

Helsinki, 19 September 2018



Decision number: CCH-D-2114440632-56-01/F

Substance name: 1-Propanaminium, 3-amino-N-(carboxymethyl)-N,N-dimethyl-, N-(C8-

18(even numbered) and C18 unsaturated acyl) derivs., hydroxides, inner salts

EC number: 931-333-8 CAS number: NS

Registration number: Submission number:

Submission date: 28/9/2017

Registered tonnage band: Over 1000

DECISION ON A COMPLIANCE CHECK

Based on Article 41 of Regulation (EC) No 1907/2006 (the REACH Regulation), ECHA requests you to submit information on:

- 1. Pre-natal developmental toxicity study (Annex X, Section 8.7.2.; test method: EU B.31./OECD TG 414) in a second species rabbit, oral route with the analogue substance (carboxymethyl)dimethyl-3-[(1-oxododecyl)amino]propylammonium hydroxide (C12 AAPB, CAS no 4292-10-8, EC no 224-292-6);
- 2. Extended one-generation reproductive toxicity study (Annex X, Section 8.7.3.; test method: EU B.56./OECD TG 443) in rats, oral route with the analogue substance (carboxymethyl)dimethyl-3-[(1-oxododecyl)amino]propylammonium hydroxide (C12 AAPB, CAS no 4292-10-8, EC no 224-292-6) specified as follows:
 - Ten weeks premating exposure duration for the parental (P0) generation;
 - Dose level setting shall aim to induce some toxicity at the highest dose level;
 - Cohort 1A (Reproductive toxicity);
 - Cohort 1B (Reproductive toxicity) without extension to mate the Cohort 1B animals to produce the F2 generation;

You may adapt the testing requested above according to the specific rules outlined in Annexes VI to X and/or according to the general rules contained in Annex XI to the REACH Regulation. To ensure compliance with the respective information requirement, any such adaptation will need to have a scientific justification, referring and conforming to the appropriate rules in the respective annex, and adequate and reliable documentation.

You have to submit the requested information in an updated registration dossier by **26 March 2021**. You also have to update the chemical safety report, where relevant. The timeline has been set to allow for sequential testing.

CONFIDENTIAL 2 (17)



The reasons of this decision are set out in Appendix 1. The procedural history is described in Appendix 2 and advice and further observations are provided in Appendix 3.

Appeal

This decision can be appealed to the Board of Appeal of ECHA within three months of its notification. An appeal, together with the grounds thereof, has to be submitted to ECHA in writing. An appeal has suspensive effect and is subject to a fee. Further details are described under: http://echa.europa.eu/regulations/appeals.

Authorised¹ by Ofelia Bercaru, Head of Unit, Evaluation E3

 $^{^1}$ As this is an electronic document, it is not physically signed. This communication has been approved according to ECHA's internal decision-approval process.



Appendix 1: Reasons

0. Grouping and read-across approach for toxicological information

Article 13(1) of the REACH Regulation provides that information on intrinsic properties of substances may be generated by means other than tests. Such other means include the use of information from structurally related substances (grouping of substances and readacross), "provided that the conditions set out in Annex XI are met".

In the registration dossier, you have adapted the standard information requirements by applying a read-across adaptation following REACH Annex XI, Section 1.5. for

- In vitro gene mutation in mammalian cells (Annex VIII, Section 8.4.3)
- Pre-natal developmental toxicity (Annex IX, 8.7.2)

Annex XI, Section 1.5. requires a structural similarity among the substances within a group or category such that relevant properties of a substance within the group can be predicted from the data on reference substance(s) within the group by interpolation. The following analysis presents your justification for the proposed grouping approach and read-across hypothesis, together with ECHA's analysis concerning the justification in both a generic and a endpoint-specific context.

A. Description of the grouping and read-across approach proposed by you

You have provided a read-across justification document entitled "

The AAPBs considered within this read-across approach include the following substances registered under REACH:

- C12 AAPB (Reference Substance Name: (carboxymethyl)dimethyl-3-[(1-oxododecyl)amino]propylammonium hydroxide), CAS number: 4292-10-8, EC number: 224-292-6
- 2. **C12-18 AAPB** (Reference Substance Name: 1-Propanaminium, 3-amino-N-(carboxymethyl)-N,N-dimethyl-, N-(C12-18(even numbered) acyl) derivs., hydroxides, inner salts), CAS number: -, EC number: 931-513-6
- 3. **C8-18 AAPB** (Reference Substance Name:1-Propanaminium, 3-amino-N-(carboxymethyl)-N,N-dimethyl-, N-C8-18(even numbered) acyl derivs., hydroxides, inner salts), CAS number: 97862-59-4, EC number: 931-296-8
- 4. **C8-18 and C18 unsatd. AAPB**, (1-Propanaminium, 3-amino-N-(carboxymethyl)-N,N-dimethyl-, N-(C8-18(even numbered) and C18 unsaturated acyl) derivs., hydroxides, inner salts), CAS number:-, EC number: 931-333-8

