

# Medicines and Healthcare products Regulatory Agency

## MANUFACTURER'S AUTHORISATION

<b>1: Authorisation Number</b>	UK MIA(IMP) 56952
<b>2: Name of authorisation holder</b>	PNR PHARMA CONSULTING LIMITED
<b>3: Address(es) of manufacturing site(s)</b>	PNR PHARMA CONSULTING LIMITED, UPPER GROUND FLOOR ROOM 2, CREUDDYN, PONTFAEN ROAD, LAMPETER, SA48 7BN, UNITED KINGDOM PNR PHARMA CONSULTING LIMITED, UNITS 5 AND 6, LIME AVENUE, EBBW VALE, NP23 6GL, UNITED KINGDOM
<b>4: Legally registered address of authorisation holder</b>	PNR PHARMA CONSULTING LIMITED, UNITS 5 AND 6, LIME AVENUE, EBBW VALE, NP23 6GL, UNITED KINGDOM
<b>5: Scope of authorisation and dosage forms</b>	ANNEX 1 and/ or ANNEX 2
<b>6: Legal Basis of authorisation</b>	Part 6 of The Medicines for Human Use (Clinical Trials) Regulations 2004 [SI 2004/1031]
<b>7: Name of responsible officer of the competent authority of the member state granting the manufacturing authorisation</b>	Confidential
<b>8: Authorisation Date</b>	10/06/2026
<b>9: Annexes attached</b>	Annex 1 and/or Annex 2

### SCOPE OF AUTHORISATION

#### Annex 2

Name and address of the site:

**PNR PHARMA CONSULTING LIMITED**, UPPER GROUND FLOOR ROOM 2, CREUDDYN, PONTFAEN ROAD, LAMPETER, SA48 7BN, UNITED KINGDOM

Human Investigational Medicinal Products
Authorised Operations
IMPORTATION OF MEDICINAL PRODUCTS (according to part 2)
Part 2 - IMPORTATION OF MEDICINAL PRODUCTS <b>[ 2.3 ] Other Importation Activities</b> [ 2.3.4 ] Other Importation of QP-certified IMPs from a country on the approved country for import list

## SCOPE OF AUTHORISATION

### Annex 2

Name and address of the site:

**PNR PHARMA CONSULTING LIMITED**, UNITS 5 AND 6, LIME AVENUE, EBBW VALE, NP23 6GL, UNITED KINGDOM

Human Investigational Medicinal Products
Authorised Operations
MANUFACTURING OPERATIONS (according to part 1) IMPORTATION OF MEDICINAL PRODUCTS (according to part 2)
<b>Part 1 - MANUFACTURING OPERATIONS</b> <b>[ 1.1 ] Sterile Investigational Medicinal Products</b> [ 1.1.3 ] Batch certification <b>[ 1.2 ] Non-sterile investigational medicinal products</b> [ 1.2.2 ] Batch certification <b>[ 1.3 ] Biological investigational medicinal products</b> [ 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 <b>[ 1.5 ] Packaging</b> [ 1.5.2 ] Secondary packaging <b>Part 2 - IMPORTATION OF MEDICINAL PRODUCTS</b> <b>[ 2.2 ] Batch certification of imported medicinal products</b> [ 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 <b>[ 2.3 ] Other Importation Activities</b> [ 2.3.1 ] Site of Physical Importation [ 2.3.2 ] Importation of Intermediate which undergoes further processing

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