ANNEX I SUMMARY OF PRODUCT CHARACTERISTICS

1. NAME OF THE VETERINARY MEDICINAL PRODUCT

Innovax-ND-IBD concentrate and solvent for suspension for injection for chickens

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Each dose of the reconstituted vaccine (0.2 ml for subcutaneous use or 0.05 ml for *in ovo* use) contains:

Active substance:

Cell-associated live recombinant turkey herpesvirus (strain HVP360), expressing the fusion protein of Newcastle disease virus and the VP2 protein of infectious bursal disease virus: $10^{3.3} - 10^{4.6} \text{ PFU}^1$.

Excipients:

Qualitative composition of excipients and other constituents	
Concentrate:	
Bovine serum	
Veggie medium	
Dimethyl sulfoxide	
Solvent:	
Sucrose	
Sodium chloride	
Disodium hydrogen phosphate dihydrate	
Phenolsulfonphthalein (Phenol red)	
Potassium dihydrogen phosphate	
Water for injections	

Concentrate: off-red to red cell concentrate.

Solvent: clear, red solution.

3. CLINICAL INFORMATION

3.1 Target species

Chickens and embryonated chicken eggs.

3.2 Indications for use for each target species

For active immunisation of one-day-old chicks or 18–19 day-old embryonated chicken eggs:

- to reduce mortality and clinical signs caused by Newcastle disease (ND) virus,
- to prevent mortality and to reduce clinical signs and lesions caused by infectious bursal disease (IBD) virus.
- to reduce mortality, clinical signs and lesions caused by Marek's disease (MD) virus.

Onset of immunity: ND: 4 weeks of age

¹PFU – plaque forming units.

IBD: 3 weeks of age

MD: 9 days

Duration of immunity: ND: 60 weeks

IBD: 60 weeks

MD: entire risk period

3.3 Contraindications

None.

3.4 Special warnings

Vaccinate healthy animals only.

3.5 Special precautions for use

Special precautions for safe use in the target species:

As this is a live vaccine, the vaccine strain is excreted from vaccinated birds and may spread to turkeys. Safety trials have shown that the strain is safe for turkeys. However, precautionary measures have to be followed in order to avoid direct or indirect contact between vaccinated chickens and turkeys.

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

The handling of liquid nitrogen should take place in a well-ventilated area.

Innovax-ND-IBD is a virus suspension packed in glass ampoules and stored in liquid nitrogen. Before withdrawing ampoules from the liquid nitrogen canister, protective equipment consisting of gloves, long sleeves and a facemask or goggles should be worn. In order to prevent serious wounds, by either the liquid nitrogen or the ampoules when removing an ampoule from the canister, hold the palm of the (gloved) hand holding the ampoule away from the body and face. Care should be exercised to prevent contaminating the hands, eyes and clothing with the ampoule content. CAUTION: The ampoules have been known to explode on exposure to sudden temperature changes. Do not thaw in hot water or ice-cold water. Thaw the ampoules in clean water at 25 $^{\circ}$ C – 27 $^{\circ}$ C.

Special precautions for the protection of the environment:

Not applicable.

3.6 Adverse events

None known.

Reporting adverse events is important. It allows continuous safety monitoring of a veterinary medicinal product. Reports should be sent, preferably via a veterinarian, to either the marketing authorisation holder or the national competent authority via the national reporting system. See section "Contact details" of the package leaflet.

3.7 Use during pregnancy, lactation or lay

The safety of the veterinary medicinal product has not been established during lay.

3.8 Interaction with other medicinal products and other forms of interaction

Safety and efficacy data are available which demonstrate that Innovax-ND-IBD can be mixed in the same solvent and administered by the subcutaneous route with Nobilis Rismavac. For this mixed use, an onset of immunity of 5 days has been demonstrated for MD.

Safety and efficacy data are available which demonstrate that Nobilis ND Clone 30 or Nobilis ND C2 or Nobilis IB Ma5 or Nobilis IB 4-91 can be administered (not mixed) to day-old chicks that are vaccinated either by the subcutaneous or by the *in ovo* route with Innovax-ND-IBD. For such associated use, an onset of immunity of 3 weeks (when used with Nobilis ND Clone 30) and 2 weeks (when used with Nobilis ND C2), has been demonstrated for ND.

