

# Anthelmin Plus XL Tablets for dogs

Authorised

- Febantel
- Pyrantel embonate
- Praziquantel

## Product identification

**Medicine name:**

Anthelmin Plus XL Tablets for dogs  
Anthelmin Plus XL compresse per cani

**Active substance:**

Febantel  
Pyrantel embonate  
Praziquantel

**Target species:**

Dog

**Route of administration:**

Oral use

## Product details

**Active substance and strength:**

Febantel

525.00 milligram(s) / 1.00 Tablet

Pyrantel embonate

504.00 milligram(s) / 1.00 Tablet

Praziquantel

175.00 milligram(s) / 1.00 Tablet

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**Pharmaceutical form:**

Tablet

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

QP52AC55

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**Legal status of supply:**

Veterinary medicinal product not subject to veterinary prescription

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**Authorisation status:**

Valid

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**Authorised in:**

Italy

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**Available in:**

Italy

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**Package description:**

OPA/Al/PVC-Al blister.Cardboard box containing 2 tablets.

OPA/Al/PVC-Al blister.Cardboard box containing 4 tablets.

OPA/Al/PVC-Al blister.Cardboard box containing 10 tablets.

OPA/Al/PVC-Al blister.Cardboard box containing 12 tablets.

OPA/Al/PVC-Al blister.Cardboard box containing 24 tablets.

OPA/Al/PVC-Al blister.Cardboard box containing 30 tablets.

OPA/Al/PVC-Al blister.Cardboard box containing 50 tablets.

OPA/Al/PVC-Al blister.Cardboard box containing 60 tablets.

OPA/Al/PVC-Al blister.Cardboard box containing 100 tablets.

OPA/Al/PVC-Al blister.Cardboard box containing 102 tablets.

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## Additional information

**Entitlement type:**

Marketing Authorisation

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**Legal basis of product authorisation:**

Generic application (Article 13(1) of Directive No 2001/82/EC)

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**Marketing authorisation holder:**

KRKA tovarna zdravil d.d. Novo mesto

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**Marketing authorisation date:**

27/01/2016

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**Manufacturing sites for batch release:**

KRKA tovarna zdravil d.d. Novo mesto  
Krka-Farma d.o.o.

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**Responsible authority:**

Ministry Of Health

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**Authorisation number:**

104844

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**Date of authorisation status change:**

27/01/2016

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**Reference member state:**

Ireland

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**Procedure number:**

IE/V/0538/002

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**Concerned member states:**

Belgium Bulgaria Finland Italy Netherlands Portugal Spain  
United Kingdom (Northern Ireland)

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To consult adverse reactions on veterinary medicinal products please go to [www.adrreports.eu/vet](http://www.adrreports.eu/vet)

## Documents

Summary of Product Characteristics

Combined File of all Documents

English (PDF)

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