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# Ubrolexin intramammary suspension for lactating dairy cows

Authorised

- Cefalexin monohydrate
- KANAMYCIN MONOSULPHATE

# Product identification

#### **Medicine name:**

Ubrolexin intramammary suspension for lactating dairy cows
UBROLEXIN SUSPENSION INTRAMAMMAIRE POUR VACHES LAITIERES EN LACTATION

#### **Active substance:**

Cefalexin monohydrate

KANAMYCIN MONOSULPHATE

#### **Target species:**

Cattle

#### Route of administration:

Intramammary use

# **Product details**

# **Active substance and strength:**

Cefalexin monohydrate

210.36 milligram(s) / 1.00 Syringe

KANAMYCIN MONOSULPHATE

120247.00 international unit(s) / 1.00 Syringe

#### **Pharmaceutical form:**

Intramammary suspension

# Withdrawal period by route of administration: Intramammary use:

## Cattle

- Meat and offal. 10 day
- Milk. 5 day

# Anatomical therapeutic chemical veterinary (ATCvet) codes:

QJ51RD01

#### Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

#### **Authorisation status:**

Valid

#### Authorised in:

France

#### Available in:

France

#### Package description:

Cardboard box with 20 single use intramammary syringes and 20 teat wipes (containing isopropanol 70%). Each 10 g syringe contains 12 ml intramammary suspension and consists of a barrel with plunger and sealed sterile tip, all made of low density polyethylene.

Cardboard box with 10 single use intramammary syringes and 10 teat wipes (containing isopropanol 70%). Each 10 g syringe contains 12 ml intramammary suspension and consists of a barrel with plunger and sealed sterile tip, all made of low density polyethylene.

# Additional information

#### **Entitlement type:**

Marketing Authorisation

#### Legal basis of product authorisation:

Well-established use application (Article 13a of Directive No 2001/82/EC)

### Marketing authorisation holder:

Boehringer Ingelheim Animal Health France

#### Marketing authorisation date:

4/08/2008

#### Manufacturing sites for batch release:

**Univet Limited** 

#### **Responsible authority:**

French Agency For Food, Environmental And Occupational Health & Safety

#### **Authorisation number:**

FR/V/1592221 0/2008

# Date of authorisation status change:

4/08/2013

#### **Reference member state:**

Ireland

#### **Procedure number:**

IE/V/0221/001

#### **Concerned member states:**

Austria Belgium Cyprus Czechia Estonia France Germany Greece Hungary Italy Latvia Lithuania Luxembourg Netherlands Poland Portugal Romania Slovakia Slovenia Spain United Kingdom (Northern Ireland) To consult adverse reactions on veterinary medicinal products please go to www.adrreports.eu/vet

# **Documents**

Summary of Product Characteristics

English (PDF)

Published on: 28/09/2025

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Combined File of all Documents

Package Leaflet and Labelling

This document does not exist in this language (English). You can find it in another language below.