficacy demonstrated are low enough to raise concerns by the evaluating Member State, the applicant should be asked to justify why the result should still be considered acceptable. Two specific reasons are discussed below: the use of ‘dummy products’ and the case of active substances not used alone but always in combination with other active substances.

4.2.4 Link to risk assessment

There is an essential link between efficacy testing and the risk assessment for human health and the environment at the active substance approval stage:

- Efficacy has to be proven for active substance concentrations used in the risk assessment
- Efficacy has to be sufficient for the use assessed in the risk assessment.

The information on efficacy is relevant in assessing the dose recommended for the use(s) applied for. The dose (or the "likely concentration(s) at which the active substance will be used" as stated in Annex II 6.4 of the BPR) is the starting point in the exposure assessment for human health and the environment.
4.3 Active substances which are not intended to be used in isolation

This section is developed to deal with active substances which are not intended to be used as the sole active substance in a product.

At the active substance approval stage, the following should be demonstrated:

- **in part A** (dedicated to the active substance), the innate activity of the active substance should be demonstrated against target organism(s) relevant for the field of use envisaged. The evaluation should demonstrate that the active substance is capable of producing an effect on its own or when formulated into a very simple product. Due to the absence of the other active substance(s), the formulation may have only a limited, rather than broad based, spectrum of activity, or a lower level of efficacy.

Evaluation of the data will be done on a case by case basis.

Some examples where limited efficacy could be acceptable:

- for wood preservatives with fungicidal activity where different fungicides are active against different groups of target fungi and therefore two or more fungicides would be included in a product to produce the full spectrum of antifungal activity;
- for insecticides that are used in combination with other active substances to improve the insecticidal performance of the latter as they exert a synergistic effect;
- for insecticides used in combination with a co-formulants (e.g. booster) that is not itself an active substance;
- the active substance is used in combination with another active substance.

However, an appropriate argumentation is always required in order to justify situations with a more restricted level of efficacy. The minimum requirements in section 4.2 have always to be fulfilled.

- **in part B** (dedicated to the accompanying/representative product), the efficacy of a product where the active substance is formulated in combination with other (active) substances should be demonstrated against target organism(s) relevant for the field of use envisaged. Relevant efficacy tests should be used and structured to allow evaluation of the contribution of the active substance to the overall efficacy. This is particularly important if efficacy data have not been submitted in part A.

Efficacy data packages for formulations containing two or more active substances are not fully suitable for determining the activity contribution from the active substance under evaluation. For that reason great attention should be paid to justify the contribution of the active substance under evaluation to the total efficacy of the product. Information about the mode of action/function of the other active substances present in the product is also requested.

The submitted data should allow the definition of an effective concentration (i.e. the concentration of active substance at the efficient application rate of the product) that can be used for the risk assessment (specified per use). If in part B a formulation is introduced with additional co-active substances, this formulation will only be considered for efficacy testing and for setting a likely in-use concentration of the active substance, not used in isolation.
A statement should be added in the BPC opinion in order to stress that the active
substance is intended to be used in combination with other active substances or
synergists.

4.4 “Dummy products”
A “dummy product” is a product that is not fully formulated. It is not intended to be placed
on the market.

In order to satisfy the requirement of the BPR, a dossier of an active substance for inclusion
in the Union list (or in Annex I of active substances referred to in Article 25a of the BPR)
may be accompanied by such a product as the associated biocidal product. To the extent
possible, data from real products are nevertheless recommended.

While some dummy products may be very similar to a fully formulated product, others may
be a very simple formulation that bears little resemblance to the product which will finally be
placed on the market. The latter may be used where the applicant has limited experience in
formulating products, for example by applicants who only manufacture active substances.

At the active substance approval stage, the following should be demonstrated:
The evaluation should demonstrate that the active substance under evaluation is capable of
producing an effect when formulated into a very simple product (active substance alone or
diluted in a solvent) and to define an application rate, which is consistent with the intended
use(s) claimed by the applicant, and that can be used for the exposure assessment.

If a dummy product is used, a more restricted level of efficacy could be acceptable if an
appropriate and detailed justification is given by the applicant. However, the minimum
requirements mentioned in section 4.2 have always to be fulfilled.

4.5 Active substances used in treated materials and treated articles
Treated articles have been included into the biocides legislation on 1 September 2013 with
the BPR (Biocidal Products Regulation). This requires different considerations and testing
approaches as compared to the previous legislation, BPD.

Guidance on treated articles is further addressed in sections 5.3 and 5.4.6.

4.5.1 Efficacy assessment for active substance approval
For biocidal products placed on the market in the EU, the authorisation requirements of the
BPR apply, including testing efficacy. For treated articles imported into the EU, there is only
the active substance approval stage to test efficacy. In this respect, it is particularly
important to evaluate and assess use in treated articles at the active substance approval
stage.

Where claims to treat articles are made for active substance or biocidal products, efficacy
data to support these claims have to be submitted (see Annex II, Title 1, 6.6 and Annex III,
Title 1, 6.6 and 6.7). If claims are made on active substance level, efficacy assessment of
the use in treated articles has to be part of the active substance evaluation.

4.5.2 Efficacy assessment for active substances in specific PTs
For active substances notified for certain PTs it is obvious that they are mainly, or
exclusively used, to treat articles/materials as for example for PTs 6, 7, 8, 9, 10 (Main group
2). Thus, efficacy testing with respect to use to treat articles/materials, is a natural part of
the active substance evaluation. In such cases use concentrations and standard use
conditions for use in treated articles have to be taken into account in assessing efficacy. The biocidal function of the PTs within Main group 2 is usually protection of specific materials from biodeterioration, in some cases odour prevention. The state of the articles treated can be solid or liquid. The use conditions can be dry, humid or wet, which can be quite crucial for the release of the active substance out of the matrix. Thus, the representative product should show the claimed effect(s) in the range of uses and use conditions which are described and in the type of matrixes applied for. Use conditions like ageing, weathering or washing should be simulated as appropriate, to demonstrate the duration of the effect in relation to the life-span of the article treated.

Active substances notified for PTs 1-5 (Main group 1) are usually used in (liquid) biocidal products as for instance hand disinfection or surface disinfection products. These products are clearly considered biocidal products. But sometimes active substances belonging to PTs 2, 3 or 4 are incorporated into textiles and other solid materials; the protection of the material itself is not intended, but a new property is introduced to an article, intended to protect its user. For such claims, testing is particularly challenging and the specific conditions of use have to be considered when designing the efficacy testing. Please read more about how to design such tests in section 5.4.6. At active substance level, the representative product should show the claimed effect(s) in a range of uses and use conditions which are described and in the type of matrixes applied for. Particularly the wet state of the use conditions (dry, humid or wet) needs to be taken into account, as this is crucial for the release of the active substance out of the matrix and thus for the efficacy of the representative product. Furthermore, use conditions like ageing, weathering or washing should be simulated as appropriate, to demonstrate the duration of the effect in relation to the life-span of the article treated. Use conditions for which no efficacy of the representative product could be demonstrated must be excluded from the approval as appropriate.

Active substances belonging to PTs 18 and 19 and used to treat (solid) articles can have different purposes. The treatment can be intended to protect the material (for instance a carpet treated with an insecticide to prevent moth damage) or it can be intended to protect humans or animals against insects (for instance clothes treated with a repellent). Again, in the latter case it has to be carefully considered whether such a product fulfils the definition of a biocidal product and has to undergo an authorisation procedure. At the active substance approval stage, any claims made should be demonstrated with appropriate efficacy tests on the representative product, taking into account the specific conditions of use (e.g. regular washing for clothes) and the availability of the active substance to the target organisms, which can differ in different matrices.
5. Product authorisation

5.1 Evaluation of efficacy at product authorisation stage

The Product Authorisation stage is the point in the evaluation process where the efficacy of the biocidal product should be looked at for the full range of claims made. More test organisms or different uses can be relevant as compared to active substance approval. At this stage, it is not the properties of the active substance which are of interest, but instead the properties of the fully formulated product, which may contain more than one active substance.

Therefore, this is the stage at which a full evaluation of the efficacy of the formulated product should be carried out, and where the efficacy is evaluated in relation to the label claims made for the product. This evaluation should include all relevant target species (or representative species), the effects of using the product, the duration and speed of effect (including ageing and weathering if relevant), any claims for residual action, together with any other specific claims.

At biocidal product authorisation, the applicant must clearly describe the uses for which the product is intended when it is used under normal conditions, at the appropriate application rate and in accordance with the use instructions. This information is required to allow a proper evaluation of the efficacy to be carried out, and must include, for every product type separately:

- The purpose of the biocide (e.g. prevent destruction of material by insect infestations, decrease of bacterial contamination on surfaces);
- The function of the product (e.g. bactericide, fungicide, rodenticide, insecticide);
- The organism(s) to be controlled;
- The effects on representative target organism(s) (e.g. attracting, killing, inhibiting);
- Any products, organisms or objects to be protected;
- The concentration at which the active substance will be used (the use concentrations for different targets should be stated for each use and method of application, if appropriate. Applicants should also indicate if the use concentrations should be different in different parts of EU);
- Description of the instructions of uses.

At the product authorisation stage, efficacy must be demonstrated against all claimed target organisms. Use against additional target organisms (i.e. which were not supported at the active substance approval stage) may be applied for at this stage.

For biocidal products used to treat articles, it is important to categorise possible wide ranges of uses into sets of similar materials and use-conditions. Please see sections 5.3, 5.4.2 and 5.5 for more details.

5.2 Product families

5.2.1 Background

A product family is a group of products with the same active substance(s) and similar use, but small differences in the formulation, which do not significantly reduce the efficacy of the
products. When authorisation is requested for a product family efficacy should be demonstrated for the whole group but not necessarily of each product. A product family can be divided in different meta SPC’s, and all products in the meta SPC have the same hazard and precautionary statements. However, it is also possible that extra meta SPC's should be added because of the efficacy assessment (e.g. some products in the family are not efficacious for some uses). It should thus be noted that the efficacy evaluation of the product family should be made in conjunction with the other parts of the evaluation (e.g. ENV, HH and phys-chem) and that an overall assessment of the division into meta SPC’s should be made taking all areas into account. This guidance is specifically aimed at an evaluation of differences in efficacy claim, which could lead to certain structures of the BPF and meta SPC’s. Therefore, some of the following examples could result in other structures of the meta SPC’s when environment, human health and phys-chem are taken into account.

5.2.2 Worst case testing

The BPF concept allows read-across of data between similar products within and across meta SPCs. Efficacy tests must be performed on the product with the lowest concentration of the active substance, under the worst case circumstances. The influence of the co-formulants on the efficacy should be taken into account. A justification should be given for the product and circumstances taken.

Tests and criteria for testing efficacy of products in a family are the same as for single products. For the data requirements and test criteria, please see the specific sections per PT. Applicants need to ensure that all products within a family have been supported, in terms of:

- target organisms;
- concentrations / application rates;
- contact time;
- influence of the co-formulants;
- application methods;
- field of use / use conditions;
- other label claims;
- formulations;
- any other relevant information.

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2 See Article 3 of the BPR for the full definition of a BPF.
3 See for the definition of a meta SPC CA-Nov15-Doc_4_3Update_note_for_guidance_on_BPF_concept.docx
Table 1: Example ready-to-use disinfectants with/without pre-cleaning*.

<table>
<thead>
<tr>
<th></th>
<th>Family A</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Concentration AS: 1-4%</td>
</tr>
<tr>
<td>meta SPC 1</td>
<td></td>
</tr>
<tr>
<td>Product 1</td>
<td>1%</td>
</tr>
<tr>
<td>Product 2</td>
<td>1%</td>
</tr>
<tr>
<td>meta SPC 2</td>
<td></td>
</tr>
<tr>
<td>Product 3</td>
<td>4%</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>concentration AS</th>
<th>target organisms</th>
<th>use conditions</th>
<th>colour</th>
</tr>
</thead>
<tbody>
<tr>
<td>Product 1</td>
<td>bacteria</td>
<td>apply after pre-cleaning</td>
<td>1</td>
</tr>
<tr>
<td>Product 2</td>
<td>yeasts</td>
<td>apply after pre-cleaning</td>
<td>2</td>
</tr>
<tr>
<td>Product 3</td>
<td>bacteria</td>
<td>apply without cleaning</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>yeasts</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>viruses</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

NOTES to Table 1

In this example one worst case for efficacy cannot be identified. Product 1 should be tested against bacteria and yeasts under clean conditions (also supporting product 2), and product 3 should be tested against bacteria, yeasts viruses, under dirty conditions.

Since these are all ready-to-use products, and presuming that 1% is not efficacious against viruses, product 1 and 2 should be in a different meta SPC than product 3 since they are not efficacious against viruses. The meta SPC of products 1 and 2 will state as target organisms bacteria and yeasts and the meta SPC of product 3 bacteria, yeasts and viruses.

* In the examples, only the information given in the table is taken into account for the deviation in meta SPC’s, presuming that all other factors are the same for the different products or of no influence. In practice other factors relating to the products will also need to be taken into account.

In some cases it is not possible to identify one worst case scenario for a combination of products and use conditions: where such a single “worst case” scenario at meta SPC level cannot be identified, an assessment of the minimum efficacy levels that might be relevant for the uses covered by a meta SPC has to be performed. For instance, the family contains products (1) and (2) with low active substance (AS) concentration which will be used as disinfectant under clean conditions and only for the control of bacteria and yeast, while another product (3) with a higher concentration of AS is used under dirty conditions for the control of bacteria, yeast, and viruses. Product (1) and (2) will not be sufficiently efficacious against viruses, so it cannot be used to demonstrate efficacy for all the uses. In this family, product (1) should be tested under clean conditions against bacteria and yeast (and cover product (2)) and product (3) should be tested under dirty conditions against bacteria and yeast and viruses (see Table 1). Tests done for a product in one meta SPC can, where relevant, be used to support a claim for a similar product in a different meta SPC, provided that variations in co-formulants have no influence on efficacy. Justification may need to be provided to allow read across.

In some product families several combinations of products and uses should be tested, to demonstrate efficacy for all combinations of products and use conditions (see Tables 2, 3, and 4).
Table 2: Example concentrated disinfectants

<table>
<thead>
<tr>
<th>Family B</th>
<th>Concentration AS: 10-40%</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>meta SPC</strong></td>
<td>Product: 10-40% AS</td>
</tr>
<tr>
<td>Dilute product to use concentration:</td>
<td></td>
</tr>
<tr>
<td>bacteria: 1% AS</td>
<td></td>
</tr>
<tr>
<td>fungi: 1% AS</td>
<td></td>
</tr>
<tr>
<td>viruses: 4% AS</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Product 1</th>
<th>Product 2</th>
<th>Product 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>concentration AS</td>
<td>10%</td>
<td>20%</td>
</tr>
<tr>
<td>target organisms</td>
<td>bacteria, fungi</td>
<td>bacteria, fungi</td>
</tr>
</tbody>
</table>

**NOTES to Table 2**

In this example all products are concentrates to be diluted before use. The applicant only claims efficacy against bacteria and fungi for product 1 and 2 and in addition viruses for product 3. Presuming all products only differ in the concentration active substance, testing can be done with either of the products at use concentration: product diluted to 1% active substance should be tested against bacteria and fungi, and product diluted to 4% active substance should be tested against viruses.

Since all concentrated products can be diluted to an efficacious concentration, when used according to the instructions on the *meta SPC*, all products can be in one *meta SPC*.

Table 3: Example surface disinfectants ready-to-use: more PT’s

<table>
<thead>
<tr>
<th>Family C</th>
<th>Concentration AS: 10%</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>meta SPC 1</strong></td>
<td></td>
</tr>
<tr>
<td>Use #1: PT3, bacteria, fungi</td>
<td></td>
</tr>
<tr>
<td>Use #2: PT4, bacteria, fungi, viruses</td>
<td></td>
</tr>
<tr>
<td><strong>meta SPC 2</strong></td>
<td></td>
</tr>
<tr>
<td>Use #2: PT4, bacteria, fungi, viruses</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Option 1</th>
<th>Option 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Product 1</td>
<td>Product 2</td>
</tr>
<tr>
<td>concentration AS</td>
<td>10%</td>
</tr>
<tr>
<td>target organisms</td>
<td>bacteria, fungi</td>
</tr>
<tr>
<td>PT</td>
<td>PT3</td>
</tr>
</tbody>
</table>

**NOTES to Table 3**

In this example all products are ready to use and have the same use concentration, they only have a different use claim (i.e. same use in different PTs). It is presumed that the products only slightly differ in their composition and that it is demonstrated that this does not influence the efficacy. In
this case either of the products can be tested under worst case conditions (justification should be given that PT3 soiling and temperature is the worst case). A representative product should be tested against the specified bacteria and fungi required for PT3, and against the specified bacteria and viruses required for PT4. Since the fungi that have to be tested for PT3 and PT4 are identical, one test performed under the worst case conditions is sufficient. Since this meta SPC can be split into 2 uses, one for PT3 and one for PT4, and all products are efficacious against all uses, it is possible to put all three products in one meta SPC, (option 1). All possible products in this meta SPC will be efficacious against use #1 and use #2. Efficacy against viruses in PT3 is not demonstrated, however, since this is not in one of the uses in the meta SPC, this is acceptable. On the product label only the specified uses, combination of PT and target organisms, can be claimed. However, an applicant might consider it easier to split the family in 2 meta SPC’s, one per PT (option 2).

Table 4: Example insecticide: take target organisms and application method into account.

<table>
<thead>
<tr>
<th>Family D</th>
<th>meta SPC 1</th>
<th>meta SPC 2</th>
<th>meta SPC 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conc. AS: 1-4%</td>
<td>Conc. AS: 1%</td>
<td>Conc. AS: 1%</td>
<td>Conc. AS: 4%</td>
</tr>
<tr>
<td>Product 1</td>
<td>Product 2</td>
<td>Product 3</td>
<td></td>
</tr>
<tr>
<td>concentration AS</td>
<td>1%</td>
<td>1%</td>
<td>4%</td>
</tr>
<tr>
<td>target organisms</td>
<td>moth</td>
<td>moth and mosquitoes</td>
<td>ants</td>
</tr>
<tr>
<td>application method</td>
<td>paper in wardrobe</td>
<td>electric device in wardrobe or room</td>
<td>bait box with sugar</td>
</tr>
</tbody>
</table>

NOTES to Table 4

In this example one worst case for efficacy testing cannot be identified and all products should be tested for all target organisms and uses.

All three products should be in different meta SPC’s because of the different application methods and organisms.

When a family contains more than one active substance it might not be sufficient to test the products to be authorised in a meta SPC, in some cases it is necessary to test a ‘dummy’ product to cover all products in one meta SPC (see Table 6). Alternatively, they could be authorised in separate meta SPC.

5.2.3 Take formulation types and chemical composition into account

While the active substance is the most important constituent for efficacy of a biocidal product, the effect of the formulation of the product on the efficacy must also be taken into account. Therefore, the justification should be given for the product used in the test, taking into account the formulation. If the product contains more than one active substance, the combined effect between different active substances will be considered.

In the case of products having different formulation types (e.g. wettable powder and water dispersible granules for PT18), bridging studies with these products can be used to substantiate that the products are equivalent in terms of their efficacy. Bridging studies should involve worst case circumstances (after appropriate justification).

Depending on the influence of the ingredients (chemical composition) on the efficacy either the product with the lowest concentration of all the ingredients should be tested or several
products, together including the whole spectrum of the formulations, should be tested (see Table 5).

### 5.2.4 Allowing for the addition of new products in a family

In general the (meta) SPC(s) of a family will give a range for the concentration of the active substance(s) and co-formulants. After authorisation of the family it is possible to add new products to the family, as long as their composition falls into the range for the (meta) SPC. For these new products no evaluation will be done. Therefore, efficacy testing should be done in such a way that efficacy against all possible new products will be demonstrated.

For instance, in the example in Table 5, a new product with 70% active substance and the lowest concentration of both acids could be added. Efficacy of this product should be demonstrated, or the two products should be put into different meta SPCs. Another example is explained in Table 6.

#### Table 5: Example disinfectant: take formulation into account.

<table>
<thead>
<tr>
<th>Family E</th>
<th>meta SPC 1</th>
<th>meta SPC 2</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Concentration AS: 70-85%</td>
<td>Concentration AS: 70-75%</td>
</tr>
<tr>
<td></td>
<td>Concentration acid 1: 1-4%</td>
<td>Concentration acid 1: 1-4%</td>
</tr>
<tr>
<td></td>
<td>Concentration acid 2: 2-5%</td>
<td>Concentration acid 2: 2-5%</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Option 1</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>meta SPC 1</td>
<td>Concentration AS: 70-75%</td>
<td>Concentration AS: 70-85%</td>
</tr>
<tr>
<td>Option 2</td>
<td>meta SPC 1</td>
<td>meta SPC 2</td>
</tr>
<tr>
<td>Product 1</td>
<td>Concentration acid 1: 1-4%</td>
<td>Concentration acid 2: 2-5%</td>
</tr>
<tr>
<td>Product 2</td>
<td>meta SPC 2</td>
<td>Product 2</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>target organisms</th>
<th>bacteria</th>
<th>fungi</th>
<th>bacteria</th>
<th>fungi</th>
<th>virus</th>
</tr>
</thead>
<tbody>
<tr>
<td>Active substance</td>
<td>70%</td>
<td>75%</td>
<td>85%</td>
<td>1%</td>
<td>1%</td>
</tr>
<tr>
<td>Acid 1</td>
<td>1%</td>
<td>4%</td>
<td>5%</td>
<td>1%</td>
<td>2%</td>
</tr>
<tr>
<td>Acid 2</td>
<td>5%</td>
<td>2%</td>
<td>5%</td>
<td>1%</td>
<td>2%</td>
</tr>
</tbody>
</table>

**NOTES to Table 5**

In this example both acids are pH regulators. It is presumed they are not considered active substances in this formulation (in some cases this should be demonstrated with tests), however, both acids might enhance the efficacy to some extent (i.e. formulation effect). Since it cannot be ruled out that there is a difference in effect between these two acids, this should be taken into account in the efficacy testing.

When product 1 and 2 are placed in one meta SPC (option 1) it should be considered that it is possible to add a new product in this meta SPC with 1% acid 1 and 2% acid 2. In that case it is not sufficient to test product 1 (with lowest concentration AS), but a ‘dummy’ product should be tested, with 70% AS, 1% acid 1 and 2% acid 2.

To prevent testing with ‘dummy’ products, it might be easier to place products 1 and 2 in separate meta SPC’s, without a range for the acids (option 2). Also in that case, read across between product 1 and 2 is not possible. Both product 1 and 2 should be tested, to rule out the effect of the formulation with different acid concentrations.
In all cases product 3 should be tested against viruses, and put in a different meta SPC (assuming 85% is necessary for viruses). The test with product 1 or the ‘dummy’ product can be used to demonstrate efficacy against bacteria and fungi for meta SPC 2 (product 3).

5.2.5 Deviation in meta SPC’s

When dividing a product family in meta SPC’s, it must be taken into account that all (possible new) products will be efficacious for all uses, target organisms, etc. Worst case testing must make sure that all possible new products will be efficacious. Where needed/possible new meta SPC’s should be made for a different group of target organisms, a different use, different application method, etc.

This means for the example family in Table 4, that all products should be in a different meta SPC.

In Table 1 product 1 and 2 should be separated from product 3, because these are not efficacious against viruses and therefore not against all target organisms in this meta SPC.

However, in some cases it might be possible to not deviate in more meta SPC’s but give a good description in the meta SPC, making sure that all products will be efficacious. For instance, in the examples in Tables 2 and 3, which are very similar to Table 1, the product with a virus claim can be in the same meta SPC. This is acceptable because all possible products are efficacious when used according to the use description in the meta SPC, either because all products can be diluted to an efficacious dose, or by making separate use numbers. In these cases some of the products in the meta SPC have a limited claim (i.e. fewer organisms, fewer PT’s).

When the different uses results in a too complicated meta SPC, with several different use numbers, it is better to divide such a meta SPC in more simpler meta SPC’s.

When dividing into meta SPC’s the applicant must make sure that the text in the meta SPC’s is unambiguous, and consider that no products can be added to the family that have not been supported in the efficacy testing (see Tables 3 and 4).

Table 6: Example anti-fouling product: Different ratio’s of two (or more) active substances.

<table>
<thead>
<tr>
<th>Family</th>
<th>Option 1</th>
<th>Option 2</th>
<th>Product 1 RTU</th>
<th>Product 2 RTU</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>meta SPC 1</td>
<td>meta SPC 2</td>
<td>target organisms</td>
<td>Macro fouling</td>
</tr>
<tr>
<td>Conc. AS 1: 5-10%</td>
<td>Conc. AS 1: 5%</td>
<td>Macro fouling</td>
<td>10%</td>
<td>5%</td>
</tr>
<tr>
<td>Conc. AS 2: 2-7%</td>
<td>Conc. AS 2: 7%</td>
<td></td>
<td>2%</td>
<td>7%</td>
</tr>
<tr>
<td></td>
<td>meta SPC 1</td>
<td>meta SPC 2</td>
<td>Active substance 1</td>
<td>10%</td>
</tr>
<tr>
<td>Conc. AS 1: 10%</td>
<td>Conc. AS 1: 5%</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Conc. AS 2: 2%</td>
<td>Conc. AS 2: 7%</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
NOTES to Table 6
In this example testing product 1 and 2 is not sufficient to cover the worst-case situation of this family. The worst-case would be a product 5% active substance 1 + 2% active substance 2. Assuming variation of co-formulants have no impact on efficacy, this ‘dummy’ product should be tested to demonstrate efficacy for this family when it consists of one meta SPC (option 1). Alternatively, product 1 and 2 can be put into different meta SPC (option 2), and efficacy test using prod 1 and 2 can be provided.

5.2.6 Minimum concentration needed
Whilst ready-to-use products authorised on their own are evaluated on their merits and not in comparison to other products, this is not the case in a product family. Since all products are presented at the same time a comparison can be made. The BPR Annex VI art. 77 of the common principles state: the recommended dose is the minimum necessary to achieve the desired effect.

For historical reasons it is possible that products on the market in one EU country contain a higher concentration of AS than another product with the same intended use in another country. When this is the case the applicant should request for authorisation for the products with the lowest concentration of AS or give a good justification why it is relevant to have different formulations.

It should be considered that there may be other products on the market which contains a lower concentration of AS and is efficacious for the same intended use.

5.3 Treated articles

⚠️ NOTE to the reader:
This section concerns treated articles and should be read in conjunction with the CA Note for Guidance “Frequently asked questions on treated articles”, CA-Sept13-Doc.5.1.e, Revision 1 December 2014.

Article 3 Definitions
1. For the purposes of this Regulation, the following definitions shall apply:
(a) ‘biocidal product’ means
- any substance or mixture, in the form in which it is supplied to the user, consisting of, containing or generating one or more active substances, with the intention of destroying, deterring, rendering harmless, preventing the action of, or otherwise exerting a controlling effect on, any harmful organism by any means other than mere physical or mechanical action,
- any substance or mixture, generated from substances or mixtures which do not themselves fall under the first indent, to be used with the intention of destroying, deterring, rendering harmless, preventing the action of, or otherwise exerting a controlling effect on, any harmful organism by any means other than mere physical or mechanical action.
A treated article that has a primary biocidal function shall be considered a biocidal product.
(l) ‘treated article’ means any substance, mixture or article which has been treated with, or intentionally incorporates, one or more biocidal products.
A treated article according to Article 3(1)(l) of the BPR is any substance, mixture or article which has been treated with or intentionally incorporates one or more biocidal products. A biocidal product, in contrast, is any substance or mixture with a biocidal function. Pursuant to Article 3(1)(a) a treated article with a primary biocidal function is considered a biocidal product.

Liquids fulfil the substance or mixture definition. Consequently, liquids may only be considered as treated articles if they do not intend to control any harmful organism. In contrast, solid treated articles are defined by their shape and function rather than by their chemical composition. Thus, solid treated articles fulfil the definition of a biocidal product if they have a primary biocidal function.

The term “primary biocidal function” is not further defined in the BPR, but in the CA document, it is described as “a biocidal function of first rank, importance, or value compared to other functions of the treated article”.

A biocidal product, in contrast, is any substance or mixture with a biocidal function. Consequently, efficacy testing and assessment is not principally different for biocidal products and treated articles. Both categories can take different forms (liquid, solid) and can concern different materials. In both cases efficacy has to be shown for normal conditions of use and against an untreated control. The untreated control should demonstrate the problem which is to be solved by the biocidal treatment.

Thus, considering the different product types for PTs 1-4, the following examples would be considered as biocidal products and not treated articles. For PT 1 or 3, disinfecting wipes would be regarded as biocidal products. For PT 2, paints and coatings intended to prevent microbial settlement and growth in order to provide a hygienic environment would likewise be regarded as biocidal products. Other PT 2 applications which could fall under either category, depending on their primary function could include for instance textiles, tissues, masks, or other articles or materials in which a biocidal product has been incorporated with the purpose of adding disinfecting properties to these articles and materials. For PT 4, examples are materials or articles which come into contact with food or feed and are treated with or incorporate a biocide; whether such articles are to be regarded as biocidal products again depends on their primary function. PT 5 applications are usually biocidal products.

Further product examples are given in Appendix 1 of the CA document.

There are some exemptions in the definition given in Art. 3(1)(a): Articles such as paper or carton, where the pulp has been treated with a biocide during manufacture, and where the biocide is not intended to have a function in the final good are not considered treated articles. Another example are articles with print on it or with glue holding it together which have been treated with an in-can preservative. However, the preservative doesn’t have any function in the final article as soon as the ink or adhesive is applied and dried. In contrast, an article like a table made of a composite material with wooden legs painted with a film preservative containing coating, is considered a treated article, as the coating still has a biocidal function in the final article.

Generally, there is no difference in efficacy testing of treated articles or biocidal products in a liquid matrix. For instance, wet state preservatives (PT 6) or a hand disinfectant (PT 1) are usually both tested in a liquid matrix, the first matrix is a treated article, the latter is a

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5 See CA document Appendix 1
6 See CA document Question 8
biocidal product; only the performance standards are different in these examples. Specific requirements apply, however, when the efficacy of solid material or articles has to be tested. A test under practical conditions of use (step 3 test) is mandatory. In contrast to preserving claims, where standard materials under certain standard conditions of use can be tested, testing for disinfecting claims has to be specific for every single article. For these types of claims, the specific conditions of use are to be considered when designing the efficacy testing; for example, a polymer coating used for a hospital bedside cabinet has to be tested for the specific contaminating situation of a hospital bedside cabinet, including cleaning schemes and soiling situation; efficacy has to be shown compared to an untreated bedside cabinet. Bactericidal effects have to take effect very quickly to show an advantage compared to an untreated cabinet, where droplets of blood or saliva will dry out quickly and not either be contaminating any more. Please read more about how to design such tests in Section5.3.

Specific requirements apply, however, when the efficacy of biocides in solid material or articles has to be tested. Treated articles with claims to protect humans or animals fall under this category. In these cases, use conditions, most importantly humidity, have to be specified. Materials can be used in articles with a wide range of use conditions, and these have an effect on efficacy. For example, for a polymer article permanently exposed to water the conditions for bacterial growth are much more favourable, and different requirements apply as compared to a polymer article which is generally dry and is only exposed to occasional splashes or to the humidity which comes from touching it. But more importantly, humidity has an effect on the availability of the active substance, because it has to be released out of the matrix somehow. Another example are clothes treated with repellents; also in this case use-conditions do influence efficacy. Wearing and tearing and washing have to be taken into account to assess the efficacy. Complete protection time needs to be defined in terms of the life-cycle of the treated clothes.

Treated articles, if not biocidal products, do not require efficacy assessment under the BPR. However, active substances and biocidal products incorporated into treated articles may require assessment of their efficacy in treated articles as part of the active substance approval and biocidal product authorisation processes (if such uses are applied for).

Consequently, if efficacy is demonstrated for a certain set of use conditions, this cannot generally be transferred to another set of use conditions. The possible limits of the use conditions have to be reflected in the approval/authorisation decision. In the following, guidance is given for the testing of (solid) materials with claims to protect humans or animals.

There are two OECD test methods available:

- Guidance Document on the Evaluation of the Efficacy of Antimicrobial Treated Articles with Claims for External Effects (OECD Series on Biocides No. 1);

5.3.1 The basic distinction between material protection and protection of humans or animals

When biocides are incorporated into materials or used in the production of treated articles they are applied with two purposes:
- To protect the materials used in the article or the properties of the article in service. The target organisms have a detrimental or other undesirable effects (e.g. biodegradation, discolouration, odour formation) on the material or article.
- To protect humans or animals from the unwanted effects of organisms. The treatment is directed towards targets organisms which have no adverse effect on the item/material treated.

The following scheme gives an overview and decision help:

**Figure 1: Decision scheme to distinguish between claims for material protection and claims for protection of humans and animals**

- **Is the treatment intended to protect the material, article or its functionality from biological deterioration in service, extend its durability or prevent odour?**

  - **Yes**
    - Main Group 2, Main group 3 (PT 18, 19) of Annex V BPR
    - Protection of material/article and its properties; sections 5.5 and specifically 5.5.7-5.5.9

  - **No**
    - Main Group 1 (PT 1-5), Main group 3 (PT 18, 19) of Annex V BPR
    - Adds properties to protect humans or animals; section 5.4.6

- **Inhibits Growth**
  - section 5.4.6.2

- **Kills, Repels**
  - section 5.4.6.3

Guidance for the testing of biocidal products with a claim to protect humans or animals is given in section 5.4.6. Guidance for material protection is given in section 5.5.
5.4 Disinfectants (Main group 1)

5.4.0 General

5.4.0.1 Introduction

This guidance describes the nature and extent of data which should be available to support the label claims for biocidal products within the Main Group 1: Disinfectants. This group covers 5 product types as described in Annex V of the BPR:

**MAIN GROUP 1: Disinfectants**

These product-types exclude cleaning products that are not intended to have a biocidal effect, including washing liquids, powders and similar products.

**Product type 1: Human hygiene**

Products in this group are biocidal products used for human hygiene purposes, applied on or in contact with human skin or scalps for the primary purpose of disinfecting the skin or scalp.

**Product type 2: Disinfectants and algacides not intended for direct application to humans or animals**

Products used for the disinfection of surfaces, materials, equipment and furniture which are not used for direct contact with food or feeding stuffs.

Usage areas include, *inter alia*, swimming pools, aquariums, bathing and other waters; air-conditioning systems; and walls and floors in private, public, and industrial areas; and in other areas for professional activities.

Products used for disinfection of air\(^7\), water not used for human or animal consumption, chemical toilets, waste water, hospital waste and soil.

Products used as algacides for treatment of swimming pools, aquariums and other waters and for remedial treatment of construction materials.

Products used to be incorporated in textiles, tissues, masks, paints and other articles or materials with the purpose of producing treated articles with disinfecting properties.

**Product type 3: Veterinary hygiene**

Products used for veterinary hygiene purposes such as disinfectants, disinfecting soaps, oral or corporal hygiene products or with anti-microbial function.

Products used to disinfect the materials and surfaces associated with the housing or transportation of animals.

**Product type 4: Food and feed area**

Products used for the disinfection of equipment, containers, consumption utensils, surfaces or pipework associated with the production, transport, storage or consumption of food or feed (including drinking water) for humans and animals.

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\(^7\) This is taken to mean the disinfection of air itself. Disinfectants sprayed or vaporised into the air (e.g. room disinfection by vaporised biocide) are normally for the purpose of disinfecting surfaces and not the air itself. Disinfectants for air conditioning systems disinfect the surfaces or liquids in these systems, not the air coming out of it.
**Product type 5: Drinking water**

Products used for the disinfection of drinking water for both humans and animals.

Products in this main group are meant for the control of micro-organisms, such as bacteria (including vegetative cells, spores and mycobacteria), fungi (including moulds and yeasts), and viruses (including bacteriophages), algae and protozoa. Control may be carried out on inanimate surfaces or skin or in liquids. Note that the term "disinfectant" used for main group 1 should be read as a generic term and not according to the definition in the glossary of terms. This means that next to disinfectants it can also include products with biostatic activity.

The most important fields of use include medical, veterinary, food, feed and drinking water sectors. Applications in public, commercial and industrial areas, where application is to inanimate surfaces without direct contact with food, are included in Product type 2. If contact between disinfected inanimate surfaces and food is possible (e.g. food industry, private and restaurant kitchens), applications are included in Product type 4.

Disinfectants for medical instruments and medical equipment that are considered medical devices are covered under the Medical Device Directive 93/42/EEC (see 3.9.1 for more information). More borderline cases with other Directives or Regulations are noted elsewhere in this Guidance Document and are defined or described in other legislation or guidance.

Cleaning products which are not intended as biocides, including liquid detergents, washing powders etc. are excluded from these product types and thus this guidance is not applicable (Annex V of BPR).

Treated articles with claimed disinfecting properties or function can also fall within PTs 1 to 5: when such articles have a primary biocidal function they are considered biocidal products (see Competent Authority (CA) document 8). These articles can include a wide variety of goods, with different applications, matrices etc. This guidance deals mainly with efficacy testing of (liquid) biocidal products; the methodology for testing (solid) treated articles can be quite different. See section 5.4.4.3 of this Guidance for details of available guidance.

A “Glossary of Terms” is at the beginning of the document.

**5.4.0.2 Dossier requirements**

The following aspects are relevant for the evaluation of the efficacy of biocidal products within PT1-5:

1. The label claim and instructions for use
2. Efficacy data of the product
3. The possible occurrence of resistance, cross-resistance or tolerance.

**5.4.0.3 Label claim**

For each product, clear label claims should be provided. When the label itself cannot contain all the necessary information, any accompanying leaflet should also be considered. To simplify the text only the term "label claim" will be used below.

The types of efficacy claims made for a disinfectant/ biocidal product depend upon, among other things, the types of micro-organisms the disinfectant targets (e.g. fungal spores, yeasts, mycobacteria, bacteria or bacterial spores) and the disinfectant’s intended use (e.g.

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in hospitals, in contact with food, in animal houses, in homes). Label claims and recommendations for use, including concentration and contact time, must be supported by the results of bactericidal, fungicidal, etc. tests appropriate to the area of application, which are normally performed on the basis of the specific standards. Complete instructions for use are an integral part of the label.

The information on the product label should fully correspond with the uses pre-defined at the authorisation stage and reflected in the corresponding version of the SPC\(^9\). Applicants must indicate clearly on the product's label the spectrum of antimicrobial activity claimed.

Examples of the common fields of applications are presented in the claims matrices which are a set of tables linked to this guidance document (see Appendix 1 for more information). The Claim Matrices are not intended to be exhaustive, but the majority of uses are included.

5.4.0.3.1 Target Organisms

The target organisms for which claims are made should be specified on the product label.

As the claimed antimicrobial efficacy for disinfectant products will encompass a large spectrum of potential target organisms, it is not necessary or indeed feasible to include all possible micro-organisms in an efficacy test designed to support a label claim. Instead the types of target organism the product is intended for are mentioned, for example, fungal spores, yeasts, viruses, algae, protozoa, (myco)bacteria or bacterial spores.

Specific species are mentioned on the label where they are the only or most relevant organisms, or where they have a different susceptibility to biocides than the rest of the group. For instance, mycobacteria are less susceptible than other bacteria and it is only relevant to control them in certain situations such as tuberculosis wards.

In general it is not possible to claim against specific single species without claiming (and demonstrating) efficacy against the group of organisms (e.g. no claim against *Mycobacterium tuberculosis* without also making a general bactericidal claim, no claims against HIV without a general claim against enveloped viruses). However, there are some cases in which it can be justified that a claim only for a single or a small number of species is made (such as bacteriophages in the milk industry, or fungi *Aspergillus fumigatus* in poultry housing.).

Claims against specific organisms or groups of organisms should not be made, if they imply a false impression of superiority of a product; for example, a claim against MRSA should not be made for a bactericidal product, because MRSA do not present a specific challenge for disinfectants.

Standard test methods normally specify one or more representative species that should be tested per group of organisms for which the claim is made. For instance, a bactericidal product should be tested on gram-positive and gram-negative bacteria, a fungicidal product should be tested on yeasts and fungal spores. The species used are representative species that take into account their relevance to practical use, susceptibility for disinfectants and adequacy for laboratory testing.

The test organisms and strains which should be used are normally stated in standard efficacy test methods, i.e. according to EN 14885 or OECD-guidance.

\(^9\) Details on how to fill out the SPC are available in the ECHA Technical Guide and SPC Editor.
When it is not possible to use standard test methods for efficacy testing and other tests are used instead, the test organisms listed in Appendix 3 should be employed. If test organisms other than those listed in Appendix 3 are used, their relevance should be justified.

Wherever possible strains should be selected from international collections (their genetic stability should be checked regularly). The preservation procedures must be clearly described (EN 12353).

Other test organisms, in addition to those specified in the test standards, can also be tested. When efficacy against specific additional species is claimed, efficacy tests with those species should also be performed. In general, claims should not be made against the specific reference species used in a standard test as this can give a misleading impression that the product shows activity beyond that covered by the general (e.g. bactericidal, fungicidal) claim.

Mentioning specific organisms on the label is still a subject of discussion between Member States. The above sections reflect the position at the time that this guidance is written.

For some areas of use there are minimum requirements for the groups of organisms for which efficacy should be demonstrated. For instance, for products used for animal transport vehicles efficacy against bacteria, yeasts and viruses should be demonstrated. For these products it is obligatory to test all required organisms. Per section, a sub-section on test organisms provides information on the minimum requirements for that use.

### 5.4.0.3.2 Areas of Use

Disinfectants are used almost everywhere that people want to “eliminate” or inhibit (for static products) micro-organisms. They are used to kill or irreversibly inactivate or inhibit bacteria, fungi and viruses on animate and inanimate surfaces and matrices, in hospitals, households, schools, restaurants, offices, swimming pools, kitchens, bathrooms, dairy farms, on medical and dental equipment, eating utensils and at many other locations.

In some cases biostatic products are used which only inhibit microorganisms (see section 5.4.0.5.3 of this guidance).

Applicants should clearly indicate the intended areas of use for the product on the label, for example, areas of use could include (not exhaustive):

- Hospital and other medical areas;
- Domestic use;
- Institutional use (offices, schools etc.);
- Industrial applications, e.g. food, cosmetic, pharmaceutical industry etc.;;
- Restaurants and large-scale/canteen kitchens;
- Veterinary areas (animal housing, animal health care, teat or hoof disinfection etc.);
- Recreational areas.

### 5.4.0.3.3 Sites of Application

In addition to the types of efficacy claimed (e.g. bactericidal, fungicidal, tuberculocidal) and the intended area of use, the applicant must specify the use patterns for which the disinfectant is recommended on the label.

Broad examples of use patterns (not exhaustive) could include areas such as:

- Use on intact skin;
- Use in hospitals, operating theatres, isolation wards, etc.;
- Use in food manufacturing, retailing, processing areas etc.;
- Use in animal housing and equipment, e.g. pigs, sheep, poultry etc.;
- Use on work surfaces, cutting boards etc.;
- Use on fabrics or textiles;
- Use on toilets, bathrooms, sinks, etc.;
- Use against micro-organisms associated with human or animal waste;
- Use in air conditioning systems;
- Use in swimming pools, spas, aquariums and bathing waters;
- Use in tanks, pipelines, equipment soak or bottle wash.

5.4.0.3.4 Directions for use (Methods of application)

The label claim must specify the application method of the product. For disinfectants there is a broad range of application methods (e.g. wiping, aerosol, spraying). The in-use concentration of the solution and the contact time, which are essential for safe and effective use, should be described on the label. Any other directions for use should also be specified, such as whether the surface should be cleaned first, and claims regarding the number of times a prepared use solution can be used (or re-used) before a fresh solution must be prepared.

The application method can have a strong influence on the efficacy of a product, therefore the testing of a product should be appropriate for the application method. If specific equipment is used for application of the product (e.g. vaporisers) this should be taken into account when testing the product for efficacy. Equipment used in laboratory tests or small scale tests may (of necessity) be different from that employed in practice. This is especially the case when biocidal active substances are generated in situ using large scale equipment, such as electrolysis. In cases where small scale tests cannot be extrapolated to actual use conditions a large scale test with the equipment should be done.

5.4.0.3.5 Other interfering parameters

Any other circumstances that can influence the efficacy of a product should be mentioned on the label (e.g. temperature or pH requirements). For example, when a surface should be cleaned before applying the biocide and a no rinsing step is involved, or that alkaline cleaning fluids should not be used with acidic biocides, and vice versa.

5.4.0.4 Efficacy testing

For efficacy testing of disinfectants in general only quantitative tests methods should be used.

5.4.0.4.1 Tiered approach

For efficacy testing of disinfectants a tiered approach is recommended. The following tiers can be distinguished (in accordance with EN 14885):

- Phase 1 tests are quantitative suspension tests to establish that a product (or an active substance) has bactericidal, fungicidal etc. activity without regard to specific conditions of intended use. Phase 1 tests cannot be used for any product claim.
- Phase 2 comprises two steps:
  o Phase 2, step 1 tests are quantitative suspension tests to establish that a product has bactericidal, fungicidal, virucidal etc. activity, simulating practical conditions appropriate to its intended use.
Phase 2, step 2 tests are quantitative laboratory tests, often using carriers or living tissues with dried-on micro-organisms, simulating practical conditions to establish that the product has bactericidal, fungicidal, virucidal etc. activity.

- Phase 3 tests are field tests under practical conditions.

### Phase 1

Phase 1 tests are laboratory suspension tests to establish the basic activity of the product or active substance. These tests may be used during the development of the product, but are not accepted for product authorisation. However, a phase 1 test can be used to demonstrate that a co-formulant does not have any biocidal activity in the product.

### Phase 2, step 1

Phase 2, step 1 tests are laboratory suspension tests in which the ultimate purpose is to establish at what concentrations the product meets specified requirements under “in-use” conditions. In these tests, in-use conditions (e.g. temperature, contact time, interfering substances) are considered in the test method.

Various laboratory methods have been developed for biocide activity testing. Although these experiments differ in their design and experimental detail, they are all based on the principle of adding a test inoculum to the disinfectant (or vice versa) and taking samples at specified times. The biocide in each sample is then neutralised and the survival of the organisms assessed. In practice, the methods can be classified into two groups, according to how the end-point of the test is determined:

- **Quantitative tests**
  
  Samples of untreated and biocide-treated cells are plated on nutrient medium after neutralisation. After incubation, the number of colony forming units is determined and the log10 reduction in viable counts is determined.

- **Capacity tests**
  
  The biocide is challenged successively with the test organism at defined time intervals. This type of test can be used for instance for swimming pools and toilet disinfectants which are challenged by new bacteria periodically. Following each inoculation, samples are taken, and after a suitable contact period has elapsed, the biocide is neutralised and the sample incubated in a suitable growth medium to determine the surviving micro-organisms. The result is expressed as the amount of the accumulated inoculum that was required to produce the “failure”.

### Phase 2, step 2

Phase 2, step 2 tests are simulated use or practical tests, performed under rigorous conditions within the laboratory, which mimic real-life conditions, for instance by pre-drying the micro-organisms onto surfaces. These tests are used in a second testing stage. After measuring the time-concentration relationship of the disinfectant in an in-vitro test (phase 2, step 1), these practical tests are performed to verify that the proposed use dilution is likely to be adequate in real-life conditions. For several uses standardised, simulated use tests exist (surface disinfection, hand wash or rub, instrument disinfection) but there are no standard tests available for many others.

Longer-lasting activity is claimed for some products. When these products are applied to surfaces, it is common that they will not be completely removed or rinsed off after application. This might lead to longer-lasting activity of the biocide on the surface. Likewise, some products are used for maintenance via continued release of low levels of biocidal product. Both effects can be determined by appropriate efficacy tests.
Phase 3 Field or in-use tests
In-use testing involves the antimicrobial evaluation of the product under actual conditions of use on specified surfaces or materials in a specified environment. As with standard and non-standard laboratory methods, representative organisms or actual organisms of concern may be used. Validated methodologies for these types of tests are currently not available, although some are in development.

The practical use conditions under which a product can be used can be very variable and are therefore difficult to standardise. Field tests, although not standardised, can however give valuable additional information on the efficacy of the product, provided that the studies are scientifically robust, well reported and provide a clear answer to the question. In these types of test, a control treatment without biocide should be included. Where this is not possible, efficacy should be judged on a comparison of the situation before and after application.

Until validated standards are prepared, the responsibility for determining the acceptability of data derived from field trials in support of the claim will lie with the CA, taking into account the guidance given in EN 14885.

5.4.0.4.2 Standard test methods

Ideally, data should be generated using internationally or nationally recognised testing methods (CEN, OECD, ISO, etc.). Several international standard test methods currently exist for disinfectant products. Recommended standard tests are presented in Appendices 2 and referenced in Appendix 4 to this guidance document.

If there are no guidelines available for the specific use of a product, or guidelines are not suitable, the applicant may use other methods (such as intra-company Standard Operating Procedures), where the studies are scientifically robust, well reported and provide a clear answer to the question. In addition, the test methods used, together with the test conditions, should be clearly and fully described and must address the efficacy claim that appears on the product label. The use of existing guidelines, with modifications to make the guideline more suitable for the specific product or use conditions, is also possible. EN 14485 provides guidance on modification of standards (EN 14485, section 4.2 version 2014).

At the time of publication of this guidance document, a broad range of CEN methods are available. OECD has several phase 2/step 2 test methods developed for the efficacy testing of disinfectants to be used on hard surfaces which have been published as Guidance Documents. Available tests are presented in Appendix 2 and referenced in Appendix 4. The use of CEN test methods is highly recommended, where these are available and relevant. However it should be noted that although this Guidance is mainly based on EN standards, there are some cases where there are discrepancies compared to the EN tests. In such cases the ECHA Guidance should be followed as the leading guidance. OECD test methods may be used if, for example no CEN standard is available.

These methods, described below, typically give a standard set of test parameters, test organisms and pass criteria. Where specific conditions apply for a field of use, such as high/low level soiling, high/low temperatures, relevant contact times etc. these conditions should be included in the efficacy tests.

CEN Standard Test Methods
A Technical Committee (TC 216) was established in the European Committee for Standardisation (CEN), to produce harmonised European methods for testing the activity of disinfectants used in medical, veterinary, food, industrial, domestic and institutional areas.
The standards are based on suspension tests (phase 1 and phase 2, step 1) and some simulated use tests like surface tests (phase 2, step 2).

European standard EN 14885 gives information on the application and interpretation of European Standards for the testing of chemical disinfectants within product types 1, 2, 3 and 4 of the Directive / Regulation.

This document outlines the various standards currently available and provides guidance as to the choice of available standards that may be used to demonstrate the effectiveness of disinfectants in particular situations (such as medical, veterinary and food hygiene) and on the interpretation of results from such tests in making and supporting efficacy claims.

In EN 14885 products intended for domestic use are grouped with products for use in food and industrial areas, and therefore the tests specified are not always relevant to domestic areas. For instance, the virus test EN 13610 only tests against bacteriophages. In these cases the test from the medical area should be used where relevant. In cases where no test method is available for one area of use (e.g. sporicidal test in medical area), a test from another area can be used instead, provided that the test parameters (soiling, temperature, etc.) are adapted to the intended use area (for further guidance on adaption of tests see EN 14885 section 4.2).

The application of disinfectants to water systems such as swimming pools, spas, and drinking water is not addressed in EN 14885. For the evaluation of activity against *Legionella* in aqueous systems (water used in cooling towers and water for general purposes, like spas, pools, showers and other uses) a quantitative suspension test is available (EN 13623).

EN 14885 includes guidance on how a phase 3 field trial should be conducted. This guidance is intended to advise on the factors to be taken into account and controlled when performing a field trial.

The use of CEN test methods is highly recommended, provided that the methods are applicable for the use of a product. In some cases, the method can be adapted (other contact times, soiling, etc.) to fit the use conditions. Any deviation from a standard must be clearly described and a justification for any deviations provided.

**OECD Standard Test Methods**

The OECD publishes practical test methods (comparable to phase 2, step 2 tests (1.4.1.3) or phase 3 (1.4.1.4)) for testing the efficacy of disinfectants on non-porous surfaces within the “Series on Testing and Assessment“ or the “Series on Biocides”, respectively. Currently, all available methods have been issued as OECD Guidance Documents. Guidance Documents are, however, not covered by the Mutual Acceptance of Data (MAD) principle and are advisory in nature. Further developed OECD Test Guidelines might become available in the future. As European Standards are not available for all types of applications yet, the use of OECD methods is recommended provided that the methods are appropriately reflecting the use of a product. Again, the methods can be adapted (other contact time, soiling, etc.) to better fit the use conditions, provided that any deviations from the standard are clearly described and justified.

Please note that in the OECD Guidance Documents on disinfectants, the volume of disinfectant solution added to the surface is very high compared to what is normally done in practice. This test protocol can only be used for uses where the volume of disinfectant solution per surface area is similar to the intended use (e.g. flooding).
Other Standard Test Methods
While CEN standards and, in case no CEN standard is available, OECD methods are highly recommended, there are circumstances in which these tests cannot be applied, i.e. they are not available, or relevant to a particular product or use pattern. In those cases other test methods can be used.

Other test methods, for example VAH (former DGHM), DVG, AFNOR, US-EPA, AOAC or ASTM methods, are available and might be used when no international standard is available for a specific application. Where these methods lack predefined test parameters, target organisms or pass criteria, the applicant has to provide evidence why the chosen parameters are appropriate for the intended application.

Where no standard tests are available, suitable test protocols may be designed (and justified) by the applicant, but these should be discussed with and agreed by the CA before testing takes place.

5.4.0.4.3 Data requirements
Label claims and recommendations must be supported by the results of tests appropriate to the area of application.

In each test the composition of the product to be tested should be clearly described, including the identity and function of the active substances specifying quality and quantity in the formulation. In addition, because the co-formulants can affect the efficacy if the product, they must also be clearly described including identity and function. Alternatively, the formulation can be identified by a retrievable reference name or number. In such cases (i.e. it may only state a code for the product for the purposes of confidentiality), the composition of the tested product should be provided separately. As the formulation may affect the efficacy of the product, the composition of the product tested should be the same as the product under consideration. If not, justifications should be provided for any differences, and these will be assessed on a case-by-case basis.

As phase 1 tests do not take practical use conditions into account, they are not considered acceptable to support claims during product authorisation. In general, phase 1 tests are used during the development of the product, for inclusion of active substances on the “Union list of approved substances” under the BPR or to prove that a co-formulant has no biocidal activity.

In general, at least phase 2, step 1 and step 2 tests are required to support label claims during product authorisation. The phase 2, step 1 test will provide basic information on the efficacy of the product (in a standard test), while phase 2, step 2 tests investigate the effects of more in-use factors (such as drying of target organisms). The combination of phase 2, step 1 and step 2 tests will generally provide a robust data package to demonstrate the efficacy of a product. Deviations from the tiered approach should be justified.

In some cases, for example when disinfection is done in suspension under real use conditions (because the target organisms are suspended in a liquid already or will be suspended during the process due to mechanical action, for example, in CIP), a phase 2, step 1 test is sufficient on its own, as this already simulates practical conditions.

In other cases a phase 2, step 2 test may be replaced by a phase 3 test where a phase 2, step 2 tests is not appropriate. In general, a phase 3 test will be done in combination with a standard phase 2, step 1 test, as phase 3 tests are often variable.

Where in-use conditions cannot be simulated, phase 3 tests are required (e.g. drinking water disinfection with ionisation equipment).
If more than one test method is available and applicable in phase 2, step 2 to substantiate a label claim for efficacy, it is sufficient to provide data from only one of the test methods. The test method selected should be one which best represents the way in which the product is used. For example, in the case of a disinfectant used for "hard, non-porous surfaces by spraying", the test method should be one for such surfaces without mechanical action and with representative conditions of use, such as contact time, soiling, temperature and test organisms.

It is not mandatory to perform the tests under obligatory test conditions of the standards if the claimed use conditions of the products are different from these obligatory tests conditions.

Tests have to be performed with relevant target organisms, which are selected in accordance with the standard and the intended use of the product. This is further discussed in Section 1.3.1 of this Guidance. A list of standard test organisms is given in Appendix 3.

The concentrations used in testing should be selected to demonstrate the threshold of product efficacy. Suspension tests should be performed with several dose rates, including at least one rate lower than the effective rate. Competent Authorities (CAs) will evaluate dose response data generated in these tests in order to assess if the recommended dose is appropriate (i.e. the concentration is not too high, or at the minimum) to achieve the desired effect.

For biocidal products which claim a biostatic effect (bacteriostatic, fungistatic, etc. i.e. the ability to inhibit growth of bacteria, fungi etc. without killing them) the efficacy should be shown by suspension tests and simulated use tests (e.g. surface tests). The suspension test and simulated use test should be performed with and without neutralisation and with a water control (where water is tested instead of the product). The results from this testing should show that the product prevents growth of the test micro-organism (i.e. a lower level of test organism compared to the water control) but does not necessarily inactivate them (the micro-organisms survive in the test without neutralisation).

Biocidal products that claim a biostatic effect bear the risk of development of organisms with temporary or permanent reduced susceptibility (resistance). For this reason, efficacy of these types of products has to be examined carefully.

In case of in situ production of the active substance or when an apparatus is used to dose the active substance in the right amount to the water, the report should contain information on safety measurements concerning over and under dosing.

Other products, which do not have biocidal or biostatic activity, might fall within the scope of the BPR, Article 3 1 (a) "with the intention of destroying, deterring, rendering harmless, preventing the action of, or otherwise exerting a controlling effect on, any harmful organism by any means other than mere physical or mechanical action". No EU standards are available for these types of product yet, so applicants should provide a method following the principles of this guidance and based on scientific evidence. During development of new tests, or when an applicant is considering using a non-standard test or using novel testing methods, they should discuss this with the CA as to the acceptability and applicability of the test.

In the following sections, guidance on the requirements per product type and use will be given.

Detailed but non-exhaustive lists of the most relevant product applications and uses of biocides, together with the required test methodology, are given in the claims matrices.
which are a set of tables linked to this guidance document (see Appendix 1 for more information).

For uses and claims that are not specifically mentioned in this document the requirements will be set on a case by case basis by the CA.

5.4.0.4.4 Relevant factors of the test procedure

Formulation of the tested product

A product authorisation is given to a single biocidal product with a defined composition or to a group of products making up a biocidal product family (BPF) and having similar uses, the same active substances, similar composition with specified variations and similar levels of risk and efficacy.

With respect to a single product the efficacy of its specific formulation should be demonstrated. Therefore it is important that the formulation tested is clearly reported in each test report (or provided alongside the test report with a statement that it is the formulation which has been tested). The formulation details should specify the active substances and co-formulants present, together with their respective concentrations, and should confirm that all tested formulations contain the same co-formulants and concentrations. Any deviations should be mentioned and justified in a statement or in the relevant efficacy reports. Where there are deviations in the formulation from that in the product for which authorisation is sought, the tests will only be considered relevant where it is evident that the deviations have no effect on efficacy. In cases where this is not evident, a confirmatory study with the organisms that is most difficult to control should be proposed.

Within the BPF the minimum level of efficacy over the whole potential range of products should be demonstrated and the permitted variations in composition and intended uses should be explicitly identified.

The test formulations should be chosen in such a way that they cover the whole potential range of products. The test formulations should include at least a product with the lowest concentration of active substance. A justification should be given whether co-formulants influence the efficacy. When co-formulants might influence the efficacy, the worst case concentration of co-formulants (i.e. low concentration of a co-formulant that might have a positive effect on efficacy, high concentration of a co-formulant that might have a negative effect on efficacy) should be tested. See also 1.5.7 for more information on testing BPF.

Hard Water Claims

The degree of hardness of the water used to dilute the disinfectant may affect its performance (by the presence of metal ions such as Ca$^{2+}$ and Mg$^{2+}$). Generally the harder the water is, the less effective the diluted disinfectant will be. Therefore, test programmes which require that products are diluted with potable water must be diluted in water of standard hardness as defined in the corresponding test standard, for the purpose of efficacy testing.

It follows that any product that carries label claims for effectiveness in hard water must be tested by the appropriate method in water with defined hardness at the level claimed.

Presence of Interfering Substances

Where disinfectants are applied to either inanimate surfaces or skin or liquids, substances may be present on the surface or in the liquid, which may affect the disinfectant’s activity.

The nature, amount and condition of the soiling present will affect the efficacy of a disinfectant.
In many cases residual contamination must be expected and in some situations (e.g. in the treatment of blood spillages) disinfectants are specifically used to decontaminate soiling, to prevent infection transfer and to assist in safe disposal.

Blood, urine, faeces, food debris, fats and oils, dust and proteinaceous materials are the most likely organic soilings to be encountered. Limescale, milkstone and soil are the most common inorganic soilings.

Where claims are made for use under soiled or dirty conditions, the use concentrations of the product must be determined from tests performed in the presence of suitable soiling materials. Soiling materials commonly used in efficacy test methods include albumin serum, blood, yeast and yeast extract.

In practice, with exception of a few situations (e.g. clean rooms), the presence of soiling on surfaces or in liquids to be disinfected cannot be ruled out. For this reason, a small amount of interfering substance should always be included during the testing of the product. In the CEN methods this is called "under clean conditions". Tests under clean conditions can be used when the surface is clean before disinfection. This is for instance the case when the label states that cleaning prior to disinfection is necessary. When a product claims combined cleaning and disinfection, the product should be tested under dirty conditions (see Appendix 4 for more information). Also, where the label only states excessive dirt should be removed, and the surface is still soiled after that (e.g. in the meat industry), soiling for dirty conditions should be used. Please note that in some cases EN 14885 is not always sufficient to meet BPR requirements.

When a product is to be recommended for certain uses where the soiling is of a specific type (such as soap film residue or hard water scum), the product must be tested in the presence of that specific soiling type. If more soiling types are relevant for the use of the product (e.g. a product must be used in the beverage industry, in meat industry or in kitchens), pre-testing should be done to determine the most challenging soiling type. Extended testing with the most challenging soiling type will be sufficient to cover all the others.

As an exception to the rule, products to be used in cleanrooms do not require additional soiling in the test. A cleanroom has a controlled level of contamination that is specified by the number of particles per cubic meter at a specified particle size. The soiling level in cleanrooms is so low that even testing under clean conditions for the EN tests is still over-dosing of soiling compared to cleanrooms. For these uses the high load of test organisms can be seen as soiling. Tests without soiling will only be accepted when the label states the specific use in clean rooms which are classified according to ISO 14644-1 in class 1 to 9 or according to GMP EU classification in Grade A to D.

Generally, soiling will reduce the efficacy of the disinfectant, and where soiling is present, longer contact times, higher concentrations, pre-cleaning or a combination of these elements may be necessary.

**Temperature**

Generally, disinfection performance increases with temperature, although this depends on the active substances and the effect on individual species may vary depending on the specific properties. Therefore, the test temperature should be representative of those encountered during the intended use of the product (e.g. low temperature in animal housing, higher temperature in CIP). Some biocides are used in chemothermal disinfection, for instance, some CIP treatments are done under temperatures of 60-80°C. Also for these uses the products should be tested at the use temperature.
If products (PT 2-4) are tested with high temperatures above 40°C heat resistant reference test organisms must be used. Enterococcus faecium must be used as the only test organism for claiming bactericidal activity. For a virucidal claim the only test organism must be Murine Parvovirus. For a sporidical claim the test organism can be spores of, for example, Bacillus cereus or Clostridium sporogenes.

For mycobacteria, yeasts and fungal spores no relevant test organisms for high temperatures are available. Most yeasts and fungal spores are already irreversibly inactivated by high temperature (>40 ºC) in the control without active substance. However, ascospores of several fungi can become heat resistant and can cause problems in, for instance, the food industry.

When standard tests with relevant temperature resistant strains become available for mycobacteria, yeasts and fungal spores, these should be used.

When efficacy against mycobacteria, yeasts and fungal spores is claimed and no temperature resistant strains are available, the standard test organisms should be tested at the maximum temperatures for which the test is validated.

For specific claims against heat resistant species (e.g. Talaromyces flavus) efficacy tests with these organisms should be provided. In these tests a control without biocide should be included which shows survival of the test organisms at the high test temperature.

It is possible that the concentration needed to pass the test is higher for the organisms tested at low temperature than for the temperature resistant organisms tested at higher temperature. In that case a justification should be given on how the test results reflect the use concentration in the use instruction on the label.

**Contact Time**

The contact time of a product on a surface etc. is an important aspect in the evaluation of the efficacy of disinfectants. In general, the longer the contact time, the more effective the disinfectant is. In trials where test organisms are taken from treated samples for further analysis, the contact time between the biocide and the test organisms should be stopped. Neutralisers, membrane filtration or subculture techniques are used to prevent residual carry-over of active substances. Neutralisation is discussed further in section 1.4.4.6 of this Guidance.

Some disinfectants act very quickly, whereas others require an extended contact time to achieve adequate performance. Mycobacteria, bacterial spores, fungal spores and non-enveloped viruses take longer to be irreversibly inactivated than most vegetative micro-organisms.

The contact time that is practical in real life use should be taken into consideration when testing. In phase 2 and phase 3 tests the product should pass the test at the contact time recommended on the product label.

**Neutralisation**

Neutralisers are used to stop the product’s activity in trials where the test organisms are taken from treated samples for further analysis, such as plate count following biocidal treatment. An effective neutraliser for the test product should be identified, and evidence demonstrating the effectiveness of the neutraliser against the product under test, and showing that the neutraliser itself does not have antimicrobial activity, must be included in a test report. Membrane filtration or subculture techniques can be used to stop the product’s activity, in combination with or instead of chemical neutralisation. These other methods are covered by the term “neutralisation” as used in this guidance.
Appropriate controls for determining the efficacy of the procedure to stop the product's activity after the contact time should be performed.

**pH**
The prevailing degree of acidity or alkalinity during disinfection can also affect the performance and choice of disinfectant. Therefore, the pH of the product at the use concentration (diluted) as used in the test must be included in the test report.

**Texture of Surfaces**
Smooth impervious surfaces are easier to disinfect (and also to clean) than rough or pitted ones. In some circumstances the micro-organisms might be protected from the action of disinfectants by being protected in porous surfaces. Clumps of micro-organisms may also be more difficult to inactivate, as cells inside are protected by dead micro-organisms on the outside. Recently porous surface tests have been developed (CEN) to test under these conditions.

Bacteria and fungi can adhere to surfaces forming biofilms. In biofilms susceptibility is decreased (the bacteria are in a different physical state) and penetration of biocide can be difficult to achieve due to the matrix surrounding the bacteria. This makes bacteria in a biofilm more difficult to inactivate.

**Repetition**
In general test results become more reliable when the tests are done in replicates (e.g. repeated in time, in more test objects). Replicates should be performed as required in the appropriate EN standards and where appropriate, internal standards or reference substances should be included.

EN14885 section 5 (parts b, c and d) state the following information on precision of the test methods (repetitions):

- For standardised tests, or adaptation of a standard test, it is recommended to repeat the test and/or include an internal standard and/or performing the test in a second and/or third laboratory. When doing the latter the second laboratory (and any further laboratory) might only repeat the test which is regarded as the most relevant one with the least susceptible test organism(s). If results from two or more laboratories are used, each laboratory has to specify one result, e.g. “R = > 5.2 lg (EN 13727-instrument disinfection)”. Then the mean of the results of all laboratories is calculated assuming each laboratory’s result as equivalent. Results with lg “more than” are set as this figure, e.g. “> 5.2 lg” is used for calculation as “5.2 lg”. All lg values are converted to real numbers, e.g. 5.2 lg to about 158.000. The mean is the arithmetic mean of these converted numbers. If one of the testing laboratories obtains a result less than the required lg reduction, the product must pass if further tests by three other laboratories demonstrate a pass. The calculations above cannot be done with tests where pass criteria are not expressed as lg reduction.

- In case of repetition of the test it is unnecessary to repeat the test with all test-organisms but only with the least susceptible to the product under test.

- If two or more tests are carried out to support a claim of performance (e.g. phase 2, step 1 and phase 2, step 2) and the ensuing recommendation for use, the tests may be ranked according to their order of relevance, i.e. their ability to predict the product’s performance under real life conditions. In case of a ranking only the result of the most relevant test may be repeated taking into account advice 3). If a ranking is not possible only the results of the test showing the highest minimum active concentration should be repeated.
5.4.0.5 General data requirements

5.4.0.5.1 Test range
Tests (phase 2, step 1) should be performed at a range of concentrations in order to verify that the use concentration is suitable for the desired effect (e.g. not too high or not at the minimum effective level).

5.4.0.5.2 Claim for several areas of use
In cases where the product is intended for several areas, it is usually acceptable to perform the tests from only one area, as long as the test is performed with the worst case test conditions (temperature, log reduction, interfering substances, etc.) and the test with the highest/most stringent pass criteria is used. In case the strains are different between the PTs all the strains must be tested.

5.4.0.5.3 Biocidal products with biostatic effect
For biocidal products with a biostatic effect (bacteriostatic, fungistatic, etc.), the efficacy should be shown by suspension tests and simulated use tests (e.g. surface tests). The suspension test and simulated use tests should be performed with and without neutralisation. The results from these tests should show that the product prevents growth of the test organism (no increase in numbers compared to the negative control) but does not necessarily inactivate them (survival of the test organism in the test without neutralisation).

5.4.0.5.4 Malodour control
There are specific requirements for products claiming control of organisms that cause malodour. Phase 2, step1 and step 2 tests should be performed with odour producing microorganisms. A justification for which bacteria, fungi, etc. are relevant to the intended use should be provided. Along with these laboratory tests, an odour test should be performed. The CA will decide on a case-by-case basis whether the product will receive authorisation.

5.4.0.5.5 Changes in ingredients
When small changes are made to the non-active ingredients in a product, it is not always necessary to repeat all the tests with the new formulation. The applicant may provide a description of the changes and the effects that they have on the efficacy of the product. In the case of a minor change, a robust justification might be sufficient (to be decided by the CA). In other cases, new efficacy tests will have to be provided. This can be either a full set of efficacy tests or a test with the least susceptible organism in the former test.

5.4.0.5.6 Treated articles
See Section 5.3 for guidance on Treated Articles.

5.4.0.5.7 Biocidal Product Families
When authorisation is requested for a product family, efficacy should be demonstrated for the whole group but not necessarily of each product. More information is available in Section 5.2 Product Families.

5.4.0.6 Resistance
See section 3.2 for guidance on resistance.

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10 For this section, the product family concept of the BPR is not yet taken into account.
5.4.0.7 Assessment of application for authorisation

5.4.0.7.1 Decision making

Biocidal Product Regulation 528/2012 (Annex VI) stipulates rules for decision making for biocides.

The test results must meet the requirements of the standards or other criteria for acceptance which are described below per type of use. Where a product does not conform to these criteria, the applicant should provide a justification in the application as to why the product should still be recommended for authorisation. The CA will decide on a case-by-case basis whether the product will receive authorisation.

5.4.0.7.2 Assessment

The CA assessor/expert assesses the performance of the product as demonstrated in the submitted efficacy tests against the label claims made for the product and the above criteria. If the product is judged to be sufficiently effective in laboratory (and, where relevant, field) tests, the product will be recommended for authorisation as far as efficacy is concerned.

In exceptional cases the applicant may provide justification as to why the specified acceptance criteria are not met but the product is still acceptable. The CA will evaluate the justification on a case-by-case basis, possibly in consultation with the other CAs, and decide whether it is acceptable or not.

The following sections give more specific dossier requirements per type of disinfectant.

5.4.1 PT1 Human hygiene biocidal products

5.4.1.1 Introduction

Product type 1 contains biocidal products used for human hygiene purposes, applied on or in contact with human skin or scalps for the primary purpose of disinfecting the skin or scalp.

Products applied on human skin may be assigned to either biocidal, medicinal or cosmetic products or even to medical devices. If the product under investigation is within the scope of the Medicinal Products Directive (2001/83/EC), the Cosmetic Products Regulation ((EC) No 1223/2009) or the Medical Devices Directive (93/42/EEC), it is excluded from the BPR for the respective use.

Products for disinfection of damaged skin (e.g. wound disinfection) or disinfection of undamaged skin before a medical treatment of a patient (e.g. pre-operative skin disinfection before surgery and disinfection before injection) and products with a claim of medicinal use, are always medicinal products (covered by the Directive 2001/83/EC on medicinal products for human use).

Biocidal products within PT1 are mainly hand disinfectants, which can include disinfection of wrist and forearm.

When applying for authorisation for a biocidal product within PT1 a detailed description of the intended use should be given, to prevent authorisation of medicinal products, or cosmetic or medicinal uses, as biocides (e.g. the claim “skin disinfection” is insufficient).

For products that fall under the BPR the data requirements described in the following sections apply.
5.4.1.2 Hand disinfectants

5.4.1.2.1 Introduction

Hand disinfectants can be divided in hygienic handwash, hygienic handrub, surgical handwash and surgical handrub products. Handwash products are intended to be used with water, handrub products are intended to be used without the addition of water. Hand disinfectants can include disinfection of wrist and forearm. Products include liquids, gels, wipes, etc.

Hand disinfectants can be used in a wide variety of areas such as hospitals and other health care institutions, food, beverage and other industry, private homes, etc.

In the sections below the requirements and acceptance criteria for most common uses are specified. For other uses and claims that are not specifically mentioned the requirements will be set on a case-by-case basis by the CAs.

5.4.1.2.2 Data requirements

Test methods

For efficacy testing of hand disinfectants, the tiered approach as described in section 5.4.0.4.1 of this Guidance is preferred. For hygienic handwash, hygienic handrub, surgical handwash and surgical handrub phase 2, step 1 and step 2 tests are required. Phase 2, step 1 tests are available for all relevant test organisms and required depending on the claim made. For a claim without specification of the area of use the phase 2, step 1 for medical area should be used.

For bacteria a phase 2, step 2 test is available for these uses and therefore mandatory. For other organisms phase 2, step 2 tests will be mandatory when they become available. For an overview of available EN tests see Appendices 2 and 4.

Disinfectant towelettes/wipes

For hand disinfectant wipes, phase 2, step 1 tests should be done preferably with the liquid extracted from the wipe or, if difficult to extract, use the liquid as it is before it is added to the wipes. Phase 2, step 2 tests for hand disinfection (modified EN 1500) should be tests with the wipe applied on volunteers hands according to the intended use. The wipes should be used on full hands and not on the fingertips only. In addition, a test must be performed that shows that either the wipe will still disinfect if the wipe dries out or that the wipe stays wet long enough to disinfect according to the claim. In addition, the use directions can address these issues, for instance, stating on the label that only wet wipes are efficacious or giving expiry dates for re-sealable packages if appropriate according to the intended use conditions.

Test organisms

Hand disinfectants intended for general hygiene purposes should be at least sufficiently effective against bacteria and yeasts. Efficacy tests with these organisms should always be provided. For hand disinfectants intended for use in industrial environments to prevent spoilage of products, in some cases also prevention of bacteria and yeast infections is of importance, for example, in food and cosmetic industry. In other industries it may be justified that only efficacy against bacteria is sufficient.

For all other groups of organisms tests have to be provided only when activity against those specific organisms is claimed.
The test organisms used in efficacy tests are normally stated in the applicable standard test methods.

Additionally to the obligatory species, other species can be used if there are valid scientific arguments to justify their use, such as a need to show activity against specific organisms of concern in a human health environment, especially emerging health risks, or in specified industries.

An overview of reference test organisms is given in Appendix 3.

**Virucidal activity**

For products used as hygienic hand disinfectants a differentiation in the virucidal activity is made.

The claims can be:

- full virucidal activity;
- limited spectrum virucidal activity;
- activity against enveloped viruses.

For each claim different test organisms should be tested.

The EN 14476\(^{11}\) test for virucidal activity gives the opportunity to either test for full or limited spectrum virucidal activity for hand disinfectants. For full virucidal activity Poliovirus, Adenovirus and Murine Norovirus should be tested. Limited spectrum virucidal activity is a claim for hygienic handrub and hygienic handwash products using Adenovirus and Murine Norovirus as test organisms, thus including activity against the test viruses and all enveloped viruses (see Appendix 5 for a list of relevant viruses). Activity against enveloped viruses is currently being discussed to be included in EN 14476\(^{12}\) (test virus: MVA = Modified Vacciniaviruses Ankara).

When only the limited spectrum virucidal activity or activity against enveloped viruses is demonstrated the label claim cannot be “virucidal”. The SPC should clearly state which of the possible virucidal claims was demonstrated.

Non-professionals may not understand the difference between a virucidal claim and a limited spectrum virucidal claim. Therefore, the instructions for non-professionals should be carefully worded. National hygiene agencies should decide how this can be communicated to the public and how the label claim (in the SPC) should be phrased to prevent misuse (e.g. limited spectrum products will not be efficacious during an outbreak of Hepatitis A virus or Enterovirus).

**Test conditions**

It is important that the tests are performed using the same contact time as claimed on the label.

The contact time can be found in the relevant EN tests. In general the contact times are:

- for hygienic handwash and handrub products the contact time is between 30 and 120 seconds;

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\(^{11}\) The current published version is EN 14476: 2013

\(^{12}\) At the time of publication of this Transitional Guidance, prEN 14476:2011 is under development and has been submitted to CEN.
• for hygienic handwash and handrub products used in medical area the contact time is usually 30 seconds for bactericidal, yeasticidal activity and activity against enveloped viruses;
• for surgical hand disinfection products the contact time should not exceed 5 minutes.

It must be assured that the disinfected hands stay wet during treatment (e.g. by applying enough product or by applying the product several times if the volume necessary is too much to apply at once).

Phase 2, step 1 tests should be carried out with soiling (interfering substances) for clean or dirty conditions depending on the intended use and according to the relevant EN tests, i.e. EN 13727 and EN 13624 (medical and veterinary area) or EN 1276 and EN 1650\(^{13}\) (industrial, domestic, institutional area). Dirty conditions in phase 2, step 1 tests are mandatory for handwash applications. For handrubs, clean conditions in phase 2, step 1 tests suffice if use instructions state that the product must be applied on visibly clean hands.

For handwash products the phase 2, step 1 tests should be performed with a dilution of the product to take into account that the product is used on wetted hands. This is described for bacteria in test EN 13727 and a similar approach should be taken for other organisms.

Phase 2, step 2 tests are performed without additional soiling according to EN 1499, EN 1500 or EN 12791. The soiling needed for clean and dirty conditions can be found in the relevant EN phase 2, step 1 tests and in the Table of Pass Criteria and EN Standards available on the ECHA Biocides Efficacy Working Group webpage (see Appendix 4 for more information). Note that dirty conditions for products used in hospitals and health care differ from those in other areas of use.

5.4.1.2.3 Acceptance criteria

A product in PT1 will be assessed to be sufficiently effective if the required laboratory tests have been carried out (using the required test organisms and test conditions), and when the pass criteria for the tests have been met.

Where pass criteria are available in the standard test, these should be met. For PT1 products the required log\(_{10}\) reductions are referenced in Appendix 4 or EN14885.

Since the test methods for these types of products are generally established, deviations are not foreseen. If, however, deviations are considered necessary, they must be justified in the application. The CA will evaluate any justification on a case-by-case basis, consulting the other CAs where appropriate, and decide whether it is acceptable or not.

5.4.1.3 Other skin and scalp disinfection

For other skin and scalp disinfection products the overlap with medicinal and cosmetic products is significant. Only products that are not covered under either of these directives can be considered as PT1 disinfectants.

5.4.1.3.1 Data requirements

Test methods

For other skin disinfection products the tiered approach as described in section 5.4.0.4.1 of this Guidance is preferred: phase 2, step 1 and step 2 tests are required.

The same phase 2, step 1 tests as required for hand disinfectants can be used.

\(^{13}\) These tests will be adapted for hand disinfectants.
Currently, there are no European phase 2, step 2 standard tests available for other skin disinfectants. However, test protocols may be designed by adapting existing standards (e.g. CEN methods involving volunteers) in a way that mirrors the respective application.

Newly designed test protocols should be timely discussed with and agreed by the CA before tests are carried out.

Deviations from the existing/future standards should always be mentioned and justified.

For an overview of available tests see Appendices 2 and 4.

**Test organisms**

Disinfectants for other skin and scalp should be at least sufficiently effective against bacteria and yeasts. Efficacy tests with these organisms should always be provided.

For phase 2, step 1 tests the standard organisms of these tests should be tested.

For phase 2, step 2 tests either the standard organisms of these tests can be tested or the normal occurring micro flora in volunteer tests.

For all other groups of organisms tests only have to be provided when activity against those specific organisms is claimed.

Justification for the used test organisms should be provided.

The test organisms used in efficacy tests are normally stated in the applicable standard test methods. An overview of reference test organisms is given in Appendix 3.

**Test conditions**

It is important that the tests are performed using the same contact time as claimed on the label. The claimed contact time has to be a realistic value and should be justified for the use.

Phase 2, step 1 and phase 2, step 2 carrier tests should be carried out with BSA as soiling (interfering substances) for clean or dirty conditions depending on the intended use. Simulated-use studies with volunteers are in general performed without additional soiling.

The soiling needed for clean and dirty conditions can be found in the relevant EN tests (see EN 14885, medical area) and referenced in Appendix 4.

5.4.1.3.2 Acceptance criteria

A product in PT1 will be assessed to be sufficiently effective if the required laboratory and simulated-use tests have been carried out (using the required test organisms and test conditions), and when the pass criteria for the tests have been met.

Where pass criteria are available in the standard test, these should be met. For PT1 products the required log_{10} reductions are referenced in Appendix 4 or EN14885.

Deviations from the pass criteria are possible, but must be justified in the application. The CA will evaluate any justification on a case-by-case basis, consulting the other CAs where appropriate, and decide whether it is acceptable or not.
5.4.2 PT2 Disinfectants and algaecides not intended for direct application to humans or animals

5.4.2.1 Introduction

Product type 2\(^{14}\) contains disinfectants and algaecides not intended for direct application to humans or animals. This includes *inter alia*:

- products used for the disinfection of surfaces, materials, equipment and furniture which are not used for direct contact with food or feeding stuffs;
- usage areas such as swimming pools, aquariums, bathing and other waters; air-conditioning systems; and walls and floors in private, public, and industrial areas and in other areas for professional activities;
- products used for disinfection of air\(^{15}\), water not used for human or animal consumption, chemical toilets, waste water, hospital waste and soil;
- products used as algaecides for treatment of swimming pools, aquariums and other waters and for remedial treatment of construction materials;
- products used to be incorporated in textiles, tissues, masks, paints and other articles or materials with the purpose of producing treated articles with disinfecting properties.

The data requirements (test standards and test organisms) and assessment criteria for the most common uses are specified below. Detailed but non-exhaustive lists of the most relevant product applications and uses of disinfectants within PT2, together with the relevant test methodologies are given in the claims matrices which are a set of tables linked to this guidance document (see Appendix 1 for more information).

All of the possible uses in this PT cannot be covered in the matrices. For other, less common, uses and claims that are not specifically mentioned, there is often no international standard tests available. Where this is the case, the applicant should provide tests that show the efficacy of the product and a justification for the use of these tests. The assessment of these products will be based on expert judgement and will be handled case-by-case.

5.4.2.2 Data requirements

There are some general data requirements that apply to all uses in PT2, and these are described below. There are also specific data requirements that apply to different types of use, and these are described in the sections covering those uses.

The intended uses of the disinfectant determine which tests will be required to support the product. Tests that most closely reproduce the practical application conditions should be selected.

In general it is not known which organisms are present on a surface or matrix to be disinfected. Therefore a disinfectant must have a broad spectrum of activity, in order to control all of the organisms that may be present.

\(^{14}\) This includes biostatic products.

\(^{15}\) This is taken to mean disinfection of air itself. Disinfectants sprayed or vaporised into the air (e.g. room disinfection by vaporised biocide) are generally for the purpose of disinfecting surfaces and not the air itself. Disinfectants for air conditioning systems disinfect the surfaces in these systems, not the air coming out of it.
5.4.2.2.1 Use in health care

For general applications in the medical sector, the products should be at least sufficiently effective against bacteria and yeasts (which are responsible for most common nosocomial infections). Additionally, efficacy against other organisms can be claimed.

Products intended to disinfect surfaces that are likely to come into contact with the patient and/or the medical staff and surfaces which are frequently touched by different people a leading to the transmission of microorganisms to the patient, must be tested with a contact time of maximum of 5 minutes. The same applies when the contact time of the product must be limited for practical reasons. Products for other surfaces than those stated above, may be tested with a contact time of a maximum of 60 minutes.

5.4.2.2 Tuberculosis departments

If the product is to be used in tuberculosis departments, the product should be efficacious as a general disinfectant used in health care (efficacy against bacteria and yeast), but tuberculocidal activity or mycobactericidal activity must also be demonstrated.

5.4.2.2.3 Cleanrooms

Products to be used in cleanrooms only differ in the data requirements for the interfering substance to be used in the tests. As an exception to the rule, products to be used in cleanrooms do not require additional soiling in the test. A cleanroom has a controlled level of contamination that is specified by the number of particles per cubic metre at a specified particle size. The soiling level in cleanrooms is so low that even testing under clean conditions for the EN tests is still over-dosing of soiling compared to cleanrooms. For these uses the high load of test organisms can be seen as soiling. Tests without soiling will only be accepted when the label states the specific use in clean rooms which are classified according to ISO 14644-1 in class 1 to 8 or according to GMP EU classification in Grade A to C.

5.4.2.2.4 Products against viruses

Products against viruses must be effective against viruses with and without an “envelope” (protein or lipid mantle). Products can claim virucidal efficacy if efficacy against non-enveloped viruses has been proven. Such products can be regarded as efficacious against enveloped and non-enveloped viruses.

The Phase 2 step 2 virus test described in EN 14476, should be used for testing products against viruses used in domestic areas. For testing products used in veterinary hospitals either EN 14476 or EN 14675 can be used. For products used in the medical area, a phase 2 step 2 test is under development, see prEN 16777.

5.4.2.3 Disinfectants for hard surfaces (in PT2)

5.4.2.3.1 Introduction

Biocides can be used to disinfect hard surfaces in areas such as hospitals (including veterinary hospitals, dental facilities etc.), industry, institutions or private homes. These surfaces may be tables, floors, walls, the outsides of machinery and hard furniture, etc. Products are often wiped or sprayed onto the surfaces and may be washed or wiped off after a certain contact time.

The testing requirements for some specific uses of hard surface disinfectants are discussed in separate sections, for example, toilets, room disinfection with vaporised biocide, immersion of equipment into the product, etc. As the areas of use can be as diverse as
private homes to operating theatres, the test requirements might vary depending on the area of use.

5.4.2.3.2 Data requirements

See general data requirements for PT2 (see sections 1.5 and 3.2 of this Guidance). A detailed, but non-exhaustive list of the most relevant product applications and uses of hard surface disinfectants and the required test methodologies are given in Claims Matrix PT2, table for “Hard surfaces”: the claims matrices are a set of tables linked to this guidance document (see Appendix 1 for more information and also Appendix 4).

Test methods

For efficacy testing of hard surface disinfectants, the tiered approach as described in section 5.4.0.4.1 of this Guidance is preferred.

The following tests are normally required for a hard surface disinfectant:

- a quantitative suspension test (phase 2, step 1);
- and a quantitative surface test (phase 2, step 2),

both simulating practical conditions appropriate to its intended use (temperature, soiling, different surfaces, contact time, etc.).

Tests in phase 3 are optional, as no validated test methods are available yet. Several methods for testing the efficacy of hard surface disinfectants are available. Appendices 2 and 4 give a list of recommended test methods.

The following documents are recommended for surface disinfection:

- EN 14885: gives an overview of which EN phase 2, step 1 and step 2 tests to use for different uses,

if CEN standards are not relevant or available for the use or organisms claimed the following documents are recommended if appropriately reflecting the application:

- OECD guidance for the testing of chemicals: Quantitative method for evaluating activity of microbiocides used on hard non-porous surfaces (these are surface tests which would be considered phase 2, step 2 tests),

The use of the specified tests is strongly recommended where they are relevant and appropriate. Where the tests are not appropriate to the product, other tests can be used, although a justification for the relevance of the tests used should also be provided.

It is preferred that tests should be selected that correspond to the use area of the product (e.g. tests from medical areas for use in hospitals and tests for industrial areas for use in cosmetic industry). Where the product is intended for use in several areas it is acceptable to perform the tests specified for only one of the areas, as long as the test with the highest/most stringent pass criteria is used. In some cases where the worse case cannot be clearly identified all areas must be tested.

Currently validated surface tests with and without mechanical action are available (EN and OECD). Validated surface tests with mechanical action have been developed, and should be used for products that are intended to be used with mechanical action (EN 16615).

Where specific conditions apply for a field of use, such as high/low level soiling, high/low temperatures, relevant contact times etc. (see section 5.4.0.4.4 of this Guidance), these conditions should be included in the efficacy testing.
**Disinfectant towelettes/wipes**

For disinfectant wipes, the phase 2, step 1 tests should be done preferably with the liquid extracted from the wipes, or if difficult to extract, use the liquid as it is before it is added to the wipes. Phase 2, step 2 tests should be tests with mechanical action. These tests are available for bacteria and yeasts. For testing other organisms surface tests can be done with liquid extracted from the wipes (not the original liquid), with a justification of the volume that is applied per square centimetre. In addition, a test must be performed that shows that either the wipe will still disinfect after the wipe dries out or that the wipe stays wet long enough to disinfect according to the claim. In addition, the use directions can address these issues, for instance, stating on the label that only wet wipes are efficacious, defining the surface area each wipe can disinfect (e.g. 0.5 m²), or giving expiry dates for re-sealable packages.

**Test organisms**

The test organisms used in efficacy tests are normally stated in the applicable standard test methods. For use in veterinary health care the target organisms in the test for the veterinary area (PT3) should be used.

If standard tests are not used (there will normally need to be a justification for this), the test organisms used to support a general claim should be demonstrated to be equivalent to the reference test organisms given in Appendix 3.

Tests with test organisms other than those mentioned in Appendix 3 are acceptable, if adequate scientific evidence is submitted on which the relevance of the test organism to the field of use can be judged.

Also see the general data requirements PT2 for specific claims and minimum requirements in health care.

**5.4.2.3.3 Acceptance criteria**

A product will be assessed to be sufficiently effective if the required laboratory and, where relevant, field tests have been performed (using the required test organisms and test conditions), and when the pass criteria for the tests have been met.

Where pass criteria are available in the standard test these should be met.

If the test does not provide these criteria, the general criteria referenced in Appendix 4 or EN 14885 can be taken as guidance for the level of reduction required. Deviations from the pass criteria are possible, but must be justified in the application. The CA will evaluate any justification on a case-by-case basis, consulting the other CAs where appropriate, and decide whether it is acceptable or not.

**5.4.2.4 Soft furnishings**

**5.4.2.4.1 Introduction**

Disinfectants for use on soft furnishings are intended to be used on fabrics in the home, institutional environment, healthcare and healthcare facilities. These can be used to treat porous soft surfaces such as curtains, sofas, upholstery, mattresses and carpets. The products are often sprayed onto the surfaces.

**5.4.2.4.2 Data requirements**

See general data requirements in sections 1.5 and 3.2 of this Guidance. A detailed, but non-exhaustive list of the most relevant product applications and uses of soft furnishing disinfectants and the required test methodology is given in Claims Matrix PT2, table for “Soft
furnishings”: the claims matrices are a set of tables linked to this guidance document (see Appendix 1 for more information).

**Test methods**

For efficacy testing of surface disinfectants for use on soft furnishings the tiered approach as described in section 5.4.0.4.1 of this Guidance is preferred.

The following tests are normally required for a surface disinfectant:

- a quantitative suspension test (phase 2, step 1);
- and a quantitative surface test (phase 2, step 2);

both simulating practical conditions appropriate to its intended use (temperature, soiling, different surfaces, contact time, etc.).

Tests in phase 3 are optional as no validated test methods are available yet.

Where possible, the phase 2, step 1 test should be selected from EN 14885 from the table that corresponds to the use area of the product (e.g. test from medical area for use in hospitals and test for domestic areas for use in private homes).

The phase 2, step 2 surface carrier test can be derived from adaptation of CEN TC 216 surface tests. Instead of a hard surface carrier, carriers could be made of suitable fabric types. ISO 20743 can also be used, or other quantitative methods including textile as carrier. EN16616 is not relevant since this is done in washing machines.

**Test organisms**

The same test organisms as given for hard surfaces should be tested. See section 5.4.2.3.2 test organisms, of this Guidance and Appendix 3.

**5.4.2.4.3 Acceptance criteria**

A product will be assessed to be sufficiently effective if the required laboratory and, where relevant, field tests have been performed (using the required test organisms and test conditions), and when the pass criteria for the tests have been met.

Where pass criteria are available in the standard tests these should be met.

If the test does not provide these criteria, the general criteria referenced in Appendix 4 can be taken as guidance for the level of reduction required. Deviations from the pass criteria are possible, but must be justified in the application. The CA will evaluate any justification on a case-by-case basis, consulting the other CAs where appropriate, and decide whether it is acceptable or not.

**5.4.2.5 Room disinfection with vaporised biocide**

**5.4.2.5.1 Introduction**

Room disinfection involves the reduction and inactivation of micro-organisms on the surfaces of the walls, floor and ceiling of the room, as well as on external surfaces of the furniture and equipment present in the treated room. The product is applied by airborne diffusion of an aerosol, a smoke, a vapour or a gas. The technical characteristics of the diffuser equipment play a central role, ensuring a homogeneous distribution of the biocide product in the volume of the room and reaching all surfaces (including ceilings and the undersides of horizontal surfaces), therefore the diffuser equipment contributes in a decisive way to the efficacy of the product. Manual spraying is not covered in this section, but under hard surface disinfection (see section 5.4.2.3 of this Guidance).
Room disinfection may not disinfect the inside parts of furniture, and will not disinfect the air itself, so these uses are not considered in this section. Room disinfection is therefore closely related to surface disinfection without mechanical action. As this causes complications in cases of organic contamination, cleaning of surfaces is necessary prior to room disinfection.

**Process**

The application of the product consists of four phases:

1. the preparation phase (required depending on type of active substance and application procedure), during which the environmental conditions (relative humidity, temperature) are modified to an optimal level for the product;
2. the conditioning phase, during which the product is diffused into the room, in order to reach the effective concentration;
3. the disinfection phase, which corresponds to the contact time required to obtain the expected level of efficacy;
4. the terminal phase, which includes aeration of the room to remove any disinfectant present in the air, or other procedures for inactivation of the active substance, before access of people or animals into the room can be permitted (see Figure 2 below).

**Figure 2: The various phases of a cycle of disinfection of an automatic process**

![Diagram of disinfection cycle](image)

**Phases of the cycle:**

1: preconditioning (optional)
2: diffusion
3: phase of contact
4: aeration

Particular attention must be given to the dispersal time and contact time. The dispersal time is the time necessary to reach a target concentration of the product in the air and on the surfaces to be disinfected in a given volume, while the contact time is the time necessary to reach the expected efficacy.
Note: the various phases of the cycle presented are theoretical and can be adapted according to the process. The maintenance of a concentration of biocide in the atmosphere may be achieved by the regular introduction of additional biocide during the contact phase.

5.4.2.5.2 Data requirements

Test methods

Airborne disinfection differs from direct application of liquids to surfaces. Therefore the EN phase 2, step 2 standards for surface disinfection are not applicable for room disinfection. The tiered approach is still possible, however, by using different test methods.

