**Source URL:** https://medicines.health.europa.eu/veterinary/en/700000140594

# Taurador 10 mg/ml Solution for Injection for cattle, sheep & pigs

Authorised

Doramectin

## Product identification

#### **Medicine name:**

Taurador 10 mg/ml Solution for Injection for cattle, sheep & pigs Taurador, 10mg/ml, Injekční roztok

#### **Active substance:**

Doramectin

#### **Target species:**

Cattle

Sheep

Pig

#### **Route of administration:**

Subcutaneous use Intramuscular use

## **Product details**

## **Active substance and strength:**

Doramectin

10.00 milligram(s) / 1.00 millilitre(s)

#### Pharmaceutical form:

Solution for injection

## Withdrawal period by route of administration:

**Subcutaneous use:** 

Cattle

- Meat and offal. 70 day
- Milk. no withdrawal period

Not authorised for use in animals producing milk for human consumption. Do not use in pregnant cows or heifers which are intended to produce milk for human consumption within 2 months of expected parturition.

#### Intramuscular use:

Sheep

- Meat and offal. 70 day
- Milk. no withdrawal period

Not authorised for use in animals producing milk for human consumption. Do not use in pregnant ewes which are intended to produce milk for human consumption within 70 days of expected parturition.

Pig

- Meat and offal. 77 day

## Anatomical therapeutic chemical veterinary (ATCvet) codes:

QP54AA03

## Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

#### **Authorisation status:**

Valid

#### **Authorised in:**

#### Package description:

(ID3) 500 millilitre(s): Behältnis (plastics) with 1 Vial (brown glass) with 500 millilitre(s), closed with (Gummi) and (Aluminium)

(ID2) 250 millilitre(s): Behältnis (plastics) with 1 Vial (brown glass) with 250 millilitre(s), closed with (Aluminium) and (Gummi)

(ID1) 100 millilitre(s): Behältnis (plastics) with 1 Vial (brown glass) with 100 millilitre(s), closed with (Gummi) and (Aluminium)

## Additional information

## **Entitlement type:**

Marketing Authorisation

## Legal basis of product authorisation:

Generic application (Article 18 of Regulation (EU) 2019/6)

## Marketing authorisation holder:

Norbrook Laboratories (Ireland) Limited

## Marketing authorisation date:

18/02/2025

## Manufacturing sites for batch release:

Norbrook Laboratories Limited

Norbrook Manufacturing Limited

## **Responsible authority:**

Institute For State Control Of Veterinary Biologicals And Medicaments

#### **Authorisation number:**

96/007/25-C

## Date of authorisation status change:

18/02/2025

#### Reference member state:

Germany

#### **Procedure number:**

DE/V/0345/001

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

Belgium Czechia France Hungary Ireland Netherlands Portugal Romania Slovakia Spain

#### **Generic of:**

60000073029

600000060573

600000085745

To consult adverse reactions on veterinary medicinal products please go to 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.