Prevention of Coronavirus Disease 2019 (COVID-19)

- Although efficacy and safety not definitively established, COVID-19 vaccine (Janssen), also known as the Johnson & Johnson COVID-19 vaccine, is available under an FDA emergency use authorization (EUA) for active immunization to prevent COVID-19 in individuals 18 years of age or older. The American Society of Health-System Pharmacists, Inc. makes no representations or warranties, express or implied, including, but not limited to, any implied warranty of merchantability and/or fitness for a particular purpose, with respect to the information contained in the accompanying monograph, and specifically disclaims all such warranties. Readers of this information are advised that ASHP is not responsible for the accuracy of the information, for any errors or omissions, and/or for any consequences arising from the use of the information contained in the monograph in any and all practice settings. Readers are advised that decisions regarding use of drugs are complex medical decisions requiring the independent, informed decision of an appropriate health care professional, and that the information contained in the monograph is provided for informational purposes only. The entire monograph for a drug should be reviewed for a thorough understanding of the drug’s actions, uses and side effects. The American Society of Health-System Pharmacists, Inc. does not endorse or recommend the use of any drug. The information contained in the monograph is not a substitute for medical care.

Uses

- The EUA for the Janssen COVID-19 vaccine was based on FDA review of safety and efficacy data from an ongoing phase 3 clinical trial in adults indicating that a single dose of the vaccine was 66.9% effective in preventing moderate to severe/critical COVID-19 occurring at least 14 days after vaccination and 66.1% effective in preventing such infections at least 28 days after vaccination. In addition, efficacy of a single dose of the vaccine in preventing severe/critical COVID-19 was 76.7 or 85.4% when cases occurring at least 14 or 28 days, respectively, after vaccination were considered.
- The EUA for the Janssen COVID-19 vaccine authorizes that distribution of the vaccine will be controlled by the US government for use consistent with the terms and conditions of the EUA. (See Restricted Distribution under Preparations.)
- To mitigate risks of this unapproved vaccine, the EUA includes certain mandatory requirements (e.g., providing the recipient or caregiver with information consistent with the EUA fact sheet for recipients and caregivers, ensuring that all vaccination administration errors and all serious adverse events potentially attributable to the vaccine are reported as specified in the EUA fact sheet for healthcare providers). (See EUA Requirements for Postvaccination Monitoring and Mandatory Vaccine Adverse Event Reporting under Cautions.)
- Consult the Janssen COVID-19 vaccine EUA letter of authorization (https://www.fda.gov/media/146303/download), EUA fact sheet for healthcare providers (https://www.fda.gov/media/146304/download), and EUA fact sheet for recipients and caregivers (https://www.fda.gov/media/146305/download) for additional information.
- There currently are 3 different COVID-19 vaccines available for use in the US under FDA EUAs, including a viral-vector vaccine (Janssen COVID-19 vaccine) and 2 nucleoside-modified mRNA vaccines (Moderna COVID-19 vaccine and Pfizer-BioNTech COVID-19 vaccine). ACIP does not state a preference for any specific currently authorized COVID-19 vaccine when the vaccines are used within the scope of their respective EUAs and states that individuals should be encouraged to receive the earliest vaccine available to them. However, currently available COVID-19 vaccines are not interchangeable with each other. (See Dosage under Dosage and Administration.)

Dosage and Administration

General

- Must have appropriate medications and supplies immediately available to assess and manage immediate allergic reactions in the event that an acute anaphylactic reaction occurs following administration of a COVID-19 vaccine, including COVID-19 vaccine (Janssen). Healthcare personnel trained and qualified to recognize signs and symptoms of anaphylaxis and administer IM epinephrine should be available at vaccination locations at all times. Vaccination locations that anticipate vaccinating large numbers of people (e.g., mass vaccination clinics) should plan adequate staffing and supplies (including epinephrine) for assessment and management of anaphylaxis. (See Hypersensitivity Reactions under Cautions.)
- Prior to administration of the Janssen COVID-19 vaccine, screen all individuals for contraindications and precautions to vaccination. Do not give the vaccine to those with a contraindication, (See Contraindications and see Warnings/Precautions under Cautions.)
- Monitor all vaccine recipients for immediate adverse reactions according to CDC (ACIP) guidelines. When administered to individuals with no contraindications to vaccination with the Janssen COVID-19 vaccine, ACIP states observe those who have a history of an immediate allergic reaction of any severity to any other vaccine or injectable therapy and those who have a history of anaphylaxis due to any cause not considered a contraindication for 30 minutes, and observe all other individuals for 15 minutes. A longer period of observation may be indicated in some individuals based on clinical concern (e.g., pruritus and swelling confined to the injection site developed during observation period). Instruct vaccine recipients to seek immediate medical care if they develop signs or symptoms of an allergic reaction after their observation period ends and they have left the vaccination site. (See Hypersensitivity Reactions under Cautions.)
- Syncope (vasovagal or vasodepressor reaction; fainting) may occur following administration of parenteral vaccines; such reactions usually occur within 15 minutes following vaccine administration and are reported most frequently in adolescents and young adults. Take appropriate measures to decrease risk of injury if a patient becomes weak or dizzy or loses consciousness (e.g., vaccinees should sit or lie down during and for 15 minutes after vaccination). If syncope occurs, observe patient until symptoms resolve.
- At the time that the Janssen COVID-19 vaccine is administered, give vaccine recipient or their caregiver a vaccination record card that provides the name of the vaccine (Janssen COVID-19 vaccine) and the date the vaccine was administered.
- Provide vaccine recipient or their caregiver with information on, and encourage participation in, CDC’s v-safe program, a voluntary smartphone-based tool that uses text messaging and web surveys to monitor for adverse effects in COVID-19 vaccine recipients. (See EUA Requirements for Postvaccination Monitoring and Mandatory Vaccine Adverse Event Reporting under Cautions.)
● Prior to vaccination, counsel vaccine recipient or their caregiver about local and systemic adverse effects that may occur following vaccination. (See Cautions and see Advice to Patients.)

● Antipyretics or analgesics (e.g., acetaminophen, NSAIDs) may be taken to treat postvaccination local or systemic symptoms, if medically appropriate. However, routine premedication for the purpose of preventing postvaccination symptoms in vaccinees is not currently recommended since information not available regarding possible impact on antibody response to COVID-19 vaccines. Premedication with antihistamines prior to vaccination to prevent allergic reactions is not recommended; antihistamines do not prevent anaphylaxis and may mask cutaneous symptoms, which could delay diagnosis and management of anaphylaxis. (See Hypersensitivity Reactions under Cautions.)

● Counsel individuals who have received the Janssen COVID-19 vaccine and are considered fully vaccinated against COVID-19 (see Dosage and Administration) to follow current CDC guidance for fully vaccinated individuals. This may include wearing a mask and physically distancing if required by federal, state, local, tribal, or territorial laws, rules, and regulations and following CDC travel guidance and any applicable workplace or school guidance. (See Limitations of Vaccine Effectiveness under Cautions.)

Administration

IM Administration

Administer only by IM injection into the deltoid.

The Janssen COVID-19 vaccine is supplied as a suspension in multiple-dose vials. Although the vaccine is initially stored frozen by the manufacturer, it is shipped and stored refrigerated at a temperature of 2–8°C. Do not refreeze the vaccine. (See Storage under Stability.)

Do not dilute the Janssen COVID-19 vaccine.

Swirl vials of vaccine gently in an upright position for 10 seconds before withdrawing a dose; do not shake.

The Janssen COVID-19 vaccine should appear as a colorless to slightly yellow, clear to very opalescent suspension; do not use if it is discolored or contains particulates.

To administer a dose, withdraw 0.5 mL of Janssen COVID-19 vaccine from the vial using aseptic technique and an appropriate syringe and needle and administer immediately.

Each multiple-dose vial of Janssen COVID-19 vaccine provides five 0.5-mL doses. Because the vaccine does not contain preservatives, it is critical that any vaccine remaining in the vial that does not constitute a full 0.5-mL dose should be discarded and not pooled with vaccine from other vials to create a dose.

