

LEVASOL 100 mg/ml

Authorised

- Levamisole hydrochloride

Product identification

Medicine name:

LEVASOL 100 mg/ml

Active substance:

Levamisole hydrochloride

Target species:

Cattle

Sheep

Pig

Chicken (broiler)

Turkey

Duck

Homing pigeon

Route of administration:

Oral use

Product details

Active substance and strength:

Levamisole hydrochloride

100.00 milligram(s) / 1.00 millilitre(s)

Pharmaceutical form:

Withdrawal period by route of administration:

Oral use:

-

Cattle

- Meat and offal. 11 day

NU este permisă utilizarea la animalele în lactație care produc lapte pentru consum uman.

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Sheep

- Meat and offal. 11 day

NU este permisă utilizarea la animalele în lactație care produc lapte pentru consum uman.

-

Pig

- Meat and offal. 10 day

-

Chicken (broiler)

- Meat and offal. 4 day

NU este permisă utilizarea la păsările ouătoare care produc ouă pentru consum uman.

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Turkey

- Meat and offal. 4 day

NU este permisă utilizarea la păsările ouătoare care produc ouă pentru consum uman.

-

Duck

- Meat and offal. 4 day

NU este permisă utilizarea la păsările ouătoare care produc ouă pentru consum uman.

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Homing pigeon

Anatomical therapeutic chemical veterinary (ATCvet) codes:

QP52AE01

Legal status of supply:

This information is not available for this product.

Authorisation status:

Valid

Authorised in:

Romania

Available in:

Romania

Package description:

Available only in Romanian

Available only in Romanian

Available only in Romanian

Available only in Romanian

Available only in Romanian

Available only in Romanian

Available only in Romanian

Available only in Romanian

Available only in Romanian

Additional information

Entitlement type:

Marketing Authorisation

Legal basis of product authorisation:

Complete application (stand-alone) - Directive No 2001/82/EC

Marketing authorisation holder:

Pasteur Filiala Filipesti S.A.

Marketing authorisation date:

26/07/2018

Manufacturing sites for batch release:

Pasteur Filiala Filipesti S.A.

Responsible authority:

Institute For Control Of Biological Products And Veterinary Medicines

Authorisation number:

230133

Date of authorisation status change:

26/07/2023

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

Documents

Summary of Product Characteristics

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