DAC-SpectroMed S.R.L. Chisinau

Declaration of EC-Conformity

Product: ASLO-DAC

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Page	1 / 1

DAC-SpectroMed SRL

Nicolae Testemitanu str. 37, MD-2025, Chisinau, Republic of Moldova

hereby declares under its own responsibility that the in-vitro diagnostics medical devices:

Product/Trade Name	Catalogue Number	Intended Purpose
ASLO-DAC	1031A100	set of reagents for serological analyses
ASLO-DAC	1031A200	set of reagents for serological analyses
ASLO-DAC	1031A250	set of reagents for serological analyses
ASLO-DAC	1031A500	set of reagents for serological analyses

Risk Class: Class B according to Rule 4 of Annex VIII of the IVDR 2017/746.

to which this declaration relates comply with the provision of the following relevant Union legislation:

• Regulation (EU) 2017/746 of the European Parliament and of the of 5 April 2017 on in vitro diagnostic medical devices.

The following conformity assessment procedure(s) has(ve) been followed

 Annex IX Conformity Assessment based on a Quality Management System and on assessment of technical documentation.

DAC-SpectroMed SRL has a Quality System in place based on EN ISO 9001:2015 and EN ISO 13485:2016, issued by the IQNet&SRAC.

Our Authorized Representative in EU is Qarad EC-REP BV, with address at Pas 257, 2440 Geel, Belgium.

General Manager Mordvinov Ghenadie 05.03.2024

Specialist of Normative-Technical Documentation and Certification Department

Anghelova Ana

05.03.2024