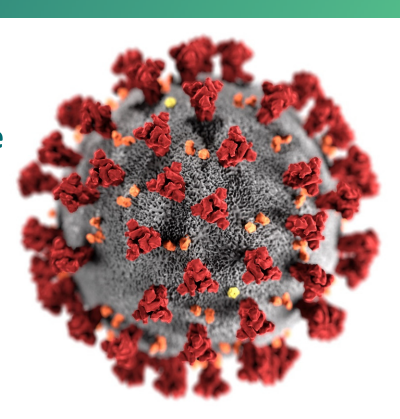


What Clinicians Need to Know About the Pfizer-BioNTech COVID-19 Vaccine

Amanda Cohn, MD Sarah Mbaeyi, MD, MPH

December 13, 2020



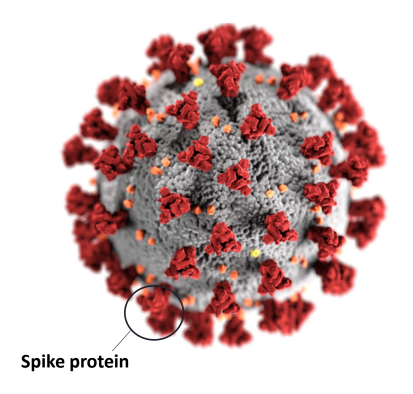


Pfizer-BioNTech COVID-19 Vaccine



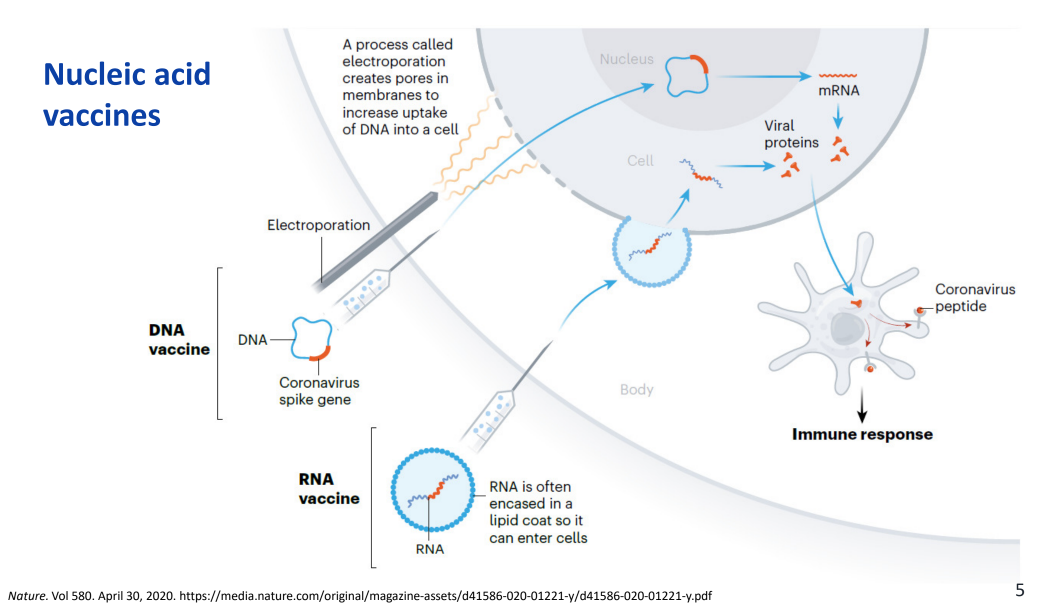
Pfizer-BioNTech COVID-19 vaccine

- Lipid nanoparticle-formulated mRNA vaccine encoding the spike protein
 - Spike protein: facilitates entry of virus into cells
- Vaccination induces antibodies that can block entry of SARS-CoV-2 into cells, thereby preventing infection
- FDA issued an Emergency Use Authorization on December 13, 2020 for use in persons aged ≥16 years



