

VETERINARY MEDICINES DIRECTORATE

On behalf of the Secretary of State for the Department for Environment, Food and Rural Affairs (Defra)

ISSUED BY:

VETERINARY MEDICINES DIRECTORATE
Woodham Lane, New Haw, Addlestone, Surrey, KT15 3LS
Tel: +44 (0)1932 336911; Website: www.gov.uk

Manufacturer's/Importer's Authorisation

SECTION 1A

1. Authorisation Number

ManA 1974

2. Name and address of Authorisation Holder

VetPlus Limited, Animal House, Boundary Road, Lytham, Lancashire, FY8 5LT, United Kingdom

3. Address(es) of Authorisation holder's manufacturing / importing site(s)

| VMD SITE NUMBER: | SITE NAME: | ADDRESS: |
|------------------|------------|---|
| S0040 | | Animal House, Boundary Road, Lytham St. Annes, Lancashire, FY8 5LT, United Kingdom |

4. Legally registered address of Authorisation Holder

As above

5. Scope of authorisation and dosage

See ANNEX 1

6. Legal basis of authorisation

See Section 1B of licence.



SECTION 1A (continued)

7. Name of responsible officer of the competent authority of the member state granting the manufacturing authorisation

CONFIDENTIAL

8. Date

14/08/2023



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Manufacturer's/Importer's Authorisation

SECTION 1B

1. This authorisation covers the manufacture, assembly and/or importation of veterinary medicinal products at the premises specified. The dosage forms authorised to be manufactured and / or products to be imported, as well as details of specific manufacturing operations permitted at each site are specified in Section 3. For veterinary medicinal products requiring a marketing authorisation, all manufacturing, assembly and/or importation operations shall be conducted so as to ensure that product strength, quality and purity meet the requirements of the marketing authorisation, or in the case of outsourced manufacture, assembly and / or importation, the specification made by the contract giver.

In relation to such products the authorisation holder shall either:

- a) provide and maintain such staff, premises and plant as are necessary to carry out any tests of the strength, quality or purity as required by the marketing authorisation / specification, or
- b) make arrangements with a person approved by the Secretary of State to carry out such tests on his behalf, and
- c) make arrangements for a Qualified Person (QP) to be available at all times for the purpose of checking that each batch of veterinary medicinal products has been manufactured and assembled in accordance with the appropriate provisions and to certify accordingly in a register.



SECTION 1B (continued)

- 2. The authorisation holder must inform the VMD, acting on behalf of the Secretary of State, in advance of any change to the details submitted or included in this authorisation. All changes must be approved by the Secretary of State prior to becoming effective. This includes any changes to named premises, persons, operations processes or structural alterations. If the business should change hands the new company or person must obtain a new authorisation prior to commencing operations. The manufacture and/or assembly and/or importation of any veterinary medicinal product pursuant to this authorisation shall not commence until a marketing authorisation has been granted naming the site named on this authorisation as being used for the manufacture of that product.
- 3. The names and addresses of holders of manufacturing authorisations for veterinary medicinal products together with addresses of authorised sites will be published on the EudraGMP database at www.eudragmp.ema.europa.eu. A register of holders of manufacturing "specials" authorisations can be found at the VMD website www.vmd.gov.uk.
- 4. Further information and specified guidelines may be obtained from the VMD website www.gov.uk
- 5. Authorisation Structure

This authorisation is divided into three sections.

- (a) <u>Section 1</u> (this section) identifies the authorisation holder and its details.
- (b) Section 2 lists variations to the authorisation.
- (c) Section 3 contains the details relating to each site named on the authorisation. Annex1 contain the details of the Authorisation holder's sites while Annex 2 lists the Qualified Person(s) and persons responsible for production and quality control. Annexes 3,4 and 5 list contracted sites (if applicable). Where the Authorisation holder has more than one manufacturing / importation site, separate versions of Annex 1 and Annex 2 along with versions of annexes 3, 4 and 5, as applicable, are included to fully capture the details for each.



