Medicines and Healthcare products Regulatory Agency

MANUFACTURER'S AUTHORISATION

1: Authorisation Number UK MIA(IMP) 40211

2: Name of authorisation holder SYMBIOSIS PHARMACEUTICAL SERVICES LIMITED

SYMBIOSIS PHARMACEUTICAL SERVICES LIMITED, SCION

HOUSE, STIRLING UNIVERSITY INNOVATION PARK, STIRLING, FK9 3: Address(es) of manufacturing site(s)

4NF, UNITED KINGDOM

SYMBIOSIS PHARMACEUTICAL SERVICES LIMITED, UNIT 10,

4: Legally registered address of authorisation holder SCION HOUSE, STIRLING UNIVERSITY INNOVATION PARK,

STIRLING, FK9 4NF, UNITED KINGDOM

5: Scope of authorisation and dosage forms ANNEX 1 and/ or ANNEX 2

Part 6 of The Medicines for Human Use (Clinical Trials) Regulations 6: Legal Basis of authorisation

2004 [SI 2004/1031]

7: Name of responsible officer of the competent authority

of the member state granting the manufacturing Confidential

authorisation

8: Authorisation Date 29/11/2024

9: Annexes attached Annex 1 and/or Annex 2

SCOPE OF AUTHORISATION

Annex 2

Name and address of the site:

SYMBIOSIS PHARMACEUTICAL SERVICES LIMITED, SCION HOUSE, STIRLING UNIVERSITY INNOVATION PARK, STIRLING, FK9 4NF, UNITED KINGDOM

Human Investigational Medicinal Products

Authorised Operations

MANUFACTURING OPERATIONS (according to part 1)

IMPORTATION OF MEDICINAL PRODUCTS (according to part 2)

Part 1 - MANUFACTURING OPERATIONS

[1.1] Sterile Investigational Medicinal Products

[1.1.1] Aseptically prepared (processing operations for the following dosage forms)

[1.1.1.2] Lyophilisates

Special Requirements

Cytotoxics

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[1.1.1.4] Small volume liquids

Special Requirements

Cytotoxics

[1.1.1.6] Other aseptically prepared products

Suspensions

Special Requirements

Cytotoxics

[1.2] Non-sterile investigational medicinal products

[1.2.1] Non-Sterile Products (processing operations for the following dosage forms)

[1.2.1.15] Other non-sterile medicinal products

Lyophilisation of non-sterile active ingredients / excipients for use as starting materials in solid oral dosage manufacturing elsewhere

Special Requirements

Cytotoxics

[1.3] Biological investigational medicinal products

[1.3.1] Biological medicinal products

[1.3.1.2] Immunological products

[1.3.1.4] Gene therapy products

[1.3.1.5] Biotechnology products

[1.3.1.8] Other biological medicinal products

Plasmid, DNA, RNA. peptides. proteins and antibodies products. This list is only indicative of the product handled.

Special Requirements

Cytotoxics

[1.4] Other investigational medicinal products or manufacturing activitiy

[1.4.2] Sterilisation of active substances/excipients/finished products:

[1.4.2.1] Filtration

[1.5] Packaging

[1.5.2] Secondary packaging

Part 2 - IMPORTATION OF MEDICINAL PRODUCTS

[2.2] Batch certification of imported medicinal products

[2.2.3] Biological medicinal products

[2.2.3.4] Gene therapy products

[2.3] Other Importation Activities

[2.3.1] Site of Physical Importation

[2.3.2] Importation of Intermediate which undergoes further processing

Issue Date: 29 Nov 2024