





GMP - GOOD MANUFACTURING PRACTICE

CERTIFICATE

No. GMP_UZ - 09:2025

Is issued on the basis of a completed pharmaceutical inspection conducted in accordance with the regulation on the procedure for conducting inspections for compliance with the requirements of good manufacturing practice (GMP).

STATE ENTITY "CENTER OF GOOD PRACTICES" APPROVES

located at

231761, Republic of Belarus, Grodno Region, Skidel, Kizevich Street, 50-4
231286, Republic of Belarus, Grodno Region, Lida, Kachana Street, 19, 19/8
220007, Republic of Belarus, Minsk, Fabrisius Street, 30
220007, Republic of Belarus, Minsk, Fabrisius Lane, 3/11
220006, Republic of Belarus, Minsk, Mayakovsky Street, 1/1
220006, Republic of Belarus, Minsk, Mayakovsky Street, 1/5
220006, Republic of Belarus, Minsk, Mayakovsky Street, 1/22

RUE "BELMEDPREPARATY"

Compliance with the requirements of
O'zDSt 2766:2018 – "Good Manufacturing Practice - GMP"

The basis for pharmaceutical inspection was application of RUE "Belmedpreparaty" No.5-62/272 dated 11th November, 2023 for pharmaceutical inspection in accordance with the requirements of O'zDSt 2766:2018 - "Good Manufacturing Practice-GMP".



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GOOD MANUFACTURING PRACTICE — GMP CERTIFICATE APPENDIX

1. Aseptically prepared (list of dosage forms):	
☐ large volume liquids	
√ small volume liquids (in vials and ampoules)	
dispersions	
√ lyophilisates	
solids	
☐ semi-solids	
☐ other aseptically prepared products:	
√ powders β-lactams (carbapenems, cephalosporins)	
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√ eye drops (the type of medicine or the type of activity is shown).	
(the type of medicine or the type of activity is shown). 2. Medicines subject to sterilization at the end of production:	
 (the type of medicine or the type of activity is shown). 2. Medicines subject to sterilization at the end of production: □ large volume liquids 	
(the type of medicine or the type of activity is shown). 2. Medicines subject to sterilization at the end of production: □ large volume liquids √ small volume liquids (in vials and ampoules)	
(the type of medicine or the type of activity is shown). 2. Medicines subject to sterilization at the end of production: □ large volume liquids √ small volume liquids (in vials and ampoules) □ solids and implants	
(the type of medicine or the type of activity is shown). 2. Medicines subject to sterilization at the end of production: □ large volume liquids √ small volume liquids (in vials and ampoules) □ solids and implants □ semi-solids	
 (the type of medicine or the type of activity is shown). 2. Medicines subject to sterilization at the end of production: □ large volume liquids 	

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II. Non-sterile products	
☐ capsules, hard shell	
☐ capsules, soft shell	
☐ chewing gums	
☐ impregnated matrices	
☐ liquids for external use	
☐ liquids for internal use	
☐ medicinal gases	
☐ other solid dosage forms	
☐ pressurised preparations	
☐ radionuclide generators	
√ semi-solids (ointment, cream, gel)	
suppositories	
$$ tablets (coated and uncoated, not classified as antibiotics; uncoated classified as antibiotics: β -la	ctams
(ampicillin) and non-β-lactams)	
☐ transdermal patches	
☐ intraruminal devices	
☐ other non-sterile medicinal product:	
(the type of medicine or the type of activity is shown).	
III. Biological medicinal products	
□ blood products	
immunobiological products	
☐ cell therapy products	
☐ gene therapy products	
☐ tissue engineered products	
☐ biotechnology products	
☐ animal extracted products	
☐ other biological medicinal products:	
☐ immunobiological products ☐ cell therapy products ☐ gene therapy products ☐ tissue engineered products	
4월 10일 12일 12일 12일 12일 12일 12일 12일 12일 12일 12	
☐ animal extracted products	
and olological medicinal products.	
(the type of medicine or the type of activity is shown).	

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IV.Other products or manufacturing activity		
√ herbal products (tablets)		
☐ homoeopathic products		
□ other product		

(the type of medicine or the type of activity is shown).

Based on the information obtained during the pharmaceutical inspection conducted on 15-19.04.2024 and 15-18.01.2025 the applicant complies with the requirements of the Good Manufacturing Practice - GMP. The certificate is valid if all its pages (both main pages and additional pages) are presented. The validity of this certificate can be checked from the database of the State entity "Center of Good Practices". If the certificate is not provided in the indicated database, it is necessary to contact the working body that issued it.

The GMP_UZ – 09:2025 Good Manufacturing Practice - GMP certificate validity period from 04.02.2025 to 03.02.2028

Director of the SE "Center of Good Practices"

Dusmatov A.F.

(full name)