IRISH MEDICINES BOARD ACT 1995

EUROPEAN COMMUNITIES (ANIMAL REMEDIES) (No. 2) REGULATIONS 2007

(S.I. No. 786 of 2007)

VPA: **10960/033/001** Case No: 7007059

The Irish Medicines Board in exercise of the powers conferred on it by Animal Remedies (No. 2) Regulations (S.I. No. 786 of 2007) hereby grants to:

Cross Vetpharm Group Ltd.

Broomhill Road, Dublin 24, Ireland

an authorisation, subject to the provisions of the said Regulations and the general conditions of the attached authorisation, in respect of the Veterinary Medicinal Product:

Bimectin 1.0 %w/v Solution for Injection

The particulars of which are set out in Part 1 and Part 2 of the said Schedule. The authorisation is also subject to any special conditions as may be specified in the said Schedule.

The authorisation, unless revoked, shall continue in force from 12/01/2010 until 11/01/2015.

Signed on behalf of the Irish Medicines Board

A person authorised in that behalf by the said Board.

(NOTE: This authorisation replaces any previous authorisation in respect of this product which is now null and void.)

Part II

Summary of Product Characteristics

1 NAME OF THE VETERINARY MEDICINAL PRODUCT

Bimectin 1% w/v Solution for Injection.

2 QUALITATIVE AND QUANTITATIVE COMPOSITION

Active Substances

Ivermectin 1.0 % w/v For a full list of excipinets, see section 6.1.

3 PHARMACEUTICAL FORM

Solution for injection.

4 CLINICAL PARTICULARS

4.1 Target Species

Cattle, sheep and pigs.

4.2 Indications for use, specifying the target species

For the effective treatment and control of the following harmful parasites of cattle, sheep and pigs:

Cattle

Gastrointestinal roundworms (adult and fourth-stage larvae):

Ostertagia spp. (including inhibited O. ostertagi)

Haemonchus placei

Trichostrongylus axei

T. colubriformis

Cooperia spp.

Bunostomum phlebotomum

Oesophagostomum radiatum

Strongyloides papillosus (adult)

Nematodirus helvetianus (adult)

N. spathiger (adult)

Trichuris spp (adult).

Lungworms (adult and fourth-stage larvae):

Dictyocaulus viviparus

Eye worms (adult):

Thelazia spp.

Warbles:

Hypoderma bovis

H. lineatum

Mange mites:

Psoroptes bovis Sarcoptes scabiei var. bovis

Suckling lice:

Linognathus vituli Haematopinus eurysternus Solenopotes capillatus

May also be used as an aid in the control of the mange mite Chorioptes bovis and biting lice Damalinia bovis, but complete elimination may not occur.

Persistent Activity:

Treatment at the recommended dose rate can control re-infection with Haemonchus placei and Cooperia spp. acquired up to 14 days after treatment, Ostertagia ostertagi and Oesophagostomum radiatum acquired up to 21 days after treatment and *Dictyocaulus viviparus* up to 28 days after treatment.

To obtain optimal benefit from the persistent activity of the product for grazing animals, it is recommended that calves which are set-stocked in the first grazing season should be treated 3, 8 and 13 weeks after the day of turn-out. This can protect the animals from parasitic gastroenteritis and lungworm disease throughout the grazing season, provided they are set-stocked, all the calves included in the programme and that no untreated cattle are added to the pasture. Treated animals should always be monitored according to good husbandry practices.

Sheep

Gastrointestinal roundworms (adult and fourth-stage larvae):

Ostertagia circumcincta including inhibited larvae

O. trifurcata

Haemonchus contortus including inhibited larvae

Trichostrongylus axei (adults)

T. colubriformis and T. vitrinus (adults)

Cooperia curticei

Oesophagostomum columbianum

O. venulosum (adults)

Nematodirus filicollis

Chabertia ovina

Trichuris ovis (adults).

Lungworms:

Dictyocaulus filaria (adult and fourth-stage larvae)

Protostrongylus rufescens (adults)

Nasal Bots (all larval stages)

Oestrus ovis

Pigs

Gastrointestinal roundworms (adult and fourth-stage larvae):

Ascaris suum

Hyostrongylus rubidus

Oesophagostomum spp.

