Vaccine Safety in the United States: Overview and Considerations for COVID-19 Vaccines

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Widespread immunization efforts have been linked to increased life expectancy and reduced illness. U.S. vaccination programs, headed by the Centers for Disease Control and Prevention (CDC) within the Department of Health and Human Services (HHS), have helped eradicate smallpox and nearly eradicate polio globally, and eliminate several infectious diseases domestically. With the Coronavirus Disease 2019 (COVID-19) pandemic now causing major health and economic impacts across the world, efforts are underway to make safe and effective vaccines available quickly to help curb spread of the virus.

Background

Federal regulation of vaccine safety began with the Biologics Control Act of 1902, which was the first federal law to require premarket review of pharmaceutical products. Since the 1902 law was enacted, federal vaccine safety activities have expanded, with the aim of minimizing the possibility of adverse events following vaccination and detecting new adverse events as quickly as possible. Today, as covered in this report, federal efforts to ensure vaccine safety include the following activities:

- **Premarket requirements**: Clinical trials, or testing of investigational vaccines in human subjects, and U.S. Food and Drug Administration (FDA) licensure or authorization.
- **Clinical recommendations**: Recommendations for the clinical use of vaccines by the Advisory Committee on Immunization Practices (ACIP), and CDC clinical guidance and resources.
- **Postmarket safety**: Manufacturing requirements and ongoing safety monitoring of vaccines administered to patients.
- **Federal research on vaccine safety**: Ongoing research to inform a better scientific understanding of vaccine safety and comprehensive scientific reviews on the safety of vaccines in use.
- **Vaccine injury compensation**: In nonemergency circumstances, the National Vaccine Injury Compensation Program (VICP) provides compensation to eligible individuals found to have been injured by a covered vaccine. In emergency circumstances, like COVID-19, a separate Countermeasures Injury Compensation Program (CICP) may be used.
- **Vaccine distribution**: Programs and requirements to ensure safety controls in vaccine distribution programs, led by CDC.

**COVID-19 Vaccine Safety Considerations**

Safety considerations for COVID-19 vaccines in development are unique in many ways. FDA has never licensed a vaccine for a coronavirus, and much remains unknown about potential safety issues related to COVID-19 vaccines. Under Operation Warp Speed (OWS)—the Trump Administration’s major medical countermeasure development initiative—COVID-19 vaccines are under an expedited development timeline. FDA may initially make the vaccine available under an Emergency Use Authorization (EUA) instead of its standard biologics licensing process—a first for the agency for a previously unapproved vaccine. In light of reported concerns from the public surrounding the safety and effectiveness of COVID-19 vaccines developed on an expedited timeline, FDA officials have sought to clarify that any vaccine candidate “will be reviewed according to the established legal and regulatory standards for medical products.” If made available within the next several months, available safety and effectiveness data would be based on months of data collection rather than on years of data collection typically used in vaccine development. In addition, efforts are underway with regard to (1) clinical guidance and prioritization of individuals to receive the likely limited initial supply of COVID-19 vaccines; (2) strengthening safety monitoring systems to collect ongoing safety surveillance data on vaccines administered to the population; and (3) preparing for safety controls in vaccine distribution and patient administration, in addition to other activities.

**Congressional Considerations**

Ever since the Biologics Control Act of 1902, Congress and the Administration (especially through FDA and CDC) have strived to ensure the safety of vaccines in the United States—from initial development to patient administration. Congress may consider how to best leverage existing requirements and programs to ensure that risk of harm from eventual COVID-19...
vaccines is mitigated and minimized. OWS, FDA, CDC, and others are working to expedite the availability of COVID-19 vaccines and to prepare for a nationwide immunization campaign. Safety has been cited as a consideration in all of these efforts. Congress may consider how to best provide oversight and make legislative changes to ensure a safe and successful COVID-19 vaccination campaign. In addition, Congress may consider and evaluate the entire federal vaccine safety system and assess whether this system warrants any policy changes to help ensure ongoing safety of all recommended vaccines.
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Introduction

Widespread immunization efforts have been linked to increased life expectancy and reduced illness.¹ In 1900, for every 1,000 babies born in the United States, 100 would die before their first birthday, often due to infectious diseases.² One study estimated that from 1993 to 2013, routine childhood immunization in the United States helped prevent 322 million illnesses, 21 million hospitalizations, and 732,000 premature deaths.³ U.S. immunization programs, headed by the Centers for Disease Control and Prevention (CDC) within the Department of Health and Human Services (HHS), have helped eradicate smallpox and nearly eradicate polio globally.⁴ U.S. immunization programs have also helped eliminate measles and rubella domestically, and have led to substantial reductions in hospitalizations linked to pneumococcus, rotavirus, and varicella (i.e., chickenpox).⁵ With the Coronavirus Disease 2019 (COVID-19) pandemic now causing major health and economic impacts across the world, efforts are underway to make safe and effective vaccines available quickly to help curb spread of the virus.

Available evidence from thousands of scientific studies shows that currently recommended vaccines are largely safe. At a population level, widespread vaccination with recommended vaccines is safer than the spread of the infectious diseases they prevent.⁶ Adverse health events for which available scientific evidence shows a causal relationship with currently recommended vaccines are rare—ranging from 1 case per million doses administered (e.g., encephalitis caused by the pertussis vaccine) to 333 cases per million doses (e.g., febrile seizures caused by the measles-mumps-rubella; MMR vaccine).⁷

Undervaccination linked to concerns about vaccine safety has been an issue in recent years. U.S. outbreaks of measles in 2019—the highest number of annual measles cases since 1992—were driven in part by geographic clusters with low vaccination rates for the MMR vaccine.⁸ U.S. surveys show that concerns about vaccine safety are a top reason for vaccine delays or refusals.⁹

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From a public health perspective, vaccines for infectious diseases often work by helping provide *herd immunity*, meaning that enough of the population has vaccine-induced immunity against the target disease to curb ongoing transmission and protect those who cannot receive vaccines (e.g., persons with compromised immune systems). Widespread vaccination can help with achieving *elimination* or *eradication* of a given disease (see text box). To effectively prevent disease spread, many vaccines must be administered to a large segment of the population. Public health practice generally aims for near 100% vaccination rates among populations recommended to receive vaccines, though the level required for herd immunity is generally lower and can vary by vaccine and population (75%-95% of the population). Nonetheless, widespread vaccination that does not meet target rates can aid in significantly curbing disease spread.

Vaccines are generally held to a higher safety standard than most other medical products for many reasons. For one, vaccines are often administered to healthy individuals to prevent disease; therefore, the expectation is that such individuals will remain healthy following vaccination. Moreover, drugs administered to healthy people are expected to have fewer side effects than drugs that treat disease, such as those for cancer or heart disease, mainly because the expected benefits differ. In addition, vaccines are often administered to vulnerable populations, including infants and pregnant women. Also, since vaccines are often mandated by state and sometimes federal law for certain groups (e.g., school children and military service members), the government has an interest in ensuring that vaccines are as safe as possible. Because vaccines are often administered to a large segment of the population, even a rare risk of adverse reactions to a vaccine could affect a sizeable number of people.

### Scope of This Report

This report provides an overview of the federal government’s role in ensuring safety of vaccines for infectious diseases. Specifically, this report

- describes federal statutory and regulatory requirements and administrative functions governing vaccine licensure (including pre- and post-licensure safety), development of clinical recommendations, and vaccine injury compensation;

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summarizes ongoing federal activities related to vaccine post-licensure safety (e.g., ongoing safety monitoring and research), as well as safety assurances in federal vaccine distribution programs; and

• discusses safety considerations in the context of developing and making available vaccine(s) for COVID-19.

This report does not provide a comprehensive scientific review on the safety of existing vaccines, nor does it specifically address vaccines for noninfectious diseases (e.g., cancer). A discussion of payment and coverage for vaccines and related health care services is outside the scope of this report.

What Is a Vaccine?

A vaccine is a biological preparation that contains small amounts of weak, dead, or modified disease-causing agents known as antigens, which can include viruses, bacteria, fractions of these agents, or the toxins they produce. Once introduced to the body, the antigen elicits a response by the immune system creating antibodies and immune memory cells that prevent future infection from the same disease. The immune response from a vaccine is similar to the immune response from acquiring an infectious disease naturally; however, since the antigen in the vaccine is weakened or dead, the vaccine usually does not cause disease. In the case of vaccines made with weakened live attenuated viruses or bacteria, the vaccine may cause a form of the disease that is usually much milder than the actual disease. In addition, the immune response triggered by any vaccine may cause some symptoms in some patients.  

Along with the antigen, vaccines contain other ingredients such as preservatives, stabilizers, and adjuvants. Preservatives, like thimerosal, can help keep the vaccine free of contamination by other germs (e.g., bacteria, fungi). Thimerosal is currently used only in multidose vials of vaccines, such as certain formulations of the influenza (flu) vaccine. Stabilizers, like sugar or gelatin, allow the vaccine to be stored for a period of time and help keep the antigen stable. Adjuvants, such as aluminum salts, help trigger the immune response to the vaccine, particularly for vaccines made with fractions of disease-causing agents. Vaccines may also contain small amounts of residual material from the manufacturing process, such as egg proteins, formaldehyde, and antibiotics.

Federal Vaccine Safety Regulation and Programs

Federal regulation of vaccine safety began with the Biologics Control Act of 1902, which was the first federal law to require premarket review of pharmaceutical products. The Biologics Control Act was enacted in response to deaths (many of them children) from tetanus contamination of smallpox vaccine and diphtheria antitoxin (a prophylaxis used for diphtheria at the time). The act


imposed requirements on the manufacturing and labeling of biological products (“biologics”) and
required inspection of manufacturing facilities before a federal license was issued for marketing
the products. The Biologics Control Act was revised and recodified when the Public Health
Service Act (PHSA) was enacted in 1944. Biologics are now subject to regulation by the U.S.
Food and Drug Administration (FDA) under the PHSA and the Federal Food, Drug, and Cosmetic
Act (FFDCA).\textsuperscript{17}

Since the 1902 law was enacted, federal vaccine safety activities have expanded to minimize the
possibility of adverse events following vaccination (such as by vaccine contamination) and to
detect new adverse events as quickly as possible, as discussed throughout this report. Major reforms to federal
vaccine safety programs were enacted as a part of the National Childhood Vaccine Injury Act of 1986 (NCVIA;
P.L. 99-660, Title III), which mandated reporting of adverse events caused by
vaccines to FDA and CDC, established the National Vaccine Program Office (NVPO) within HHS to coordinate
federal vaccine efforts, granted FDA mandatory recall authority for biological products, and established the
National Vaccine Injury Compensation Program (VICP). NCVIA was enacted after a spate of lawsuits against vaccine
manufacturers alleging safety issues. The lawsuits caused several vaccine manufacturers to exit the market,
leading to concerns about the vaccine supply and possible reintroduction of
certain diseases.\textsuperscript{18}

As covered in this report, efforts to ensure vaccine safety include several federal activities:

- **Premarket requirements:** Clinical trials and FDA licensure or authorization.
- **Clinical recommendations:** Recommendations for the safe and appropriate
clinical use of vaccines by the Advisory Committee on Immunization Practices
(ACIP), and CDC clinical guidance and resources.

