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Icelandic authorisation of the product family, IPA Family 1, authorised by a Union authorisation

Regulation (EU) No 528/2012 concerning the making available on the market and use of biocidal products, is implemented into Icelandic legislation through the Icelandic Regulation No 878/2014 on biocidal products.

The Environment Agency of Iceland (Umhverfisstofnun) refers to Commission Implementing Regulation (EU) 2023/364 of 16 February 2023, granting a Union authorisation for the biocidal product family, IPA Family 1. 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 Environment Agency of Iceland (Umhverfisstofnun) hereby accepts the European Commission's decision on granting a Union Authorisation for the biocidal [product/product family], [name of product/product family], by publishing a summary of the decision on the Agency's website (ust.is).

When placing the above-mentioned biocidal product family on the market in Iceland, the products shall be labelled according to Article 69 of Regulation (EU) No 528/2012 and if the products are classified as hazardous according to Regulation (EU) No 1272/2008 (CLP), such labelling shall be in Icelandic cf. Article 32 of the Chemicals Act No 61/2013 (see section 6 of the Summary of Product Characteristics (SPC)).

Sincerely

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