

Tricaine Pharmaq 1000 mg/g Powder for Solution for Fish Treatment

Authorised

- Tricaine mesilate

Product identification

Medicine name:

Tricaine Pharmaq 1000 mg/g Powder for Solution for Fish Treatment

Active substance:

Tricaine mesilate

Target species:

Ornamental fish
Other fish

Route of administration:

Dipping

Product details

Active substance and strength:

Tricaine mesilate
1000.00 milligram(s) / 1.00 gram(s)

Pharmaceutical form:

Powder for solution for fish treatment

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

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Ornamental fish

- Fish meat. 70 degree day

Fish must not be slaughtered for human consumption during treatment. Fish can only be harvested for human consumption 70 degree days after the last treatment.

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Other fish

- Fish meat. 70 degree day

Fish must not be slaughtered for human consumption during treatment. Fish can only be harvested for human consumption 70 degree days after the last treatment.

Anatomical therapeutic chemical veterinary (ATCvet) codes:

QN01AX93

Legal status of supply:

Veterinary medicinal product subject to veterinary prescription

Authorisation status:

Valid

Authorised in:

United Kingdom (Northern Ireland)

Package description:

High Density Polyethylene (HDPE) tamper resistant tubs closed with an integral, tamper evident, low density polyethylene cap (snap on) or polypropylene screw cap containing 1000 g.

High Density Polyethylene (HDPE) tamper resistant tubs closed with an integral, tamper evident, low density polyethylene cap (snap on) or polypropylene screw cap containing 250 g.

High Density Polyethylene (HDPE) tamper resistant tubs closed with an integral, tamper evident, low density polyethylene cap (snap on) or polypropylene screw cap containing 100 g.

High Density Polyethylene (HDPE) tamper resistant tubs closed with an integral, tamper evident, low density polyethylene cap (snap on) or polypropylene screw cap containing 25 g.

Additional information

Entitlement type:

Marketing Authorisation

Legal basis of product authorisation:

Generic application (Article 13(1) of Directive No 2001/82/EC)

Marketing authorisation holder:

Pharmaq AS

Marketing authorisation date:

7/02/2013

Manufacturing sites for batch release:

Pharmaq Limited

Responsible authority:

The Veterinary Medicines Directorate

Authorisation number:

Vm 21714/3002

Date of authorisation status change:

20/11/2024

Reference member state:

Norway

Procedure number:

NO/V/0012/001

Concerned member states:

Greece Iceland Ireland Italy Spain United Kingdom (Northern Ireland)

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

Documents

Combined File of all Documents