

EC DECLARATION OF CONFORMITY

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 including amendments

Manufacturer: **GENERI BIOTECH s.r.o.**
Machkova 587/42, 500 11 Hradec Kralove 11 – Trebes, Czech Republic

hereby declares, that **real-time PCR kits** as *In vitro diagnostic medical devices (IVD)* listed below:

gb Sarbeco E (primary test)	Cat. no. 3227-500 and 3227-100
gb SARS-CoV-2 RdRP (confirmation test)	Cat. no. 3228-500 and 3228-100
gb Sarbeco N (primary test)	Cat. no. 3229-500 and 3229-100
gb SARS-CoV-2 N (confirmation test)	Cat. no. 3230-500 and 3230-100
gb SARS-CoV-2 Multiplex	Cat. no. 3231-500, 3231-200, and 3231-050
gb SARS-CoV-2 Combi	Cat. no. 3232-500 and 3232-100
gb ONCO BRAF (V600E)	Cat. no. 3241-048 and 3241-024
gb ONCO EGFR (T790M)	Cat. no. 3245-048 and 3245-024
gb ONCO BCR-ABL DETECT	Cat. no. 3246-048
gb Human B2M mRNA	Cat. no. 3153-500 and 3153-100
gb GENETIC HLA-B*27	Cat. no. 3257-100 and 3257-025

comply with the essential requirements of *Annex I of the Directive 98/79 EC* including amendments. The conformity was established according to *Annex III* (class: other IVD products) *of the Directive 98/79 EC*. Following harmonized technical standards were used to demonstrate the compliance:

EN ISO 13485:2016
EN ISO 18113-2:2011
EN ISO 23640:2015
EN 13612:2002
EN ISO 14971:2012
EN ISO 15223-1:2016

In: Hradec Kralove
Date: 3. 8. 2020


PharmDr. Radovan Haluza, Ph.D.
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