The following tests are normally required for a room disinfectants:

- when applicable, a quantitative suspension test (phase 2, step 1);
- semi-field trial such as European standard based on NF T 72-281 (EN standard in preparation) for disinfection using airborne application (phase 2, step 2).

The CEN phase 2, step 1 tests are suitable as suspension tests under clean or dirty conditions, although only applicable for products that can be tested in suspension (e.g. not for gasses). These tests are not sufficient on their own, and should be combined with a semi-field trial for disinfection using airborne application. Where it is not possible to test the product in a suspension test, the semi-field trial will be sufficient.

NF T 72-281 was developed by AFNOR (the French standardisation body) in 1986, and updated in 2009. This standard was taken as a start to develop a new EN standard on airborne disinfection of surfaces (decision taken within the framework of CEN TC 216 in November 2012). This semi-field method evaluates the efficacy of disinfectants when vaporised in a room (automatic diffusion process) or when sprayed in the direction of a surface (manual application). Only application by vaporisation is discussed in this section. Once this method has been finalised and adopted at European level, any method variations should be taken into account.

Basic principles of room disinfection

Inert and dry carriers infected with a known number of micro-organisms (bacteria, fungi, yeasts, mycobacteria, spores and viruses including bacteriophages) are placed in a room of defined volume, temperature and relative humidity. The size of the test room should be relevant to the claims for the product. The carriers used are often stainless steel, but other relevant (generally non-porous) materials can also be used, such as glass or plastic.

When the disinfection of textiles (curtains etc.) and other materials (e.g. wallpaper, filters in flow cabinets) is claimed, appropriate carriers should be used to demonstrate efficacy.

The inoculated carriers must be placed in a vertical position with inocula facing away from the diffuser. Their distance to the diffuser depends on the room dimensions (for instance: see Appendix B of NF T 72-281). The test method defines obligatory test conditions for parameters that may influence the success of the disinfection.

This test includes the use of milk in order to maintain viability of the micro-organisms on the carriers during the test. Depending on the area of use, suitable interfering substances should be tested (e.g. blood for use in hospitals).

Similar carriers are placed in a second room nearby which is not treated with diffused product, to act as controls.

Additional tests can be performed to simulate specific conditions that are encountered in the practice and to fit with label instructions. In this case, all experimental conditions should be
described clearly in the test reports. The standard lists the information that must be included in the final report.

**Diffuser**

As mentioned earlier, the disinfection efficacy is closely related to the technical characteristics of the diffuser. Section 5 “intended uses and efficacy” of the *Guidance on information requirements for biocides*. (Volume II Efficacy, part A) requires applicants to take into account the technical equipment used, together with the product to be authorised.

A detailed description of the equipment and its characteristics must be provided in sufficient detail to distinguish it from other equipment:

- equipment name and model;
- diffusion principles (e.g. fogging, vapour, fumigation) and particles size distribution of aerosols or powder;
- description of the diffusion performance of the equipment (e.g. volume to disinfect, diffusion speed);
- description of the ambient conditions (e.g. humidity, temperature) in which the process can be used;
- diffusion time for a specific volume;
- precautions for over and under-dosing.

The product authorisation will only be granted for use with the equipment described in the application. After authorisation, any modification to the equipment should be validated and reported to the CA for evaluation.

For major modifications that can affect the efficacy (e.g. pipe, pump, nozzles), it should be demonstrated that the efficacy of the process has not been affected (e.g. by a new study on the most resistant organism).

For minor modifications that do not change the efficacy of the process, only a notification of the modifications to the equipment must be provided.

**Contact time**

As room disinfection may necessitate a long period of treatment, the contact time to be tested is not defined. The testing should demonstrate efficacy at a contact time proposed for the intended use. This should be relevant to practical use and depends on substance concentration, volume of room, power of the diffuser equipment, etc... All of these parameters should be stated on the product label or in a technical information sheet.

**Test organisms**

Since room diffusion is used to disinfect hard (and soft) surfaces, the same organisms should be tested as for hard surface disinfection (section 5.4.2.3 of this Guidance). Appendix 3 contains a table of reference test organisms.

The general data requirements for PT2 for specific claims and minimal requirements in health care also apply for room disinfection with vaporised biocide.

**5.4.2.5.3 Acceptance criteria**

A product will be assessed to be sufficiently effective if the required laboratory and, (semi) field tests have been performed (using the required test organisms and test conditions), and when the pass criteria for the tests have been met.

Where pass criteria are available in the standard tests these should be met.
If the test does not provide these criteria, the general criteria referenced in Appendix 4 can be taken as guidance for the level of reduction required. Deviations from the pass criteria are possible, but must be justified in the application. The CA will evaluate any justification on a case-by-case basis, consulting the other CAs where appropriate, and decide whether it is acceptable or not.

**5.4.2.5.4 Notes**

**Limitations**

Any limitations of the procedure should be specified in the application.

Literature has shown that disinfection by vaporised biocide may not be as effective on wet surfaces (lower concentration of the product) or inside closed cupboards and closets (where the vapour cannot penetrate). Therefore carriers should be tested under these conditions, and if efficacy is not proven, the label instructions should provide appropriate information (such as stating that cupboard doors should be opened, surfaces should be dried and wet areas (such as sinks and toilet bowls) should be disinfected with suitable alternative products.

Other factors which may influence the efficacy of the process in the practical use such as the equipment, furniture, special structures (e.g. bumps on the walls) or special materials (copper in hydrogen peroxide procedures), including environmental conditions (e.g. temperature, relative humidity) which may affect the success of the disinfection, have also to be considered. The conditions of a sufficient vaporisation should also be specified.

**5.4.2.6 Swimming pools, spas and hot tubs**

**5.4.2.6.1 Introduction**

Disinfectants are used to treat water in swimming pools, spas and hot tubs. These may be public pools (which may be used by many people daily) or household pools or tubs (which might be used only occasionally). An intermediate situation consists of facilities in hotels, housing complexes or sports clubs, where the bather load may be lower than in a fully public facility, but still high compared to private, domestic facilities.

Disinfectant products can be added to a pool continuously, intermittently, by shock dosing or through generation *in situ*. Large public facilities may have dedicated staff to maintain the pool using automated control systems, whereas smaller pools may be treated using manual methods by janitorial staff. Private pools may be treated by individual householders, supplemented in some cases by professional pool treatment personnel. Disinfection is only one aspect of pool maintenance and other activities, such as ensuring the correct pH and the removal of pollutants by oxidation, flocculation and filtration, are essential to ensure adequate water quality.

The principal purpose of the use of disinfectants is to treat the water to prevent the water-borne transmission of pathogens between pool users. Supplementary purposes are to ensure the aesthetic quality of a pool (by ensuring that algae do not result in turbid water or unsightly and slippery microbial growth on pool surfaces, such as the floor and walls of the pool) and to prevent microbial slime and biofilm formation in pipework and related equipment.

This section only deals with disinfection of the pool water and the pipework and related equipment containing pool water. The disinfection of hard surfaces surrounding the pool is covered in section 5.4.2.3 of this Guidance.
5.4.2.6.2 Data requirements

See PT2 general data requirements in sections 1.5 and 3.2 of this Guidance.

A detailed, non-exhaustive list of the most relevant applications and of appropriate test methodology is given in Claims Matrix PT2, table for “Swimming pools”: the claims matrices are a set of tables linked to this guidance document (see Appendix 1 for more information).

Test methods

For efficacy testing of pool disinfectants the tiered approach as described in section 5.4.0.4.1 of this Guidance is preferred.

The following tests are normally required for a pool disinfectant following a tiered approach:

- a quantitative suspension test (phase 2, step 1);
- simulated-use tests with pool water or a surface test (phase 2, step 2)*;
- and a field test (phase 3)**;

all simulating practical conditions appropriate to its intended use (temperature, contact time, soiling/bather load etc.).

* A phase 2, step 2 test may be appropriate in cases where a product has a specific use in surface disinfection. Otherwise, a simulated use test is appropriate for products intended to disinfect the water in a pool or spa.

** In some cases the field trial can be waived. The OECD guidance document (described below) is based on experience with hypochlorous acid/hypochlorite. Therefore, it is acceptable that for products based on hypochlorous acid/ hypochlorite the field test is waived and only laboratory test data are provided. In some other cases, waiving the phase 3 test can also be justified.

The OECD “Guidance Document for Demonstrating Efficacy of Pool and Spa Disinfectants in Laboratory and Field testing” (OECD Series of Testing and Assessment No 170, version dated 08 October 2012) describes laboratory and field test methods, conditions and criteria needed to demonstrate efficacy of a pool disinfectant. The protocol for field tests should be agreed between the applicant and CA before a field test is initiated.

For products that are used for specific purposes such as disinfecting pipework, filters and filter media, it may be more appropriate to test using the EN 14885 methods for the disinfection of surfaces in institutional applications.

Test organisms

Besides bacteria and viruses, protozoa can also be of importance in swimming pools. Fungi may pose a health hazard on wet surfaces surrounding the pool and can cause slime build up in pipework. OECD guidance lists the organisms that normally should be tested. Although algae and protozoa in pools are, in general only a problem when maintenance of the pool is not carried out properly, data against algae and/or protozoa should be provided where claims against these targets are made.

5.4.2.6.3 Acceptance criteria

A product will be assessed to be sufficiently effective if the required laboratory and simulated use tests and where relevant, field tests have been performed (using the required test organisms and test conditions) and the pass criteria for the tests have been met.

When pass criteria are available in the standard tests these should be met.
The OECD guidance document sets out criteria for laboratory and field tests. However, it may be noted that there is a current OECD project review underway to look at criteria for laboratory and field tests.

Where these criteria are not met, the applicant can provide a justification as to why the product should still be considered acceptable. However, the CA will evaluate any justifications on a case-by-case basis, consulting the other CAs as necessary, and will decide whether it is acceptable or not.

The OECD guidance document contains more details on factors to be considered.

5.4.2.7 Toilets

5.4.2.7.1 Introduction

Biocides can be used to disinfect toilet bowl surfaces in diverse environments including, hospitals, industry, institutions or households. Toilet bowl biocides are available in a wide variety of forms, such as liquids, foams, powders, gels, pastes and tablets.

These products are often applied via pouring around the inside rim of the toilet bowl surfaces with the area scrubbed after a minimum contact time.

Other products are applied in the toilet permanently. They can be attached over the rim of the toilet bowl, stuck directly onto the side of the toilet bowl, placed directly in the cistern (water reservoir), or attached by other means. These products are normally discharged when the toilet is flushed.

Hard surfaces on the inside of toilets are covered by this section. Surfaces on the outside and toilet seats, lids etc. are covered by section 5.4.2.3 “hard surfaces” of this Guidance.

The use of biocides in chemical toilets, most commonly found on airplanes, trains, and in portable toilets, is not covered in this section, (see section 5.4.2.13 of this Guidance).

5.4.2.7.2 Data requirements

See PT2 general data requirements in 5.4.0.5 and 5.4.2.2.

A detailed, non-exhaustive list of the most relevant applications and of appropriate test methodology is given in Claims Matrix PT2, table for "Toilet bowls": the claims matrices are a set of tables linked to this guidance document (see Appendix 1 for more information).

Test methods

For efficacy testing of toilet disinfectants the tiered approach as described in section 5.4.0.4.1 of this Guidance is preferred.

The following tests are normally required for a hard surface disinfectant:

- a quantitative suspension test (phase 2, step 1);
- and a quantitative surface test (phase 2, step 2);

both simulating practical conditions appropriate to its intended use (temperature, soiling, contact time, etc.).

Several test methods for quantitative suspension and surface tests are available.

Appendix 2 gives a list of recommended test methods. The following documents are recommended for surface disinfection:

- EN 14885: gives an overview of what EN phase 2, step 1 and step 2 test to use for different uses,
if CEN standards are not relevant or available for the use or organisms claimed the following documents are recommended if appropriately reflecting the application:

- OECD guidance for the testing of biocides: Quantitative method for evaluating activity of microbiocides used on hard non-porous surfaces (these are surface tests which would be considered phase 2, step 2 tests).

The use of the specified tests is strongly recommended where they are relevant and appropriate. Where tests are not appropriate to the product other tests can be used, although a justification for the relevance of the tests used should also be provided.

For products intended to be added to the water reservoir or hanging down from the rim of the bowl, the concentration of the product (or at least the active substance) in the water before, between and after flushing should be determined. This can be done by an analysis of the water under in-use conditions or, for products where all parameters are defined, by calculation. The laboratory efficacy tests should be performed using these concentrations.

Tests in phase 3 are optional.

When efficacy against biofilm is claimed a simulated-use test or field test has to be provided, next to a phase 2, step 1 test. See section 5.4.2.11 of this Guidance for test methods.

**Test organisms**

The same test organisms as for hard surfaces should be tested. See section 5.4.2.3.2 of this Guidance and Appendix 3.

Products will normally only target bacteria and, optionally, yeasts and viruses. Fungi and spores are usually not relevant in the toilet bowl but when efficacy is claimed testing is required.

**5.4.2.7.3 Acceptance criteria**

A product will be assessed to be sufficiently effective if the required laboratory and, where relevant, field tests have been performed (using the required test organisms and test conditions), and when the pass criteria for the tests have been met.

Where pass criteria are available in the standard tests these should be met.

If the test does not provide these criteria, the general criteria referenced in Appendix 4 can be taken as guidance for the level of reduction required. Deviations from the pass criteria are possible, but must be justified in the application. The CA will evaluate any justification on a case-by-case basis, consulting the other CAs where appropriate, and decide whether it is acceptable or not.

**5.4.2.8 Air-conditioning systems**

**5.4.2.8.1 Introduction**

Disinfection of air-conditioning systems is similar to hard surface disinfection since only the surfaces in the system are disinfected and not the air itself. The difference with general surface disinfection is that the surfaces are mostly hidden inside the system and cannot be reached easily without taking it apart (for instance for air-conditioning systems in cars, dismantling the system would not be desirable).

In general, disinfectants for air-conditioning systems are applied by airborne diffusion of an aerosol, a smoke, a vapour or a gas. The biocide is applied to an operating air-conditioning system at the inlet of the system. This way the biocide is sucked into and passes through the whole system.
Preservation of cooling liquids is not covered under PT2 but rather within PT11 (preservatives for liquid cooling and processing systems).

**5.4.2.8.2 Data requirements**

For products that are applied by airborne diffusion of an aerosol, smoke, vapour or gas the same test methods and test organisms should be used as for room disinfection. Therefore, the same data requirements as for section 5.4.2.5 of this Guidance (Room disinfection with vaporised biocide) are applicable here.

The following tests are normally required for a disinfectant for air-conditioning systems:

- when applicable, a quantitative suspension test (phase 2, step 1);
- semi-field trial such as NF T 72-281 for disinfection using airborne application (phase 2, step 2).

See section 5.4.2.5 of this Guidance for specifications.

In the semi-field test the carriers inoculated with test organisms are placed in the air-conditioning system at the beginning and at the end of the system. When it is not possible to put carriers in the system they should be placed between the biocide application site and the inlet of the system and at the end of the system, in the out-flowing air. If carriers at both sides fulfil the criteria it can be assumed that the surfaces in between are also disinfected sufficiently.

For products that are applied by manual spray, the test methods and test organisms used for hard surface disinfection should be employed. See section 5.4.2.3 of this Guidance (Hard surface disinfection) for data requirements.

In addition to these data, the applicant should provide a justification that the spray apparatus is capable of reaching all (hidden) surfaces of the air conditioning system.

**5.4.2.8.3 Acceptance criteria**

A product will be assessed to be sufficiently effective if the required laboratory and, where relevant, field tests have been performed (using the required test organisms and test conditions), and when the pass criteria for the tests have been met.

The same pass criteria can be used as for other surface disinfection (section 5.4.2.3.3 of this Guidance). The criteria referenced in Appendix 4 can be used as guidance for what level of log$_{10}$ reduction is normally required. Deviations from these are possible, but have to be justified in the application.

**5.4.2.9 Equipment disinfection by immersion**

**5.4.2.9.1 Introduction**

Although instrument or equipment disinfection can be considered equal to hard surface disinfection, it differs from the intended use in section 5.4.2.3 of this Guidance because it is mainly applied by immersion of the equipment or instruments in the biocide solution or by filling equipment with the solution (disinfection of inner surfaces). The products are intended for equipment used, for example, in health care facilities, laboratories and industry. The requirements for products to be used for CIP are not included in this section and can be found in section 5.4.4.3 of this Guidance.

Some of the products used for disinfection of medical instruments, which are to be used specifically for diagnostic and/or therapeutic purposes for human beings, do not fall under the scope of the BPR. Disinfectants that are specifically used for the disinfection of medical
devices or a group of medical devices (anaesthetic equipment, endoscopes, surgical instruments, incubators) are covered under the Medical Device Directive 93/42/EEC. However, some disinfectants have a broader claim, for example, disinfection of instruments and surfaces. They are so called ‘dual use products’ as their distinct claims are covered by more than one legislative instrument. The BPR states that such biocidal products should comply, in addition to the requirements laid down in the BPR with the relevant essential requirements set out in Annex I to Council Directive 90/385/EEC of 20 June 1990 on the approximation of the laws of the Member States relating to active implantable medical devices, Council Directive 93/42/EEC of 14 June 1993 concerning medical devices and Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on in vitro diagnostic medical devices.

5.4.2.9.2 Data requirements

Test methods

For efficacy testing of equipment disinfectants the tiered approach as described in section 5.4.0.4.1 of this Guidance is preferred.

The following tests are normally required for an instrument disinfectant:

- a quantitative suspension test (phase 2, step 1);
- and a quantitative carrier test (phase 2, step 2);

both simulating practical conditions appropriate to its intended use (temperature, soiling, different surfaces, contact time, etc.).

Methods for testing efficacy of equipment or instrument disinfectants are available.

Appendix 2 gives a list of recommended test methods. The following document is recommended for instrument disinfection:

- EN 14885: gives an overview of which EN phase 2, step 1 and step 2 test to use for different uses.

The use of the specified tests is strongly recommended where they are relevant and appropriate.

For use in industry and institutional areas, no specific tests for instrument disinfection are given in EN14885. Nevertheless, phase 2, step 1 suspension tests from the industry and institutional areas can be used, by employing area specific soiling. For phase 2, step 2 tests, the instrument tests for medical areas are most appropriate. Soiling specific to the area of intended use should be employed.

Test organisms

For general disinfection of medical (including dental and veterinary) equipment, instruments, and equipment and other instruments which are used in contact with skin or mucous membranes (e.g. instruments for pedicure), efficacy against bacteria, yeasts, viruses and fungal spores must be demonstrated. For instruments and equipment used in laboratory and industry the test organisms specified for hard surfaces should be tested.

See section 5.4.2.3.2 of this Guidance and Appendix 3.

5.4.2.9.3 Acceptance criteria

A product will be assessed to be sufficiently effective if the required laboratory and, where relevant, field tests have been performed (using the required test organisms and test conditions), and when the pass criteria for the tests have been met.
Where pass criteria are available in the standard tests these should be met.

If the test does not provide these criteria, the general criteria referenced in Appendix 4 can be taken as guidance for the level of reduction required. Deviations from the pass criteria are possible, but must be justified in the application. The CA will evaluate any justification on a case-by-case basis, consulting the other CAs where appropriate, and decide whether it is acceptable or not.

5.4.2.10 Textiles

5.4.2.10.1 Introduction

Biocides can be used to treat textiles and fabrics in hospitals, health care facilities, industry, institutions or private homes, when relevant micro-organisms (pathogenic, spoiling) in the textiles have to be reduced. These products can be in the form of laundry products which combine detergent and biocide or can be specialised products in the form of laundry additives which are added to the wash cycle or as finishing products (e.g. fabric softeners) which are added in the last rinsing step or as pre-treatment.

Typically contaminated clothes, linen or other washable textiles are treated in an appropriate washing machine. The biocide is added in concentrated form and diluted in the machine with water according to the specification of the manufacturer to get a defined concentration in the machine. The automated chemical-thermal process normally comprises an (optional) initial pre-treatment step for heavily soiled laundry, followed by the main washing step (at a defined temperature and defined contact time) and 3 to 4 rinsing steps with cold water.

In some cases laundry can be treated through a hand-wash process in diluted biocide, which can take the form of a pre-soak (after which, machine washing is used), a hand wash only, or through soaking to disinfect textiles before they are destroyed (e.g. in an infectious disease outbreak situation).

Biocidal laundry products, either as combined biocide/detergent/conditioner or as special additives, are available for either targeted pre-treatment of contaminated articles or for whole-wash use.

5.4.2.10.2 Data requirements

See PT2 general data requirements in sections 1.5 and 3.2 of this Guidance. A detailed, non-exhaustive list of the most relevant applications and of appropriate test methodology for is given in Claims Matrix PT2, table for “Laundry products”: the claims matrices are a set of tables linked to this guidance document (see Appendix 1 for more information).

Test methods

For efficacy testing of textile disinfectants the tiered approach as described in section 5.4.0.4.1 of this Guidance is preferred.

The following tests are normally required for a textile disinfectant:

- a quantitative suspension test (phase 2, step 1),
- a quantitative carrier test involving carriers made of test fabric (cotton, polyester) (phase 2, step 2),

Both should simulate practical conditions relevant to its intended use (concentration of the product, temperature, soiling, different fabrics, contact time, etc.). This includes the application of a normal washing procedure (including detergent) as a control.

Currently, the following types of test are available:
• phase 2, step 1 suspension tests as described in EN 14885,
• phase 2, step 2 tests involving
  o a full-scale laundry machine test (EN 16616)
  o for products not intended to be used in washing machines, small scale laboratory setting (e.g. for pre-soaking in a bucket) may be considered (e.g. ASTM E4206 or ASTM E2274).

In the phase 2, step 2 tests fabric is contaminated with test organisms and then exposed to the disinfectant.

The EN tests are strongly recommended where available and appropriate.

**Test organisms**

Textile disinfection products should be at least sufficiently effective against bacteria and yeasts. Efficacy tests with these organisms should always be provided.

For all other groups of organisms, data only have to be provided when activity against those organisms is claimed.

When disinfection is done at high temperatures (>40 °C) relevant test organisms for these temperatures should be used as described in section 5.4.0.4.4 of this Guidance.

An overview of reference test organisms, also for high temperatures, is given in Appendix 3.

**Test conditions**

For products intended to be added to washing machines, information on the following in-use conditions should be provided:

• the concentration of the product in the water during disinfecting process (i.e. washing or rinsing). The water volume used can differ between wash and rinse cycle and different washing programmes, but also between washing machines;
• the water to textile ratio in the test is an important factor that should reflect the in-use conditions;
• the temperature during the disinfection process (high when added in wash process, low in rinse process);
• the contact time (differs between various washing programmes and washing machines).

The laboratory tests should be performed under these conditions. The conditions for effective disinfection can normally only be carried out in professional washing machines.

If the exact conditions cannot be met, for example, in household machines, reasonable worst case conditions must be tested.

Worst case conditions, e.g.:

• the lowest temperature;
• the highest volume of water (i.e. maximum dilution of the product);
• the shortest contact time;
• the maximum load of laundry (i.e. smallest water to textile ratio).

When phase 2, step 2 tests involving fabric test carriers are performed, both the micro-organisms remaining on the test carriers, those released into the washing liquid and those transferred to previously uncontaminated control carriers should be assessed.
Manual soaking or pre-soaking can be done at room temperature but for some intended uses the temperature might start high and will cool down during the contact time (e.g. where hot water is used, which cools naturally). This should also be taken into account in the tests.

**Soiling**

The interfering substance most appropriate for the in-use conditions should be used. For instance, blood for products used in the medical area and protein for products used in industry, institutional and domestic areas are recommended. The soiling on a domestic product for use in pre-soak (dirty clothes) will be very much higher than the soiling present for a post-wash rinse additive (clean clothes). For products used during pre-soak and wash, tests should be done under dirty conditions. For products used during post-wash rinse, tests should be done under clean conditions.

5.4.2.10.3 **Acceptance criteria**

A product will be assessed to be sufficiently effective if the required laboratory and, where relevant, field tests have been performed (using the required test organisms and test conditions), and when the pass criteria for the tests have been met. When the product is intended to be used in combination with or directly after a detergent, a clear effect of the disinfectant alone should be demonstrated. There should be a significant difference (+log 2) between disinfectant+ detergent and the detergent alone.

EN and VAH tests provide pass criteria.

No acceptance criteria have been specified in the ASTM standards for laundry (ASTM E 2406-04 or ASTM E 2274-09).

If the test does not provide pass criteria, the general criteria referenced in Appendix 4 can be taken as guidance for the level of reduction required. Deviations from the pass criteria are possible, but must be justified in the application. The CA will evaluate any justification on a case-by-case basis, consulting the other CAs where appropriate, and decide whether it is acceptable or not.

5.4.2.11 **Biofilms**

5.4.2.11.1 **Introduction**

A biofilm is a complex aggregation of micro-organisms usually distinguished by the excretion of a protective and adhesive matrix attached to a solid surface in contact with a fluid. The matrix may incorporate other components derived from the environment.

Once the first cell succeeds in attaching to a surface and a biofilm starts to form, growth of the biofilm may become very fast, as subsequent free floating bacteria find it much easier to attach to the developing matrix.

Biofilms can grow in areas such as inside water tanks and the distribution pipelines of hospitals, hotels, industries and, in general, in any water systems which have temperatures and nutrients adequate for microbial growth.

The consequences of biofilm formation in a water system or facility may be severe depending on environmental conditions and any safety and performance requirements.

In healthcare facilities, biofilm contamination of medical equipment or water systems may increase the risk of nosocomial infections; in industrial facilities biofilm may cause microbial contamination of production (pharmaceuticals, cosmetics, etc.); in other situations biofilms may be responsible for significant reduction of the performance of water systems by obstructing normal flow or they may induce corrosion of the pipelines.
Several factors may contribute to biofilm formation, with important factors including the chemical composition and roughness characteristics of the pipe, tank or tube circuit.

Bacteria in biofilms are more resistant to disinfection than planktonic bacteria of the same species, as the presence of extracellular polymeric substances can act as a physical barrier to the biocide. This matrix may hamper biocidal penetration to the lower layers of the biofilm or may interact with the biocide and neutralize it. Additionally, the physiological state of the bacteria in the biofilm differs from bacteria in suspension, which can also influence the susceptibility of the bacteria to biocides. Complex communication systems are often also present that allow increased tolerance of members of the biofilm community to be initiated.

Two types of activities of biocides against biofilm can be identified:

1) Prevention of biofilm formation: the biocide acts on biofilm formation (i.e. in this case the biocide is present before the biofilm is formed and may affect the early adhesion of cells to the surface or the viability of the cells);

2) Biofilm disinfection (“curative”): the biocide acts on a mature biofilm (i.e. when the biofilm is already present on a surface and the biocide interacts with the biofilm-embedded cells, with a -cidal effect). Biocidal products of this type may also achieve detachment of the biofilm (possibly in conjunction with physical action).

In case where the biofilm is not removed as a result of the biocide treatment, it should be followed by mechanical removal of the biofilm.

Industry is increasingly developing new technologies for prevention, inactivation and/or detachment of biofilms and/or inactivation of biofilm embedded organisms, for example through the use of UV light, water ionization or impregnated or coated materials and new biocides which claim specific efficacy against biofilms.

5.4.2.11.2 Data requirements

There are currently no standard laboratory tests available to verify the efficacy of biocides against biofilms. As this is an area in which the science is developing rapidly, the information below should be considered as general guidance reflecting the state of knowledge at the time of writing this Guidance.

Tests to demonstrate the efficacy of disinfectants according to EN and OECD are based on simpler models than are found in biofilms. The available surface/carrier tests are not representative of biofilm models, as they do not consider the presence of extra cellular polymeric substances which act as a physical barrier to the biocide.

Other characteristics of the biofilm and biocidal product should be taken into account. For example, if biocide impregnated materials claim a preventive effect on biofilm formation, the prevention of biofilm formation should be demonstrated, taking into consideration the half-life of the impregnating substance which may differ depending on the material characteristics. The active substance may be released from the surface and/or may be inactivated by environmental factors.

A standard suspension test can only be used to confirm basic activity of the product against the claimed organisms in a tiered approach.

A suggested general approach could be:

1) a suspension test: any biocide claiming to act on biofilm, has to be first evaluated in standard suspension test (preferably EN);
2) a simulated use efficacy test to demonstrate the ability of the product to exert a controlling effect on the biofilm under either static condition or under flow conditions depending on the use pattern (claim). This controlling effect can be to destroy and detach, inhibit or prevent the formation of a biofilm;

3) a field trial, where the biofilm is formed under (simulated) use conditions.

These tests should be performed in sequence to obtain more complete information on the activity of the product on biofilm.

For biofilm disinfection (curative) a suspension test (as for (1) above) and suitable robust data from either a simulated use test (2) or field trial (3) should be performed. If there are no robust data from a simulated use test (2), a field test (3) is mandatory.

For biofilm prevention the approach is different to that for biofilm disinfection, as the biocide is present before the biofilm is formed and may affect the early adhesion of cells to the surface or the viability of the cells. In this case the suspension test (1) may not be useful since the product might not have a –cidal effect.

Test methods
Suspension tests
The first step in the tiered approach is a suspension test. The CEN phase 2, step 1 tests are suitable as suspension tests. This test is only applicable for products that can be tested in suspension and which have a –cidal effect.

Simulated use tests
Standard laboratory tests to verify the efficacy of biocides against biofilms are not currently available. Therefore, before performing a biofilm test, the methods should be agreed with the CA.

Applicants should provide a method following the principles in this guidance and based on scientific evidence. During development of the tests CAs of member states should be consulted to make sure that the tests are acceptable.

Biofilms can be formed and evaluated in static or flow conditions. The way the biofilm is formed has an effect on the susceptibility of the biofilm to biocides: biofilms formed under flow conditions are generally more resistant to biocides than biofilms formed under static conditions.

The conditions under which the biocidal products will have to operate should also be taken into account. Under static conditions the disinfectant operates without the aid of the removal effect of a fluid flow or shear stress. Under flow conditions the contact time might be shorter when shock dosing is used.

Static tests are less expensive and easier to standardise, but flow tests are generally closer to the real use scenarios.

In both cases, the reproducibility and repeatability of results over time should be ensured; so a method that allows a series of observations, rather than a single observation, should be employed.

Laboratory tests for evaluating the efficacy of biofilm disinfectants should emulate the critical factors of a real-world environment. In most instances, a biofilm will not be comprised of a single species and tests based on consortia relevant to the end use should be employed when simulating actual use.
In cases where only efficacy against biofilms formed under static conditions is claimed (e.g. use in tanks without flow) it is sufficient to only test against these biofilms.

Examples of methods for testing under flow and static conditions are described below, but other protocols are available in literature or may be under development.

**Static condition assay**

Standard laboratory tests to verify the efficacy of biocides against biofilms formed under static conditions are not currently available. However, literature describes several methods of how to create a biofilm in the laboratory under static conditions.

An example of a semi-quantitative method for biofilm evaluation is the microplate test, where a biofilm is formed in static conditions and the amount of biofilm can be quantified by spectrophotometric measurements. The amount of living cells in the biofilm before and after treatment can also be determined. In this case, the disinfectant operates without the aid of the removal effect of a fluid flow or shear stress.

A positive aspect of such an assay is that it is a low cost, easy-to-conduct test, that allows several replicates and/or the testing of several conditions (several biocide concentrations, more species, etc.) to be performed, which would provide the basis for a more accurate and closer-to-reality test.

This method consists of the formation of a biofilm by the species of interest on the bottom of 96 well plates (the material and coating of the plates should be specified); the disinfectant may be present before (preventive effect) or after (inhibition/removal effect) the biofilm is formed. The amount of biofilm (biomass) is quantified after staining of the adherent material and spectrophotometric measurement. Detecting chemicals such as ATP to measure bacterial viability may also be used.

**Flow condition assay**

Standard laboratory tests to verify the efficacy of biocides against biofilm formed under flow conditions are not currently available. However, systems to generate a standard biofilm have been developed by CEN (CEN ISO/TS 15883-5:2005 Annex F) and ASTM (ASTM E2196 and ASTM E2562). Using either of these reproducible biofilms, a method for the assessment of prevention and/or elimination of biofilm in terms of viable cells reduction and bacterial biomass reduction can be carried out.

The CEN method consists of the production of a standard *Pseudomonas aeruginosa* biofilm inside a Teflon tube, using a flowing system to simulate a real world situation.

ASTM E2196 and ASTM E2562 standards use biofilm rotating disc reactors, which are especially suited for high shear forces.

The biofilm is then treated with a disinfectant to evaluate the biocidal capacity to remove or to reduce the biofilm.

Other carrier types (e.g. silicon, steel, PVC, etc.) can be selected and used depending on the biofilm development system, and the experimental conditions can be adapted to compare the efficacy of different treatments in preventing biofilm formation.

A reference substance of known activity must be tested in parallel (e.g. chlorine dioxide, sodium hypochlorite).

**Field trials**

As for other situations in which biocides are used, only field tests (phase 3 tests) are fully representative of the activity of the biocide on biofilms, but these tests are difficult to
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standardise, and such tests should be complemented by laboratory suspension or simulated use tests, which have a higher degree of robustness and reproducibility.

A field trial should reproduce the in-use conditions of the worst-case situation of the intended uses.

Prevention and/or elimination of biofilm (in terms of viable cells reduction and bacterial biomass reduction) should be demonstrated by sampling before and after disinfection.

A field test can be waived if a suitably robust simulated use test, which adequately mimics the in-use conditions is provided. A robust test could for instance be a complex pipe system, in which natural biofilm formation takes place, either in combination with the addition of standard organisms or not.

**Test organisms**

The choice of micro-organisms for a test is relevant, since the use of only one organism per test is limiting and may not be fully representative of the real events leading to micro-organism aggregation (biofilms in settings where disinfectants are used, are normally multi-microbial, i.e. composed of several different species). Moreover, contaminants from environmental sources may be embedded in the biofilm matrix which may reduce the disinfectant’s efficacy.

Bacteria are not the only inhabitants of biofilms, as both fungi and algae may also inhabit biofilms. Protozoans that consume bacteria may feed on biofilms. Protozoan oocysts and virus particles can become entrapped in a biofilm and later detach, returning to the environment.

In a suspension test, the standard organisms per claimed group (bacteria, fungi, etc.) should be tested.

For a general claim of efficacy against biofilms, as a minimum, bacteria should be tested in laboratory biofilm tests. When action against other groups of organisms (e.g. fungi, algae etc.) is claimed these should be tested as well.

In suspension tests the standard organisms should be tested (see Appendix 3).

*Pseudomonas aeruginosa, Staphylococcus aureus* and *Legionella* spp. are acceptable test organisms for the laboratory biofilm tests. Mixtures of test organisms for producing biofilms are only acceptable as additional tests as it is difficult to standardise such methods.

In simulated use or field trials the biofilm may be formed *in vivo* with naturally occurring micro-organisms.

**5.4.2.11.3 Acceptance criteria**

A product will be assessed to be sufficiently effective if the required laboratory and, where relevant, field tests have been performed (using the required test organisms and test conditions), and when the pass criteria for the tests have been met.

Where pass criteria are available in the standard tests these should be met.

If the test does not provide these criteria, the general criteria referenced in Appendix 4 can be taken as guidance for the level of reduction required. Deviations from the pass criteria are possible, but must be justified in the application. The CA will evaluate any justification on a case-by-case basis, consulting the other CAs where appropriate, and decide whether it is acceptable or not.
5.4.2.12 Soil
Disinfection of soil and other substrates (in playgrounds) with biocides is not common (and so far not claimed for Annex I of the BPD or the “Union list of approved substances” of the BPR). This is more often done for plant protection. Therefore, plant protection guidelines and EPPO standards on soil treatments should be referred to for test methods. The use of the test methods should be justified with the application.

5.4.2.13 Other uses in PT2
Several other uses are mentioned in the description of PT2: wastewater and hospital waste disinfection, algaeicides for swimming pools and indoor/outdoor aquatic area (aquaria / garden ponds), foot baths in swimming pools, chemical toilets, disinfection of air. No data requirements and acceptance criteria for these uses are currently available.

However, the general principles for efficacy evaluation in PT2 are applicable for these other uses. Efficacy data should be adequate to demonstrate efficacy and suitability for the intended use, based on laboratory and/or practical data from existing and/or proposed new quantitative studies. If desired the design of any proposed efficacy tests may be agreed between the applicant and the CA taking into account all relevant conditions of use. Such factors include consideration of the organisms to be controlled, requirements for biocidal or biostatic effects, contact time and temperature and the nature and presence of interfering substances.

Specific requirements should also be set on a case by case basis by the CA as appropriate for specific claims.

5.4.3 PT3 Veterinary hygiene biocidal products

5.4.3.1 Introduction
Product type 3 contains biocidal products used for veterinary hygiene purposes such as disinfectants, disinfecting soaps, oral or corporal hygiene products or with anti-microbial function. Products used to disinfect the materials and surfaces associated with the housing or transportation of animals are also included.

Some of the products in PT3 are on the borderline with veterinary medicinal products or cosmetic products. If the product under investigation is within the scope of the Veterinary Medicinal Products Directive (2001/82/EC as amended by 2004/28/EC) it is excluded from the BPR for the respective use. When a product only has a cosmetic claim (e.g. cleaning skin, hoofs, paws) and no reference is made to any biocidal claim (e.g. skin disinfection, activity against microorganisms), it is excluded from the BPR.

Borderline cases are discussed in more detail in the respective sections below.

Following a decision taken at the CA meeting in May 2015 (CA-May -2015-Doc 8.3) products applied for general disinfection of surfaces in the medical area (medical practices, hospitals) as well as of surfaces in veterinary practices associated with examination and operation/treatment of the animals are assigned to PT 2, whereas products for specific veterinary hygiene purposes (e.g. products with specific claims against a target organism only relevant in the veterinary area) are considered to be in PT 3.

In the sections below the requirements and acceptance criteria for most common uses are specified. For other uses and claims that are not specifically mentioned the requirements will be set on a case-by-case basis by the CAs.
5.4.3.2 Disinfectants for hard surfaces in PT3

5.4.3.2.1 Introduction

Biocides can be used to disinfect hard surfaces, both porous and non-porous, in areas such as animal housing (stables, cages, housing for pets, etc.), animal transportation vehicles (including tyres), hatcheries, etc. These surfaces may be tables, floors, walls, the outsides of (milking) machinery (including milking robots and milking clusters/claws) and hard furniture, equipment, boots, etc. Products may be applied by spraying, wiping, foaming or soaking, and may be washed or wiped off after a certain contact time. Boots and tyres may be disinfected by walk-through, drive-through bath or mat, or even by a machine (boot wash station), etc.

The testing requirements for some specific uses of hard surface disinfectants are discussed in separate sections, for example, beehives.

Disinfection of inner surfaces of pipelines or reservoirs for milk, water or feed for animals are considered food and feed contact surfaces and are therefore considered PT4 (see section 5.4.4.3 of this Guidance). Outer surfaces of milking equipment are considered in this section.

5.4.3.2.2 Data requirements

Test methods

For efficacy testing of veterinary hard surface biocidal products, the tiered approach as described in section 5.4.0.4.1 of this Guidance is preferred.

The following tests are normally required for a hard surface disinfectant:

- a quantitative suspension test (phase 2, step 1);
- and a quantitative surface test (phase 2, step 2);

both simulating practical conditions appropriate to its intended use (temperature, soiling, porous or non-porous surfaces, contact time, etc.).

Field tests in phase 3 are optional, according to section 5.4.0.4.3 of this Guidance. No validated test methods are available yet.

Several methods for testing the efficacy of hard surface disinfectants are available.

Appendices 2 and 4 give a list of recommended test methods.

The following documents are recommended for surface disinfection:

- EN 14885: gives an overview of which EN phase 2, step 1 and step 2 tests to use for different uses;
- DVG guideline (relevant for testing against endoparasites and virucidal activity on porous or non-porous surfaces, as long as no EN tests are available).

The use of the specified tests is strongly recommended where they are relevant and appropriate. Where the tests are not appropriate to the product, other tests can be used, although a justification for the relevance of the tests used should also be provided.

Since OECD tests are not specified for veterinary use, they are not specifically recommended.

In the veterinary area very often rough, porous surfaces have to be disinfected (i.e. wood, concrete, rough plastic materials). When tests for porous surfaces are available it is recommended to use these tests for general surface disinfection in veterinary areas.
For boot, tyre, and equipment disinfection by immersion in a bath, information should be provided on how long the efficacy of a bath can be guaranteed (time period, number of boots etc. passing through). Challenging efficacy tests (capacity tests, see section 5.4.0.4.1 of this Guidance) should be done, simulating the consecutive challenge not only by microorganisms but also by soiling. A test with relevant organic soiling should be provided in order to ensure that biocidal product can be challenged successfully with the test organism until the end of the claimed period of use. Alternatively, for products with one active substance that can easily be measured, efficacy can be demonstrated using a field test in which the amount of active substance is measured several times during the test period. Efficacy (suspension) tests should be provided with the concentration of the product tested (in the suspension test) and the active substance concentration obtained in the field at the end of the claimed period of use.

A product can be applied by airborne diffusion of an aerosol, a smoke, a vapour or a gas, with the intention to disinfect the surfaces of the walls, floor and ceiling of the room, as well as external surfaces of the furniture and equipment present in the treated room. For these products the test methods are described in section 5.4.2.5 of this Guidance. These tests should be adapted to fit the conditions (soiling, etc. see section 5.4.3.2.2 of this Guidance) for veterinary use.

When efficacy against a biofilm is claimed, a simulated-use test or field test has to be performed, along with a phase 2, step 1 test. See section 5.4.2.11 of this Guidance for test methods.

Where no phase 2, step 2 or phase 3 tests are provided this must be justified in the application for authorisation and will be evaluated on a case-by-case basis.

The EN tests are strongly recommended where available and appropriate. For an overview of available EN tests see Appendices 2 and 4.

**Test organisms**

Relevant groups of organisms to be controlled in the veterinary area can be bacteria, yeasts, fungal spores, viruses, mycobacteria, bacterial spores, and endoparasites (oocysts).

Veterinary hard surface biocidal products should be at least sufficiently effective against bacteria and yeasts. Efficacy tests with these organisms should always be provided.

Products for disinfection of veterinary instruments and/or animal transportation vehicles should not only be effective against bacteria and yeasts but also against viruses.

Activity against fungi is also required for products used in hatcheries.

For all other groups of organisms, data only have to be provided when activity against those organisms is claimed.

The test organisms used in efficacy tests are normally stated in the applicable standard test methods. An overview of reference test organisms is given in Appendix 3.

**Test conditions**

It is important that the tests are performed using the same contact time as claimed on the label.

The claimed contact time has to be a realistic value, for instance:

- for surface disinfection products used on the outside of animal transport vehicles (specifically tyres) the contact time should not exceed 5 minutes;
• for disinfectants used on boots applied by spraying or walk-through bath the contact time should not exceed 1 minute;
• for disinfectants applied by dipping in bath, used on boots, materials etc. the contact time should be as claimed on the label;
• for surface disinfection products used in animal housing on floors, walls etc. the contact times as stated in the standard tests should be taken into account.

Additional contact times can be considered if appropriate and justified by the application (e.g. overnight disinfection).

Tests should be carried out with soiling for clean or dirty conditions (low or high-level soiling) in accordance with the test requirements. Tests under clean conditions will only suffice when the label instructions state that cleaning prior to disinfection is necessary. If this is not stated on the label, the test should be done under dirty conditions. The soiling needed for clean and dirty conditions can be found in the relevant EN tests and are referenced in Appendix 4. When the test does not state two levels of soiling (e.g. porous surface test), the soiling referenced in Appendix 4 should be used.

Normally PT3 products are tested at 10°C or below since the temperature in animal housings can be low. For some uses higher temperatures are acceptable (e.g. hatcheries). Deviations from this temperature requirement must be justified in the application and will be evaluated on a case-by-case basis. Any limitations on the temperatures at which the product should be used, and for which efficacy has been proven should be stated on the label.

5.4.3.2.3 Acceptance criteria

A product will be assessed to be sufficiently effective if the required laboratory and simulated-use or field tests have been performed (using the required test organisms and test conditions), and when the pass criteria for the tests have been met.

Where pass criteria are available in the standard test these should be met. For PT3 products the required log₁₀ reductions are referenced in Appendix 4.

Deviations from the pass criteria are possible, but must be justified in the application. The CA will evaluate any justification on a case-by-case basis, consulting the other CAs where appropriate, and decide whether it is acceptable or not.

5.4.3.3 Disinfection of bee hives and beekeeping equipment

5.4.3.3.1 Introduction

Disinfection of beehives is done to prevent spread of diseases from one bee population to the next.

Only disinfection of empty bee hives and beekeeping equipment, with products without a medicinal claim, is a biocidal use for general disinfection. Products used in beehives with bees, honey and brood combs are veterinary medicinal products. These products are within the scope of the Veterinary Medicinal Products Directive (2001/82/EC as amended by 2004/28/EC) and are therefore excluded from the BPR.

Important disease which can be spread via bee hives are American foulbrood (Paenibacillus larvae), European foulbrood, (Melissococcus plutonius), Nosema (Nosema apis, Nosema ceranae), chalkbrood (Ascosphaera apis) stonebrood (Aspergillus flavus) and some viral diseases. Of these diseases American foulbrood, which is an endospore-forming bacterium, is the most difficult to control.
Normal practice in case of American and European foulbrood is to clean/disinfect bee hives and beekeeping equipment and additionally disinfected by scorching with a blowtorch.

5.4.3.3.2 Data requirements

Test methods

For efficacy testing of disinfection products for beekeeping, the tiered approach as described in section 5.4.0.4.1 of this Guidance is preferred.

The following tests are normally required for disinfectants for bee hives:

- a quantitative suspension test (phase 2, step 1);
- and a quantitative carrier test (phase 2, step 2) for porous surfaces;

both simulating practical conditions appropriate to its intended use (temperature, soiling, different surfaces, contact time, etc.).

Field tests in phase 3 are optional, according to section 5.4.0.4.3 of this Guidance. No validated test methods are available yet.

There are no standard tests available specifically for use in bee hives. Phase 2, step 1 EN tests for veterinary area are suitable, and for sporicidal activity the EN 13704. EN phase 2, step 2 tests for veterinary area on porous material would be suitable but they are not available for all organisms yet. This can be either EN 16437 phase 2, step 2 test on bacteria for veterinary area on porous material or DVG guidelines on rough surfaces. These tests can be adapted for other organisms.

In these tests a reference substance must be included.

Where no phase 2, step 2 tests for veterinary area on porous material are available, the available test should be adapted for this use (e.g. EN 16437 adapted for other organisms).

When the claim for the product is to replace both the cleaning/disinfection step and the flaming with a welding torch, a field trial has to be provided in which it is demonstrated that the product is as efficacious against foulbrood infected hives as is cleaning with sodium hydroxide combined with scorching with a blowtorch.

For an overview of available EN tests see Appendices 2 and 4.

Test organisms

Disinfection products for bee hives should be at least sufficiently effective against bacteria and bacterial spores. Efficacy tests with these organisms should always be provided.

For all other groups of organisms, data only have to be provided when activity against those organisms is claimed.

For bacterial spores only a test for the food area is available (EN 13704). For disinfection products for beehives spores of two bacterial species should be tested. Next to the current standard test organism, *Bacillus subtilillus* spores, also *Bacillus cereus* should be tested.

Test conditions

It is important that the tests are performed using the same contact time as claimed on the label.

The claimed contact time has to be a realistic value.

It must be ensured that the disinfected parts stay wet during the contact time. When residual efficacy is claimed for dried products this should be demonstrated in efficacy tests.
For disinfection of bee hives and beekeeping equipment, tests should be performed under dirty conditions (high-level soiling) used for surfaces in the veterinary area. If bee hives are not cleaned before disinfection the high-level soiling for suspension tests should be used, also in the porous surface test and tests adopted from other areas of use (e.g. EN 13704). The soiling needed for dirty conditions can be found in the relevant EN tests and referenced in Appendix 4.

For disinfection of bee hives a temperature of 10ºC or lower is acceptable. Deviations from this temperature requirement must be justified in the application and will be evaluated on a case-by-case basis.

5.4.3.3 Acceptance criteria

A product will be assessed to be sufficiently effective if the required laboratory and simulated-use or field tests have been performed (using the required test organisms and test conditions), and when the pass criteria for the tests have been met. Where pass criteria are available in the standard test these should be met. For PT3 products the required log$_{10}$ reductions are referenced in Appendix 4.

Deviations from the pass criteria are possible, but must be justified in the application. The CA will evaluate any justification on a case-by-case basis, consulting the other CAs where appropriate, and decide whether it is acceptable or not.

5.4.3.4 Animal feet disinfection

5.4.3.4.1 Introduction

Animal feet disinfection includes hoof and claw disinfection. Products are applied in a bath, through which the animals can walk, or as wipes, foam, spray, etc. See section 5.4.3.1 of this Guidance for overlap with other EU directives.

5.4.3.4.2 Data requirements

Test methods

For efficacy testing of animal feet disinfection products, the tiered approach as described in section 5.4.0.4.1 of this Guidance is preferred.

The following tests are normally required for an animal feet disinfectant:

- a quantitative suspension test (phase 2, step 1);
- and a quantitative carrier test (phase 2, step 2);

both simulating practical conditions appropriate to its intended use (temperature, soiling, different surfaces, contact time, etc.).

Field tests in phase 3 are optional, according to section 5.4.0.4.3 of this Guidance. No validated test methods are available yet.

There are no standard tests available specifically for use on animal feet. Phase 2, step 1 EN tests for veterinary area are suitable. Since hoofs are made of porous material EN phase 2, step 2 tests for veterinary area on porous material would be suitable but these are not available for all organisms yet. Alternatively, DVG guideline tests on rough surfaces can be used.

The phase 2, step 2 test design must always reflect the application. When no standard test is used the test design should be discussed with, and agreed by, the CA before testing takes place.
When no phase 2, step 2 or phase 3 tests are provided this must be justified in the application and will be evaluated on a case-by-case basis.

For disinfection in a hoof bath, information should be provided on how long the efficacy of a hoof bath can be guaranteed (time period, number of animals passing through). Challenging efficacy tests (capacity tests, see section 5.4.0.4.1 of this Guidance) should be done, simulating the consecutive challenge not only by micro-organisms but also by soiling. A test with relevant organic soiling should be provided in order to ensure that the biocidal product can be challenged successfully with the test organism until the end of the claimed period of use. When a challenge test is provided the quantitative suspension test can be waived. Alternatively, for products with one active substance that can easily be measured, efficacy can be demonstrated using a field test in which the amount of active substance is measured several times during the test period. Efficacy (suspension) tests should be provided with the concentration of the product tested (in the suspension test) and the active substance concentration obtained in the field at the end of the claimed period of use.

For an overview of available EN tests see Appendices 2 and 4.

**Test organisms**

Animal feet disinfection should be at least sufficiently effective against bacteria. Efficacy tests with these organisms should always be provided.

For all other groups of organisms, data only have to be provided when activity against those organisms is claimed.

**Test conditions**

It is important that the tests are performed using the same contact time as claimed on the label.

The claimed contact time has to be a realistic value, therefore maximum contact times are set.

For animal feet disinfection products the contact time should not exceed 5 minutes.

It must be ensured that it is possible to keep the disinfected parts wet during the contact time in practice. When residual efficacy is claimed for dried products this should be demonstrated in efficacy tests.

Tests should be carried out with high-level soiling conditions in accordance with the test requirements. Soiling conditions for animal feet disinfectants are the same as for other veterinary area disinfectants. The soiling needed for clean and dirty conditions can be found in the relevant EN tests and referenced in Appendix 4.

Normally animal feet disinfection products are tested at 10°C since feet disinfection is often carried out outside animal housings at low temperatures. Deviations from this temperature requirement must be justified in the application and will be evaluated on a case-by-case basis.

**5.4.3.4.3 Acceptance criteria**

A product will be assessed to be sufficiently effective if the required laboratory and simulated-use or field tests have been performed (using the required test organisms and test conditions), and when the pass criteria for the tests have been met.

Where pass criteria are available in the standard test these should be met. For PT3 products the required log\(_{10}\) reductions are referenced in Appendix 4.
Deviations from the pass criteria are possible, but must be justified in the application. The CA will evaluate any justification on a case-by-case basis, consulting the other CAs where appropriate, and decide whether it is acceptable or not.

5.4.3.5 Teat disinfection

5.4.3.5.1 Introduction

Teat disinfection products are used to disinfect the teats of the udder of dairy animals (e.g. cows, sheep and goats) before or after milking. Products can be applied by dipping, spraying, foaming, wiping, etc.

See section 5.4.3.6.1 of this Guidance for overlap with other EU directives.

5.4.3.5.2 Data requirements

Test methods

For efficacy testing of teat disinfection products, the tiered approach as described in section 5.4.0.4.1 of this Guidance is preferred.

The following tests are normally required for a teat disinfectant:

- a quantitative suspension test (phase 2, step 1);
- and a quantitative carrier test (phase 2, step 2), or a field test;

all simulating practical conditions appropriate to its intended use (temperature, soiling, different surfaces, contact time, etc.).

Phase 2, step 1 tests for the veterinary area, with relevant soiling for teat disinfection should be used.

No European standard phase 2, step 2 tests are available for teat disinfection. To demonstrate efficacy a phase 2, step 2 tests should be provided with a test design relevant for the use. The test design must reflect the application and should be discussed with and agreed by the CA before testing takes place.

When standard tests become available, which are relevant for teat disinfectants, it is recommended to use these tests.

Alternatively a phase 3 test, field trial, may be provided with a test design relevant for the use. The test design must reflect the application, should include a control with water instead of biocide, and should be discussed with and agreed by the CA before testing takes place.

Disinfectant towelettes/wipes

For disinfectant wipes, the phase 2, step 1 tests should be done preferably with the liquid extracted from the wipe or if difficult to extract, use the liquid as it is before it is added to the wipes. Phase 2, step 2 tests should be tests with mechanical action or, when this test is not available, with liquid extracted from the wipe (not the original liquid), with a justification of the volume that is applied per square centimetre. In addition, a test must be performed that shows that either the wipe will still disinfect after the wipe dries out or that the wipe stays wet long enough to disinfect according to the claim. In addition, the use directions can address these issues, for instance, stating on the label that only wet wipes are efficacious, or giving expiry dates for re-sealable packages.

For an overview of available EN tests see Appendices 2 and 4.
Example of phase 2, step 2 tests

The phase 2, step 2 surface carrier test can be derived from adaptation of CEN TC 216 surface tests. Instead of a hard surface carrier, carriers involved could be made of material simulating the teat. Justification for the used carrier should be provided.

Cells of test organisms should be applied and fixed onto the surface in a manner which represents pre- and post-application, (dried in case of pre-milking or not dried in case of post-milking), and incubated with the product for the appropriate time (see EN phase 2, step 2 test, for example, EN 14349 or EN 16437, for growth conditions, controls, etc.). After incubation with the product the cell count reduction is evaluated and compared to a water control.

The test design should be discussed with and agreed by the CA before testing takes place.

Test organisms

Teat disinfection products should be at least sufficiently effective against bacteria and yeasts. Efficacy tests with these organisms should always be provided.

For all other groups of organisms, data only have to be provided when activity against those organisms is intended to be claimed.

Virucidal activity

For products used as teat disinfectants a differentiation in the virucidal activity is made.

The claims can be:

- full virucidal activity or
- activity against enveloped viruses.

For each claim different test organisms should be tested.

The EN 14675 test for virucidal activity in the veterinary area tests Bovine Enterovirus Type 1 (ECBO), a non-enveloped virus. When this test is passed, full virucidal activity can be claimed.

Activity against enveloped viruses can be claimed when MVA = Modified Vaccinia Ankara is tested in a (modified) EN 14675 test.

When only activity against enveloped viruses is demonstrated the label claim cannot be “virucidal”. The SPC should clearly state which of the possible virucidal claims was demonstrated.

Test conditions

It is important that the tests are performed using the same contact time as claimed on the label.

The claimed contact time has to be a realistic value, therefore maximum contact times are set.

For post-milking teat disinfection products the contact time is normally 1 minute but should not exceed 5 minutes.

The contact time for pre-milking teat disinfection products is normally 30 seconds or less and should not exceed 60 seconds.

Deviations from this contact time requirement must be justified in the application for authorisation and will be evaluated on a case-by-case basis.
Tests for pre-milking products should be carried out with either low or high-level soiling for veterinary surfaces, depending on the instructions given for pre-cleaning procedures.

Tests for post-milking products should be carried out with soiling for teat disinfectants in accordance with the test requirements. Soiling conditions for teat disinfectants are mentioned in the bactericidal test and should be used for the test with other organisms as well.

The soiling needed can be found in EN 1656 and referenced in Appendix 4.

For teat disinfection a test temperature of 30ºC or lower is acceptable. Deviations from this temperature requirement must be justified in the application and will be evaluated on a case-by-case basis.

5.4.3.5.3 Acceptance criteria

A product will be assessed to be sufficiently effective if the required laboratory and simulated-use or field tests have been performed (using the required test organisms and test conditions), and when the pass criteria for the tests have been met.

Where pass criteria are available in the standard test these should be met. For PT3 products the required log_{10} reductions are referenced in Appendix 4.

Deviations from the pass criteria are possible, but must be justified in the application. The CA will evaluate any justification on a case-by-case basis, consulting the other CAs where appropriate, and decide whether it is acceptable or not.

5.4.3.6 Other animal corporal hygiene

5.4.3.6.1 Introduction

Disinfectants for animal corporal hygiene are used to disinfect the skin of animals. This section includes all animal skin disinfectants, which are not covered in the sections on teat or animal feet disinfection below.

A product applied on animal skin could be either a biocidal or a veterinary medicinal or a product for cleaning or cosmetic purposes. If the product under investigation is within the scope of the Veterinary Medicinal Products Directive (2001/82/EC as amended by 2004/28/EC) it is excluded from the BPR for the respective use. When a product does not have a biocidal claim (e.g. skin disinfection, activity against microorganisms claimed) but only a cosmetic claim (e.g. cleaning skin, paws) it is excluded from the BPR for the respective use.

Products for disinfection of damaged skin (e.g. wound disinfection) or disinfection of undamaged skin before a medical treatment (e.g. pre-operative skin disinfection or disinfection before injection) are always veterinary medicinal products.

When applying for authorisation for an animal corporal hygiene biocidal product within PT3 a detailed description of the intended use should be given, to prevent authorisation of veterinary medicinal products or medicinal uses, as biocides (e.g. the claim “animal skin disinfection” is insufficient).

For products that fall under the BPR the data requirements described in the following sections apply.
5.4.3.6.2 Data requirements

Test methods

For efficacy testing of animal corporal hygiene products, the tiered approach as described in section 5.4.0.4.1 of this Guidance is preferred.

The following tests are normally required for an animal corporal hygiene disinfectant:

- a quantitative suspension test (phase 2, step 1);
- and a quantitative carrier test (phase 2, step 2);

both simulating practical conditions appropriate to its intended use (temperature, soiling, different surfaces, contact time, etc.).

Field tests in phase 3 are optional, according to section 5.4.0.4.3 of this Guidance. No validated test methods are available yet.

Phase 2, step 1 tests for the veterinary area can be used.

No European standard phase 2, step 2 tests are available for animal skin disinfection. To demonstrate efficacy a phase 2, step 2 tests should be provided with a test design relevant for the use. The test design must reflect the application and should be discussed with and agreed by the CA before testing takes place.

When standard tests become available, which are relevant for skin disinfectants, it is recommended to use these tests.

For an overview of available EN tests see Appendices 4 and 6.

Example of phase 2, step 2 tests

The phase 2, step 2 surface carrier test can be derived from adaptation of CEN TC 216 surface tests. Instead of a hard surface carrier, carriers could be made of material simulating animal skin\(^\text{16}\). Method are currently being developed, but their aptitude for the respective biocidal use/demonstration of efficacy for animal skin disinfectants remains to be proven. Justification for the used carrier should be provided.

Cells of test organisms could be applied to the surface, dried, and incubated with the product for the appropriate time (see EN phase 2, step 2 test, e.g. EN 14349, for growth conditions, controls, etc.). After incubation with the product the cell count reduction is evaluated and compared to a water control.

For an overview of available EN tests see Appendices 2 and 4.

Test organisms

Animal corporal hygiene products should be at least sufficiently effective against bacteria and yeasts. Efficacy tests with these organisms should always be provided.

For all other groups of organisms, data only have to be provided when activity against those organisms is claimed.

Test conditions

It is important that the tests are performed using the same contact time as claimed on the label.

\(^\text{16}\) Please take into account EU regulation 1069/2009, on animal by-products.
The claimed contact time has to be a realistic value.

It must be ensured that the test surface does not remain wet longer than the part of the animal body treated with the product, for example, by using higher (more realistic) temperatures. When residual efficacy is claimed this should be demonstrated in efficacy tests.

Tests should be carried out with high level or low level soiling conditions in accordance with the test requirements. Soiling conditions for animal corporal hygiene products are the same as for other veterinary area disinfectants. The soiling needed for clean and dirty conditions can be found in the relevant EN tests and referenced in Appendix 4.

For animal corporal hygiene products a test temperature of 30ºC or lower is acceptable. Deviations from this temperature requirement must be justified in the application and will be evaluated on a case-by-case basis.

5.4.3.6.3 Acceptance criteria

A product will be assessed to be sufficiently effective if the required laboratory and simulated-use tests have been performed (using the required test organisms and test conditions), and when the pass criteria for the tests have been met.

Where pass criteria are available in the standard test these should be met. For PT3 products the required log\(_{10}\) reductions are referenced in Appendix 4.

Deviations from the pass criteria are possible, but must be justified in the application. The CA will evaluate any justification on a case-by-case basis, consulting the other CAs where appropriate, and decide whether it is acceptable or not.

5.4.3.7 Disinfection of hatching-eggs

5.4.3.7.1 Introduction

Disinfection of hatching-eggs includes the disinfection of eggs before they hatch in hatcheries. Products are applied in a bath, as a spray, as wipes, fumigation, etc..

5.4.3.7.2 Data requirements

Test methods

For efficacy testing of disinfection products for hatching-eggs, the tiered approach as described in section 5.4.0.4.1 of this Guidance is preferred.

The following tests are normally required for disinfectant for hatching-eggs:

- a quantitative suspension test (phase 2, step 1);
- and a quantitative carrier test (phase 2, step 2);

both simulating practical conditions appropriate to its intended use (temperature, soiling, different surfaces, contact time, etc.).

Field tests in phase 3 are optional, according to section 5.4.0.4.3 of this Guidance. No validated test methods are available yet.

Phase 2, step 1 tests for the veterinary area can be used.

As long as no standard phase 2, step 2 tests are available it is not obligatory to provide these tests. Phase 2, step 2 tests have to be provided as soon as standard tests are available.
When no phase 2, step 2 or phase 3 tests are provided this must be justified in the application and will be evaluated on a case-by-case basis.

For egg disinfection in a bath, information should be provided on how long the efficacy of a bath can be guaranteed (time period, number of eggs passing through). Challenging efficacy tests (capacity tests, see section 5.4.0.4.1 of this Guidance) should be done, simulating the consecutive challenge not only by microorganisms but also by soiling. A test with relevant organic soiling should be provided in order to ensure that biocidal product can be challenged successfully with the test organism until the end of the claimed period of use. When a challenge test is provided the quantitative suspension test can be waived. Alternatively, for products with one active substance that can easily be measured, efficacy can be demonstrated using a field test in which the amount of active substance is measured several times during the test period. Efficacy (suspension) tests should be provided with the concentration of the product tested (in the suspension test) and the active substance concentration obtained in the field at the end of the claimed period of use.

For products applied by airborne diffusion of an aerosol, a smoke, a vapour or a gas, with the intention to disinfect the room, as well as on external surfaces of the eggs in the room, the test methods are described in section 5.4.2.5 of this Guidance. These tests should be adapted to fit the conditions (soiling, etc. see section 5.4.3.7.2 of this Guidance) for veterinary use.

For an overview of available EN tests see Appendices 2 and 4.

**Test organisms**

Disinfection products for hatching-eggs should be at least sufficiently effective against bacteria and fungi. Efficacy tests with these organisms should always be provided.

For all other groups of organisms, data only have to be provided when activity against those organisms is claimed.

**Test conditions**

It is important that the tests are performed using the same contact time as claimed on the label.

The claimed contact time has to be a realistic value.

It must be ensured that the disinfected parts stay wet during the contact time. When residual efficacy is claimed for dried products this should be demonstrated in efficacy tests.

Tests should be carried out with soiling for clean or dirty conditions in accordance with the test requirements. Tests under clean conditions will only suffice when the label instructions state that cleaning prior to disinfection is necessary. If this is not stated on the label, the test should be done under dirty conditions. Soiling conditions for of hatching-eggs disinfectants are the same as for other veterinary area disinfectants. The soiling needed for clean and dirty conditions can be found in the relevant EN tests and referenced in Appendix 4.

For disinfection of hatching-eggs a temperature of 30°C or lower is acceptable. Deviations from this temperature requirement must be justified in the application and will be evaluated on a case-by-case basis.
5.4.3.7.3 Acceptance criteria

A product will be assessed to be sufficiently effective if the required laboratory or, when applicable, simulated-use or field tests have been performed (using the required test organisms and test conditions), and when the pass criteria for the tests have been met.

Where pass criteria are available in the standard test these should be met. For PT3 products the required log_{10} reductions are referenced in Appendix 4.

Deviations from the pass criteria are possible, but must be justified in the application. The CA will evaluate any justification on a case-by-case basis, consulting the other CAs where appropriate, and decide whether it is acceptable or not.

5.4.3.8 Textile disinfection in PT3

5.4.3.8.1 Introduction

Textile disinfection products within PT3 are mainly used to disinfect the cloths used for teat cleaning/disinfection of dairy cattle before milking. Products are normally applied by dipping the cloth in a disinfectant solution. For other uses the requirements below should be adapted to fit the intended use.

5.4.3.8.2 Data requirements

Test methods

For efficacy testing of textile disinfection products, the tiered approach as described in section 5.4.0.4.1 of this Guidance is preferred.

The following tests are normally required for textile disinfection products:

- a quantitative suspension test (phase 2, step 1);
- a quantitative carrier test involving carriers made of test fabric (cotton, polyester) (phase 2, step 2);

both simulating practical conditions appropriate to its intended use (temperature, soiling, different surfaces, contact time, repeated challenges, etc.).

Field tests in phase 3 are optional, according to section 5.4.0.4.3 of this Guidance. No validated test methods are available yet.

Test methods for textile disinfection are described in section 5.4.2.10 of this Guidance. Currently, the following tests are available:

- phase 2, step 1 suspension tests as described in EN 14885,
- phase 2, step 2 tests involving test fabrics in:
  - a small scale laboratory setting (e.g. ASTM E2406) or;
  - a full-scale laundry machine test (pr EN 16616, or DGHM).

In the phase 2, step 2 tests fabric is contaminated with test organisms and then exposed to the disinfectant. These tests should be adapted to fit the conditions (soiling, etc. see 4.8.2.3) for veterinary use. For disinfection in washing machines a full-scale laundry machine test, according to test conditions mentioned in section 5.4.2.10.2 of this Guidance, is obligatory.

The EN tests are strongly recommended where available and appropriate. For an overview of available EN tests see Appendices 2 and 4.
Test organisms
Textile disinfection products should be at least sufficiently effective against bacteria and yeasts. Efficacy tests with these organisms should always be provided.

For all other groups of organisms, data only have to be provided when activity against those organisms is claimed.

When the product is intended to be used at high temperatures (>40 °C) relevant test organisms for these temperatures should be used as described in section 5.4.0.4.4 of this Guidance.

Test conditions
It is important that the tests are performed using the same contact time as claimed on the label.

The claimed contact time has to be a realistic value.

The contact time products intended for disinfection of textile in between milking sessions can be several hours.

Tests should be carried out with soiling for clean or dirty conditions in accordance with the test requirements for veterinary area. Tests under clean conditions will only suffice when the label instructions state that cleaning prior to disinfection is necessary. If this is not stated on the label, the test should be done under dirty conditions. Soiling conditions for milking-textile disinfectants are the same as for teat disinfectants. The soiling needed for clean and dirty conditions can be found in the relevant EN tests and referenced in Appendix 4.

For textile disinfection a test temperature should be according to the use instructions. When the textile is immersed in a bucket with warm water it should be taken into account that the water temperature will decrease during the disinfection process. This should be reflected in the test conditions.

5.4.3.8.3 Acceptance criteria
A product will be assessed to be sufficiently effective if the required laboratory and simulated-use or field tests have been performed (using the required test organisms and test conditions), and when the pass criteria for the tests have been met.

Where pass criteria are available in the standard test these should be met. For PT3 products the required log_{10} reductions are referenced in Appendix 4.

Deviations from the pass criteria are possible, but must be justified in the application. The CA will evaluate any justification on a case-by-case basis, consulting the other CAs where appropriate, and decide whether it is acceptable or not.

5.4.3.9 Disinfection of manure, litter and other substrates for veterinary use
5.4.3.9.1 Introduction
Manure mainly consists of urines and faeces (organic matters and intestinal bacteria) in which can also be mixed straw of litters in more or less big quantity, according to the breeding technique (partial slats or complete slats).

Manure has a potential for spreading infectious diseases and biocidal products are used to destroy some infective agents and also control microbial agents responsible of malodours.
Litters are usually used in animal housing (poultry, pigsties, etc.) and also for pets in private uses. They absorb urines and faeces. Biocidal products are mainly used to deodorize and neutralize bad smells.

5.4.3.9.2 Data requirements

Test methods

For efficacy testing of disinfects biocidal products used for manure and litter disinfection, the tiered approach as described in section 5.4.0.4.1 of this Guidance is preferred.

The following tests are normally required:

- a quantitative suspension test (phase 2, step 1),
- and simulated-use test, or field test

all simulating practical conditions appropriate to its intended use (temperature, soiling, different surfaces, contact time, mode of application, pH, etc.).

An example of a simulated-use test could be autoclaved manure or litter collected in animal housing and tested in the lab with inoculation of target organisms. A control without addition of disinfectants should be included. The test design should be discussed with and agreed by the CA before testing takes place.

In case of products claiming malodour control, the same requirements as mentioned in the section 5.4.0.5.4 of this Guidance, are required.

Test organisms

Generally, target organisms have to be representative of the veterinary area, as stated in EN 14885.

For specific uses in industry, an exception can be made when sound justification is provided. This will be evaluated on a case-by-case basis.

Taking into account the specificity of some kind of uses, it may be justified to test additional target organisms (e.g. Brachyspira hyodysenteriae agent of swine dysentery,), special growth conditions, etc.

In case of malodour control, tests should be performed with odour producing microorganisms. A justification for which bacteria, fungi, etc. are relevant to the intended use should be provided. Along with these laboratory tests, an odour test can be performed.

Test conditions

It is important that the tests are performed with the same contact time as claimed on the label.

The claimed contact time has to be a realistic value.

Quantitative suspension tests must be carried out with high level soiling conditions and a temperature of 10°C or less.

The test temperature should be according to the use instructions on the label and appropriate to the uses (stables, private homes, etc.).

Field and simulated-use test have to be performed according to the dose, conditions and mode of application of the product. For example, if the product is applied on top of the manure, the product does not have to be mixed with the organic matter but has to be put on top of it (to mimic the diffusion and evaluate efficacy in the same conditions as in the practice).
In case of litter, if persistence is claimed with some recommendations about the frequency of renewal, adequate simulating tests (with appropriate contribution of organic matters in the test) have to be performed.

Deviations from these requirements must be justified in the application for authorisation and will be evaluated on a case-by-case basis.

5.4.3.9.3 Acceptance criteria

A product will be assessed to be sufficiently effective if the required laboratory and, field (or simulated-use) tests have been performed (using the required test organisms and test conditions), and when the pass criteria for the tests have been met.

Where pass criteria are available in the standard test these should be met. For PT3 products the required log_{10} reductions are referenced in Appendix 4.

Deviations from the pass criteria are possible, but must be justified in the application. The CA will evaluate any justification on a case-by-case basis, consulting the other CAs where appropriate, and decide whether it is acceptable or not.

5.4.3.10 Other uses in PT3

Several uses of PT3 products have been specified in the above sections and data requirements and acceptance criteria for these uses are described. For products with other uses that do not fit in one of the described uses, it is up to the applicant to demonstrate efficacy in an appropriate way.

In general, the tiered approach as described in section 5.4.0.4.1 of this Guidance is preferred. Where possible the standard tests required for the described uses should be taken (e.g. EN phase 2, step 1 and step 2 tests for veterinary area). Where the tests are not appropriate for the product, other tests can be used. In that case, a justification for the relevance of the tests used should be provided. The test design should be discussed with and agreed by the CA before testing takes place. The evaluation will be done on a case-by-case basis by the CAs.

The guidance will be updated when new methods become available.

5.4.4 PT4 Food and feed area disinfectants

5.4.4.1 Introduction

Product type 4 contains biocidal products used for the disinfection of equipment, containers, consumption utensils, surfaces or pipework associated with the production, transport, storage or consumption of food or feed (including drinking water) for humans and animals.

Some disinfectants applied in the food or feed area can be either biocidal product or a preservative for food or feed. If the product under investigation is within the scope of Regulations (EC) 852/2004, 853/2004 and 854/2004 on food hygiene, it is excluded from the BPR. The Regulation 852/2004 is on the hygiene of foodstuffs; the Regulation 853/2004 lays down specific hygiene rules for food of animal origin; the Regulation 854/2004 lays down specific rules for the organisation of official controls on products of animal origin intended for human consumption.

In the sections below the requirements and acceptance criteria for most common uses are specified. For other uses and claims that are not specifically mentioned the requirements will be set on a case-by-case basis by the CAs.
5.4.4.2 Disinfection of hard surfaces in food and feed area PT4

5.4.4.2.1 Introduction

Biocides can be used to disinfect hard surfaces in areas such as food industry, kitchens in restaurants or homes, shops like butchers and grocery shops were food is processed etc. These surfaces may be tables, floors, walls, the outsides of machinery, equipment, reservoirs for water or feed in animal housing etc. Products are often wiped, sprayed, foamed, applied by low to high pressure etc., onto the surface, and maybe washed or wiped off after a certain contact time.

The testing requirements for some specific uses of hard surface disinfectants are discussed in separate sections, for example, CIP, equipment and dishwashing disinfectants etc.

5.4.4.2.2 Data requirements

Test methods

For efficacy testing of food and feed area biocidal products used on hard surfaces, the tiered approach as described in section 5.4.0.4.1 of this Guidance is preferred.

The following tests are normally required for a hard surface disinfectants:

- a quantitative suspension test (phase 2, step 1);
- and a quantitative surface test (phase 2, step 2);

both simulating practical conditions appropriate to its intended use (temperature, soiling, different surfaces, contact time, etc.).

Tests in phase 3 are optional, according to section 5.4.0.4.3 of this Guidance. No validated test methods are available yet.

Several methods for testing the efficacy of hard surface disinfectants are available. Tests with mechanical action might be adopted from medical area, if appropriate. Appendices 2 and 4 give a list of recommended test methods.

The following documents are recommended for surface disinfection:

- EN 14885: gives an overview of which EN phase2/step1 and step2 tests to use for different uses;

if CEN standards are not relevant or available for the use or organisms claimed the following documents are recommended if appropriately reflecting the application:

- OECD guidance for the testing of chemicals: Quantitative method for evaluating activity of microbiocides used on hard non-porous surfaces (these are surface tests which would be considered phase 2, step 2 tests).

The use of the specified tests is strongly recommended where they are relevant and appropriate.

When efficacy against biofilm is claimed a simulated-use test or field test has to be provided, next to a phase 2, step 1 test. See section 5.4.2.11 of this Guidance for test methods.

A product can be applied by airborne diffusion of an aerosol, a smoke, a vapour or a gas, with the intention to disinfect on the surfaces of the walls, floor and ceiling of the room, as well as on external surfaces of the furniture and equipment present in the treated room. For these products the test methods are described in section 5.4.2.5 of this Guidance. These tests should be adapted to fit the conditions (soiling, etc. see section 5.4.3.2.2 of this Guidance) for use in food and feed area.
**Disinfectant towelettes/wipes**

For disinfectant wipe, the phase 2, step 1 tests should be done preferably with the liquid extracted from the wipe, or if difficult to extract, use the liquid as it is before it is added to the wipes. Phase 2, step 2 tests should be tests with mechanical action. These tests are available for bacteria and yeasts. For testing other organisms surface tests can be done with liquid extracted from the wipe (not the original liquid), with a justification of the volume that is applied per square centimetre. In addition, a test must be performed that shows that either the wipe will still disinfect after the wipe dries out or that the wipe stays wet long enough to disinfect according to the claim. In addition, the use directions can address these issues, for instance, stating on the label that only wet wipes are efficacious, defining the surface area each towel can disinfect (e.g. 0.5 m$^2$), or giving expiry dates for re-sealable packages.

**Test organisms**

Food and feed hard surface biocidal products should be at least sufficiently effective against bacteria and yeasts. Efficacy tests with these organisms should always be provided.

For specific uses in industry, an exception can be made when sound justification is provided. This will be evaluated on a case-by-case basis.

For all other groups of organisms, data only have to be provided when activity against those organisms is claimed. Relevant groups of test organisms, next to bacteria and yeasts, can be fungi (fungal spores), viruses, bacteriophages, and bacterial spores. Bacteriophages are mainly of importance in the dairy industry.

The test organisms used in efficacy tests are normally stated in the applicable standard test methods. For food area disinfectants *Salmonella* Thyphimurium, *Salmonella* spp., *Listeria* spp. and *Campylobacter jejuni* are relevant target organisms. For products which claim general efficacy against bacteria, the standard test bacteria should be tested. For these products efficacy against *Salmonella* spp., *Listeria* spp. and *Campylobacter jejuni* is assumed, because they are more susceptible than the standard test bacteria.

The EN standards for food area only include a test on bacteriophages but not on other viruses. To demonstrate a general virus claim a modified EN phase 2, step 1 test (medical area test with food area soiling) can be provided with Adenovirus and Murine Norovirus as test organism and a phase 2, step 2 test (either modified EN medical test, or DVG test or, as soon as available, an EN food area test) with Murine Norovirus.

An overview of reference test organisms is given in Appendix 3.

**Test conditions**

It is important that the tests are performed with the same contact time as claimed on the label. The claimed contact time has to be a realistic value.

Tests should be carried out with soiling for clean or dirty conditions in accordance with the test requirements. Tests under clean conditions will only suffice when the label instructions state that cleaning prior to disinfection is necessary. If this is not stated on the label the test should be done under dirty conditions. Note that for use in specific industries different types of soiling for dirty conditions should be used.

The soiling needed for clean and dirty conditions can be found in the relevant EN tests or EN 14885 (version 2014 or later) and referenced in Appendix 4.

If a product is intended to be used in more than one area of use (e.g. milk industry and meat industry) it is justified, after having identified the most challenging test organism, to
test the relevant soiling types with this organism. That applies only per group of organisms (e.g. bacteria).

The test temperature should be according to the use instructions on the label. Food and feed area disinfectants are generally used at room temperature (test temperature 20 °C) but for some uses and claims (e.g. surfaces in cold storage rooms) low temperatures of 4 °C or 10 °C are relevant and should be tested.

5.4.4.2.3 Acceptance criteria

A product will be assessed to be sufficiently effective if the required laboratory and simulated-use or field tests have been performed (using the required test organisms and test conditions), and when the pass criteria for the tests have been met.

Where pass criteria are available in the standard test these should be met. For PT4 products the required log10 reductions are referenced in Appendix 4.

Deviations from the pass criteria are possible, but must be justified in the application. The CA will evaluate any justification on a case-by-case basis, consulting the other CAs where appropriate, and decide whether it is acceptable or not.

5.4.4.3 Disinfection of inner surfaces in PT4

5.4.4.3.1 Introduction

Biocides can be used to disinfect the inner surfaces of pipes, tanks, fillers, mixers, and other machines which come in contact with food or feed (including liquids). This includes food and feed industry, milking equipment on farms, large equipment in restaurants or shops were food is processed, etc. Inner surfaces in contact with water are discussed in the following sections.

These surfaces are disinfected by filling and circulating the biocide in the pipes, tanks, machines, etc. with disinfectant (Cleaning In Place, CIP). Also disinfection of inner surfaces of equipment by filling without circulation (not using CIP) is included in this section.

5.4.4.3.2 Data requirements

Test methods

For efficacy testing of food and feed area biocidal products used on inner surfaces using CIP, the following tests are normally required:

- quantitative suspension tests (phase 2, step 1), simulating practical conditions appropriate to its intended use (temperature, soiling, different surfaces, contact time, etc.).

For efficacy testing of food and feed area biocidal products used on inner surfaces by filling without circulation, the following tests are normally required for these disinfectants:

- quantitative suspension tests (phase 2, step 1);
- and a quantitative surface test (phase 2, step 2);

both simulating practical conditions appropriate to its intended use (temperature, soiling, different surfaces, contact time, etc.).

Tests in phase 3 are optional, according to section 5.4.0.4.3 of this Guidance. No validated test methods are available yet.

Several methods for testing the efficacy of inner surface disinfectants are available. Appendices 2 and 4 give a list of recommended test methods.
The following documents are recommended for inner surface disinfection using CIP:

- EN 14885 gives an overview of which EN phase 2, step 1 and step 2 tests to use for different uses;

if CEN standards are not relevant or available for the use or organisms claimed the following documents are recommended if appropriately reflecting the application:

- OECD guidance for the testing of chemicals: Quantitative method for evaluating activity of microbiocides used on hard non-porous surfaces. (These are surface tests which would be considered phase 2, step 2 tests).

The use of the specified tests is strongly recommended where they are relevant and appropriate.

When efficacy against biofilm is claimed a simulated-use test or field test has to be provided, next to a phase 2, step 1 test. See section 5.4.2.11 of this Guidance for test methods.

When the disinfection is done with vaporised biocide a simulated-use test or a field test has to be provided. See section 5.4.2.5 of this Guidance for test methods.

**Test organisms**

Food and feed hard surface biocidal products should be at least sufficiently effective against bacteria and yeasts. Efficacy tests with these organisms should always be provided.

For specific uses in industry, an exception can be made when sound justification is provided. This will be evaluated on a case-by-case basis.

For all other groups of organisms, data only have to be provided when activity against those organisms is claimed. Relevant groups of organisms, next to bacteria and yeasts, can be fungal spores, viruses, phages, and bacterial spores. Phages are mainly of importance in the dairy industry.

The test organisms used in efficacy tests are normally stated in the applicable standard test methods. For food area disinfectants *Salmonella* spp., *Listeria* spp. and *Campylobacter jejuni* are relevant target organisms. For products which claim general efficacy against bacteria, the standard test bacteria should be tested. For these products efficacy against *Salmonella* spp., *Listeria* spp. and *Campylobacter jejuni* is assumed, because they are more susceptible than the standard test bacteria.

The EN standards for food area only include a test on bacteriophages but not for other viruses. To demonstrate a general virus claim a modified EN phase 2, step 1 test (medical area test with food area soiling) can be provided with Adenovirus and Murine Norovirus as test organism and a DVG phase 2, step 2 test.

When CIP is done at high temperatures relevant test organisms for these temperatures should be used as described in section 5.4.0.4.4 of this Guidance.

An overview of test organisms, also for high temperatures, is given in Appendix 3.

**Test conditions**

It is important that the tests are performed with the same contact time as claimed on the label. The claimed contact time has to be a realistic value.

Tests should be carried out with soiling for clean or dirty conditions in accordance with the test requirements. Tests under clean conditions will only suffice when the label instructions state that cleaning prior to disinfection is necessary. If this is not stated on the label the test
should be done under dirty conditions. Note that for use in specific industries different types of soiling for dirty conditions should be used.

The soiling needed for clean and dirty conditions can be found in the relevant EN tests or EN 14885 (version 2014 or later) and referenced in Appendix 4.

The test temperature should be according to the use instructions on the label. Food and feed area disinfectants are generally used at room temperature (test temperature 20 ºC) but for some uses and claims other temperatures are relevant. For example, for surfaces in cold machinery, low temperatures of 4 ºC or 10 ºC are relevant and should be tested. CIP disinfection is often done at high temperatures of 40 to 80 ºC. When this is the intended use the test temperature should be in accordance with the use and relevant test organisms should be used (see section 5.4.4.3.2 of this Guidance).

5.4.4.3.3 Acceptance criteria

A product will be assessed to be sufficiently effective if the required laboratory and, when applicable, simulated-use or field tests have been performed (using the required test organisms and test conditions), and when the pass criteria for the tests have been met.

Where pass criteria are available in the standard tests these should be met. For PT4 products the required log_{10} reductions tests are referenced in Appendix 4.

Deviations from the pass criteria are possible, but must be justified in the application. The CA will evaluate any justification on a case-by-case basis, consulting the other CAs where appropriate, and decide whether it is acceptable or not.

5.4.4.4 Equipment disinfection by soaking

5.4.4.4.1 Introduction

Biocides can be used to disinfect dishes, equipment, crates, boxes, etc. by soaking. This can include dishwashing disinfectants, however, normal dishwashing detergents are cleaning products and not included in the BPR. Equipment disinfection in washing machines is covered in the next section.

This can be used in areas such as food industry, kitchens in restaurants or homes, shops like butchers and grocery shops were food or feed is processed, etc.

5.4.4.4.2 Data requirements

Test methods

For efficacy testing of equipment and dish washing disinfectants the tiered approach as described in section 5.4.0.4.1 of this Guidance is preferred.

The following tests are normally required for these disinfectants:

- a quantitative suspension test (phase 2, step 1);
- a quantitative surface test (phase 2, step 2);

both tests simulating practical conditions appropriate to its intended use (temperature, soiling, different surfaces, contact time, etc.).

Tests in phase 3 are optional, according to section 5.4.0.4.3 of this Guidance. No validated test methods are available yet.

Several methods for testing the efficacy of hard surface disinfectants are available.

Appendices 2 and 4 give a list of recommended test methods.
The following documents are recommended for equipment and dish washing disinfection:

- EN 14885: gives an overview of which EN phase2/step1 and step2 tests to use for different uses,

if CEN standards are not relevant or not available for the use or organisms claimed the following documents are recommended if appropriately reflecting the application:

- OECD guidance for the testing of chemicals: Quantitative method for evaluating activity of microbiocides used on hard non-porous surfaces. (These are surface tests which would be considered phase 2, step 2 tests)

The use of the specified tests is strongly recommended where they are relevant and appropriate.

When efficacy against biofilm is claimed a simulated-use test or field test has to be provided, next to a phase 2, step 1 test. See section 5.4.2.11 of this Guidance for test methods.

**Test organisms**

Equipment and dish washing disinfectants should be at least sufficiently effective against bacteria and yeasts. Efficacy tests with these organisms should always be provided.

For specific purposes in industrial uses, an exception can be made when sound justification is provided. This will be evaluated on a case-by-case basis.

For all other groups of organisms, data only have to be provided when activity against those organisms is claimed. Relevant groups of test organisms, next to bacteria and yeasts, can be fungi (fungal spores), viruses, bacteriophages, and bacterial spores. Bacteriophages are mainly of importance in the dairy industry.

The test organisms used in efficacy tests are normally stated in the applicable standard test methods. For dish washing disinfectants *Salmonella* spp., *Listeria* spp. and *Campylobacter jejuni* are relevant target organisms. For products which claim general efficacy against bacteria, the standard test bacteria should be tested. For these products efficacy against *Salmonella* spp., *Listeria* spp. and *Campylobacter jejuni* is assumed, because they are more susceptible than the standard test bacteria.

The EN standards for food area only include a test on bacteriophages but not for other viruses. To demonstrate a general virus claim a modified EN phase 2, step 1 test (medical area test with food area soiling) can be provided with Adenovirus and Murine Norovirus as test organism and a DVG phase 2, step 2 test.

When the product is intended to be used at high temperatures (>40 °C) relevant test organisms for these temperatures should be used as described in section 5.4.0.4.4 of this Guidance.

An overview of reference test organisms, also for high temperatures, is given in Appendix 3.

**Test conditions**

It is important that the tests are performed with the same contact time as claimed on the label. The claimed contact time has to be a realistic value. For manual dishwashing disinfectants the contact time will be short (seconds), while industrial equipment disinfection by soaking in a solution can be very long (hours).

In general dish washing disinfectants should be tested under dirty conditions, since these products are mainly used for combined cleaning and disinfection. Tests under clean conditions will only suffice when the label instructions state that cleaning prior to disinfection is necessary. If this is not stated on the label the test should be done under dirty conditions.
Tests should be carried out with soiling for clean or dirty conditions in accordance with the test requirements.

Note that for use in specific industries different types of soiling for dirty conditions should be used. The soiling needed for clean and dirty conditions can be found in the relevant EN tests or EN 14885 (version 2014 or later) and referenced in Appendix 4.

The test temperature should be according to the use instructions on the label.

Dish washing disinfectants for manual use are normally used at 40ºC and therefore tests should be done at this temperature. When the product is used at lower temperatures (e.g. only for rinsing after normal dish washing with hot water) tests can be done at 20ºC. When the intended use is soaking, starting with hot water and after which the solution will cool down during the contact time, this should also be taken into account in the tests.

When disinfection is done at temperatures of 40 to 80 ºC the test temperature should be in accordance with the use and relevant test organisms should be used (see section 5.4.4.4.2 of this Guidance).

5.4.4.4.3 Acceptance criteria

A product will be assessed to be sufficiently effective if the required laboratory and, when applicable, simulated-use or field tests have been performed (using the required test organisms and test conditions), and when the pass criteria for the tests have been met.

Where pass criteria are available in the standard test these should be met. For PT4 products the required log10 reductions tests are referenced in Appendix 4.

Deviations from the pass criteria are possible, but must be justified in the application. The CA will evaluate any justification on a case-by-case basis, consulting the other CAs where appropriate, and decide whether it is acceptable or not.

5.4.4.5 Disinfection in dish washing machines and crate washers

5.4.4.5.1 Introduction

Biocides can be used to disinfect dishes, equipment, crates, boxes, etc. in industrial or dishwashing machines.

This can be used in areas such as food or feed industry, kitchens in restaurants or homes, shops like butchers and grocery shops were food is processed, etc..

5.4.4.5.2 Data requirements

Test methods

For efficacy testing of equipment and dish washing disinfectants the tiered approach as described in section 5.4.0.4.1 of this Guidance is preferred.

The following tests are normally required for these disinfectants:

- a quantitative suspension test (phase 2, step 1);
- and a quantitative surface test (phase 2, step 2);
- and simulated-use or field test (phase 3) for disinfectants used in (dish)washing machines;

all tests simulating practical conditions appropriate to its intended use (temperature, soiling, different surfaces, contact time, etc.).

Several methods for testing the efficacy of hard surface disinfectants are available.
Appendices 2 and 4 give a list of recommended test methods.

The following documents are recommended for surface disinfection in dish washing machines:

- EN 14885: gives an overview of which EN phase 2, step 1 and step 2 tests to use for different uses.

The following test might be helpful for designing simulated-use or field tests:

- prDIN SPEC10534.

**Test organisms**

Equipment and dish washing disinfectants should be at least sufficiently effective against bacteria and yeasts. Efficacy tests with these organisms should always be provided.

For uses in industrial dish washers for specific purposes, an exception can be made when sound justification is provided. This will be evaluated on a case-by-case basis.

For all other groups of organisms, data only have to be provided when activity against those organisms is claimed. Relevant groups of test organisms, next to bacteria and yeasts, can be fungi (fungal spores), viruses, bacteriophages, and bacterial spores. Bacteriophages are mainly of importance in the dairy industry.

The test organisms used in efficacy tests are normally stated in the applicable standard test methods. For dish washing disinfectants *Salmonella* spp., *Listeria* spp. and *Campylobacter jejuni* are relevant target organisms. For products which claim general efficacy against bacteria, the standard test bacteria should be tested. For these products efficacy against *Salmonella* spp., *Listeria* spp. and *Campylobacter jejuni* is assumed, because they are more susceptible than the standard test bacteria.

The EN standards for food area only include a test on bacteriophages but not for other viruses. To demonstrate a general virus claim a modified EN phase 2, step 1 test (medical area test with food area soiling) can be provided with Adenovirus and Murine Norovirus as test organism and a DVG phase 2, step 2 tests.

When the product is intended to be used at high temperatures (>40 ºC) relevant test organisms for these temperatures should be used as described in section 5.4.0.4.4 of this Guidance.

An overview of reference test organisms, also for high temperatures, is given in Appendix 3.

**Test conditions**

It is important that the tests are performed with the same contact time as claimed on the label. The claimed contact time has to be a realistic value. It will depend on the contact time for the disinfection cycle in (dish)washing machines. Justification for the used contact time should be given.

In general, dish washing disinfectants should be tested under dirty conditions since these products are mainly used for combined cleaning and disinfection. Tests under clean conditions will only suffice when the label instructions state that cleaning prior to disinfection is necessary or when this is incorporated in a previous cycle of the (dish)washing machine. If this is not stated on the label the test should be done under dirty conditions.

Tests should be carried out with soiling for clean or dirty conditions in accordance with the test requirements.
Note that for use in specific industries different types of soiling for dirty conditions should be used.

The soiling needed for clean and dirty conditions can be found in the relevant EN tests or EN 14885 (version 2014 or later) and referenced in Appendix 4.

For products intended to be added to (dish)washing machines, information on the following in-use conditions should be provided:

- the concentration of the product (or at least the active substance) in the water during disinfecting process (i.e. washing or rinsing). The water volume used can differ between wash and rinse cycle and different washing programmes, but also between dish washing machines;
- the water to dishes ratio in the test is an important factor that should reflect the in-use conditions;
- the temperature during the disinfection process (high when added in wash process, low in rinse process);
- the contact time (differs between various washing programmes and washing machines).

The laboratory tests should be performed under these conditions. The conditions for effective disinfection can normally only be carried out in professional dish washing machines. If the exact conditions cannot be met, for example, in household machines, reasonable worst case conditions must be tested.

Worst case conditions, e.g.:

- the lowest temperature;
- the highest volume of water (i.e. maximum dilution of the product);
- the shortest contact time;
- the maximum load of dishes (i.e. smallest water to dishes ratio).

The test temperature should be according to the use instructions on the label.

When the product is used at lower temperatures (e.g. only for rinsing after normal dish washing with hot water) tests can be done at 20ºC. When disinfection is done at temperatures of 40 to 80 ºC the test temperature should be in accordance with the use and relevant test organisms should be used (see section 5.4.4.5.2. of this Guidance).

5.4.4.5.3 Acceptance criteria

A product will be assessed to be sufficiently effective if the required laboratory and simulated-use or field tests have been performed (using the required test organisms and test conditions), and when the pass criteria for the tests have been met.

Where pass criteria are available in the standard test these should be met. For PT4 products the required log_{10} reductions tests are referenced in Appendix 4.

Deviations from the pass criteria are possible, but must be justified in the application. The CA will evaluate any justification on a case-by-case basis, consulting the other CAs where appropriate, and decide whether it is acceptable or not.
5.4.4.6 Disinfection of inner surfaces in human drinking water systems

5.4.4.6.1 Introduction

Biocides can be used to disinfect surfaces in human drinking water systems (further referred to as drinking water. This can be large water systems in drinking water companies, transport pipes in between drinking water companies (semi-finished product), the communal piping system, collective drinking water systems (hospitals and other health care facilities, hotels, penitentiary institutions, etc.), and tanks and reservoirs for drinking water (for instance on ships).