In your read-across justification, you also include the following substance:

5. **C12-14 AAPB**, (Reference Substance Name: 1-Propanaminium, 3-amino-N-(carboxymethyl)- N,N-dimethyl-, NC12-14 acyl derivs., hydroxides, inner salts), EC: not available

ECHA notes that the latter substance is characterised by its name only, and the read-across justification document contains no other identifiers such as EC or CAS numbers that would

CONFIDENTIAL 4 (17)



allow ECHA to verify its identity and hence its suitability for the read-across. In addition, there are no experimental data available with this substance regarding its physico-chemical, environmental and toxicological properties, neither in the read-across justification document nor attached to the technical dossiers of the other 4 substances. As a consequence, since there are no source data available with this substance, ECHA does not consider it as a source or target substance for the purpose of this read-across. In conclusion, ECHA has assessed the read-across only for the first 4 substances listed above.

You have provided a hypothesis for grouping alkylbetaines on the basis of structural similarity and the presence of same functional groups.

You have provided the following hypothesis: "the substances under evaluation have similar physicochemical, toxicological and ecotoxicological properties because they share structural similarities with common functional groups: quaternary amines, amide bonds, carboxymethyl groups, and fatty acid chains, differing in length and degree of saturation. This prediction is supported by physicochemical, toxicological and ecotoxicological data on the substances themselves."

You have explained structural differences in relation to toxicological properties that could be attributed to:

- 1. Differences in the fatty acid moiety that would relate to the degree of saturation and/or alkyl chain length. In particular you indicated that the AAPBs differ by their carbon chain length distribution and the degree of unsaturation in the fatty acid moiety. However, is the major ingredient of all AAPBs. You further state that "Higher amounts of higher chain lengths and corresponding lower amounts of lower chain length could result in a rising average lipophilicity".
- 2. Different amounts of unsaturated fatty ester moieties: "Effects may be expected for e.g. physical state and for some toxicological endpoints, mainly local effects (e.g. irritation)".

You have further addressed the impact of impurities: "Due to the lack of differentiation between constituents and impurities, the terms "main constituents" and "impurities" are not regarded as relevant for UVCB substances". You have provided a table of "minor constituents" present in the composition of the substances used in the read-across approach.

You have also provided data matrix for physicochemical and (eco)toxicological properties to further support the mutual read-across of the AAPBs to one another regarding presence or absence of (eco)toxicological effects.

You further state that the read-across approach is justified due to following reasons:

- a) All AAPBs are similar in structure, since they are manufactured from similar resp. identical precursors under similar conditions and all contain the same functional groups. Thus a common mode of action can be assumed.
- b) The content of minor constituents in all products are comparable and differ to an irrelevant amount.



c) The only deviation within this group of substances is a minor variety in their fatty acid moiety, which is not expected to have a relevant impact on intrinsic toxic or ecotoxic activity and environmental fate. Potential minor impact on specific endpoints will be discussed in the specific endpoint sections.

B. ECHA analysis of the grouping and read-across approach in light of the requirements of Annex XI, 1.5.

B1. Grouping - Structural Similarity

In order to meet the provisions in Annex XI 1.5 to predict physicochemical and toxicological properties from data for a reference substance within the group by interpolation to other substances in the group, ECHA considers that structural similarity alone is not sufficient. It has to be justified why such prediction is possible in view of the identified structural differences and the provided evidence has to support such explanation. In particular, the structural similarities must be linked to a scientific explanation of how and why a prediction is possible.

ECHA understands that you intend to use a read-across approach where structurally similar substances have the same type and strength of effects.

ECHA agrees that the constituents of the four substances (i.e. C8 to C18 AAPB) share the same functional groups, namely: quaternary amines, amide bonds, carboxymethyl groups, and fatty acid chains. ECHA considers that the common functional groups support the readacross approach on the basis of structural similarity. ECHA further notes that the main constituents of the four substances exhibit the following structural differences: length of the C-chain and the degree of saturation in the fatty acid moiety.

ECHA notes that the four substances used in the read-across approach differ in their composition, i.e. in the distribution of the fatty acid moiety chain length, as shown in the table below with the information you provided in the read-across justification document.

