

for Use of COVID-19 Vaccines Currently Authorized or Approved in the United States

COVID-19 vaccine products currently approved or authorized in the United States

Pfizer-BioNTech Primary series Booster doses Vaccine Vaccine vial Label border Dilution Age indication composition color required cap color Dose **Injection volume** Dose **Injection volume** Monovalent (Use for A booster dose is not authorized 6 months-4 years Doses 1 and 2: 3 µg/0.2 m Maroon Maroon Yes 1st and 2nd Dose)* for children who received a 3-dose primary series regardless which Bivalent (Use for 3rd vaccine (monovalent or bivalent) 6 months-4 years Maroon Maroon Yes Dose 3: 3 µg/0.2 mL Dose) was administered for Dose 3. 5–11 years Monovalent Orange Orange Yes 10 µg 0.2 ml NA NA 5-11 years Bivalent Orange Orange Yes NA NA 10 µg 0.2 mL 12 years and older Monovalent Grav Grav No 30 µg 0.3 mL NA NA 12 years and older **Bivalent** No NA NA 30 µa 0.3 mL Grav Grav Moderna **Primary series Booster doses** Vaccine vial Label border Dilution Vaccine Age indication composition cap color color required Dose **Injection volume** Dose **Injection volume** Dark blue 6 months-5 years Monovalent Magenta No 25 µg 0.25 mL NA NA Bivalent* 6 months-5 years Dark pink Yellow No NA NA 10 µg 0.2 mL 6-11 years Monovalent Dark blue Purple No 50 µg 0.5 mL NA NA **Bivalent** Dark blue No NA NA 0.25 mL 6-11 years Gray 25 µg 12 years and older Monovalent Red Light blue No 100 µg 0.5 mL NA NA 12 years and older **Bivalent** Dark blue NA 0.5 mL Gray No NA 50 µg

* A monovalent Pfizer-BioNTech vaccine is used for the first and second primary doses; a bivalent Pfizer-BioNTech vaccine is used for the third primary dose.



for Use of COVID-19 Vaccines Currently Authorized or Approved in the United States

COVID-19 vaccine products currently approved or authorized in the United States Continued

Janssen	Janssen COVID-19 Vaccine is authorized for adults ages 18 years and older in certain limited situations due to safety considerations. For guidance on retrospective record review, scheduling and administration of Janssen vaccine see Interim Clinical Considerations for Use of COVID-19 Vaccines: Appendix A												
Agoindication	Vaccine		Vaccine vial		Label border	Dilution	Primary series				Booster doses		
Age indication	compos	composition		cap color		color	required	C	Dose	Injection volume		Dose	Injection volume
18 years and older	Monova	Monovalent		Blue		No Color	No	No 5×10 ¹⁰ viral particles		0.5 mL		5×10 ¹⁰ viral particles	0.5 mL
Novavax													
Age indication	Vaccine	Vaccine	vial	Label bord	er Dilution	Primary series			Booster doses ⁺		doses ⁺		
	composition	сар со	lor	color		required	Dose		Injection	volume	De	ose	Injection volume
12 years and older	Monovalent	Royal blue N		No Color	or No		5 μg rS and 50 μg of Matrix-M™ adjuvant					nd 50 <i>µg</i> of ™ adjuvant	0.5 mL

+ Booster doses are only indicated for recipients 18 years and age and older in limited situations, see: Interim Clinical Considerations for Use of COVID-19 Vaccines Currently Approved or Authorized in the United States

All currently authorized or approved COVID-19 vaccines			
COVID-19 vaccination schedule	See the Interim COVID-19 Immunization Schedule for Ages 6 Months or Older		
Pre-vaccination counseling	 Prior to vaccination: Provide the vaccine-specific Fact Sheet for Recipients and Caregivers Screen for contraindications and precautions. CDC's Prevaccination Screening Form and Guidance document can be found at, <u>U.S. COVID-19</u> <u>Vaccine Product Information CDC</u>. Inform vaccine recipients mRNA or Novavax COVID-19 vaccines are recommended over Janssen COVID-19 Vaccine. Counsel vaccine recipients, parents, or guardians about expected reactions post-vaccination (e.g., pain and swelling at the injection site, fever, fatigue, headache). Inform mRNA and Novavax vaccine recipients, especially males ages 12-39 years, of the rare risk of myocarditis and pericarditis following receipt of these COVID-19 vaccines and the benefit of COVID-19 vaccination in reducing the risk of severe outcomes from COVID-19.⁺ Counseling should also include the need to seek care if symptoms of myocarditis and pericarditis. Inform vaccine recipients interested in or receiving Janssen COVID-19 Vaccine of the risk and symptoms of thrombosis with thrombocytopenia syndrome (TTS), as well as the need to seek immediate medical care should symptoms develop after receiving Janssen COVID-19 Vaccine. 		

