Explaining Operation Warp Speed

What’s the goal?
Operation Warp Speed’s goal is to produce and deliver 300 million doses of safe and effective vaccines with the initial doses available by January 2021, as part of a broader strategy to accelerate the development, manufacturing, and distribution of COVID–19 vaccines, therapeutics, and diagnostics (collectively known as countermeasures).

How will the goal be accomplished?
By investing in and coordinating countermeasure development, OWS will allow countermeasures such as a vaccine to be delivered to patients more rapidly while adhering to standards for safety and efficacy.

Who’s working on Operation Warp Speed?
OWS is a partnership among components of the Department of Health and Human Services (HHS), including the Centers for Disease Control and Prevention (CDC), the Food and Drug Administration (FDA), the National Institutes of Health (NIH), and the Biomedical Advanced Research and Development Authority (BARDA), and the Department of Defense (DoD). OWS engages with private firms and other federal agencies, including the Department of Agriculture, the Department of Energy, and the Department of Veterans Affairs. It will coordinate existing HHS–wide efforts, including the NIH’s Accelerating COVID–19 Therapeutic Interventions and Vaccines (ACTIV) partnership, NIH’s Rapid Acceleration of Diagnostics (RADx) initiative, and work by BARDA.

What’s the plan and what’s happened so far?
**DEVELOPMENT:** To accelerate development while maintaining standards for safety and efficacy, OWS has been selecting the most promising countermeasure candidates and providing coordinated government support.

Protocols for the demonstration of safety and efficacy are being aligned, which will allow these harmonized clinical trials to proceed more quickly, and the protocols for the trials will be overseen by the federal government (NIH), as opposed to traditional public–private partnerships, in which pharmaceutical companies decide on their own protocols. Rather than eliminating steps from traditional development timelines, steps will proceed simultaneously, such as starting manufacturing of vaccines and therapeutics at industrial scale well before the demonstration of efficacy and safety as happens normally. This increases the financial risk, but not the product risk.
Select actions to support OWS vaccine and therapeutic development so far include:

- **March 30**: HHS announced $456 million in funds for Johnson & Johnson’s (Janssen) candidate vaccine. Phase 1 clinical trials began in Belgium on July 24th and in the U.S on July 27th.

- **April 16**: HHS made up to $483 million in support available for Moderna’s candidate vaccine, which began Phase 1 trials on March 16 and received a fast-track designation from FDA. This agreement was expanded on July 26 to include an additional $472 million to support late-stage clinical development, including the expanded Phase 3 study of the company’s mRNA vaccine, which began on July 27th.

- **May 21**: HHS announced up to $1.2 billion in support for AstraZeneca’s candidate vaccine, developed in conjunction with the University of Oxford. The agreement is to make available at least 300 million doses of the vaccine for the United States, with the first doses delivered as early as October 2020 and Phase 3 clinical studies beginning this summer with approximately 30,000 volunteers in the United States.

- **July 7**: HHS announced $450 million in funds to support the large-scale manufacturing of Regeneron’s COVID-19 investigational anti-viral antibody treatment, REGN-COV2. This agreement is the first of a number of OWS awards to support potential therapeutics all the way through to manufacturing. As part of the manufacturing demonstration project, doses of the medicine will be packaged and ready to ship immediately if clinical trials are successful and FDA grants EUA or licensure.

- **July 7**: HHS announced $1.6 billion in funds to support the large-scale manufacturing of Novavax’s vaccine candidate. By funding Novavax’s manufacturing effort, the federal government will own the 100 million doses expected to result from the demonstration project.

- **July 22**: HHS announced up to $1.95 billion in funds to Pfizer for the large-scale manufacturing and nationwide distribution of 100 million doses of their vaccine candidate. The federal government will own the 100 million doses of vaccine initially produced as a result of this agreement, and Pfizer will deliver the doses in the United States if the product successfully receives FDA EUA or licensure, as outlined in FDA guidance, after completing demonstration of safety and efficacy in a large Phase 3 clinical trial, which began July 27th.

