Selames 120 mg spot-on solution for dogs 10.1–20.0 kg



• Selamectin

Product identification

Medicine name:

Selames 120 mg spot-on solution for dogs 10.1–20.0 kg Selames 120 mg spot-on oplossing voor honden 10,1 – 20,0 kg

Active substance:

Selamectin

Target species:

Dog

Route of administration:

Spot-on use

Product details

Active substance and strength:

Selamectin 120.00 milligram(s) / 1.00 Pipette

Pharmaceutical form:

Spot-on solution

Withdrawal period by route of administration:

Spot-on use:

Dog

Anatomical therapeutic chemical veterinary (ATCvet) codes:

QP54AA05

Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

Authorisation status:

Valid

Authorised in:

Netherlands

Package description:

Translucent polypropylene unit-dose pipette with polyethylene or polyoxymethylene or polypropylene closure with spikepackaged into a laminated triplex bag composed of polyester, aluminium and polyethylene.Cardboard box containing 1 pipette. Translucent polypropylene unit-dose pipette with polyethylene or polyoxymethylene or polypropylene closure with spikepackaged into a laminated triplex bag composed of polyester, aluminium and polyethylene.Cardboard box containing 3 pipettes. Translucent polypropylene unit-dose pipette with polyethylene or polyoxymethylene or polypropylene closure with spikepackaged into a laminated triplex bag composed of polyester, aluminium and polyethylene.Cardboard box containing 6 pipettes. Translucent polypropylene unit-dose pipette with polyethylene or polyoxymethylene or polypropylene closure with spikepackaged into a laminated triplex bag composed of polyester, aluminium and polyethylene.Cardboard box containing 6 pipettes. Translucent polypropylene unit-dose pipette with polyethylene or polyoxymethylene or polypropylene closure with spikepackaged into a laminated triplex bag composed of polyester, aluminium and polyethylene.Cardboard box containing 15 pipettes.

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:

KRKA tovarna zdravil d.d. Novo mesto

Marketing authorisation date:

20/11/2018

Manufacturing sites for batch release:

TAD Pharma GmbH KRKA tovarna zdravil d.d. Novo mesto

Responsible authority:

Medicines Evaluation Board

Authorisation number: REG NL 122211

Date of authorisation status change:

26/01/2022

Reference member state:

Ireland

Procedure number: IE/V/0407/003

Concerned member states:

Belgium France Germany Italy Netherlands Portugal Spain

United Kingdom (Northern Ireland)

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

Documents

Combined File of all Documents

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

Summary of Product Characteristics

Source URL: https://medicines.health.europa.eu/veterinary/60000049007