

Forthyron 200 Microgram Flavoured Tablet

Authorised

- Levothyroxine sodium

Product identification

Medicine name:

Forthyron 200 Microgram Flavoured Tablet
Forthyron vet 200 mikrog tablett med smak til hund

Active substance:

Levothyroxine sodium

Target species:

Dog

Route of administration:

Oral use

Product details

Active substance and strength:

Levothyroxine sodium
200.00 microgram(s) / 1.00 Tablet

Pharmaceutical form:

Tablet

Withdrawal period by route of administration:

Oral use:

-

Dog

Anatomical therapeutic chemical veterinary (ATCvet) codes:

QH03AA01

Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

Authorisation status:

Valid

Authorised in:

Norway

Package description:

10 tablets in a blister [Aluminium (20micrometre) - PVC_PE_PVDC (250_30_90) white],
25 blisters per carton

10 tablets in a blister [Aluminium (20micrometre) - PVC_PE_PVDC (250_30_90) white],
5 blisters per carton

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:

Eurovet Animal Health B.V.

Marketing authorisation date:

14/03/2012

Manufacturing sites for batch release:

Dales Pharmaceuticals Limited

Eurovet Animal Health B.V.
Genera d.d.

Responsible authority:

Norwegian Medical Products Agency

Authorisation number:

10-7971

Date of authorisation status change:

26/10/2016

Reference member state:

Netherlands

Procedure number:

NL/V/0286/001

Concerned member states:

Austria Belgium Bulgaria Croatia Cyprus Czechia Denmark Estonia Finland
France Germany Greece Hungary Ireland Italy Latvia Lithuania
Luxembourg Norway Poland Portugal Romania Slovakia Slovenia Spain
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

Summary of Product Characteristics

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PAR 108733.pdf

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