Consultation procedure for guidance

1. Purpose

This procedure document describes the general structure of the consultation procedure for Guidance documents that are part of the process “Provision of new guidance and guidance update” (PRO-0012).

The procedure includes the organisation and management of the consultation with ECHA partners intended to ensure a broad acceptance of the guidance.

This procedure applies when a new guidance or an update to a guidance document needs to be developed.

2. Scope

The procedure begins when the Project Team has finalised and agreed on the draft of a new or updated guidance document and when the Guidance roadmap (when foreseen; TEM-0005) has been approved (PRO-0012) by the Head of Unit (HoU) responsible for the process. The procedure ends when the final guidance document (new or updated) is ready for publication on the ECHA website.

3. Description

The consultation process is organised and co-ordinated by a Project Manager (PM), who is nominated by the HoU of the Unit responsible for the process. The PM is supported by a Project Team. The Project Team is described in the Guidance Roadmap (when foreseen) and consists of the PM, a representative of the Legal Affairs Unit (LAU) and other relevant ECHA staff contributing to the development of the new guidance or guidance update.

The PM launches the Consultation Procedure according to the project description, resources and timelines and as described in the approved Guidance roadmap. It is launched under the responsibility of the PM and subject to approval by the HoU responsible for the process.

The PM ensures that the draft documents sent for consultation at the different stages are published on the ECHA website in the section dedicated to the ECHA Consultation Procedure on guidance, following the process “Handling requests for the ECHA website” (WIN-0034) in order to keep the process open and transparent. This also allows stakeholders not directly involved, such as third countries and other interested parties, to follow the progress of work closely and to comment using the standard form on the website; answers to individual comments made via this standard form (i.e. outside the formal consultation procedure described in the rest of this document) are not provided.
The HoU decides whether:

- to skip the consultation of a certain body for justified reasons;
- to choose consulting the bodies in parallel or consecutively.

The necessary steps of the Guidance Consultation Procedure will be selected, depending on the needs for the provision of new guidance or the guidance update. The rationale behind the choices will be documented in the Guidance roadmap. The relevant Competent Authorities and the European Commission always need to be consulted.

If the HoU decides not to consult the PEG (Partner Expert Group) and/or the Committees/Forum, he/she needs to ensure that all of the following criteria are met:

- The process for the provision of new guidance or the guidance update is transparent e.g. via a new alert or an information note on the ECHA website in the section dedicated to the ECHA Consultation Procedure on guidance.
- The aim of achieving a common understanding among stakeholders of the subject matter and striving for consensus on the provision of new guidance or the guidance update is not jeopardised.
- Stakeholders that have an interest/role with regard to the tasks that are addressed by the new guidance or the guidance update - are consulted either via:
  - Representation in another body that will be consulted (e.g. skipping the consultation of the PEG by ensuring that they are sufficiently represented at the later stages); or
  - Consultation of an equivalent body that represents relevant stakeholders (e.g. consultation of one of ECHA’s expert groups fulfilling the role of the PEG).

When deciding on whether to consult the different bodies in parallel or consecutively, elements such as whether it concerns a completely new guidance or whether the guidance addresses a complicated topic that has not been addressed before need to be considered as this may impact on the efficiency of the process. However, this is a case-by-case decision. It is usual to follow the steps in sequence especially if there is consultation with the PEG and the Committees/Forum.

**Step 1: Consultation of the Partner Expert Group (PEG)**

In case of an update for which it has been decided that a PEG consultation is not needed, this step is not performed. In such cases the procedure either starts at step 2 or (in case also consultation of ECHA committees/Forum is not foreseen) at step 3.

Annex 1 provides some elements that need to be considered when deciding on the involvement in the consultation of the PEG. According to what has been approved in the Guidance roadmap, the consultation of the PEG can proceed.