No information is available on the safety and efficacy of this vaccine when used with any other veterinary medicinal product except the products mentioned above. A decision to use this vaccine before or after any other veterinary medicinal product therefore needs to be made on a case by case basis.

3.9 Administration routes and dosage

Posology:

Subcutaneous use: one single injection of 0.2 ml per chick. *In ovo*: one single injection of 0.05 ml per chicken egg.

Preparation of the vaccine:

The usual aseptic precautions should be applied to all preparation and administration procedures. The handling of liquid nitrogen should take place in a well-ventilated area.

1. Use solvent for cell associated poultry vaccines for reconstitution.

For subcutaneous use reconstitute the vaccine according to the table below:

Solvent bag	Number of vaccine ampoules for subcutaneous use
Bag of 400 ml solvent	1 ampoule containing 2000 doses
Bag of 800 ml solvent	2 ampoules containing 2000 doses
Bag of 800 ml solvent	1 ampoule containing 4000 doses
Bag of 1200 ml solvent	3 ampoules containing 2000 doses
Bag of 1600 ml solvent	4 ampoules containing 2000 doses
Bag of 1600 ml solvent	2 ampoules containing 4000 doses

When this product is mixed with Nobilis Rismavac, both should be diluted in the same solvent bag in the same way (400 ml of solvent for each 2000 doses of both products or 800 ml of solvent for each 4000 doses of both products).

For *in ovo* use reconstitute the vaccine according to the table below:

Solvent bag	Number of vaccine ampoules for in ovo use
Bag of 400 ml solvent	4 ampoules containing 2000 doses
Bag of 400 ml solvent	2 ampoules containing 4000 doses
Bag of 800 ml solvent	8 ampoules containing 2000 doses
Bag of 800 ml solvent	4 ampoules containing 4000 doses
Bag of 1200 ml solvent	12 ampoules containing 2000 doses
Bag of 1200 ml solvent	6 ampoules containing 4000 doses
Bag of 1600 ml solvent	16 ampoules containing 2000 doses
Bag of 1600 ml solvent	8 ampoules containing 4000 doses

The solvent must be clear, red coloured, without sediment and at room temperature (15 $^{\circ}$ C -25 $^{\circ}$ C) at the time of mixing.

2. Preparation of the vaccine shall be planned before the ampoules are taken from the liquid nitrogen and the exact amount of vaccine ampoules and amount of solvent needed shall be

- calculated first. There is no information available on the number of doses on the ampoules once they are removed from the cane, so special care has to be taken to ensure that the mix-ups of ampoules with different number of doses is avoided and the correct solvent is used.
- 3. Before withdrawing the ampoules from the liquid nitrogen container, protect the hands with gloves, wear long sleeves and use a facemask or goggles. When removing an ampoule from the cane, hold in the palm of a gloved hand away from the body and the face.
- 4. When withdrawing a cane of ampoules from the canister in the liquid nitrogen container, expose only the ampoule(s) to be used immediately. It is recommended to handle a maximum of 5 ampoules (from one cane only) at a time. After removing the ampoule(s), the remaining ampoules should be put back immediately into the canister in the liquid nitrogen container.
- 5. Thaw the content of the ampoule(s) rapidly by immersing the ampoule in clean water at 25 °C 27 °C. Gently swirl the ampoule(s) to disperse the contents. In order to protect the cells, it is important that the ampoule content is mixed, immediately after thawing, with the solvent. Dry the ampoule, then break the ampoule at its neck and immediately proceed as described below.
- 6. Gently withdraw the contents of the ampoule into a sterile syringe fitted with an 18-gauge needle.
- 7. Insert the needle through the stopper of the solvent bag, and then slowly and gently add the contents of the syringe to the solvent. Gently swirl and invert the bag to mix the vaccine. Withdraw a small quantity from the solvent bag into the syringe and rinse the ampoule. Inject the remaining contents of the ampoule gently into the solvent bag.
- 8. Repeat steps 6 and 7 for additional ampoules, if required.
- 9. Remove the syringe and invert the bag (6–8 times) to mix the vaccine.
- 10. The vaccine is now ready for use.

