

COVID-19 VACCINE SCREENING AND CONSENT FORM

		Facility Name/Fac	-						
SECTION 1: INFORMATION ABO Name: Last:	JUI PAIIE	NT (PLEASE PRINT) First:)	Middle Initial:					
Date of Birth: Month:	Day:		Mobile Phone Numb	per (Patient or Guardian): ()				
Address:			Apt/Room #:						
City:			State:	ZIP:					
Name of Legal Guardian: La	ast:		First:	Middle Initial:					
Sex (Gender assigned at birth) Female Male	Race ☐ America ☐ Asian	an Indian or Alaska Native	☐ Native Hawaiian or Other☐ Pacific Islander☐ White		Ethnicity Hispanic or Latin Not Hispanic orl Unknown		0		
Primary Insurance Carrier	ID #:		Grp #:		_				
Insurance Company:			Insu	rance Company Phone #:					
Insured's Name:		R	elationship:	Insured's Date	e of Birth:		_		
Secondary Insurance Carri	er ID #:	_	Grp #:						
Insurance Company:			Insu	rance Company Phone #:			_		
Insured's Name:		R	elationship:	Insured's Date	e of Birth:				
Decimalism of OOVID 40			□ □ □ □ □ □ □ □	and Dane		_ +			
Designation of COVID-19 v	accination	dose number?	□ FIRST DOSE □ Sec	ond Dose ☐ Third Dose*	□Booster Dos	e"			
SECTION 2: COVID-19 SCREEN	ING QUES	TIONS							
Please check YES or NO for						Yes	No		
				ortness of breath, difficulty breathin y nose, nausea, vomiting or diarrhe					
2. Have you tested positive for ar							1		
3. Have you had a severe allergion	c reaction (fo	or example, needed ep	oinephrine or hospital care)	to a previous dose of this vaccine o	r to any of the				
ingredients of this vaccine?									
4. Have you had any COVID-19	antibody the	rapy within the last 90	days (for example, Regene	ron, COVID Convalescent Plasmae	etc.)	<u> </u>			
SECTION 3: IMMUNIZATION SCR	EENING GU	IDANCE FOR COVID	-19 VACCINE						
Please check YES or NO for each	ch question					Yes	No		
5. Do you carry an EpiPen for emergency treatment of anaphylaxis and/or have allergies or reactions to any medications, foods, vaccines or latex?									
6. For women, are you pregnant or is there a chance you could become pregnant?									
7. For women, are you currently b			-:			<u> </u>	-		
Are you immunocompromised of the second						<u> </u>	-		
10. Are you a female aged 18 to 4				OVID-19 vaccine?			-		
11. If you are under the age of 18							+		
12. Have you received a previous							1		
*13. If you meet one or more of the	ne following	;	•						
1) A third dose (d	or additional	dose if first dose was	Janssen [Johnson and Johr	nson]) for moderately to severely					
				osuppressant medications, active trea					
			-BIONTech COVID-19) or 19 OVID-19 primary series.	8 years of age (for Moderna vaccine	e) and at least 28				
				munocompromised) have passed s	ince the				
	completion of an mRNA COVID-19 vaccine primary series and you are 5 years of age or older (Pfizer-BioNTech COVID-19								
vaccine only)	or are 18 yea	ars of age or older (Mo	oderna COVID-19 vaccine).	•					
	For a booster dose of Janssen (Johnson and Johnson), at least 2 months have passed since the initial dose of your Janssen (Johnson and Johnson) COVID-19 vaccination, or at least 2 months after your additional dose if immunocompromised, and you								
(Jonnson and are 18 years o			or at least 2 months after yo	oui auditional dose il immunocompro	omiseu, and you				
			h COVID-19 vaccine may	oe administered at least 4 months a	fter the first booster	ſ			

Effective Date: 06/05/2022 DH8010-DCHP-08/2021 dose of any authorized or approved COVID-19 vaccine to individuals 50 years of age and older or those who are 12 years of age and older with certain immunocompromising conditions.

