Macromectin 0.5% w/v Pour-On Solution

Authorised

Ivermectin

Product identification

Medicine name:

Macromectin 0.5% w/v Pour-On Solution BAYMEC 0,5% SOLUTION POUR-ON

Active substance:

Ivermectin

Target species:

Cattle

Route of administration:

Topical use

Product details

Active substance and strength:

Ivermectin

5.00 milligram(s) / 1.00 millilitre(s)

Pharmaceutical form:

Pour-on solution

Withdrawal period by route of administration:

Topical use:

- . Cattle
 - Meat and offal. 28 day

Anatomical therapeutic chemical veterinary (ATCvet) codes:

QP54AA01

Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

Authorisation status:

Valid

Authorised in:

France

Package description:

The product will be supplied in: 250 ml twin-neck high density polyethylene dispensers

The product will be supplied in:1.0 L single-neck high density polyethylene dispensers

The product will be supplied in:2.5 L low density polyethylene backpacks.

The product will be supplied in:5 L low density polyethylene backpacks.

The product will be supplied in: 250 ml squeeze-measure high density polyethylene dispensers

The product will be supplied in:1.0 L squeeze-measure high density polyethylene dispensers

The product will be supplied in:250 ml single-neck high density polyethylene dispensers

The product will be supplied in:1 L high density polyethylene backpacks.

The product will be supplied in: 1.0 L twin-neck high density polyethylene dispensers

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: Norbrook Laboratories (Ireland) Limited | |
|---|--|
| Marketing authorisation date: 11/05/2006 | |
| Manufacturing sites for batch release: Norbrook Laboratories Limited Norbrook Manufacturing Limited | |
| Responsible authority: French Agency For Food, Environmental And Occupational Health & Safety | |
| Authorisation number: FR/V/0681919 0/2006 | |
| Date of authorisation status change: 11/05/2011 | |
| Reference member state: Ireland | |
| Procedure number: IE/V/0180/001 | |
| Concerned member states: | |

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

Documents

France

Summary of Product Characteristics

English (PDF)

Published on: 11/02/2022

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