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Reykjavík 1 November 2018
UST201808-135/H.I.I.
07.06.04

Major Change on request of National Authorisation for the biocidal product family HCl Family A

The Environment Agency of Iceland (Umhverfisstofnun) received your application for National Authorisation - major change on request of HCl Family A on 27th December 2017 (BC-YB036781-38). The case was accepted by the agency 23rd August 2018 and validated on 29th October 2018. The Agency based the evaluation on the application documents as well as the authorisation of the Latvian Environment, Geology and Meteorology Centre (LEGMC).

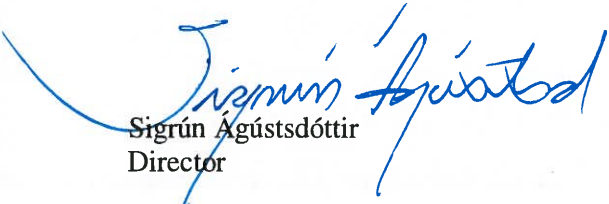
The major change of authorisation for the product family, HCl Family A, with the asset number IS-0016070-0000 is granted in exercise of the powers conferred by Articles 50(2) and (3)(c) of Regulation (EU) No 528/2012 and Article 8 (7) of Regulation (EU) No 354/2013 (on changes) in accordance with Article 1(5) and 1(6) of Icelandic Regulation No 878/2014 on biocidal products, which implements Regulation (EU) No 354/2013 into Icelandic legislation. The Product Assessment Report is accessible under the authorisation in the R4BP 3 database.

We hereby confirm that we accept the notification for a major change on request to account for the change in environmental/ecotoxicological endpoints of the substances, Ethomeen T/12 and Arquad T50, as referred to in Section 1 of Title 3 to the Annex to Regulation (EU) No 354/2013.

After evaluation of submitted changes it has been concluded that Ethomeen T/12 and Arquad T50 do not pose a risk to the environment and reclassification of the biocidal product family is not appropriate.


The Environment Agency of Iceland amends the asset number IS-0016070-0000 in the following terms:

1. The biocidal product family and restrictions outlined in document number UST201610-115 are updated by the relevant conditions and restrictions as listed in Appendix 1 to this certificate.
2. A Summary of the Product Family Characteristics, compliant with Article 22 (2) of Regulation (EU) No 528/2012 is listed in Appendix 1 – the relevant criteria for this biocidal family authorisation applies as described therein.



Sigrún Agústs dóttir
Director

Sincerely



Hafdis Inga Ingvarsdóttir
Hafdis Inga Ingvarsdóttir
Advisor

Appendix 1: Summary of Product Characteristics for a Biocidal Product Family

Appendix 2: Conditions of Authorisation