Strongyloides ransomi (adult and somatic larval stages)

Lungworms:

Metastrongylus spp. (adults)

Lice:

Haematopinus suis

Mange mites:

Sarcoptes scabiei var. suis

4.3 Contraindications

Do not use in animals with known hypersensitivity to the active ingredient. Do not use by intramuscular or intravenous administration.

4.4 Special warnings for each target species

None

4.5 Special precautions for use

(i) Special Precautions for use in animals

The product has been formulated specifically for use in cattle, sheep and pigs. It should not be used in other species as severe adverse reactions, including fatalities in dogs, may occur.

Frequent and repeated use may lead to the development of resistance. It is important that the correct dose is given in order to minimise the risk of resistance. To avoid under-dosing, animals should be grouped according to their bodyweight and dosed according to the dose of the heaviest animal in the group.

(ii) Special Precautions to be taken by the Person Administering the Product to Animals

Take care to avoid self-administration: the product may cause local irritation and/or pain at the site of injection.

Direct contact of the product with the skin should be kept to a minimum.

Do not smoke or eat while handling the product.

Wash hands after use.

(iii) Other precautions

When using the 250 ml and 500 ml pack sizes, use only automatic syringe equipment. For the 50 ml pack size, use of a multiple dose syringe is recommended. To refill the syringe, use of a draw off needle is recommended to avoid excessive broaching of the stopper.

4.6 Adverse reactions (frequency and seriousness)

Cattle

Transient discomfort has occasionally been observed in cattle following subcutaneous administration. A low incidence of soft tissue swelling at the injection site has been observed.

Sheep

Discomfort, sometimes intense but usually transient, has been observed in some sheep immediately following subcutaneous administration.

Pigs

Mild and transient discomfort has occasionally been observed in pigs following subcutaneous injection.

All these reactions disappeared without treatment.

4.7 Use during pregnancy, lactation or lay

Pregnancy

The product can be administered to beef cows, sheep and pigs at any stage of pregnancy

Lactation

Do not use in dairy cows or sheep producing milk for human consumption

Do not use in non-lactating dairy cows or sheep within 60 days of calving/lambing. The product can be used in sows during lactation.

<u>Fertility</u>

Fertility is not affected by administration of the product.

4.8 Interaction with other medicinal products and other forms of interaction

The product can be used concurrently without adverse effects with foot and mouth disease vaccine or clostridial vaccine, given at separate injection sites

4.9 Amounts to be administered and administration route

The product should be given only by subcutaneous injection at the recommended dosage level of 200 mcg ivermectin per kg bodyweight under the loose skin in front of, or behind, the shoulder in cattle and over the neck in sheep. At the recommended dosage level of 300 mcg ivermectin per kg of bodyweight, the product should be given only subcutaneously in the neck of pigs.

Each ml contains 10 mg of ivermectin sufficient to treat 50 kg of bodyweight of cattle and sheep and 33 kg of bodyweight of pigs. The injection may be given with any standard automatic or single-dose or hypodermic syringe. Use of 17 gauge x ½ inch needle is suggested. Replace with a fresh sterile needle after every 10 to 12 animals. Injection of wet or dirty animals is not recommended. If using a single-dose or hypodermic syringe, use a separate sterile needle to withdraw the product from the container. Massage the injection site after administration of the product.

In young pigs, especially those below 16 kg for which less than 0.5 ml of the product is indicated, dosing accuracy is important. The use of a syringe that can accurately deliver as little as 0.1 ml is recommended.

In young lambs weighing less than 20.0 kg give 0.1 ml per 5 kg. In these lambs the use of a syringe with can deliver as little as 0.1 ml is recommended.

4.10 Overdose (symptoms, emergency procedures, antidotes), if necessary

Cattle

Single doses of 4.0 mg ivermectin per kg (20 x the use level) given subcutaneously resulted in ataxia and depression.

Sneep

At dose levels up to 4 mg ivermectin per kg (20 x the use level) given subcutaneously resulted in ataxia and depression. No signs of systemic toxicity were observed in sheep treated with the product at up to 3 times the recommended dose rate, soft tissue swellings at the injection site were observed.

