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For much of my four-decade career, I’ve worked with biotech companies trying to solve significant health challenges, including HIV. Novavax is uniquely positioned to help end this pandemic with our protein-based vaccine. I believe in NVX-CoV2373 so much, I came out of retirement to join the fight.
Our COVID-19 Vaccine: NVX-CoV2373

We responded to the COVID-19 pandemic at an unprecedented pace, engineering a recombinant protein subunit-based vaccine, NVX-CoV2373, within one month of the SARS-CoV-2 sequence being published in early 2020.

Novavax’ proven history of vaccine experience, as well as promising preclinical studies of NVX-CoV2373, enabled us to secure critical funding in 2020 that fueled the development of our vaccine. By the end of the year, we successfully secured over $2 billion in funding from partners including the Coalition for Epidemic Preparedness Innovations (CEPI), the Bill and Melinda Gates Foundation, and the U.S. government through both Operation Warp Speed and the Department of Defense.

Our Clinical Achievements

These partnerships enabled us to rapidly advance NVX-CoV2373 through various clinical trials, including into three efficacy trials globally, as well as to scale up our manufacturing capabilities in parallel.

We have demonstrated remarkable results in terms of NVX-CoV2373’s safety and efficacy. In our UK Phase 3 clinical trial initiated in September 2020, we demonstrated 96% efficacy against the original strain of COVID-19 and 86% efficacy against the B.1.1.7 variant strain. In volunteers 65 years of age and older, 10 cases of COVID-19 were observed, with 90% of those cases occurring in the placebo group. This importantly demonstrates our ability to protect older adults who are among the groups most impacted by the disease and at high risk of complications from COVID-19.

In addition, we gathered critical data from our Phase 2b clinical trial initiated in South Africa in August 2020, when the B.1.351 variant was widely circulating. In this trial, we demonstrated efficacy of 55% in the portion of the study population that was HIV-negative, as well as efficacy of 49% in the overall trial population. There were no severe cases of disease in the vaccine arms of either of these trials. The observed efficacy against the original strain of COVID-19 and widely circulating variants drives continued confidence that NVX-CoV2373 can play a critical role in ending the COVID-19 pandemic on a global scale.

Our team works tirelessly every day to help the world return to hugging family members and seeing friends.
In the U.S. and Mexico, we initiated our PREVENT-19 pivotal Phase 3 trial in December of 2020. In early 2021, we completed enrollment of 30,000 participants. This study population represents individuals of Latin American, African American, Native American, and Asian American backgrounds, underscoring our commitment to serving diverse populations. We look forward to sharing additional updates as this trial progresses.
Our Global

Manufacturing and Supply Chain

To ready ourselves for the commercial launch of NVX-CoV2373, we took significant strides in building out our manufacturing capabilities. Throughout 2020 and into 2021, we built a global supply chain comprised of manufacturing sites and partners across over 10 countries. Through these efforts, our projected manufacturing capacity is over 2 billion annualized doses.

Global Supply Chain

This global supply chain includes Novavax owned facilities in the Czech Republic and Sweden, as well as partnerships with contract manufacturing organizations in the U.S., Canada, the UK and Spain. To ensure the widespread distribution of our vaccine globally, we also secured licensing agreements with Serum Institute of India (Serum), SK bioscience, and Takeda Pharmaceutical Company Limited (Takeda) for supply of NVX-CoV2373 in India, South Korea and Japan respectively.

In the year to come, we expect our expansive supply chain will enable us to deliver upon our supply commitments globally. We have seen continued demand for NVX-CoV2373 around the world, with supply agreements in place representing the potential for several billion dollars of revenue. As of the first quarter of 2021, we have secured advance purchase agreements for approximately 200 million doses of NVX-CoV2373, as well as committed 110 million doses to the U.S. government.
Building a global supply chain from scratch has not been easy, but the internal team, along with our amazing partners, have accomplished the impossible. We now have GMP facilities on three continents getting ready to produce NVX-CoV2373 to help end this pandemic. I could not be prouder of the team and our accomplishments.

Jose Torres-Vorshirm
VICE PRESIDENT, GLOBAL SUPPLY CHAIN & STRATEGIC SOURCING

Our Commitment to
Fair and Equitable Access of NVX-CoV2373

We continue to observe the reality that pandemics observe no borders. In light of this, we remain committed to ensuring fair and equitable access to our vaccine around the world. We have made it among our core values to ensure that those in economically disadvantaged countries have the opportunity to receive our vaccine in parallel with the rest of the world. Underscoring this commitment is our partnership with Serum, whereby we have jointly committed to increase our production capacity to over 2 billion doses annually. Our aim is that NVX-CoV2373 can address the vast global health need and reach countless individuals, regardless of country-specific income.
Our Dedication to Our Employees

Over the course of 2020, our employee base grew immensely. We began 2020 with approximately 150 employees and grew to over 800 employees during the first quarter of 2021. The tireless efforts and dedication of our employees spanning the Czech Republic, Sweden and the U.S. drove forward our achievements in 2020.

In 2020, we committed special efforts to support our employees during the COVID-19 pandemic. We encouraged all of our employees who were able to work from home to do so. Because the nature of our business required that some employees remain onsite, we adopted new safety protocols for our facilities to protect our essential employees. We also established an emergency relief fund for our employees whose immediate families had been materially and negatively impacted by the COVID-19 pandemic. After a challenging year in the midst of the pandemic, we were proud to have been recognized in the 2021 Top Workplaces USA list based upon employee surveys. We believe this award reflects our dedication to our employees and our investment in an exceptional culture.

At the heart of our culture is our commitment to diversity, equity and inclusion. We believe this enables us to create, develop and fully leverage the strengths of our workforce to meet our growth objectives. We recently completed an evidence-based analysis of the current state of diversity, equity and inclusion at Novavax. Through this analysis, our aim is to understand how to best create a culture of inclusion and build a sustainable strategy to drive continued progress moving forward.
Our Strategic Priorities

Looking to the months ahead, with so many of us longing for a return to normalcy, we believe we are at a critical juncture for the implementation of our vaccine globally. With that, we recognize that there is still much work to be done. We are focused on these three areas:

1. Obtaining regulatory authorizations for NVX-CoV2373 on a global basis
2. Bringing all of our manufacturing sites to full production capacity
3. Developing a next generation vaccine to address COVID-19 variant strains that are emerging

Our Pipeline

Although much of the past year has been committed to developing NVX-CoV2373, we recognize the continued need for innovative vaccines in other therapeutic areas, such as influenza, respiratory syncytial virus (RSV), and other emerging infectious diseases. In our NanoFlu program, we have taken steps to ensure we continue to advance NanoFlu in parallel with our COVID-19 activities. After announcing the successful completion of our pivotal Phase 3 clinical trial in the first quarter of 2020, we developed a leadership team solely dedicated to our influenza program.

Robust Vaccine Pipeline

<table>
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<th>Stage</th>
<th>Product</th>
<th>Matrix-M</th>
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<td>Phase 1</td>
<td>NanoFlu</td>
<td></td>
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<td>Phase 2</td>
<td>NanoFlu / NVX-CoV2373</td>
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<td>Phase 3</td>
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<td>NanoFlu / NVX-CoV2373 / RSV</td>
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<tr>
<td>Marketed</td>
<td>ResVax Older Adults Pediatrics</td>
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<td></td>
<td>MERS SARS Ebola</td>
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</tbody>
</table>
Our NanoFlu team has also begun exploring the potential for combination vaccines, including NanoFlu, NVX-CoV2373 and potentially RSV, which could be used in a post-pandemic setting. We see significant commercial opportunity for these combination vaccines, as well as an opportunity to drive forward our mission of providing critical solutions to address diseases plaguing the world. For other areas of our robust pipeline, we believe exciting opportunities for development lie ahead. We remain confident that our unique technology will enable us to rapidly respond to a breadth of emerging infectious diseases in the years to come.

I am incredibly proud of what we, as a company, have accomplished over this past year. I express my deepest gratitude to our shareholders for your continued support during this pivotal year for Novavax. Only through this continued support have we been able to achieve such incredible progress in 2020. We thank you and look forward to sharing significant milestones in the months and years ahead.

Stanley C. Erck
President and Chief Executive Officer

A Message From Our Chairman of the Board

To Our Shareholders:

2020 presented a critical opportunity for Novavax to execute on our mission and develop a life-saving vaccine to address the COVID-19 pandemic. That mission drove our remarkable progress throughout the year as we dedicated ourselves to addressing this vast global health threat.

We concentrated our efforts in 2020 on successfully developing our COVID-19 vaccine, NVX-CoV2373, and setting the stage for the successful delivery of our vaccine to the world. To accomplish this, we grew immensely as a company, with over 800 employees today and a truly global presence. With your support, Novavax has transitioned into a world-class biotechnology company rapidly preparing for commercialization. As a Board, we have in parallel dedicated efforts to ensure we are prepared to oversee these commercial preparations and partner effectively with our management team.

In 2020, we were thrilled to welcome David Mott, Gregg Alton, J.D., and Margaret G. McGlynn, R. Ph., to the Board as independent directors. These individuals bring to the Board extensive leadership experience across the pharmaceutical and vaccine industries. We believe these valuable additions to the Board offer the diverse perspectives and deep industry expertise required to ensure the success of Novavax, as well as the continued creation of value for our shareholders in the months and years to come.

As we reflect on this past year, we do so with our deepest gratitude for your continued support of Novavax over the years and excitement about the progress we will make in the years ahead. With your support, we look forward to delivering on key milestones in the years ahead and continuing to leverage our technology to address today’s most urgent global health needs.

James F. Young, Ph.D.
Chairman of the Board of Directors
UNITED STATES SECURITIES AND EXCHANGE COMMISSION
Washington, D.C. 20549

Form 10-K
☒ ANNUAL REPORT PURSUANT TO SECTION 13 OR 15(d) OF THE SECURITIES EXCHANGE ACT OF 1934
For the fiscal year ended December 31, 2020

OR
☐ TRANSITION REPORT PURSUANT TO SECTION 13 OR 15(d) OF THE SECURITIES EXCHANGE ACT OF 1934
For the transition period from _______ to _______

Commission File No. 000-26770

NOVAVAX, INC.
(Exact name of Registrant as specified in its charter)

Delaware 22-2816046
(State of incorporation) (I.R.S. Employer Identification No.)

21 Firstfield Road,
Gaithersburg, Maryland 20878
(Address of principal executive offices) (Zip Code)

Registrant’s telephone number, including area code: (240) 268-2000

Title of each class Trading Symbol Name of each exchange on which registered
Common Stock, Par Value $0.01 per share NVAX The Nasdaq Global Select Market

Securities registered pursuant to Section 12(b) of the Act:

Securities registered pursuant to Section 12(g) of the Act: Not Applicable

Indicate by check mark if the Registrant is a well-known seasoned issuer, defined in Rule 405 of the Securities Act. Yes ☒ No ☐

Indicate by check mark if the Registrant is not required to file reports pursuant to Section 13 or 15(d) of the Act. Yes ☐ No ☒

Indicate by check mark whether the Registrant: (1) has filed all reports required to be filed by Section 13 or 15(d) of the Securities Exchange Act of 1934 during the preceding 12 months (or for such shorter period that the Registrant was required to file such reports), and (2) has been subject to such filing requirements for the past 90 days. Yes ☒ No ☐

Indicate by check mark whether the registrant has submitted electronically every Interactive Data File required to be submitted pursuant to Rule 405 of Regulation S-T ($232.405 of this chapter) during the preceding 12 months (or for such shorter period that the registrant was required to submit such files). Yes ☒ No ☐

Indicate by check mark if disclosure of delinquent filers pursuant to Item 405 of Regulation S-K is not contained herein, and will not be contained, to the best of the Registrant’s knowledge, in definitive proxy or information statements incorporated by reference in Part III of this Form 10-K or any amendment to this Form 10-K. ☒

Indicate by check mark whether the registrant is a large accelerated filer, an accelerated filer, a non-accelerated filer, smaller reporting company, or an emerging growth company. See the definitions of “large accelerated filer,” “accelerated filer,” “smaller reporting company,” and “emerging growth company” in Rule 12b-2 of the Exchange Act. (Check one):

Large accelerated filer ☒ Accelerated filer ☐
Non-accelerated filer ☐ Smaller reporting company ☐
Emerging growth company ☐

If an emerging growth company, indicate by check mark if the registrant had elected not to use the extended transition period for complying with any new or revised financial accounting standards provide pursuant to Section 13(a) of the Exchange Act. ☐

Indicate by check mark whether the registrant has filed a report on and attestation to its management’s assessment of the effectiveness of its internal control over financial reporting under Section 404(b) of the Sarbanes-Oxley Act (15 U.S.C. 7262(b)) by the registered public accounting firm that prepared or issued its audit report. ☒

Indicate by check mark whether the Registrant is a shell company (as defined in Rule 12b-2 of the Exchange Act). Yes ☐ No ☒

The aggregate market value of the voting and non-voting common equity held by non-affiliates of the Registrant (based on the last reported sale price of Registrant’s common stock on June 30, 2020 on the Nasdaq Global Select Market) was approximately $5,078,700,000.

As of February 24, 2021, there were 73,858,882 shares of the Registrant’s common stock outstanding.

Documents incorporated by reference: Portions of the Registrant’s Definitive Proxy Statement to be filed no later than 120 days after the fiscal year ended December 31, 2020 in connection with the Registrant’s 2021 Annual Meeting of Stockholders are incorporated by reference into Part III of this Annual Report on Form 10-K to the extent indicated herein.
NOVAVAX, INC.

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CERTAIN DEFINITIONS

All references in this Annual Report on Form 10-K to “Novavax,” the “Company,” “we,” “us” and “our” refer to Novavax, Inc. and its wholly-owned subsidiaries, Novavax AB and Novavax CZ (formerly Praha Vaccines a.s.) (unless the context otherwise indicates).

NOTE REGARDING TRADEMARKS

Novavax™, NanoFlu™, Matrix-M™, Matrix™, Prepare™, Resolve™, and ResVax™ are trademarks of Novavax. Any other trademarks referred to in this Annual Report on Form 10-K are the property of their owners. All rights reserved. We do not intend our use or display of other companies’ trade names or trademarks to imply an endorsement or sponsorship of us by such companies, or any relationship with any of these companies.

FORWARD-LOOKING INFORMATION

This Annual Report on Form 10-K contains forward-looking statements that involve risks and uncertainties. As a result of many factors, such as those set forth under “Risk Factors” and elsewhere in this Annual Report on Form 10-K, our actual results may differ materially from those anticipated in these forward-looking statements. Please also see the disclaimer under the section titled “Management’s Discussion and Analysis of Financial Condition and Results of Operations.”

SUMMARY OF RISKS ASSOCIATED WITH OUR BUSINESS

Our business is subject to numerous risks which are discussed more fully under the heading “Risk Factors” in this Annual Report on Form 10-K. These risks include, but are not limited to, the following:

• We have a history of losses, and our future profitability is uncertain.

• We will continue to require significant funding to maintain our current level of operations and fund the further development of our vaccine candidates.

• Because our vaccine product development efforts depend on new and rapidly evolving technologies, we cannot be certain that our efforts will be successful.

• Although we have made rapid progress, the regulatory and commercial success of our COVID-19 vaccine candidate, NVX-CoV2373, remains uncertain. We may be unable to obtain regulatory approval or produce a successful vaccine in a timely manner, if at all.

• We are a biotechnology company and face significant risk in developing, manufacturing and commercializing our products.

• Because we depend on third-parties to conduct some of our laboratory testing and clinical trials, and a significant amount of our vaccine manufacturing and distribution, we may encounter delays in or lose some control over our efforts to develop and supply products.

• Many of our competitors have significantly greater resources and experience, which may negatively impact our commercial opportunities and those of our current and future licensees.

• There is significant competition in the development of a vaccine against COVID-19, influenza, and RSV, and we may never see returns on the significant resources we are devoting to our vaccine candidates.

• We have not completed the development of vaccine products, and we may not succeed in obtaining the FDA licensure necessary to sell such vaccine products.
The regulatory pathway for NVX-CoV2373 is continually evolving and may result in unexpected or unforeseen challenges.

We are conducting, and plan to conduct in the future, a number of clinical trials for NVX-CoV2373 at sites outside the United States, and the FDA may not accept data from trials conducted in such locations.

Even if regulatory approval is received for our vaccine candidates, the later discovery of previously unknown problems with a product, manufacturer or facility may result in restrictions, including withdrawal of the product from the market.

Our success depends on our ability to maintain the proprietary nature of our technology.

Our business may be adversely affected if we do not successfully execute our business development initiatives.

Servicing our 3.75% convertible senior unsecured notes due 2023 (the “Notes”) requires a significant amount of cash, and we may not have sufficient cash flow resources to pay our debt.

Because our stock price has been and will likely continue to be highly volatile, the market price of our common stock may be lower or more volatile than expected.

Litigation could have a material adverse impact on our results of operation and financial condition.

We or the third-parties upon whom we depend may be adversely affected by natural or man-made disasters or public health emergencies, such as the COVID-19 pandemic.
Product Pipeline

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<th>Therapeutic Area</th>
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<td>Main</td>
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<td></td>
<td>Variant Strain (Booster and or Brivet)</td>
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<td>Marketed</td>
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<td></td>
<td>NanoFlu / RSV</td>
<td>Main</td>
<td>Main</td>
<td>Main</td>
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<tr>
<td></td>
<td>NanoFlu / NVX-CoV2373 / RSV</td>
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<td><strong>ResVax®</strong> (Infants via Maternal Immunization)</td>
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<td><strong>Other Emerging Infectious Diseases</strong></td>
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</tbody>
</table>

(1) Supported by funding from the U.S. government partnership formerly known as Operation Warp Speed (“OWS”), U.S. Department of Defense (the “DoD”), Coalition for Epidemic Preparedness Innovations (“CEPI”) and the Bill & Melinda Gates Foundation (“BMGF”)

(2) Ongoing PREVENT-19, a Phase 3 clinical trial in U.S. and Mexico; Ongoing Phase 3 in UK; Ongoing Phase 2b in South Africa

(3) Supported by a grant from BMGF

Technology Overview

Recombinant Nanoparticle Vaccine Technology

Novavax’ recombinant nanoparticle vaccines combine the power and speed of genetic engineering to efficiently produce a new class of highly immunogenic vaccines that target a variety of viral pathogens.

Once a pathogenic threat has been identified, the genetic sequence encoding the antigen is selected for subsequent use in developing the vaccine construct. The genetic sequence may be optimized to enhance protein stability or confer resistance to degradation. This genetic construct is inserted into the baculovirus Spodoptera frugiperda (Sf9/BV) insect cell-expression system, which enables efficient, large-scale expression of the optimized protein. The Sf9/BV system produces proteins that are properly folded and modified – which can be critical for functional, protective immunity – as the vaccine antigen. Protein antigens are purified and organized around a polysorbate-based nanoparticle core, in a configuration that resembles their native presentation. This presentation results in a highly immunogenic nanoparticle that is ready to be formulated with Matrix-M adjuvant.

Matrix-M Adjuvant

Matrix-M is composed of 40-nanometer particles derived from saponin extracted from the bark of the Quillaja saponaria Molina tree. Once purified, these particles are fused with a unique formulation of cholesterol and phospholipid. This proprietary adjuvant has demonstrated potent and well-tolerated efficacy by stimulating the entry of antigen presenting cells (APCs) into the injection site and enhancing antigen presentation in local lymph nodes, which in turn activates T-cell, B-cell, and APC populations, thereby boosting immune response. Matrix-M has been shown to increase neutralizing antibodies and induces long-lasting memory B-cells, which increases B-cell immunity and recruits and increases the frequency of CD4+ and CD8+ T-cells to enhance T-cell immunity. The potent immune-stimulating mechanism of action is designed to enable a lower dose of antigen required to achieve the desired immune response, ultimately contributing to increased supply and manufacturing capacity. These immune-boosting and dose-sparing capabilities contribute to the adjuvant’s highly unique profile.

To date, we have formulated many of the vaccine candidates in our pipeline with Matrix-M, including NVX-CoV2373 and NanoFlu. Matrix-M has been well tolerated in human studies to date.

Pipeline Overview

As the world continues to address the global COVID-19 pandemic, we remain focused today on bringing our NVX-CoV2373 vaccine candidate to market following global regulatory approvals. In addition to this focus, NanoFlu continues to be a priority for our team, especially as it relates to a potential combined NanoFlu/NVX-CoV2373 vaccine. Although NVX-CoV2373 and NanoFlu are our near-term priorities, we remain optimistic that the additional programs in our pipeline including our vaccine candidates for RSV and other emerging infectious diseases, present viable opportunities for future development.

Coronavirus

Coronaviruses (“CoV”), so named for their “crown-like” appearance, are a large family of viruses, some of which are believed to have spread from animals to humans. These viruses cause human diseases such as Middle East Respiratory Syndrome (“MERS”), Severe Acute Respiratory Syndrome (“SARS”), and COVID-19, the disease resulting from the SARS CoV-2 coronavirus. COVID-19 first emerged in late 2019 in China, and, as of March 2020, the World Health Organization declared it a global pandemic.

NVX-CoV2373

We have successfully produced NVX-CoV2373, designed to provide protection against SARS-CoV-2. We engineered NVX-CoV2373 from the genetic sequence of SARS-CoV-2, using our recombinant nanoparticle technology to generate the antigen derived from the coronavirus spike (S) protein. NVX-CoV2373 includes our proprietary Matrix-M adjuvant.
NVX-CoV2373 Preclinical Development

In April 2020, we announced that NVX-CoV2373 demonstrated high immunogenicity in animal models measuring spike protein-specific antibodies, antibodies that block the binding of the spike protein to the receptor and high levels of wild-type virus neutralizing antibodies.

NVX-CoV2373 Clinical Development

PREVENT-19 Phase 3 U.S. and Mexico

In February 2021, we completed enrollment of our PREVENT-19 pivotal Phase 3 study in the U.S. and Mexico initiated in December 2020. PREVENT-19 is a randomized, placebo-controlled, observer-blinded study to evaluate the efficacy, safety and immunogenicity of NVX-CoV2373 with Matrix-M adjuvant that enrolled more than 30,000 participants aged 18 years or older. The trial largely reached its demographic goal of enrolling participants at high-risk for COVID-19 including adults over the age of 65, people with medical comorbidities and racial/ethnic subgroups who are at greater risk of infection and disease. The participant study population is composed of the following: 20% LatinoX, 13% African American, 6% Native American, 5% Asian American, and 13% older adults aged 65 years and older. The trial design has been harmonized to align with other Phase 3 trials conducted under the auspices of OWS, including the use of a single external independent Data and Safety Monitoring Board to evaluate safety and conduct an unblinded review when predetermined interim analysis events are reached. The trial’s primary endpoint is the prevention of PCR-confirmed, symptomatic COVID-19. The primary and secondary endpoints will be assessed at least seven days after the second study vaccination in volunteers who have not been previously infected with SARS-CoV-2. Two-thirds of the participants will be assigned to randomly receive two intramuscular injections of the vaccine, administered 21 days apart, while one-third of the trial participants will receive placebo. The primary efficacy analysis is event-driven, based on the number of participants with symptomatic mild, moderate or severe COVID-19 disease, occurring between the first and second dose for 24 months following the second injection. Dependent on the overall COVID-19 attack rate, interim data in this event-driven trial are expected in the second quarter of 2021. PREVENT-19 is being conducted with support from OWS, including a $1.7 billion agreement.

Phase 3 United Kingdom (“UK”)

In January 2021, we announced that NVX-CoV2373 demonstrated 89.3% efficacy in an interim analysis of our Phase 3 UK study initiated in September 2020. Our Phase 3 study in the UK was in partnership with the UK Government’s Vaccines Taskforce. The trial was a randomized, placebo-controlled, observer-blinded study to evaluate the efficacy, safety and immunogenicity of NVX-CoV2373 in over 15,000 enrolled participants aged 18 to 84 years, including 27 percent of participants over the age of 65. Half of the trial participants received two intramuscular injections of NVX-CoV2373 comprising 5 micrograms of antigen with 50 micrograms of Matrix-M, administered 21 days apart, while the other half of the trial participants received placebo. The primary endpoint was first occurrence of PCR-confirmed, symptomatic COVID-19 with onset at least seven days after the second study vaccination in volunteers who have not been previously infected with SARS-CoV-2. Preliminary analysis indicates that the UK variant strain that was increasingly prevalent was detected in over 50% of the PCR-confirmed symptomatic cases. Interim results from this trial showed that efficacy by strain was 95.6% against the original COVID-19 strain (i.e., non-variant), while the subsequent infections during the study were largely attributable to the variant virus. These data suggest that prior infection with COVID-19 may not completely protect against subsequent infection by the South Africa escape variant, however, vaccination with NVX-CoV2373 provided significant protection.

Phase 1/2 U.S. and Australia

In August 2020, we announced positive preliminary immunogenicity and safety results from our Phase 1 portion of the Phase 1/2 clinical trial of NVX-CoV2373 initiated in May 2020. The Phase 1 portion was a randomized, observer-blinded, placebo-controlled trial in 131 participants at two sites in Australia. The trial was designed to evaluate the immunogenicity and safety of NVX-CoV2373, both adjuvanted with Matrix-M and unadjuvanted. The protocol’s two-dose trial regimen assessed two dose levels (5 and 25 micrograms) with Matrix-M and without. Results from this trial showed that NVX-CoV2373 was generally well-tolerated with induced robust antibody responses numerically superior to that seen in human convalescent serum and induced robust polyfunctional CD4+ T-cell responses. In September 2020, the Phase 1 portion clinical results were published in The New England Journal of Medicine. In January 2021, we reported favorable six-month immunogenicity (IgG ELISA) data during the J.P. Morgan 39th Annual Healthcare Conference. In August 2020, we initiated the Phase 2 portion of the Phase 1/2 clinical trial. The Phase 2 portion is designed to evaluate the safety and immunogenicity of NVX-CoV2373 with Matrix-M in participants aged 18 to 84 years. The Phase 2 portion will assess two dose levels (5 and 25 micrograms), each with 50 micrograms of Matrix-M. We completed enrollment of 1,288 healthy volunteers in October 2020, with approximately 50% of participants 60 years of age and older, at up to 40 sites in the U.S. and Australia. In late October 2020, we reported favorable preliminary reactivity data from the Phase 2 portion of the trial during the Centers for Disease Control and Prevention Advisory Committee on Immunization Practices meeting. As of March 2021, some participants from this trial are receiving a six-month boost dose of NVX-CoV2373 to examine the functional immune response of our vaccine candidate.

NVX-CoV2373 Clinical Development Conducted by Partner

Phase 1/2 Japan

In February 2021, Takeda Pharmaceutical Company Limited (“Takeda”) initiated a Phase 1/2 clinical trial of NVX-CoV2373 in Japan. This placebo-controlled trial will evaluate the immunogenicity and safety in 200 participants aged 20 years and older. Variant Strain (Booster and/or Bivalent)

In January 2021, we initiated development of new constructs against the emerging strains of COVID-19, and in February 2021, we selected candidates for variant strain vaccines as standalone and bivalent candidates. We are currently evaluating these candidates in ongoing non-human primate studies and plan to begin clinical evaluation of variant vaccine candidates in mid-2021.

NVX-CoV2373 Regulatory and Licensure

In February 2021, we announced the initiation of a rolling submission with non-clinical data to the UK Medicines and Healthcare products Regulatory Agency (“MHRA”). We expect to file for authorization by early second quarter of 2021 after we have gathered sufficient data.
In November 2020, we announced that the U.S. Food and Drug Administration ("FDA") granted NVX-CoV2373 Fast Track designation, which is intended for products that treat serious or life-threatening diseases or conditions that demonstrate the potential to address unmet medical needs for such diseases or conditions. The Fast Track program is designed to facilitate development and expedite the review of drugs to treat serious conditions, with the intent of providing patients with earlier access to important new drugs. Specifically, Fast Track designation facilitates meetings with the FDA to discuss all aspects of development to support licensure and provides the opportunity to submit sections of a Biologics Licensing Application ("BLA") on a rolling basis as data become available. We continue to be in communication with the FDA through submissions to our open investigational new drug application ("IND") and discussions on various aspects of the program required to support the regulatory approval process. We also plan to file submissions for Emergency Use Authorization ("EUA") with the FDA and expect to complete our EUA filing in the second quarter of 2021.

In addition, we initiated the rolling review process with submissions to several regulatory agencies worldwide, including the European Medicines Agency ("EMA"), Health Canada, Australian Therapeutic Goods Administration ("TGA"), and New Zealand Medsafe. As part of the rolling review, we will continue to submit additional information, including clinical and manufacturing data as they become available. These rolling reviews are initiated to expedite the assessment of vaccines, particularly during public health emergencies.

COVID-19 Vaccine Funding

In May 2020, we signed a restated funding agreement which was amended in November 2020, with CEPI (the "CEPI Funding Agreement"), under which we are entitled to receive funding of up to $599.5 million to be used by us for the development of NVX-CoV2373. Pursuant to the CEPI Funding Agreement, if approved, a portion of the NVX-CoV2373 supply produced by us, other than vaccine manufactured under the OWS Agreement (as defined below), is expected to be purchased and allocated through the COVAX Facility component of the Access to COVID-19 Tools (ACT) Accelerator, an international equitable vaccine purchasing initiative launched by the World Health Organization, Gavi the Vaccine Alliance ("Gavi"), CEPI and other global non-governmental organizations and governmental leaders in 2020.

In June 2020, we were awarded a contract by the DOD, which was last amended in January 2021, under which we are entitled to receive funding of up to $45.7 million to support certain activities related to the development of NVX-CoV2373, including the manufacturing and delivery of 10 million doses of NVX-CoV2373 to the U.S. government.

In July 2020, we were selected to participate in OWS, a U.S. government sponsored program working to accelerate the development, manufacturing and distribution of COVID-19 vaccines, therapeutics and diagnostics. Through a Base Agreement and a Project Agreement (together, the "OWS Agreement") entered into with Advanced Technology International, Inc., the Consortium Management Firm acting on behalf of the Medical CBRN Defense Consortium in connection with OWS, which was last amended in December 2020, we have been allotted funding of $1.6 billion and are entitled to receive maximum funding up to $1.75 billion to support certain activities related to the development of NVX-CoV2373, and including the manufacture and delivery of 100 million doses of NVX-CoV2373 to the U.S. government. We expect this funding will assist in rapidly developing our large-scale manufacturing capacity and transitioning into ongoing production, including the capability to stockpile and distribute large quantities of NVX-CoV2373 for use in clinical trials and potentially for commercial sale, if authorized for emergency use or licensed. The OWS Agreement will fund the late-stage clinical studies necessary to determine the safety and efficacy of NVX-CoV2373, including PREVENT-19. Funding under the OWS Agreement is also expected to support our plans to file submissions for EUA and licensure with the FDA.

NVX-CoV2373 Manufacturing and Supply

In 2020, we established a global supply chain to support the commercialization of NVX-CoV2373. The acquisition of Praha Vaccines a.s. ("Praha Vaccines") in the Czech Republic in May 2020 demonstrated the Company’s first major step toward building out our global manufacturing capabilities. Since May 2020, we have established partnerships worldwide to amplify and solidify our global reach.

In August 2020, we expanded upon our manufacturing and supply capabilities to include partnerships with both Takeda in Japan and SK Bioscience Co., Ltd. ("SK bioscience") in South Korea and furthered these collaborations in February 2021. These additional partnerships will further increase our production capacity and are expected to support a rapid roll-out of NVX-CoV2373 globally.

To date, we have increased our projected global manufacturing production rate of NVX-CoV2373 to be over two billion annualized doses when we are at full capacity, which we expect to occur in mid-2021. Of this anticipated capacity, approximately one billion doses will be manufactured by SIIPL.

NVX-CoV2373 and its components are being manufactured at the following Novavax (in bold) and partnered sites:

- **Novavax CZ**
  - Biofabri S.L. in Spain
  - FLUFLM Deltorphin Biotechnologies ("FDB") in North Carolina and Texas in the U.S.
  - FDB in the UK
  - National Research Council’s Biologics Manufacturing Centre in Canada

- **Matrix-M Adjuvant**
  - AAGB Biologies in the U.S. and Denmark
  - PolyPeptide Group (will manufacture two key components used in Matrix-M) in the U.S. and Sweden

- **Fill/Finish Activities**
  - Baxter International Inc. in Germany
  - Jublant HollisterStier LLC in the U.S.
  - Par Pharmaceutical Companies, Inc. in the U.S.
  - Siegfried AG in Germany

- **Antigen Production, Out-licensing & Collaborations**
  - SIIPL in India
  - SK bioscience in the Republic of Korea
  - Takeda in Japan

A summary and status of key manufacturing and supply developments follows:

In February 2021, we announced a Memorandum of Understanding ("MOU") with Gavi to provide 1.1 billion doses of NVX-CoV2373 for the COVAX Facility. The vaccine doses will be manufactured and distributed globally by us and SIIPL, the latter under an existing agreement between Gavi and SIIPL. We expect to work with Gavi, which leads the design and implementation of the COVAX Facility, to finalize an advance purchase agreement for vaccine supply and global distribution via the COVAX Facility and its partners. We have licensed our NVX-CoV2373 technology to SIIPL and are jointly committed with SIIPL to deliver the 1.1 billion doses to the COVAX Facility. We expect to supply doses to primarily high-income countries, with SIIPL providing the majority of supply for low-, middle-, and upper-middle-income countries, utilizing a tiered pricing schedule.

In February 2021, we also reached a MOU with the Canadian government to produce NVX-CoV2373 in Canada. We plan to produce NVX-CoV2373 at the National Research Council’s Biologics Manufacturing Centre in Montreal once both the vaccine candidate and the facility receive Health Canada approvals. The MOU also includes a broader intention for the Government of Canada and us to work together to increase our Canadian presence. We will explore a range of partnership opportunities for us to expand vaccine production in Canada, including partnerships with Canadian contract manufacturers. We recently initiated the rolling submission process for regulatory approval to Health Canada.

