

Phenoxyphen WSP

Authorised

- Phenoxyphen WSP

Product identification

Medicine name:

Phenoxyphen WSP

Phenoxylin 325 mg/g pulver til oral oplosning

Active substance:

Phenoxyphen WSP

Target species:

Chicken

Route of administration:

In drinking water use

Oral use

Product details

Active substance and strength:

Phenoxyphen WSP

325.00 milligram(s) / 1.00 gram(s)

Pharmaceutical form:

Powder for oral solution

Withdrawal period by route of administration:**In drinking water use:**

-

Chicken

- Meat and offal. 2 day
- Eggs. 0 day

Oral use:

-

Chicken

- Meat and offal. 2 day
- Eggs. no withdrawal period zero days

Anatomical therapeutic chemical veterinary (ATCvet) codes:

QJ01CE02

Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

Authorisation status:

Valid

Authorised in:

Denmark

Available in:

Denmark

Package description:

White polypropylene bucket provided with a polypropylene lid containing 5 kg of product.

White polypropylene bucket provided with a polypropylene lid containing 2.5 kg of product.

Securitainer: white PP cylindrical container, with white HDPE/LDPE closure, with thumb tab for opening, containing 1000 g (1875 ml) of product.

Composite can: three layered rectangular container, which consists of a cardboard base with an inner lining of aluminium paper and with label on the outside containing

1 kg of product.

White polypropylene bucket provided with a polypropylene lid containing 1 kg of product.

Securitainer: white PP cylindrical container, with white HDPE/LDPE closure, with thumb tab for opening, containing 250 g (650 ml) of product.

Additional information

Entitlement type:

Marketing Authorisation

Legal basis of product authorisation:

Full application - Known active substance (Article 12(3) of Directive No 2001/82/EC)

Marketing authorisation holder:

Dopharma Research B.V.

Marketing authorisation date:

9/01/2007

Manufacturing sites for batch release:

Dopharma B.V.

Responsible authority:

Danish Medicines Agency

Authorisation number:

39520

Date of authorisation status change:

9/01/2007

Reference member state:

Netherlands

Procedure number:

NL/V/0121/001

Concerned member states:

Austria Belgium Bulgaria Czechia Denmark France Germany Greece
Hungary Ireland Italy Lithuania Poland Portugal Romania Slovakia Spain
United Kingdom (Northern Ireland)

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

Documents

Summary of Product Characteristics

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