

European Chemicals Agency

Opinion on the administrative change of the Union authorisation of the biocidal product family: SALVECO SALVESAFE PRODUCTS

UAD-C-1717922-28-00/F



Opinion of the European Chemicals Agency

on an administrative change of the Union authorisation of SALVECO SALVESAFE PRODUCTS

In accordance with Article 11(3) of Implementing Regulation (EU) No 354/2013 of the European Commission 18 May 2013 on changes of biocidal products authorised in accordance with Biocidal Product Regulation (EU) No 528/2012 (BPR), the European Chemicals Agency (ECHA) has prepared this opinion on the administrative change to the Union authorisation of:

Name of the biocidal product family: SALVECO SALVESAFE PRODUCTS

Authorisation holder: SALVECO S.A.S.

Target asset number: EU-0028967-0000

Active substance common name: L-(+)-lactic acid

Product types: 2, 3, 4

1. Process for the adoption of the opinion

The notification for the administrative change was submitted to ECHA on 5 February 2024, and recorded in R4BP 3 under case number BC-VH092958-06.

Following its acceptance by ECHA, the evaluation of the notification was initiated on 14 February 2024.

The evaluation included a check that the proposed change of an existing authorisation is of a purely administrative nature involving no change to the properties or efficacy of the biocidal product family in accordance with Article 3(1)aa of the BPR.

The scope of the assessment itself was based on and limited to the information provided by the authorisation holder in the supporting document for the notification for an administrative change of a Union or simplified authorisation under Regulation (EU) No 354/2013 supplied via R4BP 3.

ECHA prepared this opinion containing the conclusions of its assessment.

2. Detailed opinion and background

2.1. ECHA opinion

During the evaluation, ECHA has assessed whether the change made in the SPC document provided by the applicant is administrative in accordance with Implementing Regulation (EU) No 354/2013.

It is ECHA's opinion that the following change to the biocidal product family sought by the authorisation holder is a change falling under Article 3(1)(aa) of the BPR, and, after the implementation of the change, the conditions of Article 19 of the BPR will still be met:



• Title 1, section **2** of the Annex to the Regulation (EU) No 354/2013 - Formulator(s) of the biocidal product - change N° 4: Addition of a formulator of the biocidal product, where the biocidal product composition and the formulating process remain unchanged.

Accordingly, it is proposed that the Commission amends the existing authorisation with the below listed administrative change to the biocidal product family sought by the authorisation holder.

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2.2. ECHA assessment

2.2.1. Description of the changes as proposed by the authorisation holder

Changes as described by the authorisation holder in their supporting document supplied via R4BP 3.

<u>Identification</u>	<u>Description</u>		
1.	Addition of new formulator on the product SALVE FAM2_5 of meta-SPC 4: IDEAL Chimic, Route de St-Julien 34, 1227, Carouge, Switzerland.		

2.2.2. Assessment of the changes as proposed by the authorisation holder

The assessment of the change sought by the authorisation holder is presented in the following table:

Identification	Corresponding reference in the Annex to Regulation (EU) No 354/2013	<u>Evaluation</u>	Result of the evaluation	<u>Comments</u>
1.	Title 1, section 2, change n° 4	The requested change matches the description in the Regulation	Acceptable	



Annex

Draft Summary of Product Characteristics