In August 2020, we entered into a development and supply agreement with SK bioscience, and in February 2021, announced an expanded collaboration and license agreement with SK bioscience. Under these agreements, SK bioscience has been granted an exclusive license to develop, manufacture and commercialize NVX-CoV2373 in the Republic of Korea.
In August 2020, we announced a collaboration agreement with Takeda, and in February 2021, we finalized an exclusive license agreement with Takeda for the development, manufacturing and commercialization of NVX-CoV2373 in Japan. We will transfer technology and supply our Matrix-M adjuvant to Takeda, who will manufacture the vaccine antigen. Takeda will receive funding from the Government of Japan’s Ministry of Health, Labour and Welfare to support the technology transfer, establishment of infrastructure and scale-up of manufacturing. We anticipate that Takeda has a manufacturing capacity of over 250 million doses per year. We will be entitled to receive payments based on the achievement of certain development and commercial milestones, as well as a portion of net profits from vaccine sales. Takeda is responsible for regulatory submission to Japan’s Pharmaceutical and Medical Devices Agency (“PMDA”).

In July 2020, we announced a manufacturing agreement with FDB allowing for the large-scale contract production of NVX-CoV2373 in connection with our OWS Agreement, beginning at FDB’s North Carolina facility.

Also in July 2020, we entered into a supply and license agreement with SIIPL, as amended by the parties in September 2020, under which we granted exclusive and non-exclusive licenses to SIIPL for the development, co-formulation, filling and finishing, registration and commercialization by SIIPL of NVX-CoV2373. SIIPL has agreed to purchase Matrix-M adjuvant from us and we have granted SIIPL a non-exclusive license to manufacture the antigen drug substance component of NVX-CoV2373 in SIIPL’s licensed territory solely for use in the manufacture of NVX-CoV2373 under the terms of the agreement. We will equally split with SIIPL the revenue from SIIPL’s sale of NVX-CoV2373 in its licensed territory, net of agreed costs. We granted to SIIPL (i) an exclusive license in India during the agreement, and (ii) a non-exclusive license (a) during the “Pandemic Period” (as declared by the WHO), in all countries other than specified countries designated by the World Bank as upper-middle or high-income countries, with respect to which we retain rights, and (b) after the Pandemic Period, in only those countries designated as low or middle-income by the World Bank. Following the Pandemic Period, we may notify SIIPL of any bona fide opportunities for us to license NVX-CoV2373 to a third-party in such low and middle- income countries and SIIPL would have an opportunity to match or improve such third-party terms, failing which, we would have the discretion to remove one or more non-exclusive countries from SIIPL’s license. We anticipate SIIPL will manufacture approximately one billion doses of NVX-CoV2373 in 2021.

In June 2020, we entered into contract manufacturing arrangements with AGC Biologics and the Polypeptide Group to provide contract development and manufacturing services, supplying us with large-scale production of Matrix-M.

In May 2020, we announced the acquisition of Praha Vaccines, formerly part of the Cyrus Poonawalla Group, in an all cash transaction of approximately $167 million. The acquisition includes a biologics manufacturing facility and associated assets in Bohumil, Czech Republic. The acquisition included a 150,000-square foot state-of-the-art vaccine and biologics manufacturing facility and other support buildings, along with the existing employees and all related and required infrastructure. The facility is expected to provide annual capacity of over 1 billion doses of antigen starting in 2021 for NVX-CoV2373. The facility is completing a renovation that includes Biosafety Level-3 (BSL-3) capabilities. As part of the transaction, approximately 150 employees with significant experience in vaccine manufacturing and support joined Novavax. The acquisition of Praha Vaccines was supported by our funding arrangements with CEPI, which we expect will enable us to dramatically expand our manufacturing capacity.

Concurrently, SK bioscience finalized an advance purchase agreement with the Republic of Korea to supply 40 million doses of NVX-CoV2373 beginning in 2021. SK bioscience will expand its capacity to manufacture the antigen component of NVX-CoV2373 for use in the final drug product globally, including product distributed by the COVAX Facility, during the COVID-19 pandemic. SK bioscience will also purchase a certain quantity of the finished vaccine product directly from us, subject to the approval by relevant regulatory authority, and sufficient doses of our Matrix-M adjuvant to manufacture the remainder of the 40 million doses of finished vaccine product SK bioscience expects to sell to the Korean government. SK bioscience will pay a tiered royalty in the low to middle double-digit range on the sale of NVX-CoV2373 in the Republic of Korea, net of certain agreed costs.

In October 2020, we entered into a SARS-CoV-2 vaccine supply agreement with The Secretary of State for Business, Energy and Industrial Strategy, acting on behalf of the government of the UK (the “Authority”), for the purchase of 60 million doses of NVX-CoV2373, plus such additional orders as the Authority may make from time to time. We agreed to continue to conduct a UK-based Phase 3 clinical trial of NVX-CoV2373 to assess the efficacy of NVX-CoV2373 in the UK population, establish a dedicated supply chain for NVX-CoV2373 in the UK and seek regulatory approval for the NVX-CoV2373 in the UK. FDB’s UK site is expected to produce up to 180 million doses annually. Excess supply of antigen manufactured at the FDB’s site in Billingham, Stockton-on-Tees may be available for us to sell to additional markets outside the UK.

In January 2021, we finalized an APA with the Government of Canada to supply up to 76 million doses of NVX-CoV2373. Canada has committed to purchase 52 million doses of NVX-CoV2373 with the option for up to an additional 24

### NVX-CoV2373 Supply Agreements

We have entered into advance purchase agreements (referred to as “APAs” or “supply agreements” throughout this Annual Report on Form 10-K) with various countries globally that, if our product candidate is approved, are expected to result in the delivery of approximately 200 million doses of NVX-CoV2373 throughout 2021 and into the first half of 2022. The APAs typically contain terms that include upfront payments intended to assist us in funding investments related to building out and operating our manufacturing and distribution network, among other expenses, in support of our global supply commitment. Such upfront payments generally become non-refundable upon our achievement of certain development milestones. We expect to sign additional APAs that are currently in active discussions and negotiations.

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<tr>
<th>Country</th>
<th>Committed Doses &amp; Additional Detail</th>
<th>Supply Agreements by Region</th>
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<tr>
<td>UK</td>
<td>Option to purchase additional orders from time to time</td>
<td>North America</td>
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<tr>
<td>Canada</td>
<td>Option to purchase up to an additional 24 million doses</td>
<td>Europe, Middle East and Africa</td>
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<td>Switzerland</td>
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<td>Summary</td>
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million doses. Under the agreement, we expect to supply doses of NVX-CoV2373 to Canada following authorization by Canada’s regulatory agency.

In December 2020, we finalized an APA with the Australian Federal Government to supply up to 61 million doses. Australia is committed to purchase 51 million doses of NVX-CoV2373 with the option for up to an additional 10 million doses. We will work with Australia’s regulatory agency, the Therapeutics Goods Administration (TGA), to seek product approvals upon demonstrating efficacy in clinical studies.

In December 2020, we finalized an APA with the government of New Zealand for the purchase of 10.72 million doses of NVX-CoV2373. Under the terms of the agreement, we will manufacture all doses of NVX-CoV2373 delivered to New Zealand under the agreement.

In February 2021, we executed a binding Heads of Terms with the government of Switzerland to supply six million doses of NVX-CoV2373. Following this Heads of Terms, we intend to negotiate a final agreement with Switzerland, with initial delivery of vaccine doses slated to ship following successful clinical development and regulatory review.

**Seasonal Influenza**

**NanoFlu Program (Older Adults)**

Influenza is a worldwide infectious disease with serious illness generally occurring in more susceptible populations such as children under and older adults, but also occurring in the general population. According to a 2019 Global Data forecast of influenza vaccines, the market for seasonal influenza vaccines is expected to grow from approximately $4.6 billion in 2018 to approximately $6.5 billion in 2028 (in the countries comprising the eight major markets).

In March 2020, we announced positive top-line results from our Phase 3 clinical trial of NanoFlu, which includes our proprietary Matrix-M adjuvant. In October 2020, we announced the formation of a leadership team within the Company focused on advancing NanoFlu to regulatory licensure. The leadership team has established a separate NanoFlu development unit within our Company, which benefits from joint shared services with key cross-functional departments within the Company and builds on the Company’s established knowledge base in the discovery and development of innovative vaccines to prevent serious infectious diseases.

We continue to seek regulatory approval from the FDA under the accelerated approval pathway previously granted to the Company and explore the potential for a combination NanoFlu/NVX-CoV2373 vaccine to be used in a post-pandemic setting.

**Combination Vaccines**

With the ongoing development of NanoFlu, NVX-CoV2373, and respiratory syncytial virus fusion (F) protein nanoparticle vaccine candidate (“RSV F Vaccine”), a strong rationale exists for potentially developing three combination respiratory vaccines designed to protect susceptible populations against these diseases. Although testing is at an early stage, we believe that combination vaccines against influenza in combination with COVID-19, influenza in combination with RSV, and influenza in combination with both RSV and COVID-19 may be achievable since these vaccines are created using our recombinant nanoparticle technology and include our proprietary Matrix-M adjuvant.

**Respiratory Syncytial Virus (“RSV”)**

Currently, there is no approved RSV vaccine available to combat the estimated 64 million RSV infections that occur globally each year. We have identified three susceptible target populations that we believe could benefit from the development of our respiratory syncytial virus fusion (F) protein nanoparticle vaccine candidate (“RSV F Vaccine”) in different formulations: (1) infants via maternal immunization, (2) older adults (60 years and older) and (3) children six months to five years old (“pediatrics”). With our current estimates of the annual global cost burden of RSV in excess of $88 billion, we believe our RSV F Vaccine represents a multi-billion-dollar worldwide opportunity.
Ebola Virus

EBOV is a filovirus that produces severe, often fatal illness in humans. Within the last decade, it has produced two large outbreaks in Sub-Saharan Africa with high mortality. There are currently two vaccines licensed to prevent EBOV.

We developed an EBOV glycoprotein vaccine candidate ("Ebola GP Vaccine") expressed in insect cells, using our core recombinant baculovirus technology. Although not in active development, our Ebola GP Vaccine is a viable development opportunity.

Competition in COVID-19, Influenza and RSV

The vaccine market is intensely competitive, characterized by rapid technological progress. Our technology is based upon utilizing the baculovirus expression system in insect cells to make recombinant vaccines. We believe this system offers many advantages when compared to other technologies and is uniquely well-suited for developing COVID-19, influenza, and RSV vaccines, as well as vaccines against a number of other infectious diseases.

A number of vaccine manufacturers, research institutions, and other organizations are developing a vaccine for SARS-CoV-2, the virus that causes COVID-19 disease. A variety of different vaccine technologies are being studied, including nucleic acid (RNA/DNA), viral vectors, live attenuated or inactivated, and protein-based vaccines. According to a coronavirus vaccine tracker published by The New York Times, there are 69 vaccines in clinical trials and 20 have reached the final stages of testing. As of February 2021, Pfizer, Moderna, and Johnson & Johnson have each received approval under Emergency Use Authorization by the FDA in the U.S. for their COVID-19 vaccines. NVX-CoV2373 is currently being evaluated in two pivotal Phase 3 trials: a trial in the UK that completed enrollment in November and the PREVENT-19 trial in the U.S. and Mexico that began in December. It is also being tested in two ongoing Phase 2 studies that began in August: a Phase 2b trial in South Africa, and a Phase 1/2 continuation in the U.S. and Australia. Based on the interim efficacy from the Phase 3 in the UK and the Phase 2b in South Africa, our vaccine candidate has demonstrated strong efficacy and will play an important role in solving this global public health crisis. Importantly, we are the first vaccine to demonstrate clinical efficacy against the original strain of COVID-19 and both of the rapidly emerging variants in the UK and South Africa.

A number of companies are developing and selling vaccines for seasonal influenza employing both traditional (egg-based) and new vaccine technologies (cell-based). Many seasonal influenza vaccines are currently approved and marketed, and most of these are marketed by major pharmaceutical companies such as Sanofi Pasteur, GSK and Seqirus. Competition in the sale of seasonal influenza vaccines is intense. For the older adult segment, Sanofi currently supplies Fluzone-HDI® and Flublok® to the majority of older adults in the U.S. Therefore, newly developed and approved products must be differentiated from existing vaccines in order to have commercial success. In order to show differentiation in the seasonal influenza market, a product may need to be more efficacious and/or less expensive and quicker to manufacture. Many of our competitors are working on new products and new generations of current products, some by adding an adjuvant that is used to increase the immunogenicity of that product, each of which is intended to be more efficacious than currently marketed products. Despite the significant competition and advancing technologies, some of which are similar to our own, based on our completed Phase 3 trial results, we believe that NanoFlu, our adjuvanted nanoparticle seasonal influenza product could be as efficacious as, or more so than, current products or products being developed by our competitors.

In general, competition among pharmaceutical products is based in part on product efficacy, safety, reliability, availability, price and patent position. An important factor is the relative timing of the market introduction of our products and our competitors' products. Accordingly, the speed with which we can develop products, complete the clinical trials and approval processes and supply commercial quantities of the products to the market is an important competitive factor. Our competitive position also may depend upon our ability to show differentiation with a product that is more efficacious and/or less expensive and quicker to manufacture. Other factors affecting our competitive position include our ability to attract and retain qualified personnel, obtain patent protection or otherwise develop proprietary products or processes and secure sufficient capital resources for the lengthy period between technological conception and commercial sale.

We generally seek patent protection for our technology and product candidates in the U.S. and abroad. The patent position of biotechnology and pharmaceutical firms generally is highly uncertain and involves complex legal and factual questions. Our success will depend, in part, on whether we can:

- obtain patents to protect our own technologies and product candidates;
- obtain licenses to use the technologies of third-parties, which may be protected by patents;
- protect our trade secrets and know-how; and
- operate without infringing the intellectual property and proprietary rights of others.

We have intellectual property (patents, licenses, know-how) related to our vaccines, manufacturing processes and other technologies. Currently, we have or have rights to over 450 U.S. patents and corresponding foreign patents and patent applications relating to vaccines and vaccine-related technologies.

Patents related to our VLP program include U.S. Patent No. 7,763,450, which covers, in part, the use of influenza gene sequences for high-yield production of consistent influenza VLP vaccines to protect against current and future seasonal and pandemic strains of influenza viruses. Corresponding European patent, European Patent No. 1644037 also covers this technology. U.S. Patent Nos. 8,080,255, 8,551,756, 8,506,967 and 8,592,197 are directed to methods of producing VLPs and inducing substantial immunity to an influenza virus infection by administering VLPs comprising HA and NA proteins, and our M1 protein derived from the avian influenza strain, A/Indonesia/5/05. Certain claims also encompass similar methods and compositions where the M1 protein is from a different strain of influenza virus than the influenza HA protein and the influenza NA protein. Related patent protection in Europe is provided by European Patent No. 2543084, which covers, in part, vaccine compositions containing VLPs that contain M1, HA, and NA proteins. Our VLP patent portfolio contains many other patents, including U.S. Patent Nos. 8,951,337, 8,992,939, 9,144,607, 9,050,290, 9,180,180, 9,381,239, 9,464,276, 9,474,799, and other patents in multiple ex-U.S. jurisdictions.

We also have been issued patents directed to other core programs, including our RSV and influenza programs. Issued patents directed to various aspects of the RSV program include U.S. Patent Nos. 8,715,692, 9,675,685, 9,731,000, 9,717,786, 10,022,437, and 10,426,829. Additional patents in the family include EP37009 in Europe, as well as others throughout the world. Patents related to our rabies program include 9,724,405 and 10,086,065 in the U.S., and EP2635257 and EP2464619 in Europe. Related patents have been granted in other world markets. Issued patents in our influenza nanoparticle program include U.S. Patent No. 10,426,829. In addition to our focus on vaccine programs, we also pursue patent protection for our Matrix Adjuvant program. Issued U.S. Patent Nos. 7,838,019, 9,205,147, 9,901,634, 8,312,922, and 10,725,764 provide examples of patents related to our Matrix Adjuvant program.

We pursue patents related to NVX-CoV2373, our COVID-19 vaccine candidate. Our applications include PCT/US2021/015220 and U.S. Serial No. 16,997,001, which the U.S. Patent Office has allowed.

We continue to prepare, file, and prosecute patent applications to provide broad and strong protection of our proprietary rights, including next generation applications focused on our RSV Program, our influenza nanoparticle program, and our adjuvant program.

The Federal Technology Transfer Act of 1986 and related statutory guidance encourages the dissemination of science and technology innovation. While our expired contract with the U.S. Department of Health and Human Services, (“DHHS”), Biomedical Advanced Research and Development Authority (“HHS BARDA”) provided us with the right to retain ownership in our inventions that may have arisen during performance of that contract, with respect to certain other collaborative research efforts with the U.S. government, certain developments and results that may have commercial potential are to be freely published, not treated as confidential, and we may be required to negotiate a license to developments and results in order to commercialize products. There can be no assurance that we will be able to successfully obtain any such license at a reasonable cost, or that such development and results will not be made available to our competitors on an exclusive or non-exclusive basis.
Trade Secrets

We also rely significantly on trade secret protection and confidentiality agreements to protect our interests. It is our policy to require employees, consultants, contractors, manufacturers, collaborators and other advisors to execute confidentiality agreements upon the commencement of employment, consulting or collaborative relationships with us. We also require confidentiality agreements from any entity that is to receive confidential information from us. With respect to employees, consultants and contractors, the agreements generally provide that all inventions made by the individual while rendering services to us shall be assigned to us as our property.

Government Regulations

The development, production and marketing of biological products, which include the vaccine candidates being developed by Novavax or our collaborators, are subject to regulation for safety, efficacy and quality by numerous governmental authorities in the U.S. and other countries. Although we focus on the U.S. regulatory process and the standards imposed by the FDA, the International Conference on Harmonisation (“ICH”) and other agencies because we believe meeting U.S. and ICH standards generally allows us to satisfy regulatory agencies in other countries where we intend to do business; however, we are mindful that expectations in some venues, notably in the European Union, differ to some degree and we take proactive steps to address such differences by maintaining regular filings and correspondence and attending regular meetings with many other non-U.S. regulatory agencies. In the U.S., the development, manufacturing, non-marketing of human pharmaceuticals and vaccines are subject to extensive regulation under the Federal Food, Drug, and Cosmetic Act, and biological products are subject to regulation under provisions of that act and the Public Health Service Act. The FDA not only assesses the safety and efficacy of these products but it also regulates, among other things, the testing, manufacture, labeling, storage, record-keeping, advertising and promotion of such products. The process of obtaining FDA licensure for a new vaccine is costly and time-consuming.

Vaccine clinical development follows the same general regulatory pathway as drugs and other biologics. Before applying for FDA licensure to market any new vaccine candidate, we expect to first submit an investigational new drug application (“IND”) that explains to the FDA, among other things, the results of preclinical toxicity testing conducted in laboratory animals, the method of manufacture, quality control tests for release, the stability of the investigational product and what we propose to do for human testing. At this stage, the FDA decides whether it is reasonably safe to move forward with testing the vaccine candidate in humans. We must then conduct Phase 1 clinical trials and larger-scale Phase 2 and 3 clinical trials that demonstrate the safety, immunogenicity and efficacy of our vaccine candidate to the satisfaction of the FDA. Following successful completion of all three phases of clinical development, a BLA can be submitted to the FDA requesting licensure of the vaccine for marketing based on the vaccine’s safety and efficacy. Similar pathways exist in Europe and other geographies.

The FDA will only approve a BLA if the vaccine is demonstrated to be safe, pure and potent. During the FDA’s review of a BLA, the proposed manufacturing facility undergoes a pre-approval inspection during which the FDA examines in detail the production of the vaccine, the manufacturing facility and the quality documentation related to the vaccine. Vaccine licensure also requires the provision of adequate product labeling to allow health care providers to understand the vaccine’s proper use, including its potential benefits and risks, to communicate with patients and parents, and to safely deliver the vaccine to the public. Until a vaccine is given to the general population, all potential adverse events cannot be anticipated. Thus, the FDA typically requires Phase 4 post-marketing clinical trials for vaccines after licensure to continue gathering safety, and sometimes effectiveness/efficacy data in the indicated and additional populations.

The Commissioner of the FDA may, under delegated authority from the Secretary of the DHHS, and under certain circumstances, issue an EUA that would permit the use of an unapproved medical product or unapproved use of an approved medical product to diagnose, treat, or prevent serious or life-threatening diseases or conditions when there are no adequate, approved, and available alternatives. When issuing an EUA, the FDA imposes conditions of authorization, with which the EUA holder must comply. Such conditions include, but may not be limited to, compliance with labeling, distribution of materials designed to ensure proper use, reporting obligations, and restrictions on advertising and promotion. The EUA is only effective for the duration of the public health emergency. The FDA may also revoke or modify an EUA sooner if, the criteria for issuance are no longer met or other circumstances make a revision or revocation appropriate to protect the public health or safety. For example, an EUA may be revoked when the FDA determines that the underlying public health emergency no longer exists or warrants such authorization, or for reasons such as significant adverse inspectional findings, reports of adverse effects linked to or suspected of being caused by the EUA product, or newly emerging data that may demonstrate the product may not be effective.

In order to ensure continuing safety, the FDA and most other non-U.S.-based regulatory agencies continue to oversee the production of vaccines even after the vaccine and manufacturing processes are approved. For example, monitoring of the vaccine and of production activities, including periodic facility inspections, must continue as long as the manufacturer holds a license for the product. Manufacturers may also be required to submit the results of their own tests for potency, safety and purity for each vaccine lot, if requested by the relevant regulatory agency. They may also be required to submit samples of each vaccine lot to the agency for testing.

In addition to obtaining FDA licensure for each product, each domestic manufacturing establishment must be registered with the FDA, is subject to FDA inspection and must comply with current Good Manufacturing Practices (“cGMP”) regulations. To supply products for use either in the U.S. or outside the U.S., including in Harmonisation U.S.” and manufacturing establishments, including third-party facilities, must comply with GMP regulations and are subject to periodic inspection by the FDA or by corresponding regulatory agencies in their home country.

The FDA has several programs designed to expedite the development and approval of drugs and biological products intended to treat serious or life-threatening diseases or conditions, including fast track designation, breakthrough therapy designation, priority review designation, and accelerated approval. First, the FDA may designate a product for Fast Track review if it is intended, whether alone or in combination with one or more other products, for the treatment of a serious or life-threatening disease or condition and demonstrates the potential to address unmet medical needs for such a disease or condition. For Fast Track products, sponsors may have more frequent interactions with the FDA and the FDA may initiate review of sections of a Fast Track product’s application before the application is complete. The FDA granted Fast Track Designation for NVX-CoV2373 in November 2020 and for NanoFlu, our recombinant quadrivalent seasonal influenza vaccine candidate, in January 2020.

Second, a product may be designated as a Breakthrough Therapy if it is intended, either alone or in combination with one or more other products, to treat a serious or life-threatening disease or condition and preliminary clinical evidence indicates that the product may demonstrate substantial improvement over existing therapies on one or more clinically significant endpoints. The FDA may hold meetings with the sponsor throughout the development process; provide timely advice to the product sponsor regarding development and approval; involve more senior staff in the review process; assign a cross-disciplinary project lead for the review team; and take other steps to design the clinical trials in an efficient manner.

Third, the FDA may designate a product for priority review if it is a product that treats a serious disease or life-threatening condition and, if approved, would provide a significant improvement in safety or effectiveness over available therapies. Significant improvement may be illustrated by evidence of increased effectiveness in the treatment of a condition, elimination or substantial reduction of a treatment-limiting product reaction, documented enhancement of patient compliance that may lead to improvement in serious outcomes, and evidence of safety and effectiveness in a new subgroup. A priority designation is intended to direct overall attention and resources to the evaluation of such applications, and, for a drug product (including a vaccine), to shorten the FDA’s goal for taking action on a marketing application from ten months to six months.

Fourth, a product may be eligible for accelerated approval, if it treats a serious or life-threatening condition and generally provides a meaningful advantage over available therapies. In addition, it must demonstrate an effect on a surrogate endpoint that is reasonably likely to predict clinical benefit or on a clinical endpoint that can be measured earlier than irreversible morbidity or mortality, or IMO that is reasonably likely to predict an effect on IMO or other clinical benefit. As a condition of approval, the FDA may require that a sponsor of a drug or biologic receiving accelerated approval perform adequate and well-controlled post-marketing clinical trials to confirm efficacy using a clinically meaningful endpoint, thereby confirming efficacy observed pre-approval using a surrogate endpoint. In June 2019, we announced that the FDA acknowledged that the accelerated approval pathway is available for NanoFlu.

In addition to regulatory approvals that must be obtained in the U.S., an investigational product is also subject to regulatory approval in other countries in which it is intended to be marketed. No such product can be marketed in a country until the regulatory authorities of that country have approved an appropriate marketing application. FDA licensure does not
effect on our sales, results of operations and financial condition. Third-party payers may also control access to, or manage utilization of, our products with various utilization management techniques. Decreases in third-party reimbursement for our product candidates or a decision by a third-purchasers. We may be obligated to provide rebates or offer discounts under government health programs or to government and private payers are increasingly challenging the price and examining the cost-effectiveness of medical products and services. In addition, significant uncertainty exists as to the reimbursement status of newly approved healthcare products. We may need to conduct expensive pharmacoeconomic studies in order to demonstrate the cost-effectiveness of our products. There may be significant delays in obtaining coverage and reimbursement for newly approved drugs, and coverage may be more limited than the indications for which the product is approved by the FDA or similar regulatory authorities outside the United States. Our product candidates may not be considered cost-effective at certain prices. Adequate third-party reimbursement may not be available in certain markets to enable us to maintain price levels sufficient to realize an appropriate return on our investment in product development. Third-party payers may also control access to, or manage utilization of, our products with various utilization management techniques. Decreases in third-party reimbursement for our product candidates or a decision by a third-party payer to not cover our product candidates could reduce physician utilization of our products and have a material adverse effect on our sales, results of operations and financial condition.

Within the U.S., if we obtain appropriate approval in the future to market any of our product candidates, those products could potentially be covered by various government health benefit programs as well as purchased by government agencies. The participation in such programs or the sale of products to such agencies is subject to regulation. In exchange for coverage, we may be obligated to provide rebates or offer discounts under government health programs or to government and private purchasers.

The U.S. and state governments continue to propose and pass legislation designed to reform delivery of, or payment for, health care, including initiatives to reduce the cost of healthcare. For example, in March 2010, the U.S. Congress enacted the Patient Protection and Affordable Care Act and the Health Care and Education Reconciliation Act ("Healthcare Reform Act") which includes changes to the coverage and reimbursement of drug products under government health care programs. Under the Trump administration, there were several efforts to modify or repeal all or certain provisions of the Healthcare Reform Act, and some modifications were implemented. Adoption of price controls and cost-containment measures and adoption of more restrictive policies in jurisdictions with existing controls and measures could further limit our net revenue and results.

Other legislative changes have been proposed and adopted in the United States since the Healthcare Reform Act was enacted. For example, through the process created by the Budget Control Act of 2011, there are automatic reductions of Medicare payments to providers of up to 2% per fiscal year, which went into effect in April 2013 and will remain in effect through 2030 due to subsequent legislative amendments contained in the Coronavirus Aid, Relief, and Economic Security Act, commonly referred to as the "CARES Act," which became law in December 2020. This legislation, the Centers for Medicare and Medicaid Services ("CMS"), issued an interim final rule that seeks to lower prescription drug prices by paying no more for certain Medicare Part B drugs than the lowest price paid for such drugs in certain other countries (the "Most Favored Nation Rule"). Under the rule, the lower payment rates for affected drugs would be phased in over a period of four years, beginning in 2021. The rule has been challenged by industry associations on a number of grounds. On December 28, 2020, the U.S. District Court for the Northern District of California issued a nationwide preliminary injunction in Biotechnology Innovation Organization v. Azar, No. 3:20-cv-08603, which preliminarily enjoins CMS from implementing the Most Favored Nation Rule. Given this preliminary injunction, the

Most Favorable Nation Rule was not implemented on January 1, 2021 and will not be implemented without further rule-making. However, this interim final rule or any similar type of reference pricing regulation could potentially harm our business if expanded to include our products.

Recently, there has been considerable public and government scrutiny in the U.S. of pharmaceutical pricing and proposals to address the perceived high cost of pharmaceuticals. There have been several recent state legislative efforts to address drug costs, which generally have focused on increasing transparency around drug costs or limiting drug prices or price increases. Adoption of new legislation at the federal or state level could affect demand for, or pricing of, our product candidates if approved for sale. It is also possible that additional governmental action will be taken in response to the COVID-19 pandemic. We cannot predict the ultimate content, timing or effect of any federal and state reform efforts. There is no assurance that federal or state health care reform will not adversely affect our future business and financial results.

Within the U.S., we may be subject to various federal and state laws pertaining to health care “fraud and abuse,” including anti-kickback laws and false claims laws, for activities related to future sales of any of our product candidates that may in the future receive regulatory and marketing approval. Anti-kickback laws generally prohibit a pharmaceutical manufacturer from soliciting, offering, receiving or paying any remuneration to generate business, including the purchase, prescription or use of a particular drug. Although the specific provisions of these laws vary, their scope is generally broad and there may not be regulations, guidance or court decisions that apply the laws to particular industry practices. There is therefore a possibility that our practices might be challenged under such anti-kickback laws. False claims laws, including the federal False Claims Act (“FCA”), prohibit anyone from knowingly and willingly presenting, or causing to be presented, any claims for payment for reimbursed drugs or services to third-party payers (including Medicare and Medicaid) that are false or fraudulent. Our activities relating to the sale and marketing of our products may be subject to scrutiny under these laws. Violations of fraud and abuse laws may be punishable by criminal or civil sanctions, including fines and civil monetary penalties, and exclusion from federal health care programs (including Medicare and Medicaid). In the U.S., federal and state authorities are paying increased attention to enforcement of these laws within the pharmaceutical industry and private individuals have been active in alleging violations of the laws and bringing suits on behalf of the government under the FCA. If we were subject to allegations concerning, or were convicted of violating, these laws, our business could be harmed.

On November 20, 2020, the DHHS published a Final Rule entitled “Removal of Safe Harbor Protection for Rebates to Plans or PBMs Involving Prescription Pharmaceuticals and Creation of New Safe Harbor Protection,” commonly referred to as the “Rebate Rule,” which amends the federal Anti-Kickback Statute discount safe harbor by eliminating protection for price concessions, including rebates, that are offered by pharmaceutical manufacturers to plan sponsors, or pharmacy benefit managers under contract with them, the Medicare Part D program and Medicare Advantage Plans, unless the price reduction is one required by law. Effective January 1, 2022, in advance of the calendar year 2022 Part D plan year, safe harbor protection will be eliminated for manufacturer rebates paid directly (or indirectly through a pharmacy benefit manager) to Part D prescription drug plans and Medicare Advantage prescription drug plans. Effective December 30, 2020, the Rebate Rule established two new safe harbors. The first new safe harbor protects price reductions paid by manufacturers to prescription drug plans (including prescription drug plans offered by Medicare Advantage organizations) and Medicaid managed care organizations, which are fully reflected at the point-of-sale. The second new safe harbor protects fair-market-value service fees paid to pharmacy benefit managers by manufacturers. This new rule could result in a change in incentives for health plans and pharmacy benefit managers in negotiating rebates and discounts with manufacturers for preferred formulary placement. At this time we cannot predict how these changes will impact our business and operations once our product candidates are commercialized.

Within the European Union, the provision of benefits or advantages to physicians to induce or encourage the prescription, recommendation, endorsement, purchase, supply, order or use of medicinal products is prohibited. The provision of benefits or advantages to physicians is also governed by the national anti-bribery laws of EU Member States, such as the UK Bribery Act 2010. Infringement of these laws could result in substantial fines and imprisonment.

We are also subject to the U.S. Foreign Corrupt Practices Act (“FCPA”), which prohibits any U.S. individual or business from paying, offering, authorizing payment or offering anything of value, directly or indirectly, to any foreign official, political party or candidate for the purpose of influencing any act or decision of the foreign entity in order to assist the individual or business in obtaining or retaining business. The FCPA also regulates companies whose securities are listed in the U.S. to
comply with certain accounting provisions requiring the company to maintain books and records that accurately and fairly reflect all transactions of the corporation, including international subsidiaries, to devise and maintain an adequate system of internal accounting controls for international operations. Compliance with the FCPA is expensive and difficult, particularly in countries in which corruption is a recognized problem. In addition, the FCPA presents particular challenges in the pharmaceutical industry, because, in many countries, hospitals are operated by the government, and doctors and other hospital employees are considered foreign officials. Certain payments to hospitals in connection with clinical trials and other work have been deemed to be improper payments to government officials and have led to FCPA enforcement actions. Various laws, in legislations and executive orders also restrict the use and dissemination outside the U.S. or the sharing with certain non-U.S. nationals, of information classified for national security purposes, as well as certain products and technical data relating to those products. As we expand our presence outside the U.S., it will require us to dedicate additional resources to comply with these laws, and these laws may preclude us from developing, manufacturing, or selling certain products and product candidates outside the United States, which could limit our growth potential and increase our development costs. We cannot guarantee that we, our employees, our consultants, or our third-party contractors are or will be in compliance with all federal, state, and foreign regulations regarding bribery and corruption. Moreover, our strategic collaborators and third-party contractors located outside the U.S. may have inadequate compliance programs or may fail to respect the laws and guidance of the territories in which they operate. The failure to comply with laws governing international business practices may result in substantial civil and criminal penalties and suspension or debarment from government contracting. The SEC also may suspend or bar issuers from trading securities on U.S. exchanges for violations of the FCPA’s accounting provisions. Even if we are not determined to have violated these laws, government investigations into these issues typically require the expenditure of significant resources and generate negative publicity, which could also have an adverse effect on our business, financial condition, and results of operations.