 After adding the contents of the ampoule to the solvent, the ready to use product is a clear, red coloured suspension for injection.

Administration:

The vaccine is administered by subcutaneous injection in the neck or by *in ovo* injection. The bag of vaccine should be gently swirled frequently during vaccination to guarantee that the vaccine suspension remains homogenous and that the correct vaccine virus titre is administered (e.g., during long vaccination sessions).

Control of correct storage:

To allow a check on correct storage and transport the ampoules are placed upside down in the liquid nitrogen containers. If frozen content is situated in the tip of the ampoule this indicates that the content has been thawed and must not be used.

3.10 Symptoms of overdose (and where applicable, emergency procedures and antidotes)

No symptoms were observed after the administration of a 10-fold dose of vaccine when applied subcutaneously. A 3-fold overdose was tested *in ovo*, which was regarded as safe. No information is available on the safety or possible adverse reactions following a 10-fold overdose applied *in ovo*.

3.11 Special restrictions for use and special conditions for use, including restrictions on the use of antimicrobial and antiparasitic veterinary medicinal products in order to limit the risk of development of resistance

Any person intending to manufacture, import, possess, distribute, sell, supply and use this veterinary medicinal product must first consult the relevant Member State's competent authority on the current vaccination policies, as these activities may be prohibited in a Member State on the whole or part of its territory pursuant to national legislation.

Official control authority batch release is required for this product.

3.12 Withdrawal periods

4. IMMUNOLOGICAL INFORMATION

4.1 ATCvet code: QI01AD16.

The vaccine is a cell-associated live recombinant turkey herpesvirus (HVT) expressing the F protein of Newcastle disease virus and the VP2 protein of infectious bursal disease virus. The vaccine induces active immunity against Newcastle disease, infectious bursal disease (Gumboro disease) and Marek's disease in chickens.

5. PHARMACEUTICAL PARTICULARS

5.1 Major incompatibilities

Do not mix with any other veterinary medicinal product, except Nobilis Rismavac and the solvent supplied for use with the veterinary medicinal product.

5.2 Shelf life

Shelf life of the concentrate as packaged for sale: 3 years.

Shelf life of the solvent (multilayer plastic bags) as packaged for sale: 3 years.

Shelf life after reconstitution according to directions: 2 hours.

5.3 Special precautions for storage

Concentrate:

Store and transport frozen in liquid nitrogen (below -140 °C).

Solvent:

Store below 30 °C.

Container:

Store liquid nitrogen container securely in an upright position in a clean, dry and well-ventilated room separated from the hatching/chicken room.

5.4 Nature and composition of immediate packaging

Concentrate:

- One Type I glass ampoule of 2 ml containing 2000 or 4000 doses. Ampoules are stored on a cane and attached to the cane is a coloured clip displaying the dose (2000 doses: salmon-pink coloured clip, and 4000 doses: yellow coloured clip).

Solvent:

- One 400 ml multilayer plastic bag.
- One 800 ml multilayer plastic bag.
- One 1200 ml multilayer plastic bag.
- One 1600 ml multilayer plastic bag.

Not all pack sizes may be marketed.

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

Medicines should not be disposed of via wastewater or household waste.

Use of take-back schemes for the disposal of any unused veterinary medicinal product or waste materials derived thereof in accordance with local requirements and with any national collection systems applicable to the veterinary medicinal product concerned.

6. NAME OF THE MARKETING AUTHORISATION HOLDER

Intervet International B.V.

7. MARKETING AUTHORISATION NUMBER(S)

EU/2/17/213/001-002.

8. DATE OF FIRST AUTHORISATION

Date of first authorisation: 22/08/2017.

