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

Ima dovoljenje  
za promet

- Doramectin

## Informacije o zdravilu

### Ime zdravila:

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

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### Učinkovina:

Na voljo samo v [English](#)

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### Ciljne živalske vrste:

govedo

Na voljo samo v [Bulgarian](#) [Spanish](#) [Czech](#) [Danish](#) [German](#) [Estonian](#) [Greek](#) [English](#) [French](#) [Italian](#) [Latvian](#) [Lithuanian](#) [Hungarian](#) [Dutch](#) [Romanian](#) [Finnish](#) [Swedish](#) [Icelandic](#) [Norwegian](#)

Na voljo samo v [Bulgarian](#) [Spanish](#) [Czech](#) [Danish](#) [German](#) [Estonian](#) [Greek](#) [English](#) [French](#) [Italian](#) [Latvian](#) [Lithuanian](#) [Hungarian](#) [Dutch](#) [Romanian](#) [Finnish](#) [Swedish](#) [Icelandic](#) [Norwegian](#)

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### Pot uporabe:

Subkutana uporaba

intramuskularna uporaba

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## Podatki o zdravilu

### **Učinkovina / Jakost:**

Na voljo samo v English  
10.00 milligram(s) / 1.00 millilitre(s)

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### **Farmacevtska oblika:**

Raztopina za injiciranje

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### **Karenca glede na pot uporabe:**

#### **Subkutana uporaba:**

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#### **govedo**

- 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.

#### **intramuskularna uporaba:**

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#### **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.

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#### **Pig**

- Meat and offal. 77 day

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**Anatomsko-terapevtsko-kemična veterinarska oznaka klasifikacija zdravil, ki se uporabljajo v veterini (ATCvet):**

QP54AA03

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**Pravni status za dobavo/izdajo zdravila:**

Zdravilo, ki se izdaja na veterinarski recept

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**Status dovoljenja za promet z zdravilom:**

Valid

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**Ima dovoljenje za promet v:**

Na voljo samo v [Spanish](#) [Czech](#) [German](#) [Estonian](#) [English](#) [French](#) [Italian](#) [Dutch](#) [Portuguese](#) [Slovak](#) [Swedish](#) [Icelandic](#) [Norwegian](#)

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**Opis ovojnine zdravila:**

Na voljo samo v [English](#)

Na voljo samo v [English](#)

Na voljo samo v [English](#)

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## Dodatne informacije

**Vrsta dovoljenja:**

Na voljo samo v [English](#) [French](#) [Croatian](#) [Italian](#) [Latvian](#) [Finnish](#) [Swedish](#) [Icelandic](#) [Norwegian](#)

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**Pravna podlaga za izdajo dovoljenja za promet z zdravilom:**

Na voljo samo v [English](#) [French](#) [Italian](#) [Latvian](#) [Lithuanian](#) [Norwegian](#)

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**Imetnik dovoljenja za promet z zdravilom:**

Norbrook Laboratories (Ireland) Limited

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**Datum dovoljenja za promet z zdravilom:**

27/01/2025

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**Proizvodna mesta, odgovorna za sproščanje serij:**

Norbrook Laboratories Limited

Norbrook Manufacturing Limited

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**Pristojni organ:**

Directorate Of Veterinary Medicinal Products

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**Številka dovoljenja za promet z zdravilom:**

Ta podatek za to zdravilo ni na voljo.

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**Datum spremembe statusa dovoljenja za promet:**

27/01/2025

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**Referenčna država članica:**

Na voljo samo v Spanish Czech German Estonian English French Italian Dutch  
Portuguese Slovak Swedish Icelandic Norwegian

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**Številka postopka:**

DE/V/0345/001

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**Zadevne države članice:**

Na voljo samo v Spanish Czech German Estonian English French Italian Dutch  
Portuguese Slovak Swedish Icelandic Norwegian

Na voljo samo v Spanish Czech German Estonian English French Italian Latvian  
Dutch Portuguese Slovak Swedish Icelandic Norwegian

Na voljo samo v Spanish Czech German Estonian English French Italian Dutch  
Portuguese Slovak Swedish Icelandic Norwegian

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Portuguese Slovak Swedish Icelandic Norwegian

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Portuguese Slovak Swedish Icelandic Norwegian

Na voljo samo v Spanish Czech German Estonian English French Italian Dutch  
Portuguese Slovak Swedish Icelandic Norwegian

Na voljo samo v Spanish Czech German Estonian English French Italian Lithuanian  
Dutch Portuguese Romanian Slovak Swedish Icelandic Norwegian

Na voljo samo v Spanish Czech German Estonian English French Italian Dutch  
Portuguese Slovak Swedish Icelandic Norwegian

Na voljo samo v Spanish Czech German Estonian English French Croatian Italian  
Dutch Portuguese Slovak Swedish Icelandic Norwegian

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**Generic of:**

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Za poročila o domnevnih neželenih učinkih zdravil za uporabo v veterinarski medicini obiščite stran: [www.adrreports.eu/vet](http://www.adrreports.eu/vet)

## Dokumenti

Combined File of all Documents

Ta dokument ne obstaja v tem jeziku (slovenščina). Spodaj ga najdete v drugem jeziku.