

PATIENT SIGNATURE:

Employe	er / Grou	p Name:	Location:	Emplo	vee 🗆 Fa	amily	/ E

COVID-19 IgG Antibody & RT-PCR Test Request Form

Laboratory Personnel - F	OR OFFICE US	E ONLY					
Today's Date:		Location Name:					
Clinician Name:		Phone:					
Patient Information: COI	MPLETED BY P	ATIENT					
First Name:		Last Name:			Phone:		
Address:							
City:		Zip Code:		County:			
State:							
Date of Birth:		Age:		S	ex: 🗆 Male	□ Female	
Email:							
Additional Information requ	ired for testing:						
Does the patient live or wor			erm care faci	lity, shelter,	group home, pris	on, jail)	
□ YES □ NO		Facility Name:		<u>, . , , , , , , , , , , , , , , , , , ,</u>		- ,	
		Employee Occupation	ո:				
Does the patient receive dialysi	s? TES	□ NO					
CLINICAL INFORMATION:	COMPLETED	BY PATIENT					
Date of symptom onset:	□ None	Does the pat	tient have any	underlying conditi	ons?		
Symptoms Observed:	□ None			Immunocompromised			
□ Fever	se 🗆 🗆 Unknown			□ Pregnant			
□ Tiredness	Loss of sm	nell 🗆 Diabete:			Chroni	c Lung Disease	
□ Dry Cough	Diarrhea	☐ Hypertensi		sion	Chroni	c Liver Disease	
□ Body Ache	Loss of Ap	petite 🗆 Cardiac D		isease	□ Chroni	c Kidney Disease	
□ Nasal Congestion			□ Other				
LABORATORY TESTING -	COMPLETED E	BY PATIENT					
Has the patient been tested for	influenza?		□ YES	□ NO			
Result: Positive	□ Negative						
Test Type: 🗆 Rapid	□ PCR						
Has the patient been tested for	any other viral re	espiratory illness?	□ YES	□ NO			
Result:							
COVID 19 TESTING – CON	IPLETED BY PA	ATIENT					
Has the patient been tested for	COVID-19?		□ YES	□ NO			
Result: Positive	□ Negative						
Test Type: 🗆 Rapid	□ PCR						
I here	by acknowleds	ge full and complete	consent for t	testing and	make request for	:	
□ RT-PCR T	est and/or 🗆	SARS-Cov2 IgG	Antibody	Test (CH	ECK ONE O	R BOTH)	
						- ,	

release The PMH Laboratory, Inc. its principals, directors, members, employees, affiliates, suppliers, providers, subcontractors, successors, agents, their respective insurance carriers, and the location sponsoring this clinic/program, its principals, directors, employees, affiliates, successors, or agents from any and all liability, injury or damage whatsoever arising from, or in any way connected with, this SARS-CoV-2 IgG Antibody Test or the administration of same including, but not limited to, acts of negligence. I authorize my medical information herein, including tests results, to be shared with my physician/insurance/employer. The PMH Laboratory, Inc., will use and disclose your personal and health information to treat you, to receive payment for the care we provide, to public health agencies as required, and for our other health care operations which generally include those activities we perform to improve quality care. We have prepared a detailed NOTICE OF PRIVACY AND CONFIDENTIALITY PRACTICES to help you better understand our policies in regard to your personal health information. I acknowledge that I have received a copy of the Notice of Privacy and Confidentiality Practices. I agree to remain in the general area for at least 5 minutes after collection of samples. Please provide a copy of this form to your physician and/or healthcare provider for your medical records. This test is for informational purposes only and to be discussed with your health care professional. The PMH Laboratory, Inc., is not providing you with medical advice nor are they responsible for any outcome in your care or treatment. Please keep in mind that a positive result does not mean you are immune or cannot become re-infected. This test was developed, and its performance characteristics determined by PMH Laboratory, Inc. This test has not been FDA cleared or approved. This test has been authorized by FDA under an Emergency Use Authorization (EUA). This test has been validated in accordance with the FDA's Guidance Document (Policy for Diagnostics Testing in Laboratories Certified to Perform High Complexity Testing under CLIA prior to Emergency Use Authorization for Coronavirus Disease-2019 during the Public Health Emergency) issued on April 20, 2020. FDA independent review of this validation is pending. This test is only authorized for the duration of time the declaration that circumstances exist justifying the authorization of the emergency use of in vitro diagnostic tests for detection of SARS-CoV-2 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.

DATE: