

# Bupredine Multidose 0.3 mg/ml Solution for Injection for Dogs, Cats and Horses

Authorised

- Buprenorphine hydrochloride

## Product identification

### Medicine name:

Bupredine Multidose 0.3 mg/ml Solution for Injection for Dogs, Cats and Horses  
Bupredine Multidose vet 0,3 mg/ml šķīdums injekcijām suņiem, kaķiem un zirgiem

### Active substance:

Buprenorphine hydrochloride

### Target species:

Dog  
Horse  
Cat

### Route of administration:

Intramuscular use  
Intravenous use

## Product details

### Active substance and strength:

Buprenorphine hydrochloride  
0.32 milligram(s) / 1.00 millilitre(s)

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**Pharmaceutical form:**

Solution for injection

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**Anatomical therapeutic chemical veterinary (ATCvet) codes:**

QN02AE01

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**Legal status of supply:**

Veterinary medicinal product subject to veterinary prescription

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**Authorisation status:**

Valid

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**Authorised in:**

Latvia

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**Package description:**

Clear type I glass vials closed with a coated bromobutyl rubber stopper and aluminium cap in a carton box. Pack size: 20 ml.

Clear type I glass vials closed with a coated bromobutyl rubber stopper and aluminium cap in a carton box. Pack size: 100 ml

Clear type I glass vials closed with a coated bromobutyl rubber stopper and aluminium cap in a carton box. Pack size: 10 ml.

Clear type I glass vials closed with a coated bromobutyl rubber stopper and aluminium cap in a carton box. Pack size: 50 ml

Clear type I glass vials closed with a coated bromobutyl rubber stopper and aluminium cap in a carton box. Pack size: 5 ml.

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## Additional information

**Entitlement type:**

Marketing Authorisation

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**Legal basis of product authorisation:**

Generic application (Article 13(1) of Directive No 2001/82/EC)

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**Marketing authorisation holder:**

Le Vet. Beheer B.V.

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**Marketing authorisation date:**

27/10/2015

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**Manufacturing sites for batch release:**

Produlab Pharma B.V.

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**Responsible authority:**

Food And Veterinary Service

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**Authorisation number:**

V/DCP/15/0048

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**Date of authorisation status change:**

27/10/2015

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**Reference member state:**

Netherlands

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**Procedure number:**

NL/V/0314/001

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**Concerned member states:**

Austria Belgium Croatia Cyprus Czechia Denmark Estonia France Germany  
Greece Hungary Iceland Latvia Lithuania Luxembourg Norway Poland  
Portugal Romania Slovakia Slovenia Spain Sweden  
United Kingdom (Northern Ireland)

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To consult adverse reactions on veterinary medicinal products please go to  
[www.adrreports.eu/vet](http://www.adrreports.eu/vet)

## Documents

Summary of Product Characteristics

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

Package Leaflet

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.