When water systems are disinfected in closed circuits, after which the system is washed with clean water, it is considered to be disinfection of the pipework and is included in PT4. When disinfection is performed in water systems while they are in service and the water is also disinfected the application is considered to be included in PT5.

The drinking water systems may be new or rehabilitated drinking water pipes (e.g. in newly built or renovated houses) or systems that are in service for some time and have become contaminated during this period.

The main need to clean and disinfect the systems is to get a fresh start of the system. Cleaning and disinfection programs may be combined to treat these systems.

The systems that have been in service for some time contain biofilm and organisms to be controlled might hide in this biofilm. For instance, *Legionella* can multiply in the biofilm.

5.4.4.6.2 Data requirements

Test methods

For efficacy testing of biocidal products used on inner surfaces of drinking water systems, the tiered approach as described in section 5.4.0.4.1 of this Guidance is preferred.

For combined cleaning and disinfecting of drinking water pipes, the following test is normally required:

- a quantitative suspension test (phase 2, step 1).

When efficacy against *Legionella* is claimed, the following tests are normally required:

- a quantitative suspension test (phase 2, step 1);
- a simulated-use test or a field test.

all simulating practical conditions appropriate to its intended use (temperature, soiling, contact time, etc.). When ring trial validated test protocols for simulated-use tests (phase 2, step 2) become available these might replace the field trial.

When efficacy against biofilms is claimed, the following tests are normally required:

- a quantitative suspension test (phase 2, step 1);
- a simulated-use test or a field test.

Laboratory tests

EN phase 2, step 1 tests for the food industrial, domestic and institutional area are relevant for this use. Efficacy against *Legionella* can be tested in EN 13623 (phase 2, step 1).

See section 5.4.2.11 of this Guidance for biofilm test methods.

Appendices 2 and 4 give a list of recommended test methods.
Field trials

For products which claim efficacy against *Legionella*, field trials with the following requirements should be provided:

- before testing it should be established that the installation contains high numbers of *Legionella* (>100 cfu/L). A zero-time measurement should be performed. Systems must not be inoculated with micro-organisms in order to perform the efficacy test;
- a field trial should be performed in a system that has been in service for some time and has become infected during this period;
- the number of sampling points per location will depend on the number of draw-off points in the installation. The table below should be used;

**Table 7: Number of sampling points**

<table>
<thead>
<tr>
<th>Number of draw-off points (outlets)</th>
<th>Number of sampling points</th>
</tr>
</thead>
<tbody>
<tr>
<td>10-100</td>
<td>4</td>
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<tr>
<td>101 – 200</td>
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<td>401 – 800</td>
<td>10</td>
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<tr>
<td>801 – 1600</td>
<td>12</td>
</tr>
<tr>
<td>&gt; 1600</td>
<td>14</td>
</tr>
</tbody>
</table>

* a draw-off point is a point where drinking water, household water or warm water is made available for use.

- after disinfection and subsequent washing of the system with clean water (removal of disinfectant), samples should be taken and the amount of bacteria (general) and *Legionella* in the water should be determined. Samples should be taken 48 hours and 2 weeks after disinfection;
- after treatment, water from none of the sampling points should contain more than 100 colony forming units/litre *Legionella*.

Test organisms

Biocidal products for drinking water disinfection should be at least sufficiently effective against bacteria. The test organisms used in efficacy tests are stated in the applicable standard test methods. Efficacy tests with these organisms should always be provided.

For products which claim efficacy against *Legionella*, a test with *Legionella* spp. should also be performed.

For all other groups of organisms, data only have to be provided when activity against those organisms is claimed.

Test conditions

It is important that the tests are performed with the same contact time as claimed on the label. The claimed contact time has to be a realistic value.

Laboratory phase 2, step 1 tests should be carried out with soiling for clean conditions in accordance with the test requirements. The soiling needed for clean conditions can be found
in the relevant EN tests and referenced in Appendix 4. Simulated-use tests should be performed with relevant soiling.

5.4.4.3 Acceptance criteria
A product will be assessed to be sufficiently effective if the required laboratory, or when applicable, field tests have been performed (using the required test organisms and test conditions), and when the pass criteria for the tests have been met.

Where pass criteria are available in the standard test these should be met. For these products the required \( \log_{10} \) reductions in the laboratory tests are referenced in Appendix 4. The field trial should not contain more than 100 colony forming units *Legionella* per litre.

5.4.4.7 Disinfection of inner surfaces in veterinary water systems

5.4.4.7.1 Introduction
Biocides can be used to disinfect surfaces in veterinary water systems in farms, bio-industry, etc.. These are water systems provide water for animals to drink, to prepare feed, and to use for cleaning the area. Water systems that are also suitable for human drinking water are not included in this section (see previous section of this Guidance).

When water systems are disinfected in closed circuits, after which the system is washed with clean water, it is considered to be disinfection of the pipework and is included in PT4. When disinfection is performed in water systems while they are in service and the water is also disinfected the application is considered to be included in PT5.

The water of these systems can be provided by drinking water companies but can also contain well, ground, or ditch water that is pumped up at the location, or other water. Water systems in livestock farming can be used to supply food additives or antibiotics to the animals. Therefore, these veterinary water systems may be more fouled than human drinking water systems.

5.4.4.7.2 Data requirements

Test methods
For the combined cleaning and disinfecting of veterinary drinking water pipes (e.g. water tanks, water in animal housings etc. used as drinking water for animals and for other uses in stables like cleaning, preparing feed, etc.), efficacy should be demonstrated in a tiered approach as described in section 5.4.0.4.1 of this Guidance. This includes a phase 2, step 1 and step 2 test.

The following documents are recommended for disinfecting of veterinary drinking water pipes:

- EN 14885 gives an overview of which EN phase 2, step 1 and step 2 tests to use for different uses, the tests (bactericidal) for the food area are relevant for this use;
  - if CEN standards are not relevant or available for the use or organisms claimed the following documents are recommended if appropriately reflecting the application:
    - OECD guidance for the testing of chemicals: Quantitative method for evaluating activity of microbiocides used on hard non-porous surfaces. (These are surface tests which would be considered phase 2, step 2 tests).

The use of the specified tests is strongly recommended where they are relevant and appropriate.
When efficacy against biofilms is claimed, a simulated-use test or field test has to be performed, as well as a phase 2, step 1 test. See section 5.4.2.11 of this Guidance for test methods.

**Test organisms**

Biocidal products for drinking water disinfection should be at least sufficiently effective against bacteria. The test organisms used in efficacy tests are stated in the applicable standard test methods. Efficacy tests with these organisms should always be provided.

For all other groups of organisms, data only have to be provided when activity against those organisms is claimed.

**Test conditions**

It is important that the tests are performed with the same contact time as claimed on the label. The claimed contact time has to be a realistic value.

Laboratory tests should be carried out with soiling for clean or dirty conditions in accordance with the test requirements for the food area. Tests under clean conditions will only suffice when the label instructions state that cleaning of the water systems prior to disinfection is necessary. If this is not stated on the label the test should be done under dirty conditions.

**5.4.4.7.3 Acceptance criteria**

A product will be assessed to be sufficiently effective if the required laboratory, or when applicable, simulated-use or field tests have been performed (using the required test organisms and test conditions), and when the pass criteria for the tests have been met.

Where pass criteria are available in the standard test these should be met. For these products the required log_{10} reductions in the laboratory tests are referenced in Appendix 4.

**5.4.4.8 Other uses in PT4**

Several uses of PT4 products have been specified in the above sections and data requirements and acceptance criteria for these uses are described. For products with other uses, that do not fit in one of the described uses, it is up to the applicant to demonstrate efficacy in an appropriate way.

In general the tiered approach as described in section 5.4.0.4.1 of this Guidance is preferred. Where possible the standard tests required for the described uses should be taken (e.g. EN phase 2, step 1 and step 2 tests for food area). Where the tests are not appropriate for the product, other tests can be used. In that case, a justification for the relevance of the tests used should be provided. The test design should be discussed with and agreed by the CA before testing takes place. The evaluation will be done on a case-by-case basis by the CAs.

The guidance will be updated when new methods become available.
5.4.5 PT5 Drinking water disinfectants

**NOTE to the reader:**

A preliminary draft text for PT 5 is included in this section: the section is currently under review within the ECHA “Disinfectants Project”.

This project will address the efficacy assessment of PT 5 and will be the subject of an update to this guidance foreseen for 2017/2018. The finalised agreed guidance document will replace, in whole or part, this section.

In the meantime, this Section 5.4.5 is available to readers for information and if there are other suitable means of demonstrating efficacy for PT 5 products, these may be used.

### 5.4.5.1 Introduction

Product type 5 contains biocidal products used for the disinfection of drinking water for both humans and animals. Definition of drinking water is according to article 2 of Council Directive 98/83/EC of 3 November 1998 on the quality of water intended for human consumption. In this section the term drinking water for humans is not only used for water that will be consumed directly but also for other uses of water coming out of the plumbing system like showering, cooking, etc..

When disinfection is done in the water systems while it is in service also the water is disinfected and this is included in PT5. When water systems are disinfected in closed circuits, after which the system is washed with clean water, this is disinfection of the pipework and included in PT4.

Disinfectant products can be added to drinking water, intermittently, by shock dosing or continually. The purpose of disinfection is to disinfect the water to prevent transmission of water-borne diseases via drinking water. Water-borne transmitted pathogens can be bacteria, viruses, yeasts, fungal spores and protozoan parasites. Disinfection is only one aspect of drinking water treatment. Application of drinking water disinfectants is associated with the responsibility to control toxic disinfectant by products. Treatment substances should only be added for specific hygienic or technical reasons, limiting application to the minimum volumes that are absolutely necessarily for achieving the targeted effect (principle of minimisation) and only under conditions optimizing their efficacy.

Disinfection within PT5 can be divided into five groups:

1. **Disinfection in drinking water companies**
   This is disinfection of water when it enters the drinking water company, transport in between drinking water companies (semi-finished product) and prior to distribution into (part of) the communal piping system.

2. **Disinfection in collective drinking water systems**
   This is disinfection in collective drinking water systems like hospitals and other health care facilities, hotels, penitentiary institutions, etc.. In these large plumbing systems water might become contaminated with *Legionella*. When physical techniques (heating, UV treatment, etc.) are insufficient chemical disinfection is allowed in some EU countries.

3. **Disinfection of stationary water in reservoirs**
   This is disinfection of water stored in tanks and reservoirs, for instance on ships.
4. Disinfection of undefined water before drinking.
   This is disinfection of for instance individual emergency water supply or other water
   that might be contaminated in places where no clean drinking water is available.

5. Disinfection of veterinary water
   This is disinfection of water in animal housings used as drinking water for animals and
   for other uses in stables (cleaning, preparing feed, etc.).

In the sections below the requirements and acceptance criteria for most common uses are
specified. For other uses and claims that are not specifically mentioned the requirements will
be set on a case-by-case basis by the CAs.

5.4.5.2 Data requirements

5.4.5.2.1 Test methods

For an overview of available EN tests see Appendix 1.

Disinfection in drinking water companies

For product authorisation of drinking water disinfectants in drinking water companies the
tiered approach as described in section 5.4.0.4.1 of this Guidance is preferred.

Next to a suspension test (EN phase 2, step 1 test) a simulated use test should be
performed.

For the simulated use test a detailed appropriate test method is given in the test method
"Quantitative determination of the efficacy of drinking water disinfectants", available on
ECHA Biocides Efficacy Working Group webpage. The test is realized on an adapted test
rig. A disinfectant neutralizer or filter system to stop reaction between disinfectant and test
organisms is required.

Disinfection in collective drinking water systems

For product authorisation of drinking water disinfectants in collective drinking water systems
the tiered approach as described in section 5.4.0.4.1 of this Guidance is preferred.

Because the control of Legionella in collective drinking water systems is of major
importance, efficacy against Legionella should always be demonstrated.

The following requirements are set for biocides to be used as disinfectant in drinking water
systems:

Laboratory tests
Basic efficacy of the product should be demonstrated in a suspension test (phase 2, step 1). Studies should show that the product can accomplish a log reduction of 5 against bacteria
and specifically Legionella. This can be done in laboratory tests (e.g. suspension tests
EN 1276 and EN 13623). The suspension tests can be waived when simulated use or field
trials are available in which the concentration of Legionella is high enough to show log
reduction of 5 (min. $10^5$ cfu/L).

Simulated use tests
A simulated use test should be performed. For the simulated use test a detailed appropriate
test method is given in the test method "Quantitative determination of the efficacy of

drinking water disinfectants”) available on ECHA Biocides Efficacy Working Group webpage\textsuperscript{18}. The test is realized on an adapted test rig. A disinfectant neutralizer or filter system to stop reaction between disinfectant and test organisms is required.

Field trials
For products with long and continuous use (drinking water disinfection PT5) field trials with the following requirements should be provided:

\textit{Locations}
A field trial should be performed at a minimum of 3 locations. 

Only locations with 100 or more operational draw-off points (downstream of the application spot) are acceptable. A location is a collective drinking water system which is treated by the product. Also part of a collective drinking water system, for instance a wing of a building or only the cold water system can be seen a test location, as long as it contains 100 or more operational draw-off points.

The drinking water quality in the different EU countries may differ. In some EU countries disinfectants like chlorine are standard included, while in other countries disinfectants are only added during calamities. Therefore some EU countries will accept only field trials in their own country or on locations with comparable water specifications. Therefore, when tests are not performed in all countries for which authorisation is applied for, the quality of the tested drinking water should be specified and the comparability of this water is to the drinking water in each country should be justified. The CA will decided whether the test is acceptable or not.

\textit{Duration of the test}
When the apparatus is in continuous or discontinuous use (so no single applications) the duration of the test is one year per location, starting from the first sampling round after starting the apparatus. When, due to starting problems etc. the first months do not give the required result (see 2.6), the test should be extended to one year starting from the point that a stable situation is reached. In this way at least a year of test results can show that the product is capable of controlling \textit{Legionella}.

\textit{Different types of water}
It is recommended that the locations are spread over the country, this to ensure that the product is tested on different types of water. For this purpose information should be provided to the Ctgb on the quality of the provided water at the different locations. In principal this information is available through the water company.

\textit{Legionella}
Before starting a test it should be clear that the installation to be treated is contaminated with Legionella bacteria. For this purpose information should be provided to the Ctgb on (recent) problems with \textit{Legionella}, like results from sampling in the past and performed cleanings, etc..

\textit{Sampling points}
The amount of sampling points per location depends on the amount of draw-off points (taps and other outlets) in the installation. The table below should be used (taken from appendix G of the Waterleidingbesluit).

\textsuperscript{18}http://echa.europa.eu/about-us/who-we-are/biocidal-products-committee/working-groups/efficacy.
Table 8: Number of sampling points

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<tr>
<td>&gt; 1600</td>
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</table>

All sampling points should be unambiguously coded.

At each sampling round two sampling points are sampled each time (standard sampling points), preferably the sampling point next to the apparatus and the sampling point the most far away from it. These sampling points should be clearly described and the code of these points should be stated. All other sampling points should vary at each sampling round. When a sampling point shows elevated values of *Legionella* or one of the other parameters this sampling point should be sampled again the next month. The total amount of sampling points stays the same, according to the table above.

The tuning of the apparatus at the time of sampling should be recorded.

*Efficacy*

To be able to evaluate the efficacy the following measurements should be performed:

- zero measurement: measurement of *Legionella*, total hardness, pH, and active substances before the apparatus is put into action.
- *Legionella*, monthly sampling, norm value 100 cfu/l (90%-percentile with a maximum of 1000 cfu/l);
- total hardness, Ca, Mg; sampling once per four months, depending on the variation a higher frequency might be necessary; also data from the waterworks companies can be collected;
- pH, monthly sampling on both standard sampling points, or data from the waterworks companies can be collected.

*Active substances and metabolites (side effects)*

To determine the amount of active substance in the water and any harmful metabolites, the relevant products should be measured monthly.

For anodic oxidation the following products are relevant:

- available chlorine, monthly measurements; norm value 0.3 mg/l at the draw-off points (90 %-percentile with a maximum of 0.5 mg/l)
- trihalomethanes: measurement 3 and 9 months after the apparatus is put into action, always at one draw-off point which represents the worst-case situation, normally the draw-off point the most far away from the apparatus. This concerns the parameters trichloromethane (chloroform), tribromomethane (bromoform), bromdichloormethane and dibroomchloormethane. Norm value: the total of the
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trihalomethanes 25 μg/l (90%-percentiel, max. 50 μg/l). The concentration broomedichloormethane should not exceed 15 μg/l.

- halogenated acetic acids: measurement 3 and 9 months after the apparatus is put into action, always at one draw-off point which represents the worst-case situation (see trihalomethanes). This concerns the parameters monochloric acid, dichloroacetic acid and trichloroacetic acid. The norm value: the total of the haloacetic acids 25 μg/L.

For copper/silver ionisation the following products are relevant:

- copper, monthly measurements; norm value 2 mg/l;
  
  Remark: the technique cannot produce the full 2 mg/l considering the contribution of copper from other sources. An increase of the copper value of maximum 1 mg/l is considered acceptable.

- silver, monthly measurements; norm value 50 μg/l (90%-percentiel with a maximum 100 μg/l).

For chlorine dioxide-generators the following products are relevant:

- chlorite, monthly measurements at all draw-off points; norm value 0.2 mg/l;

- chlorate, monthly measurements at all draw-off points; norm value 0.2 mg/l;

- trihalomethanes: measurement 3 and 9 months after the apparatus is put into action, always at one draw-off point which represents the worst-case situation, normally the draw-off point the most far a way from the apparatus. This concerns the parameters trichloromethane (chloroform), tribromomethane (bromoform), broomedichloormethane and dibroomchloormethane. Norm value: the total of the trihalomethanes 25 μg/l (90%-percentiel, max. 50 μg/l). The concentration broomedichloormethane should not exceed 15 μg/l.

Evaluation criteria per location

For the evaluation of the results of the measurements the norm values as mentioned in 6.2.1.2.2.6 and 6.2.1.2.2.7 are used. Per location 90% of the measurements should fulfil the requirements. Over all locations together 90% of the locations should fulfil the requirements.

General requirements for study reports

Every study report should contain a good description of material (location, number of draw-off point, sampling points, history of Legionella, etc.), method (starting date, tuning of the apparatus) and results (including 0-measurement). In the study reports of the field tests the results should be interpreted per location. Remarks like for instance high values above the norm, should be mentioned and explained. The report should be closed with a conclusion.

Apparatus

In case of in situ production of the active substance or when an apparatus is used to dose the active substance in the right amount to the water, the report should contain information on safety measurements concerning over and under dosing.

Disinfection of stationary water in reservoirs

For this use it is in most cases acceptable to demonstrate efficacy with phase 2, step 1 tests only. In some cases efficacy against biofilm is of importance in this use. For testing efficacy against biofilms see section 5.4.2.11 of this Guidance.
Disinfection of undefined water used as drinking-water

For this use it is in most cases acceptable to demonstrate efficacy with phase 2, step 1 tests only.

Disinfection of water for animals

For efficacy testing of disinfectants for water for animals the tiered approach as described in section 5.4.0.4.1 of this Guidance is preferred. Next to a phase 2, step 1 test also a simulated-use test or field test (phase 3) should be performed, to provide information under in-use conditions. In some cases efficacy against biofilm is of importance in this use. For testing efficacy against biofilms see section 5.4.2.11 of this Guidance.

5.4.5.2.2 Test organisms

PT5 products should be at least sufficiently effective against bacteria. Efficacy test with these organisms should always be provided.

There are few exceptions to this rule:

- for products for disinfection of drinking water in drinking water companies also efficacy should be demonstrated against bacteria and viruses. This can be done in the simulated use test.
- for products for disinfection of drinking water in collective systems also efficacy should be demonstrated against Legionella spp..

For all other groups of organisms test only have to be provided when efficacy against the organisms are claimed.

The test organisms used in efficacy tests are normally stated in the applicable standard test methods. An overview of reference test organisms is given in Appendix 3.

5.4.5.2.3 Contact time

It is important that the tests are carried out with the same contact time as claimed on the label.

The claimed contact time has to be a realistic value. For the use as drinking water disinfectant no maximum contact times are set.

5.4.5.2.4 Soiling

The suspension test (phase 2, step 1) should be carried out with soiling for clean or dirty conditions in accordance with the test requirements. Depending on the water source that has to be disinfected the test should be performed under either clean or dirty (e.g. undefined or pumped up water) conditions.

- Dirty conditions: 3 g/L bovine albumin solution
- Clean conditions: 0.3 g/L bovine albumin solution

5.4.5.3 Acceptance criteria

A product will be assessed to be sufficiently effective if the required laboratory, or when applicable, field tests have been carried out (using the required test organisms and test conditions), and when the pass criteria for the tests have been met.

Where pass criteria are available in the standard test these should be met. For PT5 products the required log reductions in suspension tests are referenced in Appendix 4.
The pass criteria for the simulated-use test are stated in the test (see Appendix 4 for more information): after a contact time of 10 minutes at least a log reduction of 2 and after 25 minutes at least a log reduction of 4.

In the drinking water disinfection field tests the aim is to keep the Legionella concentration below 100 cfu/L. Ninety percent of all the test samples per location should show a Legionella concentration below 100 cfu/L while the maximum Legionella concentration should not exceed 1000 cfu/L. The test should be done on ten locations and 90% of these locations should meet this criterion.

Deviations from the pass criteria are possible, but must be justified in the application. The CA will evaluate any justification on a case-by-case basis, consulting the other CAs where appropriate, and decide whether it is acceptable or not.

5.4.6 Materials and Articles Treated to Protect Humans or Animals

For testing materials and articles with claims to protect humans or animals, a tailored approach is compulsory. The testing strategy entirely depends on the specific claim made. In the majority of cases, a claim can only be made for a specific type of final article, as use area and use conditions are decisive for describing the problem which the biocide must solve, and to demonstrate efficacy in exactly those conditions is necessary. Consequently, this section describes testing principles and strategies rather than recommending specific tests.

A tiered approach has to be followed in demonstrating claims for protection of humans or animals:

- **Tier 1 - Proof of principle:** Tier one tests should document the efficacy of the incorporated biocide in the relevant matrix against relevant target organism(s) under relevant conditions (e.g. humidity, temperature).
- **Tier 2 - Simulated Use:** Tier two tests should document the efficacy of the incorporated biocide in the relevant matrix under real-life conditions (e.g. way of contamination, cleaning regimes, time to take effect) and the duration of the effect.

Depending on the claim made (e.g. “kills bacteria on door-handles to prevent cross contamination”, “protects against mosquito-bites”), even Tier 3 testing can be necessary:

- **Tier 3 - In-Use Evaluation/Field studies:** To substantiate health benefit claims, treated and untreated articles would be tested via statistically designed use trials by a representative user group.

Generally, the principle applies that only claims can be made which have been demonstrated.

5.4.6.1 Determining the purpose of the Treatment

The effects of articles with a disinfection claim cannot be detected by changes in appearance, mechanical properties or odour. The precondition for demonstrating efficacy is a clear description of the purpose of the treatment. Often, claims are unclear about whether the treatment prevents growth or kills bacteria on contact. On most articles, no bacteria will grow under normal conditions of use. Nevertheless, antibacterial claims (such as ‘anti-bacterial’, ‘hygienically clean’, ‘free of bacteria’, ‘prevents the spread of hazardous bacteria’) are made, insinuating that bacteria will be killed on the material, though only growth
inhibition tests have been carried out. In most environments, the sheer presence of bacteria does not present a problem. If this is a problem, it is in most cases much more effective to use traditional disinfection methods with a liquid disinfectant. In most cases, the treatment of articles should not be used as the only measure of disinfection, but should be combined with a disinfection management regime.

**5.4.6.2 Effects Intended to Inhibit Microbial Growth**

Under the majority of indoor situations, most microorganisms will not grow on environmental surfaces due to lack of humidity. To make a claim for growth inhibition, wet or at least humid conditions are a precondition, unless otherwise justified. To demonstrate such a claim, sub-samples of treated and untreated material of the article in question could be tested using a method adapted from ISO 22196 (see Figure 3). Soiling conditions, temperature, test species and contact time have to be adapted to mimic a realistic in-use situation (Tier 1). The impact of in-use conditions like ageing or cleaning regimes on the effect would have to be included in the testing (Tier 2). The minimum requirements for disinfection are laid down in the Claims matrix for treated articles (see Appendix 1) with claims to protect humans or animals [http://echa.europa.eu/about-us/who-we-are/biocidal-products-committee/working-groups/efficacy].

**Figure 3: A Test for Antibacterial Activity in Wet Conditions**

ISO 22196

ISO 22196, Method Outline:

An aliquot (usually 400 μl) of a log phase bacterial cell suspension (ca10⁶ cells ml⁻¹) in ¹/₅₀₀ Nutrient Broth are held in intimate contact with each of 3 replicates of both treated and untreated variants of the test materials using a 40 mm x 40 mm polyethylene film (e.g. cut from a sterile Stomacher bag) for 24 hours at 35°C. Usually, *S. aureus*, *P. aeruginosa*, *Enterococcus hirae* and *E. coli* should be tested (see Appendix 3). The populations are then
recovered using a neutraliser solution and the size of the surviving populations are determined as colony forming units (CFUs) using a dilution plate count method. Additional replicate unfortified samples are also inoculated in the same manner but are analysed immediately to determine the size of microbial population present prior to incubation. The differences between the initial and final population as well as between the treated and untreated materials are used to assess the basic antibacterial properties of the test materials.

5.4.6.3 Effects intended to Kill Microorganisms through Contact

Claims made for materials and articles to kill on contact to prevent cross-contamination are not easy to demonstrate. Mostly, the effect will require the release of the active substance from the surface of the material; this release needs to be triggered somehow. In the majority of cases, water or other liquids are the crucial component to facilitate such release and transfer. If the event that caused the deposition of the target organism does not introduce moisture and the normal exposure conditions of the material or article are dry (or only subject to normal, ambient indoor humidity), the effect of the treatment will probably be limited.

Another issue is the speed of activity needed to inhibit cross-contamination. If for instance door handles in a hospital would be treated with an active substance to kill deposited pathogenic organisms, the effect would have to be sufficiently fast to prevent the next person using the door handle from cross-contamination. In combination with the little moisture which is deposited in the event, it will be challenging to demonstrate a satisfying effect. The minimum requirements for disinfection are laid down in the Claims matrix for treated articles (see Appendix 1) with a claim to protect humans or animals. Additional requirements may apply depending on the claim made.

Testing could be carried out using protocols such as those given in Figures 4, 5 and 6 below. Again, care must be taken to adapt test conditions to realistic in-use conditions. Figures 4 and 5 show the approach used for non-porous materials and for absorbent materials, respectively, both intended to simulate contamination through contact with splashes of contaminated liquids. Figure 6 illustrates a protocol intended to simulate contamination through, for example, hand/gloved hand contact.

5.4.6.4 Acceptance Criteria

The performance criteria for treated articles can be found in the Claims Matrix for treated articles (Appendix 1). For choosing test organisms please refer to the liquid disinfectants (Appendix 3). As the performance criteria for treated articles are lower than for liquid disinfectants, the treatment of articles should generally not be used as the only measure of disinfection, but should be combined with a disinfection management regime.
Figure 4: Simulated Splash Model Non-Porous Materials

CFU= colony forming units

Prepare replicate test-pieces (4 per contact time per treatment)

Inoculate Test Pieces (30 x 30 mm) with 100 µl of Cell Suspension each

Incubate without cover at 24°C for 5-60 min at 60% Relative Humidity

Cell suspension (ca $10^6$ cells ml$^{-1}$)

Constant Humidity Chamber

Transfer Test Piece to Neutraliser in Stomacher Bag

Determine CFU for each contact time
Figure 5: Simulated Splash Model Porous Materials

CFU = colony forming units, RH = relative humidity, BSA = Bovi ne Serum Albumine

Prepare replicate swatches (4 per contact time per treatment) and place on sterile grid to allow good air circulation.

Suspension (ca $10^5$ CFU ml$^{-1}$) of test species in BSA solution (15 g litre$^{-1}$)

Swatches inoculated with 100 µl

Incubated for 5 – 60 min at 24°C & 50% RH

Determine CFU

Individual swatches transferred to 10 ml neutraliser at intervals
Figure 6: Printing Model

TVC= total viable count
CFU= colony forming units

Prepare Cell suspension (ca $10^5$ cells ml$^{-1}$)

Inoculate Large Bioassay Dish containing $\frac{1}{10}$ strength Nutrient Agar with 10 ml of cell suspension

Incubate for 18 Hours at 25°C

Transfer Inoculum to Test Pieces

Recover cells in Neutraliser and Determine TVC

Incubate Inoculated Test Piece at 20°C for 5 min then Determine CFU

Transfer Inoculum to Test Pieces in Pairs 1000 g

Lift Cells from Bioassay Plate 100 g

Determine TVC Prior to Incubation

Plate replicator

Locking ring
Table 9: Protection of Humans or Animals – Example Claims, Problems and Testing Approaches

<table>
<thead>
<tr>
<th>Claim</th>
<th>PT</th>
<th>Proof required</th>
<th>Example method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bedside cabinet for use in hospitals that has been treated to reduce infections by killing ‘bacteria on contact’.</td>
<td>2</td>
<td>Data should show that microorganisms, when deposited through skin contact (even under simulated conditions) and through the deposition of fine aerosols are killed within a time-frame that would prevent the surfaces becoming a vector for cross-contamination.</td>
<td>Plaques made of the identical material used for the cabinet are employed in the test. Both treated and untreated variants are used.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Skin contact</td>
<td>The method described in Figure 6 is employed to deposit bacteria onto test plaques. A range of contact times between 5 minutes and 1 hour are used. A log reduction of 3 should be achieved.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Aerosol</td>
<td>The method described in Figure 4 is adapted for use by employing multiple droplets of 1 µl on each test plaque. A range of contact times between 5 minutes and 1 hour are used to explore activity. A log reduction of 3 should be achieved.</td>
</tr>
<tr>
<td>A plastic conveyer belt is treated to prevent the growth of bacteria between cleaning intervals in a food factory.</td>
<td>4</td>
<td>Data should show that relevant bacteria grow on an untreated conveyer belt under normal conditions of use during a 6 hour interval. Significantly reduced growth should be demonstrated on the treated belt.</td>
<td>Plaques made of the identical material used for the belt are employed in the test. Both treated and untreated variants are used.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>ISO 22196 is adapted to simulate a moist conveyor belt. A soiling agent relevant to the end use is included. A contact time and temperature equal to that encountered in practice are employed.</td>
</tr>
</tbody>
</table>
Table 10: Basic Requirements for a Valid Test Protection of Humans or Animals

<table>
<thead>
<tr>
<th>Requirement</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>i.</td>
<td>The test should be carried out on the type of final article.</td>
</tr>
<tr>
<td>ii.</td>
<td>A test which mimics the way of deposition and the type of material needs to be chosen.</td>
</tr>
<tr>
<td>iii.</td>
<td>An untreated variant of the test material must be included such that the impact of the treatment can be demonstrated.</td>
</tr>
<tr>
<td>iv.</td>
<td>Test conditions should reflect normal conditions of use in terms of humidity, temperature, soiling, contact frequency, etc.</td>
</tr>
<tr>
<td>v.</td>
<td>The test should employ organisms that are relevant to the end use of the article and the purpose being claimed.</td>
</tr>
<tr>
<td>vi.</td>
<td>Tests that employ a single species of organisms should be favoured over those that use consortia.</td>
</tr>
<tr>
<td>vii.</td>
<td>Minimum of three replicate test pieces of both treated and untreated materials should be employed (unless justified).</td>
</tr>
<tr>
<td>viii.</td>
<td>The final data should include either some indication of the impact of service conditions on the performance of the treated material/article or data from an ageing study. The intention is to demonstrate how long the claimed effect will be sustained.</td>
</tr>
<tr>
<td>ix.</td>
<td>If claims are made which require a field test, relevant data including statistical evaluations have to be provided.</td>
</tr>
</tbody>
</table>
5.5 Preservatives (Main group 2)

General
Preservatives in main group 2 are intended to prevent the biodeterioration of a material or a matrix. Wood can lose stability by the action of microorganisms or insects, fabric can be destroyed by fungi, and even polymer-based plastics are prone to biological deterioration. Plasticised PVC would soon become fouled by surface growths of fungi, lose plasticity and crack without the inclusion of a fungicide. A water-based paint, free of volatile organic compounds (VOCs), could not be stored without the use of a biocide. Polyurethane, for example as used for the soles of shoes, can become colonised by fungi and actinomycetes. The heat exchangers in cooling towers have to be kept free from microbial growth to enhance performance by treatment of the cooling liquid.

This section covers the group of preservatives (PT6 to PT13) and the following sections (5.5.1-5.5.3) apply to all PTs (or as indicated in the headings). For PT8, the guidance is more developed and includes standard tests, which is not the case for the other PTs: PT8 is the exception and section 5.5.8 is dedicated to PT8.

5.5.1 Distinction between preservation/curative treatment and disinfection
Preservatives are directed towards the protection of a material. If the material itself is not affected by the target organisms, the claim does not belong in main group 2. The aim of preservation is to prevent microbial spoilage, decay or the accumulation of biomass that is detrimental to the functionality of an item, material or system. Detrimental effects can be caused by proliferation of cells or by the metabolic activity of cells and may not necessarily involve cell multiplication. The presence of microorganisms can result in either a degradation of the matrix in which they are present or damage to the system in which they are present either due to their metabolic activities (e.g. corrosion) or by fouling or blocking pipes, forming biofilms on heat exchangers etc. It is not the intention of preservatives to transfer their effects to other materials, humans or animals, but to protect the material itself. A long-term effect is generally required. A preservative can have a reversible effect on microorganisms (e.g. by causing stress or cell damage without total loss of viability). In contrast to disinfection no level of reduction is defined for a set of predefined claims.

Curative treatments are also directed towards material protection and therefore likewise fall into main group 2. The aim of a curative action is to either cure microbial spoilage which has already occurred or to eliminate / reduce populations in materials and systems prior to them being treated with a preservative (in some instances a biocidal product can have both curative and preservative functionality).

The level required to prevent spoilage in different media/conditions will be defined by the individual claim made. This will also be the case when the treatment is intended to achieve a curative action.

The claim made will define in which of the PTs an application will fall. The following data is needed:

- A problem description: Scale, speed and type of effect required and what would happen if the biocide was not present
- The target organisms

19 See CA-Sept15-Doc.8.3 – Curative use of preservatives
- Categorisation of the material/matrix to be treated including dose-rate/concentration of the biocide in the material/matrix.
- The intended use pattern of the treated material/matrix including service-life, weathering conditions, leaching (intended or unintended).

**Figure 7: Decision scheme for the distinction between preservation/curative action and disinfection**

<table>
<thead>
<tr>
<th>Is the treatment intended to protect the material/article or its functionality from biological deterioration during storage or in service, extend its durability or prevent odour?</th>
</tr>
</thead>
<tbody>
<tr>
<td>Is the active substance/product intended for curative treatment of wood, industrial liquids, solutions, dispersions or processes?</td>
</tr>
</tbody>
</table>

| Is the treatment intended to protect humans or animals? Is remedial treatment of construction materials with algaecides intended? |

Preservative (Main Group 2)  Disinfectant (Main Group 1)

**5.5.1.1 Curative uses**

Curative uses often require rates and speeds of effects that are similar to those required for disinfectants but do not have prescribed performance standards (with the exception of some PT8 standards). Such uses are nevertheless intended to cure (eliminate or reduce) contamination in materials, matrices or systems. They therefore fall under main group 2. Performance requirements will be defined by the requirements of either the matrix or the process involved. A curative effect and a preservative effect may sometimes be achieved using the same biocidal product, only the concentration may differ. In other cases active substances with curative properties will be combined with those that have preservative. Curative and preservative effects need to be demonstrated separately and different methods need to be employed. When claims are made for curative uses, it is important to carry out the health and environmental risk assessment with any higher doses that may be required. A typical example of curative action is the treatment of a contaminated product prior to packaging and sale (in some cases in addition to a preservative – in other cases the curative product may be capable of achieving both a preservative and a curative effect). Another example is the treatment of a contaminated system by reducing the microbial population it contains to limits that are acceptable to the process (e.g. on a paper mill). Please read more about testing of curative uses in section 5.5.5.1 and 5.5.8.
5.5.1.2 Borderline case: Algaecides

If algae are expected not to destroy the material or damage its function, algaecides are not considered to be preservatives. Thus products used against algae for treatment of swimming pools, aquariums and other waters and for the remedial treatment of construction materials belong to product type 2 in main group 1, whereas products with protective function are considered as products belonging to main group 2.

For example surface coatings for outdoor use are often formulated with both a fungicide and an algaecide. The algaecide, like the fungicide, is performing a preservative function in the coating and is thus covered by PT 7. Similarly, algaecides are incorporated into plastics (e.g. electricity pylon insulation sleeves - to prevent growth that would otherwise cause arcing and system failure) and material used in aquatic and marine environments (including some cementitious materials). Algae are a problem in many water-based cooling systems and water-based process systems (e.g. paper making), where either a preservative or a curative action may be required. Such applications likewise belong to main group 2.

5.5.1.3 Borderline cases: Treated articles

Treated articles can both belong to Main Group 2 or Main Group 1 (and even to Main Group 3 or 4). Please refer to section 5.3 on treated articles and section 5.4.6 on materials and articles treated to protect humans or animals.

5.5.2 Principles for testing preservatives

The aim of any preservation is to maintain the present state/properties of a material or matrix along with its functionality. This can be done in several ways: To determine microbial activity in a biocide-free material, the method of measuring colony forming units is the most common approach to prove that a preservative is needed, i.e. the population needs to be shown to increase in size in the untreated material. The production of a biofilm or an increase in biomass may also be appropriate. Other parameters indicating metabolism can also be documented like e.g. changes in pH, in viscosity, in colour. Data needs to be recorded from the beginning of the test (incubation time 0) and before and after each new inoculation.

Showing growth / metabolism of the microorganisms in the untreated system is an essential requirement of any demonstration of effectiveness of an active substance or biocidal product. It is then assumed, if not proven in every case, that changes have taken place that were induced by microbial growth and that this can be prevented by the use of a biocide acting as a preservative. Often, when growth cannot be proven this is caused by an unnecessarily high inoculation rate. If, at the beginning of the test, an inoculum of for example $10^4$ CFU for bacteria is employed, an increase to $10^5$ - $10^6$ can often easily be shown during the test period. When a higher inoculum density for example $10^6$ is employed, growth is much harder to achieve due to limitations in the supply of nutrient etc. An important consideration is to use a model substrate that can support growth readily rather than attempt to achieve growth in a final product that is less susceptible to the non-acclimated species employed in laboratory tests (i.e. it is often nearly impossible to replicate the failure phenomena observed in practice in a laboratory).

Often a fungicidal or bactericidal claim needs to be supported. For this purpose a species can be tested singly or, as is good practice in many test protocols, in mixed suspensions of either bacterial species or fungal species. Mixing of bacteria and fungi should generally be avoided in these suspensions, but filamentous fungi (“moulds”) and non-filamentous fungi
("yeasts") can be mixed in the inoculum. However, for determining growth different methods need to be applied for yeasts and filamentous fungi.

Many microorganisms are able to form dormant cells or spores to survive unfavourable environmental conditions. These resting cells do not proliferate and show no significant metabolic activity until they find a suitable environment. It is therefore possible that vegetative and active cells, being exposed to an unfavourable environment e.g. a synthetic paint containing solvent or a preservative, are forced into dormancy. Only when a sample of the material is taken out of this environment and is spread onto a nutrient medium do the cells start to grow and to build new colonies. This underlines that the appearance of colony forming units (CFU) on a nutrient media is not necessarily sufficient evidence that growth had been occurring in the matrix used in the test. Growth can only be determined by counting CFU and demonstrating that the number of CFU increased in the untreated matrix during incubation, compared to the number measured immediately after inoculation. The same or a smaller number of CFU than measured initially demonstrates survival, but not necessarily growth. However, for testing solid material, showing growth by adding a nutrient medium to the material is not necessarily enough. It needs to be shown that the material itself is damaged or loses its functionality, or, alternatively, provides growth of microorganisms relevant for the group of organisms which have a negative impact on the stability and/or functionality of the material. Please read more in section 5.5.7.

5.5.3 Tiered approach to testing preservatives

A tiered approach should be followed for testing biocidal products:

Tier 1 - Proof of principle: Tier one tests should document the biocidal efficacy of the incorporated biocide in a relevant model matrix against the target organism(s) under relevant basic environmental conditions (e.g. temperature, humidity).

Tier 2 - Simulated Use: The biocide should demonstrate efficacy under real life conditions relevant to its anticipated service life. Factors such as weathering, UV-stability, extended ageing or leaching should be considered.

Tier 3 – In-use evaluation/field studies: to substantiate specific claims, treated and control articles/products can be tested via statistically designed in-use trials by a representative user group, or by other appropriate methods.

In a Tier 1 test, the damage should be shown in a model matrix and demonstrate how the inclusion of the biocide prevents it (often with the help of an inoculum representing the organisms that cause the damage). In a Tier 2 test, damage or impact of the target organisms under either simulated use conditions or in a manner that simulates an anticipated shelf life should be shown, and even sometimes without the use of an inoculum (soil burial). When moving up from tier 1 to tier 2, a test design has to be more tailored to the field of application envisaged. In tier 1, existing standards are often suitable when the biocide is tested in a relevant matrix with defined organisms and under relevant and reproducible conditions (which are normally only to be found in a laboratory). In tier 2, testing is more complex and often specific standards do not exist. However, sometimes the same standards can be used as for tier 1 tests, simulating use conditions by employing pre-treatment of the matrix. There may be a need for weathering cycles, wind tunnel tests, cleaning regimes etc. Similarly soiling and the influence of other microorganisms can be of more significance. Accelerated aging tests may have to be performed before microbiological testing to allow for factors such as UV, temperature changes, leaching etc. Consideration must be given to which environmental conditions are relevant for simulated aging in realistic in-use conditions. When aging is performed in the field or under in-use conditions,
reproducibility can become a difficult issue, as the aging factors such as e.g. evaporation and soiling are difficult to reproduce and can influence the results. Generally, the applicant should be able to justify how the specific conditions used in testing relate to the in-use conditions relevant to the product or active substance. Tier 3 testing entirely depends on the claim made and is generally for specific uses in case of specific claims. The results have to be relevant for that claim and to be scientifically sound.

5.5.4 Standard Test Methods

A list of the most commonly used standard test methods can be found in Appendices 8, 9 and 10; however, please note that these test methods are not necessarily appropriate to use; they are listed with comments to give an orientation for the assessor as to when and where these tests can be meaningful to prove /support a claim and when they aren’t. In contrast to disinfection, there are no specific tests allocated to the different tiers, with the exception for PT8 where standard-tests are available and tiered testing is defined, (see section 5.5.8 for more information). Often the same test can be employed for tier 1 and tier 2, and only the pre-treatment of the matrix will differ. Different factors can trigger the choice of a test: In some cases the choice of one type of method over another is related to the speed with which it generates results. Often, a method is ‘known’ to be capable of guiding the choice and concentration of a biocide for a certain material through experience within an industry. However, this may not necessarily mean that the method is suitable for demonstrating the claim made.

Care has to be taken as to whether the test method is appropriate for the testing of preservatives, or if it is intended to prove a curative/sanitising activity of a biocide. Generally, for preservative action growth needs to be shown in the untreated controls. The number of replicates required by the methodology is not necessarily 3 replicates; in such cases this needs to be explained and justified.

Nevertheless, an existing test method can form a good basis regarding the parameters of choice of microorganisms, temperature, and choice of neutraliser. If necessary, these methods need to be amended by adding untreated control samples, determining the numbers of organisms that can be recovered immediately after inoculation (0 hours incubation), use of a neutraliser, and the use of a smaller sized inoculum etc. Particularly for tier 2 and 3 testing, it is important that the chosen adaptations reflect the relevant conditions for which the claims must apply.

Specific tests which are recommended for certain uses are described under the sections for the different PTs.

5.5.4.1 Practical aspects for testing bacteria

A relevant study that proves the need for a biocide and its efficacy as a preservative against bacteria must have the following features:

a. The test must be performed in a range of relevant model matrixes that the claim of efficacy is made for (e.g. dishwasher liquid, paints, glues, textiles, etc);

b. The test has to be performed in relevant environmental conditions (temperature, type of matrix, humidity);

c. Control samples without the addition of a biocide must be included during the whole test. These control samples must be handled identically to the other samples, except that they must have no biocide included. The study must include replicate sub-
samples for each treatment (minimum of 3; if less than 3 replicates, then explain and justify).

d. For preservative uses, the control samples should typically show growth (e.g. indicated by an increased number of CFUs) during incubation and this has to be documented. If no growth in the control samples can be seen, this could indicate that only the dormant stages of bacterial cells, without active metabolism, are present in the matrix. The treated samples should show statistically significant effects as compared to the controls;

e. Only if growth cannot be proven by increase in CFU, data concerning other factors like e.g. CO₂-emission, O₂ depletion, change of pH, colour change or disintegration of the matrix should be used to demonstrate the need of preservation of a matrix by the active ingredient or preservative;

f. Relevant bacteria for the intended use have to be tested.

5.5.4.2 Practical aspects for testing fungi

A relevant study that proves the need of a biocide and its efficacy as a preservative against filamentous fungi is in many ways the same as for bacteria, but an attempt to count colony forming units of thread-like mycelia after incubation in liquid systems is bound to fail for several reasons:

- It is impossible to take a representative aliquot from the incubated test vessel since the mycelia tend to conglomerate into pellets of different sizes (often blocking the tip of a pipette).
- Different seized fragments of mycelium and spores that are dormant in the matrix form colonies on a petri dish and their origin cannot be differentiated and so their numbers do not reflect the increase in biomass that has occurred.

However, counting CFU is a practical option to measure the recovery rate of spores inoculated into liquids before spore germination (time 0 analysis) and for unicellular yeasts. At this stage, no mycelia have formed in the liquid, so no fragments will be counted as CFU and wrongly interpreted as growth. Therefore, after the control samples and the biocide-containing samples have been inoculated with spores, the recovery rate can be recorded by measuring colony forming units.

Ascomycetes and fungi imperfecti form thread-like hyphe and spores. Spores serve as dormant stages when environmental conditions are detrimental to growth. When growth conditions are favourable, the spores germinate and form a mycelium and maybe other spores. In liquids the fungal growth tends to form pellets. These can be very small or up to several millimetres in diameter. Furthermore, it is possible that a visible biofilm will accumulate at the sides of the test vessel, e.g. an Erlenmeyer flask or on the surface of the matrix. Both phenomena are visible by the naked eye and clearly demonstrate that the fungus has grown. In highly fluid materials this growth can be quantified by filtering the whole contents of the test vessel and then determining the amount of growth as dry weight. The use of replicates is an important factor in such tests. The number of replicates required by the methodology is not necessarily 3 which is the usual minimum; in such cases this needs to be explained and justified.

For testing solid materials, fungal growth is often assessed by optical appearance, using a rating scale from 0 (no growth) to 5 (>70% cover).
5.5.5 Testing conditions for specific states

5.5.5.1 Wet-state preservation and curative treatments

**Preservation (PT 6, 13)**

Challenge tests are generally employed for preservatives which must preserve liquid matrices, dispersions or fluids used in systems. The inoculum used and the strength of the inoculum depends on which claim must be supported. For preservation claims, growth needs to be shown in the untreated samples and prevention of growth in the treated samples. A larger population (generated by prior growth in an untreated matrix) may be more appropriate for demonstrating a curative effect. Some methods for wet-state preservation are compiled in Appendix 8, however, please note that these test methods are not necessarily appropriate to use; they are listed with comments to give an orientation for the assessor as to when and where these tests can be meaningful to prove/support a claim and when they aren’t.

A series of concentrations of the active substance or the biocidal product should be employed in order to investigate which concentration achieves which level of efficacy. It is likely that the application rate in practice will vary depending on the in-use conditions of a biocidal product even though the matrix is identical, e.g. in a metal working fluid, where the in-use concentration is achieved by diluting the product at the point of use.

**Curative Treatments (PT 6, 7, 11, 12, 13)**

Suspension tests are generally employed for curative treatments of liquid matrices, dispersions or systems. A curative treatment might be applied to a system to reduce a population prior to employing a maintenance regime / treatment (e.g. PTs 11, 12 and 13) or it might be used prior to the addition of a preservative in either a final product, intermediate or a raw material (e.g. PT 6). A model matrix that has been inoculated with microorganisms appropriate to the claim to achieve either growth or a stable population must be treated with the active substance / biocidal product and the effect measured after an appropriate contact time using a dilution plate count (methods described for wet state preservation can be employed to generate the model contaminated matrices / systems). The inoculum can comprise of aerobic or anaerobic bacteria, endospore forming bacteria, yeasts, fungal spores and / or mycelial growth as appropriate to the claim. A log-reduction relevant to the matrix and its use needs to be shown in the treated samples. Viability / growth should be shown to be maintained in the untreated samples. Replicate sub-samples must be employed (minimum of 3, but if the number of replicates required by the methodology is not 3 this needs to be explained and justified) and any differences that result should be shown to be statistically significant. Data from samples treated under field conditions can be used as supporting evidence provided that any effects shown can be attributed to the treatment applied.

5.5.5.2 Protection of solid material: PT 7, 9, 10

This section describes the nature and extent of data which should be made available to support the label claims for biocidal products within PT 7 through PT 10. The common denominator of these PTs is that they concern the treatment of solid material where use conditions can vary considerably, depending on the site and type of use of the material (e.g. treated wood to be used in constant contact with water compared to use in dry conditions; a film preservative to protect a bathroom sealant compared to protecting a house-façade). In contrast to liquid disinfectants or preservatives belonging to PTs 11, 12 and 13, where application often takes place on-site (that is where the target organisms occur), the treatment of materials can take place anywhere, for example where the material is manufactured or at a specific-treatment site. This may not necessarily be within the EU.
Use conditions are much more variable for these product types than they are for liquid disinfectants and liquid preservatives. Often, many different materials can be treated with the same biocide, and even more different articles can be manufactured from the treated materials, which are used in a wide variety of conditions. For instance, water absorption properties of different polymer materials vary and so does the release of the biocide. The concentration of the biocide has to be adapted accordingly. Biocides can be applied as a coating to fabrics or can be incorporated into the material by adding the biocide to the polymer before spinning or extrusion. This alters the fixation in or on the material and has an impact on performance. Materials and articles can be used indoors, outdoors, in wet, humid or dry conditions and at varying temperatures. All of this has an impact on performance. Simulating service life, as length, weathering conditions, temperature, leaching, laundering, etc. is crucial for testing of products within these PTs. Thus, efficacy testing for PT 7 through 10 requires a good description of the frame in which the biocide must perform. In many cases it will be impossible to test every material/substance combination; it might be feasible, however, to categorize different parameters: material, concentrations ranges, use (outdoor, indoor, temperature, humidity, use for load-bearing components, etc.) and to try to test representative, preferably worst-case, examples for every category. It is important though, to describe and justify which range the tested sample represents.

Model matrices
The array of possible material and biocide combinations is vast and phenomena observed in practice cannot always be reproduced in the laboratory. A model matrix has to be chosen which represents a certain type of material and which is relevant to the intended use. For example, plasticised PVC and polyurethane would be useful models for rigid or semi-rigid polymers and a room temperature vulcanised silicone would provide a useful model of a sealant etc. Relevance is the key factor. Thus, if a treatment is intended to protect natural fibres in service then a natural fibre should be employed as the model. When more than one type of material (e.g. plastics, paints and synthetic fibres) can be protected by the biocide, then representative matrices that demonstrate the range of protection should be employed. Different materials can require different biocide concentrations due to varying release behaviour. It is also important to consider what the purpose of the end use is (e.g. in one application the biocide may provide essential protection of a matrix whereas in another it may increase durability). The objective is in any case to support the claims made.

Representative species
The species employed in any test should be relevant to the intended use (i.e. fungi should be employed if the material is affected by fungal growth, odour producing bacteria to be found on the skin should be employed for odour testing, etc.). Consortia rather than individual species should be employed (although mixing bacteria with fungi, algae etc. should, in general, be avoided, see 5.5.2). In exceptional cases, it can be acceptable to use individual species when justified, however, using consortia of microorganisms can be a good option to reflect realistic use conditions but the use of individual species is also acceptable. The species employed in the tests should be relevant to the material under investigation especially where the prevention of the degradation of a material is intended. In many cases the organisms will be specified with the method. Very limited ranges of model organisms should be avoided where possible (e.g. the use of A. brasiliensis as the sole fungus). The test should include replicates (at least three) for both the treated and untreated variants.
### Table 11: Examples

<table>
<thead>
<tr>
<th>Claim</th>
<th>PT</th>
<th>Example Problem</th>
<th>Example Method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fungicide is used to treat paint to prevent causing stains by mould growth in service</td>
<td>7</td>
<td>Painted panels exposed to weather become stained by mould growth and have to be re-painted more often.</td>
<td>BS 3900 Part G6</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Painted panels inoculated with a mixture of spores of fungi known to colonise paints exposed to humid conditions for up to 12 weeks should show visual appearance of fungal growth. The treated sample should be free of it.</td>
<td></td>
</tr>
<tr>
<td>Fungicide is used to treat paper goods to prevent mould growth in service</td>
<td>9</td>
<td>Labels used on wine and beer bottles become degraded and stained by fungi and difficult to read when stored in cellars and cool stores.</td>
<td>ASTM D 2020-03</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Samples of untreated material should demonstrate a high susceptibility to fungal growth in the test. Treated samples should be free of growth.</td>
<td></td>
</tr>
<tr>
<td>Biocide with fungicidal and bactericidal properties is used to protect PVC sheet materials from spoilage and degradation in service</td>
<td>9</td>
<td>PVC sheet flooring used on solid floors can become colonised by bacteria and fungi on its under surface. This causes staining, cracking and detachment from the substrate.</td>
<td>ISO 846 Parts A and C.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Samples of untreated material should support bacterial and fungal growth. Treated material should be free of growth.</td>
<td></td>
</tr>
<tr>
<td>Growth inhibition of moulds occurring on the plasters and walling in building structures</td>
<td>10</td>
<td>Surfaces of walls exposed to weather can be infected by saprophytic molds</td>
<td>Field tests : moulds growth should be shown on untreated material. Treated material should be free of moulds growth.</td>
</tr>
</tbody>
</table>
5.5.6 PT6 Preservatives for products during storage

In-can preservatives are included in many manufactured products, including paints, adhesives and binders. They are used to control micro-organisms that may be present in the product and which may cause deterioration prior to use. They therefore help to ensure product integrity during normal shelf life. Note: Food preservatives and cosmetics preservatives, which are used exclusively for this purpose, are not included in Product Type 6.

In order to grow in a manufactured product, a micro-organism must have access to both moisture (water) and a nutrient source. An extremely wide range of substances can act as a source of nutrition. These substances may be utilised by micro-organisms as they are, or following some form of conversion or degradation.

Utilisation of nutrition sources by micro-organisms results in the loss from the product of one or more components, leading to reduced integrity and spoilage. By-products of microbial growth also contribute to spoilage. Thus vulnerable products require an in-can preservative content for protection during the wet state, prior to use.

The broad group of wet-state preservatives for the purpose of storage prior to use has been divided into the sub-categories and sub-scenarios:

- PT6.1 Washing and cleaning fluids and human hygienic products
  - 6.1.1 Washing and cleaning fluids (human hygienic products)
  - 6.1.2 Washing and cleaning fluids (general) and other detergents
- PT6.2 Paints and Coatings (PN)
- PT6.3 Fluids used in paper, textile and leather production (P)
  - 6.3.1 Fluids used in paper production (Bulk raw materials in storage)
  - 6.3.2 Fluids used in textile production (Bulk raw materials in storage)
  - 6.3.3 Fluids used in leather production (Bulk raw materials in storage)
- PT6.4 Metal working fluid
  - 6.4.1 Lubricants (P)
  - 6.4.2 Machine oils (P)
- PT6.5 Fuel
- PT6.6 Glues and Adhesives
- PT6.7 Mineral slurries and other matrices

Each of these sub-scenarios can be tested as described in 5.5.2 and 5.5.5.1. This can be summarised as follows.

- A relevant matrix must be chosen according to the intended use. This matrix should be selected in a way that it can easily support growth if no biocide is present. A reasonably high water content and organic matter (either from the matrix itself or added as a soiling agent) will allow for growth.
- If available, a standard that covers the matrix must be chosen (e.g. for glues you might choose ASTM standard D 4783). From this test protocol the test organisms, the method of cultivating the test organisms, duration of the incubation, incubation
temperature, etc. can be extracted and integrated into a test protocol that follows the principles outlined above (e.g. by reducing the size of the inoculum).

Examples for test protocols that follow these principles are listed below. Other test methods which are commonly used for PT 6 can be found in Appendix 8. However, please note that these test methods are not necessarily appropriate to use; they are listed with comments to give an orientation for the assessor when and where these tests can be meaningful to prove a claim and when they aren’t:

i. A Method for Determining the Basic Efficacy of Biocidal Active Substances used in Polymer Dispersions, IBRG PDG 16-001;

ii. A Method for Determining the Basic Efficacy of Biocidal Active Substances used in Aqueous-Based Paints, (IBRG2 P 16-001;

iii. Tier 1 Basic Efficacy Method for Biocidal Active Substances used to Preserve Aqueous-Based Products, (IBRG2, IBRG PDG 16-007.

These documents describe methods for determining the basic efficacy of biocidal active substances in an aqueous based matrix and are intended for the generation of tier 1 data. The impact of additional factors like temperature and chemical stability etc., depending on the claim, would need to be tested.

When a claim of an active is to reduce bacterial growth, all 3 methods work according to the same principles, but differ in the bacteria used as they are specific to the matrix and the strength of the inoculum (also refer to 5.5.4.1). When the active substance also claims to reduce fungal growth, it will be necessary to differentiate between unicellular yeasts and filamentous fungi as yeasts can be counted as colony forming units, whereas filamentous fungi cannot (also refer to 5.5.4.2).

The filamentous fungus Geotrichum candidum is an organism that forms filamentous chains of fragmented cells. These are special in so far as they disintegrate easily into single arthrospores. Enumeration of growth of this fungus can therefore be performed in the same way as for unicellular yeasts. Details for culturing this fungus are given in method ii (Paints).

Whereas methods i) deal with polymer dispersions and ii) deal with paints, the efficacy of preservatives in all other matrices in PT 6 are at this point tested according to a generic method shown under iii) above. It provides a unified approach and is for use with those materials that do not (yet) have a specific method available (e.g. surfactants, cleaning products, mineral slurries etc.). It is designed to satisfy the basic requirements described in this document. As with the above tests, it is based on a challenge test (multiple inoculations at weekly intervals) and has the same basic requirements.

### 5.5.7 PT 7 Film preservatives and PT 9 Fibre, rubber and polymerised materials preservatives

Uses within PT 7 (film preservatives) and PT 9 (fiber, leather, rubber and polymerized material preservatives) often overlap. Sometimes, PT 7 and 9 differ only in the manner of application: the biocide can be applied as a coating layer onto the material or it can be incorporated into the material. Thus, the described requirements and principles apply in the same way to both PTs.

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20 IBRG website for test protocols: [http://ibrg.org/Methods.aspx](http://ibrg.org/Methods.aspx)
When selecting the appropriate method, consideration must be given to the release mode characteristics of a particular biocide/material combination. Some biocides have a very low solubility in water and hence are emitted at a very low rate from a matrix. This may be sufficient to protect a material that is inherently highly susceptible and which microorganisms may penetrate and colonise. However, if a test (e.g. ISO 16869) relies on the emission of the biocide from the matrix into an agar layer to measure the effect, the test would indicate that such a biocide has no function. Other materials, which are damaged by growth on their surface (especially where soiling is present) due to the production of extracellular enzymes, may fail to be protected by a biocide with such a low emission rate. Thus, the choice of method will be highly dependent on the characteristics of the material as well as the biocide. The applicant should justify this for the product under evaluation.

5.5.7.1 Simulation Tests (Tier 1 testing)

The ideal test method would present a material to a consortium of relevant test organisms under conditions that simulate real life realistically. This would produce effects that are identical to those observed in practice and allow a treatment to be identified with precision. There are methods that come closer to this ideal than others. For example, BS 3900 Part G6 (Appendix 6) exposes painted panels that have been inoculated with a mixture of spores of fungi known to colonise paints to humid conditions, free of external nutrients (although these can be added with the inoculum if necessary) for up to 12 weeks (see Figure 8). The resulting growth on untreated coatings has a visual appearance very similar to that observed in practice. For Tier 2 pre-exposure, leaching or artificial weathering can be used to help explore service life. A comparison can be made between treated and untreated variants of a formulation. A similar test, that forms the basis of many of the military standards and specifications, is BS EN 6006-2:10:2005 (see Appendix 6); this test is applicable to a wider range of materials. Again, samples are inoculated and incubated under conditions intended to simulate real life or at least be optimal for fungal growth.
Figure 8: Example of a Simulated Growth Test

**BS 3900 Part G6, Method Overview:**

Replicate sub-samples of both treated and untreated variants of each coating are sprayed with a suspension of spores of a range of fungi known to colonise surface coatings. The samples are then transferred to a humid chamber and incubated for up to 12 weeks. The extent of growth is assessed using a rating scale and this, as well as photographs of the panels, are presented as the results.

Rating scale: 0 = no growth, 1 = trace to 1% cover, 2 = 1 - 10% cover, 3 = 10 - 30% cover, 4 = 30 - 70% cover and 5 = > 70% cover

There is no pass/fail criterion in the standard but many workers in the coatings industry consider that growth represented by a rating of 2 is the maximum that would normally be tolerated. An example of growth on an untreated coating is shown on the left.

Example for growth level 5.

Modifications of these methods have been made to allow them to study the effects on algae (the IBRG algal test method for surface coatings) and, less commonly, bacteria. Effectiveness is assessed in these tests by visual appearance, measuring loss of weight or determining changes in the physical properties of the material (e.g. resistance to bending or extension under load). As with all biological tests, some degree of replication will be essential and tests should employ, as a minimum, three replicate sub-samples of each variant. Simulation tests are indeed very useful and provide valuable information especially for specific material/biocide combinations and can be correlated in some cases to service expectations. However, they can take a long time to perform and, in many cases, need to be adapted in some manner to accommodate a specific material.

5.5.7.2 Tests based on artificial growth media (Tier 1 testing)

By far the most commonly used methods for studying the performance of biocides intended to protect materials are those based on artificial growth media such as agar plates. For example, both ISO 846: 1997 and ASTM G21-09 are used widely in the plastics industry to measure the performance of fungicides in formulations (also ISO 16869: 2008). ISO 846 allows for studies into the susceptibility of plastic formulations to fungal and bacterial deterioration by attempting to make the plastic the sole source of nutrients for the organisms used, as well as providing a variant that provides an external source. It also includes a service life simulation test variant in which samples are buried in soil and then examined for loss of weight and strength (extremely useful in industries manufacturing pipes and cables). Although making the plastic the sole source of nutrients might seem like the ideal way to examine the ability of a biocide to protect the material, in many instances it is the presence of soiling that leads to colonisation and subsequent damage to the polymer (sometimes referred to as bio-corrosion). Thus, for certain polymers, the presence of
external nutrients is essential in determining the efficacy of a biocide. In many instances a consortium of organisms is required to effect colonisation and deterioration of the material and, in general, methods that employ consortia should be selected.

Similar testing technologies as those used for plastics exist for certain textiles, paper and surface coatings. The most commonly used are listed in Appendix 9; However, please note that these test methods are not necessarily appropriate to use; they are listed with comments to give an orientation for the assessor when and where these tests can be meaningful to prove a claim and when they aren’t. A description of the basic principles of tests on artificial growth media is given in Figure 9 using ASTM G21 as an example.

The huge disadvantage of agar-plate based tests is the interference of the growth medium with the biocide. The biocide can diffuse into the agar, demonstrating an effect there but at the same time be diluted in the original matrix. A less soluble substance, which does not diffuse into the agar, may in contrast show a false negative effect. For these reasons, a simulation test is always to be preferred over an agar-plate based test.

**Figure 9: An Example of an Agar Plate Based Test**

<table>
<thead>
<tr>
<th>ASTM G21, Method Outline:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Replicate samples of both treated and untreated material are embedded in a mineral salts-based agar medium. The sample and surrounding agar are then inoculated with the spores of a mixture of fungal species known to colonise plastics. The plates are then placed into chambers in which the humidity is maintained at &gt; 85% RH for up to 28 days. The samples are then inspected for the presence of fungal growth. Typical growth on an untreated material is shown in the plate on the left.</td>
</tr>
</tbody>
</table>

**5.5.7.3 Tier 2 Testing**

Depending on the intended use, pre-exposure, leaching or artificial weathering can be used to help explore service life. The relevance of the chosen parameters should be explained. There are no special tests or designs available for tier 2 testing. Basically, the same methods as in tier 1 can be applied except that the tested material undergoes pre-treatment. In some cases, ageing norms can be employed (e.g. adaptations of EN 73:2014\(^{21}\), EN 84:1997\(^{22}\), which are both developed for treated wood). In other cases, variations of the tier 1 methods can be used (as for example the soil burial variant of ISO 846 as described above). It is

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\(^{21}\) Accelerated ageing test of treated wood prior to biological testing. Evaporative ageing procedure

\(^{22}\) Accelerated ageing tests of treated wood prior to biological testing. Leaching procedure
particularly important to show growth or damage on the untreated material under service-life conditions.

In some cases it may not be necessary to use an artificial inoculum for tier 2 tests. It may be possible to use a test medium colonised naturally so that it is representative of the organisms that are typically encountered during the use of the product. It may be valid to use lower levels of contamination such as those encountered in practice. In some cases there may be a need to include application-related test-organisms in addition to standard test-organisms. In any case, the applicant should provide a rationale as to why the test organisms are relevant for the respective application/s of the preservative. Representatives for all claimed organisms should be tested

5.5.7.4 Tier 3 Testing

In some cases, tier 3 testing might be needed to support specific claims. These can be field tests where treated materials are compared to untreated materials in use. For example, treated house facades could be compared to untreated house facades in the same area and the time until re-painting is needed could be measured. Likewise, the replacement time for untreated buried cables compared to treated ones can be studied in a field test. Care has to be taken that the conditions for the treated and untreated materials are the same or at least comparable and that other parameters than the parameters observed are not influencing the results. The validity of the conclusions may need to be reinforced by statistical analysis etc, especially if any differences observed are small.

Table 12: Basic Requirements for a Valid Test Protection

| The following summary provides a guide to the basic requirements for a valid test: |
| i. A relevant model matrix should be chosen to represent the material(s) which must be protected; |
| ii. Relevant use conditions should be chosen in terms of humidity temperature and soiling; |
| iii. An untreated variant of the test material must be included and show the pattern of growth/deterioration that the biocide is intended to prevent at the end of the test; |
| iv. The test should employ organisms that are relevant to the material/problem being addressed; |
| v. Tests that employ a consortium of organisms should be favoured over those that use single species; |
| vi. A minimum of three replicate test pieces of both treated and untreated materials should be employed; |
| vii. The final data should include either some indication of the impact of service conditions on the performance of the treated material/article or data from an ageing. |

5.5.7.5 Prevention of Odour by odour-producing microorganisms

With most of the biocidal functions within PT 7 and 9, test conditions simulate in-use conditions rather well and the effects of microbial growth or activity can be observed quite easily. With the control of odour, this is much harder to achieve in a laboratory test, as odour often cannot be measured in a simple manner.
Laboratory tests to simulate odour production are currently not available, though some work is done to develop such tests (for example a test to inhibit the bioconversion of L-leucine to iso-valeric acid, representing a dominant compound of foot-odour). Thus, at present, the prevention of odour is in most cases measured indirectly by measuring microbial inhibition.

There are two major types of test that have traditionally been used with textiles (and related materials). The first major group employs agar plates and the other major group uses suspension in an aqueous medium. In both cases, the impact of a treated textile on populations of (usually) bacteria are studied. An overview is given in Appendix 10; however please note that the test methods listed are not necessarily appropriate to use; they are listed with comments to give an orientation for the assessor when and where these tests can be meaningful to prove a claim and when they aren’t.

**Agar plate-based tests**

Agar plate-based tests are not recommended. These tests have almost no useful utility in measuring effects intended to control odour in textiles. Such tests rely on the biocide migrating from the textile into the agar medium at sufficient concentration to inhibit the growth of bacteria either seeded into the agar or placed onto it (see Figure 9). The diffusion characteristics vary hugely from one biocide to another and from one textile to another and the growth medium itself presents a large soiling load to be overcome by the biocide. Larger areas clear of growth are often associated with more potent effects but they could be attributed equally to differences in the leaching rate of a biocide from a material.

**Suspension tests**

The second major group, the suspension tests, measure changes in the size of a population following contact with a treated textile. A number of protocols are described in Appendix 10. However, most employ relatively high concentrations of nutrients in the suspending medium so that their application, like the agar diffusion methods, can lead to over-treatment of textiles. Thus, these methods should not be used. By using lower concentrations of nutrients in the suspending medium and using pre-treatments such as laundering, these methods can be adapted for use in measuring effects on odour. Such an adaptation has been applied in the OECD Tier 1 method for treated articles (porous materials\(^\text{23}\)) and the IBRG Textile Method\(^\text{24}\). These are described schematically in Figure 10 and are based on the ‘germ’ count or absorption phase of ISO 20743: 2007 where the amount of nutrients present in the cell suspension has been reduced substantially.

Many treated materials would certainly be capable of demonstrating activity in a suspension test. Activity against a consortium of bacteria (e.g. against a range of Gram Positive and Gram Negative bacterial species such as *Staphylococcus epidermidis*, *Corynebacterium xerosis*, *Proteus vulgaris*, *Escherichia coli*, etc.) would probably inhibit the production of odour. However, excess exposure of the skin of the wearer should be minimized as far as possible. Therefore, tests adapted to textile treatments such as the OECD Tier 1 method and the IBRG Textile method (Figure 10) are preferable.

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Figure 10: OECD/IBRG Tier 1 Textile Test

Tier 2 testing
In many cases, a large fraction of the active substance incorporated in a textile is lost during laundering, either through emission of loosely or only partially bound material or associated with loss of fibres (lint). This also means that there is potential for active substances to be transferred from treated materials to non-treated materials when laundered together. In general, the emission rate is rarely continuous either to the environment or to the wearer. Moreover, other chemicals from the textile treatment as well as chemicals used in the laundering process might interfere with the function of the biocide.

In general, the effects required to prevent the formation of odour in shoes and apparel are subtle. The greatest demand on them is usually in maintaining activity following multiple laundering cycles. Therefore, simulation of service life conditions by laundering and ageing are essential. Care must be taken to maintain the functionality and to minimise excess exposure of the environment through emissions of the biocide in use, during cleaning and at the time of disposal. The method described in Figure 10 (as well as chemical analysis) in combination with laundering cycles can be useful in measuring the maintenance of efficacy in service.

An active substance or a biocidal product is often intended to treat a wide range and mix of textile types with a wide variety of anticipated demands and expectations of durability. It might be difficult to address every potential combination and garment type. However, studies on typical textile blends could be used to provide appropriate efficacy. Some examples are given in Table 13 below.
**Tier 3 testing**

At present the only truly reliable methods for demonstrating anti-odour functionality is through replicated and statistically designed wearing trials. Tier 1 and 2 tests described above can provide useful data related to durability etc. but care must be taken when interpreting the data they produce. For example, a treatment may be applied to only certain parts of a garment or shoe or it may be present on only a certain number of filaments in the weave of a textile. In the bioassay, the inoculum is dispersed throughout the whole of the sub-sample of textile and any active substance released would be able to migrate throughout that inoculum whereas in use, this may not occur. The humidity produced by bodily excretions might trigger less release of the biocide than the liquid suspension the textile is covered with in the test. The bacterial populations present on the skin might be less affected by the biocide as compared to the testing consortium employed. Consequently, user trials are proposed as reliable methods to prove anti-odour effects, especially in case of textiles, but also suitable microbiological studies with relevant odour-causing microorganisms can be acceptable ways to prove anti-odour claims. A standard with human assessors which could possibly be adapted to test anti-odour claims is EN 13725.

**Table 13: Odour: Example Claims, Problems and Testing Approaches**

<table>
<thead>
<tr>
<th>Claim</th>
<th>PT</th>
<th>Proof Required</th>
<th>Example Method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carpet is treated to prevent odours caused by mould growth.</td>
<td>9</td>
<td>Data should show that the treated carpet does not support fungal growth whereas the untreated one does. The effect should be shown to be sufficiently durable.</td>
<td>A method such as AATCC 174 can be used to demonstrate resistance to fungal growth. For active substances that do not migrate from the fibres/backing a cabinet-based simulation test may be more appropriate. Activity should be shown to persist following simulated ageing.</td>
</tr>
<tr>
<td>A sports vest is treated to inhibit the production of odour.</td>
<td>9</td>
<td>Data from a field trial should show that odour is reduced in treated sports shirts when compared with untreated ones. The effect should be shown to be of sufficient durability during service life to match any claim made.</td>
<td>Wearing trial or scientifically valid odour based simulation study. A comparison of the effectiveness both before and after simulated ageing/washing should be performed. This could be performed either through field trials, simulation tests or the use of a test such as the OECD Tier 1 method. The latter could be used to demonstrate that sufficient activity is still present after washing/ageing to elicit an antimicrobial effect.</td>
</tr>
</tbody>
</table>
5.5.8 PT8 Wood preservatives

General Introduction

This document deals with the evaluation methodology of efficacy tests for wood preservatives biocidal products that are applicable in the frame of the EU Biocidal Products Regulations (BPR) for the authorisation of biocidal products (BPR Annex VI).

The document is not intended to replace standards, standardized methods or other methods used as reference for developing the required data. It is considered as scientific guidance and the reader is advised to refer to the standards themselves or appropriate literature in case details should require further clarification.

The aim of this document is to provide a common base for the assessment of the efficacy for the biocidal product authorization for PT8 products for the applicants and the Competent Authorities (CAs).

Although alternative test methods could be taken into account, this document is mainly based on the EN 599-1 standard for preventive uses and on the EN 14128 standard for curative uses.

This document covers the products used for the preventive treatments of wood (including the saw-mill stage), by the control of wood-destroying or wood-disfiguring organisms (temporary treatments of logs in the sawmill or log yards, temporary treatments of green sawn timber, treatments of sawn timber including round timber, treatments of wood based panel) and products used for the curative treatments of sawn timber in service.

For product already on the market before entering into force of the standards (in 1990 for EN 599 and in 2004 for EN 14128):

- Efficacy data on the product should be provided.
- The assessment of the product efficacy should be based on expert judgement;
- Some data taken from the literature or used in certification could be accepted on case by case basis.

When the data are not enough robust to demonstrate the efficacy of the product, new tests according to EN 599 and/or EN 14128 will be required.

At the review time of this document, it has been chosen to include the catalogue of uses in the Chapter 7 of the Technical notes for guidance (TNsG) on product evaluation (PT8). The inclusion of the catalogue of uses to this document is to provide a common basis to harmonize the claims of the product. It will facilitate in a second time the mutual recognition by listing the elements of the claim in the same order and using the same terminology. On the label, the categories related to the product should be presented as described in the following paragraphs. The codes increase the readability of this document and are not expected on the label.

Concerning the updating of this document, it should be considered as a living document and will be reviewed on a regular basis and updated if necessary, under ECHA’s procedures.

The tests should be performed according to the current version in force of this document. Any tests initiated before the endorsement of the new version remain acceptable.

5.5.8.1 Label claims

In order to harmonize the efficacy issues, it is proposed that the different uses of the product are presented following the proposal below. This should follow the order of the categories listed below.
The aim of this categorisation is to have an explicit answer on the following questions:

- Where is the product used?
- What is the product used for?
- How is the product used? To control which organisms?

The data which support the efficacy should also follow this format.

The main categories that should be present on the label are listed in Table 14 and are detailed in the following paragraphs.

**Table 14: Different categories and the related product codes**

<table>
<thead>
<tr>
<th>Categories</th>
<th>Code for product</th>
</tr>
</thead>
<tbody>
<tr>
<td>User category</td>
<td>A.xx</td>
</tr>
<tr>
<td>Wood category</td>
<td>B.xx</td>
</tr>
<tr>
<td>Wood product</td>
<td>C.xx</td>
</tr>
<tr>
<td>Application aim &amp; Field of use</td>
<td>D.xx &amp; E.xx</td>
</tr>
<tr>
<td>Method of application and rate</td>
<td>F.xx</td>
</tr>
<tr>
<td>Target organisms</td>
<td>G.xx</td>
</tr>
</tbody>
</table>

**5.5.8.1.1 User Category (Code for Product A.xx)**

Information on the intended users of the product has to be presented on the label, the different user categories are presented in Table 15.

**Table 15: User categories**

<table>
<thead>
<tr>
<th>User Category</th>
<th>Example</th>
<th>Product Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>Non-professional/general public</td>
<td>Product used at home by consumers</td>
<td>A.10</td>
</tr>
<tr>
<td>Industrial</td>
<td>Industrial applicator</td>
<td>A.20</td>
</tr>
<tr>
<td>Professional</td>
<td>Pest control operator</td>
<td>A.30</td>
</tr>
</tbody>
</table>
5.5.8.1.2 Wood Category (Code for product B.xx)
This section deals with the wood category and not the use classes as defined in EN 335 standard. From an efficacy point of view, in EN 599-1, annex D the wood timbers are divided into two categories: softwood and hardwood.

Softwood and hardwood species of timber react differently to the degree and the type of attack by certain biological agents.

In most cases, the tests are performed with softwood. In some cases it is acceptable for this data to be read across to hardwoods, but in other cases specific testing against hardwoods is required. (see EN 599-1)

**Table 16: Wood categories**

<table>
<thead>
<tr>
<th>Wood Category</th>
<th>Product Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>Softwood</td>
<td>B.10</td>
</tr>
<tr>
<td>Hardwood</td>
<td>B.20</td>
</tr>
</tbody>
</table>

5.5.8.1.3 Wood Product (Code for product C.xx)
Table 17 below describes the types of wood products that are used as building materials or in the manufacture of furniture. Wood products are divided in two main categories: solid wood and wood based panels. Based on European standards, wood based panels are divided in four categories: plywood (EN 636), OSB (EN 300), Particles (EN 309 & EN 312) and Fibers (EN 622).

**Table 17: Wood product categories**

<table>
<thead>
<tr>
<th>Wood Category</th>
<th>Product Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>Solid wood</td>
<td>C.10</td>
</tr>
<tr>
<td>Reconstituted solid wood</td>
<td>C.11</td>
</tr>
<tr>
<td>Engineered solid wood products produced by processes involving pressure, adhesives and binders</td>
<td></td>
</tr>
<tr>
<td>Panels</td>
<td>C.20</td>
</tr>
<tr>
<td>Plywood panels</td>
<td>C.21</td>
</tr>
<tr>
<td>OSB panels</td>
<td>C.22</td>
</tr>
<tr>
<td>Particles panels</td>
<td>C.23</td>
</tr>
<tr>
<td>Fibers panels</td>
<td>C.24</td>
</tr>
</tbody>
</table>

5.5.8.1.4 Application aim and field of use

5.5.8.1.4.1 Application aim (code for product D.xx)
A preventive treatment is used to prevent sound wood from being infected by wood destroying agents and/or disfiguring fungi. The curative treatment is used to kill infective organisms that have already attacked the wood, to prevent them from spreading in the rest of the wood.
The preventive treatments are most of the time used during the manufacturing process but can also be done when the wood is in its service situation (e.g. framework of the building, a bridge).

According to the fact that a product can be used in wood preventive treatments, in curative treatments and sometimes both, and according to the fact that wood preservative and curative treatments are not covered by the same treatments, it is proposed to split the application aims as presented in Table 18.

The aim of this classification is to ensure having the same classification throughout the EU.

**Table 18: Application aim**

<table>
<thead>
<tr>
<th>Application Aim</th>
<th>Kind of Treatment</th>
<th>Product Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>Preventive</td>
<td>Temporary preventive treatment / logs</td>
<td>D.10</td>
</tr>
<tr>
<td></td>
<td>Temporary preventive treatment / green sawn timber</td>
<td>D.20</td>
</tr>
<tr>
<td></td>
<td>Preventive treatment / blue stain in service</td>
<td>D.30</td>
</tr>
<tr>
<td></td>
<td>Preventive treatment-use class (cf. the following section for the field of use – code E)</td>
<td>D.40</td>
</tr>
<tr>
<td>Curative</td>
<td>Curative treatment / wood in service</td>
<td>D.50</td>
</tr>
<tr>
<td>Preventive</td>
<td>Other (for e.g. pole maintenance)</td>
<td>D.60</td>
</tr>
</tbody>
</table>

### 5.5.8.1.4.2 Field Of uses (Code For Product E.xx)

The use classes described in EN 335:2013 are defined in terms of service conditions, with reference to the generalised moisture content and the prevailing biological agents of deterioration. The different classes (and their related application codes) are presented in Table 19.

- **Use class 1:** situation in which the wood or wood based product is inside a construction, not exposed to the weather and wetting;
- **Use class 2:** situation in which the wood or wood-based product is under cover and not exposed to the weather (particularly rain and driven rain) but where occasional, but not persistent, wetting can occur;
- **Use class 3:** situation in which the wood or wood-based product is above ground and exposed to the weather (particularly rain);
- **Use class 4:** situation in which the wood or wood-based product is in direct contact with ground or fresh water;
- **Use class 5:** situation in which the wood or wood based product is permanently or regularly submerged in salt water (i.e. sea water and brackish water).

Use class 3 is split into two sub-classes:

- **3.1:** wood and wood based products will not remain wet for long periods. Water will not accumulate;
- **3.2:** wood and wood-based products will remain wet for long periods. Water may accumulate.
The use classes 4.1 and 4.2 described in the former version of the EN 335 standard (2009) have been merged into a single use class 4, including both wood in exterior, in ground and/or fresh water contact.

**Table 19: Different field of uses**

<table>
<thead>
<tr>
<th>Field of Uses</th>
<th>Product Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>Use class 1</td>
<td>E.10</td>
</tr>
<tr>
<td>Use class 2</td>
<td>E.20</td>
</tr>
<tr>
<td>Use class 3*</td>
<td>E30</td>
</tr>
<tr>
<td>Use class 3.1</td>
<td>E.31</td>
</tr>
<tr>
<td>Use class 3.2</td>
<td>E.32</td>
</tr>
<tr>
<td>Use class 4</td>
<td>E.40</td>
</tr>
<tr>
<td>Use class 5</td>
<td>E.50</td>
</tr>
</tbody>
</table>

* includes use class 3.1 and use class 3.2

### 5.5.8.1.5 Method of application and application rate (Code for product F.xx):

The various methods available can be broadly split into three groups:

- **Superficial treatments:** Such non-pressure processes include brush, spray, roller, pad application and immersion (dipping) processes (where the wood can be in contact for preservative for periods of time ranging from a few minutes to several hours). The application rates are commonly expressed in g/m², ml/m².

- **Penetrating treatments:** Such processes include the vacuum pressure, alternating oscillating pressure, double vacuum and non-pressure processes such as diffusion treatments. The application rates are commonly expressed in kg/m³.

- **Other treatment methods:** For application methods different from those described above (fumigation, injection), either specifically relevant data or some justification for non-inclusion of data (i.e. details on penetrability/retention, etc.) will need to be provided to the CA for consideration.

Some PT 8 products are designed to be used with a top coat, e.g. primers for window framing. If a top coat is needed according to the manufacturer, this must be applied with the product. When a more general use is envisaged, generic coating materials can be used according to the norms performed.

**Table 20: Method of application**

<table>
<thead>
<tr>
<th>Method of application</th>
<th>Product Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>Superficial application / brush/roller/pad treatment</td>
<td>F.10</td>
</tr>
<tr>
<td>Superficial application / spray treatment</td>
<td>F.11</td>
</tr>
<tr>
<td>Superficial application / flow coat /aspiration</td>
<td>F.12</td>
</tr>
<tr>
<td>Superficial application / foam treatment</td>
<td>F.13</td>
</tr>
</tbody>
</table>
### Method of application

<table>
<thead>
<tr>
<th>Method of application</th>
<th>Product Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>Superficial application / dipping treatment</td>
<td>F.14</td>
</tr>
<tr>
<td>Injection</td>
<td>F.20</td>
</tr>
<tr>
<td>Pressure process</td>
<td>F.30</td>
</tr>
<tr>
<td>Pressure process / vacuum pressure impregnation</td>
<td>F.31</td>
</tr>
<tr>
<td>Pressure process / double vacuum</td>
<td>F.32</td>
</tr>
<tr>
<td>Fumigation</td>
<td>F.40</td>
</tr>
<tr>
<td>Fumigation bubble</td>
<td>F.41</td>
</tr>
<tr>
<td>Pole in services fumigation</td>
<td>F.42</td>
</tr>
<tr>
<td>Mixing with glue and mortar</td>
<td>F.50</td>
</tr>
<tr>
<td>Diffusion</td>
<td>F.60</td>
</tr>
<tr>
<td>Solid pellets / rods</td>
<td>F.61</td>
</tr>
<tr>
<td>Pole bandage / wrapping / pad application</td>
<td>F.62</td>
</tr>
<tr>
<td>Other application methods</td>
<td>F.70</td>
</tr>
</tbody>
</table>

#### 5.5.8.1.6 Target organisms (Code for product G.xx)

This section describes the main categories of target organisms, in relation to the claimed uses of the product, either for treatments to prevent biological attack, or for curative treatments to disinfest or to eradicate existing attack.

Appendix 11 gives more information on the principle target organisms.

There are a number of possible effects on target organisms resulting from the proposed use of a wood preservative product. The efficacy data for a wood preservative must be suitable to demonstrate the efficacy of products applied as either pre-treatments to prevent biological attack, or as curative treatments to disinfest or to eradicate existing attack. These may be in a variety of forms; they may yield toxic values, mortality values, subjectively derived ratings or effective retention values.

On the claimed matrix, the target organisms against which an efficacy is claimed must be clearly described. For the purpose of harmonisation, it is proposed that the target organism presented in Table 21 should be used, although these should not be considered as an exhaustive list. The species presented below are the species being representative of wood attacking organisms. For specific claims, efficacy data against each named target pest will be required.
Table 21: Examples of target organisms for wood preservatives
(N.B. these examples are not intended to be exhaustive with respect to target organisms or prescriptive with respect to data to be generated).

<table>
<thead>
<tr>
<th>Common English term</th>
<th>Code F for product</th>
<th>Target organisms according to EN 1001</th>
<th>Classification</th>
<th>Scientific name</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Fungi</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wood rotting fungi</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wood rottjng basidiomycetes</td>
<td>G.10</td>
<td>Brown rot fungi</td>
<td>Basidiomycetes</td>
<td>e.g. <em>Gloeophyllum trabeum</em></td>
</tr>
<tr>
<td></td>
<td>G.11</td>
<td>White rot fungi</td>
<td>Basidiomycetes</td>
<td>e.g. <em>Coriolus versicolor</em></td>
</tr>
<tr>
<td>Soft rot fungi</td>
<td>G.12</td>
<td>Soft rot fungi</td>
<td>Ascomycetes, Deuteromycetes</td>
<td>e.g. <em>Chaetomium globosum</em></td>
</tr>
<tr>
<td>Wood discolouring fungi</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>G.21.1</td>
<td></td>
<td>Sapstain fungi (bluestain mainly)</td>
<td>Ascomycetes, Deuteromycetes</td>
<td>e.g. <em>Ophiostoma pini</em> (&lt;i&gt;Ceratocystis pilifer&lt;i&gt;)</td>
</tr>
<tr>
<td>G.21.2</td>
<td></td>
<td>Bluestain in service</td>
<td>Ascomycetes, Deuteromycetes</td>
<td>e.g. <em>Aureobasidium pullulans</em></td>
</tr>
<tr>
<td>G.22</td>
<td></td>
<td>Mould fungi</td>
<td>Ascomycetes, Deuteromycetes</td>
<td>e.g. <em>Aspergillus niger</em></td>
</tr>
<tr>
<td><strong>Insects</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Beetles</td>
<td>G.30</td>
<td>Wood boring beetles</td>
<td>Coleoptera</td>
<td>e.g. <em>Hylotrupes bajulus</em>.</td>
</tr>
<tr>
<td>G.31</td>
<td></td>
<td>House longhorn beetle</td>
<td>Coleoptera</td>
<td>e.g. <em>Anobium punctatum</em></td>
</tr>
<tr>
<td>G.32</td>
<td></td>
<td>Common furniture beetle</td>
<td>Coleoptera</td>
<td>e.g. <em>Lycus brunneus</em></td>
</tr>
<tr>
<td>G.33</td>
<td></td>
<td>Powder post beetles</td>
<td>Coleoptera</td>
<td>e.g. <em>Scolytus spp</em></td>
</tr>
<tr>
<td>G.50</td>
<td></td>
<td>Termites (genus claimed)</td>
<td>Isoptera</td>
<td>e.g. <em>Reticulitermes spp</em>, e.g. <em>Coptotermes spp</em></td>
</tr>
<tr>
<td>G.51</td>
<td></td>
<td>Subterranean termites (genus claimed)</td>
<td>Isoptera</td>
<td>e.g. <em>Cryptotermes spp</em></td>
</tr>
<tr>
<td>G.52</td>
<td></td>
<td>Drywood termites (genus claimed)</td>
<td>Isoptera</td>
<td>e.g. <em>Nasutitermes spp</em></td>
</tr>
<tr>
<td>G.53</td>
<td></td>
<td>Tree termites (genus claimed)</td>
<td>Isoptera</td>
<td>e.g. <em>Nasutitermes spp</em></td>
</tr>
<tr>
<td><strong>Wood destroying marine organisms</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>G.60</td>
<td></td>
<td>Marine borers (genus claimed)</td>
<td>Teneridae, Pholadidae</td>
<td>e.g. <em>Tereod sp</em>, Martesia sp</td>
</tr>
<tr>
<td>G.61</td>
<td></td>
<td>Mussels</td>
<td>Teneridae, Pholadidae</td>
<td>e.g. <em>Limnoria sp</em>, Chelura sp</td>
</tr>
<tr>
<td>G.62</td>
<td></td>
<td>Crustaceans</td>
<td>Isopoda, Amphipoda</td>
<td>e.g. <em>Limnoria sp</em>, Chelura sp</td>
</tr>
</tbody>
</table>
5.5.8.1.7 Examples of a claimed matrix

To illustrate the previous sections described, the following table gives an example of claimed matrix based on the categories from the catalogue of uses. This framework should be followed for the efficacy claim’s part of the label. Only the categories and the matrix wordings (not the code) are expected to be listed on the label.

This matrix allows a harmonisation of the efficacy elements presented in the dossier for product authorization. Elements in the claimed matrix must be present on the physical label.

**Table 22: Examples of claim matrix based on the application codes for product**

<table>
<thead>
<tr>
<th>Categories</th>
<th>Matrix Wording</th>
<th>Code Product for Product</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Label 1</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>User category</td>
<td>Industrial</td>
<td>A.20</td>
</tr>
<tr>
<td>Wood category</td>
<td>softwood and hardwood</td>
<td>B.10; B.20</td>
</tr>
<tr>
<td>Wood product</td>
<td>solid wood</td>
<td>C.10</td>
</tr>
<tr>
<td>Application aim and field of use</td>
<td>preventive treatment - use class 3.2</td>
<td>D.40; E.32</td>
</tr>
<tr>
<td>Method of application and rate</td>
<td>superficial application/dipping treatment</td>
<td>F.14</td>
</tr>
<tr>
<td></td>
<td>application rate: 100 g/m² in the analytical zone</td>
<td></td>
</tr>
<tr>
<td></td>
<td>a top coat must be applied.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>pressure process/vacuum impregnation</td>
<td>F.31</td>
</tr>
<tr>
<td></td>
<td>application rate: 50 kg/m³ in the analytical zone</td>
<td></td>
</tr>
<tr>
<td>Target organisms</td>
<td>wood boring beetles</td>
<td>G.30</td>
</tr>
<tr>
<td></td>
<td>termites (genus <em>Reticulitermes</em>)</td>
<td>G40</td>
</tr>
<tr>
<td></td>
<td>brown rot fungi</td>
<td>G.10</td>
</tr>
<tr>
<td></td>
<td>white rot fungi</td>
<td>G.11</td>
</tr>
<tr>
<td><strong>Label 2</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>User category</td>
<td>Industrial</td>
<td>A.20</td>
</tr>
<tr>
<td>Wood category</td>
<td>softwood and hardwood</td>
<td>B.10; B.20</td>
</tr>
<tr>
<td>Wood product</td>
<td>solid wood</td>
<td>C.10</td>
</tr>
<tr>
<td>Application aim and field of use</td>
<td>preventive treatment - use classes 2, 3 and 4</td>
<td>D.40 - E.20; E.30; E.40</td>
</tr>
<tr>
<td>Method of application and rate</td>
<td>superficial application/dipping treatment</td>
<td>F.14</td>
</tr>
<tr>
<td></td>
<td>application rate in the analytical zone:</td>
<td></td>
</tr>
<tr>
<td></td>
<td>UC 2: 80 - 120 g/m²</td>
<td></td>
</tr>
<tr>
<td></td>
<td>UC 3 (coated): 100 – 160 g/m²</td>
<td></td>
</tr>
<tr>
<td></td>
<td>pressure process/vacuum pressure impregnation</td>
<td>F.31</td>
</tr>
</tbody>
</table>
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application rate in the analytical zone:
UC2: 30 kg/m²
UC3: 40 - 70 kg/m³
UC4 (softwood): 80 – 150 kg/m³
UC4 (hardwood): 100 – 150 kg/m³

<table>
<thead>
<tr>
<th>Target organisms</th>
<th>brown rot fungi</th>
<th>G.10</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>white rot fungi</td>
<td>G.11</td>
</tr>
<tr>
<td></td>
<td>soft rot fungi</td>
<td>G.12</td>
</tr>
<tr>
<td></td>
<td>wood boring beetles</td>
<td>G.30</td>
</tr>
<tr>
<td></td>
<td>termites (genus Reticulitermes)</td>
<td>G.40</td>
</tr>
</tbody>
</table>

Label 3

| User category | Industrial | A.20 |
| Wood category | softwood   | B.10. |
| Wood product  | solid wood | C.10 |
| Application aim and field of use | temporary preventive treatment - use class 1 | D.20 E.10 |
| Method of application and rate | superficial application / dipping treatment application rate 100 g/m² in the analytical zone | F.14 |
| Target organisms | sapstain | G.21.1 |
|                  | mould fungi | G.22 |

5.5.8.2 Available data

5.5.8.2.1 Standard test methods

When considering the overall evaluation of proposed claims, CAs should ensure that the test methods (data, method of application and application/dose rates used in the tests, product tested) are appropriate to demonstrate the efficacy claimed on the label for the product.

