Medicines and Healthcare products Regulatory Agency

MANUFACTURER'S AUTHORISATION

1: Authorisation Number

2: Name of authorisation holder

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

4: Legally registered address of authorisation holder

5: Scope of authorisation and dosage forms

6: Legal Basis of authorisation

7: Name of responsible officer of the competent authority of the member

state granting the manufacturing authorisation

8: Authorisation Date

9: Annexes attached

UK MIA 44646

SMIRO QUALITAS LIMITED

SMIRO QUALITAS LIMITED, 4 FLATFORD ROAD,

HAVERHILL, CB9 7WW, UNITED KINGDOM

SMIRO QUALITAS LIMITED, 4 FLATFORD ROAD,

HAVERHILL, CB9 7WW, UNITED KINGDOM

ANNEX 1 and/ or ANNEX 2

Regulation 17 of The Human Medicines Regulations

2012 (SI 2012/1916)

Confidential

24/07/2025

Annex 1 and/or Annex 2

SCOPE OF AUTHORISATION

Annex 1

Name and address of the site:

SMIRO QUALITAS LIMITED, 4 FLATFORD ROAD, HAVERHILL, CB9 7WW, UNITED KINGDOM

Human Medicinal Products

Authorised Operations

IMPORTATION OF MEDICINAL PRODUCTS (according to part 2)

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

[2.2.3.2] Immunological products

[2.3] Other Importation Activities

Issue Date: 24 Jul 2025

[2.3.4] Products authorised under regulation 174 (supply in response to spread of pathogenic agents etc)

GMP Certification and Batch release of imported biological imported Product- Monoclonal antibodies, Vaccines, oral vaccines, recombinant proteins, MVAs and adjuvants