+ See Interim Clinical Considerations for Use of COVID-19 Vaccines Currently Approved or Authorized in the United States for detailed guidance.



for Use of COVID-19 Vaccines Currently Authorized or Approved in the United States

All currently authorized or approved COVID-19 vaccines				
Interchangeability of vaccines	 In general, the same COVID-19 monovalent vaccine product (Pfizer-BioNTech, Moderna, Novavax) should be used for all doses in the primary series. In exceptional situations when the previous product cannot be determined/not available or if a person is unable to complete a series with the same COVID-19 vaccine due to a contraindication any age-appropriate mRNA COVID-19 vaccine may be used (administer at a minimum interval of 28 days). For booster vaccination, any homologous or heterologous age-appropriate mRNA vaccine can be used. Recommendations vary based on age and primary series product. See, Timing, spacing, age transitions, and coadministration of COVID-19 vaccines CDC.[§] 			
Coadministration with other vaccines	 COVID-19 vaccines may be administered on the same day as other vaccines. Persons, particularly adolescent or young adult males, might consider waiting 4 weeks after orthopoxvirus (monkeypox) vaccination (either JYNNEOS or ACAM2000) before receiving a Moderna, Novavax, or Pfizer-BioNtech COVID-19 vaccine. Administer each injection in a different injection site. 			
Contraindications	 History of: Severe allergic reaction (e.g., anaphylaxis) after a previous dose or to a component of the COVID-19 vaccine A known diagnosed allergy to a component of the COVID-19 vaccine For the Janssen COVID-19 Vaccine, TTS following receipt of a previous Janssen COVID-19 Vaccine (or other COVID-19 vaccines not currently authorized or approved in the United States that are based on adenovirus vectors, e.g., AstraZeneca)¹ 			
Precautions	 History of anaphylaxis after any other vaccine or injectable therapy (i.e., intramuscular, intravenous, or subcutaneous vaccines or therapies [excluding subcutaneous immunotherapy for allergies, i.e., "allergy shots"]) History of multisystem inflammatory syndrome in children (MIS-C) or multisystem inflammatory syndrome in adults (MIS-A) History of an immediate (within 4 hours of exposure) non-severe allergic reaction after a dose of one type of COVID-19 vaccine have a precaution to the same type of COVID-19 vaccine Allergy-related contraindication to one type of COVID-19 vaccine have a precaution to the other types of COVID-19 vaccines.^{**} Moderate or severe acute illness, with or without fever History of myocarditis or pericarditis after a dose of an mRNA or Novavax COVID-19 vaccine For Janssen COVID-19 Vaccine, a history of Guillain-Barré syndrome⁺⁺ 			

§ For booster vaccination, homologous or heterologous mRNA booster is recommended.

¶ Additionally, people with a history of an episode of immune-mediated syndrome characterized by thrombosis and thrombocytopenia, such as spontaneous or classic HIT, should not receive Janssen COVID-19 Vaccine. These people should receive an
mRNA or Novavax COVID-19 vaccine booster dose.

** People with a known allergy to polysorbate have a contraindication to both Novavax ad Janssen COVID-19 vaccines.

++ People who develop GBS within 6 weeks after receipt of Janssen COVID-19 Vaccine should not receive another dose of Janssen COVID-19 Vaccine. These people should receive a booster dose of an mRNA COVID-19 vaccine for subsequent doses.



for Use of COVID-19 Vaccines Currently Authorized or Approved in the United States

Considerations for all FDA-authorized or -approved COVID-19 vaccines				
Persons receiving HCT and CAR-T-cell therapy	If received doses of COVID-19 vaccine prior to or during HCT or CAR-T cell therapy, should be revaccinated for any monovalent primary ser and bivalent booster doses received before or during treatment at least 3 months (12 weeks) after transplant or CAR-T-cell therapy. There is revaccination for monovalent booster doses.			
Persons who are moderately or severely immunocompromised	See the Interim COVID-19 Immunization Schedule for Ages 6 Months or Older			
Persons receiving immunosuppressive therapies	Whenever possible, COVID-19 vaccines should be administered at least 2 weeks before initiation or resumption of immunosuppressive therapies.			
 SARS-CoV-2 infection Current infection History of previous infection Exposed to an infected person 	 COVID-19 vaccination is recommended for everyone ages 6 months and older, regardless of a history of symptomatic or asymptomatic SARS-CoV-2 infection. Defer vaccination until person has recovered from acute illness and criteria have been met for them to discontinue isolation. People who recently had SARS-CoV-2 infection may consider delaying their next COVID-19 dose by 3 months from symptom onset or positive test (if infection was asymptomatic). Viral testing to assess for acute SARS-CoV-2 infection or serologic testing to assess for prior infection is not recommended for the purpose of vaccine decision-making. Additional information at: Interim Clinical Considerations for Use of COVID-19 Vaccines: COVID-19 vaccination and SARS-CoV-2 infection CDC COVID-19 vaccination is not recommended for post-exposure prophylaxis. 			
Persons with history of multisystem inflammatory syndrome (MIS-C and MIS-A) from SARS-CoV-2 infection	 Wait until clinical recovery and at least 90 days after an MIS-C or MIS-A diagnosis to administer COVID-19 vaccine. For persons who developed MIS-C or MIS-A after COVID-19 vaccination, a conversation between the vaccine recipient, guardian, and clinical team or specialist to discuss benefits and risks of receiving a COVID-19 vaccine is encouraged. Additional information at: Interim Clinical Considerations for Use of COVID-19 Vaccines COVID-19 vaccination and MIS-C and MIS-A CDC 			
Persons who received passive antibody therapy (convalescent plasma/ monoclonal antibodies)	 COVID-19 vaccination can be given at any interval following receipt of passive antibody therapy. Persons should wait 2 weeks after COVID-19 vaccination before receiving tixagevimab/cilgavimab (EVUSHELD) for pre-exposure prophylaxis. 			
Persons who are pregnant, breastfeeding, trying to get pregnant, or might become pregnant in the future	Are recommended to be vaccinated according to the recommended schedule.			



