Source URL: https://medicines.health.europa.eu/veterinary/en/700000168105

Nanotrim 464.2 mg/g + 100 mg/g powder for use in drinking water/milk for chickens, turkeys, pigs and cattle



- Sulfachlorpyridazine sodium
- Trimethoprim

Product identification

Medicine name:

Nanotrim 464.2 mg/g + 100 mg/g powder for use in drinking water/milk for chickens, turkeys, pigs and cattle

Nanotrim, 464,2 mg/g + 100 mg/g, milteliai, skirti naudoti su geriamuoju vandeniu ar pienu vištoms, kalakutams, kiaulėms ir galvijams

Active substance:

Sulfachlorpyridazine sodium

Trimethoprim

Target species:

Pig

Chicken

Turkey

Cattle (pre-ruminant)

Route of administration:

Product details

Active substance and strength:

Sulfachlorpyridazine sodium 500.00 milligram(s)/gram / 1.00 gram(s)

Trimethoprim

100.00 milligram(s)/gram / 1.00 gram(s)

Pharmaceutical form:

Powder for use in drinking water/milk

Withdrawal period by route of administration:

In drinking water/milk use:

Pig

- Meat and offal. 7 day

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Chicken

- Meat and offal. 3 day

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Turkey

- Meat and offal. 9 day

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Cattle (pre-ruminant)

- Meat and offal. 7 day

Anatomical therapeutic chemical veterinary (ATCvet) codes:

QJ01EW12

Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

Authorisation status: Valid
Authorised in: Lithuania
Package description: Pillow sachet with 100 g made of polyethylene/aluminium/polyethylene terephthalate laminate 1k g resealable block-bottom zipped sachet made of polyethylene/aluminium/polyethylene terephthalate laminate
Additional information Entitlement type: Marketing Authorisation
Legal basis of product authorisation: Hybrid application (Article 19(1) of Regulation (EU) 2019/6)
Marketing authorisation holder: HuVepharma
Marketing authorisation date: 3/02/2025
Manufacturing sites for batch release: Biovet AD
Responsible authority: State Food And Veterinary Service

Authorisation number:

LT/2/25/2860/001-002

Date of authorisation status change:

25/09/2025

Reference member state:

Netherlands

Procedure number:

NL/V/0420/001

Concerned member states:

Austria Bulgaria Croatia Cyprus Czechia Denmark Estonia Greece Hungary Iceland Ireland Italy Latvia Lithuania Luxembourg Malta Poland Portugal Romania Slovakia Slovenia Spain

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

Documents

Combined File of all Documents

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