Nobivac Respira Bb vet. suspension for injection for dogs

Authorised

• Bordetella bronchiseptica, strain Bb7 92932, fimbriae

Product identification

Medicine name:

Nobivac Respira Bb vet. suspension for injection for dogs Nobivac Respira Bb, süstesuspensioon koertele

Active substance:

Bordetella bronchiseptica, strain Bb7 92932, fimbriae

Target species:

Dog

Route of administration:

Subcutaneous use

Product details

Active substance and strength:

Bordetella bronchiseptica, strain Bb7 92932, fimbriae 88.00 enzyme-linked immunosorbent assay unit / 1.00 millilitre(s)

Pharmaceutical form:

Suspension for injection

Withdrawal period by route of administration:

Subcutaneous use:
Anatomical therapeutic chemical veterinary (ATCvet) codes: QI06AE02
Legal status of supply: Veterinary medicinal product subject to veterinary prescription
Authorisation status: Valid
Authorised in: Estonia
Package description: Available only in Danish
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: Intervet International B.V.
Marketing authorisation date: This information is not available for this product.
Manufacturing sites for batch release: INTERVET INTERNATIONAL B.V.
Responsible authority: State Agency Of Medicines
Authorisation number: 2250

Reference member state: Denmark
Procedure number: DK/V/0123/002
Concerned member states: Austria Belgium Bulgaria Croatia Cyprus Czechia Estonia Finland France Germany Greece Hungary Iceland 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
Documents
Summary of Product Characteristics
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PI.pdf

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Date of authorisation status change:

30/08/2020