ANNEX I

SUMMARY OF PRODUCT CHARACTERISTICS
1. NAME OF THE VETERINARY MEDICINAL PRODUCT

HALOCUR 0.5 mg/ml oral solution for calves

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Active substance:

Halofuginone base 0.50 mg/ml  
(as lactate salt)

Excipients:

Benzoic acid (E 210) 1.00 mg/ml  
Tartrazine (E 102) 0.03 mg/ml

For a full list of excipients, see section 6.1

3. PHARMACEUTICAL FORM

Oral solution
Canary yellow homogenous clear solution.

4. CLINICAL PARTICULARS

4.1 Target species

New born calves.

4.2 Indications for use, specifying the target species

Prevention of diarrhoea due to diagnosed Cryptosporidium parvum, in farms with history of cryptosporidiosis. Administration should start in the first 24 to 48 hours of age.

Reduction of diarrhoea due to diagnosed Cryptosporidium parvum. Administration should start within 24 hours after the onset of diarrhoea.

In both cases, the reduction of oocysts excretion has been demonstrated.

4.3 Contraindications

Do not use on an empty stomach.
Do not use in case of diarrhoea established for more than 24 hours and in weak animals.

4.4 Special warnings

None.

4.5 Special precautions for use

Special precautions for use in animals
Administer after colostrum feeding, or after milk or milk replacer feeding only, using either a syringe or any appropriate device for oral administration. Do not use on an empty stomach. For treatment of anorexic calves, the product should be administered in half a litre of an electrolyte solution. The animals should receive enough colostrum according to good breeding practice.

Special precautions to be taken by the person administering the veterinary medicinal product to animals

Repetitive contact with the product may lead to skin allergies. Avoid skin, eye or mucosal contact with the product. Wear protective gloves while handling the product. In case of skin and eye contact wash the exposed area thoroughly with clean water. If eye irritation persists, seek medical advice. Wash hands after use.

4.6 Adverse reactions (frequency and seriousness)

An increase in the level of diarrhoea has been observed in treated animals in very rare cases.

The frequency of adverse reactions is defined using the following convention:
- very common (more than 1 in 10 animals treated displaying adverse reaction(s))
- common (more than 1 but less than 10 animals in 100 animals treated)
- uncommon (more than 1 but less than 10 animals in 1,000 animals treated)
- rare (more than 1 but less than 10 animals in 10,000 animals treated)
- very rare (less than 1 animal in 10,000 animals treated, including isolated reports).

4.7 Use during pregnancy, lactation or lay

Not applicable.

4.8 Interaction with other medicinal products and other forms of interaction

None known.

4.9 Amounts to be administered and administration route

For oral use in calves after feeding.

The dosage is: 100 µg of halofuginone base / kg body weight (BW) / once a day for 7 consecutive days, i.e. 2 ml of HALOCUR / 10 kg BW / once a day for 7 consecutive days.

However, in order to make the HALOCUR treatment easier, a simplified dosage scheme is proposed:

- 35 kg < calves ≤ 45 kg: 8 ml of HALOCUR once a day during 7 consecutive days
- 45 kg < calves < 60 kg: 12 ml of HALOCUR once a day during 7 consecutive days

For smaller or higher weights, a precise calculation should be performed (2 ml/10 kg BW).
To ensure a correct dosage, the use of either a syringe or any appropriate device for oral administration is necessary.

The consecutive treatment should be done at the same time each day.

Once the first calf has been treated, all the forthcoming new-born calves must be systematically treated as long as the risk for diarrhoea due to C. parvum persists.

4.10 Overdose (symptoms, emergency procedures, antidotes), if necessary
As symptoms of toxicity may occur at twice the therapeutic dose, it is necessary to apply the recommended dosage strictly. Symptoms of toxicity include diarrhoea, visible blood in faeces, decline in milk consumption, dehydration, apathy and prostration. Should clinical signs of overdosing occur the treatment must be stopped immediately and the animal fed unmedicated milk or milk replacer. Rehydration may be necessary.

4.11 Withdrawal period(s)

Meat and offal: 13 days.

