Tulissin 100 mg/ml - Solution for injection (cattle, pigs, sheep)

Authorised

• Tulathromycin

Product identification

Medicine name:

Tulissin 100 mg/ml - Solution for injection (cattle, pigs, sheep)

Active substance:

Tulathromycin

Target species:

Cattle

Sheep

Pig

Route of administration:

Intramuscular use

Subcutaneous use

Product details

Active substance and strength:

Tulathromycin 100.00 milligram(s) / 1.00 Vial

Pharmaceutical form:

Solution for injection

Withdrawal period by route of administration: Intramuscular use:

•

Cattle

- Meat and offal. 22 day 22 days

•

Sheep

- Meat and offal. 16 day 16 days

•

Pig

- Meat and offal. 13 day 13 days

Subcutaneous use:

•

Cattle

- Meat and offal. 22 day 22 days

•

Sheep

- Meat and offal. 16 day 16 days

•

Pig

- Meat and offal. 13 day 13 days

Anatomical therapeutic chemical veterinary (ATCvet) codes:

QJ01FA94

Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

Authorisation status:

Valid

Authorised in:

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

Available in:

Austria, Cyprus, Denmark, Estonia, Finland, France, Greece, Hungary, Ireland, Italy, Latvia, Lithuania, Portugal, Romania, United Kingdom (Northern Ireland)

Package description:

Packaging: Vial (glass) with protective sleeve, Package size: 1 vial, Content: 250 ml

Packaging: Vial (glass), Package size:1 vial, Content:100 ml

Packaging: Vial (glass), Package size: 1 vial, Content: 250 ml

Packaging: Vial (glass), Package size: 1 vial, Content: 50 ml

Packaging: Vial (glass), Package size: 1 vial, Content: 20 ml

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:

Virbac S.A.

Marketing authorisation date:

24/04/2020

Manufacturing sites for batch release:

Virbac

Fareva Amboise

Responsible authority:

European Commission

Authorisation number:

This information is not available for this product.

Date of authorisation status change:

24/04/2020

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

Documents

Combined File of all Documents

English (PDF)

Published on: 18/04/2024

Download

ema-puar-tulissin-v-5073-par-en.pdf

Source URL: https://medicines.health.europa.eu/veterinary/600000004023