WE200.02-DoC Revision: 04en

EU DECLARATION OF CONFORMITY

LO DECLARAT	ion or com	
Product		
Product name/Trade name	REF	Basic UDI-DI
Substrat/ Substrate LINE	WE200.02	++E654WE20002V02JE
Manufacturer		
Virotech Diagnostics GmbH Waldstrasse 23 A2		
63128 Dietzenbach Germany		
SRN: DE-MF-000025733		
We, as the manufacturer of the device(s) take mentioned product(s) meet(s) the provisions of the Regulation EU 2017/746 on <i>In vitro</i> Diagnostic W	he following Regulation	•
	ledical Devices	
RISK CLASS ⊠ A □ B □ C □ D		
CONFORMITY ROUTE		
☐ ANNEX IX Technical Documentation Examination	EU CERTIFICATE #: Name of Notified Body: mdc medical device certification GmbH Notified Body Identification: 0483	
□ ANNEX IX Full Quality System	EU CERTIFICATE #: Name of Notified Body: mdc medical device certification GmbH Notified Body Identification: 0483	
☐ ANNEX X Type Examination	EU CERTIFICATE #: Name of Notified Body: mdc medical device certification GmbH Notified Body Identification: 0483	
□ ANNEX XI Production Quality System	EU CERTIFICATE #: Name of Notified Body: mdc medical device certification GmbH Notified Body Identification: 0483	
⊠ ANNEX II+III		
Common Specifications (CS): n/a Common Specifications have not been issued for this p	product.	
DocuSigned by:		
Marullo Salio Signature	Dietzenbach Issued in	2023-06-22 Date of issue
Marcello Salio	Managing Director	
Name	Function	