Explaining mRNA COVID-19 vaccines

- mRNA vaccines take advantage of the process that cells use to make proteins in order to trigger an immune response
 - Like all vaccines, COVID-19 mRNA vaccines have been rigorously tested for safety before being authorized for use in the United States
 - mRNA technology is new, but not unknown. They have been studied for more than a decade
 - mRNA vaccines do not contain a live virus and do not carry a risk of causing disease in the vaccinated person
 - mRNA from the vaccine never enters the nucleus of the cell and does not affect or interact with a person's DNA



Advisory Committee (Practices (ACIP) Reco



ACIP recommendations for use of COVID-19 vaccines

- On December 12, 2020, ACIP recommended use of the Pfizer-BioNTech COVID-19 vaccine in persons 16 years of age and older under the FDA's Emergency Use Authorization
- ACIP recommends that when a COVID-19 vaccine is authorized by FDA and recommended by ACIP, that 1) health care personnel and 2) residents of long-term care facilities be offered vaccination in the initial phase of the COVID-19 vaccination program

Vaccine Administration



Administration

- 2-dose series administered intramuscularly 3 weeks apart
- Administration of 2nd dose within 4-day grace period (e.g., day 17-21) considered valid
- If >21 days since 1st dose, 2nd dose should be administered at earliest opportunity (but no doses need to be repeated)
- Both doses are necessary for protection; efficacy of a single dose has not been systematically evaluated

Interchangeability with other COVID-19 vaccine products

- Pfizer-BioNTech COVID-19 vaccine not interchangeable with other COVID-19 vaccine products
 - Safety and efficacy of a mixed series has not been evaluated
- Persons initiating series with Pfizer-BioNTech COVID-19 vaccine should complete series with same product
- If two doses of different mRNA COVID-19 vaccine products inadvertently administered,
 no additional doses of either vaccine recommended at this time
 - Recommendations may be updated as further information becomes available or additional vaccine types authorized

Coadministration with other vaccines

- Pfizer-BioNTech COVID-19 vaccine should be administered alone with a minimum interval of 14 days before or after administration with any other vaccines
 - Due to lack of data on safety and efficacy of the vaccine administered simultaneously with other vaccines
- If Pfizer-BioNTech COVID-19 vaccine is inadvertently administered within 14 days of another vaccine, doses do not need to be repeated for either vaccine

Vaccination of Person CoV-2 Infection or Exp



Persons with a history of SARS-CoV-2 infection

- Vaccination should be offered to persons regardless of history of prior symptomatic or asymptomatic SARS-CoV-2 infection
 - Data from phase 2/3 clinical trials suggest vaccination safe and likely efficacious in these persons
- Viral or serologic testing for acute or prior infection, respectively, is not recommended for the purpose of vaccine decision-making

Persons with known <u>current</u> SARS-CoV-2 infection

- Vaccination should be deferred until recovery from acute illness (if person had symptoms) and <u>criteria</u> have been met to discontinue isolation
- No minimal interval between infection and vaccination
- However, <u>current evidence</u> suggests reinfection uncommon in the 90 days after initial infection, and thus persons with documented acute infection in the preceding 90 days may defer vaccination until the end of this period, if desired

Persons who previously received passive antibody therapy for COVID-19

- Currently no data on safety or efficacy of COVID-19 vaccination in persons who received monoclonal antibodies or convalescent plasma as part of COVID-19 treatment
- Vaccination should be deferred for at least 90 days to avoid interference of the treatment with vaccine-induced immune responses
 - Based on estimated half-life of therapies and evidence suggesting reinfection is uncommon within 90 days of initial infection

Persons with a known SARS-CoV-2 exposure

- Community or outpatient setting:
 - Defer vaccination until <u>quarantine period</u> has ended to avoid exposing healthcare personnel (HCP) or other persons during vaccination visit
- Residents of congregate healthcare settings (e.g., long-term care facilities):
 - May be vaccinated, as likely would not result in additional exposures. HCP are already in close contact with residents and should employ appropriate <u>infection prevention and control</u> <u>procedures</u>
- Residents of other congregate settings (e.g., correctional facilities, homeless shelters)
 - May be vaccinated, in order to avoid delays and missed opportunities for vaccination
 - Where feasible, precautions should be taken to limit mixing of these individuals with other residents or non-essential staff

