

**District Government of Cologne**

**AUTHORISATION FOR THE WHOLESALE DISTRIBUTION OF MEDICINAL PRODUCTS**

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| 1. Authorisation number/file number  | DE_NW_04_WDA_2020-0002-02  |
| 2. Name of the authorisation holder  | avenar pharma GmbH   |
| 3. Registered address of the authorisation holder  | Moltkestraße 25<br>42799 Leichlingen   |
| 4. Address of the authorisation holder's business premises   | Moltkestraße 25<br>Warehouse I E1 and Warehouse II E6<br>42799 Leichlingen   |
| 5. Scope of authorisation<br>(Please provide information for each of the business premises listed in number 4.)    | Appendix 1   |
| 6. Legal basis for the issue of the authorisation  | § 52 a (1) of the Medicinal Products Act ( <i>Arzneimittelgesetz – AMG</i> ) in the version in force from time to time |
| 7. Name of the responsible case handler of the competent authorities of the Member State issuing the authorisation | Ina Niemann, Government<br>Pharmaceuticals Director  |
| 8. Signature   | (Signature of Ina Niemann) (Stamp of the District Government of Cologne)   |
| 9. Date  | 17 August 2020   |

10. Appendices attached:

Appendix 1

Scope of the authorisation

## SCOPE OF THE AUTHORISATION

Name and address of the business premises:  
 avenar pharma GmbH  
 Warehouse I E1 and Warehouse II E6  
 Moltkestraße 25  
 42799 Leichlingen

### 1. MEDICINAL PRODUCTS

Medicinal products for human use     Medicinal products for veterinary use

- 1.1  with authorisation for placing on the market in a country of the EEA
- 1.2  which, without authorisation for placing on the market in a country of the European Economic Area (EEA), are placed on the market in the EEA (exemption from obtaining authorisation)\*
- 1.3  which, without authorisation for placing on the market in a country of the European Economic Area, are NOT placed on the market in the EEA (medicinal products for third countries)

### 2. PERMITTED ACTIVITIES

- 2.1  Procuring
- 2.2  Storing
- 2.3  Supplying (delivering)
- 2.4  Exporting
- 2.5  Other activities: (please describe)

### 3. MEDICINAL PRODUCTS WITH SPECIAL REQUIREMENTS

- 3.1  Medicinal products pursuant to Art. 83 of Directive 2001/83/EC<sup>1</sup>
- Medicinal products pursuant to 67 of Directive 2001/82/EC
- 3.1.1  Narcotic or psychotropic substances
- 3.1.2  Medicinal products derived from blood
- 3.1.3  Immunological medicinal products
- 3.1.4  Radiopharmaceuticals (including radionuclides)
- 3.2  Medical gases
- 3.3  Medicinal products requiring the maintenance of the cold chain (storage and transportation at low temperatures)
- 3.4  Other activities: (Please specify or refer to Appendix 5)

Restrictions or clarifications regarding the scope of the authorisation (publicly available)

/./.....  
 \*Art. 5 of Directive 2001/83/EC or Art. 83 of Regulation 726/2004/EC

<sup>1</sup>Without prejudice to other authorisations due to national provisions

(Stamp of the District Government of Cologne)

I hereby certify that the translation overleaf is a true and correct rendering of the German document submitted to me.

The German document is attached to this translation.



D. Elliott

Dr. Donna Elliott  
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Linz am Rhein, 31.08.2020

Sworn translator for English and German  
Authorized by the President of the Cologne Higher Regional Court

Register Number: The translation overleaf is entered in my register under the number 820202283.