

# NEWVAC LA SOTA

Authorised

- Newcastle disease virus, strain La Sota, Live

## Product identification

**Medicine name:**

NEWVAC LA SOTA

**Active substance:**

Newcastle disease virus, strain La Sota, Live

**Target species:**

Chicken (hen)

**Route of administration:**

Ocular use

In drinking water use

Nebulisation use

Subcutaneous use

## Product details

**Active substance and strength:**

Newcastle disease virus, strain La Sota, Live

6.50 log 10 50% embryo infective dose / 0.03 millilitre(s)

**Pharmaceutical form:**

This information is not available for this product.

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

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**Chicken (hen)**

- Meat and offal. 0 day
- Egg. 0 day

**In drinking water use:**

- 

**Chicken (hen)**

- Meat and offal. 0 day
- Egg. 0 day

**Nebulisation use:**

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**Chicken (hen)**

- Meat and offal. 0 day
- Egg. 0 day

**Subcutaneous use:**

- 

**Chicken (hen)**

- Meat and offal. 0 day
- Egg. 0 day

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**Anatomical therapeutic chemical veterinary (ATCvet) codes:**

QI01AD06

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**Legal status of supply:**

This information is not available for this product.

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**Authorisation status:**

Valid

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**Authorised in:**

Romania

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**Available in:**

Romania

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**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

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## Additional information

**Entitlement type:**

Marketing Authorisation

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**Legal basis of product authorisation:**

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

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**Marketing authorisation holder:**

Pasteur Filiala Filipesti S.A.

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**Marketing authorisation date:**

8/08/2005

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**Manufacturing sites for batch release:**

Pasteur Filiala Filipesti S.A.

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**Responsible authority:**

Institute For Control Of Biological Products And Veterinary Medicines

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**Authorisation number:**

150485

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**Date of authorisation status change:**

11/04/2022

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To consult adverse reactions on veterinary medicinal products please go to [www.adrreports.eu/vet](http://www.adrreports.eu/vet)

## Documents

Summary of Product Characteristics

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