Ubrostar Red 100 mg / 280 mg / 100 mg Intramammary Suspension for cattle

- Penethamate hydriodide
- Framycetin sulfate
- Benethamine penicillin

Product identification

Medicine name:

Ubrostar Red 100 mg / 280 mg / 100 mg Intramammary Suspension for cattle Ubrostar Dry Cow 100 mg / 280 mg / 100 mg intramamarna suspenzija za krave

Active substance:

Penethamate hydriodide

Framycetin sulfate

Benethamine penicillin

Target species: Cattle

Route of administration: Intramammary use

Product details

Active substance and strength:

Authorised

Penethamate hydriodide 100.00 milligram(s) / 1.00 Syringe Framycetin sulfate 100.00 milligram(s) / 1.00 Syringe Benethamine penicillin 280.00 milligram(s) / 1.00 Syringe

Pharmaceutical form:

Intramammary suspension

Withdrawal period by route of administration:

Intramammary use:

Cattle

- Meat and offal. 10 day
- Milk. 37 day

Anatomical therapeutic chemical veterinary (ATCvet) codes:

QJ51RC25

Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

Authorisation status:

Valid

Authorised in:

Slovenia

Package description:

Cardboard box or plastic container containing 20 single useintramammary syringes and 20 teat wipes (containing isopropanol 70%).Each 4.5 g syringe (cylinder with piston and cap, all made of low densitypolyethylene) contains 5 ml intramammary suspension.

Cardboard box or plastic container containing 60 single useintramammary syringes and 60 teat wipes (containing isopropanol 70%).Each 4.5 g syringe (cylinder with piston and cap, all made of low densitypolyethylene) contains 5 ml intramammary suspension. Cardboard box or plastic container containing 120 single useintramammary syringes and 120 teat wipes (containing isopropanol 70%).Each 4.5 g syringe (cylinder with piston and cap, all made of low densitypolyethylene) contains 5 ml intramammary suspension.

Additional information

Entitlement type: Marketing Authorisation

Legal basis of product authorisation: Generic application (Article 13(1) of Directive No 2001/82/EC)

Marketing authorisation holder:

Boehringer Ingelheim Vetmedica GmbH

Marketing authorisation date:

18/01/2012

Manufacturing sites for batch release:

Lohmann Pharma Herstellung GmbH Haupt Pharma Latina S.r.l.

Responsible authority:

Agency For Medicinal Products And Medical Devices Of The Republic Of Slovenia

Authorisation number: DC/V/0374/001

Date of authorisation status change:

18/01/2012

Reference member state:

Ireland

Procedure number: IE/V/0271/001

Concerned member states:

Austria Belgium Bulgaria Cyprus Czechia Estonia France Germany Greece Hungary Latvia Lithuania Netherlands Poland Portugal Romania Slovakia Slovenia Spain United Kingdom (Northern Ireland)

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

Documents

Summary of Product Characteristics

English (PDF) Published on: 11/02/2022 Download

Package Leaflet

This document does not exist in this language (English). You can find it in another language below.

Labelling

This document does not exist in this language (English). You can find it in another language below.

Source URL: https://medicines.health.europa.eu/veterinary/60000052236