ECHA agrees that the C12 (C12 carbon chain le	ength distribution) is the main common fatty
acid molety for all substances ranging from	%, with the remaining constituents
composing mostly of higher chain lengths in the	
concentrations (%) and (%) % of C8 an	d C10. The unsaturated fatty acid moieties
are mostly present in the C8-18 AAPB (< 2 %) and C8-18 and C18 unsaturated AAPB
(%).	

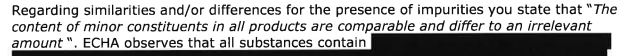


Carbon chain length distribution of Alkylamidopropyl betaines (AAPBs) as described in the read-across justification document submitted by you

C12 AAPB	C12-18 AAPB	C8-18 AAPB	C8-18 and C18 unsatd. AAPB	C12-14 AAPB
C10: < %, C12: > %, C14: < %	C8 + C10: = ,<br C12: %, C14: %, C16: %, C18: %, C18 unsatd.: < %	C8: <= %, C10: <= %, C12: %, C14: %, C16: %, C18: %, C18 unsatd.: <=	C6: <= %, C8: <= %, C10: <= %, C12: %, C14: %, C16: %, C18: %, C18 unsatd.:	C10: < %, C12: %, C14: %

You have addressed the differences in the structure of the constituents of the four substances and state that "The only deviation within this group of substances is a minor variety in their fatty acid moiety, which is not expected to have a relevant impact on intrinsic toxic or ecotoxic activity and environmental fate." Furthermore, you have addressed the differences in the composition of the four substances and state that "The content of minor constituents in all products are comparable and differ to an irrelevant amount."

ECHA observes that the differences in composition are covered with experimental data on **C8-18 AAPB** and **C8-18 and C18 unsatd. AAPB** addressing the impact of carbon chain length and unsaturation in the toxicological profile of the four substances used in the readacross approach.



The impurity profile of **C8-18 AAPB** differs from the other substances used in the readacross approach as it contains also ECHA considers that this difference is unlikely to affect the toxicological properties of the substance.

Based on the above ECHA considers that the structural similarity and the dissimilarities of the analogues are sufficiently explained with a view to considering the possibility of prediction.

B2. Predictions for toxicological properties

ECHA considers that the experimental studies conducted with the substances used in a read-across approach need to sufficiently cover the structural differences of the substances with regard to carbon chain length and unsaturation. This is needed to present a robust justification which meets the requirements of Annex XI, Section 1.5. that toxicological properties may be predicted from data for target substances. ECHA has therefore assessed the adequacy and reliability of the experimental studies provided and how the structural differences are covered by these studies.



As support for the proposed predictions for the read-across approach, you have provided:

- In vivo toxicokinetic data conducted with C12 AAPB (oral and dermal route) and in vitro dermal absorption study with C8-18 and C18 unsatd. AAPB;
- Experimental physico-chemical data conducted with C12 AAPB, C8-18 AAPB and C8-18 and C18 unsatd. AAPB. You state that "Similar physicochemical properties are expected for the other members of this group for which no experimental data are available based on structural similarity with differences only in the fatty acid chain length distribution";
- Experimental data on toxicological properties and conclude that the fatty acid moiety is not expected to "be relevant to the intrinsic systemic toxicity of the compounds", and not to have any influence on sensitisation. You have used C8-18 and C18 unsatd. AAPB as a worst case for skin and eye irritation and genotoxicity because it contains short chain fatty acid moieties and unsaturated fatty acid moieties. In particular, you have provided experimental data from C8-18 AAPB and C8-18 and C18 unsatd. AAPB regarding acute toxicity, skin and eye irritation, skin sensitisation and genotoxicity. You have also provided two sub-chronic toxicity (90-day) studies conducted with C8-18 AAPB and C8 unsatd. AAPB and a sub-acute (28-day) study conducted with C8-18 and C18 unsatd. AAPB, and a pre-natal developmental toxicity study in rats with C8-18 AAPB. You use this data to predict the toxicological properties of the other substances in the read-across approach.

You further conclude that "The read-across hypothesis is based on structural similarity of target and source substances. Based on the available experimental data, including key physico-chemical properties and data from toxicokinetic, acute toxicity, irritation, sensitisation, genotoxicity and repeated dose toxicity studies, the read-across strategy is supported by a quite similar toxicological profile of all five substances".

ECHA observes that the experimental studies provided in the read-across approach have been conducted with **C8-18 AAPB** and **C8-18 and C18 unsaturated AAPB** (with one supporting skin sensitisation study conducted with **C12 AAPB**).

