

# Calcibel 240/60/60 mg/ml solution for infusion for horses, cattle, sheep, goats and pigs

Authorised

- Boric acid
- Calcium gluconate
- Magnesium chloride hexahydrate

## Product identification

### Medicine name:

Calcibel 240/60/60 mg/ml solution for infusion for horses, cattle, sheep, goats and pigs

Calcibel, 240/60/60 mg/mL, otopina za infuziju, za konje, goveda, ovce, koze i svinje

### Active substance:

Boric acid

Calcium gluconate

Magnesium chloride hexahydrate

### Target species:

Cattle

Goat (adult female)

Sheep

Horse

Pig

### Route of administration:

Intravenous use

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## Product details

### Active substance and strength:

Boric acid

60.00 milligram(s) / 1.00 millilitre(s)

Calcium gluconate

240.00 milligram(s) / 1.00 millilitre(s)

Magnesium chloride hexahydrate

60.00 milligram(s) / 1.00 millilitre(s)

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### Pharmaceutical form:

Solution for infusion

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### Anatomical therapeutic chemical veterinary (ATCvet) codes:

QA12AX

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### Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

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### Authorisation status:

Valid

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### Authorised in:

Croatia

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### Available in:

Croatia

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### Package description:

Bottle for infusion made of polypropylene with scaling, closures made of bromobutyl rubber stopper, aluminium caps with tear-off. Pack size 6 x 500 ml.

Bottle for infusion made of polypropylene with scaling, closures made of bromobutyl rubber stopper, aluminium caps with tear-off. Pack size 12 x 500 ml.

Bottle for infusion made of polypropylene with scaling, closures made of bromobutyl rubber stopper, aluminium caps with tear-off. Pack size 1 x 500 ml.

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## Additional information

**Entitlement type:**

Marketing Authorisation

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**Legal basis of product authorisation:**

Hybrid application (Article 13(3) of Directive No 2001/82/EC)

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**Marketing authorisation holder:**

Bela-Pharm GmbH & Co. KG

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**Marketing authorisation date:**

27/10/2016

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**Manufacturing sites for batch release:**

Bela-Pharm GmbH & Co. KG

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**Responsible authority:**

Ministry Of Agriculture Veterinary And Food Safety Directorate

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**Authorisation number:**

UP/I-322-05/21-01/703

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**Date of authorisation status change:**

25/06/2025

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**Reference member state:**

Netherlands

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**Procedure number:**

NL/V/0197/001

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**Concerned member states:**

Austria Croatia Cyprus Czechia Denmark Finland Greece Hungary Ireland  
Poland Portugal Romania Slovakia Slovenia Spain Sweden  
United Kingdom (Northern Ireland)

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## Documents

Summary of Product Characteristics

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