The Federal Health Insurance Portability and Accountability Act of 1996 ("HIPAA"), created additional federal criminal statutes that prohibit, among other actions, knowingly and willfully executing, or attempting to execute, a scheme to defraud any healthcare benefit program, including private third-party payors, knowingly and willfully embezzling or stealing from a healthcare benefit program, willfully obstructing a criminal investigation of a healthcare offense, and knowingly and willfully falsifying, concealing or covering up a material fact or making any materially false, fictitious or fraudulent statement in connection with the delivery of or payment for healthcare benefits, items or services. HIPAA, as amended by the Health Information Technology for Economic and Clinical Health Act ("HITECH") and their implementing regulations, imposes requirements regarding the privacy and security of individually identifiable health information, including mandatory contractual terms, for covered entities, or certain healthcare providers, health plans, and healthcare clearinghouses, and their business associates that provide services to the covered entity that involve individually identifiable health information. HIPAA and HITECH also increased the civil and criminal penalties that may be imposed against covered entities and business associates, and created a new general authority to file civil actions for damages or injunctions in federal courts to enforce HIPAA. While pharmaceutical and biotechnology companies are typically not directly regulated by HIPAA, our business may be indirectly impacted by HIPAA in our interactions with providers, payors, and others that have HIPAA compliance obligations. We are also subject to state and foreign laws governing the privacy and security of health or personal information such as the European Union General Data Protection Regulation ("GDPR") and the California Consumer Privacy Act of 2018 ("CCPA").

There has been a recent trend of increased federal and state regulation of payments made to physicians and other healthcare providers. The Physician Payments Sunshine Act imposes annual reporting requirements on certain manufacturers of drugs, devices, biologics, and medical supplies for which payment is available under Medicare, Medicaid, or the Children’s Health Insurance Program, with specific exceptions, for payments made by them to physicians (defined to include doctors, dentists, optometrists, podiatrists and chiropractors) and teaching hospitals, as well as ownership and investment interests held by physicians and their immediate family members. Beginning in 2022, applicable manufacturers will also be required to report information related to payments and other transfers of value provided in the previous year to physician assistants, nurse practitioners, clinical nurse specialists, certified registered nurse anesthetists, and certified midwives.

Within the European Union, payments made to physicians in certain EU Member States must be publicly disclosed. Moreover, agreements with physicians often must be the subject of prior notification and approval by the physician’s employer, their competent professional organization, or the regulatory authorities of the individual EU Member States. These requirements are provided in the national laws, industry codes, or professional codes of conduct, applicable in the European Union Member States. Failure to comply with these requirements could result in reputational risk, public reprimands, administrative penalties, fines, or imprisonment.

Laws and regulations have been enacted by the federal government and various states to regulate the sales and marketing practices of pharmaceutical manufacturers with marketed products. The laws and regulations generally limit financial interactions between manufacturers and health care providers and/or require disclosure to the government and public of such interactions. Many of these laws and regulations contain ambiguous requirements or require administrative guidance for implementation. Given the lack of clarity in laws and their implementation, any future activities (if we obtain approval and/or reimbursement from federal healthcare programs for our product candidates) could be subject to challenge.

Given the significant global impact of the COVID-19 pandemic, it is possible that one or more government entities may take actions, including the U.S. Government under the Defense Production Act of 1950, as amended, which could directly or indirectly have the effect of diminishing some of our rights or opportunities with respect to NVX-CoV2373 and the economic value of a COVID-19 vaccine to us could be limited. In addition, during a global health crisis, such as the COVID-19 pandemic, where the spread of a disease needs to be controlled, closed or heavily regulated national borders will create challenges and potential delays in our development and production activities and may necessitate that we pursue strategies to develop and produce our vaccine candidates within self-contained national or international borders, at potentially much greater expense and with longer timeframes for public distribution.

We face an inherent risk of product liability as a result of the clinical testing of our product candidates and will face an even greater risk if we commercialize any products. For example, we may be sued if any product we develop allegedly causes injury or is found to be otherwise unsuitable during product testing, manufacturing, marketing or sale. Any such product liability claims may include allegations of defects in manufacturing, defects in design, a failure to warn of dangers inherent in the product, negligence, strict liability and a breach of warranties. Claims could also be asserted under state consumer protection acts. If we cannot successfully defend ourselves against product liability claims, we may incur substantial liabilities or be required to limit commercialization of our product candidates. In the United States, the Public Readiness and Emergency Preparedness Act (the “PREP Act”), provides immunity for manufacturers from all claims under state or federal law for “loss or injury arising out of the administration or use of a ‘covered countermeasure.” However, injured persons may still bring a suit for “willful misconduct” against the manufacturer under some circumstances. “Covered countermeasures” include security countermeasures and “qualified pandemic or epidemic products”, including products intended to diagnose or treat pandemic or epidemic disease, such as pandemic vaccines, as well as treatments intended to address conditions caused by such products. For these immunities to apply, the Secretary of DHHS must issue a declaration in cases of public health emergencies or “credible risk” of a future public health emergency. On March 17, 2020, the Secretary of DHHS issued a declaration under the PREP Act and has issued subsequent amendments thereto since then to provide liability immunity for activities related to certain countermeasures against the ongoing COVID-19 pandemic. While we believe our products would be covered under the provisions of the PREP Act, this cannot be assured.

Also, there can be no assurance that the Secretary of the HHS will make other declarations in the future that cover any of our other product candidates or that the U.S. Congress will not act in the future to reduce coverage under the PREP Act or to repeal it altogether. If product liability lawsuits are brought against us, we may incur substantial liabilities and may be required to limit commercialization of our product candidates.

**HUMAN CAPITAL**

**Employees**

As of February 24, 2021, we have 791 full-time employees, of whom 90 hold M.D. or Ph.D. degrees and 189 of whom hold other advanced degrees. Of our total workforce, 653 are engaged primarily in research, development and manufacturing activities and 139 are engaged primarily in executive, business development, finance and accounting, legal and administrative functions. None of our U.S. and Czech employees are represented by labor unions or covered by collective bargaining agreements; 68 of our 69 Swedish employees are covered by typical collective bargaining agreements. To nurture, grow, and treat our employees fairly is imbued in our culture. We were recognized as one of the Top 50 employers to work for in the U.S. by our employees.
We are proud to have been recognized in the 2021 Top Workplaces USA list based on employee surveys. We believe this award reflects our investment in an exceptional culture.

COVID-19 Employee Safety and Benefits

With the emergence of the COVID-19 global pandemic, we took precautions to reduce the risk of virus exposure for all employees. We encouraged all of our employees who were able to work from home to do so, and we reduced the number of people in our offices significantly with the remote work option. Because of our business, it was necessary for essential employees to remain on-site. For those employees and any other employees who entered our offices, we adopted new safety protocols including, social distancing, face mask requirements, temperature screening and health questionnaires.

In March 2020, we recognized the severity of COVID-19 and offered a special enrollment period to our employees to provide them with an additional opportunity to participate in our health insurance plan. We have also established an emergency relief fund for our employees whose immediate families have been materially and negatively impacted by the COVID-19 emergency.

Compensation and Benefits; Health and Wellness

Our total rewards are designed to attract, motivate, and retain top talent in the industry. We strive to provide pay, comprehensive benefits and services that help meet the varying needs of our employees. Our generous total rewards package includes competitive market pay, fully covered healthcare benefits for employees, a health savings account, a 401(k) retirement savings plan, paid vacation, family leave, flexible work schedules, an employee assistance program, and on-site and online concierge services. In addition, we offer every employee, the benefit of equity ownership in the company through stock option and restricted stock unit grants and our employee stock purchase plan.

Recruitment, Development and Training

The attraction, development, and retention of employees is a critical success factor for Novavax. We utilize a variety of recruitment vehicles to source top talent, including strategic partnerships with search firms, leveraging social media channels, and a robust employee referral program. Since March 2020, we have hired over 400 full-time and part-time employees to address the global COVID-19 pandemic and bring our NVX-CoV2373 vaccine candidate to market following global regulatory approvals.

To support the growth and advancement of our employees, we offer tuition and continuing education reimbursement, and an array of training and professional development opportunities, including on-the-spot coaching with executive coaches and access to the LinkedIn Learning library of over 16,000 on-demand video tutorials that address skill, knowledge and behaviors related to business, leadership, technology, and creativity. In the last 12 months, videos were viewed and completed over 7,000 times by our employees. We provide an Executive Development Program for employees identified as having high potential and for potential successors to leadership positions, executive coaching engagements, and leadership development programs to strengthen our leadership bench and accelerate and prepare our top talent for future growth.

Internal Communications

We employ a variety of tools to facilitate open and direct communication, including global forums with executives, employee surveys, and engagement through forums and committees. Our executive leadership team continues to recognize the importance of increased employee engagement.

Diversity and Inclusion

Our culture of diversity, equity and inclusion enables us to create, develop and fully leverage the strengths of our workforce to meet our growth objectives. We recently completed an evidence-based analysis of current state on diversity, equity and inclusion to understand how best to create a culture of inclusion and diverse workforce and how to build a sustainable strategy to drive diversity and inclusion at Novavax. We are very fortunate to have a diverse workforce and we believe our DEI strategy will enable us to continuously improve and excel.

Corporate Social Responsibility

We are endeavoring to develop relationships, give back to our communities and engage in corporate social responsibility and sustainability initiatives. As we grow our employee base, we are focused on extending our efforts in these areas.

Availability of Information

Our website address is www.novavax.com. We make available, free of charge and through our website, our Annual Reports on Form 10-K, Quarterly Reports on Form 10-Q, Current Reports on Form 8-K and our other filings with the Securities and Exchange Commission (“SEC”), and any amendments to any such reports filed or furnished pursuant to Section 13(a) or 15(d) of the Securities Exchange Act of 1934, as amended, as soon as reasonably practicable after filing with or furnished to the SEC. The SEC maintains an Internet site that contains reports, proxy and information statements, and other information regarding issuers that file electronically with the SEC at www.sec.gov.

We use our website (www.novavax.com) as a means of disclosing material non-public information and for complying with our disclosure obligations under Regulation Fair Disclosure promulgated by the SEC. These disclosures are included on our website (www.novavax.com) in the “Investors” or “News” sections. Accordingly, investors should monitor these portions of our website (www.novavax.com), in addition to following our press releases, SEC filings and public conference calls and webcasts.

Also available on our website is information relating to corporate governance at Novavax and our Board of Directors, including our Code of Business Conduct and Ethics. We intend to disclose on our website any future amendments to and waivers from this code that apply to our Chief Executive Officer, Principal Financial Officer, Principal Accounting Officer and Controller, and persons performing similar functions, as promptly as practicable, as may be required under applicable SEC and Nasdaq rules.

We webcast our earnings calls and certain events we participate in or host with members of the investment community on the investor relations section of our website. Additionally, we provide notifications of news or announcements regarding press and earnings releases as part of the investor relations section of our website. The contents of our website are not part of this Annual Report on Form 10-K, or any other report we file with, or furnish to, the SEC.
Item 1A. RISK FACTORS

You should carefully consider the following risk factors in evaluating our business. A number of risk factors could cause our actual results to differ materially from those that are indicated by forward-looking statements. Some risks relate principally to our business and the industry in which we operate. Others relate principally to the securities market and ownership of our common stock. The risks and uncertainties described below are not the only ones we face. Additional risks and uncertainties of which we are unaware, or that we currently deem immaterial, also may become important factors that affect us. If any of the following risks occur, our business, financial condition or results of operations could be materially and adversely affected. You also should consider the other information included in this Annual Report on Form 10-K.

Risks Related to Our Financial Condition and Capital Requirements

We have a history of losses and our future profitability is uncertain.

Our expenses have exceeded our revenue since our formation in 1987, and our accumulated deficit at December 31, 2020 was $1.9 billion. Our revenue for the last three fiscal years was $475.6 million in 2020, $187.8 million in 2019 and $34.3 million in 2018. We may not be successful in entering into collaborations, strategic alliances and marketing, distribution or licensing arrangements with other companies or government agencies that result in significant revenue to offset our expenses. Our net losses for the last three fiscal years were $418.3 million in 2020, $132.7 million in 2019 and $184.7 million in 2018.

Historically, our losses have resulted predominantly from research and development expenses for our vaccine candidates, manufacturing-related expenses, costs related to protection of our intellectual property and other general and administrative operating expenses, a significant portion of which have been noncash. Our expenses have exceeded our revenue since inception, and we believe our expenses will fluctuate over time, and may substantially increase in some years, as a result of continuing research and development efforts to support our vaccine development efforts, and, if our product candidates are approved, commercialization efforts.

As of the end of fiscal year 2020, our investment in the development and manufacture of NVX-CoV2373 has been significant and we expect such levels of investment to continue through 2021 and beyond, although the precise magnitude of our total investment will be subject to clinical trial data results, the duration of the COVID-19 pandemic and other factors, including our competitive landscape and regulatory outcomes. If we are unable to timely commercialize a vaccine against COVID-19, we may never recoup this investment. We expect to continue to incur significant operating expenses and anticipate significant losses over time as we seek to:

- conduct clinical trials and seek regulatory approval for NVX-CoV2373 and other potential vaccine candidates;
- conduct preclinical studies for other potential vaccine candidates;
- expand our global manufacturing and distribution capacity; and
- maintain, expand and protect our intellectual property portfolio.

As a result, we expect our cumulative operating losses to increase until such time, if ever, that product sales, licensing fees, royalties, milestones, contract research and other sources generate sufficient revenue to fund our operations. We may never achieve profitability and may not sustain profitability, if achieved.

We will continue to require significant funding to maintain our current level of operations and fund the further development of our vaccine candidates.

We do not currently generate sufficient revenue from product sales, licensing fees, royalties, milestones, contract research or other sources to fully fund our operations. We will therefore use our cash resources, and expect to require additional funds, to maintain our operations, continue our research and development programs, commence future preclinical studies and clinical trials, seek regulatory approvals and manufacture and market our products.

To date, we have financed our operations primarily through the sale of equity and debt securities, government funding and grant agreements, and we cannot be certain that additional such funding will be available to us on favorable terms, or at all. Although we have recently entered into supply agreements for NVX-CoV2373 that include prepayments from the purchasers, until we can generate sufficient product revenue in amounts sufficient to fully fund our operations, which we may never do, we expect to finance our future cash needs through a combination of additional public or private equity or debt financings, as well as potential collaborations, strategic alliances and marketing, distribution or licensing arrangements and non-dilutive funding from governmental and non-governmental funding entities, as well as other sources. While we may continue to apply for contracts or grants from academic institutions, non-profit organizations and governmental entities, we may not be successful. Adequate additional funding may not be available to us on acceptable terms, if at all. Furthermore, any negative clinical trial data or setbacks, or perceived setbacks, with respect to our vaccine candidates, particularly NVX-CoV2373, could impair our ability to raise additional financing on favorable terms, or at all. If we cannot raise the additional funds required for our anticipated operations, we may be required to delay significantly, reduce the scope of or eliminate one or more of our research or development programs, downsize our organization, or seek alternative measures to avoid insolvency, including arrangements with collaborative partners or others that may require us to relinquish rights to certain of our technologies or vaccine candidates. If we raise additional funds through future offerings of shares of our common stock or other securities, such offerings would cause dilution of current stockholders’ percentage ownership in the Company, which could be substantial. Future offerings also could have a material and adverse effect on the price of our common stock.

Economic uncertainty may adversely affect our access to capital, cost of capital and ability to execute our business plan as scheduled.

Generally, worldwide economic conditions remain uncertain, particularly due to the COVID-19 pandemic. Access to capital markets is critical to our ability to operate. Traditionally, biotechnology companies have funded their research and development expenditures through raising capital in the equity markets. Declines and uncertainties in these markets in the past have severely limited raising new capital and have affected companies’ ability to continue to expand or fund existing research and development efforts. We require significant capital for research and development for our vaccine candidates and clinical trials. The general economic and capital market conditions, both in the U.S. and worldwide, have been volatile in the past and at times have adversely affected our access to capital and increased the cost of capital. There is no certainty that the capital and credit markets will be available to raise additional capital on favorable terms, particularly given the ultimate impact of the COVID-19 pandemic on the economies of countries worldwide is unknown. If economic conditions become worse, as a result of the COVID-19 pandemic or otherwise, our future cost of equity or debt capital and access to the capital markets could be adversely affected. In addition, if we are unable to access the capital markets on favorable terms, our ability to execute our business plan as scheduled would be compromised. Moreover, we rely and intend to rely on third-parties, including clinical research organizations, contract manufacturing organizations and other important vendors and consultants. Global economic conditions may result in a disruption or delay in the performance of our third-party contractors and suppliers. If such third-parties are unable to adequately satisfy their contractual commitments to us in a timely manner, our business could be adversely affected.
Our existing funding and supply agreements do not assure success of our vaccine candidates or that we will be able to fully fund our vaccine candidates.

The OWS Agreement, the DoD Agreement and the CEPI Funding Agreement each reimburse a portion of the expenses associated with the development and commercialization of NVX-CoV2373. To the extent funding commitments in such agreements are conditioned on our meeting certain milestones or conditions, including regulatory approval in applicable jurisdictions, we may not ultimately receive the full amount of committed funds and could be exposed to urgent needs for additional funding to support our NVX-CoV2373 development, manufacturing and distribution activities. There can be no assurance that we will be able to timely obtain additional government or private funding, if at all. Additionally, we have entered into, and plan to continue entering into, supply agreements for NVX-CoV2373 that include prepayments from the purchasers. In the event we are unable to successfully develop and commercialize NVX-CoV2373 or fail to meet certain product volume or delivery timing obligations under our supply agreements, we may be required to refund significant portions of the prepayments, which could have a material and adverse effect on our financial condition. Our inability to succeed with key clinical or development activities could jeopardize our ability to obtain licensure from the FDA or other regulatory authorities to sell NVX-CoV2373. We can make no assurance that the OWS Agreement, the DoD Agreement and the CEPI Funding Agreement will be sufficient to fund our vaccine candidate development or our supply agreements will be sufficient to fund our commercial launch.

Similarly, the Grant Agreement with BMGF reimburses a portion of specified expenses associated with the development of ResVax. The Grant Agreement with BMGF does not assure success of ResVax or that the vaccine candidate will be licensed by the FDA. Additional development activities likely will be needed and BMGF may not reimburse us for any portion of these activities.

Risks Related to Product Development and Commercialization

Because our vaccine product development efforts depend on new and rapidly evolving technologies, we cannot be certain that our efforts will be successful.

Our vaccine development efforts depend on, rapidly evolving technologies and on the marketability and profitability of our products. Our development efforts and, if those are successful, commercialization of our vaccines could fail for a variety of reasons, and include the possibility that:

- our recombinant nanoparticle vaccine technologies, any or all of the products based on such technologies or our proprietary manufacturing process will be ineffective or unsafe, or otherwise fail to receive necessary regulatory approvals or achieve commercial viability;
- we or our third-party manufacturer facilities will be unable or unwilling to scale-up manufacturing capabilities for our products in a cost-effective manner;
- the products, if safe and effective, will be difficult to manufacture on a large-scale or uneconomical to market;
- our in-house or third-party manufacturing facilities will fail to continue to pass regulatory inspections;
- proprietary rights of third-parties will prevent us or our collaborators from exploiting technologies, and manufacturing or marketing products; and
- third-party competitors will gain greater market share due to superior products or marketing capabilities.

Although we have made rapid progress, the regulatory and commercial success of our COVID-19 vaccine candidate, NVX-CoV2373, remains uncertain. We may be unable to obtain regulatory approval or produce a successful vaccine in a timely manner, if at all.

In response to the outbreak of COVID-19, we are pursuing the development and manufacture of our vaccine candidate, NVX-CoV2373, which is currently in Phase 3 of clinical testing. Even though we have reported positive data from Phase 1, 2 and 3 clinical trials, our development of NVX-CoV2373 is ongoing and such results may not be predictive of future clinical trial results or whether future clinical trial results will be sufficient to support regulatory authorization or approval, accelerated or otherwise. We may be unable to produce a vaccine that successfully prevents COVID-19 in a timely manner, if at all.

Additionally, even if NVX-CoV2373 receives regulatory approval, our ability to successfully commercialize it depends on our ability to effectively scale up manufacturing capabilities at our own locations and those of our manufacturing partners and contractors. In May 2020, we acquired Novavax CZ (formerly Praha Vaccines a.s.) including its vaccine manufacturing facility in Bohumil, Czech Republic and approximately 150 of its employees. We are also actively entering into agreements with third-parties to manufacture the antigen component of NVAX-CoV2373 and our proprietary Matrix-M adjuvant, as well as to distribute NVX-CoV2373. Because of contractual restraints and the limited number of third-party manufacturers with the expertise, required regulatory approvals and facilities to manufacture NVX-CoV2373 on a fundamentally commercial scale, replacement of a manufacturer may be expensive and time-consuming and may cause interruptions in production. Manufacturing of NVX-CoV2373 involves a complicated process that will require significant investments of time and financial resources to implement. We cannot guarantee that we will be able to timely and effectively produce NVX-CoV2373 in adequate quantities to meet global demand.

The Company has not previously had a commercial launch of any vaccine product and doing so in a pandemic environment with an urgent, critical global need creates additional challenges. In addition to scaling up our manufacturing capabilities, we will need to develop global distribution channels and form partnerships with third-parties worldwide, as well as hire, train and integrate additional management, administrative and sales and marketing personnel. Rapid and significant growth may strain our administrative and operational infrastructure, imposing significant additional responsibilities on our organization, and our efforts to establish these capabilities may not meet initial expectations as to timing, scale-up, yield, cost or quality. If we are unable to successfully manage our growth and the increased complexity of our operations, our business, financial position, results of operations and prospects may be materially and adversely affected.

We are a biotechnology company and face significant risk in developing, manufacturing and commercializing our products.

We focus our research and development activities on vaccines, an area in which we believe we have particular strengths and a technology that appears promising. The outcome of any research and development program is highly uncertain. Only a small fraction of biopharmaceutical development programs ultimately results in commercial products or even product candidates and a number of events could delay our development efforts and negatively impact our ability to obtain regulatory approval for, and to manufacture, market and sell, a vaccine. Vaccine candidates that initially appear promising often fail to yield successful products. In many cases, preclinical studies or clinical trials will show that a product candidate is not efficacious or that it raises safety concerns or has other side effects that outweigh its intended benefit. Success in preclinical or early clinical trials may not translate into success in large-scale clinical trials. Further, success in clinical trials often leads to increased investment, accelerating cumulative losses. Even if clinical trial results appear positive, regulatory approval may not be obtained if the FDA does not agree with our interpretation of the results, and we may face challenges when scaling-up the production process to commercial levels. Even after a product is approved and launched, general usage or post-marketing clinical trials may identify safety or other previously unknown problems with the product, which may result in regulatory approvals being suspended, limited to narrow the scope of the approval, or
revoked, which may otherwise prevent successful commercialization. Intense competition in the vaccine industry could also limit the successful commercialization of any products for which we receive commercial approval.

Because we depend on third-parties to conduct some of our laboratory testing and clinical trials, and a significant amount of our vaccine manufacturing activities and distribution, we may encounter delays in or lose some control over our efforts to develop and supply products.

We are highly dependent on third-party organizations to conduct some of our laboratory testing and clinical trials and a significant amount of our vaccine manufacturing activities and distribution. If we are unable to obtain any necessary services on acceptable terms, we may not complete our product development efforts in a timely manner. We may lose some control over these activities or become too dependent upon those parties. These third-parties may not complete testing, manufacturing or distribution activities on schedule, or in satisfaction of regulatory or commercial requirements. Certain of our facilities are also contracted for defined time frames and through association with OWS and CEPI and may not be available for sufficient periods of time to adequately supply our products.

We are responsible for confirming that each of our clinical trials is conducted in accordance with its general investigational plan and protocol. Moreover, the FDA and foreign regulatory agencies require us to comply with regulations and standards, commonly referred to as good clinical practices, for conducting, recording and reporting the results of clinical trials to assure that data and reported results are credible and accurate and that the rights, safety and welfare of clinical trial participants are adequately protected. The FDA and foreign regulatory agencies also require us to comply with good manufacturing practices. Our reliance on third-parties does not relieve us of these responsibilities and requirements. These third-parties may not successfully carry out their contractual duties or regulatory obligations. Furthermore, if our third-party manufacturer is producing materials or products for themselves or other companies, our third-party manufacturer may be exposed to regulatory risks for the production of such materials and products. As a result, failure to meet the regulatory requirements for the production of those materials and products may generally affect the regulatory status of the third-party manufacturer’s facility, which could impact its ability to produce our materials and products. Any of our third-party service providers may need to be replaced, the quality or accuracy of the data they obtain may be compromised, or the product they manufacture may be contaminated due to the failure to adhere to our clinical and manufacturing protocols, regulatory requirements or for other reasons. In any such event, our preclinical development activities or clinical trials may be extended, delayed, suspended or terminated, and we may not be able to obtain regulatory approval of, or successfully commercially manufacture, our vaccine candidates.

The results from the Prepare trial, including that ResVax failed to meet the primary endpoint of the trial, will likely create challenges, some of which may be significant, around further development of that vaccine.

While the Prepare results suggest that ResVax is safe and is likely efficacious in more serious manifestations of RSV disease, the trial failed to achieve its primary clinical endpoint. Not achieving the primary clinical endpoint has been viewed negatively by our investors. Although the failure to achieve the primary endpoint in the trial is not evidence that the vaccine is ineffective, it means that regulatory agencies like the FDA and EMA are likely to require additional clinical trial data prior to licensure. This development may be viewed negatively by our potential collaborators and partners, which may make the ongoing development of ResVax, and any other RSV F Vaccine candidates, more challenging.

We may have product liability exposure.

The administration of drugs or vaccines to humans, whether in clinical trials or after marketing approval, can result in product liability claims. We maintain product liability insurance coverage for our current clinical programs, including our NVX-CoV2373 trials. If and when we obtain marketing approval for any vaccine candidate, we intend to expand our insurance coverage to include the sale of commercial products; however, we may not be able to obtain or maintain insurance coverage on commercially reasonable terms, at a reasonable cost or in sufficient amounts to protect us against losses due to liability. Furthermore, such insurance coverage and our resources may not be sufficient to satisfy all liabilities that result from product liability claims. A successful claim may prevent us from obtaining adequate product liability insurance in the future on commercially desirable terms, if at all. Even if a claim is not successful, defending such a claim would be time-consuming and expensive, may damage our reputation in the marketplace and would likely divert management’s attention.

In addition, because we are developing NVX-CoV2373 in response to the outbreak of COVID-19, a global pandemic, we may have a widely used vaccine in the U.S. and other countries as an investigational vaccine or a product authorized for temporary or emergency use prior to our receipt of marketing approval. Unexpected safety issues in these circumstances could lead to product liability claims and our existing insurance may not be adequate for such claims.

Regardless of merit or eventual outcome, liability claims may result in:
- decreased demand for our products;
- withdrawal of regulatory approvals;
- voluntary or mandatory recalls of our products;
- necessity for additional nonclinical or clinical studies, changes in labeling, or changes to manufacturing processes, specifications and/or facilities;
- impairment of our business reputation and negative media attention;
- withdrawal of clinical trial participants;
- costs of related litigation;
- substantial monetary awards to participants or other claimants;
- loss of revenue; and
- inability to commercialize our vaccine candidates.

In the United States, the PREP Act, provides immunity for manufacturers from all claims under state or federal law for “loss” arising out of the administration or use of a “covered countermeasure.” However, injured persons may still bring a suit for “willful misconduct” against the manufacturer under some circumstances. “Covered countermeasures” include security countermeasures and “qualified pandemic or epidemic products”, including products intended to diagnose or treat pandemic or epidemic disease, such as pandemic vaccines, as well as treatments intended to address conditions caused by such products. For these immunities to apply, the Secretary of DHHS must issue a declaration in cases of public health emergency or “credible risk” of a future public health emergency. On March 17, 2020, the Secretary of DHHS issued a declaration under the PREP Act and has issued subsequent amendments thereto since then to provide liability immunity for activities related to certain countermeasures against the ongoing COVID-19 pandemic. While we believe our products would be covered under the provisions of the PREP Act, this cannot be assured. Also, there can be no assurance that the Secretary of the HHS will make other declarations in the future that cover any of our other product candidates or that the U.S. Congress will not act in the future to reduce coverage under the PREP Act or to repeal it altogether. If product liability lawsuits are brought against us, we may incur substantial liabilities and may be required to limit commercialization of our product candidates.

If we are unable to manufacture our vaccines in sufficient quantities, at sufficient yields or are unable to obtain regulatory approvals for a manufacturing facility for our vaccines, we may experience delays or an adverse impact on product development, clinical trials, regulatory approval and commercial distribution.

Completion of our clinical trials and commercialization of our vaccine candidates require access to, or development of, facilities to manufacture our vaccine candidates at sufficient yields and at commercial-scale. We have limited experience manufacturing any of our vaccine candidates in the volumes that will be necessary to support large scale clinical trials or commercial sales. While we have recently increased our projected global manufacturing capacity for NVX-CoV2373, our efforts to establish manufacturing capabilities may not meet expectations as to scheduling, scale-up, reproducibility, yield, purity, cost, potency or quality. The antigen component of NVX-CoV2373 is currently being manufactured by our vaccine candidates.
manufactured at Novavax CZ, as well as numerous partnered manufacturing sites, including FUJIFILM in the United States, SIIPL in India and Takeda in Japan, among others.

Manufacturing our vaccine candidates involves a complicated process with which we have limited experience. We are highly dependent on third-party organizations to conduct a significant amount of our vaccine manufacturing activities. If we and our third-party manufacturing organizations are unable to manufacture our vaccine candidates in clinical quantities or, when necessary, in commercial quantities and at sufficient yields, then we will need to identify and reach supply arrangements with additional third-parties. Third-party manufacturers must also receive FDA or equivalent foreign regulatory body approval before they can produce clinical material or commercial products. Our vaccines may be in competition with other products for access to these third-party facilities and may find that the facilities or third-parties give other products higher priority. We may not be able to enter into any necessary additional third-party manufacturing arrangements on acceptable terms, or on a timely basis. In addition, we have to enter into technical transfer agreements and share our know-how with the third-party manufacturers, which can be time-consuming and may result in delays.

Because of contractual restraints and the limited number of third-party manufacturers with the expertise, required regulatory approvals and facilities to manufacture our bulk vaccines on a commercial-scale, replacement of a manufacturer may be expensive and time-consuming and may cause interruptions in the production of our vaccine. We and our third-party manufacturers may also encounter difficulties in production. These problems may include:

- difficulties with production costs, scale up and yields;
- availability of raw materials and supplies;
- quality control and assurance;
- shortages of qualified personnel;
- compliance with strictly enforced and evolving federal, state and foreign regulations that vary in each country where products might be sold including nationalization or other territory restrictions placed on our owned and third-party manufacturing sites; and
- lack of capital funding.

As a result, any delay or interruption could have a material adverse effect on our business, financial condition, results of operations and cash flows.

We must identify vaccines for development with our technologies and establish successful third-party relationships.

The near and long-term viability of our vaccine candidates will depend in part on our ability to successfully establish new strategic collaborations with pharmaceutical and biotechnology companies, non-profit organizations and government agencies. Establishing strategic collaborations and obtaining government funding is difficult and time-consuming. Potential collaborators may reject collaborations based upon their assessment of public need, the public interest, our products’ ability to address these areas, or other reasons beyond our expectations or control. Past success in establishing strategic collaborations with pharmaceutical and biotechnology companies, non-profit organizations and government agencies in the past is no guarantee of future success. If we fail to establish a sufficient number of collaborations or government relationships on acceptable terms, we may not be able to commercialize our vaccine candidates or generate sufficient revenue to fund further research and development efforts.

There is no guarantee that the collaborations we have established or will establish will result in the successful development or commercialization of any vaccine candidates for several reasons, including the fact that:

- we may not have the ability to control the activities of our partners and cannot provide assurance that they will fulfill their obligations to us, including with respect to the license, development and commercialization of vaccine candidates, in a timely manner or at all;
- such partners may not devote sufficient resources to our vaccine candidates or properly maintain or defend our intellectual property rights;
- our partners could independently develop, or develop with third-parties, products that compete directly or indirectly with our vaccine candidates if such partners believe that competitive products are more likely to be successfully developed or can be commercialized under terms that are more economically attractive than ours;
- any failure on the part of our partners to perform or satisfy their obligations to us could lead to delays in the development or commercialization of our vaccine candidates and affect our ability to realize product revenue; and
- disagreements, including disputes over the ownership of technology developed with such collaborators, could result in litigation, which would be time consuming and expensive, and may delay or terminate research and development efforts, regulatory approvals and commercialization activities.

If we or our collaborators fail to maintain our existing agreements or in the event, we fail to establish agreements as necessary, we could be required to undertake research, development, manufacturing and commercialization activities solely at our own expense. These activities would significantly increase our capital requirements and given our lack of sales, marketing and distribution capabilities, significantly delay the commercialization of our vaccine candidates.

Even if licensed to market, our vaccine products may not be initially or ever profitable.

Whether Novavax makes a profit from the sale of its vaccine products is dependent on a number of variables, including the costs we incur manufacturing, testing and releasing, packaging and shipping such vaccine product. Additionally, the CEPI Funding Agreement necessitates that we allocate a certain number of doses of NVX-CoV2373 to certain middle- and lower-income countries and the Grant Agreement with BMGF necessitates that we commit to a specific amount of sales in certain specified middle- and lower-income countries, which may impact our ability to make profits. We cannot predict when, if at all, our approved vaccine products will be profitable to the Company.

Even if we successfully commercialize any of our vaccine candidates, either alone or in collaboration, we face uncertainty with respect to pricing, third-party reimbursement and healthcare reform, all of which could adversely affect any commercial success of our vaccine candidates.

Our ability to collect revenue from the commercial sale of our vaccines may depend on our ability, and that of any current or potential future collaboration partners or customers, to obtain adequate levels of approval, coverage and reimbursement for such products from third-party payers such as:

- government health administration authorities such as the Advisory Committee for Immunization Practices of the Centers for Disease Control and Prevention;
- private health insurers;
- managed care organizations;
- pharmacy benefit management companies; and
- other healthcare related organizations.