Many standard protocols currently exist to test wood preservatives; the lists of standards for the efficacy assessment of wood preservatives are available on the ECHA Biocides Efficacy Working Group webpage [http://echa.europa.eu/about-us/who-we-are/biocidal-products-committee/working-groups/efficacy]. For PT8, the CEN standards are highly recommended.

Two main categories of treatment are described:

- Preventive treatments, which are covered by EN 599-1;
- Curative treatments, which are covered by EN 14128.

Some other treatments (C.20: green sawn timber) are covered by other standards (e.g. CEN TS 15082).

It is highly recommended to perform the studies according to these standards. If the standards are not applicable or suitable, the applicant may adapt the methodology or use
another method (including his own method). When a standard is modified or when a non CEN standard is used, a robust justification and description have to be provided. For very specific cases, tests or ageing procedures could be waived with a robust justification. The study submitted has to provide a clear answer to the issue.

In the general part of the TNsG on data requirements it is mentioned that the test (and the data generated) should be based on sound scientific principles and practices. Compliance with quality standards is highly recommended.

In the TNsG on product evaluation, it is mentioned that for efficacy testing, the principles of Good Laboratory Practice (GLP) are not required by the legislation. However this guidance indicates that the spirit of such principles should be applied for the testing of efficacy.

Particular attention should be paid to:

- what information is needed to substantiate a 'claim matrix';
- the Quality Assurance procedures which should be adopted (cf. ISO 17025 for testing and certification);
- the overall evaluation of the data package when the completeness and adequacy of the data are compared with the label claim.

For products intended for application as solids, pastes or encapsulated forms and those intended for curative (in-situ) use, modification of the relevant protocols/testing strategies may be done or other direct evidence may be submitted on their potential efficacy against the claimed target organisms (e.g. for pastes such evidence could be in the form of penetrability and retention characteristics).

The test methods used to provide data should be relevant to the target organisms and application processes claimed on the label (see EN 599-1 and individual test standards).

It has to be noted that in some cases, a different formulation from which an authorization is sought could be tested. The results could be accepted by the RMS in a case by case approach (see section 5.5.8.3 of this guidance and Annex A of the EN 599-1 and EN 14128). A full composition of the tested product and a robust justification why the test is relevant should be provided.

For EN113, where the protocol states that several organisms have to be tested in order to fulfil the efficacy criteria, it is recommended that all testing is done in the same laboratory at the same time. The sponsor must have the right to provide his rational for justification why the simultaneous testing may have not been followed. Derogation (inter alia) is acceptable i.e. in the following cases:

- where the test was performed with limited organisms and later completed with additional organisms which could be tested in another laboratory (extension of claim);
- where the laboratory cannot run the test with specific targets;
- where the laboratory has ceased to provide services;
- in the case where a ‘simultaneous test’ is not available, but valid tests (according to the criteria in the standard) are available.

Table 21 and Table 23 below are informative for the test methods used. The user should also refer to EN 599-1 or EN 14128 depending on the claims.
Table 23: Preventive treatments: List of available standards and others methods used in wood preservation

<table>
<thead>
<tr>
<th>Organisms</th>
<th>Code for product</th>
<th>Temporary treatment of logs</th>
<th>Temporary treatment</th>
<th>Treatment of solid wood (List of standards mentioned in the tables 1 to 5 of EN 599-1)</th>
<th>Treatment of wood based panels</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Note 1: In some conditions, ageing tests (EN 84, EN 73) or natural weathering are required (see EN 599-1)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Note 2: It is highly recommended to refer to EN 599-1 to determine the tests to be done in accordance with table 1 to 5 of EN 599-1</td>
<td></td>
</tr>
<tr>
<td>Brown rot fungi</td>
<td>G.10</td>
<td></td>
<td></td>
<td>EN 113, EN 839, EN 330, EN 113, EN 252, EN 113, ENV 12038</td>
<td></td>
</tr>
<tr>
<td>White rot fungi</td>
<td>G.11</td>
<td></td>
<td></td>
<td>EN 113, EN 839, EN 330, EN 113, EN 252, EN 113, ENV 12038</td>
<td></td>
</tr>
<tr>
<td>Soft rot fungi</td>
<td>G.12</td>
<td></td>
<td></td>
<td>ENV 807, EN 252, ENV 807</td>
<td></td>
</tr>
<tr>
<td>Sapstain fungi</td>
<td>G.21.1</td>
<td>No CEN standard*</td>
<td>No CEN standard*</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bluestain fungi</td>
<td>G.21.2</td>
<td>No CEN standard*</td>
<td></td>
<td>EN 152, EN 152, EN 152, EN 152</td>
<td></td>
</tr>
<tr>
<td>Mould fungi</td>
<td>G.22</td>
<td>No CEN standard*</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wood boring beetles</td>
<td>G.30</td>
<td></td>
<td></td>
<td>EN 46, EN 47, EN 46, EN 47, EN 47, EN 47, EN 47, EN 47, EN 47, EN 49-2, EN 49-2, EN 20-2</td>
<td></td>
</tr>
</tbody>
</table>

25 For wood based panels, the reader is aware that standards can be adapted in specific cases (e.g. CEN/TS 15083-2 for soft rot fungi, EN 20-2 for powder post-beetle and EN 117 and EN 118 for termites)
<table>
<thead>
<tr>
<th>Organisms</th>
<th>Code for product</th>
<th>Temporary treatment of logs</th>
<th>Temporary treatment</th>
<th>Treatment of solid wood (List of standards mentioned in the tables 1 to 5 of EN 599-1)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Note 1: In some conditions, ageing tests (EN 84, EN 73) or natural weathering are required (see EN 599-1)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Note 2: It is highly recommended to refer to EN 599-1 to determine the tests to be done in accordance with table 1 to 5 of EN 599-1</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Use Class 1</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>EN 49-2</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>EN 20-1</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>EN 20-2</td>
</tr>
<tr>
<td>House longhorn beetle</td>
<td>G.31</td>
<td></td>
<td></td>
<td>EN 46</td>
</tr>
<tr>
<td>Common furniture beetle</td>
<td>G.32</td>
<td></td>
<td></td>
<td>EN 49-1</td>
</tr>
<tr>
<td>Powder post-beetle</td>
<td>G.33</td>
<td></td>
<td></td>
<td>EN 20-1</td>
</tr>
<tr>
<td>Fresh wood insect</td>
<td>G.40</td>
<td></td>
<td></td>
<td>No CEN standard*</td>
</tr>
<tr>
<td>Termites</td>
<td>G.50</td>
<td></td>
<td></td>
<td>EN 118</td>
</tr>
<tr>
<td>Marine borers</td>
<td>G.60</td>
<td></td>
<td></td>
<td>EN 117</td>
</tr>
</tbody>
</table>

Blank cell: Not applicable;

* National standards available (see the ECHA Biocides Efficacy Working Group webpage [http://echa.europa.eu/about-us/who-we-are/biocidal-products-committee/working-groups/efficacy]).
Table 24: Curative treatments: List of available standards used in wood curative treatments (based on EN 14128)

<table>
<thead>
<tr>
<th>Organisms</th>
<th>Code for Product</th>
<th>Curative treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brown rot fungi</td>
<td>G.10</td>
<td></td>
</tr>
<tr>
<td>White rot fungi</td>
<td>G.11</td>
<td></td>
</tr>
<tr>
<td>Soft rot fungi</td>
<td>G.12</td>
<td></td>
</tr>
<tr>
<td>Sapstain fungi</td>
<td>G.21.1</td>
<td></td>
</tr>
<tr>
<td>Blue stain fungi</td>
<td>G.21.2</td>
<td></td>
</tr>
<tr>
<td>Mould fungi</td>
<td>G.22</td>
<td></td>
</tr>
<tr>
<td>Wood boring beetles</td>
<td>G.30</td>
<td></td>
</tr>
<tr>
<td>House longhorn beetle</td>
<td>G.31</td>
<td>ENV 1390</td>
</tr>
<tr>
<td>Common furniture beetle</td>
<td>G.32</td>
<td>EN 48 or EN 370</td>
</tr>
<tr>
<td>Powder post beetles</td>
<td>G.33</td>
<td>No CEN standard available</td>
</tr>
<tr>
<td>Fresh wood insect</td>
<td>G.40</td>
<td></td>
</tr>
<tr>
<td>Termites (genus claimed)</td>
<td>G.50</td>
<td>No CEN standard available</td>
</tr>
<tr>
<td>Marine borers (genus claimed)</td>
<td>G.60</td>
<td></td>
</tr>
</tbody>
</table>

*Blank cell: Not applicable

### 5.5.8.2.2 Preventive treatments

Most of the available data are laboratory generated and related to the organisms for which biocidal efficacy is claimed.

Field tests, although desirable in cases where the product is intended for use in the more severe service environments (e.g. in ground contact (use class 3, 4 and 5)) are considered mandatory to fulfill the minimum performance criteria, according to the tests required in the paragraphs related to the use classes. As this could lead to a significant delay before a new product could be introduced to the market, literature, monitoring or other methods provided to support the derived application rate could be accepted in case by case by the CAs (see also notes in sections 5.5.8.2.2.3 and 5.5.8.2.2.4).

The assessment of the preventive efficacy of wood preservative formulations has to be made from values derived from a relevant biological test. These values are either the actual quantitative amounts of the product established in the test as causing the appropriate level of mortality of the target organism, or they represent the threshold limits, the so-called 'toxic values'. These toxic values are two concentrations in the series used in the test, the first which just permits continued attack and the second which just prevents it.

### 5.5.8.2.2.1 Temporary treatments of logs (in the sawmill or in storage area)

This kind of treatment is used to prevent the degradation of logs which do not immediately have their bark removed. Indeed, some microscopic fungi (e.g. stain) infect the wood and/or some species of insects belonging to the family of Scolytidae and Bostrychidae (named “Fresh wood insect” in Table 21) lay their eggs between the bark and the wood.
To prevent these damages, the logs may be treated with a biocidal product. As the treatment is temporary, use class is not relevant in this case.

**5.5.8.2.2 Temporary treatment of green timber**

This kind of treatment is used for the protection of freshly felled green lumber against colonization by blue stain and other discolouring micro-organisms (often named ‘sapstain’ as there are more than 200 fungi which can caused discoloration of the sapwood) and surface mould.

A technical specification (CEN/TS 15082) is available.

1. **blue stain fungi and other discolouring sapwood fungi**
   - Blue stain is caused by microscopic fungi that only infect the sapwood. They can cause blue or grey discoloration of the sapwood, but have no impact on its strength. Blue stain reduces the value of the wood.
   - Typical blue stain fungi are: *Ceratocystis spp, Ophiostoma spp Aureobasidium spp*
   - Typical other discolouring fungi are: *Stereum spp*
   - In the final stage of processing in a sawmill, treatment with a biocidal product (commonly applied by dipping to prevent blue stain fungi) may be carried out.

2. **moulds growing often on the wood surface**
   - The major problems caused by moulds fungi are discoloration on surfaces, and sometimes health problems. They do not affect the strength properties of wood.
   - Typical mould fungal genera on wood are: *Alternaria, Aspergillus, Penicillium, Trichoderma*.
   - A dose rate / dipping time is part of the efficacy assessment. The label claim must mention the dose rate and the dipping time.

**5.5.8.2.3 Treatments of solid wood (EN 599-1 Standard)**

When the purpose is to protect the wood, a preventive treatment is often applied to prevent the degradation of wood by micro-organisms (for example fungi) and/or by insects (for example wood boring insects). The treatment type is related to the organisms against which the wood has to be protected and to the use class. EN 599-1 specifies what test should be done for each use class claimed.

Different target organisms may preferentially attack either softwood or hardwood. Tests must be conducted on softwood and/or on hardwood as appropriate to the target organisms and following the requirements presented in the relevant test procedures.

It must be noted that Use Class 1 requires only insecticide products and, starting from Use Class 2, products are fungicide alone or combine fungicide and insecticide activities.

**Use Class 1**

Required data
Refer to EN 599 -1 table 1.

Data will include suitable laboratory data using treated test blocks to determine the toxic values against insects as appropriate.

Data should be presented on test blocks subjected to pre-conditioning by an evaporative ageing process (e.g. EN 73).

Test species
The insect species tested will depend on whether a general or a specific efficacy claim is made. Data should demonstrate activity against one or more of the following specific
insects as indicator species: *Hylotrupes bajulus, Anobium punctatum, Lyctus brunneus*, and where appropriate, termites.

**Note**

CASs should evaluate the available data to determine whether they are sufficient for label claims as follows:

a) for general claims against "wood boring beetles"\(^{26}\)

All relevant beetle species (*Hylotrupes bajulus, Anobium punctatum* and *Lyctus brunneus*) should be tested except if data (relevant and robust literature data where the materials and methods are detailed; certification data\(^{27}\) on a case by case basis) are provided which demonstrate that one of the targets is the less sensitive or that the product has an equivalent activity against all beetle species (refer to EN599-1:2014, section 5.2.3)

b) for claims against a specific beetle species

If claims against individual beetle species are detailed on a product label, then suitable efficacy data against those named target pests will be required.

c) for claims against termites

Some data on efficacy against termites will only be required when the product is to be marketed for use as a termiticidal product or where local requirements demand such activity.

For a product claiming activity against termites, suitable data demonstrating preventive efficacy against a European *Reticulitermes* species will be required.

For a product claiming efficacy against overseas tropical termites, suitable data demonstrating preventive efficacy against relevant species will be required.

**Use Class 2**

**Required data**

Refer to EN 599-1:2009 table 2.

Data will include suitable laboratory data using treated test blocks to determine the toxic values against the fungi and insects as appropriate.

**Test species**

The test species used will depend upon the label claims and will include as a minimum the brown rot fungi and insects if appropriate (as in Use Class 1).

**Note**

The CASs evaluate the available data to determine if they are sufficient for label claims as follows:

a) For claims against wood rotting fungi the following data have to be available:

Suitable laboratory data demonstrating efficacy against brown rot fungi after ageing test in accordance with EN 73.

b) For claims against wood discolouring fungi the following data have to be available:

- Suitable laboratory data on the protective efficacy of the product against blue stain in service after ageing test in accordance with EN 73 or after a natural or

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\(^{26}\) This correction has been made for an error in drafting and should be considered to be effective immediately and not subject to the standard transitional period of 2 years for new guidance.

\(^{27}\) This certification ensures that products are fit for purpose and defines a capacity in the use of products taking into account among others the durability in the function (efficiency of the treatment). The efficacy part of the certification scheme is (in France) generated according the requirement of the EN 599.
artificial weathering cycle as given in EN 152;
  o The application process used in the tests (i.e. whether by superficial or penetrative treatment) has to be in accordance with label claims.
c) For claims against insect pests the following data have to be available:
As outlined in Use Class 1.

Use Class 3

Required data
Refer to EN 599-1:2009 table 3a and table 3b.

Data will include suitable laboratory data using treated test blocks to determine the toxic values against the fungi and insects as appropriate.

Test species
The test species used will depend upon the label claims and will include as a minimum the brown rot fungi and insects if appropriate (as in Use Class 1).

Note
The CAs should evaluate the available data to determine if they are sufficient for claims matrix as follows:
a) For claims against wood rotted fungi, the following data have to be available:
  o Suitable laboratory tests as outlined for Use Class 2 and in addition, the efficacy will be demonstrated following pre-conditioning of the treated test blocks by a suitable leaching procedure according to EN 84
b) For claims against wood discoloring fungi the following data have to be available:
  o Suitable laboratory data on the protective efficacy of the product against blue stain in service after a natural weathering or an artificial weathering as given in EN 152.
  o The application process used in the tests (i.e. whether by superficial or penetrative treatment) should be in accordance with label claims.
c) For claims against insect pests (if relevant) the following data have to be available:
As outlined in Use Class 1, and in addition the efficacy will be demonstrated following pre-conditioning of the treated test blocks by a suitable leaching procedure according to EN 84 if technically possible (i.e. this is not the case for EN 20-1 and 20-2 due to methodological constraints).

According to EN 599-1 field test results, according to EN 330 may be used by the applicant instead of certain EN 113 test results, after EN 84 leaching test to derive the brown rot fungi. They are not needed to derive the minimum retention requirements. Moreover EN 330 may be used as an alternative to basidiomycetes laboratory tests (EN 113 + EN 84) for product under coating.

Use Class 4

Required data
Refer to EN 599-1:2009 table 4.

Data will include suitable laboratory data using treated test blocks to determine the toxic values against the fungi and insects as appropriate. In this situation available data should only include application of the preservative by penetrative treatments.
Test species

Test species used will depend upon the label claims and will likely include the following target organisms: brown and white rot fungi, soft rot micro-fungi and if relevant to label claims, blue stain fungi and insects as appropriate.

Note

The CAs should evaluate the available data to determine if they are sufficient for matrix claims as follows:

a) For claims against wood rott ing fungi, the following data have to be available
   - Suitable laboratory data as outlined for Use Class 3 with the following supplements:
     - all laboratory data should derive from impregnated treated test blocks (i.e. a penetrative treatment) with the test formulation to determine the toxic values against both brown and white rot fungi separately;
     - a suitable laboratory test to determine the toxic efficacy against soft rot fungi and other soil inhabiting microorganisms is required;

b) For claims against wood discolouring fungi, the following data have to be available:
   - A suitable laboratory test determining the protective efficacy of the product against blue stain for wood in service as given in EN 152.

c) For claims against insect pests, the following data have to be available:
   - As outlined for Use Class 1 and in addition, efficacy will be demonstrated following pre-conditioning of the treated test blocks by a suitable procedure according to EN 73 and to EN 84 separately).

In Use Class 4 data (e.g. EN 252, literature, monitoring or other methods) will be provided to support the derived application rate.

Use Class 5

Required data

Refer to EN 599-1 table 5.

The principal agent of decay in this situation is the marine borers. Therefore in this Use Class available data must include evidence of efficacy in a relevant marine field trial carried out for a minimum of 5 years (e.g. to EN 275 or an equivalent test).

The decay in this situation by basidiomycetes fungi does occur but marine soft rot fungi are more common causing surface softening of timber. Assessment of products against marine fungi is not normally conducted using routinely laboratory tests because of the difficulties for providing conditions which appropriately model the marine environment. There is, at present, not a recognised standard laboratory test for assessment of timber intended for use in salt water.

Test species

Test species used will depend upon the label claims. The principal agent of decay in the marine environment is the marine borers although claims against fungi can also be made.

The CAs evaluate the data to determine if they are sufficient for label claims as follows:

For claims against wood rott ing fungi and marine borers, the following data have to be available:

- For fungi available data as outlined in Use Class 4 as a surrogate has to be acceptable.
- For marine borers, a relevant marine field trial data has to be carried out for a minimum of 5 years according to EN 275.
5.5.8.2.2.4 Treatments of wood-based panels

The biocidal treatment of wood-based panels is achieved either during or after the manufacturing process.

During the manufacturing process, product can be included into the glue prior to application or directly by wood treatment.

The evaluation of the durability of wood-based panels against brown rot fungi and white rot fungi should be carried out according to the ENV 12038 test method.

There is no specific standardized methodology allowing the evaluation of the resistance of treated wood-based panels against soft rot or insects such as *Lyctus spp.* or termites. However, some of the existing standards usually applied to solid wood can be adapted to the evaluation of wood-based panels: CEN/TS 15083-2 (natural durability to soft rot fungi), EN 20-2 (*Lyctus spp.*), EN 117 and EN 118 (termites).

For post-manufacturing treatment, product can be applied by using a surface application process or pressure process.

In that case, the EN 599-1 is appropriate for determining the retention of post manufacture treatment.

5.5.8.2.2.5 Barrier treatment against *Serpula lacrymans*

The dry rot fungus (*Serpula lacrymans* = true dry rot fungus) occurs in buildings, causing brown rot in timber. The fungus can develop at relatively low wood moisture contents and is able to penetrate damp masonry over long distances in order to infect further timber or to develop its fruit-bodies.

In general, in case of an infestation of *Serpula lacrymans*, the infected wood is cut away. To prevent the infection of the new placed wood with fungi coming from the surrounding masonry, a curative treatment against dry rot in walls (mortar) will result in creating a ‘preventive’ barrier in/on walls hindering the fungus to grow through.

There is a specific Technical Specification (CEN/TS 12404) for determining the performance of a preservative applied to the upper surface of the mortar in preventing the growth of dry rot through the treated mortar when exposed to the fungus. This method is only applicable to masonry fungicides applied as a true solution of preservative. It is not applicable to rods, pastes and other similar preservative types. This method is applicable to preservatives applied to masonry by brushing, spraying and/or injection techniques or mixed into rendering and plastering mortar for masonry.

5.5.8.2.2.6 Determination of preventive product application rate with regard to service life

The evaluation of PT8 products efficacy is based on the retention of the product as determined in standard test methods, e.g. according to standards listed in EN 599-1. The values determined in this way are critical values (CV’s) for a particular formulation. The application rates derived from the CV’s are deemed to provide only a baseline efficacy and no conclusion on service life can be made. Indeed, neither is the term service life an absolute measure and no uniform mathematical model exists to derive such from CV’s, nor is determination / claim of a distinct service life part of the BPR. Estimation of service life (ESL) is based on the assumption, that different parameters have an impact on the service life of wood. This is explained in ISO 1586-1 and ISO 15686-2.

An estimated service life of wooden products is influenced e.g. by local exposure conditions, maintenance, consumer expectation and long term experiences from field testing or industrial experiences. This can provide justification for setting higher or lower retention rates as derived from CV’s only.
Because the concept of ESL is not part of the BPR and claims for a specific service life is consequently solely the applicant’s responsibility, the applicant must have the right to apply for lower or higher retentions than just the CV up to the retention rate which is limited by the human health and environmental risk assessments.

In order to support his claim, for UC3 claims, the applicant should submit data from e.g. literature, EN 330. For UC4, the applicant will provide, EN 252 (applicable to UC4 claims) and/or other methods for justification.

Particular specification for use class 4:

The field tests sites (minimum two) or the data extracted from literature must be representative for climatic zones with regards to the markets targeted by the product. The selected sites must allow the evaluation of the product’s efficacy on all the biological organisms covered by the label claim.

5.5.8.2.3 Curative treatment

EN 14128 is the lead standard providing detailed insight into the minimum testing requirements for wood preservatives claiming curative activity. It must be noted, that testing standards concerning PT8 products are only available for testing against wood boring insects.

It is important to understand that conducting curative treatments may comprise series/combinations of different steps and application methods/techniques in order to achieve the desired result and quite often result in providing preventive and curative efficacy at the same time.

5.5.8.2.3.1 Wood boring insects

Data required to support label claims for curative efficacy may include some tests generated using existing EN standards for the relevant beetle species or other alternative supporting data.

A number of EN standard tests exist for curative treatments for insecticides against *Hylotrupes bajulus* (ENV 1390) and *Anobium punctatum* (EN 48). The curative activity against *Lyctus* is not tested separately but is derived from results from testing against *Anobium punctatum* and *Hylotrupes bajulus*.

5.5.8.2.3.2 Termites

The control of termites enters into the scope of the PT8 and the PT18 depending of the use of the product. The definition of the product type is related to the use/mode of application of the product.

The reader is also invited to refer to the PT18 efficacy (section 5.6.4).

The curative treatments against termites are designed most of the time to kill the termite colony and prevent degradation of wood.

We can distinguish treatment applied to wood, for example treatment of art furniture, wood rubble from treatment applied to other support than wood for example soil or masonry.

If the product is applied on wood, then this product is covered by the requirement of the PT8. If the product is applied on another support than wood then it is covered by PT18.

We can distinguish three groups of termites:

- **Drywood termites (Cryptotermes, Kalotermes):** Drywood termites live inside of the wood which is attacked. The curative treatments applied to the wood consequently destroy the entire colony.
- **Subterranean termites (Reticulitermes, Coptotermes, Heterotermites):** The core of the subterranean termite colony is located in the soil. Termite workers
built tunnels to reach wood and destroy it. The treatment applied on infested wood kills the termites present inside of the wood but not the other members of the colony.

- **Tree termites (Nasutitermes):** Tree termites built epigeous (above-ground) nests, frequently on living trees. As a part of the colony has a subterranean location, termites infestations of wood in building may originate either from the nestmates located in the ground or in the epigeous nests. The treatment applied on infested wood kills the termites presents inside the wood but not the others members of the colony.

5.5.8.2.3.3 Fungi

Any claims for curative activity against wood rotting fungi will be supported by suitable efficacy data. No EN standard test protocols presently exist for curative treatments applied to wood. In general, as curative treatment, the infected wood is cut away.

In all cases CAs evaluate the data available to determine if they are sufficient for supporting the label claims.

5.5.8.2.4 Resistance

Information on resistance and the likelihood of its development is required for BPR Annex I inclusion and is also demanded for product authorisation.

At this point, no target organism resistance in field of chemical wood preservatives is known.

More information on resistance can be found in Chapter 6.2 of this TNsG on Product Evaluation, in the Chapter 10 on the TNsG on the BPR Annex I inclusion and on the website of the Insecticide Resistance Action Committee and the Fungicide Resistance Action Committee (FRAC: [http://www.frac.info](http://www.frac.info)).

5.5.8.3 Biological re-testing after changing the product formulation

While EN599-1 and EN 14128 provide the baseline for the testing requirements of new products, the corresponding annexes to both standards provide guidance on testing requirements when a formulation variation is caused by the addition, the substitution or removal of an active substance. Not all changes are subjected to re-testing and the informative sections of the standards do allow the consideration and taking into account of other data on a case by case expert judgment basis without additional testing. These data sources are not defined in detail but could include:

- Literature data;
- Certification of the product by recognised national quality scheme systems e.g. CTBP+RAL;
- National registrations;
- Others.

For any other changes in the formulation, refer to the informative annex A of EN599-1 and EN 14128. An explanation of Annex A of EN599-1 can be found in Appendix 12.

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28 Appendix 12 is under development and due to be finalised at the March WG meeting: the Appendix will be added by an update procedure planned for April/May 2017 with publication by September 2017.
5.5.9 PT9 Fibre, rubber and polymerised materials preservatives
The text for this section is under section 5.5.7 with PT7.

5.5.10 PT10 Construction material preservatives
Please refer to the General sections 1-3 and the Preservatives general sections (i.e. 5.5.1- 5.5.3) of this guidance.

5.5.11 PT11 Preservatives for liquid-cooling and processing systems
Please refer to the General sections 1-3 and the Preservatives general sections (i.e. 5.5.1- 5.5.3) of this guidance.

5.5.12 PT12 Slimicides
Please refer to the General sections 1-3 and the Preservatives general sections (i.e. 5.5.1- 5.5.3) of this guidance and the TNsG.

5.5.13 PT13 Working or cutting fluid preservatives
PT13 deals with preservatives for metal working fluids during their use in industrial processes. The general principles for evaluating PT13 products can be found in section 5.5.2 to 5.5.5. IBRG developed a method that allows to test the efficacy of active substances in a model matrix (“A Method for Determining the Basic Efficacy of Biocidal Active Substances used in Aqueous-Based Metal Working Fluids for their Protection in Use, IBRG FFG 16-001. This method should be used, unless it is justified that the method is not relevant for this specific product.

29 International biodeterioration research group (IBRG): www.ibrg.org
5.6 Pest Control (Main group 3)

5.6.1 General

The text for this section is under development and will be added at a future update.

Humaneness

According to the BPR (Article 19(1)(b) criterion ii and common principles point 49 and 76 in Annex VI) biocidal products should cause no unacceptable effects on the target organisms, including unnecessary suffering and pain for vertebrates (humaneness). This criterion is relevant for biocides in the Pest Control PTs14, 15, 17, 19 (repelling or attracting vertebrates) and PT20.

For these biocides an assessment must be made to demonstrate that the biocidal product does not cause unnecessary suffering in its effect on target vertebrates. This must include an evaluation of the mechanism by which the effect is obtained and the observed effects on the behaviour and health of the target vertebrates; where the intended effect is to kill the target vertebrate, the time necessary to obtain the death of the target vertebrate and the conditions under which death occurs must be evaluated.

A biocidal product intended to control vertebrates must not normally be regarded as satisfying criterion (ii) under point (b) of Article 19(1) unless:

- death is synchronous with the extinction of consciousness, or
- death occurs immediately, or
- vital functions are reduced gradually without signs of obvious suffering.

For repellent products, the intended effect must be obtained without unnecessary suffering and pain for the target vertebrate.

Guidance on the assessment of humaneness is currently not included in Volume II Efficacy Part B/C: Efficacy Assessment and Evaluation, but some general guidance can be found in the TNsG on Product Evaluation Chapter 6.

5.6.2 PT14 Rodenticides

General introduction

This section provides guidance on the methodology for the evaluation of the efficacy of rodenticide biocidal products according to the common principles laid down in Annex VI of the BPR in order to demonstrate that the condition for granting an authorisation in Article 19(1)(b)(1) of the BPR is fulfilled (i.e. the rodenticide is sufficiently effective).

5.6.2.1 Introduction

Depending on its intended purpose, a rodenticide may be regulated as a biocidal product or as a plant protection product. This document covers the rodenticides under the BPR.

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30 Biocidal product (PT14): Rodenticides used for the control of mice, rats or other rodents (by means other than repulsion or attraction) outside plant growing areas, for example in farms, cities, industrial premises etc, and inside plant growing areas not to protect plant or plant products.

Plant protection product: Rodenticides applied in plant growing areas (agricultural field, greenhouse, forest) to protect plants or plant products temporarily stored in the plant growing areas in the open without using storage facilities.

Where a product is used in both situations (as PPP and BP), it will need dual authorisation for the relevant use in accordance with the last subparagraph of Article 2(2) of the BPR. See also http://ec.europa.eu/food/plant/protection/evaluation/borderline_en.htm
which are used predominantly for the control of the house mouse (*Mus musculus*), brown rat (*Rattus norvegicus*) and the roof rat (*Rattus rattus*). Also other target species such as water voles (*Arvicola amphibius*), bank vole (*Myodes glareolus*), common voles (*Microtus arvalis*), field or wood mice (*Apodemus* spp.) and the grey squirrel (*Sciurus carolinensis*) are considered.

The four standard fields of use are given below with examples of possible fields of use:

- **in and around buildings**
  - in and around residential homes and other places in which people are accommodated;
  - in and around rooms intended for the preparation, processing or storage of food and beverages;
  - in and around stores, ships’ holds, factories and silos;
- **at waste dumps**;
- **in sewers**
  - in moist/wet environments such as sewers and watersides;
- **open areas**
  - open areas such as airports or leisure areas.
  - on animal husbandry farms (pigs, poultry, cattle, etc.);

Since the majority of rodenticides are bait products, most of this guidance deals with the evaluation of the efficacy of baits. In the text it is indicated where it specifically concerns bait products or concerns other types of rodenticides.

### 5.6.2.1.1 Aim

The aim of this document is to provide guidance on how to assess the efficacy of rodenticides, in order to ensure that only sufficiently effective products are authorised and therefore placed on the market for use. Animal welfare considerations are also taken into account.

### 5.6.2.1.2 Global structure of the assessment

Full assessment of efficacy is conducted on applications for product authorisations.

Information on effectiveness and intended use(s) of the product, together with its active substance(s), must be sufficient to permit an evaluation of the product and to define its conditions of use.

Efficacy studies (see section 2 below for the type of testing required) should be performed with the product to evaluate whether the product is effective for the intended use(s) at the specified doses. Efficacy tests should be performed with the product (in its final formulation) for which the authorisation is sought, and the composition of the test-product should be provided in the efficacy reports (especially for field tests and palatability tests). Any efficacy data from scientific literature are considered only as supportive data and should not replace efficacy data obtained from efficacy tests, which should be performed according to recognised standards. Data on the mortality and, in case of bait products palatability of the bait, resulting from these studies are compared with the specified criteria. The basis for the evaluation is the uses specified in the application (i.e. draft SPC) submitted by the applicant.

### 5.6.2.2 Dossier Requirements

Data on efficacy are required for every application for authorisation. The following information on effectiveness is required for each biocidal product in accordance with Annex III of the BPR:

1. Function (e.g. rodenticide) and mode of control (e.g. killing);
2. Representative organism(s) to be controlled and products, organisms or objects to be protected;
3. Effects on representative target organisms;
4. Intended concentration at which the active substance will be used and application rate;
5. Mode of action (including time delay);
6. The intended uses for the product;
7. Efficacy data to support these intended uses, including any available standard protocols, laboratory tests or field trials used including performance standards where appropriate and relevant;
8. Any known limitations on efficacy:
   8.1. Information on the occurrence or possible occurrence of the development of resistance and appropriate management strategies;
   8.2. Observations on undesirable or unintended side effects for example, on beneficial and other non-target organisms.

**Efficacy testing**

It should be noted that any efficacy testing conducted in the European Union on rodents should be in accordance with the principles set under Directive 2010/63/EU\(^{31}\) on the protection of animals used for scientific purposes. However, field trials with rodenticide products to control wild rodent infestations under actual use conditions that are carried out to demonstrate the results of already obtained data on palatability, mortality and humaneness are not considered animal procedures for the purposes of Directive 2010/63/EU.

For all types of rodenticides, efficacy has to be demonstrated in a laboratory trial and a field trial or alternatively in a semi-field trial and a field trial for each target organism submitted in the application, unless specified otherwise in this guidance. For roof rats it is also acceptable to demonstrate efficacy:

- in two or more well-conducted semi-field trials (for description see section 2.6 below), since in some regions infestations of roof rats are quite rare; or
- Two (or more) well-conducted field trial(s) in regions with infestations of roof rats.

In general it applies that tests should be of high quality to be considered for evaluation. For animal welfare reasons, in laboratory tests, the number of animals per test should be restricted to a minimum.

Positive results in field trials may outweigh negative results\(^{32}\) in laboratory studies, but only under the following conditions:

- there is at least one other laboratory study (or semi-field trial) with positive results for each study with negative results and;
- there is at least one field trial of high quality with positive results.

Positive results in laboratory studies **cannot** outweigh negative results in field and semi-field trials.

In case of testing only in semi-field or field trials (roof rats):

- at least two well-conducted semi-field tests or one field trial should have positive

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\(^{32}\) Negative results are those showing insufficient efficacy against the evaluation criteria (see section 4.1 of this Guidance).
results, respectively.

The following guidance is designed to be flexible and does not specify rigid protocols to which tests must be conducted. Published or unpublished data from any source will be considered provided the data are scientifically valid and relevant to the application. In all cases, the methods have to be described in sufficient detail to make the data reproducible. Ideally, data should be generated using national or internationally recognised testing methods and in accordance with the principles set under Directive 2010/63/EU on the protection of animals used for scientific purposes. However, applicants can also submit data generated using their own testing strategies where these are conducted and well reported to a sound scientific standard. In all cases, the data must allow a specific assessment of efficacy and, in case of bait products, palatability of the product. Anecdotal evidence will not be acceptable.

Assessment will be made in relation to the effectiveness of the product for the intended uses in the draft SPC submitted with the application. This assessment will take into account the animals that are considered to be harmful and are to be controlled (target species), indoor or outdoor use, the method(s) of application, application rates, use patterns of the product, maximum storage period (shelf life) of the product, together with any other specific terms and conditions concerning the use of the product.

The target species selected for efficacy testing should be appropriate to the geographic regions in which the product will be used. They should be named in the draft SPC for the product (either common or generic names may be used). Please note that in some countries specific rodent species are protected and no control action against them is permitted.

**Intended uses**

Examples of intended uses given in the draft SPC associated with the target organisms are:

- for use against house mice:
  - this will require testing against *Mus musculus*.

- for use against rats
  - this will require testing against *Rattus norvegicus* and *Rattus rattus*.

- for use against brown rats
  - this will require testing against *Rattus norvegicus*.

- for use against rats and house mice
  - this will require testing against *Rattus norvegicus, Rattus rattus* and *Mus musculus*.

- for use against rats in sewers
  - this will require testing against *Rattus norvegicus* with specifically treated bait (see section 2.4 below)

- for use against voles
  - this will require testing against at least two vole species which differ in size and behaviour, for example, water voles (*Arvicola amphibius*), bank vole (*Myodes glareolus*) and common voles *Microtus arvalis*.

- for use against a field mice (wood mice) species
  - this will require testing against the specified target species, for example the long-tailed field mouse/wood mouse (*Apodemus sylvaticus*) or yellow-necked field mouse (*Apodemus flavicollis*).

- for use against [name of target species]
  - this will require testing against the given target species. An example could be the grey squirrel (*Sciurus carolinensis*).
General intended uses given in the draft SPC, such as ‘for use as a rodenticide’ or ‘for use against mice’, with no further clarification of the target species are not acceptable. This is because it would allow use against rodent species for which the product is not tested and/or not intended. Concerning the target species, intended uses have to be species-specific (both for products authorised for professional and non-professional users).

Testing has to be species-specific, and for each target organism that is given in the draft SPC, a study should be conducted. This is because the biology, behaviour and susceptibility of target species, even within taxonomic groups such as rats, voles or mice, may differ considerably. For example, the brown rat (*R. norvegicus*) is more sensitive for anticoagulants than the roof rat (*R. rattus*), whereas it has been observed that the roof rat is more neophobic and will be less likely to accept baits than the brown rat. Mice are taxonomically very unspecific and may be applied to a broad range of species (e.g. *Mus musculus*, or various *Apodemus* species) with different biology, behaviour and susceptibility against the active substances. Vole species differ considerably in their size and habitat. Therefore, all target organisms given in the draft SPC have to be tested. If the authorisation of a rodenticide with a less specific intended use, such as ‘for use against voles’ or ‘for use against mice’ is applied for, the product has to be tested at least against all representative species of the respective taxonomic group. For voles there are products authorised under the plant protection products (PPP) legislation, but under some circumstances, there can be a need for biocidal product approvals (e.g. in case of invasions near buildings and disease spreading).

Resistance claims are allowed for products based on actives with a mode of action other than anticoagulants. For products based on anticoagulants there is differing opinions of permitting claims by Member States and therefore, until further discussions and decisions are made, such intended resistance claims must be considered on a case by case basis in discussion with the Member States. An intended use such as ‘for use against rats and/or mice resistant to the first generation anticoagulants’, is generally not possible, because test animals which are resistant to first generation anticoagulants are difficult to define and their degree of susceptibility may vary. Moreover, when a case of resistance is recognised in a field situation, it is generally advisable to use non-chemical methods like mechanical or electronic traps, rodenticide with non-anticoagulant mode of action, or the most potent anticoagulant rodenticides, and the use instructions in the draft SPC should generally contain a paragraph about resistance management. Therefore, a general intended use concerning resistance on an anticoagulant product may not be regarded as informative, since resistance generally refers to the active substance rather than a specific product.

### 5.6.2.2.1 Test animals

Although laboratory testing should preferably be performed on second generation wild animals housed in groups, the difficulty and constraints associated with obtaining and maintaining them for testing purposes is recognised. Therefore for tests conducted within the laboratory, animals sourced from recognised commercially available strains are acceptable.

In accordance with Directive 2010/63/EU, Articles 7 and 9 and Section A, 3.2. of Annex III, semi-field trials should preferably be conducted using wild rodents or their offspring. Although not preferred, it is possible to use strains that resemble wild strains in semi-field trials as an alternative. These strains should be outbred strains (e.g. Long Evans or Lister Hooded rats) which retain the behavioural characteristics of wild rodents, which includes neophobia, anxiety, and fully capable sensory organs (no impairment of

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33 This issue is under review and discussion and the guidance will be updated if the situation regarding resistance claims for anticoagulants changes.
seeing, hearing, smelling or taste). When laboratory strains that resemble wild strains are used, a short description of the behavioural characteristics as well as reasoning for the choice of the respective strain as test animals should be provided. Generally, the diet which rodents (laboratory and wild strain) receive prior to the tests can be crucial for their behaviour towards bait products. It is therefore important that, as far as possible, the study reports should also include information on the dietary history of the test animals. It is recommended that test animals should receive a rather broad diet during breeding. Where wild animals are used in laboratory or semi-field studies, these may be live trapped from the wild, reared in either outdoor colonies or under laboratory conditions such that it permits the animals to retain much of their natural physiological and behavioural characteristics. Breeding stock used for rearing wild rodents should not be selected for docile qualities or other characteristics that significantly alter their wild tendencies.

OECD Guidance Document on the recognition, assessment and use of clinical signs as humane endpoints for experimental animals used in safety evaluation (OECD, 2002) must be considered. Unnecessary suffering must be avoided (e.g. excessive weight loss/severe dehydration, persistent convulsions, cannibalism/self-mutilation, etc.) and animals should be checked regularly. Moribund animals should be euthanized in line with the requirements to apply humane end-points by using clinical signs to determine impending death.

Field trials should be conducted on wild rodent infestations and are not considered animal experiments provided the respective tests on efficacy, palatability and humaneness have been confirmed under controlled laboratory studies.

The purpose of Article 62 of the BPR is to minimise the number of tests on animals and not duplicate any studies on vertebrates that might be required by the BPR. While the objective is clear for laboratory tests and semi-field trials, for which animals are used on purpose, for field trials the situation can be seen from a different perspective. Where a field trial is carried out under real life conditions and the rodents subject to such field trial would have been to be killed/controlled in any case by using other authorised products, then it is considered that such field trial does not involve any duplication of testing. Therefore, field trials for PT 14 would be exempted from Article 62 of the BPR.

Concerning laboratory tests and semi-field trials, the objectives of Article 62 (of BPR) would be achieved by data waiving where there were already tests with a fully comparable bait containing an active substance with similar or lower toxicity (see Table 25 in section 5.6.2.2.7 below). In such cases read-across could be accepted provided that, where relevant, a LoA (Letter of Access) is presented by the applicant.

5.6.2.2.2 Laboratory studies for bait products

For testing the efficacy of bait products, two types of laboratory studies are available, mortality tests (i.e. no-choice feeding tests) and choice feeding tests. Since mortality tests give very little information in addition to data from the bait choice feeding testing and in order to reduce the number of animal experiments, mortality tests (i.e. no-choice feeding tests) are not recommended and are not required. However, many applicants may have no-choice studies on their products as they have been conducted in the past. These can still be submitted as part of the data package but no new studies should be conducted.

Tests conducted to EPPO or the specimen protocol (Appendix 12 of this Guidance) are preferable but other data will be considered on their merits. The study must be representative for the treatment. Depending on the intended aim of the product, the house mouse, roof rat, brown rat or other species should be used as the test animal. Wild strain testing is preferable and is most important for the bait-choice test. However, since this is probably impractical for some applicants, an outbreed lab strain (e.g. CD rats) which is likely to exhibit traits of the wild strain is accepted as surrogate.
Rodenticides with special indications, for instance foam products, which are taken up orally but are not bait products since they adhere to the rodent fur, require separate laboratory trials, where the conditions are properly simulated (see section 5.6.2.2.3 below).

**The bait choice feeding trials**

The aim of the bait choice feeding trials is to determine the palatability of the product for the test animal. If conducted on both fresh and aged product it may provide information on efficacy after a long period of storage of the product (see section 5.6.2.2.5 below). This test is preferably done with wild strain animals. In this test design, animals have the choice between a non-toxic food source (challenge diet) and the bait containing the active substance. Either the amount of bait consumed, in which the active substance is incorporated, or the mortality of the rodents is an indication that the bait is sufficiently palatable for a lethal dose to be ingested. Results are compared with the specified criterion (see section 5.6.2.4.1 below).

Make sure that the challenge diet is a product that the rodent is accustomed to.

Full details of the methods used should be provided and data should be presented to show the daily intake of both untreated diet and product, the palatability ratio (amount of product: amount of challenge diet) or product acceptance (amount of product eaten expressed as a percentage of total (product + challenge diet) consumption) for different sexes of rodent, any signs of poisoning and days to death, with appropriate statistical analysis. When no significant differences exist between the sexes, the data from the two sexes may be combined. Clinical observations should be conducted to determine mode of action, degree of suffering, duration of toxicosis prior to unconsciousness, etc. These data are optional but provide useful information, especially on new active substances.

In some cases comparison with normal food intake is inappropriate. For instance when fast-acting rodenticides cause a reduction in feeding activity or when only very small quantities of bait are required to cause effect. Therefore, the main criterion is not the percentage of consumed bait but the mortality resulting from poison uptake.

**Bait choice feeding trials with voles**

The test protocol for choice test against voles in the laboratory should be principally the same as for rats and house mice.

### 5.6.2.3 Laboratory studies related to contact rodenticides and gassing agents

**Contact rodenticides**

The information that should be available in order to demonstrate efficacy will include:

i) Estimates of time to death from individually or group caged rodents exposed to the product for stated periods of time. Reference to EPPO Guidelines (EPPO, 1986) should be made.

ii) Evidence from the laboratory that the target rodents will pick up the required dose from the application method is recommended.

**Gassing agents**

Rodenticidal gassing agents are typically used in gas-tight buildings, ships, airplanes, containers and storage locations or for burrow fumigation. The type of information that should be available in order to demonstrate efficacy will include estimates of the potency of the active substance and product by inhalation when applied as described in the use instructions in the draft SPC for the product.

There are no internationally recognised standardised test protocols for testing efficacy of rodenticidal gassing agents. In general, the dossier requirements are the same as with bait products. No-choice tests are not necessary. The dossier should include simulated...
use-tests as well as field tests. Simulated use tests should be conducted in gastight containers. The size of the container, duration of exposure as well as the concentration of the fumigant in the container should reflect a real-usage situation.

It has to be noted that the use of gassing agents in sealed rooms, buildings, ships, airplanes or containers (generally denoted here as “rooms”) is different from use in burrows (generally denoted here as “rodent burrows”). Hence, it has to be declared for which use an authorisation is applied for. For each type of use a field study must be conducted.

Generally, during each experiment the concentration of gas has to be monitored. The test reports should contain a detailed description of gas concentration, position of measurement points as well as the analytical method. The absence or presence of sorptive materials has to be documented.

Field tests for burrow fumigants should follow the protocol for rodent baits. It has to be demonstrated that rodent populations in infected objects can be eliminated. The study has to include a description of the burrow (location in the infested object, position of entrance holes), for example, Ross, (1986), and Méthode CEB n°254 (2013) listed in Appendix 14 of this Guidance. The methods for a population census before and after application as well as the mortality criteria are the same as for bait products (see Appendix 13 of this Guidance).

Field tests for rooms should include an estimation of the population size, but it is recognised that a feeding census is often not possible (e. g. in containers). In these cases, cages with the respective target organisms (mice, rats) should be introduced to the field object. Their placement should reflect the expected distribution of rodents in the object. It is important that some cages should be placed at spots which would represent “worst case scenarios”, i.e. places with air draft (since a room or container may not be perfectly airtight) or in hideouts. The test report should contain a detailed description of placement of the cages, as well as number, age and sex of the test rodents. Exposure time should be according to the use instructions in the draft SPC. After exposure, the number of dead rodents within the sealed room/compartment and/or inside the cages must be determined. Field tests with no scientifically comprehensible data on population reduction or mortality will not be accepted. In cases where a sufficient number of caged rodents have been introduced to field objects for efficacy testing, simulated use tests can be waived. The mortality criteria are the same as for baits.

Considering the risks linked to the presence of rodents in an airplane, an efficacy of 100% is necessarily required. Indeed rats and mice (these latter being able to hide in places of low volume and completely inaccessible in airplanes) can cause damage, besides the problems of public health, which affect the safety of the airplane and the passengers. Besides possible damage linked to the urine on the electronics, these rodents possess incisors with continuous growth which oblige them to eat away permanently at any type of materials (threads, girdles, steering cables, printed circuits.). There is therefore no tolerance threshold, because a single rodent can cause irreversible damage. In order to make sure that the dose administered according to recommendations and within the framework of fumigation under actual conditions, achieves the required mortality concentrations, the following requirements have to be carried out:

- during fumigation, the measurements of the “CT” (measured effective concentration x time of fumigation) must be systematically taken. The aircraft to be fumigated may not be completely airtight and gas leaks may occur, therefore measures need to be taken for the required 100% efficiency;
- for every trial, the data for the calculation of the “CT” are to be collected from the start of fumigation with statements of concentration (two minimum test points according to the type of airplanes) made at regular intervals (frequency of five
minutes) for the duration of fumigation as claimed by the applicant. It is suggested that these data should be collected for two operations of fumigation;

- to make sure that there is good distribution of the gas at lethal concentrations in the entire airplane, rats in individual cages (five rats per test point) must be placed next to all the concentration test points. This will allow estimation of the relation between the measurements, the “CT” and the mortality of the rodents;
- a statement of temperature and humidity should be made.

In case a gassing agent is used in combination with a specific device or is part of a device (e.g., traps), results from laboratory choice tests as well as (semi-)field tests should be submitted. A no-choice test is not necessary; (semi-)field tests should have the same protocol as field tests for baits. A population census like in bait tests before and after application is needed. The mortality criteria are also the same as for baits.

5.6.2.2.4 Laboratory studies related to specific efficacy claims regarding suitability of bait products for use in damp conditions

Where it is claimed that a product is suitable for use in sewers or under damp conditions, the retention of palatability (such as the effect of the heat and humidity on palatability) should be tested in a choice test against all claimed target species, using product that has been specifically pre-treated to simulate such conditions. Please note that sewers are generally only infested by the brown rat.

For this purpose, the bait product must be exposed to a warm and humid surrounding for at least five days. Bait which is pre-treated in such conditions, may be tested either with experimental animals or, preferably, in a semi-natural test system (pen test). The total number of animals should be 10 to 20.

Below a preferred test protocol is described. Other test protocols will be considered on their merits and are acceptable provided they are scientifically justified.

The bait portions/blocks must be weighed before treatment and then exposed to preferably 30°C to 35 °C and 80 to 99% RH for five days. Stable conditions can best be achieved in a climate chamber. The bait should be placed in a water-permeable clay bowl, which itself is placed in a water-tight clay dish. The clay dish contains water, which permeates through the wall of the clay bowl with the bait, so that the surface of the clay bowl is permanently wet to simulate the moist surface of sewer walls. Each pre-treated bait portion/block is applied to the test animals for one day. The bait portions/blocks are then removed and replaced with new pre-treated bait. Since bait exposure to warm and humid conditions is for five days, the baits must be pre-treated stepwise, so that for each testing day, bait with exactly the same pre-treatment time will be applied. The test chamber or test cage is not acclimatised, i.e. the test animals do not experience specifically warm or humid conditions. The bait is replaced daily with freshly pre-treated bait and is offered in a wet clay bowl to maintain surface moisture, so that the bait remains wet and does not dry out during the 24 h exposure to the test animals. Specific acclimatisation of test chambers/cages to high temperatures and humidity is therefore unnecessary and not advisable, as the test animals will most likely originate from laboratory colonies which are kept under normal conditions (i.e. moderate humidity and temperature). High temperatures and humidity may cause them to react with behavioural disturbances.

To determine the bait consumption, bait is removed from the test chambers/cages each day and weighed back. After this, the bait should be dried, preferably by placement in a drying oven at 30 to 36 °C (note: since most bait blocks contain a significant portion of

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34 Field tests may be accepted in case of a controlled situation without re-entry of rats, but laboratory studies are preferred.
paraffin, the temperature for drying must not be too high). Bait portions/blocks are then weighed until no further weight decrease can be measured (i.e. the bait lost all water and is dry).

To calculate the bait uptake, it must be taken into account that the initial weight of the bait is fresh weight, whereas the final weight after bait application to the rats and subsequent drying is the dry weight. Thus, the difference between both is not exactly the amount of bait consumed by the rats, since fresh baits may contain moisture (which adds to the fresh weight at the beginning of the experiment, but is removed after drying for the final weight determination). Hence, the water content of bait must be determined by placing five untreated bait portions for each product in a drying oven until no further weight decrease is determined. The difference between the fresh and dry weight is then taken into account for the determination of the amount of bait uptake (Equation [1]):

$$b = f - \frac{d}{(1 - w)}$$

Where:

- $b$ is the amount of bait taken up
- $f$ is the fresh weight of the bait prior to heat and humidity exposure
- $d$ is the dry weight after bait application, consumption and drying
- $w$ is the proportion of water content of the bait (determined through drying of untreated bait).

The relative portion of bait taken up by the test animals in relation to overall food consumption can be then calculated as (Equation [2]):

$$c = \frac{\sum b}{\sum b + \sum a} \times 100$$

Where:

- $c$ is the percentage of consumed bait during the test
- $b$ the amount of bait taken up (corrected after Equation [1])
- $a$ is the amount of challenge diet taken up.

5.6.2.2.5 Studies related to specific efficacy claims regarding to the shelf life of bait products

When a bait product is claimed to be effective after a long period of storage, it is necessary to demonstrate that the product will still be effective and palatable after the stated storage period (i.e. shelf life). Analytical studies on active substance content are therefore not sufficient to support shelf life claims of bait products.

Based on expert opinion, most bait products have been found to be effective and palatable for 24 months (with preservatives). Efficacy testing should therefore only be provided for:

- bait products with preservatives that claim a shelf life of longer than 24 months;
- bait products without preservatives that claim a shelf life of longer than 12 months;
- bait products for which the degradation of the active content is >10% and assessment of the degradation on the efficacy is needed to substantiate the shelf life claim.

For bait products with a shorter shelf life claim than stated above, no efficacy tests on aged bait (i.e. product at the end of maximum storage) have to be provided. For these products it is sufficient to provide tests on fresh bait (i.e. newly produced product).
For bait products with a longer shelf life claim, the applicant must deliver data on the palatability of the product at the end of maximum storage for all target organisms claimed. The palatability of the aged product preferably is tested in field trials, but can be tested in field trials, provided these tests are scientifically valid (see section 2.6 below). Accelerated ageing studies, i.e. palatability studies in which the product tested is stored under challenging conditions, are not acceptable as these cannot simulate longer storage periods.

5.6.2.2.6 Field trial and semi field trial

The following text describes the field and semi-field testing of bait products, but is also largely valid for other rodenticide products.

Field trials

The aim of the field trial is to demonstrate the results on the effectiveness (palatability, mortality and humaneness) obtained during laboratory studies of the rodenticide product containing active substance under actual use conditions for the purposes of marketing authorisation. Field trials should only be performed once efficacy, palatability and humaneness have been confirmed in laboratory (semi-field) studies under Directive 2010/63/EU.

Tests conducted to EPPO or the specimen protocols (Appendices 13 and 14) are preferable but other data will be considered on their merits. Depending on the intended use(s) of the product, populations of the respective target organisms (house mice, brown rats, roof rats or others) are used for this trial.

Ideally, sites chosen for field trials should be representative of the range of locations where the rodenticide is to be used (indoor/outdoor), and should be infested with sufficient numbers of the target rodents so that the effectiveness of the product can be clearly demonstrated. It is advantageous if the rodent infestations on the sites chosen are, as far as possible, discrete and not subject to potential rapid re-invasion. Rodent activity on the site should be determined before and after treatments using at least two standard techniques.

Sketch maps of the sites approximately to an indicated scale showing all the important features including signs of infestation and location of rodenticide application should be provided. The amount of bait applied at each bait point and the distance range between bait points should correspond to those given in the draft SPC. Replenishment of the bait should follow intervals given in the draft SPC. Bait exposure should normally be for 4 days for acute products and 30-40 days for multi-dose products after the first bait uptake or less when full control is achieved. Data should be presented to indicate levels of rodent activity both before and after treatment, amounts of bait consumed and all relevant information regarding treatment details.

Semi-field trials

As an alternative or addition to 'field' trials, evidence of the efficacy of a rodenticide product may be obtained with semi-field trials (otherwise referred to as pen trials). A semi-field trial simulates field conditions under controlled laboratory conditions. Bait acceptance and bait uptake in the field is strongly influenced by the social behaviour of the target species. Both rat species (R. norvegicus and R. rattus) as well as house mice (M. musculus) are social animals, and food exploration is largely social in these species. Hence, the most important field condition to be simulated is the presence of conspecifics, i.e. the semi-field trial has to be conducted with groups of rodents. Group size should be at least 10 animals in tests with both rat species and at least 10 animals in tests with house mice. Sex ratio should be approximately 1:1 although single sex groups may be used with robust justification, e.g. to avoid unacceptable levels of aggression. Groups should consist of related animals to avoid intraspecific aggression. The test animals should either be directly caught in the field, or be bred from wild catches, as only wild-
strain rodents show the typical behaviour of the target species which could be expected in the field. A test with laboratory strain rodents cannot be regarded as a proper simulation of field conditions.

The test arena should provide shelter for the animals, as well as sufficient space for the animals to roam. The minimum space requirement would be ≥ 0.5 m² per rat and 0.25 m² per mouse. If possible, cage enrichment such as branches, ladders, tunnels and wooden nest boxes with nest material may be provided and details on this should be given in the test report. Cage enrichment should be designed in a way that daily inspection for dead rodents and spilled bait material and feed causes only minimum disturbance.

The rodents have to be familiarised for at least three days with the test arena prior to bait exposure. The semi-field trial is always a choice test, and a suitable challenge diet must be provided together with the bait. The amount of bait applied should correspond to the amount given in the draft SPC. Bait exposure should normally be for 4 days for acute products and 30-40 days for multi-dose products. Bait exposure must be followed by a 14 day post baiting observation period.

Field tests with voles

For efficacy testing of products against voles, the test protocols for house mice and rats are only suitable when the infestation is inside a building. Efficacy testing outside of buildings should be conducted with a specific protocol. In contrast to rats and house mice, voles excavate and inhabit galleries (tunnels beneath the surface) for food exploration and nesting.

For each field test with voles, one test plot and one control plot should be investigated. Principally, the test protocol is the same for oral baits and gassing tablets/pellets. The pre-treatment and post-treatment censuses are conducted by counting occupied galleries. For this, at least ten galleries should be opened on each plot (treatment and control). After 24 h, the number of refilled galleries is then counted. The number of refilled single openings is set into relation to the number of openings as an indicator for vole activity. Depending on the vole species, an alternative census method could be the closing of burrow openings. Reopening of burrows is then counted as a sign for activity. During the treatment, vole activity should be controlled after 5 and 10 days with the same method.

Application of the rodenticidal product should follow the use instructions in the draft SPC. Normally, one bait portion has to be placed in each gallery. Replenishment of the bait should follow intervals given in the use instructions in the draft SPC. Bait exposure should be for 14 days. The efficacy is then calculated as (Equation [3]):

\[
E = 100 \times \left(1 - \frac{t2 \times c1}{t1 \times c2}\right)
\]

Where:

\(E\) is the efficacy,
\(t\) are treated plots
\(c\) are control plots,
\(t1\) and \(c1\) are the ratios of refilled galleries/open galleries before treatment
\(t2\) and \(c2\) are the ratios of refilled galleries/open galleries after treatment.

Treatment and trials with oral bait should be undertaken in spring or autumn, as in the winter not much activity is to be expected, and in summer other food sources than the bait are too abundant.
5.6.2.2.7 Waivers

Waiving of laboratory trials or semi-field trials will reduce animal testing. For bait products, because the composition of the bait determines the palatability and hence efficacy of the product, even small changes in ingredients may affect the attractiveness. This may differ between target organisms and is difficult to predict in advance.

Semi-field trials

Laboratory testing of bait products (bait choice test or semi-field trial) should always be requested for new active substances, or if a product was altered regarding the active substance concentration and/or bait formulation. One exception would be if there were already test data with a fully comparable bait, i.e. containing a different active substance but otherwise the same or similar formulation with the same mode of action and similar or lower toxicity; (see Table 25 below for a ranking of toxicity of existing active substances), in such cases read-across could be accepted; however if the two formulations contained the same active substance, then the concentration of the active substance would need to be the same.

Field trials

Field trials are always required when the composition of a product is changed. Exceptions could possibly include changes of minor importance in ingredients that are likely not to have an effect on palatability or efficacy, such as change in colour of a product. In case of waiving, the applicant needs to provide a robust justification why no testing was performed.

Read-across between species is generally unacceptable unless the applicant can demonstrate that there is no significant difference in the susceptibility and behavior of the species.

Table 25: Toxicity ranking of known active substances used in anticoagulant rodenticides based on LD 50 (acute) data of brown rats and house mice compiled from CA-Reports, ranking from high (1) to lower toxicity (3)

<table>
<thead>
<tr>
<th>Rank of toxicity</th>
<th>Active substance</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Flocoumafen, brodifacoum, difethialone</td>
</tr>
<tr>
<td>2</td>
<td>Bromadiolone, difenacoum</td>
</tr>
<tr>
<td>3</td>
<td>Chlorophacinone, warfarin, coumatetralyl</td>
</tr>
</tbody>
</table>

5.6.2.2.8 Biocidal Product Families (BPF)

A BPF of rodenticide baits may contain several bait products with different formulations, for example, various grain, block, paste and gel products. Each bait formulation should be allocated to a different meta-SPC\textsuperscript{35}. Each bait formulation within the BPF has to be tested, because it cannot be predicted which form is the least palatable. It would also be difficult to select one product that could be regarded as a ‘worst case scenario’ for testing all the formulations. Within a given meta-SPC, an individual product should only be tested to consider the minimum level of efficacy within the concentration ranges of the active substance in that meta-SPC.

\textsuperscript{35} See Q&A pair number 6 in Annex IV of the Note for guidance "Implementing the new concept of biocidal product families" (CA-Nov14-Doc.5.8 – Final.rev2). [https://circabc.europa.eu/w/browse/c309ae58-bdd7-421d-a678-8d8ac361d4e0]
5.6.2.3 Methodology of assessment

There are many standard test methods currently available that may be appropriate for the assessment of the effectiveness of rodenticides. A list of such test standards is presented in Appendix 14 of this Guidance.

In addition to the standard test methods presented in Appendix 14, specimen protocols for a Choice Test and a Field Test are presented in Appendices 12 and 13 respectively. These Appendices are intended only to provide further information regarding the types of studies that may be utilised to assess the efficacy of some rodenticides, and some of the factors that should be taken into account.

Any known limitations on efficacy (including resistance) should be considered during the assessment. Possible restrictions, risk mitigation measures, or recommendations concerning the use of the product in specific environmental or other conditions can be considered. Possible factors that can reduce the efficacy, for instance hot, cold or humid environments or the presence of other substances, in addition to the grounds for these should be stated. Possible recommendations concerning the avoidance of the continuous use of the product in order to prevent the selection and spread of resistant strains and the grounds for these (see TnSG on Product Evaluation and a report on risk mitigation measures for anticoagulant rodenticides as biocidal products 36). State if the product cannot be mixed with, for example, other biocidal products or if the use of the product with other biocidal products is recommended. The guidance given on resistance for the corresponding data requirement of the active substance also applies here. The study results are compared directly with the criteria for efficacy (see section 4.1 below).

5.6.2.4 Assessment of authorisation

5.6.2.4.1 Norms and criteria

In accordance with Article 19(1)(b)(1) of the BPR, a biocidal product may only be authorised if it is sufficiently effective. This is implemented in the following way.

In general rodenticide products are normally considered to be sufficiently effective if the following results can be achieved:

- required results in laboratory test and semi-field test:
  - ≥90% mortality within a relevant time frame
- required results in field test:
  - Monitoring of the test population should show a ≥90% decrease of the population

Rodenticide bait products are considered to be sufficiently effective if the following results can be achieved:

- required results in the bait choice feeding test, semi-field test and sewer test (if claimed):
  - ≥90% mortality. The percentage of ingested bait containing the product should be normally ≥20%, but it may be lower because a mortality of ≥90% the product would still be effective. In case of a bait ingestion <20%, justification should be provided.
- required results in field test:
  - feeding on census bait after treatment should be reduced by at least 90% from the levels of feeding on census baits before treatment. When other types of quantitative monitoring of the test population are used, such as

36 “Risk mitigation measures for anticoagulant rodenticides as biocidal products” [https://circabc.europa.eu/sd/a/343a61cd-b8d4-40af-9e5c-4f763aea3240/CA-Nov14-Doc.5.1%20-%20draft_final_report_RMM.docx6].
tracking activity measurement and census by trapping, they should sufficiently show the decrease of the population (≥90%).

The efficacy of the product after a specified storage time (e.g. shelf life as claimed in the use instructions in the draft SPC) is also taken into account when assessing efficacy of a rodenticide bait.

Deviations from the norms are possible, but must be justified in the application. The Competent Authority will evaluate any justification on a case-by-case basis, consulting the other CAs where appropriate and decide whether it is acceptable or not.

In order to promote the development of new types of products (less toxic, more humane), a mortality <90% could be acceptable when the product is used as an accompanying method, (i.e. used with another product to demonstrate efficacy). but not as a stand alone product. However, mortality of these new type of products should not be <50%. The use of a product as an accompanying method should be reflected in the use instructions in the draft SPC.

For the assessment of resistance, reference is made to TNsG on Product Evaluation. Information on resistance testing techniques is also available from the Rodenticide Resistance Action Committee (RRAC) and Prescott et al. (2007).

5.6.2.5 References for PT14


5.6.3 PT15 Avicides, PT16 Molluscicides, vermicides and products to control other invertebrates & PT17 Piscicides

Please refer to the General sections 1-3 of this guidance and the TNsG.

For product-type 16, EPPO guidelines for efficacy testing are highly recommended (e.g. EPPO guidelines 95 for molluscicides in terrestrial environment).

5.6.4 PT18 Insecticide, Acaricides & other Biocidal Products against Arthropods+ PT 19 Repellents & Attractants (arthropods)

5.6.4.1 Introduction

Depending on its field of use a product to control, repel or attract insects and other arthropods may be classified as a biocidal product or plant protection product. This section covers the products to control, repel or attract insects and other arthropods in the category of biocides, which are products against all pest arthropods except those that are plant parasitic.

Attractants used in monitoring traps to assess the necessity and the success of pest management measures are considered outside the scope of Biocides Directive (Manual of Decisions, “Traps for monitoring purposes”).
This first section gives a general introduction. The following sections describe per insect or per type of use what the requirements for efficacy testing are. Information is missing on some of the organisms to be controlled with these products and also some of the uses and types of products. For instance, little information is provided on attractants (e.g. sex pheromones etc.) and treated articles (e.g. insecticide treated mosquito nets etc.). These data gaps will be filled in a future update of this guidance.

5.6.4.1.1 Aim

The aim is to assess the efficacy of biocidal products, to ensure that only effective products enter the market.

5.6.4.1.2 Global structure of the assessment

A full assessment of efficacy is conducted for applications for product authorisations. Factors, which are taken into consideration during assessment of the efficacy for a biocidal product to control, repel or attract insects and other arthropods for which authorisation is sought, are:

- the target organism to be controlled, repelled or attracted;
- the physical state in which the product is applied (e.g. liquid/powder/bait);
- the areas of use, these may be:
  - in and around residential homes and other spaces in which people are accommodated;
  - in and around spaces in which animals are accommodated
  - in spaces intended for the preparation, processing or storage of food and beverages;
  - in empty stores, ship’s holds, factories and silos.

Information on effectiveness and intended uses of the product, together with its active substances, must be sufficient to permit an evaluation of the product, including the nature and benefits that accrue following use of the product in comparison to suitable reference products or damage thresholds, and to define its conditions of use.

A combination of laboratory studies, rigorous simulated-use laboratory studies, or field studies can be used to evaluate whether the product is effective for the requested use(s) at the specified doses. Data from these studies are compared with the specified criteria.

Assessment will be made mainly in relation to the claims for the effectiveness of the product made on the product label. This assessment will take into account the pest(s) to be controlled, indoor or outdoor use, the method(s) of application, application rates and use patterns of the product, maximum storage period of the product, together with any other specific claims made for the product. More information on different aspects of the label claim can be found in Appendix 1. Appendix 16 shows examples of possible label claims.

5.6.4.1.3 Dossier requirements

Data on efficacy are required for every application for authorisation.

The following guidance is designed to be flexible and does not specify rigid protocols to which tests must be conducted. Published or unpublished data from any source will be considered provided the data are valid and relevant to the application. In all cases, the methods and results have to be described in sufficient detail to make the data reproducible and to allow a full assessment. Anecdotal evidence will not be acceptable.

Ideally, data should be generated using internationally recognised testing methods (ISO, CEN, OECD, WHO etc.). Several international standard test methods currently exist for insecticide/acaricide products. A list of these is presented in Appendix 17 to this document.
If there are no guidelines available or guidelines are not suitable, the applicant may use their own methods (intra-company Standard Operating Procedures), on condition however, that the studies are scientifically robust, well reported and provide a clear answer to the question. In addition, the test methods applied and the test conditions should be clearly and fully described and must address the efficacy claim that appears on the product label. The use of existing guidelines, with revisions to make the guideline more suitable for the specific product or company conditions, is also possible.

For each test information such as the following should be available:

- the names of actives substances and their respective concentration in the tested formulation;
- as the formulation may be very important for the efficacy, if the test item differs from the product to be authorised, its composition should be provided;
- a statement about what is expected from the test, what should be determined and with which precision. Power and sample size considerations should be included as well;
- description of the test conditions (size of cage, floor area, presence of harbourages, presence of (alternative) food, water, temperature, photoperiod, location, weather conditions);
- are the test organisms allowed to acclimatise to the test conditions before the test? For how long?
- how many test organisms are present (sample size)?
- describe population composition (males, gravid or non-gravid females, nymphs, larvae, age of the population or generation number F1, fed or unfed) noting that the feeding behaviour of some insects (i.e. Blattella) changes during their life;
- are the test organisms starved prior to the test?
- are field strains or known insecticide-resistant strains tested (claim “effective against strains resistant to x“)?
- a description of the history and origin of the test strain;
- is bait consumption determined? If so, a covered bait should be included to determine weight loss due to evaporation to correct weight loss of the exposed bait for actual consumption;
- are one or more alternative baits (e.g. registered reference products) or alternative food source present in the same test container or protocol?
- raw data should be available for each study, rather than just a summary of the results;
- show the results of both tests (with biocide) and control (without biocide) treatment, preferably in a table;
- size of the test population in the field before and after the test;
- description of the monitoring methods used before, during and after the test;
- statistical methods, if appropriate.

5.6.4.1.3.1 Test design

Although in general nationally or internationally recognised testing methods are preferred it is not always possible to use these. For some products no standard methods are suitable. In that case a test has to be designed.

Various factors must be considered when designing the tests, for example the number of test individuals (insects, mites, other arthropods) needed. The ultimate aim of relevant considerations should be to design experiments that economise on test individuals, but on the other hand generate sufficient power to detect effects of a magnitude considered important to demonstrate. To save test individuals, replicate tests are conducted.
Another argument for using replicates is to account for the variation among test individuals in susceptibility and responses to the biocides. Numbers of test individuals per replicate group and dose level (treatment group) as well as the number of replicates in the entire study need to be established prior to conducting the tests. As the improvement in power wears off substantially as the number of replicates increases beyond five, it is usually sufficient to conduct four or five replicate tests at each dose level, employing 10 (or 20) test individuals each. The precise needs will depend on the size of the variances, relative and absolute, between and within the replicates. This can differ between insect species and test design. Sample size should be adequate to detect differences among groups (untreated vs treated) with a statistical power of at least 80%. Some details on these issues are outlined at the end of each section.

Useful information on the principles of test design, analyses end evaluation of efficacy trials can be found in the EPPO standards pp1/152(3) and pp1/181(3).

5.6.4.1.3.2 Test examples

In the following sections (5.6.4.1.3.2 to .15) examples are given of what kind of tests can be expected for efficacy testing. Sometimes these examples are a summary of a standard test, in other cases a company test is described or a general idea of what the test should be like is given. There is a great variation in how specific the description is. For instance, the number of replicates is given only when this was determined in the test described.

In all cases these tests are only meant as examples, not obligatory requirements. Since products against insects and other arthropods are so diverse in application method, mode of action etc. the guidance cannot possibly cover all possible ways of controlling arthropods.

5.6.4.1.3.3 Laboratory versus (semi) field trials

Laboratory and field trials with the test arthropods are normally needed to assess the efficacy of the product. Field trials are not mandatory in some cases, as outlined in the sections on specific groups of arthropods below. In some cases when robust field studies are available, laboratory studies can be waived. If the product is applied as a bait, the entire bait, including the bait-box if applicable, should be tested, not only the product which is contained in the bait. When efficacy against several insects or other arthropods is claimed not all organisms have to be tested when appropriate bridging studies are available.

In the case of field trials where true replication is almost certainly impossible to achieve, a full description of any factors that might be expected to influence product performance should be given. These may include the risk of re-invasion from adjacent areas, general levels of sanitation, treatment history etc. and are intended to provide the authorities with information to assist with the interpretation of the results obtained.

In the following sections (5.6.4.1.3.2 to .12) more specific dossier requirements are given per pest species. In most cases a general description of a proposed method is provided. This is only to give an idea of what kind of tests should be provided. More detailed descriptions of tests can be found in the standard test methods (norms) listed in Appendix 17. This is a list of all available methods (as far as we know now) without distinction on usefulness, repeatability, order of acceptability or robustness. Some norms might have a different approach than described in the section for that insect. If this approach is more suitable for the product under investigation the norm should be used.

5.6.4.1.3.4 The importance of controls on efficacy studies

The importance of control experiments for efficacy studies must be stressed with regard to the efficacy evaluation. Studies should be conducted alongside negative controls wherever possible to provide a reference point for the treatment results. A useful definition of this term is given: "A negative control situation may be one in which the
The experimental design of the study is identical to that of the biocide challenge test except that the biocidal agent is not applied in the control study. A biocidal agent may be considered as the formulation or as the actual biocidal active gradient itself.”

The negative control trial should normally be of similar size (i.e. number of replications) as the test itself, to make statistical comparison possible and to get a fair impression of control mortality.

A relevant reference product (authorised, commercially available) can often be included at label rates in a protocol for laboratory and/or field studies as positive control. Unfortunately at this moment no standard reference products are available, however, an authorised reference can be included.

It is recognised that generation of such control data can be relatively straightforward in well-defined test situations such as laboratory and simulated-use tests. However, it is also recognised that this can present a problem in field situations, where control sites may not be environmentally equivalent to the treatment site.

In such instances, there may be an alternative means of generating reference data other than collecting data from an untreated site. This method may involve pre-treatment monitoring of the site in question. This monitoring must be quantitative, e.g., assessment of numbers of trapped insects. In these instances, a ‘baseline’ infestation level would be established through such monitoring and then the effect of treatment on this baseline can be assessed. Post-treatment monitoring is required for this method.

5.6.4.1.3.5 Specific data to support label claims

In assessing the efficacy of a biocidal product to control, repel or attract insects and other arthropods competent authorities should in particular take the following parameters into account:

- target organisms/spectrum of activity;
- mode of action/effect;
- use patterns/methods of application;
- dose rate.

The data provided in support of the efficacy claims must be sufficient to cover these key parameters.

5.6.4.1.3.6 Examples of specific label claims with respect to target organisms

For specific target pests where only efficacy against one insect/arachnid order or a certain family within that order is claimed, data against only a limited number of pest species will normally be required. To illustrate this point, a number of examples are given below:

- FOR USE AGAINST FLEAS - Data against the cat flea (*Ctenocephalides felis*) or the dog flea (*C. canis*) should normally be available;
- FOR USE AGAINST COCKROACHES - Data against two key species such as German cockroach (*Blattella germanica*) and the oriental cockroach (*Blatta orientalis*) should normally be available;
- FOR USE AGAINST DUST MITES - Data against *Dermatophagoides sp.* should normally be available.

In the European tropical overseas regions, the most common genus encountered could be different. A specific claim should therefore be proposed, with referred target organisms. This special request could concern for examples termites, cockroaches or mosquitoes.
5.6.4.1.3.7 Examples of broad label claims with respect to target organisms

Broad label claims, such as "crawling insect killer" or "flying insect killer", should be accompanied by qualification of the range of pests against which the product may be used. When broad claims are made, data on representative pest species will need to be provided for the range of pest orders against which efficacy is claimed.

Representative pests from these orders will have to be appropriate to the use pattern of the biocidal product i.e. the environment of the areas to which the biocide is to be applied and the nature of the application (e.g. whether it is a space application or surface application) will define the most appropriate pests to be tested.

For each order stated, at least the principal target species will need to be tested for public hygiene use, before a general claim is likely to be supported. In more specific areas, such as use against stored product pests, data on at least two major representatives of the orders in question will normally be needed before a general claim is likely to be supported.

Where such a claim covers a diverse range of pest habitats and pest morphology and biology, data from a greater number of representative species will need to be provided. Appendix 16 shows examples of possible label claims and the test species required.

When cockroaches are used as a reference species, it can only be used for the general claim "crawling insects". If efficacy against other insects are claimed specifically (e.g. crawling insects including bed bugs) tests against these other insect should also be provided. Also if a company wants authorisation for more specific use with the same product they have to present specific data on the specific pest they are claiming. This is a consequence of the use of “reference species”, which should not be a way of short-circuiting the evaluation for efficacy.

5.6.4.1.3.8 The distinction between professional and consumer products

In some cases the dossier requirements and norms and criteria for the evaluation may differ between professional and consumer products. Products used by professionals must have a high level of efficacy since the objective is to eradicate the infestation. For consumer products an immediate knockdown or repellence is often more important than eradication, of course depending on the claim. For instance a spray against cockroaches does not necessarily have to eradicate the whole population but it should work fast. Consumers want to see that the insect/arthropod dies/knocks down immediately after they spray. For consumers it is difficult to eradicate a whole cockroach population since reinvasion from other premises will take place, therefore eradication does not always have to be proven. For each pest group it will be listed whether requirements differ for consumer and professional products.

5.6.4.1.3.9 The distinction between principal target and secondary/incidental target pests

Screening tests (see sections below for details) can be used as bridging studies, showing similar effect of the product to different pest species, after which in some cases field studies can be waived for secondary target pest species.

5.6.4.1.3.10 Claims for residual efficacy

Most insect/arthropod pests are cryptic and/or nocturnal in behaviour and are unlikely to be contacted directly by a spray during application. For this reason many control programmes involve the use of relatively stable active substances applied to buildings and other surfaces to leave residual deposits. These compounds are intended to remain chemically active and therefore effective for periods of weeks up to several months following treatment, i.e. they have a high residuality. Residual life is a term to describe the period during which the biocide will be present in sufficient quantity to kill target pests, which walk upon it for a sufficient period of time to pick up a lethal dose.
Thus the amount of biocide residue deposited on treated surfaces is critical to the effectiveness of many treatments against crawling (and flying) pests. Ideally, the amount of residue deposited should be determined for instance by calculation or under actual or simulated use conditions. The method(s) of determination must be available with the test data.

Residual efficacy must be proven in tests. Usually, laboratory testing is performed to establish the efficacy direct after application and at the end of the residual life of the product.

The types of surfaces to which residual products are applied must be reported since surface type has a pronounced effect on the amount of active residue available to pests. In general a selection of both absorptive and non-absorptive surfaces, related to the label claim, should be tested when supporting a residuality claim for crawling (and flying) pests. These could include vinyl tile or linoleum, stainless steel, painted and unpainted wood, carpet, concrete and ceramic tile.

Efficacy data submitted to the competent authority in support for residual treatments should indicate the appropriate dosage and the utility of the formulation when used as directed.

5.6.4.1.3.11 Residual treatments may also involve the use of palatable baits.

When a bait product is claimed to be effective after a long period of storage, it is necessary to demonstrate that the product will still be effective and attractive after the stated storage period. The applicant must either submit data for palatability of the product at the end of maximum storage or alternatively (in case of a new product) data for a stress test with ‘accelerated ageing’, i.e. a palatability test with the product which is stored under challenging conditions (see FAO accelerated test).

5.6.4.1.3.12 Claims relating to outdoor use

When products are intended for outdoor use, tests should normally demonstrate efficacy under outdoor conditions. Changes in temperature and rainfall can have effect on the efficacy of the products. In general field trials cover this outdoor use. In some cases a field trial can be waived when a laboratory test can be done under worse case conditions.

5.6.4.1.3.13 Mode of action

There are a variety of modes of action and possible effects on target organisms derived from the proposed use of a product to control, repel or attract insects and other arthropods. The available data should give brief details to indicate the route and nature of the action (e.g. whether action is by contact or stomach poison), and the nature of the effect (e.g. cholinesterase inhibitor, chitin synthesis inhibition, juvenile hormone analogue giving rise to sexually immature adults or supernumerary nymphs).

A variety of molecules exist which control invertebrate pests by preventing successful completion of the insect’s life cycle, rather than being acutely toxic to the insect. Examples of such molecules include chitin synthesis inhibitors (CSI) and juvenile hormone analogues (JHα). The CSI act by disrupting the deposition of chitin during the formation of the insect’s larval cuticle after moult, whereas JHα aim to interfere with the hormone based control of metamorphosis and reproduction. These two types of molecules are often referred to as insect growth regulators (IGR) to distinguish them from conventional insecticides with neurotoxic action.

Consequently molecules that affect the developmental cycle of insects may be effective without resulting in the immediate death of the insect and therefore efficacy trials should be designed to address the most appropriate life cycle stage of the insect sensitive to the molecule of interest and also to measure any long term effects (e.g. on the fertility and fecundity of females or any effects on the embryonic development in the egg stage).
For example, in measuring the effectiveness of JHa, trials should be designed to record the number of adults produced from treated nymphs/larvae, the number of adults with deformed wings or terminalia and the mortality of insects prior to and at metamorphosis. Additionally a number of newly moulted females should be selected randomly from each treatment dose/formulation and their ability to produce viable eggs/oothecae after pairing with untreated males should be recorded.

IRAC, the Insecticide Resistance Action Committee, has developed a classification of insecticides based on mode of action (www.irac-online.org).

5.6.4.1.3.14 Resistance

Information on resistance and the likelihood of its development is required for BPR Annex I inclusion and is also important for product authorisation.

For insecticides resistance can be a problem. Some pests are more capable of building up resistance than others. For instance flies, with multiple generations and multiple females that can lay many eggs, resistance can be expected to build up easily. In ants on the other hand, with one or few queens who lay eggs for a long period, and a biocide that kills the whole colony most of the time, it is not to be expected that resistance will build up. Therefore, a resistance management strategy has to be provided for flies but not for ants for evaluation at product authorisation.

A resistance management strategy is generally based on the use of two modifiers, the frequency of use and the rotation with other active substances. For instance, for products against house flies, a label could state that the product should not be used more than five times per year and should only be used in rotation with at least one other product with a different mode of action.

For consumer products it is necessary to make clear that there might be a risk of building up resistance and that this can be reduced. Since consumers have no knowledge of resistance the label claim should contain information to prevent it. For instance, the following sentence could be added to the label: “When the product is not used according to the label resistance of insects might occur. When the infestation persists contact a professional.”

More information on resistance can be found in Chapter 6.2 of this TNG on Product Evaluation and the Insecticide Resistance Action Committee (IRAC: www.irac-online.org).

5.6.4.1.4 Methodology of assessment

Methods of application and dose rates

When considering the overall evaluation of a proposed label claim competent authorities should ensure that the data presented are relevant not only to biological challenge and treatment environment but also that the method of application and application/dose rate(s) used in the test(s) are appropriate to the label claims and proposed use of the product.

The application technique should therefore reflect the claims proposed on the label, whether crack and crevice, spot, space spray, contact spray or total release.

General considerations

The efficacy data submitted should demonstrate that the biocidal product, when used as directed by the product label, will result in a measurable beneficial effect. The data supplied should demonstrate that an acceptable, consistent level and duration of control or other intended effect will result from the use of the product at the recommended dose rate.
This may, depending on the individual product, be measured as a reduction of the pest population to an acceptable level or a reduction in damage. The acceptable level may vary depending on the purpose of the proposed use.

Competent authorities should evaluate available data to determine whether they are sufficient to support a label claim.

The competent authority will examine the submitted data package and a judgment will be made as to whether any data omissions are considered significant as to delay assessment. Those so identified will be communicated back to the applicant. The applicant can then supply additional data or modify their label claims in line with whatever has been supported.

Any known limitations on efficacy (including resistance) should be considered during the assessment.

- possible restrictions or recommendations concerning the use of the product in specific environmental or other conditions. State possible factors that can reduce the efficacy, for instance hot, cold or humid environments or the presence of other substances, in addition to the grounds for these. Possible recommendations concerning the avoidance of the continuous use of the product in order to prevent the development of resistant strains and the grounds for these (see also TNsG on product authorisation Chapter 6.2). State if the product cannot be mixed with, for example, other biocidal products or if the use of the product with other biocidal products is recommended;

- the guidance given on resistance for the corresponding data requirement of the active substance also applies here.

5.6.4.1.4.1 Assessment of specific claims

Sometimes a claim will include specific properties of the product, for instance:

- kills within 15 minutes;
- residual effect up to 3 months;
- storage period up to 5 years;
- control of tropical ants.

Where a particular property is claimed the data submitted to support the product should show that the product actually has these properties. If data do not support this claim, the product may still gain authorisation with amended label claims, provided that the product still shows acceptable efficacy.

For example: If a product claims complete control of ants within 2 weeks of application, the data submitted must show a high level of mortality (approximately 100%) within two weeks of application in order for these claims to be acceptable.

However, if the submitted data showed 90% mortality within 2 weeks and 100% mortality within 3 weeks, the product may still gain authorisation provided that the product claims were amended to ‘complete control of ants within 3 weeks of application’.

Situations such as the example above will require each study to be evaluated on its own merits, taking into account what the data is actually showing. Evaluators must use scientific judgement to determine when authorisation would not be acceptable.

For example:

If a product claims to kill ants within 15 minutes of application, the data submitted must show sufficient mortality within 15 minutes of application in order for these claims to be acceptable.

However, if the submitted data showed 50% mortality within 15 minutes but 90% mortality within 2 hours, the product would still not be granted authorisation on the
basis that for claims such as ‘kills ants’, the average user would expect a rapid visual effect following application (unless the product label clearly states how long the product takes to have an effect).

5.6.4.1.5 Assessment of authorisation

When considering the overall evaluation of proposed label claims, competent authorities should ensure that the data and the method of application and application/dose rates used in the tests are appropriate to the label claims and proposed use of the product.

5.6.4.1.5.1 Norms and criteria

The test results are compared directly with the norms and criteria for efficacy described below per insect/arthropod pest. The performance criteria set in this guidance ask for high levels of efficacy, which is of course what we aim for. However, some products that do not fully meet the criteria can still be valuable in some cases.

When a product does not perform to the criteria it should be justified in the application why this product is still recommended for authorisation. For example, in a field trial the criteria may not be met because of immigration of insects from untreated areas (e.g. flies, mosquitoes). When this is explained well in a justification the product might still be accepted for authorisation, depending on the results of other field trials, simulated use and laboratory trials.

Special attention should be paid to resistance, since under low insecticide pressure resistance can build up more easily. Moreover, it should be taken care of that no placebo’s or misleading products are registered. If the efficacy level is significantly lower than the criteria state it should be mentioned on the label.

The justification will be evaluated case by case. The product should not be authorised, unless there is a good reason for having a product of lower effectiveness.

5.6.4.1.5.2 Assessment

The assessor/expert assesses on the basis of the label claim and the above criteria. If the product was assessed to be sufficiently effective in laboratory and/or field tests, it will be authorised as far as efficacy is concerned.

5.6.4.2 General Claims: Crawling Insects, Flying Insects, Acaricide

5.6.4.2.1 Introduction

Some products have a very broad claim: against crawling insects, against flying insects, insecticide-acaricide spray, etc. In these cases it is not possible to test the product against all claimed target pests. For each group claimed tests should be performed on a few relevant species, of significant importance, and on the species specifically claimed on the label.

General claims (e.g. insecticide, crawling insects) cannot be used for bait products, since the bait differs per insect species.

5.6.4.2.1.1 Crawling insects

A crawling insect is defined as an insect that generally moves on the ground. These include amongst others cockroaches, ants, fleas, crickets, silver fish, bed bugs and carpet beetle larvae. The effect of biocides on these insects is primarily based upon contact. The products involved can be sprays, dusts, etc. Amongst the crawling insects, cockroaches are the most difficult to control.
5.6.4.2.1.2 Flying insects
A flying insect is defined as an insect that generally flies from one spot to the other. These include flies, mosquitoes, wasps and moths. The products involved can be sprays, strips, paints, etc.

5.6.4.2.1.3 Insecticide, acaricide and other arthropods
A general claim for insecticides includes all insects. A general claim for acaricides includes ticks and mites. Other arthropods could include spiders (Araneae), harvestmen (Opiliones), centipedes (Chilopoda), millipedes (Diplopoda), woodlice (Isopoda) and scorpions (Scorpiones).

5.6.4.2.2 Dossier requirements
A clear label claim should be submitted. The study results of trials should demonstrate the efficacy of the product based on the submitted label claim. Laboratory, simulated-use tests and field trials with the test organisms are needed to assess the efficacy of the product. Ideally, the studies should be performed according to established guidelines where these are available. These may be international, EU or national guidelines. Ideally, data should be generated using national or international recognised testing methods (ISO, CEN, OECD, etc.) where available and appropriate. See Appendix 17 for a list of available guidelines. If there are no guidelines available or guidelines are not suitable, the applicant may use their own methods (intra-company Standard Operating Procedures), on condition however, that the study is scientifically robust, well reported, provides a clear answer to the question and demonstrates the efficacy claimed. In addition, the test methods applied and the test conditions should be clearly and fully described and must address the efficacy claim that appears on the product label. A control treatment without biocide (negative control) should be included in all laboratory trials.

In the case of field trials where true replication is almost certainly impossible to achieve and where normal control methods are not restricted to use of a single biocidal product, a full description of any factors that might be expected to influence product performance should be given. These may include the risk of re-invasion from adjacent areas, general levels of sanitation, treatment history, season, etc. and are intended to provide the authorities with information to assist with the interpretation of the results obtained.

5.6.4.2.2.1 Test species
Claim: crawling insects. In case of an application for authorisation of a product with a claim of “killing crawling insects” a product, which has demonstrated sufficient effectiveness against cockroaches, may also be authorised to control other crawling insects. However, if also population control and/or nest kill is claimed both cockroaches and ants have to be tested.

Tests with cockroaches should normally be performed with two key species, one small, one large, such as the German cockroach (Blattella germanica) and either the oriental cockroach (Blatta orientalis) or the American cockroach (Periplaneta americana). Tests with ants should normally be performed with the Black garden ant (Lasius niger).

Claim: flying insects. In case of an application for authorisation of a product with a claim of “killing flying insects” tests should be provided with flies, mosquitoes and wasps. Tests with flies should normally be performed with the house fly, Musca domestica. Tests with mosquitoes should normally be performed with Culex spp. Test with wasps should normally be performed with Vespula spp.

Claim: acaricide. If a product is claimed to be an acaricide tests should be provided with mites and ticks. What species should be used depends on the area of use (house dust mites in homes, flour mites in storage rooms, etc., for instance: Dermatophagoides
pteronyssinus, *Tyrophagus putrescentiae*, *Acarus siro*). For mites and ticks relevant species can be found in sections 7 and 8.

**Claim: other arthropods.** For this claim the applicant should provide information on what organisms are relevant for the intended use. At least some example should be given and these should be tested.

**Specific claim next to general claim:**
Whenever efficacy against a specific organism is claimed next to a general claim or as specification of a general claim (e.g. crawling insects, including bedbugs), tests against this organism should be provided.

**5.6.4.2.2 Laboratory tests and field trials**
Test requirements for each test species can be found at the following sections dedicated to these insects/acomids. For other arthropods a field trial should be provided or a good justification why this is not appropriate.

**5.6.4.2.3 Assessment of authorisation**

**5.6.4.2.3.1 Norms and criteria**
A biocidal product may only be authorised if it “possesses a sufficient level of efficacy”. For products with general claims the performance criteria per tested organism are the same as those for products with a specific claim for the test species. I.e. for crawling insects the criteria are the same as for cockroaches and ants, for flying insect the same as flies, mosquitoes and wasps, etc. The criteria can be found in the sections dedicated to these insects/acomids.

**5.6.4.3 Cockroaches**

**5.6.4.3.1 Introduction**
Cockroaches are a common and persistent problem in many households. These crawling insects (although several species can also fly) are scavengers allowing them to readily adapt to changing food availability. Cockroaches can carry bacteria such as Salmonella in areas co-inhabited by humans. Cockroaches are also identified as a major cause of allergies and asthma, particularly in children. Amongst the crawling insects, cockroaches are the most difficult to control.

The effect of biocides on these insects is mainly based on either contact, both dermal and tarsal, or the ingestion of bait products.

**5.6.4.3.1.1 Biology**
Cockroaches belong to the (sub) order Blattodea. There are over 3500 species of cockroaches, but only a few are considered domestic pests in the EU. The German cockroach, *Blattella germanica*, is the most common.

Upon hatching from an egg capsule, cockroaches begin their nymphal stage (smaller version of adults minus fully developed wings and sex reproduction organs) and moult through various instars until reaching the adult stage. Time of development can take weeks or months depending upon the species and the surrounding environmental conditions. For instance the eggs of German cockroaches hatch after 3 to 5 weeks (depending on the temperature), the nymphal stage (5 to 7 moultings) can be 40 days to 6 months and the adults live about 6 month (longer under lab conditions).

In temperate European countries most cockroach species will almost never be found outside, with foraging activities almost entirely within human-made structures.

**5.6.4.3.2 Dossier requirements**
A clear label with comprehensive claims should be submitted. The study results of trials should demonstrate the efficacy of the product based on the submitted label claim.
Requirements can differ for products for professional use and for consumer products. For professional use a field trial is always required, for consumer products in some cases laboratory and simulated-use tests are sufficient. If the product is applied as a bait, the entire bait (formulated, including the bait box if applicable) should be tested, not only the active substance which is contained in the bait.

Ideally, the studies should be performed according to established guidelines where these are available. These may be international, EU or national guidelines. See Appendix 17 for a list of available guidelines. Appendix 18 gives an example of a test guideline that can be used. If the available guidelines are not suitable, industry standard or a company’s own protocols are acceptable, on condition however, that the study is scientifically robust, well reported and provides a clear answer to the question. In addition, the test methods applied and the test conditions should be clearly and fully described and must address the efficacy claim that appears on the product label. A control treatment without biocide (negative control) should be included in all laboratory trials.

In the case of field trials where true replication is almost certainly impossible to achieve and where normal control methods are not restricted to use of a single insecticidal product, a full description of any factor that might be expected to influence product performance should be given. These may include the risk of re-invasion from adjacent areas, pest activity before the trial is initiated, general levels of sanitation, treatment history, season, etc. and are intended to provide the authorities with information to assist with the interpretation of the results obtained.

5.6.4.3.2.1 Test species

For use against cockroaches data against two key species, one small species normally German cockroach (*Blattella germanica*) and one large species either the Oriental cockroach (*Blatta orientalis*) or the American cockroach (*P. americana*), should normally be available for spray products (aerosol, space spray, residual spray) to support general claims against cockroaches. For bait products, the label can only claim efficacy against species that have been treated under field conditions.

5.6.4.3.2.2 Laboratory tests and field trials

For the evaluation of biocides against cockroaches different types of laboratory, simulated-use tests and field tests can be used. Examples of test are listed below.

**Screening Studies (No-Choice Test)**

The product is applied to representative surfaces or via direct cuticle application, in an arena with cockroaches, to assess inherent contact toxicity or knockdown effects of the active substance. Specify whether adults (male or female) or nymphs are used. Tests may be used to demonstrate basic efficacy or efficacy against insects, resistance to specific chemicals (LD50 versus a susceptible field strain) or insect growth regulator effects (nymphs are treated and subsequent effects are recorded such as inhibition of moulting, deformities, sterile adults).

Results support descriptions related to the mode of action (symptomology) or “effective against strains resistant to “x” class of insecticides”, or similar efficacy claims.

