

Biocidal Products Committee (BPC)

Opinion on the application for approval of the active substance:

**Reaction products of paraformaldehyde and 2-hydroxypropylamine
(ratio 3:2)**

Product type: 2

ECHA/BPC/334/2022

Adopted

08 June 2022

Opinion of the Biocidal Products Committee

on the application for approval of the active substance Reaction products of paraformaldehyde and 2-hydroxypropylamine (ratio 3:2) for product type 2

In accordance with Article 90(2) of Regulation (EU) No 528/2012 of the European Parliament and of the Council 22 May 2012 concerning the making available on the market and use of biocidal products (BPR), the Biocidal Products Committee (BPC) has adopted this opinion on the approval in product type 2 of the following active substance:

Common name:	Formaldehyde released from the reaction products of paraformaldehyde and 2-hydroxypropylamine (ratio 3:2)
	RP 3:2
Chemical name:	Reaction products of paraformaldehyde and 2-hydroxypropylamine (ratio 3:2)
EC No.:	not applicable
CAS No.:	not applicable
Existing active substance	

This document presents the opinion adopted by the BPC, having regard to the conclusions of the evaluating Competent Authority. The assessment report, as a supporting document to the opinion, contains the detailed grounds for the opinion.

Process for the adoption of BPC opinions

Following the submission of an application by Task Force Lubrizol Deutschland GmbH and Schülke & Mayr GmbH. on 1 August 2007, the evaluating Competent Authority Austria submitted an assessment report and the conclusions of its evaluation to the European Chemicals Agency on 29 September 2016. In order to review the assessment report and the conclusions of the evaluating Competent Authority, the Agency organised consultations via the BPC (BPC-21) and its Working Groups (WG-II-2017). Revisions agreed upon were presented and the assessment report and the conclusions were amended accordingly.

Information on the fulfilment of the conditions for considering the active substance as a candidate for substitution was made publicly available at <https://echa.europa.eu/de/addressing-chemicals-of-concern/biocidal-products-regulation/potential-candidates-for-substitution-previous-consultations/-/substance-rev/15204/term> on 4 November 2016, in accordance with the requirements of Article 10(3)

of Regulation (EU) No 528/2012. Interested third parties were invited to submit relevant information by 3 January 2017.

In May 2018 a request from DG SANTE according to Article 75(1)(g) of Regulation (EU) No 528/2012 to the BPC regarding an ED-assessment according to the scientific criteria set out in Commission Delegated Regulation (EU) 2017/2100 was forwarded by ECHA to the eCA. Therefore, the evaluation of "RP 3:2" was amended accordingly with an ED-assessment based on the available data. The revised evaluation report including the ED assessment was submitted on 15 September 2021 in ECHA Process flow 43 (WG-I-2022 and BPC-43) for review and discussion.

Adoption of the BPC opinion

Rapporteur: Austria

The BPC opinion on the approval of the active substance reaction products of paraformaldehyde and 2-hydroxy-propylamine (ratio 3:2) in product type 2 was adopted on 8 June 2022.

The BPC opinion takes into account the comments of interested third parties provided in accordance with Article 10(3) of BPR.

The BPC opinion was adopted by consensus.

The opinion is published on the ECHA webpage at:
<http://echa.europa.eu/regulations/biocidal-products-regulation/approval-of-active-substances/bpc-opinions-on-active-substance-approval>.

Detailed BPC opinion and background

1. Overall conclusion

Since RP 3:2 fulfils the criteria set in Article 5(1) of Regulation (EU) No 528/2012, the overall conclusion of the BPC is that RP 3:2 in product type 2 should normally not be approved, unless one of the conditions for derogation in Article 5(2) is met. The process related to the demonstration of whether the conditions for derogation set in Article 5(2) are met, is not in the remit of the BPC¹. The detailed grounds for the overall conclusion are described in the assessment report.

2. BPC Opinion

2.1. BPC Conclusions of the evaluation

Presentation of the active substance including the classification and labelling of the active substance

This evaluation covers the use of Reaction products of paraformaldehyde and 2-hydroxypropylamine (ratio 3:2), furthermore addressed as RP 3:2 in product type 2. RP 3:2 was originally notified as 3,3'-methylene-bis(5-methyl-oxazolidine) – MBO². RP 3:2 is a formaldehyde-releaser. Specifications for the reference source are established.