The PM invites Accredited Stakeholder Organisations (ASOs) and institutional interested parties to nominate experts and, if relevant, invites individual experts to participate in the PEG. Experts whose nomination or confirmation of individual willingness to participate is received by the specified deadline (circa 3 weeks from the invitation) are included in the consultation of the PEG.
The PEG and its respective mandate are established by sending the confirmation of nomination to each expert by email. The PM, at the same time, informs the PEG members about the Consultation procedure, the timeline and the indicative date of the PEG meeting (if foreseen). The main task of the PEG is to comment and strive for consensus on the draft proposed by the Project Team, with a view to ensure that it should be acceptable to all interested parties. The outcome of the consultation of the PEG serves as the basis for next draft version of the guidance text for further consultation steps.

When the draft guidance update or new guidance is internally agreed, the PM sends it to the members of the PEG together with a cover note and a blank template for comments. The PEG is invited to submit comments on the guidance document within a specified timeframe of a minimum of four weeks.

The PM with the support of the Communications Unit, ensures that the draft guidance document (that has been sent to the PEG) and the composition of the PEG, are published on the ECHA website (in the section dedicated to the ECHA Consultation Procedure on guidance).

After the deadline for comments has expired, the PM, with the support of the Project Team analyses the comments received. The PM drafts the agenda for the PEG meeting from this analysis.

In exceptional cases when a physical PEG meeting is not planned, the following activities are not performed and the process continues from outcome A or B (below).

The PEG meeting can be a physical meeting taking usually place at ECHA or can be in the format of a WebEx. If it is not a meeting at ECHA and for any unforeseen situation arises or for logistic considerations, the venue of the meeting may be changed or the event may be organised via a remote connection. The meeting is chaired by a representative of the ECHA Secretariat and the ECHA Secretariat (or an appointed ECHA contractor) is/are also responsible for taking the minutes.

The draft minutes are prepared and sent to the PEG members after the meeting. However the revisions to the draft guidance document take priority in order to comply with the timetable for the next consultation step. PEG members are given an appropriate time to provide comments (minimum 5 working days) and clarifications before finalising the minutes.

From the meeting, one of two outcomes may occur:

**Outcome A: Consensus achieved**

The PM, in collaboration with the Project Team, assesses the comments received from the consultation round, including the discussions at the PEG meeting and decides on their relevance and implementation. The PM in collaboration with the Project Team, where necessary, prepares a consolidated text, seeking further advice from the PEG members if necessary. A brief response to all the comments is provided to the PEG members through a summary spreadsheet either indicating that the comment was accepted or specifying the reasons why comments are not implemented or are implemented in part through alternative changes. The PM also sends the revised draft guidance document and consolidated comments with responses to the PEG members to verify the correct understanding of the input received, giving the PEG members approximately 10 working days to indicate if their
own comments have been misunderstood. No new (i.e not previously raised) comments from PEG members are expected to be submitted to ECHA at this point. The draft guidance document can be further modified by the PM, taking into consideration written communications received from PEG members that one or more comments have been misinterpreted or overlooked. In this case, the consolidated comments with responses as well as the draft text, if appropriate, are revised accordingly by the PM.

The final version of the consolidated draft guidance document and summary of the comments received from the PEG with the Project Team’s reply are reviewed by the LAU representative. It is good practice to consult the LAU representative during the implementation of the comments, for efficiency reasons so that this step is only a formality.

**Outcome B: Consensus is not achieved**

If consensus on one or more main issues is not achieved during the meeting, the majority position of the PEG is taken. If there is no clear majority or in exceptional cases, such as a large number of difficult, contentious or unforeseen issues, the PM in consultation with the responsible HoU may decide that a second PEG consultation round and possibly a (second) PEG meeting needs to be organised. This is a case-by-case decision. The PM (in consultation with the HoU) establishes new timelines for this second round and informs the PEG via email that a second consultation is foreseen and of the timelines for it.

**Step 2: Consultation of the experts from the ECHA Committees and/or the Forum**

In case of an update for which a consultation of the ECHA Committees and/or Forum is not foreseen, this step is not applied. In this case the procedure continues or (in case step 1 did not apply) starts at step 3.

