Medicines and Healthcare products Regulatory Agency

MANUFACTURER'S AUTHORISATION

1:	Authorisation	Number

2: Name of authorisation holder

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

UK MIA 4351

PENN PHARMACEUTICAL SERVICES LIMITED

PENN PHARMACEUTICAL SERVICES LIMITED, UNITS 23-24, TAFARNAUBACH INDUSTRIAL ESTATE, TAFARNAUBACH, TREDEGAR, NP22 3AA, UNITED KINGDOM

- 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

PENN PHARMACEUTICAL SERVICES LIMITED, UNITS 23-24, TAFARNAUBACH INDUSTRIAL ESTATE, TAFARNAUBACH, TREDEGAR, NP22 3AA, UNITED KINGDOM

ANNEX 1 and/ or ANNEX 2

Regulation 17 of The Human Medicines Regulations 2012 (SI 2012/1916)

Confidential

12/03/2025 Annex 1 and/or Annex 2

SCOPE OF AUTHORISATION

Annex 1 Name and address of the site:

PENN PHARMACEUTICAL SERVICES LIMITED, UNITS 23-24, TAFARNAUBACH INDUSTRIAL ESTATE, TAFARNAUBACH,

TREDEGAR, NP22 3AA, UNITED KINGDOM

Human Medicinal Products

Authorised Operations

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

Part 1 - MANUFACTURING OPERATIONS

[1.1] Sterile Products

[1.1.3] Batch certification

[1.2] Non-sterile products

[1.2.1] Non-Sterile Products (processing operations for the following dosage forms)

- [1.2.1.1] Capsules, hard shell **Special Requirements** Inclusion of Potent, Toxic Compounds
- [1.2.1.5] Liquids for external use **Special Requirements** Inclusion of Potent, Toxic Compounds
- [1.2.1.6] Liquids for internal use **Special Requirements** Inclusion of Potent, Toxic Compounds
- [1.2.1.11] Semi-solids **Special Requirements** Inclusion of Potent, Toxic Compounds
- [1.2.1.12] Suppositories **Special Requirements** Inclusion of Potent, Toxic Compounds
- [1.2.1.13] Tablets **Special Requirements** Inclusion of Potent, Toxic Compounds
- [1.2.1.17] Other non-sterile medicinal products Filling of Powders
- [1.2.2] Batch certification
- [1.3] Biological medicinal products
 - [1.3.2] Batch certification
 - [1.3.2.2] Immunological products
 - [1.3.2.5] Biotechnology products
 - [1.3.2.6] Human or animal extracted products

[1.5] Packaging

- [1.5.1] Primary packaging
 - [1.5.1.1] Capsules, hard shell **Special Requirements** Inclusion of Potent, Toxic Compounds
 - [1.5.1.5] Liquids for external use **Special Requirements** Inclusion of Potent, Toxic Compounds
 - [1.5.1.6] Liquids for internal use **Special Requirements** Inclusion of Potent, Toxic Compounds
 - [1.5.1.11] Semi-solids **Special Requirements** Inclusion of Potent, Toxic Compounds
 - [1.5.1.12] Suppositories Special Requirements Inclusion of Potent, Toxic Compounds
 - [1.5.1.13] Tablets **Special Requirements** Inclusion of Potent, Toxic Compounds
 - [1.5.1.17] Other non-sterile medicinal products Filling of powders

[1.5.2] Secondary packaging

[1.6] Quality control testing

[1.6.2] Microbiological: non-sterility

[1.6.3] Chemical/Physical

Part 2 - IMPORTATION OF MEDICINAL PRODUCTS

[2.1] Quality control testing of imported medicinal products

[2.1.2] Microbiological: non-sterility

[2.1.3] Chemical/Physical

[2.2] Batch certification of imported medicinal products

[2.2.2] Non-sterile products

- [2.3] Other Importation Activities
 - [2.3.1] Site of Physical Importation
 - [2.3.2] Importation of Intermediate which undergoes further processing

