

# *National Authority Of Medicines And Medical Devices*

Report No: *NCF/001/2024/RO*

## **STATEMENT OF NON-COMPLIANCE WITH GMP**

***Exchange of information between National Competent Authorities (NCAs) of the EEA following the discovery of serious GMP non-compliance at a manufacturer<sup>1</sup>***

### **Part 1**

Issued following an inspection in accordance with Art. 111(7) of Directive 2001/83/EC as amended

The competent authority of Romania confirms the following:

The manufacturer: ***Linde Gaz Romania S.R.L.***

Site address: ***Strada Comertului Nr 7, Comuna Domnesti, Domnesti, 077090, Romania***

OMS Organisation Id. / OMS Location Id.: ***ORG-100002989 / LOC-100090376***

Other  
routine inspection

From the knowledge gained during inspection of this manufacturer, the latest of which was conducted on ***2024-07-25***, it is considered that **it does not comply with the Good Manufacturing Practice** requirements referred to in

- The principles and guidelines of Good Manufacturing Practice laid down in Directive (EU) 2017/1572 and/or Commission Delegated Regulation (EU) 2017/1569, as reflected by the product categories stated in Part 2.

Note to receiving authorities: Please contact the issuing authority within 20 working days in case there are critical(2) medicinal products potentially affected by this statement.

Manufacturing Authorisation Holders directly affected by this statement have failed to comply with their obligations under Art. 46 of Directive 2001/83/EC or Art. 93(1)(j) to (l) of Regulation (EU) 2019/6 and as a consequence the Qualified Person referred to in Art. 48 of Directive 2001/83/EC and Art. 97(1) of Regulation (EU) 2019/6 is unable to perform the batch certification referred to in Art. 51 of Directive 2001/83/EC and Art. 97 (6) and (7) of Regulation (EU) 2019/6.

In exceptional circumstances there may be no objection to the Qualified Person certifying affected batches thereby allowing their release provided all of the following conditions are fulfilled:

1. Batch certification is performed in order to maintain supply of critical medicinal products only.
2. A documented risk assessment has been performed by, or on behalf of, the Qualified Person and additional actions have been implemented by the manufacturing and/or batch release site to mitigate the risks posed by the non-compliance. Note: Repeated testing alone is not normally sufficient risk mitigation but, together with other actions, can form part of a strategy commensurate with the nature and the level of risk.
3. A thorough risk-benefit evaluation has been performed for the acceptance of risk and a report prepared that takes full account of the nature of the non-compliance with the involvement of:

- The Manufacturing Authorisation Holder and the Qualified Person of the site responsible for batch certification.
- The manufacturing site subject to this Statement of Non-Compliance, if different from the above.
- The relevant Marketing Authorisation Holder(s).

The report has been shared with the National Competent Authorities of the countries in which distribution of the affected batches is anticipated and that any comments from those authorities have been taken into account.

4. Written confirmation has been obtained from the National Competent Authorities in whose territories the affected batches are intended to be distributed that the product is considered critical on its territory, and that there is no objection to distribution.
5. The Supervisory Authority has been informed, if different from the above, and it has not suspended or revoked the relevant Manufacturing Authorisation.
6. The affected Marketing Authorisations have not been revoked or suspended.
7. Any further conditions imposed by the Supervisory Authority and other involved National Competent Authorities are met.

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<sup>1</sup>The statement of non-compliance referred to in paragraph 111(7) of Directive 2001/83/EC and Art. 94(2) of Regulation (EU) 2019/6, as amended, is also applicable to importers.

<sup>2</sup>See Appendix 3 of the relevant procedure in the Compilation of Union Procedures.

## Part 2

Human Medicinal Products	
<b>1 NON-COMPLIANT MANUFACTURING OPERATIONS</b>	
Include total and partial manufacturing (including various processes of dividing up, packaging or presentation), batch release and certification, storage and distribution of specified dosage forms unless informed to the contrary;	
<b>1.2</b>	<b>Non-sterile products</b>
	1.2.2 <i>Batch certification</i>
<b>1.5</b>	<b>Packaging</b>
	1.5.1 <i>Primary Packaging</i> 1.5.1.7 Medicinal gases
	1.5.2 <i>Secondary packaging</i>
<b>1.6</b>	<b>Quality control testing</b>
	1.6.3 <i>Chemical/Physical</i>

## Part 3

<b>1.Nature of non-compliance:</b>
Following the re-certification inspection on 25.07.2024, a total of 4 critical deficiencies, 6 major deficiencies and 8 other deficiencies were found. The manufacturer does not have sufficient and adequate staff to carry out the activity and also carries out transport and distribution activities through unauthorized and unqualified partners.
<b>Action taken/proposed by the NCA</b>
<b>Suspension of the manufacturing authorisation No. 28F in Part</b> suspension of Manufacturing Authorisation no.28F only for site address Str. Comerului, nr.7, Sat Domnesti, Comuna Domnesti, Județul Ilfov, cod poștal 077090, România
<b>Withdrawal, of current valid GMP certificate No. 052/2024/RO</b> Withdrawal of current valid GMP certificate no.052/2024/RO

2024-07-30

Name and signature of the authorised person of the  
Competent Authority of Romania

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**Confidential**  
**National Authority Of Medicines And Medical Devices**  
Tel: **Confidential**  
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EudraGMP

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