\textbf{Federal Agencies Involved in Vaccine Safety}

Within the Department of Health and Human Services (HHS):

- FDA regulates the safety, effectiveness, and quality of vaccines through premarket review and postmarket
requirements (e.g., adverse event reporting).
- CDC supports cross-cutting immunization programs that include, as relevant to vaccine safety: safety monitoring,
clinical guidance for vaccines, vaccine safety research, and
efforts to ensure safety in public vaccine distribution.
- The National Institutes of Health (NIH) is the primary federal agency that supports medical and health research,
including vaccine research.
- The Centers for Medicare & Medicaid Services (CMS) monitors vaccine safety among the Medicare population.
- The Agency for Healthcare Research and Quality (AHRQ) conducts vaccine safety reviews.
- The Health Resources and Services Administration (HRSA) administers the VICP.

The Department of Veterans Affairs (VA) conducts some vaccine research and monitors vaccine safety among veterans
who receive care in the VA system.

The Department of Defense (DOD) conducts some vaccine research and has a database for monitoring adverse events from
vaccination among military service members and their families.

\textsuperscript{17} Until 1972, biologics, including vaccines, were regulated by the National Institutes of Health (NIH, or its precursors)
under the Biologics Control Act of 1902. In 1972, regulatory responsibility over biologics was transferred from NIH to
the U.S. Food and Drug Administration (FDA). See David M. Dudzinski, “Reflections on Historical, Scientific, and
Legal Issues Relevant to Designing Approval Pathways for Generic Versions of Recombinant Protein-Based
Therapeutics and Monoclonal Antibodies,” Food and Drug Law Journal, 2005, vol. 60, no. 2, pp. 143-260. See also
CRS Report R44620, Biologics and Biosimilars: Background and Key Issues.

\textsuperscript{18} Geoffrey Evans, “Update on Vaccine Liability in the United States: Presentation at the National Vaccine Program
Office on Strengthening the Supply of Routinely Recommended Vaccines in the United States, 12 February 2002,”
1655-1658.
- **Postmarket safety**: Manufacturing requirements and ongoing safety monitoring of vaccines administered to patients.

- **Federal research on vaccine safety**: Ongoing research to inform a better scientific understanding of vaccine safety, and comprehensive scientific reviews on the safety of vaccines.

- **Vaccine injury compensation**: In nonemergency circumstances, the VICP can provide compensation to eligible individuals found to have been injured by a covered vaccine.

- **Vaccine distribution**: Programs and requirements to ensure safety controls in vaccine distribution programs, led by CDC.

### Vaccine Safety Basics

As defined by FDA regulations, safety is “the relative freedom from harmful effect to persons affected, directly or indirectly, by a product when prudently administered, taking into consideration the character of the product in relation to the condition of the recipient at the time.”

Vaccine safety is distinct from efficacy and effectiveness; however, it is useful to consider vaccine safety in the context of efficacy and effectiveness, which are defined as follows:

- **Vaccine efficacy** is defined as the reduction in disease incidence in a vaccinated group compared with an unvaccinated group under optimal conditions (i.e., healthy individuals and proper administration).

- **Vaccine effectiveness** is defined as the reduction in disease incidence in a vaccinated group compared with an unvaccinated group under real-world conditions.

Like all pharmaceutical products, vaccines are not 100% safe for all patients. Vaccine safety programs continually assess the benefits and risks of vaccination. Adverse events following vaccination can be classified in many ways:

- **Frequency**—is the adverse event common or rare?

- **Severity**—is the adverse event mild, such as minor pain or swelling, or severe, such as leading to hospitalization, disability, or death?

- **Causality**—can a causal relationship be established with the vaccine with clinical, laboratory, or epidemiologic evidence? (see *text box* below)

- **Preventability**—is the adverse event intrinsic to the vaccine (i.e., provoked by the immune response caused by the vaccine), or related to faulty production or administration of the vaccine?

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19 C.F.R. §600.3(p).


Some adverse events following vaccination may be linked directly to the antigen in the vaccine, such as paralytic poliomyelitis (i.e., paralysis), which is rarely caused by the live oral polio vaccine. Other adverse events are precipitated by the vaccine, such as febrile seizures that occur following a vaccine-induced fever. Some adverse events can be linked to improper vaccine administration; for example, a vaccine administered too high on the arm of an adult can cause deltoid bursitis (inflammation of the shoulder joint). In the past, improper vaccine manufacturing has been tied to large-scale adverse health events. In 1955, one polio vaccine manufacturer failed to completely inactivate the poliovirus in the manufacturing process. As a result, 40,000 people developed mild polio from the vaccine, 200 became paralyzed, and 10 died.

In some cases, establishing a causal connection between a vaccine and an adverse event is difficult. Vaccination may co-occur with an adverse health event. For example, early childhood—a time when several recommended pediatric vaccines are typically administered—coincides with the same period when signs and symptoms of developmental disorders, such as autism, may begin to appear. Available evidence rejects a causal relationship between childhood vaccines and autism. To determine causality between a vaccine and a given health event, scientists and public health experts evaluate many kinds of evidence, including the time period between vaccination and the event; the biologic plausibility that the health event was caused by vaccination; clinical or laboratory evidence that supports causation by the vaccine; and population-based epidemiological analyses that assess whether vaccinated individuals are more likely to develop a certain health outcome within a certain time period following vaccination compared to individuals who did not receive the vaccine in that time period. Several of the programs covered in this report generate data or other evidence that can allow for causality assessments to link certain adverse events with vaccination (see text box).

### What Is a Causality Assessment?

Immune systems are arguably among the most complex biological systems—therefore, studying vaccines and their effect on the human body can be difficult. Individual studies may provide suggestive evidence of adverse health effects linked to vaccines. For example, an analysis of health data on a population of thousands of individuals could find that vaccination with a certain vaccine is statistically associated with higher rates of a certain adverse health event that occurred following vaccination. Yet, another similar study could conduct a similar analysis among a different population and find no such evidence. In addition, further evidence based on the research in the laboratory, such as with animals or human tissue samples, might find that a certain adverse event following vaccination is or is not likely based on an understanding of biological systems. Therefore, in order to determine if all the available evidence favors a causal relationship between a vaccine and a subsequent adverse health event, researchers will combine evidence across many types of studies as a part of a causality assessment. Good quality systematic causality assessments usually include the following attributes:

- Search methods to identify all possible studies of interest within all relevant areas of research.
- A selection process to determine which studies are actually relevant and used rigorous scientific methods that provide quality evidence based on defined criteria.

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- A review process to compare evidence across studies, considering differences such as study populations, study design, and the quality of each study.
- Methods to weigh different types of evidence and combine evidence across studies in order to determine whether all the evidence, in total, supports or does not support a causal relationship between vaccination with a specific vaccine and a subsequent adverse event, or yields inconclusive results.

For a further discussion, see the “Federal Research on Vaccine Safety” section. Causality assessments may also be conducted on an ongoing basis using data and information from postmarket monitoring systems (see the “Postmarket Safety” section).


Premarket Safety

Vaccines generally follow the same clinical development and approval process as drugs and other biologics (i.e., therapeutics derived from living organisms).\(^{27}\) To be marketed in the United States, a new vaccine must first receive licensure (i.e., approval) from FDA. Licensure is based on a determination by FDA that the vaccine and the facility in which it is manufactured, processed, packed, or held meet standards to ensure that the product is safe, pure, and potent (effective).\(^{28}\) Except under very limited circumstances, FDA requires data from clinical trials—formally designed, conducted, and analyzed studies of human subjects—to provide evidence of a vaccine’s safety and effectiveness. These requirements apply to all vaccines marketed in the United States, regardless of whether the manufacturing facility is located domestically or in a foreign country.

Clinical Trials

Vaccines are typically tested in several stages of human clinical trials. Before beginning clinical testing, a vaccine’s sponsor must file an investigational new drug (IND) application, which is a request for FDA authorization to administer an investigational biologic (or drug) to humans.\(^{29}\) The IND must include information about the proposed clinical study design, completed animal test data, and the lead investigator’s qualifications.\(^{30}\) The investigator also must provide assurance that an Institutional Review Board (IRB) will provide initial and continuous review and approval of each of the studies in the clinical investigation to ensure that participants are aware of the drug’s investigational status, and that any risk of harm will be necessary, explained, and

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\(^{27}\) Biological products include vaccines, monoclonal antibodies, and cytokines, among other examples. For additional information about biologics, see CRS Report R44620, Biologics and Biosimilars: Background and Key Issues.

\(^{28}\) PHSA §351(a)(2)(C) [42 U.S.C. §262(a)(2)(C)]. FDA approves drugs that are safe and effective; the equivalent terminology for biologics is safe, pure, and potent. FDA has interpreted potency to include effectiveness. See the FDA Guidance for Industry, Scientific Considerations in Demonstrating Biosimilarity to a Reference Product, https://www.fda.gov/media/82647/download.


\(^{30}\) 21 C.F.R. 312 Subpart B.
minimized. FDA has 30 days to review an IND, after which a manufacturer may begin clinical testing if FDA has not objected and imposed a clinical hold.

Clinical trials for an IND may be sponsored by the drug company seeking to commercially market the vaccine, a university or nonprofit organization, a government agency, or a combination or partnership of all the above. The funder(s) may differ for each stage of testing. In typical circumstances, the public sector (e.g., federal agencies, nonprofit organizations) generally finances more of the earlier stages of clinical trials, such as Phase 1 clinical trials. Later-stage testing, such as Phase 3 clinical trials, are typically funded more so by drug companies than government agencies.

The sponsor of the trial is responsible for selecting qualified investigators, maintaining an effective IND, and ensuring proper monitoring of the investigations, including that they are conducted in accordance with the IND. In certain cases, the sponsor may establish an independent Data and Safety Monitoring Board (DSMB) of relevant experts with no relevant financial or other ties to the sponsor to oversee the investigations. The DSMB often advises the sponsor on the ongoing safety of trial subjects and the continuing validity and scientific merit of the trial. One DSMB may be responsible for overseeing multiple clinical trials.

In general, vaccine clinical trials occur in three sequential phases:

- **Phase 1** trials are the first in-human studies of a vaccine candidate, and they assess safety and immunogenicity in a small number of volunteers.
- **Phase 2** trials assess side effects and the dosing at which the investigational vaccine may have a protective effect and may enroll hundreds of volunteers.
- **Phase 3** trials assess effectiveness and continue to monitor safety and typically enroll thousands of volunteers.

