



BATCH VALIDITY STATEMENT EUROPEAN PHARMACOPOEIA REFERENCE STANDARDS (CRS) & (BRP)

This Batch Validity Statement has to be used in conjunction with Ph. Eur. general chapter 04/2015:51200 Reference Standards.

European Directorate for the Quality of Medicines & HealthCare (EDQM) - Council of Europe

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Name

Lincomycin hydrochloride

Catalogue

L0650000

code

Batch

number*

Assigned value

95.6%

Validity

Batch 4 is valid at the printing date: 2017-8-10

Additional information

Storage

The standard is intended for immediate use.

conditions

Recommended EDQM storage conditions for unopened containers : +5°C \pm 3°C

Safety data Safety Data Sheet is available from the detailed view or upon request. Click on the hyperlink to download the leaflet containing the instructions for use, if

available (Adobe Acrobat Reader version 5 or higher, or the corresponding browser plug-

Leaflet

Origin

in is needed to open the file)

click to download the leaflet

Click on the hyperlink to download the origin to check if import permit is required in your

country, if available (Adobe Acrobat Reader version 5 or higher, or the corresponding

browser plug-in is needed to open the file)

click to download Origin Of Goods.pdf

* Sub-batches 1.1, 1.2,1.3, etc., are obtained from the same batch of bulk material. Notice: the previous classification of the sub-batches 1a,1b, 1c will be gradually replaced with 1.1, 1.2, 1.3 etc.

This statement is valid at the date of printing: 2017-8-10

Legal notice: The Council of Europe (EDQM) makes no representation or warranty with respect to the accuracy, completeness, or currentness, of this The Council of Europe (EDQM) shall not be liable on account of any potential errors or omissions.

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INFORMATION LEAFLET Ph. Eur. Reference Standard

Lincomycin hydrochloride CRS batch 4

1. Identification

Catalogue code: L0650000

Unit Quantity: ca 100 mg

2. Scientific Information

2.1 Intended use

Reference Standard for laboratory tests as prescribed in the European Pharmacopoeia only. Established for use with the monograph(s): 0583, 0582, 0996.

2.2 Analytical information related to intended use, when applicable

The "as is" content is

: 95.6 % C18H35CIN2O6S (for 0583)

2.3 Uncertainty of the assigned value, when applicable

The uncertainty of the assigned value is not stated since it is considered to be negligible in relation to the defined limits of the method-specific assays for which the reference standard is used. Please also refer to Ph. Eur. chapter 5.12.

2.4 Validity

Ph. Eur. RS are periodically tested to ensure their continuous fitness for purpose. For each valid Ph. Eur. RS, a Batch Validity Statement at the time of use can be downloaded and printed from the EDQM website (Reference Standards Database).

2.5 Instructions for use

The container should not be opened until required for use. Allow the closed container to equilibrate at ambient temperature before opening to avoid uptake of moisture. Use "as is". Do not dry/desiccate before use. Ph. Eur. RS are for immediate use. Once the container has been opened, its entire content must be used immediately. Any further storage and re-use are not warranted.

3. Storage conditions

In the original container at $+5^{\circ}\text{C} \pm 3^{\circ}\text{C}$, protected from light. Re-instate promptly upon receipt.

4. Safety

For scientific research, development and analysis only. Handle in accordance with good occupational hygiene, safety and laboratory practices and take precautions to avoid exposure. More information is available at the EDQM website (Reference Standards Database): Safety Data Sheet for hazardous chemicals and Safety Data Statement for other materials.

5. Shipping conditions

Each Ph. Eur. RS is shipped under conditions that preserve its suitability for use and comply with the Each Ph. Eur. RS is shipped under conditions that preserve its suitability for use and comply with the Each Ph. Eur. RS is shipped under conditions that preserve its suitability for use and comply with the Each Ph. Eur. RS is shipped under conditions that preserve its suitability for use and comply with the Each Ph. Eur. RS is shipped under conditions that preserve its suitability for use and comply with the Each Ph. Eur. RS is shipped under conditions that preserve its suitability for use and comply with the Each Ph. Eur. RS is shipped under conditions that preserve its suitability for use and comply with the Each Ph. Eur. RS is shipped under conditions that preserve its suitability for use and comply with the Each Ph. Eur. RS is shipped under conditions that preserve its suitability for use and comply with the Each Ph. Eur. RS is shipped under conditions that preserve its suitability for use and comply with the Each Ph. Eur. RS is shipped under conditions that the Each Ph. Eur. RS is shipped under conditions that the Each Ph. Eur. RS is shipped under conditions that the Each Ph. Eur. RS is shipped under conditions that the Each Ph. Eur. RS is shipped under conditions that the Each Ph. Eur. RS is shipped under conditions that the Each Ph. Eur. RS is shipped under conditions that the Each Ph. Eur. RS is shipped under conditions that the Each Ph. Eur. RS is shipped under conditions that the Each Ph. Eur. RS is shipped under conditions that the Each Ph. Eur. RS is shipped under conditions that the Each Ph. Eur. RS is shipped under conditions that the Each Ph. Eur. RS is shipped under conditions that the Each Ph. Eur. RS is shipped under conditions that the Each Ph. Eur. RS is shipped under conditions that the Each Ph. Eur. RS is shipped under conditions that the Each Ph. Eur. RS is shipped under conditions that the Each Ph. Eur. RS is shipped under conditions that the Each Ph. Eur. RS is shipped under conditions relevant regulations. For more details see EDQM website (Reference Standards Database)

6. Warranties, Liabilities and responsibility

In the event of any safety concerns, please read carefully the safety data sheets of safety data statements available for each product. It is for Purchasers to determine independently the risks associated with the items and to take appropriate safety measures, including the provision of appropriate information, equipment and training of those persons coming into contact with the item.

- Warranties

Except for the use of Reference Standards in tests and assays carried out in accordance with the official methods of the European Pharmacopoeia and by professionals with the necessary technical skills and at their own discretion and risk, the EDQM makes no representation, contractual statement, or expression of opinion concerning the quality or safety of any item supplied, the presence of any defect in it, or its fitness for any particular purpose except that as described above.





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8. Citation

Users shall ensure that any reference made to an EDQM Reference Standard in any publication, presentation or public document (ex. scientific articles, data sheets for kits) bears the exact name, and catalogue code of the Reference Standard and the exact name and address of EDQM as given on the first page of this information leaflet.

The suitability for intended use has been officially adopted by the European Pharmacopoeia Commission.

10. Signature

This document is electronically signed by:

Dr Pierre Leveau Head of the Quality, Safety and Environment Division

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