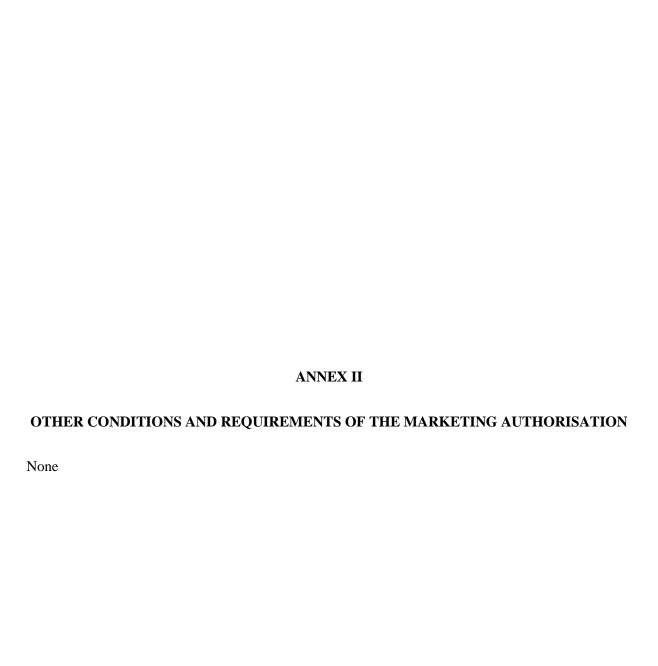
9. DATE OF THE LAST REVISION OF THE SUMMARY OF THE PRODUCT CHARACTERISTICS

 $\{MM/YYYY\}$

10. CLASSIFICATION OF VETERINARY MEDICINAL PRODUCTS

Veterinary medicinal product subject to prescription.

Detailed information on this veterinary medicinal product is available in the <u>Union Product Database</u> (<u>https://medicines.health.europa.eu/veterinary</u>).



ANNEX III LABELLING AND PACKAGE LEAFLET

A. LABELLING

AMPOULE 2000/4000 doses (2 ml glass)		
1. NAME OF THE VETERINARY MEDICINAL PRODUCT		
Innovax-ND-IBD		
2. QUANTITATIVE PARTICULARS OF THE ACTIVE SUBSTANCES		
HVP360		
3. BATCH NUMBER		
Lot {number}		
4. EXPIRY DATE		

Exp. {mm/yyyy}

MINIMUM PARTICULARS TO APPEAR ON SMALL IMMEDIATE PACKAGING UNITS

PARTICULARS TO APPEAR ON THE IMMEDIATE PACKAGE		
SOLVENT BAG 400/800/1200/1600 ml		
1. NAME OF THE VETERINARY MEDICINAL PRODUCT		
Solvent for cell associated poultry vaccines		
2. CONTENTS BY WEIGHT, BY VOLUME OR BY NUMBER OF DOSES		
400 ml 800 ml 1200 ml 1600 ml		
3. ROUTE(S) OF ADMINISTRATION		
Read package leaflet before use.		
4. STORAGE CONDITIONS		
Store below 30°C.		
5. BATCH NUMBER		
Lot {number}		
6. EXPIRY DATE		
EXP {MM/YYYY}		
7. THE WORDS "FOR ANIMAL TREATMENT ONLY"		
For animal treatment only.		

B. PACKAGE LEAFLET

PACKAGE LEAFLET

1. Name of the veterinary medicinal product

Innovax-ND-IBD concentrate and solvent for suspension for injection for chickens

2. Composition

Each dose of the reconstituted vaccine (0.2 ml for subcutaneous use or 0.05 ml for *in ovo* use) contains:

Cell-associated live recombinant turkey herpesvirus (strain HVP360), expressing the fusion protein of Newcastle disease virus and the VP2 protein of infectious bursal disease virus: $10^{3.3} - 10^{4.6} \text{ PFU}^1$.

¹ PFU – plaque forming units.

Concentrate and solvent for suspension for injection.

Concentrate: off-red to red cell concentrate.

Solvent: clear, red solution.

3. Target species

Chickens and embryonated chicken eggs.