Most clinical trials for vaccines include a control group, such as a placebo or alternative vaccine, to compare outcomes for those who received the target vaccine compared with those who did not. Phase 3 clinical trial data are typically needed to fully assess the safety and effectiveness of an investigational vaccine. Typically, only the Phase 3 clinical trials are large enough to allow for robust scientific evidence on the safety and effectiveness of the investigational vaccine among different population segments (e.g., children, older adults). Under typical circumstances, a vaccine candidate moves through each phase of clinical testing upon successful completion of the prior phase.

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31 21 C.F.R. §312.23(a)(1)(iv) and 21 C.F.R. Part 56.
34 *Immunogenicity* refers to the extent to which a substance is able to stimulate an immune response. An immune response to a pharmaceutical product may affect its safety and effectiveness. See Jonathan Law and Elizabeth Martin, ed., *Concise Medical Dictionary* (Oxford University Press).
Phase 3 clinical trials are typically longer (usually at least a year) than the other phases in order to fully assess the safety and effectiveness of the investigational vaccine. Adequate time is often needed to give trial participants a chance to be exposed to the target disease in the community and to assess if infection rates vary between the vaccine recipient and control groups. In some cases, an experimental vaccine that showed promise in Phase 1 and Phase 2 clinical trials was found to be ineffective in Phase 3 trials. For example, an experimental vaccine for herpes simplex virus type 2 (HSV-2) showed safety and preliminary evidence of an immune response to the virus in Phase 2 clinical trials (i.e., HSV-2 antibodies in the bloodstream). However, during the Phase 3 clinical trials, by a year after vaccination, there was no difference in rates of acquired HSV-2 infections between the recipient and control groups, despite vaccine recipients showing a preliminary immune response.37

In addition to providing insights into the effectiveness of investigational vaccines, long-term Phase 3 studies can uncover important safety data. For example, three years of safety data on the vaccine for dengue virus produced by Sanofi Pasteur (Dengvaxia) found an issue of antibody-mediated enhancement of infections, where the antibodies raised in response to vaccination could worsen the severity of dengue for those without a prior dengue infection. Data on the vaccine showed a higher rate of hospitalizations for dengue three years after vaccination in young children compared with children who were unvaccinated.38

For some vaccines, Phase 3 clinical trials are very large to detect rare adverse events. For instance, two second-generation rotavirus vaccines (RotaTeq and RotaRix) were subject to Phase 3 clinical trials involving over 60,000 infants in order to ascertain the risk of intussusception (intestinal obstruction) following vaccine administration (estimated to be about 1 in 10,000 in the first-generation vaccine).39 However, such large trials involve higher costs and increased time to licensure.

**Biologics License Application (BLA) and Licensure Requirements**

After completing clinical trials, a sponsor may submit a Biologics License Application (BLA) to FDA’s Center for Biologics Evaluation and Research (CBER). A BLA is a request for permission to market the vaccine and must contain certain information, including data from nonclinical laboratory and clinical studies demonstrating that the product meets requirements of safety, purity, and potency.40 For each nonclinical laboratory study, the BLA must include either (1) a statement that the study was conducted in compliance with FDA regulations governing Good Laboratory Practice (GLP) for nonclinical laboratory studies41 or (2) if the study was not conducted in compliance with GLP regulations, a brief statement explaining the reason for noncompliance. In addition, for each clinical investigation involving human subjects, the BLA must contain statements that each clinical investigation either was conducted in compliance with

37 FDA, 22 Case Studies Where Phase 2 and Phase 3 Trials had Divergent Results, January 2017.
40 FDA regulations at 21 C.F.R. §601.2(a) specify the required contents of a BLA.
41 21 C.F.R. Part 58 “Good Laboratory Practice for Nonclinical Laboratory Studies.”
the requirements for institutional review set forth in FDA regulations, or that it was not subject to such requirements and was conducted in compliance with requirements for informed consent. The BLA also must contain “a full description of manufacturing methods; data establishing stability of the product through the dating period; sample(s) representative of the product for introduction or delivery for introduction into interstate commerce; summaries of results of tests performed on the lot(s) represented by the submitted sample(s); specimens of the labels, enclosures, and containers;” and the address of each location involved in the manufacture of the vaccine. If applicable, a BLA must contain any medication guide proposed to be used for the product. Finally, the BLA must include a financial certification or disclosure statement(s) or both for clinical investigators.

As noted above, a vaccine manufacturer must submit proposed vaccine labeling as part of a BLA. FDA reviews the proposed labeling to determine whether it is scientifically accurate and that it conforms to regulatory requirements. As for prescription drugs and other biologics, vaccine labeling must include warnings and precautions, contraindications, dosage and administration, storage and handling conditions, and adverse reactions, among other information. Labeling for vaccines must specifically contain a statement describing how suspected adverse reactions can be reported. In addition, the labels affixed to each container or package of a vaccine must include the name of the manufacturer, the lot number or other lot identification, and the recommended individual dose (for multiple dose containers), among other information. Vaccines require special processing and handling, such as refrigeration and proper storage, and information about storage temperature and other handling instructions must be on the label affixed to each package containing a vaccine.

FDA regulations also provide for biological product manufacturing establishment standards. Such standards cover personnel, the physical establishment in which a product is manufactured, records maintenance, retention of samples, reporting of product deviations, and product temperature during shipment. Most of these requirements apply broadly to biologics, but several provisions are vaccine-specific, including requirements for live vaccine work areas and live vaccine processing, as well as product-specific maintenance temperatures. In addition, FDA regulations establish requirements for testing product potency, sterility, purity, and identity, as well as requirements for constituent materials used in licensed products, including preservatives, diluents, and adjuvants. Vaccines, like other biological products, are subject to lot release requirements, which provide that “[n]o lot of any licensed product shall be released by the manufacturer prior to the completion of tests for conformity with standards applicable to such

43 21 C.F.R. Part 50 “Protection of Human Subjects.”
44 21 C.F.R. §§201.56 and 201.57.
45 21 C.F.R. §201.57(a)(11)(iii).
46 “Lot” refers to “that quantity of uniform material identified by the manufacturer as having been thoroughly mixed in a single vessel.” 21 C.F.R. § 600.3(x).
47 21 C.F.R. §§610.60 and 610.61.
48 21 C.F.R. §610.61.
49 21 C.F.R. Part 600.
50 21 C.F.R. §600.10(c)(4).
51 21 C.F.R. §600.11(c)(4).
52 21 C.F.R. §600.15.
53 21 C.F.R. Part 610.
product.”54 FDA may require that samples of any lot of any licensed product and the protocols and applicable test results be submitted to CBER. In such case, a manufacturer may not distribute a lot of a vaccine until it is released by FDA.55

**Expedited Pathways and Access to Unapproved Vaccines**

Because clinical testing and the FDA review process typically take several years, FDA and Congress have established mechanisms to expedite the premarket development and review processes for pharmaceutical products, including vaccines, coming onto the market, as well as to expand access to products that are still under investigation. Historically, certain FDA expedited pathways such as Emergency Use Authorization (EUA) have been used infrequently for vaccines. However, a public health emergency, such as a pandemic, may affect the risk assessment in making a vaccine available before full long-term safety data are available.

**Expedited Development and Review**

To address unmet medical needs in the treatment or prevention of serious or life-threatening diseases or conditions, FDA can expedite the development and review processes for drugs and biologics, including vaccines, through four programs:

- fast track product designation,
- breakthrough therapy designation,
- accelerated approval, and
- priority review.56

Vaccines may be designated to more than one program. Fast track product designation and breakthrough therapy are both intended to streamline the clinical development process, but the qualifying criteria and features of these programs differ.

To qualify for *fast track product designation*, a vaccine must be intended for a serious condition, and nonclinical or clinical data must demonstrate its potential to address an unmet medical need.57 The sponsor of a fast track-designated product is eligible for frequent interactions with the FDA review team, priority review, and rolling review (in which FDA reviews portions of a BLA before a complete application is submitted).58

To qualify for *breakthrough designation*, a vaccine must be intended for a serious condition, and preliminary clinical evidence must indicate that it demonstrates potential substantial improvement on a clinically significant endpoint(s) over available therapies. Features of breakthrough therapy designation include rolling review; intensive FDA guidance on designing an efficient drug development program; involvement of “senior managers and experienced review and regulatory health project management staff in a proactive, collaborative, cross-disciplinary review” to expedite the development and review of a breakthrough therapy; and eligibility for other expedited programs.

54 21 C.F.R. §610.1.
55 21 C.F.R. §610.2.
57 FFDCA §506(b) [21 U.S.C. §356(b)].
58 FFDCA §506(a) [21 U.S.C. §356(a)].
Interested sponsors must submit to FDA a request for fast track product designation or breakthrough therapy designation. The request may be submitted with either the IND or any time after, as further specified in FDA guidance.

The accelerated approval pathway allows a vaccine to be licensed based on its effect on a surrogate endpoint (e.g., a laboratory measurement such as development of neutralizing antibodies) that predicts effectiveness, or on a clinical endpoint that can be measured earlier than irreversible morbidity or mortality. To qualify for accelerated approval, a vaccine must (1) be intended for a serious condition, (2) generally provide a meaningful advantage over available therapies, and (3) demonstrate an effect on an endpoint that is reasonably likely to predict clinical benefit. Postmarketing confirmatory studies generally must be completed to demonstrate actual effectiveness. Because surrogate endpoints for vaccines are often difficult to characterize, owing to the complexity of protective immune responses, accelerated approval may not be a relevant licensure pathway for many vaccines.

A priority review designation signifies that FDA’s goal is to take action on an application within 6 months of its filing, compared with 10 months for standard review. A BLA may qualify for priority review designation if, for example, it is for a vaccine intended for a serious condition and, if approved, would provide a significant improvement in safety or effectiveness. A BLA also may qualify for priority review if submitted with a priority review voucher.

Animal Rule

As mentioned above, FDA typically requires substantial evidence of effectiveness from adequate and well-controlled trials conducted in humans prior to licensing a vaccine. However, in certain cases, evaluating a vaccine’s efficacy or effectiveness through human trials is not possible. For example, it would not be ethical to expose human subjects to lethal toxic substances in order to test an investigational vaccine.

