BIO-RAD	GLOBAL FORM	04.01.GLB.FRM.00125
	EU DECLARATION OF CONFORMITY	
Division/Group: RAQA		Revision: 1

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MANUFACTURER: Bio-Rad ADDRESS: 3 Boulevard Raymond Poincaré, 9243	0 Mames-la-Coquette, France		
EUROPEAN AUTHORIZED REPRESENTATIVE	1		
PRODUCT(S) NAME(S) and CATALOG NUMBE	R(S): Genscreen™ Ultra HIV Ag-Ab -	- cat# 72386-72388	
GENERIC DEVICE GROUP CODE (GMDN nome	nclature): 48445		
GENERIC DEVICE GROUP TERM (GMDN Nom (EIA)	enclature): HIV1/HIV2 antigen/antiboo	dy, kit, enzyme immunoassay	
We hereby declare that the above mentioned prod	uct(s) meet(s) the provisions of the follo	wing Directive	
■ Directive 98/79/EC of the European Parliament medical devices	and of the Council of 27 October 1998	on in vitro Diagnostic	
CLASSIFICATION:			
☑ ANNEX II-A □ ANNEX II-B	☐ DEVICE FOR SELF TESTING☐ OTHER DEVICE		
CONFORMITY ROUTE			
☐ ANNEX III ☑ ANNEX IV.3 Full Quality System	EC CERTIFICATE No.: 9150 Name of Notified Body: GMED Notified Body Identification No.: 0	459	
☑ ANNEX IV.4 Product Design Examination	Expiration Date: May 26th, 2025 EC CERTIFICATE No.: 8977 Name of Notified Body: GMED Notified Body Identification No.: 0 Expiration Date: May 26th 2024	1459	
☐ ANNEX V Type Examination	Expiration Date . Iviay 20° 2024		
☐ ANNEX VII Production Quality System			
NEW PRODUCT(S) (Notification according to artic	e 10 point 4)	⊠ NO	
Date of the first issuance of the EU Declaration	of Conformity: July 22, 2004		
Jan D	Marnes-la-Coquet		
Signature	Issued in	Date	

Sylvie Fernez Associate Director Regulatory Affairs Name Function