# Enrotron Flavour 50 mg Tablets for dogs

Authorised

Enrofloxacin

## Product identification

#### **Medicine name:**

Enrotron Flavour 50 mg Tablets for dogs Enrotron Flavour 50 mg

#### **Active substance:**

Enrofloxacin

### **Target species:**

Dog

#### Route of administration:

Oral use

## **Product details**

## **Active substance and strength:**

Enrofloxacin 50.00 milligram(s) / 1.00 Tablet

#### **Pharmaceutical form:**

**Tablet** 

## Withdrawal period by route of administration:

#### Oral use:

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Dog

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

**QI01MA90** 

#### Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

#### **Authorisation status:**

Valid

#### **Authorised in:**

Germany

#### Package description:

(ID3) 30 Tablet: unspecified outer container with 1 Blister (Aluminium; Aluminium) with 30 Tablet

(ID9) 50 Tablet: unspecified outer container with 1 Blister (Aluminium; PolyVinyl Chloride) with 50 Tablet

(ID6) 10 Tablet: unspecified outer container with 1 Blister (Aluminium; PolyVinyl Chloride) with 10 Tablet

(ID2) 20 Tablet: unspecified outer container with 1 Blister (Aluminium; Aluminium) with 20 Tablet

(ID5) 100 Tablet: unspecified outer container with 1 Blister (Aluminium; Aluminium) with 100 Tablet

(ID7) 20 Tablet: unspecified outer container with 1 Blister (Aluminium; PolyVinyl Chloride) with 20 Tablet

(ID8) 30 Tablet: unspecified outer container with 1 Blister (Aluminium; PolyVinyl Chloride) with 30 Tablet

(ID4) 50 Tablet: unspecified outer container with 1 Blister (Aluminium; Aluminium) with 50 Tablet

(ID1) 10 Tablet: unspecified outer container with 1 Blister (Aluminium; Aluminium) with 10 Tablet

(ID10) 100 Tablet: unspecified outer container with 1 Blister (Aluminium; PolyVinyl Chloride) with 100 Tablet

## 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:

aniMedica GmbH

#### Marketing authorisation date:

22/09/2010

#### Manufacturing sites for batch release:

aniMedica GmbH

#### **Responsible authority:**

Federal Office Of Consumer Protection And Food Safety

#### **Authorisation number:**

401296.00.00

## Date of authorisation status change:

5/04/2016

#### Reference member state:

Germany

#### **Procedure number:**

DE/V/0137/001

#### **Concerned member states:**

Austria Belgium Denmark Finland Greece Hungary Iceland Ireland Luxembourg Netherlands Poland Slovenia United Kingdom (Northern Ireland)

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

## **Documents**

Summary of Product Characteristics	
2401296-paren-20101101.pdf	

**Source URL:** https://medicines.health.europa.eu/veterinary/600000064863