- **July 31**: HHS announced approximately $2 billion in funds to support the advanced development, including clinical trials and large scale manufacturing, of Sanofi and GlaxoSmithKline’s (GSK) investigational adjuvanted vaccine. By funding the manufacturing effort, the federal government will own the approximately 100 million doses expected to result from the demonstration project. The adjuvanted vaccine doses could be used in clinical trials or, if the FDA authorizes use, as outlined in agency guidance, the doses would be distributed as part of a COVID-19 vaccination campaign.

- **August 5**: HHS announced approximately $1 billion in funds to support the large-scale manufacturing and delivery of Johnson & Johnson’s (Janssen) investigational vaccine candidate. Under the terms of the agreement, the U.S. Government will own the resulting 100 million doses of vaccine, and will have the option to acquire more. The company’s investigational vaccine relies on Janssen’s recombinant adenovirus technology, AdVac, a technology used to develop and manufacture Janssen’s Ebola vaccine with BARDA support; that vaccine received European Commission approval and was used in the Democratic Republic of the Congo (DRC) and Rwanda during the 2018–2020 Ebola outbreak that began in the DRC.
• **August 11:** HHS announced up to $1.5 billion in funds to support the large-scale manufacturing and delivery of Moderna’s investigational vaccine candidate. Under the terms of the agreement, the U.S. Government will own the resulting 100 million doses of vaccine, and will have the option to acquire more. The vaccine, called mRNA-1273, has been co-developed by Moderna and scientists from the National Institute of Allergy and Infectious Diseases (NIAID), part of the National Institutes of Health. NIAID has continued to support the vaccine’s development including nonclinical studies and clinical trials. Additionally, BARDA has supported phase 2/3 clinical trials, vaccine manufacturing scale up and other development activities for this vaccine. The Phase 3 clinical trial, which began July 27, is the first government-funded Phase 3 clinical trial for a COVID-19 vaccine in the United States.

• **August 23:** As part of the agency’s efforts to combat COVID-19, the FDA issued an emergency use authorization (EUA) for investigational convalescent plasma. Based on available scientific evidence, the FDA determined convalescent plasma may be effective in lessening the severity or shortening the length of COVID-19 illness in hospitalized patients, and that the known and potential benefits of the product outweigh the known and potential risks. The EUA authorizes the distribution of convalescent plasma in the U.S. as well as its administration by health care providers, as appropriate, to treat suspected or confirmed cases of COVID-19. Click here to learn more about EUAs.

As announced on May 15, the vaccine development plan is as follows, subject to change as work proceeds:

• **Fourteen** promising candidates have been chosen from the 100+ vaccine candidates currently in development—some of them already in clinical trials with U.S. government support.

• **The 14** vaccine candidates are being narrowed down to about seven candidates, representing the most promising candidates from a range of technology options (nucleic acid, viral vector, protein subunit), which will go through further testing in early-stage clinical trials.

• **Large-scale** randomized trials for the demonstration of safety and efficacy will proceed for the most promising candidates.

**MANUFACTURING:** The federal government is making investments in the necessary manufacturing capacity at its own risk, giving firms the confidence to invest aggressively in development which will allow faster distribution of an eventual vaccine. Manufacturing capacity for selected candidates will be advanced while they are still in development, rather than scaled up after approval or authorization. Manufacturing capacity developed will be used for whatever vaccine is eventually successful, if possible given the nature of the successful product, regardless of which firms have developed the capacity.

Select actions to support OWS manufacturing efforts so far include:

• **The May 21, April 16, and March 30** HHS agreements with AstraZeneca, Moderna, and Johnson & Johnson respectively include investments in manufacturing capabilities.