Data not available regarding concomitant administration with other vaccines. (See Vaccines under Interactions.)

Dosage

Administer the Janssen COVID-19 vaccine as a single 0.5-mL dose. Each 0.5-mL dose contains 5 × 10^10 virus particles of recombinant, replication-incompetent Ad26.

The Janssen COVID-19 vaccine is not interchangeable with any other COVID-19 vaccine.

A single dose of the Janssen COVID-19 vaccine is considered a complete, valid vaccination series. Individuals should not receive more than one single, valid vaccination series for active immunization against COVID-19 (i.e., single dose of the Janssen COVID-19 vaccine or 2-dose regimen of an mRNA vaccine [Moderna COVID-19 vaccine or Pfizer-BioNTech COVID-19 vaccine]).

Individuals are considered fully vaccinated against COVID-19 if ≥2 weeks have elapsed since they received a single dose of the Janssen COVID-19 vaccine or completed a 2-dose vaccination series of an mRNA vaccine (Moderna COVID-19 vaccine or Pfizer-BioNTech COVID-19 vaccine). Those who have a contraindication to vaccination or who otherwise cannot complete a vaccination series are not considered fully vaccinated.

Safety and efficacy regarding use of the Janssen COVID-19 vaccine after a dose of an mRNA COVID-19 vaccine not established. However, ACIP states that, in limited, exceptional situations when an individual received the first dose of an mRNA COVID-19 vaccine but is unable to complete the vaccination series with either the same or different mRNA COVID-19 vaccine (e.g., due to a contraindication), may consider giving a single dose of the Janssen COVID-19 vaccine at least 28 days after the dose of mRNA COVID-19 vaccine. (See Hypersensitivity Reactions under Cautions.) In such exceptional circumstances, consider the individual to have received valid, single-dose vaccination with the Janssen COVID-19 vaccine (not a mixed vaccination series) and consider the individual to be fully vaccinated against COVID-19 if ≥2 weeks have elapsed since the dose of Janssen COVID-19 vaccine.

Report all vaccine administration errors and deviations from currently recommended dosage and vaccination schedule to the vaccinee and the Vaccine Adverse Event Reporting System (VAERS). Information on preventing and reporting COVID-19 vaccine administration errors and recommendations for specific actions to take if an administration error or deviation from recommended vaccination schedule occurs are available at the CDC website at https://www.cdc.gov/vaccines/covid-19/info-by-product/clinical-considerations.html.
component elicited the reaction: ACIP considers this a precaution, but not a contraindication, to COVID-19 vaccination.

History of allergic reactions (including severe allergic reactions) not related to COVID-19 vaccines or other vaccines or injectable therapies: ACIP states that allergic reactions to food, pets, insects, venom, or environmental allergies and allergic reactions to oral medications (including the oral equivalents of injectable medications) are not a contraindication or precaution to COVID-19 vaccination. Latex allergy is not a contraindication or precaution since viai stoppers of COVID-19 vaccines are not made with natural rubber latex. Allergies to eggs or gelatin are not a contraindication or precaution since COVID-19 vaccines do not contain eggs or gelatin.

If a precaution for COVID-19 vaccination is identified, ACIP recommends performing a risk assessment to help decide whether the individual should be vaccinated. The risk assessment should consider risk of exposure to SARS-CoV-2 (e.g., because of residence in a congregate setting such as a long-term care facility, occupation), risk of severe disease or death due to COVID-19 (e.g., because of age or underlying medical conditions), unknown risk of anaphylaxis (including fatal anaphylaxis) following COVID-19 vaccination in individuals with a history of immediate allergic reactions to other vaccines or injectable therapies, and ability to be vaccinated in a setting where appropriate medical care is immediately available to treat anaphylaxis if it occurs.

When a COVID-19 vaccine, including the Janssen COVID-19 vaccine, is administered to individuals without a contraindication to such vaccines, ACIP states observe those with a history of an immediate allergic reaction of any severity to any other vaccine or injectable therapy and those with a history of anaphylaxis due to any cause not considered a contraindication for 30 minutes after the vaccine dose and observe all other individuals for 15 minutes. Instruct vaccine recipients to seek immediate medical care if they develop signs or symptoms of an allergic reaction after their observation period ends and they have left the vaccination site.

Appropriate medications and supplies to assess and manage immediate allergic reactions (e.g., sufficient quantities of epinephrine in prefilled syringes or autoinjectors) must be immediately available in the event that an acute anaphylactic reaction occurs following administration of a COVID-19 vaccine. Early recognition of clinical signs and symptoms of anaphylaxis is important since such reactions require immediate treatment. Immediately treat individuals with suspected anaphylaxis with IM epinephrine.

ACIP interim guidance regarding early recognition of clinical signs and symptoms of anaphylaxis and guidance regarding preparation for and management of anaphylaxis at COVID-19 vaccination sites, including recommendations for medications and supplies to have immediately available and specific recommendations regarding therapeutic management of anaphylaxis, are available at the CDC website at https://www.cdc.gov/vaccinesafety/enhancing-safety/monitoring/cisa/index.html.

Thrombotic Events
At the time of FDA’s safety analysis of data from the ongoing phase 3 trial for the EUA, there had been 6 reports of deep-vein thrombosis in individuals who received the Janssen COVID-19 vaccine (2 serious; 5 within 28 days) and 2 such events in placebo recipients (1 serious; 2 within 28 days). In addition, there were 4 reports of pulmonary embolism (3 serious; 2 within 28 days) in vaccine recipients versus 1 report in placebo recipients (serious; within 28 days) and 1 report of transverse sinus thrombosis (serious; within 28 days of vaccination) in vaccine recipients versus none in placebo recipients. A causal relationship between these thromboembolic events and the vaccine was not established based on the phase 3 trial data. However, following issuance of the FDA EUA for the Janssen COVID-19 vaccine, thrombosis involving large blood vessels occurring with thrombocytopenia were reported rarely, and evaluation of these cases suggest that a causal relationship between the vaccine and thrombosis with thrombocytopenia is plausible.

Thrombosis with Thrombocytopenia
Thrombosis involving large blood vessels (e.g., the cerebral venous sinuses, portal vein, lower extremity veins, pulmonary artery) occurring with thrombocytopenia (also known as thrombosis with thrombocytopenia syndrome [TTS]) and with onset of symptoms approximately 1–2 weeks after vaccination reported rarely in recipients of the Janssen COVID-19 vaccine during post-authorization surveillance.

In response to reports of cerebral venous sinus thrombosis (CVST) with thrombocytopenia in a few recipients of the Janssen COVID-19 vaccine, FDA and CDC temporarily paused use of the vaccine in the US out of an abundance of caution to allow time for investigation of these cases, inform vaccine recipients and healthcare providers of possible symptoms, and inform healthcare providers about appropriate management of this extremely rare adverse event. The temporary pause was lifted 10 days later after completion of a thorough safety review. At the time the pause was initiated (April 13, 2021), 6 cases of CVST had been reported in vaccine recipients, including one fatality; all 6 cases were reported in females 18–49 years of age and occurred within 6–13 days after vaccination.

After extensive data evaluation and analyses of risks and benefits of vaccination with the Janssen COVID-19 vaccine (both population- and individual-level risks and benefits), FDA and CDC determined that known and potential benefits of the vaccine outweigh its known and potential risks in adults ≥18 years of age. At the time of CDC’s safety data analysis (April 21, 2021), a total of 15 cases of TTS had been reported and confirmed, including 3 fatalities. These post-authorization cases all occurred in females 18–59 years of age (median age 37 years), 6 of the 15 cases occurred in women 18–49 years of age, symptom onset was 6–15 days after vaccination (median 8 days), and clinical course shared features with autoimmune heparin-induced thrombocytopenia. Some of these women had underlying medical conditions or risk factors for hypercoagulability (e.g., obesity, oral contraceptive use, hypothyroidism, hypertension); none had a documented history of previous thrombotic events, known diagnosis of an underlying clotting disorder, or family or personal history of clotting disorders. A single case of CVST with thrombocytopenia was reported during the phase 3 clinical trial of the vaccine (a male in the 18–49 years of age group). At the time of CDC’s safety analysis, highest rates of TTS per vaccine doses administered were in women <50 years of age (reporting rates to VAERS were 7 cases of TTS per million doses of the vaccine administered to women 18–49 years of age and 0.9 cases per million doses administered to women ≥50 years of age).