Under the Animal Rule, if human efficacy studies are not ethical, and if field trials (i.e., trials conducted outside of the clinical setting) are not feasible, FDA may license a vaccine based on adequate and well-controlled animal efficacy studies if those studies establish that the vaccine is likely to produce clinical benefit in humans. The Animal Rule is intended for drugs and biologics that would treat or prevent serious or life-threatening conditions caused by chemical, biological, radiological, or nuclear substances (e.g., nerve agents, emerging infectious pathogens, snake venom, and industrial chemicals). For FDA to rely on evidence from animal studies to provide evidence of effectiveness, four criteria must be met:

59 FFDCA §506(a)(2) & (b)(2) [21 U.S.C. §356(a)(2) & (b)(2)].
61 FFDCA §506(c) [21 U.S.C. §356(c)].
63 Three priority review voucher programs are currently authorized in the FFDCA: (1) the tropical disease priority review program, (2) the rare pediatric disease priority review program, and (3) the material threat MCM priority review voucher program. Under each of these programs, the sponsor of an NDA or BLA that meets the statutory requirements of the specific program is eligible to receive, upon approval, a transferable voucher, and the sponsor may either use that voucher for the priority review of another application or sell it to another sponsor to use.
There is a reasonably well-understood pathophysiological mechanism of the toxicity of the substance and its prevention or substantial reduction by the product;

The effect is demonstrated in more than one animal species expected to react with response predictive for humans, unless the effect is demonstrated in a single animal species that represents a sufficiently well-characterized animal model for predicting the response in humans;

The animal study endpoint is clearly related to the desired benefit in humans, generally the enhancement of survival or prevention of major morbidity; and

The data or information on the kinetics and pharmacodynamics of the product or other relevant data or information, in animals and humans, allows selection of an effective dose in humans.65

Drugs and vaccines evaluated for efficacy under the Animal Rule are evaluated for safety under the existing requirements for drugs and biologics. Postmarketing studies, such as field studies, must be conducted once feasible, and the sponsor of the vaccine must prepare certain patient-specific information explaining that the approval was based on efficacy studies conducted in animals alone. FDA also may impose postmarketing restrictions on distribution of the product if necessary to ensure safety (e.g., restricting distribution to certain facilities or practitioners with special training or experience).66 To date, FDA has licensed one vaccine under the Animal Rule: BioThrax (Anthrax Vaccine Adsorbed [injection]). Specifically, in 2015, the Animal Rule was used to approve a new use—post-exposure prophylaxis of disease—of a previously licensed anthrax vaccine.67

**Emergency Use Authorization (EUA)**

In general, a vaccine may be provided to patients only if FDA has licensed its marketing under a BLA or authorized its use in a clinical trial under an IND. In certain circumstances, however, FDA may allow patients to access investigational vaccines outside this framework, including through emergency use authorization (EUA).

FDA may enable access to an unapproved vaccine by granting an EUA, if the HHS Secretary declares that circumstances exist to justify the emergency use of an unapproved product or an unapproved use of an approved medical product.68 The HHS Secretary’s declaration must be based on one of four determinations; for example, a determination that an actual or significant potential exists for a public health emergency that affects or has significant potential to affect national security or the health and security of U.S. citizens living abroad.69 Following the HHS Secretary’s declaration, FDA, in consultation with the Assistant Secretary for Preparedness and Response (ASPR), the National Institutes of Health (NIH), and CDC, may issue an EUA authorizing the emergency use of a vaccine, provided that the following criteria are met:

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66 21 C.F.R. §601.91.


68 FFDCA §564 [21 U.S.C. §360bbb-3]. For additional information, see CRS In Focus IF10745, *Emergency Use Authorization and FDA’s Related Authorities*.

69 FFDCA §564(b)(1) [21 U.S.C. §360bbb-3(b)(1)].
• the agent that is the subject of the EUA can cause a serious or life-threatening
disease or condition;
• based on the totality of the available scientific evidence, it is reasonable to
believe that the product may be effective in diagnosing, treating, or preventing
such disease or condition, and that the known and potential benefits of the
product outweigh its known and potential risks; and
• there is no adequate, approved, or available alternative to the product.\textsuperscript{70}

The standard of evidence for an EUA is different than that for approval. EUA issuance, as noted
above, is based on FDA’s determination that the totality of the available scientific evidence
suggests that a product may be effective in diagnosing, treating, or preventing a disease or
condition, and that the known and potential benefits of the product outweigh its known and
potential risks. This standard of evidence is different from the one required for full FDA approval
or licensure, which is based on \textit{substantial evidence} of effectiveness derived from adequate and
well-controlled studies.\textsuperscript{71}

FDA must impose certain conditions as part of an EUA to the extent practicable (e.g., distributing
certain information to health care providers and patients) and may impose additional discretionary
conditions where appropriate.\textsuperscript{72} FDA may waive or limit current good manufacturing practices
(e.g., storage and handling) and prescription dispensing requirements for products authorized
under an EUA. In addition, FDA may establish conditions on advertisements and other
promotional printed matter that relates to the emergency use of a product. An EUA remains in
effect for the duration of the emergency declaration made by the HHS Secretary under FFDCA
Section 564, unless revoked at an earlier date.

To date, FDA has not granted an EUA for an unapproved (i.e., unlicensed) vaccine. However, in
2005, FDA had issued an EUA for the \textit{unapproved use} of a previously licensed vaccine.\textsuperscript{73}

\section*{Advisory Committee Consultation}

FDA consults with a federal advisory committee on various vaccine-related matters. Specifically,
the Vaccines and Related Biological Products Advisory Committee (VRBPAC) is made up of
non-FDA medical and scientific experts who inform FDA’s regulation of vaccines and related
biological products. The committee “reviews and evaluates data concerning the safety,
effectiveness, and appropriate use of vaccines and related biological products” and “considers the
quality and relevance of FDA’s research program which provides scientific support for the
regulation of these products and makes appropriate recommendations” to the FDA
Commissioner.\textsuperscript{74} VRBPAC may, for example, meet to discuss approaches for demonstrating

\textsuperscript{70} FFDCA \$564(c) [21 U.S.C. \$360bbb-3(c)]. These criteria are explained in more detail in the FDA guidance
media/97321/download.
\textsuperscript{71} FFDCA \$505(d) [21 U.S.C. \$355(d)].
\textsuperscript{72} FFDCA \$564(e) [21 U.S.C. \$360bbb-3(e)].
\textsuperscript{73} Authorization of Emergency Use of Anthrax Vaccine Adsorbed for Prevention of Inhalation Anthrax by Individuals
at Heightened Risk of Exposure Due to Attack With Anthrax, 70 \textit{Federal Register} 5452, February 2, 2005.
\textsuperscript{74} Vaccines and Related Biological Products Advisory Committee, https://www.fda.gov/advisory-committees/blood-
vaccines-and-other-biologics/vaccines-and-related-biological-products-advisory-committee.
effectiveness of a particular vaccine in a specific population.\textsuperscript{75} VRBPAC is subject to the requirements of the Federal Advisory Committee Act.\textsuperscript{76}

**Clinical Recommendations**

Official HHS/CDC clinical recommendations for vaccination—such as the age and population groups recommended to receive each vaccine, as well as the number of doses and interval between doses—are informed by the Advisory Committee on Immunization Practices (ACIP), a federal advisory committee composed of medical and public health experts who make policy recommendations for the use of licensed vaccines and related agents for the control of vaccine-preventable diseases in the civilian population of the United States.\textsuperscript{77} ACIP may also develop guidance for use of unlicensed vaccines “if circumstances warrant.” ACIP was established by the U.S. Surgeon General in 1964, under authority provided by Public Health Service Act (PHSA) Section 222.\textsuperscript{78}

After FDA licenses a new vaccine or licenses an existing vaccine for a new indication, ACIP typically makes one of two types of clinical recommendations:

- **Full recommendation:** The vaccine is recommended for all people in an age- or risk-based group, except for those with a contraindication (i.e., a condition that would make the vaccine harmful, such as a condition that compromises the immune system). For example, ACIP has issued a full recommendation for two doses of the measles-mumps-rubella (MMR) vaccine routinely for children, with the first dose administered at 12-15 months and the second dose administered before school entry at four to six years of age.\textsuperscript{79}

- **Clinical Decisionmaking:** The vaccine is recommended for certain subpopulations, and its use is based on clinical decisionmaking.\textsuperscript{80} For example, ACIP recommends the two Serogroup B Meningococcal vaccines for persons 10 years of age or older who have certain health conditions or are at increased risk of exposure to the disease, as specified.\textsuperscript{81}


\textsuperscript{76} For additional information about the Federal Advisory Committee Act (FACA) and FACA committees, see CRS Report R44253, *Federal Advisory Committees: An Introduction and Overview*.


To make its vaccine recommendations, ACIP considers disease epidemiology and burden of disease, vaccine efficacy and effectiveness, the quality of evidence reviewed, economic analyses, and implementation issues. Recommendations made by ACIP are reviewed by the CDC Director and, if adopted, published as official CDC/HHS recommendations. ACIP recommendations inform which vaccines are provided through the CDC’s Vaccines for Children program, as well as which vaccines must be covered by private health care insurance plans subject to the preventive health services requirement as added by the Patient Protection and Affordable Care Act (ACA).

ACIP recommendations are used to establish the CDC-recommended child and adult immunization schedules (for children, birth to 18 years of age; for adults, 19 years of age and older), which are used by health care providers, parents, and others to understand which vaccines should be administered at various ages. The immunization schedules distinguish between vaccines recommended to all people in a certain age group and vaccines recommended only for certain high-risk groups. As a part of the immunization schedules, CDC also publishes a specific table of vaccine recommendations by common contraindications, such as persons with HIV, immunocompromised individuals, and pregnant individuals. The table includes when recommended vaccines should not be administered to individuals with these contraindications.

Once clinical recommendations are made, CDC develops and provides resources and training for health care providers on current vaccine recommendations, best practices for vaccine administration, and patient education. CDC develops Vaccination Information Statements (VIS) on the risks and benefits of vaccinations; these statements are required to be given to vaccine recipients and their parents or legal guardians whenever vaccines recommended for routine use among children and pregnant women are administered. VISs are developed by CDC in consultation with the Advisory Commission on Childhood Vaccines (ACCV; a committee of health care professionals, attorneys, and parents of vaccine-injured children), health care providers, and FDA, and are published in the Federal Register for public comment.

**Postmarket Safety**

Although pre-licensure clinical trials and research are designed to identify common safety risks associated with a vaccine, such trials may not identify all long-term or rare adverse effects (similar to all pharmaceutical products). As such, vaccines may be subject to additional postmarket study requirements, called Phase 4 studies, or other safety monitoring to provide...
additional information about a vaccine’s risks, benefits, and optimal use. FDA may require a vaccine manufacturer to conduct a postapproval study or clinical trial to assess a known serious risk or signals of serious risk related to use of the vaccine, or to identify an unexpected serious risk when available data indicate the potential for a serious risk. In addition, because vaccines require special manufacturing processes to avoid contamination, post-licensure safety programs are designed to ensure safety in vaccine manufacturing. Post-licensure safety requirements and programs are also intended to identify long-term or rare adverse health events that result from vaccination, and FDA may require vaccine manufacturers to revise vaccine product labeling if new information becomes available after licensure.