5. PHARMACOLOGICAL PROPERTIES

Pharmacotherapeutic group: Quinazolinone derivate
ATCvet code:QP51AX08

5.1 Pharmacodynamic properties

The active substance, halofuginone, is an antiprotozoal agent of the quinazolinone derivatives group (nitrogenous polyheterocycles). Halofuginone lactate (RU 38788) is a salt whose antiprotozoal properties and efficacy against Cryptosporidium parvum have been demonstrated both in in vitro conditions and in artificial and natural infections. The compound has a cryptosporidiostatic effect on Cryptosporidium parvum. It is mainly active on the free stages of the parasite (sporozoïte, merozoïte). The concentrations to inhibit 50 % and 90 % of the parasites, in an in vitro test system, are IC$_{50}$ < 0.1 µg/ml and IC$_{90}$ of 4.5 µg/ml respectively.

5.2 Pharmacokinetic particulars

The bioavailability of the drug in the calf, following single oral administration, is about 80 %. The time necessary to obtain the maximum concentration T$_{max}$ is 11 hours. The maximum concentration in plasma C$_{max}$ is 4 ng/ml. The apparent volume of distribution is 10 l/kg. The plasmatic concentrations of halofuginone after repeated oral administrations are comparable to the pharmacokinetic pattern after single oral treatment. Unchanged Halofuginone is the major component in the tissues. Highest values have been found in the liver and the kidney. The product is mainly excreted in the urine. The terminal elimination half-life is 11.7 hours after intravenous administration and 30.84 hours after single oral administration.

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

Benzoic acid (E 210)
Tartrazine (E 102)
Lactic acid (E 270)
Water, purified

6.2 Major incompatibilities

None known.

6.3 Shelf life

Shelf-life of the veterinary medicinal product as packaged for sale: 3 years. Shelf-life after first opening the container: 6 months.
6.4 Special precautions for storage

This medicinal product does not require any special storage conditions.

6.5 Nature and composition of immediate packaging

High-density polyethylene portable bottle of 500 ml containing 490 ml of the oral solution. Not all pack sizes may be marketed.

6.6 Special precautions for the disposal of unused veterinary medicinal product or waste materials derived from the use of such products

Any unused veterinary medicinal product or waste materials derived from such veterinary medicinal products should be disposed of in accordance with the local requirements. HALOCUR should not enter water courses, as this may be dangerous for fish and other aquatic organisms.

7. MARKETING AUTHORISATION HOLDER

Intervet International B.V. 
Wim de Körverstraat 35 
5831 AN Boxmeer 
The Netherlands

8. MARKETING AUTHORISATION NUMBER(S)

EU/2/99/013/001-002

9. DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORISATION

Date of first authorisation: 29 October 2004 
Date of last renewal: 23 November 2009

10 DATE OF REVISION OF THE TEXT

Detailed information on this veterinary medicinal product is available on the website of the European Medicines Agency http://www.emea.europa.eu/.

PROHIBITION OF SALE, SUPPLY AND/OR USE

Not applicable.
ANNEX II

A. MANUFACTURER RESPONSIBLE FOR BATCH RELEASE

B. CONDITIONS OR RESTRICTIONS REGARDING SUPPLY AND USE

C. STATEMENT OF THE MRLs
A. MANUFACTURER RESPONSIBLE FOR BATCH RELEASE

Name and address of the manufacturer responsible for batch release

Intervet Productions SA
Rue de Lyons
F-27460 Igoville
France

B. CONDITIONS OR RESTRICTIONS OF THE MARKETING AUTHORISATION REGARDING SUPPLY OR USE

To be supplied only on veterinary prescription.

C. STATEMENT OF THE MRLs

The active substance in HALOCUR is an allowed substance as described in table 1 of the annex to Commission Regulation (EU) No 37/2010:

<table>
<thead>
<tr>
<th>Pharmaco-logically active substance</th>
<th>Marker residue</th>
<th>Animal species</th>
<th>MRLs</th>
<th>Target tissues</th>
<th>Other provisions</th>
<th>Therapeutic classification</th>
</tr>
</thead>
<tbody>
<tr>
<td>Halofuginone</td>
<td>Halofuginone</td>
<td>Bovine</td>
<td>30 µg/kg</td>
<td>Liver</td>
<td>Not for use in animals from which milk is produced for human consumption.</td>
<td>Antiparasitic agents/Agents acting against protozoa</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>30 µg/kg</td>
<td>Kidney</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>10 µg/kg</td>
<td>Muscle</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>25 µg/kg</td>
<td>Fat</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The excipients listed in section 6.1 of the SPC are allowed substances for which table 1 of the annex to Commission Regulation (EU) No 37/2010 indicates that no MRLs are required.
ANNEX III

