

3EC International a.s., Hraničná 18, 821 05 Bratislava, Slovak Republic Notified body No. 2265

EU QUALITY MANAGEMENT SYSTEM CERTIFICATE No. 2024-MDR/QS-022

UNIMED PHARMA spol. s r.o.

Oriešková 11, 821 05 Bratislava, Slovak Republic

SRN č.: SK-MF-000003111

This EU Quality Management System Certificate issued in accordance with the Regulation (EU) 2017/745 of the European Parliament and of the Council on medical devices as amended confirms, that quality management system of medical device:

OCUflash® eye drops, solution Models / variants: see Annex I Intended purpose: see Annex II

MD class IIb

(detailed list is stated in the annex(es) if applicable)

meets the requirements on quality management system according to the Chapter I and III of Annex IX of the Regulation (EU) 2017/745 of the European Parliament and of the Council on medical devices as amended.

Conditions for or limitations to the validity of the certificate: N/A

Validity of the certificate is conditional upon positive results of regular surveillance audits.

Notified body No. 2265 has performed assessment of the quality management system of the abovementioned medical device and found that it meets the requirements stated above. The outcome of the assessment of the quality management system of the abovementioned medical device is stated in the MD Technical Documentation Assessment Report No. MDR287_2023 from 15.02.2024, MD Clinical Evaluation Report No. MDR287_2023 from 15.02.2024 and MD Audit Report No. SK-0297-23 from 15.02.2024. Information on all examinations and tests performed is stated in the abovementioned reports and is available on request.

This **EU Quality Management System Certificate** applies only to the quality management system of the abovementioned medical device. The certificate validity is conditional upon fulfilment of relevant legal requirements by the manufacturer.



Valid from: 22.02.2024 Valid until: 22.02.2029

First issue: 22.02.2024 Revision: 00

History: see Annex III



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In Bratislava, Slovakia, 22.02.2024