Data accumulated as of May 7, 2021 indicated >6.7 million doses of the Janssen COVID-19 vaccine had been administered in the US and a total of 28 cases of TTS had been reported to VAERS and confirmed (22 cases in females and 6 cases in males). Analyses of all 28 cases indicated that the median age was 40 years (range: 18–59 years of age), median time from vaccination to symptom onset was 9 days (range 4–15 days), total of 3 deaths had occurred, CVST occurred in 19 of the 28 cases, nadir platelet counts were <50,000/mm³ in 18 of the 28 cases, and results of enzyme-linked immunosorbent assay (ELISA) for platelet-activating antibodies against platelet factor 4 (PF4) were positive in 24 of the 26 patients tested.

FDA and CDC are continuing to closely monitor reports of TTS in recipients of the Janssen COVID-19 vaccine. Based on currently available evidence, FDA states that a causal relationship between TTS and the vaccine is plausible. Specific risk factors for TTS following vaccination with the Janssen COVID-19 vaccine and level of potential excess risk due to vaccination still under investigation.

ACIP states that women <50 years of age can receive any FDA-authorized COVID-19 vaccine; however, they should be informed about the rare risk of TTS after receipt of the Janssen COVID-19 vaccine and the availability of other FDA-authorized COVID-19 vaccines (Moderna COVID-19 vaccine, Pfizer-BioNTech COVID-19 vaccine). In addition, until more information becomes available about the etiology of TTS associated with the Janssen COVID-19 vaccine, some experts advise offering Moderna COVID-19 vaccine or Pfizer-BioNTech COVID-19 vaccine to individuals with a history of an immune-mediated syndrome characterized by thrombosis and thrombocytopenia (see Individuals with a History of Thrombosis or Risk Factors for Thrombosis under Cautions).

Healthcare providers should be alert to and maintain a high index of suspicion for signs and symptoms of TTS (e.g., severe headache, backache, new neurologic symptoms, severe abdominal pain, shortness of breath, leg swelling, petechiae, new or easy bruising) occurring approximately 1–2 weeks after vaccination in individuals who have received the Janssen COVID-19 vaccine.

Based on data regarding the US patients who developed TTS after receiving the Janssen COVID-19 vaccine and data regarding patients in Europe who developed immune thrombotic thrombocytopenia after receiving a different adenoviral-vector COVID-19 vaccine not available in the US (i.e., AstraZeneca COVID-19 vaccine), the clinical course shares features with autoimmune heparin-induced thrombocytopenia and may be associated with platelet-activating antibodies against PF4.

When managing thrombotic events and thrombocytopenia in patients who recently received vaccination with the Janssen COVID-19 vaccine, avoid use of heparin and its derivatives (may be harmful); consider use of other anticoagulants and high-dose immune globulin IV (IGIV). Consultation with hematology specialists strongly recommended. Information regarding diagnosis and management of suspected cases of TTS is provided in the CDC Health Alert Network (HAN) notification at https://emergency.cdc.gov/han/2021/han00442.asp and is available from the American Society of Hematology (ASH) at https://www.hematology.org/covid-19/vaccine-induced-immune-thrombocytopenia.

Instruct vaccinees to seek immediate medical attention if they develop shortness of breath, chest pain, leg swelling, persistent abdominal pain, neurologic symptoms (including severe or persistent headaches or blurred vision), easy bruising, or a diffuse rash consisting of petechiae (pinpoint-like spots) beyond the vaccination site within a few weeks after receiving the Janssen COVID-19 vaccine. These symptoms are distinct from commonly reported adverse effects that may be experienced in the first few days following vaccination (e.g., headache, fatigue, muscle aches, nausea) that usually are mild to moderate in severity and last 1–2 days.

If TTS occurs following vaccination, report the case to VAERS. (See EUA Requirements for Postvaccination Monitoring and Mandatory Vaccine Adverse Event Reporting Under Cautions.)

Other Adverse Events
Other adverse effects with a numerical imbalance between vaccine recipients and placebo recipients include 4 reports of seizures in vaccine recipients (1 serious; 4 within 28 days of vaccination) versus 1 seizure event in placebo recipients and 6 reports of tinnitus in vaccine recipients (not serious; all 6 within 28 days, including 3 events within 2 days) versus none in
placode recipients. A causal relationship between these events and the vaccine could not be established.

### Concomitant Illness

- **Function)**, personal risk of severe acute COVID-19 (e.g., age, underlying conditions), level of decisions regarding COVID-19 vaccination in those with a history of MIS-A or MIS-C, and timing of any immunomodulatory therapies.

Current evidence suggests that the risk of reinflection with SARS-CoV-2 is low in the months after initial infection, but may increase with time due to waning immunity. ACIP states that individuals with a history of MIS-A or MIS-C should consider deferring COVID-19 vaccination until they have recovered from their illness and for 90 days after the date MIS-A or MIS-C was diagnosed, recognizing that the risk of reinfection and, therefore, the benefit from vaccination might increase with time following the initial infection.

If MIS-A or MIS-C associated with a confirmed SARS-CoV-2 infection develops after receipt of a COVID-19 vaccine, consider referral to a specialist in infectious diseases, rheumatology, or cardiology. Healthcare providers and health departments can also request a clinical consultation from the Clinical Immunization Safety Assessment COVIDvax project (https://www.cdc.gov/vaccinesafety/ensuring-safety/monitoring/cisa/index.html).

If MIS-A or MIS-C occurs following COVID-19 vaccination, report the case to VAERS. (See EUA Requirements for Postvaccination Monitoring and Mandatory Vaccine Adverse Event Reporting under Cautions.)

### Individuals with Underlying Medical Conditions

ACIP states that individuals with altered immunocompetence or certain underlying medical conditions may receive COVID-19 vaccine if they have no contraindications to vaccination. ACIP does not state a preference for any specific COVID-19 vaccine in such individuals. Clinical trials of COVID-19 vaccines have demonstrated safety and efficacy profiles in individuals with some underlying medical conditions, including those that place them at increased risk for severe COVID-19, that are similar to safety and efficacy profiles in individuals without comorbidities.

US healthcare personnel and health departments can request a consultation from the Clinical Immunization Safety Assessment COVIDvax project (https://www.cdc.gov/vaccinesafety/ensuring-safety/monitoring/cisa/index.html) if they have concerns about vaccinating individuals with certain underlying medical conditions.

### Individuals with Altered Immunocompetence

Individuals with altered immunocompetence, including those receiving immunosuppressive therapy (see Specific Drugs under Interactions), may have diminished immune responses to vaccines, including the Janssen COVID-19 vaccine.

Although some individuals with altered immunocompetence (e.g., stable HIV infection) were included in the ongoing phase 3 trial evaluating the Janssen COVID-19 vaccine, the number of such individuals was insufficient to evaluate safety of the vaccine in such populations.

ACIP states that individuals with HIV infection or other immunocompromising conditions and individuals receiving immunosuppressive therapies may be at increased risk for severe COVID-19 and, although data are not currently available to establish safety and efficacy in such individuals, they may receive any authorized COVID-19 vaccine if they have no contraindications to vaccination. However, counsel such individuals about the unknown safety profile and effectiveness of COVID-19 vaccines in immunocompromised populations and the potential for reduced immune responses and need to continue following all current guidelines to protect themselves from COVID-19.

