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Primarius Pathology

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| Laboratory Use Only | | | - | | | | |
| Accession Number | | Date Received | | | Time Received | | |
| Practice Name | | Practice ID_ | | | Practice Contact In | formation | |
| Ordering Physicians | | | | | Address | | |
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| | (Currently | | | rance Informa | ation Plans - Contract Pendin | a) | |
| First Name | | | | - | Middle Initia | | nder* |
| Address Line 1 | | | | | | | |
| DOB | | Cell/Home | | | - | | · · |
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| Insured's Name | | | | | | | |
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| | | Re | espirator | y Test Panels | | | |
| Nasal Swab OR Nasopharyngeal Swab: Genesis Respiratory Pathogen Panel (Including COVID-19) (UTM Swab) (See back for details.) COVID-19 + Influenza A & B, and RSV Panel (UTM Swab) COVID-19 ONLY (SARS-CoV-2 PCR Assay) (UTM Swab) Genesis Expanded Respiratory Pathogen Panel** (UTM Swab) (See back for details.) **Please note that an immunodeficiency diagnosis code must be selected in order for this test to be performed. If an appropriate immunodeficiency code is not selected, testing will automatically be reflexed to the "Genesis Respiratory Pathogen Panel". | | | | | | | Streptococcus mies/eSwab) |
| ICD-10 Codes | | | | | | | |
| COVID-19 CODES ARE LISTED | BELOW AND M | IUST BE CHECKED | OFF | | Immunodeficiency I | CD-10 Codes | |
| J06.9 Acute upper respiratory infection, unspecified J11.1 Flu like symptoms R05 Cough R06.02 Shortness of breath R06.03 Acute respiratory distr | Pess | R50.9 Fever unspecified U07.1 COVID-19 virus identified D81.9 Combined immunodeficiency, unspecified D81.9 Combined immunodeficiency, unspecified D83.8 Other common variable immunodeficiencies D83.9 Common variable immunodeficiency, unspecified D84.89 Other immunodeficiencies | | | | | unspecified nodeficiencies iciency, unspecified |
| **For Genesis Expanded Respiratory Pathogen Panel, there must be <u>at least one</u> of the immunodeficiency codes in addition to the primary diagnosis code. | | | | | | | |
| This test is medically necessary for impairment, symptom, syndrome of medical management and treatme provider is authorized by law to order | r disorder. The resunt decisions. The p | ults will determine my erson listed as the ord | patient's | | zation: I hereby authorize G n to my insurance company y behalf. | | |
| | | | | Signature of Pa | tient (REQUIRED) | | Date |
| Signature of Physician or Other Au | thorized NPI Prov | ider (REQUIRED) | Date | | Accessioner l | Initials 1 | 2 |

Genesis Respiratory Pathogen Panel

Viruses

Adenovirus
COVID-19 (SARS-CoV-2)
Human Metapneumovirus
Influenza A virus/Influenza B virus Pan

Respiratory syncytial virus

Bacteria

Legionella pneumophila

Bordetella pan (bronchiseptica, parapertussis, pertussis) Bordetella pertussis Chlamydia pneumoniae Mycoplasma pneumoniae

| COVID-19 + Influenza A & B, and RSV Panel | COVID-19 ONLY | Group A Streptococcus |
|---|-----------------------|------------------------|
| COVID-19 (SARS-CoV-2) Influenza A | COVID-19 (SARS-CoV-2) | Streptococcus pyogenes |
| Influenza B | | |
| Respiratory Syncytial Virus (RSV) | | |

Genesis Expanded Respiratory Pathogen Panel

Viruses

Adenovirus Influenza A virus A/H3 Coronavirus 229E Influenza A virus A/H1-2009 Influenza B virus Coronavirus HKU1 Coronavirus NL63 Parainfluenza virus 1 Parainfluenza virus 2 Coronavirus OC43 COVID-19 (SARS-CoV-2) Parainfluenza virus 3 Parainfluenza virus 4 Human Metapneumovirus Human Rhinovirus/Enterovirus Respiratory syncytial virus

Bacteria

Bordetella pan (bronchiseptica, parapertussis, pertussis) Bordetella pertussis Chlamydia pneumoniae Mycoplasma pneumoniae

Influenza A virus
Influenza A virus A/H1

**For Genesis Expanded Respiratory Pathogen Panel, there must be <u>at least one</u> of the immunodeficiency codes in addition to the primary diagnosis code.

COVID-19

Information about COVID-19 (SARS-CoV-2 PCR Assay)

PLEASE NOTE: Testing was performed using the TaqPath COVID-19 Combo Kit and/or Panther Fusion® SARS-CoV-2 Assay Kit by real-time (RT) PCR. Positive results do not rule out co-infection with organisms not included in the TaqPath COVID-19 Combo Kit and/or Panther Fusion® SARS-CoV-2 Assay Kit by real-time (RT) PCR. The agent detected may not be the definite cause of this disease. This assay is for in vitro diagnostic use under the FDA Emergency Use Authorization (EUA). This test is only authorized for the duration of time of the declaration and for the detection of SARSCoV-2 RNA virus and/or diagnosis of COVID-19 infection under section 564 (b)(1) of the act, 21 U.S.C.360bbb-3(b)(1) unless the authorization is terminated or revoked sooner.

A Positive result indicates the presence of SARS-CoV-2 RNA but does not exclude bacterial infection and/or coinfection with other viruses. The agent detected may not be the definitive cause of disease. This result should be combined with clinical observation, patient history, and epidemiological information for patient management decisions.

Negative results do not preclude SARS-CoV-2 infection and should not be used as the sole basis for treatment and other patient management decisions. Negative results must be used in conjunction with clinical observation, patient history, and epidemiological information for patient management decisions.