



SQI Diagnostics Announces New Corporate Website

New Website Provides Essential Information on Company's Strategic Initiative

TORONTO, February 18, 2021 -- SQI Diagnostics Inc. (the "Company" or "SQI") (TSX-V: SQD; OTCQB: SQIDF), a precision medicine company that discovers, develops, and commercializes innovative rapid diagnostic testing for healthcare professionals, patients and consumers worldwide, today is proud to announce the launch of its newly redesigned website at <https://www.sqidiagnostics.com>. The revamped site features a streamlined, modern design, improved functionality and easy access to essential information to help educate all stakeholders about the Company and its new strategic vision, including the latest product development and investor information.

"We are thrilled to debut our new Company website to our customers, partners, investors and visitors who are looking to understand the breadth of SQI's products and services," said Rob Chioini, SQI's CEO. "This new comprehensive website redesign truly encompasses our Company vision and enables each visitor to have the same experience and access to our new product development for our novel COVID-19 and lung transplantation diagnostics, along with our home testing platform and custom pharmaceutical diagnostic services."

About SQI Diagnostics

SQI Diagnostics, Inc. is a precision medicine company that discovers, develops, and commercializes innovative rapid diagnostic testing for healthcare providers, patients, and consumers worldwide. The Company's proprietary advanced diagnostics target organ transplant, autoimmune disease and COVID-19 testing which includes the developmental direct-to-consumer COVID-19 HOME Antibody Test, the RALI-Dx™ COVID-19 Severity Triage Test and the COVID-19 RALI-fast™ Severity Triage Point-of-Care (POC) Test. SQI's rapid diagnostic tests are intended to be sold to healthcare professionals so that patients can get accurate results and fast effective treatment, and direct-to-consumers so that individuals can be empowered to improve their health outcomes from the comfort of home.

SQI is fast-tracking the development of three COVID-19 diagnostic tests: a direct-to-consumer COVID-19 Antibody Test and two COVID-19 Severity Triage tests. The COVID-19 HOME Antibody Test identifies the presence of IgM, IgA and IgG antibodies of SARS-CoV-2 in individuals suspected to have been infected with COVID-19 and asymptomatic individuals wanting to know if they have been exposed. The test is > 99% accurate with results delivered in 24-48 hours. The Company currently expects to apply to the U.S. Food and Drug Administration ("FDA") for Emergency Use Authorization ("EUA") for its COVID-19 HOME Antibody Test in the second quarter of calendar year 2021. Should the COVID-19 HOME Antibody Test receive regulatory approval, the test is expected to be available direct-to-consumer which would allow individuals to avoid travelling to a clinic or hospital to be tested for the presence of the SARS-CoV-2 antibody.

The RALI-Dx™ COVID-19 Severity Triage Test and the RALI-fast™ COVID-19 Severity Triage POC Test each help clinicians identify which patients with SARS-CoV-2 will have a severe inflammatory response and should be admitted to the hospital or not. Both tests measure the critical biomarker IL-6 which plays a key role in the cytokine storm phase of COVID-19. The RALI-Dx™ delivers results from the lab in about 50 minutes while the RALI-fast™ delivers results at the patient point-of-care in about 15 minutes. The Company currently expects to apply for EUA to the FDA and for an Interim Order with Health Canada for both tests in the first and second quarters of calendar year 2021, respectively.

Under organ transplant, SQI is pioneering the development of an advanced diagnostic test that increases the chance of successful lung transplant by assessing the health of the donor organ prior to transplant surgery. The Company's

developmental TORdx™ Lung Test can detect inflammation at the molecular level to assess the health of the donor lung, enabling surgeons to transplant healthy lungs which otherwise would have been rejected; there is currently no other such test. SQI has partnered clinical development with the University Health Network (UHN) Hospitals, one of the largest health and medical research organizations in North America. Upon regulatory approval of the TORdx™ Lung Test, clinical development is planned for diagnostic tests designed to increase the chance of successful kidney and liver transplant.

Under autoimmune disease testing, SQI has a direct-to-consumer Celiac Disease and a Rheumatoid Arthritis (RA) Test that enable people to screen for the diseases from the comfort of their home. The direct-to-consumer RA Test can help identify and confirm RA symptoms for timely care and treatment. The direct-to-consumer Celiac Test confirms disease and validates the effectiveness of dietary and lifestyle changes to confirm the autoimmune response is improving.

The Company is not making any express or implied claims that its products can eliminate, cure or contain COVID-19 (or SARS-2 Coronavirus) at this time. For its research and development, the Company is collaborating with UHN Hospitals, one of the largest health and medical research organizations in North America.

For more information, please visit www.sqidiagnostics.com.

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FORWARD-LOOKING INFORMATION

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