Annex 2 provides some elements that need to be considered when deciding on the involvement in the consultation of the ECHA Committees and/or the Forum. The PM launches the consultation with the relevant bodies. The consultation takes place via a written procedure with the PM sending the consolidated draft guidance document together with a blank template for comments and the consolidated summary of PEG comments directly to the members of the Committees, and/or the Forum through the Forum Secretariat with a request to provide comments within a deadline of approximately four weeks.

The PM, with the support of the Communications Unit, ensures that the consolidated draft guidance document sent to the Committees/Forum and the consolidated summary of PEG comments with ECHA responses to PEG comments, are published on the ECHA website (in the section dedicated to the ECHA Consultation Procedure on guidance).

The PM, in collaboration with the Project Team, assesses the comments received from the consultation round and decides on their relevance and implementation. The PM then prepares a consolidated text, seeking advice from the Project Team whenever necessary.

A summary of the responses to all the comments is provided to the Committees and/or the Forum members by way of the consolidated document indicating either that the comments were implemented or the reasons why ECHA disagrees with a proposed change where comments are not implemented. The PM sends the revised guidance document and consolidated comments with responses to the Committee members and/or the Forum members (through the Forum Secretariat) to inform about the implementation of the input
Step 3: Consultation of the European Commission and the Competent Authorities / Designated National Authorities

The consultation with the European Commission and the Competent Authorities (CAs) / Designated National Authorities (DNAs) constitutes the final step of every consultation procedure. It ensures that any new guidance, or updated guidance, will find as wide support as possible with harmonised implementation by all the authorities.

The PM launches the consultation via a written procedure by submitting a request to the Commission services to upload the final version of the consolidated draft guidance, a cover invitation letter, and a consolidated summary of PEG and Committee/Forum comments (if applicable) with the Project Team’s replies to those comments, in CIRCABC or equivalent platform. The consultation is carried out according to the “silence gives consent” principle: parties that do not provide any comments by the deadline (approximately 4 weeks) will be deemed to have agreed with the proposed draft text.

The PM, with the support of the Communications Unit, ensures that the final version of the consolidated draft guidance is sent to the CAs/DNAs and that the consolidated summary of the Committee/Forum comments with ECHA responses to the comments, are published on the ECHA website (in the section dedicated to the ECHA Consultation Procedure on guidance).

From this consultation, one of two situations may occur:

Outcome A: The written consultation is conclusive

The PM, in collaboration with the Project Team, implements relevant comments and prepares a final version of the draft guidance document based on the outcome of the consultation.

A summary of the ECHA responses to all the comments is provided to the CAs/DNAs either indicating that they have been implemented or specifying the reasons why the proposed changes have not been made. The PM sends the request to Commission services to upload in CIRCABC or otherwise communicates via an agreed mechanism the final version of the guidance document as well as the consolidated comments received from the CAs/DNAs, giving them approximately 10 working days to check that their own comments have not been misinterpreted. No new (i.e. not previously raised) comments from CAs/DNAs are expected to be submitted to ECHA at this point.
The PM may further modify the draft guidance document to take into consideration written communications from CAs/DNAs or the Commission informing that one or more comments have been misinterpreted or overlooked. The consolidated comments with responses are aligned with the modified draft by the PM before publication and upload of the comments to the website.

**Outcome B: A consensus cannot be achieved through a written procedure**

The controversial issues are discussed at the first possible meeting of the CAs/DNAs. The PM coordinates the process, requesting to add a discussion point on those issues to the agenda of the meeting.

**Option 1:** If the consultation at the meeting is conclusive, the PM implements relevant comments and prepares a final version of the draft guidance document based on the outcome of the consultation.

**Option 2:** If a consensus at the meeting cannot be achieved, the majority and minority opinions and their justifications are recorded in the meeting minutes, which are made public. The PM in collaboration with the Project Team prepares a note to be signed by the ED to make the reader of the guidance specifically aware of a lack of consensus and provide a cross-reference to the meeting minutes. This note is made as the new first page of the PDF-file containing the guidance.

