Versiguard Rabies, Suspension for injection

Authorised

• Rabies virus, strain SAD Vnukovo-32, Inactivated

Product identification

Medicine name:

Versiguard Rabies, Suspension for injection Versiguard Rabies, suspensão injetável

Active substance:

Rabies virus, strain SAD Vnukovo-32, Inactivated

Target species:

Cat

Cattle

Pig

Sheep

Goat

Horse

Ferret

Dog

Route of administration:

Intramuscular use

Subcutaneous use

Product details

Active substance and strength:

Rabies virus, strain SAD Vnukovo-32, Inactivated 2.00 international unit(s) / 1.00 Dose

Pharmaceutical form:

Suspension for injection

Withdrawal period by route of administration:

Intramuscular use:

- . Cat
- . Cattle
 - Milk. 0 hour
 - Meat and offal. 0 day
- Pig
 - Meat and offal. 0 day
- Sheep
 - Milk. 0 hour
 - Meat and offal. 0 day
- Goat
 - Milk. 0 hour
 - Meat and offal. 0 day
- Horse
 - Milk. 0 hour
 - Meat and offal. 0 day
- Ferret

Subcutaneous use:

- . Dog
- . Cat
- . Cattle
 - Milk. 0 hour

- Meat and offal. 0 day
- . Pig
 - Meat and offal. 0 day
- Sheep
 - Milk. 0 hour
 - Meat and offal. 0 day
- . Goat
 - Milk. 0 hour
 - Meat and offal. 0 day
- Horse
 - Milk. 0 hour
 - Meat and offal. 0 day
- Ferret

Anatomical therapeutic chemical veterinary (ATCvet) codes:

OI07AA02

Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

Authorisation status:

Valid

Authorised in:

Portugal

Package description:

Glass Vial 1 x 1.0 Dose

Glass Vial 10 x 10.0 Dose

Glass Vial 10 x 1.0 Dose

Additional information

Entitlement type:

Marketing Authorisation

Legal basis of product authorisation:

Full application (Article 12(3) of Directive No 2001/82/EC)

Marketing authorisation holder:

Zoetis Portugal Lda.

Marketing authorisation date:

24/04/2013

Manufacturing sites for batch release:

Bioveta a.s.

Responsible authority:

Directorate General For Food And Veterinary

Authorisation number:

830/10RIVPT

Date of authorisation status change:

24/04/2013

Reference member state:

Czechia

Procedure number:

CZ/V/0100/001

Concerned member states:

Austria Belgium Bulgaria Croatia Cyprus Denmark Estonia Finland France Germany Greece Hungary Ireland Italy Latvia Lithuania Luxembourg Netherlands 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

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