4. Indications for use

For active immunisation of one-day-old chicks or 18–19 day-old embryonated chicken eggs:

- to reduce mortality and clinical signs caused by Newcastle disease (ND) virus,
- to prevent mortality and to reduce clinical signs and lesions caused by infectious bursal disease (IBD) virus,
- to reduce mortality, clinical signs and lesions caused by Marek's disease (MD) virus.

Onset of immunity: ND: 4 weeks of age

IBD: 3 weeks of age

MD: 9 days

Duration of immunity: ND: 60 weeks

IBD: 60 weeks

MD: entire risk period

5. Contraindications

None.

6. Special warnings

Special warnings:

Vaccinate healthy animals only.

Special precautions for safe use in the target species:

As this is a live vaccine, the vaccine strain is excreted from vaccinated birds and may spread to turkeys. Safety trials have shown that the strain is safe for turkeys. However, precautionary measures

have to be followed in order to avoid direct or indirect contact between vaccinated chickens and turkeys.

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

The handling of liquid nitrogen should take place in a well-ventilated area.

Innovax-ND-IBD is a virus suspension packed in glass ampoules and stored in liquid nitrogen. Before withdrawing ampoules from the liquid nitrogen canister, protective equipment consisting of gloves, long sleeves and a facemask or goggles should be worn. In order to prevent serious wounds, by either the liquid nitrogen or the ampoules when removing an ampoule from the canister, hold the palm of the (gloved) hand holding the ampoule away from the body and face. Care should be exercised to prevent contaminating the hands, eyes and clothing with the ampoule content. CAUTION: The ampoules have been known to explode on exposure to sudden temperature changes. Do not thaw in hot water or ice-cold water. Thaw the ampoules in clean water at 25 $^{\circ}$ C – 27 $^{\circ}$ C.

Special precautions for the protection of the environment:

Not applicable.

Laying birds:

The safety of the veterinary medicinal product has not been established during lay.

Interaction with other medicinal products and other forms of interaction:

Safety and efficacy data are available which demonstrate that Innovax-ND-IBD can be mixed in the same solvent and administered by the subcutaneous route with Nobilis Rismavac. For this mixed use, an onset of immunity of 5 days has been demonstrated for MD.

Safety and efficacy data are available which demonstrate that Nobilis ND Clone 30 or Nobilis ND C2 or Nobilis IB Ma5 or Nobilis IB 4-91 can be administered (not mixed) to day-old chicks that are vaccinated either by the subcutaneous or by the *in ovo* route with Innovax-ND-IBD. For such associated use, an onset of immunity of 3 weeks (when used with Nobilis ND Clone 30) and 2 weeks (when used with Nobilis ND C2), has been demonstrated for ND.

No information is available on the safety and efficacy of this vaccine when used with any other veterinary medicinal product except the products mentioned above. A decision to use this vaccine before or after any other veterinary medicinal product therefore needs to be made on a case by case basis.

Overdose:

No symptoms were observed after the administration of a 10-fold dose of vaccine when applied subcutaneously. A 3-fold overdose was tested *in ovo*, which was regarded as safe. No information is available on the safety or possible adverse reactions following a 10-fold overdose applied *in ovo*.

Special restrictions for use and special conditions for use:

Any person intending to manufacture, import, possess, distribute, sell, supply and use this veterinary medicinal product must first consult the relevant Member State's competent authority on the current vaccination policies, as these activities may be prohibited in a Member State on the whole part of its territory pursuant to national legislation.

Official control authority batch release is required for this product.

Major incompatibilities:

Do not mix with any other veterinary medicinal product except Nobilis Rismavac and the solvent supplied for use with the veterinary medicinal product.

7. Adverse events

None known.

Reporting adverse events is important. It allows continuous safety monitoring of a product. 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 contact, in the first instance, your veterinarian. You can also report any adverse events to the marketing authorisation holder using the contact details at the end of this leaflet, or via your national reporting system: {national system details}.

8. Dosage for each species, routes and method of administration

After dilution, administer 1 dose of 0.2 ml vaccine per chicken by subcutaneous injection in the neck or 1 dose of 0.05 ml per egg by *in ovo* injection.