For bait products dietary bioassay studies can be conducted using the biocidal bait as a food source. Replicate groups of test insects are exposed to either a continuous toxic diet, or a toxic diet for 24 hours and then a non-toxic diet for the rest of test period.

In all laboratory studies a treatment without biocide should be conducted as a negative control, with insects from the same insect population and with the same number of replicates.

Screening tests are not always necessary. When efficacy is demonstrated in residual tests, palatability tests or similar tests, this is deemed sufficient. Screening tests can
sometimes be used as bridging studies: if tests involving a product result in similar effects in different target species, field studies can be waived for some insect species.

**Determination of residual efficacy**

Formulated product (spray, powder, dust, etc.) is applied to representative surfaces at a specified dose rate, or rates, including the recommended label rate(s). Cockroaches (adults) are exposed to the deposit at several time intervals after application (including the day of treatment and at the end of the claimed residual period). Exposure time should, preferably, be comparable to the time the cockroaches might reasonably be expected to be in contact with a treated surface under natural conditions (e.g. 10 min - 1 hour) and assessors will take this factor into consideration when evaluating the data. Treated surfaces should include at least one porous and one non-porous substrate (or according to the label claim) representing surfaces that might, typically, be treated for cockroach control (e.g. ceramic tile, plywood, painted plywood, stainless steel, concrete). Mortality is normally assessed 1 day and up to 7 days post-exposure.

To substantiate a *knockdown claim* the number of cockroaches on their backs is counted at stated times after exposure (typically at 5 minute intervals until +30 min, then again at 45 and 60 min). The time until 50% (KT50) and 95% (KT95) of the insects are knocked down is derived statistically.

For *insect growth regulators*, exposure conditions can be as described above, but selection of the developmental stage (nymph, adult) and post-exposure assessment (deformities, moulting success, sterility, mortality) must be adapted to suit the mode of action of the active substance. Hence, assessments may continue to be made several weeks after exposure (sub-lethal or non-lethal effects on fertility, sterility for example may contribute to long term population control without short term mortality).

Groups of cockroaches of the target species should be of specified age/sex and number. Normally tests are performed with 5 or more replicates, with at least 10 cockroaches per replicate. When only 3 replicates are used, at least 20 insects per replicate should be used. Replicates should be conducted per applied dose, time point, surface, and a reference product (at registered rate) and untreated surfaces should be included as negative controls.

Environmental conditions must be specified for the test itself, and during storage of the treated substrates (temperature, humidity, photoperiod). Temperature would be expected to fall in the range 19-29°C. When efficacy at high temperatures is claimed 40°C would be a good test temperature.

**Palatability tests with bait products**

The aim of the bait choice feeding trials is to determine the palatability of the product for the test insect. If conducted on both fresh and aged product it may provide information on the storage stability of the product. In this test design, nymphs and adults of German and Oriental cockroaches have the choice between a non-toxic food source (challenge diet, either the non-toxic bait or a non-toxic food source known to be a strong feeding source for the test species) and the bait containing the active substance. Normally tests are performed with 5 or more replicate tests, with at least 10 cockroaches per replicate. When only 3 replicates are performed, at least 20 insects per replicate should be used. In all laboratory studies a treatment without biocide should be conducted with insects from the same insect population, as a negative control.

The test should demonstrate acceptable toxicity in competition with the alternative food source.

The population composition (males, gravid non-gravid females, nymphs) in these tests is of importance. Preferably mature insects should be used since immature stages do not need to feed every 24 hours. It should be noted that the feeding behaviour of German
cockroach females, changes during ‘pregnancy’ and that early instar nymphs tend to forage less than older instars.

**Simulated use**

These tests are designed to mimic the practical use situation. The insects must have a choice to be in contact with the biocide or not. For example, cockroaches (*B. orientalis*, *B. germanica*) can be introduced into choice boxes with one half of the base surface being sprayed with a test formulation. Food and water is always on the non-treated area to be reached by the animals without crossing the treated area. Variations on this test would be to expose insects (voluntary contact) to a variety of different treated surfaces, e.g. plywood, cement, vinyl, ceramic tiles, glass etc.

For products claiming “population control” (eradicates cockroach population) an entire population or at least different life stages should be tested while there is a possibility that only a few individuals get in contact with the biocide.

For “secondary kill” (kills cockroaches that do not visit the bait, however, not always the whole population) claims at least different life stages should normally be tested where only a few individuals get in contact with the biocide directly. Life stage is dependent on a specific mode of action (necrophagy versus coprophagy) and the claim. Either nymphs or adults could be used.

In all laboratory studies a treatment without biocide should be conducted with insects from the same insect population, as a negative control.

**Field trial**

In field trials the product is tested in actual use situation, for instance in an infested home or warehouse and applied according to the direction for use on the label. An example of the results to be achieved in a field trial can be found in Appendix 18.

5.6.4.3.2.3 Requirements per type of claim

Per type of claim the requirements will be listed.

**Products intended for use as general surface treatment or aerosol for consumers:**

- a laboratory test showing mortality and/or knockdown and/or residual efficacy, depending on the claim;
- a simulated-use test showing mortality and knockdown according to the claim.

**Products intended for use as general surface treatment or aerosol for professionals:**

- a laboratory test showing mortality and/or knockdown and/or residual efficacy, depending on the claim;
- a simulated-use test showing mortality and knockdown according to the claim;
- a field trial according to the directions for use.

**Products intended for use as general surface treatment or aerosol with a claim of population control or secondary kill:**

- a laboratory test showing residual efficacy;
- a simulated-use test showing mortality according to the claim;
- a field trial according to the directions for use.

**Products intended for use as baits:**

- due to the specificity of baits, only effects against species of cockroach that have been tested in the field can be claimed on the product label;
- a laboratory test showing palatability, of fresh product and product at the end of the claimed maximum storage period;
- a simulated-use test showing mortality according to the claim;
• a field trial according to the directions for use and with the claimed cockroach species.

Simulated-use tests can be waived if a robust field trial is submitted

5.6.4.3.3 Assessment of authorisation

5.6.4.3.3.1 Norms and criteria

A biocidal product may only be authorised if it “possesses a sufficient level of efficacy” (BPD). This is implemented in the following way.

An insecticidal product intended for the control of cockroaches is normally considered to be sufficiently “effective” if the following results can be achieved:

**Products intended for use as general surface treatment or aerosol for consumers:**

- required results in laboratory tests and simulated-use tests:
  - \( \geq 90\% \) knockdown within a few minutes after contact with the product (or according to the claim), direct after spray and at the end of the residual period claimed;
  - mortality according to the label claim, preferably \( \geq 90\% \) in 24 hour.

**Products intended for use as general surface treatment or aerosol for professionals:**

- required results in laboratory tests:
  - direct application: \( 100\% \) mortality within 1 hour after spraying the cockroaches, mortality between 90 and 100\% can be accepted provided a qualified explanation is given for the lack of total control;
  - residual test: \( 100\% \) mortality within 24 hours after placing the cockroaches in the test area, direct after spray and at the end of the claimed residual period. Mortality between 90 and 100\% can be accepted provided a qualified explanation is given for the lack of total control.

- required results in field tests:
  - after a period of 2-10 weeks, the population reduction exceeds \( \geq 90\% \) relative to either untreated sites or pre-treatment levels. If retreatment is necessary 100\% mortality should then be achieved.

**Products intended for use as general surface treatment or aerosol with a claim of population control or secondary kill:**

- required results in laboratory tests and simulated-use tests:
  - \( \geq 90\% \) mortality within the test period, direct after spray and at the end of the residual period claimed;

**Products intended for use as baits:**

- required results in laboratory palatability choice test (bait and alternative food):
  - at least 95\% of the test insects have been killed at a given time point;

- required results in simulated-use tests:
  - \( \geq 90\% \) reduction of the population within a few weeks;

- required results in field tests:
  - after a period of 2-10 weeks, the population reduction exceeds 80\% relative to either untreated sites or pre-treatment levels.
Products based in insect growth regulators (IGR):
- required results in laboratory tests:
  - at least 95% of the insects does not develop to the next instar;
- required results in simulated-use tests:
  - ≥ 90% reduction of the population within a few weeks;
- required results in field tests:
  - after a period of 6 -14 weeks, the population reduction exceeds 80% relative to either untreated sites or pre-treatment levels.

Deviation from these norms is possible but should be justified in the application.

Field trial data at the label application rate(s) must preferably be evaluated by an experienced assessor since performance can vary considerably, even from apartment to apartment in the same building. Number of trials, the complexity of the trials sites, the use (or not) of additional measures that can contribute to effective control, treatment history, etc. can all have a substantial effect upon the level of control that is achieved. The data must provide evidence of suitable levels of efficacy during the residual period claimed, relative to pre-treatment population assessments and/or performance of reference products under similar conditions, and/or assessments of cockroach populations in untreated areas under similar conditions. Where mean population reduction exceeds 90% relative to either untreated sites or pre-treatment levels, the product is considered effective, but the assessor has the discretion to view each data set on its merits and consider all factors before concluding whether the data support the claimed level of performance or not.

5.6.4.4 Ants

5.6.4.4.1 Introduction

Ants may cause inconvenience both indoors and outdoors.

In Europe the following ant species are common:

- Black garden ant, Lasius spp, most common L. niger
- Pavement ant, Tetramorium caespitum
- Red ant, Myrmica rubra
- Erratic ant, Tapinoma erraticum.

Next to these native ant species tropical ants can cause inconvenience, mainly indoors.

Of the tropical ant species there are two species that are most commonly found causing inconvenience in buildings in Europe:

- Pharaoh ant, Monomorium pharaonis
- Argentine ant, Linepithema humile.

5.6.4.4.1.1 Biology

Ant development involves a complete metamorphosis that includes distinct egg, larval, pupal and adult stages. Most ant species form colonies comprised of complicated social structures that include infertile female workers, one or more specialised fertile queens and (at certain stages in nest development) sexually mature males. Some species have developed additional specialised workers that are responsible for guarding the nest and attacking intruders, whilst others perform domestic and foraging duties. These workers will actively forage on a wide range of foods including sweet substances, seeds, insects and aphid secretions. A successful foraging ant also has the ability to communicate where to find food to her co-workers, using chemical signals (trail pheromones).
5.6.4.4.2 Dossier requirements

A clear label claim should be submitted. The study results of field trials should demonstrate the efficacy of the product, based on the submitted label claim.

Requirements can differ for products for professional use and for consumer products. For professional use products a field trial is always required, while laboratory and simulated use tests might be considered sufficient in some cases for consumer products. Requirements also depend on the use: for “nest kill” and bait products alike, both laboratory and field trials with the test insects are needed; for products that only claim to kill individual insects that are in contact with the biocide, laboratory and simulated-use tests are sufficient. If the product is applied as a bait, the entire bait (formulated, including the bait box if applicable) should be tested, not only the active substance which is contained in the bait.

Ideally, the studies should be performed according to established guidelines where these are available. These may be international, EU or national guidelines. See Appendix 17 for a list of available guidelines.

If there are no guidelines available or the guidelines are not suitable, the applicant may use their own methods (intra-company Standard Operating Procedures), on condition however, that the study is scientifically robust, well reported and provides a clear answer to the question. In addition, the test methods applied and the test conditions should be clearly and fully described and must address the efficacy claim that appears on the product label. In the case of field trials where true replication is almost certainly impossible to achieve and where normal control methods are not restricted to the use of a single insecticidal product, a full description of any factors that might be expected to influence product performance should be given. These may include the risk of re-invasion from adjacent areas, general levels of sanitation, treatment history etc. and are intended to provide the authorities with information to assist with the interpretation of the results obtained.

5.6.4.4.2.1 Test species

Table 26 below shows for this group of insecticides the possible combinations of target organisms, and the corresponding test organisms on which efficacy is tested in both laboratory and field tests. The selection of test species should be relevant to the label claim.

Table 26: Target organisms versus test organisms

<table>
<thead>
<tr>
<th>Target organisms of the insecticide:</th>
<th>Test organisms:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ants</td>
<td>Garden ant (<em>Lasius niger</em>)</td>
</tr>
<tr>
<td>Tropical ants</td>
<td>Pharaoh ant (<em>Monomorium pharaonis</em>), Argentine ant (<em>Linepithema humile</em>)</td>
</tr>
</tbody>
</table>

5.6.4.4.2.2 Laboratory tests and field trials

For the evaluation of biocides against ants different types of laboratory, simulated-use tests and field tests can be used.

**Screening studies for direct spray or general surface treatments**

In all laboratory studies a treatment without biocide should be conducted with insects from the same insect population, as a negative control. Examples of tests are:

Direct spray: 20 ants placed within a Petri dish and directly sprayed with material. Knockdown, time to death and total mortality is recorded. For insecticides with a “nest kill” claim the time to death will be longer (>1 day) since these ants have to live long
enough to take the insecticide into the nest. Normally at least 5 replications and 5 non-treated controls should be used. Controls are very important in this case, as it often turns out to be very difficult to keep ants active in trials.

**Residual spray:** 20 ants placed on a surface treated with the product. Ants are placed in the arena directly after application, at several time intervals after application and also at the end of the period claimed for residual effect. The time to death of the ants and total mortality is recorded.

A control treatment without biocide should be included in all laboratory trials. Normally at least 5 replications and 5 non-treated controls should be used.

**Palatability tests with bait products**

The important factors relating to testing bait products are to establish the appropriate dosage and intrinsic palatability of the formulation in laboratory tests. Claims made for bait products should distinguish between ants and tropical ants, since the latter can be attracted by completely different baits than the more common European ant, *L. niger*. Data should be provided for all species, for which claims are made.

The most important factor involved in laboratory testing is to provide a free choice alternative food source to the test insects. This may be sugar-based materials for European ants and protein-based materials (meat, eggs, dead insects) for some tropical ants. The formulation should demonstrate acceptable toxicity in competition with the alternative food source. A control treatment without biocide of similar size as the test itself (i.e. number of replications) should be included in all laboratory trials.

When a product is claimed to be effective after a long period of storage, it is also necessary to demonstrate that the product will still be effective, and attractive, after the stated storage period. The applicant must either provide data on the palatability of the product at the end of maximum storage period or alternatively (in case of a new product) data gained in a stress test with 'accelerated ageing', i.e. a palatability test with the product which is stored under challenging conditions.

**Simulated use studies**

These tests are designed to mimic the practical use situation. The tests should be relevant to the use and label claims. A control treatment without biocide should be included in all laboratory tests. Control trials should be of similar size (i.e. number of replications) as the test itself, to make statistical comparison possible and to get a fair impression of control mortality.

Examples of tests are:

**Direct general surface treatments without nest kill:**
Ants (normally at least 20 worker ants) can be introduced into choice boxes/arenas with one half of the base surface being sprayed with a test formulation, at the correct application rate according to the product label. Food and water is always on the non-treated area to be reached by the animals without crossing the treated area. Variations on this test would be to expose insects (voluntary contact) to a variety of different treated surfaces, e.g. plywood, cement, vinyl, ceramic tiles, glass etc. Mortality is recorded.

Normally tests should be performed in triplicate.

**Direct general surface treatments with nest kill:**
In a double chamber trial an ant’s nest (normally at least 20 (worker) ants) is placed within one arena, which is connected to another arena. Part of the second arena is treated with the insecticide at the correct application rate according to the product label. Adequate food and water is placed on the non-treated surface of this second arena. Ants must be able to reach the food without contacting the treated surface. Normally tests
should be performed in triplicate. Efficacy is assessed e.g. length of time taken to result in control of the ant population (e.g. no foraging ants).

The nest should be opened at the end of the trial (e.g. 1 week), to check whether all ants within the nest are dead, especially the queen(s).

**Bait products:**
The efficacy of the entire formulated bait is tested, hence not only the active component within the bait. An ant's nest is placed within an arena trial under controlled conditions (e.g. with respect to temperature, relative humidity, photoperiod, etc.). Adequate food (bait without the active substance or and alternative food source) and water are placed opposite the nest. Insects are allowed to acclimatise for 7 days before introduction of bait. An additional fasting period of 4 days, providing them with water only, is recommended. At regular time intervals (in hours), the attractiveness of the bait for the ants is recorded (by observing whether they approach the bait or avoid it). Ant mortality is recorded at regular time intervals (in days). At the end of the trial the nest could be opened to check whether all ants within the nest, including the queen(s), are dead.

**Field trials for all claims**
The tests should be relevant to the use and label claims. Tests with *Lasius niger* are done preferably during the early spring. In the end of summer population decline might be due to natural causes instead of the insecticide. Non-treated nests should be used as a negative control, to test nest activity.

Monitor ant numbers at various locations around a building and locate the entrances of nests and “ant-trails” (routes taken by ants). Apply the insecticide according to the label instructions.

The efficacy tests against ants should normally be performed in a minimum of three objects. An object can be a place in or near the house, where ants cause inconvenience for the inhabitants. This may be in a house, on a balcony, a terrace or in a garden, depending on the field of use of the product. If the test is performed outdoors, records of temperature and rainfall should be kept.

Monitoring should be conducted at the same locations (as the pre-treatment) and at similar times during the entire trial (e.g. at 12.30, 13.00, etc.). Monitoring should continue (e.g. 1 day after treatment, 1 week after treatment, etc. at least once weekly) until control is seen. If no ants are seen during a post-treatment monitoring visit then the site should be re-visited once to ensure that re-infestation does not occur.

The effect on the ant population can be determined by counting. For this purpose, a fixed position on the ‘ant-trail’ is to be used and a count of the number of any ants that pass is made in 1 minute, at several time intervals during the test.

### 5.6.4.4.2.3 Requirements per type of claim

**Per type of claim the requirements will be listed.**

**Products intended for use as general surface treatment for consumers:**
- a laboratory test showing mortality and/or knockdown and/or residual efficacy, depending on the claim;
- a simulated-use test showing mortality and knockdown according to the claim.

**Products intended for use as general surface treatment for professionals:**
- a laboratory test showing mortality and/or knockdown and/or residual efficacy, depending on the claim;
- a simulated-use test showing mortality and knockdown;
- a field trial according to the directions for use.
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Products intended for use as general surface treatment with a claim of nest kill:
- a laboratory test showing residual efficacy;
- a simulated-use test showing mortality;
- a field trial according to the directions for use.

Products intended for use as baits:
- Due to the specificity of baits, only effects against ant species that have been tested in the field can be claimed on the product label;
- a laboratory test showing palatability;
- a simulated-use test showing mortality;
- a field trial according to the directions for use and with the claimed ant species.

Simulated-use tests can be waived if a robust field trial is submitted.

5.6.4.4.3 Assessment of authorisation

5.6.4.4.3.1 Norms and criteria

A biocidal product may only be authorised if it “possesses a sufficient level of efficacy” (BPD). This is implemented for ants in the following way.

An insecticide against ants is normally considered to be sufficiently “effective” if the following results can be achieved:

Products intended for use as general surface treatment for consumers:
- required results in laboratory mortality tests and simulated-use tests:
  - ≥ 90% knockdown in 5 -10 minutes (or according to the claim), direct after spraying the ants and at the end of the residual period;
  - mortality according to the label claim, preferably ≥90% after 24 hour.

Products intended for use as general surface treatment for professionals:
- required results in laboratory tests:
  - direct application: 100% mortality within 24 hours after spraying the ants, mortality between 90 and 100% can be accepted provided a qualified explanation is given for the lack of total control;
  - residual tests: ≥ 90% mortality within 24 hours after placing the ants in the test area, direct after spray and at the end of the residual period;
- required results in field tests:
  - after a period of 2-8 weeks, the population reduction exceeds 90% relative to either untreated sites or pre-treatment levels.

Products intended for use as general surface treatment with a claim of nest kill:
- laboratory tests:
  - 100% mortality within the test period, direct after spray and at the end of the residual period;
- required results in simulated-use tests:
  - slow knockdown, ants must be able to reach the nest;
  - ≥ 90% mortality within the test period, including ants in the nest;
- required results in field tests:
  - after a period of 2-8 weeks, the population reduction 100% relative to either untreated sites or pre-treatment levels, in case of lower efficacy it has to be shown that the queen(s) in the test nests is killed.
Products intended for use as baits:

- required results in laboratory palatability choice test (bait and alternative food):
  - at least 95% of the test insects have been killed at a given time point;
- required results in simulated-use tests:
  - ≥ 90% reduction of the population within a few weeks;
- required results in field tests:
  - after a period of 2-4 weeks, the population reduction exceeds 90% relative to either untreated sites or pre-treatment levels.

Deviations from these norms is possible but should be justified in the application.

5.6.4.5 Termites

5.6.4.5.1 Introduction

Termites, in natural settings, work as beneficial insects by breaking down cellulose-containing materials, such as dead trees. However, termites can cause damage to living trees and many crop plants, but the fact that they can use dead wood makes them a major pest for timber used both outdoors and inside buildings. Termites become a problem to humans when they infest timber used in constructions (i.e. wood structures) in risk areas. Owing to their high moisture requirements, they usually nest in soils, but can invade buildings from underneath through cracks and seams or by building shelter tubes connecting the wood to their nest in the soil. In Europe and in the European tropical overseas regions, there are three main types of termites: subterranean, tree and drywood termites, the subterranean being the most destructive termites in construction. Due to their biological characteristics (subterranean termites), they live in the soil and must maintain contact with the ground or some other moisture source to survive.

Insecticides against termites can be divided into PT8 products, preventive treatments to protect the wood and curative treatments on the wood, and PT18 products, which are considered in this section.

5.6.4.5.1.1 Biology

Termites belong to the order of Isoptera. In Europe and in the European tropical overseas regions there are three main termite families; subterranean (Rhinotermitidae), drywood termites (Kalotermitidae) and tree termites (Nasutitermitidae).

*Reticulitermes* is the most common genus encountered from the Rhinotermitidae family in Europe. The main species registered are: *R. flavipes* (former *R. santonensis*), *R. lucifugus*, *R. lucifugus corsicus*, *R. grassei*, *R. banyulensis*, *R. balkanensis*.

They are widespread around the Mediterranean (Spain, France, Italy, Portugal, Balkans, and Greece) and Black Sea (Turkey, Rumania), though some termite spots in the UK and Germany have been reported. Several unanswered questions remain about the origin of these termites. While some Reticulitermes are native to Europe, others may be related to species from eastern North America and the Middle East (Israel, Asian Turkey, etc.).

*Coptotermes* sp. and *Heterotermes* sp. are the main two species belonging to the *Rhinotermitidae* family found in European tropical overseas regions.

*Nasutitermes* sp. are the main species belonging to the *Termitidae* family (tree termites) encountered in the European tropical overseas regions.

*Kalotermes flavicollis* and *Cryptotermes brevis* are the main two species of drywood termites present in Europe (especially in the coastal areas of Mediterranean countries and Canary Islands). *Cryptotermes* sp. is a main genus belonging to drywood termites encountered in the European tropical overseas regions.

A brief explanation of the life cycle (figure 11) may help to clarify the difficulties involved in control of termites. There is a split after the larval stages into two lines, the sexual
and the worker line. Individuals going down the sexual line develop into nymphs and then into either alates (which are the reproductive form most people are familiar with) or neotenics (supplementary reproductives). The alates do form queens (physogastrics), however, these are much more mobile than those found in tropical species. The alternative line of development, the neutral line, is the development of larvae into workers, which in turn can either remain workers or develop into neotenics or soldiers. Workers are approximately 4 to 6 mm in length. An important feature in the biology of termites that makes them very difficult to control is the ability of individuals in both lines to form sexual reproductives and, hence, give rise to a new, viable colony. In addition, supplementary reproductives can be produced in very large numbers.

Figure 11: Life cycle of subterranean termites

5.6.4.5.1.2 Control methods

Preventive treatments

Traditionally, the methods used to fight termites were based upon treating infested or exposed wood with wood preservatives. This is valid for all termite types (subterranean, tree and drywood). Those products are included in product type 8 (wood preservatives) of the BPD, and are not considered in this section.

In addition to the preventive treatment of timber, a barrier can be used to isolate the paths used by subterranean termites to access the building from underneath where the nest is located. Barriers systems usually consist of a polymer membrane or other material and an insecticide (product type 18). The system is installed between the soil and the construction to keep subterranean termites outside and to eliminate those that come into contact with the insecticide.

Remedial treatments

Different methods are currently used in Europe:

Chemical barriers

Methods based on treating the infested wood with wood preservatives are included in product type 8 (wood preservative) of the BPD, and are not considered in this section.

In addition to the wood treatment, two types of chemical barriers are used to impregnate the walls of the construction and the soil around.

Considering the subterranean termites, this method aims to eliminate insects inside the construction and to protect it for several years. This method does not eliminate the nest (which is located in the soil).
Bait system
It consists typically of a cellulose-based matrix treated with a slow acting insecticide, which is consumed by workers and is spread through the colony by trophallaxis (one individual is fed by another). Consequently, this method may be useful to eradicate the whole colony.

Treatment of waste
In order to prevent termite contamination by waste infested and transported into an area not infested, it could be relevant to treat the waste with biocidal products.

5.6.4.5.2 Dossier requirements
A clear label claim should be submitted.

Laboratory and field trials with termites are needed to assess the efficacy of the products. Ideally, the studies should be performed according to established guidelines where these are available. These may be EU or national guidelines. European standardisation work is being conducted by several termite experts in Europe. At this moment, no European standard has been published yet, only French standards are available. However, due to the greater significance of termites as structural pests in countries outside Europe, such as the United States and Australia, a variety of standard test methods are published, together with extensive reports in the scientific literature which may prove useful references. Account should be taken of results obtained using such methods, especially where the same termite species are present as those in Europe including the French overseas territories. See Appendix 17 for a list of available guidelines (guidelines outside EU not included yet).

If there are no guidelines available or guidelines are not suitable to evaluate the termiticide (e.g. if new products are developed), the applicant may use their own methods, on condition however, that the study is scientifically robust, well reported and provides a clear answer to the question. In addition, the test methods applied and the test conditions should be clearly and fully described and must address the efficacy claim that appears on the product label. In the case of field trials a full description of any factors that might be expected to influence product performance should be given. These may include the risk of re-invasion from adjacent areas, treatment history, etc. and are intended to provide the authorities with information to assist with the interpretation of the results obtained.

A control treatment without biocide should be included when testing any termite products in laboratory trials.

5.6.4.5.2.1 Test species
A product against termites in Europe should normally be tested on termites belonging to the genus Reticulitermes.

For European tropical overseas regions, the product should normally be tested at least against termites belonging to the genus Coptotermes and on every genus claimed by the applicant.

Remarks:

a) In any case, the termite species needs to be identified and all useful information about the colony collected (locality of origin, laboratory rearing conditions, characteristics of their natural environment if termites are collected in field);

b) For the evaluation of termite baits, the species referred to in the label claims should be used. If the claim refers generally to Reticulitermes species (without specifying the species), it is recommended to test, at least, two different European species in lab tests;
Due to the specificity of baits, only effects against species of termites that have been tested should be claimed on the product label.

5.6.4.5.2.2 Laboratory tests and field trials

The tests specified below are mainly for bait products. While laboratory tests can be conducted for all the termiticide products, field tests are addressed specially for bait products. For soil/wall barrier products and for physico-chemical systems the tests should be designed to mimic the practical use situation. The test should be performed according to the label claim.

Due to the specificity of baits, only effects against species of termites that have been tested should be claimed on the product label.

The important factors relating to testing bait products are to:

a) establish the appropriate dosage of the formulation in laboratory tests. This can be done in a mortality test (evaluation of the toxicity of the insecticidal formulation in a force-feed environment). The formulation should demonstrate acceptable toxicity;

b) test the palatability of the bait. The aim of the bait choice feeding trials is to determine the palatability of the product for the test insect. In this test design, insects have the choice between a non-poisoned food source (challenge diet) and the bait containing the active substance;

c) the test should demonstrate acceptable toxicity in competition with the alternative food source;

d) assess if a contaminated group of termites can transfer the insecticide to a group of termites that have never been exposed to it before. This transfer study should demonstrate acceptable toxicity of termites not exposed directly to the baits.

Laboratory/screening tests

No-choice test (A): test the termiticidal efficacy and the delayed effect of an insecticide formulation on a group of subterranean termites:

A group of termites is put into contact with an insecticide formulation. When testing baits, bait is the only source of food. For other types of termiticides the termites are exposed to the product according to the intended use (e.g. spray the surface and add the termites to the surface. The test is performed in assay containers. Mortality of the insects is assessed.

From this test the time “te” can be determined, necessary to perform the test B (te=time of exposure of the termites to the insecticide formulation which is required to observe a significant mortality compared with termites in an untreated control).

Transfer test (B): the transmission of the insecticide used in the baiting system to an uninfected group of termites:

Termites are exposed to the tested bait long enough to be contaminated with the active substance (time te). A group of termites is removed from the colony and put in contact with a healthy uncontaminated group. The mortality rate of both groups of termites (contaminated and uncontaminated) is assessed separately.

Choice test / palatability test (C): the suppression of a group of termites reared in laboratory under conditions of food competition; with the use of the same insecticidal bait formulation:

Add the insecticidal bait formulation to a group of termites already exploiting another source of food. The test is performed in assay containers. The aim is to assess the mortality after a given period of time.
Field trial

In field trials the product is tested in actual use situation and applied according to the direction for use on the label. The test method should evaluate the efficacy of the baits or barrier products in an experimental site where termite activity is reported.

The repellent termite barriers can be disposed in walls or soils, according to the claim. A common claim for a barrier product is the duration of “protection”. This is normally in terms of a number of years and should be demonstrated by long-duration soil tests in field plots.

For bait products consumption of the tested bait must be registered at least in the first 6 months after the introduction of the baits. The elimination of termites in the experimental site should be registered maximum after 18 months (counted since the introduction of the first tested bait), excluding the winter period.

Table 27 gives an overview of available (French) guidelines for termites and how to use them.

**Table 27: Overview guidelines on termites**

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<th>Preventive treatment / Physico-chemical barrier</th>
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<th>Ageing Test</th>
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<td>ENV 1250-2 (effect of water)</td>
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<td>Field test</td>
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<td>Field test</td>
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<td>Wall chemical barrier</td>
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5.6.4.5.3 Assessment of authorisation

5.6.4.5.3.1 Norms and criteria

A biocidal product may only be authorised if it “possesses a sufficient level of efficacy” (BPD). This is implemented in the following way.

An insecticidal product intended for the control of termites is normally considered to be sufficiently “effective” if the following results can be achieved (derived from standards NF XPX41-551, NF XPX41-543-3 and FCBA-BIO-E-041):

Products intended for use as baits:
- no-choice test: 100% mortality before the end of the test (16 weeks). Besides, if the 100% mortality is achieved too fast (less than 48 hours) the test bait should be rejected;
- transfer test: 100% mortality of all the termites, which have not been exposed directly with the tested bait;
- choice test / palatability test: more than 95% mortality;
- bait field test: No termite activity should be reported within the test period (max. 18 months, excluding the winter period). No termite activity should be reported in at least the following 3 months.

Products intended for use as termite barriers
- laboratory test: 100% mortality after the test (only for barriers with lethal activity);
- field test:
  - In soil barrier products, termites should not penetrate the soil more than 10 mm;
  - In wall barriers (i.e. thermoplastic films), termites should not be able to perforate the film after the duration of the test;
  - In other type of repellent barriers, termites should not be able to access to the other side of the barrier. Furthermore, any carrying of termite material (i.e. soil) to the other side of the barrier should not be reported.

5.6.4.6 Bed Bugs

5.6.4.6.1 Introduction

Bedbugs are small, wingless blood feeding insects. Of the many recognized species, only three are known to feed on humans. In temperate climate regions of the EU, *Cimex lectularius* is the dominant species. Bedbugs are not known to transmit disease in Europe, but infestations can cause painful and irritating bites on the skin while humans sleep. Once infested, treatment and control is very difficult.

A sign of bedbug presence include bites on the exposed skin (small red itchy bumps) of humans during sleep. If observed, confined locations such as mattress linings or furniture folds should be inspected for faecal spotting and the presence of bedbugs.

5.6.4.6.1.1 Biology

Bedbugs belong to the order of Hemiptera, Family Cimicidae.

Bedbugs harbour themselves in very confined areas in wall cracks, furniture joints, along lining of mattresses, behind pictures and in seams of furnishings. These insects generally confine themselves to these areas and leave them only to feed. Bedbugs are negatively phototactic and not usually seen outside the harbourage in the day or when the lights are on.
Female bedbugs can lay up to 500 eggs during their lifetime. Depending on frequency of blood meals, bedbugs can live for more than a year. They are able to survive for months without feeding (dependent upon temperature: at 16°C survival can be a year). The first nymph hatch from small white eggs after 7-10 days at room temperature (around 20°C) and earlier at higher temperatures. Each of the 5 nymphal stages need a blood meal to complete development to the next instar. The whole life-cycle from egg to egg takes a minimum of 28 days at 27°C or around 42 days at 22°C.

5.6.4.6.2 Dossier requirements

A clear label claim should be submitted. The study results of field trials should demonstrate the efficacy of the product, based on the submitted label claim.

Laboratory and field trials with bedbugs are needed to assess the efficacy of the product. Ideally, the studies should be performed according to established guidelines where these are available. These may be international, EU or national guidelines. See Appendix 17 for a list of available guidelines.

If there are no guidelines available or guidelines are not suitable, the applicant may use their own methods, on condition however, that the study is scientifically robust, well reported and provides a clear answer to the question. In addition, the test methods applied and the test conditions should be clearly and fully described and must address the efficacy claim that appears on the product label. In the case of field trials where true replication is almost certainly impossible to achieve and where normal control methods are not restricted to use of a single insecticidal product, a full description of any factors that might be expected to influence product performance should be given. These may include the risk of re-invasion from adjacent areas, general levels of sanitation, treatment history etc. and are intended to provide the authorities with information to assist with the interpretation of the results obtained.

5.6.4.6.2.1 Test species

A product against bedbugs should normally be tested on the common bedbug (Cimex lectularius) or tropical bedbug (Cimex hemipterus).

5.6.4.6.2.2 Laboratory tests and field trials

For the evaluation of biocides against bedbugs different types of laboratory, simulated-use tests and field test can be used. Examples of tests are listed below.

Screening studies (no-choice test)

Testing should include application of the product to representative surfaces (e.g. plywood, painted plywood, textile fabric, wallpaper) or direct cuticle application of the product to bedbugs to assess inherent contact toxicity of the active substance. It should be specified whether adults or nymphs are used. A test may be used to demonstrate basic efficacy or efficacy against insects resistant to specific chemicals (LD50 versus a susceptible field or laboratory strain) or insect growth regulator effects (nymphs are treated and subsequent effects are recorded such as inhibition of moulting, deformities, sterile adults).

Results must support description related to the mode of action (symptomology) or “effective against strains resistant to “x” class of insecticides”, or similar efficacy claims.

Screening tests are not always necessary. It is sufficient to demonstrated efficacy in residual tests or similar tests.

Determination of residual efficacy

Good residual efficacy is essential for insecticides used in bedbug control, as is impossible to treat all bedbugs directly or reach all of their hiding.
For the determination of residual efficacy, the formulated product (spray, powder, dust, etc.) should be applied to representative surfaces at the recommended label rate. Bedbugs (adults) should be exposed to the deposit at several time intervals after the deposit has dried (including the day of treatment, but after the deposit has dried completely and at the end of the claimed residual period). Exposure time should, preferably, be comparable to the time the bedbugs might reasonably be expected to be in contact with a treated surface under practical conditions (e.g. 10 min - 6 hours) and assessors will take this factor into consideration when evaluating the data. Treated surfaces should include at least two porous and one non-porous substrate, representing surfaces that might, typically, be treated for bedbug control (e.g. plywood, painted plywood, textile fabric, wallpaper, according to the label claim). Mortality is normally assessed after 1 day up to 14 days post-exposure.

For insect growth regulators, exposure conditions can be as described above, but selection of the developmental stage (nymph, adult) and post-exposure assessment (deformities, moulting success, sterility, mortality) must be adapted to suit the mode of action of the active substance. Hence, assessments may continue to be made several weeks after exposure (sub-lethal or non-lethal effects on fertility, sterility for example may contribute to long term population control without short term mortality).

Groups of bedbugs should be of specified age/sex and number. Tests should be performed in triplicate, with at least 20 bedbugs per replicate. When 5 or more replicates are used, 10 insects per replicate are adequate. Replicates should preferably be conducted per applied dose, time point, and surface. Untreated surfaces must be included as negative controls.

Environmental conditions must be specified for the test itself, and during storage of the treated substrates (temperature, humidity, photoperiod). Temperature would be expected to fall in the range 19-29°C. For use in Southern European countries higher temperatures (up to 40°C) might be necessary.

A control treatment without biocide should be included in all laboratory trials. The control trial should be of adequate size (i.e. number of replications and individuals), providing sufficient statistical power and a fair impression of control mortality.

**Simulated use**

These tests are designed to mimic the practical use situation. The insects must have a choice to be in contact with the biocide or not. Due to the normal behaviour of the bedbugs, it seems to be very difficult to design simulated-use tests for the evaluation of products for bedbug control. Bedbugs do not leave their harbourage during daytime and without a host which attracts them.

**Field trials**

In field trials the product is tested in actual use situations, for instance in an infested home or hotel and applied according to the direction for use on the label.

It has to be considered that in bedbug infestations the aim of professional control operations must be the eradication of the population. It is not acceptable to have even very small remaining populations. Usually, pest control operations against bedbugs have to combine different measures. The documentation of the trial has to give all information on the products or other measures used.

**5.6.4.6.2.3 Requirements per type of claim**

Appropriate efficacy tests are needed for each claim.

**Products intended for use as general surface treatment for consumers:**

- a laboratory test showing mortality and/or knockdown and/or residual efficacy, depending on the claim.
Products intended for use as general surface treatment for professionals:

- a laboratory test showing mortality and/or knockdown and/or residual efficacy, depending on the claim;
- a simulated-use test showing mortality and knockdown according to the claim and/or;
- a field trial according to the directions for use;
- Simulated-use tests can be waived if a robust field trial is submitted.

5.6.4.6.3 Assessment of authorisation

5.6.4.6.3.1 Norms and criteria

A biocidal product may only be authorised if it “possesses a sufficient level of efficacy” (BPD). This is implemented in the following way.

An insecticidal product intended for the control of bedbugs is considered to be sufficiently “effective” if the following results are achieved:

Products intended for use as general surface treatment for consumers:

- required results in laboratory tests (and simulated-use tests):
  - ≥ 90% knockdown within a few minutes after contact with the product (or according to the claim), direct after application and at the end of the residual period;
  - mortality according to the label claim, preferably ≥90% in 1 hour.

Products intended for use as general surface treatment for professionals:

- required results in laboratory tests:
  - direct application: 100% mortality within 24 hours after spraying the bedbugs;
  - residual test: ≥ 95% mortality within 24 hours after placing the bedbugs in the test area, direct after spray and at the end of the residual period.
- required results in field test:
  - after a period of 6-10 weeks, the population reduction exceeds 90% relative to either untreated sites or pre-treatment levels.

Treatment repeats usually are necessary in bedbug control. At the end of a treatment, 100% efficacy should be achieved.

Deviations from these norms is possible but should be justified in the application.

Data from field trials at the label application rate must preferably be evaluated by an experienced assessor since performance can vary considerably, even from apartment to apartment in the same building. The number of trials, the complexity of the trials sites, the use (or not) of additional measures that can contribute to effective control, treatment history etc. can all have a substantial effect upon the level of control that is achieved. The data must provide evidence of suitable levels of efficacy during the residual period claimed, relative to pre-treatment population assessments and/or performance of reference products under similar conditions, and/or assessments of bedbug populations in untreated areas under similar conditions. Where mean population reduction exceeds 90% relative to either untreated sites or pre-treatment levels, the product is considered effective, but the assessor has the discretion to view each data set on its merits and consider all the factors before concluding whether the data support the claimed level of performance.
5.6.4.7 Ticks

5.6.4.7.1 Introduction

Ticks are small arthropods classed along with mites and spiders in the Class Arachnida. All ticks are blood feeders. Certain tick species are known for carrying and transmitting many different pathogenic microorganisms including bacteria, viruses, parasites and fungi. Diseases associated with tick transmission in Europe include Lyme disease, tick-borne encephalitis, and human anaplasmosis, all transmitted by *Ixodes ricinus*. The tick *Hyalomma marginatum* can transmit Crimean-Congo haemorrhagic fever, a viral disease common in East and West Africa. Mediterranean spotted fever is transmitted by the brown dog tick (*Rhipicephalus sanguineus*). Ticks also have an important role in animal health. They can cause anaemia, reduction of milk production and bodyweight gain of animals.

5.6.4.7.1.1 Biology

Ticks differ from insects morphologically having two main body parts (insects have three) and eight legs as nymphs and adults (six legs for insects). Ticks go through four stages to complete their lifecycle: egg, larva, nymph, and adult. Feeding will occur in both the immature and adult stages. After mating female hard ticks will feed once more followed by oviposition of hundreds to even thousands of eggs.

Ticks can be differentiated on their host choices:
- one host: developing stages and adults feed on one host (e.g. *Boophilus*);
- two hosts: larvae and nymphs feed on the same host, adults feed on another host (e.g. *Rhipicephalus*);
- three hosts: larvae, nymphs and adults feed on three different hosts. (e.g. *Ixodes, Haemophysalis, Dermacentor*).

Ticks can be classified into two main families: soft ticks (*Argasidae*) and hard ticks (*Ixodidae*). The hard ticks consist of many commonly known species such as the sheep tick (*Ixodes ricinus*), the brown dog tick (*R. sanguineus*) and *Dermacentor sp*. *H. marginatum* is also a hard tick. Hard ticks vary in host-tick relationship. Species may have one host, two different hosts or three different hosts. After mating female hard ticks will feed once more followed by oviposition of hundreds to even thousands of eggs.

Soft ticks have similar body parts as the hard ticks. Key differences are that soft ticks lack the sclerotized outer cuticle found in hard ticks and the mouthparts of soft ticks are located below the end of the body (hard tick mouthparts stick out the front of the protected hood). For example the bird ticks, *Argas reflexus* and *A. persicus*, are soft ticks which can be a pest in for instance poultry farms.

Hard ticks have to be fixed to their hosts and the meal can last five days, while soft ticks are not fixed and the meal is finished in 20 to 50 minutes.

When searching for a possible host, ticks generally remain stationary until a host passes by. Once attached, ticks crawl to locate a place to feed. Commonly, ticks will attach to human skin along pant or sock lines or other tight locations which are warm and humid. Feeding can take hours to days depending on the species.

The bird ticks, *Argas persicus* and *A. reflexus* have worldwide distribution in warm climates. *A. persicus* occurs in small poultry farms and feeds blood on chicken and other domestic fowls. *A. reflexus* occurs in pigeon farms and on urban pigeons and their surroundings in towns. They can get from the nests of pigeons to lofts and attic rooms and feed on sleeping humans for blood. *A. reflexus* is an urban pest parasitizing urban pigeons and may cause a wide range of allergic reactions.

*Argas* spp. hide in cracks and crevices of chicken houses, nests, wooden equipments etc. during the day and come out to blood feed at night. Males and females are both blood
sucking. They are able to survive starvation for two years, which is why the protection against these mites is very difficult.

5.6.4.7.2 Dossier requirements

A clear label claim should be submitted. The study results of field trials should demonstrate the efficacy of the product, based on the submitted label claim.

Laboratory and, for some claims, field trials with ticks are needed to assess the efficacy of the product. The studies should normally be performed according to established guidelines where these are available. These may be international, EU or national guidelines. See Appendix 17 for a list of available guidelines. If no guidelines are available, the applicant may use their own methods, on condition however, that the study is scientifically robust, well reported and provides a clear answer to the question. In addition, the test methods applied and the test conditions should be clearly and fully described and must address the efficacy claim that appears on the product label. In the case of field trials where true replication is almost certainly impossible to achieve and where normal control methods are not restricted to use of a single acaricidal product, a full description of any factors that might be expected to influence product performance should be given. These may include the risk of re-invasion from adjacent areas, general levels of sanitation, treatment history etc. and are intended to provide the authorities with information to assist with the interpretation of the results obtained.

5.6.4.7.2.1 Test species

A product against ticks should normally be tested on the sheep tick *Ixodes ricinus*. When control or repellence of dog or bird ticks is claimed, tests with these ticks should be performed too (*Rhipicephalus sanguineus*, *A. reflexus*). When efficacy in the tropics is claimed or efficacy against *H. marginatum*, this tick should be tested too. *H. marginatum* behaves differently than *I. ricinus* since it is aggressive and it actively seeks the host to feed on and moves quickly on the ground. When the product is intended for use in poultry farms tests should be performed against *A. persicus*.

5.6.4.7.2.2 Laboratory tests and field trials

For the evaluation of biocides against ticks different types of laboratory and simulated-use tests can be used. Examples of tests are listed below.

**Laboratory test to evaluate knockdown and kill effect (no-choice test)**

A clear label claim should be submitted. The study results of field trials should demonstrate the efficacy of the product, based on the submitted label claim.

Laboratory and, for some claims, field trials with ticks are needed to assess the efficacy of the product. The studies should normally be performed according to established guidelines where these are available. These may be international, EU or national guidelines. See Appendix 17 for a list of available guidelines. If no guidelines are available, the applicant may use their own methods, on condition however, that the study is scientifically robust, well reported and provides a clear answer to the question. In addition, the test methods applied and the test conditions should be clearly and fully described and must address the efficacy claim that appears on the product label. In the case of field trials where true replication is almost certainly impossible to achieve and where normal control methods are not restricted to use of a single acaricidal product, a full description of any factors that might be expected to influence product performance should be given. These may include the risk of re-invasion from adjacent areas, general levels of sanitation, treatment history etc. and are intended to provide the authorities with information to assist with the interpretation of the results obtained.

**Laboratory test for repellents**

Candidate repellents are applied to human forearms from wrist to elbow. As a negative control untreated arm will be tested, preferably the other arm of the test person. A line
is drawn 3 cm above the wrist. Disease free ticks are placed on the back of the hand with a forceps, 5 ticks per test. Fresh, starved ticks are required for each exposure. The arm is inverted to promote upward movement toward the treated surface (ticks are negatively geotropic). The first exposure is one hour post treatment and continues once an hour for four hours. Each exposure is 5 minutes. Several criteria for repellence can be used. Either, a tick is considered non-repelled if it crosses the line 3 cm above the wrist or a tick is considered repelled when it drops down from the arm. A negative control treatment on an untreated arm, preferably the other arm of the same test person, should be performed.

Percentage repellence is calculated by recording the number of ticks crossing the line or dropping down from a treated arm as opposed to a control arm.

Normally, per repellent at least 10 persons are tested since repellence/attractiveness to ticks varies considerably between human individuals.

An alternative method could be using animals instead of humans to test the repellence against ticks. *I. ricinus, R. sanguineus* and *H. marginatum* will bite both humans and animals.

### Simulated use tests

To prevent disease transmission ticks must be knocked down, killed or repelled before attaching to the skin. For repellents the test described in 5.6.4.7.2.2 is a “worse case” test, therefore there is no need to do a field trial with repellents. For products that knockdown and kill ticks a simulated-use tests should be performed in which the product is applied according to the instruction for use and then tested in the presence of a person or an arm or foot or animal. For some products this can be a similar test set up as described in 5.6.4.7.2.2. Then it has to be established that the ticks are knocked down or killed before they can attach to the skin and start feeding. This is compared to a control test.

### 5.6.4.7.2.3 Requirements per type of claim

- Repellent: laboratory test for repellents;
- Insecticide with knockdown or kill effect: laboratory and simulated-use tests.

Simulated-use tests can be waived if a robust field trial is submitted.

### 5.6.4.7.3 Assessment of authorisation

#### 5.6.4.7.3.1 Norms and criteria

A biocidal product may only be authorised if it “possesses a sufficient level of efficacy” (BPD). This is implemented for ticks in the following way.

An insecticide against ticks is normally considered to be sufficiently “effective” if the following results can be achieved:

- Repellent:
  - \( \geq 90\% \) repellence during the claimed efficacy period;
- Product with knockdown effect:
  - \( 100\% \) knockdown before ticks start feeding and;
  - \( \geq 80\% \) kill within 24 hours;
- Product with kill effect:
  - \( \geq 95\% \) kill before ticks start feeding.

Deviations from these norms is possible but should be justified in the application.
5.6.4.8 Mites

5.6.4.8.1 Introduction

Mites, along with ticks, belong to the subclass Acarina (also known as Acari) and the class Arachnida. Mites are among the most diverse and successful of all the invertebrate groups. They have exploited an incredible array of habitats, and because of their small size (most are microscopic) most go totally unnoticed. Perhaps the best-known mite, is the house dust mite (family Pyroglyphidae), which can cause asthma and allergic symptoms. Mites are also important as vectors of microorganisms, transmitting rickettsiae and bartonellae. Flour mites (Acarus siro) and mould or storage mites (Tyrophagus putrescentiae, T. longior) are important pests in stored goods. Mites like the red mite, Dermanyssus gallinae, can be a pest in bird cages and poultry farms. The red mite can also feed on some species of mammals, including humans, but need an avian host to reproduce.

Part of the control of mites is covered in section 5.6.4.11 on stored goods. Often mites are only mentioned on a label as a secondary pest, while insects are the main pests.

5.6.4.8.1.1 Biology

The house dust mite is widespread in human habitation. House dust mites thrive in the indoor environment provided by homes, specifically in bedrooms and kitchens. Dust mites survive well in mattresses, carpets, furniture and bedding, with figures around 188 animals/g dust. Dust mites feed on organic detritus such as flakes of shed human skin and flourish in the stable environment of dwellings. The European house dust mite (Dermatophagoides pteronyssinus) and the American house dust mite (Dermatophagoides farinae) are two different species, but are not necessarily confined to Europe or North America; a third species Euroglyphus maynei also occurs widely. The average life cycle for a male house dust mite is 10 to 19 days. A mated female house dust mite can live for 70 days, laying 60 to 100 eggs in the last 5 weeks of her life.

The flour mite, A. siro, is the most common species of mite in foodstuffs. The males are 0.33 mm to 0.43 mm long and female are 0.36 mm to 0.66 mm in length. Flour mites contaminate grain and flour by allergens and they transfer pathogenic microorganisms. Foodstuffs acquire a sickly sweet smell and an unpalatable taste. When fed infested foodstuff, animals show reduced feed intake, diarrhoea, inflammation of the small intestine and impaired growth.

The red mite, Dermanyssus gallinae, is an ectoparasite of poultry and birds. They can be found in houses of laying hens, chickens and other fowls. The mites are blood feeders and attack resting birds at night. The optimal temperature is 27-28 °C. After feeding they hide in cracks and crevices away from daylight, where they mate and lay about 30-35 eggs in their lifetime. Their maximal lifetime is 8 weeks without starving and 6-10 months with starving. In spite of that these mites are ectoparasites, the main method of control is treating of the walls, bird cages, nests and hidden places in poultry farms with biocides.

5.6.4.8.2 Dossier requirements

A clear label claim should be submitted. The study results of laboratory or field trials should demonstrate the efficacy of the product, based on the submitted label claim.

Laboratory and/or field trials with mites are needed to assess the efficacy of the product. The studies should normally be performed according to established guidelines where these are available. These may be international, EU or national guidelines. See Appendix 17 for a list of available guidelines. If no guidelines are available or guidelines are not suitable, the applicant may use their own methods, on condition however, that the study is scientifically robust, well reported and provides a clear answer to the question. In addition, the test methods applied and the test conditions should be clearly and fully described and must address the efficacy claim that appears on the product
label. In the case of field trials where true replication is almost certainly impossible to achieve and where normal control methods might not be restricted to use of a single acaricidal product, a full description of any factors that might be expected to influence product performance should be given. These may include the risk of re-invasion from adjacent areas, general levels of sanitation, treatment history etc. and are intended to provide the authorities with information to assist with the interpretation of the results obtained.

5.6.4.8.2.1 Test species

Which test species should be used depends on the intended area of use and the label claim. In homes the European house dust mites, D. pteronyssinus, is the most important. In storage rooms the flour mite or storage mites, etc., for instance T. putrescentiae, A. siro. For use on poultry farms D. gallinae should be tested. When specific mite species are mentioned in the claim these should be tested.

5.6.4.8.2.2 Laboratory tests and field trials

For the evaluation of biocides against mites different types of laboratory and simulated-use tests can be used. Examples are given below.

Laboratory test to evaluate knockdown and kill effect (no-choice test)

The product is applied to representative surfaces or via direct cuticle application, in a container with mites, to assess inherent contact toxicity or knockdown effect of the active substance. For instance spray on a filter paper and put the filter paper in an aluminium dish. Specify whether adults (male or female) or nymphs are used. Normally tests are performed with 3 or more replicates, with normally 20 to 30 mites per replicate. Tests are done at 25°C and 70-75% R.H.. In all laboratory studies a treatment without biocide should be conducted with mites from the same population, as a negative control. The number of dead mites is counted at 24 hours after treatment.

Residual effect

For determination of residual efficacy, the formulated product should be applied to representative surfaces at a specified dose rate, or rates, including the recommended label rate. Mites should be exposed to the deposit at several time intervals after the deposit has dried (including the day of treatment, but after the deposit has dried completely and at the end of the claimed residual period). Exposure time should, preferably, be comparable to the time the mites might reasonably be expected to be in contact with a treated surface under practical conditions and assessors will take this factor into consideration when evaluating the data. Treated surfaces should include at least two porous and one non-porous substrate, representing surfaces that might, typically, be treated for mite control (e.g. plywood, painted plywood, textile fabric, according to the label claim). Mortality is normally assessed after 1 day up to 14 days post-exposure.

Simulated use tests

These tests are designed to mimic the practical use situation. For products that knockdown and kill mites simulated-use tests should be performed in which the product is applied according to the instruction for use. When products for general surface treatment are tested the mites must have a choice to be in contact with the biocide or not. The results should be compared to a control test, without biocide.

5.6.4.8.2.3 Requirements per type of claim

Specific mites: when specific mite species are mentioned in the claim (e.g. dust mite, red mite) both laboratory and simulated-use tests are required with the target species.

Mites as secondary pest: When mites are mentioned on the label claim only as a secondary pest, only laboratory tests with one mite species are required.
Acaricides: When mites are the main pest to control both laboratory and simulated-use tests are required with more than one mite species.

Space and structural treatments: requirements for these products are covered in section 5.6.4.11 on stored goods.

Simulated-use tests can be waived if a robust field trial is submitted.

5.6.4.8.3 Assessment of authorisation

5.6.4.8.3.1 Norms and criteria

A biocidal product may only be authorised if it “possesses a sufficient level of efficacy” (BPD). This is implemented for mites in the following way.

A biocide against mites is normally considered to be sufficiently "effective" if the following results can be achieved:

- laboratory tests: ≥90% mortality in 24 hours;
- simulated-use tests: ≥90% mortality in 1 week;
- field trials for space and structural treatments: requirements for these products are covered in section 5.6.4.11 on stored goods.

Deviations from these norms is possible but should be justified in the application.

5.6.4.9 Fleas

5.6.4.9.1 Introduction

This section covers the assessment of efficacy of products used for treatment against cat and dog fleas. The application of these products is indoors on surfaces.

These biocides are divided into two groups, namely the adulticidal and ovicidal/larvicidal products. Adulticidal products are intended for use against fleas in the adult growth stage, and the ovicidal/larvicidal products for use against fleas in the egg and larval stages. This distinction is based on the very different modes of action of the product, which result in different criteria for assessment.

It should be emphasized that products against fleas, which are applied directly on dogs and cats and have a medical claim are covered by legislation on Veterinary Medical Products. The reader may refer to the borderline dossier available on the ECB website (www.ecb.jrc.it/biocides).

5.6.4.9.1.1 Biology

Of the over 2000 species of fleas (Siphonaptera), the cat flea (Ctenocephalides felis) and the dog flea (C. canis) are the most common in-home pests in the EU. Fleas undergo complete metamorphosis (egg, larva, pupa, adult) and the lifecycle begins when an adult female finds a suitable host. Once found, the female flea will remain on this host for the rest of its life. Females produce several eggs after each blood feeding and can produce several hundred eggs in its lifetime. Once laid, the eggs fall off the animal host and develop in the areas where the host animal spends its time. The eggs tend to accumulate in the lowest areas such as deep in fibres of carpets, cracks in the floor, or crevices in furniture and furnishings.

Larvae require high protein food for their survival. This protein comes from feeding on the dry faeces of the adult fleas. The adult flea takes in more blood from the host than necessary for nourishment and excretes the remaining blood in almost pure form. Once dried, the faeces falls off the host animal where the larvae can feed. The larvae spin a cocoon and begin the pupal state.

An adult flea emerges from the pupae after stimulation from external cues that indicate an animal host is near. Once emerged, a flea must usually find a host (located using visual and thermal cues) within a week, or it risks death due to desiccation. Complete
development from egg to adult occurs in as little as two weeks, but this can take much longer depending on environmental conditions.

5.6.4.9.2 Dossier requirements

A clear label claim should be submitted. The study results of laboratory, simulated-use tests and field trials should demonstrate the efficacy of the product, based on the submitted label claim.

Laboratory and field trials with fleas are needed to assess the efficacy of the product. Ideally, the studies should be performed according to established guidelines where these are available. These may be international, EU or national guidelines. See Appendix 17 for a list of available guidelines. If there are no guidelines available or the guidelines are not suitable, the applicant may use their own methods, on condition however, that the study is scientifically robust, well reported and provides a clear answer to the question. In addition, the test methods applied and the test conditions should be clearly and fully described and must address the efficacy claim that appears on the product label. In the case of field trials where true replication is almost certainly impossible to achieve and where normal control methods are not restricted to use of a single insecticidal product, a full description of any factors that might be expected to influence product performance should be given. These may include the risk of re-invasion from adjacent areas, general levels of sanitation, treatment history etc. and are intended to provide the authorities with information to assist with the interpretation of the results obtained.

5.6.4.9.2.1 Test species

A product against fleas should normally be tested on the cat flea (*Ctenocephalides felis*) or the dog flea (*C. canis*).

5.6.4.9.2.2 For claims made for products intended for use as general surface treatments

For the evaluation of biocides against fleas different types of laboratory, simulated-use tests and field tests can be used.

Examples of the types of data that may be available when considering the efficacy of insecticide products intended for use as surface treatments are given below.

Laboratory studies

The product is applied to representative surfaces (e.g. carpet discs). Information on the fibre length and density should be provided, as this has a bearing onto flea survival. Long fibres enable fleas to hide and, thus, protect fleas from getting their share of the insecticide applied. Fleas are transferred to the surface, either before (direct contact) or after (residual performance) application of the product, to assess inherent contact toxicity or knockdown effect of the active substance.

Alternatively, ovicidal or larvicidal products can be tested in flea rearing medium containing flea eggs or larvae and the active substance in a range of concentrations, including the intended use concentration. Preferably, tests should be done in five replicates per treatment.

A control treatment without biocide with the same number of replicates should be included in all laboratory trials.

Simulated use studies

These tests are designed to mimic the practical use situation. The test should be performed according to the label claim.

Simulated-use tests can be waived if a robust field trial is submitted.
5.6.4.9.2.3 For claims made for products intended to be used as space spray treatments

Some insecticides against fleas can be used in foggers. For the evaluation of these insecticides different types of laboratory, simulated-use tests and field tests can be used. The efficacy test design should be defined for the available treatment method.

5.6.4.9.3 Assessment of authorisation

5.6.4.9.3.1 Norms and criteria

A biocidal product may only be authorised if it “possesses a sufficient level of efficacy”. This is normally implemented for fleas in the following way.

For laboratory and simulated use:

An adulticidal product against fleas is considered to be sufficiently “effective” if:

- within 24 hours 100% knockdown of the adult fleas should occur (this norm only applies if the test fleas are sprayed directly or are placed immediately on a treated carpet) and;
- within 48 hours ≥90% mortality of adult fleas should occur.

An ovicidal/larvicidal product against fleas is considered to be sufficiently “effective” if:

- ≥80% inhibition should occur of the development of produced eggs/larvae into adult fleas during the claimed ovicidal/larvicidal duration of action of the product.

Deviations from these norms is possible but should be justified in the application.

5.6.4.10 Litter Beetles

5.6.4.10.1 Introduction

There are several species of "litter beetles" that inhabit poultry droppings and litter. Litter beetles belong to the order Coleoptera, family Tenebrionidae. The most important are the lesser mealworm (other names: darkling beetle), Alphitobius diaperinus, and two species in the dermestid genus Dermestes; the hide beetle (D. maculatus) and the larder beetle (D. lardarius). Other species of beetles that occasionally cause damage to poultry housing are Dermestes ater, Tenebrio molitor, Alphitobius laevigatus, and Trox spp.

Litter beetles are of particular importance as a vector and competent reservoir of several poultry pathogens and parasites. The transmission of bacteria, (Salmonella, Escherichia coli) and protozoa (several Eimeria species which can cause coccidiosis) and different viruses can cause problems in livestock. This pest can also cause damage to poultry housing and is suspected to be a health risk to humans in close contact with larvae and adults. Adults can become a nuisance when they move en masse toward artificial lights generated by residences near fields where beetle-infested manure has been spread.

Often these beetles are only mentioned on a label as a secondary pest, while other insects are the main pests (control of flies, cockroaches, and litter beetles in poultry houses). But when they are mentioned specifically on the label they should be tested.

5.6.4.10.1.1 Biology

Lesser mealworm adults lay their eggs in cracks and crevices in the poultry house, in manure or litter, and in grain hulls. Larvae hatch and complete development to the adult stage in 40-100 days depending on temperature and food quality. The larvae consume spilled feed, manure and, to a lesser extent, dead birds and cracked eggs. Beetle populations in broiler and turkey houses often are concentrated around lines of feeders, which provide the beetles with shelter and an opportunity to feed on spilled bird feed. Mature larvae disperse when they are crowded to find isolated pupation sites, and this behaviour is responsible for much of their destructive activity. Crowded larvae leave the
litter and tunnel into thermal insulation materials where they construct pupal cells. Both larval and adult stages are omnivorous. The lesser mealworm is nocturnal, with greatest activity of both larvae and adults occurring shortly after dark. Populations of lesser mealworm often reach high densities, especially in deep-litter broiler and turkey houses and in high-rise caged layer operations. It is not unusual for the litter of a broiler house to move from beetle activity or for 70% of the surface of manure in a high-rise house to be covered with adult beetles.

5.6.4.10.2 Dossier requirements

A clear label claim should be submitted. The study results of laboratory, simulated-use tests and field trials should demonstrate the efficacy of the product, based on the submitted label claim.

Laboratory and simulated field trials with litter beetles are needed to assess the efficacy of the product. Ideally, the studies should be performed according to established guidelines where these are available. These may be international, EU or national guidelines. See Appendix 17 for a list of available guidelines. If there are no guidelines available or the guidelines are not suitable, the applicant may use their own methods, on condition however, that the study is scientifically robust, well reported and provides a clear answer to the question. In addition, the test methods applied and the test conditions should be clearly and fully described and must address the efficacy claim that appears on the product label. In the case of field trials where true replication is almost certainly impossible to achieve and where normal control methods are not restricted to use of a single insecticidal product, a full description of any factors that might be expected to influence product performance should be given. These may include the risk of re-invasion from adjacent areas, general levels of sanitation, treatment history etc. and are intended to provide the authorities with information to assist with the interpretation of the results obtained.

5.6.4.10.2.1 Test species

A product against litter beetles should normally be tested on the lesser mealworm, *A. diapernus*.

5.6.4.10.2.2 For claims made for products intended for use as general surface treatments

Examples of the types of data that may be available when considering the efficacy of insecticide products intended for use as surface treatments are given below.

**Laboratory studies**

The product is applied to representative surfaces, either before (persistence test) or after (direct contact) the insects are transferred to the surface, to assess inherent contact toxicity or knockdown effect of the active substance.

Preferably, test should be done in five replicates per treatment.

A control treatment without biocide should be included in all laboratory trials.

**Simulated use studies**

These tests are designed to mimic the practical use situation. The test should be performed according to the label claim.

Simulated-use tests can be waived if a robust field trial is submitted.

5.6.4.10.3 Assessment of authorisation

5.6.4.10.3.1 Norms and criteria

A biocidal product may only be authorised if it “possesses a sufficient level of efficacy”. This is normally implemented for litter beetles in the following way.
A product against litter beetles is considered to be sufficiently “effective” if:
For laboratory and simulated use:
- adulticide: ≥ 95% mortality;
- larvacide: ≥ 95% mortality;
- insect growth regulator: ≥ 90% mortality.

Deviations from these norms is possible but should be justified in the application.

5.6.4.11 Textile-attacking Insects (including fur and fabric attaching insects)

5.6.4.11.1 Introduction

Insecticides against textile-attacking insects can be used by professionals and non-professionals, use against beetle or moth larvae infested carpets for example.

Home user products may be used in vapour phase to prevent moth contact with stored clothing (via killing, repelling or attracting moth in traps) or insecticides may be applied to the surface of clothing to kill landing moths on contact.

Insecticides against textile-attacking insects can also be incorporated in the textile by industry for preventive treatments.

Other products made from textiles treated with insecticides are the so-called treated articles with an external claim (e.g. carpet with an insecticide not to protect the carpet but against fleas that are in contact with the carpet). These treated articles will not be considered specifically in this section since other than textile-attacking insects are the target insects.

5.6.4.11.1.1 Biology

The two main orders containing textile attacking insect species are Lepidoptera (moths) and Coleoptera (beetles). The webbing clothes moth (*Tineola bisselliella*), fur moth (*Tinea pellionella*), brown house moth (*Hofmannophila pseudospretella*) and carpet beetles (*Anthrenus* sp., *Anthrenocerus* sp.) are common in-house pests that feed on clothing, drapery, carpet and other natural hair fibres. The larvae of these insects have a diet consisting of natural hair fibres, which provide protein from keratin in the hair. These insects have adapted to be able to digest keratin, which is not easily digested by other insects.

Clothes moths are distributed worldwide. They feed during the larval cycle within a silken cocoon attached to hair fibre. Clothes moths larvae that feed only on natural hair fibres such as wool, will not feed on, silk, cotton, linens or synthetic fibres. Adult clothes moths do not feed. These adults mate and the females lays eggs directly on the natural fibre food source.

Carpet beetle larvae (e.g. *Anthrenus* sp., *Anthrenocerus* sp.) attack woollens, rugs and upholstered furniture, etc.. The adult beetles, which feed on nectar and pollen, can usually enter the home on plants, flowers or other vegetation. Eggs are then laid on lint in protected areas such as behind baseboards. Once hatched, larvae begin feeding on a number of natural textiles or displays (animal horns, hoofs, insect collections, etc).

5.6.4.11.2 Dossier requirements

A clear label claim should be submitted. The study results of simulated-use tests or field trials should demonstrate the efficacy of the product, based on the submitted label claim.

For vapour based products the label should provide information on the volume that can be covered with the product (closet of x m³, room of y m³).
Laboratory and simulated-use trials with textile-attacking insects are normally needed to assess the efficacy of the product. Ideally, the studies should be performed according to established guidelines where these are available. These may be international, EU or national guidelines. See Appendix 17 for a list of available guidelines. If there are no guidelines available or the guidelines are not suitable, the applicant may use their own methods, on condition however, that the study is scientifically robust, well reported and provides a clear answer to the question. In addition, the test methods applied and the test conditions should be clearly and fully described and must address the efficacy claim that appears on the product label. In the case of field trials where true replication is almost certainly impossible to achieve and where normal control methods are not restricted to use of a single insecticidal product, a full description of any factors that might be expected to influence product performance should be given. These may include the risk of re-invasion from adjacent areas, general levels of sanitation, treatment history etc. and are intended to provide the authorities with information to assist with the interpretation of the results obtained.

5.6.4.11.2.1 Test species

A product against textile-attacking insects should normally be tested on:

- one of the following moth species:
  - the clothes moth (*Tineola bisselliella*);
  - the fur moth (*Tinea pellionella* L.);
  - the brown house moth (*Hofmannophila pseudospretella*);
- one of the following carpet beetle species:
  - *Anthrenus* sp;
  - *Anthrenocerus* sp.

Whether adults or larvae or both should be tested depends on the label claim.

5.6.4.11.2.2 Laboratory tests and field trials

For the evaluation of biocides against textile attacking insects different types of laboratory and simulated-use tests can be used. Examples of tests, mainly for clothes moth, are listed below.

**Laboratory tests**

**Mortality test**

Webbing clothes moths, adults, larvae (2nd-3rd instar) or eggs may be placed in a jar (e.g. 240 ml glass jars, brass-screened lid) containing a treated textile sample (e.g. circular, 4cm diameter, 100% wool sample).

Jars are periodically evaluated by recording mortality, egg laying and hatch (optional), and larval damage. A moth is considered inactivated when it is not able to walk or fly, in a spontaneous way or when stimulated with a brush or pin.

New moths are introduced into the jars periodically to test residual effects (depending on the label claims). Tests should normally be done in five replicates. A control treatment without biocide with the same number of replicates should be included in all laboratory trials.

**Repellency test**

Moth repellency can be tested in a choice test. Moths are placed in a clear tunnel between two dark boxes, both containing wool. One of the boxes contains the repellent product. The adult moths are released in the tunnel after which they can choose the treated or untreated box. The ratio of moths found in the treated vs. untreated box is a measure for the efficacy of the product.
Simulated use

These tests are designed to mimic the practical use situation. The study results should provide a clear picture of the efficacy of the product.

An example of tests that might match the proposed intended use of the product:

Simulated-use tests with moths added to drawers (minimum air volume: 0.016 m$^3$) or closets (minimum air volume: 0.5 m$^3$) can provide good information on home user products. In tests with vapour based products the door should be opened with a frequency resembling normal opening of a closet, to show that this does not reduce efficacy: once a day during completion of the assay, 5 seconds for drawers and 10 seconds for closets. Assessments of mortality would form the basis for efficacy claims. Additionally damage to the test material can be assessed. The damage will depend upon the number of insects, their developmental stage, the exposure time and the size and quality of the piece of carpet, etc. Therefore, damage should always be assessed in comparison to the control treatment.

Simulated-use tests can be waived if a robust field trial is submitted.

Test similar to the ones mentioned above can also be used to show efficacy against carpet beetles and the larvae of carpet beetles.

5.6.4.11.3 Assessment of authorisation

5.6.4.11.3.1 Norms and criteria

A biocidal product may only be authorised if it “possesses a sufficient level of efficacy”. This is implemented for textile attacking-insects normally in the following way.

At the end of an exposure period (e.g. 1 week):

- more than 90% of the adults and larvae should be killed (unless claimed different);
- a repellent should perform according to the label claim, preferably >90%.

Deviations from these norms is possible but should be justified in the application.

5.6.4.12 Stored Goods-attacking Insects and Mites

5.6.4.12.1 Introduction

The purpose of biocidal products against stored goods-attacking insects and mites is to control pests in storerooms, freight and alternative transport containers for products of plant origin etc. They should also protect the actual stored goods against insects and mites. The term “stored” in this regard refers specifically to: stored products (of plant origin) for human consumption, animal feed, industrial processing and propagation.

Products against stored goods-attacking insects can either be biocides or plant protection products. In general, where the stored products are protected, prior to processing, the use falls under plant protection and is not relevant in this guideline.

There are a number of different insects that attack stored goods. Common beetle invaders include grain beetles (*Tribolium castaneum*, *Oryzaephilus surinamensis*, etc.), confused flour beetles (*Tribolium confusum*), and rice weevils (*Sitophilus oryzae*). Indian meal moth (*Plodia interpunctella*) and flour mite (*Acarus siro*) are also very common pest. Infestations of these pests can occur at the packaging plant, the store, or in the home, making it difficult to determine where the source of the problem is. Sometimes these infestations are only noticed by the consumer once the insect leaves the food product and enters the home environment.

For professional and industrial use there are two classifications of such products:

- fumigation with gases, which is used for controlling pests in rooms used for the storage of products of plant origin (storerooms, freight structures and means of
transport, gassing installations etc.);

- products other than gases, which are used for controlling pests in empty or full storerooms (including products which are applied by means of vaporisers).

5.6.4.12.2 Dossier requirements

A clear label claim should be submitted. The study results of simulated-use tests and field trials should demonstrate the efficacy of the product, based on the submitted label claim.

Laboratory and field trials with stored goods-attacking insects are needed to assess the efficacy of the product. Ideally, the studies should be performed according to established guidelines where these are available. These may be international, EU or national guidelines. See Appendix 17 for a list of available guidelines. EPPO standards PP 201 to 204 are recommended (Appendix 17). If these guidelines are not suitable, the applicant may use their own methods, on condition however, that the study is scientifically robust, well reported and provides a clear answer to the question. In addition, the test methods applied and the test conditions should be clearly and fully described and must address the efficacy claim that appears on the product label.

In the case of field trials where true replication is almost certainly impossible to achieve and where normal control methods are not restricted to use of a single insecticidal product, a full description of any factors that might be expected to influence product performance should be given. These may include the risk of re-invasion from adjacent areas, general levels of sanitation, treatment history etc. and are intended to provide the authorities with information to assist with the interpretation of the results obtained.

5.6.4.12.2.1 Test species

A product against stored goods-attacking insects may be tested on beetles, moths or mites (more specifically mentioned in the relevant EPPO guidelines), or insects that are specifically identified in the label claim.

5.6.4.12.2.2 Laboratory tests and field trials

Depending on the application and the purpose of the product, one of the trials below (or equivalent trials) normally should be performed.

**Consumer products**

For consumer products laboratory or simulated-use tests are required. A direct spray test method can be used to evaluate performance against stored goods-attacking insects. A simulated use test can be a test, performed in a laboratory, where insects (either cultured or natural populations) are in contact with the stored goods (e.g. breakfast cereal, flour) and the biocide is applied according to the instructions for use.

Simulated-use tests can be waived if a robust field trial is submitted.

A control treatment without biocide with the same number of replicates should be included in all laboratory trials.

**Gases for use in storerooms, freight and transport rooms and gassing installations with stored products present**

Additional laboratory studies are not required, only field trials.

A field trial should normally be conducted according to the EPPO guideline PP 1/201(1) “Fumigants to control insect and mite pests of stored plant products”.

The field of use of the gas are places where large supplies are stored, in particular cereal products, but also other food products such as dried nuts, processed vegetables, spices or meals.
The use of gas can be intended for controlling/fighting pests in spaces but also for controlling/fighting pests in or on the product itself.

**Products other than gases for storerooms with or without stored products**

Additional laboratory studies are not required, only field trials.

A field trial normally should be performed according to the EPPO guideline PP 1/202 (1) "Space and structural treatments of storerooms".

The products concerned exclude gases, but do include those applied by means of vaporisers (fogs, smokes, vapours, space sprays).

This trial focuses on the control of pests in full or empty storerooms (walls, cracks, etc.). The trial does not serve to test the efficacy of the treatment on pests in the stored products themselves.

The trial can be performed in two ways.

- the first possibility is conducting the trial in rooms where there is already an infestation. Using a trapping system, the effectiveness is determined by scoring the number of insects caught in the traps before and after the treatment;
- the second possibility is conducting the trial in a room where test organisms have been introduced artificially (usually in small cages). The effectiveness is determined by scoring the number of alive, ‘knocked down’ and dead organisms in comparison with an untreated room.

### 5.6.4.12.3 Assessment of authorisation

**5.6.4.12.3.1 Norms and criteria**

A biocidal product may only be authorised if it “possesses a sufficient level of efficacy”. This is implemented for stored goods attacking-insect in the following way.

- consumer products: normally 100% mortality in direct spray tests, in simulated-use tests >90% knockdown and >70% mortality after 24 hours would be sufficient;
- gases: the duration of gassing (as specified in the label claim) should be such that at the end of gassing 100% of the insects/mites are dead or dying. It is possible to distinguish between dead and dying insects, which will not recover anymore, so these should also be counted;
- the duration of gassing should not be longer than necessary;
- all non-gases: the effect should be achieved within the duration of the treatment, as specified in the label claim. Normally >90% would be sufficient.

Deviations from these norms is possible but should be justified in the application.

### 5.6.4.13 Flies

**5.6.4.13.1 Introduction**

Flies are common pests in and around the house and in animal rearing facilities. Some of these insect species are merely a nuisance, others provide discomfort from irritating bites, and some potentially carry and transmit diseases.

The possible fields of use of the insecticides include: residential and other types of accommodation, public spaces, hospitals, storerooms, kitchens, waste dumps and stables and manure storage facilities.

**5.6.4.13.1.1 Biology**

House flies (*Musca domestica*) and other nuisance flies are common non-biting pests in the EU. The house fly lifecycle goes through four stages: egg, larvae (maggots), pupa, and adult. Eggs are laid on organic debris including faeces, decaying vegetation, etc.
Once hatched, larvae feed by burrowing into the organic debris and filter decaying organic matter. In the pupal stage the fly is transformed into the adult. During this transformation, no feeding takes place. At the adult stage, house flies feed by regurgitating on food, then lap up the food in liquid form. The life cycle of house flies, from egg to fly, is 1 to 3 weeks, depending on the climate conditions. Males die soon after mating, females live temperature dependent normally one to several weeks in the field.

Flies regularly fly into and out of man-made structures. Outside, flies land on faecal material and other debris. Inside, flies land on human food and contact other substrates regularly touched by humans. Here, potential pathogens can be transferred on the flies’ body (legs) or from inside the body (vomiting on potential food in order to feed) which are picked up in faecal or other decaying material. More than 100 germs have been documented as being transferred by house flies. Among them are *Salmonella* sp. and *E. coli* have been documented as being transferred by house flies.

The stable fly (*Stomoxys calcitrans*) is a pest often found in stables alone or together with the housefly. Rather unusual for a member of the family Muscidae is that it sucks blood from mammals. Under favourable conditions the stable flies develop from egg to fly in 3 weeks. The adults live several weeks.

Other biting flies include black fly (Simuliidae) and deer and horse flies (*Chrysops* and *Tabanids*), are also common pests in the EU. These insects can inflict a painful bite leaving an itchy welt. Some are also known to transmit disease. Apart from these species blow-flies can be of significance in a number of localities, including food producing facilities (Carrion flies, blue bottle fly, green bottle flies).

**5.6.4.13.2 Dossier requirements**

A clear label claim should be submitted. The study results of laboratory and simulated-use tests and field trials should demonstrate the efficacy of the product based on the submitted label claim.

Laboratory, simulated-use tests and field trials with the test insects are needed to assess the efficacy of the product, depending on the label claim. Ideally, the studies should be performed according to established guidelines where these are available. These may be international, EU or national guidelines. See Appendix 17 for a list of available guidelines. If there are no guidelines available or the guidelines are not suitable, the applicant may use their own methods (intra-company Standard Operating Procedures), on condition however, that the study is scientifically robust, well reported and provides a clear answer to the question. In addition, the test methods applied and the test conditions should be clearly and fully described and must address the efficacy claim that appears on the product label. A control treatment without biocide with the same number of replicates should be included in all laboratory trials.

In the case of field trials where true replication is almost certainly impossible to achieve and where normal control methods are not restricted to use of a single insecticidal product, a full description of any factors that might be expected to influence product performance should be given. These may include the risk of re-invasion from adjacent areas, general levels of sanitation, treatment history, season, etc. and are intended to provide the authorities with information to assist with the interpretation of the results obtained.

**5.6.4.13.2.1 Test species**

In case of an authorisation against flies the prescribed test insect is the housefly (*M. domestica*). When the product claim includes use in stables and animal housings (except poultry), for a general claim against flies both the housefly and the stable fly (*S. calcitrans*) should be tested. If efficacy against blow-flies is claimed tests have to be done with a blow-fly species (Calliphoridae). Skin repellents against flies should be
tested against biting flies, for instance the stable fly. Spatial repellents against flies should be tested against the housefly or, when used in stables, against both housefly and stable fly. Products intended for use as repellent on horses (recreational and/or sport horses) should be tested against the claimed target organisms (see Appendix 16 Species grid PT19 3H to 3T).

5.6.4.13.2.2 Laboratory testing simulated use tests and field trials

For evaluation of biocides against flies different types of laboratory, simulated-use tests and field test can be used. Examples of tests are listed below.

Laboratory tests

Flies can be tested in the laboratory in small jars or Petri dishes. The surface can be treated or granules can be placed, after which insects can be added at different time intervals. Alternatively, the flies can be sprayed directly. The knockdown percentages and mortality are determined.

A control treatment without biocide with the same number of replicates should be included in all laboratory trials.

Simulated use tests

For assessment of efficacy simulated-use tests should be conducted in a test chamber, for instance the Peet-Grady chamber. This is an airtight room of 1.8 x 1.8 x 1.8 m³, into which a certain amount of product is introduced. Other chambers of similar or bigger size are acceptable, either airtight or with air exchanges. The chamber should be washed and dried between each replicate to avoid chemical contamination.

Environmental conditions must be specified during the test (temperature, humidity, photoperiod). Temperature would be expected to fall in the range 19-29°C, may be lower for use in stables. A control treatment without biocide with the same number of replicates should be included in all laboratory trials.

Simulated-use tests can be waived if a robust field trial is submitted.

Examples of tests for different products are listed below. For other types of products similar test can be performed.

Space treatment

In the case of an application of a liquid for space treatment, the aerosol test method is performed in the test chamber in the laboratory. A known number (50-100) test insects, including males and females, are exposed to the space treatment. The dose sprayed in the chamber should be comparable to the label directions. The test is performed in quadruplicate. A control treatment without biocide should be included. The knockdown percentages and mortality of flies in both insecticide treatment and negative control are determined.

Surface treatment

Products for surface treatment (including window stickers) act on the insect via contact with or feeding from the treated surface. The product can be applied by spraying, brushing, painting, etc. according to the label. These products are also tested in the test chamber.

In the test chamber the product is applied on a small surface or on the whole chamber, in a dose rate appropriate to the label claim. After the surfaces have been left to dry the test can commence. The insects are released in the test chamber at several time points after application (or at least at the maximum residual time claimed at the label), to show residual efficacy. At a suitable period of exposure (e.g. 24 hours) after each test time point mortality of the test insects is recorded. It is mandatory to report temperature and air humidity in the test room. These should agree as much as possible with practical use conditions.
Products to be vaporized or fogged

Only a French recognized guideline (NF T 72-321) is available for efficacy studies with products against flies that should be vaporized (heating element that heats a tablet or liquid, coils, fan driven devices, etc.) or products that should be applied in a fogging treatment. Recently WHO published a guideline for these types of products against mosquitoes. This guideline might be adapted for fly products. Further, the “Large room test” is generally accepted. Other methods are also acceptable if they are scientifically sound and provide a clear picture of the efficacy of the product.

The “Large room test” test can be performed in a non-ventilated room of 20 to 30 m$^3$. When a ventilated room is used (mimics in some cases reality better) the air exchange should be measured (e.g. one air chamber renovation per hour). The product is applied according to the intended use, allowing it to evaporate over a specified time period (depending on the label claim e.g. 9 hours).

House flies (M. domestica) are exposed to the vapour/fog at different time points, e.g. at 0, 2, 4, 6 and 8 hours. The test insect to be used depends on the requested application. At every time point a known number of test insects (e.g. 50), including males and females, are exposed to the vapour. The test is performed in quadruplicate. A control treatment without biocide should be included.

The knockdown percentages (KD50, KD95, KD100), mortality and, if possible, the concentration of the active substance in the room are determined.

Larvicides

Larvicides are often applied to the floor of stables and to manure to prevent maggots and pupa from developing into the next stage. These products can be tested in naturally or artificially infested manure, in boxes covered with gauze. Adult flies emerging from the manure are counted and the difference between treated and untreated manure is analysed. Where IGRs (insect growth regulators) are used as larvicides, it is possible to additionally assess the deformation of larvae and pupae.

Bait products

For products formulated as baits the product should also be tested to establish the intrinsic palatability of the formulation.

The most important factor involved in laboratory testing is to provide a free choice alternative food source to the test insects. The formulation should demonstrate acceptable toxicity in competition with the alternative food source. A control treatment without biocide of similar size as the test itself (i.e. number of replications) should be included in all laboratory trials.

If conducted on both fresh and aged product it may provide information on the storage stability of the product.

Repellents

For products with a repellent effect against flies no agreed protocols are available. The tests should be designed to mimic the practical use situation. The study results should provide a clear picture of the efficacy of the product. The submitted data from studies are checked for completeness, based on the applied dose per treated area. It is also checked whether the duration of exposure is sufficient. If the formulation alone i.e. without the carrier (e.g. a product with a tissue as carrier) has been tested, data on release from the carrier are also required. The study data should provide a clear picture of the efficacy of the formulated product.

Field trials

For application in cattle houses, pigsties and/or treatment of pig and cattle manure for controlling flies, field trials are normally required, both for insecticides and repellents.
Tests are done preferably during spring and beginning of summer. At the end of summer and autumn population decline might be due to natural causes instead of the insecticide treatment. Apply the insecticide according to the label instructions.

During field trials in stables, special consideration should be given to the choice of the building material (concrete, wood etc.) of the walls and floors of the stables, as well as to the ventilation (number of total air changes per 24 hours), because the conditions should be representative of a practical situation. This can differ per EU country. It is possible to assess whether extrapolation to other types of accommodation is justified. If for example a general registration for poultry houses is requested, but studies conducted in a house for laying hens have been submitted, a rational should be provided that extrapolation is justified.

The effect on the fly population can be determined by counting the numbers of flies (estimation of population size) before, during and after the treatment, or by the differences between treated and untreated objects in the same area. Various assessment methods are acceptable including visual assessments (fly density on a surface or animals is assigned to a category) or quantified measures such as using sticky fly papers, digital photographs of marked areas on walls, collecting dead flies from a defined floor or aisles area etc..

5.6.4.13.2.3 Requirements per type of claim

Per type of claim the requirements will be listed.

Products intended for use as general surface treatment, space treatment or vaporisers in houses:

- a simulated-use test showing mortality and knockdown and/or residual efficacy according to the claim.

Products intended for use as general surface treatment, space treatment or vaporisers in stables and waste dumps:

- a laboratory test showing mortality and/or knockdown and/or residual efficacy, depending on the claim;
- a field trial according to the directions for use.

Products intended for use as larvicides:

- a laboratory test showing larva mortality;
- a simulated-use test showing decrease in number of emerging flies.

Products intended for use as repellent:

- a laboratory or simulated-use test showing repellence;
- field test showing repellence (only required in some cases, for instance when a repellent is used to prevent flies from entering stables).

Products intended for use as repellent on horses (recreational and/or sport horses only):

- a laboratory test demonstrating repellence;
- a simulated use/ field test demonstrating repellence against the specific target fly species on target animals.

Simulated-use tests can be waived if a robust field trial is submitted.

5.6.4.13.3 Assessment of authorisation

5.6.4.13.3.1 Norms and criteria

A biocidal product may only be authorised if it “possesses a sufficient level of efficacy”. An insecticide against flies is considered to be sufficiently “effective” if the following results can be achieved:
Products intended for use as general surface treatment, space treatment or vaporisers in houses:

- required results in simulated-use tests:
  - the level of knockdown efficacy should be \( \geq 80\% \);
  - mortality after 24 hour should be \( >90\% \).

Products intended for use as general surface treatment, space treatment or vaporisers in stables and waste dumps:

- required results in laboratory tests:
  - the level of knockdown efficacy should be \( \geq 80\% \);
  - mortality after 24 hour should be \( \geq 90\% \);
- required results in field trials:
  - reduction in the amount of flies according to the claim (or compared to the control situation).

Products intended for use as larvicides:

- required results in laboratory tests:
  - \( >90\% \) larva mortality;
  - showing decrease in number of emerging flies.

Products intended for use as skin repellent:

- required results in tests:
  - showing repellence, preferably \( >90\% \).

Products intended for use as spatial repellent:

- required results in tests:
  - showing repellence, preferably \( >80\% \).

Products intended for use as repellent on horses (recreational and/or sport horses only):

- a laboratory test demonstrating repellence;
- a simulated use/field test demonstrating sufficient repellence against the specific target fly species on target animals.

Deviations from these norms is possible but should be justified in the application.

5.6.4.14 Mosquitoes

5.6.4.14.1 Introduction

Mosquitoes, including species in the Culex, Aedes, and Anopheles Genera are common pests in parts of the EU. As well as their annoying behaviour and itching bites, mosquitoes are well-known for transmitting diseases such as Malaria (Anopheles spp.), yellow fever, Dengue (Aedes spp.), West Nile (e.g. Culex spp.), blue tongue virus in animals, and various encephalitis. Although none of these diseases are endemic in Europe, occasional outbreaks occur and European travellers might encounter them, either in European tropical overseas regions or in the rest of the world. Biocides against mosquitoes can only claim to kill or repel the mosquitoes, not to prevent the diseases.

5.6.4.14.1.1 Biology

Like all Diptera, mosquitoes also go through four stages of development. The egg, larval and pupal stages take place in still aquatic environments such as floodplains, drainage ditches, natural and artificial water containers. Depending on the species, female mosquitoes will lay eggs directly in these aquatic environments or adjacent to locations in mud which typically have fresh water or tidal flooding events. Depending on the genera, eggs are laid individually or in clumps called rafts.

Once larvae hatch, filter feeding begins near the top of the water. Typically, mosquitoes go through 4 larval instars before beginning the pupal stage. Once completed, mosquito
adults emerge from the aquatic and enter the aerial environments. Mating usually begins a few hours to days after emergence. Once mated, the females begin to search for a blood meal. Humans and domestic animals are included as potential blood hosts, with some mosquito species preferring human blood to other animals.

Adult female mosquitoes locate potential blood hosts by detecting attractants such as carbon dioxide and skin emanations. Once located, the mosquito will attempt to bite, taking in a blood meal. This blood meal is partially digested and used for the development of eggs.

5.6.4.14.2 Dossier requirements

A clear label claim should be submitted. The study results of trials should demonstrate the efficacy of the product based on the submitted label claim.

Laboratory, simulated-use tests and field trials with the test insects are needed to assess the efficacy of the product. Ideally, the studies should be performed according to established guidelines where these are available. These may be international, EU or national guidelines. See Appendix 17 for a list of available guidelines. Several WHO tests are available for mosquito testing. If the available guidelines are not suitable, the applicant may use their own methods (intra-company Standard Operating Procedures), on condition however, that the study is scientifically robust, well reported and provides a clear answer to the question. In addition, the test methods applied and the test conditions should be clearly and fully described and must address the efficacy claim that appears on the product label. A control treatment without biocide should be included in all laboratory trials.

In the case of field trials where true replication is almost certainly impossible to achieve and where normal control methods are not restricted to use of a single insecticidal product, a full description of any factors that might be expected to influence product performance should be given. These may include the risk of re-invasion from adjacent areas, general levels of sanitation, treatment history, season, etc. and are intended to provide the authorities with information to assist with the interpretation of the results obtained.

5.6.4.14.2.1 Test species

In case of an authorisation against mosquitoes insecticide testing should be performed with the house mosquito (Culex spp.) since this is the most common in Europe and a large mosquito, which makes it one of the most difficult to kill. Since Aedes spp. are the most aggressive mosquitoes repellents should be tested on this species too.

When use in tropical areas is claimed it should be specified against which mosquito spp. the product is effective and these should be tested (e.g. malarial mosquitoes: Anopheles).

Products intended for use as repellent on horses (recreational and/or sport horses) should be tested against the claimed target organisms (see Appendix 16 Species grid PT 19 3H to 3T).

5.6.4.14.2.2 Laboratory studies

For the evaluation of biocides against mosquitoes different types of laboratory, simulated-use tests and field test can be used. Examples of test are listed below. Mosquitoes used in all tests should be disease free.

Laboratory tests against adults

Insecticides against mosquitoes should normally be tested in the laboratory in WHO cones or WHO cylinders by force tarsal contact. The test is well described in WHO guidelines (methodology, number, age, nutritional status of the specimens and insecticide susceptibility of the strains). Only females have to be tested. First laboratory
test (bio assay) can be conducted on a laboratory strain of well-known insecticide susceptibility. A second test can be conducted on field populations obtained by larval collection. Tests should be conducted on F1 generation adults. Mosquitoes are exposed during a few minutes to a treated surface and their evolution (knock down, death) is followed during 24 hours. The knockdown percentages and mortality are determined.

The cone tests can also be used to evaluate the efficacy of insecticide treated net. For netting evaluation the exposure time is only 3 minutes and mortality is also checked after 24 hours.

Tunnel tests baited with birds or little mammals could be conducted to assess the feeding inhibition, the repellent effect and the insecticide effect.

A control treatment without biocide with an adequate number of replicates should be included in all laboratory trials.

**Laboratory tests: Larvicides**

Larvicides are applied to water to prevent larva to develop into adult mosquitoes. These products can be tested in naturally or artificially infested water, in boxes covered with gauze. Tests are normally not performed in tap water but in water containing organic particles, especially where a claim for residual performance is made. Test is normally performed on late 3rd-early 4th larval stages only. Mortality is usually checked after 24 hours. For slow acting insecticides and insect growth regulators mortality has to be checked for several days. In that case food has to be supplied to larval stages. A control population susceptible to insecticide should be used as control in all bio-assays (positive control). A control treatment without biocide should be included as negative control. Adult mosquitoes emerging from the water are counted and the differences between treated and untreated boxes are analysed. The methodology of this bio-assay is described in WHO guidelines (WHO/CDS/WHOPES/GCDPP/2005.13).

**Simulated use tests**

For assessment of efficacy simulated-use tests should be conducted in a test chamber, for instance the Peet-Grady chamber. This is an airtight room of 1.8*1.8*1.8 m, into which a certain amount of product is introduced. Other chambers of similar or bigger size are acceptable.

The chamber should be washed and dried between each replicate to avoid chemical contamination.

Next to chambers experimental huts can be used. These huts are small buildings, several build next to each other, in which wild mosquitoes can enter but they have no way to escape. Volunteers are in the huts as attractants for mosquitoes. In each hut, the treatment of the hut (space or surface treatment) or the volunteers (skin repellents) should be different: test product, negative control (no biocide) or positive control (standard product e.g. DEET for repellents). At the end of a test period (e.g. one night) number of mosquitoes are counted by species, by status (death or alive), by engorgement (fed or unfed) and by position in the hut (hut or exit traps). Advantage of the hut is that wild populations can be used and that it is ventilated (mimics reality better in some cases).

Environmental conditions must be specified at the beginning and during the test (temperature, humidity, photoperiod). Temperature would be expected to fall in the range 19-29°C. When efficacy at high temperatures is claimed (use in the tropics) test at temperatures >30°C should be provided. A control treatment without biocide should be included in all laboratory trials.

Simulated-use tests can be waived if a robust field trial is submitted.
Space treatment simulated use tests
In the case of an application for a liquid for space treatment, the aerosol test method is performed in the test chamber in the laboratory. A known number (e.g. 50-100) test insects (females) are exposed to the space treatment. The dose sprayed in the chamber should be comparable to the label directions. The test is replicated 3 or more times. The knockdown percentages and mortality of mosquitoes in both insecticide treatment and negative control are determined. Ideally, a ventilated room should be used to mimic the intended use better.

Surface treatment simulated use tests
Products for surface treatment act on the insect by tarsal contact with the treated surface. The product can be applied by spraying, brushing, painting, etc. according to the label. These products are also tested in a test chamber or an experimental hut. The WHO guideline for testing mosquito adulticides describes such a test.

