

# Pergosafe 0.25 mg film-coated tablets for horses

Authorised

- Pergolide mesilate

## Product identification

**Medicine name:**

Pergosafe 0.25 mg film-coated tablets for horses

Pergosafe 0,25 mg comprimidos revestidos por película para cavalos

**Active substance:**

Pergolide mesilate

**Target species:**

Horse

**Route of administration:**

Oral use

## Product details

**Active substance and strength:**

Pergolide mesilate

0.33 milligram(s) / 1.00 Tablet

**Pharmaceutical form:**

Film-coated tablet

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

QN04BC02

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**Legal status of supply:**

Veterinary medicinal product subject to veterinary prescription

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**Authorisation status:**

Valid

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**Authorised in:**

Portugal

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**Package description:**

PVC-PE-PVDC-aluminium blisters, containing 10 tablets each, in a carton box (160 tablets)

PVC-PE-PVDC-aluminium blisters, containing 10 tablets each, in a carton box (90 tablets)

PVC-PE-PVDC-aluminium blisters, containing 10 tablets each, in a carton box (60 tablets)

PVC-PE-PVDC-aluminium blisters, containing 10 tablets each, in a carton box (30 tablets)

PVC-PE-PVDC-aluminium blisters, containing 10 tablets each, in a carton box (240 tablets)

PVC-PE-PVDC-aluminium blisters, containing 10 tablets each, in a carton box (120 tablets)

PVC-PE-PVDC-aluminium blisters, containing 10 tablets each, in a carton box (100 tablets)

PVC-PE-PVDC-aluminium blisters, containing 10 tablets each, in a carton box (10 tablets)

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## Additional information

**Entitlement type:**

Marketing Authorisation

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**Legal basis of product authorisation:**

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

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**Marketing authorisation holder:**

Alfasan Nederland B.V.

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**Marketing authorisation date:**

14/11/2025

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**Manufacturing sites for batch release:**

Alfasan Nederland B.V.

Lelypharma B.V.

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**Responsible authority:**

Directorate General For Food And Veterinary

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**Authorisation number:**

1750/01/25DFVPT

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**Date of authorisation status change:**

14/11/2025

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**Reference member state:**

Netherlands

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**Procedure number:**

NL/V/0357/004

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**Concerned member states:**

Austria Belgium Bulgaria Croatia Cyprus Czechia Denmark Estonia Finland  
France Germany Greece Hungary Iceland Ireland Italy Latvia Lithuania  
Luxembourg Norway Poland Portugal Romania Slovakia Slovenia Spain  
Sweden United Kingdom (Northern Ireland)

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## Documents

Combined File of all Documents

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