

## CE Technical Documentation Review Report

**Applicant:** **Artron Laboratories Inc.**  
3938 North Fraser Way, Burnaby,  
BC, V5J 5H6, Canada

**Report Number:** **50353284 001**

**Examination intent:** Examination the completeness of the Technical Documentation according to the requirements of the In Vitro Diagnostic Medical Devices Directive 98/79/EC Annex III

**Product(s):** COVID-19 IgM/IgG Antibody Test

**Type(s)/Model(s):** Cassette

**Classification:** Other IVD products  
(according to manufacturer's declaration)

**Examination period:** Mar.18.2020

**Date of expiry:** May.26.2024

**Review result:** During the examination of the provided Technical Documentation (CE-COVID-19-A03-51-322, Revision 01, Dated 2020-Mar-15) no Non-compliance according to the requirements of the In Vitro Diagnostic Medical Devices Directive 98/79/EC Annex III was detected.

  
Yuhong CHEN  
Vice General Manager Medical Greater China  
TÜV Rheinland (China) Ltd.



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or please visit our official website: <http://www.tuv.com>