ECHA notes that the composition of the test substances in the available experimental studies (namely: **C8-18 AAPB** and **C8-18 and C18 unsaturated AAPB**) are similar. The only difference is the concentration of the constituent C18 unsaturated, which is reported to be < % and % in these substances, respectively. ECHA further notes that in addition to the C12 fatty acid moiety these substances contain both the lower (C8 and C10) and higher (C14, C16, C18) carbon chain lengths and unsaturated C18 carbon chains.

ECHA has assessed the experimental data available and considers them adequate and reliable.

ECHA considers that structural and compositional variations of all the read-across substances are sufficiently covered with experimental data from **C8-18 AAPB** and **C8-18 and C18 unsatd. AAPB** regarding acute toxicity, skin and eye irritation, skin sensitisation, genotoxicity, repeated dose and prenatal developmental toxicity. ECHA notes that although no experimental studies are available for the **C12 AAPB** and **C12-18 AAPB** substances, the toxicological properties can be predicted from the common constituents with the **C8-18**



AAPB and **C8-18 and C18 unsatd. AAPB** substances that have adequate experimental data.

Conclusion on the grouping and read-across approach for toxicological properties:

Based on the reasons presented above, ECHA considers that the available studies and information are adequate and reliable and support the read-across approach as presented in the justification document for the endpoints that are not addressed with requests in this decision.

ECHA concludes that the read-across approach for these endpoints is plausible taking into account the toxicokinetic data (absorption, distribution, metabolism, elimination) and similar physico-chemical properties of the substances and the analysis of structural similarity presented in Section B1 above.

In accordance with Article 10(a)(ii) of the REACH Regulation, the technical dossier must contain information on the identity of the substance as specified in Annex VI, Section 2 to the REACH Regulation. In accordance with Annex VI, Section 2 the information provided has to be sufficient to enable the identification of the registered substance.

1. Pre-natal developmental toxicity study (Annex X, Section 8.7.2.) in a second species

In accordance with Articles 10(a) and 12(1) of the REACH Regulation, a technical dossier registered at more than 1000 tonnes per year must contain, as a minimum, the information specified in Annexes VII to X to the REACH Regulation. The information to be generated for the dossier must fulfil the criteria in Article 13(4) of the same regulation.

Pre-natal developmental toxicity studies (test method EU B.31./OECD TG 414) on two species are part of the standard information requirements for a substance registered for 1000 tonnes or more per year (Annex IX, Section 8.7.2., column 1, Annex X, Section 8.7.2., column 1, and sentence 2 of introductory paragraph 2 of Annex X of the REACH Regulation).

The technical dossier contains information on a pre-natal developmental toxicity study in rats by the oral route using the analogue substance **C8-18 AAPB** (CAS no 97862-59-4, EC no 931-296-8) as test material.

You have sought to adapt this information requirement according to Annex X, Section 8.7.2., column 2 and Annex XI, Section 1.2. You provided the following justification for the adaptation

"In accordance with Annex X column 2 of the REACH Regulation (EC) No 1907/2006, the performance of a Prenatal developmental toxicity study in a second species (non-rodent) is not required. AAPB is of low systemic toxicity as indicated by a LD50 > 2000 mg/kg bw. No indication of any systemic toxicity of AAPBs relevant in view of a potential health risk for humans was found in the sub-chronic studies, including reproductive organs. From developmental toxicity data, there is no evidence for teratogenic effects. AAPBs have no genotoxic properties as proven in the full data set including in vivo data. The use profile of the substance indicates that relevant exposure to humans occurs via the dermal route. Reliable, relevant and adequate toxicokinetic data from an in vitro study on human skin

CONFIDENTIAL 9 (17)



showed a dermal resorption rate of 0 %. Based on the above specified toxicological and toxicokinetic data, it can be proven that the substance is of low toxicological activity and that no systemic absorption occurs via the relevant route of exposure. Therefore, further reproductive toxicity studies do not need to be conducted.

Further, in accordance with Annex XI, section 1.2 of the REACH Regulation (EC) No 1907/2006, the performance of a Prenatal developmental toxicity study in a second species (non-rodent) is scientifically unjustified. As indicated above there is no indication of any systemic toxicity of AAPBs relevant in view of a potential health risk for humans, neither from sub-chronic data nor from developmental toxicity data. In conclusion, further testing on vertebrate animals in a Prenatal developmental toxicity study in a second species (non-rodent), is unjustified".