- B) A second booster dose of the **Moderna COVID-19 vaccine** may be administered at least 4 months after the first booster dose of any authorized or approved COVID-19 vaccine to individuals 50 years of age and older <u>or</u> those who are 18 years of age and older with the same certain kinds of immunocompromising conditions.
- I certify that I am: (a) the patient and at least 18 years of age; (b) the legal guardian of the patient and confirm that the patient is at least 5 years of age (for Pfizer vaccine consent only); or (c) legally authorized to consent for vaccination for the patient named above. Further, I hereby give my consent to the Florida Department of Health (DOH) or its agents to administer the COVID-19 vaccine.
- Pfizer BioNTech COVID-19 vaccine product has been fully approved and licensed by the U.S. Food and Drug Administration (FDA). This FDA approval and license is for use in individuals 16 years of age and older only. The Moderna COVID-19 vaccine product has also been fully approved and licensed by FDA. This FDA approval and license is for use in individuals 18 years of age and older only.
- I understand that this product (other than Pfizer and Moderna for usage in ages mentioned above only) has not been approved or licensed by FDA, but has been authorized for emergency use by FDA, under an Emergency Use Authorization (EUA) to prevent Coronavirus Disease 2019 (COVID-19) for use in individuals either 5–15 years of age (Pfizer only) or 18 years of age and older (Johnson and Johnson); and the emergency use of this product is only authorized for the duration of the declaration that circumstances exist justifying the authorization of emergency use of the medical product under Section 564(b)(1) of the Food, Drug, and Cosmetic Act unless the declaration is terminated or authorization revoked sooner.
- I understand that it is not possible to predict all possible side effects or complications associated with receiving vaccine(s). I understand the risks and benefits associated with the above vaccine and have received, read and/or had explained to me the Emergency Use Authorization Fact Sheet on the COVID-19 vaccine I have elected to receive. I also acknowledge that I have had a chance to ask questions and that such questions were answered to my satisfaction.
- I acknowledge that I have been advised to remain near the vaccination location for approximately 15 minutes (or more in specific cases) after administration for observation. If I experience a severe reaction, I will call 9-1-1 or go to the nearest hospital.
- On behalf of myself, my heirs and personal representatives, I hereby release and hold harmless the State of Florida, the Florida Department of Health (DOH),
 the Florida Division of Emergency Management (FDEM) and their staff, agents, successors, divisions, affiliates, subsidiaries, officers, directors, contractors
 and employees from any and all liabilities or claims whether known or unknown arising out of, in connection with, or in any way related to the administration of
 the vaccine listed above.
- I acknowledge that: (a) I understand the purposes/benefits of Florida SHOTS, Florida's immunization registry and (b) DOH will include my personal
 immunization information in Florida SHOTS and my personal immunization information will be shared with the Centers for Disease Control (CDC) or other
 federal agencies.
- I further authorize DOH, FDEM or its agents to submit a claim to my insurance provider or Medicare Part B without supplemental coverage payment for me for the above requested items and services. I assign and request payment of authorized benefits be made on my behalf to DOH, FDEM or its agents with respect to the above requested items and services. I understand that any payment for which I am financially responsible is due at the time of service or if DOH invoices me after the time of service, upon receipt of such invoice.
- I acknowledge receipt of the DOH Notice of Privacy Practices.

Signature of Patient or Authorized Representative:					Date:						
Print Name of Representative and Relationship to Person Receiving Vaccine:											
Site (LD/RD)	Route	Manufacturer (MVX)		Lot # Unit of Use/ Unit of Sale	Expiration Date	Date of EUA Fact Sheet					
	IM										
				I							
Administer name/ID	ed at l	ocation: Facility									
Administer	ed at l	ocation: Type									
Administra	tion Ac	Idress:									
CVX (prod	uct)										
Sending or	ganiza	tion:									
Vaccinator Prin	t Name:			Signature:		Date:					
Vaccine Administering Provider Suffix:											

Effective Date: 06/05/2022 DH8010-DCHP-08/2021