

Hormon ReVet RV13 - Injektionslösung für Tiere

Authorised

- Glandula suprarenalis suis C9
- Glandula thymi suis C9
- Hypophysis suis C9
- Ovarium suis C9
- Pancreas suis C9
- Testis suis C9
- Glandula thyreoidea suis c9

Product identification

Medicine name:

Hormon ReVet RV13 - Injektionslösung für Tiere

Active substance:

Glandula suprarenalis suis C9

Glandula thymi suis C9

Hypophysis suis C9

Ovarium suis C9

Pancreas suis C9

Testis suis C9

Glandula thyreoidea suis c9

Target species:

Pigeon

Reptile
Ornamental bird
Dog
Cat
Rabbit
Ferret
Small rodents

Route of administration:

Intramuscular use
Intravenous use
Subcutaneous use

Product details

Active substance and strength:

Glandula suprarenalis suis C9
0.29 gram(s) / 2.00 millilitre(s)
Glandula thymi suis C9
0.29 gram(s) / 2.00 millilitre(s)
Hypophysis suis C9
0.29 gram(s) / 2.00 millilitre(s)
Ovarium suis C9
0.29 gram(s) / 2.00 millilitre(s)
Pancreas suis C9
0.29 gram(s) / 2.00 millilitre(s)
Testis suis C9
0.29 gram(s) / 2.00 millilitre(s)
Glandula thyreoidea suis c9
0.29 gram(s) / 2.00 millilitre(s)

Pharmaceutical form:

Solution for injection

Withdrawal period by route of administration:

Intramuscular use:

-

Pigeon

- Meat and offal. 0 day

-

Rabbit

- Meat and offal. 0 day

Intravenous use:

-

Rabbit

- Meat and offal. 0 day

Subcutaneous use:

-

Pigeon

- Meat and offal. 0 day

-

Rabbit

- Meat and offal. 0 day

Anatomical therapeutic chemical veterinary (ATCvet) codes:

QV03AX

Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

Authorisation status:

Valid

Authorised in:

Austria

Package description:

Available only in German

Additional information

Entitlement type:

Marketing Authorisation

Legal basis of product authorisation:

Homeopathic medicinal products (Article 19 of Directive No 2001/82/EC)

Marketing authorisation holder:

Pharmazeutische Fabrik Dr. Reckeweg & Co. GmbH

Marketing authorisation date:

6/07/1998

Manufacturing sites for batch release:

Pharmazeutische Fabrik Dr. Reckeweg & Co. GmbH

Responsible authority:

Austrian Agency For Health And Food Safety

Authorisation number:

8-30038

Date of authorisation status change:

6/07/1998

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

Documents

Package Leaflet

This document does not exist in this language (English). You can find it in another language below.

Summary of Product Characteristics

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.