Antibody testing to assess for immunity to SARS-CoV-2 following COVID-19 vaccination in individuals with altered immunocompetence is not recommended. (See Interpretation of SARS-CoV-2 Testing in Vaccinated Individuals under Cautions.)

### Individuals with Autoimmune Conditions

ACIP states that individuals with autoimmune conditions may receive any authorized COVID-19 vaccine, unless they have a contraindication to vaccination. Individuals with autoimmune conditions were not excluded from clinical trials evaluating the mRNA COVID-19 vaccines and these trials showed no imbalances in the occurrence of symptoms consistent with autoimmune conditions or inflammatory disorders in trial participants who received COVID-19 vaccine compared with those who received placebo.

### Individuals with a History of Thrombosis or Risk Factors for Thrombosis

TTTS reported rarely in recipients of the Janssen COVID-19 vaccine (see Thrombosis with Thrombocytopenia under Cautions). Although etiology of TTTS associated with the Janssen COVID-19 vaccine unclear, it appears to be similar to heparin-induced thrombocytopenia. Until more information becomes available, some experts advise that an mRNA COVID-19 vaccine (Moderna COVID-19 vaccine or Pfizer-BioNTech COVID-19 vaccine) should be offered to individuals who have had an episode of an immune-mediated syndrome characterized by thrombosis and thrombocytopenia (e.g., heparin-induced thrombocytopenia) that resolved within the past 90 days.

Consider that venous thromboembolism (VTE; defined as DVT and/or PE) is common and that biologic mechanisms for VTE (as well as arterial thrombosis) differ from underlying immune-mediated mechanisms for heparin-induced thrombocytopenia. Based on current knowledge, experts believe that individuals with risk factors for VTE (e.g., inherited or acquired thrombophilia including factor V Leiden; prothrombin gene 20210A mutation; antiphospholipid syndrome; protein C, protein S, or antithrombin deficiency) or history of other types of thrombosis (including CVST) not associated with thrombocytopenia are unlikely to be at increased risk for TTTS.

In addition, although the risk of thrombosis is increased during pregnancy and the postpartum period and with certain hormonal contraceptives (e.g., estrogen-progesterin oral contraceptiv...
transdermal system, or vaginal ring), experts believe that these factors do not make individuals more susceptible to TTS after receipt of the Janssen COVID-19 vaccine. ACIP states that such individuals can receive any FDA-authorized COVID-19 vaccine, including the Janssen COVID-19 vaccine.

ACIP states that premedication with anticoagulants or aspirin prior to vaccination with the Janssen COVID-19 vaccine is not recommended. (See Specific Drugs under Interactions.)

**Individuals with Liver Disease**

American Association for the Study of Liver Diseases (AASLD) released a consensus statement regarding use of COVID-19 vaccines in individuals with chronic liver disease or a liver transplant.

Although safety and efficacy data regarding use of COVID-19 vaccines in individuals with chronic liver disease are limited and additional studies needed, safety and efficacy of the vaccines in such individuals expected to be similar to the general population. AASLD states that individuals with chronic liver disease receiving antiviral treatment for HBV or HCV infection and those receiving medical therapy for primary biliary cholangitis or autoimmune hepatitis should not discontinue such therapy when receiving COVID-19 vaccination. In addition, consider COVID-19 vaccination for patients with hepatocellular carcinoma undergoing locoregional or systemic therapy without interruption of treatment.

Although solid organ transplant recipients, including liver transplant recipients, not included in clinical trials of COVID-19 vaccines and efficacy and safety in such individuals not known, AASLD states that liver transplant candidates should receive COVID-19 vaccination prior to transplantation, whenever possible, to help ensure an adequate immune response. The best time for COVID-19 vaccination in previously unvaccinated liver transplant recipients is likely to be ≥3 months after transplant; however, vaccination may be given as early as 6 weeks after transplant if indicated based on ongoing community spread of SARS-CoV-2, especially in those at highest risk with other comorbid factors associated with severe COVID-19.

Consult AASLD consensus statement for additional guidance on use of COVID-19 vaccines in individuals with chronic liver disease.

**Individuals with a History of Guillain-Barré Syndrome**

At the time of FDA’s safety analysis of data from the ongoing phase 3 trial for the EUA, a single case of Guillain-Barré syndrome (GBS) occurred in a vaccine recipient 16 days after the vaccine dose and a single case occurred in a placebo recipient 10 days after the dose. FDA stated that, although the case of GBS was unlikely to be related to the vaccine, a causal relationship cannot be definitively excluded.

ACIP states that individuals with a history of GBS may receive COVID-19 vaccination, unless they have a contraindication to the vaccine. A history of GBS is not usually considered a contraindication or precaution to vaccination with most vaccines.

If GBS occurs following COVID-19 vaccination, report the case to VAERS. (See EUA Requirements for Postvaccination Monitoring and Mandatory Vaccine Adverse Event Reporting under Cautions.)

**Individuals with a History of Bell’s Palsy**

At the time of FDA’s safety analysis of data from the ongoing phase 3 trial for the EUA, there had been 2 reported cases of Bell’s palsy (facial paralysis) in the vaccine group and 2 cases in the placebo group. FDA stated that, although the cases of Bell’s palsy were unlikely to be related to the vaccine, a causal relationship cannot be definitively excluded.

ACIP states that, in the absence of a causal relationship between COVID-19 vaccines and Bell’s palsy, individuals with a history of Bell’s palsy may receive COVID-19 vaccination, unless they have a contraindication to the vaccine.

If Bell’s palsy occurs following COVID-19 vaccination, report the case to VAERS. (See EUA Requirements for Postvaccination Monitoring and Mandatory Vaccine Adverse Event Reporting under Cautions.)

**Individuals with Increased Bleeding Risk**

Advises individuals who have bleeding disorders or are receiving anticoagulant therapy and/or their caregiver about the risk of hematoma from IM injections.

ACIP states that IM vaccines may be given to individuals who have bleeding disorders if a clinician familiar with the patient’s bleeding risk determines that the preparation can be administered IM with reasonable safety. In these cases, use a fine needle (23 gauge or smaller) to administer the vaccine and apply firm pressure to the injection site (without rubbing) for ≥2 minutes. In individuals receiving therapy for hemophilia, schedule IM vaccines for administration shortly after a dose of such therapy.

Individuals receiving anticoagulation therapy presumably have the same bleeding risk as those with clotting factor disorders and should follow the same guidelines for IM administration. If possible, schedule IM vaccines prior to use of an anticoagulant so that the patient’s risk of bleeding is not increased by the drug’s therapeutic action.

**Individuals Vaccinated Outside the US**

Some individuals in the US may have previously received vaccination against COVID-19 in another country using a vaccine not authorized by FDA and/or not listed for emergency use by WHO. For the purposes of public health guidance, ACIP states that only individuals who have received all recommended doses of a COVID-19 vaccine authorized by FDA or listed by WHO for emergency use are considered fully vaccinated.

Data not available regarding safety and efficacy of administering an FDA-authorized COVID-19 vaccine to individuals who previously received a COVID-19 vaccine not authorized in the US. However, ACIP states that such individuals may be offered revaccination with an FDA-authorized COVID-19 vaccine in certain circumstances. If an FDA-authorized COVID-19 vaccine is administered to an individual who previously received a vaccine not authorized by FDA, the minimum interval between last dose of a non-FDA-authorized COVID-19 vaccine and dose of an FDA-authorized COVID-19 vaccine is 28 days.

**Fully or Partially Vaccinated with an FDA-authorized COVID-19 Vaccine**

Individuals vaccinated outside the US with an FDA-authorized COVID-19 vaccine do not need any additional doses in the US if they previously received all the recommended doses of the vaccine.

If an individual in the US received the first dose of an FDA-authorized COVID-19 vaccine outside the US and a 2-dose regimen is required, ACIP states the vaccination series does not need to be restarted; administer the second dose of the vaccine as close to the recommended interval as possible.

