

# Medicines and Healthcare products Regulatory Agency

## MANUFACTURER'S AUTHORISATION

**1: Authorisation Number** UK MIA 18532

**2: Name of authorisation holder** SEQIRUS VACCINES LIMITED

**3: Address(es) of manufacturing site(s)** SEQIRUS VACCINES LIMITED , GASKILL ROAD,  
SPEKE, LIVERPOOL, L24 9GR, UNITED KINGDOM

**4: Legally registered address of authorisation holder** SEQIRUS VACCINES LIMITED, GASKILL ROAD,  
SPEKE, LIVERPOOL, L24 9GR, 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** 13/10/2025

**9: Annexes attached** Annex 1 and/or Annex 2

### SCOPE OF AUTHORISATION

#### Annex 1

Name and address of the site:

**SEQIRUS VACCINES LIMITED , GASKILL ROAD, SPEKE, LIVERPOOL, L24 9GR, UNITED KINGDOM**

|                                                                                                                                                                                                                                                                                                                                                                                                                                     |
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| Human Medicinal Products                                                                                                                                                                                                                                                                                                                                                                                                            |
| Authorised Operations                                                                                                                                                                                                                                                                                                                                                                                                               |
| MANUFACTURING OPERATIONS (according to part 1)<br>IMPORTATION OF MEDICINAL PRODUCTS (according to part 2)                                                                                                                                                                                                                                                                                                                           |
| <b>Part 1 - MANUFACTURING OPERATIONS</b><br><b>[ 1.1 ] Sterile Products</b><br>[ 1.1.1 ] Aseptically prepared (processing operations for the following dosage forms)<br>[ 1.1.1.6 ] Other aseptically prepared products<br>Vaccines<br><b>[ 1.3 ] Biological medicinal products</b><br>[ 1.3.1 ] Biological medicinal products<br>[ 1.3.1.2 ] Immunological products<br>[ 1.3.1.8 ] Other biological medicinal products<br>Vaccines |

- [ 1.3.2 ] Batch certification
  - [ 1.3.2.2 ] Immunological products
  - [ 1.3.2.8 ] Other biological medicinal products
    - Vaccines

**[ 1.4 ] Other products or manufacturing activity**

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

**[ 1.5 ] Packaging**

- [ 1.5.1 ] Primary packaging
  - [ 1.5.1.17 ] Other non-sterile medicinal products
    - Vaccines
- [ 1.5.2 ] Secondary packaging

**[ 1.6 ] Quality control testing**

- [ 1.6.1 ] Microbiological: sterility
- [ 1.6.2 ] Microbiological: non-sterility
- [ 1.6.3 ] Chemical/Physical
- [ 1.6.4 ] Biological

**Part 2 - IMPORTATION OF MEDICINAL PRODUCTS**

**[ 2.1 ] Quality control testing of imported medicinal products**

- [ 2.1.1 ] Microbiological: sterility
- [ 2.1.2 ] Microbiological: non-sterility
- [ 2.1.3 ] Chemical/Physical
- [ 2.1.4 ] Biological

**[ 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
- [ 2.2.3 ] Biological medicinal products
  - [ 2.2.3.2 ] Immunological products
  - [ 2.2.3.8 ] Other biological medicinal products
    - Vaccines

**[ 2.3 ] Other Importation Activities**

- [ 2.3.1 ] Site of Physical Importation
- [ 2.3.2 ] Importation of Intermediate which undergoes further processing
- [ 2.3.3 ] Biological Active Substance
- [ 2.3.4 ] Products authorised under regulation 174 (supply in response to spread of pathogenic agents etc)
  - Importation, testing and release of Neuraminidase inhibitor