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RESPIRATORY PANEL AND SARS-COVID-19 TEST REQUISITION FORM

PATIENT INFORMATION			PROVIDER INFORMATION		
Last Name Fi	rst Name	MI	Facility/Group	Referring Physician	
Gender: □ M □ F □ M/F □ F	/M Date of Birth/	_/	NPI Provider Nr:		
Patient Address			Physician Address		
City, State, ZIP code Contac	t Information (E-mail & F	Phone)	City, State, ZIP code		
Occupation/Exposure setting	g:				
Pregnancy Status: □ Yes □ No □ N/A			Diagnostic Codes (ICD-10 codes*** see Page 3)		
			Clinical Information** (date of onset/exposure, travel		
Race: ☐ Amer Ind/Alaskan ☐ White ☐ Black/Afr Amer ☐ Asian ☐ Native Hawaiian/Pacific Islander ☐ Other			history, previous lab results – attach additional info.)		
Ethnicity: □ Hispanic/Latino	□ Non-Hispanic/Non-Latir	no □ Otl	her		
Billing information: ☐ Self-P	ay (see Page 2) ☐ Com	mercial	Insurance (attach copy)	☐ Medicare (attach copy)	
RESPIRATORY PANEL and/ Influenza A/B Respiratory Human Metapneumovirus (Previous Coronavirus strain Current Coronavirus "SA	y Syncytial Virus (RSV) A/hMPV) □ Parainfluenza (Ins (HKU1, NL63, 229E, OCRS-CoV-2": ORF1ab (Rdl	/B □ Rh PIV1-4) C43) □ Rp/nsp1 LE HANI □ The □ Clir	inovirus/Enterovirus □ I □ Adenovirus □ <i>Mycop Chlamydophila pneumo</i> 0), E gene and N gene DLING following <i>MUST</i> be complete nical Information provided.	olasma pneumoniae (*) oniae (*) d (check all that apply):	
Collected by:			□ Nasal, Oro-/Nasopharyngeal swab placed in transport medium and in biohazard bag (labeled with patient information – First/Last Name, DOB). If submitting to lab within 24 hours place swab back into sleeve and then in biohazard bag.		
INFORMED CONSENT		PF	ROVIDER INFORMATION		
I consent to the collection of spetesting, and certify that the tests or by an authorized health care provordered by a qualified provider will be stored indefinitely and used for interidentifiers have been removed. Insurance provider and to receive ordered by my physician. I furthe physician to release to my insinformation necessary to process the may be an out-of-network facility of the store of the stor	dered have been explained to n ider. I understand that only test be performed. This sample may be rnal test validation after person I also authorize lab to bill n payment of benefits for the test rauthorize lab and the ordering surance provider any medic his claim. I acknowledge that lavith my insurance provider.	I attended based complete for the comple	st that the requested testing is d on the patient's diagnosis a leted the diagnosis codes ab is patient.	s medically necessary and appropriate and treatment plan. I have personally ove to indicate the accurate diagnosis	
Signature of Patient or Legal Guardian: If Guardian, Print Name:		Autho	orized Provider Signature:		
		Date:			

BILLING INFORMATION					
☐ Insurance Bill ☐ Accour	nt Bill 🗌 Patient Bill 🔲 Pre-Pay (Pa	yment Information must be completed)			
		age Determinations for further information concerning rogram requirements (ICD10-codes required) or the cla			
Bill Ordering Institution:		Bill Insurance:			
		(Provide legible photocopy of front & back of insurance card)			
Name of Insured:	Relation to Patient:	Insurance Company:	Membe		
Social Security #:	Member Group #:	Insurance Address:	Member Policy #		
Insurance Phone:					
PAYMENT INFORMATION	N (PRE-PAY)				
Check Card Used for Payme	ent:	□ VISA □ MasterCard □ American Expr	ress Discover		
Card Number:		Card Security Code:			
Signature:		Exp. Date:			

(*) Designates Bacterial targets

(**) Priority COVID-19 Testing Groups if mild/moderate symptoms observed (Fever, Cough, etc.):

- Evidence of lower respiratory disease without alternative diagnosis, especially if hospitalized
- Any resident of a senior living facility, including skilled nursing facilities or assisted living facilities
- Persons who care for the elderly
- Persons living in congregate setting (homeless shelters, etc.)
- Health care workers, first responders, and other emergency workers

Test information:

NxTAG® Respiratory Pathogen Panel (RPP) for In Vitro Diagnostic Use (IVD), incorporates multiplex Reverse Transcriptase Polymerase Chain Reaction (RT-PCR) with the Luminex® proprietary universal tag sorting system on the Luminex platform to easily detect respiratory pathogen targets. Extracted total nucleic acid is added to pre-plated, Lyophilized Bead Reagents (LBRs), and mixed to resuspend the reaction reagents. The reaction is amplified via RT-PCR and the reaction product undergoes near simultaneous bead hybridization within the sealed reaction well. The hybridized, tagged beads are then sorted and read on the MAGPIX® instrument, and the signals are analyzed using the NxTAG Respiratory Pathogen Panel Assay File for SYNCT™ Software, providing a reliable, qualitative call for each of the 20 targets and internal controls within each reaction well.