**Manufacturing Safety**

FDA continues to inspect vaccine manufacturing facilities post-licensure. The HHS Secretary may authorize any HHS officer, agent, or employee to “during all reasonable hours enter and inspect any establishment for the propagation or manufacture and preparation of any biological product [e.g., vaccine].” If FDA determines that a batch, lot, or other quantity of a vaccine “presents an imminent or substantial hazard to the public health,” the agency must issue an order immediately recalling the batch, lot, or other quantity of the vaccine.

Manufacturers of vaccines listed in the Vaccine Injury Table (see the “National Vaccine Injury Compensation” section) or mandated to be state-administered must maintain records related to the safety and quality of each batch of vaccines produced, and must report any identified public health hazards to FDA. Specifically, vaccine manufacturers are required to maintain records documenting the manufacturing, processing, testing, and reworking of each batch, lot, or other quantity of a vaccine, including whether any significant problems were identified during these processes, and to report if any safety test on such batch, lot, or other quantity indicates a potential imminent or substantial public health hazard.

In addition, vaccine manufacturers are required to report adverse events to FDA. This includes the submission of 15-day alert reports and periodic safety reports. A 15-day alert report is required for each serious and unexpected adverse experience and must be submitted to FDA as soon as possible but no later than 15 days from initial receipt of the information by the manufacturer. The manufacturer must “promptly investigate” such adverse event and submit follow-up reports within 15 days of receiving new information or as requested by FDA. Periodic safety reports are required for each adverse experience not reported in a 15-day alert report and must be submitted to FDA at quarterly intervals for three years from the date of issuance of the

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94 PHSA §351(c) [42 U.S.C. §262(c)].

95 PHSA §351(d)(1) [42 U.S.C. §262(d)(1)].

96 PHSA §2128 [42 U.S.C. §300aa–28]. This authority has been delegated from the HHS Secretary to the FDA Commissioner, per the FDA Staff Manual Guide 1410.10, item 31, effective date August 26, 2016, https://www.fda.gov/media/81983/download.

97 PHSA §2128(a) [42 U.S.C. §300aa–28(a)].

98 21 C.F.R §600.80(c).
vaccine’s license, and at annual intervals thereafter. Individual case safety reports for vaccines submitted to FDA must include specified information about the patient who is the subject of the report (e.g., name, age, gender) and the vaccine (e.g., manufacturer, lot number). If a vaccine manufacturer fails to establish and maintain records or report adverse events, FDA can take enforcement action, including revocation of the BLA for that vaccine.

Surveillance

CDC and FDA are the primary federal agencies that conduct surveillance (i.e., data monitoring) activities on the safety of administered vaccines. Other federal agencies such as the Department of Defense (DOD) and the Centers for Medicare & Medicaid Services (CMS) also operate databases on vaccine safety events among their covered populations. The NVPO within the HHS Office of Infectious Disease and HIV/AIDS Policy (OIDP) is tasked with coordinating vaccine safety monitoring across federal agencies.

FDA and CDC monitor and conduct research on vaccine safety through various mechanisms. As discussed below, each of the programs or systems has strengths and limitations, but together they provide various ways of assessing vaccines to ensure their safety. Each of the systems allows for monitoring of adverse events linked to specific lots of manufactured vaccines. This lot-specific monitoring enables distinctions between adverse events linked to improper manufacturing, compared with adverse events linked to a particular type of vaccine.

Vaccine Adverse Event Reporting System (VAERS)

VAERS, established in 1990 and operated jointly by FDA and CDC, is a monitoring system for adverse events related to vaccines. Using the VAERS system, anyone, including physicians, nurses, and the general public, can submit an online report of an adverse event following vaccination. Pursuant to PHSA Section 2125, health care providers and vaccine manufacturers are required to report the occurrence of any adverse event in the Vaccine Injury Table (see the “National Vaccine Injury Compensation” section), the occurrence of a contraindicating reaction specified on the vaccine label, and other serious and unexpected events as required through regulations. Scientists at CDC and FDA monitor VAERS reports and use the information to conduct further investigations on the reported

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99 21 C.F.R §600.80(g).
100 21 C.F.R §600.80(l).
cases. Consolidated data on reported adverse events in the VAERS system are publicly available online.

VAERS is a passive reporting system. Its data represent reports of adverse health events related to vaccines, rather than validated cases. In addition, data in the system lack information on total vaccines administered in the covered populations. Therefore, VAERS data are often inadequate for epidemiological analyses of adverse health events at a population level. VAERS is useful, however, for helping identify new and unusual clusters of cases of adverse health events linked to vaccination. VAERS also can provide some of the first postmarket safety data on newly introduced vaccines. In addition, VAERS can help identify extremely rare and unusual adverse health events that occur following vaccination. Researchers can use VAERS reports to generate hypotheses about vaccine safety and then use other sources of data (such as from the databases discussed below) and clinical evidence to assess their hypotheses.

**Vaccine Safety Datalink (VSD)**

VSD, established in 1990 and operated by CDC, is an active surveillance system that allows for population-level scientific analyses of adverse events that follow vaccination. VSD is a collaborative project for conducting studies on vaccine safety between CDC and eight integrated health care organizations (i.e., combined payer and provider organizations) around the country. VSD uses electronic patient and medical records from participating sites, which allows for large-scale and controlled analyses of medical events (e.g., hospitalizations, diagnoses) that occur after vaccination to identify associated risks. VSD studies may supplement these records with other sources of information, such as patient surveys, medical charts, and pharmacy, laboratory, and radiology data, to validate vaccination data and outcomes. Health data on about 9 million people are included annually in VSD.

VSD allows for near real-time detection of large-scale adverse events linked to vaccination. Researchers have developed methods to use VSD data to study the health effects of vaccines, such as whether the measles-mumps-rubella (MMR) vaccine is associated with autism (studies have found no such association). Among its limitations, the population represented by VSD, while large, is not completely representative of the entire U.S. population in terms of geography, race, socioeconomic status, and other factors, particularly because the participating organizations are private health plans which generally over-represent people of higher socioeconomic status and

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non-minority groups.\textsuperscript{111} In addition, VSD’s population size may not be adequate for detecting extremely rare adverse events linked to vaccination.\textsuperscript{112}

**Sentinel Initiative**

FDA established the Sentinel Initiative in 2008, fulfilling a statutory directive to collaborate with public, academic, and private entities to develop methods for obtaining access to disparate data sources and to validate means of linking and analyzing safety data from multiple sources.\textsuperscript{113} As part of the Sentinel Initiative, FDA has established two programs that address vaccines: (1) the Post-Licensure Rapid Immunization Safety Monitoring (PRISM) program, and (2) the Biologics Effectiveness and Safety (BEST) system.

PRISM is an active surveillance program that uses electronic health records from insurance providers and state immunization registries to monitor adverse events following vaccination. It was established in 2009 and deployed during the H1N1 influenza pandemic.\textsuperscript{114} PRISM has been the largest linked database for monitoring vaccine safety in the United States, involving data on over 100 million people.\textsuperscript{115} PRISM, similar to the CDC VSD program, can allow for population-level scientific analyses of adverse events following vaccination. Because of the larger population covered, PRISM can detect rarer adverse events than VSD and enable stratified analyses of vaccine-linked adverse events by subpopulation (e.g., by race/ethnicity).\textsuperscript{116} As of 2012, VSD allowed for more rapid analyses than PRISM due to data-sharing agreements between the participating health organizations and CDC that allow for near real-time data collection.\textsuperscript{117}

PRISM has been used to inform FDA-required postmarket labeling changes.\textsuperscript{118} For example, after some studies found an association between risk of intussusception (i.e., intestinal blockage) and administration of two rotavirus vaccines (RotaTeq and Rotarix), FDA launched a study in PRISM to assess whether infants faced a similar risk.\textsuperscript{119} The PRISM study identified an increased, but


\textsuperscript{113} The Sentinel system was implemented as an “Active Post-Market Risk Identification and Analysis program” under FFDCA §505(k)(3), as amended by §905 of the FDA Amendments Act, P.L. 110-85.

\textsuperscript{114} PRISM is the vaccine component of FDA’s Sentinel Initiative.


rare, risk of intussusception with RotaTeq among infants, which led to FDA-required labeling changes for the licensed vaccine.\textsuperscript{120}

In 2017, CBER initiated the BEST system as part of Sentinel to assure the safety and effectiveness of vaccines and other biologics. It is broader than PRISM in that it also covers blood and blood products, tissue products, and other advanced therapeutic biologics.\textsuperscript{121} The goal of BEST is to “leverage high-quality data, analytics and innovation to enhance surveillance, real-world evidence generation, and clinical practice that benefits patients.” Like other Sentinel components, BEST uses electronic health record, administrative, and claims-based data for active surveillance and research. BEST fulfills the FDAAA requirements for an active postmarket risk and analysis system covering at least 100 million persons.\textsuperscript{122}

**Other Safety Monitoring Systems**

As mentioned above, federal agencies other than FDA and CDC conduct vaccine safety monitoring. CMS has a database for vaccine safety among the Medicare population; the database represents vaccines administered to persons aged 65 and older. DOD has a database for monitoring adverse events from vaccination among military service members and their families, and the Department of Veterans Affairs (VA) has a database for veterans who receive care in the VA system. In addition, the Indian Health Service (IHS) operates a database for vaccine safety monitoring among the IHS-covered population.\textsuperscript{123}

<table>
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<tr>
<th>Safety Monitoring Using Multiple Surveillance Systems: A Case Study</th>
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<td>Researchers have used information from multiple vaccine safety monitoring systems to draw associations between vaccines and subsequent adverse health events. For example, during the 2010-2011 influenza season, VAERS received an increased number of reports of febrile seizures following vaccination with Fluzone™. FDA then initiated a PRISM study to investigate febrile seizures after vaccination with Fluzone™ and other trivalent inactivated influenza vaccines (TIVs). The study found no statistically significant association between TIVs and increased risk of febrile seizures.</td>
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\textsuperscript{120} According to FDA, “The Mini-Sentinel PRISM study is the largest study of intussusception after rotavirus vaccines to date and identified an increased risk of intussusception in the 21 day time period after the first dose of RotaTeq, with most cases occurring in the first 7 days after vaccination. No increased risk was found after the second or third doses. These findings translate into 1 to 1.5 additional cases of intussusception per 100,000 first doses of RotaTeq.” See “FDA Safety Communication: FDA Approves Required Revised Labeling for RotaTeq Based Final Study Results of a Mini-Sentinel Postlicensure Observational Study of Rotavirus Vaccines and Intussusception,” July 22, 2013, [https://www.sentinelinitiative.org/communications/fda-safety-communications/fda-safety-communication-fda-approves-required-revised](https://www.sentinelinitiative.org/communications/fda-safety-communications/fda-safety-communication-fda-approves-required-revised).


