Press Release

Business Partnership with Nihon University, Gunma University, and Tokyo Medical University for a Rapid Diagnostic Methods for Viruses in the Field of Infectious Diseases, Including Novel Coronavirus - A Rapid Diagnostic Method with High Sensitivity that can be Visually Determined without the Need for a Detection Device in 25 Minutes of Reaction from Samples such as Saliva -

Osaka, Japan, June 22, 2020 – Shionogi & Co., Ltd. (President & CEO: Isao Teshirogi, Ph.D., hereafter: “Shionogi”) announced today that Shionogi has agreed with Nihon University, Gunma University, and Tokyo Medical University on a license agreement regarding a new rapid diagnostic method for viruses including the novel coronavirus (SARS-CoV-2). We will work in collaboration with public institutions, academia including the above three universities, and partner companies to put this diagnostic method into practical use.

With continued social disruption caused by the worldwide spread of SARS-CoV-2, the spread of infection from not only patients with symptoms but also non-progressor and the infected person in the incubation period has become a major issue in the control of infectious disease. At present, as the test methods for diagnosing patients with novel coronavirus infection (COVID-19), PCR (polymerase chain reaction) are used for diagnosing COVID-19 for detecting nucleic acids in viruses from samples collected from the nasal cavity, pharynx and saliva as well as antigen test kits. However, many challenges remain with these testing methods, such as the need for specific detector, the ease and speed of measurement, and the risk of infection of healthcare workers during sample collection. Therefore, there is a need for highly sensitive and inexpensive diagnostic methods that overcome these challenges.

The Joint Research Team, consisting of Nihon University, Gunma University, and Tokyo Medical University, has succeeded in developing a unprecedented innovative viral rapid diagnostic method using signal amplification by ternary initiation complexes method (SA TIC method). The SA TIC method is a technology that can measure not only specific genes, but also mutant genes, as well as in-vivo molecules such as proteins and metabolites, using a simple method with high specificity. This rapid diagnostic method using SATIC methods has all of the following features.

- The presence or absence of SARS-CoV-2 or influenza virus can be easily determined visually without a detector.
- Virus determination can be made in about 25 minutes after sample collection.
- There is no non-specific reaction such as false positive reaction, and high sensitivity equivalent to that of PCR.
- The ability to detect viruses from saliva and sputum as well as nasopharyngeal swabs reduces the invasiveness of the patient and the risk of infection among healthcare workers associated with specimen collection as much as possible. In the case of saliva, specimens can be collected by the patient themselves.
On June 3, 2020, Shionogi launched IgG/IgM Antibody-test Kit for COVID-19 as a research reagent aimed at identifying the number of SARS-CoV-2 infected patients. On the other hand, this diagnostic method allows us to easily know the presence or absence of infection at the time of testing for COVID-19 and influenza virus infections disease, etc. in a short time. As indications for the practical use of this diagnostic method, it is assumed that the presence or absence of infections at medical facilities and quarantines, and in the future, the screening of infections by travellers from abroad will be considered. Since no testing method combines all of the above-mentioned features, the use of this test method enables the diagnosis of SARS-CoV-2 infected individuals, including those without symptoms, quickly and conveniently, and provides a timely understanding of the trends in domestic infected individuals. In addition, it offers the benefits of early diagnosis-based preventive measures against aggravation and early administration of treatment drugs.

Shionogi is committed to “Protect people worldwide from the threat of infectious diseases” as our key focus. We are not limiting ourselves to the research and development of therapeutic medications, but are also focused on the total care of infectious disease, through awareness building, prevention, diagnosis, and treating exacerbations, as well as the infection itself. Shionogi will work closely with government, industry, and academia including the above three universities, and will continue to strive to fulfill the early commercialization of this rapid diagnostic method to fulfill our social responsibility and to contribute to re-establishing the safety and security of society to support early containment of the pandemic.

About Signal Amplification by Ternary Initiation Complexes method (SATIC method)

Figures 1-4 show the principle of the SATIC method and the results of tests of the actual samples provided by COVID-19 patients.

Figure 1: Basic principle of SATIC method
(P: Primer, CT: Circular Template, THT-HE: ThT derivative, dNTP: deoxyribonucleoside triphosphate)
Figure 2: Flow to judgment (procedure)

Figure 3: Test results by PCR method
Forward-Looking Statements
This announcement contains forward-looking statements. These statements are based on expectations in light of the information currently available, assumptions that are subject to risks and uncertainties which could cause actual results to differ materially from these statements. Risks and uncertainties include general domestic and international economic conditions such as general industry and market conditions, and changes of interest rate and currency exchange rate. These risks and uncertainties particularly apply with respect to product-related forward-looking statements. Product risks and uncertainties include, but are not limited to, completion and discontinuation of clinical trials; obtaining regulatory approvals; claims and concerns about product safety and efficacy; technological advances; adverse outcomes of important litigation; domestic and foreign healthcare reforms and changes of laws and regulations. Also for existing products, there are manufacturing and marketing risks, which include, but are not limited to, inability to build production capacity to meet demand, unavailability of raw materials and entry of competitive products. The company disclaims any intention or obligation to update or revise any forward-looking statements whether as a result of new information, future events
or otherwise.

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References
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   Succeeded in developing rapid diagnostic method for COVID-19 using innovative nucleic acid
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2. Press release on June 3, 2020
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3. H. Fujita et al. Novel one-tube-one-step real-time methodology for rapid transcriptomic biomarker
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