**Previously Received a COVID-19 Vaccine not Authorized by FDA but Listed for Emergency Use by WHO**

ACIP states that individuals who completed a COVID-19 vaccination series with a vaccine listed for emergency use by WHO do not need any additional doses using an FDA-authorized COVID-19 vaccine.

May offer vaccination with an FDA-authorized COVID-19 vaccine to individuals who partially completed a COVID-19 vaccination series outside the US with a vaccine listed for emergency use by WHO.

**Previously Received a COVID-19 Vaccine not Authorized by FDA or Listed for Emergency Use by WHO**

May offer vaccination with an FDA-authorized COVID-19 vaccine to individuals who completed or partially completed a COVID-19 vaccination series outside the US with a vaccine not authorized by FDA or listed for emergency use by WHO.

**Limitations of Vaccine Effectiveness**

May not protect all vaccine recipients against COVID-19.

Use of COVID-19 vaccines for breakthrough management or for postexposure prophylaxis to prevent SARS-CoV-2 infection in individuals with a specific known exposure to the virus is unlikely to be effective and is not currently recommended. ACIP states that, because the median incubation period of SARS-CoV-2 infection is 4–5 days, it is unlikely that a dose of a COVID-19 vaccine would provide an adequate immune response within the incubation period for effective postexposure prophylaxis.

FDA states that data are too limited to date to assess the effect of the Janssen COVID-19 vaccine for prevention of asymptomatic SARS-CoV-2 infection; additional evaluations are needed, including data from clinical trials and from use of the vaccine after issuance of the EUA.

FDA states that data are too limited to date to assess the effect of the Janssen COVID-19 vaccine against transmission of SARS-CoV-2 from individuals who become infected despite vaccination. Demonstrated high efficacy against symptomatic COVID-19 may translate to overall prevention of transmission in populations with high enough vaccine uptake; however, it is possible that if efficacy against asymptomatic infection were lower than efficacy against symptomatic infection, asymptomatic cases in combination with reduced mask-wearing and social distancing could result in significant continued transmission of the virus. Additional evaluations are needed, including data from clinical trials and from use of the vaccine after issuance of the EUA, to assess the effect of the vaccine in preventing virus shedding and transmission, particularly in individuals with asymptomatic infection.

Based on unknown duration of vaccine-induced protection and unknown efficacy against emerging SARS-CoV-2 variants, counsel individuals who receive COVID-19 vaccination and are considered fully vaccinated to follow current guidance for fully vaccinated individuals to protect themselves and others. This may include wearing a mask and physically distancing in certain settings and venues if required by federal, state, local, tribal, or territorial laws, rules, and regulations and following CDC travel guidance and any applicable workplace or school guidance. CDC issued interim public health recommendations for individuals who are fully vaccinated against COVID-19 (defined as ≥2 weeks after a single dose of the Janssen COVID-19 vaccine or ≥2 weeks after completion of a 2-dose vaccination series of the Moderna COVID-19 vaccine or Pfizer-BioNTech COVID-19 vaccine); consult these recommendations (available at the CDC website at https://www.cdc.gov/coronavirus/2019-ncov/vaccines/fully-vaccinated-guidance.html) for information on precautionary measures that fully vaccinated individuals should take in certain social situations and/or following exposure to someone with suspected or confirmed COVID-19.

Withholding COVID-19 vaccination due to concerns about efficacy against current or future SARS-CoV-2 viral variants not recommended.

If COVID-19 vaccine breakthrough infection occurs in an individual who is fully vaccinated against COVID-19 (i.e., RNA or antigen detected in a respiratory specimen collected ≥14 days after an individual completed all recommended doses of an FDA-authorized COVID-19 vaccine),
healthcare providers, local health departments, and clinical laboratories are encouraged to request that the respiratory specimen be held for further testing and the case reported to the state health department for further investigation and reporting to the national system. If COVID-19 vaccine breakthrough infection results in hospitalization or death, report the case to VAERS. (See EUA Requirements for Postvaccination Monitoring and Mandatory Vaccine Adverse Event Reporting under Cautions.)

If an individual is fully vaccinated and tests positive for SARS-CoV-2, healthcare providers and local health departments are encouraged to request that the specimen be held and the case reported to the state health department. CDC will work with the state health department to collect information about the case. In addition, report information about such cases to VAERS. (See EUA Requirements for Postvaccination Monitoring and Mandatory Vaccine Adverse Event Reporting under Cautions.)

Duration of Immunity

Duration of protection against SARS-CoV-2 infection following vaccination with a single dose of the Janssen COVID-19 vaccine not fully evaluated.

Because available trial data have a limited length of follow-up to date, it is not possible at this time to assess sustained efficacy over a period longer than 2 months. ACIP states that the need for and timing of booster doses of COVID-19 vaccines not established. Additional vaccine doses beyond those recommended for a complete, valid vaccination series (see Dosage under Dosage and Administration) are not recommended at this time. Recommendations on revaccination or additional doses of COVID-19 vaccines may be updated when additional information is available.

Improper Storage and Handling

Improper storage or handling of vaccines may reduce or destroy vaccine potency resulting in inadequate or no immune response in vaccinees. Inspect all vaccines on delivery and monitor during storage to ensure that recommended storage temperatures are maintained.

The Janssen COVID-19 vaccine must be shipped, stored, and handled under specific conditions at all times, according to specifications in the EUA fact sheet for healthcare providers and guidance from the manufacturer and CDC. Do not administer vaccine that has been mishandled or has not been stored at the recommended temperatures. (See Storage under Stability.)

Contact the manufacturer at 800-565-4008 for guidance if there are concerns about mishandling or defective or damaged vaccine.

EUA Requirements for Postvaccination Monitoring and Mandatory Vaccine Adverse Event Reporting

Safety and efficacy not established. FDA issued an EUA that permits use of the Janssen COVID-19 vaccine for prevention of COVID-19 in individuals ≥18 years of age when administered according to the single-dose regimen specified in the EUA. (See Prevention of Coronavirus Disease 2019 [COVID-19] under Uses.)

Some data available regarding adverse effects associated with use of the vaccine. (See Common Adverse Effects under Cautions.) Additional adverse effects, some of which may be serious, may become apparent with more widespread use.

Monitor all vaccine recipients for immediate adverse reactions according to CDC (ACIP) guidelines. (See General under Dosage and Administration.)

Provide vaccine recipients or their caregivers with information on, and encourage participation in, CDC’s voluntary smartphone-based tool (v-safe) that uses text messaging and web surveys to check in with individuals who have received a COVID-19 vaccine to identify potential adverse effects. Reports to v-safe that indicate a medically important health impact are followed up by the CDC v-safe call center to collect additional information to complete a VAERS report. Information on v-safe is available at https://www.cdc.gov/vsafe.

It is mandatory that vaccination providers administering the Janssen COVID-19 vaccine report all vaccine administration errors (even if not associated with an adverse event) and serious adverse events (irrespective of attribution to vaccination) that occur following vaccination and also report all cases of multisystem inflammatory syndrome (MIS) and COVID-19 that result in hospitalization or death in vaccine recipients to VAERS. Can complete and submit VAERS reports online at https://vaers.hhs.gov/reportevent.html or by faxing to 877-721-0366; include the words “Janssen COVID-19 Vaccine EUA” in description field. If TST or IGRA required according to administrative policies (e.g., healthcare employment, admission to long-term care facilities), perform such testing before or during same visit that COVID-19 vaccine is administered. If such testing cannot be done prior to or at the same time as vaccine administration, ACIP recommends delaying testing until ≥4 weeks after completion of COVID-19 vaccination. If a tuberculosis testing requirement or policy cannot be modified to accept a delay in TST or IGRA testing during the COVID-19 pandemic, it should be understood that a false-negative TST or IGRA cannot be excluded; in such situations, consider repeating a negative TST or IGRA test ≥4 weeks after completion of COVID-19 vaccination. In addition, if TST was performed as the initial test, consider the possibility that boosting could be a factor if results of a repeat TST are positive.