Clinical Assessment

The Clinical Immunization Safety Assessment (CISA), a CDC program established in 2001, is a network of clinical scientists who conduct clinical studies (i.e., studies with patients) on vaccine safety. Scientists in the network can conduct studies on complex individual patient cases of possible adverse health events that followed vaccination. Using CISA, scientists can assess the biological mechanisms that cause adverse health events after vaccination. In addition, CISA manages a repository of biospecimen samples from patients who experience unusual adverse events following vaccination. These samples can be systemically analyzed to inform a mechanistic understanding of such adverse events.

Federal Research on Vaccine Safety

Postmarket surveillance systems and clinical assessments provide important data and evidence on potential adverse events following vaccination. To further understand and determine whether vaccines cause or could plausibly cause certain adverse health events, scientists conduct various types of research that inform a scientific understanding of vaccine safety (separate from the clinical trials under an IND). Such activities are supported primarily by HHS agencies, mainly CDC and the National Institutes of Health (NIH). In addition, FDA supports regulatory research related to methods for evaluating vaccine safety. Major areas of research related to vaccines can include the following:

- **Biological research**: Research often with animals, cell cultures, or biological specimens (e.g., human tissue samples) to explore the mechanisms by which vaccines act in biological systems, informing an understanding of how adverse events may occur. (Also referred to as basic biomedical research).
- **Epidemiological research**: A form of statistical research involving health data collected among defined human populations (such as postmarket surveillance data) to explore whether statistical associations exist between vaccination and subsequent adverse events, and any related risk factors for those events among those populations.
- **Clinical research**: Research with patients to understand the clinical features of adverse health events among patients that are hypothesized to be connected to vaccination.

Research can also explore the underlying methodologies used to assess vaccine safety through any of these forms of research.

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CDC Research

CDC conducts and supports many types of research on vaccine safety, including epidemiological and clinical studies. Many of CDC’s research publications rely on data and findings from its safety monitoring systems, as listed above, including VAERS, VSD, and CISA. CDC research often focuses on the use of specific vaccines in specific populations, as well as hypothesized side effects and adverse events potentially attributable to vaccination. For example, a recent CDC study published in 2020 explored probability-based methods of determining which vaccine or combination of vaccines were linked to an adverse event following vaccination (in this case, a seizure) when multiple vaccines were administered at once.

NIH Research

In addition to CDC research, biological research related to immunology or infectious disease supported by NIH informs an understanding of vaccine safety. NIH tends to support more biological research than CDC, in that NIH research focuses on the fundamental biological mechanisms underlying vaccine safety, as well as research methodologies for examining it. For the past several years, NIH, in collaboration with CDC and NVPO, has issued annual funding opportunity announcements for “Research on Vaccine Safety.” Research projects can include scientific investigations into physiological and immunological responses to vaccines; explorations of how genetic variation affects responses to vaccines; investigations into risk factors for adverse responses to vaccination; exploration and validation of statistical methods for analyzing data on vaccine safety; and the application of genomic and molecular technologies to assess vaccine safety.

The National Institute of Allergy and Infectious Diseases (NIAID, which is one of 27 NIH Institutes and Centers) also supports the Human Immunology Project Consortium (HIPC), a program established in 2010 that collects in-depth biological data over time on the immune systems of a diverse cohort of patients. The program consolidates data on the cohort into centralized databases for use by researchers. Researchers are using HIPC to study certain aspects of vaccine safety, such as whether a relationship exists between short-term adverse events caused by vaccination and long-term health effects. When combined with postmarket surveillance data and studies, NIH-supported research can contribute to robust evaluations on the safety of vaccines.

FDA Research

FDA conducts regulatory science research to facilitate its evaluation of vaccine safety and effectiveness, and to support the development of new vaccines. For example, CBER scientists have published studies on the agency’s effort to develop and evaluate assays and animal models

for studying the safety and efficacy of vaccines against specific pathogens, as well as to characterize biomarkers of vaccine safety and efficacy. In addition, FDA has studied certain adjuvants and preservatives added to vaccines, including thimerosal and the impact of aluminum in vaccines on infants. FDA research efforts have also focused on vaccine availability, specifically on influenza vaccine production and ensuring a sufficient supply of a safe vaccine.

**Other Federal Research**

Other federal agencies conduct or support research related to vaccine safety. For example, the NVPO has issued Funding Opportunity Announcements (FOA) for grants to support vaccine safety research. The Agency for Healthcare Research and Quality (AHRQ) has conducted vaccine safety reviews. The Department of Defense (DOD) and the Department of Veterans Affairs (VA) also support some vaccine safety research.

Periodically, federal agencies (particularly HHS) conduct or commission comprehensive scientific reviews on the safety of recommended vaccines. As described in the text box on page 6, these reviews often evaluate and combine evidence from a large number of studies and a range of research types to make assessments about the safety of vaccines that are as conclusive as possible. For example, in 2011, under HHS contract, the National Academy of Medicine (NAM) conducted a comprehensive review of the scientific evidence regarding the safety of eight pediatric vaccines. The resulting NAM report, *Adverse Effects of Vaccines: Evidence and Causality*, was used to inform an update of the Vaccine Injury Table for the National Vaccine Injury Compensation Program (see the “National Vaccine Injury Compensation” section). This review was subsequently updated in 2014 with additional research by AHRQ, supported by the NVPO; AHRQ is currently in the process of updating this review.

**Challenges of Vaccine Safety Reviews**

As discussed earlier, causality assessments that combine evidence across many studies allow for researchers to assess if all the available evidence favors a causal relationship between a vaccine and a subsequent adverse health event. In general, establishing true causal linkages between a

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136 See, for example, BetaSam.gov, “Research, Monitoring and Outcomes Definitions for Vaccine Safety,” https://beta.sam.gov/fal/c8125303527f478981f6b7395c528788/view.


138 NAM was named the Institute of Medicine when the Immunization Safety Review Committee was formed.


vaccine and certain subsequent adverse health events can be challenging; however, researchers draw conclusions using multiple forms of evidence. The clinical trials required for vaccine licensure are well-controlled scientific experiments that allow researchers to draw conclusions about the safety of products. Postmarket safety studies, on the other hand, can face a variety of methodological challenges. For one, the population of vaccinated individuals is often much larger than and demographically different from the population of unvaccinated individuals, making it difficult to draw comparisons in health outcomes between the two groups. Researchers therefore often rely on time intervals between vaccination and an adverse health event—assessing whether a certain adverse health event is more likely to occur within a defined time interval after vaccination compared with other time periods. While this approach can work for short-term health effects caused by vaccines, it can be less effective for hypothesized long-term effects of vaccines or adverse health events that are otherwise common in the population. Statistical association between vaccination and an adverse health event is often necessary but not sufficient to establish causality. As discussed earlier, to make a causality assessment about whether a particular vaccine causes an adverse health event, experts use evidence and results from many scientific studies, including epidemiological evidence, clinical evidence, and biological laboratory evidence, usually with methods to weigh, compare, and combine evidence across studies. Such causality assessments may be conducted as a part of a comprehensive scientific review by federal or academic scientists, or by independent scientific advisory bodies, such as the NAM.

**National Vaccine Injury Compensation**

The National Vaccine Injury Compensation Program (VICP) provides compensation to individuals who file a petition and are found to have been injured by a covered vaccine. VICP is based in the Health Resources and Services Administration (HRSA) and was established by the National Childhood Vaccine Injury Act of 1986 (P.L. 99-660). VICP publishes a “Vaccine Injury Table” that lists vaccines covered by the program and the injuries associated with those vaccines for which claims may be filed, developed based on the causality assessments conducted by IOM and AHRQ. Claimants may submit claims for injuries that are not listed on the table, but they must present evidence that the vaccine caused the injury. In addition to HHS/HRSA, VICP involves the Department of Justice (DOJ) and the U.S. Court of Federal Claims. The Advisory Committee on Childhood Vaccines (ACCV) also provides oversight of VICP by making recommendations to the HH Secretary, including those related to the Vaccine Injury Table. ACCV is a nine-member federal advisory committee made up of health and legal representatives, as well as parents or legal representatives of children who have been injured by vaccines.

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145 HHS, “Charter- Advisory Commission on Childhood Vaccines,” https://www.hrsa.gov/sites/default/files/hrsa/advisory-committees/vaccines/acvccharter.pdf. For the parents or legal representatives of children who have suffered a vaccine-related injury or death, HH specifies that to be considered for appointment, “there must have been a finding (i.e., a decision) by the U.S. Court of Federal Claims or a civil court that a VICP-covered vaccine caused, or was presumed to have caused, the represented child’s injury or death.” From HRSA, “Advisory Commission on Vaccines: Frequently Asked Questions,” 2018, https://www.hrsa.gov/sites/default/files/hrsa/vaccine-compensation/job-
VICP is funded by the Vaccine Compensation Trust Fund, which is funded by an excise tax on vaccines paid by manufacturers. VICP was established in response to vaccine shortages that occurred after hundreds of injury lawsuits were filed against vaccine manufacturers in the 1980s, leading to halts in vaccine production and creating instability in the vaccine market. VICP is a no-fault system to compensate individuals who were injured as a result of vaccination. It serves to protect manufacturers from injury lawsuits. As of October 1, 2020, over 22,272 petitions have been filed with VICP, and 7,611 were determined to be compensable, with total compensation paid of about $4.4 billion since the program was established in 1988.\(^\text{146}\)

During an emergency situation such as the COVID-19 pandemic, vaccines may be covered under a different injury compensation program—the Countermeasures Injury Compensation Program (CICP), as discussed in the “Injury Compensation and Patient Safety Information” section.\(^\text{147}\)

### Safety in Vaccine Distribution

Managing vaccine supply and distribution requires temperature control, safety controls, and regular monitoring of expiry dates due to the limited shelf life of products.\(^\text{146}\) Given that public dollars (federal and state) pay for over 50% of vaccines (by volume) in the United States, federal agencies play a role in the supply and distribution of vaccines.\(^\text{149}\) CDC, in particular, conducts activities to help improve management of the vaccine supply chain. Vaccine storage practices especially have implications for a vaccine’s potency (i.e., effectiveness).\(^\text{150}\)

Vaccines are distributed through a decentralized network of health care providers, health centers, pharmacies, and health departments. State requirements vary regarding the types of entities that can be licensed or authorized to administer various vaccines. In the CDC’s Vaccines for Children (VFC) program, health care providers can apply to receive and provide VFC-covered vaccines through state or local coordinators, who ensure that the provider meets program requirements (e.g., ability to properly store and handle vaccines).\(^\text{151}\) Any provider that is licensed or otherwise authorized to administer pediatric vaccines can apply to participate in a state’s VFC program and receive and administer a supply of vaccine.\(^\text{152}\)

Vaccine programs are expected to make vaccines widely available, while ensuring that they are safely stored, properly administered, and used or discarded before their expiry date. However, this requirement is a challenge for many vaccine programs. A 2012 HHS Inspector General report found that many VFC providers did not meet vaccine management requirements, either by exposing vaccines to improper temperatures, storing expired and nonexpired vaccines together, or failing to maintain documentation. CDC agreed with the report recommendations and committed

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\(^{147}\) CRS Legal Sidebar LSB10443, The PREP Act and COVID-19: Limiting Liability for Medical Countermeasures.


