

# GABBROVET MULTI 140 MG/ML SOLUTION FOR USE IN DRINKING WATER/ MILK FOR PRE-RUMINANT CATTLE AND PIGS

Authorised

- Paromomycin sulfate

## Product identification

**Medicine name:**

GABBROVET MULTI 140 MG/ML SOLUTION FOR USE IN DRINKING WATER/ MILK FOR PRE-RUMINANT CATTLE AND PIGS

Gabbrovet Multi 140 mg/ml oplossing voor gebruik in drinkwater/melk voor niet-herkauwende kalveren en varkens

**Active substance:**

Paromomycin sulfate

**Target species:**

Cattle (calf)

Pig

**Route of administration:**

Oral use

## Product details

**Active substance and strength:**

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

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**Pharmaceutical form:**

Solution for use in drinking water/milk

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**Withdrawal period by route of administration:****Oral use:**

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**Cattle (calf)**

- Meat and offal. 110 day

Cryptosporidiosis: Dosage: 150 mg/kg/day for 5 days.

- Meat and offal. 20 day

Colibacillosis: Dosage: 25-50 mg/kg/day for 3 to 5 days.

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**Pig**

- Meat and offal. 3 day

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

QA07AA06

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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:**

Netherlands

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**Package description:**

Cardboard box containing 1 bottle of 250 ml (HDPE/EVOH/HDPE) containing a dosing device of 30 ml

Cardboard box containing 1 bottle of 500 ml (HDPE/EVOH/HDPE) containing a dosing device of 30 ml

Cardboard box containing 1 bottle of 1 L (HDPE/EVOH/HDPE) containing a dosing device of 30 ml

Cardboard box containing 1 bottle of 125 ml (HDPE) containing a dosing device of 30 ml

Cardboard box containing 1 bottle of 250 ml (HDPE) containing a dosing device of 30 ml

Cardboard box containing 1 bottle of 500 ml (HDPE) containing a dosing device of 30 ml

Cardboard box containing 1 bottle of 1 L (HDPE) containing a dosing device of 30 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:**

Ceva Sante Animale

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**Marketing authorisation date:**

19/07/2022

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**Manufacturing sites for batch release:**

Ceva Sante Animale

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**Responsible authority:**

Medicines Evaluation Board

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**Authorisation number:**

REG NL 128521

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**Date of authorisation status change:**

28/03/2024

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**Reference member state:**

France

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**Procedure number:**

FR/V/0429/001

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**Concerned member states:**

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

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To consult adverse reactions on veterinary medicinal products please go to  
[www.adrreports.eu/vet](http://www.adrreports.eu/vet)

## Documents

Combined File of all Documents

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