9. Advice on correct administration

The bag of vaccine should be gently swirled frequently during vaccination to guarantee that the vaccine suspension remains homogenous and that the correct vaccine virus titre is administered (e.g., during long vaccination sessions).

Preparation of the vaccine:

The usual aseptic precautions should be applied to all preparation and administration procedures. The handling of liquid nitrogen should take place in a well-ventilated area.

1. Use solvent for cell associated poultry vaccines for reconstitution.

For subcutaneous use reconstitute the vaccine according to the table below:

Solvent bag	Number of vaccine ampoules for subcutaneous use
Bag of 400 ml solvent	1 ampoule containing 2000 doses
Bag of 800 ml solvent	2 ampoules containing 2000 doses
Bag of 800 ml solvent	1 ampoule containing 4000 doses
Bag of 1200 ml solvent	3 ampoules containing 2000 doses
Bag of 1600 ml solvent	4 ampoules containing 2000 doses
Bag of 1600 ml solvent	2 ampoules containing 4000 doses

When this product is mixed with Nobilis Rismavac, both should be diluted in the same solvent bag in the same way (400 ml of solvent for each 2000 doses of both products or 800 ml of solvent for each 4000 doses of both products).

For *in ovo* use reconstitute the vaccine according to the table below:

Solvent bag	Number of vaccine ampoules for in ovo use
Bag of 400 ml solvent	4 ampoules containing 2000 doses
Bag of 400 ml solvent	2 ampoules containing 4000 doses
Bag of 800 ml solvent	8 ampoules containing 2000 doses
Bag of 800 ml solvent	4 ampoules containing 4000 doses
Bag of 1200 ml solvent	12 ampoules containing 2000 doses
Bag of 1200 ml solvent	6 ampoules containing 4000 doses
Bag of 1600 ml solvent	16 ampoules containing 2000 doses
Bag of 1600 ml solvent	8 ampoules containing 4000 doses

The solvent must be clear, red coloured, without sediment and at room temperature (15 °C -25 °C) at the time of mixing.

2. Preparation of the vaccine shall be planned before the ampoules are taken from the liquid nitrogen and the exact amount of vaccine ampoules and amount of solvent needed shall be

- calculated first. There is no information available on the number of doses on the ampoules once they are removed from the cane, so special care has to be taken to ensure that the mix-ups of ampoules with different number of doses is avoided and the correct solvent is used.
- 3. Before withdrawing the ampoules from the liquid nitrogen container, protect hands with gloves, wear long sleeves and use a facemask or goggles. When removing an ampoule from the cane, hold in the palm of a gloved hand away from the body and the face.
- 4. When withdrawing a cane of ampoules from the canister in the liquid nitrogen container, expose only the ampoule(s) to be used immediately. It is recommended to handle a maximum of 5 ampoules (from one cane only) at a time. After removing the ampoule(s), the remaining ampoules should be put back immediately into the canister in the liquid nitrogen container.
- 5. Thaw the content of the ampoule(s) rapidly by immersing the ampoule in clean water at 25 °C 27 °C. Gently swirl the ampoule(s) to disperse the contents. In order to protect cells, it is important that the ampoule content is mixed, immediately after thawing, with the solvent. Dry the ampoule, then break the ampoule at its neck and immediately proceed as described below.
- 6. Gently withdraw the contents of the ampoule into a sterile syringe, fitted with an 18-gauge needle.
- 7. Insert the needle through the stopper of the solvent bag, and then slowly and gently add the contents of the syringe to the solvent. Gently swirl and invert the bag to mix the vaccine. Withdraw a small quantity from the solvent bag into the syringe and rinse the ampoule. Inject the remaining contents of the ampoule gently into the solvent bag.
- 8. Repeat steps 6 and 7 for additional ampoules, if required.
- 9. Remove the syringe and invert the bag (6–8 times) to mix the vaccine.
- 10. The vaccine is now ready for use.

