## Prascend 1 mg tablets for horses



• Pergolide mesilate

## Product identification

Medicine name: Prascend 1 mg tablets for horses Prascend 1 mg tabletten voor paarden

Active substance: Pergolide mesilate

**Target species:** Horse (non food-producing)

Route of administration:

Oral use

## **Product details**

#### Active substance and strength:

Pergolide mesilate 1.31 milligram(s) / 1.00 Tablet

#### **Pharmaceutical form:**

Tablet

Withdrawal period by route of administration: Oral use:

#### Horse (non food-producing)

- Meat and offal. no withdrawal period

Not authorised for use in horses intended for human consumption. The horse must have been declared as not intended for human consumption under national horse passport legislation. Not authorised for use in mares producing milk for human consumption.

- Milk. no withdrawal period

Not authorised for use in horses intended for human consumption. The horse must have been declared as not intended for human consumption under national horse passport legislation. Not authorised for use in mares producing milk for human consumption.

**Anatomical therapeutic chemical veterinary (ATCvet) codes:** QN04BC02

#### Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

#### Authorisation status:

Valid

#### Authorised in:

Netherlands

#### Package description:

(ID5): 1 Box with 3 Box with 16 Blister (PolyAmide, Aluminium, PolyVinyl Chloride) with 10 Piece (480.0 Piece)
(ID4): 1 Box with 13 Blister (PolyAmide, Aluminium, PolyVinyl Chloride) with 7 Piece (91.0 Piece)
(ID3): 1 Box with 1 Blister (Aluminium, PolyVinyl Chloride, PolyAmide) with 160 Piece (160.0 Piece)
(ID2): 1 Box with 1 Blister (Aluminium, PolyVinyl Chloride, PolyAmide) with 60 Piece (60.0 Piece)
(ID1): 1 Box with 1 Blister (Aluminium, PolyVinyl Chloride, PolyAmide) with 100 Piece (100.0 Piece)

## Additional information

#### **Entitlement type:**

Marketing Authorisation

#### Legal basis of product authorisation:

Full application (Article 12(3) of Directive No 2001/82/EC)

#### Marketing authorisation holder:

Boehringer Ingelheim Vetmedica GmbH

# Marketing authorisation date: 24/05/2012

## Manufacturing sites for batch release:

Boehringer Ingelheim Vetmedica GmbH Haupt Pharma Amareg GmbH

#### **Responsible authority:**

Medicines Evaluation Board

#### Authorisation number: REG NL 110598

#### Date of authorisation status change:

7/02/2022

#### **Reference member state:**

Germany

## Procedure number:

DE/V/0130/001

#### **Concerned member states:**

Austria Belgium Denmark Finland France Iceland Ireland Italy Luxembourg Netherlands Norway Sweden United Kingdom (Northern Ireland) To consult adverse reactions on veterinary medicinal products please go to www.adrreports.eu/vet

### Documents

Combined File of all Documents

English (PDF) Published on: 6/08/2024 <u>Download</u>

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