

VIRBACTAN 150 MG INTRAMAMMARY OINTMENT

Authorised

- Cefquinome sulfate

Product identification

Medicine name:

VIRBACTAN 150 MG INTRAMAMMARY OINTMENT

VIRBACTAN 150 mg pomata per uso intramammario

Active substance:

Cefquinome sulfate

Target species:

Cattle (dry cow)

Route of administration:

Intramammary use

Product details

Active substance and strength:

Cefquinome sulfate

177.80 milligram(s) / 1.00 Syringe

Pharmaceutical form:

Intramammary ointment

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

•

Cattle (dry cow)

- Meat and offal. 2 day
- Milk. 36 day

Milk: 36 days after treatment when dry period is 5 weeks or less.

- Milk. 1 day

Milk: 1 day after calving when dry period is more than 5 weeks

Anatomical therapeutic chemical veterinary (ATCvet) codes:

QJ51DE90

Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

Authorisation status:

Valid

Authorised in:

Italy

Available in:

Italy

Package description:

Box of 1 sachet of 4 applicators and 4 cleaning towels
Box of 30 sachets of 4 applicators and 120 cleaning towels
Box of 15 sachets of 4 applicators and 60 cleaning towels
Box of 6 sachets of 4 applicators and 24 cleaning towels
Box of 5 sachets of 4 applicators and 20 cleaning towels

Additional information**Entitlement type:**

Marketing Authorisation

Legal basis of product authorisation:

Full application - Known active substance (Article 12(3) of Directive No 2001/82/EC)

Marketing authorisation holder:

Virbac

Marketing authorisation date:

25/02/2005

Manufacturing sites for batch release:

Haupt Pharma Latina S.r.l.

Virbac

Responsible authority:

Ministry Of Health

Authorisation number:

103706

Date of authorisation status change:

28/02/2005

Reference member state:

France

Procedure number:

FR/V/0148/001

Concerned member states:

Austria Belgium Cyprus Czechia Estonia Germany Greece Ireland Italy

Latvia Lithuania Luxembourg Netherlands Poland Portugal Slovakia Slovenia

Spain United Kingdom (Northern Ireland)

To consult adverse reactions on veterinary medicinal products please go to

www.adrreports.eu/vet

Documents

Combined File of all Documents

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