<u>SARS-CoV-2 (COVID-19):</u> The NxTAG® CoV Extended Panel (NxTAG CoV) is an EUA-approved (In Vitro Diagnostic Use Under Emergency Use Authorization) test authorized by the FDA under an EUA for use by authorized laboratories. This test has been authorized only for the detection of nucleic acid from SARS-CoV-2, not for any other viruses or pathogens. incorporates multiplex Reverse Transcriptase Polymerase Chain Reaction (RT-PCR) with the Luminex® proprietary universal tag sorting system on the Luminex platform to easily detect SARS-CoV-2s. Extracted total nucleic acid is added to pre-plated, Lyophilized Bead Reagents (LBRs) and mixed to resuspend the reaction reagents. The reaction is amplified via RT-PCR and the reaction product undergoes near simultaneous bead hybridization within the sealed reaction well. The hybridized, tagged beads are then sorted and read on the MAGPIX® instrument. Sensitivity: The Limit of Detection (LoD) study established the lowest SARS-CoV-2 viral concentration (Genomic Copy Equivalents or GCE = 5.00E+03 GCE/mL) that can be detected by the NxTAG® CoV Extended Panel Assay at least 95% of the time using viral genomic RNA. 100% at 3X LoD and 5X LoD. Specificity: 100% PPV and NPV.

*** ICD10 Codes:

SARS-CoV-2:

U07.1 COVID-19

Z20.828 Contact with and (suspected) exposure to other viral communicable diseases

General Respiratory Virus:

B30.2 Viral pharyngoconjunctivitis

B34.0 Adenovirus infection, unspecified

B34.2 Coronavirus infection, unspecified

B97.0 Adenovirus as the cause of diseases classified elsewhere

B97.21 SARS-associated coronavirus. cause of diseases classified elsewhere

B97.29 Other coronavirus as the cause of diseases classified elsewhere

B97.4 Respiratory syncytial virus as the cause of diseases classified elsewhere

B97.81 Human metapneumovirus as the cause of diseases classified elsewhere

B97.89 Other viral agents as the cause of diseases classified elsewhere

J00 Acute nasopharyngitis [common cold]

J05.0 Acute obstructive laryngitis [croup]

J06.9 Acute upper respiratory infection, unspecified

J09.X1 Influenza due to identified novel influenza A virus with pneumonia

J09.X2 Influenza due to identified novel influenza A virus with other respiratory manifestations

J09.X3 Influenza due to identified novel influenza A virus with gastrointestinal manifestations

J09.X9 Influenza due to identified novel influenza A virus with other manifestations

J10.00 Influenza due to other identified influenza virus with unspecified type of pneumonia

J10.01 Influenza due to other identified influenza virus with the same other identified influenza virus pneumonia

J10.08 Influenza due to other identified influenza virus with other specified pneumonia

J10.1 Influenza due to other identified influenza virus with other respiratory manifestations

J10.2 Influenza due to other identified influenza virus with gastrointestinal manifestations

J10.81 Influenza due to other identified influenza virus with encephalopathy

 $\ensuremath{\mathsf{J10.82}}$ Influenza due to other identified influenza virus with myocarditis

J10.83 Influenza due to other identified influenza virus with otitis media

J10.89 Influenza due to other identified influenza virus with other manifestations

J11.00 Influenza due to unidentified influenza virus with unspecified type of pneumonia

J11.08 Influenza due to unidentified influenza virus with specified pneumonia

J11.1 Influenza due to unidentified influenza virus with other respiratory manifestations

J11.2 Influenza due to unidentified influenza virus with gastrointestinal manifestations

J11.81 Influenza due to unidentified influenza virus with encephalopathy

J11.82 Influenza due to unidentified influenza virus with myocarditis

J11.83 Influenza due to unidentified influenza virus with otitis media

J11.89 Influenza due to unidentified influenza virus with other manifestations

J12.0 Adenoviral pneumonia

J12.1 Respiratory syncytial virus pneumonia

J12.2 Parainfluenza virus pneumonia

J12.3 Human metapneumovirus pneumonia

J12.81 Pneumonia due to SARS-associated coronavirus

J12.9 Viral pneumonia, unspecified

J20.4 Acute bronchitis due to parainfluenza virus

J20.5 Acute bronchitis due to respiratory syncytial virus

J20.6 Acute bronchitis due to rhinovirus

J21.0 Acute bronchiolitis due to respiratory syncytial virus

J21.9 Acute bronchiolitis, unspecified

J22 - Unspecified acute lower respiratory infection

Z11.59 Encounter for screening for other viral diseases

Pneumonia:

A49.3 Mycoplasma infection, unspecified site

B96.0 Mycoplasma pneumoniae [M. pneumoniae] as the cause of diseases classified elsewhere

J15.7 Pneumonia due to Mycoplasma pneumoniae

J20.0 Acute bronchitis due to Mycoplasma pneumoniae

Z11.2 Encounter for screening for other bacterial diseases