\(^{152}\) Social Security Act §1928(c); 42 U.S.C. §1396s(c).
to improving management among providers. Following the report, CDC changed VFC program requirements and issued recommendations to providers and immunization program managers.

CDC’s immunization programs include several efforts among state and local partners to improve the vaccine supply chain and vaccine distribution:

- The **Vaccine Management Business Improvement Project** (VMBIP) is an effort among CDC and state and local partners to improve the management of the vaccine supply chain, particularly for vaccines distributed through VFC. Since the project began in 2003, it has changed funding mechanisms, forecasting for supply needs, provider distribution, and inventory tracking among vaccine providers.

- The **Vaccine Tracking System** (VTrckS) is an information technology platform for managing the publicly funded vaccine supply chain available to CDC, state and local health departments, and providers.

### Safety Considerations for COVID-19 Vaccines

*The COVID-19 vaccine development, approval, and distribution planning situation is evolving. Readers should note the date of this publication and be aware that this report may not reflect events or actions that occurred after that date.*

Much remains unknown about potential safety issues related to COVID-19 vaccines. FDA has never licensed a vaccine for a coronavirus. Several COVID-19 vaccines in development use novel vaccine technologies, some of which have never before been used in licensed FDA vaccines. Among the few mass emergency vaccination efforts in the past century, there have been some unexpected safety issues. For example, in 1976, the federal government attempted a rapid mass influenza (flu) vaccination campaign in response to a novel swine flu strain. The vaccines were later found to lead to higher rates of Guillain-Barre Syndrome (a neurological disorder) among those vaccinated, ending the campaign.

U.S. vaccine development efforts have been supported and coordinated by Operation Warp Speed (OWS), the nation’s major COVID-19 vaccine, therapeutic, and diagnostic (medical countermeasures) development initiative. OWS has chosen to support 14 potential COVID-19 vaccine candidates from a pool of 93, with the stated goal of reducing the number of candidates to 7 as additional results from clinical trials and research become available. As of October 29,

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2020, eight investigational vaccines were supported within OWS’s portfolio. OWS and CDC are planning for a federally coordinated nationwide COVID-19 vaccine distribution campaign.

Making safe and effective COVID-19 vaccines available within a year represents an unprecedented scientific and public health effort. The safety considerations and applicability of the requirements, processes, and programs described in this report will likely differ when applied to COVID-19 vaccines in several key ways, particularly with respect to (1) vaccine development, (2) FDA marketing authorization, (3) clinical recommendations and prioritization, (4) surveillance and safety monitoring, (5) injury compensation and patient safety information, and (6) vaccine distribution. Each of these is described in more detail below.

Vaccine Development and Current Status

Typically, the vaccine development and testing process is linear, with an investigational vaccine progressing through each phase of clinical testing upon completion of the prior phase. As mentioned above, the first stage is basic research, and if laboratory and animal test data indicate that a vaccine candidate appears safe and effective against a pathogen, then a first-in-human Phase 1 trial generally follows. If the Phase 1 trial indicates that the vaccine is safe in humans, then Phase 2 testing commences, further examining safety and at what dosage the vaccine has an effect. Finally, if those studies are successful, then a large, placebo-controlled Phase 3 trial follows. This sequential process helps minimize potential health risks to study participants and financial risks to the company sponsoring the investigations. The OWS COVID-19 vaccine development process is not following this phased approach. Instead, it is conducting some of these steps simultaneously to generate safety and effectiveness data in a shorter period.

Several COVID-19 vaccines are currently in Phase 3 clinical trials, and initial results are available from several vaccines that have completed Phase 2 clinical trials. Federal officials have indicated that OWS expects to have initial results from Phase 3 clinical trials in late 2020 and early 2021. Results from several Phase 1 and Phase 2 clinical trials of COVID-19 candidate vaccines have demonstrated short-term safety and some evidence of efficacy. Initial safety data on vaccines supported by Moderna, Pfizer/BioNTech, AstraZeneca, and Johnson & Johnson found no serious safety issues, although more participants who received the vaccine in the trials experienced mild or moderate side effects (e.g., fatigue, fever) compared with the control groups. In addition, all four vaccines show initial evidence of immunogenicity, including antibodies (immune proteins) and other blood cells that neutralized the virus in blood samples of

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those who received the candidate vaccines.\textsuperscript{165} Much remains unknown about COVID-19 immunity, and these results are considered preliminary.

As covered in this report, critical data related to the safety and efficacy of vaccines are generally collected in Phase 3 clinical trials. Given that no vaccine for a coronavirus has been previously tested in Phase 3 clinical trials, much remains unknown about the safety issues that may arise. In particular, experts are concerned about the potential for vaccine enhanced disease, in which vaccination could worsen the health effects of COVID-19 infections, as seen with the dengue and other vaccines. Animal studies of other coronavirus vaccines have found some potential for vaccine enhanced disease, and experts recommend rigorous monitoring in clinical trials to detect this safety issue.\textsuperscript{166} One scientific review noted that the scientific and clinical evidence with COVID-19, thus far, provides limited evidence with respect to the issue of enhanced disease. Along with other evidence, the authors explore evidence from treatment of COVID-19 patients with convalescent plasma (a treatment involving antibodies) and note that distinguishing antibody enhanced disease from worsening of symptoms is difficult, and therefore the potential for this issue should be studied further.\textsuperscript{167}

OWS reports that it is providing scientific support for COVID-19 vaccine clinical trials, in collaboration with other federal agencies like NIH. According to a medical journal publication authored by OWS leaders, OWS is coordinating many components of the vaccine development process. With regard to efficacy data, “OWS will maximize the size of phase 3 trials (30,000 to 50,000 participants each) and optimize trial-site location by consulting daily epidemiologic and disease-forecasting models to ensure the fastest path to an efficacy readout.” Phase 3 trial endpoints have been coordinated between the trials, in collaboration with NIAID.\textsuperscript{168} NIH has leveraged some of its existing clinical trials networks for testing certain COVID-19 vaccines participating in Operation Warp Speed, named the COVID-19 Prevention Trials Network (COVPN) that, among other things, works to harmonize clinical endpoints for the trials and recruit study participants.\textsuperscript{169}

All of the COVID-19 vaccines supported by OWS that are in Phase 3 clinical trials have a Data and Safety Monitoring Board (DSMB) that independently reviews safety and effectiveness data on the investigational vaccine to determine if the trial should continue, be modified, be terminated, or be considered for FDA marketing authorization (see the “FDA Marketing Authorization” section).\textsuperscript{170} Three Phase 3 clinical trials of candidate vaccines supported by

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OWS—those of Moderna, AstraZeneca, and Johnson & Johnson—are overseen by a common DSMB developed in consultation with NIH’s Accelerating COVID-19 Therapeutic Interventions and Vaccines partnership as a part of the COVPN. As the Pfizer/BioNTech vaccine has a separate DSMB.

On September 8, 2020, it was reported that AstraZeneca paused its Phase 3 clinical trial in response to a potential safety issue; such pauses are not uncommon in any drug or biologic development effort. On October 12, 2020, it was reported that Johnson & Johnson paused its Phase 3 trial due to “an unexplained illness in a study participant” and that a DSMB has been convened to review the case. As of October 23, 2020, these trials have resumed. In response to calls for transparency, several vaccine developers, including Moderna, Pfizer/BioNTech, AstraZeneca, and Johnson & Johnson, have made their Phase 3 clinical trial protocols for COVID-19 vaccines publicly available.

As shown in the protocols, the trials are using an event-driven design, meaning that efficacy of the vaccines are to be evaluated once a certain number of “events” occur among the study population—in this context, COVID-19 cases with symptoms. Once a certain number of COVID-19 cases are detected, the DSMB is to evaluate the data and conduct a statistical analysis to determine if the difference in cases between the vaccine recipient group and the control group meet the FDA’s standard for effectiveness for a COVID-19 vaccine. For example, Moderna has determined that 151 COVID-19 cases among its study population would provide enough statistical power to determine whether the vaccine is 60% effective, with interim analyses of the data by the DSMB planned at 35% and 70% of the total target cases. The DSMB may recommend that the vaccine companies end the trials if interim analyses indicate safety issues or do not show adequate evidence of effectiveness. Vaccine expert groups, such as the Coalition for Epidemic


Preparedness, have advocated for the event-driven approach to COVID-19 vaccine trials in order to expedite vaccine availability without compromising scientific rigor. Other vaccine experts have voiced concerns about this approach, arguing that it “may make statistical sense, but it defies common sense.” These experts argue that the vaccines should be assessed for whether they protect against moderate and severe forms of COVID-19 and that the trials should be fully completed to generate adequate data.

FDA Marketing Authorization

The development and testing process for a COVID-19 vaccine is designed to be significantly shorter compared with the usual timeline for vaccine development. This shortened process may make it difficult to detect potential unexpected adverse events that may not manifest right away. Moreover, because the review process is to be shorter than the typical 6 to 10 months needed for a Biologics License Application (BLA) review, FDA scientists would have had less time to review the safety and effectiveness data. Although FDA uses various formal mechanisms to expedite the development and review of medical products intended to address unmet medical need, FDA has not yet granted Emergency Use Authorization (EUA) for a previously unapproved (i.e., unlicensed) vaccine. Thus, if a COVID-19 vaccine is first made available under an EUA rather than a BLA, it will be a first for the agency.

In light of reported concerns from the public surrounding the safety and effectiveness of COVID-19 vaccines developed on an expedited timeline, FDA officials have sought to clarify that any vaccine candidate “will be reviewed according to the established legal and regulatory standards for medical products.” In addition, FDA officials have indicated that the amount of safety and effectiveness data needed to support EUA issuance will be similar to the data that would be appropriate for a BLA. As mentioned above, the level of evidence required by statute for EUA issuance is different from licensure, although both require the submission of safety and effectiveness data to FDA. For licensure under a BLA, a vaccine would need to be proven safe and have substantial evidence of effectiveness to receive full licensure under a BLA. In the case that a vaccine is first made available under an EUA, substantial evidence of effectiveness would not be required by statute. Rather, the totality of the available scientific evidence would need to indicate that the vaccine may be effective in preventing COVID-19, and that the known and potential benefits of the vaccine outweigh its known and potential risks.

