Medicines and Healthcare products Regulatory Agency

WHOLESALE DISTRIBUTION AUTHORISATION

(MEDICINAL PRODUCTS FOR HUMAN USE)

1. Authorisation Number	UK WDA(H) 101
2. Name of Authorisation Holder	NOVARTIS PHARMACEUTICALS UK LIMITED
3. Legally registered address of Authorisation Holder	NOVARTIS PHARMACEUTICALS UK LIMITED, 2ND FLOOR, THE WEST WORKS BUILDING, WHITE CITY PLACE, 195 WOOD LANE, LONDON, W12 7FQ, UNITED KINGDOM
4. Address(es) of Site(s)	NOVARTIS PHARMACEUTICALS UK LIMITED, 2ND FLOOR, THE WEST WORKS BUILDING, WHITE CITY PLACE, 195 WOOD LANE, LONDON, W12 7FQ, UNITED KINGDOM
5. Scope of authorisation (complete for each site under 4)	ANNEX 1
6. Legal b <mark>asis of au</mark> thorisation	Regulation 18 of the Human Medicines Regulations 2012
7. Name of responsible officer of the competent authority of the member state granting the wholesaling authorisation	Confidential
8. Date	06/11/2024
9. Annexes attached	Annex 1 Scope of wholesale distribution authorisation Annex 2 (Optional) Address(es) of contract wholesale distribution sites and their authorisation number Annex 3 (Optional) Name(s) of responsible person(s) Annex 4 (Optional) Date of Inspection on which authorisation was

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ANNEX 1

SCOPE OF WHOLESALE DISTRIBUTION AUTHORISATION

Name and address of the site:

NOVARTIS PHARMACEUTICALS UK LIMITED, 2ND FLOOR, THE WEST WORKS BUILDING, WHITE CITY PLACE, 195 WOOD LANE, LONDON, W12 7FQ, UNITED KINGDOM

1. MEDICINAL PRODUCTS

1.1 With "an authorisation" (a UK, Great Britain or Northern Ireland Marketing Authorisation, an Article 126a authorisation, a certificate of registration or traditional herbal registration)

1.2 Without "an authorisation" (a UK, Great Britain or Northern Ireland Marketing Authorisation, an Article 126a authorisation, a certificate of registration or traditional herbal registration) in GB or EEA and intended for the UK market

2. AUTHORISED WHOLESALE DISTRIBUTION OPERATIONS

2.1 Procurement

2.3 Supply

2.4 Export

2.6 Products imported from countries on a list

2.6a Products certified under Article 51 of Directive 2001/83/EC

2.6b Products not certified under Article 51 of Directive 2001/83/EC

3. MEDICINAL PRODUCTS WITH ADDITIONAL REQUIREMENTS

3.1.2 Medicinal products derived from blood

3.1.3 Immunological medicinal products

3.1.4 Radiopharmaceutical (including radionuclide kits)

3.3 Cold chain products (requiring low temperature handling)

3.4 Other Products

Products in the category Advanced Therapy Medicinal Product (ATMP)

Any restrictions or clarifying remarks (for all users)

4 Categories of Products Handled at this Site: 4.1 Prescription Only Medicines, 4.2 General Sales List, 4.4 Pharmacy