

National Authority Of Medicines And Health Products

UNION FORMAT FOR A WHOLESALE DISTRIBUTION AUTHORISATION

Human medicinal products

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| 1. Authorisation Number | : A012/17/H/007/2025 |
| 2. Name of Authorisation Holder | : Originpharma S.A.
(ORG-100036909 / LOC-100058308) |
| 3. Address(es) of Site(s) | : Centro Empresarial De Alverca, Rua Casal Das Ervas
Fracao 4, Alverca Do Ribatejo, 2615-187, Portugal
ACTIVE |
| 4. Legally registered address of Authorisation Holder | : Centro Empresarial De Alverca, Rua Casal Das Ervas
Fracao 4, Alverca Do Ribatejo, 2615-187 |
| 5. Scope of authorisation (complete for each site under 4) | : ANNEX 1 |
| 6. Legal basis of authorisation | : Art. 77(1) of Directive 2001/83/EC
National legal basis of authorisation: Art. 94.º of
Decree-law 176/2006, of 30th August. |
| 7. Name of responsible officer of the competent authority of the member state granting the wholesaling authorisation | : Maria Fernanda Ralha Henriques Matos, +351 21
7987278 |
| 8. Signature | : |
| 9. Date | : 2025-08-21 |
| 10. Annexes attached | : 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 (Optional) Additional provisions based on
national requirements |

ANNEX 1

SCOPE OF WHOLESALE DISTRIBUTION AUTHORISATION

Name and address of the site: Originpharma S.A.

(ORG-100036909 / LOC-100058308)

, Centro Empresarial De Alverca, Rua Casal Das Ervas Fracao 4, Alverca Do Ribatejo, 2615-187, Portugal **ACTIVE**

Human medicinal products

1. MEDICINAL PRODUCTS

- 1.1. with a Marketing Authorisation or registration in EEA country(s)
- 1.3. without a Marketing Authorisation or registration in the EEA and intended for exportation

2. AUTHORISED WHOLESALE DISTRIBUTION OPERATIONS

- 2.1. Procurement
- 2.2. Holding
- 2.3. Supply
- 2.4. Export

3. MEDICINAL PRODUCTS WITH ADDITIONAL REQUIREMENTS

- 3.1. Narcotic or psychotropic products**
- 3.2. Products requiring low temperature handling
 - 3.2.1. Temperatures between 2 to 8 °C
- 3.3. Other products: (please specify here or make a reference to Annex 5) Medicinal products derived from blood and Immunological.

Any restrictions or clarifying remarks (for all users): WDA scope activity - Wholesale Distributor and 3 PL (Third Party Logistics Providers).

Updating WDA, regarding a renewal of the latest Inspection Date of the Certificate of GDP Compliance.

**Without prejudice to further authorisations as may be required according to national legislation.

Anexo 3 / Annex 3

Nome(s) do(s) Director(es) Técnico(s) / Name(s) of Responsible Person(s)

Dr (a). Adriana Sofia Cabral Reis da Silva