Vaccination of Special



Persons with underlying medical conditions

- Vaccine may be administered to persons with underlying medical conditions who have no contraindications to vaccination
- Phase 2/3 clinical trials demonstrate similar safety and efficacy profiles in persons with underlying medical conditions, including those that place them at <u>increased risk for</u> <u>severe COVID-19</u>, compared to persons without comorbidities

Immunocompromised persons

- Persons with HIV infection, other immunocompromising conditions, or who take immunosuppressive medications or therapies might be at increased risk for severe COVID-19
- Data not currently available to establish safety and efficacy of vaccine in these groups
- These individuals may still receive COVID-19 vaccine unless otherwise contraindicated
- Individuals should be counseled about:
 - Unknown vaccine safety and efficacy profiles in immunocompromised persons
 - Potential for reduced immune responses
 - Need to continue to follow all current guidance to protect themselves against COVID-19

Pregnant women

- There are no data on the safety of COVID-19 vaccines in pregnant women
 - Animal developmental and reproductive toxicity (DART) studies are ongoing
 - Studies in humans are ongoing and more planned
- mRNA vaccines and pregnancy
 - Not live vaccines
 - They are degraded quickly by normal cellular processes and don't enter the nucleus of the cell
- COVID-19 and pregnancy
 - Increased risk of severe illness (ICU admission, mechanical ventilation and death)
 - Might be an increased risk of adverse pregnancy outcomes, such as preterm birth
- If a woman is part of a group (e.g., healthcare personnel) who is recommended to receive a COVID-19 vaccine and is pregnant, she may choose to be vaccinated. A discussion with her healthcare provider can help her make an informed decision.

Pregnant women

- Considerations for vaccination:
 - level of COVID-19 community transmission (risk of acquisition)
 - her personal risk of contracting COVID-19 (by occupation or other activities)
 - the risks of COVID-19 to her and potential risks to the fetus
 - the efficacy of the vaccine
 - the known side effects of the vaccine
 - the lack of data about the vaccine during pregnancy
- Pregnant women who experience fever following vaccination should be counseled to take acetaminophen as fever has been associated with adverse pregnancy outcomes
- Routine testing for pregnancy prior to receipt of a COVID-19 vaccine is not recommended.

Breastfeeding/Lactating women

- There are no data on the safety of COVID-19 vaccines in lactating women or the effects of mRNA vaccines on the breastfed infant or milk production/excretion
- mRNA vaccines are not considered live virus vaccines and are not thought to be a risk to the breastfeeding infant
- If a lactating woman is part of a group (e.g., healthcare personnel) who is recommended to receive a COVID-19 vaccine, she may choose to be vaccinated

Patient Vaccine Couns



Reactogenicity

- Before vaccination, providers should counsel vaccine recipients about expected local and systemic post-vaccination symptoms
- Unless a person develops a contraindication to vaccination, they should be encouraged to complete the series even if they develop post-vaccination symptoms in order to optimize protection against COVID-19
- Antipyretic or analgesic medications may be taken for treatment of postvaccination symptoms
 - Routine prophylaxis for the purposes of preventing symptoms is not recommended at this time, due to lack of information on impact of use on vaccine-induced antibody responses

Vaccine efficacy

- Two doses required to achieve high efficacy
 - Efficacy after 2nd dose: 95.0% (95% CI: 90.3%, 97.6%)
- Patients should be counseled on importance of completing the 2-dose series in order to optimize protection

Public health recommendations for vaccinated persons

- Protection from vaccine is not immediate; vaccine is a 2-dose series and will take 1 to 2
 weeks following the second dose to be considered fully vaccinated
- No vaccine is 100% effective
- Given the currently limited information on how well the vaccine works in the general population; how much it may reduce disease, severity, or transmission; and how long protection lasts, vaccinated persons should continue to follow all <u>current guidance</u> to protect themselves and others, including:
 - Wearing a mask
 - Staying at least 6 feet away from others
 - Avoiding crowds
 - Washing hands often
 - Following CDC travel guidance
 - Following quarantine guidance after an exposure to someone with COVID-19
 - Following any applicable workplace or school guidance

