Nuflor Minidose 450 mg/ml solution for injection for cattle

• Florfenicol

Product identification

Medicine name:

Nuflor Minidose 450 mg/ml solution for injection for cattle NUFLOR 450 SOLUTION INJECTABLE POUR BOVINS

Active substance:

Florfenicol

Target species:

Cattle

Route of administration: Subcutaneous use Intramuscular use

Product details

Active substance and strength:

Florfenicol 450.00 milligram(s) / 1.00 millilitre(s)

Pharmaceutical form:

Solution for injection

Withdrawal period by route of administration: Subcutaneous use: • Cattle

- Milk. no withdrawal period

Not permitted for use in lactating animals producing milk for human consumption.

- Meat and offal. 64 day

Intramuscular use:

Cattle

- Milk. no withdrawal period

Not permitted for use in lactating animals producing milk for human consumption.

- Meat and offal. 37 day

Anatomical therapeutic chemical veterinary (ATCvet) codes:

QJ01BA90

Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

Authorisation status:

Valid

Authorised in:

France

Package description:

(ID3) 250 millilitre(s): unspecified outer container with 1 Vial (Glass) with 250 millilitre(s), closed with Stopper (bromobutyl rubber`)
(ID2) 100 millilitre(s): unspecified outer container with 1 Vial (Glass) with 100 millilitre(s), closed with Stopper (bromobutyl rubber`)
(ID1) 50 millilitre(s): unspecified outer container with 1 Vial (Glass) with 50 millilitre(s), closed with Stopper (bromobutyl rubber`)

Additional information

Entitlement type:

Marketing Authorisation

Legal basis of product authorisation:

Generic application (Article 13(1) of Directive No 2001/82/EC)

Marketing authorisation holder: Intervet International B.V.

Marketing authorisation date:

17/09/2008

Manufacturing sites for batch release:

Intervet International GmbH Trirx Segre Vet Pharma Friesoythe GmbH

Responsible authority:

French Agency For Food, Environmental And Occupational Health & Safety

Authorisation number: FR/V/7051353 4/2008

Date of authorisation status change:

17/09/2013

Reference member state:

Germany

Procedure number: DE/V/0122/001

Concerned member states:

Belgium Bulgaria Cyprus Czechia Denmark Finland France Greece Hungary Ireland Italy Luxembourg Netherlands Portugal Romania 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: 28/01/2022 <u>Download</u>

Combined File of all Documents

Source URL: https://medicines.health.europa.eu/veterinary/60000060539