LABELLING AND PACKAGE LEAFLET
A. LABELLING
<table>
<thead>
<tr>
<th>PARTICULARS TO APPEAR ON THE IMMEDIATE PACKAGE</th>
</tr>
</thead>
<tbody>
<tr>
<td>HDPE container (500 ml and 1000 ml presentations)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>1. NAME OF THE VETERINARY MEDICINAL PRODUCT</th>
</tr>
</thead>
<tbody>
<tr>
<td>HALOCUR 0.5 mg/ml oral solution for calves</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>2. STATEMENT OF ACTIVE SUBSTANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Halofuginone base (as lactate salt) 0.5 mg/ml</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>3. PHARMACEUTICAL FORM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oral solution</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>4. PACKAGE SIZE</th>
</tr>
</thead>
<tbody>
<tr>
<td>500 ml bottle containing 490 ml of oral solution</td>
</tr>
<tr>
<td>1000 ml bottle containing 980 ml of oral solution</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>5. TARGET SPECIES</th>
</tr>
</thead>
<tbody>
<tr>
<td>New born calves</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>6. INDICATION(S)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Read package leaflet before use.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>7. METHOD AND ROUTE(S) OF ADMINISTRATION</th>
</tr>
</thead>
<tbody>
<tr>
<td>For oral use in new born calves after feeding.</td>
</tr>
<tr>
<td>Read the package leaflet before use.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>8. WITHDRAWAL PERIOD(S)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Withdrawal period: Meat and offal: 13 days.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>9. SPECIAL WARNING(S), IF NECESSARY</th>
</tr>
</thead>
<tbody>
<tr>
<td>Read the package leaflet before use.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>10. EXPIRY DATE</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
</tr>
</tbody>
</table>
EXP: month/year
Once broached, use within 6 months.

11. SPECIAL STORAGE CONDITIONS

12. SPECIAL PRECAUTIONS FOR THE DISPOSAL OF UNUSED PRODUCT OR WASTE MATERIALS, IF ANY

Disposal: read package leaflet.

13. THE WORDS “FOR ANIMAL TREATMENT ONLY” AND CONDITIONS OR RESTRICTIONS REGARDING SUPPLY AND USE, IF APPLICABLE

For animal treatment only. To be supplied only on veterinary prescription.

14. THE WORDS “KEEP OUT OF THE SIGHT AND REACH OF CHILDREN”

Keep out of the sight and reach of children.

15. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER

Intervet International B.V.
Wim de Körverstraat 35
NL-5831 AN Boxmeer
The Netherlands

16. MARKETING AUTHORISATION NUMBER(S)

EU/2/99/013/001-002

17. MANUFACTURER’S BATCH NUMBER

<Batch> <Lot> <BN> number.
B. PACKAGE LEAFLET
1. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER AND OF THE MANUFACTURING AUTHORISATION HOLDER RESPONSIBLE FOR BATCH RELEASE, IF DIFFERENT

Marketing authorisation holder:
Intervet International B.V.
Wim de Körverstraat 35
5831 AN Boxmeer
The Netherlands

Manufacturer responsible for batch release:
Intervet Productions S.A.
Rue de Lyons
27460 Igoville
France

2. NAME OF THE VETERINARY MEDICINAL PRODUCT

HALOCUR 0.5 mg/ml oral solution for calves

3. STATEMENT OF THE ACTIVE SUBSTANCE AND OTHER INGREDIENTS

The veterinary medicinal product is a canary yellow oral solution. HALOCUR contains 0.5 mg/ml halofuginone base (as lactate salt).

4. INDICATION(S)

Prevention of diarrhoea due to diagnosed Cryptosporidium parvum, in farms with history of cryptosporidiosis. Administration should start in the first 24 to 48 hours of age.

Reduction of diarrhoea due to diagnosed Cryptosporidium parvum. Administration should start within 24 hours after the onset of diarrhoea.