The physico-chemical properties of the active substance and biocidal product have been evaluated and are deemed acceptable for the appropriate use and materials suitable for storage and transport of the active substance and biocidal product. Regarding the explosive properties the justification for non-submission of data has not been accepted. An experimental test has to be conducted as the substance contains of unknown constituents and therefore the waiving cannot be justified by structural considerations.

Validated analytical methods are available for the active substance as manufactured and for the relevant and significant impurities. With regard to the methods submitted for determination of the hydrolysis product 2-hydroxypropylamin in water and soil the data has been considered as not sufficient. The classification and labelling for RP 3:2 according to Regulation (EC) No 1272/2008 (CLP Regulation) as agreed by RAC-35 (December 2015)³ and published in Regulation (EC) No 2017/776 (10th adaption to technical progress):

Classification according to the CLP Regulation	
Hazard Class and Category Codes	Acute Tox. 4, H302 Acute Tox. 3, H311 Acute Tox. 4, H332 Skin Corr. 1B, H314 Skin Sens. 1A, H317 Eye dam. 1, H318

¹ See document: Further guidance on the procedures related to the examination of the exclusion criteria and the conditions for derogation under Article 5(2) (CA-Nov14-Doc.4.5-Final).

² The renaming of 3,3'-methylene-bis(5-methyl-oxazolidine) – MBO into Reaction products of paraformaldehyde and 2-hydroxypropylamine (ratio 3:2) is not regarded as a redefinition according to Article 11 of Regulation (EU) No 1062/2014.

³<https://echa.europa.eu/documents/10162/96051139-f376-4da3-b25a-130668d6db45>

⁴https://echa.europa.eu/documents/10162/16908203/esd_pt_13_final_en.pdf/cc11b52f-0c3e-4a56-883d-2abebcb0d5fc

	STOT RE 2, H373 Muta 2, H341* Carc. 1B, H350** Aquatic Chronic 2, H411
	* The classification as a mutagen need not apply if it can be shown that the maximum theoretical concentration of releasable formaldehyde, irrespective of the source, in the mixture as placed on the market is less than 1%. ** The classification as a carcinogen need not apply if it can be shown that the maximum theoretical concentration of releasable formaldehyde, irrespective of the source, in the mixture as placed on the market is less than 0.1%.
Labelling	
Pictograms	GHS 05, GHS 06, GHS 08, GHS 09
Signal Word	Danger
Hazard Statement Codes	H302: Harmful if swallowed H311: Toxic in contact with skin H332: Harmful if inhaled H314: Causes severe skin burns and eye damage H317: May cause an allergic skin reaction H373 May cause damage to organs (gastrointestinal tract and respiratory tract) H341: Suspected of causing genetic defects H350: May cause cancer H411: Toxic to aquatic life with long-lasting effects
Suppl. Hazard Statement Code	EUH071: Corrosive to the respiratory tract
Specific Concentration limits, M-Factors	M = not applicable

a) Intended use, target species and effectiveness

RP 3:2 containing biocidal products are used as disinfectant system cleaner of metal working systems for the disinfection of inner surfaces of vessels and tubes. Therefore, this application can be actually assigned to product type 2 as it is considered to be a disinfectant. Nevertheless, system cleaners are used in the same industrial area such as metalworking fluids.

The biocidal product tested inhibited growth of gram negative bacteria such as *P. putida*, *E. coli* and gram positive bacteria such as *Staphylococcus aureus* and *Mycobacterium immunogenum*, yeasts such as *Candida albicans* and fungi such as *Fusarium oxysporum*.

Overall, a microbicidal effect of the biocidal product was demonstrated with concentrations ranging from 0.25% (=2500 ppm) up to 2% (20000 ppm) depending on the exposure time.

The active substance is a formaldehyde-releaser. The biocidal activity of the active substance is due to the interaction of the released formaldehyde with protein, DNA and RNA. The interaction with protein results from a combination with the primary amide and the amino groups. It reacts with carboxyl, sulfhydryl and hydroxyl groups.

As formaldehyde is not specific for one cellular target, the development of resistance is unlikely, if sufficiently high formaldehyde concentrations are guaranteed that exceed the capacity of the innate detoxification systems.

b) Overall conclusion of the evaluation including need for risk management measures

A common core dossier was developed for formaldehyde, which was agreed at a Biocides Technical Meeting. This core dossier forms the basis of the hazard assessment of formaldehyde for all formaldehyde releasing active substances.

