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Primarius Pathology

	Respirator	y Requisition					
Date Specimen Collected:	Time Specimen Collected:						
Laboratory Use Only							
Accession Number	Date Received		Time Received				
Practice Name	Practice ID		ractice Contact				
Ordering Physicians			Address				
			City, State, Zip Phone				
		PI					
Fax Patient and Insurance Information (Currently Not Accepting Medicaid or Managed Medicaid Plans - Contract Pending)							
First Name					Gende	r*	
	Address Line 2						
	Cell/Home Phone*		-	il		-	
		Ethnicity* Sexual Orientat					
			Social Security #				
Home Phone							
Primary Insurance		Secondary Insurance					
Group # ID#		Group #	ID#				
Address		Address					
City State	Zip	City	State _	Zip			
	Respirator	y Test Panels					
Genesis Respiratory Pathogen) (UTM Swab) (See b	back for details.)	Group A (Liquid A				
ICD-10 Codes							
R05.1 Acute cough < 3 weeks	J02.9 Acute pharyngitis	, unspecified	Immunodefic	iency ICD-10 Co	des		
R05.2 Subacute cough > 3 weeks	J04.0 Acute laryngitis	D84.821 Immunodeficiency due to drugs					
R06.02 Shortness of breath	 J06.9 Acute upper respiratory infection, 		D81.9 Combined immunodeficiency,				
R06.03 Acute respiratory distress	unspecified		unspecified				
R06.2 Wheezing	J12.89 Other viral pneumonia		D83.8 Other common variable				
R07.1 Chest pain on breathing	J18.8 Other pneumonia, unspecified		immunodeficiencies				
R43.9 Unspecified disturbances of	•	D83.9 Common variable immunodeficiency,					
smell and taste		J20.8 Acute bronchitis due to other specified unspecifie					
R50.9 Fever unspecified				Other immunodeficiencies			
R53.1 Weakness	□ J22 Unspecified acute lower respiratory □D84.9 Immunode			munodeficien	cy, unspe	ecified	
R68.83 Chills (without fever)	infection	D81.89 Other combined immunodeficiencies					
J00 Acute nasopharyngitis		Z20.828 Exposure to other viral					
(common cold)	communicable disea						
This test is medically necessary for the diagnosis impairment, symptom, syndrome or disorder. The medical management and treatment decisions. T provider is authorized by law to order the test(s) r	or detection of a disease, illness, results will determine my patient's he person listed as the ordering	Patient Authorizatio to submit a claim to necessary on my beh Signature of Patient	my insurance comp nalf.		ices and app		
Signature of Physician or Other Authorized NPI	Provider (PEQUIPED) Data		Accession	ner Initials 1	2		

*Gender, Cell/Home Phone, Gender Identity, Race, Ethnicity, and Sexual Orientation are required by certain states and the CDC. ICD-10 Codes are listed for information purposes only. It is the provider's responsibility to order tests that are medically necessary and in the best interest of the patient. For specimen pick up please call our Courier Line at 732-508-9154.

Viruses

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

Bacteria

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

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

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.