Medicines and Healthcare products Regulatory Agency

MANUFACTURER'S AUTHORISATION

1: Authorisation Number UK MIA 19819

2: Name of authorisation holder BIOTEC SERVICES INTERNATIONAL LIMITED

BIOTEC SERVICES INTERNATIONAL LIMITED, UNITS 2100, 2110, 2120, 2130, 2010, 2430 AND 2500, PHASE 18, CENTRAL PARK, BRIDGEND INDUSTRIAL ESTATE, BRIDGEND, CF31 3TY, UNITED KINGDOM

3: Address(es) of manufacturing site(s)

BIOTEC SERVICES INTERNATIONAL LIMITED, BIOTEC HOUSE,

CENTRAL PARK, WESTERN AVENUE, BRIDGEND INDUSTRIAL ESTATE,

BRIDGEND, CF31 3RT, UNITED KINGDOM

BIOTEC SERVICES INTERNATIONAL LIMITED, BIOTEC HOUSE,

4: Legally registered address of authorisation holder CENTRAL PARK, WESTERN AVENUE, BRIDGEND INDUSTRIAL ESTATE,

BRIDGEND, CF31 3RT, UNITED KINGDOM

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

6: Legal Basis of authorisation Regulation 17 of The Human Medicines Regulations 2012 (SI 2012/1916)

7: Name of responsible officer of the competent

authority of the member state granting the

manufacturing authorisation

Confidential

8: Authorisation Date 12/01/2021

9: Annexes attached Annex 1 and/or Annex 2

SCOPE OF AUTHORISATION

Annex 1

Name and address of the site:

BIOTEC SERVICES INTERNATIONAL LIMITED, UNITS 2100, 2110, 2120, 2130, 2010, 2430 AND 2500, PHASE 18, CENTRAL PARK, BRIDGEND INDUSTRIAL ESTATE, BRIDGEND, CF31 3TY, UNITED KINGDOM

Human 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 Products

[1.1.3] Batch certification

[1.2] Non-sterile products

Issue Date: 12 Jan 2021

SCOPE OF AUTHORISATION

[2.3.1] Site of Physical Importation

Annex 1

Name and address of the site:

Manufacturer's Authorisation: UK MIA 19819

Page 2 of 3

Issue Date: 12 Jan 2021

BIOTEC SERVICES INTERNATIONAL LIMITED, BIOTEC HOUSE, CENTRAL PARK, WESTERN AVENUE, BRIDGEND INDUSTRIAL ESTATE, BRIDGEND, CF31 3RT, UNITED KINGDOM

Human 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 Products

[1.1.3] Batch certification

[1.2] Non-sterile products

[1.2.2] Batch certification

[1.3] Biological medicinal products

[1.3.2] Batch certification

[1.3.2.1] Blood products

[1.3.2.2] Immunological products

[1.3.2.3] Cell therapy products

[1.3.2.4] Gene therapy products

[1.3.2.5] Biotechnology products

[1.3.2.6] Human or animal extracted products

[1.3.2.7] Tissue Engineered Products

Special Requirements

Tissue engineered product

[1.3.2.8] Other biological medicinal products

Tissue engineered products

[1.5] Packaging

[1.5.2] Secondary packaging

Part 2 - IMPORTATION OF MEDICINAL PRODUCTS

[2.2] Batch certification of imported medicinal products

[2.2.1] Sterile Products

[2.2.1.1] Aseptically prepared

[2.2.1.2] Terminally sterilised

[2.2.2] Non-sterile products

[2.2.3] Biological medicinal products

[2.2.3.1] Blood products

[2.2.3.2] Immunological products

[2.2.3.3] Cell therapy products

[2.2.3.4] Gene therapy products

[2.2.3.5] Biotechnology products

[2.2.3.6] Human or animal extracted products

[2.2.3.7] Tissue Engineered Products

[2.3] Other Importation Activities

[2.3.1] Site of Physical Importation

Issue Date: 12 Jan 2021