Human health

The toxicity of the active substances is dominated by skin sensitization and local irritation and local (in vitro) genotoxicity (but negative systemic in vivo genotoxicity) and the hydrolysis study and efficacy mode of action support that the equilibrium within the RP 3:2 quickly shifts towards formaldehyde and 2-hydroxypropylamine by dilution and by the reaction of formaldehyde with biological media. This is essentially the basis for reading across the classification of formaldehyde for germ cell mutagenicity category 2 and carcinogenicity category 1B. However the risk assessment provided below for local and for systemic effects includes also the potential for carcinogenic effects.

The risk from the application of RP 3:2 as PT2 within system cleaners within industrial processes is characterised in this CAR. Application concentrations are typically 10%. Risk is estimated for the formulation and use of system cleaners. Due to the high concentration of RP 3:2 in the systems cleaner incomplete hydrolysis to formaldehyde and 2-hydroxypropylamin is expected. Therefore dermal risk estimates are provided for 2 situations: non-hydrolysed RP 3:2 and full hydrolysis to formaldehyde and 2-hydroxypropylamin. The second hydrolysis product 2-hydroxypropylamine is not further considered since the respective AEL is much higher and the exposure potential is not higher compared to formaldehyde. However for respiratory exposure just the situation of complete hydrolysis is considered: the vapour pressures of the products and the composition of the gaseous phases are expected to be already determined inter alia by hydrolysis products. Therefore it is considered to be appropriate to estimate the respiratory exposure for the situation of complete hydrolysis of RP 3:2 to formaldehyde. The second hydrolysis product 2- hydroxypropylamine is not further considered since the respective AEL is much higher and volatility, i.e. exposure potential, is lower compared to formaldehyde.

The table below summarises the exposure scenarios assessed.

Summary table: human health scenarios			
Scenario	Primary or secondary exposure and description of scenario	Exposed group	Conclusion
PT2: Formulation of system cleaners	<p>Primary exposure covering the following tasks: mixing of components, sampling, filling and bottling, cleaning</p> <p>Exposure to active substance and system cleaners</p> <p>Acceptable with local exhaust ventilation (LEV), automatic dosing system, sampling with practically closed system*, technical and organisational RMM and PPE (like gloves, coveralls, masks) for high local hazard category (dosing, mixing, sampling) and RMM and PPE for medium local hazard category (cleaning)</p> <p>* use of closed bottles and appropriate LEV, if system is opened</p>	Industrial workers	Acceptable with RMM and PPE
PT2: Use of system cleaners	<p>Primary exposure covering the following tasks: loading, dosing, sampling, cleaning, maintenance of the system</p> <p>Exposure to system cleaners and treatment solutions</p> <p>Acceptable with automatic dosing system, technical and organisational RMM and PPE (like gloves, coveralls, masks) for high local hazard category (loading) and RMM and PPE for medium local hazard category for other tasks.</p>	Industrial workers	Acceptable with RMM and PPE.

Risk for mixing and loading of RP 3:2 to system cleaner formulation vessels, sampling for formulation control, filling, bottling and cleaning of vessels in formulation plants is considered. Due to the corrosive and sensitizing hazard of 100% RP 3:2 as well as the 10% dilution in system cleaners, closed systems (including sampling task) and high industrial organisational and technical RMM have to be used in order to allow concluding that the risk for local respiratory and local dermal effects is acceptable. In this case also risk for systemic effects is acceptable. Considering that local irritation is a condition for the development of tumours and applying a deterministic threshold AEC and AEL, also the risk for potential carcinogenic effects appears acceptable.

No exposure of general public, no exposure of pets and no dietary exposure is expected due to the intended PT2 use. Dermal contact against dried concentrates in dirty clothes in home laundry of working clothes is assumed to be not relevant as RP 3:2 residues will quickly

hydrolyse and generate gaseous formaldehyde, which is transferred to the gaseous phase and will not remain on the clothes.