To help companies develop a vaccine to prevent COVID-19, and to increase transparency regarding the FDA’s expectations for safety and effectiveness data, the agency has issued two guidance documents. The first guidance, issued in June 2020, aims to clarify FDA’s expectations

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regarding the data and information necessary to support licensure under a BLA. 182 The guidance notes, among other things, that with respect to effectiveness, FDA expects a COVID-19 vaccine to prevent disease or decrease disease severity in at least 50% of people who are vaccinated. On October 6, 2020, FDA issued a second guidance, which focuses on the agency’s expectations for the data and information needed to support an EUA for a COVID-19 vaccine. 183 The recommendations outlined in the October 2020 guidance have been characterized as more stringent than what typically may be required for an EUA. 184 For example, the guidance indicates that data from Phase 3 trials submitted to the agency should include a median follow-up duration of at least two months after completion of the full vaccination regimen to help provide adequate information to assess a vaccine’s benefit-risk profile. FDA also expects clinical testing of an EUA-authorized vaccine to continue to support eventual licensure under a BLA. As such, the guidance recommends that sponsors submit, as part of the EUA request, strategies that will be implemented to (1) address loss of follow-up information for participants who choose to withdraw from the study to receive the vaccine under an EUA, and (2) ensure that ongoing clinical trials of the vaccine are able to assess long-term safety and effectiveness (e.g., evaluating for vaccine-associated ERD, decreased effectiveness over time) in sufficient numbers to support vaccine licensure.

The FDA Vaccines and Related Biological Products Advisory Committee (VRBPAC) met on October 22, 2020, to discuss generally the development, authorization, and licensure of vaccines to prevent COVID-19. 185 The VRBPAC discussed, among other things, FDA’s approach to safety and effectiveness as outlined in the agency’s guidance documents; expectations for the data that must be submitted for licensure or EUA, including information about the manufacturing process; and plans for postmarket surveillance, including use of existing systems such as VAERS and BEST. FDA also is reportedly developing master protocols to guide its safety and effectiveness oversight, to be made publicly available on its website. 186 To further provide transparency, FDA has indicated it will convene additional VRBPAC meetings to discuss specific vaccine candidates ready for an EUA or licensure. 187

Clinical Recommendations and Prioritization

ACIP has begun to weigh considerations related to COVID-19 vaccine clinical recommendations and prioritization, and has made information available for its public meetings in June, July, August, and September 2020. Many of these deliberations note unknowns with regard to COVID-19 vaccines, in particular, as related to clinical trial data on the safety and efficacy of vaccines. 188

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In addition, at the direction of NIH and CDC, the National Academies of Science, Engineering, and Medicine (NASEM) set up an ad hoc committee to develop a framework for equitably allocating COVID-19 vaccines, both domestically and globally. NASEM published its draft framework on September 1 and published its final report with recommendations on October 2. The framework establishes a prioritization methodology that recommends who should be the first to receive COVID-19 vaccines when they become available. Recommended recipients include high-risk workers in health care facilities, first responders, older adults, people with underlying conditions known to be associated with severe outcomes, critical risk workers (workers who are in essential industries and are at substantially higher risk of exposure), and teachers and school staff (see Figure 1).

**Figure 1. NASEM-Recommended Phased Approach to COVID-19 Vaccine Allocation**

![Figure 1](https://www.nationalacademies.org/graphics/mynacademies/figures/s2.jpg)

As of September 2020, ACIP has begun to publicly weigh NASEM’s recommendations and compare them to the recommendations from other groups, in particular, from the WHO Strategy Advisory Group of Experts (SAGE) and from the Johns Hopkins Bloomberg School of Public Health. ACIP is considering these recommendations in the context of its own proposed ethical principles of (1) maximizing benefits and minimizing harm, (2) equity, (3) justice, (4) fairness, and (5) transparency. In addition, ACIP has considered how to prioritize vaccine allocation

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191 ACIP COVID-19 Vaccines Work Group, “Overview of Vaccine Equity and Prioritization Frameworks,” September
within groups recommended for Phase 1 allocation, should limited vaccine supply require such choices. According to media reporting, these discussions continued at a public meeting in late October (meeting materials are not yet available on ACIP’s website).

Both the ACIP and NASEM groups are advisory; they do not establish binding policy. Although HHS has historically followed ACIP’s recommendations and often considers NASEM recommendations, it is unclear whether and to what extent these recommendations will inform HHS’s COVID-19 vaccine prioritization policies. A report to Congress from Operation Warp Speed on September 16, 2020 noted the NASEM and ACIP roles in prioritizing eventual vaccines but stated that final decisions about prioritization will not be made until closer to implementation.

Safety in Vaccine Distribution

CDC has begun to establish requirements for vaccine management, including requirements related to storage and transportation. As announced on August 14, McKesson Corporation is to act as a central distributor for the COVID-19 vaccine campaign—the same distributor that managed the federally coordinated H1N1 influenza pandemic vaccine campaign. States, localities, territories, and tribes (hereinafter, jurisdictions) are to have much of the responsibility for tracking vaccines provided and for local transportation of vaccines within the jurisdiction.

COVID-19 vaccines in development have different temperature control requirements: some must be refrigerated (2 to 8 degrees Celsius), some must be stored frozen (-15 to -25 degrees Celsius) and some must be kept ultra-cold (-60 to -80 degrees Celsius). CDC’s planning guidance to jurisdictions takes these different temperature requirements into account and seeks to minimize potential breaks in the cold chain during vaccine distribution. According to CDC, “certain COVID-19 vaccine products, such as those with ultra-cold temperature requirements, will be shipped directly from the manufacturer to the vaccination provider site,” while others will be distributed by CDC’s distributor directly to the provider sites or secondary depots for distribution (e.g., chain drug store’s central distribution). The guidance then further explains how these vaccines should be stored onsite until usage.

CDC, in collaboration with jurisdictions, is planning trainings for newly registered providers regarding safe storage, handling and administration of the vaccines. Providers who seek to participate in the COVID-19 vaccination program must be credentialed/licensed in the jurisdiction where vaccination takes place, and sign and agree to the conditions in the CDC COVID-19 Vaccination Program Provider Agreement. Jurisdictions’ immunization programs and


health care providers administering COVID-19 vaccines are to be responsible for many aspects of vaccine tracking, storage, and handling to ensure that vaccine safety and effectiveness are maintained, as outlined in CDC’s preliminary guidance.\(^{197}\) This guidance is likely to evolve as more information is available regarding the vaccines.

**Surveillance and Safety Monitoring**

Given the condensed nature of the COVID-19 development programs, FDA may require additional clinical studies to be conducted post-licensure to allow for continued monitoring of adverse events.\(^{198}\) In guidance, FDA further recommends that at the time of a BLA submission for a COVID-19 vaccine, a Pharmacovigilance Plan (PVP) be submitted to address known and potential risks of the vaccine. FDA may recommend that a PVP include expedited or more frequent adverse event reporting or the establishment of a pregnancy exposure registry to collect information on associated pregnancy and infant outcomes. As mentioned above, manufacturers of BLA-licensed vaccines typically must report adverse events to FDA within 15 days of becoming aware of them. In the event that a COVID-19 vaccine is first made available under an EUA rather than a BLA, FDA is expected to impose, as a condition of an EUA, requirements for health care providers and vaccine manufacturers to report and track any adverse events associated with administration of the vaccine.\(^{199}\)

Several federal vaccine safety databases are to be used to monitor postmarket safety for COVID-19 vaccines. A CDC presentation from the August ACIP meeting identifies that CMS, VA, DOD, IHS, FDA, and CDC databases will be leveraged to provide ongoing monitoring of COVID-19 vaccine safety. It was also reported that “FDA plans to develop new electronic data sources through [electronic health record] EHR partners.”\(^{200}\)

CDC has reported several efforts to enhance its safety monitoring systems in anticipation of the COVID-19 vaccination program. For health care providers participating in the COVID-19 vaccination program, per the **CDC COVID-19 Vaccination Program Provider Agreement**, providers are required to report adverse events following vaccination through VAERS and are advised to report such events even if the providers are not sure that vaccination caused the adverse event.\(^{201}\) A preliminary list of “adverse events of special interest” has been developed for monitoring attention in VAERS reports.\(^{202}\) As communicated to CRS, CDC is strengthening its existing safety monitoring systems in several ways, including by adding additional clinicians to the CISA network, by adding staff to the VAERS network, and by preparing these systems to

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provide rapid analyses on COVID-19 vaccine safety data. CDC plans to implement smartphone-based active vaccine safety monitoring of early recipients of COVID-19 vaccines, called Vaccine Safety Assessment for Essential Workers, or v-safe. This process would involve text, text-to-web survey, and email-to-web survey monitoring of healthcare workers and essential workers who might be prioritized to receive early doses of vaccine when it becomes available (see Figure 2).

**Figure 2. Graphical Presentation of Vaccine Safety Assessment for Essential Workers**

Presented at Advisory Committee on Immunization Practices meeting, September 22, 2020

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**Injury Compensation and Patient Safety Information**

Vaccine injury compensation for COVID-19 vaccines will likely differ from usual injury compensation under VICP. The Public Readiness and Emergency Preparedness Act (PREP Act) declaration issued on March 10, 2020, established certain immunity from legal liability related to the “manufacture, testing, development, distribution, administration, and use” of covered countermeasures as part of the public health response to COVID-19. Persons who suffer serious injury or death from a covered countermeasure may seek compensation through the Covered Countermeasure Process Fund as a part of the Countermeasures Injury Compensation Program (CICP). The HHS Secretary may transfer funds available in the Public Health and Social Services

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Emergency Fund (PHSSEF) in several coronavirus supplemental appropriations acts to this fund.\textsuperscript{205}

Because COVID-19 vaccines will likely not be added to the Vaccine Injury Table used for VICP (at least initially), CDC is not required to develop Vaccine Information Statements (VIS) for COVID-19 vaccines. CDC may choose to do so. Separately, if a vaccine is made available under an EUA, FDA has stated it will make fact sheets available for vaccine recipients (or their parents/legal guardians) and vaccine providers.\textsuperscript{206} CDC and vaccine manufacturers are also developing other educational material regarding the vaccines.\textsuperscript{207}

**Congressional Considerations**

Since enactment of the Biologics Control Act of 1902, Congress and the Administration (especially through FDA and CDC) have strived to ensure the safety of vaccines in the United States—from initial development to patient administration. With the COVID-19 pandemic causing considerable health and economic consequences, there is significant interest in developing safe and effective vaccines to help curb transmission of the disease. Congress may consider how to best leverage existing requirements and programs to ensure that risk of harm from eventual COVID-19 vaccines is mitigated and minimized. Several efforts are underway through OWS, FDA, and CDC to expedite the availability of COVID-19 vaccines and to prepare for a nationwide immunization campaign. Safety has been cited as a primary concern in all of these efforts. Congress may consider how to best provide oversight and make legislative changes to ensure a safe and successful COVID-19 vaccination campaign. In addition, Congress may consider and evaluate the entire federal vaccine safety system and assess whether this system warrants any policy changes to help ensure the safety of all recommended vaccines.

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\textsuperscript{205} CRS Legal Sidebar LSB10443, *The PREP Act and COVID-19: Limiting Liability for Medical Countermeasures*.  
\textsuperscript{207} Ibid., p. 23.
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