PRONESTESIC 40 mg/ml + 0.036 mg/ml solution for injection for horses, cattle, swine and sheep

Authorised

- ADRENALINE TARTRATE PH. EUR.
- Procaine hydrochloride

Product identification

Medicine name:

PRONESTESIC 40 mg/ml + 0.036 mg/ml solution for injection for horses, cattle, swine and sheep

Pronestesic 40 mg/ml + 0,036 mg/ml Injektionslösung für Pferde, Rinder, Schweine und Schafe

Active substance:

ADRENALINE TARTRATE PH. EUR.

Procaine hydrochloride

Target species:

Cattle

Sheep

Pig

Horse

Route of administration:

Perineural use

Subcutaneous use

Product details

Active substance and strength:

ADRENALINE TARTRATE PH. EUR. 0.04 milligram(s) / 1.00 millilitre(s)

Procaine hydrochloride 40.00 milligram(s) / 1.00 millilitre(s)

Pharmaceutical form:

Solution for injection

Withdrawal period by route of administration:

Perineural use:

- . Cattle
 - Meat and offal. 0 day
 - Milk. 0 day
- Sheep
 - Meat and offal. 0 day
 - Milk. 0 day
- Pig
 - Meat and offal. 0 day
- Horse
 - Meat and offal. 0 day
 - Milk. 0 day

Subcutaneous use:

- . Cattle
 - Meat and offal. 0 day
 - Milk. 0 day
- . Sheep
 - Meat and offal. 0 day
 - Milk. 0 day
- . Pig

- Meat and offal. 0 day
- . Horse
 - Meat and offal. 0 day
 - Milk. 0 day

Anatomical therapeutic chemical veterinary (ATCvet) codes:

ON01BA52

Legal status of supply:

Medicinal product on medical prescription for renewable delivery

Authorisation status:

Valid

Authorised in:

Austria

Available in:

Austria

Package description:

box containing 10 vials of 100 ml

box containing 1 vial of 250 ml

box containing 1 vial of 100 ml

box containing 1 vial of 50 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:

Fatro S.p.A.

Marketing authorisation date:

This information is not available for this product.
Manufacturing sites for batch release: Fatro S.p.A.
Responsible authority: Austrian Agency For Health And Food Safety
Authorisation number: 836934
Date of authorisation status change: 24/02/2021
Reference member state: Spain
Procedure number: ES/V/0238/001
Concerned member states: Austria Belgium Croatia Cyprus Czechia Denmark Estonia Finland France Germany Greece Iceland Ireland Italy Latvia Lithuania Luxembourg Netherlands Norway Poland Portugal Slovakia Slovenia Sweden United Kingdom (Northern Ireland)
To consult adverse reactions on veterinary medicinal products please go to www.adrreports.eu/vet
Documents
Package Leaflet
English (PDF)

Published o	on:	25/12/2023
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Summary of Product Characteristics

English (PDF)

Published on: 22/12/2023

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Labelling

English (PDF)

Published on: 25/12/2023

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