Environment

The risk characterization was based on the hydrolysis products 2-hydroxypropylamine and formaldehyde as during the disinfection and use of the metal working system RP 3:2 has almost completely hydrolysed. The parent compound itself is therefore not expected to reach any environmental compartment. 2-Hydroxypropylamine and formaldehyde are expected to be readily biodegradable in the environment and are unlikely to bioaccumulate in biota. For acute toxicity algae is the most sensitive species with a 72h- ErC_{50} of 5.7 mg/L (geometr. Mean, *Desmodesmus subspicatus*) for formaldehyde and a 96h- EcC_{50} of 118.4 mg/L (nominal, buffered *Pseudokirchneriella subcapitata*) for 2-hydroxypropylamine. Both compounds are not classifiable towards environmental hazards based on the available data.

Possible releases into the environment during use of system cleaners are identical with releases caused by metalworking fluids, i.e. local releases during use of system cleaners are covered by the emission scenarios for metalworking fluids (Refinement of the Emission Scenario Document (ESD) for Product Type 13, May 2015⁴). This approach was agreed at WG-II-2017 and it must be emphasised that only this specific use as system cleaners of metal working systems is covered by the ESD of PT13.

The table below summarises the exposure scenarios assessed.

Summary table: environment scenarios		
Scenario	Description of scenario including environmental compartments	Conclusion
PT2: Private area and public health area disinfectant and other biocidal products Emulsifiable fluids Waste treatment: - End-user, on-site: splitting techniques of emulsion based on K_{ow}	<p>The cleaner is not discharged but diluted in the metalworking fluid (MWF) and will then be used as preservative. Therefore, possible releases into the environment during use of system cleaners are covered by the emission scenario for metalworking fluids (PT13).</p> <p>Waste treatment is done by end-users who treat their waste on-site and apply commonly used emulsion splitting techniques based on the partition coefficient K_{ow} (e.g. chemical splitting, ultrafiltration). All waste waters resulting from the use of water miscible MWF will be led to biological treatment before discharge into the environment.</p> <p>Affected environmental compartments:</p> <p>STP, surface water, sediment, soil, groundwater</p>	Not acceptable for STP, surface water and the terrestrial compartment for 2-hydroxypropylamine and formaldehyde at an in-use concentration of 0.25% of RP 3:2.

⁴https://echa.europa.eu/documents/10162/16908203/esd_pt_13_final_en.pdf/cc11b52f-0c3e-4a56-883d-2abebcb0d5fc

Summary table: environment scenarios		
Scenario	Description of scenario including environmental compartments	Conclusion
<p>Emulsifiable fluids</p> <p>Waste treatment:</p> <p>- End-user, on-site: splitting via evaporation (based on vapour pressure)</p>	<p>Waste treatment is done by end-users who treat their waste on-site. The splitting of the emulsion is based on the on vapour pressure and done via evaporation. All waste waters resulting from the use of water miscible MWF will be led to biological treatment before discharge into the environment.</p> <p>Affected environmental compartments:</p> <p>STP, surface water, sediment, soil, groundwater</p>	<p>Acceptable for 2-hydroxypropylamine at an in-use concentration of 0.25% of RP 3:2.</p> <p>Not acceptable for formaldehyde for STP, surface water and soil at an in-use concentration of 0.25% of RP 3:2.</p>
<p>Emulsifiable fluids</p> <p>Waste treatment:</p> <p>- Waste management company: splitting techniques of emulsion based on K_{ow}</p>	<p>Waste treatment is managed by a waste management company which receives waste from smaller MWF-using companies. The splitting of the emulsion is based on the partition coefficient K_{ow}. All waste waters resulting from the use of water miscible MWF will be led to biological treatment before discharge into the environment.</p> <p>Affected environmental compartments:</p> <p>STP, surface water, sediment, soil, groundwater</p>	<p>Acceptable for 2-hydroxypropylamine at an in-use concentration of 0.25% of RP 3:2.</p> <p>Not acceptable for formaldehyde for STP, surface water and soil at an in-use concentration of 0.25% of RP 3:2.</p>
<p>Emulsifiable fluids</p> <p>Waste treatment:</p> <p>- Waste management company: splitting via evaporation (based on vapour pressure)</p>	<p>Waste treatment is managed by a waste management company which receives waste from smaller MWF-using companies. The splitting of the emulsion is based on the on vapour pressure and done via evaporation. All waste waters resulting from the use of water miscible MWF will be led to biological treatment before discharge into the environment.</p> <p>Affected environmental compartments:</p> <p>STP, surface water, sediment, soil, groundwater</p>	<p>Acceptable for 2-hydroxypropylamine at an in-use concentration of 0.25% of RP 3:2.</p> <p>Not acceptable for formaldehyde for STP, surface water and the terrestrial compartment at an in-use concentration of 0.25% of RP 3:2.</p>

The risk assessment was calculated for the life cycle stage "waste treatment after refreshment".