The PM prepares a final version of the guidance document for publication on the ECHA website. In the case of option 2, the ED (on a case by case decision) has informed the Management Board (MB).

A summary of the ECHA responses to all the comments is provided to the CAs/DNAs either indicating that they have been implemented or specifying the reasons why the proposed changes have not been made. The PM sends the request to Commission services to upload in CIRCABC or otherwise communicates via an agreed mechanism the final version of the guidance document as well as the consolidated comments received from the CAs/DNAs, with the aim to inform them about the implementation of the input received before and during the meeting.

**Step 4: Finalisation of the consultation procedure**

After review of the content by LAU and approval by the ECHA management, the final version of the guidance document and the consolidated comments received from the CAs/DNAs with ECHA responses to them are published on the ECHA website (this corresponds to step 3.6 of PRO-0012).
4. Flowchart

Consultation procedure for Guidance

Draft update guidance or new guidance internally agreed (Step 3.1 to 3.4 of PRO-0012)

New guidance or normal update

Preparation of second consultation round

PEG meeting

Number of (controversial) issues/absence of clear majority requires additional round?

Consolidation of the draft

PEG written consultation

Yes

No

Consentation of the CAs (written consultation)

Discussion at meeting of the CAs

Consensus achieved?

Majority and minority opinions recorded and ED informs the MB

Consolidation of the draft

Number of (controversial) issues/absence of clear majority requires additional round?

PEG consultation

Consentation of the Committees and/or Forum (written consultation)

Yes

No

Consolidation of the draft

PEG consultation necessary?

Committee/Forum consultation necessary?

Consultation of the CAs (written consultation)

Yes

No

PEG consultation necessary?

See Annex 2

See Annex 2

Committee/Forum consultation necessary?

Consultation of the CAs (written consultation)

Yes

No

PEG consultation necessary?

Consultation of the CAs (written consultation)

Yes

No

Finalisation of the consultation procedure

Publication process (Step 3.6 of PRO-0012)

Committee/Forum consultation necessary?

Consentation of the CAs (written consultation)

Yes

No

PEG consultation necessary?

Consentation of the CAs (written consultation)

Yes

No

PEG consultation necessary?
### 5. Definitions

<table>
<thead>
<tr>
<th>Term or abbreviation</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>ASO</td>
<td>Accredited Stakeholder Organisation, organisations with a legitimate interest in the work of ECHA who applied for the Accredited Stakeholder role and are selected based on eligibility criteria adopted by the Management Board</td>
</tr>
<tr>
<td>CA</td>
<td>Competent Authority</td>
</tr>
<tr>
<td>CIRCABC</td>
<td>Communication and Information Resource Centre for Administrations, Businesses and Citizens</td>
</tr>
<tr>
<td>DNA</td>
<td>Designated National Authorities</td>
</tr>
<tr>
<td>Guidance document</td>
<td>Formal documents of a highly technical nature that require interpretation of the underlying regulation(s) and a common understanding on how to fulfil the obligation that they place on industry and authorities. Therefore, these documents when developed or updated will be the subject of consultation as described in this PRO.</td>
</tr>
<tr>
<td>ECHA partners</td>
<td>Combination of both “institutional interested parties” and “stakeholders”.</td>
</tr>
<tr>
<td>ED</td>
<td>Executive Director</td>
</tr>
<tr>
<td>Institutional interested parties</td>
<td>Member State Competent Authorities, the European Commission as well as third country representatives</td>
</tr>
<tr>
<td>Forum</td>
<td>Forum for Exchange of Information on Enforcement</td>
</tr>
<tr>
<td>HoU</td>
<td>Head of Unit</td>
</tr>
<tr>
<td>LAU</td>
<td>Legal Affairs Unit</td>
</tr>
<tr>
<td>MB</td>
<td>Management Board</td>
</tr>
<tr>
<td>Normal update</td>
<td>See PRO-0012 “Provision of new guidance and guidance update”</td>
</tr>
<tr>
<td>PM</td>
<td>Project manager</td>
</tr>
<tr>
<td>PEG</td>
<td>Partner Expert Group</td>
</tr>
<tr>
<td>PRO</td>
<td>Procedure</td>
</tr>
<tr>
<td>Stakeholders</td>
<td>Non-institutional interested partners (industry, trade unions, environmental and consumer NGOs, academia, etc.)</td>
</tr>
</tbody>
</table>
# 6. Records