ACIP states COVID-19 vaccines can be given to individuals who have active tuberculosis disease or an illness being evaluated as active tuberculosis disease; however, consider that a moderate or severe acute illness usually is a precaution for vaccination (see Concomitant Illness under Caution). If TST or IGRA is being considered for medical diagnosis of latent tuberculosis infection (e.g., during contact investigation after exposure to contagious tuberculosis disease), a decision to delay such testing until ≥4 weeks after completion of COVID-19 vaccination is at the discretion of the responsible medical provider and local tuberculosis program overseeing the contact investigation. If a decision is made to delay TST or IGRA testing (e.g., in individuals at high risk for progression to tuberculosis disease) and test results are negative, ACIP states consider retesting ≥4 weeks after completion of the vaccination series.

Specific Populations

Pregnancy

Data insufficient to date regarding use of the Janssen COVID-19 vaccine in pregnant women to inform vaccine-associated risks during pregnancy. A developmental toxicity study in female rabbits did not reveal evidence of vaccine-related adverse effects on female fertility, embryofetal development, or postnatal development.

Observational data suggest that, while the absolute risk is low, pregnant women with COVID-19 are at increased risk of severe illness, including illness resulting in admission to an intensive care unit (ICU), mechanical ventilation, extracorporeal membrane oxygenation (ECMO), or death. Additionally, such women are at increased risk of perinatal birth and may be at an increased risk of adverse pregnancy complications or outcomes, such as pre eclampsia, coagulopathy, and stillbirth.

Although data are limited regarding safety of the Janssen COVID-19 vaccine during pregnancy, a different adenovirus-vector vaccine (i.e., Ebola virus vaccine not available in the US) has been used in a large-scale vaccination trial that included pregnant women who were vaccinated during any trimester and no adverse pregnancy-related outcomes, including infant outcomes, were identified that were determined to be related to the vaccine. The Janssen COVID-19 vaccine contains a replication-incompetent adenovirus viral vector and cannot cause SARS-CoV-2 infection in the pregnant woman or her fetus.

FDA states that pregnancy is not a contraindication for use of the Janssen COVID-19 vaccine; pregnant women should discuss their options with their healthcare providers.

ACIP states that pregnant women are eligible for and can receive COVID-19 vaccination and does not state a preference for any specific COVID-19 vaccine in such women. Based on current knowledge, COVID-19 vaccines are unlikely to pose a risk to pregnant women or the fetus. Although potential risks of COVID-19 vaccines administered during pregnancy unknown, clinical trials to evaluate the safety and efficacy of the vaccines in pregnant women are underway or planned.

ACOG recommends that COVID-19 vaccines not be withheld from pregnant women. In the interest of patient autonomy, these experts recommend that pregnant women be free to make their own decision regarding COVID-19 vaccination.

ACIP and ACOG state that pregnant and postpartum women <50 years of age should be informed about the rare risk of TTS after receipt of the Janssen COVID-19 vaccine (see Thrombosis with Thrombocytopenia under Cautions) and the availability of other FDA-authorized COVID-19 vaccines (Moderna COVID-19 vaccine, Pfizer-BioNTech COVID-19 vaccine). ACOG states that women who choose not to receive the Janssen COVID-19 vaccine should be strongly encouraged to receive a different FDA-authorized COVID-19 vaccine.
ACIP and ACOG state that a conversation between the pregnant woman and her clinical team may assist with decisions regarding use of COVID-19 vaccines available under an EUA; however, such a conversation is not required and written permission is not needed prior to vaccination. When making a decision, pregnant women and their healthcare providers should consider the level of COVID-19 transmission in the community, the individual's personal risk of contracting COVID-19, the increased risk of severe COVID-19 in the pregnant woman and potential risks to the fetus, the known and potential benefits of vaccination, efficacy of the vaccine, adverse effects of the vaccine, and limited but growing data about use of the vaccine during pregnancy.

Adverse effects similar to those reported in non-pregnant individuals can occur following COVID-19 vaccination in pregnant women. Advise pregnant women who experience fever following COVID-19 vaccination to take acetaminophen; may also offer acetaminophen as an option for pregnant women experiencing other postvaccination symptoms.

Defer administration of other vaccines (e.g., diphtheria and tetanus toxoids and acellular pertussis vaccine adsorbed [DTaP], influenza vaccine) in pregnant women for 14 days after COVID-19 vaccination. (See Vaccines under Interactions.) ACOG states do not withhold Rh(D) immune globulin when indicated in an individual who is planning to receive or recently received a COVID-19 vaccine. (See Specific Drugs under Interactions.)

Pregnancy exposure registry established to monitor pregnancy outcomes in women exposed to the Janssen COVID-19 vaccine during pregnancy. Encourage women who are vaccinated with the Janssen COVID-19 vaccine during pregnancy to enroll in the registry at https://c-viper.registry.com.

Encourage women who receive a COVID-19 vaccine during pregnancy and those who become pregnant within 30 days after receiving a COVID-19 vaccine to participate in CDC's v-safe program. (See EUA Requirements for Postvaccination Monitoring and Mandatory Vaccine Adverse Event Reporting under Cautions.)

Females and Males of Reproductive Capacity
Routine pregnancy testing not recommended before receiving a COVID-19 vaccine.

ACIP states that women trying to become pregnant do not need to avoid pregnancy after COVID-19 vaccination.

ACOG recommends COVID-19 vaccination for all eligible individuals, including those who may consider future pregnancy.

Unfounded claims linking COVID-19 vaccines to infertility have been scientifically disproven. Because the Janssen COVID-19 vaccine contains a replication-incompetent adenoviral vector that cannot cause infection or alter the DNA of vaccine recipients, it cannot cause infertility.

Lactation
Not known whether the Janssen COVID-19 vaccine administered to a woman who is breastfeeding has any effects on the breast-fed infant or milk production. FDA states breastfeeding is not a contraindication for use of the Janssen COVID-19 vaccine; women who are breastfeeding should discuss their options with their healthcare providers.

ACOG states that FDA-authorized COVID-19 vaccines administered to breastfeeding women cannot cause SARS-CoV-2 infection in women or their infants; therefore, breastfeeding women can receive COVID-19 vaccination. ACIP does not state a preference for any specific COVID-19 vaccine in such women. However, ACIP states that lactating women <50 years of age should be informed about the rare risk of TTS after receipt of the Janssen COVID-19 vaccine (see Thrombosis with Thrombocytopenia under Cautions) and the availability of other FDA-authorized COVID-19 vaccines (Moderna COVID-19 vaccine, Pfizer-BioNTech COVID-19 vaccine).

ACOG states offer COVID-19 vaccines to lactating women, similar to other individuals. ACOG also states that theoretical concerns regarding safety of vaccinating lactating women do not outweigh potential benefits of the vaccine; there is no need for individuals who receive a COVID-19 vaccine to avoid initiating breastfeeding or to discontinue breastfeeding.

Pediatric Use
Safety and efficacy of the Janssen COVID-19 vaccine not assessed in individuals <18 years of age.

FDA EUA permits use of the Janssen COVID-19 vaccine only in individuals ≥18 years of age.

Geriatric Use
Individuals ≥65 years of age included in clinical trials evaluating the Janssen COVID-19 vaccine, and data from such individuals contribute to overall assessment of safety and efficacy of the vaccine.

At the time of FDA's safety and efficacy analysis of data from the ongoing phase 3 trial for the EUA, 19.5% of participants were ≥65 years of age and 3.7% were ≥75 years of age. No overall differences in safety or efficacy observed between those ≥65 years of age and younger adults.

Common Adverse Effects
Solicited local adverse effects reported in vaccine recipients included injection site pain (48.6%), erythema (7.3%), and swelling (5.3%). Onset usually within first 1–2 days after vaccination with median duration of 2 days. However, local adverse effects reported to last >7 days in some vaccine recipients.