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Manufacturer's/Importer's Authorisation

SECTION 3

ANNEX 1 - AUTHORISATION HOLDER'S SITE INFORMATION

SCOPE OF AUTHORISATION

| NAME AND ADDRESS OF SITE: | | |
|---------------------------|---|--|
| SITE NAME: | VetPlus Limited | |
| ADDRESS: | Animal House, Boundary Road, Lytham, Lancashire, FY8 5LT, United Kingdom | |
| VMD SITE NUMBER: | S0040 | |

| TYPE OF PRODUCTS HANDLED | |
|-------------------------------|--|
| Veterinary Medicinal Products | |

| AUTHORISED OPERATIONS | | |
|---|---------------------|--|
| Manufacturing Operations (according to Part 1) | Authorised | |
| Importation of Medicinal Products (according to Part 2) | Not Authorised Food | |



ANNEX 1 – AUTHORISATION HOLDER'S SITE INFORMATION (continued)

Part 1 – MANUFACTURING OPERATIONS

- authorised manufacturing operations include total and partial manufacturing (including various processes of dividing up, packaging or presentation), batch release and certification, importation, storage and distribution of specified dosage forms unless informed to the contrary
- quality control testing and/or release and batch certification activities without manufacturing operations should be specified under the relevant items
- if the company is engaged in manufacture of products with special requirements i.e. B-lactam antibiotics, other highly sensitising antibiotics, live cells, pathogenic organisms (biosafety Level 3 or 4), radiopharmaceuticals, ectoparasiticides or other highly potent or highly toxic products this should be stated under the relevant product type and dosage

| 1.1 | Sterile Products | Manufacture |
|-------|---|-------------------------|
| 1.1.1 | Aseptically Prepared (processing operations for the following dosage forms) | |
| | 1.1.1.1 Large volume liquids | Not Authorised |
| | 1.1.1.2 Lyophilisates | Not Authorised |
| | 1.1.1.3 Semi-solids | Not Authorised |
| | 1.1.1.4 Small volume liquids | Not Authorised |
| | 1.1.1.5 Solids and implants | Not Authorised |
| | 1.1.1.6 Other aseptically prepared products | Not Authorised NT, FOOD |



| 1.1.2 | Terminally Sterilised (processing operations for the following dosage forms) | |
|-------|--|----------------|
| | 1.1.2.1 Large volume liquids | Not Authorised |
| | 1.1.2.2 Semi-solids | Not Authorised |
| | 1.1.2.3 Small volume liquids | Not Authorised |
| | 1.1.2.4 Solids and implants | Not Authorised |
| | 1.1.2.5 Other terminally sterilised prepared products | Not Authorised |
| 1.1.3 | Batch certification | Not Authorised |





| 1.2 | Non-sterile products | Manufacture |
|-------|---|---------------------|
| 1.2.1 | Non-sterile products (processing operations for the following dosage forms) | |
| | 1.2.1.1 Capsules, hard shell | Authorised |
| | 1.2.1.2 Capsules, soft shell | Not Authorised |
| | 1.2.1.3 Chewing gums | Not Authorised |
| | 1.2.1.4 Impregnated matrices | Not Authorised |
| | 1.2.1.5 Liquids for external use | Authorised |
| | 1.2.1.6 Liquids for internal use | Not Authorised |
| | 1.2.1.7 Medicinal gases | Not Authorised |
| | 1.2.1.8 Other solid dosage forms | Not Authorised |
| | 1.2.1.9 Pressurised preparations | Not Authorised |
| | 1.2.1.10 Radionuclide generators | Not Authorised |
| | 1.2.1.11 Semi-solids | Authorised |
| | 1.2.1.12 Suppositories | Not Authorised |
| | 1.2.1.13 Tablets | Not Authorised |
| | 1.2.1.14 Transdermal patches | Not Authorised |
| | 1.2.1.15 Intraruminal devices | Not Authorised |
| | 1.2.1.16 Veterinary premixes | Not Authorised |
| | 1.2.1.17 Other non-sterile medicinal product | Not Authorised NARY |
| 1.2.2 | Batch certification | Authorised CTORATE |



