

COVID-19 Immunization Screening and Consent Form

for Moderately to Severely Immunocompromised People

Updated: April 1, 2022

Reci	pient Name (please print)	Preferred Name					
DOE	Current Gender ID Key: W – Woman/Girl TW – Transgender Woman/Girl M – Man/Boy TM – Transgender Man/Boy NB – Non-Binary Person GNC – Gender Non-Conforming Q – Not Sure/Questioning NR – Chose not to Respond GNL - Gender not Listed (write-in) * Gender Pronouns: write-in by client's name						
	Assigned at Birth Key: cate Sex Below:	Marital Status Indicate Status Below: S – Single W – Widowed V – Civil Union U – Unknown					
Pare	ent/Guardian/ Surrogate (if applicable, please print)	Phone	Preferred L	anguage			
	cate Ethnicity Below: DECL — Declined HIS — Hispanic Origin NHL — Non-Hispanic Origin UNK - Unknown	DECL – D	ative American or Alaskan ASN – Asian African American or Black Declined Native Hawaiian or Pacific Islander				
Prim	nary Insurance Name	Primary Insurance ID#	Subscriber Name/DOB Subscrib to Patier			criber Relation atient	
Prim	nary Insurance Address	Primary Insurance Group #	Primary Ins	isurance Phone #			
Seco	ondary Insurance Name	Secondary Insurance ID#	r Name/DOB Subscriber Relation to Patient				
Seco	ondary Insurance Address	Secondary Insurance Group #	/ Insurance Phone #				
Clinic/Office Site Where Vaccine is Administered Primary Care Physician Address/Phone Number							
	Scree	ning Questionnaire					
1.	Will you be under the age of 5 years old for the Pfizer vaccine, or under 18 years old for the Moderna vaccine, on the day of your appointment?					No	
2.	Are you feeling sick today?					No	
3.	In the last 10 days, have you had a COVID-19 test because you had symptoms and are still awaiting your test results or been told by a health care provider or health department to isolate orquarantine at home due to COVID-19 infection or exposure?					No	□ Unknown
4.	Have you been treated with antibody therapy or convalescent plasma for COVID-19 in the past 90 days (3 months)? <i>If yes, when did you receive the last dose?</i> Date:					No	□ Unknown
5.	Have you ever had an immediate allergic reaction (e.g., hives, facial swelling, difficulty breathing, anaphylaxis) to any vaccine, injection, or shot or to any component of the COVID-19 vaccine, or a severe allergic reaction (anaphylaxis) to anything?					No	□ Unknown
6.	Are you pregnant or considering becoming pregnant?					No	□ Unknown

7.	Are you moderately or severely immunocompromised due to one or more of the medical conditions or receipt of immunosuppressive medications or treatments listed below?		Yes		No	□ Unknown
	1) Active treatment for solid tumor and hematologic malignancies, 2) Receipt of solid-organ transplant and taking immunosuppressive therapy, 3) Receipt of CAR-T-cell or hematopoietic stem cell transplant (within 2 years of transplantation or taking immunosuppression therapy), 4) Moderate or severe primary immunodeficiency (e.g., DiGeorge syndrome, Wiskott-Aldrich syndrome), 5) Advanced or untreated HIV infection, 6) Active treatment with high-dose corticosteroids (i.e., 8805;20mg prednisone or equivalent per day), alkylating agents, antimetabolites, transplant-related					
	immunosuppressive drugs, cancer chemotherapeutic agents classified as severely immunosuppressive, tumor-necrosis (TNF) blockers, and other biologic agents that are immunosuppressive or immunomodulatory					
8.	Do you have a bleeding disorder, a history of blood clots or are you taking a blood thinner?		Yes		No	□ Unknown
9.	Do you have a history of myocarditis (inflammation of the heart muscle) or pericarditis (inflammation of the lining around the heart)?		Yes		No	□ Unknown
10.	Have you had Guillain-Barre Syndrome after receipt of the Janssen vaccine?		Yes		No	□ Unknown
11.	Do you have a history of MIS-C or MIS-A (multisystem inflammatory syndrome in children or multisystem inflammatory syndrome in adults)?		Yes		No	□ Unknown
12.	Have you received 2 previous doses of the Pfizer or Moderna COVID-19 vaccine, and was your last dose at least 28 days ago?		Yes		No	Date:
						(if applicable)
13.	Have you received a previous dose of the Janssen (Johnson & Johnson) COVID-19 vaccine at least 28 days ago?		Yes		No	
14*	Are you 12 years old or older and have received 3 doses of the Pfizer or Moderna COVID-19 vaccine, and was your last dose at least 3 months ago?		Yes		No	Date:
						(if applicable)
15*	Have you received 2 doses of a Janssen (Johnson & Johnson) COVID-19 vaccine, or one dose of Janssen (Johnson & Johnson) followed by an mRNA vaccine (Pfizer or Moderna), and was your last		Yes		No	Date:
	dose at least 2 months ago?					(if applicable)
16**	Are you 12 years old or older and have received 4 doses of the Pfizer or Moderna COVID-19 vaccine, and was your last dose at least 4 months ago?		Yes		No	Date:
						(if applicable)
17**	Have you received any combination of Janssen (Johnson & Johnson) COVID-19 vaccine and mRNA vaccine (Pfizer or Moderna) totaling 3 doses, and was the last dose at least 4 months ago?				No	Date:
						If applicable)
18.	Have you received a previous dose or doses of a non-FDA authorized or approved COVID-19 vaccine (AstraZeneca – VAXZEVRIA, Sinovac – CORONAVAC, Serum Institute of India –		Yes		No	Date(s):
	COVISHIELD, Sinopharm/BIBP, COVAXIN, Novavax – Covovax or Nuvaxovid)? ¹					(if applicable)

