INTENDED USE
The RHINOstic™ Nasal Swab Collection Device provides a method for clinical collection, stabilization, and transport of an anterior nasal sample collected in a clinical setting and provided to a designated laboratory for testing. Specimens collected using the RHINOstic™ Nasal Swab Collection Device are transported at ambient temperature for testing at a qualified testing laboratory. Testing is limited to laboratories certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA), 42 U.S.C. §263a, to perform high complexity tests.

SUMMARY AND EXPLANATION
SARS-CoV-2, also known as the COVID-19 virus, was first identified in Wuhan, Hubei Province, China December 2019. This virus, as with the novel coronavirus SARS-1 and MERS, is thought to have originated in bats, however the SARS-CoV-2 may have had an intermediary host such as pangolins, pigs or civets. The WHO declared that COVID-19 was a pandemic on March 11, 2020, and human infection has spread globally, with hundreds of thousands of confirmed infections and deaths. The pandemic has put unprecedented pressure on testing laboratories on a global basis, including supply chains and the ability to source collection devices as well as laboratory throughput. The Rhinostics collection devices bring easy to manufacture materials with properties that allow for dry shipment and sample concentration in addition to automation and rapid accessioning to allow for home collection as well as increase laboratories’ throughput and lower costs.

The RHINOstic™ Nasal Swab Collection Device is for the collection, stabilization and transport of an anterior nasal sample collected on a hydrophobic polymer nasal swab from patients. The RHINOstic™ Nasal Swab Collection Device consists of an anterior nasal swab with a threaded lid attached for transport in a 1 ml storage tube. Once the sample is collected, the swab is placed into the storage tube, so the threaded lid aligns with the threading on the open portion of the storage tube. The thread swab cap is screwed shut onto the tube. The tube is placed into a specimen bag for transport to the testing laboratory. Testing will be performed in laboratories certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA), 42 U.S.C. §263a, to perform high or moderate complexity tests, or by similarly qualified non-U.S. laboratories. When the collection device is delivered to an approved testing site, the technician reconstitutes the sample in accordance with instructions provided.

MATERIALS PROVIDED
The RH100001 RHINOstic™ Nasal Swab Collection Device includes:
1 sterile polypropylene nasal swab
1 capped and unlabeled collection tube

The LV-2D00001/LV-2D1D001/LV-2D1D002 RHINOstic™ Nasal Swab Collection Devices include
1 sterile polypropylene nasal swab
1 uncapped barcoded collection tube

MATERIALS NEEDED BUT NOT PROVIDED
Specimen bag for collected sample
No reconstitution buffers or assay materials are included with the RHINOstic™ Nasal Swab Collection Device
PRECAUTIONS
All clinical specimens should be considered biohazards and handled with care. Wear appropriate personal protective equipment and follow laboratory and biosafety guidelines when handling clinical specimens.

Do not use the collection device if the sterile package containing the swab is damaged or not sealed completely. Do not use if the swab is visibly damaged.

Do not use the device beyond the expiration date printed on the label.

This product is for single use only; reuse may cause a risk of infection and inaccurate results.

Gently collect the nasal sample following the RHINOstic™ Nasal Swab Collection Device collection instructions without using excessive force in the nostril. Refer to the Centers for Disease Controls and Prevention’s instructions on how to collect an anterior nares sample.

The swab should be screwed tightly into the collection tube to ensure it stays in the tube during transport to the laboratory for testing.

Dispose of the used collection device materials according to biohazard disposal regulations.

COLLECTION DEVICE STORAGE
For optimum performance, store at 2-25 °C. Avoid freezing and excessive heat.

SAMPLE TRANSPORT AND STORAGE
Samples should be transported at ambient temperature and tested within 72 hours. If not tested within 72 hours, the samples can be frozen at -70°C until able to test.

SPECIMEN COLLECTION
The swab portion for sample collection should be removed from the polybag without touching the collection end of the swab.

The transport tube should be removed from the package so that after collection it is ready for the sample to be placed into the transport tube.

