

CERTIFICATE OF NOTIFICATION

This is to certify that, according to the council directive 98/79/EC, SUNGO performed all notification duties and responsibilities as the European authorized representative of:

Applicant: Address: Xiamen AmonMed Biotechnology Co.,Ltd. Unit 503, 120 Xinyuan Road, Haicang District, Xiamen, Fujian, China.

The Manufacturer has provided SUNGO with all the appropriate declarations according to the 98/79/EC Directive requirements including the EC Declaration of Conformity confirming that his In vitro diagnostic medical device, as stipulated here below, is fulfilling the applicable requirements of the European Council Directive 98/79/EC.

Product(s):	COVID-19 IgM/IgG test kit (Colloidal Gold);
	COVID-19/Influenza A virus/Influenza B virus test kit (Rare earth
	nano fluorescence immunochromatography)
Type(s):	See Annex
Product Classification:	IVDD Other

Where the manufacturer affix's the CE marking to the product listed they must ensure that all the requirements of the appropriate EU directive(s) have and continue to be met.

The notification of aforementioned device has been completed by the European Representative in Netherlands. The Netherlands Competent Authority is notified of the manufacturer's medical devices and has allocated registration. NOTIS number is CIBG- 20200534.



Issued: Mar. 19 2020 Cert. No.: EU201518 Expiration Date: Mar. 18 2025



This is not a CE mark and is only provided as a template for informational purpose.



Annex to Cert. No.: EU201518

Product Name	Туре
COVID-19 IgM/IgG test kit (Colloidal Gold)	25 tests/kit,50 tests/kit
COVID-19/Influenza A virus/Influenza B virus test	
kit (Rare earth nano fluorescence	25 tests/kit,50tests/kit
immunochromatography)	

END OF THE ANNEX