Emergency Use Authorization

The FDA has made the COVID-19 vaccine available under an emergency use authorization (EUA). The EUA is used when circumstances exist to justify the emergency use of drugs and biological products during an emergency, such as the COVID-19 pandemic. This vaccine has not undergone the same type of review as an FDA-approved or cleared product. However, the FDA's decision to make the vaccine available is based on the totality of scientific evidence available, showing that known and potential benefits of the vaccine outweigh the known and potential risks. Please note: FDA approved the Pfizer-BioNTech COVID-19 vaccine as a two-dose series in individuals 16 years of age and older. The vaccine continues to be available under an EUA for certain populations, including for those individuals 5 through 15 years of age and for the administration of a third dose in the populations set forth in the consent section below.

Emergency Use Instruction

Emergency Use Instructions (EUIs) are issued by the CDC to provide information about emergency use of FDA-approved medical products that may not be included in or differ in some way from the information provided in the FDA-approved labeling (package insert). The COVID-19 vaccine by Pfizer-BioNTech is an FDA-approved COVID-19 vaccine (brand name Comirnaty, mRNA) to prevent COVID-19 in persons 16 years of age and older. CDC is issuing EUI to provide information about use of this vaccine as an additional primary dose in certain immunocompromised persons (12 years of age and older) and a booster dose in certain adults (18 years of age and older) who received certain non-FDA authorized or approved COVID-19 vaccine (e.g., certain vaccines available outside of the United States or from clinical trial participation).

¹ As set forth in CDC's Emergency Use Instructions (EUI) "a non-FDA authorized or approved COVID-19 vaccine includes such vaccines "listed for emergency use by the World Health Organization, or is included in CDC's Technical Instructions for Implementing Presidential Proclamation Advancing Safe Resumption of Global Travel During the COVID-19 Pandemic and CDC's Order, or that is a non-placebo part of a clinical trial within or outside the United States that is a WHO-EUL COVID-19 vaccine or a vaccine that is not listed for emergency use by WHO but for which a U.S. data and safety monitoring board or equivalent has independently confirmed efficacy in the United States (hereinafter 'non-FDA authorized or approved COVID-19 vaccines')."

^{*}Questions 12 and 13 pertain to the first booster dose eligibility.

^{**}Questions 13 and 14 pertain to the second booster dose eligibility.

Consent

* Use of this form is optional.

I hereby certify under penalty of law that I am of an age and, if applicable, immunocompromised (e.g., moderate to severe immune compromise due to a medical condition or receipt of immunosuppressive medications or treatments) as authorized by an EUA or in accordance with an EUI, as applicable, to receive this vaccine, or, the person for whom I am legally authorized to make health care decisions is of an age and, if applicable, immunocompromised as authorized by an EUA or in accordance with an EUI, as applicable, to receive this vaccine. I have had a chance to ask questions which were answered to my satisfaction (and ensured the person named above for whom I am authorized to provide surrogate consent was also given a chance to ask questions). I understand the benefits and risks of the vaccination as described.

I request that the COVID-19 vaccination be given to me (or the person named above for whom I am authorized to make this request and provide surrogate consent). I understand there will be no cost to me for this vaccine. I understand that any monies or benefits for administering the vaccine will be assigned and transferred to the vaccinating provider, including benefits/monies from my health plan, Medicare or other third parties who are financially responsible for my medical care. I authorize release of all information needed (including but not limited to medical records, copies of claims and itemized bills) to verify payment and as needed for other public health purposes, including reporting to applicable vaccine registries.

Recipient/Surrogate/Guardian (Signature) recipient			Date / Time	Priı	nt Name		Relationship to Patient (if other than recipient)	
Telephonic Interpreter's ID # OR			Date / Time					
Signature: Interpreter			Date/ Time	Prir	nt: Interpreter	's Name and Rela	ationship to Patient	
Area Below to be Completed by Vaccinator								
Which vaccine is the patient receiving today?								
Vaccine Name			Administ	ration		EUA Fact Sheet Date	Manufacturer & Lot Number	
Pfizer/ BioNTech	□ First Dose	□ Second Dose	□ Third Dose	□ First Booster	□ SecondBooster			
Moderna	□ First Dose	□ Second Dose	□ Third Dose	□ First Booster	SecondBooster			
Janssen (Johnson & Johnson)	□ First Dose	□ Second Dose	□ First Booster	□ Second Booster				
Administration Site Left De			eltoid 🗆	Right Deltoid	□ Left	Thigh 🗆	Right Thigh	
Dosage		□ 0.5 m	l o	0.3 ml	□ 0.25	5 ml \Box	0.2 ml	
□ I have provided the patient (and/or parent, guardian or surrogate, as applicable) with information about the vaccine and consent to vaccination was obtained.								
Vaccinator Signature:								

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