KETABEL 100 MG/ML SOLUTION FOR INJECTION

Authorised

• Ketamine hydrochloride

Product identification

Medicine name:

KETABEL 100 MG/ML SOLUTION FOR INJECTION Ketabel 100 mg/ml šķīdums injekcijām

Active substance:

Ketamine hydrochloride

Target species:

Cattle

Pig

Rat

Guinea pig

Rabbit

Cat

Sheep

Goat

Dog

Horse

Hamster

Mouse

Route of administration:

Intramuscular use

Product details

Active substance and strength:

Ketamine hydrochloride 115.34 milligram(s) / 1.00 millilitre(s)

Pharmaceutical form:

Solution for injection

Withdrawal period by route of administration: Intramuscular use:

•

Cattle

- Milk. 0 day
- Meat and offal. 1 day

•

Pig

- Meat and offal. 1 day

•

Rat

•

Guinea pig

•

Rabbit

•

Cat

•

Sheep

- Meat and offal. 1 day
- Milk. 0 day

• Goat - Meat and offal. 1 day - Milk. 0 day
• Dog
Intravenous use:
•
Cattle - Meat and offal. 1 day
- Milk. 0 day
· Uarra
Horse - Meat and offal. 1 day
•
- Milk. 0 day
•
Sheep
- Meat and offal. 1 day
- Milk. 0 day
•
Goat
- Meat and offal. 1 day
- Milk. 0 day
Intraperitoneal use:
• Hamster

Rat

Mouse

Legal status of supply:	
Veterinary medicinal product subject to veterinary prescription	
Authorisation status:	
Valid	
Authorised in:	
Latvia	
Available in:	
Latvia	
Package description:	
Carton with 1 vial of 10 mL	
Carton with 10 vials of 25 mL	
Carton with 1 vial of 25 mL	
Carton with 10 vials of 10 mL	
Additional information	
Entitlement type:	
Marketing Authorisation	
Legal basis of product authorisation:	
Generic application (Article 13(1) of Directive No 2001/82/EC)	
Marketing authorisation holder:	
Bela-Pharm GmbH & Co. KG	
Marketing authorisation date: 5/07/2020	

Responsible authority:

Food And Veterinary Service	
Authorisation number: V/DCP/20/0038	
Date of authorisation status change: 6/07/2020	
Reference member state: France	
Procedure number: FR/V/0338/001	
Concerned member states: Austria Belgium Bulgaria Cyprus Czechia Denmark Estonia Finland Germany Greece Hungary Iceland Ireland Italy Latvia Lithuania Luxembourg Netherlands Norway 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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Package Leaflet	

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Labelling

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