Version: 2.0

Date: 2022-05-26

Declaration of Conformity

for Immersion Oil

Regulation (EU) 2017/746 of the European Parliament and of the Council of 5 April 2017 concerning In Vitro Diagnostic Medical Devices

The undersigned, under their sole responsibility, declares that the products described in this document meet the Council provisions that apply to them and the CE Mark may be affixed.

General Product Name:	Immersion Oil	
Legal Manufacturer: (Name on Label)	Pro-Lab Diagnostics 3 Bassendale Road Bromborough CH62 3QL UK	
SRN:	GB-MF-000003833.	
Basic UDI-DI:	506047246ImmersionOilA7.	
Variants:	As per Appendix II (This document) – Product Listing/Schedule.	
Intended Purpose:	As an accessory to IVD medical devices in the microscopic examination of prepared slides from clinical specimens.	
IVDR Classification:	ssification: Class A [Rule 5].	
Notified Body:	Not applicable for Class A devices.	
CE Certificate:	te: Not applicable for Class A devices.	
EU Authorised Representative:	Advena Limited. Tower Business Centre, 2 nd Flr., Tower Street, Swatar, BKR 4013 Malta.	
EU Authorised Representative SRN:	MT-AR-00000234	
IVDR Assessment Route:	Issuing of the Declaration of Conformity in accordance with Article 17 after drawing up the technical documentation in Annexes II and III of the EU IVDR 2017/746.	

Name	Mike Owen	Position	Quality Management System Coordinator			
Signed	maun	Date	2022-05-26	Place	Pro-Lab Diagnostics	

Who is the natural and legal person with responsibility for the design, manufacture, packaging and labelling before the device is placed on the market under this manufacturer's name regardless of whether these operations are carried out by the manufacturer or on his behalf by a third party.

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Appendix I – Applicable Standards

This present declaration is also in conformity with the following European standards and Common Specifications (CS):

Standard/CS/Document Name	Description		
	Regulation (EU) 2017/746 of the European Parliament and of the		
2017/746	Council of 5 April 2017 concerning In Vitro Diagnostic Medical		
	Devices		
EN ISO 13485:2016+A11:2021	Medical Devices – Quality Management Systems – Requirements		
EN 130 13485.2010+A11.2021	for Regulatory Purposes		
EN ISO 14971:2019+A11:2021	Medical Devices – Application of Risk Management to Medical		
EN 130 14971.2019+A11.2021	Devices		
EN ISO 15223-1:2021	Medical devices. Symbols to be used with information to be		
EN 130 13223-1.2021	supplied by the manufacturer - General requirements		
EN ISO 20417:2021	Medical devices. Information to be supplied by the manufacturer		
	In vitro diagnostic medical devices — Information supplied by the		
EN ISO 18113-1:2011	manufacturer (labelling) — Part 1: Terms, definitions and		
	general requirements.		

Appendix II - Product Listing/Schedule

Catalogue Number	Device Description	EMDN Code	UDI-DI	
PL.396	Immersion Oil	W010399	05060472463517	

Version History

Version	Compiled by	Date	Description
1.0	M.Owen	2019 12 06	First issue
2.0	M.Owen	2022 05 26	Updated for EU IVDR 2017/746