

Kaltetan forte, 458.4 mg/ml + 125 mg/ml + 20 mg/ml solution for infusion for horses, cattle and pigs

- CALCIUM GLUCONATE FOR INJECTION
- Magnesium chloride hexahydrate
- Sodium glycerophosphate pentahydrate

Product identification

Medicine name:

Kaltetan forte, 458.4 mg/ml + 125 mg/ml + 20 mg/ml solution for infusion for horses, cattle and pigs

Kaltetan forte, 458,4 mg/ml + 125 mg/ml + 20 mg/ml, infuzinis tirpalas arkliams, galvijams ir kiaulėms

Active substance:

CALCIUM GLUCONATE FOR INJECTION

Magnesium chloride hexahydrate

Sodium glycerophosphate pentahydrate

Target species:

Horse

Cattle

Piq

Route of administration:

Product details

Active substance and strength:

CALCIUM GLUCONATE FOR INJECTION 458.40 milligram(s) / 1.00 millilitre(s)

Magnesium chloride hexahydrate 125.00 milligram(s) / 1.00 millilitre(s)

Sodium glycerophosphate pentahydrate 20.00 milligram(s) / 1.00 millilitre(s)

Pharmaceutical form:

Solution for infusion

Withdrawal period by route of administration: Intravenous use:

Horse

- Milk. 5 week Meat and offal: Zero days. Milk: Zero hours

Cattle

- Meat and offal. 5 week Meat and offal: Zero days. Milk: Zero hours

Pig

- Meat and offal. 5 week Meat and offal: Zero days

Anatomical therapeutic chemical veterinary (ATCvet) codes:

QA12AX

Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

| Authorisation status: Valid |
|---------------------------------------------------------------------------------------------------------------------------------------------------------|
| Authorised in: Lithuania |
| Available in: Lithuania |
| Package description: Polypropylene (PP) bottles closed with a bromobutyl rubber stopper, type I and secured with an aluminium cap. Package size: 500 ml |
| Additional information |
| Entitlement type: Marketing Authorisation |
| Legal basis of product authorisation: Hybrid application (Article 13(3) of Directive No 2001/82/EC) |
| Marketing authorisation holder: Vet-Agro Multi-Trade Company Sp. z o.o. |
| Marketing authorisation date: 25/01/2022 |
| Manufacturing sites for batch release: Przedsiebiorstwo Wielobranzowe Vet-Agro Sp. z o.o. |
| Responsible authority: State Food And Veterinary Service |
| Authorisation number: LT/2/22/2697/001 |
| Date of authorisation status change: 25/01/2022 |

| Reference member state: Poland |
|--------------------------------------------------------------------------------------------------|
| Procedure number: PL/V/0110/002 |
| Concerned member states: Bulgaria Czechia Greece Hungary Italy Lithuania Portugal Romania Spain |
| To consult adverse reactions on veterinary medicinal products please go to www.adrreports.eu/vet |
| Documents |

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RV2697.pdf