

# CYDECTIN TRICLAMOX 1 MG/ML + 50 MG/ML ORAL SOLUTION FOR SHEEP

Ima dovoljenje  
za promet

- Triclabendazole
- Moxidectin

## Informacije o zdravilu

### Ime zdravila:

CYDECTIN TRICLAMOX 1 MG/ML + 50 MG/ML ORAL SOLUTION FOR SHEEP

### Učinkovina:

Na voljo samo v [English](#)

Na voljo samo v [English](#)

### Ciljne živalske vrste:

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

### Pot uporabe:

Peroralna uporaba

## Podatki o zdravilu

### Učinkovina / Jakost:

Na voljo samo v [English](#)  
50.00 milligram(s) / 1.00 millilitre(s)

Na voljo samo v [English](#)  
1.00 milligram(s) / 1.00 millilitre(s)

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**Farmacevtska oblika:**

Peroralna raztopina

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**Karenca glede na pot uporabe:**

**Peroralna uporaba:**

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**Sheep**

- Milk. no withdrawal period

Not authorised for use in ewes producing milk intended for human consumption including during the dry period. Do not use within 1 year prior to the first lambing in ewes intended to produce milk for human consumption.

- Meat and offal. 31 day

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

QP54AB52

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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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**Na voljo v:**

Germany

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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](#) [Italian](#) [Latvian](#) [Norwegian](#)

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### **Imetnik dovoljenja za promet z zdravilom:**

Zoetis Deutschland GmbH

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### **Datum dovoljenja za promet z zdravilom:**

8/02/2010

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

Zoetis Manufacturing & Research Spain S.L.

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### **Pristojni organ:**

Federal Office Of Consumer Protection And Food Safety

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### **Številka dovoljenja za promet z zdravilom:**

401201.00.00

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### **Datum spremembe statusa dovoljenja za promet:**

10/03/2015

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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:**

FR/V/0201/001

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### **Zadevne države članice:**

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

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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). Lahko ga najdete spodaj v drugem jeziku.