

New York State Department of Health Bureau of Immunization

COVID-19 Immunization Screening and Consent Form*

Rec	ipient Name (please print)	Preferred Name					
DO	W – Womar Indicate ID Below: TM – Trans Q – Not Sur GNL - Gend	aan/Girl TW – Transgender Woman/Girl M – Man/Boy ansgender Man/Boy NB – Non-Binary Person GNC – Gender Non-Conforming aure/Questioning NR – Chose not to Respond nder not Listed (write-in) [.] Pronouns: write-in by client's name					
Indi	Assigned at Birth Key: cate Sex Below: M – Male F – Female I – Intersex NR – Chose not to Respond	Marital Status Key: Indicate Status Below: S – Single D – Divorced M – Married W – Widowed V – Civil Union U – Unknown					
Add	ress City	State Zip	Email Address	5			
Pare	ent/Guardian/ Surrogate (if applicable, please print)	Phone	Preferred Lan	guage			
Ethnicity Ethnicity Key: Race Race Key: Indicate Ethnicity Below: DECL – Declined Indicate Race Below: AIA – Native American HIS – Hispanic Origin NHL – Non-Hispanic Origin BAA – African American UNK – Unknown NHP – Native Hawaiia WHT – White				or Blac or Pacif	k ic Island	N – Asian Ier er or Multiracial	
Prin	nary Insurance Name	Primary Insurance ID#	Subscriber Na		B Sub	oscriber Relation Patient	
Prin	nary Insurance Address	Primary Insurance Group #	Primary Insur	ance Ph	ione #		
Sec	ondary Insurance Name	Secondary Insurance ID#	Subscriber Na	me/DO		oscriber Relation Patient	
Sec	ondary Insurance Address	Secondary Insurance Group #	Secondary Ins	surance	Phone #	ŧ	
Clin	ic/Office Site Where Vaccine is Administered	Primary Care Physician Addres	s/Phone Numbe	er			
	Scree	ening Questionnaire					
1.	Are you feeling sick today?			Yes	🗆 No		
2.	In the last 10 days, have you had a COVID-19 tes awaiting your test results or been told by a heal isolate or quarantine at home due to COVID-19 info	th care provider or health depa		Yes	□ No	🗆 Unknown	
days (3 months)? If yes, when did you receive the last dose? Date:						🗆 Unknown	
4.	Have you ever had an immediate allergic reaction (e anaphylaxis) to any vaccine, injection, or shot or to a severe allergic reaction (anaphylaxis) to anything?			Yes	□ No	🗆 Unknown	
5. Are you pregnant or considering becoming pregnant?				Yes	🗆 No	🗆 Unknown	

6.	Do you have cancer, leukemia, HIV/AIDS or any other condition that weakens the immune system?		Yes		No	Unknown
7.	Do you take any medications that affect your immune system, such as cortisone, prednisone or other steroids, anticancer drugs, or have you had any radiation treatments?		Yes		No	Unknown
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.*	Are you 12 years old or older, and have you received 2 doses of the Pfizer vaccine, the second dose being at least 5 months ago?		Yes		No	Date of 2 nd dose:
						(if applicable
13.*	Have you received 2 doses of the Moderna vaccine, the second dose being at least 5 months ago?		Yes		No	Date of 2 nd dose:
						(if applicable
14.*	Have you received a previous dose of the Janssen vaccine, at least 2 months ago?		Yes		No	Date of 1 st dose:
						(if applicable
15.*	If you had a previous dose of Janssen (Johnson & Johnson), did you develop thrombosis with thrombocytopenia syndrome (TTS)?		Yes		No	Unknown
16.**	Are you 50 years old or older, and have you received 3 doses of the Pfizer or Moderna vaccine, the third dose being at least 4 months ago?		Yes		No	Date of 3 rd dose:
						(if applicable)
17.**	Have you received 2 doses of the Janssen (Johnson & Johnson) vaccine, or 1 dose of the Janssen vaccine and 1 dose of mRNA vaccine, the second dose being at least 4 months ago?		Yes		No	Date of 2 nd dose (if applicable)
18. ¹	Have you received a previous dose of a non-FDA authorized or approved COVID-19 vaccine authorized by the WHO ¹ but not by the FDA (AstraZeneca – VAXZEVRIA, Sinovac – CORONAVAC, Serum Institute of India – COVISHIELD, Sinopharm/BIBP, COVAXIN, Novavax – Covovax, Nuvaxovid)?		Yes		No	Unknown

*Questions #10 - 14 pertain to the first booster dose eligibility.

**Questions #14 and 15 pertain to the second booster dose eligibility.

¹ As set forth in the <u>CDC's Emergency Use Instructions (EUI)</u>, a non-FDA authorized or approved COVID-19 vaccine such as those 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').

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).

Consent

I have read, or had explained to me, the information sheet about the COVID-19 vaccination. I understand that if my vaccine requires two doses, I will need to be administered (given) two doses to be considered fully vaccinated. Further, I understand that a booster dose of COVID-19 vaccine is recommended at least 2 months following the first dose of Janssen vaccine (if I am age 18 or older), or at least 5 months following the second dose of Pfizer-BioNTech (if I am age 5 or older) or Moderna COVID-19 vaccine (if I am age 18 or older), to increase my protection.

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)	Date / Time	Print Name	Relationship to Patient(if other than recipient)			
Telephonic Interpreter's ID # OR	Date / Time					
Signature: Interpreter	Date/ Time	Print: Interpreter's	Name and Relationship to Patient			

	A		e Completed by is the patient receiving		or	
Vaccine Name		Adminis	EUA Fact Sheet Date	Manufacturer & Lot #		
Pfizer/BioNTech	🗆 First Dose	Second Dose	 First Booster Dose 	 Second Booster Dose 		
Moderna	First Dose	Second Dose	 First Booster Dose 	□ Second Booster Dose		
Janssen	□ Single Dose	 First Booster Dose 	 Second Booster Dose 	N/A		
Administration Site Dosage	 Left Deltoid 0.5 ml 	□ Right Delt □ 0.3 ml	oid □ Left Thi □ 0.25 ml	gh □	Right Thigh	

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:

* Use of this form is optional.

Updated April 1, 2022