# Medicines and Healthcare products Regulatory Agency

## WHOLESALE DISTRIBUTION AUTHORISATION

# (MEDICINAL PRODUCTS FOR HUMAN USE)

1. Authorisation Number UK WDA(H) 44056

2. Name of Authorisation Holder B.N.O.S. MEDITECH LIMITED

3. Legally registered address of

**Authorisation Holder** 

B.N.O.S. MEDITECH LIMITED, UNIT 9, FIFTH AVENUE, BLUEBRIDGE INDUSTRIAL

ESTATE, HALSTEAD, CO9 2SZ, UNITED KINGDOM

B.N.O.S. MEDITECH LIMITED, UNIT 13, SIXTH AVENUE, BLUEBRIDGE INDUSTRIAL

ESTATE, HALSTEAD, CO9 2FL, UNITED KINGDOM

4. Address(es) of Site(s)

B.N.O.S. MEDITECH LIMITED, UNIT 9, FIFTH AVENUE, BLUEBRIDGE INDUSTRIAL

ESTATE, HALSTEAD, CO9 2SZ, UNITED KINGDOM

B.N.O.S. MEDITECH LIMITED, UNIT 23, FIFTH AVENUE, BLUEBRIDGE INDUSTRIAL

ESTATE, HALSTEAD, CO9 2SZ, UNITED KINGDOM

5. Scope of authorisation (complete for

each site under 4)

ANNEX 1

6. Legal basis of authorisation 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

9. Annexes attached

Confidential

8. Date 18/08/2025

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 granted Annex 5 Additional provisions

#### **ANNEX 1**

#### SCOPE OF WHOLESALE DISTRIBUTION AUTHORISATION

Name and address of the site:

B.N.O.S. MEDITECH LIMITED, UNIT 13, SIXTH AVENUE, BLUEBRIDGE INDUSTRIAL ESTATE, HALSTEAD, CO9 2FL, 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)

### 2. AUTHORISED WHOLESALE DISTRIBUTION OPERATIONS

Issue Date: 18 Aug 2025

	2.1 Procurement
	2.2 Holding
	2.3 Supply
	2.4 Export
	3. MEDICINAL PRODUCTS WITH ADDITIONAL REQUIREMENTS
	3.2 Medicinal gases
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#### Any restrictions or clarifying remarks (for all users)

4 Categories of Products Handled at this Site: 4.2 General Sales List, 4.4 Pharmacy Name and address of the site:

B.N.O.S. MEDITECH LIMITED, UNIT 9, FIFTH AVENUE, BLUEBRIDGE INDUSTRIAL ESTATE, HALSTEAD, CO9 2SZ, UNITED KINGDOM

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#### 3. MEDICINAL PRODUCTS WITH ADDITIONAL REQUIREMENTS

3.2 Medicinal gases

#### Any restrictions or clarifying remarks (for all users)

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B.N.O.S. MEDITECH LIMITED, UNIT 23, FIFTH AVENUE, BLUEBRIDGE INDUSTRIAL ESTATE, HALSTEAD, CO9 2SZ, UNITED KINGDOM

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- 2.3 Supply
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- 3. MEDICINAL PRODUCTS WITH ADDITIONAL REQUIREMENTS
- 3.2 Medicinal gases

Any restrictions or clarifying remarks (for all users)

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



# Medicines and Healthcare products Regulatory Agency

Certificate No: UK WDA(H) 44056 Insp GDP 44056/11410101-0005

# CERTIFICATE OF GDP COMPLIANCE OF A WHOLESALE DISTRIBUTOR

Issued following an inspection in accordance with Regulation 331 (4) (b) of the Human Medicines Regulations 2012

The wholesale distributor: B.N.O.S. MEDITECH LIMITED

Site address: B.N.O.S. MEDITECH LIMITED, UNIT 9, FIFTH AVENUE, BLUEBRIDGE INDUSTRIAL ESTATE, HALSTEAD, CO9 2SZ,

**UNITED KINGDOM** 

Has been inspected under the national inspection programme in connection with authorisation number UK WDA(H) 44056 Insp GDF 44056/11410101-0005 in accordance with regulation 18 of the Human Medicines Regulations 2012

From the knowledge gained during inspection of this wholesale distributor, the latest of which was conducted on 05/08/2024, it is considered that it complies with the principles of good distribution practice requirements referred to in regulation C17 of the Human Medicines Regulations 2012

This certificate reflects the status of the premises at the time of the inspection noted above and should not be relied upon to reflect the compliance status if more than five years have elapsed since the date of that inspection. However this period of validity may be reduced using regulatory risk management principles, by an entry in the Restrictions or Clarifying Remarks field.

This certificate is valid only when presented with all pages.

The authenticity of this certificate may be verified in MHRA-GMDP. If it does not appear please contact the issuing authority.

16/01/2025	Name and signature of the authorised person of the Competent Authority of United Kingdom
	Confidential
	Medicines and Healthcare products Regulatory Agency
	Confidential