

**UNILABS GROUP TEMPLATE** 

Public Declaration

Code: FORM-HQ-RA-01

Revision AA

Effective Date: 23-May-2024

# **Public Declaration for In-House devices**

### Part A - Information about the manufacturing Health Institution

Legal Entity of the Health Institution			
Name	Address		
Unilabs laboratoire d'analyses médicale	Chemin des Perrières 2, 1296 Coppet		

List of Health Institution Sites under scope of this Public Declaration					
Name	ame Department		Contact Person		
Unilabs, laboratoire	N/A	Chemin des Perrières 2,	Françoise Morel		
d'analyses médicales SA		1296 Coppet			

Contact Data of Responsible Person for the Legal Entity					
Name	Job Title	Telephone Number	E-mail		
Françoise Morel	Quality Manager	+41 79 636 75 24	francoise.morel2@unilabs.com		

#### Declaration

Above mentioned Health Institution states for all tests listed in part B:

- i) The device is manufactured or modified and used in sites within the same legal entity under the scope of this Public Declaration.
- ii) The device is manufactured or modified by this Health Institution under compliant EN ISO 15189 quality management system.
- iii) The device is compliant with relevant General Safety and Performance Requirements (GSPR) as depicted in Annex I of *in vitro* Medical Device Regulation (EU) 2017/746 (IVDR) and according to Article 5 (5).
- iv) A reasoned justification is provided in case applicable General Safety and Performance Requirements are not fully met.
- v) Argumentation for use of in-house tests is documented in the quality management system, through technical file documentation.

I, the undersigned, in my capacity as Laboratory Director certify that the information given is true, accurate, present and complete to the best of my knowledge. I understand that if I have deliberately given any false information regarding any situation, I am liable against the law

Signed on behalf the organisation					
Name of the responsible of the lab (Technical Director or equivalent technical role)	Date	Signature  Firmato da:			
Lorena Miele	08-Oct-2024   10:30 CEST	Lowe / ws			
		B130907C28A541A			



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## Part B - Description of the In-House devices

List of	List of assays or in-house devices within the Family						
#	In-house reference	In-house name	Device Type	Risk class	Intended Purpose	Applicable GSPR fully met?	Information on and justification for applicable GSPR that are not fully met (using the numbering as in Annex I of the IVDR/MDR)
1	Constitutional Cytogenetics	Constitutional Cytogenetics family	IVD	С	See IMS reference: 1057594	Yes	See IMS reference: 1058183
2	Onco Hematology Cytogenetics	Onco Hematology Cytogenetics family	IVD	С	See IMS reference: 1057593	Yes	See IMS reference: 1058152
3	FISH	FISH family	IVD	С	See IMS reference: 1058228	Yes	See IMS reference: 1058230