ECHA has analysed the conditions as specified in Annex X, column 2, 8.7.

a) Low toxicological activity:

As the read-across approach is considered acceptable (see Section 0 above) ECHA considers that data from the substances used in the read-across approach can be used.

ECHA agrees that the acute oral and dermal toxicity of the category members is low (LD50 > 2000 mg/kg bw/day) and no major systemic adverse effects were observed in the subchronic toxicity (90-day, in diet, OECD TG 408) and sub-acute toxicity (28-day, gavage) studies with the registered substance and sub-chronic toxicity (90-day, gavage, OECD 408) with **C8-18 AAPB** (CAS no 97862-59-4, EC no 931-296-8). However, ECHA notes that the highest doses used in these studies are 300 (90-day, gavage) and 247/300 mg/kg bw/day (90-day in diet/28-day, gavage) and thus it cannot be excluded that toxicity would be seen with higher doses.

ECHA further notes that in the pre-natal developmental toxicity study (OECD TG 14) conducted with **C8-18 AAPB** (CAS no 97862-59-4, EC no 931-296-8) effects on foetuses have been observed.

ECHA considers that the effects observed in the foetuses cannot be explained solely due to maternal toxicity. The available evidence indicates that the effects can also be attributed to the substance and therefore indicative for toxicological activity of the substance. Hence ECHA considers that the criteria of Annex IX, Column 2, 8.7. "low toxicological activity (no evidence of toxicity seen in any of the tests available)" are not met.

a) Toxicokinetic data

In your justification you state that "in vitro study on human skin showed a dermal resorption rate of 0 %" and "no systemic absorption occurs via the relevant route of exposure". ECHA notes that in the chemical safety report you also conclude that "Absorption after oral or dermal exposure in the described reliable experimental study on rats reached a maximum of 10 %. In an reliable in vitro study on dermal resorption on human skin, the resorption rate for Coco AAPB was even 0 %".

ECHA agrees that based on the *in vitro* dermal absorption study conducted with the registered substance dermal absorption is indeed 0 %. However, ECHA notes that *in vivo* dermal absorption study conducted with **C12 AAPB** (CAS no 4292-10-8, EC no 224-292-6) shows 3.5-6% (females) and 2-3.5% (males) absorption. Further, based on the *in vivo*



toxicokinetic study the same substance (**C12 AAPB**) is absorbed vial oral route ("approximately 5 % of the 14C dose was excreted in urine and < 2 % in expired air and < 2 % remained in the carcass").

ECHA therefore considers that there is evidence from reliable toxicokinetic data that systemic absorption occurs via relevant routes of exposure, e.g. dermal and oral and thus the criteria of Annex IX, Column 2, 8.7. "no systemic absorption occurs via relevant routes of exposure (e.g. plasma/blood concentrations below detection limit using a sensitive method and absence of the substance and of metabolites of the substance in urine, bile or exhaled air)" are not met.

ECHA observes that you further refer to the adaptation based on Annex XI, Section 1.2. Weight of Evidence: "no indication of any systemic toxicity of AAPBs relevant in view of a potential health risk for humans, neither from sub-chronic data nor from developmental toxicity data".

ECHA notes that according to Annex XI, Section 1.2. "There may be sufficient weight of evidence from several independent sources of information leading to the assumption/conclusion that a substance has or has not a particular dangerous property, while the information from each single source alone is regarded insufficient to support this notion".

As stated above in section a) Low toxicological activity, there is evidence from the pre-natal developmental toxicity study conducted with **C8-18 AAPB** that the substance(s) have toxicological activity.

ECHA observes that the information from the Chemical Safety Report and the exposure scenarios indicate potential for exposure from the oral, dermal and inhalation routes.

ECHA concludes that the substance(s) cannot be considered as having low toxicological activity and that no systemic exposure occurs.

Therefore, ECHA notes that your adaptation neither meets the specific rules for adaptation of Annex IX, Section 8.7., column 2 nor those of the general rules for adaptation of Annex XI; Section 1.2.

Therefore, your adaptation of the information requirement is rejected. As explained above, the information provided on this endpoint for the registered substance in the technical dossier does not meet the information requirement. Consequently there is an information gap and it is necessary to provide information for this endpoint.

The test in the first species was carried out by using a rodent species (rat). According to the test method EU B.31./OECD 414, the rabbit is the preferred non-rodent species. On the basis of this default assumption, ECHA considers that the test should be performed with rabbit as a second species.

