## **MANUFACTURER'S AUTHORISATION** 1 - 2

1. Authorisation Number 2023\_159\_1\_2

2. Name of authorisation holder Solvias France (ORG-100012001 / LOC-100018755)

3. Address(es) of manufacturing site(s) Solvias France (ORG-100012001 / LOC-100018755), Zone Industrielle, Rue Du Canal D Alsace, Hombourg, 68490, France

3.a Additional details on units inspected of manufacturing site(s) address(es)

4. Legally registered address of authorisation Zone Industrielle, Rue Du Canal D Alsace, Hombourg, 68490, France

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

6. Legal Basis of authorisation Art. 40 of Directive 2001/83/EC
Art. 61 of Regulation (EU) No 536/2014

7. Name of responsible officer of the competent confidential authority of the member state granting the manufacturing authorisation

8. Signature

holder

9. Date 2023-06-15

10. Annexes attached Annex 1 and/or Annex 2

Optional Annexes as required:

 $Annex\ 3 (Addresses\ of\ Contract\ Manufacturing\ Site(s))$ 

Annex 4(Addresses of Contract laboratories)

Annex 5(Name of Qualified Person)
Annex 6(Name of responsible persons)

Annex 7(Date of inspection on which authorisation granted, scope of last

inspection)

Annex 8(Manufactured/imported products authorised)<sup>3</sup>

<sup>&</sup>lt;sup>1</sup>The authorisation referred to in paragraph 40(1) of Directive 2001/83/EC and 44(1) of Directive 2001/82/EC, as amended, shall also be required for imports coming from third countries into a Member State.

 $<sup>^2</sup>$ Guidance on the interpretation of this template can be found in the Help menu of EudraGMDP database.

<sup>&</sup>lt;sup>3</sup>The Competent Authority is responsible for appropriate linking of the authorisation with the manufacturer's application (Art. 42(3) of Directive 2001/83/EC and Art. 46(3) of Directive 2001/82/EC as amended).

#### **SCOPE OF AUTHORISATION**

ANNEX 1

Name and address of the site: Solvias France, Zone Industrielle, Rue Du Canal D Alsace,

Hombourg, 68490, France

Additional Details:

**Human Medicinal Products** 

### **Authorised Operations**

MANUFACTURING OPERATIONS(according to part 1)

IMPORTATION OF MEDICINAL PRODUCTS (according to part 2)

Part 1	- MANUFACTURING OPERATIONS
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

Any restrictions or clarifying remarks related to the scope of these Manufacturing operations (for Public users)

Manufacturer (Article R.5124-2 1° of the French Public Health Code)

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		

# Any restrictions or clarifying remarks related to the scope of these Importation operations (for Public users)

Importer (Article R.5124-2 2° of the French Public Health Code) --- Signatory: Mrs Mélanie Cachet, deputy director, Inspection division --- The ANSM does not issue hard copy of this authorisation.

### **SCOPE OF AUTHORISATION**

ANNEX 2

Name and address of the site: Solvias France, Zone Industrielle, Rue Du Canal D Alsace,

Hombourg, 68490, France

**Human Investigational Medicinal Products** 

### **Authorised Operations**

MANUFACTURING OPERATIONS (according to part 1)
IMPORTATION OF MEDICINAL PRODUCTS (according to part 2)

Part 1 - MANUFACTURING OPERATIONS			
1.6	<b>Quality control testing</b>		
	1.6.1 Microbiological: sterility		
	1.6.2 Microbiological: non-sterility		
	1.6.3 Chemical/Physical		
	1.6.4 Biological		

Any restrictions or clarifying remarks related to the scope of these Manufacturing operations (for Public users)

Manufacturer (Article R.5124-2 1° of the French Public Health Code)

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		

Any restrictions or clarifying remarks related to the scope of these Importation operations (for Public users)

Importer (Article R.5124-2 2° of the French Public Health Code) --- Signatory: Mrs Mélanie Cachet, deputy director, Inspection division --- The ANSM does not issue hard copy of this authorisation.