**Source URL:** https://medicines.health.europa.eu/veterinary/en/700000134551

# PULSIX 250 mg/1250 mg spot-on solution for dogs over 10 kg up to 25 kg

Authorised

- Imidacloprid
- Permethrin (40:60)

# Product identification

### **Medicine name:**

PULSIX 250 mg/1250 mg spot-on solution for dogs over 10 kg up to 25 kg Pulsix (250 mg + 1250 mg)/pipetkę 2,5 ml Roztwór do nakrapiania

### **Active substance:**

**Imidacloprid** 

Permethrin (40:60)

# **Target species:**

Dog

### Route of administration:

Spot-on use

# **Product details**

# **Active substance and strength:**

Imidacloprid

250.00 milligram(s) / 1.00 Pipette

Permethrin (40:60)

1250.00 milligram(s) / 1.00 Pipette

### **Pharmaceutical form:**

Spot-on solution

# Anatomical therapeutic chemical veterinary (ATCvet) codes:

QP53AC54

## Legal status of supply:

Veterinary medicinal product not subject to veterinary prescription

### **Authorisation status:**

Valid

### Authorised in:

Poland

## Package description:

HISTORICAL Cardboard box containing 1 pipette of 2.5 ml without pouch Cardboard box containing 2 pipettes of 2.5 ml with pouch HISTORICAL Cardboard box containing 2 pipettes of 2.5 ml without pouch Cardboard box containing 3 pipettes of 2.5 ml with pouch HISTORICAL Cardboard box containing 3 pipettes of 2.5 ml without pouch Cardboard box containing 4 pipettes of 2.5 ml with pouch HISTORICAL Cardboard box containing 4 pipettes of 2.5 ml without pouch Cardboard box containing 6 pipettes of 2.5 ml with pouch HISTORICAL Cardboard box containing 6 pipettes of 2.5 ml with pouch Cardboard box containing 1 pipette of 2.5 ml with pouch Cardboard box containing 24 pipettes of 2.5 ml with pouch HISTORICAL Cardboard box containing 24 pipettes of 2.5 ml without pouch HISTORICAL Cardboard box containing 12 pipettes of 2.5 ml without pouch Cardboard box containing 12 pipettes of 2.5 ml with pouch

# Additional information

# **Entitlement type:**

# Marketing Authorisation

# Legal basis of product authorisation:

Hybrid application (Article 13(3) of Directive No 2001/82/EC)

# Marketing authorisation holder:

Ab7 Sante

## Marketing authorisation date:

21/01/2025

# Manufacturing sites for batch release:

Ab7 Sante

# **Responsible authority:**

Office For Registration Of Medicinal Products Medical Devices And Biocidal Products

### **Authorisation number:**

3380

# Date of authorisation status change:

21/01/2025

### **Reference member state:**

Ireland

### **Procedure number:**

IE/V/0667/003

### **Concerned member states:**

France Germany Italy Netherlands Poland Portugal Spain

To consult adverse reactions on veterinary medicinal products please go to <a href="https://www.adrreports.eu/vet">www.adrreports.eu/vet</a>

# **Documents**

Summary of Product Characteristics
This document does not exist in this language (English). You can find it in another language below.
Package Leaflet
This document does not exist in this language (English). You can find it in another language below.
Labelling
This document does not exist in this language (English). You can find it in another language below.
Combined File of all Documents