| 1.3 | Biological medicinal products | Manufacture |
|-------|---|-----------------------|
| 1.3.1 | Biological medicinal products (list of product types) | |
| | 1.3.1.1 Blood products | Not Authorised |
| | 1.3.1.2 Immunological products | Not Authorised |
| | 1.3.1.3 Cell therapy products | Not Authorised |
| | 1.3.1.4 Gene therapy products | Not Authorised |
| | 1.3.1.5 Biotechnology products | Not Authorised |
| | 1.3.1.6 Human or animal extracted products | Not Authorised |
| | 1.3.1.7 Tissue engineered products | Not Authorised |
| | 1.3.1.8 Other biological medicinal products | Not Authorised |
| 1.3.2 | Batch certification (list of product types) | |
| | 1.3.2.1 Blood products | Not Authorised |
| | 1.3.2.2 Immunological products | Not Authorised |
| | 1.3.2.3 Cell therapy products | Not Authorised |
| | 1.3.2.4 Gene therapy products | Not Authorised |
| | 1.3.2.5 Biotechnology products | Not Authorised |
| | 1.3.2.6 Human or animal extracted products | Not Authorised |
| | 1.3.2.7 Tissue engineered products | Not Authorised |
| | 1.3.2.8 Other biological medicinal products | Not Authorised NARY |
| | | MEDICINES DIRECTORATE |



| 1.4 | Other products or manufacturing activity (any other relevant manufacturing activity/product type that is not covered above e.g. sterilisation of active substances, manufacture of biological active starting materials (when required by national legislation), herbal or homeopathic products, bulk or total manufacturing, etc). | Manufacture |
|-------|---|----------------|
| 1.4.1 | Manufacture of: | |
| | 1.4.1.1 Herbal products | Not Authorised |
| | 1.4.1.2 Homoeopathic products | Not Authorised |
| | 1.4.1.3 Other | Not Authorised |
| 1.4.2 | Sterilisation of active substances/excipients/finished product: | |
| | 1.4.2.1 Filtration | Not Authorised |
| | 1.4.2.2 Dry heat | Not Authorised |
| | 1.4.2.3 Moist heat | Not Authorised |
| | 1.4.2.4 Chemical | Not Authorised |
| | 1.4.2.5 Gamma irradiation | Not Authorised |
| | 1.4.2.6 Electron beam | Not Authorised |
| 1.4.3 | Others | Not Authorised |





| 1.5 | Packaging | Manufacture |
|-------|---|-----------------------------|
| 1.5.1 | Primary packing | |
| | 1.5.1.1 Capsules, hard shell | Authorised |
| | 1.5.1.2 Capsules, soft shell | Not Authorised |
| | 1.5.1.3 Chewing gums | Not Authorised |
| | 1.5.1.4 Impregnated matrices | Not Authorised |
| | 1.5.1.5 Liquids for external use | Authorised |
| | 1.5.1.6 Liquids for internal use | Not Authorised |
| | 1.5.1.7 Medicinal gases | Not Authorised |
| | 1.5.1.8 Other solid dosage forms | Not Authorised |
| | 1.5.1.9 Pressurised preparations | Not Authorised |
| | 1.5.1.10 Radionuclide generators | Not Authorised |
| | 1.5.1.11 Semi-solids | Authorised |
| | 1.5.1.12 Suppositories | Not Authorised |
| | 1.5.1.13 Tablets | Not Authorised |
| | 1.5.1.14 Transdermal patches | Not Authorised |
| | 1.5.1.15 Intraruminal devices | Not Authorised |
| | 1.5.1.16 Veterinary premixes | Not Authorised |
| | 1.5.1.17 Other non-sterile medicinal products | Not Authorised |
| 1.5.2 | Secondary packing | MEDICINES AuthorisedCTORATE |



| 1.6 | Quality control testing | Manufacture |
|-----|--------------------------------------|----------------|
| | 1.6.1 Microbiological: sterility | Not Authorised |
| | 1.6.2 Microbiological: non-sterility | Not Authorised |
| | 1.6.3 Chemical/Physical | Not Authorised |
| | 1.6.4 Biological | Not Authorised |

Any restrictions or clarifying remarks related to the scope of these Manufacturing operations:

None

