

# NOBILIS GUMBORO 228 E LYOPHILISAT POUR ADMINISTRATION DANS L'EAU DE BOISSON POUR POULES

Authorised

- Infectious bursal disease virus, strain 288E, Live

## Product identification

### **Medicine name:**

NOBILIS GUMBORO 228 E LYOPHILISAT POUR ADMINISTRATION DANS L'EAU DE BOISSON POUR POULES

### **Active substance:**

Infectious bursal disease virus, strain 288E, Live

### **Target species:**

Chicken (layer hen)

Chicken (for reproduction)

Chicken (broiler)

### **Route of administration:**

In drinking water use

## Product details

### Active substance and strength:

Infectious bursal disease virus, strain 288E, Live  
2.00 log 10 50% embryo infective dose / 1.00 Dose

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### Pharmaceutical form:

Lyophilisate for suspension

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### Withdrawal period by route of administration:

#### In drinking water use:

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#### Chicken (layer hen)

- All relevant tissues. 0 day

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#### Chicken (for reproduction)

- All relevant tissues. 0 day

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#### Chicken (broiler)

- All relevant tissues. 0 day

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

QI01AD09

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### Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

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### Authorisation status:

Valid

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### Authorised in:

France

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### Package description:

Available only in French

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## Additional information

### **Entitlement type:**

Marketing Authorisation

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### **Legal basis of product authorisation:**

Legal basis not covered by Directive 2001/82/EC

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### **Marketing authorisation holder:**

Intervet

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### **Marketing authorisation date:**

27/11/2002

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### **Manufacturing sites for batch release:**

Intervet International B.V.

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### **Responsible authority:**

French Agency For Food, Environmental And Occupational Health & Safety

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### **Authorisation number:**

**Date of authorisation status change:**

27/11/2012

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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

### Summary of Product Characteristics

This document does not exist in this language (English). You can find it in another language below.

### Package Leaflet and Labelling

This document does not exist in this language (English). You can find it in another language below.