

SYNULOX RTU ενέσιμο εναιώρημα για βοοειδή, χοίρους, σκύλους και γάτες

Geautoriseerd

- Amoxicillin trihydrate
- Potassium clavulanate

Product identification

Naam van het geneesmiddel:

SYNULOX RTU ενέσιμο εναιώρημα για βοοειδή, χοίρους, σκύλους και γάτες

Werkzame stof:

Alleen beschikbaar in [English](#)

Alleen beschikbaar in [English](#)

Doeldiersoort(en):

Rund

Varken

Kat

Hond

Toedieningsweg:

Intramusculair gebruik

Subcutaan gebruik

Product details

Werkzame stof / Sterkte:

Alleen beschikbaar in [English](#)
140.00 milligram(s) / 1.00 millilitre(s)

Alleen beschikbaar in [English](#)
35.00 milligram(s) / 1.00 millilitre(s)

Farmaceutische vorm:

Suspensie voor injectie

Withdrawal period by route of administration:

Intramusculair gebruik:

• **Rund**

- Meat and offal. 42 day
- Milk. 60 hour

• **Varken**

- Meat and offal. 26 day

• **Kat**

- Not applicable. no withdrawal period

• **Hond**

- Not applicable. no withdrawal period

Subcutaan gebruik:

• **Kat**

- Not applicable. no withdrawal period

• **Hond**

- Not applicable. no withdrawal period
-

Anatomisch therapeutisch chemische veterinaire classificatie (ATCvet code)

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QJ01CR02

Afleverstatus:

Alleen beschikbaar in [Czech](#) [Estonian](#) [English](#) [French](#) [Italian](#) [Latvian](#) [Portuguese](#)
[Slovenian](#) [Finnish](#) [Swedish](#) [Icelandic](#) [Norwegian](#)

Status toelating:

Valid

Authorised in:

Griekenland

Package description:

Alleen beschikbaar in [Greek](#)

Alleen beschikbaar in [Greek](#)

Additional information

Entitlement type:

Alleen beschikbaar in [English](#) [French](#) [Croatian](#) [Italian](#) [Latvian](#) [Finnish](#) [Swedish](#)

[Icelandic](#) [Norwegian](#)

Rechtsgrondslag productvergunning:

Alleen beschikbaar in [English](#)

Handelsvergunninghouder:

Zoetis Hellas S.A.

Marketing authorisation date:

8/06/1999

Productielocaties partijvrijgifte:

Haupt Pharma Latina S.r.l.

Verantwoordelijke instantie:

National Organization For Medicines

Toelatingsnummer:

31804/09-04-2021/K-0077403

Wijzigingsdatum status toelating:

9/04/2021

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

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