Solicited systemic adverse effects reported in vaccine recipients included headache (38.9%), fatigue (38.2%), myalgia (33.2%), nausea (14.2%), and fever (9%). Onset usually within first 1–2 days after vaccination with median duration of 2 days.

Use of antipyretics/analgesics within 7 days following vaccination reported in 19.9% of vaccine recipients versus 5.7% of placebo recipients.

Solicited adverse local and systemic reactions reported more frequently in vaccinees 18–59 years of age than in vaccinees ≥60 years of age.

At the time of FDA's safety analysis of data from the ongoing phase 3 trial for the EUA, serious adverse events (excluding those related to confirmed COVID-19) reported in 0.4% of vaccine recipients and 0.4% of placebo recipients.

Severe allergic reactions, including anaphylaxis, reported rarely. (See Hypersensitivity Reactions under Cautions.)

Interactions

Vaccines
Data not available to date to assess concomitant administration of COVID-19 vaccines, including the Janssen COVID-19 vaccine, with other vaccines.

Although ACIP previously recommended giving COVID-19 vaccines alone, with a minimum interval of 14 days before or after administration of any other vaccines, these experts currently state that COVID-19 vaccines and other vaccines may be administered without regard to timing, including on the same day or within 14 days of each other.

Extensive experience with non-COVID-19 vaccines demonstrated that immunogenicity and adverse event profiles are generally similar whether vaccines are administered concomitantly or alone. However, it is not known whether reactogenicity of COVID-19 vaccines is increased when administered concomitantly with other vaccines, including those known to be more reactogenic (e.g., adjuvanted vaccines, live vaccines). Base decisions to administer a COVID-19 vaccine concomitantly with other vaccine(s) on whether routine immunizations with the other vaccines have been delayed or missed, the individual's risk of vaccine-preventable disease (e.g., during an outbreak or occupational exposures), and reactogenicity profiles of the vaccines.

If a COVID-19 vaccine is administered concomitantly with other vaccines, give each parenteral vaccine at a different injection site and, if possible, separate injection sites by ≥1 inch. ACIP states that, although >1 vaccine can be given IM into the deltoid muscle in adolescents and adults, give COVID-19 vaccines and vaccines likely to cause a local reaction (e.g., tetanus toxoid-containing vaccines, adjuvanted vaccines) in different limbs, if possible.

Specific Drugs

<table>
<thead>
<tr>
<th>Drug</th>
<th>Interaction</th>
<th>Comments</th>
</tr>
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<tbody>
<tr>
<td>Anticoagulants and aspirin</td>
<td>Premedication with an anticoagulant or aspirin not recommended prior to vaccination</td>
<td>Do not need to discontinue or alter dosage of anticoagulants or aspirin prior to COVID-19 vaccination in individuals receiving the drugs routinely</td>
</tr>
<tr>
<td>Antiviral agents</td>
<td>Antiviral agents given at any interval before or after COVID-19 vaccination unlikely to impair development of vaccine-induced protective antibody responses</td>
<td></td>
</tr>
<tr>
<td>COVID-19 convalescent plasma</td>
<td>Data not available; not known whether prior receipt of such antibody therapy interferes with immune response to the vaccine</td>
<td>To avoid potential interference with vaccine immune response, ACIP recommends deferring COVID-19 vaccination for &gt;90 days after such antibody therapy based on estimated half-life of SARS-CoV-2 antibody therapies and evidence suggesting reinfection uncommon in first 90 days after initial infection</td>
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</table>
Hormonal contraceptives (e.g., oral contraceptive, transdermal system, or vaginal ring)

Although certain hormonal contraceptives may increase overall general risk of thrombosis, experts believe such contraceptives do not make individuals more susceptible to TTS after receipt of the Janssen COVID-19 vaccine. Discontinuance or change in hormonal contraceptive methods not needed in women who have received or plan to receive the Janssen COVID-19 vaccine.

Immune globulin and antibody therapies not specific for SARS-CoV-2 (e.g., immune globulin IV [IGIV], Rh[D] immune globulin)

May give COVID-19 vaccine concurrently with or at any interval before or after immune globulin or antibody therapies not specific for SARS-CoV-2.

Immunosuppressive agents (e.g., cancer chemotherapy, corticosteroids, radiation)

Possible decreased or suboptimal antibody responses to vaccines, including the Janssen COVID-19 vaccine. Data insufficient to date to inform optimal timing of COVID-19 vaccination for individuals planning to receive immunosuppressive therapies. ACIP states that individuals receiving immunosuppressive therapy may receive COVID-19 vaccine if they have no contraindications to vaccination. Based on general best practices for vaccination of immunocompromised individuals, ACIP states COVID-19 vaccination should ideally be completed ≥2 weeks before initiation of immunosuppressive therapies; if this is not possible, individuals on immunosuppressive therapy can still receive COVID-19 vaccination; consider individual’s risks related to their underlying condition if making decisions to delay immunosuppressive therapy to complete COVID-19 vaccination. Revaccination after immune competence regained not recommended in individuals who received COVID-19 vaccine during chemotherapy or treatment with other immunosuppressive agents. Corticosteroids given topically or by local injection (e.g., intra-articular, intrabursal).

If COVID-19 subsequently develops in a vaccinated individual, ACIP states prior receipt of COVID-19 vaccine should not affect treatment decisions, including use of corticosteroids, or timing of such treatment. Discontinuance or change in hormonal contraceptive methods not needed in women who have received or plan to receive the Janssen COVID-19 vaccine.

SARS-CoV-2-specific monoclonal antibodies (bamlanivimab and etesevimab, casirivimab and imdevimab)

Data not available; not known whether prior receipt of such antibody therapy interferes with immune response to the vaccine.

Storage

Parenteral

Suspension, for IM Use

Supplied as a suspension in multiple-dose vials that are initially stored frozen by the manufacturer and are then shipped at a temperature of 2–8°C.

After receipt, store multiple-dose vials of the vaccine in a refrigerator at 2–8°C, protect from light, and do not freeze.

If the vaccine is still frozen upon receipt, thaw in a refrigerator at 2–8°C. If the vaccine needs to be used immediately, may thaw appropriate number of vials at room temperature (up to 25°C). If thawing at room temperature is necessary, a carton of 10 vials will take approximately 2 hours to thaw and an individual vial will take approximately 1 hour to thaw. Do not refreeze thawed vaccine.

Consult EUA fact sheet for healthcare providers for the Janssen COVID-19 vaccine and information provided by CDC and the manufacturer for information on storage, handling, and stability of the vaccine. Storage and handling information contained in the EUA fact sheet for healthcare providers supersedes storage and handling information on vaccine vials and carton labels.

After removal from refrigerator, may store unused (i.e., unpunctured) vials for up to 12 hours at room temperature (9–25°C).

After withdrawing first dose of vaccine from a multiple-dose vial, vial may be held in a refrigerator (2–8°C) for up to 6 hours or at room temperature (up to 25°C) for up to 2 hours and must be discarded if not used within these times after first vial entry.

Expiration dates are not marked on vials and cartons of the Janssen COVID-19 vaccine. To avoid potential interference with vaccine immune response, ACIP recommends deferring COVID-19 vaccination for ≥90 days after such antibody therapy based on estimated half-life of SARS-CoV-2 antibody therapies and evidence suggesting reinfection uncommon in first 90 days after initial infection.

If COVID-19 subsequently develops in a vaccinated individual, ACIP states prior receipt of COVID-19 vaccine should not affect treatment decisions, including use of SARS-CoV-2 antibody therapies, or timing of such treatment.

Stability

Suspension, for IM Use

Supplied as a suspension in multiple-dose vials that are initially stored frozen by the manufacturer and are then shipped at a temperature of 2–8°C.

After receipt, store multiple-dose vials of the vaccine in a refrigerator at 2–8°C, protect from light, and do not freeze.

If the vaccine is still frozen upon receipt, thaw in a refrigerator at 2–8°C. If the vaccine needs to be used immediately, may thaw appropriate number of vials at room temperature (up to 25°C). If thawing at room temperature is necessary, a carton of 10 vials will take approximately 2 hours to thaw and an individual vial will take approximately 1 hour to thaw. Do not refreeze thawed vaccine.