The risk for 2-hydroxypropylamine was acceptable for the cleaning of metal working fluid lines (PT2), in-use concentrations 0.25% of RB 3:2, if spent fluids were treated by specialized waste management companies. For an in-use concentration of 2% of RB 3:2 a risk for the

scenario waste management companies (splitting based on Kow) was identified for STP, surface water and the terrestrial compartment.

Risks of formaldehyde in the four scenarios covering PT2 as well as risks of 2- hydroxypropylamine for PT2 from spent metal working fluids (MWFs) for end-user on-site treatment are acceptable only:

- If formaldehyde concentrations in the water phase after on-site or off-site treatment is below 40 mg/L, unless further evidence is provided within product authorisation that measurements are not necessary and
- End-user who treat the MWFs on-site apply evaporation splitting techniques

Formaldehyde is a very reactive compound so it can be assumed that during and after application of the product the concentration of formaldehyde will decrease by a chain of chemical reactions. Degradation since last dosing, i.e. operating time since last biocide dosing, storage at end-user site or transport to waste management facility and storage at waste management site was not taken into account in the exposure calculations (as there is no standard algorithm that can be used to derive such degradation rates). However, for an estimation of the exposure reduction, a supporting pilot study on the degradation of formaldehyde in used MWF was submitted. In this pilot study the formaldehyde concentration of an used emulsion of a metalworking company was measured, which was delivered to a waste treatment company. The measurements showed that after distillation, the content of formaldehyde in the distillate is below the limit of quantification (<40 mg/L). The pilot study results are considered as supportive information due to several limitations. Regarding a concentration of 40 mg formaldehyde per litre waste water in the influent of a municipal STP, no unacceptable risks concerning the affected environmental compartments are indicated. For a refinement of the exposure calculations additional robust information on degradation of formaldehyde between the last dosing and the start of waste treatment would be necessary. This additional information enables the use of an elimination fraction that includes degradation during use, storage at end-user site, transport to waste management facility and storage at waste management site.

It has to be highlighted, that the environmental exposure calculations are very conservative as no degradation of formaldehyde in the system during use was assumed. However the above mentioned pilot study provides evidence for high degradation of formaldehyde during use (83 % between dosing and removal of the metal working fluid).

As the exceedance of the PEC/PNEC_{STP} ratio regarding formaldehyde is around 5, the required new data will allow to refine the exposure calculations also in this respect, which will most likely result in an acceptable risk.

A risk management option for the terrestrial compartment is a restriction of the use of sewage sludge in agriculture depending on national legislations.

The measurements of formaldehyde in effluents take into consideration aggregate exposure from multiple uses and product types as well as exposure from other formaldehyde sources, at least for end-user on-site treatment aggregate exposure of PT2 and PT13 would be covered.

Concerning risks to surface water intended for the abstraction of drinking water the parametric value of 0.1 µg/L of the Drinking Water Directive 98/83/EC may be exceeded. This

may be considered by the relevant national authorities when issuing permits for recovery plants.

Overall conclusion

Overall a safe use has been identified for the use of RP 3:2 as a system cleaner in PT2 provided adequate RMM and PPE are considered for human health and environment.

2.2. Exclusion, substitution and POP criteria

2.2.1. Exclusion and substitution criteria

The table below summarises the relevant information with respect to the assessment of exclusion and substitution criteria:

