

## Sample-to-Answer in 3 **Easy Steps**

















Direct Sample. Extraction-Free. Simple yet Superior.

## Multi-Platform, High-Throughput COVID-19 Testing



Complete Lyophilized Test in a Single Vial No other reagent required



No sample preparation at all Direct sample addition



**High-Throughput Workflow** 80 Minutes for 96 to 384 samples



Includes two sets of controls RNase P gene detection as SPC RNA IAC as RNA amplification control

BioGX Xfree™ COVID-19 Direct RT-PCR and FDA recommended comparator method clinical concordance study.

QuantStudio™ 5 Direct Sample Addition	FDA recommended comparator method	
Xfree™ COVID-19 Direct RT-PCR	Positive	Negative
Positive	57	0
Negative	3*	35
Positive Percent Agreement (95%CI)	95.0% (86.3-98.3%)	
Negative Percent Agreement (95%CI)	100.0% (90.1-100.0%)	

<sup>\*</sup>The three direct samples with discordant results were identified as positive by the comparator method with Ct values between 37.6 & 40.5.

Specimen Types: Nasopharyngeal swab, Nasal swab, Mid-turbinate swab, Oropharyngeal swab transported in Copan ESwab™, Copan UTM®, BD UVT, VTM and dry swab (resuspended in saline)

This product has not been FDA cleared or approved, but has been authorized for emergency use by FDA under an EUA for use by

This product has been authorized only for the detection of nucleic acid from SARS-CoV-2, not for any other viruses or pathogens The emergency use of this product is only authorized for the duration of the declaration that circumstances exist justifying the authorization of emergency use of in vitro diagnostics for detection and/or diagnosis of COVID-19 under Section 564(b)(1) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. § 360bbb3(b)(1), unless the declaration is terminated or authorization is revoked sooner.

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