Pigs

A dose of 30 mg ivermectin per kg (100 x the recommended dose of 0.3 mg per kg) injected subcutaneously to pigs caused lethargy, ataxia, bilateral mydriasis, intermittent tremors, laboured breathing and lateral recumbancy.

In the case of overdosage, symptomatic treatment should be given.

4.11 Withdrawal Period(s)

Cattle must not be treated within 49 days of slaughter for human consumption. Do not use in lactating cows producing milk for human consumption. Do not use in non-lactating dairy cows including pregnant dairy heifers within 60 days of calving.

Sheep must not be treated within 42 days of slaughter for human consumption. Do not use in lactating sheep producing milk for human consumption. Do not use in sheep within 60 days of lambing where milk is to be used for human consumption.

Pigs must not be treated within 28 days of slaughter for human consumption.

5 PHARMACOLOGICAL or IMMUNOLOGICAL PROPERTIES

ATC vet code: QP54AA01 Therapeutic group: Endectocide

5.1 Pharmacodynamic properties

Ivermectin is a member of the macrocyclic lactone class of endectocides which have a unique mode of action. Compounds of the class bind selectively and with high affinity to glutamate-gated chloride ion channels which occur in invertebrate nerve and muscle cells. This leads to an increase in the permeability of the cell membrane to chloride ions with hyperpolarization of the nerve or muscle cell, resulting in paralysis and death of the parasite. Compounds of this class may also interact with other ligand-gated chloride channels, such as those gated by the neurotransmitter gamma-aminobutyric acid (GABA).

The margin of safety for compounds of this class is attributable to the fact that mammals do not have glutamate-gated chloride channels, the macrocyclic lactones have a low affinity for other mammalian ligand-gated chloride channels and they do not readily cross the blood-brain barrier.

5.2 Pharmacokinetic properties

Maximum plasma concentration:

Cattle:

At a dose level of 0.2 mg ivermectin per kg a maximum plasma concentration of 35-50 ng/ml is reached in \pm 2 days and the half-life in plasma is 2.8 days. It is also established that ivermectin is carried mainly in the plasma (80%). This distribution between plasma and blood cells remains relatively constant.

Sheep:

At a dose of 0.3 mg ivermectin per kg an average peak of 16 ng/ml is reached one day after injection.

Pigs:

During trials carried out at a dose rate of 0.3 mg ivermectin per kg bodyweight, peak plasma concentrations were reached in 3 ± 0.5 days and the drug persisted in plasma for up to 28 days.

Excretion: length of time and route

<u>Cattle:</u>

Only about 1-2% is excreted in the urine the remainder is excreted in the faeces, approximately 60% of which is excreted as unaltered drug. The remainder is excreted as metabolites or degradation products.

Sheep

Radioactive ivermectin was administered to sheep at a dose rate of 0.3 mg per kg. Analyses of the faeces showed that about 99% of the drug and its metabolites are excreted in the faeces, +/- 1% being excreted in the urine.

<u>Pigs</u>:

Biliary excreation is also the major route of ivermectin excretion in pigs.

6 PHARMACEUTICAL PARTICULARS

6.1 List of excipients

Glycerin formal

6.2 Incompatibilities

Do not mix with other medicinal products.

6.3 Shelf-life

Shelf-life of the veterinary medicinal product as packaged for sale: 3 years. Shelf-life after first opening the immediate packaging: 28 days.

6.4 Special precautions for storage

None.

6.5 Nature and composition of immediate packaging

Multiple-dose polyethylene bottles of 50 ml, 250 ml and 500 ml sealed with bromobutyl seals and aluminium overseals, each containing a clear, colourless sterile solution.

6.6 Special precautions for the disposal of unused veterinary medicinal products or waste materials

Any unused veterinary product or waste material derived from the product should be disposed of in accordance with local requirements. The product should not enter water courses as this may be dangerous to fish and other aquatic organisms.

7 MARKETING AUTHORISATION HOLDER

Cross Vetpharm Group Limited, Broomhill Road, Tallaght, Dublin 24, Ireland

8 MARKETING AUTHORISATION NUMBER(S)

VPA 10960/033/001

9 DATE OF THE FIRST AUTHORISATION/RENEWAL OF THE AUTHORISATION

19th May 2005

10 DATE OF REVISION OF THE TEXT

13th January 2010