Property		Conclusions	
CMR properties	Carcinogenicity (C)	Cat 1B	RP 3:2 does fulfil criterion (a) of Article 5(1)
	Mutagenicity (M)	Cat 2	
	Toxic for reproduction (R)	no classification required	
PBT and vPvB properties	Persistent (P) or very Persistent (vP)	not P or vP	RP 3:2 does not fulfil criterion (e) of Article 5(1) and does not fulfil criterion (d) of Article 10(1)
	Bioaccumulative (B) or very Bioaccumulative (vB)	not B or vB	
	Toxic (T)	T	
Endocrine disrupting properties	Section A of Regulation (EU) 2017/2100: ED properties with respect to humans	<p>An assessment of the endocrine disrupting properties was conducted:</p> <ul style="list-style-type: none"> - the ED criteria for the T modality are not met; - for EAS modalities no conclusion can be drawn based on the available data. <p>However, considering the known severe hazard properties of this substance and based on scientific reasons, further data will not be requested in this special case.</p>	No conclusion can be drawn whether RP 3:2 fulfils criterion (d) of Article 5(1) and/or criterion (e) of Article 10(1).
	Section B of Regulation (EU) 2017/2100: ED properties with respect to non-target organisms	<p>An assessment of the endocrine disrupting properties was conducted: for EAS modalities as well as for T-modality no conclusion can be drawn based on the available data.</p> <p>However, considering the hazard profile of this</p>	

Property		Conclusions	
		substance and the anticipated difficulties to determine the mode of action, further data will not be requested in this special case based on scientific reasons.	
	Article 57(f) and 59(1) of REACH	No	
	Intended mode of action that consists of controlling target organisms via their endocrine system(s).	No	
Respiratory sensitisation properties	No classification required. RP 3:2 does not fulfil criterion (b) of Article 10(1).		
Concerns linked to critical effects	Based on the available data it cannot be concluded if RP: 3:2 does fulfil criterion (e) of Article 10(1).		
Proportion of non-active isomers or impurities	The substance does not contain a significant proportion of non-active isomers or impurities. RP 3:2 does not fulfil criterion (f) of Article 10(1).		

Consequently, the following is concluded:

Reaction products of paraformaldehyde and 2-hydroxypropylamine (ratio 3:2) does meet the exclusion criteria laid down in Article 5(1) of Regulation (EU) No 528/2012 by the released formaldehyde being a carcinogen Cat 1B.

Reaction products of paraformaldehyde and 2-hydroxypropylamine (ratio 3:2) does meet the conditions laid down in Article 10(1)(a) of Regulation (EU) No 528/2012, and is therefore considered as a candidate for substitution by meeting the exclusion criteria.

The exclusion and substitution criteria were assessed in line with the "Note on the principles for taking decisions on the approval of active substances under the BPR"⁵ and in line with "Further guidance on the application of the substitution criteria set out under article 10(1) of the BPR"⁶ and "Implementation of scientific criteria to determine the endocrine –disrupting properties of active substances currently under assessment"⁷ agreed at the 54th, 58th and 77th meeting respectively, of the representatives of Member States Competent Authorities for the implementation of Regulation 528/2012 concerning the making available on the market

⁵ See document: Note on the principles for taking decisions on the approval of active substances under the BPR (available from <https://circabc.europa.eu/d/a/workspace/SpacesStore/c41b4ad4-356c-4852-9512-62e72cc919df/CA-March14-Doc.4.1%20-%20Final%20-%20Principles%20for%20substance%20approval.doc>)

⁶ See document: Further guidance on the application of the substitution criteria set out under article 10(1) of the BPR (available from [https://circabc.europa.eu/d/a/workspace/SpacesStore/dbac71e3-cd70-4ed7-bd40-fc1cb92cfe1c/CA-Nov14-Doc.4.4%20-%20Final%20-%20Further%20guidance%20on%20Art10\(1\).doc](https://circabc.europa.eu/d/a/workspace/SpacesStore/dbac71e3-cd70-4ed7-bd40-fc1cb92cfe1c/CA-Nov14-Doc.4.4%20-%20Final%20-%20Further%20guidance%20on%20Art10(1).doc))

⁷ See document: Implementation of scientific criteria to determine the endocrine –disrupting properties of active substances currently under assessment (<https://circabc.europa.eu/sd/a/48320db7-fc33-4a91-beec-3d93044190cc/CA-March18-Doc.7.3a-final-%20EDs-%20active%20substances%20under%20assessment.docx>)

and use of biocidal products. This implies that the assessment of the exclusion criteria is based on Article 5(1) and the assessment of substitution criteria is based on Article 10(1)(a, b, d, e and f).