In the test chamber the product is applied on small surface, or on the whole chamber, in a dose rate appropriate to the label claim. A negative control should be included. After the surfaces have been left to dry the test can commence. The insects are released in the test chamber at several time points after application (or at least at the maximum residual time claimed at the label), to show residual efficacy. After 24 hours mortality of the test insects is recorded. It is mandatory to report temperature and air humidity in the test room. These should agree as much as possible with practical use conditions.

Products to be vaporized or fogged simulated use tests
No officially recognized guidelines are available for efficacy studies with products that should be vaporized (heating element that heats a tablet of liquid, coils, fan driven dives, etc.) or products that should be applied in a fogging treatment. The “Large room test” is generally accepted. Other methods are also acceptable if they are scientifically sound and provide a clear picture of the efficacy of the product.

The “Large room test” test can be performed in a non-ventilated room of 20 to 60 m³. When a ventilated room is used (mimics reality better in some cases) the air exchange should be measured (e.g. one air chamber renovation per hour). The product is applied according to the intended use, allowing it to evaporate over a specified time period (depending on the label claim e.g. 9 hours).

Mosquitoes are exposed to the vapour/fog at different time points, e.g. at 0, 2, 4, 6 and 8 hours. At every time point a known number of female test insects (50-100) are exposed to the vapour. The test is replicated 3 or more times. A negative control should be included.

The knockdown percentages (KD50, KD95, KD100), mortality and, if possible, the concentration of the active substance in the room are determined.

When the label claim says that the product should be used in ventilated rooms the opening of windows and doors should be simulated in the test.

Repellents
Repellents are products with a repellent effect and can drive away mosquitoes. These products are either applied on human or animal skin or on clothes (topical or skin repellent) or release the active ingredient to the air (spatial repellents). This is based on the biological activity of the evaporated active substance.

Products with a repellent effect, which are applied on the human skin or clothes, can be tested in an “arm-in-cage” simulated-use test. The repellent is applied in the specified dose on the (bare) forearm of the test person. The forearm is subsequently exposed to test mosquitoes in a cage for 5 minutes. This should be repeated every hour, at least up to the claimed efficacy period. If one bite is received during an exposure followed by another bite in the next exposure (confirming the first bite), the test should be stopped and the time of the first confirmed bite recorded as the length of repellence. If bites are
not received in succession, then the test is continued and the first bite should be
corrected ‘unconfirmed’. The same test is repeated with untreated forearms of,
preferably, the same test persons. For the untreated forearm, a minimum of 5 lands in 5
minutes is required to qualify the test. Once 5 lands are received, the arm should be
removed to prevent excess biting. If less than 5 lands are counted in 5 minutes, then the
test should not proceed and the mosquito cage should be replaced with ‘fresh’
mosquitoes. The results of treated and untreated forearms are compared.
Alternative methods using rabbits are developed. The repellent solution could be applied
directly on the skin of a rabbit on which a cage containing female mosquitoes is placed.
Skin repellents can also be tested in and experimental hut, as long as the number of
mosquitoes entering the hut is not too low.

Similar tests can be used for cloth in which a repellent is incorporated (treated article).
Repellent effectiveness is based on protection time, that is, the time between repellent
application and the time of 2 or more bites on the treated arm, or the first confirmed bite
(a bite followed by another within 30 min.).
For products with a repellent effect, which are applied in another way (not on the human
skin or clothes, for instance spatial repellents), no common protocols are available.
These products can be tested in a simulated use test, for instance in an experimental
hut. The submitted data from studies are checked for completeness, based on the
applied dose per treated area. It is also checked whether the duration of exposure is
sufficient. If the formulation alone i.e. without the carrier (e.g. a product with a tissue as
carrier) has been tested, data on release from the carrier are also required. The study
data should provide a clear picture of the efficacy of the product.

When the label claim says that the product should be used in ventilated rooms the
opening of windows and doors should be simulated in the test.

Larvicides simulated use tests
In small scale simulated-use tests, insecticide formulation can be tested in natural
breeding sites or simulated larval breeding sites. When natural larval populations are
used pre-treatment assessments of the population should done at the site (larval count
by dipping technique). Depending on the protocol, eggs or larvae can be regularly
introduced in the treated sites to evaluate the residual efficacy. Breeding sites are kept
uncovered to allow wild adults to lay their eggs. The methodology of this test is

Field trials
For some products against mosquitoes, field trials are not required. Especially when field
populations are used in the lab or in an experimental hut. However, for some products
and uses a simulated-use test cannot mimic the practical situation sufficiently (e.g.
larvicides used in large swamps and lakes, aerial applications). Especially with aerial
applications the way the product is dispersed can make a difference for efficacy. In these
cases the competent authorities should require a field test.
Tests are done preferably during spring and beginning of summer. In autumn population
decline might be due to natural causes instead of the insecticide. Larvicides should
normally be tested in July-August when sufficient levels of Culex spp. and Aedes spp.
can be found. In any field trial, the assessment of efficacy requires pre- and post-
treatment assessments of the population. CDC light traps are one commonly used
method to trap mosquitoes and can provide both quantitative (how many mosquitoes)
and qualitative (which species are present) data. Other methods (exhauster, aspirator)
can be used too. Apply the insecticide according to the label instructions.
5.6.4.14.2.3 Requirements per type of claim

Per type of claim the requirements will be listed.

Products intended for use as general surface treatment, space treatment or vaporisers in houses:
- a laboratory test showing adult mortality;
- a simulated-use test showing mortality and knockdown and/or residual efficacy according to the claim.

Products intended for use as larvicides:
- a laboratory test showing larva mortality;
- a simulated-use test showing decrease in number of emerging mosquitoes;
- depending on the claim (mandatory for use in natural waters) field test showing larval mortality or decrease in number of emerging mosquitoes.

Products intended for use as repellent on skin or clothes:
- a simulated-use test (arm-in-cage) showing repellence;
- a field study showing repellence in the field.

Products intended for use as repellent not on skin or clothes:
- a laboratory and/or simulated-use test showing repellence;
- depending on the claim field test showing repellence.

Products intended for use as repellent on horses (recreational and/or sport horses):
- a laboratory test demonstrating repellence;
- a simulated use/field test demonstrating repellence against the specific target mosquito species on target animals.

Simulated-use tests can be waived if a robust field trial is submitted.

5.6.4.14.3 Assessment of authorisation

5.6.4.14.3.1 Norms and criteria

A biocidal product may only be authorised if it "possesses a sufficient level of efficacy". An insecticide against mosquitoes is considered to be sufficiently "effective" if the following results can be achieved:

Products intended for use as general surface treatment, space treatment or vaporisers in houses:
- required results in simulated-use tests:
  - the level of knockdown efficacy should be >80%;
  - mortality after 24 hour should be >90%.

Products intended for use as larvicides:
- required results in laboratory tests:
  - 100% mortality after 24 hours of contact is usually required. For slow acting insecticide 100% mortality after 48, 72 hours or more could be considered. Exceptionally a larval mortality >90% can be acceptable if all the surviving larvae died before or during emergence;
- required results in simulated-use or field tests:
  - >90% larva mortality;
  - showing decrease in number (usually 80%) of emerging mosquitoes.

Products intended for use as repellent on skin or clothes:
- required results in simulated-use test:
  - during the claimed protection period the protection should be ~100% (i.e. period to the second bite or the first confirmed bite is the claimed period);
if the claimed protection is less restrictions should be placed on these products preventing marketing as a way to prevent disease transmission.

Products intended for use as repellent not on skin or clothes:
- a laboratory and/or simulated-use test showing repellence;
- depending on the claim field test showing repellence (e.g. ~80% for repellents that are dispensed to protect an outdoor space (vaporisers, coils, etc).

Products intended for use as repellent on horses (recreational and/or sport horses):
- a laboratory test demonstrating repellence;
- a simulated use or field test demonstrating sufficient repellency over the time period claimed, preferably 90% or provision of data that allow calculation of the ‘complete protection time’, i.e. the time till the first confirmed bite/landing.

Deviations from these norms is possible but should be justified in the application.

5.6.4.15 Wasps

5.6.4.15.1 Introduction

There are two types of wasp control: control of the wasps’ nest and control of single flying wasps entering a home. The control of wasps’ nests may be performed both indoors (in cavity walls or attics), as well as outdoors (in trees, under roof gutters).

5.6.4.15.1.1 Biology

The major pest wasps (Hymenoptera) are the social wasps in the family Vespidae. Yellow-jackets (\((Para)\)Vespula spp., Dolichovespula spp.), paper wasps (Polistes spp.), and hornets (Vespa spp.) all belong to this family and are the greatest pests to homeowners. Wasps can be easily differentiated from bees by the fact that a wasp’s body appears to be hairless and their hind legs thinner than a bees.

The vespid or social wasp lives in colonies in nests built of a paper-like material. Each nest is begun in the spring by a single queen who has mated the previous autumn. The queen builds a small nest in which she begins to lay eggs. It is only non-fertile female worker wasps that emerge from these initial eggs. These workers take over the nest building duties and forage for food to feed the larvae that emerge from subsequent eggs. Some of these eggs are fertile females and some are males.

Mature colonies are divided into a social order consisting of the queen, workers, males, and fertile females. In the autumn, the males and newly produced queens leave the nest to mate. The male’s sole purpose is to inseminate the fertile females, which will become next year’s queens. The newly inseminated queens will then find a sheltered place where they will hibernate to begin the cycle with building a new nest the following spring.

Unprovoked, wasps are not aggressive stingers but will protect themselves and their nests making them an undesirable occupant of properties and buildings. Wasps commonly infiltrate in and around homes in search of nest sites and areas to hibernate causing problems for the homeowner. Some people are allergic to wasp venom, and can have life-threatening allergic reactions. Unlike bees, wasps can sting repeatedly.

For effective control of wasps, the entire wasps’ nest should be treated. The control is aimed at exterminating all wasps that are within the nest that can fly. If this is achieved, the eggs and larvae that are still present cannot be taken care of and fed anymore, resulting in the elimination of the entire nest.

5.6.4.15.2 Dossier requirements

A clear label claim should be submitted. The study results of field trials should demonstrate the efficacy of the product based on the submitted label claim.

Laboratory and field trials with the test insects are needed to assess the efficacy of the product. Ideally, the studies should be performed according to established guidelines
where these are available. These may be international, EU or national guidelines. See Appendix 17 for a list of available guidelines. If there are no guidelines available or the guidelines are not suitable, the applicant may use their own methods (intra-company Standard Operating Procedures), on condition however, that the study is scientifically robust, well reported and provides a clear answer to the question. In addition, the test methods applied and the test conditions should be clearly and fully described and must address the efficacy claim that appears on the product label. A control treatment without biocide should be included in all laboratory trials.

In the case of field trials where true replication is almost certainly impossible to achieve and where normal control methods are not restricted to use of a single insecticidal product, a full description of any factors that might be expected to influence product performance should be given. These are intended to provide the authorities with information to assist with the interpretation of the results obtained.

5.6.4.15.2.1 Test species

A product for use against wasps should be tested on colonies and/or workers of *Vespula* spp. or *Dolichovespula* spp.

5.6.4.15.2.2 Laboratory simulated use tests and field studies

For the evaluation of biocides against wasps different types of laboratory and field test can be used. Examples of test are listed below.

**Laboratory tests**

Wasps can be tested in the laboratory in small jars or Petri dishes. The individual wasps should have sufficient access to food (e.g. sugar solution), since they can starve to death within hours when isolated from their nest without food. The surface can be treated, after which insects can be added at different time intervals. Alternatively, the wasps can be sprayed directly. Concentrations used must be in accordance with the claim. The knockdown percentages and/or mortality and/or residual effect are determined.

A control treatment without biocide with a similar number of replications should be included in all laboratory trials.

**Repellents/attractants**

For products with a repellent or attracting effect against wasps no agreed protocols are available. The tests should be designed to mimic the practical use situation. The study results should provide a clear picture of the efficacy of the product. Methods should be described well. The submitted data from studies are checked for completeness, based on the applied dose per treated area. It is also checked whether the duration of exposure is sufficient. If the formulation alone i.e. without the carrier (e.g. a product with a tissue as carrier) has been tested, data on release from the carrier are also required.

**Field trials**

Insecticides with a claim to kill wasps’ nests should be tested in a field trial. The efficacy of the product should be tested in at least 5 nests. Depending on the label claim different nests (locations) should be tested (e.g. free hanging in trees or on buildings, hidden in the soil or in wall cavities, etc.). A few like size nests should be monitored over the same test period as untreated controls. A pre-treatment activity count should be taken over a pre-determined time interval of both treated and untreated nests. A well-established parameter for wasp colony activity is the traffic rate, which is defined as the number of wasps entering and leaving the colony in a given time. The traffic rate can be determined 7 days before the treatment for at least 5 minutes at two different times of day as well as on the day of treatment in order get a picture of the colony activity and development. The time interval between both observations must be at least 2 h. Treatment should be consistent with label instructions. When the nest is visible it can be treated directly. In some cases the nest is hidden, for instance in between walls or
ceiling of houses. In those cases normally all the openings through which the wasps enter the space in which the nest is hidden should be treated. Nest position, number of entrances as well as wasp species must be described.

After 24 hours, one week and two weeks post-treatment the activity or lack thereof should be recorded by determination of the traffic rate at the treated and untreated nests. The check after one and two weeks is required since it is possible that, when pupae are not eliminated, wasps emerging from pupae can take over the duties of feeding the larvae.

**5.6.4.15.2.3 Requirements per type of claim**

**Products intended for the control of the wasps’ nest:**
- field trial with at least 5 treated nests.

**Products intended for the control of flying wasps:**
- laboratory or simulated-use test.

**Products intended for repelling wasps**
- simulated-use or field trials.

**5.6.4.15.3 Assessment of authorisation**

**5.6.4.15.3.1 Norms and criteria**

A biocidal product may only be authorised if it “possesses a sufficient level of efficacy”. For wasps this is implemented in the following way.

**Products intended for the control of the wasps’ nest:**
- required results in a field test:
  - in 80% of the treated nests mortality of the flying wasps should be 100% within 24 hours and all of the treated nests must have 100% mortality (i.e. no visible signs of nest activity) after one and two weeks.

**Products intended for the control of flying wasps:**
- required results in a laboratory or simulated-use test:
  - ≥ 90% knockdown within a 5 -10 minutes after contact with the product (or according to the claim), direct after spray and at the end of the residual period;
  - mortality according to the label claim, preferably 90% in 1 hour.

**Products intended for repelling wasps**
- required results in a simulated-use or field test:
  - a simulated-use test showing repellence;
  - depending on the claim field test showing repellence.

**5.6.5 PT19 Repellents & Attractants (non-arthropods)**

Please refer to the General sections 1-3 of this guidance and the TNsG.

EPPO guidelines 199 and 200 are available for efficacy testing of rodent repellents intended for plant protection. These might be modified for biocidal use.

**5.6.6 PT20 Other vertebrates**

Please refer to the General sections 1-3 of this guidance and the TNsG.
5.7 Other biocidal products (Main group 4)

5.7.1 PT21 Antifouling products

5.7.1.1 General Introduction

This section deals with the methodology for the evaluation of efficacy tests for antifouling products that is applicable for the authorisation of products under the EU Biocidal Products Regulation (BPR, Regulation (EU) 528/2012).

5.7.1.1.1 Introduction

This chapter describes the nature and extent of data which should be available to support the label claims for biocidal products within Product Type 21 - Antifouling Products. These are defined in the BPR as “Products used to control the growth and settlement of fouling organisms (microbes and higher forms of plant or animal species) on vessels, aquaculture equipment, or other structures used in water”.

5.7.1.1.2 Types of Coating

The antifouling products currently available can be categorised into the following broad coating types:

- Soluble matrix
- Insoluble matrix
- Self polishing

The categorisation of coating types outlined above is general. It should be noted that some antifouling products do not necessarily rely on one single coating technology and combinations of different technologies have been developed by antifouling formulators to suit customer specifications and environmental requirements. A description of the main coating types can be found in Appendix 19.

It should be noted that the protection periods described in the appendix for each coating type are typical life times that may be achieved by using products within these very broad groups. The efficacy of an antifouling coating will heavily depend upon use, for instance a vessel's operational pattern (such as dry-docking interval, sailing speed, and idle times as well as the temperature, fouling intensity, and other environmental characteristics where the vessel is trading). It also depends on the extent to which the antifouling paint specification has been tailored to meet these specific conditions. Surface preparation, primers, quality of work, dry film thickness, etc. may also affect the quality and/or duration of the protection.

5.7.1.1.3 Mode of Action

Antifouling products form paint films that act as release vehicles for the active substance(s) contained in the paints. The active substance(s) will be released over the specified lifetime of the products, creating a microlayer of biocide rich water at the paint surface. Here, in this water microlayer, the concentration should be sufficient to deter the settlement and/or growth of fouling organisms. A more detailed description of the respective modes of action and physical characteristics of the various coating types are outlined in Appendix 19 of this document.

5.7.1.1.4 Categorisation of antifouling products

Antifouling paints are made available for different use types. Typically they are prescribed for yachts, commercial vessels (such as bulk carriers, tankers, container ships, car carriers, passenger ships, etc.), and aquaculture.

The three broad categories of products (in Appendix 19) can be defined by the way in which the products control the release the active substance(s). Given the fact that a
single active substance may not have a sufficiently broad spectrum activity to control the wide range of fouling organisms, antifouling products often contain more than one active substance.

### 5.7.1.1.5 Spectrum of activity

Target organisms belong to very different taxonomic groups. There are many organisms that can live within a fouling community, but only a few cause severe fouling problems. Which organisms will present a problem depends on the local conditions and the operation of the individual vessel. For example, typical target organisms in European waters may include, but are not limited to, various species of the following genus: 

- *Pseudomonas* (light slime),
- *Amphora* (dense slime),
- *Ulva* (macro-algae), and
- *Semibalanus* (animals).

Fouling organisms and growth rates differ between tropical and temperate regions. The fouling intensity and the species that dominate a fouling community may vary locally and seasonally. While it is not normally feasible to claim efficacy against specific target organisms, applicants may choose to supplement their label claim that the product is an "antifouling product" with an indication as to whether the product will be effective against one or more of the following fouling groups:

- Slime
- Weed (macro-algae)
- Animals

### 5.7.1.1.6 Dossier requirements

The following aspects are required for the efficacy evaluation of antifouling products:

1. The label claims and instructions for use including the technical data sheet
2. Efficacy data on the product

### 5.7.1.1.7 Label claims

For each product a set of label claims should be provided as part of the dossier submitted. Claims for the activity of the product include those made on a technical data sheet or other associated documentation, as well as those on the label itself. To simplify the text, only the term 'label claim' will be used below.

In general the claim for antifouling products can be rather unspecific, for instance 'antifouling product for professional application'. The label should also indicate to which fouling groups (see 5.7.1.1.5) the product is effective and whether it can be used in marine or fresh water.

The label claim for products used in areas other than on vessels, such as products used for aquaculture, in the inlet and outflow pipes of cooling systems, or for other "non-vessel" uses should be more precise, and clearly describe purposes for which the product can be used.

According to Article 69(2)(f) of the BPR the label must clearly and indelibly show the uses for which a biocidal product is authorised.

### 5.7.1.1.7.1 Areas Of Use

The product label, technical data sheet or other associated documentation should contain information on the main use categories for the product, for example use on vessels and larger boats, yachts, stationary installations, or aquaculture equipment, etc. This will normally also include information on whether the product is intended (primarily or exclusively) for use in either marine or fresh water.

As the fouling challenge is more severe under static conditions, installations and recreational boats (which are normally tied up in marinas) will foul more quickly than
commercial vessels that spend most of their time in motion. Therefore, if a product is intended specifically for static or recreational use, this should be specified in the label claims.

(For human risk assessment purposes, it is important that a label claim specifies if a product is intended for amateur use or if is for application by professionals only.)

**5.7.1.1.7.2 Application method/dose rate**

Antifouling coatings may be applied using methods such as airless and conventional spray, brush and roller, or dipping and immersion (aquaculture). The specified total dry film thickness will vary depending on the intended dry-docking interval, activity of the vessel (such as sailing speed and idle times), and on the temperature, fouling intensity, and other environmental characteristics where the vessel is operating. Furthermore, larger vessels will normally have different antifouling products and different paint film thicknesses specified for different parts of the underwater hull depending on, for instance, water flow and light conditions. Some areas, such as those with less frequent maintenance intervals than those for the rest of the underwater hull, and those with strong water throughput (e.g. inside thrusters) may require higher film thicknesses to minimize the risk of transmigration of non-indigenous species in these areas.

It is important to note that the paint thickness does not affect the efficacy of a product, which will control fouling regardless of the thickness of the paint applied. Instead, the film thickness will define the in-service life of the product.

For antifouling paints there is no direct relationship between the applied dose (paint film thickness applied) and the efficacy of the product (unlike agrochemicals, for example, where applying more pesticide increases the concentration of the pesticide and therefore the magnitude of the controlling effect on the pest).

Recommended dry film thicknesses are given to ensure that enough paint is applied to the vessel to avoid the coating being ‘polished through’ during service, exposing the underlying anticorrosive paint which will be susceptible to fouling. When paint is applied by spray, more than one coat of paint is normally applied to protect against possible application defects, such as ‘pin holing’, where small areas of the anticorrosive are left exposed.

As the three major types of antifouling coatings (Appendix 19) vary in their ability to maintain a sufficient release of active(s), this is reflected in their different typical lifetimes.

**5.7.1.1.8 Efficacy tests**

**5.7.1.1.8.1 Laboratory tests (including in-vitro screening tests)**

Laboratory tests are typically conducted on a single active substance and with a limited number of test organisms, and may provide information about the specific action of a substance against a known fouling species. It is acknowledged that model target organisms may be used in these tests as well as those that may successfully be cultivated in a laboratory (e.g. juvenile barnacles). Consideration should be given to the use of species known to be critical fouling species.

Laboratory tests are routinely used to demonstrate efficacy of an individual active substance, often at a very early stage during research in order to screen new active substances.

Laboratory testing of individual paints is not undertaken as it is not considered to be a realistic evaluation of the product. Field testing is routinely undertaken instead (described below).
5.7.1.1.8.2 Simulated field tests (static raft testing)

These may be studies that are conducted with the candidate product or with the active substance(s) incorporated into a model coating type. Such tests involve the immersion of panels treated with the test coating on static rafts for a period of months or years at an appropriate location. For aquaculture products this could be nets or (sections of) cages treated with the test product and immersed at an appropriate site.

Efficacy data on antifouling coatings should normally be generated by testing over at least six months of peak fouling activity. As far as is practical the test location(s) should be representative of the intended uses of the product. When testing in locations with seasonal variation in fouling challenge, the test period should cover the full fouling season. The length of a season will vary depending on the location of the test site. When choosing the test location(s), factors such as shelter (from strong waves and ship traffic) and access have to be balanced against water exchange conditions and other characteristics determining whether the water at a site is representative for the end use conditions.

Since raft testing is carried out in natural environments, the same product may perform differently at the same site in different years. This variability in fouling intensity, and thus the test results, is due to weather conditions, availability of nutrients, and other uncontrollable factors that may affect the type and extent of fouling and its rate of settlement and growth. Therefore, a negative control (a surface which has no antifouling effect) should be included in all tests, which will indicate the degree of fouling that would be present under static conditions if the tested coatings were totally ineffective. A reference coating of proven or known efficacy (a positive control) may also be used. The absolute amount of fouling present on a test coating may not be reproducible at the same site from year to year.

Efficacy studies include regular assessments of fouling throughout the period. These assessments usually describe the major types of fouling (e.g. slime, algae and other weeds, and barnacles or other fouling organisms), but describing these as to genus and species is unnecessary. As sharp edges on test panels may be difficult to protect, fouling that is not growing on the front of panels (i.e. attached along the edges) should be disregarded.

The presentation of data should include the assessment method (the rating/scoring for the test panels and how these are interpreted), together with photographs and/or diagrams of the test panels.

5.7.1.1.8.3 Field tests/In-service monitoring

Since field tests involve long-term exposure to practical conditions, they can be regarded as in-service tests. Field tests permit antifouling products to be tested under similar operating conditions and stresses as those encountered when the antifouling product are in service. Possible examples of these tests include:

- Panel tests where coated panels have been attached to a vessel during parts of or during a complete dry-docking interval
- Patch tests where vessels have been painted with the test coating as a strip or patch on the hull
- In-service monitoring of aquaculture nets, cages, etc.

Any field data generated in support of an application should be conducted on the candidate product or representative products that closely resemble the fully formulated commercial product. A robust justification should be provided to support bridging of data from a similar (but not identical) product.

It is recognised that it may not be possible to run concurrent untreated panels or patches during field trials. Therefore information on the performance of the main
antifouling coating over the test period should be provided instead. Monitoring reports of the performance of an antifouling product on a fully treated vessel may also be submitted, where these are available. It is also recognised that data generation from field trials may require many years to carry out and are more likely to be available for well-known technologies than for products containing new active substances (or new combinations or concentrations of active substances) or for coating types based on new technologies.

Where field data are not available, the applicant has the option to provide data on other existing formulation(s) where appropriate, and read across to the current application through scientific reasoned cases and arguments. Such arguments may include:

- The composition of the 'old' (and well documented) and the 'new' antifouling product
- Simulated field tests of the ‘old’ and the ‘new’ antifouling product
- Possible field data on the ‘old’ antifouling formulations
- Further justification, such as why bridging is appropriate (e.g. in-service monitoring)

It is understood that extensive field data or bridging data may not be available when established biocides have been introduced into products based on new technology or new active substances are being developed. Field tests from different ships have limited value for the purpose of comparing efficacy due to the diversity of operational patterns and trading routes and the likeliness for unforeseen circumstances or incidents not recorded. This, together with the complexity with respect to application and monitoring and the long exposure times required, explain why in-service tests are normally not available for new antifouling products. However, when data on in-service/field tests are available, these should be submitted as additional information.

However, field data are required at renewal of a product authorisation, as the product will have been on the market for several years by this point. Further guidance on how to perform and assess these data will be developed in the future and incorporated into this guidance.

5.7.1.1.8.4 Replication of efficacy tests

Antifouling paints are normally tested in series during product development, where panels treated with a range of formulations, with only small variations between them, are tested to assess the effects of exposure on other paint properties, as well looking at the efficacy of the formulations.

Since the testing takes place in a natural environment, the variation in fouling propagation and intensity between different years at the same test site will vary. A variable natural environment, the differences in fouling activity between years, and the criteria for establishing efficacy (the general nature of a label claim) make very detailed evaluations unnecessary.

However, to increase the scientific rigour of the evaluation, the results of three replicate plates should be submitted.

It is acknowledged that it is not common practice to test multiple replicates of individual formulations, however panels treated with similar formulations containing the same combination and concentration of active substances may be considered replicates when these are supported by a suitably robust reasoned case explaining the relevance of these formulations to the candidate product. The results from such panels should be submitted, along with details of the formulations used, as well as the reasoned case.
5.7.1.1.9 Standard test methods
5.7.1.1.9.1 Simulated use test methods

The standard test methods available for the generation of simulated field data through raft testing of antifouling coatings are:

1. Efficacy evaluation of antifouling products. Conduct and reporting of antifouling efficacy evaluation trials. CEPE Antifouling Working Group, June 2012. This methodology has also been adopted by the International Paint and Printing Ink Council – IPPIC and presented at Technical Meeting I 2013 PT 21 efficacy workshop (Appendix 20).


Reports based on both the above methods should be accepted.

However, it should be noted that the ASTM methods were primarily developed to satisfy the detailed requirements of the US Navy and are not commonly used by the general antifouling industry. The main reasons for this are that they are resource intensive (in terms of the level of detail required in both the materials used as well as the analysis and reporting of the fouling species [including the number and diameter of individual organisms), thereby exceeding the requirements for substantiating a general product label claim (since normally specify only the general types of fouling and their extent are reported for regulatory purposes]) and that they specify relatively dated materials (paints), for which better and more applicable alternatives are available. Notwithstanding, the methods may provide a good basis for biological research.

5.7.1.1.9.2 Field/In-service tests

There are currently no national or international standards that cover field evaluation of antifouling products. Field tests (application on ships) are rarely used to screen formulations and establish the basis for an efficacy claim since they are time consuming and costly and since the results are heavily dependent upon the operations of individual vessels. To the extent field trials are used, their purpose is normally to determine relative differences in efficacy between already commercial formulations during different use conditions (such as vessel speed, idle times, etc.).

Typically a new antifouling paint represents an incremental improvement or an adaptation to a specific user requirement. Normally, therefore, the experience from similar commercial products will contribute to the confidence the manufacturer has with respect to the efficacy of a new product.

However, at the point of renewal of a product authorisation, a product will have been on the market for several years and field data should be generated to demonstrate the actual performance of the product in use.

5.7.1.1.10 Resistance

Resistance is discussed in the general part of the TNsG on Product Evaluation in Chapter 6. A review of resistance is part of the evaluation at product authorisation. If new information is available which was not reviewed during the approval of active substance, this information should be provided at the time of product authorisation.

In general development of resistance is not to be expected for marine use, as ships are treated with several antifouling paint products containing different active substances. However, this may not be the case for use in fresh water and aquaculture.
5.7.1.11 Reports of development of resistance should always be mentioned.

**Service life**

Amateur antifouling products for recreational crafts are normally claimed to last for one yachting season, and are recommended to be retreated annually. Commercial vessels will have extensive tailor made paint specifications depending on their dry-docking interval and operational pattern. Different products and film thicknesses are frequently used at different parts of the vessel due to different light conditions and hydrodynamic forces. In the case where a label claim includes different types of use (e.g. both vessels and static installations), the corresponding protection times may differ.

With respect to the ability of fouling organisms to settle and attach, static conditions are much more favourable than the conditions on vessels that are only idle for relatively short periods at the time. This together with the greatest levels of marine growth occurring in near shore conditions (as described in 2.1), explain why static raft testing is a worst case test. For recreational craft, however, the use conditions may be very different. Therefore, tests are frequently carried out for the same number of fouling seasons as the recommended use.

It is not obligatory to state on the label what the service life of a product will be.

5.7.1.2 Products intended for marine use

5.7.1.2.1 Introduction

Raft tests represent worst case conditions with respect to fouling intensity due to their static nature and because the tests are carried out in near shore environments. As the release of active substances from antifouling paints is assisted by hydrodynamic forces (i.e. through polishing), fouling will be more severe on static surfaces compared with moving boats and ships.

Coastal waters are known to have the highest fouling intensity. The littoral zone along coasts constitutes a tiny part of the world’s oceans, but contributes markedly to the total marine production. The reason is that benthic production (per unit surface area) exceeds pelagic production by a factor of ten. Coastal macrophytes account for two-thirds of the total biomass of marine photo-synthetic organisms although they can only inhabit less than 0.5% of the surface area of the oceans\(^\text{37}\). Therefore, when efficacy is demonstrated in coastal waters (the worst case situation), a product is also assumed to be effective in open sea and brackish conditions, and the data can be used to support these uses.

5.7.1.2.2 Dossier requirements

A report of the results from efficacy testing may also include the following about the test site, the test procedures, and the data reported:

- Method of application and information on the panel type and panel preparation;
- Location, geography, and water exchange conditions;
- Water temperature and salinity, including seasonal variations;
- Orientation, dimensions, and exposure depth of the test surface;
- Dimensions and type of material of test panels;
- Identity of the tested product and the control(s);
- Details on the panel preparation (application technique, possible primer paint, paint film thickness, number of coats);
- Date and duration of test;
- Date and raw data from each individual assessment of a test panel;

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• Photos of test panel and control(s);
• The overall fouling assessment rating at each inspection during the exposure period;
• A description of the reporting company’s weighting system used to provide the overall fouling assessment rating. This should include how fouling coverage has been weighted in order to provide an overall efficacy assessment. The description should be transparent and explicitly explain the calculations carried out. (See example in Appendix 21);
• An interpretation of the data including a conclusion and a discussion of the validity of the results relative to the unprotected reference and the label claim for the product tested.

5.7.1.2.2.1 Testing and field trials

The recommended method for demonstrating efficacy of marine antifouling products is static raft testing. Raft testing allows a high number of formulations to be tested at worst case conditions.

At least one raft test in European coastal waters should be provided. Test in Atlantic or Northern European Seas are preferred; however, other European waters are acceptable too. It is preferable to also provide the reports from additional tests, although these additional tests can be performed in other locations (e.g. in Europe or elsewhere in the world). At least three replicate panels should be provided per product (see section 5.7.1.1.8.4 for more information on replication of tests). Tests should be performed for at least one fouling season, which is at least six months covering the period of peak fouling activity.

5.7.1.2.3 Assessment of authorisation

The ability a product has to produce an antifouling effect is determined by a combination of the activity of the active substance(s) and the mechanical/physico-chemical properties of the paint. Parameters that will define the efficacy of an antifouling product include:

• The potency and release rate of the active substance(s)
• Operational patterns (e.g. speed, idle times, dry-docking interval, etc.)
• Physico-chemical conditions of the water and other climatic, seasonal, or local factors affecting fouling intensity (e.g. concentration of nutrients, hours of daylight, salinity, temperature, presence of ice, turbidity, etc.)

The efficacy data submitted in support of an application represent part of the information assessed to establish if the product has the claimed level of efficacy. It is recognised that the actual in-service performance of an antifouling product will be dependent on a range of factors, which may include how and where a boat or vessel is operated, seasonal and annual variations, as well as the specifics of the antifouling coating itself. Commercial vessels receive tailor-made product specifications in order to meet various planned (and unforeseen) operational conditions. Thus, the general efficacy of a product under typical fouling conditions according to criteria in paragraph 2.3.1 should be demonstrated.

5.7.1.2.3.1 Norms and criteria

The purpose of an efficacy test is to support the label claim. Efficacy is evaluated by comparing the extent of fouling on the test substrate with the fouling on a similar, but unprotected substrate which has been exposed simultaneously and at the same site.

Fouling coverage is frequently evaluated based on the coverage of the typical marine fouling species such as slimes, algae and animals (barnacles, mussels, etc.).

The three types of fouling species (slime, macro-algae and animals) may be rated differently when merged to an overall fouling assessment for the tested product since slime fouling is less significant compared to macro-fouling (for instance for the fuel
consumption and manoeuvrability of a ship). An overall fouling assessment may describe the efficacy of a panel in categories such as for instance: 'Excellent', 'Good', 'Fair', and 'Poor'. An example to illustrate how the coverage of the main categories of fouling may be combined to provide an overall fouling assessment is given in Appendix 21.

Since different companies may use different overall fouling assessment systems and interpretation of the result may vary with the type of product (what is 'poor' efficacy for marine water vessels might be 'good' for fresh water yachts), these ratings are not used as the pass/fail criterion for authorisation. Instead, the percentage fouling on the control and test panels is used.

Normally, when tested in marine waters, the negative control will have at least 75 % fouling coverage at the end of the test. In this case, the result from a product under test should be acceptable if the coverage of macro-fouling on the panels is below 25 %. Macro-fouling is defined as large, distinct multicellular organisms visible to the human eye such as barnacles, tubeworms, or fronds of algae. Algae shorter than 5 mm should be regarded as micro-fouling, together with slimes.

If the 25 % criterion is not met, a justification should be provided for why the product may still be regarded as sufficiently efficacious for the intended use.

5.7.1.3 Products for freshwater use

5.7.1.3.1 Introduction

Fresh and brackish waters are known to represent a less severe fouling challenge compared to marine waters. Effective antifouling protection may be environmentally important even where the general fouling challenge is low. For example, to reduce the risk of translocating invasive species (such as zebra mussels) into or between inland waterways, lakes, or brackish seas.

5.7.1.3.2 Dossier requirements

See 5.7.1.2 for the requirements on reporting the test procedure and data.

5.7.1.3.2.1 Testing and field trials

For products intended for use in both fresh water and marine waters, a raft test in marine coastal water is sufficient and a separate efficacy test under fresh water conditions is not normally carried out for. Since fresh and brackish waters are known to represent a less severe fouling challenge compared to marine waters, it is common practice to use the bridging principle and refer to tests conducted in marine waters.

For products only intended to be used in fresh water, at least one raft test in fresh water should be provided. When raft tests are carried out in fresh water, the test site should be one known to have relatively high fouling levels, preferably in an area where zebra mussels are present. However, it is preferable to also provide the reports from additional tests. At least three replicate panels should be provided per product (see section 5.7.1.1.8.4 for more information on replication of tests). Tests should be performed for at least one fouling season, which is at least six months covering the period of peak fouling activity.

5.7.1.3.3 Assessment of authorisation

See section 5.7.1.2.3.

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38 IMO’s 2011 Guidelines for the Control and Management of Ship’s Biofouling to Minimize the Transfer of Invasive Aquatic Species, Section 2.1. Definitions.
5.7.1.3.3.1 Norms and criteria

The purpose of an efficacy test is to support the label claim. Efficacy is evaluated by comparing the extent of fouling on the test substrate with the fouling on a similar, but unprotected substrate which has been exposed simultaneously and at the same site.

In the case that an efficacy test is carried out in fresh water, it should be noted that as the fouling challenge is low, a 75 % or more coverage of fouling organisms on a negative control test panel cannot be expected. Therefore, if a test is carried out where micro-fouling is predominant and the coverage of macro-fouling is less than 75 %, the test may still be valid. In the case where less than 75 % of the surface of the negative control is covered with fouling, an explanation should be provided for why the test should be considered valid.

It is also possible that in freshwater, macro-fouling (such as freshwater hydrozoans or zebra mussels) may completely cover a negative control.

For tests in fresh water where the control panel has 75 % or more coverage of fouling organisms, the result from a product under test should be considered acceptable if the coverage of macro-fouling on the panels is below 25 %.

For tests in marine water see Section 5.7.1.2.3 for criteria.

5.7.1.4 Products for use in aquaculture

5.7.1.4.1 Introduction

In aquaculture use, antifouling products are used to treat infrastructure, including immersed structures such as cages, nets, ropes, buoys and pontoons, as well as equipment such as pipelines, pumps, filters, and holding tanks.

5.7.1.4.2 Dossier requirements

See 5.7.1.2.2 for the requirements on reporting the test procedure and data.

5.7.1.4.2.1 Testing and field trials

Relevant field or simulated use trials should be provided to demonstrate the efficacy under in-use conditions. Static testing closely resembles real life conditions for aquaculture use. Test surfaces may include panels and net/cage samples suspended securely from the raft.

At least one field test should be provided. However, it is preferable to also provide the reports from additional tests. At least three replicates should be provided per product (see section 5.7.1.1.8.4 for more information of replication of tests). Tests should be performed for at least one fouling season, which is at least six months covering the period of peak fouling activity.

5.7.1.4.3 Assessment of authorisation

The ability a product has to produce an antifouling effect is governed by mechanical and physico-chemical properties of the paint. Relevant parameters to be taken into account when assessing the efficacy of an antifouling product include:

- The potency and release rate of the active substance(s) in the paint
- Physico-chemical conditions of the water and other climatic, seasonal or local factors affecting fouling intensity (e.g. concentration of nutrients, hours of daylight, salinity, temperature, presence of ice, turbidity, etc.)

A report of results from efficacy testing should include the following information about the test site, the test procedures, and the data reported:

- Method of application (e.g. dipping of nets) and type of test substrate
- Location, geography, and water exchange conditions
- Water temperature and salinity
• Orientation, dimensions, exposure depth of test surface, and date and duration of the test
• The extent and main categories of fouling and an interpretation of this relative to an unprotected surface and the label claim for the product tested

5.7.1.4.3.1 Norms and criteria

The purpose of an efficacy test is to defend the label claim. Efficacy is evaluated by comparing the extent of fouling on the test substrate (panel, cage, net, etc.) with the fouling on a similar, but unprotected substrate which has been exposed simultaneously and at the same site. Efficacy is demonstrated if fouling on the treated surface is considerably reduced compared to the fouling on the unprotected surface.

Fouling coverage is frequently evaluated based on the coverage of typical fouling species. These ratings are then merged to provide a consolidated figure for the three major types of fouling species: slime, macro-algae and animals (Appendix 21, Table 2). The three types may be rated differently when combined to an overall fouling assessment for the tested product. For example, slime fouling is less significant compared to macro-algae and large hard animals for the water exchange through nets and cages.

If a product for aquaculture use is tested on panels, the pass/fail criteria for the test may be the same as in paragraph 5.7.1.2.3.

5.7.2 PT22 Embalming and taxidermist fluids

5.7.2.1 General introduction

Annex V of BPR defines Product Type 22 products as follows: "Embalming and taxidermist fluids. Products used for the disinfection and preservation of human or animal corpses, or parts thereof". Embalming for this purpose only aims at the temporary preservation of the deceased person, before burial. Taxidermy fluids and those intended for long-term preservation (e.g. repatriation as shipping cases) are not covered by this guidance document. These particular cases will be taken into account in a future update and inclusion into Volume II Part B of the new BPR guidance structure.

This guidance document is intended for applicants to assist them in compiling an authorisation request dossier regarding the efficacy aspect, and thus specifies the general conditions for carrying out efficacy assessments of biocidal products for marketing authorisations.

This guidance document may be reviewed in the event of regulatory changes or technical advances.

5.7.2.2 Use of the products

5.7.2.2.1 The issue of bodily decomposition

5.7.2.2.1.1 Physical, chemical and microbiological post-mortem activities

A body starts to decompose as soon as the blood ceases to circulate and oxygen is no longer supplied to the tissues. Under conditions favourable to decay, the body cools in the first few hours after death, dehydration sets in (lividity) together with rigor mortis resulting from anaerobic hydrolysis of muscle glycogen. The first stages of cell degradation can be seen with the onset of lividity.

The natural degradation of the body's organic matter results from the action of enzyme, tissue and microbial processes. The ecosystem whose characteristics determine the succession of physical, chemical and microbiological changes that occur post mortem can be defined as the set of interactions between ambient factors (temperature,
hygrometry), individual factors, especially the body's water, muscle and fat composition, and the body's own microbial flora, both external (skin) and internal (digestive and respiratory). Together, these conditions affect the establishment, acclimatisation and development of the dominant indigenous flora, separately or in association, and thus steer the metabolism towards either speed or slow decomposition.

The activity of the microflora, initially latent, intensifies; the first stages of mineralisation of the organic matter, stages of the nitrogen, carbon, oxygen and hydrogen cycles, constitute both superficial and profound decomposition. This decay is defined partly by the decomposition of the organic tissues, mainly under the influence of the bacteria hosted by the individual, especially those in the intestinal flora, and then by fungi, and partly by the decomposition of the organic matter and the bacteria responsible for mineralisation that gradually invade the body, via the body fluids.

As the proteins, lipids and certain carbohydrates that provide the substrate degrade; they produce malodorous soluble and gaseous substances, containing sulphur, nitrogen and carboxylates. Depending on the specific activities developed by the flora in place, the resulting foul odours can vary in nature and intensity. It is increased by higher temperatures and by interference between chemical groups. As degradation progresses, the source of foul odours moves gradually from the body itself to the fluid products of decay, which rapidly become the principal source of foul odours. As the organic matter becomes hydrolysed into more soluble compounds it becomes easier for microorganisms to assimilate them, facilitating the production of foul odours.

5.7.2.2.1.2 The microorganisms involved

- in the early stages of decomposition of the liquids and soft tissues (with production of gases), only the following species are found: *Pseudomonas fluorescens* and *Micrococcus ureae*;
- at a later stage of lipid transformation, the following appear: *Pseudomonas* sp. and then *Pyogenes* sp.

The initial wave consists of aerobic bacteria while those following are anaerobic (*Diplococcus magnus*, *Streptococcus* sp., *Serratia liquefaciens*, *Bacteriodes* sp. etc.). This decomposition of the body due to bacteria and saprotrophic fungi gradually leads to autolysis of the remains, which is pursued later and over time by the bacteria active in the mineralisation of the organic matter, although this last stage is related to the level of humidity. Various factors concerning the environment of the body intervene (humidity, temperature, aeration) as well as its size, age, causes of death and place of storage.

The decay is predominantly influenced by the bacteria that had been hosted by the individual, especially those in the intestinal flora. The bacterial species frequently found in decomposing bodies are:

- of intestinal origin: enterobacteria, especially *Escherichia coli*; clostridia, especially *Clostridium tetani*, *C. welchii* and *C. difficile*; and faecal *Streptococcus*;
- of dermal origin: *Staphylococcus* spp.;
- of environmental origin: *Bacillus* spp.

The saprotrophic fungi and yeasts succeed one another in specific groups and the flora changes in line with the gradual alteration of the substrate, which thus provides a choice habitat for certain species of mycota at one moment and not at others.

The decomposition of the body due to bacteria and saprotrophic mycota accelerates the alteration started by autolysis, before the mineralising bacteria that invade the body later bring it into the cycle of waste material in the biosphere.

There may also be other pathogenic microorganisms, such as the tuberculosis bacillus (*Mycobacterium tuberculosis*) or other mycobacteria, or again viruses such as hepatitis or Human Immunodeficiency Virus (HIV), which can persist in the body.
5.7.2.2 Products for preserving human bodies and their uses

5.7.2.2.1 Types of application

The embalmer begins by physically working the limbs to reduce lividity and facilitate the flow of the preserving fluid. This is used for two separate purposes and at different concentrations:

- **arterial fluid:** an aqueous solution injected under pressure into the vascular system (the embalmer adjusts the final concentration to the condition of the body). This liquid is injected in the arterial system via the carotid or the femoral artery (sometimes at several points if diffusion is poor). The injection is made under pressure (by pump) or by gravity. This result in venous drainage: replaced by the injected product, the blood leaves the body via the jugular vein. Six to ten litres are injected and four litres (of blood and other body fluids) are removed by suction;

- **cavity fluids:** these are usually used at high concentration to preserve the thoracic and abdominal cavities, which cannot be irrigated by arterial injection. Using a trocar connected to a pump, about two litres of the pure undiluted solution are injected into the peritoneal cavity through an incision close to the navel.

There are also preparations for dermal use. These are gels designed to limit the decomposition of the body by treating bedsores. For this type of product, applicants must complete the appropriate section of the assessment grid, demonstrating the efficacy of the product.

In addition to its biocidal active substance(s), such a formulation could include the following co-formulants, which must have no biocidal activity:

- **anticoagulants:** to fluidify the product and ensure correct diffusion (sodium chloride and sodium citrate);

- **hydrating and moistening agents:** to slow the drying out of the body by hydrating the tissues and making them more supple (glycerine, ethylene glycol, propylene glycol, hexylene glycol, urea);

- **surface-active agents:** to facilitate adsorption of the fluid and penetration of the membranes and to maintain the solubility of the other components of the formulation, which are generally cations, as these surfactants are often also antimicrobials;

- **colouring agents:** to ensure that the fluid is of a colour similar to blood; synthetic colouring agents are generally used (eosin, erythrosine or food colouring agents);

- **perfumes.**

5.7.2.2.2 Products used for aesthetic purposes

Preservation may be supplemented with aesthetic treatment involving remodelling the face (modelling wax), sewing or bonding together the upper and lower jaws, placing eye caps under the eyelids to keep the eyes closed (or possibly gluing them shut). Finally, when all other treatment has been completed, cosmetic make-up may be applied, partly to give a more agreeable appearance but also partly to delay dehydration.

These products are not considered during assessment of the efficacy of the preservation product. However, if these products contain substantial amount of active substance and claim an effect on bodily composition, they should be considered as biocide.

5.7.2.3 Data required

5.7.2.3.1 Claims and labelling

When an application for the approval of a PT 22 substance is being assessed, the evaluation of the efficacy is focused on the efficacy of the biocidal product and not on the
other products (as cosmetic) which can be also included in an embalming treatment, so this aspect must be demonstrated unambiguously in laboratory tests and tests on human bodies, the details of which must be available on request.

As a minimum, a PT 22 product must claim to be active against a broad spectrum of bacteria; yeasts, fungi and viruses are considered as an additional spectrum. As explained above, bacteria are the principal microorganisms targeted by PT 22 products. Yeast, fungi and viruses have less relevance in the early stages of bodily decomposition.

Nonetheless, an active substance with a broad spectrum on different types of microorganism would provide better protection for users (e.g. against tuberculosis bacilli, hepatitis viruses or HIV, etc.).

5.7.2.3.2 Efficacy tests

5.7.2.3.2.1 Laboratory tests

As there is currently no standardised method recognised at European level targeting the scope covered by PT 22 products, and as no technical reference documents were found either in France or throughout the world, it is important that methods used should achieve two different yet complementary goals:

- the rapid destruction of bacteria, representative of the bacterial sphere, in the presence of a strongly interfering organic load simulating the bodily fluids;
- to maintain this antibacterial activity for several days, thus demonstrating that there is no subsequent proliferation of these microorganisms.

5.7.2.3.2.2 Determining bactericidal activity

As already mentioned above, the minimum claim is a bactericidal activity. Other additional activities, such as fungicide or virucide activities must be supported by relevant tests.

From among the techniques available, the selection was made based on the following criteria:

- a method that has been standardised at least at European level – the bacterial "suspension" test used in the medical sector
- the presence of a standardised strong organic load accurately simulating organic bodily fluids.

In compliance with the classification of European standards (EN 14885), the two tests selected belong to the categories of tests in Phase 2, Step 1 which include quantitative suspension tests for establishing that a biocidal product has a bactericidal activity by simulating its use under real conditions:

a) tests according to the EN 13727 standard: this mandatory test determines the minimum bactericidal concentration of a product on the basis of a 5-log reduction in titre of a bacterial suspension, at a temperature of 20°C, for 60 minutes of contact, in the presence of a strong organic load (bovine albumin 3 g/L + ovine erythrocytes 3 ml/L), on three species of bacteria (Staphylococcus aureus ATCC 6538, Pseudomonas aeruginosa ATCC 15442, Enterococcus hirae ATCC 10541);

b) tests according to the EN 14348 standard: this additional test must be taken into account if the applicant advances any claim concerning activity against agents responsible for tuberculosis, or if complementary tests prove necessary to cover this particular need. This test has a methodology similar to that for the previous test, determining the minimum tuberculocidal concentration of a product on the basis of a 4-log reduction in titre of a bacterial suspension, at a temperature of 20°C, for 60 minutes of contact, in the presence of a strong organic load (bovine albumin 3 g/L + ovine erythrocytes 3 ml/L), on the bacterium Mycobacterium terrae ATCC 15755.
Any claim by applicants that a product targets a specific microorganism must be supported by supplementary studies. For example, a claim of activity against the agents responsible for tuberculosis must be verified in compliance with the EN 14348 standard. If there is no recognised standard for a specific microorganism, the EN 14348 standard may be used for the microorganism in question.

The most recent version of standards in force at the time of the tests must be used.

Furthermore, in accordance with the conclusions in Annex VI (77) the level, consistency and duration of protection, control or other intended effects must, as a minimum, be similar to those resulting from suitable reference products, where such products exist, or to other means of control. Where no reference products exist, the biocidal product must give a defined level of protection or control in the areas of proposed use.

Considering the history of the use of formaldehyde, it may therefore be worthwhile to include with the application information about the bactericidal efficacy of formaldehyde, if available.

In France, formaldehyde is most commonly used at concentrations of about 28% for cavity fluid and 1.5% for arterial fluid. As formaldehyde is currently under assessment in the review programme, efficacy data may become available when the assessment report is published by the evaluating Competent Authority (eCA). The standards proposed above for validating claims may be reviewed at a later stage in the context of the review of this guidance document as a result of the conclusions published by the eCA on the efficacy of formaldehyde, or in the event of other data for this same substance becoming available in the future.

5.7.2.3.2.3 Verifying that antibacterial activity is maintained

When embalming, the biocidal product must remain effective over several days, until burial. The persistence indicated on the label must be proven, e.g. by challenge tests. The following protocol may be used, adapted from the French NF X30-503 standard (Healthcare waste - Reduction by disinfection pre-treatment appliances in microbiological and mechanical risks involving infections and other comparable healthcare waste).

- In order to ensure that bacteria are destroyed and not merely subjected to stress or inhibition by the biocidal product, and to confirm the absence of bacterial revival, the bacterial suspension, treated according to the EN 13727 standard, is held at ambient temperature for four to six days and then the bacteria are counted. In the laboratory, it is held at 20°C until analysis.
- The bacteria in the bacterial suspension are counted on the day of treatment and again after four to six days.
- Lasting disinfection is shown by the absence of bacterial revival, i.e. the bacterial count on day 4-6 must not be increased by more than one log compared to the bacterial load measured in the sample taken on the day of treatment (Day 0).
- The "effective" dose of the product must be in a range bounded by upper and lower limits, which are:
  - a lower concentration for which bacterial recrudescence is observed after 4-6 days;
  - a higher concentration.

5.7.2.3.2.4 Tests on human bodies

To complement in vitro efficacy tests for the biocidal product used for the preservation of human bodies, tests on bodies are necessary to assess product performance.

Because of the number of factors that can influence the efficacy of a biocidal product, such as the cause of death or the time lapsed or the condition of the body before embalming begins, a sufficient number of bodies (at least 20) satisfying the requirements of the grid in Appendix 22 and the claims for the product, must be
available for optimum assessment of the results in terms of preservation of the body for viewing by families.

**NOTE to the reader:**
The applicant has to inquire about the legislation in force in the Member State (MS) where the tests on human bodies are performed (e.g. current French regulations only allow bodies donated to science to be used to test a product that has not yet been approved.

Every centre for the donation of bodies participating in these tests on human bodies must declare the number of bodies undergoing tests in its establishment. This declaration is supplied to the applicant and must be submitted with the application.

In all cases, whatever the legislation in force in each MS, tests on human bodies with good quality and in line with this guidance will be accepted by MS when the dossier will be submitted for authorisation.

The assessment grid for specific biocidal products is shown in Appendix 23. Its purpose is not to assess the overall embalming treatment but only the biocidal product for which authorisation is being requested.

The grid consists of:

- general information: date and place of the treatment, identification of the deceased (gender, age), weight, corpulence, adiposity, date and causes of death, etc.;
- the preoperative body examination: bodily integrity, autopsy, external prostheses, surgery, visible anomalies (decomposition, rigidity, dehydration, lividity, colouring of tissues, dermal lesions, distension of the abdomen, bruising, etc.). The bodies used must be representative of the range of criteria listed in this section;
- the techniques used to inject the biocidal product: timetable, sites and types of injection, biocidal product used, drainage and puncture;
- observations concerning the injection of the biocidal product: observations during treatment, 48 hours after treatment and after different periods in accordance with the applicant's claims;
- where necessary, the use of other products during the preservation process: products for cosmetic purposes, humidifiers and other products.

The embalmer thus assesses the efficacy of the embalming product on a series of human bodies, using the grid provided. The efficacy is judged for the duration claimed by the manufacturer according to observations concerning odour, colouring and the suppleness of the skin after injection of the biocidal product. In the event that the tests on these human bodies have to be interrupted for any reason, the results already obtained remain valid for three years following the official decision to halt the tests.

**5.7.2.3.2.5 Choice of dose**
The usage dose\(^{39}\) claimed is a matter for the applicant. Indeed, related to the body conditions, it can be necessary to test several doses above the dose determined in laboratory and then define a range of doses, adapted to difficult cases. They must choose the usage dose claimed according to the efficacy sought and the precautions for use that will be imposed on embalming technicians by their employers, depending on the health risks created by the full preparation (active substance at the chosen concentration plus excipients and solvents). In cases where little is known about the pathogenic microorganisms that might present a risk to the embalmer, it is essential that protective

\(^{39}\) Concentration and volume injected
measures be taken during the preservation process. These measures should not be primary criteria for choosing the biocidal product used for the treatment.

If the applicant chooses a range of doses instead of a single value, the lower must be justified with appropriate tests, as defined in the preceding section (and also the higher dose in the case where different doses have been tested in the human body tests to cover difficult cases). The applicant may also request approval for two different doses, one of them more concentrated for special or difficult cases (bodies found some time after death or in contact with water, for example).

5.7.2.4 Assessing the application for authorisation

The assessment of the embalming product shall be favourable if it satisfies the following efficacy criteria:

- laboratory test: bactericidal properties (EN 13727 and/or EN 14348 standards): obligatory test conditions;
- laboratory test: e.g. challenge test: no bacterial recrudescence for at least 4-6 days by more one log compared to the bacterial load measured in the sample taken on the day of treatment (Day 0), with the bacterial suspension being held at ambient temperature;
- field test: 80% of the bodies must meet the satisfaction criteria at T+48 hours. Satisfaction criteria are according to the grid: normal or fair odour, colouring and suppleness of the skin, related to the initial conditions of the body.

**SUMMARY OF THE PARAMETERS ASSESSED**

**EFFICACY CLAIMS ON THE LABEL SUBMITTED**

1. Does the applicant make any specific claims? Y/N
2. Have the efficacy claims on the label been judged and dealt with according to the parameters described in this guidance document for this type of product? Y/N

**ASSESSING THE DATA**

3. Has each study (or supplementary item) been assessed individually for robustness? Y/N
4. Has each study (or supplementary item) been assessed individually for quality assurance? Y/N
5. Has each study (or supplementary item) been assessed individually for suitability (i.e. for reliability and relevance concerning the claims)? Y/N

**DECISION-MAKING**

Considering all the available data:

6. Are the claims on the label sufficiently supported? Y/N
7. Do the claims on the label require modifications? Y/N
8. On the basis of the efficacy data submitted, can authorisation for the use of the product be recommended? Y/N
Appendix 1. Claims Matrices

The claims matrices are a set of tables linked to this guidance document: these documents are available on the ECHA Biocides Efficacy Working Group webpage [http://echa.europa.eu/about-us/who-we-are/biocidal-products-committee/working-groups/efficacy].

The claims matrices linked to this document are intended to cover biocidal products covered under the scope of Product Type 1, 2, 3 and 4 and for Treated Articles.

The claims matrix is a tool for the applicant and CAs. It is intended to capture the information that is needed in the authorisation dossier, to adequately describe typical combinations of products, formats of application of the products, as well as target sites. It also includes the claims made and the requirements for testing these claims (in terms of methodology and appropriate performance standards) for a product to be used in this way.

The reader should note that the matrices are not exhaustive in terms of use patterns, scenarios and test methods.

The claims matrix must be used together with the relevant sections within the efficacy guidance document so as to provide both applicants and CAs alike with clear direction as to the nature and extent of the efficacy data required to support a claimed effect. The claims matrix acts as a guide to the information required when compiling an efficacy dataset for a PT1, PT2, PT3 or PT4 biocidal product and for Treated Articles.

To note:

- Each row (entry) within the matrices is not independent and can be linked to other entries.
- These matrices only address biocidal claims made for these products.
- The claim matrix will be updated regularly according to the state-of-the-art.
Appendix 2. Standards and testing methods for efficacy-testing of disinfectant biocidal products (PT 1-5)

The methods for testing efficacy referenced within this guidance document are enlisted below. The use of European Standards (Table 28) is highly recommended if available and appropriate for the respective application. Should no European Standard for an application be available yet and an adaption of an existing standard is not possible according to the rules laid down in EN 14885, other test methods and guidance documents (Table 29) may be used. In cases where the below mentioned methods are inappropriate to demonstrate efficacy of a product for special applications, methods from other national or international standardisation bodies may also be employed. These include for example, OECD, ASTM or ISO methods. It is recommended to agree such testing strategies with the evaluating CA before tests are performed.

Tests should be carried out according to the respective latest edition of a standard. Please check the respective web sites for the latest information.

**Table 28: CEN European standards**

<table>
<thead>
<tr>
<th>Reference</th>
<th>Title</th>
<th>PT</th>
<th>Scope/Remarks</th>
</tr>
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<tbody>
<tr>
<td>EN 1276</td>
<td>Chemical disinfectants and antiseptics - Quantitative suspension test for the evaluation of bactericidal activity of chemical disinfectants and antiseptics used in food, industrial, domestic, and institutional areas - Test method and requirements (phase 2, step 1)</td>
<td>1,2,4</td>
<td>This European Standard specifies a method for testing bactericidal activity by assessing reduction in the number of viable bacterial cells in suspension under defined conditions. The approach can be applied to formulated products or to biocidal active substances.</td>
</tr>
<tr>
<td>EN 1499</td>
<td>Chemical disinfectants and antiseptics - Hygienic handwash - Test method and requirements (phase 2, step 2)</td>
<td>1</td>
<td>This European Standard specifies a test method simulating practical conditions for establishing whether a hygienic handwash product reduces the transmission of transiently contaminating microorganisms when used to wash the artificially contaminated hands of volunteers.</td>
</tr>
<tr>
<td>EN 1500</td>
<td>Chemical disinfectants and antiseptics - Hygienic handrub - Test method and requirements (phase 2, step 2)</td>
<td>1</td>
<td>This European Standard specifies a test method simulating practical conditions for establishing whether a hygienic handrub product reduces the transmission of transiently contaminating microorganisms when rubbed onto the artificially contaminated hands of volunteers.</td>
</tr>
<tr>
<td>EN 1650</td>
<td>Chemical disinfectants and antiseptics - Quantitative suspension test for the evaluation of fungicidal or yeasticidal activity of chemical disinfectants and antiseptics used in food, industrial, domestic, and institutional areas - Test method and requirements (phase 2, step 1)</td>
<td>1,2,4</td>
<td>This European Standard specifies a method for testing fungical or yeasticidal activity by assessing reduction in the number of viable mould spores and/or yeast cells in suspension under defined conditions. The approach can be applied to formulated products or to biocidal active substances.</td>
</tr>
</tbody>
</table>

40 The CEN does not sell or distribute standards or any other deliverable. All European Standards (EN) and drafts (prEN) as well as other approved documents are directly available for purchase from the CEN national standardisation bodies.
<table>
<thead>
<tr>
<th>Reference</th>
<th>Title</th>
<th>PT</th>
<th>Scope/Remarks</th>
</tr>
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<tbody>
<tr>
<td>EN 1656</td>
<td>Chemical disinfectants and antiseptics - Quantitative suspension test for the evaluation of bactericidal activity of chemical disinfectants and antiseptics used in the veterinary area - Test method and requirements (phase 2, step 1)</td>
<td>3</td>
<td>This European Standard specifies a method for testing bactericidal activity by assessing reduction in the number of viable bacterial cells in suspension under defined conditions. The approach can be applied to formulated products or to biocidal active substances.</td>
</tr>
<tr>
<td>EN 1657</td>
<td>Chemical disinfectants and antiseptics - Quantitative suspension test for the evaluation of fungicidal or yeasticidal activity of chemical disinfectants and antiseptics used in the veterinary area - Test method and requirements (phase 2, step 1)</td>
<td>3</td>
<td>This European Standard specifies a method for testing fungicidal or yeasticidal activity by assessing reduction in the number of viable mould spores and/or yeast cells in suspension under defined conditions. The approach can be applied to formulated products or to biocidal active substances.</td>
</tr>
<tr>
<td>EN 12353</td>
<td>Chemical disinfectants and antiseptics - Preservation of test organisms used for the determination of bactericidal (including Legionella), mycobactericidal, sporidical, fungicidal and virucidal (including bacteriophages) activity</td>
<td>1,2,3,4,5</td>
<td>This method specifies how to keep test organisms used and defined in European Standards for the determination of bactericidal, mycobactericidal, sporidical, fungicidal and virucidal (incl. bacteriophages) activity of chemical disinfectants and antiseptics drawn up by CEN/TC 216.</td>
</tr>
<tr>
<td>EN 12791</td>
<td>Chemical disinfectants and antiseptics - Surgical hand disinfection - Test method and requirements (phase 2, step 2)</td>
<td>1</td>
<td>This European Standard specifies a test method simulating practical conditions for establishing whether a product for surgical hand disinfection reduces the transmission of the microbial flora on hands when used for the treatment of clean hands of volunteers.</td>
</tr>
<tr>
<td>EN 13610</td>
<td>Chemical disinfectants - Quantitative suspension test for the evaluation of virucidal activity against bacteriophages of chemical disinfectants used in food and industrial areas - Test method and requirements (phase 2, step 1)</td>
<td>4</td>
<td>This European Standard specifies a method for testing virucidal activity against bacteriophages by assessing reduction in the number of infectious bacteriophage particles in suspension under defined conditions. The approach can be applied to formulated products or to biocidal active substances.</td>
</tr>
<tr>
<td>EN 13623</td>
<td>Chemical disinfectants and antiseptics - Quantitative suspension test for the evaluation of bactericidal activity against Legionella of chemical disinfectants for aqueous systems - Test method and requirements (phase 2, step 1)</td>
<td>2,4,5</td>
<td>This European Standard specifies a method for testing bactericidal activity against Legionella by assessing reduction in the number of viable Legionella cells in suspension under defined conditions. The approach can be applied to formulated products or to biocidal active substances.</td>
</tr>
<tr>
<td>EN 13624</td>
<td>Chemical disinfectants and antiseptics - Quantitative suspension test for the evaluation of fungicidal and yeasticidal activity in the medical area - Test method and requirements (phase 2, step 1)</td>
<td>1,2</td>
<td>This European Standard specifies a method for testing fungicidal or yeasticidal activity by assessing reduction in the number of viable mould spores and/or yeast cells in suspension under defined conditions. The approach can be applied to formulated products or to biocidal active substances.</td>
</tr>
<tr>
<td>Reference</td>
<td>Title</td>
<td>PT</td>
<td>Scope/Remarks</td>
</tr>
<tr>
<td>-------------</td>
<td>----------------------------------------------------------------------</td>
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<td>--------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>EN 13697</td>
<td>Chemical disinfectants and antisepsics - Quantitative non-porous surface test for the evaluation of bactericidal and/or fungicidal activity of chemical disinfectants used in food, industrial, domestic and institutional areas - Test method and requirements without mechanical action (phase 2, step 2)</td>
<td>2,4</td>
<td>This European Standard specifies a method for testing bactericidal and/or fungicidal or yeasticidal activity by assessing reduction in the number of viable bacterial cells and/or mould spores and/or yeast cells dried on a steel carrier under defined conditions. The approach can be applied to formulated products or to biocidal active substances.</td>
</tr>
<tr>
<td>EN 13704(^1)</td>
<td>Chemical disinfectants - Quantitative suspension test for the evaluation of sporicidal activity of chemical disinfectants used in food, industrial, domestic and institutional areas - Test method and requirements (phase 2, step 1)</td>
<td>4</td>
<td>(1, 2, 3) This European Standard specifies a method for testing sporicidal activity by assessing reduction in the number of viable bacterial endospores in suspension under defined conditions. The approach can be applied to formulated products or to biocidal active substances.</td>
</tr>
<tr>
<td>EN 13727</td>
<td>Chemical disinfectants and antisepsics - Quantitative suspension test for the evaluation of bactericidal activity in the medical area - Test method and requirements (phase 2, step 1)</td>
<td>1,2</td>
<td>This European Standard specifies a method for testing bactericidal activity by assessing reduction in the number of viable bacterial cells in suspension under defined conditions. The approach can be applied to formulated products or to biocidal active substances.</td>
</tr>
<tr>
<td>EN 14204</td>
<td>Chemical disinfectants and antisepsics - Quantitative suspension test for the evaluation of mycobactericidal activity of chemical disinfectants and antisepsics used in the veterinary area - Test method and requirements (phase 2, step 1)</td>
<td>3</td>
<td>This European Standard specifies a method for testing mycobactericidal activity by assessing reduction in the number of viable mycobacterial cells in suspension under defined conditions. The approach can be applied to formulated products or to biocidal active substances.</td>
</tr>
<tr>
<td>EN 14347</td>
<td>Chemical disinfectants and antisepsics - Basic sporicidal activity - Test method and requirements (phase 1)</td>
<td>1,2</td>
<td>(3, 4) This European Standard specifies a method for testing sporicidal activity by assessing reduction in the number of viable bacterial endospores in suspension under defined conditions. The method is declared as a phase 1 test but, based on its requirements, it can serve as a suspension test (comparable to phase 2, step 1) until revised/additional CEN methodology for testing sporicidal activity becomes available. The approach can be applied to formulated products or to biocidal active substances.</td>
</tr>
<tr>
<td>EN 14348</td>
<td>Chemical disinfectants and antisepsics - Quantitative suspension test for the evaluation of mycobactericidal activity of chemical disinfectants in the medical area including instrument disinfectants - Test methods and requirements (phase 2, step 1)</td>
<td>1,2</td>
<td>This European Standard specifies a method for testing mycobactericidal activity by assessing reduction in the number of viable mycobacterial cells in suspension under defined conditions. The method is also applicable to demonstrate tuberculocidal activity only. The approach can be applied to formulated products or to biocidal active substances.</td>
</tr>
</tbody>
</table>

\(^1\) EN 13704 is under review and the revised standard will include veterinary and human health care areas.
<table>
<thead>
<tr>
<th>Reference</th>
<th>Title</th>
<th>PT</th>
<th>Scope/Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>EN 14349</td>
<td>Chemical disinfectants and antiseptics - Quantitative surface test for the evaluation of bactericidal activity of chemical disinfectants and antiseptics used in the veterinary area on non-porous surfaces without mechanical action - Test method and requirements (phase 2, step 2)</td>
<td>3</td>
<td>This European Standard specifies a method for testing bactericidal activity by assessing reduction in the number of viable bacterial cells dried on a steel carrier under defined conditions. The approach can be applied to formulated products or to biocidal active substances.</td>
</tr>
<tr>
<td>EN 14476</td>
<td>Chemical disinfectants and antiseptics - Quantitative suspension test for the evaluation of virucidal activity in the medical area - Test method and requirements (phase 2, step 1)</td>
<td>1,2 (4)</td>
<td>This European Standard specifies a method for testing virucidal activity by assessing reduction in the number of infectious virus particles in suspension under defined conditions. The approach can be applied to formulated products or to biocidal active substances.</td>
</tr>
<tr>
<td>EN 14561</td>
<td>Chemical disinfectants and antiseptics - Quantitative carrier test for the evaluation of bactericidal activity for instruments used in the medical area - Test method and requirements (phase 2, step 2)</td>
<td>2</td>
<td>This European Standard specifies a method for testing bactericidal activity by assessing reduction in the number of viable bacterial cells dried on a frosted glass carrier under defined conditions. The approach can be applied to formulated products or to biocidal active substances.</td>
</tr>
<tr>
<td>EN 14562</td>
<td>Chemical disinfectants and antiseptics - Quantitative carrier test for the evaluation of fungicidal or yeasticidal activity for instruments used in the medical area - Test method and requirements (phase 2, step 2)</td>
<td>2</td>
<td>This European Standard specifies a method for testing fungicidal or yeasticidal activity by assessing reduction in the number of viable mould spores and/or yeast cells dried on a frosted glass carrier under defined conditions. The approach can be applied to formulated products or to biocidal active substances.</td>
</tr>
<tr>
<td>EN 14563</td>
<td>Chemical disinfectants and antiseptics - Quantitative carrier test for the evaluation of mycobactericidal or tuberculocidal activity of chemical disinfectants used for instruments in the medical area - Test method and requirements (phase 2, step 2)</td>
<td>2</td>
<td>This European Standard specifies a method for testing mycobactericidal activity by assessing reduction in the number of viable mycobacterial cells dried on a frosted glass carrier under defined conditions. The method is also applicable to demonstrate tuberculocidal activity only. The approach can be applied to formulated products or to biocidal active substances.</td>
</tr>
<tr>
<td>EN 14675</td>
<td>Chemical disinfectants and antiseptics - Quantitative suspension test for the evaluation of virucidal activity of chemical disinfectants and antiseptics used in the veterinary area - Test method and requirements (phase 2, step 1)</td>
<td>3</td>
<td>This European Standard specifies a method for testing virucidal activity by assessing reduction in the number of infectious virus particles in suspension under defined conditions. The approach can be applied to formulated products or to biocidal active substances.</td>
</tr>
<tr>
<td>Reference</td>
<td>Title</td>
<td>PT</td>
<td>Scope/Remarks</td>
</tr>
<tr>
<td>-------------</td>
<td>-----------------------------------------------------------------------</td>
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<td>---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>EN 14885</td>
<td>Chemical disinfectants and antiseptics - Application of European Standards for chemical disinfectants and antiseptics</td>
<td>1,2,3,4,5,</td>
<td>This European Standard specifies the European Standards, i.e. test methods, to which products have to conform in order to support the claims for microbicidal activity which are referred to in this document. It also specifies terms and definitions which are used in European Standards. It is applicable to products for which activity is claimed against the following microorganisms: vegetative bacteria (incl. mycobacteria and Legionella), bacterial spores, yeasts, fungal spores and viruses (incl. bacteriophages).</td>
</tr>
<tr>
<td>EN 16437</td>
<td>Chemical disinfectants and antiseptics - Quantitative surface test for the evaluation of bactericidal activity of chemical disinfectants and antiseptics used in veterinary area on porous surfaces without mechanical action - Test method and requirements (phase 2, step 2)</td>
<td>3</td>
<td>This European Standard specifies a method for testing bactericidal activity by assessing reduction in the number of viable bacterial cells dried on a wood carrier under defined conditions. The approach can be applied to formulated products or to biocidal active substances.</td>
</tr>
<tr>
<td>EN 16438</td>
<td>Chemical disinfectants and antiseptics - Quantitative surface test for the evaluation of fungicidal or yeasticidal activity of chemical disinfectants and antiseptics used in the veterinary area on non-porous surfaces without mechanical action - Test method and requirements (phase 2, step 2)</td>
<td>3</td>
<td>This European Standard specifies a method for testing fungicidal or yeasticidal activity by assessing reduction in the number of viable mould spores and/or yeast cells dried on a steel carrier under defined conditions. The approach can be applied to formulated products or to biocidal active substances.</td>
</tr>
<tr>
<td>EN 16615</td>
<td>Chemical disinfectants and antiseptics - Chemical-thermal test method for the evaluation of bactericidal and yeasticidal activity on non-porous surfaces with mechanical action employing wipes in the medical area (4-field test) - Test method and requirements (phase 2, step 2)</td>
<td>2(4)</td>
<td>This European Standard specifies a method for testing bactericidal and/or yeasticidal activity by assessing reduction in the number of viable bacterial and/or yeast cells dried on a PVC carrier under defined conditions. The test applies to products that are used for disinfecting non-porous surfaces by wiping and includes 'ready-to-use wipes' which are impregnated with a microbicidal solution.</td>
</tr>
<tr>
<td>EN 16616</td>
<td>Chemical disinfectants and antiseptics - Chemical-thermal textile disinfection - Test method and requirements (phase 2, step 2)</td>
<td>2(3,4)</td>
<td>This European Standard specifies a method for testing microbicidal activity of a disinfection process for the treatment of contaminated textile. The procedure is carried out by using a washing machine and microbicidal activity is assessed as the reduction in the number of viable test organisms, such as bacterial, mycobacterial or yeast cells and mould spores, dried on a cotton carrier under defined conditions.</td>
</tr>
</tbody>
</table>
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Version 1.0  February 2017

<table>
<thead>
<tr>
<th>Reference</th>
<th>Title</th>
<th>PT</th>
<th>Scope/Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>EN 16777</td>
<td>Chemical disinfectants and antiseptics - Quantitative non-porous surface test without mechanical action for the evaluation of virucidal activity of chemical disinfectants used in the medical area - Test method and requirements (phase 2, step 2)</td>
<td>2  (4)</td>
<td>This European Standard specifies a method for testing virucidal activity by assessing reduction in the number of infectious virus particles dried on a steel carrier under defined conditions. The approach can be applied to formulated products or to biocidal active substances.</td>
</tr>
</tbody>
</table>

#### Table 29: Other test methods and guidance documents

<table>
<thead>
<tr>
<th>Reference</th>
<th>Title</th>
<th>PT</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>ASTM E2196</td>
<td>Standard Test Method for Quantification of <em>Pseudomonas aeruginosa</em> Biofilm Grown with Medium Shear and Continuous Flow Using Rotating Disk Reactor</td>
<td>2,3,4</td>
<td>This test method is used for growing a reproducible <em>Pseudomonas aeruginosa</em> biofilm in a continuously stirred tank reactor (CSTR) under medium shear conditions. In addition, the test method describes how to sample and analyse biofilm for viable cells. Available via: <a href="http://www.astm.org/Standard/">http://www.astm.org/Standard/</a> or the national standardisation bodies</td>
</tr>
<tr>
<td>ASTM E2274</td>
<td>Standard Test Method for Evaluation of Laundry Sanitizers and Disinfectants</td>
<td>2,3</td>
<td>This test method is designed to evaluate sanitizing/disinfectant laundry detergents/additives for use in top-loading automatic clothes washing operations. This test method is designed predominantly to provide testing with representative vegetative bacteria but can also be designed to accommodate the testing of fungi and viruses.</td>
</tr>
<tr>
<td>ASTM E2406</td>
<td>Standard Test Method for Evaluation of Laundry Sanitizers and Disinfectants for Use in High Efficiency Washing Operations</td>
<td></td>
<td>This test method is designed to evaluate sanitizing/disinfectant laundry detergents/additives for use in high efficiency (HE) automatic clothes washing operations that typically utilize very low wash water volumes. This test method is designed to provide testing with representative vegetative bacteria but can also be designed to accommodate the testing of fungi and viruses.</td>
</tr>
<tr>
<td>ASTM E2562</td>
<td>Standard Test Method for Quantification of <em>Pseudomonas aeruginosa</em> Biofilm Grown with High Shear and Continuous Flow using CDC Biofilm Reactor</td>
<td>2,3,4</td>
<td>This test method specifies the operational parameters required to grow a reproducible <em>Pseudomonas aeruginosa</em> biofilm under high shear. The resulting biofilm is representative of generalized situations where biofilm exists under high shear rather than being representative of one particular environment. Available via: <a href="http://www.astm.org/Standard/">http://www.astm.org/Standard/</a> or the national standardisation bodies</td>
</tr>
<tr>
<td>Reference</td>
<td>Title</td>
<td>PT</td>
<td>Remarks</td>
</tr>
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<td>---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>DIN SPEC 10534</td>
<td>Food hygiene - Commercial dishwashing - Hygiene requirements, testing</td>
<td>4</td>
<td>This document is a summary of the standards DIN 10510, DIN 10511, DIN 10512 and DIN 10522. It specifies hygiene requirements relating to the design, construction and operation of commercial warewashers and in particular provides information on their hygienic and proper operation, on cleaning and disinfection of wash ware and on care and maintenance of the machinery. It describes the methods for testing hygienic operation. Available via: <a href="http://www.beuth.de/en/">http://www.beuth.de/en/</a> or the national standardisation bodies</td>
</tr>
<tr>
<td>DVG Guidelines</td>
<td>Guidelines for the testing of disinfection procedures and chemical disinfectants; Original title: Richtlinien für die Prüfung von Desinfektionsverfahren und chemischen Desinfektionsmitteln</td>
<td>3,4</td>
<td>DVG Guidelines specify methods for testing activity of chemical disinfectants against bacteria, yeasts and fungal spores, viruses, and parasites. They apply to the veterinary and the food sector, such as animal husbandry, veterinary practices, meat production/food of animal origin, and large-scale/canteen kitchens (except ward kitchens catering patients). DVG Guidelines are published by the German Veterinary Medical Society (DVG). Available in German via: <a href="http://www.desinfektion-dvg.de">http://www.desinfektion-dvg.de</a></td>
</tr>
<tr>
<td>ISO/TS 15883-5</td>
<td>Washer-disinfectors - Part 5: Test soils and methods for demonstrating cleaning efficacy</td>
<td>2,3,4</td>
<td>ISO 15883 relates to a series of standards that specify the required performance levels of Washer-Disinfectors. Part 5, the Technical Specification (TS), describes a method to generate biofilm formed by <em>Pseudomonas aeruginosa</em>. Available via: <a href="http://www.iso.org/iso/home.htm">http://www.iso.org/iso/home.htm</a> or the national standardisation bodies</td>
</tr>
<tr>
<td>NF T72-281</td>
<td>Methods of airborne disinfection of surfaces - Determination of bactericidal, fungicidal, yeasticidal, mycobactericidal, tuberculocidal, sporicidal and virucidal activity, including bacteriophages; Original title: Procédés de désinfection des surfaces par voie aérienne - Détermination de l'activité bactéricide, fongicide, levuricide, mycobactéricide, tuberculocide sporicide et virucide incluant les bactériophages</td>
<td>2,3,4</td>
<td>This French standard specifies a method for testing microbicidal activity of airborne disinfection processes. The tested product is diffused, e.g. in gaseous form or as an aerosol, to reduce the number of relevant test organisms, such as bacteria, bacterial spores, yeasts, and fungal spores. Available in French via: <a href="http://www.afnor.org/en">http://www.afnor.org/en</a> or the national standardisation bodies</td>
</tr>
<tr>
<td>Nordic Working Paper</td>
<td>Efficacy Assessment of Treated Articles: A guidance</td>
<td>1,2,3,4</td>
<td>The document provides guidance on efficacy testing of biocides used in treated articles. The presence and relevance of existing standard test methods is described and, where they do not exist or where they do not provide sufficient support, the nature of the data required will be described. The document was published by the Nordic Council of Ministers. Open access via: <a href="http://www.norden.org/en/publications/publikationer/2014-904/">http://www.norden.org/en/publications/publikationer/2014-904/</a></td>
</tr>
<tr>
<td>Reference</td>
<td>Title</td>
<td>PT</td>
<td>Remarks</td>
</tr>
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</tr>
<tr>
<td>OECD Series on Biocides No. 4</td>
<td>Guidance Document for Demonstrating Efficacy of Pool and Spa Disinfectants and Field Testing (Series on Testing and Assessment No. 170 and Series on Biocides No. 4)</td>
<td>2</td>
<td>The document provides guidance on setting up a strategy for efficacy testing of pool and spa disinfectants in a laboratory scale testing phase and a field testing phase in a full-size swimming or spa pool. Open access via: <a href="http://www.oecd.org/env/ehs/pesticides-biocides/biocidestestguidelinesandguidencedocuments.htm">http://www.oecd.org/env/ehs/pesticides-biocides/biocidestestguidelinesandguidencedocuments.htm</a></td>
</tr>
<tr>
<td>OECD Series on Biocides No. 8</td>
<td>Guidance Document for Quantitative Method for Evaluating Antibacterial Activity of Porous and Non-Porous Antibacterial Treated Materials (Series on Testing and Assessment No. 202 and Series on Biocides No. 8)</td>
<td>1,2,3,4</td>
<td>The document provides guidance for testing the basic antibacterial performance of porous (textile) and non-porous (plastic) materials that have been treated with a biocide with the intention of introducing antibacterial/hygienic properties into that material. Open access via: <a href="http://www.oecd.org/env/ehs/pesticides-biocides/biocidestestguidelinesandguidencedocuments.htm">http://www.oecd.org/env/ehs/pesticides-biocides/biocidestestguidelinesandguidencedocuments.htm</a></td>
</tr>
<tr>
<td>VAH Standard methods</td>
<td>VAH certification of chemical disinfection procedures; Original title: VAH-Zertifizierung chemischer Desinfektionsverfahren</td>
<td>1,2</td>
<td>VAH Standard methods specify methods for testing activity of chemical disinfectants against bacteria (incl. mycobacteria), yeasts, and fungal spores. They apply to testing products used for disinfection in public facilities (medical and other) and, in the event of substantiated medical indications, also in the private home. VAH Standard methods are published by the Association for Applied Hygiene (VAH). Available in German via: <a href="http://www.mhp-verlag.de/en/home/">http://www.mhp-verlag.de/en/home/</a></td>
</tr>
</tbody>
</table>
Appendix 3. Table of Reference Test Organisms (PT 1-5)

This table (Table 30) is given as a general overview of relevant test organisms for testing disinfectants in accordance with the BPR.

This table comprises mainly those reference test organisms that are included in the EN norms covered by EN 14885. Furthermore, strains are listed that are recommended for some uses (e.g. endoparasites from DVG standard).

The reader can check the website of the CEN (European Standardization Organizations): www.cen.eu for new and updated standards.

Since the EN systematics of WG’s 1 to 3 does not fit exactly to the BPR PT scheme, in borderline cases an indicated reference test organism might be used for other PTs as well. In cases where there are discrepancies between this ECHA guidance and the guidance in EN 14885, the ECHA Guidance should be followed as the leading guidance. However, EN 14885 can be followed with satisfactory justification to meet the requirements of the BPR.

Tests with test organisms in addition to those mentioned below are acceptable, if adequate scientific evidence is submitted on which the relevance of the test organism to the field of use can be judged.

Table 30: Reference Test Organisms

<table>
<thead>
<tr>
<th>Micro-organisms</th>
<th>PT1*</th>
<th>PT2*</th>
<th>PT3*</th>
<th>PT4*</th>
<th>PT5*</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Staphylococcus aureus</em> ATCC 6538</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td><em>Pseudomonas aeruginosa</em> ATCC 15442 (not for teat disinfection)</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td><em>Enterococcus hirae</em> ATCC 10541 (not for teat disinfection)</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td><em>Escherichia coli</em> ATCC 10536 (teat disinfection)</td>
<td>X</td>
<td>X</td>
<td>O</td>
<td>X</td>
<td>O</td>
</tr>
<tr>
<td><em>Escherichia coli</em> K12 NCTC 10538</td>
<td>X</td>
<td>X</td>
<td>O</td>
<td>O</td>
<td>O</td>
</tr>
<tr>
<td><em>Salmonella Typhimurium</em> ATCC 13311</td>
<td>X</td>
<td>O</td>
<td>O</td>
<td>O</td>
<td>O</td>
</tr>
<tr>
<td><em>Lactobacillus brevis</em> DSM6235</td>
<td>X</td>
<td>O</td>
<td>O</td>
<td>O</td>
<td>O</td>
</tr>
<tr>
<td><em>Enterobacter cloacae</em> DSM 6234</td>
<td>X</td>
<td>O</td>
<td>O</td>
<td>O</td>
<td>O</td>
</tr>
<tr>
<td><em>Enterococcus faecium</em> ATCC 6057 (for T ≥40°C)</td>
<td>X</td>
<td>O</td>
<td>O</td>
<td>O</td>
<td>O</td>
</tr>
<tr>
<td><em>Proteus vulgaris</em> ATCC 13315 (not for teat disinfection)</td>
<td>X</td>
<td>(X)</td>
<td>O</td>
<td>O</td>
<td>O</td>
</tr>
<tr>
<td><em>Streptococcus uberis</em> ATCC 19436 (teat disinfection)</td>
<td>X</td>
<td>(X)</td>
<td>O</td>
<td>O</td>
<td>O</td>
</tr>
<tr>
<td><em>Legionella pneumophila</em> ATCC 33152 (PT2: pools, hot tubs; PT4: drinking water systems, PT5: in collective drinking water systems)</td>
<td>X</td>
<td>X</td>
<td>O</td>
<td>O</td>
<td>O</td>
</tr>
</tbody>
</table>

Key for Table 28:
* X = basic requirement to claim activity against micro-organism;
(X) = basic requirement for specific use as described in the table below in brackets;
O = optional;
** Aspergillus brasiliensis is the name of Aspergillus niger after reclassification in 2008;
*** for a limited spectrum virus claim in PT1 Poliovirus does not have to be tested.
## Micro-organisms

<table>
<thead>
<tr>
<th>Yeasts</th>
<th>PT1*</th>
<th>PT2*</th>
<th>PT3*</th>
<th>PT4*</th>
<th>PT5*</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Candida albicans</em> ATCC 10231</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td><em>Saccharomyces cerevisiae</em> ATCC 9763 (breweries)</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>(X)</td>
<td>(X)</td>
</tr>
<tr>
<td><em>Saccharomyces cerevisiae</em> DSM 70487 (breweries)</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>(X)</td>
<td>(X)</td>
</tr>
</tbody>
</table>

### Fungal spores

| Aspergillus brasiliensis** ATCC 16404 | X    | X    | X    | X    | X    |

### Viruses

| Polio virus type 1, LSc-2ab (Picornavirus) | X*** | X    |     |     |     |
| Adenovirus, type 5, strain Adenoid 75, ATCC VR-5. | X    | X    | X    | (X)  | (X)  |
| Murine norovirus, strain S99 Berlin | X    | X    | X    | (X)  | (X)  |
| Murine Parvovirus, strain Crawford, ATCC VR-1346 (for T ≥40°C) | (X)  |     |     |     |     |
| Bovine Enterovirus Type 1, ECBO - Virus ATCC VR-248 | (X)  |     |     |     |     |
| Rotavirus (pools, hot tubs) | (X)  |     |     |     |     |

### Enveloped Viruses

| MVA = Modified Vacciniavirus Ankara (teat disinfection) | X    | (X)  |     |     |     |

### Bacteriophages

| Bacteriophage P001 DMS 4262 (milk industry) | X    |     |     |     |     |
| Bacteriophage P008 DMS 10567 (milk industry) | X    |     |     |     |     |

### Mycobacteria

| Mycobacterium terrae ATCC 15755 | X    | X    |     |     |     |
| Mycobacterium avium ATCC 15769 | X    | X    | X    |     |     |

(PT1 and PT2 claim for mycobactericidal: both, tuberculocidal: *M. terrae* only)

### Bacterial spores

| Spores of *Bacillus cereus* ATCC 12826 (bee hives) | O    | (X)  | O    |     |     |
| Spores of *Bacillus subtilis* ATCC 6633 (bee hives) | X    | O    | (X)  | X    |     |
| Spores of *Clostridium sporogenes* ATCC 7955 | O    | O    | O    |     |     |
| Spores of *Geobacillus stearothermophilus* (for T ≥60°C) | O    | O    | O    |     |     |

### Endoparasites

| Oocysts of *Eimeria tenella* strain Houghton (chicken farms) | (X)  |     |     |     |     |
Appendix 4. Overview of standards, test conditions and pass criteria (PT 1-5)

The overview is presented in a number of tables which are available on the ECHA Biocides Efficacy Working Group webpage [http://echa.europa.eu/about-us/who-we-are/biocidal-products-committee/working-groups/efficacy].

These tables provide an overview of available phase 2,1 and 2,2 EN standards which are applicable for testing the efficacy of disinfectant biocides. This overview is not exhaustive. For other or more specific uses and tests other than EN standards, reference should be made to the relevant sections of this guidance.

It should be noted that although this Guidance is mainly based on EN standards, there are some cases where there are discrepancies amongst the EN tests and in such cases the ECHA Guidance should be followed as the leading guidance. Where noted these are identified in the table.

The reader is strongly advised to check whether there are new versions of the standards on the website of the CEN: www.cen.eu.

It should be noted that if tests other than CEN standards (notably when no CEN tests are available) are used, and pass criteria are available, these should be met (unless stated differently in this guidance). When the test does not provide pass criteria, the criteria in this table can be taken into account as guidance for what level of reduction is normally required.

In all cases, deviations from these standards are possible but should be justified in the application.

Regarding the table for PT05, it should be noted that the text in Section 5.4.5 (PT 5) of this Guidance document is only “preliminary draft text” and has not been reviewed or revised to address written PEG comments received and the section is currently under review within the “Disinfectants Project”. In the meantime, the “preliminary draft text” is available to readers for information and it is for this reason that a table for PT05 is included, but this will be reviewed when Section 5.4.5 of the Guidance is reviewed.
Appendix 5. Examples of viruses sorted according to their presence in the human body in case of virus infection

These viruses may contaminate hands, instruments, other surfaces and textiles.

NOTE 1 This list is not exhaustive.

NOTE 2 Enveloped viruses are in **bold**.