In both cases, the reduction of oocysts excretion has been demonstrated

5. CONTRAINDICATIONS

Do not use on an empty stomach.
Do not use in case of diarrhoea established for more than 24 hours and in weak animals.

6. ADVERSE REACTIONS

An increase in the level of diarrhoea has been observed in treated animals in very rare cases.

The frequency of adverse reactions is defined using the following convention:
- very common (more than 1 in 10 animals treated displaying adverse reaction(s))
- common (more than 1 but less than 10 animals in 100 animals treated )
- uncommon (more than 1 but less than 10 animals in 1,000 animals treated)
- rare (more than 1 but less than 10 animals in 10,000 animals treated)
- very rare (less than 1 animal in 10,000 animals treated, including isolated reports).

If you notice any side effects, even those not already listed in this package leaflet or you think that the medicine has not worked, please inform your veterinary surgeon.

7. TARGET SPECIES

New born calves.

8. DOSAGE FOR EACH SPECIES, ROUTE(S) AND METHOD OF ADMINISTRATION

For oral use in calves after feeding.

The dosage is: 100 µg of halofuginone base / kg body weight (BW) / once a day for 7 consecutive days, i.e. 2 ml of HALOCUR / 10 kg BW / once a day for 7 consecutive days.

However, in order to make the HALOCUR treatment easier, a simplified dosage scheme is proposed:

- 35 kg < calves ≤ 45 kg: 8 ml of HALOCUR once a day during 7 consecutive days
- 45 kg < calves < 60 kg: 12 ml of HALOCUR once a day during 7 consecutive days

For smaller or higher weights, a precise calculation should be performed (2 ml/10 kg BW).

9. ADVICE ON CORRECT ADMINISTRATION

To ensure a correct dosage, the use of either a syringe or any appropriate device for oral administration is necessary.

The consecutive treatment should be done at the same time each day.

Once the first calf has been treated, all the forthcoming new-born calves must be systematically treated as long as the risk for diarrhoea due to *C. parvum* persists.

10. WITHDRAWAL PERIOD(S)

Meat and offal: 13 days.

11. SPECIAL STORAGE PRECAUTIONS

Keep out of the sight and reach of children.

This medicinal product does not require any special storage conditions.

Shelf-life after first opening the container: 6 months.

12. SPECIAL WARNING(S)

Special precautions for use in animals:
Administer after colostrum feeding, or after milk or milk replacer feeding only, using either a syringe or any appropriate device for oral administration. Do not use on an empty stomach. For treatment of anorexic calves, the product should be administered in half a litre of an electrolyte solution. The animals should receive enough colostrum according to good breeding practice.

**Special precautions to be taken by the person administering the veterinary medicinal product to animals:**

Repetitive contact with the product may lead to skin allergies. Avoid skin, eye or mucosal contact with the product. Wear protective gloves while handling the product.

In case of skin and eye contact wash the exposed area thoroughly with clean water. If eye irritation persists, seek medical advice.

Wash hands after use.

**Overdose (symptoms, emergency procedures, antidotes):**

As symptoms of toxicity may occur at twice the therapeutic dose, it is necessary to apply the recommended dosage strictly. Symptoms of toxicity include diarrhoea, visible blood in faeces, decline in milk consumption, dehydration, apathy and prostration. Should clinical signs of overdosing occur the treatment must be stopped immediately and the animal fed unmedicated milk or milk replacer. Rehydration may be necessary.

**13. SPECIAL PRECAUTIONS FOR THE DISPOSAL OF UNUSED PRODUCT OR WASTE MATERIALS, IF ANY**

Medicines should not be disposed of via wastewater.
Ask your veterinary surgeon how to dispose of medicines no longer required. These measures should help to protect the environment.
HALOCUR should not enter water courses as this may be dangerous for fish and other aquatic organisms.

**14. DATE ON WHICH THE PACKAGE LEAFLET WAS LAST APPROVED**

DD/MM/YYYY
Detailed information on this veterinary medicinal product is available on the website of the European Medicines Agency (http://www.ema.europa.eu/).

**15. OTHER INFORMATION**

High-density polyethylene portable bottle of 500 ml containing 490 ml of the oral solution. High-density polyethylene portable bottle of 1000 ml containing 980 ml of the oral solution.

Not all pack sizes may be marketed.