Contraindications and



Contraindications and precautions

- Package insert:
 - Severe allergic reaction (e.g., anaphylaxis) to any component of the Pfizer-BioNTech COVID-19 vaccine is a contraindication to vaccination
 - Appropriate medical treatment used to manage immediate allergic reactions must be immediately available in the event an acute anaphylactic reaction occurs following administration of the vaccine
- Because of reports of anaphylactic reactions in persons vaccinated outside of clinical trials, the additional following guidance is proposed:
 - A severe allergic reaction to any vaccine or injectable therapy (intramuscular, intravenous, or subcutaneous) is a precaution to vaccination at this time
 - Vaccine providers should observe patients after vaccination to monitor for the occurrence of immediate adverse reactions:
 - Persons with a history of anaphylaxis: 30 minutes
 - All other persons: 15 mins

Algorithm for the triage of persons presenting for Pfizer-**COVID-19** vaccine

PROCEED WITH VACCINATION

PRECAUTION TO VACCINATION

CONTRAINDICATION TO VACCINATION

CONDITIONS

- •Immunocompromising conditions
- Pregnancy
- Lactation

ACTIONS

- •Additional counseling*
- •15-minute observation period

CONDITIONS

•Moderate/severe acute illness

ACTIONS

- •Risk assessment
- •Potential deferral of vaccination
- •15-minute observation period if vaccinated

CONDITIONS

•None

ACTIONS

•N/A

ALLERGIES

- •History of food, pet, insect, venom, environmental, latex, etc., allergies
- •History of allergy to oral medications (including the oral equivalent of an injectable medication)
- •Non-serious allergy to vaccines or other injectables (e.g., no anaphylaxis)
- •Family history of anaphylaxis

ACTIONS

•15-minute observation period

ALLERGIES

- •History of severe allergic reaction (e.g., anaphylaxis) to another vaccine (not including Pfizer-BioNTech vaccine)
- •History of severe allergic reaction (e.g., anaphylaxis) to an injectable medication

ACTIONS:

- Risk assessment
- •Potential deferral of vaccination
- •30-minute observation period if vaccinated

ALLERGIES

•History of severe allergic reaction (e.g., anaphylaxis) to any component of the Pfizer-BioNTech vaccine

ACTIONS

•Do not vaccinate

^{*} See Special Populations section for information on patient counseling in these group

Interpretation of SAR in Vaccinated Persons



SARS-CoV-2 tests

 Viral tests: Prior receipt of the Pfizer-BioNTech COVID-19 vaccine will not affect the results of SARS-CoV-2 nucleic acid amplification or antigen tests

Antibody tests:

- Currently available antibody tests for SARS-CoV-2 assess IgM and/or IgG to spike or nucleocapsid proteins
- Pfizer-BioNTech COVID-19 vaccine contains mRNA that encodes the spike protein; thus, a
 positive test for spike protein IgM/IgG could indicate either prior infection or vaccination
- To evaluate for evidence of prior infection in an individual with a history of Pfizer-BioNTech COVID-19 vaccination, a <u>test</u> specifically evaluating IgM/IgG to the nucleocapsid protein should be used

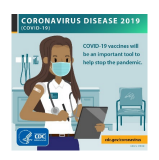
Clinical Resources



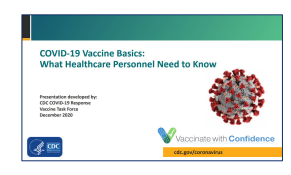
COVID-19 vaccine communication resources

- Engaging in Effective COVID-19
 Vaccine Conversations
 - https://www.cdc.gov/vaccines/covid-19/hcp/engaging-patients.htm
- Toolkit for Medical Centers, Clinics, and Clinicians
 - https://www.cdc.gov/vaccines/covid-19/healthsystems-communication-toolkit.html
- More toolkits coming soon
 - Long-term care facilities
 - Health departments
 - Community-based organizations
 - Employers of essential workers