**Table 31: Examples of viruses**

<table>
<thead>
<tr>
<th><strong>Blood</strong></th>
<th><strong>Respiratory tract</strong></th>
<th><strong>Neuronal tissue, ear, nose &amp; eye</strong></th>
<th><strong>Gastro-intestinal</strong></th>
<th><strong>Skin, breast and/or milk</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Enterovirus</td>
<td>Filoviridae</td>
<td>Influenza Virus</td>
<td>Adenovirus (Mast-)</td>
<td>Adenovirus(Mast-)</td>
</tr>
<tr>
<td>Flavivirus</td>
<td>Flaviviridae</td>
<td>Paramyxoviridae</td>
<td>Coronavirus</td>
<td>Caliciviridae</td>
</tr>
<tr>
<td>Herpesviridae</td>
<td>Herpesviridae</td>
<td>Rubella Virus</td>
<td>Enterovirus</td>
<td>Human Immunodeficiency Virus (HIV)</td>
</tr>
<tr>
<td>Hepatitis A Virus (HAV)</td>
<td>Measles Virus</td>
<td>Human Immunodeficiency Virus (HIV)</td>
<td>Hepatitis A Virus (HAV)</td>
<td>Enterovirus</td>
</tr>
<tr>
<td>Hepatitis B virus (HBV)</td>
<td>Poxviridae</td>
<td></td>
<td>Hepatitis E Virus (HEV)</td>
<td></td>
</tr>
<tr>
<td>Spleen and lymph nodes (see also blood)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>----------------------------------------</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Human T Cell Leukaemia Virus (HTLV)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Human Immunodeficiency Virus (HIV)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Dental procedure</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adenovirus (Mast-)</td>
</tr>
<tr>
<td>Enterovirus</td>
</tr>
<tr>
<td><strong>Herpesviridae</strong></td>
</tr>
<tr>
<td><strong>Hepatitis B virus (HBV)</strong></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Urogenital tract</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Hepatitis B Virus (HBV)</strong></td>
</tr>
<tr>
<td><strong>Herpesviridae</strong></td>
</tr>
<tr>
<td><strong>Human Immunodeficiency Virus (HIV)</strong></td>
</tr>
</tbody>
</table>

**Reference:**
Appendix 6. Selection of recommended tests for solid materials (excluding wood-preservatives)\(^42\)

Table 32: Selection of recommended tests for solid materials (excluding wood-preservatives)

<table>
<thead>
<tr>
<th>Standard Method + section reference</th>
<th>Title</th>
<th>Description</th>
<th>Possible application area</th>
</tr>
</thead>
<tbody>
<tr>
<td>ISO 22196, Section 5.4.2.2</td>
<td>Measurement of antibacterial activity on plastics and other non-porous surfaces</td>
<td>Test to measure inhibition of bacterial growth on plastic material used in wet or humid conditions.</td>
<td>Treated articles in PT 2, 3, 4, with a claim to protect people/animals by inhibition of bacterial growth.</td>
</tr>
<tr>
<td>Section 5.4.2.3, Figure 4</td>
<td>Simulated Splash Model Non-Porous Materials</td>
<td>Test to measure killing on contact for non-porous material when the contaminant is spread by splashes. Speed of required effect (5-60 min) depends on claim.</td>
<td>Treated articles in PT 2, 3, 4, with a claim to protect people/animals by killing on contact to prevent cross-contamination</td>
</tr>
<tr>
<td>Section 5.4.2.3, Figure 5</td>
<td>Simulated Splash Model Porous Materials</td>
<td>Test to measure killing on contact for porous material when the contaminant is spread by splashes. Speed of required effect (5-60 min) depends on claim.</td>
<td>Treated articles in PT 2, 3, 4, with a claim to protect people/animals by killing on contact to prevent cross-contamination</td>
</tr>
<tr>
<td>Section 5.4.2.3, Figure 6</td>
<td>Printing Model</td>
<td>Test to measure killing on contact for non-porous material when the contaminant is spread by e.g. hand-contact. Speed of required effect (5-60 min) depends on claim.</td>
<td>Treated articles in PT 2, 3, 4, with a claim to protect people/animals by killing on contact to prevent cross-contamination</td>
</tr>
<tr>
<td>BS 3900 Part G6, Section 5.5.8.1</td>
<td>Methods of test for paints. Part G6: Assessment of resistance to fungal growth</td>
<td>Painted panels inoculated with a mixture of spores of fungi known to colonise paints exposed to humid conditions for up to 12 weeks should show</td>
<td>PT 7</td>
</tr>
</tbody>
</table>

\(^42\) These tests are not necessarily appropriate for all claims and materials. Tests have to be chosen depending on the claim made, the materials used and the conditions of use foreseen for the treated material/article.
<table>
<thead>
<tr>
<th>Standard Method + section reference</th>
<th>Title</th>
<th>Description</th>
<th>Possible application area</th>
</tr>
</thead>
<tbody>
<tr>
<td>ASTM G21-09, Section 5.5.8.2</td>
<td>Standard Practice for Determining Resistance of Synthetic Polymeric Materials to Fungi</td>
<td>The synthetic polymer portion of plastic materials is usually fungus-resistant in that it does not serve as a carbon source for the growth of fungi. It is generally the other components, such as plasticizers, cellulosics, lubricants, stabilizers, and colorants, that are responsible for fungus attack on plastic materials.</td>
<td>PT 7, 9</td>
</tr>
<tr>
<td>ISO 846: 1997, Section 5.5.8.2</td>
<td>Plastics - Evaluation of the action of microorganisms</td>
<td>Method for determining the deterioration of plastics due to the action of fungi and soil microorganisms by visual appearance, changes in mass or changes in physical properties. The aim is not to determine the biodegradability of plastics. Includes even a soil burial variant. Note: the section covering bacteria is not considered to be useful.</td>
<td>PT 7, 9</td>
</tr>
<tr>
<td>ISO 16869:2008, Section 5.5.8.2</td>
<td>Plastics - Assessment of the effectiveness of fungistatic compounds in plastics formulations</td>
<td>Method for determining the effectiveness of fungistatic compounds in protecting susceptible ingredients like plasticizers, stabilizers, etc., in plastics formulations. A minimum diffusion of the fungicide out of the matrix is necessary as the spores are added in an agar-layer. Evaluation by visual examination.</td>
<td>PT 7, 9</td>
</tr>
<tr>
<td>BS EN 60068-2-10:2005, Section 5.5.8.1</td>
<td>Environmental testing. Tests. Test J and</td>
<td>Test for fungal and microbial resistance applicable to a wider range of materials</td>
<td>PT 7, 9</td>
</tr>
<tr>
<td>Standard Method + section reference</td>
<td>Title</td>
<td>Description</td>
<td>Possible application area</td>
</tr>
<tr>
<td>-----------------------------------</td>
<td>-------</td>
<td>-------------</td>
<td>---------------------------</td>
</tr>
<tr>
<td>OECD (OECD ENV/JM/MONO(2014)18 Section 5.5.8.5.2</td>
<td>Guidance Document for Quantitative Method for Evaluating Antibacterial Activity of Porous and Non-Porous Antibacterial Treated Materials.</td>
<td>Method for measuring the inhibition of bacterial growth or metabolism of porous and non-porous materials that have been treated with a biocide.</td>
<td>Anti-odour testing for textiles, PT 9</td>
</tr>
<tr>
<td>IBRG TEX13-005.4, Section 5.5.8.5.2</td>
<td>Tier 1 Textile Method Antibacterial Properties</td>
<td>Method to determine the basic antibacterial properties of textiles and porous materials and articles treated with a biocide.</td>
<td>Anti-odour testing for textiles, PT 9</td>
</tr>
</tbody>
</table>
## Appendix 7. Selection of recommended tests for liquid materials

### Table 33: Selection of recommended tests for liquid materials

<table>
<thead>
<tr>
<th>Reference + section reference</th>
<th>Title</th>
<th>Description</th>
<th>Possible application area</th>
</tr>
</thead>
<tbody>
<tr>
<td>IBRG P 16-001.2, Section 5.5.7</td>
<td>Tier 1 Wet State Paint Method</td>
<td>A Method for Determining the Basic Efficacy of Biocidal Active Substances in aqueous based paints.</td>
<td>PT 6</td>
</tr>
<tr>
<td>IBRG PDG 16-001.2, Section 5.5.7</td>
<td>Tier 1 Polymer dispersion Method</td>
<td>A Method for Determining the Basic Efficacy of Biocidal Active Substances used in polymer dispersions.</td>
<td>PT 6</td>
</tr>
<tr>
<td>IBRG PDG 16-007.2, Section 5.5.7</td>
<td>Tier 1 Basic Efficacy Method for Biocidal Active Substances used to Preserve Aqueous-Based Products</td>
<td>Method for determining the basic efficacy of biocidal active substances for in-can preservation in aqueous based products</td>
<td>PT 6</td>
</tr>
<tr>
<td>IBRG FFG 16-001.4, Section 5.5.13</td>
<td>Tier 1 Metal Working Fluids Method</td>
<td>Method for determining the basic efficacy of biocidal active substances in aqueous based metalworking fluids.</td>
<td>PT 13</td>
</tr>
</tbody>
</table>

43 These tests are not necessarily appropriate for all claims and materials. Tests have to be chosen depending on the claim made, the materials used and the conditions of use foreseen for the treated material/article.
Appendix 8. Commonly Used Methods to Measure the Effects of Preservative/Curative Action in Liquid Matrices

Table 34: Commonly Used Methods to Measure the Effects of Preservative/Curative Action in Liquid Matrices

<table>
<thead>
<tr>
<th>Reference</th>
<th>Title</th>
<th>Description</th>
<th>PT</th>
</tr>
</thead>
<tbody>
<tr>
<td>ASTM D2574-06</td>
<td>Standard Test Method for Resistance of Emulsion Paints in the Container to Attack by Microorganisms</td>
<td>This test method covers the determination of the relative resistance of emulsion paints to attack in the container by microorganisms.</td>
<td>6</td>
</tr>
<tr>
<td>ASTM D4783-01e1</td>
<td>Standard Test Methods for Resistance of Adhesive Preparations in Container to Attack by Bacteria, Yeast, and Fungi</td>
<td>Determination of the resistance of liquid adhesive preparations to microbial attack in the container by challenging adhesive specimens with cultures of bacteria, yeast, or fungi, and checking for their ability to return to sterility. These test methods return qualitative results.</td>
<td>6</td>
</tr>
<tr>
<td>ASTM E1259-05</td>
<td>Standard Practice for Evaluation of Antimicrobials in Liquid Fuels Boiling Below 390°C</td>
<td>The procedure should be used to evaluate the relative efficacy of microbicides in liquid fuels boiling below 390°C. The effect of environmental conditions, such as a variety of fuel additives, metal surfaces, and climatology, are variables that can be included in specific tests using this protocol.</td>
<td>6</td>
</tr>
<tr>
<td>SABS 1102 (1987)</td>
<td>Bacterial efficacy of biocides used in water-based emulsion paints</td>
<td>Efficacy test for in can preservatives in paints (emulsion) against bacteria.</td>
<td>6</td>
</tr>
<tr>
<td>NF X41-520 March 1968</td>
<td>Protection. Testing method for resistance of paints to microorganisms and their protective power.</td>
<td></td>
<td>6</td>
</tr>
<tr>
<td>ASTM E2275-03e1 (replaces D3946 and E686)</td>
<td>Standard Practice for Evaluating Water-Miscible Metalworking Fluid. Bioresistance and Antimicrobial Pesticide Performance</td>
<td>Laboratory procedures for rating the relative inherent bioresistance of water-miscible metalworking fluids, the bioresistance attributable to augmentation with antimicrobial pesticides or both, for determining the need for microbicide addition prior to or during fluid use in metalworking systems and for evaluating microbicide performance. Relative bioresistance is determined by challenging metalworking fluids with a biological inoculum that may either be characterized (comprised of one or more known biological cultures) or uncharacterized (comprised of biologically</td>
<td>13</td>
</tr>
</tbody>
</table>

44 Please note: The methods listed are not necessarily appropriate in all cases. Their applicability depends on the claim made, the materials used and the conditions of use for the treated material/article. These methods are listed to give an overview for the assessor when and where a method is meaningful to demonstrate a claim and where its limits are.
<table>
<thead>
<tr>
<th>Reference</th>
<th>Title</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ASTM E979-91(2004)</td>
<td>Standard Test Method for Evaluation of Antimicrobial Agents as Preservatives for Invert Emulsion and Other Water Containing Hydraulic Fluids</td>
<td>This laboratory test method is designed to evaluate the utility and effectiveness of antimicrobial agents intended to control microbial growth in invert emulsion and other water containing hydraulic fluids.</td>
</tr>
<tr>
<td>ASTM WK8252</td>
<td>New Standard Test Method for Determining Resistance of Aqueous Metalworking Fluids towards Non-Tuberculous, Rapidly Growing Mycobacteria</td>
<td>Determines the relative bioresistance of aqueous metalworking fluids towards non-tuberculous (NTM), rapidly growing (RGM), environmental mycobacteria by challenging them with a mycobacterial inoculum isolated from actual spoiled metalworking fluid field samples from the user’s site. In order to simulate field conditions, another challenge inoculum consisting of a mixture of common metalworking fluid spoilage microorganisms originating from actual MWF field samples is also used</td>
</tr>
<tr>
<td>Rawlinson and Shennan, 1987.</td>
<td>A recirculating test rig for the investigation of metal-working fluid spoilage. In Industrial microbiological testing 1987 pp. 227-231. Edited by Hopton and, J.W.; Hill, E.C.</td>
<td>The method described, which attempts to simulate the conditions under which a metal working fluid will be used in service, has been used extensively for the testing of new product formulations and the evaluation of biocides.</td>
</tr>
<tr>
<td>UK MOD 91-70 issue (1990)</td>
<td>Cutting fluid, soluble, biostable joint service designation ZX-9</td>
<td></td>
</tr>
</tbody>
</table>
Appendix 9. Commonly Used Methods to Measure the Effects of Protecting Material\(^{45}\)

<table>
<thead>
<tr>
<th>Reference</th>
<th>Title</th>
<th>Description</th>
<th>Major Principle/Use</th>
</tr>
</thead>
<tbody>
<tr>
<td>EN 14119:2003</td>
<td>Testing of textiles – Evaluation of the action of microfungi</td>
<td>The test is designed to determine the susceptibility of textiles to fungal growth. Assessment is by visual rating and measurement of tensile strength.</td>
<td>Agar plate test</td>
</tr>
<tr>
<td>AATCC 30-2004</td>
<td>Antifungal activity, Assessment on textile materials: mildew and rot resistance of textile materials</td>
<td>The two purposes of the test are to determine the susceptibility of textiles to microfungi and to evaluate the efficacy of fungicides on textiles.</td>
<td>Agar plate test</td>
</tr>
<tr>
<td>DIN 53931</td>
<td>Testing of textiles; determination of resistance of textiles to mildew; growth test</td>
<td>The test determines the efficacy of treatments for prevention of fungal growth on/in textiles. It also allows the performance testing of a treatment after UV irradiation, leaching etc.</td>
<td>Agar plate test</td>
</tr>
<tr>
<td>MIL-STD-810F</td>
<td>Environmental Engineering considerations and laboratory tests; Method 508.5 FUNGUS</td>
<td>The purpose of the method is to assess the extent to which a material will support fungal growth and how performance of that material is affected by such growth.</td>
<td>Humid chamber test (90 to 99% humidity)</td>
</tr>
<tr>
<td>BS 6085:1992</td>
<td>Determination of the resistance of textiles to microbial deterioration</td>
<td>The purpose of the method is to assess the extent to which a material will support fungal/bacterial growth and how performance of the material is affected by such growth. Visual Assessment and measurement of tensile strength.</td>
<td>a) soil burial test; b) agar plate test; c) humid chamber test</td>
</tr>
<tr>
<td>EN ISO 11721-1 (2001)</td>
<td>Textiles - Determination of resistance of cellulose-containing textiles to micro-organisms: Soil burial test Part 1: Assessment of rot retarding finishing</td>
<td>The test is designed to determine the susceptibility of cellulose containing textiles against deterioration by soil micro-organisms. Preserved and unpreserved textiles are compared. Visual Assessment and measurement of tensile strength.</td>
<td>Soil burial test</td>
</tr>
</tbody>
</table>

\(^{45}\) Please note: The methods listed are not necessarily appropriate in all cases. Their applicability depends on the claim made, the materials used and the conditions of use for the treated material/article. These methods are listed to give an overview for the assessor when and where a method is meaningful to demonstrate a claim and where its limits are.
<table>
<thead>
<tr>
<th>Reference</th>
<th>Title</th>
<th>Description</th>
<th>Major Principle/Use</th>
</tr>
</thead>
<tbody>
<tr>
<td>EN ISO 11721-2</td>
<td>Textiles - Determination of resistance of cellulose-containing textiles to micro-organisms: Soil burial test Part 2: Identification of long-term resistance of a rot retardant finish</td>
<td>The test identifies the long-term resistance of a rot-retardant finish against the attack of soil inhabiting micro-organisms. It allows to make a distinction between regular long-term resistance and increased long-term resistance. Visual Assessment and measurement of tensile strength</td>
<td>Soil burial test</td>
</tr>
<tr>
<td>(2003)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>BS 2011 :</td>
<td>Basic environmental testing procedures</td>
<td>Mould growth test to show the susceptibility of a material towards colonization by fungi.</td>
<td>Humid chamber test (90 to 99% humidity)</td>
</tr>
<tr>
<td>Part 2.1J</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(IEC 68-2-10)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AS 1157.2 -</td>
<td>Australian Standard - Methods of Testing Materials for Resistance to Fungal Growth Part 2: Resistance of Textiles to Fungal Growth. Section 1 - Resistance to Surface Mould Growth.</td>
<td>Test specimens are inoculated with a suspension of spores of <em>Aspergillus niger</em> and then incubated on the surface of a mineral salts based agar for 14 days and then assessed for growth. Both leached and unleached specimens are examined. Glass rings are employed to hold the specimens in intimate contact with agar when necessary. Specimens are examined for the presence of surface mould growth.</td>
<td>Agar plate test</td>
</tr>
<tr>
<td>1999</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AS 1157.3 -</td>
<td>Australian Standard - Methods of Testing Materials for Resistance to Fungal Growth Part 2: Resistance of Cordage and Yarns to Fungal Growth.</td>
<td>Test specimens are inoculated with a suspension of spores of <em>Chaetomium globosum</em> and then incubated on the surface of a mineral salts based agar for 14 days and then assessed for growth. Both leached and unleached specimens are examined and exposed samples are subjected to a tensile strength test. Glass rings are employed to hold the specimens in intimate contact with agar when necessary.</td>
<td>Agar plate test</td>
</tr>
<tr>
<td>1999</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AS 1157.4 -</td>
<td>Australian Standard - Methods of Testing Materials for Resistance to Fungal Growth Part 2: Resistance of Textiles to Fungal Growth. Section 2 - Resistance to Cellulolytic Fungi.</td>
<td>Test specimens are inoculated with a suspension of spores of <em>Chaetomium globosum</em> and then incubated on the surface of a mineral salts based agar for 14 days and then assessed for growth. Both leached and unleached specimens are examined and exposed samples are subjected to a tensile strength test. Glass rings are employed to hold the specimens in intimate contact with agar when necessary.</td>
<td>Agar plate test</td>
</tr>
<tr>
<td>1999</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### Table II: Methods used to Examine the Resistance to Biodeterioration: Geotextile

<table>
<thead>
<tr>
<th>Reference</th>
<th>Title</th>
<th>Description</th>
<th>Major Principle</th>
</tr>
</thead>
<tbody>
<tr>
<td>EN 12225:2000</td>
<td>Geotextiles and Geotextiles-related products - Method for determining the microbiological resistance by a soil burial test</td>
<td>The test is designed to determine the susceptibility of geotextiles and related products to deterioration by soil microorganisms. Visual Assessment and measurement of tensile strength.</td>
<td>Soil burial test</td>
</tr>
</tbody>
</table>

### Table III: Methods used to Examine the Antimicrobial Activity and Microbial Resistance of Paper etc.

<table>
<thead>
<tr>
<th>Reference</th>
<th>Title</th>
<th>Description</th>
<th>Major Principle</th>
</tr>
</thead>
<tbody>
<tr>
<td>DIN EN 1104 - 05</td>
<td>Paper and board intended to come into contact with foodstuffs Determination of transfer of antimicrobial constituents</td>
<td>A minimum of 20 replicate sub-samples (each 10 - 15 mm in diameter) taken from 10 samples of a batch of paper are placed in intimate contact with nutrient agar plates inoculated with either <em>Bacillus subtilis</em> or <em>Aspergillus niger</em> and incubated at 30° C for 7 days and at 25° C for 8 - 10 days respectively.</td>
<td>Zone Diffusion Assay.</td>
</tr>
<tr>
<td>ASTM D 2020-03</td>
<td>Standard Test Methods for Mildew (Fungus) Resistance of Paper and Paperboard - Direct Inoculation</td>
<td>Replicate samples (3) are inoculated with a suspension of fungal spores and then incubated on the surface of a minimal mineral-salts medium to determine if they support fungal growth.</td>
<td>Biodeterioration Test.</td>
</tr>
<tr>
<td>ASTM D 2020-03</td>
<td>Standard Test Methods for Mildew (Fungus) Resistance of Paper and Paperboard - Soil Burial</td>
<td>Replicate samples (5) are buried in soil for 14 days and then examined for the deterioration compared with unburied samples for both physical deterioration and loss of tensile strength.</td>
<td>Biodeterioration/ Biodegradation Test.</td>
</tr>
<tr>
<td>AS 1157.7 - 1999</td>
<td>Australian Standard - Methods of Testing Materials for Resistance to Fungal Growth Part 6: Resistance of Papers and Paper Products to Fungal Growth.</td>
<td>Test specimens are placed on the surface of a mineral-salts based agar and then both the specimen and the agar are inoculated with a suspension of spores of a range of fungi. They are then incubated for 14 days and then assessed for growth. Growth on the specimen is assessed.</td>
<td>Agar plate test</td>
</tr>
<tr>
<td>AS 1157.5 - 1999</td>
<td>Australian Standard - Methods of Testing Materials for Resistance to Fungal Growth Part 5: Resistance of Timber to Fungal Growth.</td>
<td>Test specimens are placed on the surface of a mineral salts based agar and then both the specimen and the agar are inoculated with a suspension of spores of a range of fungi. They are then incubated for 14 days and then assessed for growth. Growth on the specimen is assessed.</td>
<td>Agar plate test</td>
</tr>
</tbody>
</table>

Test specimens are placed on the surface of a mineral salts based agar and then both the specimen and the agar are inoculated with a suspension of spores of a range of fungi. They are then incubated for 14 days and then assessed for growth. Both leached and unleached specimens are examined. Growth on specimens is assessed. Sucrose containing media is employed where true controls cannot be obtained.

### Table IV: Methods used to Examine the Resistance to Biodeterioration: Plastics

<table>
<thead>
<tr>
<th>Reference</th>
<th>Title</th>
<th>Description</th>
<th>Major Principle</th>
</tr>
</thead>
<tbody>
<tr>
<td>ASTM D 5338 - 92</td>
<td>Humid chamber test (90 to 99% humidity)</td>
<td>Humid chamber test (90 to 99% humidity)</td>
<td>Biodegradability test</td>
</tr>
<tr>
<td>ASTM E 1428 - 99</td>
<td>Humid chamber test (90 to 99% humidity)</td>
<td>Humid chamber test (90 to 99% humidity)</td>
<td>Agar plate test</td>
</tr>
<tr>
<td>ASTM G 22 - 76</td>
<td>Agar plate test</td>
<td>Agar plate test</td>
<td>Agar plate test</td>
</tr>
<tr>
<td>ASTM G 21 - 96</td>
<td>Agar plate test</td>
<td>Agar plate test</td>
<td>Agar plate test</td>
</tr>
<tr>
<td>ASTM G 29 - 96</td>
<td>Agar plate test</td>
<td>Agar plate test</td>
<td>Biofouling test</td>
</tr>
<tr>
<td>EN 14047:2002</td>
<td>Agar plate test</td>
<td>Agar plate test</td>
<td>Biodegradability test</td>
</tr>
<tr>
<td>EN 14048:2002</td>
<td>Humid chamber test (90 to 99% humidity)</td>
<td>Humid chamber test (90 to 99% humidity)</td>
<td>Biodegradability test</td>
</tr>
<tr>
<td>ISO 846:1997</td>
<td>Humid chamber test (90 to 99% humidity)</td>
<td>Humid chamber test (90 to 99% humidity)</td>
<td>Agar plate test; soil burial test</td>
</tr>
<tr>
<td>EUROCAE ED-14B/ RTCA DO 160B</td>
<td>Agar plate test</td>
<td>Agar plate test</td>
<td>Humid chamber test (90 to 99% humidity)</td>
</tr>
<tr>
<td>MIL-STD-810F</td>
<td>Environmental Engineering considerations and laboratory tests; Method 508.5 FUNGUS</td>
<td>The purpose of the method is to assess the extent to which a material will support fungal growth and how performance of the material is affected by such growth.</td>
<td>Humid chamber test (90 to 99% humidity)</td>
</tr>
<tr>
<td>BS 2011 : Part 2.1J (identical)</td>
<td>Basic environmental testing procedures</td>
<td>Mould growth test to show the susceptibility of a material towards the colonization by fungi.</td>
<td>Humid chamber test (90 to 99% humidity)</td>
</tr>
</tbody>
</table>
### Guidance on the BPR: Volume II Parts B+C
#### Version 1.0  February 2017

### Table V: Methods used to Examine the Antimicrobial Activity and Microbial Resistance of Surface Coatings & Adhesives

<table>
<thead>
<tr>
<th>Reference</th>
<th>Title</th>
<th>Description</th>
<th>Major Principle</th>
</tr>
</thead>
<tbody>
<tr>
<td>with IEC 68-2-10)</td>
<td>Plastics - Assessment of the effectiveness of fungistatic compounds in plastics formulations</td>
<td>A specimen is placed on a nutrient-salt-agar (without additional carbon source) in a petri dish and overlayed with the same agar containing fungal spores. Rate of growth on the specimen is visually assessed.</td>
<td>Agar plate test</td>
</tr>
<tr>
<td>ISO 16869:2008</td>
<td>Australian Standard - Methods of Testing Materials for Resistance to Fungal Growth Part 4: Resistance of Coated Fabrics and Electronic Boards to Fungal Growth</td>
<td>Test specimens are inoculated with a suspension of spores of <em>Chaetomium globosum</em> and then incubated on the surface of a mineral salts based agar for 14 days and then assessed for growth. Both leached and unleached specimens are examined and exposed samples are subjected to a tensile strength test. Glass rings are employed to hold the specimens in intimate contact with agar when necessary.</td>
<td>Agar plate test</td>
</tr>
<tr>
<td>AS 1157.4 - 1999</td>
<td>Australian Standard - Methods of Testing Materials for Resistance to Fungal Growth Part 4: Resistance of Coated Fabrics and Electronic Boards to Fungal Growth</td>
<td>Test specimens are inoculated with a suspension of spores of a range of fungi and then incubated on the surface of a mineral salts based agar for 14 days and then assessed for growth. Both leached and unleached specimens are examined. Glass rings are employed to hold the specimens in intimate contact with agar when necessary.</td>
<td>Agar plate test</td>
</tr>
<tr>
<td>BS3900 Part G6</td>
<td>Assessment of resistance to fungal growth</td>
<td>Replicate test panels coated with the test coating are inoculate with a suspension of spores of fungi known to grow on the surface of paints and related materials. The samples are then incubated under conditions suitable to support fungal growth (23 ± 2°C and high humidity/surface condensation). In the</td>
<td>Biodeterioration Test</td>
</tr>
<tr>
<td>Reference</td>
<td>Title</td>
<td>Description</td>
<td>Major Principle</td>
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</tr>
<tr>
<td>ASTM D3273-12</td>
<td>Standard Test Method for Resistance to Growth of Mold on the Surface of Interior Coatings in an Environmental Chamber</td>
<td>Replicate test panels coated with the test coating are inoculated with a suspension of spores of fungi known to grow on the surface of paints and related materials. The samples are then incubated under conditions suitable to support fungal growth.</td>
<td>Biodeterioration Test</td>
</tr>
<tr>
<td>WK4201</td>
<td>Standard Test Method for Resistance to Mold Growth on Building Products in an Environmental Chamber</td>
<td>Replicate test panels coated with the test coating are inoculated with a suspension of spores of fungi known to grow on the surface of paints and related materials. The samples are then incubated under conditions suitable to support fungal growth.</td>
<td>Biodeterioration Test</td>
</tr>
<tr>
<td>ASTM D5590-94</td>
<td>Standard Test Method for Determining the Resistance of Paint Films and Related Coatings to Fungal Defacement by Accelerated Four-Week Agar Plate Assay</td>
<td></td>
<td>Agar Plate Test</td>
</tr>
<tr>
<td>SS345 Appendix 9</td>
<td>Formal Title Missing at Present</td>
<td>The bottom of glass petri dishes are coated with paint. After drying, a culture of algae in a suitable growth liquid medium is placed into the dish and incubated under conditions suitable for algal growth.</td>
<td>Biodeterioration Test.</td>
</tr>
<tr>
<td>EN 15457:2007</td>
<td>Paints and varnishes – Laboratory method for testing the efficacy of film preservatives in a coating against fungi</td>
<td>Coatings are applied to glass fibre discs and then placed in intimate contact with the surface of nutrient agar plates. The coatings and surrounding media are then inoculated with a mixed suspension of spores of 4 fungal species selected from a list of 10. The plates are then incubated at 24°C for 21 days and then assessed for growth using a rating scale. The test is intended to support claims that a biocide</td>
<td>Zone Diffusion Assay</td>
</tr>
<tr>
<td>Reference</td>
<td>Title</td>
<td>Description</td>
<td>Major Principle</td>
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</tr>
<tr>
<td>AS 1157.10 - 1999</td>
<td>Australian Standard - Methods of Testing Materials for Resistance to Fungal Growth Part 10: Resistance of Dried or Cured Adhesives to Fungal Growth</td>
<td>Test materials coated onto glass microscope slides are inoculated with a suspension of spores of a range of fungal species and then incubated on the surface of a mineral salts based agar for 14 days and then assessed for growth.</td>
<td>Agar plate test</td>
</tr>
<tr>
<td>EN 15458:2007</td>
<td>Paints and varnishes – Laboratory method for testing the efficacy of film preservatives in a coating against algae</td>
<td>Coatings are applied to glass fibre discs and then placed in intimate contact with the surface of nutrient agar plates. The coatings and surrounding media are then inoculated with a mixed suspension of 3 algal species selected from a list of 5. The plates are then incubated at 23°C under illumination (16 hour day length, 1000 Lux) for 35 days and then assessed for growth using a rating scale. The test is intended to support claims that a biocide can have an effect in a surface coating in support of its listing in the relevant use category within the EU BPD. It is not intended to assess the performance of surface coatings.</td>
<td>Zone Diffusion Assay</td>
</tr>
<tr>
<td>VdL RL06</td>
<td>Guideline to Evaluate the Resistance of Coating Materials against Mold Growth</td>
<td>Coatings are applied to paper discs and then placed in intimate contact with the surface of nutrient agar plates. The coatings and surrounding media are then inoculated with a mixed suspension of spores of <em>A niger</em> and <em>Penicillium funiculosum</em>. The plates are then incubated at 28°C for 3 weeks and assessed for growth using a rating scale after 1, 2 and 3 weeks. Coatings for exterior use and ‘wet’ applications are leached in water prior to testing.</td>
<td>Zone Diffusion Assay/Humid Chamber Test</td>
</tr>
<tr>
<td>VdL RL07</td>
<td>Guideline to Evaluate the Resistance of Coating Materials against Mold Growth</td>
<td>Coatings are applied to paper discs and then placed in intimate contact with the surface of nutrient agar plates. The coatings and surrounding media are then inoculated with a mixed suspension of <em>Scenedesmus vacuolaris</em> and <em>Stichococcus bacillaris</em>. The plates are then incubated at 23°C for 3 weeks under illumination (16 hour day length, 1000 Lux) and assessed for growth using a rating scale after 1, 2 and 3 weeks. Coatings for exterior use and ‘wet’ applications are leached in water prior to testing.</td>
<td>Zone Diffusion Assay/Humid Chamber Test</td>
</tr>
</tbody>
</table>
### Table VI: Methods used to Examine the Antimicrobial Activity of Textiles (fabric, yarn or pile/wadding)

<table>
<thead>
<tr>
<th>Reference</th>
<th>Title</th>
<th>Description</th>
<th>Major Principle</th>
</tr>
</thead>
<tbody>
<tr>
<td>JIS L 1902: 2008</td>
<td>Testing Method for Antibacterial Activity of Textiles Qualitative Test</td>
<td>Three replicate samples of fabric, yarn or pile/wadding are placed in intimate contact with the surface of agar plates that have been inoculated with a cell suspension of either <em>Staph aureus</em> or <em>K. pneumoniae</em> and incubated at 37° C for 24 - 48 hours. The presence of and size of any zone of inhibition around the samples is then recorded.</td>
<td>Zone diffusion assay. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>JIS L 1902: 2008</td>
<td>Testing Method for Antibacterial Activity of Textiles Quantitative Test</td>
<td>Replicate samples of fabric (6 of the control and 3 of the treated) are inoculated with individual bacterial species (<em>e.g. Staph. aureus</em> and <em>K. pneumoniae</em>) suspended in a heavily diluted nutrient medium. The samples are incubated under humid conditions at 37° C for a specified contact time. Activity is assessed by comparing the size of the initial population in the control with that present following incubation. No neutraliser is employed during cell recovery.</td>
<td>Cell suspension intimate contact test. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>EN ISO 20645 - 2004</td>
<td>Textile Fabrics - Determination of the antibacterial activity - Agar plate test (ISO/FDIS 20645:2004)</td>
<td>Four replicate samples of fabric (25 ± 5 mm) are placed in intimate contact with a solid nutrient medium in a petri dish. The samples are then overlaid with molten solid nutrient media which has been inoculated with a cell suspension of either <em>Staph. aureus</em>, <em>Escherichia coli</em> or <em>K. pneumoniae</em>. The plates are then incubated for between 18 and 24 hours and the plates are then assessed for growth based on either the presence of a zone of inhibition of &gt; 1 mm or the absence/strength of the growth in the media overlaying the test specimen.</td>
<td>Zone diffusion assay. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>SN 195920</td>
<td>Examination of the Antibacterial Effect of Impregnated Textiles by the Agar Diffusion Method</td>
<td>Four replicate samples of fabric (25 ± 5 mm) are placed in intimate contact with a solid nutrient medium in a petri dish. The samples are then overlaid with molten solid nutrient media which has been inoculated with</td>
<td>Zone diffusion assay. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>Reference</td>
<td>Title</td>
<td>Description</td>
<td>Major Principle</td>
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</tr>
<tr>
<td></td>
<td></td>
<td>a cell suspension of either <em>Staph. aureus</em> or <em>E. coli</em>. The plates are then incubated for between 18 and 24 hours and the plates are then assessed as described in BS EN ISO 20645 above.</td>
<td></td>
</tr>
<tr>
<td>SN195924</td>
<td>Textile Fabrics - Determination of the Antibacterial Activity: Colony Plate Count Method</td>
<td>Fifteen replicate samples (each replicate is comprised of sufficient specimens of 25 ± 5 mm to absorb 1 ml of test inoculum) are inoculated with cells of either <em>E. coli</em> or <em>Staph. aureus</em> suspended in a liquid nutrient medium and incubated in sealed bottles for up to 24 hours at 27° C. After 0, 6 and 24 hours, 5 replicate samples are analysed for the size of the viable population present. A neutraliser is employed. An increase of 2 orders of magnitude of the population exposed to a control sample is required to validate the test. The method defines a textile as antibacterial if no more than a specified minimum level of growth is observed after 24 hours in 4 of the 5 replicate groups of samples.</td>
<td>Cell suspension intimate contact test. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>SN195921</td>
<td>Textile Fabrics - Determination of Antimycotic Activity: Agar Diffusion Plate Test</td>
<td>Replicate (4) samples of sterilised fabric (25 ± 5 mm diameter) are placed in intimate contact with a solid nutrient medium in a petri dish. Each petri dish has been prepared as a double layer. The first layer consists of 10 ml nutrient agar, the second layer of another 10 ml of the same nutrient agar to which 0.1 ml spore suspension (10⁷ ml⁻¹) of either <em>Candida albicans</em>, <em>Aspergillus niger</em>, <em>Cladosporium sphaerospermum</em> or <em>Trichophyton mentagrophytes</em> had been added. The plates are then incubated at 28° C either 2 days (<em>C. albicans</em>) or 7 days (<em>A. niger, C. sphaerospermum</em> and <em>T. mentagrophytes</em>). The test is valid when control specimens of the same material without biocide, or of a biocide-free standard specified cotton material are fully overgrown. Good antimycotic efficacy is considered to be demonstrated.</td>
<td>Zone diffusion assay. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>Reference</td>
<td>Title</td>
<td>Description</td>
<td>Major Principle</td>
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</tr>
<tr>
<td>ISO 20743</td>
<td>Textiles - Determination of antibacterial activity of antibacterial finished products: Absorption method</td>
<td>Replicate (6) samples of textile are inoculated with a standardised broth culture of either <em>Staph. aureus</em> or <em>K. pneumoniae</em> in individual tubes and then incubated at 37° C for 18 - 24 hours in closed containers. Samples are analysed for the presence of viable bacteria both before and after incubation by either total viable count or the determination of total ATP. Samples are sterilised prior to testing and a neutraliser is employed during recovery. The test is validated by growth of ( \geq 1 ) order of magnitude during the incubation period.</td>
<td>Cell suspension intimate contact test. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>ISO 20743</td>
<td>Textiles - Determination of antibacterial activity of antibacterial finished products: Transfer method</td>
<td>Replicate (6) samples of test material are placed in contact with an agar plate that has been inoculated with a specified volume of a known cell suspension of either <em>Staph. aureus</em> and <em>K. pneumoniae</em> using a 200 g weight for 1 minute. The samples are then removed. Replicate (3) samples are analysed for either the number of viable bacteria or the total ATP content both before and after incubation under humid conditions at 37° C for 24 hours. Samples are sterilised prior to testing and a neutraliser is employed during cell recovery. The test is validated by either growth of ( \geq 1 ) order of magnitude during the incubation period or by a measure of the variability of the data obtained.</td>
<td>Cell suspension intimate contact test. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>ISO 20743</td>
<td>Textiles - Determination of antibacterial activity of antibacterial finished products: Printing method</td>
<td>Replicate (6) samples of test material are either <em>Staph. aureus</em> and <em>K. pneumoniae</em> by ‘printing’ cells collected on a membrane filter onto their surface in a standardised manner. The samples are then incubated under humid conditions for 18 - 24 hours at 20° C for a specified contact time(s). Replicate (3) samples</td>
<td>‘Dry’ inoculum intimate contact test. The transfer method of inoculation could be adapted to provide some simulation data.</td>
</tr>
<tr>
<td>Reference</td>
<td>Title</td>
<td>Description</td>
<td>Major Principle</td>
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</tr>
<tr>
<td>ISO/FDIS 13629-1</td>
<td>Textiles - Determination of Antifungal Activity of Textile Products: Part 1 - Luminescence Method</td>
<td>Samples of textiles are inoculated with a suspension of fungal spores either by direct application or transfer from an agar surface and then incubated. Germination and growth of the spores is followed by measuring the ATP concentration associated with the samples. The presence of an antifungal treatment is expected to show either an inhibition of germination or a reduction in the rate of growth as indicated by reduced concentrations of ATP associated with the treated material in comparison with the untreated material.</td>
<td>Basic efficacy test that has limited use as a simulation of final use of a treated material. The transfer method of inoculation could be adapted to provide some simulation data.</td>
</tr>
<tr>
<td>ISO/WD 13629-1</td>
<td>Textiles - Determination of Antifungal Activity of Textile Products: Part 2 - Plate Count Method</td>
<td>Samples of textiles are inoculated with a suspension of fungal spores either by direct application or transfer from an agar surface and then incubated. Germination and growth of the spores is followed by measuring the number of colony forming units. The presence of an antifungal treatment is expected to show either an inhibition of germination or a reduction in the rate of growth as indicated by reduced numbers of colony forming units associated with the treated material in comparison with the untreated material.</td>
<td>Basic efficacy test that has limited use as a simulation of final use of a treated material. The transfer method of inoculation could be adapted to provide some simulation data.</td>
</tr>
</tbody>
</table>
### Table VII: Methods used to Examine the Antimicrobial Activity of Carpets

<table>
<thead>
<tr>
<th>Reference</th>
<th>Title</th>
<th>Description</th>
<th>Major Principle</th>
</tr>
</thead>
<tbody>
<tr>
<td>AATCC 174-2011</td>
<td>Antimicrobial Activity Assessment of Carpets Qualitative Antibacterial Activity</td>
<td>Petri dishes with nutrient media are inoculated with a single, diagonal streak (approx. 7.5 cm) of either <em>Staph. aureus</em> or <em>K. pneumoniae</em>. An unsterilized test specimen (25 mm x 50 mm) is placed in intimate contact and transversely across the inoculum on the agar surface. The plates are then inoculated at 37° C for 18 - 24 hours. The front and back of the carpet are tested separately. After incubation, the plates are inspected for the presence of growth both below the specimens and for any zone of inhibition surrounding the specimens. The test can also be used to test the effect of cleaning regimes. An untreated control is optional.</td>
<td>Qualitative assessment of rate of kill and zone diffusion test. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>AATCC 174-2011</td>
<td>Antimicrobial Activity Assessment of Carpets Quantitative Antibacterial Activity</td>
<td>Unsterilized specimens of carpet are pre-wetted with either sterile water or a wetting agent before being inoculated with individual suspensions of either <em>Staph. aureus</em> or <em>K. pneumoniae</em> in either a low or a high nutrient solution. The samples are then incubated in a tightly closed jar at 37° C for a specified contact time. Cells are recovered in 100 ml of a neutraliser after 0 and 6 - 24 hours of incubation. Activity is assessed by comparing the size of the initial population in the control (if used) with that present following incubation. A control is optional. When not employed, viable counts following incubation of the treated specimens alone are considered. The test can also be used to test the effect of cleaning regimes.</td>
<td>Cell suspension intimate contact test. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>AATCC 174-2011</td>
<td>Antimicrobial Activity Assessment of Carpets Quantitative Antifungal Activity</td>
<td>Petri dishes containing Sabouraud Dextrose Agar are inoculated with 1 ml of a spore suspension of <em>Aspergillus niger</em>. Immediately afterwards, specimens (38 mm diameter) of unsterile test material are placed into intimate contact with the agar. An additional 0.2 ml of the</td>
<td>Zone diffusion test/surface growth test. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>Reference</td>
<td>Title</td>
<td>Description</td>
<td>Major Principle</td>
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</tr>
<tr>
<td>WIRA Test F</td>
<td>Test Method for Assessing the Survival of Test Organisms on Floor Coverings</td>
<td>same spore suspension is also employed to inoculate the test pieces directly. The samples are then incubated at 28°C for 7 days. The back and front of the discs of carpet are tested in separate dishes. The zone of inhibition and the growth of fungus on the upper surface of the specimens are reported (no growth, microscopic growth, macroscopic growth). The test can also be used to test the effect of cleaning regimes.</td>
<td>Cell suspension intimate contact test. Potential to demonstrate the effectiveness of an antimicrobial treatment if appropriate incubation conditions are selected and addition species employed.</td>
</tr>
</tbody>
</table>

Specimens (850 mm x 350 mm) are conditioned at 20°C and 65% RH before being subjected to 2 wet and 2 dry passes using a commercial spray extraction machine or a test rig. After 24 h drying, 12 specimens (each 60 mm diameter) are cut from the carpet. An aliquot (1 ml) of a suspension of cells of *E. coli* in nutrient broth is poured onto filter paper (7 cm diameter). The filter paper is then pressed for 1 min onto the surface of the carpet using a 1 kg weight. The filter paper is then discarded. After 0, 6 and 24 hours incubation at a specified temperature the carpet’s surface is pressed onto contact plates of McConkey agar. After 24h replicate (3) plugs (10 mm) are taken from each specimen and suspended in 10 ml nutrient broth for 30 seconds and then analysed for the presence of *E. coli* by total viable count.
# Table VIII: Methods used to Examine the Antimicrobial Activity of Non-Porous Surfaces

<table>
<thead>
<tr>
<th>Reference</th>
<th>Title</th>
<th>Description</th>
<th>Major Principle</th>
</tr>
</thead>
<tbody>
<tr>
<td>JIS Z 2801:2000</td>
<td>Antimicrobial products - Test for antibacterial activity and efficacy</td>
<td>The surface of replicate sample (3 for each treatment and 6 for the blank reference material - usually 50 mm x 50 mm) are inoculated with a suspension of either <em>E. coli</em> or <em>Staph. aureus</em> in a highly diluted nutrient broth. The cell suspension is then held in intimate contact with the surface by the use of a sterile polyethylene film (usually 40 mm x 40 mm) for 24 hours at 35° C under humid conditions. The size of the population on the treated surface is then compared with the size on the control surface both prior to and after incubation. A neutraliser for certain biocide types is employed. Antibacterial activity is certified if the difference between the Log\textsubscript{10} of the population on the treated sample and that on the control surface is &gt; 2.</td>
<td>Cell suspension intimate contact test. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>ISO 22196:2011</td>
<td>Plastics - Measurement of antibacterial activity on plastics surfaces.</td>
<td>This is the current New Work Proposal at ISO created from JIS Z 2801 by the SIAA of Japan. Modification and validation is in progress in collaboration with the IBRG. Some changes are expected.</td>
<td>Cell suspension intimate contact test. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>XP G 39-010</td>
<td>Propriétés des étoffes - Étoffes et surfaces polymérisées à propriétés antibactériennes - Caractérisation et mesure de l’activité antibactérienne</td>
<td>Four replicate samples of test material are placed in contact with an agar plate that has been inoculated with a specified volume of a known cell suspension of either <em>Staph. aureus</em> and <em>K. pneumoniae</em> using a 200g weight for 1 minute. The samples are then removed. Duplicate samples are analysed for the number of viable bacteria both before and after incubation under humid conditions at 37°C for 24 hours. A neutraliser is employed during cell recovery.</td>
<td>Cell suspension intimate contact test. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>ASTM E2180-07</td>
<td>Standard Test Method for Determining the</td>
<td>Replicate (3) samples of material are inoculated with cells of either <em>Staph. aureus</em> or</td>
<td>Immobilised cell suspension intimate contact test.</td>
</tr>
<tr>
<td>Reference</td>
<td>Title</td>
<td>Description</td>
<td>Major Principle</td>
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</tr>
<tr>
<td></td>
<td>Activity of Incorporated Antimicrobial Agent(s) in Polymeric or Hydrophobic Materials</td>
<td><em>K. pneummoniae</em> suspended in molten semi-solid isotonic saline/agar. This attempts for form an ‘artificial biofilm’ which holds the suspension in intimate contact with the test surface of inherently hydrophobic materials. Samples are then incubated at a temperature similar to that intended for the final use for a specified period (usually 24 hours) under humid conditions. The size of the viable bacterial populations on the control and treated surfaces is then determined using a dilution plate count. Any effect is recorded using percent reduction calculated from the geometric means of the data. A neutraliser may be employed and sonication is used to separate the ‘biofilm’ from the test surfaces and suspend the agar gel. Subsequent imprinting of the test surface onto solid nutrient media can be performed to look for the presence of adherent viable cells.</td>
<td>Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>ASTM E2149-10</td>
<td>Standard Test Method for Determining the Antimicrobial Activity of Immobilized Antimicrobial Agents Under Dynamic Contact Conditions</td>
<td>Dynamic shake flask test. Test material is suspended in a buffer solution containing a known number of cells of <em>Klebsiella pneumoniae</em> and agitated. Efficacy is determined by comparing the size of the population both before and after a specified contact time.</td>
<td>Relies on either diffusion of antimicrobial agents from treated material into the cell suspension or due to interaction between the population and the surface of the material in suspension. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
</tbody>
</table>
## Appendix 10. Commonly Used Methods to Measure Antimicrobial Activity

### Table VI: Methods used to Examine the Antimicrobial Activity of Textiles (fabric, yarn or pile/wadding)

<table>
<thead>
<tr>
<th>Reference</th>
<th>Title</th>
<th>Description</th>
<th>Major Principle</th>
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</thead>
<tbody>
<tr>
<td>ASTM E2149-10</td>
<td>Standard Test Method for Determining the Antimicrobial Activity of Immobilized Antimicrobial Agents Under Dynamic Contact Conditions</td>
<td>Dynamic shake flask test. Test material is suspended in a buffer solution containing a known number of cells of <em>Klebsiella pneumoniae</em> and agitated. Efficacy is determined by comparing the size of the population both before and after a specified contact time.</td>
<td>Relies on either diffusion of antimicrobial from treated material into the cell suspension. Some activity may be due to interaction between the population and the surface of the material in suspension. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>AATCC 147-2011</td>
<td>Antibacterial Activity Assessment of Textile Materials: Parallel Streak Method</td>
<td>Agar plates are inoculated with 5 parallel streaks (60 mm long) of either <em>Staphylococcus aureus</em> or <em>K. pneumoniae</em>. A textile sample is then placed over the streaks and in intimate contact with the surface of the agar and incubated. Activity is assessed based on either the mean zone of inhibition over the 5 streaks or the absence of growth behind the test specimen.</td>
<td>Zone diffusion assay. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>AATCC 100-2012</td>
<td>Antibacterial Finishes on Textile Materials: Assessment of...</td>
<td>Replicate samples (sufficient to absorb 1 ml of test inoculum) of fabric are inoculated with individual bacterial species (<em>e.g.</em> <em>Staph. aureus</em> and <em>K. pneumoniae</em>) suspended in a nutrient medium. The samples are incubated under humid conditions at 37° C for a specified contact time. Activity is assessed by comparing the size of the initial population with that present following incubation. A</td>
<td>Cell suspension intimate contact test. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
</tbody>
</table>

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46 Please note: The methods listed are not necessarily appropriate in all cases. Their applicability depends on the claim made, the materials used and the conditions of use for the treated material/article. These methods are listed to give an overview for the assessor when and where a method is meaningful to demonstrate a claim and where its limits are.
<table>
<thead>
<tr>
<th>Reference</th>
<th>Title</th>
<th>Description</th>
<th>Major Principle</th>
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<tbody>
<tr>
<td>XP G 39-010</td>
<td>Propriétés des étoffes - Étoffes et surfaces polymériques à propriétés antibactériennes - Caractérisation et mesure de l’activité antibactérienne</td>
<td>Four replicate samples of test material are placed in contact with an agar plate that has been inoculated with a specified volume of a known cell suspension of either \textit{Staph. aureus} and \textit{K. pneumoniae} using a 200 g weight for 1 minute. The samples are then removed. Duplicate samples are analysed for the number of viable bacteria both before and after incubation under humid conditions at 37° C for 24 hours. A neutraliser is employed during cell recovery.</td>
<td>Cell suspension intimate contact test. The transfer method of inoculation could be adapted to provide some simulation data.</td>
</tr>
<tr>
<td>JIS L 1902: 2008</td>
<td>Testing Method for Antibacterial Activity of Textiles Qualitative Test</td>
<td>Three replicate samples of fabric, yarn or pile/wadding are placed in intimate contact with the surface of agar plates that have been inoculated with a cell suspension of either \textit{Staph. aureus} or \textit{K. pneumoniae} and incubated at 37° C for 24 - 48 hours. The presence of and size of any zone of inhibition around the samples is then recorded.</td>
<td>Zone diffusion assay. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>JIS L 1902: 2008</td>
<td>Testing Method for Antibacterial Activity of Textiles Quantitative Test</td>
<td>Replicate samples of fabric (6 of the control and 3 of the treated) are inoculated with individual bacterial species (\textit{e.g. Staph. aureus} and \textit{K. pneumoniae}) suspended in a heavily diluted nutrient medium. The samples are incubated under humid conditions at 37° C for a specified contact time. Activity is assessed by comparing the size of the initial population in the control with that present following incubation. No neutraliser is employed during cell recovery.</td>
<td>Cell suspension intimate contact test. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>EN ISO 20645 - 2004</td>
<td>Textile Fabrics - Determination of the antibacterial activity - Agar plate test (ISO/FDIS 20645:2004)</td>
<td>Four replicate samples of fabric (25 ± 5 mm) are placed in intimate contact with a solid nutrient medium in a petri dish. The samples are then overlaid with molten solid nutrient media which has been inoculated with a cell suspension of either \textit{Staph. aureus}, \textit{Escherichia coli} or \textit{K. pneumoniae}. The plates are then incubated for between 18 and 24 hours and the plates are then assessed for growth</td>
<td>Zone diffusion assay. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>Reference</td>
<td>Title</td>
<td>Description</td>
<td>Major Principle</td>
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<tr>
<td>SN 195920</td>
<td>Examination of the Antibacterial Effect of Impregnated Textiles by the Agar Diffusion Method</td>
<td>Four replicate samples of fabric (25 ± 5 mm) are placed in intimate contact with a solid nutrient medium in a petri dish. The samples are then overlaid with molten solid nutrient media which has been inoculated with a cell suspension of either <em>Staph. aureus</em> or <em>E. coli</em>. The plates are then incubated for between 18 and 24 hours and the plates are then assessed as described in BS EN ISO 20645 above.</td>
<td>Zone diffusion assay. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>SN195924</td>
<td>Textile Fabrics - Determination of the Antibacterial Activity: Colony Plate Count Method</td>
<td>Fifteen replicate samples (each replicate is comprised of sufficient specimens of 25 ± 5 mm to absorb 1 ml of test inoculum) are inoculated with cells of either <em>E. coli</em> or <em>Staph. aureus</em> suspended in a liquid nutrient medium and incubated in sealed bottles for up to 24 hours at 27° C. After 0, 6 and 24 hours, 5 replicate samples are analysed for the size of the viable population present. A neutraliser is employed. An increase of 2 orders of magnitude of the population exposed to a control sample is required to validate the test. The method defines a textile as antibacterial if no more than a specified minimum level of growth is observed after 24 hours in 4 of the 5 replicate groups of samples.</td>
<td>Cell suspension intimate contact test. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>SN195921</td>
<td>Textile Fabrics - Determination of Antimycotic Activity: Agar Diffusion Plate Test</td>
<td>Replicate (4) samples of sterilised fabric (25 ± 5 mm diameter) are placed in intimate contact with a solid nutrient medium in a petri dish. Each petri dish has been prepared as a double layer. The first layer consists of 10 ml nutrient agar, the second layer of another 10 ml of the same nutrient agar to which 0.1 ml spore suspension (10⁷ ml⁻¹) of either <em>Candida albicans</em>, <em>Aspergillus niger</em>, <em>Cladosporium sphaerospermum</em> or <em>Trichophyton mentagrophytes</em> had been added. The plates are then incubated at 28° C either 2 days (<em>C. albicans</em>) or</td>
<td>Zone diffusion assay. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>Reference</td>
<td>Title</td>
<td>Description</td>
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<tr>
<td>ISO 20743</td>
<td>Textiles - Determination of antibacterial activity of antibacterial finished products: Absorption method</td>
<td>Replicate (6) samples of textile are inoculated with a standardised broth culture of either <em>Staph. aureus</em> or <em>K. pneumoniae</em> in individual tubes and then incubated at 37° C for 18 - 24 hours in closed containers. Samples are analysed for the presence of viable bacteria both before and after incubation by either total viable count or the determination of total ATP. Samples are sterilised prior to testing and a neutraliser is employed during recovery. The test is validated by growth of &gt;1 order of magnitude during the incubation period.</td>
<td>Cell suspension intimate contact test. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>ISO 20743</td>
<td>Textiles - Determination of antibacterial activity of antibacterial finished products: Transfer method</td>
<td>Replicate (6) samples of test material are placed in contact with an agar plate that has been inoculated with a specified volume of a known cell suspension of either <em>Staph. aureus</em> and <em>K. pneumoniae</em> using a 200 g weight for 1 minute. The samples are then removed. Replicate (3) samples are analysed for either the number of viable bacteria or the total ATP content both before and after incubation under humid conditions at 37° C for 24 hours. Samples are sterilised prior to testing and a neutraliser is employed during cell recovery. The test is validated by either growth of &gt;1 order of magnitude during the incubation period or by a measure of the variability of the data obtained.</td>
<td>Cell suspension intimate contact test. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>ISO 20743</td>
<td>Textiles - Determination of antibacterial activity of antibacterial finished products: ‘Dry’ inoculum intimate contact test.</td>
<td>Replicate (6) samples of test material are either <em>Staph. aureus</em></td>
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<tr>
<td>Reference</td>
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<td>Description</td>
<td>Major Principle</td>
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<tr>
<td>ISO/FDIS 13629-1</td>
<td>Textiles - Determination of Antifungal Activity of Textile Products: Part 1 - Luminescence Method</td>
<td>Samples of textiles are inoculated with a suspension of fungal spores either by direct application or transfer from an agar surface and then incubated. Germination and growth of the spores is followed by measuring the ATP concentration associated with the samples. The presence of an antifungal treatment is expected to show either an inhibition of germination or a reduction in the rate of growth as indicated by reduced concentrations of ATP associated with the treated material in comparison with the untreated material.</td>
<td>Basic efficacy test that has limited use as a simulation of final use of a treated material. The transfer method of inoculation could be adapted to provide some simulation data.</td>
</tr>
<tr>
<td>ISO/WD 13629-1</td>
<td>Textiles - Determination of Antifungal Activity of Textile Products: Part 2 - Plate Count Method</td>
<td>Samples of textiles are inoculated with a suspension of fungal spores either by direct application or transfer from an agar surface and then incubated. Germination and growth of the spores is followed by measuring the number of colony forming units. The presence of an antifungal treatment is expected to show either an inhibition of germination or a reduction in the rate of growth as indicated by reduced numbers of colony forming units associated with the treated material in comparison with the untreated material.</td>
<td>Basic efficacy test that has limited use as a simulation of final use of a treated material. The transfer method of inoculation could be adapted to provide some simulation data.</td>
</tr>
</tbody>
</table>
### Table VII: Methods used to Examine the Antimicrobial Activity of Carpets

<table>
<thead>
<tr>
<th>Reference</th>
<th>Title</th>
<th>Description</th>
<th>Major Principle</th>
</tr>
</thead>
<tbody>
<tr>
<td>AATCC 174-2011</td>
<td>Antimicrobial Activity Assessment of Carpets Qualitative Antibacterial Activity</td>
<td>Petri dishes with nutrient media are inoculated with a single, diagonal streak (approx.7.5 cm) of either <em>Staph. aureus</em> or <em>K. pneumoniae</em>. An unsterilized test specimen (25 mm x 50 mm) is placed in intimate contact and transversely across the inoculum on the agar surface. The plates are then inoculated at 37° C for 18 - 24 hours. The front and back of the carpet are tested separately. After incubation, the plates are inspected for the presence of growth both below the specimens and for any zone of inhibition surrounding the specimens. The test can also be used to test the effect of cleaning regimes. An untreated control is optional.</td>
<td>Qualitative assessment of rate of kill and zone diffusion test. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>AATCC 174-2011</td>
<td>Antimicrobial Activity Assessment of Carpets Quantitative Antibacterial Activity</td>
<td>Unsterilized specimens of carpet are pre-wetted with either sterile water or a wetting agent before being inoculated with individual suspensions of either <em>Staph. aureus</em> or <em>K. pneumoniae</em> in either a low or a high nutrient solution. The samples are then incubated in a tightly closed jar at 37° C for a specified contact time. Cells are recovered in 100 ml of a neutraliser after 0 and 6 - 24 hours of incubation. Activity is assessed by comparing the size of the initial population in the control (if used) with that present following incubation. A control is optional. When not employed, viable counts following incubation of the treated specimens alone are considered. The test can also be used to test the effect of cleaning regimes.</td>
<td>Cell suspension intimate contact test. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>AATCC 174-2011</td>
<td>Antimicrobial Activity Assessment of Carpets Quantitative Antifungal Activity</td>
<td>Petri dishes containing Sabouraud Dextrose Agar are inoculated with 1 ml of a spore suspension of <em>Aspergillus niger</em>. Immediately afterwards, specimens (38 mm diameter) of unsterile test material are placed into intimate contact with the agar. An additional 0.2 ml of the same spore suspension is also employed to inoculate the test.</td>
<td>Zone diffusion test/surface growth test. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>Reference</td>
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<td>Description</td>
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<td>pieces directly. The samples are then incubated at 28°C for 7 days. The back and front of the discs of carpet are tested in separate dishes. The zone of inhibition and the growth of fungus on the upper surface of the specimens are reported (no growth, microscopic growth, macroscopic growth). The test can also be used to test the effect of cleaning regimes.</td>
<td></td>
</tr>
<tr>
<td>WIRA Test F</td>
<td>Test Method for Assessing the Survival of Test Organisms on Floor Coverings</td>
<td>Specimens (850 mm x 350 mm) are conditioned at 20°C and 65% RH before being subjected to 2 wet and 2 dry passes using a commercial spray extraction machine or a test rig. After 24 h drying, 12 specimens (each 60 mm diameter) are cut from the carpet. An aliquot (1 ml) of a suspension of cells of <em>E. coli</em> in nutrient broth is poured onto filter paper (7 cm diameter). The filter paper is then pressed for 1 min onto the surface of the carpet using a 1 kg weight. The filter paper is then discarded. After 0, 6 and 24 hours incubation at a specified temperature the carpet’s surface is pressed onto contact plates of McConkey agar. After 24h replicate (3) plugs (10 mm ) are taken from each specimen and suspended in 10 ml nutrient broth for 30 seconds and then analysed for the presence of <em>E. coli</em> by total viable count.</td>
<td>Cell suspension intimate contact test. Potential to demonstrate the effectiveness of an antimicrobial treatment if appropriate incubation conditions are selected and addition species employed.</td>
</tr>
</tbody>
</table>
## Table VIII: Methods used to Examine the Antimicrobial Activity of Non-Porous Surfaces

<table>
<thead>
<tr>
<th>Reference</th>
<th>Title</th>
<th>Description</th>
<th>Major Principle</th>
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<tbody>
<tr>
<td>JIS Z 2801: 2000</td>
<td>Antimicrobial products - Test for antibacterial activity and efficacy</td>
<td>The surface of replicate sample (3 for each treatment and 6 for the blank reference material - usually 50 mm x 50 mm) are inoculated with a suspension of either <em>E. coli</em> or <em>Staph. aureus</em> in a highly diluted nutrient broth. The cell suspension is then held in intimate contact with the surface by the use of a sterile polyethylene film (usually 40 mm x 40 mm) for 24 hours at 35° C under humid conditions. The size of the population on the treated surface is then compared with the size on the control surface both prior to and after incubation. A neutraliser for certain biocide types is employed. Antibacterial activity is certified if the difference between the Log$_{10}$ of the population on the treated sample and that on the control surface is &gt; 2.</td>
<td>Cell suspension intimate contact test. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>ISO 22196:2011</td>
<td>Plastics - Measurement of antibacterial activity on plastics surfaces.</td>
<td>This is the current New Work Proposal at ISO created from JIS Z 2801 by the SIAA of Japan. Modification and validation is in progress in collaboration with the IBRG. Some changes are expected.</td>
<td>Cell suspension intimate contact test. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>XP G 39-010</td>
<td>Propriétés des étoffes - Étoffes et surfaces polymères à propriétés antibactériennes - Caractérisation et mesure de l'activité antibactérienne</td>
<td>Four replicate samples of test material are placed in contact with an agar plate that has been inoculated with a specified volume of a known cell suspension of either <em>Staph. aureus</em> and <em>K. pneumoniae</em> using a 200g weight for 1 minute. The samples are then removed. Duplicate samples are analysed for the number of viable bacteria both before and after incubation under humid conditions at 37°C for 24 hours. A neutraliser is employed during cell recovery.</td>
<td>Cell suspension intimate contact test. Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>ASTM E2180-07</td>
<td>Standard Test Method for Determining the Activity of</td>
<td>Replicate (3) samples of material are inoculated with cells of either <em>Staph. aureus</em> or <em>K. pneumoniae</em> suspended in molten semi-solid</td>
<td>Immobilised cell suspension intimate contact test.</td>
</tr>
<tr>
<td>Reference</td>
<td>Title</td>
<td>Description</td>
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<td></td>
<td>Incorporated Antimicrobial Agent(s) in Polymeric or Hydrophobic Materials</td>
<td>isotonic saline/agar. This attempts for form an ‘artificial biofilm’ which holds the suspension in intimate contact with the test surface of inherently hydrophobic materials. Samples are then incubated at a temperature similar to that intended for the final use for a specified period (usually 24 hours) under humid conditions. The size of the viable bacterial populations on the control and treated surfaces is then determined using a dilution plate count. Any effect is recorded using percent reduction calculated from the geometric means of the data. A neutraliser may be employed and sonication is used to separate the ‘biofilm’ from the test surfaces and suspend the agar gel. Subsequent imprinting of the test surface onto solid nutrient media can be performed to look for the presence of adherent viable cells.</td>
<td>Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
<tr>
<td>ASTM E2149-10</td>
<td>Standard Test Method for Determining the Antimicrobial Activity of Immobilized Antimicrobial Agents Under Dynamic Contact Conditions</td>
<td>Dynamic shake flask test. Test material is suspended in a buffer solution containing a known number of cells of <em>Klebsiella pneumoniae</em> and agitated. Efficacy is determined by comparing the size of the population both before and after a specified contact time. Relies on either diffusion of antimicrobial agents from treated material into the cell suspension or due to interaction between the population and the surface of the material in suspension.</td>
<td>Basic efficacy test that has limited use as a simulation of final use of a treated material.</td>
</tr>
</tbody>
</table>
Appendix 11. Information on the principle target organisms for PT 8 as outlined in the document (5.5.8)

Fungi

Wood rotting fungi

White rot/ brown rot fungi (Basidiomycetes):
Fungi responsible for brown rot (e.g. Serpula lacrymans, Coniophora puteana) and white rot (e.g. Coriolus versicolor, Donkioporia expansa)

Soft rot fungi (mainly Ascomycetes, Deuteromycetes):
Fungi responsible for a type of rot characterised by surface softening of the wood although they also cause rot at depth (e.g. Chaetomium globosum). They are specifically significant for wood in ground contact.

Wood discolouring fungi

Sapstain:
The blue-black and brown discolouration of freshly felled logs or sawn timber have an economic importance. Sapstain causing fungi can only colonise wood as long as the sap wood contains enough water to provide solved sugars as a nutrient for these fungi ("green" wood). Therefore, these fungi can be controlled by rapid drying of the wood after felling, chemical treatments are sometimes used.

Common sapstain species include e.g. Stereum spp, blue staining species.

Blue stain cause blue to black permanent colour of variable intensity and depth mainly in the sapwood depending on the wood species. This does not result in appreciable alteration of the mechanical properties but can increase the permeability of the wood and thereby makes it more susceptible to fungal degradation.

Common blue staining species include e.g. Aureobasidium spp, Ceratocystis spp

Mould fungi:
Fungi, e.g. Aspergilus spp, Penicillium spp being evident as spots of various colours on the surface of moist wood. (for instance, as a result of high relative humidity or of condensation of water vapour). They do not significantly alter the mechanical properties of the wood but have a special significance for wood in service if discoloration is undesirable or unacceptable.

For green sawn timber, the moulds are covered by the CEN TS 15082 standard. But for the preservation of solid wood against mould, the EN 152 does not cover mould and no CEN standard is available. In that case the applicant is invited to submit relevant data (in house method, literature data...) which could be accepted by expert judgement.

Insects

Fresh wood insects
A number of insects bore and tunnel into fresh logs after they are cut and debarked. These fresh wood insects feed upon the starch reserves and can cause damages to the wood. Most of them belong to the families of Scolytidae (genus Scolytus), Cerambycidae (genus Phematodes), Lyctidae (genus Lyctus), Anobiidae (genus Anobium), Bostrychidae (genus Bostrychus).

Some other groups, belonging to the Scolytidae family, bore the fresh logs and introduce 'Ambrosia' fungi inside the gallery, resulting in wood staining (as a consequence of the development of the dark hyphae).
**Wood boring beetles (Coleoptera)**

Insects which lay their eggs in wood pores or cracks and whose larvae feed upon wood. They are present throughout Europe but the risk of attack varies greatly and is ranged from high to insignificant. The most important are *Hylotrupes bajulus*, *Anobium punctatum* and *Lyctus brunneus*.

*Hylotrupes bajulus* (House longhorn beetle)

This beetle attacks many softwood species and can cause significant structural damage. Many softwood species are affected, whereas hardwoods are not attacked. Larvae damage both the sapwood and the heartwood of non durable species.

This insect occurs throughout Europe, but is of less importance in the north and northwest of Europe. The vitality and longevity of larvae depend principally on ambient temperature and the wood moisture content.

*Anobium punctatum* (Common furniture beetle)

The larvae attack the sapwood of certain softwood and hardwood species. The damage can extend to the heartwood in some wood species and can have occasionally a structural significance impact. Its presence is particularly noted in coastal climates and where damp conditions prevail.

*Lyctus brunneus* (Powder post beetle)

The larvae attack sapwood of certain starch-containing hardwoods and have a significant impact throughout Europe for both European and imported hardwood timbers.

**Termites (Isoptera)**

Termites belong to the order Isoptera. In Europe and in the European tropical overseas regions there are three main termite families; subterranean termites (Rhinotermitidae), drywood termites (Kalotermitidae) and tree termites (Nasutitermitidae):

- *Reticulitermes* is the most common genus encountered from the Rhinotermitidae family in Europe. The main species registered are: *R. flavipes* (former *R. santonensis*), *R. grassei*, *R. lucifugus*, *R. banyulensis*, *R. balkanensis*, *R. urbis*.

They are widespread around the Mediterranean basin (Spain, France, Italy, Portugal, Balkans, and Greece) and Black Sea (Turkey, Romania), though some termite spots in the UK or Germany have been reported. Several unanswered questions remain about the origin of these termites. While some *Reticulitermes* are native to Europe, others may be related to species from eastern North America and the Middle East (Israel, Asian Turkey, etc.).

*Coptotermes* and *Heterotermes* are the main two genera belonging also to the Rhinotermite family located in the European tropical overseas regions.

- *Kalotermes flavicollis* and *Cryptotermes brevis* are the main two species of drywood termites present in Europe (especially in the coastal areas of Mediterranean countries and Canary Islands). *Cryptotermes* is a main genus belonging to drywood termites encountered in the European tropical overseas regions.

- *Nasutitermes* is the main genus belonging to the Termitidae family (tree termites) encountered in the European tropical overseas regions.
Marine borers

This term is applied to marine invertebrates such as *Limnoria* spp and *Teredo* spp which need a certain salinity of water and which hollow out extensive tunnels and cavities in wood. These organisms can cause serious damage to fixed or floating structures.

In European waters the most common marine borers are shipworm (*Teredo navalis*) and gribble (*Limnoria* spp.). Shipworm is a bivalve mollusc related to the sea snails and mussels. It is a soft, worm like animal with its shell modified into hard grinding jaws. The larvae are part of the microscopic zooplankton and swim freely in the sea until they settle on timber. They develop a shell with which they bore into the wood and lodge there, growing into large worms in holes up to 5 mm in diameter. They destroy the wood by making a massive network of galleries throughout the timber. Gribble is a small shrimp-like crustacean about 4 mm in length. It bores into the surface of the wood and lodges near the surface making numerous side burrows. The combination of this boring and wave action causes rapid erosion of marine timbers.
Appendix 12. Laboratory studies for rodenticides: bait choice test

This appendix describes a protocol of a laboratory study to determine the efficacy of an as yet unauthorised product (rodenticide) against the house mouse, brown rat and roof rat containing a bait formulation. This protocol can be applied to other target organisms (e.g. voles).

A feeding test is conducted to determine the extent to which rodents will eat the product when they are given a free choice between that and their normal food. This type of palatability test is most suited to slow-acting toxicants. The test consists of an acclimatisation period, followed by a pre-test diet take assessment, then a test period of normally\(^{47}\) 3-5 days and at least 14 days of post-treatment observation.

Pre-test period

For the test, normally 10 wild or laboratory strain rodents (5 males and 5 females) are required. Laboratory rodents should be healthy, non-pregnant adults of known strain (STATE). Preferably wild adult rodents are used. They should be healthy and obtained from free-living populations (STATE WHERE) in accordance with Directive 2010/63/EU, Articles 7 and 9 and Section A, 3.2 of Annex III. On arrival at the laboratory, the wild strains should be treated with an appropriate insecticide to kill ectoparasites and then be housed in small groups (no more than five per cage) of the same sex and treatment group if no aggressive behaviour is expected, preferably in solid floor cages with appropriate environmental enrichment. Animals may be housed individually only if scientifically justified. With wild rats especially, it is advisable to place all items (i.e. food pots) required for the test in the cage before each animal is released into it. Wild rodents should be acclimatised to laboratory conditions for at least 3 weeks to ensure that no females are pregnant when the test begins. During this time they should be offered a laboratory animal diet and water should be freely available. To encourage variation in response, animals with body weights throughout the range normally expected for the species should be used as far as possible.

Before the test period begins, it is necessary to ensure that the animals are feeding normally. Following acclimatisation, two food pots, placed either side at the front of the cage, are filled with cereals, such as wheat, broken wheat, or a wheat-based mixture or ground laboratory diet or EPA meal. All other food is removed, but water remains freely available. The quantity of food placed in each pot (STATE) should be sufficient to meet each animal's daily needs. Food uptake should be determined, therefore all unused food (i.e. food left in the pot) and scattered food must be collected and taken into account by weighing to determine how much of the food has not been eaten. All unused diet (i.e. food left in the pot and scattered food) should be discarded and the pot refilled with a fresh supply, to ensure it is palatable. This procedure should be repeated for a further 3 days and on the last day (of this pre-treatment period) the animals should be weighed. Also on the last day, the diet remaining in each pot and scattered food, is weighed and the total amount of food eaten by each rodent calculated (STATE). Any rodent not eating normally by the last day should be discarded.

Test period

The palatability test commences with 2 clean bait containers, one filled with a quantity of the test product and the other with a suitable challenge diet (e.g. an EPPO challenge

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\(^{47}\) Deviation from this norm is possible but should be explained in the application.
diet\textsuperscript{48} or standard laboratory diet). Again, the quantity in each pot should exceed the normal daily requirement for each animal. After 24 hours, the diet remaining in each pot is weighed and the total amount of food eaten by each rodent calculated. All used test and challenge diet is discarded and fresh quantities of each diet are placed in clean pots. In placing the pots back in the cage, the positions of the rodenticide and the challenge diet should be interchanged to avoid place preference. This procedure should be repeated every day during the choice period. After day 4 (3 or 5 is also acceptable) the animals should be returned to the standard laboratory diet.

Observation period

During the observation period the rodents are observed at least once per day and any signs of toxicity and mortality are recorded. Humane end-points should be applied in line with Directive 2010/63/EU to all animals showing clinical signs that can determine impending death.

Guidance Document on the recognition, assessment and use of clinical signs as humane endpoints for experimental animals used in safety evaluation (OECD, 2002) must be considered.

Results

Results should be shown as the percentage intake of rodenticide and the percentage intake of challenge diet (see section 2.2.1 for further details). Also the percentage mortality and any other symptoms should be mentioned.

Liquid bait formulations

The test must be carried out as above with the following exceptions:

- a suitable compounded laboratory diet shall be freely available;
- tap water must be used as the control bait;
- all procedures relating to the solid control and test baits must be applied instead and as appropriate to the liquid control and test baits;
- when the positions of the test and control baits are interchanged the positions of the drinking tubes, if used, should not be interchanged;
- liquid baits must be provided in containers with non-drip nozzles or suitable open pots;
- a filled container must be placed out of reach of the animals in order to monitor weight loss due to evaporation.

\textsuperscript{48} EPPO guideline PP1/113 for the efficacy of rodenticides, Laboratory tests for evaluation of the toxicity and acceptability of rodenticides and rodenticides preparations. Revised 1998.
Appendix 13. Field trial for rodenticide baits

This appendix describes a protocol and factors to be taken into account when conducting a field trial to determine the efficacy of an as yet unauthorised rodenticide bait product against the house mouse, brown rat or roof rat. This protocol can be applied to other target organisms (e.g. voles).

Ideally field trials should:

- be conducted with separate rat and mice populations (as appropriate to the intended uses in the draft SPC);
- be carried out at sites that are representative of the intended uses in the draft SPC (for example industrial, commercial, domestic);
- include sites with ‘known’ anticoagulant resistant populations (if appropriate to the intended uses in the draft SPC);
- have had no rodenticide treatments over the past 6 weeks;
- Incorporate lag phases before and after the treatment phase;
- for testing concentrates, cover a range of bait bases;
- for product that is sold with a specific bait station, include the whole device (the bait and its station) in the test;
- be carried out at 2 or 3 locations (i.e. a trial site sufficiently far away from the next, dependent on the roaming pattern of the test organism; e.g. Sites >30 m apart for Norway rats (Buckle and Smith 2015).

The following suggested method for bait formulations details the extent of the data required, but the methods may be replaced or supplemented by new techniques as appropriate.

Suggested procedure for bait formulations

**Trial sites**

Each trial site should, as far as possible, comprise a discrete infestation of one target species, with little chance of rapid reinvasion from adjoining areas.

During the entire trial, the baiting sites should be at exactly the same locations, taking into account distances as specified in the intended use, local structure and rodent activity as established prior to the trial. See also the Good Practice Document released by Cefic (http://www.cefic.org/Documents/Industry%20sectors/EBPF/Guideline-on-Best-Practice-in-the-Use-of-Rodenticides-in-the-EU.pdf), and the field trial protocol released by the RRAC (www.rrac.info/releases/technical-monographs/).

At each baiting site, a bait container is placed, the top of which is closed/covered, to protect the bait from weather and avoid spillage. When selecting baiting sites, it is important that the animals can feed without being disturbed.

The amount of bait applied in each feeding point should correspond to the amount given in the use instructions in the draft SPC. In general, for mice, the amount of bait applied in each feeding point is less than for brown or roof rats. In other respects, the test design is identical for both groups. It is important that there is always enough fresh food or bait containing the active substance present.

Before the trial begins, draw a sketch map showing all significant features of the site including signs of infestation.

Data on field efficacy is likely to be more reliable if infestations of brown rats and house mice are selected on the basis that a stable level of activity is obtained during the pre-treatment assessment. The level of activity can be determined by two of the following (as appropriate to the situation, species etc.):
• census baiting;
• tracking techniques;
• census by live trapping;
• electronic methods of census.

**Pre-treatment activity measurement/estimation of numbers**

Indices of the target species population should be obtained both before and after the test treatment normally by at least 2 of the following quantitative methods. Other methods, such as electronic remote detection systems, can be used as additional information for example, in combination with bait census.

**Pre-treatment bait census**

The position of the census bait points should be indicated on the site sketch plan. Census bait should be laid for at least 4 days to cover the whole infestation in quantities at each bait point which as far as possible exceed the maximum daily take by rodents. The number of census baits should be approximately the same as the planned number of test bait points. Census points should not be located at the same place chosen to lay poison points but should be at different (intermediate) positions. Census bait should be different to the bait base used in the test product.

The number of points where take has occurred and the amount of the take of the census bait, should be recorded daily. An indication of the change in weight of the bait due to moisture loss or uptake should be included.

At the end of the bait census all baits and containers should be removed from the trial site. The total amount of census bait consumed will give an index of population size.

**Tracking activity measurement**

This is recommended for both rats and mice, and should be measured over at least 3 days, simultaneously with the bait census, using tracking patches/boards laid around the site in numbers similar to the census bait points but as far as possible, not in the same locations. The locations of the patches/boards should be indicated on the plan.

The patches/boards should be inspected for signs of activity and resurfaced daily. A simple scoring system can be devised to assess the number of rodent footprints per patch/board: summing the individual scores gives a daily activity index. When the pre-treatment assessment is complete, the tracking patches/boards may be removed from the site or maintained to provide supplementary information on rodent activity.

**Census by trapping**

This is recommended for mice only, and should be carried out for a period of at least 3 days using rodenticide-free bait in the live traps. Live traps should be laid around the site in numbers appropriate to the situation and likely population size.

Animals caught should be marked by fur clipping and subsequently released. The numbers caught should be recorded and used to estimate the size of the population.

The live traps should then be removed from the test site during the rodenticide treatment.

**Lag period**

Once the pre-treatment population measurement has been conducted there should be a lag period, normally 3-14 days (or longer for acute poisons where no pre-baiting is recommended) with no experimental interference (other than tracking) on the site.
Test treatment

The test formulation must be applied in accordance with the draft SPC for an appropriate period (normally 49 days for acute products and 30-40 days for multi-dose products). The locations of test bait points should, as far as possible, be different from those of the census bait points, traps, and tracking patches/boards.

Where applicable the following items should be recorded:

- the locations of the bait points on the plan;
- the amount of bait deposited at each point at each visit and the amount retrieved, including details of the type of container used;
- the number and species of rodents and other animals found dead, and the dates on which they were found;
- the dates of all observations, treatments and censuses;
- any other information deemed relevant. This may include, for example weather conditions, temperature data, site changes instituted by the occupier (including improvements in hygiene and proofing), or supplementary information on rodent tracking activity.

On termination of the treatment all poisoned baits and bait containers should be removed from the trial sites. Similarly rodent bodies should be searched for, removed and disposed of in the appropriate way for example, burial or burning.

Post-treatment lag period

On completion of the treatment there should be a lag period sufficient to allow poisoned animals to die or survivors to recover from the sub-lethal effects of the rodenticide. This period may be 3-14 days, depending on previous observations of time to death or full recovery. During this period there should be no experimental interference with the site other than tracking.

Post-treatment activity measurement/estimation of numbers

Once the post-treatment lag period is completed, the methods employed to measure pre-treatment activity should be conducted in exactly the same way. Traps, baits and tracking patches should be laid in exactly the same places as in the pre-treatment census.

After each field trial, a comparison of population indices before and after treatment determines how successful the product has been in controlling the target population. The degree of control is expressed as a percentage reduction in the pre-treatment index.

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49 Deviation from this norm is possible but should be explained in the application.
Appendix 14. List of currently available standard test methods for rodenticides

This list may not be exhaustive, and makes no comment on the suitability of particular test methods for efficacy testing.

Table 35: List of standards

<table>
<thead>
<tr>
<th>Standard</th>
<th>Title</th>
<th>Target Organism(s)</th>
<th>Mode of Application</th>
</tr>
</thead>
<tbody>
<tr>
<td>EPA/OPP Protocol Number 1.201</td>
<td>Standard Norway Rat and Roof Rat Anticoagulant Liquid Bait Laboratory Test Method</td>
<td>Brown Rat/Roof Rat</td>
<td>Liquid bait</td>
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<tr>
<td>EPA/OPP Protocol Number 1.202</td>
<td>Standard House Mouse Anticoagulant Liquid Bait Laboratory Test Method</td>
<td>House Mouse</td>
<td>Liquid bait</td>
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<td>Standard Norway Rat and Roof Rat Anticoagulant Dry Bait Laboratory Test Method</td>
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<td>Dry Bait</td>
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<td>EPA/OPP Protocol Number 1.205</td>
<td>Standard Norway Rat/Roof Rat Anticoagulant Tracking Powder Efficacy Laboratory Test Method</td>
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<td>Tracking Powder</td>
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<tr>
<td>EPA/OPP Protocol Number 1.212</td>
<td>Standard House Mouse Anticoagulant Tracking Powder Efficacy Laboratory Test Method</td>
<td>Brown Mouse</td>
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<td>EPA/OPP Protocol Number 1.213</td>
<td>Standard Norway Rat/Roof Rat Anticoagulant Wax Block and Wax Pellet Laboratory Test Method</td>
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<td>Wax Block and Wax Pellet</td>
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<td>EPA/OPP Protocol Number 1.214</td>
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<td>Wax Block and Wax Pellet</td>
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<td>Standard Norway Rat and Roof Rat Anticoagulant Placepack Laboratory Test Method</td>
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<td>Placepack dry bait</td>
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<td>Standard House Mouse Anticoagulant Placepack Penetration Laboratory Test Method</td>
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<td>Placepack penetration</td>
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<td>Proposed Norway Rat Anticoagulant Technical and Concentrated Dry Bait Laboratory Test Method</td>
<td>Brown Rat</td>
<td>Technical and Concentrated Dry Bait</td>
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<td>Technical and Concentrated Dry Bait</td>
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<td>EPA/OPP Protocol Number: 1.207</td>
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<td>Brown Rat/Roof Rat</td>
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<td>Brown Rat/Roof Rat</td>
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<td>Standard</td>
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<td>Placepack penetration</td>
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<td>EPA/OPP Protocol Number: 1.220</td>
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<td>Placepack dry bait</td>
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<td>EPA/OPP Protocol Number: 1.222</td>
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<td>Norway rat</td>
<td>Technical and Concentrated Dry Bait</td>
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<td>EPA/OPP Protocol Number: 1.226</td>
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<td>Tracking Powder</td>
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<td>BBA 9 - 3.1</td>
<td>Richtlinie für die Prufung Prüfung von Nagetierbekämpfungsmitteln gegen Hausmause</td>
<td>House Mouse</td>
<td>Dry and liquid bait, wax block and pellets, contact rodenticides</td>
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<td>BBA 9- 3.2</td>
<td>Richtlinie für die Prüfung von Nagetierbekämpfungsmitteln gegen Wanderratten</td>
<td>Brown Rat</td>
<td>Dry and liquid bait, wax block and pellets, contact rodenticides</td>
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<tr>
<td>EPPO 1982</td>
<td>Guidelines for the Biological Evaluation of Rodenticides No1. Laboratory Tests for Evaluation of the Toxicity and Acceptability of Rodenticides and Rodenticide Preparations</td>
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<td>EPPO 1982</td>
<td>Guidelines For the Biological Evaluation of Rodenticides. Field Tests Against Synanthropic Rodents (Mus musculus, Rattus norvegicus, Rattus rattus)</td>
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<td>EPPO 1986</td>
<td>Guidelines for the Biological Evaluation of Rodenticides. Laboratory and Field Tests for the Evaluation of Rodenticidal Dusts</td>
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<td>EPPO Standards/97(2)</td>
<td>Laboratory and field tests for the evaluation of rodenticidal dusts</td>
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<td>EPPO Standards /113(2)</td>
<td>Laboratory tests for evaluation of the toxicity and acceptability of rodenticides and rodenticide preparations</td>
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<td>Field tests against synanthropic rodents</td>
<td>Brown rat/Roof rat/ House mouse</td>
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<td>Standard</td>
<td>Title</td>
<td>Target Organism(s)</td>
<td>Mode of Application</td>
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<tr>
<td>EPPO Standards /169(2)</td>
<td>Efficacy trials with rodenticide baits under practical conditions against Voles (<em>Arvicola terrestris</em> and <em>Microtus</em> spp.) in their subterranean galleries&quot;</td>
<td>Voles <em>(Microtus, Arvicola)</em></td>
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<td>EPPO Standards /197(1)</td>
<td>Non-target effects of rodenticides</td>
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<tr>
<td>EPPO Standards /198(1)</td>
<td>Testing rodents for resistance to anticoagulant rodenticides</td>
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<tr>
<td>RRAC rat field trial protocol 2013</td>
<td>Field Trial to Evaluate the Efficacy of Rodenticide Baits for the Control of Rats (<em>Rattus norvegicus</em>)</td>
<td>Brown Rat/Roof Rat</td>
<td>Dry Bait</td>
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<td>OECD</td>
<td>OECD Guidance Document on the Recognition, Assessment and Use of Clinical Signs as Human Endpoints for Experimental Animals Used in Safety Evaluation (2002)</td>
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<td>BBA (1963)</td>
<td>Richtlinie 9-2, Richtlinien für die Prüfung von Nagetierbekämpfungsmitteln gegen Schermaus (in German)</td>
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<td>BBA (1980)</td>
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<td>Méthode CEB n°254 (2013)</td>
<td>Méthode d’essai d’efficacité pratique de générateurs de gaz fumigants pour lutter contre la taupe (<em>Talpa europaea</em>) et le campagnol terrestre (<em>Arvicola terrestris</em>) dans leurs galeries souterraines au champ</td>
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<td>Gassing agent</td>
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<td>Méthode CEB n°257 (2014)</td>
<td>Méthode d’essai d’efficacité pratique d’appâts rodenticides pour lutter contre les campagnols (<em>Arvicola terrestris, Microtus spp.</em>) dans leurs galeries souterraines au champ</td>
<td>Voles, moles</td>
<td>Bait</td>
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</table>
Appendix 15. Additional information on label claims

Assessing the efficacy of biocidal products

The evaluation of the efficacy of biocidal products differs greatly from that of active substances.

Whilst the efficacy assessment of an active substance for Annex I inclusion requires only a minimal assessment, sufficient to show an innate level of activity for the active substance, the assessment needed for a biocidal product at the product authorisation stage is much more detailed.

Rather than looking at innate effects, the efficacy assessment of a biocidal product is based on substantiating the efficacy claims made for a product. The assessment is made on the product in its normal conditions of use.

This principle is set out in paragraph 51 of Annex VI of the Directive (Common Principles for the Evaluation of Dossiers for Biocidal Products), which states:

5.1 Data shall be submitted and evaluated to ascertain if the efficacy claims of the biocidal product can be substantiated. Data submitted by the Applicant or held by the Member State must be able to demonstrate the efficacy of the biocidal product against the target organism when used normally in accordance with the conditions of authorisation.

The label claims for the product must be submitted as part of the common core data set, as set out in Annex IIB (Common Core Data Set for Biocidal Products), which requires:

V. INTENDED USES AND EFFICACY

5.10. The proposed label claims for the product and efficacy data to support these claims, including any available standard protocols used, laboratory tests, or field trials, where appropriate

As the label claims are central to the efficacy evaluation for a biocidal product, it is important to understand exactly what is an efficacy claim, and be able to identify the individual components of a claim.

Label claims for biocidal products

As efficacy claims are assessed against the product ‘when used normally in accordance with the conditions of authorisation’, then it is important to define the ‘normal use’ of the product.

There are several pieces of information which will form part of the conditions of authorisation which relate to the efficacy assessment. These are:

1. The Formulation Type

This is determined by the product itself – e.g. a solvent based ready-for-use, a water based concentrate, a dusting powder, a gel bait, etc.

2. Application Method

This is the method by which the product is intended to be applied. e.g. coarse spray, ultra low volume (ULV) spray, bait station, skin lotion, etc.

The application method may also describe a specific pattern of treatment. This is particularly common for spray applications, but may also apply to other formulation types. General descriptions of some common treatment patterns are given below.
(i) Surface treatments
These are treatments where the product is applied over surfaces such as walls, floors and ceilings, or as a treatment to outdoor surfaces. These treatments may involve treating a large area of surface or may only involve application to a narrow band. Surface treatments can also include application to temporary or permanent bodies of water (e.g. in mosquito control) and to solid and semi-solid manure.

(ii) Crack and crevice treatments
These are treatments where products are applied into cracks and crevices where insects hide and harbourage, or through which they may enter the building. Such openings commonly occur at expansion joints, between different elements of construction and between equipment and floors. These openings may lead to voids such as hollow walls, equipment legs and bases, conduits and junction or switch boxes.

(iii) Contact (direct) spray treatments
These involve application directly onto insects, and are normally only possible when the insects are visible and available to be sprayed. In practice this often restricts direct application methods to controlling flying insects (such as adult moths and houseflies), although some limited control of minor infestations of crawling insects (such as ants or beetles) may be possible.

(iv) Space treatments
These are treatments where the product is applied into the air rather than onto a surface. They are intended to disperse small droplets or particles into the atmosphere of a room or other open space, where they will normally stay for a period of time (very small particles may stay in the air for several hours under still conditions).

(v) Spot treatments
These are treatments where products are applied to limited areas on which insect pests are likely to occur, but which will not be in contact with food or utensils and will not ordinarily be contacted by workers. These areas may occur on floors, walls and bases or undersides of equipment.

(vi) Baits
Bait treatments use products that are intended to be ingested by the target. This is normally through the insect feeding on the product directly, but may also include products which the target will come into contact with and later ingest during grooming/cleaning. The attractiveness of these products is through the use of a palatable food base, however they may also incorporate an attractant (e.g. a pheromone) which is intended to attract the target pests over a greater distance.

3. Application Rate
This is the rate at which the product will be applied in use, e.g. apply 100 ml of product per square metre, apply at a rate of 1 bait station per 3 m², spray for 20 seconds, etc.

For efficacy assessment purposes, it is useful to consider the application rate as the amount of active substance applied to surface area or volume. Unlike a human health or environmental risk assessment which look at the maximum amounts of product which are considered to be acceptable (i.e. if the amount of active or application rate increase, the risks to man or the environment will be unacceptable), an efficacy evaluation looks at the minimum application/dose rate which will be effective (i.e. if the application rate decreases, the product may not work).

4. Frequency of treatment and any specific interval between applications
Some products will be used in a way that will require more than one treatment. These products will give information on the treatment schedule which should be followed (e.g. insecticide re-treatment intervals or rodenticide re-baiting periods).
Together, these pieces of information define the ‘normal use’ of the product (e.g. a solvent based ready-for-use product to be applied as a coarse spray at a rate of 100 ml product m⁻²), and efficacy must be demonstrated for the product when it is used in this way.

Whilst information on the application method and rate etc. will normally be clearly defined, the claims made for the effects of the product are much more difficult to identify.

5. Other specific conditions to be taken into account

Occasionally, the “normal use” of a product will involve the use of the product in conjunction with other activities. This will include the cleaning of an area prior to treatment. The contributions made by other components of an Integrated Pest Management procedure may also have to be taken into account.

Product labels and label claims

The product label is the major source of information on a product. It will give the use pattern to help determine the ‘normal use’ of the product, but will also make claims about the effectiveness of the product.

These label claims form the core of any efficacy evaluation. Efficacy is assessed mainly in relation to the claims made for the product. The norms and criteria set per insect pest will further guide the evaluation.

Whilst the phrase ‘label claims’ is generally used, this phrase actually encompasses all claims made for the product, not just those made on the label itself. Claims may also be made for a product with any accompanying information (such as leaflets) or on advertising material.

For efficacy purposes, all of these claims also have to be justified before they can be allowed onto a label.

What is a label claim?

A label claim is anything on the product label that makes a claim about what the product does or the benefits that will result from its use. At this moment there is no standard format for making claims about the effects and benefits of using the product, and the type and style of label claims can vary widely between different Member States.

For example, a product which claims to be ‘For the control of cockroaches’ in one Member State may claim that it ‘Kills cockroaches fast!!’ in another.

To aid in the evaluation process, a standardised method for identifying the main components of a label claim is set out below.

Label claims – understanding the components

A set of label claims will consist of 2 types of information which describe what the product will do when it is used (in accordance with its ‘normal use’). These are:

1. The target species which the product will be effective against and

2. The effect (or effects) which the use of the product will have on the target species and the benefits which may result from this effect

Target species

The product label will give details about which species the product is to be used against. This information will often be quite specific (e.g. ‘for the control of pharaohs ants’ or ‘kills ants, cockroaches, fleas and bed bugs or repels mosquitoes’). In these cases it is easy to identify what are the target species.
However there can also be instances where a more general claim is made, such as for use against ‘crawling insects’. In these cases, it is difficult to require data on every crawling insect.

They will need to supply efficacy data on relevant representative species, which may be those used in standard test methods or those that the Applicant argues are representative of the use pattern of the biocide and the nature of the application (e.g. whether it is a space application or a surface application).

In some instances it is possible to allow a compromise on the label. For example, members of the general public may not know what species of fly is in their home, but the regulators will need to know what the product is effective against. In this particular instance it may be possible to allow a claim such as ‘Effective against flying insects such as the housefly, mosquitoes and midges’.

**The effects of using the product**

The remaining parts of the label claim will describe the effects on the target organisms and benefits of using the product.

The major effects which are generally claimed are that a product will:

- kill, knock down, repel, attract, reduce the numbers of or inhibit a target organism
- control, reduce or prevent the build-up of a population
- prevent or reduce an undesirable effect.

For insecticide products, the following claims are the ones that are frequently encountered:

‘*Kill*’ claims generally refer to the death of an individual or a number of individuals (the death of an entire population is more generally found under a ‘control’ claim) and generally refer to an existing infestation.

‘*Knockdown*’ claims are generally restricted to insecticides and acaricides. A knockdown effect is one where a target insect becomes unable to carry out coordinated movement, but has not been killed.

Knockdown effects are often included in an insecticide product to produce a rapid, visible effect on a target in order to satisfy user expectations. These effects can be reversible, with insects able to recover after a period of time. Recovery is often dependent upon dose administered.

Knockdown claims may be found in conjunction with a kill claim, and many ‘dual action’ insecticide products contain two active substances - with one active substance producing a quick knockdown effect (such as a flying insect falling out of the air) whilst a second, slower acting, active substance produces the killing effect. Combined claims may be along the lines of ‘knockdown within 10 minutes and kill within 2 hours’.

When it comes to efficacy testing, some companies use the two terms interchangeably, so you will get products or test reports mentioning ‘knockdown’ where a killing effect is actually meant. For evaluation purposes, knockdown and kill are considered to be separate effects.

‘*Complete control’, ‘colony kill*’ or ‘nest kill’ claims will generally refer to the elimination of an entire infestation or population - i.e. use of the product will essentially ‘remove the problem’.

As stated above, the mortality of individuals (rather than populations) is considered to be a ‘kill’ effect.
To highlight the difference between ‘kill’ and ‘control’, we can take the example of an ant nest outside of a house, close to the back door. The queen (which does all of the reproduction) remains hidden away in the nest and produces new ants for the colony, and the only ants seen outside of the nest are the sterile female workers.

An aerosol product which is intended to be sprayed onto ants wandering around in your kitchen to kill them will only be having a ‘kill’ effect. Killing off individuals or numbers of workers will have little effect on the nest and the colony as a whole, as the queen and fertile males will remain unaffected in the nest.

In order to remove the problem, you actually have to kill off the colony. So a product claiming to ‘control’ an infestation of ants would have to eliminate the queen or disrupt the ability of the colony to reproduce.

‘Reduce’ claims will generally refer to reducing the numbers of (but not completely eliminating) a target population. Whilst not eliminating an infestation may seem to be an odd claim to make, there are situations where it would be practically impossible to totally control a target population and where the best result is to reduce the scale of the problem.

An example of this would be reducing the fly burden in a poultry house or intensive animal house. However, the issue of resistance must always be kept in mind when considering treatments which do not fully control a population.

**More complex label claims**

Whilst a label claim is, at its most basic, a target and an effect, most claims are more complex, introducing further elements beyond the basic target/effect combination described above.

These additional parts of a label claim more fully describe the effects on the target organisms and benefits to be gained from using the product.

Claims for the effects and benefits of using the product can generally be broken down into 6 major components, which are described in Table 26.

The examples given in the table cannot be exhaustive, but are given to illustrate the type of information which appears in label claims.