ECHA considers that the oral route is the most appropriate route of administration for substances except gases to focus on the detection of hazardous properties on reproduction as indicated in ECHA *Guidance on information requirements and chemical safety assessment*



(version 5.0, December 2016) Chapter R.7a, Section R.7.6.2.3.2. Since the substance to be tested is a solid, ECHA concludes that testing should be performed by the oral route.

ECHA further considers that the test needs to be performed with the analogue substance **C12 AAPB** (CAS number 4292-10-8, EC number: 224-292-6), taking into account animal welfare considerations as well as because:

- 1. The **C12 AAPB** is the major constituent of all AAPBs used in the read-across approach
- 2. The C12 AAPB has the highest concentration of this constituent,
- 3. The C12 AAPB does not have experimental data covering systemic toxicity, developmental/reproductive toxicity
- 4. The higher and lower molecular weight constituents are covered by the available toxicity studies with the other substances used in the read-across approach.

In addition, **C12 AAPB** is considered suitable to be tested since the tests can be used as bridging studies to further strengthen the read-across approach.

Therefore, pursuant to Article 41(1) and (3) of the REACH Regulation, you are requested to submit the following information derived with the analogue substance (carboxymethyl)dimethyl-3-[(1-oxododecyl)amino]propylammonium hydroxide), (C12 AAPB, CAS no 4292-10-8, EC no 224-292-6) subject to the present decision: Pre-natal developmental toxicity study (test method: EU B.31./OECD TG 414) in a second species rabbit by the oral route.

Notes for your consideration

ECHA notes that a revised version of OECD TG 414 was adopted this year by the OECD. This revised version contains enhancements of certain endocrine disrupting relevant parameters. You should test in accordance with the revised version of the guideline as published on the OECD website for adopted test guidelines (https://www.oecd-ilibrary.org/environment/oecd-guidelines-for-the-testing-of-chemicals-section-4-health-effects 20745788).

2. Extended one-generation reproductive toxicity study (Annex X, Section 8.7.3.)

In accordance with Articles 10(a) and 12(1) of the REACH Regulation, a technical dossier registered at more than 1000 tonnes per year must contain, as a minimum, the information specified in Annexes VII to X to the REACH Regulation. The information to be generated for the dossier must fulfil the criteria in Article 13(4) of the same regulation.

The basic test design of an extended one-generation reproductive toxicity study (test method EU B.56./OECD TG 443 with Cohorts 1A and 1B, without extension of Cohort 1B to include a F2 generation, and without Cohorts 2A, 2B and 3) is a standard information requirement as laid down in column 1 of 8.7.3., Annex X. If the conditions described in column 2 of Annex X are met, the study design needs to be expanded to include the



extension of Cohort 1B, Cohorts 2A/2B, and/or Cohort 3. Further detailed guidance on study design and triggers is provided in the ECHA Guidance on information requirements and chemical safety assessment Chapter R.7a, Section R.7.6 (version 5.0, December 2016).

Adequate information on this endpoint needs to be present in the technical dossier for the registered substance to meet this information requirement.

a) The information provided

You have sought to adapt this information requirement according to Annex X, Section 8.7., column 2 and Annex XI, section 1.2. You provided the following justifications for the adaptation:

"In accordance with Annex X column 2 of the REACH Regulation (EC) No 1907/2006, the performance of an EOGRTS is not required. AAPB is of low systemic toxicity as indicated by a LD50 > 2000 mg/kg bw. No indication of any systemic toxicity of AAPBs relevant in view of a potential health risk for humans was found in the sub-chronic studies, including reproductive organs. From developmental toxicity data, there is no evidence for teratogenic effects. AAPBs have no genotoxic properties as proven in the full data set including in vivo data. The use profile of the substance indicates that relevant exposure to humans occurs via the dermal route. Reliable, relevant and adequate toxicokinetic data from an in vitro study on human skin showed a dermal resorption rate of 0 %. Based on the above specified toxicological and toxicokinetic data, it can be proven that the substance is of low toxicological activity and that no systemic absorption occurs via the relevant route of exposure. Therefore, further reproductive toxicity studies do not need to be conducted. Further, in accordance with Annex XI, section 1.2 of the REACH Regulation (EC) No 1907/2006, the performance of an EOGRTS is scientifically unjustified. As indicated above there is no indication of any systemic toxicity of AAPBs relevant in view of a potential health risk for humans, neither from sub-chronic data nor from developmental toxicity data. In conclusion, further testing on vertebrate animals in an EOGRTS is unjustified", and

"Further, in accordance with Annex XI, section 1.2 of the REACH Regulation (EC) No 1907/2006, the performance of a two-generation reproductive toxicity study is scientifically unjustified. As indicated above there is no indication of any systemic toxicity of AAPBs relevant in view of a potential health risk for humans, neither from sub-chronic data nor from developmental toxicity data. In conclusion, further testing on vertebrate animals in a 2-generation reproductive toxicity study or extended one generation reproductive toxicity study is unjustified".

