

DECLARATION OF CONFORMITY

Expiration 2028-12-31

MANUFACTURER: OASIS® Medical, Inc.
510-528 S. Vermont Ave.
Glendora, CA USA 91741

SRN: US-MF-000034540

Basic UDI-DI: 08458190PVA5XX13704ND

EUROPEAN REPRESENTATIVE: Donawa Lifescience Consulting Srl
Piazza Albania, 10
00153 Roma, Italy

PRODUCT: SOFT CELL® PVA Foam Surgical Spears
REF #: 0525

INTENDED PURPOSE: The PVA Surgical Spear is indicated for short-term transient use in ophthalmic surgical procedures for absorbing excess fluids from the operative field.

CLASSIFICATION: Class IIa, Rule 6 according to Annex IX of Directive 93/42/EEC

CONFORMITY ASSESSMENT ROUTE: Annex II of Directive 93/42/EEC

OASIS® MEDICAL, INC. DECLARES THAT THE ABOVE MENTIONED PRODUCTS CONFORM TO THE PROVISIONS OF THE COUNCIL DIRECTIVE 2023/ 607 AND EU MDR ARTICLE 120 FOR MEDICAL DEVICES. ALL TECHNICAL DOCUMENTATION IS RETAINED UNDER THE PREMISES OF THE MANUFACTURER.


TO THE BEST OF ITS KNOWLEDGE, INFORMATION AND BELIEF, OASIS® MEDICAL, INC. IS IN COMPLIANCE, IN ALL MATERIAL RESPECTS WITH ITS QUALITY MANAGEMENT SYSTEM ACCORDING TO EN ISO 13485:2016. THIS DECLARATION IS ISSUED UNDER THE SOLE RESPONSIBILITY OF OASIS® MEDICAL, INC.

NOTIFIED BODY: DQS 0297 – DQS Medizinprodukte GmbH
August-Schanz-Strasse 21
60433 Frankfurt a.M.
Germany

EC CERTIFICATE(S): EC Certificate Number: 288050 MR2
Issued: 2020-10-22
Valid Until: 2023-09-15

CONFIRMATION LETTER EU 2023-607: Issued: 2023-06-23
(MDD CERTIFICATE EXTENSION GRANTED BY DQS)



 11 Sept 2024

Ishan Patil Date
Director of Regulatory Affairs, OASIS® Medical, Inc.