# Peptizole 370 mg/g Oral Paste for Horses

Authorised

• Omeprazole

## Product identification

#### **Medicine name:**

Peptizole 370 mg/g Oral Paste for Horses Peptizole, 370 mg/g suukaudne pasta hobustele

### **Active substance:**

Omeprazole

#### **Target species:**

Horse

#### Route of administration:

Oral use

# **Product details**

## **Active substance and strength:**

Omeprazole 370.00 milligram(s) / 1.00 gram(s)

#### **Pharmaceutical form:**

Oral paste

## Withdrawal period by route of administration:

#### Oral use:

- Horse
  - Meat and offal. 1 day

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

**QA02BC01** 

#### Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

#### **Authorisation status:**

Valid

#### Authorised in:

Estonia

#### Package description:

Immediate package Immediate packaging: 7 ml oral syringe containing 7.57 g of paste composed of polyethylene barrel, plunger and end cap, with polypropylene dosing rings. Outer package and sales presentations - Bucket of 72 oral syringes Immediate package Immediate packaging: 7 ml oral syringe containing 7.57 g of paste composed of polyethylene barrel, plunger and end cap, with polypropylene dosing rings. Outer package and sales presentations - Carton box of 7 oral syringes Immediate package Immediate packaging: 7 ml oral syringe containing 7.57 g of paste composed of polyethylene barrel, plunger and end cap, with polypropylene dosing rings. Outer package and sales presentations - Carton box of 1 oral syringe

## 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

Manufacturing sites for batch release:				
Norbrook Laboratories Limited				
Norbrook Manufacturing Limited				
Responsible authority:				
State Agency Of Medicines				
Authorisation number:				
1787				
Date of authorisation status change:				
26/09/2013				
Reference member state:				
Ireland				
Procedure number:				
IE/V/0307/001				
Concerned member states:				
Austria Belgium Czechia Estonia Finland		9	•	
Lithuania Luxembourg Malta Netherlands	Poland	Portugal	Slovakia	Spain
United Kingdom (Northern Ireland)				

## **Documents**

Summary of Product Characteristics

English (PDF)

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