Third-party payers are increasingly challenging the prices charged for medical products and may deny coverage or offer inadequate levels of reimbursement if they determine that a prescribed product has not received appropriate clearances from the FDA, or foreign equivalent, or other government regulators, is not used in accordance with cost-effective treatment methods as determined by the third-party payer; or is experimental, unnecessary or inappropriate. Prices could also be driven down by managed care organizations that control or significantly influence utilization of healthcare products.
In both the U.S. and some foreign jurisdictions, there have been a number of legislative and regulatory proposals and initiatives to change the health care system in ways that could affect our ability to sell vaccines and could adversely affect the prices that we receive for our vaccine candidates, if approved. Some of the proposals and implemented reforms could result in reduced pharmaceutical pricing or reimbursement rates for medical products, and while we have no current vaccines available for commercial sale, the impact of such reform could nevertheless adversely affect our business strategy, operations and financial results. For example, the Healthcare Reform Act contained several cost containment measures that could adversely affect our future revenue, including, for example, increased drug rebates under Medicaid for brand name prescription drugs, extension of Medicaid rebates to Medicaid managed care organizations, and expansion of so-called 340B discounted pricing on pharmaceuticals sold to certain healthcare providers. Additional provisions of the healthcare reform laws that may negatively affect our future revenue and prospects for profitability include the assessment of an annual fee based on our proportionate share of sales of brand name prescription drugs to certain government programs, including Medicare and Medicaid. The Healthcare Reform Act also established a Medicare Part D coverage gap discount program, in which manufacturers must agree to offer 70% point-of-sale discounts off negotiated prices of applicable branded on drugs (including vaccines) to eligible beneficiaries during their coverage gap period (the so-called "donut hole"), as condition for the manufacturer’s outpatient drugs to be covered under Medicare Part D. Other aspects of healthcare reform, such as expanded government enforcement authority and heightened standards that could increase compliance-related costs, could also affect our business. In addition, we face uncertainties because there are ongoing federal legislative and administrative efforts to repeal, substantially modify or invalidate some or all of the provisions of the Affordable Care Act, and the Trump Administration has taken actions to reduce the ACA’s impact. In December 2019, the U.S. Court of Appeals for the D.C. Circuit ruled that the individual mandate was unconstitutional and remanded the case back to the District Court to determine whether the remaining provisions of the Affordable Care Act are invalid as well. On March 2, 2020, the U.S. Supreme Court granted the petitions for writs of certiorari to review the case, and oral arguments were heard on November 10, 2020. It is unclear when a decision will be made or how the U.S. Supreme Court will rule.

Other legislative changes have been proposed and adopted since the Healthcare Reform Act was enacted. These changes include aggregate reductions to Medicare payments to providers of 2% per fiscal year pursuant to the Budget Control Act of 2011 and subsequent laws, which began in 2013 and, due to subsequent legislative amendments, will stay in effect through 2030 unless additional Congressional action is taken. In January 2013, the American Taxpayer Relief Act of 2012 was signed into law, which, among other things, further reduced Medicare payments to several types of providers, including hospitals, imaging centers and cancer treatment centers, and increased the statute of limitations period for the government to recover overpayments to providers from three to five years. New laws may result in additional reductions in Medicare and other healthcare funding, which may materially adversely affect customer demand and affordability for our products and, accordingly, the results of our financial operations. Additionally, the pharmaceutical industry has also been the subject of significant publicity in recent years regarding the pricing of pharmaceutical products, including publicity and pressure resulting from prices charged by pharmaceutical companies for new products as well as price increases by pharmaceutical companies on older products that some people have deemed excessive. As a result, pharmaceutical product prices have been the focus of increased scrutiny by the U.S. government, including certain state attorneys general, members of congress, presidential candidates and the United States Department of Justice. If reforms in the health care industry make reimbursement for our potential products less likely, the market for our potential products will be reduced, and we could lose potential reimbursement. The existence or threat of cost containment measures could cause our competitors to be less willing or able to pursue research and development programs related to our vaccine candidates. Further, it is also possible that additional governmental action is taken in response to the COVID-19 pandemic. We cannot predict the ultimate content, timing or effect of any healthcare reform legislation or the impact of potential legislation on us.

We have limited marketing capabilities, and if we are unable to enter into collaborations with marketing partners or develop our own sales and marketing capability, we may not be successful in commercializing any approved products.

Although we have initiated preliminary activities in anticipation of commercialization of our vaccine candidates, we currently have limited dedicated sales, marketing or distribution capabilities. As a result, we depend on collaborations with third-parties that have established distribution systems and sales forces, including our collaboration with SIPL, among others. To the extent that we enter into co-promotion or other licensing arrangements, our revenue will depend upon the efforts of third-parties, over which we may have little or no control. If we are unable to reach and maintain agreements with one or more pharmaceutical companies or collaborators, we may be required to market our products directly. Developing a marketing and sales force is expensive and time-consuming and could delay a product launch. We may not be able to attract and retain qualified sales personnel or otherwise develop this capability.

Our vaccine candidates may never achieve market acceptance even if we obtain regulatory approvals.

Even if we receive regulatory approvals for the commercial sale of our vaccine candidates, the commercial success of these vaccine candidates will depend on, among other things, their acceptance by physicians, patients and third-party payers, such as health insurance companies and other members of the medical community, as a vaccine and cost-effective alternative to competing products. If our vaccine candidates fail to gain market acceptance, we may be unable to earn sufficient revenue to continue our business. Market acceptance of, and demand for, any product that we may develop and commercialize will depend on many factors, including:

- our ability to provide acceptable evidence of safety and efficacy;
- the prevalence and severity of adverse side effects;
- whether our vaccines are differentiated from other vaccines;
- availability, relative cost and relative efficacy of alternative and competing treatments;
- the effectiveness of our marketing and distribution strategy;
- publicity concerning our products or competing products and treatments; and
- our ability to obtain sufficient third-party insurance coverage or reimbursement.

If our vaccine candidates do not become widely accepted by physicians, patients, third-party payers and other members of the medical community, our business, financial condition and results of operations could be materially and adversely affected.

We may not be able to secure sufficient supplies of a key component of our adjuvant technology.

Because an important component of our adjuvant technology is extracted from a species of soap-bark tree (Quillaja saponaria) grown in Chile, we need long term access to quillaja extract with a consistent and sufficiently high quality. We need a secure supply of raw material, as well as back-up suppliers, or our adjuvant products may be delayed, and we may not be able to meet our obligations under our various collaboration and supply agreements.

Current or future regional relationships may hinder our ability to engage in larger transactions.

We have entered into regional collaborations to develop, manufacture and distribute our vaccine candidates in certain parts of the world, and we anticipate entering into additional regional collaborations. Our relationships with SIPL, Cadila and BMGF are examples of these regional relationships. These relationships often involve the licensing of our technology to our partner or entering into a distribution agreement, frequently on an exclusive basis. Generally, exclusive agreements are restricted to certain territories. Because we have entered into exclusive license and distribution agreements,
larger companies may not be interested, or able, to enter into collaborations with us on a worldwide scale. Also, these regional relationships may make us an unattractive target for an acquisition.

Our product candidates are sensitive to shipping and storage conditions, which could subject our vaccine candidates to risk of loss or damage.

Our vaccine candidates are sensitive to storage and handling conditions. Loss in vaccine candidates could occur if the product or product intermediates are not stored or handled properly. It is possible that our vaccine candidates could be lost due to expiration prior to use. If we do not effectively maintain our supply logistics, then we may experience an unusual number of returned or out of date products. Failure to effectively maintain our supply logistics, by us or third-parties, could lead to additional manufacturing costs and delays in our ability to supply required quantities for clinical trials or otherwise.

Our vaccine candidates could become subject to a product recall which could harm our reputation, business, and financial results.

The FDA and similar foreign governmental authorities have the authority to require the recall of certain vaccine candidates. Manufacturers may, under their own initiative, recall a product if any material deficiency in a product is found. A government-mandated or voluntary recall by us or our strategic collaborators could occur as a result of manufacturing errors, design or labeling defects or other deficiencies and issues. Recalls of any of our vaccine candidates would divert managerial and financial resources and have an adverse effect on our financial condition and results of operations. A recall announcement could harm our reputation with customers and negatively affect our sales, if any.

Risks Related to Our Industry and Competition

Many of our competitors have significantly greater resources and experience, which may negatively impact our commercial opportunities and those of our current and future licensees.

The biotechnology and pharmaceutical industries are subject to intense competition and rapid and significant technological change. We have many potential competitors, including major pharmaceutical companies, specialized biotechnology firms, academic institutions, government agencies and private and public research institutions. Many of our competitors have significantly greater financial and technical resources, experience and expertise in:

- research and development;
- preclinical testing;
- designing and implementing clinical trials;
- regulatory processes and approvals;
- production and manufacturing; and
- sales and marketing of approved products.

Principal competitive factors in our industry include:

- the quality and breadth of an organization’s technology;
- management of the organization and the execution of the organization’s strategy;
- the skill and experience of an organization’s employees and its ability to recruit and retain skilled and experienced employees;
- an organization’s intellectual property portfolio;
- the range of capabilities, from target identification and validation to drug discovery and development to manufacturing and marketing; and
- the availability of substantial capital resources to fund discovery, development and commercialization activities.

Large and established companies, such as Merck & Co., Inc., GlaxoSmithKline plc, CSL Ltd, Sanofi Pasteur, SA, Pfizer Inc. and AstraZeneca, among others, compete in the vaccine market. In particular, these companies have greater experience and expertise in securing government contracts and grants to support their research and development efforts, conducting testing and clinical trials, obtaining regulatory approvals to market products, manufacturing such products on a broad scale and marketing approved products.

Regardless of the size, smaller or early-stage companies and research institutions also may prove to be significant competitors, particularly through collaborative arrangements with large and established pharmaceutical companies. As these companies develop their technologies, they may develop proprietary positions, which may prevent or limit our product development and commercialization efforts. We will also face competition from these parties in recruiting and retaining qualified scientific and management personnel, establishing clinical trial sites and participant registration for clinical trials and in acquiring and in-licensing technologies and products complementary to our programs or potentially advantageous to our business. If any of our competitors succeed in obtaining approval from the FDA or other regulatory authorities for their products sooner than we do or for products that are more effective or less costly than ours, our commercial opportunity could be significantly reduced.

In order to effectively compete, we will have to make substantial investments in development, testing, manufacturing and sales and marketing or partner with one or more established companies. We may not be successful in gaining significant market share for any vaccine, our technologies and vaccines also may be rendered obsolete or non-competitive as a result of products introduced by our competitors to the marketplace more rapidly and at a lower cost.

There is significant competition in the development of a vaccine against COVID-19, influenza, and RSV and we may never see returns on the significant resources we are devoting to our vaccine candidates.

We may be unable to produce a successful COVID-19 vaccine and establish a competitive market share for our vaccine before a competitor, or before the COVID-19 outbreak is contained or significantly diminished. A large number of vaccine manufacturers, academic institutions and other organizations have developed COVID-19 vaccines or are developing COVID-19 vaccine candidates. In particular, Moderna, Pfizer/BioNTech, and Johnson & Johnson have received emergency use authorizations for their COVID-19 vaccines in the U.S., among other countries, and many other companies, including AstraZeneca, Sinovac Biotech, Sinopharm, and Inovio are in various stages of developing COVID-19 vaccine candidates. Despite funding provided to us to date, many of our competitors pursuing vaccine candidates have significantly greater product candidate development, manufacturing and marketing resources than we do. Larger pharmaceutical and biotechnology companies have extensive experience in clinical testing and obtaining regulatory approval for their products and may have the resources to heavily invest in the development of their vaccine candidates. Our business could be materially and adversely affected if competitors develop and commercialize one or more COVID-19 vaccines before we can complete development and seek approval for our vaccine candidate, or if they develop and commercialize one or more COVID-19 vaccines that are safer, more effective, have fewer or less severe side effects, have broader market acceptance, are more convenient or are less expensive than any vaccine candidate that we may develop. Furthermore, if any competitors are successful in producing a more efficacious vaccine or other treatment for COVID-19, or if any competitors are able to manufacture and distribute any such vaccines or treatments with greater efficiency, there may be a diversion of potential governmental and other funding away from us and toward such other parties.

We are allocating significant financial and personnel resources to the development of NVX-CoV2373, which may cause delays in or otherwise negatively impact our other development programs. Our business could be materially impacted by our allocation of significant resources to combating a global health threat that is unpredictable or against which our vaccine, if developed, may not be partially or fully effective, and may ultimately prove unsuccessful or unprofitable.

Many seasonal influenza vaccines are currently approved and marketed. Competition in the sale of these seasonal influenza vaccines is intense. Therefore, newly developed and approved products must be differentiated and competitive as a result of products introduced by our competitors to the marketplace more rapidly and at a lower cost.
than those currently marketed. Our nanoparticle seasonal influenza vaccine candidate may not prove to be more efficacious than current products or products under development by our competitors. Further, our in-house or third-party manufacturing arrangements may not provide enough savings of time or money to provide the required differentiation for commercial success.

We are also aware that there are multiple companies with active RSV vaccine programs at various stages of development. Thus, while there is no RSV vaccine currently on the market, there is likely to be significant and consistent competition as these active programs mature. Different RSV vaccines may work better for different segments of the population, so it may be difficult for a single RSV vaccine manufacturer to provide vaccines that are marketable to multiple population segments. Geographic markets are also likely to vary significantly, which may make it difficult to market a single RSV vaccine worldwide. Even if a manufacturer brings an RSV vaccine to license, it is likely that competitors will continue to work on new products that could be more efficacious and/or less expensive. Our RSV vaccine candidate may not be as far along in development as other active RSV vaccine programs about which we are not aware, nor as efficacious as products under development by competing companies. Even if our RSV vaccine candidate receives regulatory approval, it may not achieve significant sales if other, more effective vaccines under development by our competitors are also approved.

Risks Related to Regulatory and Compliance Matters

We have not completed the development of vaccine products and we may not succeed in obtaining the FDA licensure necessary to sell such vaccine products.

The development, manufacture and marketing of our pharmaceutical and biological products are subject to government regulation by the U.S. FDA and regulatory authorities in other countries, including the European Medicines Agency EMA, the State Institute for Drug Control (SUKL) with respect to our manufacturing facility in the Czech Republic and the Swedish Medical Products Agency (Läkemedelsverket, LV) with respect to our adjuvant product being developed in Sweden, as well as other country authorities into which active pharmaceutical ingredients and excipients are imported and/or manufactured by us or our sub-contracted manufacturers. In the U.S. and most foreign countries, we must complete rigorous preclinical testing and extensive clinical trials that demonstrate the safety and efficacy of a product in order to apply for regulatory approval to market the product. None of our vaccine candidates has yet gained regulatory approval in the U.S. or elsewhere. We also have vaccine candidates in clinical trials and preclinical laboratory or animal studies. The steps generally required by the FDA before our proposed investigational products may be marketed in the U.S. include:

- performance of preclinical (animal and laboratory) tests;
- submission to the FDA of an IND, which must become effective before clinical trials may commence;
- performance of adequate and well controlled clinical trials to establish the safety and efficacy of the investigational product in the intended target population;
- performance of a consistent and reproducible manufacturing process at commercial scale capable of passing FDA inspection;
- submission to the FDA of a BLA or a NDA; and
- FDA approval of the BLA or NDA before any commercial sale or shipment of the product.

These processes are expensive and can take many years to complete, and we may not be able to demonstrate the safety and efficacy of our vaccine candidates to the satisfaction of regulatory authorities. The start of clinical trials can be delayed or take longer than anticipated for many and varied reasons, many of which are out of our control. Safety concerns may emerge that could lengthen the ongoing clinical trials or require additional clinical trials to be conducted. Promising results in early clinical trials may not be replicated in subsequent clinical trials. Regulatory authorities may also require additional testing, and we may be required to demonstrate that our proposed products represent an improved form of treatment over existing therapies, which we may be unable to do without conducting further clinical trials. Moreover, if the FDA grants regulatory approval of a product, the approval may be limited to specific indications or limited with respect to its distribution. Expanded or additional indications for approved products may not be approved, which could limit our revenue. Foreign regulatory authorities may apply similar limitations or may refuse to grant any approval. Consequently, even if we believe that preclinical and clinical data are sufficient to support regulatory approval for our vaccine candidates, the FDA and foreign regulatory authorities ultimately may not grant approval for commercial sale in any jurisdiction or may impose regulatory requirements that make further pursuit of approval uneconomical in one or more jurisdictions. If our vaccine candidates are not approved, our ability to generate revenue will be limited and our business will be adversely affected.

We may fail to obtain regulatory approval for our products on a timely basis or comply with our continuing regulatory obligations after approval is obtained.

Delays in obtaining regulatory approval can be extremely costly in terms of lost sales opportunities, loss of any potential marketing advantage of being early to market and increased clinical trial costs. The speed with which we begin and complete our preclinical studies necessary to begin clinical trials, clinical trials and our applications for marketing approval will depend on several factors, including the following:

- our ability to manufacture or obtain sufficient quantities of materials for use in necessary preclinical studies and clinical trials;
- regulatory agency review and approval of proposed clinical trial protocols;
- approval of clinical trials protocols and informed consent forms by institutional review boards responsible for overseeing the ethical conduct of the trial;
- the rate of participant enrollment and retention, which is a function of many factors, including the size of the participant population, the proximity of participants to clinical sites, the eligibility criteria for the clinical trial and the nature of the protocol;
- unfavorable test results or side effects experienced by clinical trial participants;
- analysis of data obtained from preclinical and clinical activities, which are susceptible to varying interpretations and which interpretations could delay, limit, result in the suspension or termination of, or prevent further conduct of clinical studies or regulatory approval;
- the availability of skilled and experienced staff to conduct and monitor clinical trials and to prepare the appropriate regulatory applications; and
- changes in the policies of regulatory authorities for drug or vaccine approval during the period of product development.

We have limited experience in conducting and managing the preclinical studies and clinical trials necessary to obtain regulatory marketing approvals. We may not be permitted to continue or commence additional clinical trials. We also face the risk that the results of our clinical trials may be inconsistent with the results obtained in preclinical studies or clinical trials of similar products or that the results obtained in later phases of clinical trials may be inconsistent with those obtained in earlier phases. A number of companies in the biotechnology and product development industry have suffered significant setbacks in advanced clinical trials, even after experiencing promising results in early animal and human testing.

Regulatory agencies may require us or our collaborators to delay, restrict or discontinue clinical trials on various grounds, including a finding that the participants are being exposed to an unacceptable health risk. In addition, we or our collaborators may be unable to submit applications to regulatory agencies within the time frame we currently expect. Once submitted, applications must be approved by various regulatory agencies before we or our collaborators can commercialize the product described in the application. All statutes and regulations governing the conduct of clinical trials are subject to change in the future, which could affect the cost of such clinical trials. Any unanticipated costs or delays in our clinical trials could delay our ability to generate revenue and harm our financial condition and results of operations.
Failure to obtain regulatory approval in foreign jurisdictions would prevent us from marketing our products internationally.

We intend to have our vaccine candidates marketed outside the U.S. In furtherance of this objective, we have entered into supply agreements with various foreign governments and international distribution agreements with commercial entities. In order to market our products in the European Union, United Kingdom, India, Asia and many other non-U.S. jurisdictions, we must obtain separate regulatory approvals and comply with numerous and varying regulatory requirements. The approval procedure varies among countries and can involve additional testing and data review. The time required to obtain foreign regulatory approval may differ from that required to obtain FDA approval. The foreign regulatory approval process may include all of the risks associated with obtaining FDA approval. We may not obtain foreign regulatory approvals on a timely basis, if at all. Approval by a regulatory agency, such as the FDA, does not ensure approval by regulatory agencies in other foreign countries. However, a failure or delay in obtaining regulatory approval in one jurisdiction may have a negative effect on the regulatory approval process in other jurisdictions, including approval by the FDA. The failure to obtain regulatory approval in foreign jurisdictions could harm our business.

The regulatory pathway for NVX-CoV2373 is continually evolving and may result in unexpected or unforeseen challenges.

The regulatory pathway for NVX-CoV2373 is evolving and failure by us to comply with any laws, rules and standards, some of which may not exist yet or are subject to interpretation and may be subject to change, could result in a variety of adverse consequences, including penalties, fines and delays in vaccine licensure. Efforts to comply with evolving laws, regulations and standards have resulted in, and are likely to continue to result in, increased general and administrative expenses and a diversion of management time and attention to regulatory compliance activities. For example, the rules, regulations and standards governing OWS are uncertain and may evolve as the program progresses. Such rules or standards may adversely affect our plans to develop NVX-CoV2373 and failure by us to comply with any laws, rules or standards, some of which may not exist yet or may change, could result in a range of adverse consequences, such as penalties, fines or failure to receive funding.

The speed at which multiple stakeholders are moving to create, test and approve a vaccine for COVID-19 is highly unusual and may increase the risks associated with traditional vaccine development, which typically takes between eight and ten years. Given this accelerated timeline, we and regulators, such as the FDA, the EMA, and the MHRA, may make decisions more rapidly than is typical. Evolving or changing plans or priorities at the FDA or other regulatory bodies, including based on new knowledge of COVID-19 and how the disease affects the human body, may significantly affect the regulatory pathway for NVX-CoV2373. Results from clinical testing may raise new questions and require us to redesign proposed clinical trials, including revising proposed endpoints or adding new clinical trial sites or cohorts of subjects. In addition, the FDA’s or other regulators’ analysis of clinical data may differ from our interpretation, or regulators’ requirements and expectations for vaccine authorization or approval may change over time, with the result that the FDA or other regulators may require that we conduct additional clinical trials or non-clinical studies. There can be no guarantee that the evolving regulatory pathway will not impede the development, commercialization and/or licensure of NVX-CoV2373.

In addition, because the path to licensure of any vaccine against COVID-19 is unclear, we may have a widely used vaccine in circulation in the U.S. or another country as an investigational vaccine or a product authorized for temporary or emergency use prior to our receipt of marketing approval. Unexpected safety issues in these circumstances could lead to significant reputational damage for Novavax and our technology platform going forward and other issues, including delays in our other programs, the need for re-design of our clinical trials and the need for significant additional financial resources.

Participants or prospective participants in our clinical trials of NVX-CoV2373 could receive one of multiple COVID-19 vaccines that have been granted emergency use authorizations or approvals in the United States or other countries, which could impact or delay our clinical development program for NVX-CoV2373.

Multiple COVID-19 vaccines have received temporary or emergency use authorization in the U.S. or in other countries. Moderna, Pfizer/BioNTech, and Johnson & Johnson have received emergency use authorizations for their COVID-19 vaccines in the U.S., among other countries, and AstraZeneca, Sinovac Biotech, Sinopharm, and others have been authorized in some manner in at least one country. Participants in our clinical trials could choose to receive a COVID-19 vaccine authorized or approved in the United States or other countries. Some participants in current studies of NVX-CoV2373 could choose to receive an authorized or approved COVID-19 vaccine or drop out of our studies altogether, particularly if they believe they may be in the placebo arm in one of our trials. The availability of authorized COVID-19 vaccines could affect our clinical trial results and impede our ability to collect sufficient data from previously enrolled participants, which could require additional enrollment or trials, either of which would be costly and time-consuming and could delay or permanently halt our development of NVX-CoV2373.

We are conducting, and plan to conduct in the future, a number of clinical trials for NVX-CoV2373 at sites outside the United States, and the FDA may not accept data from trials conducted in such locations.

We are currently conducting several clinical trials of NVX-CoV2373 at sites outside the U.S., including a Phase 3 trial in the UK, a Phase 2b trial in South Africa, and a Phase 1/2 trial partially in Australia. We also plan in the future to conduct (or collaborate to conduct) a Phase 2/3 trial in India, Phase 2 trial in Czech Republic, and Phase 1/2 trial in Japan. Although the FDA may accept data from clinical trials conducted outside the U.S., acceptance of these data is subject to conditions imposed by the FDA. For example, the clinical trial must be well designed and conducted and be performed by qualified investigators in accordance with ethical principles. The trial population must also adequately represent the U.S. population, and the data must be applicable to the U.S. population and U.S. medical practice in ways that the FDA deems clinically meaningful. In addition, while these clinical trials are subject to the applicable local laws, FDA acceptance of the data will depend on its determination that the trials also complied with all applicable U.S. laws and regulations. If the FDA does not accept the data from any trial that we conduct outside the U.S., it could result in delay pending completion of our trials conducted in the U.S. or result in the need for additional trials, which would be costly and time-consuming and could delay or permanently halt our development of NVX-CoV2373.

Even if regulatory approval is received for our vaccine candidates, the later discovery of previously unknown problems with a product, manufacturer or facility may result in restrictions, including withdrawal of the product from the market.

Even after a product gains regulatory approval, the product and the manufacturer of the product will be subject to continuing regulatory review, including adverse event reporting requirements and the FDA’s general prohibition against promoting products for unapproved uses. Failure to comply with any post-approval requirements can, among other things, result in warning letters, product seizures, recalls, substantial fines, injunctions, suspensions or revocations of marketing licenses, operating restrictions and criminal prosecutions. Any such enforcement actions, any unanticipated changes in existing regulatory requirements or the adoption of new requirements, or any safety issues that arise with any approved products, could adversely affect our ability to market products and generate revenue and thus adversely affect our ability to continue our business.

We also may be restricted or prohibited from marketing or manufacturing a product, even after obtaining product approval, if previously unknown problems with the product or its manufacture are subsequently discovered. We cannot provide assurance that newly discovered or developed safety issues will not arise following regulatory approval. With the use of any vaccine by a wide patient population, serious adverse events may occur from time to time that did not arise in the clinical trials of the product or that initially appeared to be unrelated to the vaccine itself and only with the collection of subsequent information were found to be causally related to the product. Any such safety issues could cause us to suspend or cease marketing of our approved products, possibly subject us to substantial liabilities, and adversely affect our ability to generate revenue and our financial condition.
Our ability to produce a successful vaccine may be curtailed by one or more government actions or interventions, which may be more likely during a global health crisis such as COVID-19.

Given the significant global impact of the COVID-19 pandemic, it is possible that one or more government entities may take actions, including the U.S. government under the Defense Production Act of 1950, as amended, which could directly or indirectly have the effect of diminishing some of our rights or opportunities with respect to NVX-CoV2373 and the economic value of a COVID-19 vaccine to us could be limited. In addition, during a global health crisis, such as the COVID-19 pandemic, where the spread of a disease needs to be controlled, closed or heavily regulated national borders will create challenges and potential delays in our development production and distribution activities and may necessitate that we pursue strategies to develop, produce and distribute our vaccine candidates within self-contained national or international borders or with additional safety measures or checks in place, at potentially much greater expense and with longer timeframes for public distribution.

Inadequate funding for the FDA, the SEC and other government agencies could hinder their ability to hire and retain key leadership and other personnel, or otherwise perform their normal functions on which the operation of our business may rely, which could negatively impact our ability to develop or commercialize new products or services, access capital markets, or otherwise operate our business.

The ability of the FDA to review and approve new products can be affected by a variety of factors, including government budget and funding levels, ability to hire and retain key personnel and accept the payment of user fees, and statutory, regulatory and policy changes. Average review times at the agency have fluctuated in recent years as a result. In addition, government funding of the SEC and other government agencies on which our operations may rely, including those that fund research and development activities, is subject to the political process, which is inherently fluid and unpredictable.

Disruptions at the FDA and other agencies may also slow the time necessary for new drugs to be reviewed and approved by necessary government agencies, which would adversely affect our business. For example, over the last several years, the U.S. government has shut down several times and certain regulatory agencies, such as the FDA and the SEC, have had to furlough employees and stop or slow the pace of critical activities. If a prolonged government shutdown occurs, it could significantly impact the ability of the FDA to timely review and process our regulatory submissions, which could have a material adverse effect on our business. Further, in our operations as a public company, future government shutdowns could impact our ability to access the public markets and obtain necessary capital in order to properly capitalize and continue our operations.

Fast Track Designation by the FDA or other regulatory acceleration options may not actually lead to a faster development or regulatory review or approval process and does not assure approval.

If a drug is intended for the treatment of a serious or life-threatening condition and the drug demonstrates the potential to address an unmet medical need for this condition, the drug sponsor may apply for FDA Fast Track Designation or similar fast track processes with other regulatory agencies, such as conditional marketing authorizations from the EMA. However, Fast Track Designation does not ensure that the drug sponsor will receive marketing approval or that approval will be granted within any particular timeframe. The FDA granted Fast Track Designation for NVX-CoV2373 in November 2020 and for NanoFlu, our recombinant quadivalent seasonal influenza vaccine candidate, in January 2020. We may also seek Fast Track Designation for more of our other vaccine candidates. If we do seek Fast Track Designation for our other vaccine candidates, we may not receive it, and even if we receive Fast Track Designation, we may not experience a faster development process, review or approval compared to conventional FDA procedures. In addition, the FDA may withdraw Fast Track designation if it believes that the designation is no longer supported by data from our clinical development program. Fast Track Designation alone does not guarantee qualification for the FDA’s priority review procedures.

Obtaining a Fast-Track Designation does not change the standards for product approval but may expedite the development or approval process. Even though the FDA has granted such designation for NVX-CoV2373 and NanoFlu, it may not actually result in faster clinical development or regulatory review or approval. Furthermore, such a designation does not increase the likelihood that NVX-CoV2373 or NanoFlu will receive marketing approval in the U.S.

Because we are subject to environmental, health and safety laws, we may be unable to conduct our business in the most advantageous manner.

We are subject to various laws and regulations relating to safe working conditions, laboratory and manufacturing practices, the experimental use of animals, emissions and wastewater discharges, and the use and disposal of hazardous or potentially hazardous substances used in connection with our research, including infectious disease agents. We also cannot accurately predict the extent of regulations that might result from any future legislative or administrative action. Any of these laws or regulations could cause us to incur additional expense or restrict our operations.

Our facilities in Maryland are subject to various local, state and federal laws and regulations relating to safe working conditions, laboratory practices, the experimental use of animals and the use and disposal of hazardous or potentially hazardous substances, including chemicals, microorganisms and various hazardous compounds used in connection with our research and development activities. In the U.S., these laws include the Occupational Safety and Health Act, the Toxic Substances Control Act and the Resource Conservation and Recovery Act. Similar national and local regulations govern our facilities in Sweden and the Czech Republic. We cannot eliminate the risk of accidental contamination or discharge or injury from these materials. Federal, state and local laws and regulations govern the use, manufacture, storage, handling and disposal of these materials. We could be subject to civil damages in the event of an improper or unauthorized release of, or exposure of individuals to, these hazardous materials. In addition, claimants may sue us for injury or contamination that results from our use or the use by third-parties of these materials, and our liability may exceed our total assets. Compliance with environmental laws and regulations may be expensive, and current or future environmental regulations may impair our research, development or production efforts.

Although we have general liability insurance, these policies contain exclusions from insurance against claims arising from pollution by chemicals or pollution from conditions arising from our operations. Our collaborators are working with these types of hazardous materials in connection with our collaborations. In the event of a lawsuit or investigation, we could be held responsible for any injury we or our collaborators cause to persons or property by exposure to, or release of, any hazardous materials. However, we believe that we are currently in compliance with all material applicable environmental and occupational health and safety regulations.

For our product candidates, we will be subject to additional healthcare laws and our failure to comply with these laws could have a material adverse effect on our results of operations and financial conditions.

Within the U.S. (and within foreign countries), if we obtain approval for any of our product candidates and begin commercializing them, our operations may be directly, or indirectly through our arrangements with third-party payors and customers, subject to additional healthcare regulation and enforcement by the federal and state governments (or the regulatory bodies or governments of foreign countries), which may constrain the business or financial arrangements and relationships through which we sell, market and distribute our products. These laws and regulations may restrict or prohibit a wide range of price, discounting, marketing and promotion, structuring and commission(s), certain customer incentive programs and other business arrangements generally. Activities subject to these laws also involve the improper use of information obtained in the course of patient recruitment for clinical trials. The applicable U.S. federal and state healthcare laws and regulations (which may be comparable to foreign laws existing in foreign countries) that may affect our ability to operate include:

- the Federal Food, Drug and Cosmetic Act, which among other things, strictly regulates drug product marketing and promotion and prohibits manufacturers from marketing such products for unapproved uses;
- the federal Anti-Kickback Statute, which prohibits, among other things, persons from knowingly and willfully soliciting, receiving or providing remuneration, directly or indirectly, to induce the referral for an item or service or the purchasing or ordering of a good or service, for which payment may be made under federal healthcare programs such as Medicare and Medicaid;
• federal false claims laws, including the FCA, which prohibit, among other things, individuals or entities from knowingly presenting, or causing to be presented, information or claims for payment from Medicare, Medicaid, or other third-party payers that are false or fraudulent;
• manufacturers can be held liable under the FCA even when they do not submit claims directly to government payors if they are deemed to “cause” the submission of false or fraudulent claims; the FCA also permits a private individual acting as whistleblower to bring actions on behalf of the federal government alleging violations of the FCA and to share in any monetary recovery;
• federal laws that require pharmaceutical manufacturers to report certain calculated product prices to the government or provide certain discounts or rebates to government authorities or private entities, often as a condition of reimbursement under government healthcare programs;
• the federal Physician Payment Sunshine Act and its implementing regulations, which require manufacturers of drugs, devices, biologicals, and medical supplies for which payment is available under Medicare, Medicaid or the Children’s Health Insurance Program (with certain exceptions) to report annually to the DHHS information related to payments or other transfers of value made to physicians (defined to include doctors, dentists, optometrists and chiropractors) and teaching hospitals, as well as ownership and investment interests held by physicians and their immediate family members; effective January 1, 2022, these reporting obligations will extend to include transfers of value made to certain non-physician providers such as physician assistants and nurse practitioners;
• the federal law known as HIPAA, which, in addition to privacy protections applicable to healthcare providers and other entities, prohibits executing a scheme to defraud any healthcare benefit program or making false statements relating to healthcare matters;
• federal consumer protection and unfair competition laws, which broadly regulate marketplace activities and activities that potentially harm consumers;
• state law equivalents of the above federal laws, such as anti-kickback and false claims laws which may apply to items or services reimbursed by any third-party payer, including commercial insurers, and state gift ban and transparency laws, many of which state laws differ from each other in significant ways and often are not preempted by federal laws, thus complicating compliance efforts; and
• state laws restricting interactions with healthcare providers and other members of the healthcare community or requiring pharmaceutical manufacturers to implement certain compliance standards.

