Prednicortone 20 mg tablets for dogs and cats

Authorised

Prednisolone

Product identification

Medicine name:

Prednicortone 20 mg tablets for dogs and cats Prednicortone, 20 mg, tableta, za pse i mačke

Active substance:

Prednisolone

Target species:

Dog

Cat

Route of administration:

Oral use

Product details

Active substance and strength:

Prednisolone

20.00 milligram(s) / 1.00 Tablet

Pharmaceutical form:

Tablet

Withdrawal period by route of administration: Oral use:

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Dog

•

Cat

Anatomical therapeutic chemical veterinary (ATCvet) codes:

OH02AB06

Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

Authorisation status:

Valid

Authorised in:

Croatia

Package description:

Cardboard box of 50 Aluminium - PVC/PE/PVDC blisters of 10 tablets.

Cardboard box of 5 Aluminium - PVC/PE/PVDC blisters of 10 tablets.

Cardboard box of 4 Aluminium - PVC/PE/PVDC blisters of 10 tablets.

Cardboard box of 3 Aluminium - PVC/PE/PVDC blisters of 10 tablets.

Cardboard box of 25 Aluminium - PVC/PE/PVDC blisters of 10 tablets.

Cardboard box of 2 Aluminium - PVC/PE/PVDC blisters of 10 tablets.

Cardboard box of 15 Aluminium - PVC/PE/PVDC blisters of 10 tablets.

Cardboard box of 10 Aluminium - PVC/PE/PVDC blisters of 10 tablets.

Cardboard box of 7 Aluminium - PVC/PE/PVDC blisters of 10 tablets.

Cardboard box of 1 Aluminium - PVC/PE/PVDC blister of 10 tablets.

Cardboard box of 9 Aluminium - PVC/PE/PVDC blisters of 10 tablets.

Cardboard box of 6 Aluminium - PVC/PE/PVDC blisters of 10 tablets.

Cardboard box of 8 Aluminium - PVC/PE/PVDC blisters of 10 tablets.

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:

Dechra Regulatory B.V.

Marketing authorisation date:

13/10/2016

Manufacturing sites for batch release:

Lelypharma B.V.

Genera d.d.

Responsible authority:

Ministry Of Agriculture Veterinary And Food Safety Directorate

Authorisation number:

UP/I-322-05/20-01/337

Date of authorisation status change:

26/03/2025

Reference member state:

Netherlands

Procedure number:

NL/V/0190/002

Concerned member states:

Austria Belgium 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

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Combined File of all Documents

Summary of Product Characteristics

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