ECHA observes that you have provided the same justification for the pre-natal developmental toxicity and the extended one-generation reproductive toxicity endpoints.

As explained in section 1 above, your adaptation neither meets the specific rules for adaptation of Annex X, Section 8.7., column 2 nor those of the general rules for adaptation of Annex XI, Section 1.2.

You further conclude the following:

"In the repeated dose toxicity studies in rats conducted with C8-18 and C18 unsatd. AAPB and C8-18 AAPB, there were no histopathological changes in reproductive organs (seminal vesicles, prostate, epididymides, testes, mammary glands, ovaries and fallopian tubes,



uterus, cervix, vagina) and no effects on reproductive organs weights (testes, ovaries). Taking into account the overall low toxic activity of the AAPBs, particularly with regard to the missing adverse effects on reproductive organs or tissues in the 28-day and 90-day studies as well as in the developmental toxicity study, the missing teratogenic activity, the fact that embryotoxic effects were found only at the maternal toxic dose level and the toxicodynamic of AAPBs, which is primarily based on its irritancy, fertility-specific effects are highly unlikely".

You claim that the available information from the repeated dose toxicity studies in the rat confirm that the reproductive organs are not affected after repeated exposure to the registered substance. ECHA notes that histopathological data alone does not adequately address all relevant elements with respect to sexual function and fertility.

ECHA further notes that your adaptation justification does not fully address the effects on offspring. The study according to OECD TG 414 in the rat provide information only on effects observable pre-natally and not effects on offspring observable and/or due to postnatal exposure. In particular, essential information on offspring toxicity observable and/or due to the peri-and postnatal exposure up to the adulthood is missing.

Thus, the information you provided does not adequately address all relevant elements with respect to effects on fertility and offspring. As explained above, the information you provided is not sufficient to support your conclusion that the substance does not have a dangerous property with respect to sexual function and fertility.

Therefore, your adaptation of the information requirement is rejected.

As explained above, the information provided on this endpoint for the registered substance in the technical dossier does not meet the information requirement. Consequently there is an information gap and it is necessary to provide information for this endpoint. Thus, an extended one-generation reproductive toxicity study according Annex X, Section 8.7.3. is required. The following refers to the specifications of this required study.

b) The specifications for the required study

Premating exposure duration and dose-level setting

To ensure that the study design adequately addresses the fertility endpoint, the duration of the premating exposure period and the selection of the highest dose level are key aspects to be considered. According to the ECHA *Guidance on information requirements and chemical safety assessment* Chapter R.7a, Section R.7.6 (version 6.0, July 2017), the starting point for deciding on the length of the premating exposure period should be ten weeks to cover the full spermatogenesis and folliculogenesis before the mating, allowing meaningful assessment of the effects on fertility.

Ten weeks premating exposure duration is required because there is no substance specific information in the dossier supporting shorter premating exposure duration as advised in the ECHA *Guidance on information requirements and chemical safety assessment* Chapter R.7a, Section R.7.6 (version 6.0, July 2017).

The highest dose level shall aim to induce some toxicity to allow comparison of effect levels

CONFIDENTIAL 14 (17)



and effects of reproductive toxicity with those of systemic toxicity. The dose level selection should be based upon the fertility effects with the other cohorts being tested at the same dose levels.

If there is no existing relevant data to be used for dose level setting, it is recommended that results from a range-finding study (or range finding studies) are reported with the main study. This will support the justifications of the dose level selections and interpretation of the results.

Species and route selection

According to the test method EU B.56/ OECD TG 443, the rat is the preferred species. On the basis of this default assumption, ECHA considers that testing should be performed in rats.

ECHA considers that the oral route is the most appropriate route of administration for substances except gases to focus on the detection of hazardous properties on reproduction as indicated in ECHA Guidance on information requirements and chemical safety assessment (version 6.0, July 2017) Chapter R.7a, Section R.7.6.2.3.2. Since the substance to be tested is a [liquid/solid/dust], ECHA concludes that testing should be performed by the oral route.

