

Riga

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COLKIM S.r.l.

Via Piemonte, 50 40064 Ozzano Emilia Italy

On an authorisation of the biocidal product BRODIM PASTA

Latvian Environment, Geology and Meteorology Centre (LEGMC) has evaluated an application submitted by **COLKIM S.R.L.** on 21 September 2018 concerning an authorisation of **BRODIM PASTA** through mutual recognition in sequence.

LEGMC has agreed with Product Assessment Report and Summary of Product Characteristics for **BRODIM PASTA** developed by the reference Member States – Italy.

Therefore, in accordance with Article 33 of Regulation (EU) No 528/2012 of the European Parliament and of the Council of 22 May 2012 concerning the making available on the market and use of biocidal products (Regulation (EU) No 528/2012), LEGMC authorises the **BRODIM PASTA** on the basis of mutual recognition process.

The authorisation holder for **BRODIM PASTA** in Latvia is:

COLKIM S.r.l.

The biocidal product **BRODIM PASTA** contains **0.005%** of *brodifacoum* (CAS No 56073-10-0, EC No 259-980-5) as an active substance.

LEGMC assigns the authorisation number for biocidal product BRODIM PASTA:

LV/2019/MR/013

The authorisation is valid until 31 December 2022.

The authorisation number shall be indicated on the label of the biocidal product.

The authorisation of **BRODIM PASTA** through mutual recognition is granted on the following terms:

- Product type: 14 Rodenticides;
- Target organisms: house mouse juveniles and adults *Mus musculus*, brown rat juveniles and adults *Rattus norvegicus*, roof rat juveniles and adults *Rattus rattus*;
- Users: professional and trained professional;
- Product description: ready to use bait;
- Product stability: shelf life 2 years;
- Use area: indoor and outdoor areas;
- Pack sizes and packaging material: as indicated in Summary of Product Characteristics.



The authorisation through mutual recognition applies only to the product **BRODIM PASTA** in the composition, form and packing for which the authorisation is granted by reference Member State.

The information on the label (and if applicable an enclosed instruction of use) of the **BRODIM PASTA** should be as it is indicated in the authorisation of above mentioned product, taking into account also the information which is stated in the Product Assessment Report and Summary of Product Characteristics issued by reference Member State.

The information on the label shall be in Latvian.

Notwithstanding content of the label specified above, requirements stated in:

- Article 69 Regulation (EU) No 528/2012;
- Regulation (EC) No. 1272/2008 of the European Parliament and of the Council of 16
 December 2008 on classification, labelling and packaging of the substances and
 mixtures, amending and repealing Directives 67/548/EEC and 1999/45/EC, and
 amending Regulation (EC) No 1907/2006;
- all other relevant legislation shall be applied.

COLKIM S.r.l. shall inform LEGMC about any changes in accordance with Commission Implementing Regulation (EU) No 354/2013 of 18th April 2013 on changes of biocidal products authorised in accordance with Regulation (EU) No 528/2012 of the European Parliament and of the Council.

If the first authorisation issued by reference Member State is amended or revoked, the authorisation of **BRODIM PASTA** through mutual recognition may be re-opened for review before 31 December 2022.

Application on renewal of an authorisation shall be submitted according to Commission Delegated Regulation (EU) No 492/2014 of 7 March 2014 supplementing Regulation (EU) No 528/2012 of the European Parliament and of the Council as regards the rules for the renewal of authorisations of biocidal products subject to mutual recognition.

Additionally LEGMC would like to inform that COLKIM S.r.l is fully responsible of the content of the biocidal product **BRODIM PASTA** as well as its classification, labelling, instruction of use and safety data sheet.

LEGMC would like to ask COLKIM S.r.l. to notify the above mentioned information down to supply chain.

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