

# V.H. + H-120 liofilizāts ūdens šķīduma pagatavošanai vistām

Authorised

- Newcastle disease virus, strain V.H., Live
- Infectious bronchitis virus, type Massachusetts, strain H120, Live

## Product identification

### **Medicine name:**

V.H. + H-120 liofilizāts ūdens šķīduma pagatavošanai vistām

---

### **Active substance:**

Newcastle disease virus, strain V.H., Live

Infectious bronchitis virus, type Massachusetts, strain H120, Live

---

### **Target species:**

Chicken (layer hen)

Chicken (broiler)

Chicken (for reproduction)

---

### **Route of administration:**

In drinking water use

Intraocular use

Nebulisation use

---

## Product details

### **Active substance and strength:**

Newcastle disease virus, strain V.H., Live

Infectious bronchitis virus, type Massachusetts, strain H120, Live

---

### **Pharmaceutical form:**

Lyophilisate for use in drinking water

---

### **Withdrawal period by route of administration:**

#### **In drinking water use:**

- 

#### **Chicken (layer hen)**

- Not specified. 0 day

- 

#### **Chicken (broiler)**

- Not specified. 0 day

- 

#### **Chicken (for reproduction)**

- Not specified. 0 day

#### **Intraocular use:**

- 

#### **Chicken (broiler)**

- Not specified. 0 day

- 

#### **Chicken (for reproduction)**

- Not specified. 0 day

- 

#### **Chicken (layer hen)**

- Not specified. 0 day

#### **Nebulisation use:**

- 

**Chicken (broiler)**

- Not specified. 0 day

- 

**Chicken (for reproduction)**

- Not specified. 0 day

- 

**Chicken (layer hen)**

- Not specified. 0 day

---

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

QI01AD

---

**Legal status of supply:**

Veterinary medicinal product subject to veterinary prescription

---

**Authorisation status:**

Valid

---

**Authorised in:**

Latvia

---

**Package description:**

Available only in [Latvian](#)

---

## Additional information

**Entitlement type:**

Marketing Authorisation

---

**Legal basis of product authorisation:**

Immunological veterinary medicinal product application (Article 13d of Directive No 2001/82/EC)

---

**Marketing authorisation holder:**

Phibro Animal Health (Poland) Sp. z o.o.

---

**Marketing authorisation date:**

16/12/2010

---

**Manufacturing sites for batch release:**

Synoptis Industrial Sp. z o.o.

---

**Responsible authority:**

Food And Veterinary Service

---

**Authorisation number:**

V/NRP/10/0033

---

**Date of authorisation status change:**

16/12/2010

---

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

## Documents

Package Leaflet

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

## Summary of Product Characteristics

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

## Labelling

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