ECHA further considers that the test needs to be performed with the analogue substance **C12 AAPB** (CAS number 4292-10-8, EC number: 224-292-6), taking into account animal welfare considerations as well as because:

- The C12 AAPB is the major constituent of all AAPBs used in the read-across approach
- 2. The C12 AAPB has the highest concentration of this constituent,
- The C12 AAPB does not have experimental data covering systemic toxicity, developmental/reproductive toxicity
- 4. The higher and lower molecular weight constituents are covered by the available toxicity studies with the other substances used in the read-across approach.

In addition, **C12 AAPB** is considered suitable to be tested since the tests can be used as bridging studies to further strengthen the read-across approach.

b) Outcome

Based on the available information, pursuant to Article 41(1) and (3) of the REACH Regulation, you are requested to submit the following information derived with the analogue substance (carboxymethyl)dimethyl-3-[(1-oxododecyl)amino]propylammonium hydroxide (C12 AAPB, CAS no 4292-10-8, EC no 224-292-6): Extended one-generation reproductive toxicity study (test method EU B.56./OECD TG 443), in rats, oral route, according to the following study-design specifications:

- Ten weeks premating exposure duration for the parental (P0) generation;
- Dose level setting shall aim to induce some toxicity at the highest dose level;

CONFIDENTIAL 15 (17)



- Cohort 1A (Reproductive toxicity);
- Cohort 1B (Reproductive toxicity) without extension to mate the Cohort 1B animals to produce the F2 generation;

While the specifications for the study design are given above, you shall also submit with the new endpoint study record a scientific justification on each of the following aspects: 1) length of the premating exposure duration and dose level selection, 2) reasons for why or why not Cohort 1B was extended, 3) termination time for F2 generation, and 4) reasons for why or why not Cohorts 2A/2B and/or Cohort 3 were included.

The conditions to include the extension of Cohort 1B are currently not met. Furthermore, no triggers for the inclusion of Cohorts 2A and 2B (developmental neurotoxicity) and Cohort 3 (developmental immunotoxicity) were identified. However, you may expand the study by including the extension of Cohort 1B, Cohorts 2A and 2B and/or Cohort 3 if relevant information becomes available after this decision is issued to justify such an inclusion. Inclusion is justified if the available information, together with the new information shows triggers which are described in column 2 of Section 8.7.3., Annex IX and further elaborated in ECHA Guidance on information requirements and chemical safety assessment R.7a, chapter R.7.6 (version 6.0, July 2017). You may also expand the study to address a concern identified during the conduct of the extended one-generation reproduction toxicity study and also due to other scientific reasons in order to avoid a conduct of a new study. The justification for the expansion must be documented.



Appendix 2: Procedural history

For the purpose of the decision-making, this decision does not take into account any updates of your registration after the date when the draft decision was notified to you under Article 50(1) of the REACH Regulation.

The compliance check was initiated on 29 August 2017.

The decision making followed the procedure of Articles 50 and 51 of the REACH Regulation, as described below:

ECHA notified you of the draft decision and invited you to provide comments.

ECHA did not receive any comments by the end of the commenting period.

ECHA notified the draft decision to the competent authorities of the Member States for proposal(s) for amendment.

As no amendments were proposed, ECHA took the decision according to Article 51(3) of the REACH Regulation.



Appendix 3: Further information, observations and technical guidance

- 1. This compliance check decision does not prevent ECHA from initiating further compliance checks on the present registration at a later stage.
- 2. Failure to comply with the requests in this decision, or to otherwise fulfil the information requirements with a valid and documented adaptation, will result in a notification to the enforcement authorities of your Member State.
- 3. In relation to the information required by the present decision, the sample of the substance used for the new tests must be suitable for use by all the joint registrants. Hence, the sample should have a composition that is suitable to fulfil the information requirement for the range of substance compositions manufactured or imported by the joint registrants.

It is the responsibility of all joint registrants who manufacture or import the same substance to agree on the appropriate composition of the test material and to document the necessary information on their substance composition. In addition, it is important to ensure that the particular sample of the substance tested in the new tests is appropriate to assess the properties of the registered substance, taking into account any variation in the composition of the technical grade of the substance as actually manufactured or imported by each registrant.

If the registration of the substance by any registrant covers different grades, the sample used for the new tests must be suitable to assess these grades. Finally there must be adequate information on substance identity for the sample tested and the grades registered to enable the relevance of the tests to be assessed.

4. If the required tests are conducted with an analogue substance in the context of a read-across approach, the identity of the test material used to perform the test should be specified in line with ECHA's Practical Guide on "How to use alternatives to animal testing to fulfil your information requirements" (chapter 4.4). This is required to show that the test material is representative of the analogue substance identified in the read-across approach and used to predict the properties of the registered substance.