1. NAME OF THE VETERINARY MEDICINAL PRODUCT

GALLIMUNE 302 ND+IB+EDS emulsion for injection

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Each 0.3 ml dose of vaccine contains:

Active substances:

corresponding to an antibody titre of 1. HI: haemagglutination inhibiting

Adjuvant:

Excipients:

Qualitative composition of excipients and other constituents	Quantitative composition if that information is essential for proper administration of the veterinary medicinal product
Thiomersal	30 μg
Formaldehyde	at most 43.2 µg
Ester of fatty acids and of polyols	
Ester of fatty acids and of ethoxylated polyols	
Water for injections	

Whitish homogeneous emulsion.

3. CLINICAL INFORMATION

3.1 Target species

Chickens (breeder and layer pullets).

3.2 Indications for use for each target species

Booster immunisation of breeder and layer pullets after vaccination with live vaccines against:

- Newcastle disease virus in order to reduce egg drop linked to Newcastle disease infection,
- Infectious bronchitis virus in order to reduce egg drop linked to infectious bronchitis infection caused by the Mass41 strain,

Active immunisation of breeder and layer pullets in order to reduce egg drop linked to infection with egg drop syndrome virus (EDS76) without priming.

Onset of immunity: 4 weeks after vaccination. Duration of immunity: one laying period.

¹ Minimum protective dose according to monograph 0870 of Ph. Eur.

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:

Not applicable.

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

To the user:

This veterinary medicinal product contains mineral oil. Accidental injection/self-injection may result in severe pain and swelling, particularly if injected into a joint or finger, and in rare cases could result in the loss of the affected finger if prompt medical attention is not given. If you are accidentally injected with this product, seek prompt medical advice even if only a very small amount is injected and take the package insert with you. If pain persists for more than 12 hours after medical examination, seek medical advice again.

To the physician:

This veterinary medicinal product contains mineral oil. Even if small amounts have been injected, accidental injection with this product can cause intense swelling, which may, for example, result in ischaemic necrosis and even the loss of a digit. Expert, PROMPT, surgical attention is required and may necessitate early incision and irrigation of the injected area, especially where there is involvement of finger pulp or tendon.

Special precautions for the protection of the environment:

Not applicable.

3.6 Adverse events

Chickens:

Very common	Abnormal histology ¹
(> 1 animal / 10 animals treated):	

¹ In clinical studies, lesions at the injection site linked to the oily adjuvant were observed histologically three weeks after injections in 87 % of cases, e.g. small quantities of oily residues and occasional aseptic micro-abscesses. No palpable reactions were observed.

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 its local representative or the national competent authority via the national reporting system. See the package leaflet for respective contact details.

3.7 Use during pregnancy, lactation or lay

Do not use in birds in lay and within 2 weeks before the start of the laying period.

3.8 Interaction with other medicinal products and other forms of interaction

No information is available on the safety and efficacy of this vaccine when used with any other veterinary medicinal product. 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

Administer one dose (0,3 ml) by intramuscular route from the age of 18 weeks and at least 4 weeks after the priming with live vaccines against Newcastle disease (strain Hitchner B1 or VG/GA-AVINEW) and infectious bronchitis (strain Mass H120).

Shake well before use.

Apply usual aseptic procedures.

Do not use syringes with natural rubber or butyl elastomer pistons.

Equipment including needles and syringes must be sterile before use.

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

In addition to the adverse events mentioned in paragraph "Adverse events", transitory apathy and slight oedema at injection site may occur after the administration of a double dose of vaccine.

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

Not applicable.

3.12 Withdrawal periods

Zero days.

4. IMMUNOLOGICAL INFORMATION

4.1 ATCvet code: QI01AA13

Inactivated vaccine in oily adjuvant against Newcastle disease, Infectious bronchitis and egg drop syndrome (EDS76).

The vaccine stimulates active immunity of breeder and layer pullets against egg drop syndrome (EDS76) (without priming), Newcastle disease and infectious bronchitis, subsequent to priming with live vaccines against these diseases.

5. PHARMACEUTICAL PARTICULARS

5.1 Major incompatibilities

Do not mix with any other veterinary medicinal product.

5.2 Shelf life

Shelf-life of the veterinary medicinal product as packaged for sale: 18 months. Shelf life after first opening the immediate packaging: use immediately.

5.3 Special precautions for storage

Store and transport refrigerated (2 $^{\circ}$ C – 8 $^{\circ}$ C). Protect from light. Do not freeze.

5.4 Nature and composition of immediate packaging

Nature of primary packaging elements:

- Polypropylene bottle
- Nitrile elastomer closure
- Aluminium cap

Sales presentations:

- 150-ml (500-dose) bottle.
- 150-ml (500-dose) bottle, box of 10 bottles.
- 300-ml (1 000-dose) bottle.
- 300-ml (1 000-dose) bottle, box of 10 bottles.

Not all pack sizes may be marketed.

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

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 national collection systems applicable to the veterinary medicinal product concerned.

6. NAME OF THE MARKETING AUTHORISATION HOLDER

Boehringer Ingelheim Vetmedica GmbH

7. MARKETING AUTHORISATION NUMBER(S)

VPA10454/051/001

8. DATE OF FIRST AUTHORISATION

25/11/2005

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

14/09/2023

10. CLASSIFICATION OF VETERINARY MEDICINAL PRODUCTS

Veterinary medicinal product subject to prescription.

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