

## EC DECLARATION OF CONFORMITY

According to Directive 98/79/EC, Annex III, on *in vitro* Diagnostic Medical Devices

**Manufacturer:** BTNX Inc., 570 Hood Rd. #23, Markham, Ontario, L3R 4G7 Canada  
**Product Name:** Rapid Response™ COVID-19 Antigen Test Cassette  
**Product Code(s):** COV-19C25  
**EDMA / GMDN:** 15.70.90.90  
**GMDN:** 64912  
**EDMA Description:** Other Virology - RT & POC  
**Risk Classification:** Other  
**Authorized EC Rep.:** MDSS GmbH, Schiffgraben 41, 30175 Hannover, Germany

We, the manufacturer, declare under our sole responsibility that the above-mentioned products meet all the provision of the council directive 98/79/EC for *in vitro* Diagnostic Medical Devices that apply to it. This self-declaration is according to Annex III (excluding Section 6) of the Directive.

Markham, Ontario, Canada, as of 2020-06-24 for BTNX Inc.,



**Khasim Ali Khan**  
QA/RA Manager  
BTNX Inc., 570 Hood Road, Unit 23  
Markham, ON, L3R 4G7, CANADA

BTNX Inc. is a medical device  
manufacturer certified under:  
ISO 13485:2016  
by Intertek Testing Services Canada



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