



SCC GmbH (on behalf of CVAS Development GmbH)  
Am Grenzgraben 11  
55545  
Bad Kreuznach  
Germany

Oslo, 21.01.2019

Your ref.:  
[Your ref.]

Our ref. :  
2019/753

Contact person:  
Hilde Karin Midthaug

## Norwegian Authorisation of the Union Authorised product family Teat disinfectants biocidal product family of CVAS

Regulation (EU) No. 528/2012 concerning the making available on the market and use of biocidal products (the Biocidal Products Regulation, BPR), is implemented in Norwegian law through the Norwegian Biocides Regulation of 18 April 2017 No. 480.

The Norwegian Environment Agency refers to Commission Implementing Regulation (EU) 2018/1853 of 27 November 2018, granting a Union authorisation for the biocidal product family Teat disinfectants biocidal product family of CVAS.

When the Commission grants a Union authorisation or decides that a Union authorisation has not been granted, the EFTA states will, according to the EEA agreement Annex II Chapter XV point 12n (e), simultaneously and within 30 days of the Commission Act take corresponding decisions.

The Norwegian Environment Agency hereby grants authorisation for the biocidal product family Teat disinfectants biocidal product family of CVAS, issued in accordance with the Commission Regulation (EU) 2018/1853 of 27 November 2018, cf. the Norwegian Biocides Regulation § 3-1.

### The authorisation concerns:

Product family name:	Teat disinfectants biocidal product family of CVAS
EU Authorisation number:	EU-0018724-0000
Active substances:	Iodine and Polyvinylpyrrolidone iodine
Product type:	Veterinary hygiene (Disinfectants) - PT 3
Authorisation date:	21.01.19
Expiry date:	30.11.2028

Additionally, the conditions provided in the Norwegian Summary of Product Characteristics (SPC) apply. The SPC is uploaded to R4BP3.

### Label

The information on the label, and, if relevant, in the Material Safety Data Sheet and Technical Data Sheet, shall be in accordance with the conditions provided in the attached SPC. Furthermore, Article 69(1), (2) and Article 70 of the BPR also apply.

The authorisation holder is responsible for ensuring that the information given in the above mentioned documents is accurate, and is translated to Norwegian, cf. Article 69(3) of the BPR.

For each meta-SPC an electronic copy of a representative label from one of the individual products belonging to that meta-SPC with the EU authorisation number EU-0018724-0000, shall be provided. The labels are to be submitted to the Norwegian Environment Agency within three months from the authorisation date, using the email address [biocides@miljodir.no](mailto:biocides@miljodir.no).

#### **Phase-out period of existing stocks, when relevant**

In line with Article 89(4), existing products that do not comply with the conditions of this authorisation, shall not be made available on the market with effect from 180 days after the date of this letter. Furthermore, the use of existing stocks of the biocidal product may continue for up to 365 days after the date of this letter. During this period, all advertising material related to products that not comply with the new conditions, should also be removed from the market.

#### **Yearly fee**

For authorised biocidal products, a yearly fee will be charged. Please see appendix 1B of the Norwegian Biocide Regulation for details.

#### **Registration in the Norwegian Product Register**

All biocidal products must be registered in the Norwegian Product Register. In addition, all biocidal products which are classified as hazardous must be fully declared if they are sold in amounts of 100 kg or more per year. Further information can be found at <http://miljodirektoratet.no/en/Areas-of-activity1/Chemicals/The-Product-Register/>.

Best regards  
Norwegian Environment Agency

*This document has been signed electronically*

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