

# DIMEXAN 200 Solubile, 200 g/1000 g, polvere solubile per uso orale per vitelli da latte, suini (fino a sei mesi), polli da carne, conigli

Not  
authorised

- Sulfadimethoxine

## Product identification

### Medicine name:

DIMEXAN 200 Solubile, 200 g/1000 g, polvere solubile per uso orale per vitelli da latte, suini (fino a sei mesi), polli da carne, conigli

### Active substance:

Sulfadimethoxine

### Target species:

Chicken (broiler)  
Rabbit  
Pig (suckling piglet)  
Cattle (suckling calf)

### Route of administration:

In drinking water/milk use

## Product details

### Active substance and strength:

Sulfadimethoxine

200.00 gram(s) / 1000.00 gram(s)

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### Pharmaceutical form:

Powder for use in drinking water/milk

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### Withdrawal period by route of administration:

#### In drinking water/milk use:

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#### Chicken (broiler)

- Meat and offal. 15 day

Non somministrare a galline che producono uova destinate al consumo umano

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#### Rabbit

- Meat and offal. 21 day

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#### Pig (suckling piglet)

- Meat and offal. 28 day

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#### Cattle (suckling calf)

- Meat and offal. 28 day

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### Anatomical therapeutic chemical veterinary (ATCvet) codes:

QJ01EQ09

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### Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

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### Authorisation status:

Surrendered

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**Authorised in:**

Italy

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**Package description:**

Available only in [Italian](#)

Available only in [Italian](#)

Available only in [Italian](#)

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## Additional information

**Entitlement type:**

Marketing Authorisation

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**Legal basis of product authorisation:**

Full application (Article 12(3) of Directive No 2001/82/EC)

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**Marketing authorisation holder:**

Ceva Salute Animale S.p.A.

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**Marketing authorisation date:**

8/05/2002

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**Manufacturing sites for batch release:**

Vetem S.p.A.

Unione Commerciale Lombarda S.p.A.

Ceva Salute Animale S.p.A.

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**Responsible authority:**

Ministry Of Health

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**Authorisation number:**

102689

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**Date of authorisation status change:**

3/11/2023

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To consult adverse reactions on veterinary medicinal products please go to [www.adrreports.eu/vet](http://www.adrreports.eu/vet)

## Documents

Combined File of all Documents

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