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

Authorised

- 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, 500+100mg/g, Prášek pro podání v pitné vodě/mléce

## **Active substance:**

Sulfachlorpyridazine sodium Trimethoprim

## **Target species:**

Pig Chicken Turkey Cattle (pre-ruminant)

# Route of administration:

In drinking water/milk use

# **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

•

# Chicken

- Meat and offal. 3 day

# Turkey

- Meat and offal. 9 day

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:

Czechia

# 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:

24/03/2025

## Manufacturing sites for batch release:

Biovet AD

## **Responsible authority:**

Institute For State Control Of Veterinary Biologicals And Medicaments

# Authorisation number:

96/008/25-C

# Date of authorisation status change:

24/03/2025

## **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

**Source URL:** https://medicines.health.europa.eu/veterinary/700000168095