Consult EUA fact sheet for healthcare providers for the Janssen COVID-19 vaccine and information provided by CDC and the manufacturer for information on storage, handling, and stability of the vaccine. Storage and handling information contained in the EUA fact sheet for healthcare providers supersedes storage and handling information on vaccine vials and carton labels.

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Expiration dates are not marked on vials and cartons of the Janssen COVID-19 vaccine. To avoid potential interference with vaccine immune response, ACIP recommends deferring COVID-19 vaccination for ≥90 days after such antibody therapy based on estimated half-life of SARS-CoV-2 antibody therapies and evidence suggesting reinfection uncommon in first 90 days after initial infection.

If COVID-19 subsequently develops in a vaccinated individual, ACIP states prior receipt of COVID-19 vaccine should not affect treatment decisions, including use of SARS-CoV-2 antibody therapies, or timing of such treatment.
is possible that expiration dates may be extended as more stability data become available, contact the manufacturer prior to discarding vaccine to determine if the expiration date has been extended.

**Actions**

- Viral vector vaccine composed of a recombinant, replication-incompetent, human adenovirus type 26 (Ad26) vector encoding the SARS-CoV-2 spike (S) protein in a stabilized conformation.
- Following IM injection, the Ad26 vector in the vaccine enters human cells and expresses the S antigen of SARS-CoV-2 (without virus propagation). The S antigen elicits an immune response to provide protection against SARS-CoV-2 infection.
- Data from clinical trial in healthy adults ≥18 years of age indicate that a single dose of Janssen COVID-19 vaccine containing $5 \times 10^{10}$ virus particles (i.e., recombinant Ad26) elicited SARS-CoV-2-neutralizing antibody when tested against wild-type virus and a SARS-CoV-2-S-binding antibody response (detected by day 15 after the dose and increased by day 57). The vaccine dose elicited cellular responses in study participants consistent with a Th-1 phenotype. A second dose of the vaccine given 56 days after the first dose resulted in increased naturalizing antibody titters.
- COVID-19 vaccine (Janssen) available for use under the FDA EUA is provided as a suspension in multiple-dose vials. Each 0.5-mL dose of COVID-19 vaccine (Janssen) contains $5 \times 10^{10}$ virus particles (i.e., recombinant Ad26). Each dose also contains citric acid monohydrate, trisodium citrate dihydrate, ethanol, 2-hydroxypropyl-β-cyclodextrin (HBCD), polysorbate 80, and sodium chloride and may contain residual amounts of host cell proteins and/or host cell DNA.
- Does not contain preservatives; vial stoppers are not made with natural rubber latex.

**Advice to Patients**

- Prior to administration of COVID-19 vaccine (Janssen), the vaccine recipient or their caregiver must be provided with information consistent with the Fact Sheet for Recipients and Caregivers: Emergency Use Authorization (EUA) of the Janssen COVID-19 Vaccine to Prevent Coronavirus Disease 2019 (COVID-19) in Individuals 18 Years of Age or Older and given a copy of the fact sheet or directed to the manufacturer’s website at http://www.janssencovid19vaccine.com to obtain the fact sheet.
- Give the vaccine recipient or their caregiver a vaccination card that provides the name of the vaccine (Janssen COVID-19 vaccine) and the date the vaccine was administered.
- Provide the vaccine recipient or their caregiver with information on, and encourage participation in, CDC’s voluntary smartphone-based tool (v-safe) that uses text messaging and web surveys to check in with individuals who have received a COVID-19 vaccine to identify potential adverse effects; live telephone follow-up is provided if a medically important health impact is reported. Information on v-safe is available at https://www.cdc.gov/vsafe.
- Inform vaccine recipients or their caregivers that FDA authorized the emergency use of the Janssen COVID-19 vaccine, which is an investigational vaccine that has not received FDA approval, for use in individuals ≥18 years of age. Advise them that an ongoing clinical trial has shown that a single dose of the vaccine can prevent COVID-19; however, the duration of protection following vaccination is unknown and the vaccine may not protect everyone who receives it.
- Inform vaccine recipients or their caregivers that the vaccination provider cannot charge them for the vaccine dose, any out-of-pocket vaccine administration fees, or any other fees for COVID-19 vaccination. However, vaccination providers may seek appropriate reimbursement from a program or plan that covers COVID-19 vaccine administration fees for the vaccine recipient (e.g., private insurance, Medicare, Medicaid, US Health Resources & Services Administration [HRSA] COVID-19 uninsured program for non-insured recipients). Individuals who become aware of any potential violations of these requirements are encouraged to report them to the Office of the Inspector General, US Department of Health and Human Services by phone (800-HHS-TIPS) or online (https://tips.oig.hhs.gov).
- Inform vaccine recipients or their caregivers that they have the option to accept or refuse the vaccine.
- Provide vaccine recipients or their caregivers with information on available alternative vaccines and the risks and benefits of those alternatives.
- Inform vaccine recipients or their caregivers about the significant known and potential risks and benefits of the Janssen COVID-19 vaccine, and the extent to which such risks and benefits are unknown. Inform them that local adverse effects (injection site pain, swelling, redness) and systemic adverse effects (headache, fatigue, muscle aches, nausea, fever) have been reported in recipients of the Janssen COVID-19 vaccine.
- Importance of vaccine recipient informing the vaccination provider of any allergies or fever. Advise vaccine recipients or their caregivers that there is a remote chance that the vaccine could cause a severe allergic reaction and such reactions would usually occur within a few minutes to 1 hour after receiving the vaccine dose and may include difficulty breathing, swelling of the face and throat, fast heartbeat, bad rash all over the body, and dizziness and weakness.
- Inform vaccine recipients or their caregivers that blood clots involving blood vessels in the brain, abdomen, and legs along with low platelet counts have occurred rarely in individuals who received the Janssen COVID-19 vaccine and that symptoms began approximately 1–2 weeks following vaccination. Although the chance of this occurring is remote, most individuals who developed these blood clots and low platelet counts were females 18–49 years of age. Advise vaccine recipients or their caregivers to immediately seek medical attention if shortness of breath, chest pain, leg swelling, persistent abdominal pain, severe or persistent headaches or blurred vision, easy bruising, or tiny blood spots under the skin at sites beyond the vaccine injection site occur following vaccination with the Janssen COVID-19 vaccine.
- Importance of vaccine recipient informing the vaccination provider if they previously received any other COVID-19 vaccine, have any medical conditions (e.g., bleeding disorders, immunocompromising diseases), or are receiving anticoagulants or immunosuppressive therapy.
- Importance of women informing clinicians if they are or plan to become pregnant or plan to breast-feed.

**Preparations**

Excipients in commercially available drug preparations may have clinically important effects in some individuals; consult specific product labeling for details.

COVID-19 vaccine (Janssen) is not commercially available. FDA issued an emergency use authorization (EUA) for the Janssen COVID-19 vaccine that permits use of the vaccine for the prevention of COVID-19 in individuals ≥18 years of age. Allocation of the vaccine for use under the EUA is being directed by the US government. The vaccine will be supplied directly from the manufacturer or authorized US distributor(s) and distributed to emergency response stakeholders as directed by the US government, including the CDC and/or other designee.

**COVID-19 Vaccine, Viral Vector (Janssen)**

**Parenteral**

Suspension, for IM use

<table>
<thead>
<tr>
<th>Janssen COVID-19 Vaccine, Janssen</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 × 10^{10} virus particles</td>
</tr>
<tr>
<td>(recombinant Adenovirus type 26)</td>
</tr>
<tr>
<td>per 0.5-mL dose</td>
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* Use is not currently included in the labeling approved by the US Food and Drug Administration. Selected Revisions May 20, 2021, © Copyright, March 15, 2021, American Society of Health-System Pharmacists, Inc.