An ED assessment for RP 3:2 has been carried out taken the EFSA/ECHA (2018) guidance for the identification of endocrine disruptors into account. The ED assessment has been also discussed in the ED expert group via a written procedure, followed by an ad hoc meeting in June 2021. The advice of the experts was considered for the ED assessment.

For the T-modality the ED criteria are not met for human health. For non-target organisms no conclusion can be drawn on T-modality based on the available data. For EAS modalities no conclusion can be drawn for human health and non-target organisms based on the available data. Further testing with the active substance is not considered appropriate in that specific case because 'testing does not appear scientifically necessary' (first heading of Annex IV of Regulation (No) 528/2012 and because 'testing is technically challenging' (referring to second heading on Annex IV), as detailed below.

- It is uncertain, if further mechanistic studies, particularly with mammals with RP 3:2 would allow establishing a mode of action, keeping in mind that endocrine mediated endpoints may be impacted secondary to general, non-endocrine toxicity and that in vivo apical endpoints can be triggered by several modes of action, including endocrine and non-endocrine modalities. Also for aquatic species it would be challenging to get meaningful results in further tests as correct dose setting and detangling the ED mode of action from non-ED modes of action are hampering the performance and interpretation of such tests. For birds no agreed and adequate study protocols are available to determine endocrine modes of action.
- Due to the properties of RP 3:2 as skin corrosive, skin sensitising and local acting genotoxic carcinogen and the corresponding low effect concentration(s), it is difficult to select an appropriate test system to get meaningful results, at least for mammals.
- The main hydrolysis product formaldehyde of RP 3:2 is an endogenously formed substance with a high turn-over rate in mammals and potentially also other non-target organisms. Exogenous FA due to biocidal product use might be a minor contributor to total systemic exposure.

Hence, a final conclusion on the exclusion criteria related to Article 5(1)(d), and on whether RP 3:2 shall be considered a candidate for substitution related to possible ED effect to Article 10(1)(e) is not possible for RP 3:2.

2.2.2. POP criteria

A PBT assessment was performed for RP 3:2 and its hydrolysis products. Based on the available data RP 3:2, 2-hydroxypropylamine and formaldehyde are neither vPvB, nor PBT substances. Furthermore, none of the 3 substances meets two of the PBT criteria. Therefore, neither the parent nor its hydrolysis products meet the criteria for POPs either.

2.2.3. Public consultation for potential candidates for substitution and alternative substances or technologies

As RP 3:2 is considered a candidate for substitution ECHA launched the public consultation in accordance with Article 10(3) of Regulation (EU) No 528/2012. The public consultation took

place from 4 November 2016 to 3 January 2017. Four contributions were submitted: one by an industry stakeholder association, two by individual companies and one by a member state. The same contributions were submitted in the consultation on the structurally and toxicologically related substance RP 1:1.

In the member state contribution it is stated that no information on alternatives is available as the product types are not covered by their national authorisation scheme. This may be the case for more member states.

In the three industry contributions information is submitted on the importance of formaldehyde releasers in the control of microbial growth in water-containing products or equipment. In addition, information on alternatives is submitted for all product types.

Three general observations are made in the industry contributions:

- First, it is stated that other formaldehyde releasers are not considered as alternatives as it can be foreseen that these will also be classified as carcinogen category 1B and subsequently meet the exclusion criteria. In total 10 other formaldehyde releasers are under evaluation and one (formaldehyde released from N,N-Methylenebis(morpholine) or MBM for PT6 and 13) is already approved.
- Second, it is stated that for an effective preservation of many water-based products a bactericide and fungicide is needed. Subsequently, fungicide active substances cannot be regarded as suitable alternatives.
- Last, it is stated that another class of bactericides are the isothiazolinones. Although these are not meeting the substitution criteria it should be considered that these are all classified as strong skin sensitisers. This triggers several obligations for the user making this class of active substances not suitable alternatives.

For PT2 with respect to the use as a system cleaner for metal working fluid systems it is stated in one of the industry contributions that the pH of the system cleaner is 9.5 to 12 to guarantee corrosion protection against steel. This limits the availability of alternatives. The only alternative indicated is glutaraldehyde which is however also a candidate for substitution as it is a respiratory sensitiser.

Several other active substances are already approved for PT2. However, for none of these active substances the intended use was similar to RP 3:2.

The limited information available is insufficient to conclude on the availability of suitable alternatives for the intended uses assessed.