Because of the breadth of these laws and the narrowness of the statutory exceptions and safe harbors available, it is possible that some of our business activities could be subject to challenge under one or more of such laws. If our operations are found to be in violation of any of such laws or any other governmental regulations that apply to us, we may be subject to, on a corporate or individual basis, penalties, including civil and criminal penalties, damages, fines, the curtailment or restructuring of our operations, the exclusion from participation in federal and state healthcare programs and even imprisonment, any of which could materially adversely affect our ability to operate our business and our financial results. In addition, the cost of implementing sufficient systems, controls, and processes to ensure compliance with all of the aforementioned laws could be significant. Any action for violation of these laws, even if successfully defended, could cause us to incur significant legal expenses and divert management’s attention from the operation of the company’s business. If any of the physicians or other healthcare providers or entities with whom we expect to do business is found not to be in compliance with applicable laws, that person or entity may be subject to criminal, civil or administrative sanctions, including exclusions from government funded healthcare programs. Prohibitions or restrictions on sales or withdrawal of future marketed products could materially affect business in an adverse way.

It is not always possible to identify and deter employee misconduct, and the precautions we take to detect and prevent inappropriate conduct may not be effective in controlling unknown or unmanaged risks or losses or in protecting us from governmental investigations or other actions or lawsuits stemming from a failure to be in compliance with such laws or regulations. Efforts to ensure that our business arrangements will comply with applicable healthcare laws may involve substantial costs. It is possible that governmental and enforcement authorities will conclude that our business practices may not comply with current or future statutes, regulations or case law interpreting applicable fraud and abuse or other healthcare laws and regulations. If any such actions are instituted against us and we are not successful in defending ourselves or asserting our rights those actions, our business may be impaired.

Risks Related to our Intellectual Property

Our success depends on our ability to maintain the proprietary nature of our technology.

Our success in large part depends on our ability to maintain the proprietary nature of our technology and other trade secrets. To do so, we must protect and maintain existing patents, obtain new patents and pursue trade secret and other intellectual property protection. We also must operate without infringing the proprietary rights of third-parties or allowing third-parties to infringe our rights. We currently have or have rights to over 450 U.S. patents and corresponding foreign patents and patent applications covering our technologies. However, patent issues relating to pharmaceuticals and biologics involve complex legal, scientific and factual questions. To date, no consistent policy has emerged regarding the breadth of biotechnology patent claims that are granted by the U.S. Patent and Trademark Office (“USPTO”) or enforced by the federal courts. Therefore, we do not know whether any particular patent applications will result in the issuance of patents, or that any patents issued to us will provide us with any competitive advantage. We also cannot be sure that we will develop additional proprietary products that are patentable. Furthermore, there is a risk that others will independently develop or duplicate similar technology or products or circumvent the patents issued to us.

There is a risk that third-parties may challenge our existing patents or claim that we are infringing their patents or proprietary rights. We could incur substantial costs in defending patent infringement suits or in filing suits against others to have their patents declared invalid or claim infringement. It is also possible that we may be required to obtain licenses from third-parties to avoid infringing third-party patents or other proprietary rights. We cannot be sure that such third-party licenses would be available to us on acceptable terms, if at all. If we are unable to obtain required third-party licenses, we may be delayed in or prohibited from developing, manufacturing or selling products requiring such licenses.

Although our patent filings include claims covering various features of our vaccine candidates, including composition, methods of manufacture and use, our patents do not provide us with complete protection against the development of competing products. Some of our know-how and technology is not patentable. To protect our proprietary rights in unpatentable intellectual property and trade secrets, we require employees, consultants, advisors and collaborators to enter into confidentiality agreements. These agreements may not provide meaningful protection for our trade secrets, know-how or other proprietary information.

Third-parties may claim we infringe their intellectual property rights.

Our research, development and commercialization activities, including any vaccine candidates resulting from these activities, may be found to infringe patents owned by third-parties and to which we do not hold licenses or other rights. There may be rights we are not aware of, including applications that have been filed, but not published that, when issued, could be asserted against us. These third-parties could bring claims against us, and that may cause us to stop manufacturing and selling our product or require license fees or royalty payments. We cannot prevent third-parties from seeking to have us enjoined under federal or state laws restricting interactions with healthcare providers and other members of the healthcare community or requiring pharmaceutical manufacturers to implement certain compliance standards.

As a result of patent infringement claims, or in order to avoid potential claims, we may choose or be required to seek a license from the third-party. These licenses may not be available on acceptable terms, or at all. Even if we are able to obtain a license, the license would likely obligate us to pay license fees or royalties or both, and the rights granted to us might be non-exclusive, which could result in our competitors gaining access to the same intellectual property. Ultimately, we could be prevented from commercializing a product, or be forced to cease some aspect of our business operations, if, as a result of actual or threatened patent infringement claims, we are unable to enter into licenses on acceptable terms. All of

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the issues described above could also impact our collaborators, which would also impact the success of the collaboration and therefore us.

There has been substantial litigation and other proceedings regarding patent and other intellectual property rights in the pharmaceutical and biotechnology industries.

We may become involved in litigation to protect or enforce our patents or the patents of our collaborators or licensors, which could be expensive and time-consuming.

Competitors may infringe our patents or the patents of our collaborators or licensors. As a result, we may be required to file suit to counter infringement for unauthorized use. This can be expensive and time-consuming. In addition, an infringement proceeding, a court may decide that a patent of ours is not valid or is unenforceable, or may refuse to stop the other party from using the technology at issue on the grounds that our patents do not cover its technology. An adverse determination of any litigation or defense proceeding could put one or more of our patents at risk of being invalidated or interpreted narrowly and could put our patent applications at the risk of not issuing.

Even if we are successful, litigation may result in substantial costs and distraction to our management. Even with a broad portfolio, we may not be able, alone or with our collaborators and licensors, to prevent misappropriation of our proprietary rights, particularly in countries where the laws may not protect such rights as fully as in the U.S.

Furthermore, because of the substantial amount of discovery required in connection with intellectual property litigation, there is a risk that some of our confidential information could be compromised by disclosure during this type of litigation. In addition, during the course of litigation, there could be public announcements of the results of hearings, motions or other interim proceedings or developments. If investors perceive these results to be negative, the market price for our common stock could be significantly harmed.

The scope, validity, and ownership of our patent claims may be challenged in various venues and, if we do not prevail, our ability to exclude competitors may be harmed, potentially reducing our ability to succeed commercially.

We may be subject to a variety of challenges from third-parties that relate to the scope of the claims or to their validity. Such challenges can be mounted in post-grant review, ex parte re-examination, and inter partes review proceedings before the USPTO, or similar adversarial proceedings in other jurisdictions. If we are unsuccessful in any such challenge, the scope of our claims could be narrowed or could be invalidated. Any such outcome could impair our ability to exclude competitors from the market in those countries, potentially impacting our commercial success.

Our patents may be subject to various challenges related to ownership and inventorship, including interference or derivation proceedings. Third-parties may assert that they are inventors on our patents or that they are owners of the patents. While we perform inventorship analyses to insure that the correct inventors are listed on our patents, we cannot be certain that a court of competent jurisdiction would arrive at the same conclusions we do. If we are unsuccessful in defending against ownership or inventorship challenges, a court may require us to list additional inventors, may invalidate the patent, or may transfer ownership of the patent to a third-party. Any of these outcomes may harm our ability to exclude competitors and potentially impact our commercial success. Further, if ownership is transferred to a third-party, we may be required to seek a license to those rights to preserve our exclusive ability to practice the invention. Such a license may not be available on commercially reasonable terms, or at all. If we are unable to obtain a license, we may be required to expend time, effort, and other resources to design around the patent. Any such license may be non-exclusive and if a competitor is able to obtain a license from the third-party, our ability to exclude that competitor from the market may be negatively impacted.

Even if we are ultimately successful, defending any such challenges may cause us to incur substantial expenses and may require us to divert substantial financial and management resources that we would otherwise be able to devote to our business.

We may need to license intellectual property from third-parties and, if our right to use the intellectual property we license is affected, our ability to develop and commercialize our vaccine candidates may be harmed.

We have in the past, and we expect in the future to license intellectual property from third-parties and that these licenses will be material to our business. We will not own the patents or patent applications that underlie these licenses, and we may not control either the prosecution or the enforcement of the patents. Under such circumstances, we may be forced to rely upon our licensors to properly prosecute and file those patent applications and prevent infringement of those patents.

While many of the licenses under which we have rights provide us with rights in specified fields, the scope of our rights under these and other licenses may be subject to dispute by our licensors or third-parties. In addition, our rights to use these technologies and practice the inventions claimed in the licensed patents and patent applications are subject to our licensors abiding by the terms of those licenses and not terminating them. Any of our licenses may be terminated by the licensor if we are in breach of a term or condition of the license agreement, or in certain other circumstances.

Further, any disputes regarding obligations in licenses may require us to take expensive and time-consuming legal action to resolve, and, even if we are successful, may delay our ability to commercialize products and generate revenue. Further, if we are unable to resolve license issues that arise, we may lose rights to practice intellectual property that is required to make, use, or sell products. Any such loss could compromise our development and commercialization efforts for current or future product candidates and/or may require additional effort and expense to design around.

Our vaccine candidates and potential vaccine candidates will require several components that may each be the subject of a license agreement. The cumulative license fees and royalties for these components may make the commercialization of these vaccine candidates uneconomical.

If patent laws or the interpretation of patent laws change, our competitors may be able to develop and commercialize our discoveries.

Important legal issues remain to be resolved as to the extent and scope of available patent protection for biopharmaceutical products and processes in the U.S. and other important markets outside the U.S., such as Europe and Japan. In addition, foreign markets may not provide the same level of patent protection as provided under the U.S. patent system. Litigation or administrative proceedings may be necessary to determine the validity and scope of patent rights. Any such litigation or proceeding may result in a significant commitment of time and resources in the future and could force us to do one or more of the following: cease selling or using any of our products that incorporate the challenged intellectual property, which would adversely affect our revenue; obtain a license from the holder of the intellectual property right alleged to have been infringed, which license may not be available on reasonable terms, if at all; and redesign our products to avoid infringing the intellectual property rights of third-parties, which may be time-consuming or impossible to do. In addition, changes in, or different interpretations of, patent laws in the U.S. and other countries may result in patent laws that allow others to use our discoveries or develop and commercialize our products. We cannot provide assurance that the patents we obtain or the unpatented technology we hold will afford us significant commercial protection.

If we do not obtain patent term extension and/or patent term adjustment in the United States under the Hatch-Waxman Act and similar extensions in foreign countries, our ability to exclude competitors may be harmed.

In the United States, the patent term is 20 years from the earliest U.S. non-provisional filing date. Extensions of patent term may be available under certain circumstances. Depending upon the timing, duration and conditions of FDA marketing approval of our product candidates, we may be able to extend the term of one patent that covers a marketed product under the Drug Price Competition and Patent Term Restoration Act of 1984, (the "Hatch-Waxman Amendments") and similar legislation in the European Union.
The Hatch-Waxman Amendments permit patent term extension of up to five years for a patent covering an approved product as compensation for effective patent term lost during product development and the FDA regulatory review process. We may not receive any extension if we fail to apply within applicable deadlines, fail to apply prior to expiration of the patent term, or otherwise fail to satisfy applicable requirements. Moreover, the length of the extension could be less than we request. If we are unable to obtain patent term extension or the term of any such extension is less than we request, the period during which we can enforce our patent rights for that product will be shortened and our competitors may obtain approval to market competing products sooner.

Patent term covering our products may also be extended for time spent during the prosecution of the patent application in the USPTO. This extension is referred to as Patent Term Adjustment (“PTA”). The laws and regulations governing how the USPTO calculates the PTA is subject to change and changes in the law can reduce or increase any such PTA. Further, the PTA granted by the USPTO may be challenged by a third-party. If we do not prevail under such a challenge, the PTA may be reduced or eliminated, shortening the patent term, which may negatively impact our ability to exclude competitors.

Risks Related to Employee Matters, Managing Growth and Information Technology

Our business may be adversely affected if we do not successfully execute our business development initiatives.

We anticipate growing through both internal development projects, as well as external opportunities, which include the acquisition, partnering and in-licensing of products, technologies and companies or the entry into strategic alliances and collaborations. The availability of high-quality opportunities is limited, and we may fail to identify candidates that we and our stockholders consider suitable or complete transactions on terms that prove advantageous. In order to pursue such opportunities, we may require significant additional financing, which may not be available to us on favorable terms, if at all. Even if we are able to successfully identify and complete acquisitions, like our business combinations with Novavax CZ (formerly Praha Vaccines a.s.) and Novavax AB, strategic transactions involve many risks, including, among others, those related to relevant patent or other rights’ failure to satisfy applicable requirements, unanticipated expenses and liabilities, and increased complexity of our operations, which could prevent us from effectively exploiting acquired facilities, successfully integrating the acquired business and personnel, or fully realizing expected synergies.

To effectively manage our current and future potential growth, we will need to continue to enhance our operational, financial and management processes and to effectively expand, train and manage our employee base. Supporting our growth will require us to successfully integrate the acquired business and personnel, or fully realize expected synergies.

Security breaches and other disruptions could compromise our information and expose us to liability, and our failure to comply with data protection laws and regulations could lead to government enforcement actions, which would cause our business and reputation to suffer.

In the ordinary course of our business, we collect and store sensitive data, including intellectual property, our proprietary business information and data about our clinical participants, suppliers and business partners and personally identifiable information. The secure maintenance of this information is critical to our operations and business strategy. Some of this information could be an attractive target of criminal attack by malicious third-parties with a wide range of motives and expertise, including organized criminal groups, “hacktivists,” patient groups, disgruntled current or former employees and others. Hacker attacks are of ever-increasing levels of sophistication, and despite our security measures, our information technology will require significant expenditures and resources, including investment in personnel and infrastructure to be vulnerable to such attacks or may be breached due to employee error or malfeasance. Any such breach could compromise our networks and the information stored there could be accessed, publicly disclosed, lost or stolen. Furthermore, if our systems become compromised, we may not promptly discover the intrusion. Like other companies in our industry, we have experienced attacks to our data and systems, including malware and computer viruses. Attacks could have a material impact on our business, operations or financial results. Any access, disclosure or other loss of information could result in legal claims or proceedings, liability under laws that protect the privacy of personal information, disrupt our operations, and damage our reputation, which could adversely affect our business. In addition, privacy and data protection laws may be interpreted and applied differently from country to country and may create inconsistent or conflicting requirements, which can increase the costs incurred by us in complying with such laws. The European Union’s GDPR, which greatly increases the jurisdictional reach of European Union law and became effective in May 2018, adds a broad array of requirements for handling personal data including the public disclosure of significant data breaches, and imposes substantial penalties for non-compliance of up to the greater of €20 million or 4% of global annual revenue for the preceding financial year. Our efforts to comply with GDPR and other privacy and data protection laws may impose significant costs and challenges that are likely to increase over time, and we could incur substantial penalties or litigation related to violations of existing or future data privacy laws and regulations.

Additionally, the CCPA, which became effective January 1, 2020, substantially expands privacy obligations of many businesses. The CCPA requires new disclosures to California consumers, imposes new rules for collecting or using information about minors, and affords consumers new abilities, such as the right to know whether the data is sold or disclosed and to whom, the right to request that a company delete personal information collected, the right to opt-out of the sale of personal information and the right to non-discrimination in terms of price or service when a consumer exercises a privacy right. If we fail to comply with these regulations, we could be subject to civil sanctions, including fines and penalties for noncompliance. The CCPA provides for civil penalties for violations, as well as a private right of action for data breaches that is expected to increase data breach litigation. Moreover, a newly passed ballot initiative, the California Privacy Rights Act (“CPRA”), which will become operational in 2023, expands on the CCPA, creating new consumer rights and protections, including the right to correct personal information, the right to opt out of the use of personal information in automated decision making, the right to opt out of “sharing” consumer’s personal information for cross-context behavioral advertising, and the right to restrict use of and disclosure of sensitive personal information, including geolocation data to third-parties. We will need to evaluate and potentially update our privacy program to ensure compliance with the CPRA and may incur additional costs and expenses in our effort to comply.

Collaborations and contracts of our wholly owned subsidiaries Novavax AB and Novavax CZ, with regional partners, such as SIIPL and Cadila, as well as with international providers, expose us to additional risks associated with doing business outside the U.S.

Swedish-based Novavax AB and Czech Republic-based Novavax CZ are wholly owned subsidiaries of Novavax, Inc. We also have entered into a manufacturing and distribution agreement with SIIPL., formed a joint venture with Cadila in India, and have entered into other agreements and arrangements with foreign governments and companies in other countries. We plan to continue to enter into collaborations or partnerships with companies, non-profit organizations and local governments in various parts of the world. Risks of conducting business outside the U.S. include negative consequences of: • the costs associated with seeking to comply with multiple regulatory requirements that govern our ability to develop, manufacture and sell products in local markets; • failure to comply with anti-bribery laws, such as the U.S. Foreign Corrupt Practices Act and similar anti-bribery laws in other jurisdictions; • new or changes in interpretations of existing trade protections measures, including tariffs, embargoes and import and export licensing requirements; • difficulties in and costs of staffing, managing and operating our international operations; • changes in environmental, health and safety laws; • fluctuations in foreign currency exchange rates; • new or changes in interpretations of existing tax laws; • political instability and actual or anticipated military or potential conflicts;
Risks Related to Our Convertible Senior Notes

If we are unable to attract or retain key management or other personnel, our business, operating results and financial condition could be materially adversely affected.

We depend on our senior executive officers, as well as key scientific and other personnel. The loss of these individuals could harm our business and significantly delay or prevent the achievement of research, development or business objectives. Turnover in key executive positions resulting in lack of management continuity and long-term history with our Company could result in operational and administrative inefficiencies and added costs.

We may not be able to attract qualified individuals for key positions on terms acceptable to us. Competition for qualified employees is intense among pharmaceutical and biotechnology companies, and the loss of qualified employees, or an inability to attract, retain and motivate additional highly skilled employees could hinder our ability to complete clinical trials successfully and otherwise develop marketable products.

We also rely from time to time on outside advisors who assist us in formulating our research and development and clinical strategy. We may not be able to attract and retain these individuals on acceptable terms, which could delay our development efforts.

Risks Related to Our Convertible Senior Notes

Servicing our 3.75% convertible senior unsecured notes due 2023 (the “Notes”) requires a significant amount of cash, and we may not have sufficient cash flow to pay our debt.

In 2016, we issued $325 million aggregate principal amount of Notes. Our ability to make scheduled payments of the principal of, to pay interest on, or to refinance our indebtedness, including the Notes, depends on our future performance, which is subject to economic, financial, competitive and other factors beyond our control. We do not expect our business to be able to generate cash flow from operations sufficient to service our debt and make necessary capital expenditures and other cash requirements. Our future operating performance and the success of our future business plans will depend on many factors, any one of which could result in a default on our debt obligations and limit our flexibility in planning for and reacting to changes in our business.

We may not have the ability to raise the funds necessary to repurchase the Notes as required upon a fundamental change, and our future debt may contain limitations on our ability to repurchase the Notes.

Holders of the Notes will have the right to require us to repurchase their Notes for cash upon the occurrence of a fundamental change at a fundamental change repurchase price equal to 100% of the principal amount of the Notes to be repurchased, plus accrued and unpaid interest, if any. A fundamental change may also constitute an event of default or prepayment under, and result in the acceleration of the maturity of, our then-existing indebtedness. We cannot assure that we will have sufficient financial resources, or will be able to arrange financing, to pay the fundamental change repurchase price in cash with respect to any Notes surrendered by holders for repurchase upon a fundamental change. In addition, restrictions in our then-existing credit facilities or other indebtedness, if any, may not allow us to repurchase the Notes upon a fundamental change. Our failure to repurchase the Notes upon a fundamental change when required would result in an event of default with respect to the Notes which could, in turn, constitute a default under the terms of our other indebtedness.

If any. If the repayment of the related indebtedness were to be accelerated after any applicable notice or grace periods, we may not have sufficient funds to repay the indebtedness and repurchase the Notes.

Capped call transactions entered into in connection with our Notes may affect the value of our common stock.

In connection with our Notes, we entered into capped call transactions (the “capped call transactions”) with certain financial institutions. The capped call transactions are expected to generally reduce the potential dilution upon conversion of the Notes into shares of our common stock.

In connection with establishing their initial hedges of the capped call transactions, these financial institutions or their respective affiliates entered into various derivative transactions with respect to our common stock and/or to purchase our common stock. The financial institutions, or their respective affiliates, may modify their hedge positions by entering into or unwinding various derivatives with respect to our common stock and/or purchasing or selling our common stock or other securities of ours in secondary market transactions prior to the maturity of the Notes. This activity could also cause or avoid an increase or a decrease in the market price of our common stock or the Notes, which could affect the value of our common stock.

Risks Related to Ownership of Our Common Stock

Because our stock price has been and will likely continue to be highly volatile, the market price of our common stock may be lower or more volatile than expected.

Our stock price has been highly volatile. From January 1, 2020 through December 31, 2020, the closing sale price of our common stock has been as low as $6.31 per share and as high as $178.51 per share. The stock market in general and the market for biotechnology companies in particular have experienced extreme volatility that has often been unrelated to the operating performance of particular companies. These broad market fluctuations may cause the market price of our common stock to be lower or more volatile than expected.

Furthermore, given the global focus on the COVID-19 pandemic and our investment in developing a COVID-19 vaccine, information in the public arena on this topic, whether or not accurate, has had and will likely continue to have an outsized impact (positive or negative) on our stock price. Information related to our development, manufacturing, regulatory and commercialization efforts with respect to NVX-CoV2373, or information regarding such efforts by competitors with respect to their COVID-19 vaccines and vaccine candidates, may meaningfully impact our stock price. As a result of this volatility, you may not be able to sell your common stock at or above your initial purchase price. The market price of our common stock may be influenced by many other factors, including:

- future announcements about us or our collaborators or competitors, including the results of testing, technological innovations or new commercial products;
- clinical trial results;
- depletion of our cash reserves;
- sale of equity securities or issuance of additional debt;
- announcement by us of significant strategic partnerships, collaborations, joint ventures, capital commitments or acquisitions;
- changes in government regulations;
- impact of competitor successes and in particular development success of vaccine candidates that compete with our own vaccine candidates;
- developments in our relationships with our collaboration partners;
announcements relating to health care reform and reimbursement levels for new vaccines and other matters affecting our business and results, regardless of accuracy; sales of substantial amounts of our stock by us or existing stockholders (including stock by insiders or 5% stockholders); development, spread or new announcements related to pandemic diseases; litigation; public concern as to the safety of our products; significant setbacks or concerns with the industry or the market as a whole; regulatory inquiries, reviews and potential action, including from the FDA or the SEC; recommendations by securities analysts or changes in earnings estimates; and the other factors described in this Risk Factors section.

In the past, following periods of volatility in the market price of a company’s securities, securities class-action litigation often has been instituted against that company. Such litigation, if instituted against us, could cause us to incur substantial costs to defend such claims and divert management’s attention and resources, which could seriously harm our business, financial condition, and results of operations, and prospects.

Raising additional capital by issuing securities or through collaboration and licensing arrangements may cause dilution to existing stockholders or require us to relinquish rights to our technologies or vaccine candidates.

If we are unable to partner with a third-party to advance the development of one or more of our vaccine candidates, we will need to raise money through additional debt or equity financings. To the extent that we raise additional capital by issuing equity securities, our stockholders will experience immediate dilution, which may be significant. There is also a risk that such equity issuances may cause an ownership change under the Internal Revenue Code of 1986, as amended, and similar state provisions, thus limiting our ability to use our net operating loss carryforwards and credits. To the extent that we raise additional capital through licensing arrangements or arrangements with collaborative partners, we may be required to relinquish, on terms that may not be favorable to us, rights to some of our technologies or vaccine candidates that we would otherwise seek to develop or commercialize ourselves. In addition, economic conditions may also negatively affect the desire or ability of potential collaborators to enter into transactions with us. They may also have to delay or cancel research and development projects or reduce their overall budgets.

Provisions of our Second Amended and Restated Certificate of Incorporation and Amended and Restated By-Laws and Delaware law could delay or prevent the acquisition of the Company, even if such acquisition would be beneficial to stockholders, and could impede changes in our Board.

Provisions in our organizational documents could hamper a third-party’s attempt to acquire or discourage a third-party from attempting to acquire control of the Company. Stockholders who wish to participate in these transactions may not have the opportunity to do so. Our organizational documents also could limit the price investors are willing to pay in the future for our securities and make it more difficult to change the composition of our Board in any one year. For example, our organizational documents provide for a staggered board with three classes of directors serving staggered three-year terms and advance notice requirements for stockholders to nominate directors and make proposals.

As a Delaware corporation, we are also afforded the protections of Section 203 of the Delaware General Corporation Law, which will prevent us from engaging in a business combination with a person who acquires at least 15% of our common stock for a period of three years from the date such person acquired such common stock, unless advance board or stockholder approval was obtained.

Any delay or prevention of a change of control transaction or changes in our Board or management could deter potential acquirers or prevent the completion of a transaction in which our stockholders could receive a substantial premium over the then current market price for their shares.

We have never paid dividends on our capital stock, and we do not anticipate paying any such dividends in the foreseeable future.

We have never paid cash dividends on our common stock. We currently anticipate that we will retain all of our earnings for use in the development of our business and do not anticipate paying any cash dividends in the foreseeable future. As a result, capital appreciation, if any, of our common stock would be the only source of gain for stockholders until dividends are paid, if at all.

General Risk Factors

Litigation could have a material adverse impact on our results of operation and financial condition.

In addition to intellectual property litigation, from time to time, we may be subject to other litigation. Regardless of the merits of any claims that may be brought against us, litigation could result in a diversion of management’s attention and resources and we may be required to incur significant expenses defending against these claims. If we are unable to prevail in litigation, we could incur substantial liabilities. Where we can make a reasonable estimate of the liability relating to pending litigation and determine that it is probable, we record a related liability. As additional information becomes available, we assess the potential liability and revise estimates as appropriate. However, because of uncertainties relating to litigation, the amount of our estimates could be wrong.

We or the third-parties upon whom we depend may be adversely affected by natural or man-made disasters or public health emergencies, such as the COVID-19 pandemic.

Our operations, and those of our clinical research organizations, contract manufacturing organizations, vendors of materials needed in manufacturing, collaboration partners, distributors and other third-parties upon whom we depend, could be subject to fires, extreme weather conditions, earthquakes, power shortages, telecommunications failures, water shortages, floods, hurricanes, typhoons, war, political unrest, sabotage or terrorism and other natural or man-made disasters, as well as public health emergencies, such as the COVID-19 pandemic. The occurrence of any of these business disruptions could prevent us from using all or a significant portion of our facilities, and it may be difficult or impossible for us to continue certain activities for a substantial period of time. The disaster recovery and business continuity plans we have in place may prove inadequate in the event of a serious disaster or similar event, and we may incur substantial expenses and delays as a result. Our ability to manufacture our product candidates and obtain necessary clinical supplies for our product candidates could be disrupted if the operations of our contract manufacturing organizations or suppliers are affected by a natural or man-made disaster, or a public health emergency.

The outbreak of COVID-19 may materially and adversely affect our business and our financial results.

The COVID-19 pandemic continues to present substantial global economic and public health challenges, which may materially and adversely impact our business, financial condition and results of operations. In response to COVID-19, various aspects of our business operations have been, and could continue to be, disrupted. We continue to implement a work from home policy, with our administrative employees working outside of our offices, and on-site staff restricted to only those required to execute certain laboratory and related support activities. Working remotely could increase our cybersecurity risk, create data accessibility concerns, and make us more susceptible to communication disruptions, any of which could adversely impact our business operations. In addition, as a result of state or local restrictions, our on-site staff conducting research and development may not be able to access our laboratories, and these core activities may be significantly limited or curtailed, possibly for extended periods of time. Travel restrictions and other governmental measures may also result in a disruption or delay in the performance of our third-party contractors and suppliers. If such third-parties
are unable to adequately satisfy their contractual commitments to us in a timely manner, our business could be adversely affected.

Our clinical trials, whether planned or ongoing, may be affected by the COVID-19 pandemic. Study procedures (particularly any procedures that may be deemed non-essential), site initiation, participant recruitment and enrollment, participant dosing, shipment of our product candidates, distribution of clinical trial materials, study monitoring, site inspections and data analysis may be paused or delayed due to changes in hospital or research institution policies, federal, state or local regulations, prioritization of hospital and other medical resources toward efforts to treat or prevent COVID-19, or other reasons related to the pandemic. In addition, there could be a potential effect of COVID-19 to the operations of the FDA or other health authorities, which could result in delays of reviews and approvals, including with respect to our product candidates. Any prolongation or de-prioritization of our clinical trials or delay in regulatory review resulting from such disruptions could materially affect the development and study of our product candidates.

The trading prices for our common stock and that of other biopharmaceutical companies have been highly volatile due to the COVID-19 pandemic, especially as a result of investor concerns and uncertainty related to the impact of the outbreak on the economies of countries worldwide. These broad market and industry fluctuations, as well as general economic, political and market conditions, may negatively impact the market price of shares of our common stock.

The COVID-19 pandemic continues to rapidly evolve. The extent to which the outbreak impacts our business, preclinical studies and clinical trials will depend on future developments, which are highly uncertain and cannot be predicted with confidence, such as the ultimate geographic spread of the disease, the duration of the pandemic, travel restrictions and social distancing in the U.S. and other countries, business closures or business disruptions and the effectiveness of actions taken in the U.S. and other countries to contain and treat the disease.

The United Kingdom’s withdrawal from the European Union could result in increased regulatory and legal complexity, which may make it more difficult for us to do business in the UK and/or Europe and impose additional challenges in securing regulatory approval of our product candidates in the UK and/or Europe.

The United Kingdom’s exit from the European Union as of January 31, 2020, with a transitional period up to December 31, 2020, commonly referred to as “Brexit”, has caused political and economic uncertainty, including in the regulatory framework applicable to our operations and vaccine candidates in the United Kingdom and the European Union, and this uncertainty may persist for years. Brexit could, among other outcomes, disrupt the free movement of goods, services and people between the United Kingdom and the European Union, and result in increased legal and regulatory complexities, as well as potential higher costs of conducting business in Europe. As one of the Brexit consequences, the EMA has relocated from the United Kingdom to the Netherlands. This has led to a significant reduction of the EMA workforce, which has resulted and could further result in significant disruption and delays in its administrative procedures, such as granting clinical trial authorization or opinions for marketing authorization, disruption of importation and export of active substance and other components of new drug formulations, and disruption of the supply chain for clinical trial product and final authorized formulations. As any European Union marketing authorization for NVX-CoV2373 would be issued after January 1, 2021, if at all, it would not be grandfathered in the UK. We therefore must seek to obtain a separate marketing authorization for the UK, increasing our regulatory burden.

The cumulative effects of the disruption to the regulatory framework may add considerably to the development lead time to marketing authorization and commercialization of products in the European Union and/or the United Kingdom. It is possible that there will be increased regulatory complexities, which can disrupt the timing of our clinical trials and regulatory approvals. In addition, changes in, and legal uncertainty with regard to, national and international laws and regulations may present difficulties for our clinical and regulatory strategy. Any delay in obtaining, or an inability to obtain, any marketing approvals, as a result of Brexit or otherwise, would prevent us from commercializing our product candidates in the United Kingdom and/or the European Union and restrict our ability to generate revenues and achieve and sustain profitability.

In addition, as a result of Brexit, other European countries may seek to conduct referenda with respect to their continuing membership with the European Union. Given these possibilities and others we may not anticipate, as well as the absence of comparable precedent, it is unclear what financial, regulatory and legal implications the withdrawal of the United Kingdom from the European Union will have, how such withdrawal will affect us, and the full extent to which our business could be adversely affected.

We are increasingly a target for public scrutiny, and our business may be impacted by unfavorable publicity.

Given that COVID-19 represents an unprecedented urgent public health crisis, that we are developing NVX-CoV2373 as a COVID-19 vaccine candidate, and that we have received significant funding from the U.S. and foreign governments and other sources to support the development and potential commercialization of NVX-CoV2373, we have observed and are likely to continue to face significant public attention and scrutiny over the complex decisions we have made and will be making regarding the development, testing, manufacturing, allocation and pricing of NVX-CoV2373. If we are unable to successfully manage these risks, we could face significant reputational harm, which could negatively affect our stock price. The intense public interest, including speculation by the media, in the development of NVX-CoV2373 has caused significant volatility in our stock price, which we expect to continue as data and other information from our ongoing clinical trials become publicly available. If concerns should arise about the actual or anticipated efficacy or safety of any of our product candidates, such concerns could adversely affect the market’s perception of these candidates, which could lead to a decline in investors’ expectations and a decline in the price of our common stock.

The increasing use of social media platforms presents new risks and challenges to our business.

Social media is increasingly being used to communicate about pharmaceutical companies’ research, product candidates, and the diseases such product candidates are being developed to prevent. Social media practices in the pharmaceutical industry continue to evolve and regulations relating to such use are not always clear. This evolution creates uncertainty and risk of noncompliance with regulations applicable to our business, resulting in potential regulatory actions against us. For example, subjects may use social media channels to comment on their experience in an ongoing blinded clinical trial or to report an alleged adverse event. When such events occur, there is a risk that we fail to monitor and comply with applicable adverse event reporting obligations or we may not be able to defend our business or the public’s legitimate interests in the face of the political and market pressures generated by social media due to restrictions on what we may say about our investigational product candidates. There is also a risk of inappropriate disclosure of sensitive information or negative or inaccurate posts or comments about us on any social media or networking website. If any of these events were to occur or we otherwise fail to comply with applicable regulations, we could incur liability, face regulatory actions, or incur reputational or other harm to our business.
Item 1B. UNRESOLVED STAFF COMMENTS

None.