Infection prevention and control recommendations for persons with post-vaccination symptoms

Healthcare personnel

Long-term care facility residents

Infection prevention and control considerations for residents of long-term care facilities with systemic signs and symptoms following COVID-19 vaccination

Note: Strategies are needed by long-term care facilities to appropriately evaluate and manage postvaccination signs and symptoms among their residents. The approach described in this document is intended to balance:

Infection prevention and control considerations for healthcare personnel with systemic signs and symptoms following COVID-19 vaccination

Note: Strategies are needed for healthcare facilities to appropriately evaluate and manage postvaccination signs and symptoms among healthcare personnel (HCP). The approach described in this document is intended to reduce the risks for disruptions in care and pathogen (e.g., SARS-CoV-2) transmission resulting from:

- · unnecessarily excluding HCP with only post-vaccination signs and symptoms from work, and
- inadvertently allowing HCP with SARS-CoV-2 or another transmissible infection to work.

These considerations are based on the current understanding of signs and symptoms following COVID-19 vaccination, including timing and duration, and might change as experience with the vaccine accumulates.

Overview

Systemic signs and symptoms, such as fever, fatigue, headache, chills, myalgia, and arthralgia, can occur following COVID-19 vaccination. Preliminary data from mRNA COVID-19 vaccine trials indicate that most systemic post-vaccination signs and symptoms are mild to moderate in severity, occur within the first three days of vaccination (the day of vaccination and following two days, with most occurring the day after vaccination), resolve within 1-2 days of onset, and are more frequent and severe following the second dose and among younger persons compared to those who are older (>55 years). Cough, shortness of breath, rhinorrhea, sore throat, or loss of taste or smell are **not** consistent with post-vaccination symptoms, and instead may be symptoms of SARS-CoV-2 or another infection.

Because systemic post-vaccination signs and symptoms might be challenging to distinguish from signs and symptoms of COVID-19 or other infectious diseases, HCP with postvaccination signs and symptoms

Based Precautions for f

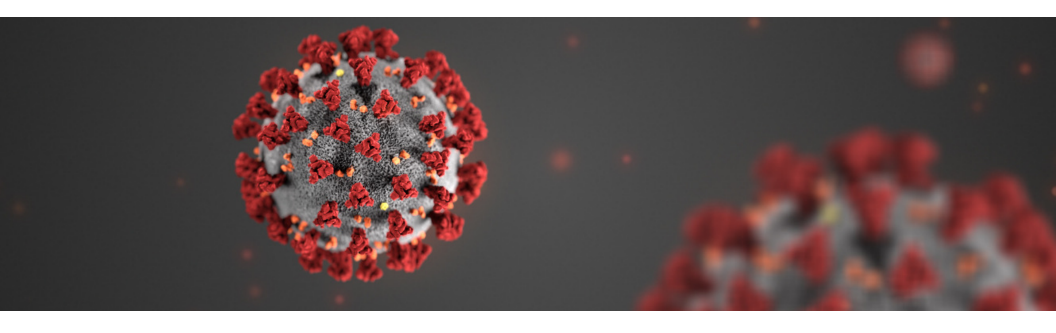
f
r transmissible infectious

applied to patients in other Inding of signs and Ind might change as

ia, and arthralgia, can occur ine trials indicate that most ity, occur within the first h most occurring the day nd severe following the (>55 years). Cough, consistent with post-

FDA EUA resources

- FDA COVID-19 EUA
 - https://www.fda.gov/media/144412/download
- FDA COVID-19 Information
 - https://www.fda.gov/emergency-preparedness-and-response/counterterrorism-and-emergingthreats/coronavirus-disease-2019-covid-19
- FDA EUA Guidance
 - https://www.fda.gov/emergency-preparedness-and-response/mcm-legal-regulatory-and-policy-framework/emergency-use-authorization#covid19euas



For more information, contact CDC 1-800-CDC-INFO (232-4636) TTY: 1-888-232-6348 www.cdc.gov

Thank you

The findings and conclusions in this report are those of the authors and do not necessarily represent the official position of the Centers for Disease Control and Prevention.