2.3. BPC opinion on the application for approval of the active substance RP 3:2 in product type 2

As the exclusion criteria are met, RP 3:2 should normally not be approved unless one of the conditions for derogation set in Article 5(2) of Regulation (EU) No 528/2012 is met.

If RP 3:2 is approved and included in the Union list of approved active substances, the approval shall be subject to the following specific conditions:

1. Specification:

The active substance has to be considered as substance of Unknown or Variable composition or Complex reaction products (UVC). Therefore the minimum purity is 1000 g/kg (100% by wt).

2. RP 3:2 is considered a candidate for substitution in accordance with Article 10(1)(a) of Regulation (EU) No 528/2012.

3. The authorisations of biocidal products are subject to the following condition:

- a. Products shall only be authorised for use in Member States where at least one of the conditions set in Article 5(2) of Regulation (EU) No 528/2012 is met. Measures shall ensure that exposure of the user and the environment is minimised as far as possible.
- b. The product assessment shall pay particular attention to the exposures, the risks and the efficacy linked to any uses covered by an application for authorisation, but not addressed in the Union level risk assessment of the active substance. In addition, pursuant to point 10 of Annex VI to Regulation (EU) No 528/2012, the product assessment shall include an evaluation as to whether the conditions of Article 5(2) of Regulation (EU) No 528/2012 can be satisfied.
- c. In view of the risks identified for the uses assessed, the product assessment shall pay particular attention to:
 - i. Professionals and industrial workers
 - ii. Sewage Treatment Plant, surface water and the terrestrial compartment

4. The placing on the market of treated articles is subject to the following condition:

The person responsible for the placing on the market of a treated article treated with or incorporating RP 3:2 shall ensure that the label of that treated article provides the information listed in the second subparagraph of Article 58(3) of the Regulation (EU) No 528/2012.

RP 3:2 does not fulfil the criteria according to Article 28(1) to enable inclusion in Annex I of Regulation (EU) 528/2012 as it is classified as Acute Tox. 3, Skin Corr. 1B, Skin Sens. 1A, STOT RE 2, Muta 2, Carc. 1B.

2.4. Elements to be taken into account when authorising products

- 1. The active substance RP 3:2 is considered as a candidate for substitution, and consequently the competent authority shall perform a comparative assessment as part of the evaluation of an application for national authorisation.
- 2. The following recommendations and risk mitigation measures have been identified for the uses assessed. Authorities should consider these risk mitigation measures when authorising products, together with possible other risk mitigation measures, and decide whether these measures are applicable for the concerned product:
 - a. The use of a biocidal product containing RP 3:2 shall be subject to appropriate risk-mitigation measures to ensure that exposure of humans, animals and the environment is minimised as far as possible.

- b. If an unacceptable risk for professional users is identified for the product, safe operational procedures and appropriate organisational measures shall be established. Where exposure cannot be reduced to an acceptable level by other means, products should be used with appropriate personal protective equipment.
- c. Unacceptable risks are identified for the Sewage Treatment Plant, surface water and the terrestrial compartment. If the risk cannot be reduced to an acceptable level by appropriate risk mitigation measures or by other means, these uses should not be authorised.
- d. Surface water intended for the abstraction of drinking water may exceed the parametric value of 0.1 µg/L of the Drinking Water Directive 98/83/EC. This may be considered by the relevant national authorities when issuing permits for recovery plants.

2.5. Requirement for further information

Sufficient data have been provided to verify the conclusions on the active substance, permitting the proposal for the approval of RP 3:2. However, further data shall be required as detailed below. Data must be provided as soon as possible but not later than 6 months before the date of approval to the evaluating Competent Authority (Austria):

- a. Regarding the explosive properties of the active substance an experimental test has to be conducted as the substance contains of unknown constituents and therefore the waiving cannot be justified by structural considerations.
- b. A specific or highly specific and fully validated analytical method for the determination of the hydrolysis product 2-hydroxypropylamin in water.
- c. For the refinement of the environmental risk assessment: monitoring of the concentration of formaldehyde in three representative spent fluids between the last dosing and the emulsion splitting (using techniques based on Kow and via evaporation) and before the biological waste treatment. If further refinement, is necessary an OECD STP simulation test 303 A or 314 should be conducted.