for Use of COVID-19 Vaccines Currently Authorized or Approved in the United States

Considerations for mRNA vaccines and Novavax						
Persons with a history of myocarditis or pericarditis	 Development of myocarditis or pericarditis after a dose of an mRNA or Novavax COVID-19 vaccine is a precaution to a subsequent dose of any COVID-19 vaccine. If after a risk assessment the decision is made to administer a subsequent COVID-19 vaccine dose, the person should wait until after their episode has resolved. For information on potential use of Janssen COVID-19 Vaccine in this situation, see Interim Clinical Considerations for Use of COVID-19 Vaccines: Appendix A CDC Persons who have a history of myocarditis or pericarditis unrelated to mRNA or Novavax COVID-19 vaccination may receive any age-appropriate COVID-19 vaccine after the episode of myocarditis or pericarditis has resolved. For more information, see Interim Clinical Considerations for Use of COVID-19 Vaccines: COVID-19 vaccination and myocarditis and pericarditis CDC 					
Considerations for Janssen COVID-	19 Vaccine					
Janssen COVID-19 Vaccine is authorized for adults ages 18 years and older in certain limited situations due to safety considerations. For more information, see Interim Clinical Considerations for Use of COVID-19 Vaccines: Appendix A CDC						
Persons with a history of Guillain- Barré syndrome (GBS)	 Guillain- A history of GBS is a precaution for receipt of Janssen COVID-19 Vaccine. An mRNA or Novavax vaccine is recommended Persons who develop GBS within 6 weeks of Janssen COVID-19 vaccination should only receive an mRNA COVID-19 vaccine (monovalent or bilvalent vaccine as indicated) for subsequent doses. 					
Persons with a history of thrombosis with thrombocytopenia syndrome (TTS)	 It is contraindicated to administer Janssen COVID-19 Vaccine to persons with a history of TTS following receipt of the Janssen COVID-19 Vaccine or any other adenovirus vector-based COVID-19 vaccines (e.g., AstraZeneca's COVID-19 Vaccine). These persons should receive a dose of an mRNA COVID-19 vaccine as a booster dose at least 2 months (8 weeks) following their dose of the Janssen COVID-19 Vaccine and after their clinical condition has stabilized. 					
Persons with a history of heparin- induced thrombocytopenia (HIT)	 Persons with a history of an episode of an immune-mediated syndrome characterized by TTS, such as a spontaneous or classic HIT, should not receive Janssen COVID-19 Vaccine. These persons should receive an mRNA or Novavax COVID-19 vaccine. 					



for Use of COVID-19 Vaccines Currently Authorized or Approved in the United States

General COVID-19 Vaccination Information						
Persons vaccinated outside the United States	The recommendations for people vaccinated outside the United States depend on the number and type of vaccine(s) received for the primary series and booster doses. Current guidance can be found at: Interim Clinical Considerations for Use of COVID-19 Vaccines: Appendix B CDC					
Post-vaccination observation periods	 15 minutes: Vaccination providers, particularly when vaccinating adolescents, should consider observing vaccine recipients for 15 minutes after vaccination because of the risk of syncope. 30 minutes: Vaccination providers should consider observing persons with the following medical histories for 30 minutes after vaccination to monitor for allergic reactions: An allergy-related contraindication to a different type of COVID-19 vaccine Non-severe, immediate (onset within 4 hours) allergic reaction after a previous dose of COVID-19 vaccine Anaphylaxis after non-COVID-19 vaccines or injectable therapies 					
SARS-CoV-2 antibody testing	Antibody testing is not recommended for vaccine decision-making or to assess immunity following vaccination.					
Reporting requirements	 Adverse events that occur following COVID-19 vaccination should be reported to <u>VAERS</u>. COVID-19 providers are required to report: Vaccine administration errors Serious adverse events Myocarditis or pericardiitis after mRNA or Novavax COVID-19 Vaccine Cases of Multisystem Inflammatory Syndrome Cases of COVID-19 that result in hospitalization or death 					