 After adding the contents of the ampoule to the solvent, the ready to use product is a clear, red coloured suspension for injection.

Control of correct storage:

To allow a check on correct storage and transport the ampoules are placed upside down in the liquid nitrogen containers. If frozen content is situated in the tip of the ampoule, this indicates that the content has been thawed and must not be used.

10. Withdrawal periods

Zero days.

11. Special storage precautions

Keep out of the sight and reach of children.

Concentrate: Store and transport frozen in liquid nitrogen (below -140 °C).

Solvent: Store below 30 °C.

Container: Store liquid nitrogen container securely in an upright position in a clean, dry and well-ventilated room separated from the hatching/chicken room.

Do not use this veterinary medicinal product after the expiry date which is stated on the label. The expiry date refers to the last day of that month.

Shelf life after reconstitution according to directions: 2 hours.

12. Special precautions for disposal

Medicines should not be disposed of via wastewater or household waste.

Use take-back schemes for the disposal of any unused veterinary medicinal product or waste materials derived thereof in accordance with local requirements and with any applicable national collection systems. These measures should help to protect the environment.

Ask your veterinary surgeon or pharmacist how to dispose of medicines no longer required.

13. Classification of veterinary medicinal products

Veterinary medicinal product subject to prescription.

14. Marketing authorisation numbers and pack sizes

EU/2/17/213/001-002.

Pack sizes:

1 ampoule, containing 2000 or 4000 doses. Ampoules are stored on a cane and attached to the cane is a coloured clip displaying the dose (2000 doses: salmon-pink coloured clip, and 4000 doses: yellow coloured clip).

Bag of 400 ml solvent, bag of 800 ml solvent, bag of 1200 ml solvent or bag of 1600 ml solvent.

Not all pack sizes may be marketed.

15. Date on which the package leaflet was last revised

{MM/YYYY}

Detailed information on this veterinary medicinal product is available in the <u>Union Product Database</u> (https://medicines.health.europa.eu/veterinary).

16. Contact details

<u>Marketing authorisation holder and manufacturer responsible for batch release and contact details to</u> report suspected adverse reactions:

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

België/Belgique/Belgien

Tél/Tel: + 32 (0)2 370 94 01

Република България

Тел: + 359 28193749

Česká republika

Tel: +420 233 010 242

Danmark

Tlf: +45 44 82 42 00

Deutschland

Tel: +49 (0)8945614100

Lietuva

Tel: + 37052196111

Luxemburg/Luxemburg

Tél/Tel: + 32 (0)2 370 94 01

Magyarország

Tel.: + 36 1 439 4597

Malta

Tel: + 39 02 516861

Nederland

Tel: + 32 (0)2 370 94 01

Eesti

Tel: + 37052196111

Ελλάδα

 $T\eta\lambda$: + 30 210 989 7452

España

Tel: + 34 923 19 03 45

France

Tél: +33 (0)241228383

Hrvatska

Tel: +385 1 6611339

Ireland

Tel: + 353 (0) 1 2970220

Ísland

Sími: + 354 535 7000

Italia

Tel: + 39 02 516861

Κύπρος

Tηλ: +30 210 989 7452

Latvija

Tel: + 37052196111

Norge

Tlf: +47 55 54 37 35

Österreich

Tel: +43 (1) 256 87 87

Polska

Tel.: + 48 22 18 32 200

Portugal

Tel: + 351 214 465 700

România

Tel: +40 21 311 83 11

Slovenija

Tel: + 385 1 6611339

Slovenská republika

Tel: +420 233 010 242

Suomi/Finland

Puh/Tel: + 358 10 2310 750

Sverige

Tel: +46 (0)8 522 216 60

United Kingdom (Northern Ireland)

Tel: + 353 (0) 1 2970220

17. Other information

The vaccine is a cell-associated live recombinant turkey herpesvirus (HVT) expressing the F protein of Newcastle disease virus and the VP2 protein of Infectious bursal disease virus. The vaccine induces active immunity against Newcastle disease, infectious bursal disease (Gumboro disease) and Marek's disease in chickens.