

CE DECLARATION OF CONFORMITY

Part no.	Product name	Content	
RT2951	AMP Rapid Test SARS-CoV-2 Ag	10 cassettes	
RT2952	AMP Rapid Test SARS-CoV-2 Ag	25 cassettes	
RT2952-C	AMP Rapid Test SARS-CoV-2 Ag w. controls	25 cassettes + pos. & neg. control	
RT2952-S	AMP Rapid Test SARS-CoV-2 Ag	25 cassettes	
IVD-classification acc. to directive 98/79/EC: other IVD			

We, AMEDA Labordiagnostik GmbH, Krenngasse 12, 8010 Graz, Austria, declare under sole responsibility that the products described above are in compliance with Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on in-vitro diagnostic medical devices and that the following harmonized standards have been resp. are applied in development, design and manufacturing.

Instructions for Use	EN ISO 18113-2:2011	In vitro diagnostic medical devices - Information	
		supplied by the manufacturer (labelling) - Part 2: In	
		vitro diagnostic reagents for professional use	
Performance	EN 13612:2002	Performance evaluation of IVD medical devices	
Stability	EN ISO 23640:2015	In vitro diagnostic medical devices – Evaluation of	
		stability of in vitro diagnostic reagents	
Symbols	EN ISO 15223-1:2016	Medical devices – Symbols to be used with medical	
		device labels, labelling and information to be supplied	
		- Part 1: General requirements	
Risk analysis	EN ISO 14971:2012	Medical devices - Application of risk management to	
		medical devices	

The products are CE marked.

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Graz - 07.01.2021

Gerald Herfort

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AMEDA Labordiagnostik GmbH