WHO Emergency Use Assessment Coronavirus disease (COVID-19) IVDs
PUBLIC REPORT

Product: Novel Coronavirus (2019-nCoV) IgM/IgG Antibody Rapid Test
EUL Number: EUL 0553-219-00
Outcome: Not Recommended.

The EUL process is intended to expedite the availability of in vitro diagnostics needed in public health emergency situations and to assist interested UN procurement agencies and Member States in determining the acceptability of using specific products in the context of a Public Health Emergency of International Concern (PHEIC), based on an essential set of available quality, safety and performance data. The EUL procedure includes the following:


Novel Coronavirus (2019-nCoV) IgM/IgG Antibody Rapid Test with product code MGRT20200630, manufactured by Bioscience (Tianjin) Diagnostic Technology Co., Ltd, No.201.10 Si Wei Road, Dongli District, Tianjin, China, is not eligible for WHO procurement.

Quality Management Systems Review

To establish the eligibility for WHO procurement, Bioscience (Tianjin) Diagnostic Technology Co., Ltd was asked to provide up-to-date information about the status of their quality management system.

Based on the review of the submitted quality management system documentation by WHO staff and external technical experts (assessors), it was established that insufficient information was provided by Bioscience (Tianjin) Diagnostic Technology Co., Ltd to fulfil the requirements described in the “Instructions for Submission Requirements: In vitro diagnostics (IVDs) Detecting SARS-CoV-2 Nucleic Acid, PQDx_347.”