<table>
<thead>
<tr>
<th>Record name</th>
<th>Security level</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Invitation to nominate experts to the PEG</td>
<td>Public</td>
<td></td>
</tr>
<tr>
<td>Expert nomination</td>
<td>Internal</td>
<td></td>
</tr>
<tr>
<td>PEG establishment</td>
<td>Internal</td>
<td></td>
</tr>
<tr>
<td>Final draft update/new guidance internally agreed and for PEG written procedure</td>
<td>Public</td>
<td></td>
</tr>
<tr>
<td>Comments on draft guidance document</td>
<td>Internal</td>
<td></td>
</tr>
<tr>
<td>PEG meeting minutes</td>
<td>Public</td>
<td></td>
</tr>
<tr>
<td>Summary spreadsheet with PEG comments with ECHA's responses</td>
<td>Public</td>
<td></td>
</tr>
<tr>
<td>Information about a second PEG consultation round and a possible PEG meeting when consensus is not achieved</td>
<td>Internal</td>
<td></td>
</tr>
<tr>
<td>Draft update/new guidance to Committees/Forum for written procedure</td>
<td>Public</td>
<td></td>
</tr>
<tr>
<td>Committees/ Forum comments</td>
<td>Internal</td>
<td></td>
</tr>
<tr>
<td>Consolidated draft update/new guidance after Committees/Forum consultation</td>
<td>Public</td>
<td></td>
</tr>
<tr>
<td>Summary spreadsheet with Committees/Forum comments with ECHA's responses</td>
<td>Public</td>
<td></td>
</tr>
<tr>
<td>Draft update/new guidance to CAs/DNAs for written procedure</td>
<td>Public</td>
<td></td>
</tr>
<tr>
<td>CAs'/DNA’s comments</td>
<td>Internal</td>
<td></td>
</tr>
<tr>
<td>Note to the reader by ED</td>
<td>Public</td>
<td></td>
</tr>
<tr>
<td>Consolidated final update/new guidance after CAs/DNAs consultation</td>
<td>Public</td>
<td></td>
</tr>
<tr>
<td>Summary spreadsheet with CAs'/DNAs’ comments with ECHA’s replies</td>
<td>Public</td>
<td></td>
</tr>
</tbody>
</table>
7. References

<table>
<thead>
<tr>
<th>Associated document code</th>
<th>Document name</th>
</tr>
</thead>
<tbody>
<tr>
<td>MB/34/2011</td>
<td>Revised eligibility criteria for ECHA’s Accredited Stakeholders</td>
</tr>
<tr>
<td></td>
<td>List of ECHA’s Accredited Stakeholder Organisations: [ECHA’s webpage]</td>
</tr>
</tbody>
</table>

8. Annexes

**Annex 1**: Criteria indicating when it is recommended to consult the PEG.

- Provision of a new guidance concerning technical issues.
- The update addresses a completely new and complicated topic that has not been addressed before.
- The update introduces new or modifies existing obligations whose implementation requires stakeholders consultation to ensure feasibility.
- The scientific and technical expertise needed for providing guidance can only be accessed via a PEG as it is not represented in the Committees and/or the Forum, nor in the final consultation of the European Commission and the Competent Authorities/Designated National Authorities.
- The stakeholder organisations that are impacted by the provision of new guidance or by the guidance update can only be accessed via the establishment of a PEG.

**Annex 2**: Criteria indicating when it is recommended to consult the ECHA Committees and/or the Forum.

- The update requires very specific expertise only accessible via the Committees and/or via the Forum (e.g. EOGRTS or the enforceability of a provision).
- The provision of new guidance or the guidance update relates to the processes of the Committees and/or the Forum.