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Zanil Fluke Drench

Not authorised

- Oxyclozanide

Product identification

Medicine name:

Zanil Fluke Drench

Active substance:

Oxyclozanide

Target species:

Cattle

Sheep

Route of administration:

Oral use

Product details

Active substance and strength:

Oxyclozanide

3.40 gram(s) / 100.00 millilitre(s)

Pharmaceutical form:

Oral suspension

Withdrawal period by route of administration:

Oral use:

-

Cattle

- Milk. 72 hour

Anatomical therapeutic chemical veterinary (ATCvet) codes:

QP52A

Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

Authorisation status:

Surrendered

Authorised in:

Ireland

Package description:

High density polyethylene back-packs of 5 litre nominal volume. Closure: Polypropylene or urea formaldehyde screw cap with wads of PVDC-faced paper on a pulpboard liner.

High density polyethylene back-packs of 2.5 litre nominal volume. Closure: Polypropylene or urea formaldehyde screw cap with wads of PVDC-faced paper on a pulpboard liner.

High density polyethylene back-packs of 1 litre nominal volume. Closure: Polypropylene or urea formaldehyde screw cap with wads of PVDC-faced paper on a pulpboard liner.

High density polyethylene bottles of 10 litre nominal volume. Closure: Polypropylene or urea formaldehyde screw cap with wads of PVDC-faced paper on a pulpboard liner.

High density polyethylene bottles of 5 litre nominal volume. Closure: Polypropylene or urea formaldehyde screw cap with wads of PVDC-faced paper on a pulpboard liner.

High density polyethylene bottles of 2.5 litre nominal volume. Closure: Polypropylene or urea formaldehyde screw cap with wads of PVDC-faced paper on a pulpboard liner.

High density polyethylene bottles of 1 litre nominal volume. Closure: Polypropylene or urea formaldehyde screw cap with wads of PVDC-faced paper on a pulpboard liner.

Additional information

Entitlement type:

Marketing Authorisation

Legal basis of product authorisation:

Complete application (stand-alone) - Council Directive 81/851/EEC

Marketing authorisation holder:

Intervet (Ireland) Limited

Marketing authorisation date:

1/10/1999

Manufacturing sites for batch release:

Trirx Segre

Responsible authority:

Health Products Regulatory Authority

Authorisation number:

VPA10996/262/001

Date of authorisation status change:

14/05/2025

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

Documents

Summary of Product Characteristics