File downloaded on 2025-12-21

Source URL: https://medicines.health.europa.eu/veterinary/en/60000005520

Parofor 140 mg/ml Solution for use in drinking water/milk

Authorised

• Paromomycin sulfate

Product identification

Medicine name:

Parofor 140 mg/ml raztopina za uporabo v vodi za pitje, mleku ali mlečnem nadomestku za teleta, ki še ne prežvekujejo in prašiče Parofor 140 mg/ml Solution for use in drinking water/milk

Active substance:

Paromomycin sulfate

Target species:

Cattle

Pig

Route of administration:

In drinking water/milk use

Product details

Active substance and strength:

Paromomycin sulfate 200.00 milligram(s) / 1.00 millilitre(s)

Pharmaceutical form:

Solution for use in drinking water/milk

Withdrawal period by route of administration:

In drinking water/milk use:

Cattle

- Meat and offal. 20 day 20 days for pre-ruminant cattle

Pig

- Meat and offal. 3 day 3 days

Anatomical therapeutic chemical veterinary (ATCvet) codes:

QA07AA06

Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

Authorisation status:

Valid

Authorised in:

Slovenia

Available in:

Slovenia

Package description:

Parofor 140 mg/ml sol. for drinking water/milk 1000 ml

Parofor 140 mg/ml sol. for drinking water/milk 500 ml

Parofor 140 mg/ml sol. for drinking water/milk 250 ml

Parofor 140 mg/ml sol. for drinking water/milk 125 ml

Additional information

Entitlement type:

Marketing Authorisation

Legal basis of product authorisation:

Hybrid application – change in strength (Article 19(1)(a) of Regulation (EU) 2019/6)

Marketing authorisation holder:

HuVepharma

Marketing authorisation date:

27/07/2017

Manufacturing sites for batch release:

Biovet AD

Responsible authority:

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

Authorisation number:

DC/V/0480/002

Date of authorisation status change:

27/07/2017

Reference member state:

Belgium

Procedure number:

BE/V/0027/002

Concerned member states:

Austria Bulgaria Cyprus Czechia Denmark Estonia France Germany Greece Hungary Ireland Italy Latvia Lithuania Luxembourg Malta 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
This document does not exist in this language (English). You can find it in another language below.
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.
Combined File of all Documents