**Table 36: Components Making Up a Label Claim**

<table>
<thead>
<tr>
<th>Group</th>
<th>Label Claim</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Against what target organism(s) will the product be used?</td>
</tr>
<tr>
<td></td>
<td>• Specific insect (e.g. ants)</td>
</tr>
<tr>
<td></td>
<td>• Several insects (e.g. ants and wasps)</td>
</tr>
<tr>
<td></td>
<td>• General claim (e.g. flying and crawling insects)</td>
</tr>
<tr>
<td>B</td>
<td>What effect will the use of the product have on the target?</td>
</tr>
<tr>
<td></td>
<td>Examples include:</td>
</tr>
<tr>
<td></td>
<td>• Kill</td>
</tr>
<tr>
<td></td>
<td>• Knockdown</td>
</tr>
<tr>
<td></td>
<td>• Control</td>
</tr>
<tr>
<td></td>
<td>• Flushing</td>
</tr>
<tr>
<td></td>
<td>• Attracting</td>
</tr>
<tr>
<td></td>
<td>• Repelling</td>
</tr>
<tr>
<td>C</td>
<td>How long will the product take to produce the effect?</td>
</tr>
<tr>
<td>Group</td>
<td>Label Claim</td>
</tr>
<tr>
<td>-------</td>
<td>-------------</td>
</tr>
<tr>
<td>D</td>
<td>the effect</td>
</tr>
<tr>
<td></td>
<td>Area of use</td>
</tr>
<tr>
<td></td>
<td>Duration of the effect</td>
</tr>
<tr>
<td></td>
<td>User</td>
</tr>
<tr>
<td></td>
<td>Other specific claims</td>
</tr>
</tbody>
</table>

A label claim will not always contain all 7 components. For example, where no residual activity is being claimed, section E will not be represented, and where no specific other claims are being made, claims in section G will not be present.

The target organism (A), the type of effect (B) and area of use (D) and the user (F) should always be given.

On some labels, the time taken to produce the effect (C) will not have been given (e.g. ‘for the control of cockroaches’) or is not a specific value (e.g. ‘kills flies fast’). In these cases, the evaluator will use the norms and criteria given per insect for the evaluation of the data.
Linking the components of the label claim

When initially trying to understand how the components of the label claims fit together, it can help to place the assorted claims into a table in order to identify how the various elements interact. For example:

**Table 37: Example of linking label claims**

<table>
<thead>
<tr>
<th>Label claim (B)</th>
<th>Effect time (C)</th>
<th>Area of use (D)</th>
<th>Duration of effect (E)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Knocks down</td>
<td>within 5 minutes</td>
<td>- on hard porous and non-porous surfaces</td>
<td>for 6 weeks</td>
</tr>
<tr>
<td>Kills</td>
<td>within 1 hour</td>
<td>- on soft furnishings</td>
<td></td>
</tr>
</tbody>
</table>

The beneficial effect of the product (B) will be accompanied by the timescale in which the effect will happen (C). In these cases, it must be demonstrated that the product will be efficacious within the stated time.

In the above example, it must be demonstrated that the product is capable of both knocking down the target insects *within 5 minutes* AND killing them *within 1 hour*.

The area of use (D) gives information about the conditions in which the product will be used and the type of surfaces it will be used on. The efficacy data supplied should demonstrate that the product will be efficacious in the areas specified or on representative surfaces of the types described.

In the example, it would have to be demonstrated that the product would produce its knockdown and kill effects *within the times stated* AND on *both hard surfaces and soft furnishings*.

The duration of effect (E) specifies the length of residual activity which must be demonstrated.

In the example, it must be demonstrated that the product is still capable of producing the effects on the specified surfaces 6 weeks after treatment (although not necessarily to the same degree as a fresh treatment).

Other claims can be linked into this process in the same way. For example, if claims were being made that the product was to be used against resistant individuals, then all of the above elements would have to be proved using a resistant test population to generate the data.

Once the various elements making up the label claims have been identified then the evaluation of the efficacy data submitted can proceed.

General guidance on the assessment of label claims is included in the paper “Broad principles of assessing efficacy in relation to claims made on the label for biocidal products”, which was agreed at the Technical Meeting TM III 05 in October 2005, and at the subsequent CA meeting.

Guidance on type of and amount of data which would normally be required to support many of the major label claims is given for the main pest species elsewhere in this guidance.
### Appendix 16. Species grid

#### Table 38: PT 18 Crawling Insects

<table>
<thead>
<tr>
<th>Action</th>
<th>SITE</th>
<th>APPLICATION METHOD</th>
<th>CLAIM</th>
<th>TEST SPECIES</th>
<th>RATIONALE</th>
<th>NOTES</th>
<th>INSECT STAGE</th>
</tr>
</thead>
<tbody>
<tr>
<td>1A</td>
<td>Flushing</td>
<td>Indoor</td>
<td>Crack &amp; Crevice</td>
<td>&quot;Flushes cockroaches out of hidden places&quot;</td>
<td><em>Blattella germanica</em> or <em>Periplaneta</em></td>
<td>Data show Periplaneta flush before <em>Blattella</em> N.B This is true with pyrethroids, the case may be different with other actives. Fast acting pyrethroids may knockdown <em>Blattella</em> faster than they can be flushed, use <em>Periplaneta</em> in this case.</td>
<td>Any additional species need specific data.</td>
</tr>
<tr>
<td>1B</td>
<td>Knockdown</td>
<td>Indoor</td>
<td>Direct Spray</td>
<td>&quot;Knocks down cockroaches&quot;; &quot;Knocks down cockroaches in x seconds&quot;</td>
<td><em>Blattella germanica</em> and either <em>Periplaneta</em> species or <em>Blatta orientalis</em></td>
<td>These species are representative of all domestic cockroaches found in Europe and around the world. Behavioural differences between species do not come into play when testing aerosols for direct spray efficacy.</td>
<td>We see little or no value in producing nymph/immature data in aerosol direct spray tests. Testing with only adults provides a very clear picture of product activity for registration studies. More than one life stage is an unnecessary burden.</td>
</tr>
<tr>
<td>1C</td>
<td>Kills</td>
<td>Indoor</td>
<td>Direct Spray</td>
<td>&quot;Kills cockroaches&quot;; &quot;Kills cockroaches in x seconds&quot;</td>
<td><em>Blattella germanica</em> and either <em>Periplaneta</em> species or <em>Blatta orientalis</em></td>
<td>See B.</td>
<td>See B</td>
</tr>
<tr>
<td>1D</td>
<td>Kills</td>
<td>Indoor</td>
<td>Direct Spray</td>
<td>&quot;Kills ants&quot;; &quot;Kills in x seconds&quot;</td>
<td><em>Lasius</em> sp.</td>
<td><em>Monomorium</em> ants are much smaller and more sensitive so would be covered by data for <em>Lasius</em></td>
<td>Adults</td>
</tr>
<tr>
<td>1E</td>
<td>Kills</td>
<td>Outdoor</td>
<td>Direct Spray</td>
<td>&quot;Kills ants&quot;</td>
<td><em>Lasius</em> sp.</td>
<td></td>
<td>Adults</td>
</tr>
<tr>
<td>Action</td>
<td>SITE</td>
<td>APPLICATION METHOD</td>
<td>CLAIM</td>
<td>TEST SPECIES</td>
<td>RATIONALE</td>
<td>NOTES</td>
<td>INSECT STAGE</td>
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<td>-------------</td>
</tr>
<tr>
<td>1F</td>
<td>Knockdown</td>
<td>Indoor</td>
<td>Direct Spray</td>
<td>&quot;Kills in x seconds&quot;</td>
<td>C + D and a variety of other common species e.g. <em>Forficula auricularia</em>, <em>Acheta domesticus</em>, <em>Cimex lectularius</em>, <em>Attagenus</em>, <em>Dermestes</em> sp, fleas, silverfish, booklice, carpet beetles, woodlice, ticks, centipedes, spiders</td>
<td>Multiple species are common world-wide. Test species will depend upon seasonal and local availability. See also B.</td>
<td>Adults</td>
</tr>
<tr>
<td>1G</td>
<td>Knockdown</td>
<td>Indoor</td>
<td>Space spray; aerosols, gases, fogs, smokes</td>
<td>Knocks down crawling insects</td>
<td>Wood borers, carpet beetles, stored product beetles, other small crawling insects. Data required for claims on cockroaches (C) and fleas as surrogates for others</td>
<td></td>
<td>Adults, immatures</td>
</tr>
<tr>
<td>1H</td>
<td>Kills</td>
<td>Indoor</td>
<td>Space spray; aerosols, gases, fogs, smokes</td>
<td>Kills crawling insects</td>
<td>Wood borers, carpet beetles, stored product beetles, other small crawling insects. Data required for claims on cockroaches (C) and fleas</td>
<td></td>
<td>Adults, immatures and if claimed eggs</td>
</tr>
<tr>
<td>I</td>
<td>Residual Kill</td>
<td>Indoor</td>
<td>Surface or Crack &amp; Crevice Spray, Powders</td>
<td>&quot;Kills cockroaches&quot;; &quot;Kills cockroaches up to x weeks or months&quot;</td>
<td><em>Blattella germanica</em> and either <em>Periplaneta</em> species or <em>Blatta orientalis</em></td>
<td>Consider substrate and ageing period in the method</td>
<td>Adults and or immature stages. Specify realistic exposure period followed by</td>
</tr>
<tr>
<td>Action</td>
<td>SITE</td>
<td>APPLICATION METHOD</td>
<td>CLAIM</td>
<td>TEST SPECIES</td>
<td>RATIONALE</td>
<td>NOTES</td>
<td>INSECT STAGE</td>
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</tr>
<tr>
<td>1J</td>
<td>Residual</td>
<td>Indoor</td>
<td>Surface or Crack &amp; Crevice Spray, Powder</td>
<td>Kills ants”; &quot;Kills ants for x weeks or months”</td>
<td>Lasius sp. and/or Monomorium pharaonis as option (see 4)</td>
<td>Consider substrate and ageing period in the method</td>
<td>Adults Specify realistic exposure period followed by reasonable “recovery” period.</td>
</tr>
<tr>
<td>1K</td>
<td>Residual</td>
<td>Indoor</td>
<td>Surface or Crack &amp; Crevice Spray, Powder</td>
<td>&quot;Kills crawling insects and arthropods”; &quot;Kills for x weeks or months”</td>
<td>K + L and a variety of other common species e.g. Forficula auricularia, Acheta domesticus, Cimex lectularius, Attagenus, Dermestes sp, fleas, silverfish, booklice, carpet beetles, woodlice, ticks, centipedes, spiders</td>
<td>We propose only roaches be tested for full period.</td>
<td>Adults and immature stages. Consider substrate and ageing period in method. Specify realistic exposure period followed by “reasonable” recovery period.</td>
</tr>
<tr>
<td>1L</td>
<td>Residual</td>
<td>Indoor</td>
<td>Bait</td>
<td>&quot;Kills cockroaches”; &quot;Kills cockroaches for x weeks or months”;</td>
<td>Blattella germanica; Periplaneta americana and Blatta orientalis</td>
<td>Either the claim is limited to a specific species or the three species are tested</td>
<td>Nymphs Adults. Consider ageing period in method. Provide harbourage and alternative food and water.</td>
</tr>
<tr>
<td>Action</td>
<td>SITE</td>
<td>APPLICATION METHOD</td>
<td>CLAIM</td>
<td>TEST SPECIES</td>
<td>RATIONALE</td>
<td>NOTES</td>
<td>INSECT STAGE</td>
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<td>--------------</td>
</tr>
<tr>
<td>1M</td>
<td>Secondary kill</td>
<td>Indoor</td>
<td>Bait</td>
<td>&quot;Kills cockroaches that do not visit the bait (secondary kill)&quot;</td>
<td>Blattella germanica; Periplaneta americana and Blatta orientalis</td>
<td>Life stage to be tested depends upon a specific mode of action (necrophagy versus coprophagy). Either nymphs or adults could be used.</td>
<td>Either the claim is limited to a specific species or the three species are tested</td>
</tr>
<tr>
<td>1N</td>
<td>Nest kill</td>
<td>Indoor</td>
<td>Bait</td>
<td>control of entire population of cockroaches</td>
<td>Blattella germanica; Periplaneta americana and Blatta orientalis</td>
<td>Either the claim is limited to a specific species or the three species are tested</td>
<td>Either the claim is limited to a specific species or the three species are tested</td>
</tr>
<tr>
<td>1O</td>
<td>Kill</td>
<td>Indoor</td>
<td>Bait</td>
<td>&quot;Kills ants&quot;; &quot;Kills ants for x weeks or months&quot;;</td>
<td>Monomorium pharaonis and /or Lasius niger.</td>
<td>Either the claim is limited to a specific species or the two species are tested. Provide harbourage and alternative food and water.</td>
<td>Either the claim is limited to a specific species or the two species are tested. Provide harbourage and alternative food and water.</td>
</tr>
<tr>
<td>1P</td>
<td>Colony kill</td>
<td>Indoor</td>
<td>Bait</td>
<td>&quot;Kills the queen and the colony&quot;</td>
<td>Monomorium pharaonis and /or Lasius niger.</td>
<td>Either the claim is limited to a specific species or the two species are tested. Provide harbourage and alternative food and water.</td>
<td>Either the claim is limited to a specific species or the two species are tested. Provide harbourage and alternative food and water.</td>
</tr>
<tr>
<td>1Q</td>
<td>Kills</td>
<td>Indoor</td>
<td>Spray, powder</td>
<td>&quot;Kills dust mites&quot;</td>
<td>Dermatophagoides sp.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1R</td>
<td>Residual Kill</td>
<td>Indoor</td>
<td>Spray, powder</td>
<td>&quot;Kills dust mites for x weeks/months&quot;</td>
<td>Dermatophagoides sp.</td>
<td>Consider substrate and ageing period in method. Specify</td>
<td></td>
</tr>
<tr>
<td>Action</td>
<td>SITE</td>
<td>APPLICATION METHOD</td>
<td>CLAIM</td>
<td>TEST SPECIES</td>
<td>RATIONALE</td>
<td>NOTES</td>
<td>INSECT STAGE</td>
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<td>-------</td>
<td>--------------</td>
</tr>
<tr>
<td>1S</td>
<td>Kill</td>
<td>Outdo or</td>
<td>Baits, Dusts, powders</td>
<td>Kills ants</td>
<td>Lasius sp</td>
<td></td>
<td>realistic insect exposure period followed by reasonable “recovery” period.</td>
</tr>
<tr>
<td>1T</td>
<td>Kill</td>
<td>Outdo or</td>
<td>Baits, Dusts, powders</td>
<td>&quot;Kills the queen and the colony&quot;</td>
<td>Lasius sp and /or Monomorium pharaonis</td>
<td>Either the claim is limited to a specific species or the two species are tested. Provide harbourage and alternative food and water.</td>
<td></td>
</tr>
<tr>
<td>1U</td>
<td>Kill</td>
<td>Outdo or</td>
<td>Sprays, liquid drenches</td>
<td>Kills ants</td>
<td>Lasius sp</td>
<td></td>
<td>Add colony kill</td>
</tr>
<tr>
<td>1V</td>
<td>Kill</td>
<td>Outdo or</td>
<td>Sprays, liquid drenches</td>
<td>&quot;Kills the queen and the colony&quot;</td>
<td>Monomorium pharaonis and /or Lasius niger.</td>
<td>Either the claim is limited to a specific species or the two species are tested. Provide harbourage and alternative food and water.</td>
<td></td>
</tr>
<tr>
<td>1W</td>
<td>Kill or repellent</td>
<td>Outdo or</td>
<td>Physico-chemical barrier. Installation between the soil and the future construction</td>
<td>Preventive Pre- construction treatment Prevent construction attack</td>
<td>All subterranean termites Reticulitermes sp. Coptotermes sp. Heterotermes sp.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1X</td>
<td>Kill or repellent</td>
<td>Outdo or</td>
<td>Chemical barrier Injection in wall and soil</td>
<td>Preventive Pre-construction treatment Prevent construction attack</td>
<td>All subterranean termites Reticulitermes sp. Coptotermes sp. Heterotermes sp.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1Y</td>
<td>Kill or repellent</td>
<td>Outdo or</td>
<td>Chemical barrier Injection in wall and soil</td>
<td>Curative Post-construction treatment</td>
<td>All subterranean termites Reticulitermes sp. Coptotermes sp. Heterotermes sp.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
## Table 39: PT 18 Flying Insects

<table>
<thead>
<tr>
<th>Action</th>
<th>SITE</th>
<th>APPLICATION METHOD</th>
<th>CLAIM</th>
<th>TEST SPECIES</th>
<th>RATIONALE</th>
<th>NOTES</th>
</tr>
</thead>
<tbody>
<tr>
<td>1Z</td>
<td>Kill</td>
<td>Outdoors</td>
<td>Baits system</td>
<td>Curative Post-construction treatment Colony elimination</td>
<td>Reticulitermes sp. Coptotermes sp.</td>
<td>Due to the specificity of baits, only species tested should be claimed on the product label</td>
</tr>
<tr>
<td>1AA</td>
<td>Kill</td>
<td>Indoor</td>
<td>Curative (Prevention is PT 8)</td>
<td>Kills dry wood termites</td>
<td>e.g. Cryptotermes sp.</td>
<td></td>
</tr>
<tr>
<td>1AB</td>
<td>Barrier treatment</td>
<td>Indoor / Outdoor</td>
<td>Sprays, Powders</td>
<td>Prevents entry of crawling insects for x weeks or months</td>
<td>Blattella germanica and either Periplaneta species or B. orientalis, Lasius sp.</td>
<td>See list above (&quot;F&quot;) for selection, but expect roaches and ants to be the main claim</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Action</th>
<th>SITE</th>
<th>APPLICATION METHOD</th>
<th>CLAIM</th>
<th>TEST SPECIES</th>
<th>RATIONALE</th>
<th>NOTES</th>
</tr>
</thead>
<tbody>
<tr>
<td>2A</td>
<td>Kills / Knocks down</td>
<td>Indoor</td>
<td>Direct spray or room treatment</td>
<td>“Knocks down and/or Kills flies, mosquitoes”;</td>
<td>Musca domestica; Culex sp. or Aedes sp.</td>
<td>These two species are representative of most urban species.</td>
</tr>
<tr>
<td>2B</td>
<td>kills</td>
<td>Indoor/Outdoor</td>
<td>Aerosol, Coils, mats or liquid electrics; Plaquettes or similar devices</td>
<td>Kills mosquitoes for up to x hours</td>
<td>Culex sp. or Aedes sp.</td>
<td>All insects, for which claims are made, should be tested. adults</td>
</tr>
<tr>
<td>2C</td>
<td>Outdoor</td>
<td>Nuisance flying insects (landfill area)</td>
<td>Kills “XYZ”</td>
<td>Musca domestica Culex sp. or Aedes sp.</td>
<td>All insects, for which claims are made, should be tested. adults</td>
<td></td>
</tr>
<tr>
<td>2D</td>
<td>Outdoors</td>
<td>Direct and residual sprays</td>
<td>Kills “XYZ”</td>
<td>Claimed insects need to be tested</td>
<td></td>
<td>adult and larvae</td>
</tr>
<tr>
<td>Action</td>
<td>SITE</td>
<td>APPLICATION METHOD</td>
<td>CLAIM</td>
<td>TEST SPECIES</td>
<td>RATIONALE</td>
<td>NOTES</td>
</tr>
<tr>
<td>--------</td>
<td>------------</td>
<td>--------------------</td>
<td>-------------------------</td>
<td>---------------------------</td>
<td>-----------</td>
<td>----------------------------------------------------------------------</td>
</tr>
<tr>
<td>2E</td>
<td>Indoor</td>
<td>Fumigants</td>
<td>Kills &quot;XYZ&quot;</td>
<td>Claimed insects need to be tested</td>
<td></td>
<td>All insects and insect stages for which claims are made, should be tested.</td>
</tr>
<tr>
<td>2F</td>
<td>Indoor</td>
<td>Direct spray or room treatment</td>
<td>&quot;Kills flying moths&quot;</td>
<td><em>Plodia interpunctella</em> or <em>Tineola bisselliella</em></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2G</td>
<td>Indoor / Outdoor</td>
<td>Direct spray</td>
<td>&quot;Kills wasps&quot;</td>
<td><em>Vespa sp.</em></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2H</td>
<td>Outdoor</td>
<td>Nest treatment (all methods)</td>
<td>&quot;Kills wasp nests&quot;; &quot;Kills the queen&quot;</td>
<td><em>Vespula sp.</em> or <em>Dolichovespula sp</em></td>
<td></td>
<td>Test on whole nests</td>
</tr>
<tr>
<td>2I</td>
<td>Indoor</td>
<td>Closet or confined space treatments</td>
<td>&quot;Kills clothes moths and larvae&quot;; &quot;Kills for x weeks or months&quot;</td>
<td><em>Tineola bisselliella</em></td>
<td></td>
<td>All insects, for which claims are made, should be tested.</td>
</tr>
<tr>
<td>2J</td>
<td>Indoor</td>
<td>Baits</td>
<td>Kills &quot;XYZ&quot;flies</td>
<td><em>Specific species claimed on the label</em></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2K</td>
<td>Outdoor</td>
<td>Mosquitoes</td>
<td>Kills mosquito larvae</td>
<td><em>Culex sp.</em> Or <em>Aedes sp.</em></td>
<td></td>
<td>for IGRs the larval stage needs to be selected according to the mode of action.</td>
</tr>
<tr>
<td>2L</td>
<td>Indoor / Outdoor</td>
<td>Fly larvicides</td>
<td>Kills &quot;XYZ&quot;flies</td>
<td><em>Specific species claimed on the label</em></td>
<td></td>
<td>for IGRs the larval stage needs to be selected according to the mode of action.</td>
</tr>
</tbody>
</table>

**Table 40: PT 19 – Repellents & Attractants**

<table>
<thead>
<tr>
<th>Action</th>
<th>SITE</th>
<th>APPLICATION METHOD</th>
<th>CLAIM</th>
<th>TEST SPECIES</th>
<th>RATIONALE</th>
<th>NOTES</th>
<th>INSECT STAGE</th>
</tr>
</thead>
<tbody>
<tr>
<td>3A</td>
<td>Personal repellent</td>
<td>Outdoor</td>
<td>Aerosol spray, pump spray,</td>
<td>&quot;Protects for a minimum/average&quot;</td>
<td><em>Aedes sp.</em> and <em>Culex sp.</em></td>
<td>Aedes is widely used as it is an aggressive</td>
<td>Any additional pest claimed needs to be tested (Sandflies</td>
</tr>
</tbody>
</table>

Note: The table provides guidance on the BPR for specific actions, sites, and insect stages, along with the necessary test species and rationale for claims made on product labels. The notes column highlights the importance of testing all stages for which claims are made, and the use of specific species in repellents and attractants.
<table>
<thead>
<tr>
<th>Action</th>
<th>SITE</th>
<th>APPLICATION METHOD</th>
<th>CLAIM</th>
<th>TEST SPECIES</th>
<th>RATIONALE</th>
<th>NOTES</th>
</tr>
</thead>
<tbody>
<tr>
<td>3B</td>
<td>Personal repellent</td>
<td>outdoor</td>
<td>Aerosol spray, pump spray, lotion, cream, towels etc.</td>
<td>Protects for a minimum/average of “x” hours against ticks</td>
<td>Ixodes sp. or Dermacentor</td>
<td>wasps) If biting flies are claimed they need to be tested; (Stomoxys) If malaria mosquitoes are claimed tests need to be carried out on Anopheles sp. If nuisance flies are claimed: Musca domestica</td>
</tr>
<tr>
<td>3C</td>
<td>Area repellent</td>
<td>Indoor / Outdoor</td>
<td>Coils, mats or liquid electrics or other devices</td>
<td>“Protects for up to x hours against mosquitoes”</td>
<td>Aedes sp. or Culex sp.</td>
<td></td>
</tr>
<tr>
<td>3D</td>
<td>Insecticide for Fabric</td>
<td>Indoor/ outdoor Fabrics, Apparel, Bednets</td>
<td>Entire materials / sprays or liquids to impregnate these materials</td>
<td>Protects for up to x weeks against “XYZ”</td>
<td>Mosquitoes (Aedes spec, Culex spec); ticks (Ixodes sp or Dermacentor spec.)</td>
<td>The following claims need to be verified by appropriate test data: malaria mosquitoes – Anopheles spec; biting flies – Stomoxys; nuisance</td>
</tr>
<tr>
<td>Action</td>
<td>SITE</td>
<td>APPLICATION METHOD</td>
<td>CLAIM</td>
<td>TEST SPECIES</td>
<td>RATIONALE</td>
<td>NOTES</td>
</tr>
<tr>
<td>--------</td>
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<td>-------</td>
</tr>
<tr>
<td>3E</td>
<td>Attracts Indoor/outdoor</td>
<td>Coils, mats or liquid electrics or other devices</td>
<td>“Protects for up to x hours against &quot;XYZ&quot;</td>
<td>Aedes sp. and Culex sp. etc (Plodia sp. Vespula sp. Musca domestica)</td>
<td></td>
<td>Any pest claimed needs to be tested.</td>
</tr>
<tr>
<td>3F</td>
<td>Attracts and traps Indoor/outdoor</td>
<td>Sex pheromone</td>
<td>Attracts male insects and catches them in a (sticky) trap</td>
<td>Specific insects for which claims are made</td>
<td>Sex pheromones are species specific and should therefore be tested on the claimed target species.</td>
<td></td>
</tr>
<tr>
<td>3G</td>
<td>Repels Indoor</td>
<td>All</td>
<td>Protects against moths for up to x days/weeks.</td>
<td>Plodia interpunctella or Tineola bisselliella</td>
<td></td>
<td>adult males</td>
</tr>
<tr>
<td>3H</td>
<td>Repels flying insects on horses Indoor/outdoor</td>
<td>sprays, creams, gels, balms etc.</td>
<td>Repels flying insects, such as .... on horses</td>
<td></td>
<td>Claim to be accompanied by a specification of the range of species; for all of these appropriate efficacy data should be provided</td>
<td></td>
</tr>
<tr>
<td>3I</td>
<td>Repels mosquitoes on horses Indoor/outdoor</td>
<td>sprays, creams, gels, balms etc.</td>
<td>Repels mosquitoes on horses</td>
<td>against two species, namely Culex spec and Aedes spec</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3J</td>
<td>Repels mosquitoes on horses Indoor/outdoor</td>
<td>sprays, creams, gels, balms etc.</td>
<td>Repels tropical mosquitoes on horses</td>
<td>against Culex spec, Aedes spec, AND Anopheles spec.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3K</td>
<td>Repels 'gnats &amp; biting midges' on horses Indoor/outdoor</td>
<td>sprays, creams, gels, balms etc.</td>
<td>Repels gnats &amp; biting midges on horses</td>
<td>species prevalent in the region (Culicoides spec.)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3L</td>
<td>Repels flies on horses Indoor/outdoor</td>
<td>sprays, creams, gels, balms etc.</td>
<td>Repels horse flies (e.g. Tabanus bovines) on horses</td>
<td>against Tabanid species prevalent in the region, e.g. Tabanus bovinus</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3M</td>
<td>Repels flies Indoor/</td>
<td>sprays, creams,</td>
<td>Repels deer flies</td>
<td>Chrysops caecutiens</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Action</td>
<td>SITE</td>
<td>APPLICATION METHOD</td>
<td>CLAIM</td>
<td>TEST SPECIES</td>
<td>RATIONALE</td>
<td>NOTES</td>
</tr>
<tr>
<td>--------</td>
<td>------</td>
<td>--------------------</td>
<td>-------</td>
<td>--------------</td>
<td>-----------</td>
<td>-------</td>
</tr>
<tr>
<td>3N</td>
<td>on horses</td>
<td>indoor/outdoor</td>
<td>sprays, creams, gels, balms etc.</td>
<td>Repels stable flies on horses</td>
<td>Stomoxys calcitrans</td>
<td></td>
</tr>
<tr>
<td>3O</td>
<td>on horses</td>
<td>indoor/outdoor</td>
<td>sprays, creams, gels, balms etc.</td>
<td>Repels black flies on horses</td>
<td>Simulium spec., e.g. Simulium equinum</td>
<td></td>
</tr>
<tr>
<td>3P</td>
<td>on horses</td>
<td>indoor/outdoor</td>
<td>sprays, creams, gels, balms etc.</td>
<td>Repels sand flies on horses</td>
<td>Phlebotominae</td>
<td></td>
</tr>
<tr>
<td>3Q</td>
<td>on horses</td>
<td>indoor/outdoor</td>
<td>sprays, creams, gels, balms etc.</td>
<td>Repels warble flies on horses</td>
<td>Hypoderma spec. (e.g. H. bovis or H. lineatum)</td>
<td></td>
</tr>
<tr>
<td>3R</td>
<td>on horses</td>
<td>indoor/outdoor</td>
<td>sprays, creams, gels, balms etc.</td>
<td>Repels horn flies on horses</td>
<td>Haematobia irritans</td>
<td></td>
</tr>
<tr>
<td>3S</td>
<td>on horses</td>
<td>indoor/outdoor</td>
<td>sprays, creams, gels, balms etc.</td>
<td>Repels Face flies, House flies on horses</td>
<td>appropriate Musca spec (e.g. M. domestica, M. autumnalis, et cetera).</td>
<td></td>
</tr>
<tr>
<td>3T</td>
<td>on horses</td>
<td>indoor/outdoor</td>
<td>sprays, creams, gels, balms etc.</td>
<td>Repels biting flies on horses</td>
<td>at least 1 Tabanid and 1 Culicoides species, prevalent to the region</td>
<td></td>
</tr>
<tr>
<td>3U</td>
<td>on horses</td>
<td>indoor/outdoor</td>
<td>sprays, creams, gels, balms etc.</td>
<td>Repels Deer/sheep ticks on horses</td>
<td>Ixodes scapularis/ricinus</td>
<td></td>
</tr>
<tr>
<td>3V</td>
<td>on horses</td>
<td>indoor/outdoor</td>
<td>sprays, creams, gels, balms etc.</td>
<td>Repels ticks such as... on horses</td>
<td>Two tick species prevalent in the region, e.g. Ixodes scapularis and I. Ricinus. Claim to be accompanied by a specification of the range of species; for all of which efficacy data need to be presented.</td>
<td></td>
</tr>
</tbody>
</table>
Appendix 17. List of currently available standard test methods for product type 18 insecticides/acaricides and product type 19 repellents/attractants (as far as they concern insects and other arthropods)

Recognised standard methods for the efficacy testing of biocidal products intended for the control of insects, acarides and other arthropods. This list is derived from A-S Wernersson, 2008 (Efficacy testing of biocidal products. FB Engineering AB, Skärgårdsgatan 1, Göteborg, Sweden) with some changes and additions.

This is a list of available standard methods (as far as we know now of) without distinction on suitability, usefulness, repeatability, order of acceptability or robustness.

Table 41: Acronyms and Abbreviations

<table>
<thead>
<tr>
<th>Acronym</th>
<th>Full name</th>
</tr>
</thead>
<tbody>
<tr>
<td>AFPP</td>
<td>Association française de protection des plants</td>
</tr>
<tr>
<td>AATCC</td>
<td>American Association of Textile Chemists and Colors</td>
</tr>
<tr>
<td>AFNOR</td>
<td>Association française de normalisation (NF standards)</td>
</tr>
<tr>
<td>AOAC</td>
<td>Association of Official Analytical Chemists</td>
</tr>
<tr>
<td>ASTM</td>
<td>American Society of Testing and Materials</td>
</tr>
<tr>
<td>ATCC</td>
<td>American Type Culture Collection</td>
</tr>
<tr>
<td>BBA</td>
<td>Federal Biological Research Centre for Agriculture and Forestry (Biologische Bundesanstalt für Land - Und Forstwirtschaft Bundesrepublik Deutschland)</td>
</tr>
<tr>
<td>BP</td>
<td>Biocidal Product</td>
</tr>
<tr>
<td>BPD</td>
<td>Biocidal Product Directive (referring to 98/8/EG)</td>
</tr>
<tr>
<td>BSI</td>
<td>British Standards Institute (BS standards)</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Description</td>
</tr>
<tr>
<td>--------------</td>
<td>-------------</td>
</tr>
<tr>
<td>CA</td>
<td>Competent Authority</td>
</tr>
<tr>
<td>CEB</td>
<td>Commission Des Essais Biologiques</td>
</tr>
<tr>
<td>CEFIC</td>
<td>European Chemical Industry Council</td>
</tr>
<tr>
<td>CEN</td>
<td>European Committee for Standardisation</td>
</tr>
<tr>
<td>CEPE</td>
<td>European council of paint, printing inks and artist’s colours industry</td>
</tr>
<tr>
<td>CSMA</td>
<td>Chemical Specialties Manufactures Association</td>
</tr>
<tr>
<td>EBPF</td>
<td>European Biocidal Product Forum</td>
</tr>
<tr>
<td>EPA</td>
<td>United States Environmental Protection Agency</td>
</tr>
<tr>
<td>EPPO</td>
<td>European and Mediterranean Plant Protection Organization</td>
</tr>
<tr>
<td>ISO</td>
<td>International Standards Organisation</td>
</tr>
<tr>
<td>MAFF</td>
<td>Ministry of Agriculture Fisheries and Foods</td>
</tr>
<tr>
<td>MS</td>
<td>Malaysian Standards</td>
</tr>
<tr>
<td>NF</td>
<td>NF standards, Association française de normalisation</td>
</tr>
<tr>
<td>OECD</td>
<td>Organisation for Economic Co-operation and Development</td>
</tr>
<tr>
<td><strong>OCSPP</strong></td>
<td>Office of Chemical Safety and Pollution Prevention (old name OPPTS)</td>
</tr>
<tr>
<td>-----------</td>
<td>---------------------------------------------------------------</td>
</tr>
</tbody>
</table>
| **OPPTS** | Office of Prevention, Pesticides and Toxic Substances, United States Environmental Protection Agency  
New name: Office of Chemical Safety and Pollution Prevention (OCSPP) | www.epa.gov/internet/oppts/ |
| **PT**    | Product Type                                                  |                                                                      |
| **SABS**  | South African Bureau of Standards                             | www.sabs.co.za                                                      |
| **AFPP**  | Association française de protection des plants                |                                                                      |
| **AATCC** | American Association of Textile Chemists and Colors           | www.aatcc.org/                                                      |

**REFERENCE LISTS FOR TABLES 42-48**

- GENERAL
- CRAWLING INSECTS
  - Cockroaches
  - Termites
  - Other crawling insects
- FLYING INSECTS
- INSECTICIDES AGAINST TEXTILE AND STORED PRODUCT PESTS
- REPELLENTS & ATTRACTANTS
<table>
<thead>
<tr>
<th>Reference</th>
<th>Title</th>
<th>PT</th>
<th>Short test description (if test method available or information provided from elsewhere)</th>
<th>Type of Reference Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>OPPTS 810.3000</td>
<td>General Considerations for Efficacy of Invertebrate Control Agents</td>
<td>18</td>
<td>General guide</td>
<td>Manufacturer; UK guidelines</td>
</tr>
<tr>
<td>CEB 196 (1997)</td>
<td>Trial method to evaluate the efficacy of insecticidal bait products against common species</td>
<td>18</td>
<td></td>
<td>TM II05 (Fr)</td>
</tr>
<tr>
<td>EPPO pp1/152 (3)</td>
<td>Design and analysis of efficacy evaluation trials</td>
<td></td>
<td>This standard provides detailed advice on the design and analysis of efficacy evaluation trials. Primarily intended for use in plant protection but also very useful for biocides.</td>
<td>EPPO web site</td>
</tr>
<tr>
<td>EPPO pp1/181 (3)</td>
<td>Conduct and reporting of efficacy evaluation trials, including good experimental practice</td>
<td></td>
<td>This standard provides guidance on how to organize trials, and how to plan, conduct and assess them, then record and interpret them, so as to obtain comparable and reliable results. It is also based on the principle that trials should be performed according to Good Experimental Practice (GEP).</td>
<td>EPPO website</td>
</tr>
<tr>
<td>OPPTS 810.3200</td>
<td>Livestock, poultry, fur- and wool-bearing animal treatments</td>
<td>18</td>
<td></td>
<td>Own searches</td>
</tr>
<tr>
<td>OPPTS 810.3300</td>
<td>Treatments to control pests of humans and pets</td>
<td>18</td>
<td></td>
<td>UK guidelines</td>
</tr>
<tr>
<td>Reference</td>
<td>Title</td>
<td>PT</td>
<td>Short test description (if test method available or information provided from elsewhere)</td>
<td>Type of Reference Source</td>
</tr>
<tr>
<td>-------------------</td>
<td>------------------------------------------------------------------------</td>
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<td>-----------------------------------------------------------------------------------------</td>
<td>-------------------------------------------</td>
</tr>
<tr>
<td>OPPTS 810.3500</td>
<td>Premises treatments</td>
<td>18</td>
<td>General guideline</td>
<td>Manufacturer; UK guidelines</td>
</tr>
<tr>
<td>SABS 233 1st rev</td>
<td>Pesticides: Biological evaluation of mists and fogs - first revision</td>
<td>18</td>
<td></td>
<td>Manufacturer</td>
</tr>
<tr>
<td>SABS 576</td>
<td>Pesticides – Biological evaluation of insecticidal oil-based space spray in low-pressurized dispensers - first revision</td>
<td>18</td>
<td></td>
<td>Manufacturer</td>
</tr>
<tr>
<td>SABS 583</td>
<td>Pesticides – Biological evaluation of the contact efficacy of liquid residual insecticides - first revision</td>
<td>18</td>
<td></td>
<td>Manufacturer</td>
</tr>
<tr>
<td>SABS 6136 (2003)</td>
<td>Pesticides – Biological evaluation of materials that release an insecticide upon heating</td>
<td>18</td>
<td></td>
<td>Manufacturer</td>
</tr>
<tr>
<td>SABS 690 (DRAFT)</td>
<td>Pesticides: biological evaluation of the properties of solid fly baits - DRAFT</td>
<td>18</td>
<td></td>
<td>Manufacturer</td>
</tr>
<tr>
<td>SABS 807</td>
<td>Methods for testing insecticides against flying and crawling insects.</td>
<td>18</td>
<td></td>
<td>TNsG on Prod Evaluation; Manufacturer; UK guidelines</td>
</tr>
<tr>
<td>SABS 899 (1987)</td>
<td>Insecticidal space spray in pressurized dispensers</td>
<td>18</td>
<td></td>
<td>Manufacturer</td>
</tr>
</tbody>
</table>
### Table 43: Crawling Insects: Cockroaches

<table>
<thead>
<tr>
<th>Reference</th>
<th>Title</th>
<th>PT</th>
<th>Short test description (if test method available or information provided from elsewhere)</th>
<th>Type of Reference Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>AFPP</td>
<td>Methode d’essai d’efficacite, en laboratoire et en conditions pratiques d’utilisation, d’appats insecticides destines a la lutte contre les blattes dans les locaux Efficacy trials method, in laboratory or in practical conditions of use, for insecticide baits intended to control cockroaches in premises</td>
<td>18</td>
<td></td>
<td>French guideline</td>
</tr>
<tr>
<td>ASTM E654-96(2003)</td>
<td>Standard Test Method for Effectiveness of Aerosol and Pressurized Spray Insecticides Against Cockroaches</td>
<td>18</td>
<td>Test of insecticides against crawling insects: cockroaches Determines the relative efficiency of aerosol and pressurised spray formulations against cockroaches, but test data by this test method may also be adequate to support claims for use of the product to control the exposed or accessible stages of silverfish, ants, centipedes, spiders, and certain stored product pests. Applied as direct sprays for 30 s. on last instar nymphs. Observation period: 48h. The test is not designed to measure the residual action.Ten groups with 20 organisms in each. The test is run in conjunction with the Official Test Aerosol II (OTA II) (or Tentative Official Aqueous Pressurized Spray (TOAPS)</td>
<td>UK guidelines; Test institute</td>
</tr>
<tr>
<td>Reference</td>
<td>Title</td>
<td>PT</td>
<td>Short test description (if test method available or information provided from elsewhere)</td>
<td>Type of Reference Source</td>
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<tr>
<td>CEB 159 (1992)</td>
<td>Trial method to evaluate the efficacy of insecticidal products for the control of cockroaches in buildings under practical conditions</td>
<td>18</td>
<td>as the standard basis of comparison. The mortality after 24h should be between 50 and 75% when testing with the OTA. The test specimens meet the standard if average % dead and moribund is equal to, above or within 10% points less than average % dead of the OTA series after 48h. Precision or bias is not specified, only states whether conforms to efficacy criteria.</td>
<td>TM II05 (Fr)</td>
</tr>
<tr>
<td>SABS 458</td>
<td>Pesticides – Rearing and handling of the German cockroach (Blatella germanica (L.)) - second revision</td>
<td>18</td>
<td></td>
<td>Manufacturer</td>
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<tr>
<td>WHO/VBC/75.593 (1981)</td>
<td>Instructions for determining the susceptibility or resistance of cockroaches to insecticides</td>
<td>18</td>
<td></td>
<td>TNsG on Prod Evaluation; UK guidelines</td>
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</table>
### Table 44: Crawling Insects: Termites

**CRAWLING INSECTS: Termites**

<table>
<thead>
<tr>
<th>Reference</th>
<th>Title</th>
<th>PT</th>
<th>Short test description (if test method available or information provided from elsewhere)</th>
<th>Type of Reference Source</th>
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</thead>
<tbody>
<tr>
<td>CTBA-BIO-E-001</td>
<td>Epreuve de vieillissement naturel des murs traités.</td>
<td>18</td>
<td>Test institute</td>
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<tr>
<td>CTBA-BIO-E-002</td>
<td>Epreuve de vieillissement naturel des sols traités.</td>
<td>18</td>
<td>Test institute</td>
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<tr>
<td>CTBA-BIO-E-007</td>
<td>Evaluation de l'efficacité anti-termite d'une barrière placée en milieu alcalin.</td>
<td>18</td>
<td>Test institute</td>
<td></td>
</tr>
<tr>
<td>CTBA-BIO-E-008/2</td>
<td>Evaluation de l'efficacité anti-termite d'une barrière physico-chimique - Essai de terrain - Dispositif sans dalle de béton.</td>
<td>18</td>
<td>Test institute</td>
<td></td>
</tr>
<tr>
<td>CTBA-BIO-E-008/3</td>
<td>Evaluation de l'efficacité anti-termite d'une barrière - Essai de terrain - Dispositif avec dalle de béton.</td>
<td>18</td>
<td>Test institute</td>
<td></td>
</tr>
<tr>
<td>CTBA-BIO-E-016</td>
<td>Version 2 : Exposition de barrières physico-chimiques anti-termites aux rayonnements solaires.</td>
<td>18</td>
<td>Test institute</td>
<td></td>
</tr>
<tr>
<td>FCBA-BIO-E-038</td>
<td>Evaluation de l'efficacité d'un traitement insecticide des déchets de démolition infestés par les termites - Essai de laboratoire.</td>
<td>18</td>
<td>Test institute</td>
<td></td>
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<tr>
<td>FCBA-BIO-E-039</td>
<td>Evaluation de l'efficacité d'un traitement insecticide des déchets de démolition infestés par les termites - Essai de terrain.</td>
<td>18</td>
<td>Test institute</td>
<td></td>
</tr>
<tr>
<td>Reference</td>
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<tr>
<td>FCBA-BIO-E-041</td>
<td>Critères de performance des méthodes d’essais CTBA-BIO-E-xx et FCBA-BIO-E-xx</td>
<td>18</td>
<td></td>
<td>Test institute</td>
</tr>
<tr>
<td>ENV 1250-2</td>
<td>Wood preservatives - Methods for measuring losses of active ingredients and other preservative ingredients from treated timber - Part 2: Laboratory method for obtaining samples for analysis to measure losses by leaching into water or synthetic sea water</td>
<td>18</td>
<td></td>
<td>International guideline</td>
</tr>
<tr>
<td>NF X 41-542</td>
<td>Produits de préservation du bois - Produit de traitement antitermites des sols, murs, fondations et maçonneries - Epreuve de vieillissement accéléré des matériaux traités avant essais biologiques - Epreuve de percolation.</td>
<td>8+1</td>
<td>French guideline. Wood preservatives - Anti-termite treatment product for floors, walls, foundations, and masonry work - Accelerated ageing test of treated materials prior to biological testing - Percolation test.</td>
<td>French guideline</td>
</tr>
<tr>
<td>NF X 41-543-2, 2008</td>
<td>Produits de préservation du bois - Détermination de l’efficacité d’un système de pièges-appâts - Partie 2 : Efficacité du système - Méthode de terrain</td>
<td>8+1</td>
<td>Wood preservatives — Determination of the efficacy of a bait-trap system — Part 2: Efficacy of the insecticide formulation — Field method. This test method is intended to evaluate the efficacy of the baits in an experimental site where termite activity is reported. Consumption of the tested bait must be registered at least in the first 6 months after the</td>
<td>French guideline</td>
</tr>
<tr>
<td>Reference</td>
<td>Title</td>
<td>PT</td>
<td>Short test description (if test method available or information provided from elsewhere)</td>
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<tr>
<td>NF X 41-543-3, 2009</td>
<td>Critères de performance des essais pièges-appâts</td>
<td>8+</td>
<td>introduction of the baits. The elimination of termites in the experimental site should be registered maximum after 18 months (counted since the introduction of the first tested bait), excluding the winter period.</td>
<td>French guideline</td>
</tr>
<tr>
<td>NF X 41-550</td>
<td>Termites - Determination of the effectiveness against termites of products or materials used as barrier designed for ground and/or wall - Laboratory method</td>
<td>8+</td>
<td></td>
<td>French guideline</td>
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<tr>
<td>NF X 41-551</td>
<td>Termites - Determination of the effectiveness against termites of products or material used as barrier designed for ground and/or wall - Performance criteria</td>
<td>8+</td>
<td></td>
<td>French guideline</td>
</tr>
<tr>
<td>OPPTS 810.3800</td>
<td>Methods for efficacy testing of termite baits</td>
<td>8+</td>
<td></td>
<td>Own searches</td>
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</table>

**Table 45: Crawling Insects: Other Crawling Insects**

**CRAWLING INSECTS: Other crawling insects**

<table>
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<th>Reference</th>
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<th>Type of Reference Source</th>
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<tr>
<td>AATCC 194-2006</td>
<td>Assessment of the Anti-House Dust Mite Properties of Textiles under Long-Term Test Conditions</td>
<td>18</td>
<td>Applied to textiles</td>
<td>Manufacturer</td>
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<tr>
<td>OCSPP 810.3900</td>
<td>Draft Product Performance Test Guidelines Laboratory Testing</td>
<td>PT</td>
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### Guidance on the BPR: Volume II Parts B+C

**Version 1.0 February 2017**

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#### Table 46: Flying Insects

**FLYING INSECTS**

<table>
<thead>
<tr>
<th>Reference</th>
<th>Title</th>
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<th>Short test description (if test method available or information provided from elsewhere)</th>
<th>Type of Reference Source</th>
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<tr>
<td>OPPTS 810.3100</td>
<td>Soil treatments for imported fire ants</td>
<td>18</td>
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<td>Own searches</td>
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<tr>
<td>US AATCC Technical Manual Method 24 (1992)</td>
<td>Test method for textiles to determine resistance to insects (e.g. moths, carpet beetles)</td>
<td>18</td>
<td>Efficacy test against larvae</td>
<td>TNSG on Prod Evaluation; UK guidelines</td>
</tr>
<tr>
<td>WHO/VBC/81.809 (1981)</td>
<td>Instructions for determining the susceptibility or resistance of adult bed-bugs to insecticides</td>
<td>18</td>
<td></td>
<td>TNSG on Prod Evaluation; UK guidelines</td>
</tr>
<tr>
<td>WHO/VBC/81.814 (1981)</td>
<td>Instructions for determining the susceptibility or resistance of adult ticks to insecticides</td>
<td>18</td>
<td></td>
<td>TNSG on Prod Evaluation; UK guidelines</td>
</tr>
<tr>
<td>WHO/VBC/81.815 (1981)</td>
<td>Instructions for determining the susceptibility or resistance of fleas to insecticides</td>
<td>18</td>
<td></td>
<td>TNSG on Prod Evaluation; UK guidelines</td>
</tr>
<tr>
<td>ASTM E652-91(2009)</td>
<td>Standard Test Method for Nonresidual Liquid Household Insecticides Against Flying Insects</td>
<td>18</td>
<td>Determines the relative efficiency of household and industrial-use, contact insecticides dissolved in base oils and applied in spray formulations. It is developed to test insecticides against house flies (<em>Musca domestica</em>, L), but test data may also be adequate to support label claims for the use of the products against mosquitoes, gnats, flying moths, wasps, and certain</td>
<td>Own searches</td>
</tr>
<tr>
<td>Reference</td>
<td>Title</td>
<td>PT</td>
<td>Short test description (if test method available or information provided from elsewhere)</td>
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<tr>
<td>ASTM E653-91</td>
<td>Standard Test Method for Effectiveness of Aerosol and Pressurized Space Spray Insecticides Against Flying Insects</td>
<td>18</td>
<td>other small flying insects. Not designed to measure the residual action of the spray formulation. For Liquids, dose: 12 cm³, 100 flies, test chambers: Peet grady Chambers (6.02 m³), Test conditions: 27ºC, 50%H.R. It has been superseded by ASTM</td>
<td>Test institute</td>
</tr>
<tr>
<td>BS 4172-1:1999</td>
<td>Hand-held pressurized aerosol dispensers against houseflies.</td>
<td>18</td>
<td>The test determines the relative efficacy of aerosol and pressurized space spray insecticide formulations against house flies (Musca domestica, L) strains and, with modifications in dosage, other flying insects. Test data obtained by this test method may also be adequate to support label claims for the use of the product against mosquitoes, gnats, flying moths, wasps, and certain other small flying insects. This test method is not designed to measure the residual activity. The test may be conducted using approximately 100 house flies per test (small group) or 500 flies per test (large group). Selected reference standards are the Official Test Aerosol II (OTA II) for oil based aerosol products and Tentative Official Aqueous Pressurized Spray (TOAPS) for water based aerosol products. Aerosol test knockdowns: % down of total flies at 5, 10, 15 minutes after application. Aerosol test knock down mortality: dead knocked down x100/total flies. These numbers should on average be equal to, greater than or no more than 5% points below the corresponding numbers of the reference in order to meet the standard. No statement on precision or bias, only whether conformance to criteria for success specified in the procedure. For Sprays, dose: 3g/28m³, 100 flies, test chambers: Peet grady Chambers (6.02 m³), Test conditions: 27ºC, 50%H.R. It has been superseded by ASTM</td>
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<td></td>
<td></td>
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<td></td>
<td>TNsG on Prod Evaluation; TM II05</td>
</tr>
<tr>
<td>Reference</td>
<td>Title</td>
<td>PT</td>
<td>Short test description (if test method available or information provided from elsewhere)</td>
<td>Type of Reference Source</td>
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</tr>
<tr>
<td>BS 4172-2:1 1999</td>
<td>Specification for insecticidal performance</td>
<td></td>
<td>For Sprays, dose: 35.3g/50m3, 100 flies, test chambers: 25 – 60 m3, Test conditions: 26°C, 45-75% H.R. The reference product is very well described, and easy to manufacture.</td>
<td>(Fr); UK guidelines</td>
</tr>
<tr>
<td>CEB 107 (1985)</td>
<td>Hand-held pressurized aerosol dispensers against houseflies</td>
<td>18</td>
<td></td>
<td>TNsG on Prod Evaluation; TM II05 (Fr); UK guidelines; Test institute</td>
</tr>
<tr>
<td>MS 1398 part 2</td>
<td>Trial method to evaluate the efficacy of insecticidal products for</td>
<td>18</td>
<td></td>
<td>Manufacturer</td>
</tr>
<tr>
<td>(1996)</td>
<td>the control of stable flies in premises for the rearing of domestic</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>animals under practical conditions</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>MS 1398 part 3</td>
<td>Specification for mosquito electric liquid vapourizer: part 2:</td>
<td>18</td>
<td></td>
<td>Manufacturer</td>
</tr>
<tr>
<td>(1996)</td>
<td>method for evaluation of biological efficacy - glass chamber method</td>
<td></td>
<td></td>
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</tr>
<tr>
<td></td>
<td>method for evaluation of biological efficacy - glass cylinder method</td>
<td></td>
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<tr>
<td>MS 23 part 1</td>
<td>Methods of biological evaluation of the efficacy of repellent -</td>
<td>18</td>
<td></td>
<td>Manufacturer</td>
</tr>
<tr>
<td>(1998)</td>
<td>bioassay method for mosquito repellent on human skin</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>MS 23 part 2</td>
<td>Specification for mosquito coils: Part 1: physical and chemical</td>
<td>18</td>
<td></td>
<td>Manufacturer</td>
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<tr>
<td>(1996)</td>
<td>requirements (third revision)</td>
<td></td>
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<td></td>
<td>Specification for mosquito coils: Part 1: method for evaluation of</td>
<td>18</td>
<td></td>
<td>Manufacturer</td>
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<td></td>
<td>biological efficacy - glass chamber</td>
<td></td>
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<tr>
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</tr>
<tr>
<td>NF T72-320 March 1977</td>
<td>Insecticides for flying insects. Insecticide distributed under pressure (&quot;aerosol&quot; type). Determination of the efficiency rating.</td>
<td>18</td>
<td>For Aerosols, dose: 1seg/10m3, 100 flies, test chambers 25-50 cubic meters, Test conditions:25ºC, 60%H.R.</td>
<td>TNsG on Prod Evaluation; TM II05 (Fr); UK guidelines; Test institute</td>
</tr>
<tr>
<td>NF T72-321 March 1977</td>
<td>Insecticides for flying insects. Permanent insecticide distributor. Determination of the efficiency rating and the regularity rating.</td>
<td>18</td>
<td>For Vaporizers, 100 flies, test chambers 25-50 cubic meters, Test conditions:25ºC, 60%H.R.</td>
<td>TM II05 (Fr); Test institute</td>
</tr>
<tr>
<td>OPPTS 810.3400</td>
<td>Mosquito, black fly, and biting midge (sand fly) treatments</td>
<td>18</td>
<td>Test of insecticides against flying insects: Mosquito, Black Fly and Biting Midge (Sand Fly)</td>
<td>UK guidelines</td>
</tr>
<tr>
<td>Verwey &amp; Sosa, 2007</td>
<td>Liquid Electric test method</td>
<td>18</td>
<td>For testing pyrethroids (draft method) and natural actives (Pyrethrum extract) on mosquitoes (knockdown). Efficacy criteria: &quot;effective against mosquitoes for X hours&quot;. Knockdown is measured repeatedly for 2h and mortality after 24h. Control (no treatment) knockdown: maximum 10%. 2-4 chamber replicates, 50 organisms in each. Mean and Standard Deviations for each time calculated as well as KT50 and KT80 (Mean time to 50% and 80% knockdown respectively).</td>
<td>Manufacturer</td>
</tr>
<tr>
<td>Reference</td>
<td>Title</td>
<td>PT</td>
<td>Short test description (if test method available or information provided from elsewhere)</td>
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<td>WHO/VBC/81.212 (1981)</td>
<td>Instructions for determining the susceptibility or resistance of mosquito larvae to insect development inhibitors</td>
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<td>TNsG on Prod Evaluation</td>
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<tr>
<td>WHO/VBC/81.806</td>
<td>Instructions for determining the susceptibility or resistance of adult mosquitoes to organochlorine, organophosphate and carbamate insecticides - diagnostic test</td>
<td>18</td>
<td></td>
<td>TNsG on Prod Evaluation; UK guidelines</td>
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<tr>
<td>WHO/VBC/81.807 (1981)</td>
<td>Instructions for determining the susceptibility or resistance of mosquito larvae to insecticides</td>
<td>18</td>
<td></td>
<td>TNsG on Prod Evaluation; UK guidelines</td>
</tr>
<tr>
<td>WHO/VBC/81.811 (1981)</td>
<td>Instructions for determining the susceptibility or resistance of blackfly larvae to insecticides</td>
<td>18</td>
<td></td>
<td>TNsG on Prod Evaluation; UK guidelines</td>
</tr>
<tr>
<td>WHO/VBC/81.812 (1981)</td>
<td>Instructions for determining the susceptibility or resistance of mosquito larvae to insect development inhibitors</td>
<td>18</td>
<td></td>
<td>UK guidelines</td>
</tr>
<tr>
<td>WHO/VBC/81.813 (1981)</td>
<td>Instructions for determining the susceptibility or resistance of houseflies, tsetse flies, stable flies, blowflies etc. to insecticides</td>
<td>18</td>
<td></td>
<td>TNsG on Prod Evaluation; UK guidelines</td>
</tr>
<tr>
<td>WHO/CVB/81.5</td>
<td>Instruction for the bio-assay of insecticidal deposits on wall surfaces</td>
<td>18</td>
<td>For Vaporizers, 100 flies, test chambers 25-50 cubic meters, Test conditions:25ºC, 60%H.R.</td>
<td>Test institute</td>
</tr>
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</table>
### Reference Title

<table>
<thead>
<tr>
<th>Reference</th>
<th>Title</th>
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<tbody>
<tr>
<td>WHO/CDS/WHOPE S/GCDPP/2003.5</td>
<td>Space spray application of insecticides for vector and public health pest control – a practitioner’s guide</td>
<td>18</td>
<td>Brief description of the main types of space spray equipment as well as the operational guidelines for space spray application of insecticides.</td>
<td>TM II05 (Fr)</td>
</tr>
<tr>
<td>WHO/CDS/WHOPE S/GCDPP/2005.13</td>
<td>Guidelines for laboratory and field testing of mosquito larvicides</td>
<td>18</td>
<td>This document provides specific and standardized procedures and guidelines for testing larvicides, including bacterial larvicides and insect growth regulators against mosquitoes.</td>
<td>WHO 2005</td>
</tr>
<tr>
<td>WHO/CDS/NTD/W HOPES/GCDPP/2006.3</td>
<td>Guidelines for testing mosquito adulticides for indoor residual spraying and treatment of mosquito nets</td>
<td>18</td>
<td>This document provides specific and standardized procedures and guidelines for testing mosquito adulticides for indoor residual spraying and for treatment of mosquito nets.</td>
<td>WHO 2006</td>
</tr>
<tr>
<td>WHO/HTM/NTD/WHOPES/2009.2</td>
<td>Guidelines for efficacy testing of insecticides for indoor and outdoor ground-applied space spray applications</td>
<td>18</td>
<td>The document provides guidance and stepwise procedures on laboratory studies, field testing and evaluation leading to the determination of efficacy, and application rates of insecticides for operational use in indoor and outdoor ground-applied space spray applications. While most examples provided pertain to mosquitoes, with some modifications the guidelines can be used to determine efficacy against other flying vectors and pests.</td>
<td>WHO 2009</td>
</tr>
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### Table 47: Insecticides Against Textile and Stored Product Pests

**INSECTICIDES AGAINST TEXTILE AND STORED PRODUCT PESTS**

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<thead>
<tr>
<th>Reference</th>
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<th>PT</th>
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<th>Type of Reference Source</th>
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<tr>
<td>CEB 135bis (1996)</td>
<td>Laboratory test method to evaluate the efficacy of insecticidal products in premises for the storage, industrial processing and sale of</td>
<td>18</td>
<td>Space treatments</td>
<td>TM II05 (Fr)</td>
</tr>
<tr>
<td>Reference</td>
<td>Title</td>
<td>PT</td>
<td>Short test description (if test method available or information provided from elsewhere)</td>
<td>Type of Reference Source</td>
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<tr>
<td>CEB 213 (1999)</td>
<td>Trial method to evaluate the efficacy of a fumigant for insect control in premises for the storage, processing and production of food</td>
<td>18</td>
<td></td>
<td>TM II05 (Fr)</td>
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<tr>
<td>CEB 224 (2001)</td>
<td>Trial method to evaluate the efficacy of fumigants for insect control in stored products</td>
<td>18</td>
<td></td>
<td>TM II05 (Fr)</td>
</tr>
<tr>
<td>EPPO PP 1/201(1)</td>
<td>Fumigants to control insect and mite pests of stored plant products</td>
<td>18 + 20</td>
<td></td>
<td>TM II05 (Fr); UK guidelines</td>
</tr>
<tr>
<td>EPPO PP 1/202(1)</td>
<td>Space and structural treatments of store rooms</td>
<td>18</td>
<td></td>
<td>TM II05 (Fr); UK guidelines</td>
</tr>
<tr>
<td>EPPO PP 1/203(1)</td>
<td>Admixture of plant protection products to stored plant products to control insects and mites</td>
<td>18 + 20</td>
<td></td>
<td>TM II05 (Fr)</td>
</tr>
<tr>
<td>EPPO PP 1/204(1)</td>
<td>Laboratory testing of plant protection products against insect and mite pests of stored plant products</td>
<td>18</td>
<td></td>
<td>UK guidelines</td>
</tr>
<tr>
<td>NF G39-011 April 2001</td>
<td>Properties of textiles - Textiles and polymeric materials having</td>
<td>18</td>
<td></td>
<td>Manufacturer; TM II05 (Fr)</td>
</tr>
<tr>
<td>Reference</td>
<td>Title</td>
<td>PT</td>
<td>Short test description (if test method available or information provided from elsewhere)</td>
<td>Type of Reference Source</td>
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<tr>
<td>NF X41-516 January 1980</td>
<td>Protection of textiles. Protection against certain insect pests. Methods of testing.</td>
<td>18</td>
<td></td>
<td>TM II05 (Fr)</td>
</tr>
<tr>
<td>SABS 332</td>
<td>Pesticides – Rearing and handling of the common clothes moth (Tineola bisselliella Hummel) - second revision</td>
<td>18</td>
<td></td>
<td>Manufacturer</td>
</tr>
<tr>
<td>ISO 3998</td>
<td>Determination of resistance to certain insect pests</td>
<td>18</td>
<td>For treated materials. Comparing the resistant material against a non-resistant material.</td>
<td>Test institute</td>
</tr>
<tr>
<td>US AATCC Technical Manual Method 24 (1992)</td>
<td>Test method for textiles to determine resistance to insects (e.g. moths, carpet beetles)</td>
<td>18</td>
<td>Efficacy test against larvae</td>
<td>TNSG on Prod Evaluation; UK guidelines</td>
</tr>
<tr>
<td>Reference</td>
<td>Title</td>
<td>PT</td>
<td>Short test description (if test method available or information provided from elsewhere)</td>
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<tr>
<td>ASTM E939-94(2012)</td>
<td>Standard Test Method of Field Testing Topical Applications of Compounds as Repellents for Medically Important and Pest Arthropods (Including Insects, Ticks, and Mites): Mosquitoes</td>
<td>19</td>
<td>Evaluates the repellency of promising compounds that have undergone primary laboratory studies and approved for skin application for secondary testing. The method is designed for the study of mosquito repellents, but can be modified to determine the repellency of candidate compounds for other flying insects that attack humans.</td>
<td>Own search</td>
</tr>
<tr>
<td>ASTM E951-94(2006)</td>
<td>Standard Test Methods for Laboratory Testing of Non-Commercial Mosquito Repellent Formulations On the Skin</td>
<td>19</td>
<td>Can be used to test the efficacy of repellent compounds that can be diluted with ethanol, acetone etc. Both biological effectiveness and persistence of the repellent can be assessed. ED50 and ED95 are determined for comparative and practical purposes respectively. Precision of the test can be evaluated (confidence intervals).</td>
<td>Own search</td>
</tr>
<tr>
<td>Dautel H, Kahl O, Siems K, Oppenrieder M, Müller-Kuhr L, Hilker M. Ent Exp Appl. 1999;91:431–441</td>
<td>A novel test system for detection of tick repellents</td>
<td>19</td>
<td>The so-called Moving Object Bioassay is described, a tool for testing the strength of potential tick repellents quantitatively. Endpoint measured is the attachment rate of Ixodes ticks.</td>
<td>Dossier</td>
</tr>
<tr>
<td>Fradin &amp; Day, July 2002, N Engl J Med vol 347 vol 13-18</td>
<td>Comparative efficacy of insect repellents against mosquito bites</td>
<td>19</td>
<td>Human subjects: Arm in cage studies (15 volunteers, 10 mosquitoes (Aedes aegypti) in each cage. Endpoint: elapsed time to first bite. Category of protection A-H (significantly different mean complete protection time; ANOVA &amp; Tukey's). No need to recalculate the results to &quot;real condition&quot; (simulate real condition)</td>
<td>Dossier</td>
</tr>
<tr>
<td>Hummel, E., Kleeberg, H. 1997.</td>
<td>Effect of the neem extract formulation neemazal-t/s on the</td>
<td>19</td>
<td></td>
<td>Dossier</td>
</tr>
<tr>
<td>Reference</td>
<td>Title</td>
<td>PT</td>
<td>Short test description (if test method available or information provided from elsewhere)</td>
<td>Type of Reference Source</td>
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<tr>
<td>in: Practice orientated results on use and production of Neem-Ingredients and Pheromones V. Proceedings of the 5th workshop, Wetzlar, Germany, January 22-25, 1996</td>
<td>green pea aphid acyrthosiphon pism in the laboratory (1995), in: Practice orientated results on use and production of Neem-Ingredients and Pheromones V</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SABS 695</td>
<td>Pesticides – Biological evaluation of the efficacy of mosquito repellents - first revision</td>
<td>19</td>
<td></td>
<td>Manufacturer</td>
</tr>
<tr>
<td>US EPA Guideline, OPPTS 810.3700 (2010); EPA 712-C-10-001</td>
<td>Insect repellents to be applied to human skin</td>
<td>19</td>
<td></td>
<td>UK guidelines</td>
</tr>
<tr>
<td>WHO/HTM/NTD/W HOPES/2009.4</td>
<td>Guidelines for efficacy testing of mosquito repellents for human skin</td>
<td>19</td>
<td>The purpose of these guidelines is to provide specific and standardized procedures and criteria for efficacy testing and evaluation of mosquito repellents for human skin. Their aim is to harmonize the testing procedures carried out in different laboratories and institutions in order to generate comparable data for registering and labelling such products by the national regulatory authorities.</td>
<td>WHO</td>
</tr>
</tbody>
</table>
Appendix 18. Efficacy guideline with Cockroach; field trial

This guidance describes an example of a field trial to determine efficacy of a product against the German cockroach (*Blattella germanica*).

**Global design**

In a pre-test it is established whether the population of cockroaches in an object is large enough for a field trial. An indication of the population size is obtained in the pre-test by using a spray with expelling action or by setting glue traps.

If the population size is large enough, a pest control operation is performed. The efficacy of the product is determined by measuring the population size again 8 weeks later and comparing it to the initial value.

During these 8 weeks the effect of the control operation should be checked at least 4 times at regular intervals (possibly using glue traps). The investigator himself should perform these checks during the trial.

**Requirements for the practical use situation in order to be suitable as test object.**

The field trial is performed in three separate objects.

Recommendations for the practical use situation to produce a good field trial for control of the German cockroach are as follows:

1. History of insecticide use should be described with as much detail as possible (which product, active ingredient, when ...). Object with recent insecticide use should not be included in the test.

2. The test object should preferably and where possible be hermetically sealed off from the surrounding buildings. If there are adjacent buildings, all cracks and crevices on the outside of the test object should be treated with an authorised biocidal product with residual action.

3. The test object should preferably contain at least a kitchen or kitchen unit, with one or more refrigerators or freezers.

4. Cockroaches should be present in the test object, both in the kitchen or kitchen unit as elsewhere.

5. In the preceding 8 weeks no other chemical control of cockroaches should have taken place in the test object.

**Field trial**

**The pre-test**

**Aim:** To determine whether the population is large enough for a field trial.

**Execution:** Within 1 week before the control operation.

The pre-test can be conducted in two different ways.

1. By using a spray liquid with an expelling action (e.g. pyrethrins):
   - Spray under the refrigerator and one other place in the kitchen where there are probably many cockroaches.
   - Spray for 3 seconds and count the cockroaches that emerge during 1 minute.
2. By using glue traps
   Place glue traps at places where many cockroaches are expected.
   Number per unit area: 5 glue traps per 100 m²
   Describe clearly where the glue traps are placed, and record the number of trapped cockroaches after an appropriate period, usually either overnight, or after up to 3 days (e.g. weekend), depending upon the scale of the infestation (shorter trap periods for heavier infestations to avoid traps becoming saturated and failing to catch cockroaches later during the monitoring period; longer periods when infestation level is low and few cockroaches are trapped each night).

Criteria for a suitable test object
- When a trap is placed for 48 hours in the kitchen or in the kitchen unit behind the refrigerator, it should contain at least 10 adult cockroaches at the end of this time, as well as several nymphs.
- Several cockroaches should be caught on at least one glue trap, which is placed at another place in the kitchen or kitchen unit and on one trap, which is placed outside the kitchen or kitchen unit, within 48 hours.

Or:
- When using a spray with expelling action, at least 5-10 cockroaches per sprayed site should be counted.

The test
Duration of the control period until measurement of efficacy is about 8 weeks.
The pest control is performed according to the directions for use of the product. During these 8 weeks the investigator will check the progress of the control at least 4 times.

Directions for use of an insecticide in the form of a spray liquid:
- It should be clear how much product is used, on average 1 L/20 m² is sprayed;
- Treatment of cracks and crevices should be done where necessary;
- If stated on the label, a second treatment can be performed.

Directions for use of an insecticide in the form of a powder:
- It should be clear how much product is used.

Directions for use of an insecticide in the form of bait:
- Number of baits placed per unit area should be according to directions for use;
- Precise descriptions of where the baits are placed should be given;
- The baits that are placed remain in situ for 8 weeks continuously, unless stated differently on the label.

Required results
At least 4 times during the test and at the end of the test (about 8 weeks after the start), an estimate of the population size is obtained in the same manner as during the pre-test. The difference in population size before and 8 weeks after the control operation provides the degree of efficacy of the product.
Appendix 19. Current Antifouling Coatings

The current major types of antifouling coatings are outlined below, together with a brief description of their properties. This list is not exhaustive, and product applications may not fall within these categories. Applicants may submit novel coating types not covered by this list.

Table 49: Current Antifouling Coatings

<table>
<thead>
<tr>
<th>Coating Type</th>
<th>Description, mode of action and properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>Soluble matrix</td>
<td>In coatings of this type the active substance(s) has (have) been physically mixed ('freely associated') into a resin matrix. Upon exposure to seawater the slightly acidic matrix slowly dissolves releasing the active substance(s) into the water. (Seawater is slightly alkaline (pH 8) and the acidic matrix dissolves). Continuous dissolution of the coating surface will occur resulting in fresh actives being released until eventually the film is exhausted. Soluble matrix antifouling products typically show a biocide release rate curve which decays exponentially. The soluble matrix coatings have reduced mechanical properties that limit their film thickness. The paint film thickness of these coatings depletes over time in a fairly imprecise manner and the film does not show smoothing characteristics on ships in service. Such coatings are normally specified for lifetimes of typically 12-36 months.</td>
</tr>
<tr>
<td>Insoluble matrix</td>
<td>This type of coating contains a mixture of resins that together form an insoluble binder phase. One or more active substances are physically mixed into this matrix. As seawater enters the paint film, the biocides are released by dissolution and diffusion from within the insoluble matrix. After active substance have been released from the film, the binder remains intact and an empty ‘honeycomb’ structure (the leached layer) remains at the paint surface. This type of coating has a high initial release rate, which decreases exponentially with time as the active substance(s) have further distance to travel through the paint film. The rate of diffusion of biocide from within the film then becomes a limiting factor in maintaining an effective biocide release rate and hence preventing fouling. Insoluble matrix antifouling coatings do not show film-depletion or polishing as the resin is insoluble. The biocide release process continues until exhaustion of the coating. The higher mechanical strength obtained with these coatings allows for applications of thicker systems and coating lifetimes of typically 12-36 months are attainable.</td>
</tr>
<tr>
<td>Self polishing</td>
<td>This group is currently the most common and covers a range of different technologies that deliver the active substance through a gradual depletion/ablation of the paint film throughout the lifetime of the coating. These coatings use binder systems which control polishing behaviour by different mechanisms. A broad range of binder technologies are found in this group and these have replaced TBT copolymer based paints which have been withdrawn from use. Binder systems range from those based on the dissolution of metal carboxylates and polymers relying on ion-exchange to polymers relying on hydrolysis to control the rate of polishing. Modification of the binder systems and pigment phases of products within this</td>
</tr>
</tbody>
</table>
group can be used to tailor the products towards different end uses. The requirements for protection of a fast moving and very active vessel can be very different from that of a slow moving less active one. Such modifications can also be used to tailor performance to accommodate the potential intensity of fouling.

The different binder technologies can be used alone or in combination and result in products with varying levels of antifouling protection. Other binder components may also be added in order to modify the overall properties of the paint film. Typical dry-docking intervals for vessels coated with self polishing antifouling paints range from 24 to 60 months, however these systems may also be specified for lifetimes beyond this period.

<table>
<thead>
<tr>
<th>Coating Type</th>
<th>Description, mode of action and properties</th>
</tr>
</thead>
</table>

Typical dry-docking intervals for vessels coated with self polishing antifouling paints range from 24 to 60 months, however these systems may also be specified for lifetimes beyond this period.
Appendix 20. Published paper (CEPE Antifouling Working Group)

NOTE to the reader:
In the following CEPE methodology there are several issues that contradict with the requirements in the guidance document (e.g. number of trial panels, period of testing). The CEPE methodology can be used as long as the agreements of the guidance are respected.

TMI2013-PT21_efficacy_workshop-CEPE Efficacy Methodology for BPR - Revised 19 June 2012.doc
The European Council of producers and importers of paints, printing inks and artists’ colours - CEPE
Guidance developed by the CEPE Antifouling Working Group

Efficacy evaluation of antifouling products

Conduct and reporting of static raft tests for antifouling efficacy

Specific scope
This document provides a baseline methodology for evaluating and reporting the efficacy of antifouling coatings. Efficacy is assessed by static raft testing relative to a negative control and, if used, a positive control coating. Efficacy may be indicative of, but has no direct one-to-one relationship with the actual performance of a product under real life conditions.

Document version
First approved in 2011-04.
Revised in 2012-06

1. Scope

Overview: The purpose of this document is to provide a methodology for determining efficacy of antifouling coatings by panel testing on static floating rafts. The document provides guidance on how to conduct, assess, record, and report results from efficacy evaluations.

Efficacy is evaluated relative to a suitable inert, negative control. A positive control of proven antifouling performance may also be included. This static exposure methodology for natural environments is not suitable for establishing absolute performance characteristics of antifouling coatings in service.

Objective: This methodology may be used by industry to obtain efficacy data during the development of new antifouling coatings. This methodology may also be used to provide national registration authorities with the information required to support the label claim of antifouling products. Efficacy is demonstrated when the extent of fouling is visibly less than on a blank panel.

The methodology is especially useful for:
- the persons responsible for writing the protocols for antifouling efficacy trials
- the persons responsible for conducting trials including the evaluation and recording of results
the persons responsible for assembling and submitting dossiers for the registration of antifouling paints
• the national authorities which are responsible for the assessment of registration dossiers.

Reproducibility and accuracy: In static raft testing the fouling intensity will vary significantly between different geographical locations, between positions on the same rafts, and from season to season. More importantly, fouling will vary from one year to the next even for identical panels where exposure starts around the same date in different years. This variability in fouling intensity, and thus the test results, is due to weather conditions, availability of nutrients, and other uncontrollable factors that may affect the type and extent of fouling and its rate of settlement and growth. Therefore, the absolute amount of fouling present on the test coating and controls may not be reproducible at the same site from year to year.

Interpretation of results: The results obtained by this methodology demonstrate the ability of antifouling coatings to prevent settlement of fouling organisms under static conditions relative to a suitable negative control and, if used, a positive control tested simultaneously at the same site. An evaluation of the relative antifouling effect of an antifouling coating compared to the negative control and, if used, the positive control is used as a tool to indicate the potential of a tested coating to protect underwater structures. The results can be used to support appropriate label claims of the antifouling coating tested and to screen for new candidate products.

Efficacy testing on raft panels represents a worst case scenario compared to real life conditions. The main reason is that the exposure is static with limited opportunity for organisms to be removed by hydrodynamic forces. Ships' and boats' movement through water also aid the release of active ingredients from their antifouling. Furthermore, fouling intensity is generally recognised as being greater near the coast relative to the open seas.

2. Definitions

Antifouling coating: A material which, when applied as a surface coating, is used to control the settlement and/or growth of fouling organisms on submerged surfaces including ships, boats, aquaculture equipment, offshore oil installations, and other man made structures.

Negative control: An inert reference surface that does not control fouling, e.g. an anticorrosive coating.

Positive control: A reference surface coated with an antifouling coating of appropriate efficacy relevant to the intended end use of the test coating.

Fouling season: The months of the year during which significant settlement and growth of fouling organisms typically occur on a negative control at the test site.

3. Apparatus

The following equipment will be required to undertake efficacy testing according to this methodology.

Panels: Panels are typically made of plastic (e.g. PVC), reinforced polyester, steel, aluminium, marine grade plywood, or other material suitable for extended immersion in natural waters. (Metal panels must be adequately protected with an anticorrosive paint system.)

Panels should be designed to allow them to be securely fixed to the test raft, for example via a suitable panel rack. Where the design requires fixing holes through panels, these holes should be drilled prior to the application of the coating to prevent damage.
The panels may be designed to allow one or more coatings and/or controls to be tested on each individual panel. The total immersed area of each coating or control should be no less than 100 cm².

**Raft:** A free floating platform which has been designed to allow test panels to be affixed and immersed at a constant depth in natural waters. The design of the raft should enable panels to be readily removed for inspection.

The minimum depth of water below the raft at low tide should generally be 2.5 m.

The floating raft should be of sufficiently rigid construction to withstand prolonged exposure to weather and wave action and prevent excessive flexing or movement of test panels. It should be designed to ensure the occupational safety of users.

The raft should be designed to ensure that all test coatings and controls of the same test series are exposed to similar levels of sunlight and water flow to minimise variation. To increase the testing capacity, panels may be affixed to the raft in rows at the same depth. Where relevant the spacing between parallel rows at the same depth should generally be at least 20 cm to allow sufficient water circulation and illumination.

Generally, the raft design should ensure that panels are fully and permanently immersed. Panels should normally be exposed vertically and at a fixed depth from 0-3 m below the water surface. The lower edge of the panel should always be at least 0.5 m above the sea bed.

The raft may also be designed to allow coatings that are intended for use in darker or lighter areas to be tested under relevant conditions where the coating receives less or more sunlight. In such cases panels may be mounted on the raft facing partly down or up. Shade may also be provided by covering parts of the raft.

4. **Safety**

This test methodology does not address possible safety, health and environmental concerns associated with its use. All operations should be performed in accordance with all relevant local and national regulations.

**Personal protection:** Antifouling coatings may contain hazardous materials that could cause skin and eye irritation on contact and adverse physiological effects if inhaled. Thus, application and drying should take place in a well ventilated area and appropriate personal protective equipment should be worn during application. Product safety data sheets should be consulted when available.

**Environmental protection:** Unused paint and other contaminated material as well as panels after exposure should be disposed of as hazardous waste.

5. **Procedure**

All controls and test antifouling coatings should be tested under equivalent conditions. The exposure (immersion) of controls and test antifouling should start simultaneously (around the same date) and the exposure should be at the same location at the same depth and orientation.

**Panel preparation:** The test coating and positive control should be applied to panels according to the manufacturer’s guidelines to ensure adhesion during the period of the study. Appropriate drying and recoating intervals and temperature and ventilation requirements for application of the coatings should be followed.

An appropriate means of application should be used. Typical methods include spray, roller, brush, or specialised application equipment like a bar type applicator. Sufficient film thickness, taking the expected polishing and leaching rate characteristics of the product into account, should be applied to last for the planned duration of the test. Unless both sides of a panel are used as test substrates, the back of the panel may be
coated with an antifouling of proven efficacy to prevent fouling on the back. Edges may be painted with the coating under test or with a different coating of proven efficacy. All panels should be marked indelibly with a suitable reference code to aid identification.

**Replicates:** In cases where the purpose of the test is simply to demonstrate the efficacy of a test coating relative to a negative control, the use of single panels may provide data of sufficient quality. When replication is used, the number of replicates should be appropriate for the specific purpose of the test and should have the same orientation as the test panels and controls. Read-across to efficacy data from other test panels in a test series of similar formulations with the same content of active ingredients may also be used when justified and reasonable to support the results obtained for the test coating.

**Exposure time:** To verify efficacy, the minimum immersion time for testing is six months. In locations where the fouling season is shorter than six months this period may be reduced. The efficacy test should cover at least one continuous and complete fouling season where appropriate. Since raft panel exposure is static, fouling intensity is high, and the tests may be regarded as an accelerated test for products for vessels.

### 6. Evaluation

**Frequency:** Antifouling coatings under test and controls should be regularly inspected and evaluated for surface fouling, typically about every two months during the fouling season. Evaluations are not necessary during periods where there is minimal settlement and growth of fouling organisms (e.g. in cold and temperate regions where winter conditions do not support fouling settlement). Generally, the panels will be removed from the water for evaluation and, except at the end of the test period, returned to the water immediately after evaluation.

**Rinsing:** Optionally, panels may be rinsed gently with water from the site in order to reduce the influence of non-sessile organisms (that would be removed by low shear forces). Rinsing may also be carried out to remove possible sedimentary material (clay or silt). If utilised, rinsing must be performed on all panels equally and at each inspection. The method chosen, or if panels are not rinsed, must be specified in the final report.

**Evaluation procedure:** The type and severity of fouling that is present on the test coating and controls shall be assessed at each inspection. Evaluation may be made by visual assessment on site or any other appropriate method e.g. image analysis. The three major types of fouling observed on the test coating or controls; Slime, algae, and animals, should be separately assessed since the same percentage of coverage may have very different economical penalties during actual in-service use (e.g. effect on the friction of a vessel through water). Also fouling organisms that are known not to attach on moving vessels, but may be frequent on static surfaces, should be assessed separately (e.g. amphipods).

Further classification of the fouling organisms present may, in addition to slime (biological film of microfouling including bacteria, diatoms, micro-algae, and extracellular biopolymers), generally be restricted to main categories such as green, red, and brown macro-algae, bryozoa, hydrozoa, barnacles, tube worms, ascidians, and mussels. A more detailed determination is generally not necessary since products shall prevent attachment of fouling irrespective of species (or other taxonomic ranking).

As the assessment is based on a visual inspection, it is advised that this is done by a trained operator. This will help to improve consistency and data quality.

Assessment for the severity of fouling for each type of organism should be semi-quantitative, for example using a scale from 0-4, where 0 indicates the absence, and 4 indicates complete coverage of the class of organism in question. Optionally an estimation of the percentage coverage can be used.
The assessment of the coverage of algae and other soft fouling (e.g. arborescent bryozoans, and hydroids), should be based on the area covered by the "hold fast" (the attached base of the organisms) and not by the area covered by the "fronds" (leaves of macro-algae) or offshoot colonies.

**Overall fouling assessment:** The individual assessments of the fouling coverage of each type of organism may be combined to provide an overall fouling assessment. To generate this, a weighting of the coverage of the different types of fouling may be applied to rate and characterise the severity of the fouling present.

When the coating under test is intended for use on ships, fouling never seen on active vessels (e.g. amphipods) may be disregarded during the weighting. Biofouling attached to other fouling organisms (secondary fouling) should also be excluded from the overall fouling assessment.

Only the fully immersed surface area (if parts of the panel are subject to splash only) should be included in the determination of the fouling rating. Fouling attached within 1 cm from all edges of the test panel and fouling around the cable ties/studs/etc. may be disregarded in cases where an edge effect is seen. (Fouling around edges is normally attributed to insufficient antifouling paint film thickness around sharp panel edges.)

Fouling caused by physical defects or damages in the substrate or accidental damages of the antifouling should be disregarded. Fouling on exposed anticorrosive paints or other substrates (except where these are used as negative controls) or on other antifouling paints that may be used to coat panel edges, should be excluded from the assessment.

Physical defects (detachment, blistering, cracking, etc.) attributed to the inherent properties of the antifouling paint itself should be recorded and reported.

**Photos:** Inspection reports should include panel photos from each inspection.

### 7. Reporting

The report should contain all relevant information obtained from the efficacy trial for a given product. This may include:

- The name of the reporting company (and client if the test is carried out on assignment)
- The geographical location of the test raft(s) (including longitude and latitude)
- The geography (e.g. open sea, bay, estuary, etc.), depth of water, and water exchange conditions (tide, currents) at the raft site
- Typical local conditions. E.g. water temperature, salinity, and pH at the raft site
- Relevant information on the typical fouling community at the test site and seasonal influences where applicable.
- A discussion of any special conditions or variables that may have arisen particular to the specific test
- Orientation and exposure depth of test panels
- Dimensions and type (material) of test panels
- Identification of the tested product and control(s)
- Details on the panel preparation for the product under test and the control(s) (No. of coats, film thickness, application technique, etc.)
- Number of replicates if used
- Initial date of immersion and the cumulative exposure time (in months) for subsequent inspections
- Raw data from each individual assessment of a test panel
- The overall fouling assessment rating at each inspection during the exposure period
- Photos of test and control panels
- A systematic appraisal of the efficacy of the test product in relation to the negative control and, if used, any positive controls and the method by which that appraisal has been conducted
- A description of the reporting company’s weighting system used to provide the overall fouling assessment rating
- A discussion on the validity and acceptability of the test result relative to the intended label claim for the product tested when commercialised [e.g. recommended use area (recreational yachts, ships' niche areas, ships' flat bottoms, ships' water line, etc.) protection time/dry-docking interval, fouling conditions in targeted markets, etc.].

An interpretation of the test data generated and a conclusion on the efficacy of the coating under test.
Appendix 21. Example Of How An Overall Fouling Assessment May Be Carried Out For Panel Testing In Marine Waters

In order to assess panels out in the field, an effective and simple system is needed. Very detailed assessments of fouling coverage do not increase the quality of the test, as field conditions are highly variable and static raft tests can only provide an indication of products’ real life performance.

Individual companies have different ways of assessing the coverage of the main categories of fouling into an overall description of the efficacy of test panels. However, the principles of the example should apply to most assessment systems. Transparency of how the overall assessment is carried out is important in order to evaluate an efficacy report.

The fouling coverage on raft panels will be assessed based on coverage intervals. Each interval will be recorded by a different 'rating'.

Table 50: Example of categorisation of fouling coverage into ratings from 0 to 4

<table>
<thead>
<tr>
<th>Fouling Coverage (examples of company specific intervals for coverage of fouling)</th>
<th>Rating</th>
</tr>
</thead>
<tbody>
<tr>
<td>Company 1</td>
<td>Company 2</td>
</tr>
<tr>
<td>0-10%</td>
<td>0%</td>
</tr>
<tr>
<td>10-30%</td>
<td>&gt;0-25%</td>
</tr>
<tr>
<td>30-50%</td>
<td>25-50%</td>
</tr>
<tr>
<td>50-80%</td>
<td>50-75%</td>
</tr>
<tr>
<td>80-100%</td>
<td>75-100%</td>
</tr>
</tbody>
</table>

As different fouling species can contribute to different impacts on a vessel (e.g. fuel consumption of a ship), the coverage ratings may be weighted in several ways to take this into account. The applicant may provide references to literature that provide more detail on the assessment and weighting factors.

Table 51: Example of weighting of ratings

<table>
<thead>
<tr>
<th>Type of fouling</th>
<th>Weighting (of ratings from 1-4)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Trace (1)</td>
</tr>
<tr>
<td>Light slime</td>
<td>0</td>
</tr>
<tr>
<td>Dense slime</td>
<td>3</td>
</tr>
<tr>
<td>Macro-algae</td>
<td>5</td>
</tr>
<tr>
<td>Animals</td>
<td>5</td>
</tr>
</tbody>
</table>

---

50 e.g. IMO MEPC/60/4/21, 2010 from IPPIC
A score may be calculated by adding up the weightings. In this example, that value is then subtracted from 100. Zero growth (apart from traces of light slime) gives the fouling resistance rating 100 (100-0) and heavy fouling of both algae and animals gives the rating 0 [100-(50+50)]. The rating is then allocated to descriptions of the overall efficacy.

**Table 52: Example of categorisation of overall efficacy**

<table>
<thead>
<tr>
<th>Fouling resistance rating</th>
<th>Efficacy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Company specific score intervals, each with a corresponding characterisation of the efficacy</td>
<td>Excellent</td>
</tr>
<tr>
<td></td>
<td>Good</td>
</tr>
<tr>
<td></td>
<td>Fair</td>
</tr>
<tr>
<td></td>
<td>Poor</td>
</tr>
</tbody>
</table>

Description of types of fouling:

**Slime:** Bacteria, micro-algae, and protozoa.

- **Light slime** is easily removed from the surface.
- **Dense slime** is not easily removed from the surface.

**Algae (weed):** Green algae, red algae, and brown algae.

**Animals:** Barnacles, tubeworms, mussels, hydroids, and bryozoans.

RELATING COMPANY FOULING ASSESSMENTS TO THE NORMS AND CRITERIA FOR PRODUCT AUTHORIZATION.

When applying for authorisation of an antifouling product, the applicant should provide their overall fouling assessment of the product, together with the raw data and photographs/diagrams of the panel tests.

This guidance document only takes into account the percentage of macro-fouling on the raft panels as pass/fail criterion, not the classification in the applicant’s assessment system.

As the percentage coverage per rating may differ between different company’s assessment systems (see Table 1), some systems might not record 25 % coverage (the pass/fail criterion) in their rating system (e.g. in Table 1 Company 1 has a borderline at 30 % not at 25 %). Therefore, not only the ratings and end category of the product should be provided but also the raw data of the panel tests. The percentage coverage with macro-fouling per panel can then be identified from the raw data. This percentage is used to see if the product is sufficiently effective (i.e. <25 % macro-fouling)
## Appendix 22. PT 22 active substances in the review programme

### Table 53: PT 22 active substances in the review programme

<table>
<thead>
<tr>
<th>Active Substance</th>
<th>RMS</th>
<th>CAS No</th>
</tr>
</thead>
<tbody>
<tr>
<td>Formaldehyde</td>
<td>DE</td>
<td>50-00-0</td>
</tr>
<tr>
<td>Bronopol</td>
<td>ES</td>
<td>52-51-7</td>
</tr>
<tr>
<td>Iodine</td>
<td>SE</td>
<td>7553-56-2</td>
</tr>
<tr>
<td>Quaternary ammonium compounds, benzyl-C12-18-alkyldimethyl, chlorides</td>
<td>IT</td>
<td>68391-01-5</td>
</tr>
<tr>
<td>Quaternary ammonium compounds, benzyl-C 12-16-alkyldimethyl, chlorides (ADBAC)</td>
<td>IT</td>
<td>68424-85-1</td>
</tr>
<tr>
<td>Quaternary ammonium compounds, benzyl-C12-14-alkyldimethyl, chlorides</td>
<td>IT</td>
<td>85409-22-9</td>
</tr>
<tr>
<td>Quaternary ammonium compounds, C12-14-alkyl[(ethylphenyl)methyl]dimethyl, chlorides</td>
<td>IT</td>
<td>85409-23-0</td>
</tr>
<tr>
<td>Polyvinylpyrrolidone iodine</td>
<td>SE</td>
<td>25655-41-8</td>
</tr>
</tbody>
</table>
Appendix 23. Assessment grid for tests on human bodies

This grid is for use in the assessment of the biocidal product itself, but not for assessing the overall embalming process with its hygiene and cosmetic aspects.

Number of the report:
Name and signature of the embalming professional:
Company:
Address of company:

1. General information

Date of the operation:
Place:
Type of place:
- Funeral parlour
- Morgue
- Establishment without a morgue (fewer than 200 deaths per year)
- Home or other (please specify):

Identification:

Gender: □ Male □ Female
Age:
Estimated weight (kg):
Estimated corpulence: □ cachectic □ thin □ medium □ stout
Adiposity: □ low □ medium □ high
Date of death (if known):
Date and time of treatment:
Body refrigerated: □ yes □ no. If "yes", for how long:
Temperature:
Causes of death (if known):
Therapeutic treatment (if known):

2. Preoperative examination of the body

Body intact: □ yes □ no, description:
Autopsy before treatment: □ yes □ no
Presence of external prostheses: □ yes □ no
Surgical intervention before death (if apparent or known): □ yes □ no
If yes, type of intervention:

Other visible anomaly(ies):

Decomposition: □ none □ commencing □ problematic
Rigidity: □ none □ minimal □ moderate □ problematic
Dehydration: □ none □ normal □ high
Lividity: □ none □ minimal □ moderate □ problematic, location:
Coloration of tissues (yellowing): □ no □ slight □ moderate □ intense, description:
Dermal lesions (sores, blisters, wounds, etc.): □ yes □ no, description:
Distension of the abdomen: □ no, □ slight □ moderate □ intense, □ liquid □ gas
Bruising: □ yes □ no, □ abdomen □ thorax □ leg, □ arm, □ face, specify degree and place:
Comments:

3. **Techniques used for injection of the biocidal product**

Time of start of treatment:
Time of end of treatment:

**Site(s) of injection:**
Carotid(s): □ right □ left.
Femoral(s): □ right □ left
Axillary(ies): □ right □ left
Other(s), description:
Ease of finding: □ easy □ normal □ deep
Condition: □ good □ atheromatous / □ hardened
Injection: □ manual □ by electric pump □ by gravity
Diffusion: □ good □ fair □ bad
Puncture before treatment: □ yes □ no. If "yes", type:

**Biocidal product used:**
Pre-injection: □ yes □ no

**Injection:**

Hypodermic:  

*product:*
Site:  

Topical:  

Name of the biocidal product:  
Active substance(s):  
Duration of efficacy claimed:  
Number of litres:  

Arterial fluid:  
Name of fluid:  
  % of dilution:  
Number of litres injected:  
Start time for the injection:  
End time for the injection:  

Cavity treatment:  
Name of fluid:  
  % of dilution:  
Number of litres injected:  
Start time for the injection:  
End time for the injection:  

Corrective injection:  
□ yes □ no  

Drainage method:  
□ Cardiac, □ Venous  
Vein(s) chosen: □ jugular, □ femoral, □ axillary  
Volume drained by circulatory system (litres):  
Total volume drained (litres):  

Type of drainage: □ drain tube(s) □ forceps □ intermittent / □ continual  
Quality of drainage: □ considerable clotting □ medium □ slight □ no clotting  

General puncture:  
Quantity:  

...........................................................................................................................................
4. **Observations concerning the injection of the biocidal product**

Observations during the treatment:

Odour: □ normal □ fair □ bad  
Colouring: □ good □ fair □ bad  
Suppleness of the skin: □ good □ fair □ bad

Observations following the treatment:

Odour: □ good □ fair □ bad  
Colouring: □ good □ fair □ bad  
Suppleness of the skin: □ good □ fair □ bad

Mandatory observation 48 hours after the treatment:

Odour: □ good □ fair □ bad  
Colouring: □ good □ fair □ bad  
Suppleness of the skin: □ good □ fair □ bad

Optional observation (at times relevant to the manufacturer's claims):

Time after treatment:

Odour: □ good □ fair □ bad  
Colouring: □ good □ fair □ bad  
Suppleness of the skin: □ good □ fair □ bad

Other products used during the preservation process:

Reasons for their use:

Description:

**Products for cosmetic purposes:**

□ yes, □ no. If yes: □ normal, □ make-up, □ significant, □ restorative

**Other restoration:** ___________________________

Moisteners and other products used (cauterising agents, disinfectants, skin tone correctors, etc.):

Name of the fluid: __________________________, % dilution: _____, litres injected: ___

Name of the fluid: __________________________, % dilution: _____, litres injected: ___

Explanations:

5. **Comments**