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# Prascend 1 mg tablets for horses

Ima dovoljenje za promet

- Pergolide mesilate

## Informacije o zdravilu

### Ime zdravila:

Prascend 1 mg tablets for horses

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### Učinkovina:

Na voljo samo v [English](#)

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### 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](#)

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### Pot uporabe:

Peroralna uporaba

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## Podatki o zdravilu

### Učinkovina / Jakost:

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

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### Farmacevtska oblika:

Tableta

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**Karenca glede na pot uporabe:**

**Peroralna uporaba:**

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**Horse (non food-producing)**

- Meat and offal. no withdrawal period

Not authorised for use in horses intended for human consumption. The horse must have been declared as not intended for human consumption under national horse passport legislation. Not authorised for use in mares producing milk for human consumption.

- Milk. no withdrawal period

Not authorised for use in horses intended for human consumption. The horse must have been declared as not intended for human consumption under national horse passport legislation. Not authorised for use in mares producing milk for human consumption.

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

QN04BC02

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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](#)

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](#)

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

Boehringer Ingelheim Vetmedica GmbH

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

21/05/2010

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

Boehringer Ingelheim Vetmedica GmbH

Haupt Pharma Amareg GmbH

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### **Pristojni organ:**

Health Products Regulatory Authority

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

VPA10454/011/001

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

21/05/2010

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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/0130/001

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### **Zadevne države članice:**

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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.

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