The swab portion of the collection device should be held by the cap to collect the sample as follows:

*The tip of the swab is inserted into one (1) nostril until pressure is felt in the nose. The swab should be placed just inside the nostril.*
  a. The swab is rotated around the inside of the nostril three (3) times being sure you are making firm contact with the inside of the nose.
  b. The swab should be gently slid up and down against the inside of the nose one time.
  c. The swab should be held against the inside of the nostril for ten (10) seconds.

*Repeat collection steps (a-d) in the second nostril using the same swab.*

The cap should be removed from the tube and discarded. The cap should be replaced with the swab into the tube portion of the collection device.

The swab cap should be carefully screwed into the tube, so that it is completely closed. Completely screwing the swab is critical to ensuring that the swab does not come out during transport.
The collection tube should be placed into the specimen bag and shipping envelope provided and sent to the laboratory for testing. The testing laboratory will provide you with instructions on how to receive the test result.

SAMPLE PREPARATION

Nasal specimens should be collected according to the RHINOstic™ Nasal Swab Collection Device instructions using the RHINOstic™ Nasal Swab Collection Device. The polypropylene swab is used to collect an anterior nares sample which is placed dry into the transport tube and shipped at ambient temperature to the laboratory for testing within 72 hours. If samples cannot be tested within 72 hours of collection, they should be frozen in the laboratory at -70°C or colder.

When the sample arrives in the laboratory, a properly collected swab specimen should have a single swab attached to the threaded cap inserted into the tube. Incoming specimen sample tubes with no swab or with two swabs have not been collected according to the instructions in their respective collection kit Instructions for Use and should not be tested. Some samples may contain excessive mucus which may result in pipetting errors or result in errors when tested on diagnostic instruments. If the sample appears to have significant mucus, vortex the sample for 30 seconds.

The dry swab should be reconstituted by using a pipette to place 350 ul of 0.9% physiological saline into the collection tube after carefully opening and lifting the cap with the swab attached. The cap with the attached swab should be placed back into the tube containing the saline and should be mixed by flicking the sample by hand several times or gently and quickly vortex or through automated spin of the swab. From the 350 ul sample volume, the appropriate amount necessary for the assay protocol should be transferred into the reaction tube or well that will be used for the assay following the manufacturer’s
• Unscrew the primary sample tube cap with swab attachment.
• Lift the cap/swab and insert pipette into the sample tube.
• Transfer the appropriate amount of the sample into the prepared secondary tube or well and dilute (if necessary) with physiological saline of 0.9% for the final volume required in the manufacturer’s instructions for use.
• Ensure complete mixing after addition of sample to the secondary tube or well and test in accordance with manufacturer’s instructions for use.
• Close the primary sample tube cap with swab attachment and store sample at -70 C.

LIMITATIONS
Reliable specimen collection and transport depends on many factors, including collection and handling techniques, specimen condition and volume, and timing. Best results are achieved when specimens are processed within 72 hours after the time of collection.

The RHINOstic™ Nasal Swab Collection Device is designed for dry shipment. Shipment in viral transport media (VTM) may dilute sample, cause variable results, and potentially leak during shipment.

Refer to the corresponding reference standard and procedures for optimum collection techniques.

Use of the RHINOstic™ Nasal Swab Collection Device must be validated prior to use by the laboratory that is certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA), 42 U.S.C. §263a, to perform high complexity tests.

The use of tubes from any other source are not qualified for use with the RHINOstic™ Nasal Swab Collection Device and could affect the performance of the product and laboratory test results.

QUALITY CONTROL
Each lot of the RHINOstic™ Nasal Swab is sterilized with gamma irradiation. A representative sample of each lot is evaluated for bioburden and all testing procedures are established per the definitions in Clinical Laboratory Standards Institute M40-A2.

PATENTS AND TRADEMARKS
RHINOstic™ and Rhinostics™
Patent applications 63/051,263 and 63/019,620

SYMBOLS

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