Item 2. PROPERTIES

We lease three facilities in Gaithersburg, Maryland, Novavax AB leases a facility in Uppsala, Sweden and Novavax CZ owns a facility in Bohumil, Czech Republic. A summary of our current facilities is set forth below. Although we believe that our facilities are suitable and adequate for our present needs, the Company’s management continues to review and assess real property requirements that may be necessary to address our current business plan.

<table>
<thead>
<tr>
<th>Property Location</th>
<th>Approximate Square Footage</th>
<th>Brief Property Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>700QO Gaithersburg, MD</td>
<td>170,000</td>
<td>Manufacturing, research and development facility and offices</td>
</tr>
<tr>
<td>21FF Gaithersburg, MD</td>
<td>53,000</td>
<td>Research and development facility and offices</td>
</tr>
<tr>
<td>22FF Gaithersburg, MD</td>
<td>40,000</td>
<td>Executive, administrative, clinical and regulatory offices</td>
</tr>
<tr>
<td>Uppsala, Sweden</td>
<td>37,300</td>
<td>Adjuvant manufacturing and research and development facility and offices</td>
</tr>
<tr>
<td>Bohumil, Czech Republic</td>
<td>138,400</td>
<td>Manufacturing facility and offices</td>
</tr>
<tr>
<td>Total square footage</td>
<td>438,700</td>
<td></td>
</tr>
</tbody>
</table>

Item 3. LEGAL PROCEEDINGS

We currently have no material pending legal proceedings.

Item 4. MINE SAFETY DISCLOSURES

Not applicable.

PART II

Item 5. MARKET FOR REGISTRANT’S COMMON EQUITY AND RELATED STOCKHOLDER MATTERS

Our common stock trades on the Nasdaq Global Select Market under the symbol “NVAX.” Our common stock was held by approximately 129 stockholders of record as of February 24, 2021, one of which is Cede & Co., a nominee for Depository Trust Company (“DTC”). All of the shares of common stock held by brokerage firms, banks and other financial institutions as nominees for beneficial owners are deposited into participant accounts at DTC, and are therefore considered to be held of record by Cede & Co. as one stockholder. We do not anticipate declaring or paying any cash dividends in the foreseeable future.

Securities Authorized for Issuance under our Equity Compensation Plans

Information regarding our equity compensation plans, including both stockholder approved plans and non-stockholder approved plans, is included in Item 12 of this Annual Report on Form 10-K.

Performance Graph

The graph below compares the cumulative total stockholders return on our common stock for the last five fiscal years with the cumulative total return on the Nasdaq Composite Index and the Russell 2000 Growth Biotechnology Index (which includes Novavax) over the same period, assuming the investment of $100 in our common stock, the Nasdaq Composite Index and the Russell 2000 Growth Biotechnology Index on December 31, 2015, and reinvestments of all dividends.

* $100 invested on 12/21/15 in stock or index, including reinvestment of dividends. Fiscal year ending December 31.
Value of $100 invested on December 31, 2015 in stock or index, including reinvestment of dividends, for fiscal years ended December 31:

<table>
<thead>
<tr>
<th></th>
<th>12/31/15</th>
<th>12/31/16</th>
<th>12/31/17</th>
<th>12/31/18</th>
<th>12/31/19</th>
<th>12/31/20</th>
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<tr>
<td>Novavax, Inc.</td>
<td>$100</td>
<td>$15.02</td>
<td>$14.78</td>
<td>$21.93</td>
<td>$2.37</td>
<td>$66.45</td>
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<tr>
<td>NASDAQ Composite</td>
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<td>$108.87</td>
<td>$141.13</td>
<td>$137.12</td>
<td>$187.44</td>
<td>$271.64</td>
</tr>
<tr>
<td>Russell 2000 Growth Biotechnology</td>
<td>$100</td>
<td>$79.71</td>
<td>$127.4</td>
<td>$105.07</td>
<td>$153.35</td>
<td>$238.36</td>
</tr>
</tbody>
</table>

This graph is not “soliciting material,” is not deemed “filed” with the SEC and is not to be incorporated by reference in any filing of the Company under the Securities Act of 1933, as amended, or the Securities Exchange Act of 1934, as amended, whether made before or after the date hereof and irrespective of any general incorporation language in any such filing.

Item 6. SELECTED FINANCIAL DATA

The following table sets forth selected financial data for each of the years in the five-year period ended December 31, 2020, which have been derived from our audited consolidated financial statements. The information below should be read in conjunction with "Results of Operations" included elsewhere in this Annual Report. These historical results are not necessarily indicative of results for future periods.

<table>
<thead>
<tr>
<th></th>
<th>Year Ended December 31</th>
<th>2020(1)</th>
<th>2019(2)</th>
<th>2018(3)</th>
<th>2017(4)</th>
<th>2016</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(in thousands, except per share amounts)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Revenue</strong></td>
<td></td>
<td>$475,598</td>
<td>$18,662</td>
<td>$34,288</td>
<td>$31,176</td>
<td>$15,353</td>
</tr>
<tr>
<td><strong>Net loss</strong></td>
<td></td>
<td>(418,259)</td>
<td>(132,694)</td>
<td>(184,748)</td>
<td>(183,769)</td>
<td>(279,966)</td>
</tr>
<tr>
<td><strong>Basic and diluted net loss per share</strong></td>
<td>(7.27)</td>
<td>(5.51)</td>
<td>(9.99)</td>
<td>(12.56)</td>
<td>(20.68)</td>
<td></td>
</tr>
<tr>
<td>Weighted average shares used in computing basic and diluted net loss per share</td>
<td>57,554</td>
<td>24,100</td>
<td>18,488</td>
<td>14,633</td>
<td>13,540</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th>As of December 31</th>
<th>2020(1)</th>
<th>2019(2)</th>
<th>2018(3)</th>
<th>2017(4)</th>
<th>2016</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Balance Sheet Data:</strong></td>
<td>(in thousands)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cash and cash equivalents, marketable securities and restricted cash</td>
<td>$806,387</td>
<td>$82,180</td>
<td>$103,939</td>
<td>$186,427</td>
<td>$270,383</td>
<td></td>
</tr>
<tr>
<td>Total current assets</td>
<td>1,248,203</td>
<td>97,247</td>
<td>119,276</td>
<td>203,311</td>
<td>287,830</td>
<td></td>
</tr>
<tr>
<td>Working capital(6)</td>
<td>668,531</td>
<td>71,452</td>
<td>73,737</td>
<td>129,636</td>
<td>221,424</td>
<td></td>
</tr>
<tr>
<td>Total assets(7)</td>
<td>1,582,479</td>
<td>172,957</td>
<td>207,978</td>
<td>302,493</td>
<td>394,301</td>
<td></td>
</tr>
<tr>
<td>Long-term debt</td>
<td>322,035</td>
<td>320,611</td>
<td>319,178</td>
<td>317,763</td>
<td>316,339</td>
<td></td>
</tr>
<tr>
<td>Accumulated deficit</td>
<td>(1,874,199)</td>
<td>(1,431,801)</td>
<td>(1,299,107)</td>
<td>(1,114,259)</td>
<td>(929,996)</td>
<td></td>
</tr>
<tr>
<td>Total stockholders’ (deficit) equity</td>
<td>627,209</td>
<td>(186,017)</td>
<td>(167,935)</td>
<td>(101,732)</td>
<td>(5,546)</td>
<td></td>
</tr>
</tbody>
</table>

(1) In 2020, we had sales of 32.4 million shares of common stock and we sold 0.4 million shares of preferred stock, which were converted to 4.4 million shares of common stock in the fourth quarter 2020, resulting in total net proceeds of approximately $1.1 billion.

(2) In 2019, we had sales of 13.0 million shares of common stock, resulting in net proceeds of approximately $98 million.

(3) In 2018, we had sales of 2.9 million shares of common stock, resulting in net proceeds of approximately $100 million.

(4) In 2017, we had sales of 2.5 million shares of common stock, resulting in net proceeds of approximately $63 million.

(5) All share and per share amounts have been retroactively restated for all periods presented to reflect the Reverse Stock Split (see Note 12 to the accompanying consolidated financial statements).

(6) Working capital is computed as the excess of current assets over current liabilities.

(7) In 2019, the Company adopted ASU 2016-02, Leases (Topic 842), under which the Company recorded right-of-use assets associated with its leases on the consolidated balance sheet (see Note 7 to the accompanying consolidated financial statements).

Item 7. MANAGEMENT’S DISCUSSION AND ANALYSIS OF FINANCIAL CONDITION AND RESULTS OF OPERATIONS

Any statements in the discussion below and elsewhere in this Annual Report about expectations, beliefs, plans, objectives, assumptions or future events or performance of Novavax, Inc. ("Novavax," together with its wholly owned subsidiaries Novavax AB and Novavax CZ, the "Company," "we" or "us") are not historical facts and are forward-looking statements. Such forward-looking statements include, without limitation, statements about our capabilities, goals, expectations regarding future revenue and expense levels and capital raising activities; potential market sizes and demand for our product candidates; the efficacy, safety and intended utilization of our product candidates; the development of our clinical-stage product candidates and our recombination vaccine and adjuvant technologies; the development of our preclinical product candidates; the conduct, timing and potential results from clinical trials and other preclinical studies; plans for and potential timing of regulatory filings; our expectation of manufacturing capacity, timing, production and delivery for NVX-CoV2373; our expectations with respect to the anticipated ongoing development and potential commercialization or licensure of NVX-CoV2373 and NanoFlu™; the expected timing and content of regulatory actions; funding from the U.S. government partnership formerly known as Operation Warp Speed ("OWS"), the U.S. Department of Defense ("DoD") and the Coalition for Epidemic Preparedness Innovations ("CEPI"); and payments from the Bill & Melinda Gates Foundation ("BMGF"); our available cash resources and usage and the availability of financing generally; plans regarding partnering activities, business development initiatives; and other matters referenced herein. Generally, forward-looking statements can be identified through the use of words or phrases such as "believe," "may," "could," "will," "would," "possible," "can," "estimate," "continue," "ongoing," "consider," "anticipate," "intend," "seek," "plan," "project," "expect," "should," "would," or "assume," the negative of these terms or other comparable terminology, although not all forward-looking statements contain these words.

Forward-looking statements involve estimates, assumptions and uncertainties that could cause actual results to differ materially from those expressed or implied in the statements. Any or all of our forward-looking statements in this Annual Report may turn out to be inaccurate or materially different from actual results.

Because the risk factors discussed in this Annual Report and other risk factors of which we are not aware could cause actual results or outcomes to differ materially from those expressed or implied in any forward-looking statements made by or on behalf of us, you should not place undue reliance on any such forward-looking statements. These statements are subject to risks and uncertainties, known and unknown, which could cause actual results and developments to differ materially from those expressed or implied in such statements. We have included important factors that could cause results to differ in the customary statements included in this Annual Report, particularly those identified in Part I, Item 1A “Risk Factors” of this Annual Report. These and other risks may also be detailed and modified or updated in our reports and other documents filed with the Securities and Exchange Commission ("SEC") from time to time. You are encouraged to read these filings as they are made.

We cannot guarantee future results, events, level of activity, performance or achievement. Further, any forward-looking statement speaks only as of the date on which it is made, and we undertake no obligation to update or revise any forward-looking statements, whether as a result of new information, future events or otherwise, unless required by law. New factors emerge from time to time, and it is not possible for us to predict which factors will arise. In addition, we cannot assess the impact of each factor on our business or the extent to which any factor, or combination of factors, may cause actual results to differ materially from those contained in any forward-looking statements.
Overview

We are a biotechnology company promoting improved global health through the discovery, development and commercialization of innovative vaccines to prevent serious infectious diseases and address urgent, global health needs. Our vaccine candidates, including both our coronavirus vaccine candidate, NVX-CoV2373, and our influenza vaccine candidate, NanoFlu, are genetically engineered, three-dimensional nanostructures of recombinant proteins. We believe that our protein-subunit-based candidates elicit differentiated immune responses that may be more efficacious than naturally occurring immunity or other, more traditional vaccine approaches. Our technology may be used to target a variety of infectious diseases. Our unique technology is paired with our proprietary immune-stimulating adjuvants, developed at Novavax AB, our wholly owned Swedish subsidiary. To date, we have formulated many of the vaccine candidates in our pipeline with our lead adjuvant, Matrix-M™, including NVX-CoV2373 and NanoFlu. Matrix-M has been shown to enhance functional immune responses and has been well-tolerated in multiple clinical trials. Matrix-M also enables dose-sparing properties.

As the world continues to address the global COVID-19 pandemic, we remain focused today on bringing our NVX-CoV2373 vaccine candidate to market. We have begun rolling reviews with five regulatory authorities worldwide and have initiated submissions to the U.S. Food and Drug Administration (“FDA”) for our open investigational new drug application. In addition, NanoFlu continues to be a priority for our team, and we are exploring the potential for a combined NanoFlu/NVX-CoV2373 vaccine. Our dedicated NanoFlu team continues to seek approval from the FDA under an accelerated approval pathway. Although NVX-CoV2373 and NanoFlu are our near-term priorities, we remain optimistic that the additional programs in our pipeline, including our vaccine candidates for respiratory syncytial virus (“RSV”) and other emerging infectious diseases, present viable opportunities for future development.

Near-term Clinical Development Focus

Our development pipeline encompasses vaccine candidates addressing therapeutic areas including coronavirus, seasonal influenza, RSV and other emerging infectious diseases. At the forefront of our pipeline, we have evaluated our COVID-19 vaccine candidate, NVX-CoV2373, in various preclinical and clinical trials, including two Phase 3 trials, one Phase 2b trial, and one Phase 1/2 trial. Through our clinical development program, we have demonstrated the safety and efficacy of NVX-CoV2373. Additionally, in February 2021, we selected candidates for COVID-19 variant strain vaccines as standalone and bivalent candidates. We plan to initiate clinical testing of these new variant vaccine candidates in mid-2021. Outside of our COVID-19 vaccine candidate, we have advanced our NanoFlu program through a Phase 3 clinical trial, demonstrating positive top-line results and achieving statistical significance across secondary endpoints. We continue to evaluate the viability of certain combination vaccines, including combinations of our NanoFlu, NVX-CoV2373 and respiratory syncytial virus fusion (F) protein nanoparticle vaccine candidate (“RSV F Vaccine”).

A summary and status of our clinical and preclinical development program follows:

<table>
<thead>
<tr>
<th>Therapeutic Area</th>
<th>Name</th>
<th>Preclinical</th>
<th>Phase 1</th>
<th>Phase 2</th>
<th>Phase 3</th>
<th>Marketed</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coronavirus</td>
<td>NVX-CoV2373</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Variant Strain</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>(Booster and/or Bivalent)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Seasonal Flu</td>
<td>NanoFlu (Older Adults) (Pre-BlA)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Combination Vaccines</td>
<td>NanoFlu / NVX-CoV2373</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>NanoFlu / RSV</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>NanoFlu / NVX-CoV2372 / RSV</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

1. Supported by funding from the OWS, DoD, CEPI and BMGF
2. Ongoing PREVENT-19, a Phase 3 clinical trial in U.S. and Mexico; Ongoing Phase 3 in UK; Ongoing Phase 2b in South Africa

COVID-19 Vaccine Funding

Funding for our NVX-CoV2373 clinical development program encompasses over $2 billion from sources including CEPI, the DoD, and OWS.

A summary and status of our key COVID-19 funding developments follows:

<table>
<thead>
<tr>
<th>Funding Partner</th>
<th>Amount</th>
<th>Additional Detail</th>
</tr>
</thead>
</table>
| CEPI               | $399.5 million | • Entitled to receive up to $399.5 million of funding to support the development of NVX-CoV2373  
|                    |              | • To supply NVX-CoV2373 through the COVAX Facility                                 |
| DoD                | $45.7 million | • Entitled to receive up to $45.7 million of funding to support the development of NVX-CoV2373  
|                    |              | • To manufacture and deliver 10 million doses of NVX-CoV2373 to the U.S. government |
| U.S. Government (OWS)| $1.75 billion | • Allotted funding of $1.6 billion and are entitled to receive maximum funding up to $1.75 billion to support the development of NVX-CoV2373  
|                    |              | • To manufacture and deliver 100 million doses of NVX-CoV2373 to the U.S. government |
NVX-CoV2373 Manufacturing and Supply

With respect to the global manufacturing and supply of NVX-CoV2373, we have secured manufacturing for our antigen component and Matrix-M adjuvant, as well as secured fill/finish activities for NVX-CoV2373 at several sites globally. Through our various manufacturing partnerships, we expect our projected global manufacturing production rate of NVX-CoV2373 to be over two billion doses annually when we are at full capacity, which we expect to occur in mid-2021. Of this anticipated capacity, approximately one billion doses will be manufactured by Serum Institute of India Private Limited (“SIIPL”).

NVX-CoV2373 and its components are being manufactured at the following Novavax (in bold) and partnered sites:

### Antigen Component of NVX-CoV2373
- Novavax CZ
- SanoBiologics in Spain
- FUJIFILM Diosynth Biotechnologies (“FDB”) in North Carolina and Texas in the U.S.
- FDB in the UK
- National Research Council’s Biologics Manufacturing Centre in Canada

### Matrix-M Adjuvant
- Novavax AB
- AGC Biologics in the U.S. and Denmark
- PolyPeptide Group (will manufacture two key components used in Matrix-M) in the U.S. and Sweden

### Fill/Finish Activities
- Baxter International Inc. in Germany
- Janssen HollisterStier LLC in the U.S.
- Par Pharmaceutical Companies, Inc. in the U.S.
- Siegfried AG in Germany

### Antigen Production, Out-licensing & Collaborations
- SIIPL in India
- SK bioscience in the Republic of Korea
- Takeda in Japan

### Additional Details

A summary and status of key manufacturing and supply developments follows:

<table>
<thead>
<tr>
<th>Partner</th>
<th>Nature of Agreement</th>
<th>Additional Detail</th>
</tr>
</thead>
<tbody>
<tr>
<td>Canada</td>
<td>Memorandum of Understanding</td>
<td>To produce NVX-CoV2373 at the National Research Council’s Biologics Manufacturing Centre</td>
</tr>
<tr>
<td>SK bioscience</td>
<td>License and Collaboration Agreement</td>
<td>SK bioscience to manufacture antigen component of NVX-CoV2373 for use in final drug product globally</td>
</tr>
<tr>
<td>SIIPL</td>
<td>Exclusive License Agreement</td>
<td>Takeda to have manufacturing capacity of over 250 million doses of NVX-CoV2373 per year</td>
</tr>
<tr>
<td>FDB</td>
<td>Manufacturing Partnership Agreement</td>
<td>Enables large-scale production of NVX-CoV2373 at multiple FDB facilities</td>
</tr>
<tr>
<td>Takeda</td>
<td>Supply and License Agreement</td>
<td>Granted SIIPL non-exclusive license to manufacture the antigen component of NVX-CoV2373</td>
</tr>
<tr>
<td>AGC Biologics</td>
<td>Contract Manufacturing Arrangement</td>
<td>AGC Biologics to provide contract development and manufacturing services, supplying us large-scale production of Matrix-M</td>
</tr>
<tr>
<td>Polypeptide Group</td>
<td>Contract Manufacturing Arrangement</td>
<td>Polypeptide Group to provide contract development and manufacturing services, supplying us large-scale production of two key components used in Matrix-M</td>
</tr>
<tr>
<td>Praha Vaccines</td>
<td>Acquisition</td>
<td>Acquired Praha Vaccines in all cash transaction of approximately $167 million</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Acquisition includes a biologics manufacturing facility to provide annual capacity of over 1 billion doses of antigen starting in 2021</td>
</tr>
</tbody>
</table>
NVX-CoV2373 Supply Agreements

Through the date of filing this Form 10-K, we have entered into several supply agreements with various countries globally that, if our COVID-19 product candidate is approved, are expected to result in the delivery of approximately 200 million doses of NVX-CoV2373, throughout 2021 and into the first half of 2022. In addition to these supply agreements, we have committed 110 million doses of NVX-CoV2373 to the U.S. government in relation to the funding received from OWS and the DoD.

A summary of our current supply agreements follows:

<table>
<thead>
<tr>
<th>Country</th>
<th>Committed Doses &amp; Additional Detail</th>
</tr>
</thead>
<tbody>
<tr>
<td>UK</td>
<td>60 million Option to purchase additional orders from time to time</td>
</tr>
<tr>
<td>Canada</td>
<td>52 million Option to purchase up to an additional 24 million doses</td>
</tr>
<tr>
<td>Australia</td>
<td>51 million Option to purchase up to an additional 10 million doses</td>
</tr>
<tr>
<td>New Zealand</td>
<td>~11 million</td>
</tr>
<tr>
<td>Switzerland</td>
<td>6 million</td>
</tr>
<tr>
<td><strong>Summary</strong></td>
<td>~180 Million Doses</td>
</tr>
</tbody>
</table>

Sale of Assets

In July 2019, we closed a transaction under an asset purchase agreement (the “Purchase Agreement”) with Catalent, pursuant to which we sold to Catalent certain assets related to our biomanufacturing and development activities located at the facilities situated at each of 20 Firstfield Road in Gaithersburg, MD 20878 and 9920 Belward Campus Drive in Rockville, MD 20850 for approximately $18 million and recorded a gain on the disposition of such assets of $9.0 million. Pursuant to the transactions contemplated by the Purchase Agreement, approximately 100 Novavax manufacturing and quality employees transferred to Catalent, and we assigned two facility leases to Catalent. We also entered into other ancillary agreements upon the closing of the transaction, including a Non-Commercial GMP Manufacturing Services Agreement pursuant to which we were required to purchase $6.0 million in certain services from Catalent set forth therein, through July 31, 2020. The transaction was treated as an asset disposition for accounting purposes.

Sale of Preferred Stock

In June 2020, we entered into a redeemable Series A Convertible Preferred Stock Subscription Agreement, pursuant to which we sold and issued in a private placement 438,885 shares of our newly designated redeemable Series A Convertible Preferred Stock, par value $0.01 per share (“Preferred Stock”), at a purchase price of $455.70 per share, for total gross proceeds of $200.0 million. During the fourth quarter of 2020, all outstanding shares of the Preferred Stock was converted, and we issued 4,388,850 shares of common stock, par value $0.01 per share and reclassified $199.8 million from preferred stock to additional paid in capital. We recognized a beneficial conversion feature of approximately $24.1 million at the time of issuance of the Preferred Stock that was recorded in additional paid-in capital and accumulated deficit as the Preferred Stock issuance was contingently redeemable and convertible at any time at the option of the holder.

Sales of Common Stock

In January 2021, we entered into an At Market Issuance Sales Agreement (“January 2021 Sales Agreement”), which allows us to issue and sell up to $500 million in gross proceeds of our common stock. From January 22 through February 24, 2021, we sold $1.7 million shares of common stock under the January 2021 Sales Agreement, resulting in $452.0 million in net proceeds, leaving $42.2 million remaining.

During 2020, we entered into various At Market Issuance Sales Agreements, which allowed us to issue and sell up to $1.0 billion in gross proceeds of our common stock. During 2020, we sold a total of 25.2 million shares of common stock under such Sales Agreements, resulting in $835.6 million in net proceeds (this amount excludes $3.2 million received in the first quarter of 2021 for shares traded in late December 2020). From January 1, 2021 through January 20, 2021, we sold $0.9 million shares of common stock from our At Market Issuance Sales Agreement entered into in November 2020 (“November 2020 Sales Agreement”), resulting in $113.0 million in net proceeds, leaving $27.2 million remaining under its November 2020 Sales Agreement. We terminated the November 2020 Sales Agreement by mutual agreement upon entering into the January 2021 Sales Agreement.

Critical Accounting Policies and Use of Estimates

The discussion and analysis of our financial condition and results of operations are based upon our consolidated financial statements, which have been prepared in accordance with generally accepted accounting principles in the United States.

The preparation of our consolidated financial statements requires us to make estimates, assumptions and judgments that affect the reported amounts of assets, liabilities and equity and disclosure of contingent assets and liabilities as of the date of the financial statements and the reported amounts of revenue and expenses during the reporting period. These estimates, particularly estimates relating to accounting for revenue, lease accounting and accounting for research and development expenses, have a material impact on our consolidated financial statements and are discussed in detail throughout our analysis of the results of operations discussed below.
We base our estimates on historical experience and various other assumptions that we believe are reasonable under the circumstances, the results of which form the basis for making judgments about the carrying value of assets, liabilities and equity that are not readily apparent from other sources. Actual results and outcomes could differ from these estimates and assumptions.

Revenue Recognition

We perform research and development under government funding, grant, license and clinical development agreements. Our revenue primarily consists of funding under U.S. government contracts and other arrangements to advance the clinical development and manufacturing of NVX-CoV2373. Our U.S. government contracts include the DoD Contract and the OWS Agreement. Other funding arrangements primarily include a grant and forgivable loan funding from CEPI.

At contract inception, we analyze our revenue arrangements to determine the appropriate accounting under U.S. GAAP. Currently, our revenue arrangements represent customer contracts within the scope ofASC Topic 606, Revenue from Contracts with Customers (Topic 606) (“ASC 606”) or are subject to the contribution guidance in Accounting Standards Codification (ASC) Topic 958-605, Not-for-Profit Entities – Revenue Recognition (“ASC 958-605”), which applies to business entities that receive contributions within the scope of ASC 958-605. We recognize revenue from arrangements within the scope of ASC 606 following the five-step model: (i) identify the contract(s) with a customer; (ii) identify the performance obligation(s) in the contract; (iii) determine the transaction price; (iv) allocate the transaction price to the performance obligations in the contract; and (v) recognize revenue when (or as) we satisfy a performance obligation. We only apply the five-step model to contracts when it is probable that we will collect the consideration we are entitled to in exchange for the goods or services we transfer to our customer. We recognize contribution revenue within the scope of ASC 958-605 when the funder-imposed conditions have been substantially met. Contributions are recorded as deferred revenue until the period in which research and development activities are performed that satisfy the funder-imposed conditions.

Under our U.S. government contracts, we are entitled to receive funding, on a reimbursable-cost or reimbursable-cost-plus fixed fee basis to support certain activities related to the development, manufacture and delivery of NVX-CoV2373 to the U.S. government. We analyzed these contracts and determined that they are within the scope of ASC 606. Our obligations under each of the contracts are not distinct in the context of the contract as they are highly interdependent or interrelated and, as such, are accounted for as a single performance obligation. The transaction price under these arrangements is the consideration we expect to receive and consists of the funded contract amount and the unfunded variable amount to the extent that it is probable that a significant reversal of revenue will not occur. We recognize revenue for these contracts over time as we transfer control over the goods and services and satisfy our performance obligation. We measure progress toward satisfaction of our performance obligation using an Estimate-at-Completion (“EAC”) process, which is a cost-based input method that reviews and monitors the progress toward the completion of our performance obligation. Under this process, we update the costs that we consider as incurred to-date, as well as projections to completion using various inputs and assumptions, including, but not limited to, progress toward completion, labor costs and level of effort, material and subcontractor costs, indirect administrative costs, and other identified risks. Estimating the total allowable cost at completion of our performance obligation under a contract is subjective and requires us to make a number of assumptions and judgments, and if significant, may impact the timing of revenue and fee recognition on our contracts. Allowable contract costs include direct costs incurred on the contract and indirect costs that are applied in the form of rates to the direct costs. Progress billings under the contracts are initially based on provisional indirect billing rates, agreed upon between us and the U.S. government. These indirect rates are subject to audit on an annual basis. The impact of changes in the indirect billing rates are recorded in the period when such changes are identified and reflect the differences between the estimated amounts used to determine the provisional indirect billing rates agreed upon with the U.S. government. We recognize revenue on our U.S. government contracts based on reimbursable allowable contract costs incurred in the period up to the transaction price. For our reimbursable-cost-plus fixed fee contracts, we recognize the fixed fee based on the proportion of reimbursable contract costs incurred to total estimated allowable contract costs expected to be incurred on completion of the underlying performance obligation as determined under the EAC process. Changes in estimates related to the EAC process are recognized in the period when such changes are made on a cumulative catch-up basis. We include the transaction price comprising both funded and unfunded portions of customer contracts, in this estimate. We have not experienced any material difference as a result of change in estimate arising from the EAC process.

Our other funding arrangements primarily include the CEPI Grant Funding and CEPI Forgivable Loan Funding (each as defined in “Note 2—Summary of Significant Accounting Policies” included in our Notes to Consolidated Financial Statements). The CEPI Forgivable Loan Funding is designated for the prepayment of certain manufacturing activities. We analyzed these other funding arrangements and determined that they are not within the scope of ASC 606 as they do not provide a direct economic benefit to the grantor. Payments received under the grant funding arrangements are considered conditional contributions under the scope of ASC 958-605 and are recorded as deferred revenue until the period in which an economic benefit is provided and development activities are actually performed that satisfy the funder-imposed conditions. Payments received under the CEPI Forgivable Loan Funding agreements are only repayable if the proceeds of sales to one or more third-parties of NVX-CoV2373 cover our costs of manufacturing such vaccine candidate, not including manufacturing costs funded by CEPI. As the financial risk remains with CEPI, we have determined that the use of the CEPI Forgivable Loan Funding is outside the scope of ASC Topic 470, Debt. The research and development risk is considered substantive, such that it is not yet probable that the development will be successful. Therefore, we have concluded that ASC Topic 730, Research and Development is considered applicable and most appropriate. Given the financial risk associated with the research and development activities lies with CEPI because repayment of any funds provided by CEPI depends solely on the results of the research and development activities having future economic benefit, we will account for our obligation under the CEPI Forgivable Loan Funding as a contract to perform research and development for others. We have determined that payments received under these agreements should be recorded as revenue under ASC 958-605 rather than a reduction to research and development expenses. This is consistent with our policy of presenting such amounts as revenue. In reaching this determination, we considered a number of factors, including whether we are principal under the arrangement, and whether the arrangement is significant to, and part of, our core operations. We will record revenue as we perform the contractual research and development services.

We have manufacturing and supply arrangements that include a license to use our intellectual property. The licensing arrangements include sales-based royalties, as well as the CEPI Grant Funding and CEPI Forgivable Loan Funding. The CEPI Grant Funding is deemed to be the predominant item to which the milestone payments and sales-based royalties relate. The fulfillment of our obligation for the license is subject to a constraint, the achievement of the development and commercial milestone or the royalty-related sales under the arrangement. For milestone payments, the constraint is overcome, and we recognize revenue when the development and commercial milestone is achieved.

Lease Accounting

We determine at the inception or modification of a contract if an arrangement is, or contains, a lease, which exists when the contract conveys the right to control the use of an underlying asset for a period of time in exchange for consideration. In determining if a contract contains a lease, we evaluate whether the contract, either explicitly or implicitly, is for the use of an identified asset and whether we have the right to control the use of, and obtain substantially all of the benefit from, the identified asset. Depending on the contract, the lease commencement date, defined as the date on which the lessee makes the underlying asset available for use by the lessee and is the date on which the Company is required to accrue lease expenses, may be different than the inception date of the contract. We evaluate changes to the terms and conditions of a lease contract to determine if they result in a new lease or a modification of an existing lease. For lease modifications, we reassess the remaining lease term and reassess the classification and the effective date of the modification. We classify leases as either operating or finance leases based on the economic substance of the arrangement.

We enter into non-cancelable lease agreements for facilities and certain equipment. We also enter into manufacturing supply arrangements with CMOs and CDMOs to manufacture our vaccine candidates. Certain of these manufacturing supply agreements include the use of identified manufacturing facilities and equipment that are controlled by us and for which we obtain appropriate. Given the financial risk associated with the research and development activities lies with CEPI because repayment of any funds provided by CEPI depends solely on the results of the research and development activities having future economic benefit, we will account for our obligation under the CEPI Forgivable Loan Funding as a contract to perform research and development for others. We have determined that payments received under these agreements should be recorded as revenue under ASC 958-605 rather than a reduction to research and development expenses. This is consistent with our policy of presenting such amounts as revenue. In reaching this determination, we considered a number of factors, including whether we are principal under the arrangement, and whether the arrangement is significant to, and part of, our core operations. We will record revenue as we perform the contractual research and development services.

For leases that have a lease term of more than 12 months at the lease commencement date, we recognize lease liabilities, which represent our obligation to make lease payments arising from the lease, and corresponding right-of-use (“ROU”) assets, which represent the right to use an underlying asset for the lease term, based on the present value of the fixed future payments over the lease term. We calculate the present value of future payments using the discount rate implicit in the lease, if available, or our incremental borrowing rate. For all leases that have a lease term of 12 months or less at the commencement date (referred to as “short-term” leases), we have elected to apply the practical expedient in ASC Topic 842. Under (“ASC 842”) to not recognize a lease liability or ROU asset but instead, recognize lease payments as an expense on a straight-line basis over the lease term and variable lease payments that do not depend on an index or rate, as an expense in the period in which the variable lease costs are incurred based on performance or usage in accordance with contractual agreements. In determining the lease period, we evaluate facts and circumstances that could affect the period over which we are reasonably certain to use the underlying asset.
while taking into consideration the non-cancelable period over which we have the right to use the underlying asset and any option period to extend or terminate the lease if we are reasonably certain to exercise the option. We re-evaluate short-term leases that are modified and if they no longer meets the requirements to be treated as short-term leases, we recognize and measure the liability and ROU asset as if the date of the modification is the lease commencement date (see Note 7 to the accompanying consolidated financial statements).

For operating leases, we recognize lease expense related to fixed payments on a straight-line basis over the lease term and lease expense related to variable payments as incurred based on performance or usage in accordance with the contractual agreements. For finance leases, we recognize the amortization of the ROU asset over the shorter of the lease term or useful life of the underlying asset. We expense ROU assets acquired for research and development activities under ASC Topic 730, Research and Development, if they do not have an alternative future use, in research and development projects or otherwise.

