

Bovilis INtranasal RSP Live nasal spray, lyophilisate and solvent for suspension for cattle

Authorised

- Bovine respiratory syncytial virus, strain Jencine-2013, Live
- Bovine parainfluenza virus 3, strain INT2-2013, Live

Product identification

Medicine name:

Bovilis INtranasal RSP Live nasal spray, lyophilisate and solvent for suspension for cattle

Bovilis INtranasal RSP Live, frostþurrkað nefúðaduft og leysir, dreifa handa nautgripum

Active substance:

Bovine respiratory syncytial virus, strain Jencine-2013, Live

Bovine parainfluenza virus 3, strain INT2-2013, Live

Target species:

Cattle

Route of administration:

Nasal use

Product details

Active substance and strength:

Bovine respiratory syncytial virus, strain Jencine-2013, Live
100000.00 unit(s) / 1.00 Dose

Bovine parainfluenza virus 3, strain INT2-2013, Live
63095.70 unit(s) / 1.00 Dose

Pharmaceutical form:

Nasal spray, suspension

Withdrawal period by route of administration:

Nasal use:

• **Cattle**

- Meat and offal. 0 day

Anatomical therapeutic chemical veterinary (ATCvet) codes:

QI02AD07

Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

Authorisation status:

Valid

Authorised in:

Iceland

Package description:

Cardboard box with 50 doses of lyophilisate and cardboard box with 100 ml of solvent

Cardboard box with 25 doses of lyophilisate and cardboard box with 50 ml of solvent

Cardboard box 5 x 5 doses of lyophilisate and 5 x 10 ml of solvent

Cardboard box 5 x 10 doses of lyophilisate and 5 x 20 ml of solvent

Cardboard box with 20 doses of lyophilisate and cardboard box with 40 ml of solvent

Cardboard box with 1 dose of lyophilisate and 2 ml of solvent.

Cardboard box 5 x 1 dose of lyophilisate and 5 x 2 ml of solvent

Cardboard box 5 doses of lyophilisate and 10 ml of solvent.

Cardboard box 10 doses of lyophilisate and 20 ml of solvent.

Additional information

Entitlement type:

Marketing Authorisation

Legal basis of product authorisation:

Full application - Known active substance (Article 12(3) of Directive No 2001/82/EC)

Marketing authorisation holder:

Intervet International B.V.

Marketing authorisation date:

This information is not available for this product.

Manufacturing sites for batch release:

INTERVET INTERNATIONAL B.V.

Responsible authority:

Icelandic Medicines Agency

Authorisation number:

IS/2/19/009/01

Date of authorisation status change:

17/05/2019

Reference member state:

Netherlands

Procedure number:

NL/V/0257/001

Concerned member states:

Austria Belgium Bulgaria Croatia Cyprus Czechia Denmark Estonia Finland
France Germany Greece Hungary Iceland Ireland Italy Latvia Lithuania
Luxembourg Norway Poland Portugal Romania Slovakia Slovenia Spain
Sweden United Kingdom (Northern Ireland)

To consult adverse reactions on veterinary medicinal products please go to www.adrreports.eu/vet

Documents

Combined File of all Documents

Source URL: <https://medicines.health.europa.eu/veterinary/600000041416>