

HIPRABOVIS-4

Authorised

- Bovine herpesvirus 1, strain LA, Inactivated
- Bovine parainfluenza virus 3, strain SF-4, Inactivated
- Bovine viral diarrhoea virus, strain NADL, Inactivated
- Bovine respiratory syncytial virus, strain Lym-56, Live

Product identification

Medicine name:

HIPRABOVIS-4

Active substance:

Bovine herpesvirus 1, strain LA, Inactivated

Bovine parainfluenza virus 3, strain SF-4, Inactivated

Bovine viral diarrhoea virus, strain NADL, Inactivated

Bovine respiratory syncytial virus, strain Lym-56, Live

Target species:

Cattle

Route of administration:

Intramuscular use

Product details

Active substance and strength:

Bovine herpesvirus 1, strain LA, Inactivated

39810700.00 50% tissue culture infectious dose / 3.00 millilitre(s)

Bovine parainfluenza virus 3, strain SF-4, Inactivated

500.00 haemagglutinating units / 3.00 millilitre(s)

Bovine viral diarrhoea virus, strain NADL, Inactivated

3981070.00 50% tissue culture infectious dose / 3.00 millilitre(s)

Bovine respiratory syncytial virus, strain Lym-56, Live

1995260.00 50% tissue culture infectious dose / 3.00 millilitre(s)

Pharmaceutical form:

Powder and solvent for suspension for injection

Withdrawal period by route of administration:

Intramuscular use:

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Cattle

- Meat and offal. 0 day

- Milk. 0 day

Anatomical therapeutic chemical veterinary (ATCvet) codes:

QI02AH

Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

Authorisation status:

Valid

Authorised in:

Ireland

Package description:

Freeze-dried fraction: the container is composed by neutral colourless glass flasks of 10 ml (30 doses) classified as Type I (Eur. Phar.), grey elastomer Type I closures (Eur. Phar.) and detachable aluminium caps. Liquid fraction: the container is composed by coloured glass flasks of 100 ml (30 doses) Type II (Eur. Phar.), grey elastomer Type II closures (Eur. Phar.) and capsules made of anodised aluminium.

Freeze-dried fraction: the container is composed by neutral colourless glass flasks of 10 ml (5 doses) classified as Typel (Eur. Phar.), grey elastomer Type I closures (Eur. Phar.) and detachable aluminium caps. Liquid fraction: the container is composed by coloured glass flasks of 20 ml (5 doses) Type I (Eur. Phar.), grey elastomer Type II closures (Eur. Phar.) and capsules made of anodised aluminium.

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:

Laboratorios Hipra S.A.

Marketing authorisation date:

6/06/2003

Manufacturing sites for batch release:

Laboratorios Hipra S.A.

Responsible authority:

Health Products Regulatory Authority

Authorisation number:

VPA10846/003/001

Date of authorisation status change:

6/06/2003

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

Documents

Summary of Product Characteristics