We use significant assumptions and judgment in evaluating our lease contracts and other agreements under ASC 842, including the determination of whether an agreement is or contains a lease, whether a change in the terms and conditions of a lease contract represent a new or modified lease, whether a lease represents an operating or finance lease, the discount rate used to determine the present value of lease obligations and the term of embedded leases in our manufacturing supply agreements.

Accounting for Research and Development Expenses

We estimate our prepaid and accrued expenses related to our research and development activities using a process that involves reviewing contracts and purchase orders, communicating with our project managers and service providers to identify services that have been performed on our behalf, and estimating the level of service performed and the associated cost incurred for the service when we have not yet been invoiced or for which we have been invoiced in advance of the service. This estimation process includes a review of:

- expenses incurred under agreements with contract research organizations ("CROs") that conduct our clinical trials and third-party consultants; and
- the cost of developing and manufacturing vaccine components under third-party CMOs and CDMOs agreements, including expenses incurred for the procurement of raw materials, laboratory supplies and equipment.

We base our expenses on our estimates of the services provided and efforts expended pursuant to contracts, statements of work and related change orders with the service provider, as well as discussion with internal personnel and external service providers as to the progress of the services and the agreed-upon fee to be paid for such services. The financial terms of these agreements are based on negotiated terms, vary from contract to contract and may result in an uneven level of activity over time. There may be instances in which payments made to our vendors will exceed the level of services provided and result in a prepayment of the expense. Additionally, invoicing from third-party service providers may not coincide with actual work performed and can result in a prepaid or an accrual position at the end of the period. The estimation process requires us to make significant judgments and estimates in determining the services incurred as of the balance sheet date, which may result in either a prepaid or an accrual balance. As actual costs become known, we adjust our estimates. Although we do not expect our estimates to be materially different from amounts actually incurred, our understanding of the status and timing of services performed may vary from the related estimates and could result in us reporting amounts that are too high or too low in a particular period. Our prepaid and accrued expenses are dependent, in part, upon the receipt of timely and accurate reporting from CROs, CMOs, CDMOs and third-party service providers. Due to the nature of the estimation process, there may be a difference between estimated costs and actual costs incurred. Historically, we have not experienced any material differences in prior periods.

Recent Accounting Pronouncements

See "Note 2—Summary of Significant Accounting Policies" included in our Notes to Consolidated Financial Statements (under the caption "Recent Accounting Pronouncements").

Results of Operations for Fiscal Years 2020 and 2019

The following is a discussion of our historical consolidated financial condition and results of operations and should be read in conjunction with the consolidated financial statements and notes thereto set forth in this Annual Report. Additional

information concerning factors that could cause actual results to differ materially from those in our forward-looking statements is described under Part I, Item 1A, “Risk Factors” of this Annual Report.

For our discussion of the year ended December 31, 2019, compared to the year ended December 31, 2018, please read Item 7. Management’s Discussion and Analysis of Financial Condition and Results of Operations located in Annual Report on Form 10-K for the year ended December 31, 2019.

Revenue:

<table>
<thead>
<tr>
<th>Revenue (in thousands):</th>
<th>2020</th>
<th>2019</th>
<th>Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>Government contracts</td>
<td>$ 217,246</td>
<td>$ 7,500</td>
<td>$ 209,746</td>
</tr>
<tr>
<td>Grants and other</td>
<td>258,352</td>
<td>11,162</td>
<td>250,852</td>
</tr>
<tr>
<td>Total revenue</td>
<td>$ 475,598</td>
<td>$ 18,662</td>
<td>$ 460,958</td>
</tr>
</tbody>
</table>

Revenue for 2020 was $475.6 million as compared to $18.7 million for 2019, an increase of $460.6 million. The significant increase in revenue in 2020 was a result of our development activities related to NVX-CoV2373 and was primarily comprised of revenue for services performed under the OWS Agreement and the CEPI Funding Agreement. Revenue for the year ended December 31, 2019 was primarily comprised of revenue for services performed under the BMGF Grant Agreement and recovery of costs on the close-out of our contract with HHS BARDA.

We expect revenue in 2021 to significantly increase due to our NVX-CoV2373 program, which we anticipate will continue to be funded by OWS and CEPI and/or other revenue sources. Further, we anticipate bringing our NVX-CoV2373 vaccine candidate to market following global regulatory approvals which, if achieved, should significantly impact revenue (also see below under "Liquidity and Capital Resources" in this Management’s Discussion and Analysis). In anticipation, we have entered into various APAs with government customers that are expected to result in the delivery of approximately 200 million doses of NVX-CoV2373 throughout 2021 and into the first half of 2022. We also entered into multiple supply and license agreements with strategic partners to supply NVX-CoV2373 in their specified territories under which we are entitled to receive royalty revenue from the sale of NVX-CoV2373 by such partners.

Expenses (in thousands):

<table>
<thead>
<tr>
<th>Expenses (in thousands):</th>
<th>2020</th>
<th>2019</th>
<th>Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>Research and development</td>
<td>$ 747,027</td>
<td>$ 113,842</td>
<td>$ 633,185</td>
</tr>
<tr>
<td>Gain on sale of assets</td>
<td>(9,016)</td>
<td>(9,016)</td>
<td></td>
</tr>
<tr>
<td>General and administrative</td>
<td>145,290</td>
<td>34,417</td>
<td>110,873</td>
</tr>
<tr>
<td>Total expenses</td>
<td>$ 892,317</td>
<td>$ 139,243</td>
<td>$ 753,074</td>
</tr>
</tbody>
</table>

Research and Development Expenses

During 2020, our research and development activities were primarily focused on the development of NVX-CoV2373. During 2020, direct external research and development expenses related to NVX-CoV2373 were $690.4 million and comprised of costs related to the following:

- expenses incurred under agreements with CROs that conduct our clinical trials and third-party consultants related to the development of NVX-CoV2373;
- developing and manufacturing the antigen drug substance and Matrix-M components of NVX-CoV2373 under agreements that we established with third-party CMOs and CDMOs;
- expenses incurred for the procurement of raw materials, laboratory supplies and equipment; and
- other costs related to preclinical studies and regulatory consulting, as well as related program management activities to support our growing global operations.
In 2020, we also incurred significant costs related to developing our NVX-CoV2373 manufacturing and supply network, including the immediate expense recognition of $245.9 million of ROU assets associated with such manufacturing supply agreements.

Research and development expenses increased to $747.0 million for 2020 as compared to $113.8 million for 2019, an increase of $633.2 million primarily due to the development of NVX-CoV2373, as shown in the table below.

The following summarizes our research and development expenses for the years ended December 31, 2020 and 2019 (in millions):

<table>
<thead>
<tr>
<th></th>
<th>2020</th>
<th>2019</th>
<th>Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>NVX-CoV2373</td>
<td>$ 609,401</td>
<td>$</td>
<td></td>
</tr>
<tr>
<td>NanoFlu</td>
<td>14,802</td>
<td>23,851</td>
<td></td>
</tr>
<tr>
<td>Other vaccine development programs</td>
<td>2,651</td>
<td>27,016</td>
<td></td>
</tr>
<tr>
<td>Total direct external research and development expense</td>
<td>$626,854</td>
<td>50,867</td>
<td></td>
</tr>
<tr>
<td>Employee expenses</td>
<td>45,882</td>
<td>33,389</td>
<td></td>
</tr>
<tr>
<td>Stock-based compensation expense</td>
<td>55,954</td>
<td>8,436</td>
<td></td>
</tr>
<tr>
<td>Facility expenses</td>
<td>7,232</td>
<td>9,243</td>
<td></td>
</tr>
<tr>
<td>Other expenses</td>
<td>11,105</td>
<td>11,907</td>
<td></td>
</tr>
<tr>
<td>Total research and development expenses</td>
<td>$747,027</td>
<td>$113,842</td>
<td></td>
</tr>
</tbody>
</table>

We do not provide forward-looking estimates of costs and time to complete our research programs due to the many uncertainties associated with vaccine development. As we obtain data from preclinical studies and clinical trials, we may elect to discontinue or delay clinical trials in order to focus our resources on more promising vaccine candidates. Completion of clinical trials may take several years or more, but the length of time can vary substantially depending upon the phase, size of clinical trial, primary and secondary endpoints and the intended use of the vaccine candidate. The cost of clinical trials may vary significantly over the life of a project as a result of a variety of factors, including:

- the number of participants who participate in the clinical trials;
- the number of sites included in the clinical trials;
- if clinical trial locations are domestic, international or both;
- the time to enroll participants;
- the duration of treatment and follow-up;
- the safety and efficacy profile of the vaccine candidate; and
- the cost and timing of, and the ability to secure, regulatory approvals.

As a result of these uncertainties, we are unable to determine the duration and completion costs of our research and development projects or when, and to what extent, we will generate future cash flows from our research projects.

For 2021, we expect research and development expenses to increase significantly over 2020 expenses due to our continued development activities for our NVX-CoV2373 program (see discussion on our NVX-CoV2373 program above) and increases in employee-related costs. Following regulatory approval of NVX-CoV2373, we expect products sales will result in certain types of costs recorded as research and development in 2020 being capitalized as inventory and expensed as cost of goods sold when product is delivered in 2021 and beyond. Cost of goods sold expenses could be significant depending on our commercial shipment levels.

Gain on Sale of Assets

As a result of the sale of assets transaction in 2019, we recorded a gain of $8.9 million.

General and Administrative Expenses

General and administrative expenses increased to $145.3 million for 2020 from $34.4 million for 2019, an increase of $110.9 million. The increase in general and administrative expenses is primarily due to increased employee-related costs, primarily stock-based compensation expense, and increased professional fees to support our NVX-CoV2373 program and relating to the acquisition and integration of Novavax CZ. As of December 31, 2020, we had 116 employees dedicated to general and administrative functions versus 41 employees as of December 31, 2019.

For 2021, we expect general and administrative expenses to increase significantly over 2020 expenses due to increased activities related to supporting our NVX-CoV2373 program and increases in employee-related costs.

Other Income (Expense):

We had total other expense, net of $1.5 million for 2020 compared to total other expense, net of $12.1 million for 2019, an increase of $10.6 million. In the year ended December 31, 2020, we recorded a $12.6 million gain on the intercompany loan with Novavax CZ due to changes in the exchange rates, and additional net interest expense of $1.5 million attributable to finance leases.

Net Loss:

Net loss for 2020 was $418.3 million, or $7.27 per share, as compared to $132.7 million, or $5.51 per share, for 2019, an increase of $285.6 million. The increase in net loss was primarily due to increased development activities relating to NVX-CoV2373, including the immediate expense recognition of $245.8 million of ROU assets associated with our manufacturing supply agreements for NVX-CoV2373 and increased employee-related costs, primarily stock-based compensation expense, partially offset by increased revenue under the CEPI Funding Agreement and the OWS Agreement.

The increase in weighted average shares outstanding for 2020 is primarily a result of the sale of 32.4 million shares of our common stock in the fourth quarter of 2020, and to a lesser degree, the conversion of our Series A Convertible Preferred Stock to 4.4 million shares of our common stock in the fourth quarter of 2020, weighted for the period the shares were outstanding during the year.

Liquidity Matters and Capital Resources

Our future capital requirements depend on numerous factors, including but not limited to our projected activities related to the development of NVX-CoV2373, including significant commitments under various CRO, CMO and CDMO agreements,
the progress of preclinical studies and clinical trials, the time and costs involved in obtaining regulatory approvals, the costs of filing, prosecuting, defending and enforcing patent claims and other intellectual property rights and other manufacturing, sales and distribution costs. We plan to continue developing other vaccines and product candidates, such as NanoFlu and potential combination vaccines candidates, which are in various stages of development. We believe our operating expenses and capital requirements will fluctuate depending upon the timing of events, such as the progress of our NVX-CoV2373 clinical trials and approval for the use of NVX-CoV2373 in the U.S. and internationally, as well as the scope, initiation and progress of our preclinical studies and clinical trials related to other research and development activities.

We have entered into APAs or supply agreements with various countries globally that, if our product candidate is approved, are expected to result in the delivery of approximately 200 million doses of NVX-CoV2373 throughout 2021 and into the first half of 2022. The APAs or supply agreements typically contain terms that include upfront payments intended to assist us in funding investments related to building out and operating our manufacturing and distribution network, among other expenses, in support of our global supply commitment. Such upfront payments generally become non-refundable upon our achievement of certain development milestones. We expect to sign additional APAs or supply agreements that are currently in active discussions and negotiations.

We have also entered into supply and license agreements with strategic partners to supply NVX-CoV2373 in their specified territories under which we are entitled to receive royalty revenue primarily from the sale of NVX-CoV2373 by our partners.

We funded our operations in 2020 with proceeds from the sale of common stock and preferred stock in equity offerings together with revenue under our CEPI Funding Agreement and the OWS Agreement that support our NVX-CoV2373 vaccine development activities. We anticipate our future operations to be funded by our cash, cash equivalents and marketable securities, revenue under our OWS, CEPI, DoD agreements, upfront payments under our APAs, and following any potential global development activities. We expect to sign additional APAs or supply agreements that are currently in active discussions and negotiations.

As of December 31, 2020, we had $806.4 million in cash and cash equivalents, marketable securities and restricted cash as compared to $82.2 million as of December 31, 2019. These amounts consisted of $553.4 million in cash and cash equivalents, $157.6 million in marketable securities and $95.3 million in restricted cash as of December 31, 2020 as compared to $78.8 million in cash and cash equivalents and $3.4 million in restricted cash as of December 31, 2019.

The following table summarizes cash flows for 2020 and 2019:

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Net cash (used in) provided by:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Operating activities</td>
<td>$(42,541)</td>
<td>$(136,623)</td>
<td>$94,082</td>
</tr>
<tr>
<td>Investing activities</td>
<td>(777,778)</td>
<td>38,492</td>
<td>$(416,270)</td>
</tr>
<tr>
<td>Financing activities</td>
<td>984,762</td>
<td>98,384</td>
<td>886,378</td>
</tr>
<tr>
<td>Effect on exchange rate on cash, cash equivalents and restricted cash</td>
<td>2,115</td>
<td>(32)</td>
<td>2,147</td>
</tr>
<tr>
<td>Net increase in cash, cash equivalents and restricted cash</td>
<td>566,558</td>
<td>221</td>
<td>566,337</td>
</tr>
<tr>
<td>Cash, cash equivalents and restricted cash at beginning of year</td>
<td>82,180</td>
<td>81,959</td>
<td>221</td>
</tr>
<tr>
<td>Cash, cash equivalents and restricted cash at end of year</td>
<td>$648,738</td>
<td>$82,180</td>
<td>$566,558</td>
</tr>
</tbody>
</table>

Net cash used in operating activities decreased to $42.5 million for 2020, as compared to $136.6 million for 2019. The decrease is primarily due to payments received under the CEPI Funding Agreement and OWS Agreement, and the timing of payments to third-parties.

During 2020, our investing activities primarily consisted of capital expenditures, purchases and maturities of marketable securities and our acquisition of Novavax CZ. During 2019, our investing activities primarily consisted of purchases and maturities of marketable securities and proceeds from the sale of assets. Capital expenditures for the year ended December 31, 2020 and 2019 were $54.6 million and $1.9 million, respectively, and the increase was primarily due to the build out of our facilities and related capital expenditures to support NVX-CoV2373. For 2021, we expect an increase in our capital expenditures due to further development activities for our NVX-CoV2373 program, including the additional build out of research and development and manufacturing facilities and related equipment, and the build-out of our new corporate office facility to accommodate anticipated increases in headcount.

Our financing activities consisted primarily of sales of our common stock under our At Market Issuance Sales Agreements and, to a much lesser extent, stock option exercises and purchases under our employee stock purchase plan. In 2020, we received net proceeds of $874.1 million (this amount excludes $3.2 million received in the first quarter of 2021 for shares traded in late December 2020) from the sale of shares of common stock through our At Market Issuance Sales Agreements and $200.0 million through the issuance of preferred stock in a private placement. In 2019, we received net proceeds of $97.5 million from selling shares of common stock through our At Market Issuance Sales Agreements.

**Contractual Obligations**

The following table summarizes our contractual obligations as of December 31, 2020 (in thousands):

<table>
<thead>
<tr>
<th>Contractual Obligations:</th>
<th>Total</th>
<th>Less than 1 Year</th>
<th>1 – 3 Years</th>
<th>3 – 5 Years</th>
<th>More than 5 Years</th>
</tr>
</thead>
<tbody>
<tr>
<td>Operating leases</td>
<td>$19,135</td>
<td>$5,392</td>
<td>$6,902</td>
<td>$4,762</td>
<td>$2,079</td>
</tr>
<tr>
<td>Finance leases obligation</td>
<td>153,800</td>
<td>112,625</td>
<td>41,175</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Convertible notes (a)</td>
<td>325,000</td>
<td>—</td>
<td>325,000</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Contractual obligations recognized as of December 31, 2020</td>
<td>497,935</td>
<td>118,017</td>
<td>373,077</td>
<td>4,762</td>
<td>2,079</td>
</tr>
<tr>
<td>Purchase commitments (b)</td>
<td>420,166</td>
<td>383,754</td>
<td>36,412</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Facilities lease agreement (c)</td>
<td>103,141</td>
<td>—</td>
<td>12,263</td>
<td>78,727</td>
<td>—</td>
</tr>
<tr>
<td>Total contractual obligations</td>
<td>$1,021,242</td>
<td>$501,771</td>
<td>$421,640</td>
<td>$17,025</td>
<td>$80,806</td>
</tr>
</tbody>
</table>

(a) See Note 11 to the consolidated financial statements included in this Annual Report regarding our Notes, which will mature on February 1, 2023, and bear cash interest of 3.75%, payable February 1 and August 1 of each year.

(b) This amount represents our non-cancelable fixed payment obligations under certain CMO and CDMO agreements that we are not contractually able to terminate for convenience. Certain agreements provide for termination rights subject to termination fees. Under such agreements, we are contractually obligated to make payments to vendors, mainly to reimburse them for their estimated unrecoverable expenses incurred. As of December 31, 2020, these agreements are active ongoing arrangements, and the Company expects to receive value from these arrangements in the future. The exact amount of such obligations is dependent on the timing of termination, and the exact terms of the relevant agreement, and cannot be reasonably estimated.

(c) This relates to the lease of 700 Quince Orchard that did not commence as of December 31, 2020 (see Note 7 to the consolidated financial statements).

In addition to the above obligations, we enter into a variety of agreements and financial commitments in the normal course of business. The terms generally allow us the option to cancel, reschedule, and adjust our requirements based on our business needs, prior to the delivery of goods or performance of services. It is not possible to predict the maximum potential amount of future payments under these agreements due to the conditional nature of our obligations and the unique facts and circumstances involved in each particular agreement.
Off-Balance Sheet Arrangements

We are not involved in any off-balance sheet agreements that have or are reasonably likely to have a material future effect on our financial condition, changes in financial condition, revenue or expenses, results of operations, liquidity, capital expenditures or capital resources.

Item 7A. QUANTITATIVE AND QUALITATIVE DISCLOSURES ABOUT MARKET RISK

We are subject to certain risks that may affect our results of operations, cash flows and fair values of assets and liabilities, including volatility in foreign currency exchange rates and interest rate movements.

Foreign Currency Exchange Risk

Although we are headquartered in the U.S., where we conduct the vast majority of our business activities, our results of operations are subject to foreign currency exchange rate fluctuations, including our foreign subsidiaries’ operations. We have two foreign consolidated subsidiaries, Novavax AB, which is located in Sweden, and Novavax CZ, which is located in the Czech Republic. While the financial results of our global activities are reported in U.S. dollars, the functional currency for our foreign subsidiaries is their respective local currency. Fluctuations in the foreign currency exchange rates of the countries in which we do business will affect our operating results, often in ways that are difficult to predict. A 10% decline in the exchange rate between the U.S. dollar and Swedish Krona would result in a decline of stockholders' equity (deficit) of approximately $4.7 million as of December 31, 2020. A 10% decline in the exchange rate between the U.S. dollar and Czech Koruna would result in a decline of stockholders' equity (deficit) of approximately $9.9 million as of December 31, 2020.

Interest Rate Risk

Our exposure to interest rate risk is primarily confined to our investment portfolio. As of December 31, 2020, our investments were classified as available-for-sale. We do not believe that a change in the market rates of interest would have any significant impact on the realizable value of our investment portfolio. Changes in interest rates may affect the investment income we earn on our marketable securities when they mature and the proceeds are reinvested into new marketable securities and, therefore, could impact our cash flows and results of operations. Our Notes have a fixed interest rate, and we have no additional material debt. As such, we do not believe that we are exposed to any material interest rate risk as a result of our borrowing activities.

Item 8. FINANCIAL STATEMENTS AND SUPPLEMENTARY DATA

The information required by this item is set forth on pages F-2 to F-38.

Item 9. CHANGES IN AND DISAGREEMENTS WITH ACCOUNTANTS ON ACCOUNTING AND FINANCIAL DISCLOSURE

None.

Item 9A. CONTROLS AND PROCEDURES

Evaluation of Disclosure Controls and Procedures

The term “disclosure controls and procedures” (defined in SEC Rule 13a-15(e)) refers to the controls and other procedures of a company that are designed to ensure that information required to be disclosed by a company in the reports that it files under the Securities Exchange Act of 1934 (the “Exchange Act”) is recorded, processed, summarized and reported, within time periods specified in the rules and forms of the Securities and Exchange Commission. “Disclosure controls and procedures” include, without limitation, controls and procedures designed to ensure that information required to be disclosed by a company in the reports that it files or submits under the Exchange Act is accumulated and communicated to the company’s management, including its principal executive and financial officers, or persons performing similar functions, as appropriate to allow timely decisions regarding required disclosure.

The Company’s management, with the participation of the chief executive officer and the chief financial officer, has evaluated the effectiveness of the Company’s disclosure controls and procedures as of the end of the period covered by this Annual Report (the “Evaluation Date”). Based on that evaluation, the Company’s chief executive officer and chief financial officer have concluded that, as of the Evaluation Date, such controls and procedures were effective at the reasonable assurance level.

Management’s Report on Internal Control over Financial Reporting

Our management is responsible for establishing and maintaining adequate internal control over financial reporting. Internal control over financial reporting is defined in Rules 13a-15(f) and 15d-15(f) promulgated under the Exchange Act, as a process designed by, or under the supervision of, the Company’s principal executive officer and principal financial officer and effected by the Company’s board of directors, management and other personnel, to provide reasonable assurance regarding the reliability of financial reporting and the preparation of financial statements for external purposes in accordance with generally accepted accounting principles in the United States (“GAAP”). Such internal control includes those policies and procedures that:

• pertain to the maintenance of records that in reasonable detail accurately and fairly reflect the transactions and
  dispositions of the assets of the Company;
  • provide reasonable assurance that transactions are recorded as necessary to permit preparation of financial
    statements in accordance with GAAP, and that receipts and expenditures of the Company are being made only in
    accordance with authorizations of management and directors of the Company; and
  • provide reasonable assurance regarding prevention or timely detection of an unauthorized acquisition, use or
    disposition of the Company’s assets that could have a material effect on the financial statements.

Because of its inherent limitations, internal control over financial reporting may not prevent or detect misstatements. Projections of any evaluation of effectiveness to future periods are subject to the risk that controls may become inadequate because of changes in conditions, or that the degree of compliance with the policies or procedures may deteriorate.

Our management assessed the effectiveness of our internal control over financial reporting as of December 31, 2020. In making this assessment, our management used the criteria set forth in the 2013 Internal Control-Integrated Framework issued by the Committee of Sponsoring Organizations of the Treadway Commission. Based on its assessment, our management has determined that, as of December 31, 2020, our internal controls over financial reporting are effective based on those criteria.

On May 27, 2020, we completed our acquisition of Novavax CZ. We are in the process of evaluating the existing controls and procedures of Novavax CZ and integrating it into our internal control over financial reporting. In accordance with SEC Staff guidance permitting a company to exclude an acquired business from management’s assessment of the effectiveness of internal control over financial reporting for the year in which the acquisition is completed, we have excluded the business that we acquired in the Novavax CZ acquisition from our assessment of the effectiveness of internal control over financial reporting as of December 31, 2020. The business that we acquired in the Novavax CZ acquisition represented 15% of the Company’s total assets as of
December 31, 2020, none of the Company’s revenue and less than 3% of the Company’s net loss for the year ended December 31, 2020.

Ernst & Young, LLP has issued a report on our internal control over financial reporting. This report is included in the Reports of Independent Registered Public Accounting Firm in Item 15. (a)(1).

Changes in Internal Control over Financial Reporting

Our management, including our chief executive officer and chief financial officer, has evaluated any changes in our internal control over financial reporting that occurred during the quarterly period ended December 31, 2020 and has concluded that there was no change that occurred during the quarterly period ended December 31, 2020 that materially affected, or is reasonably likely to materially affect, our internal control over financial reporting.

Management’s assessment of and conclusion on the effectiveness of disclosure controls and procedures and internal controls over financial reporting did not include the internal controls related to the operations acquired in the acquisition of Novavax CZ that are included in our December 31, 2020 consolidated financial statements. Our audit of internal control over financial reporting also did not include an evaluation of the internal control over financial reporting of Novavax CZ.

Equity Compensation Plan Information

<table>
<thead>
<tr>
<th>Plan Category</th>
<th>Number of Securities to be Issued Upon Exercise of Outstanding Options, Warrants and Rights</th>
<th>Weighted-Average Exercise Price of Outstanding Options, Warrants and Rights</th>
<th>Number of Securities Remaining Available for Future Issuance Under Equity Compensation Plans (Excluding Securities Reflected in Column (e))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Equity compensation plans approved by security holders</td>
<td>6,679,629</td>
<td>39.96</td>
<td>2,729,512</td>
</tr>
<tr>
<td>Equity compensation plans not approved by security holders</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
</tbody>
</table>

(1) Includes our 2015 Stock Incentive Plan, 2005 Stock Incentive Plan and ESPP. The weighted-average exercise price in column (b) excludes restricted stock units, which are not subject to an exercise price.

Item 9B. OTHER INFORMATION

None.

PART III

Item 10. DIRECTORS, EXECUTIVE OFFICERS AND CORPORATE GOVERNANCE

The information required by this item is incorporated by reference from our definitive Proxy Statement for our 2021 Annual Meeting of Stockholders scheduled to be held in June 2021 (the “2021 Proxy Statement”). We expect to file the 2021 Proxy Statement within 120 days after the close of the fiscal year ended December 31, 2020.

Item 11. EXECUTIVE COMPENSATION

We incorporate herein by reference the information required by this item concerning executive compensation to be contained in the 2021 Proxy Statement.

Item 12. SECURITY OWNERSHIP OF CERTAIN BENEFICIAL OWNERS AND MANAGEMENT AND RELATED STOCKHOLDER MATTERS

We incorporate herein by reference the information required by this item concerning security ownership of certain beneficial owners and management and related stockholder matters to be contained in the 2021 Proxy Statement.

The following table provides our equity compensation plan information as of December 31, 2020. Under these plans, our common stock may be issued upon the exercise of stock options and purchases under our Employee Stock Purchase Plan (“ESPP”). See also the information regarding our stock options and ESPP in Note 13 to the consolidated financial statements included herewith.

Item 13. CERTAIN RELATIONSHIPS AND RELATED TRANSACTIONS, AND DIRECTOR INDEPENDENCE

We incorporate herein by reference the information required by this item concerning certain related party transactions set forth in Note 16 to our consolidated financial statements included herewith. We incorporate herein by reference other information required by this item concerning certain other relationships and related transactions and director independence to be contained in the 2021 Proxy Statement.

Item 14. PRINCIPAL ACCOUNTING FEES AND SERVICES

We incorporate herein by reference the information required by this item concerning principal accountant fees and services to be contained in the 2021 Proxy Statement.

PART IV

Item 15. EXHIBITS AND FINANCIAL STATEMENT SCHEDULES

(a) The following documents are filed as part of the Annual Report:

(1) Index to Financial Statements

<table>
<thead>
<tr>
<th>Financial Statement Schedule</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reports of Independent Registered Public Accounting Firm</td>
<td>F-2</td>
</tr>
<tr>
<td>Consolidated Balance Sheets as of December 31, 2020 and 2019</td>
<td>F-7</td>
</tr>
<tr>
<td>Consolidated Statements of Operations and Statements of Comprehensive Loss for the years ended December 31, 2020, 2019 and 2018</td>
<td>F-9</td>
</tr>
<tr>
<td>Consolidated Statements of Stockholders’ Equity (Deficit) for the years ended December 31, 2020, 2019 and 2018</td>
<td>F-10</td>
</tr>
<tr>
<td>Consolidated Statements of Cash Flows for the years ended December 31, 2020, 2019 and 2018</td>
<td>F-11</td>
</tr>
<tr>
<td>Notes to Consolidated Financial Statements</td>
<td>F-12</td>
</tr>
</tbody>
</table>

(2) Financial Statement Schedules

Financial statement schedules are omitted because they are not applicable, not required under the instructions or all the information required is set forth in the financial statements or notes thereto.
### Exhibits

Exhibits marked with a single asterisk (*) are filed herewith.

Exhibits marked with a double dagger (††) refer to management contracts, compensatory plans or arrangements.

Confidential treatment has been granted for portions of exhibits marked with a double asterisk (**).

Confidential information contained in exhibits marked with a caret (^) has been omitted because it (i) is not material and/or (ii) would be competitively harmful if publicly disclosed.

All other exhibits listed have previously been filed with the SEC and are incorporated herein by reference.

