Amoxibactin 250 mg tablets for dogs

Authorised

• Amoxicillin trihydrate

Product identification

Medicine name:

Amoxibactin 250 mg tablets for dogs Amoxibactin vet 250 mg tabletes suniem

Active substance:

Amoxicillin trihydrate

Target species:

Dog

Route of administration:

Oral use

Product details

Active substance and strength:

Amoxicillin trihydrate 287.50 milligram(s) / 1.00 Tablet

Pharmaceutical form:

Tablet

Withdrawal period by route of administration:

Oral use:

. Dog

Anatomical therapeutic chemical veterinary (ATCvet) codes:

QJ01CA04

Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

Authorisation status:

Valid

Authorised in:

Latvia

Package description:

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

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

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

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

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

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 containing 10 separate cardboard boxes, each containing 1 blister 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:

Le Vet. Beheer B.V.

Marketing authorisation date:

31/10/2014

Manufacturing sites for batch release:

Lelypharma B.V.

Responsible authority:

Food And Veterinary Service

Authorisation number:

V/DCP/14/0060

Date of authorisation status change:

2/11/2014

Reference member state:

Netherlands

Procedure number:

NL/V/0186/002

Concerned member states:

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

To consult adverse reactions on veterinary medicinal products please go to www.adrreports.eu/vet

Documents

Summary of Product Characteristics

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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.

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