• **June 1:** HHS announced a task order with Emergent BioSolutions to advance domestic manufacturing capabilities and capacity for a potential COVID-19 vaccine as well as therapeutics, worth approximately $628 million, using Emergent’s BARDA-supported Center for Innovation in Advanced Department and Manufacturing.
• **July 27:** HHS announced a task order with Texas A&M University and FUJIFILM to advance domestic manufacturing capabilities and capacity for a potential COVID–19 vaccine, worth approximately $265 million, using another BARDA–supported CIADM.

• **August 4:** Grand River Aseptic Manufacturing Inc., (GRAM) Grand Rapids, Michigan, was awarded a $160 million firm–fixed–price contract for domestic aseptic fill and finish manufacturing capacity for critical vaccines and therapeutics in response to the COVID–19 pandemic.

**DISTRIBUTION:** Before the countermeasures are approved or authorized, the program will build the necessary plans and infrastructure for distribution.

HHS plans for a tiered approach to vaccine and therapeutic distribution, which will build on allocation methodology developed as part of pandemic flu planning and be adjusted based on experience from the COVID–19 response so far, data on the virus and its impact on populations and the performance of a given countermeasure, and the needs of the essential workforce. OWS will expand domestic manufacturing and supplies of specialized materials and resources, such as glass vials, that can be necessary for distribution. DoD’s involvement will enable faster distribution and administration than would have otherwise been possible.

Select actions to support OWS vaccine and therapeutic development so far include:

• **May 12:** DoD and HHS announced a $138 million contract with ApiJect for more than 100 million prefilled syringes for distribution across the United States by year–end 2020, as well as the development of manufacturing capacity for the ultimate production goal of over 500 million prefilled syringes in 2021.

• **June 9:** HHS and DoD announced a joint effort to increase domestic manufacturing capacity for vials that may be needed for vaccines and treatments.

• **June 11:** HHS announced $204 million in funds to Corning to expand the domestic manufacturing capacity to produce approximately 164 million Valor Glass vials per year if needed. Valor Glass provides chemical durability to minimize particulate contamination. The specialized glass allows for rapid filling and capping methods that can increase manufacturing throughput by as much as 50 percent compared with conventional filling lines, which in turn can reduce the overall manufacturing time for vaccines and therapies.

• **June 11:** HHS announced $143 million to SiO2 Materials Science to ramp up capacity to produce the company’s glass–coated plastic container, which can be used for drugs and vaccines. The new lines provide the capacity to produce an additional 120 million vials per year if needed.

• **August 14:** HHS and DoD announced that McKesson Corporation will be a central distributor of future COVID–19 vaccines and related supplies needed to administer the pandemic vaccinations. The CDC is executing an existing contract option with McKesson to support vaccine distribution. The company also distributed the H1N1 vaccine during the H1N1 pandemic in 2009–2010. The current contract with McKesson, awarded as part of a competitive bidding process in 2016, includes an option for the distribution of vaccines in the event of a pandemic. Detailed planning is underway to ensure rapid distribution as soon as the FDA authorizes one or more vaccines. Once these decisions are made, McKesson will work under CDC’s guidance to ship COVID–19 vaccines to administration sites.
Who’s leading Operation Warp Speed?
HHS Secretary Alex Azar and Defense Secretary Mark Esper oversee OWS, with Dr. Moncef Slaoui designated as chief advisor and General Gustave F. Perna confirmed as the chief operating officer. To allow these OWS leaders to focus on operational work, in the near future the program will be announcing separate points of contact, with deep expertise and involvement in the program, for communication with Congress and the public.

What are you doing to make these products affordable for Americans?
The Administration is committed to providing free or low-cost COVID-19 countermeasures to the American people as fast as possible. Any vaccine or therapeutic doses purchased with US taxpayer dollars will be given to the American people at no cost.

How is Operation Warp Speed being funded?
Congress has directed almost $10 billion to this effort through supplemental funding, including the CARES Act. Congress has also appropriated other flexible funding. The almost $10 billion specifically directed includes more than $6.5 billion designated for countermeasure development through BARDA and $3 billion for NIH research.