<table>
<thead>
<tr>
<th>Exhibit Number</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.1</td>
<td>Second Amended and Restated Certificate of Incorporation of the Registrant (Incorporated by reference to Exhibit 3.1 to the Registrant’s Quarterly Report on Form 10-Q for the quarter ended June 30, 2015, filed on August 10, 2015 (File No. 000-26770))</td>
</tr>
<tr>
<td>3.2</td>
<td>Certificate of Amendment to the Second Amended and Restated Certificate of Incorporation of the Registrant (Incorporated by reference to Exhibit 3.1 to the Registrant’s Current Report on Form 8-K filed on May 9, 2019 (File No. 000-26770))</td>
</tr>
<tr>
<td>3.3</td>
<td>Amended and Restated By-Laws of the Registrant (Incorporated by reference to Exhibit 3.2 to the Registrant’s Annual Report on Form 10-K for the year ended December 31, 2012, filed on March 12, 2013 (File No. 000-26770))</td>
</tr>
<tr>
<td>3.4</td>
<td>Certificate of Designation of Series A Convertible Preferred Stock of the Registrant (Incorporated by reference to Exhibit 3.1 to the Registrant’s Current Report on Form 8-K filed June 19, 2020 (File No. 000-26770))</td>
</tr>
<tr>
<td>4.1</td>
<td>Specimen stock certificate for shares of common stock of the Registrant, par value $.01 per share (Incorporated by reference to Exhibit 4.1 to the Registrant’s Registration Statement on Form S-3, filed on December 31, 2019 (File No. 333-235761))</td>
</tr>
<tr>
<td>4.2</td>
<td>Indenture (including form of Notes) with respect to Novavax, Inc.’s 3.75% Convertible Senior Notes due 2023, dated as of January 29, 2016, between Novavax, Inc. and The Bank of New York Mellon Trust Company, N.A., as trustee (Incorporated by reference to Exhibit 4.1 to the Registrant’s Current Report on Form 8-K, filed on January 29, 2016 (File No. 000-26770))</td>
</tr>
<tr>
<td>4.3</td>
<td>Form of Series A Convertible Preferred Stock Certificate of the Registrant (Incorporated by reference to Exhibit 4.1 to the Registrant’s Current Report on Form 8-K filed June 19, 2020 (File No. 000-26770))</td>
</tr>
<tr>
<td>4.4*</td>
<td>Description of Registrant’s Securities</td>
</tr>
<tr>
<td>10.1††</td>
<td>Novavax, Inc. Amended and Restated 2005 Stock Incentive Plan (Incorporated by reference to Exhibit 10.2 to the Registrant’s Annual Report on Form 10-K for the year ended December 31, 2012, filed on March 12, 2013 (File No. 000-26770))</td>
</tr>
<tr>
<td>10.2††</td>
<td>Amendment to Amended and Restated 2005 Stock Incentive Plan (Incorporated by reference to Appendix 1 of the Registrant’s Definitive Proxy Statement filed on April 30, 2014 in connection with the Annual Meeting held on June 12, 2014 (File No. 000-26770))</td>
</tr>
<tr>
<td>10.3††</td>
<td>Form of Non-Statutory Stock Option Award Agreement granted under the Novavax, Inc. Amended and Restated 2005 Stock Incentive Plan (Incorporated by reference to Exhibit 10.4 to the Registrant’s Annual Report on Form 10-K for the year ended December 31, 2014, filed on February 27, 2015 (File No. 000-26770))</td>
</tr>
<tr>
<td>10.4††</td>
<td>Form of Incentive Stock Option Award Agreement granted under the Novavax, Inc. Amended and Restated 2005 Stock Incentive Plan (Incorporated by reference to Exhibit 10.5 to the Registrant’s Annual Report on Form 10-K for the year ended December 31, 2014, filed on February 27, 2015 (File No. 000-26770))</td>
</tr>
<tr>
<td>10.5††</td>
<td>Amended and Restated 2013 Employee Stock Purchase Plan (Incorporated by reference to Exhibit 10.1 to the Registrant’s Quarterly Report on Form 10-Q for the quarter ended March 30, 2020, filed on May 11, 2020 (File No. 000-26770))</td>
</tr>
<tr>
<td>10.6††</td>
<td>Amended and Restated Novavax, Inc. 2015 Stock Incentive Plan (Incorporated by reference to Appendix A of the Registrant’s Definitive Proxy Statement filed on May 13, 2020 in connection with the Annual Meeting held on June 25, 2020 (File No. 000-26770))</td>
</tr>
<tr>
<td>10.7††</td>
<td>Form of Non-Statutory Stock Option Award Agreement granted under the Novavax, Inc. 2015 Stock Incentive Plan (Incorporated by reference to Exhibit 10.3 to the Registrant’s Quarterly Report on Form 10-Q for the quarter ended June 30, 2015, filed on August 10, 2015 (File No. 000-26770))</td>
</tr>
</tbody>
</table>
10.38* Base Agreement between Novavax, Inc. and Advanced Technology International, dated June 25, 2020 (Incorporated by reference to Exhibit 10.1 to the Registrant’s Quarterly Report on Form 10-Q for the quarter ended September 30, 2020 (File No. 000-26770))

10.39* Undefinitized Project Agreement No. 1 between Novavax, Inc. and Advanced Technology International, dated July 6, 2020 (Incorporated by reference to Exhibit 10.2 to the Registrant’s Quarterly Report on Form 10-Q for the quarter ended September 30, 2020 (File No. 000-26770))

10.40* Modification No. 01 to Undefinitized Project Agreement No. 1 between Novavax, Inc. and Advanced Technology International, dated July 9, 2020 (Incorporated by reference to Exhibit 10.3 to the Registrant’s Quarterly Report on Form 10-Q for the quarter ended September 30, 2020 (File No. 000-26770))

10.41* Modification No. 02 to Undefinitized Project Agreement No. 1, entered into September 10, 2020, between the Company and Advanced Technology International

10.42* Modification No. 03 to Undefinitized Project Agreement No. 1, entered into September 18, 2020, between the Company and Advanced Technology International

10.43* Modification No. 04 to Undefinitized Project Agreement No. 1, entered into December 23, 2020, between the Company and Advanced Technology International

10.44* Modification No. 05 to Undefinitized Project Agreement No. 1, dated January 12, 2021, between the Company and Advanced Technology International

10.45* Amendment of Solicitation/Modification of Contract between Novavax, Inc. and the U.S. Department of Defense Joint Program Executive Office for Chemical, Biological, Radiological and Nuclear Defense, dated June 8, 2020 (Incorporated as reference to Exhibit 10.4 to the Registrant’s Quarterly Report on Form 10-Q for the quarter ended June 30, 2020 (File No. 000-26770))

10.46 Amendment of Solicitation/Modification of Contract between Novavax, Inc. and the U.S. Department of Defense Joint Program Executive Office for Chemical, Biological, Radiological and Nuclear Defense, dated June 8, 2020 (Incorporated as reference to Exhibit 10.4 to the Registrant’s Quarterly Report on Form 10-Q for the quarter ended June 30, 2020 (File No. 000-26770))

10.47 Amendment of Solicitation/Modification of Contract, Modification No. 2, entered into December 1, 2020, between the Company and the U.S. Department of Defense Joint Program Executive Office for Chemical, Biological, Radiological and Nuclear Defense

10.48* Amendment of Solicitation/Modification of Contract, Modification No. 3, entered into January 5, 2021, between the Company and the U.S. Department of Defense Joint Program Executive Office for Chemical, Biological, Radiological and Nuclear Defense

10.49* Base Call Option Transaction Confirmation, dated as of January 25, 2016, between Novavax and JPMorgan Chase Bank, National Association, London Branch (Incorporated by reference to Exhibit 10.51 to the Registrant’s Current Report on Form 8-K, filed on January 29, 2016 (File No. 000-26770))

10.50 Base Call Option Transaction Confirmation, dated as of January 25, 2016, between Novavax and Morgan Stanley & Co. LLC (Incorporated by reference to Exhibit 10.2 to the Registrant’s Current Report on Form 8-K, filed on January 29, 2016 (File No. 000-26770))

10.51 Additional Base Call Option Transaction Confirmation, dated as of February 2, 2016, between Novavax and JPMorgan Chase Bank, National Association, London Branch (Incorporated by reference to Exhibit 10.52 to the Registrant’s Annual Report on Form 10-K for the year ended December 31, 2015, filed on February 29, 2016 (File No. 000-26770))

10.52 Additional Base Call Option Transaction Confirmation, dated as of February 2, 2016, between Novavax and Morgan Stanley & Co. LLC (Incorporated by reference to Exhibit 10.52 to the Registrant’s Annual Report on Form 10-K for the year ended December 31, 2015, filed on February 29, 2016 (File No. 000-26770))


10.54 Restated Funding Agreement, entered into on May 11, 2020, between Novavax, Inc. and the Coalition for Epidemic Preparedness Innovations (Incorporated as reference to Exhibit 10.1 to the Registrant’s Quarterly Report on Form 10-Q for the quarter ended June 30, 2020 (File No. 000-26770))

10.55 Amendment Number 1 to the IPDP and Budget of the Outbreak Response Funding Agreement (Step 2), entered into on November 2, 2020, between Novavax, Inc. and the Coalition for Epidemic Preparedness Innovations

10.56** Letter Contract between Novavax, Inc. and the U.S. Department of Defense Joint Program Executive Office for Chemical, Biological, Radiological and Nuclear Defense, dated September 16, 2020 (Incorporated as reference to Exhibit 10.3 to the Registrant’s Quarterly Report on Form 10-Q for the quarter ended June 30, 2020 (File No. 000-26770))

10.57 Share Purchase Agreement between Novavax, Inc. (solely as guarantor), Novavax AB, De Bilt Holdings B.V., Poonawalla Science Park B.V., Bilthoven Biologicals B.V. and Serum Institute International B.V. (solely as guarantor), dated May 27, 2020 (Incorporated as reference to Exhibit 10.3 to the Registrant’s Quarterly Report on Form 10-Q for the quarter ended June 30, 2020 (File No. 000-26770))

10.58 Code of Business Conduct and Ethics (Incorporated by reference to Exhibit 14 to the Registrant’s Quarterly Report on Form 10-Q for the quarter ended June 30, 2011, filed on August 9, 2011 (File No. 000-26770))

11* Subsidiaries of the Registrant

23* Consent of Ernst & Young, LLP Independent Registered Public Accounting Firm

24* Certification of Chief Executive Officer pursuant to Rule 13a-14(a) or 15d-14(e) of the Securities Exchange Act

24* Certification of Chief Financial Officer pursuant to Rule 13a-14(a) or 15d-14(e) of the Securities Exchange Act

32* Certification of Chief Executive Officer pursuant to 18 U.S.C. Section 1350, as adopted pursuant to Section 906 of the Sarbanes-Oxley Act of 2002

32* Certification of Chief Financial Officer pursuant to 18 U.S.C. Section 1350, as adopted pursuant to Section 906 of the Sarbanes-Oxley Act of 2002

101 The following financial information from our Annual Report on Form 10-K for the year ended December 31, 2020, formatted in Extensible Business Reporting Language (XBRL): (i) the Consolidated Balance Sheets as of December 31, 2020 and 2019, (ii) the Consolidated Statements of Operations for the three years in the period ended December 31, 2020, (iii) the Consolidated Statements of Comprehensive Loss for the three years in the period ended December 31, 2020, (iv) the Consolidated Statements of Changes in Stockholders’ Equity (Deficit) for the three years in the period ended December 31, 2020, (v) the Consolidated Statements of Cash Flows for the three years in the period ended December 31, 2020, and (vi) the Notes to Consolidated Financial Statements.

Item 16. FORM 10-K SUMMARY
Not applicable.

SIGNATURES
Pursuant to the requirements of Section 13 or 15(d) of the Securities Exchange Act of 1934, the registrant has duly caused this report to be signed on its behalf by the undersigned, thereunto duly authorized.

NOVAVAX, INC.

By: /s/ Stanley C. Erck

Stanley C. Erck

President and Chief Executive Officer

Date: March 1, 2021
Pursuant to the requirements of the Securities Exchange Act of 1934, this report has been signed below by the following persons on behalf of the Registrant and in the capacities and on the dates indicated:

<table>
<thead>
<tr>
<th>Name</th>
<th>Title</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>/s/ Stanley C. Erck</td>
<td>President and Chief Executive Officer and Director (Principal Executive Officer)</td>
<td>March 1, 2021</td>
</tr>
<tr>
<td>Stanley C. Erck</td>
<td></td>
<td></td>
</tr>
<tr>
<td>/s/ Gregory F. Covino</td>
<td>Executive Vice President, Chief Financial Officer (Principal Financial and Accounting Officer)</td>
<td>March 1, 2021</td>
</tr>
<tr>
<td>Gregory F. Covino</td>
<td></td>
<td></td>
</tr>
<tr>
<td>/s/ James F. Young</td>
<td>Chairman of the Board of Directors</td>
<td>March 1, 2021</td>
</tr>
<tr>
<td>James F. Young</td>
<td></td>
<td></td>
</tr>
<tr>
<td>/s/ Gregg H. Alton</td>
<td>Director</td>
<td>March 1, 2021</td>
</tr>
<tr>
<td>Gregg H. Alton</td>
<td></td>
<td></td>
</tr>
<tr>
<td>/s/ Richard H. Douglas</td>
<td>Director</td>
<td>March 1, 2021</td>
</tr>
<tr>
<td>Richard H. Douglas</td>
<td></td>
<td></td>
</tr>
<tr>
<td>/s/ Gary C. Evans</td>
<td>Director</td>
<td>March 1, 2021</td>
</tr>
<tr>
<td>Gary C. Evans</td>
<td></td>
<td></td>
</tr>
<tr>
<td>/s/ Rachel K. King</td>
<td>Director</td>
<td>March 1, 2021</td>
</tr>
<tr>
<td>Rachel K. King</td>
<td></td>
<td></td>
</tr>
<tr>
<td>/s/ Margaret G. McGlynn</td>
<td>Director</td>
<td>March 1, 2021</td>
</tr>
<tr>
<td>Margaret G. McGlynn</td>
<td></td>
<td></td>
</tr>
<tr>
<td>/s/ Michael A. McManus</td>
<td>Director</td>
<td>March 1, 2021</td>
</tr>
<tr>
<td>Michael A. McManus</td>
<td></td>
<td></td>
</tr>
<tr>
<td>/s/ Rajiv I. Modi</td>
<td>Director</td>
<td>March 1, 2021</td>
</tr>
<tr>
<td>Rajiv I. Modi</td>
<td></td>
<td></td>
</tr>
<tr>
<td>/s/ David M. Mott</td>
<td>Director</td>
<td>March 1, 2021</td>
</tr>
<tr>
<td>David M. Mott</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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Years ended December 31, 2020, 2019 and 2018

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- Consolidated Statements of Changes in Stockholders' Equity (Deficit) for the years ended December 31, 2020, 2019 and 2018 F- 10
- Consolidated Statements of Cash Flows for the years ended December 31, 2020, 2019 and 2018 F- 11
- Notes to Consolidated Financial Statements F- 12
The Company measures and mined that certain of these arrangements contain information of allowable costs to be incurred for cision with clinical research and

We have audited, in accordance with the standards of the Public Company Accounting Oversight Board (United States) (PCAOB), the Company’s internal control over financial reporting as of December 31, 2020, based on criteria established in Internal Control—Integrated Framework issued by the Committee of Sponsoring Organizations of the Treadway Commission (2013 framework) and our report dated March 1, 2021, expressed an unqualified opinion thereon.

These financial statements are the responsibility of the Company’s management. Our responsibility is to express an opinion on the Company’s financial statements based on our audits. We are a public accounting firm registered with the PCAOB and are required to be independent with respect to the Company in accordance with the U.S. federal securities laws and the applicable rules and regulations of the Securities and Exchange Commission and the PCAOB.

We conducted our audits in accordance with the standards of the PCAOB. Those standards require that we plan and perform the audit to obtain reasonable assurance about whether the financial statements are free of material misstatement, whether due to error or fraud. Our audits included performing procedures to assess the risks of material misstatement of the financial statements, whether due to error or fraud, and performing procedures that respond to those risks. Such procedures included examining, on a test basis, evidence regarding the amounts and disclosures in the financial statements. Our audits also included evaluating the accounting principles used and significant estimates made by management, as well as evaluating the overall presentation of the financial statements. We believe that our audits provide a reasonable basis for our opinion.

The critical audit matters communicated below are matters arising from the current period audit of the financial statements that were communicated or required to be communicated to the audit committee and that: (1) relate to accounts or disclosures that are material to the financial statements and (2) involved our especially challenging, subjective or complex judgments. The communication of the critical audit matters does not alter in any way our opinion on the consolidated financial statements, taken as a whole, and we are not, by communicating the critical audit matters below, providing a separate opinion on the critical audit matters or on the account or disclosures to which they relate.

Revenue recognition related to the cost-based input method for U.S. government contracts

As described in Note 2 to the consolidated financial statements, the Company recorded $217.2 million of revenue from U.S. government contracts to advance the clinical development and manufacturing of NVX-CoV2373 on a reimbursable-cost or reimbursable-cost-plus fixed fee basis. The Company measures progress toward satisfaction of its performance obligations using a cost-based input method that requires an estimate of total allowable cost at completion. Estimating the total allowable costs at completion is highly subjective. Changes in the estimated total allowable cost at completion could materially impact the timing of revenue recognition. Allowable contract costs include direct costs incurred on the contract and indirect costs that are applied in the form of rates to the direct costs.

Auditing revenue recognition based on the cost-based input method involved subjective auditor judgment. The estimates of costs at completion are based on management’s assessment of the costs necessary to fulfill its performance obligations under the contracts. Auditing allowable contract costs was complex due to the specialized knowledge needed to evaluate the costs included in the calculation of indirect rates and the contract terms.

We obtained an understanding, evaluated the design, and tested the operating effectiveness of controls over recognition of revenue under the cost-based input method. For example, we tested controls over the appropriateness of significant assumptions regarding the estimation of allowable costs to be incurred for the performance obligations and controls over the appropriateness of the indirect rate calculation.

To test the recognition of revenue under the cost-based input method, our audit procedures included among others, reviewing management’s estimate of total allowable costs at completion for consistency with contract terms, obtaining an understanding of the stage of completion through review of project deliverables, evidencing of stage of completion including discussion with clinical research and manufacturing teams, and comparing actual results to prior management estimates. To test the recognition of revenue related to indirect rates, our audit procedures included among others, testing the allowability of the underlying costs used in the Company’s calculation of indirect rates. We utilized specialists to evaluate the treatment of significant indirect cost types.

Identification of embedded leases related to manufacturing supply agreements

As described in Note 7 to the consolidated financial statements, the Company entered into multiple manufacturing supply agreements with contract manufacturing organizations and contract development and manufacturing organizations. The Company determined that certain of these arrangements contain embedded leases as it has the exclusive use of, and control over, a portion of the manufacturing facility or equipment of the contract manufacturing organization during the contractual term of the arrangements. As a result of identifying embedded leases in certain of these arrangements, the Company immediately expensed $245.9 million, which represented the right of use assets related to these arrangements that currently do not have alternative future use.

Auditing embedded leases within manufacturing supply agreements was complex due to the judgment required to evaluate whether each arrangement included a lease and the related lease term. This significant auditor judgment involves the assessment of whether the Company has the right to obtain substantially all of the economic benefits from the use of identified assets and an assessment of the lease term, including whether the Company is reasonably certain not to exercise its termination provisions within the arrangements.
How We Addressed the Matter in Our Audit

We obtained an understanding, evaluated the design, and tested the operating effectiveness of controls over the identification of embedded leases in supply agreements. For example, we tested controls over management’s review of the supply agreements that evaluated whether management was entitled to substantially all of the economic benefits, as well as management’s assessment of the various termination provisions.

To test the Company’s identification of embedded leases, our audit procedures included among others, reviewing the terms of manufacturing supply agreements with contract manufacturing organizations and contract development and manufacturing organizations, obtaining an understanding of the facilities and equipment subject to the arrangements through discussions with representatives of the counterparties, and evaluating the identification of embedded leases and determination of the lease term.

/s/ Ernst & Young, LLP

We have served as the Company’s auditor since 2014.

Tysons, Virginia

March 1, 2021

Report of Independent Registered Public Accounting Firm

To the Board of Directors and Stockholders of Novavax, Inc.

Opinion on Internal Control over Financial Reporting

We have audited Novavax, Inc.’s internal control over financial reporting as of December 31, 2020, based on criteria established in Internal Control—Integrated Framework issued by the Committee of Sponsoring Organizations of the Treadway Commission (2013 framework) (the COSO criteria). In our opinion, Novavax, Inc. (the Company) maintained, in all material respects, effective internal control over financial reporting as of December 31, 2020, based on the COSO criteria.

As indicated in the accompanying Management’s Report on Internal Control over Financial Reporting included in item 9A, management’s assessment of and conclusion of the effectiveness of internal control over financial reporting did not include the internal controls of Novavax CZ (formerly Praha Vaccines a.s.), which is included in the 2020 consolidated financial statements of the Company and constituted 15% of total assets, as of December 31, 2020, and 0% and 3% of revenue and net loss, respectively, for the year then ended. Our audit of internal control over financial reporting of the Company also did not include an evaluation of the internal control over financial reporting of Novavax CZ.

We also have audited, in accordance with the standards of the Public Company Accounting Oversight Board (United States) (PCAOB), the consolidated balance sheets of the Company as of December 31, 2020 and 2019, the related consolidated statements of operations, comprehensive loss, changes in stockholders’ equity (deficit), and cash flows for each of the three years in the period ended December 31, 2020, and the related notes and our report dated March 1, 2021, expressed an unqualified opinion thereon.

Basis for Opinion

The Company’s management is responsible for maintaining effective internal control over financial reporting and for its assessment of the effectiveness of internal control over financial reporting included in the accompanying Management’s Report on Internal Control over Financial Reporting included in Item 9A. Our responsibility is to express an opinion on the Company’s internal control over financial reporting based on our audit. We are a public accounting firm registered with the PCAOB and are required to be independent with respect to the Company in accordance with the U.S. federal securities laws and the applicable rules and regulations of the Securities and Exchange Commission and the PCAOB.

We conducted our audit in accordance with the standards of the PCAOB. Those standards require that we plan and perform the audit to obtain reasonable assurance about whether effective internal control over financial reporting was maintained in all material respects.

Our audit included obtaining an understanding of internal control over financial reporting, assessing the risk that a material weakness exists, testing and evaluating the design and operating effectiveness of internal control based on the assessed risk, and performing such other procedures as we considered necessary in the circumstances. We believe that our audit provides a reasonable basis for our opinion.

Definition and Limitations of Internal Control Over Financial Reporting

A company’s internal control over financial reporting is a process designed to provide reasonable assurance regarding the reliability of financial reporting and the preparation of financial statements for external purposes in accordance with generally accepted accounting principles. A company’s internal control over financial reporting includes those policies and procedures that (1) pertain to the maintenance of records that, in reasonable detail, accurately and fairly reflect the transactions and dispositions of the assets of the company; (2) provide reasonable assurance that transactions are recorded as necessary to permit preparation of financial statements in accordance with generally accepted accounting principles, and that receipts and expenditures of the company are being made only in accordance with authorizations of management and directors of the company; and (3) provide reasonable assurance regarding prevention or timely detection of unauthorized acquisition, use, or disposition of the company’s assets that could have a material effect on the financial statements.
Because of its inherent limitations, internal control over financial reporting may not prevent or detect misstatements. Also, projections of any evaluation of effectiveness to future periods are subject to the risk that controls may become inadequate because of changes in conditions, or that the degree of compliance with the policies or procedures may deteriorate.

/s/ Ernst & Young, LLP
Tysons, Virginia
March 1, 2021

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### NOVAVAX, INC.

#### CONSOLIDATED BALANCE SHEETS

<table>
<thead>
<tr>
<th></th>
<th>December 31,</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>2020</td>
<td>2019</td>
</tr>
<tr>
<td><strong>ASSETS</strong></td>
<td>(in thousands, except share and per share information)</td>
<td></td>
</tr>
<tr>
<td><strong>Current assets:</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cash and cash equivalents</td>
<td>$553,398</td>
<td>$78,823</td>
</tr>
<tr>
<td>Marketable securities</td>
<td>$157,649</td>
<td>—</td>
</tr>
<tr>
<td>Restricted cash</td>
<td>$93,880</td>
<td>$2,947</td>
</tr>
<tr>
<td>Accounts receivable</td>
<td>$262,012</td>
<td>$7,500</td>
</tr>
<tr>
<td>Prepaid expenses and other current assets</td>
<td>$181,264</td>
<td>$7,977</td>
</tr>
<tr>
<td><strong>Total current assets</strong></td>
<td>$1,248,203</td>
<td>$97,247</td>
</tr>
<tr>
<td>Restricted cash</td>
<td>$1,460</td>
<td>$410</td>
</tr>
<tr>
<td>Property and equipment, net</td>
<td>$179,954</td>
<td>$11,445</td>
</tr>
<tr>
<td>Intangible assets, net</td>
<td>$5,725</td>
<td>$5,581</td>
</tr>
<tr>
<td>Goodwill</td>
<td>$135,379</td>
<td>$51,154</td>
</tr>
<tr>
<td>Other non-current assets</td>
<td>$11,758</td>
<td>$7,120</td>
</tr>
<tr>
<td><strong>Total assets</strong></td>
<td>$1,582,479</td>
<td>$172,957</td>
</tr>
</tbody>
</table>

|                     |              |       |
| **LIABILITIES AND STOCKHOLDERS’ EQUITY (DEFICIT)** |       |       |
| **Current liabilities:** |           |       |
| Accounts payable      | $54,332     | $2,910 |
| Accrued expenses      | $137,390    | $14,867 |
| Accrued interest      | $5,078      | $5,078 |
| Deferred revenue      | $273,228    | $1,678 |
| Current portion of finance lease liabilities | $105,862 | — |
| Other current liabilities | $3,782    | $1,262 |
| **Total current liabilities** | $579,672 | $25,795 |
| Convertible notes payable | $322,035 | $320,611 |
| Non-current finance lease liabilities | $40,083 | — |
| Other non-current liabilities | $13,480 | $12,568 |
| **Total liabilities** | $955,270    | $358,974 |

| **Preferred stock, $0.01 par value, 2,000,000 shares authorized at December 31, 2020 and 2019; no shares issued and outstanding at December 31, 2020 and 2019** |       |       |

---
STOCKHOLDERS' EQUITY (DEFICIT)

Common stock, $0.01 par value, 600,000,000 shares authorized at December 31, 2020 and 2019; and 71,350,365 shares issued and 70,953,739 shares outstanding at December 31, 2020 and 32,399,352 shares issued and 32,352,416 shares outstanding at December 31, 2019

<table>
<thead>
<tr>
<th></th>
<th>2020</th>
<th>2019</th>
<th>2018</th>
</tr>
</thead>
<tbody>
<tr>
<td>Additional paid-in capital</td>
<td>2,535,476</td>
<td>1,260,551</td>
<td></td>
</tr>
<tr>
<td>Accumulated deficit</td>
<td>(1,874,199)</td>
<td>(1,431,801)</td>
<td></td>
</tr>
<tr>
<td>Treasury stock, 396,626 shares, cost basis at December 31, 2020 and 46,936 shares, cost basis at December 31, 2019</td>
<td>(41,806)</td>
<td>(2,583)</td>
<td></td>
</tr>
<tr>
<td>Accumulated other comprehensive income (loss)</td>
<td>7,024</td>
<td>(12,508)</td>
<td></td>
</tr>
<tr>
<td>Total stockholders' equity (deficit)</td>
<td>627,209</td>
<td>(186,017)</td>
<td></td>
</tr>
<tr>
<td>Total liabilities and stockholders' equity (deficit)</td>
<td>$ 1,582,479</td>
<td>$ 172,957</td>
<td></td>
</tr>
</tbody>
</table>

The accompanying notes are an integral part of these financial statements.
NOVAVAX, INC.
CONSOLIDATED STATEMENTS OF CHANGES IN STOCKHOLDERS' EQUITY (DEFICIT)
Year Ended December 31, 2020, 2019 and 2018

<table>
<thead>
<tr>
<th>Common Stock</th>
<th>Additional Paid-in Capital</th>
<th>Accumulated Deficit</th>
<th>Treasury Stock</th>
<th>Accumulated Other Comprehensive Income (Loss)</th>
<th>Total Stockholders' Equity (Deficit)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shares</td>
<td>Amount</td>
<td>(in thousands)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Balance at December 31, 2017</td>
<td>16,184,241 $</td>
<td>162 $</td>
<td>$1,023,532 (1,314,259) $</td>
<td>(2,456) $</td>
<td>(8,617) $</td>
</tr>
<tr>
<td>Non-cash stock-based compensation</td>
<td>—</td>
<td>—</td>
<td>18,314</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Stock issued under incentive programs</td>
<td>120,561</td>
<td>1</td>
<td>2,744</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Restricted stock cancelled</td>
<td>(936)</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Issuance of common stock, net of issuance costs of $4,265</td>
<td>2,941,438</td>
<td>29</td>
<td>100,031</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Unrealized gain on marketable securities</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>12</td>
<td>12</td>
</tr>
<tr>
<td>Foreign currency translation adjustment</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>(2,586)</td>
<td>(2,586)</td>
</tr>
<tr>
<td>Net loss</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>(184,748)</td>
<td>(184,748)</td>
</tr>
<tr>
<td>Balance at December 31, 2018</td>
<td>19,245,302 $</td>
<td>192</td>
<td>1,144,621 (1,299,187) $</td>
<td>(2,456)</td>
<td>(11,191) $</td>
</tr>
<tr>
<td>Non-cash stock-based compensation</td>
<td>—</td>
<td>—</td>
<td>17,048</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Stock issued under incentive programs</td>
<td>173,873</td>
<td>2</td>
<td>1,122</td>
<td>(132)</td>
<td>—</td>
</tr>
<tr>
<td>Fractional shares purchased in stock split</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>(1)</td>
<td>(1)</td>
</tr>
<tr>
<td>Issuance of common stock, net of issuance costs of $1,655</td>
<td>12,980,177</td>
<td>130</td>
<td>97,780</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Unrealized gain on marketable securities</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>Foreign currency translation adjustment</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>(1,322)</td>
<td>(1,322)</td>
</tr>
<tr>
<td>Net loss</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>(132,694)</td>
<td>(132,694)</td>
</tr>
<tr>
<td>Balance at December 31, 2019</td>
<td>32,199,392 $</td>
<td>324</td>
<td>1,269,551 (1,431,801) $</td>
<td>(2,583)</td>
<td>(12,508) $</td>
</tr>
<tr>
<td>Preferred stock beneficial conversion</td>
<td>—</td>
<td>—</td>
<td>24,139 (24,139)</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Conversion of preferred stock</td>
<td>4,388,950</td>
<td>44</td>
<td>199,778</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Non-cash stock-based compensation</td>
<td>—</td>
<td>—</td>
<td>128,035</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Stock issued under incentive programs</td>
<td>2,166,725</td>
<td>22</td>
<td>44,447 (39,223)</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Issuance of common stock, net of issuance costs of $11,416</td>
<td>32,193,438</td>
<td>324</td>
<td>878,526</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Unrealized gain on marketable securities</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>9</td>
<td>9</td>
</tr>
<tr>
<td>Foreign currency translation adjustment</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>(418,259)</td>
<td>(418,259)</td>
</tr>
<tr>
<td>Net loss</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>(1,424,790)</td>
<td>(1,424,790)</td>
</tr>
<tr>
<td>Balance at December 31, 2020</td>
<td>71,359,365 $</td>
<td>714</td>
<td>3,535,476 (1,874,199) $</td>
<td>(41,806)</td>
<td>(7,624) $</td>
</tr>
</tbody>
</table>

The accompanying notes are an integral part of these financial statements.

NOVAVAX, INC.
CONSOLIDATED STATEMENTS OF CASH FLOWS
Year Ended December 31, 2020, 2019 and 2018

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>(in thousands)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Operating Activities:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Net loss</td>
<td>(418,259)</td>
<td>(132,694)</td>
<td>(184,748)</td>
</tr>
<tr>
<td>Reconciliation of net loss to net cash used in operating activities:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Depreciation and amortization</td>
<td>4,885</td>
<td>5,676</td>
<td>8,139</td>
</tr>
<tr>
<td>Gain on sale of assets</td>
<td>(9,016)</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Non-cash impact of lease termination</td>
<td>—</td>
<td>—</td>
<td>(4,381)</td>
</tr>
<tr>
<td>Amortization of debt issuance costs</td>
<td>1,424</td>
<td>1,424</td>
<td>1,424</td>
</tr>
<tr>
<td>Right-of-use assets expense</td>
<td>245,861</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Non-cash stock-based compensation</td>
<td>128,035</td>
<td>17,048</td>
<td>18,314</td>
</tr>
<tr>
<td>Other</td>
<td>(16,504)</td>
<td>4,957</td>
<td>(2,451)</td>
</tr>
<tr>
<td>Changes in operating assets and liabilities:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Accounts receivable, prepaid expenses and other assets</td>
<td>(422,689)</td>
<td>(4,202)</td>
<td>1,212</td>
</tr>
<tr>
<td>Accounts payable and accrued expenses</td>
<td>161,161</td>
<td>(11,485)</td>
<td>(6,744)</td>
</tr>
<tr>
<td>Deferred revenue</td>
<td>271,245</td>
<td>(8,331)</td>
<td>(15,610)</td>
</tr>
<tr>
<td>Net cash used in operating activities</td>
<td>(42,541)</td>
<td>(136,623)</td>
<td>(184,823)</td>
</tr>
<tr>
<td>Investing Activities:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Capital expenditures</td>
<td>(54,623)</td>
<td>(1,857)</td>
<td>(1,372)</td>
</tr>
<tr>
<td>Acquisition of Novavax CZ, net of cash acquired</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Proceeds from sale of assets</td>
<td>18,333</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Purchases of marketable securities</td>
<td>(363,202)</td>
<td>(17,446)</td>
<td>(120,150)</td>
</tr>
<tr>
<td>Proceeds from maturities of marketable securities</td>
<td>205,162</td>
<td>39,800</td>
<td>150,118</td>
</tr>
<tr>
<td>Net cash (used in) provided by investing activities</td>
<td>(377,778)</td>
<td>38,492</td>
<td>28,896</td>
</tr>
<tr>
<td>Financing Activities:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Net proceeds from sale of preferred stock</td>
<td>199,822</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Net proceeds from sales of common stock</td>
<td>875,623</td>
<td>97,392</td>
<td>100,060</td>
</tr>
<tr>
<td>Proceeds from the exercise of stock-based awards</td>
<td>44,609</td>
<td>992</td>
<td>2,745</td>
</tr>
<tr>
<td>Treasury stock related to tax withholding on stock-based awards</td>
<td>(39,087)</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Finance lease payments</td>
<td>(96,065)</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Net cash provided by financing activities</td>
<td>984,762</td>
<td>98,384</td>
<td>102,805</td>
</tr>
<tr>
<td>Effect of exchange rate on cash, cash equivalents and restricted cash</td>
<td>2,115</td>
<td>(52)</td>
<td>(48)</td>
</tr>
<tr>
<td>Net increase (decrease) in cash, cash equivalents and restricted cash</td>
<td>566,358</td>
<td>221</td>
<td>(3,472)</td>
</tr>
<tr>
<td>Cash, cash equivalents and restricted cash at beginning of year</td>
<td>82,180</td>
<td>81,959</td>
<td>135,431</td>
</tr>
<tr>
<td>Cash, cash equivalents and restricted cash at end of year</td>
<td>648,738</td>
<td>82,180</td>
<td>81,959</td>
</tr>
<tr>
<td>Supplemental disclosure of non-cash activities:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sale of common stock under the Sales Agreement not settled at year-end</td>
<td>3,227</td>
<td>497</td>
<td>—</td>
</tr>
<tr>
<td>Capital expenditures included in accounts payable and accrued expenses</td>
<td>9,255</td>
<td>49</td>
<td>519</td>
</tr>
<tr>
<td>Right-of-use assets from new lease agreements</td>
<td>247,399</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Supplemental disclosure of cash flow information:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cash interest payments, net of amounts capitalized</td>
<td>13,705</td>
<td>12,188</td>
<td>12,188</td>
</tr>
</tbody>
</table>

The accompanying notes are an integral part of these financial statements.
Note 1 – Organization

Novavax, Inc. ("Novavax," and together with its wholly owned subsidiaries, Novavax AB and Novavax CZ (formerly Praha Vaccines a.s.), the "Company") is a late-stage biotechnology company that promotes improved global health through the discovery, development and commercialization of innovative vaccines to prevent serious infectious diseases and address urgent, global health needs. The Company’s vaccine candidates, including both its coronavirus vaccine candidate, NVX-CoV2373, and its lead influenza vaccine candidate, NanoFlu™, are genetically engineered, three-dimensional nanostructures of recombinant proteins critical to disease pathogenesis and may elicit differentiated immune responses, which may be more efficacious than naturally occurring immunity or traditional vaccines.

Note 2 – Summary of Significant Accounting Policies

Basis of Presentation

The consolidated financial statements include the accounts of Novavax, Inc. and its wholly owned subsidiaries, Novavax AB and Novavax CZ. All intercompany accounts and transactions have been eliminated in consolidation.

Use of Estimates

The preparation of the consolidated financial statements in conformity with generally accepted accounting principles in the United States ("U.S. GAAP") requires management to make estimates and assumptions that affect the reported amounts of assets and liabilities and disclosure of contingent assets and liabilities at the date of the consolidated financial statements and the reported amounts of revenue and expenses during the reporting period. Actual results could differ materially from those estimates.

Cash and Cash Equivalents

Cash and cash equivalents consist of highly liquid investments with maturities of three months or less from the date of purchase. Cash and cash equivalents consist of the following on December 31 (in thousands):

<table>
<thead>
<tr>
<th></th>
<th>2020</th>
<th>2019</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cash</td>
<td>$122,312</td>
<td>$15,863</td>
</tr>
<tr>
<td>Money market funds</td>
<td>96,116</td>
<td>42,960</td>
</tr>
<tr>
<td>Government-backed securities</td>
<td>44,250</td>
<td>20,000</td>
</tr>
<tr>
<td>Treasury securities</td>
<td>44,052</td>
<td>—</td>
</tr>
<tr>
<td>Corporate debt securities</td>
<td>246,668</td>
<td>—</td>
</tr>
<tr>
<td>Cash and cash equivalents</td>
<td>$553,398</td>
<td>$78,823</td>
</tr>
</tbody>
</table>

Cash equivalents are recorded at cost, which approximate fair value due to their short-term nature.

 Marketable Securities

Marketable securities consist of debt securities with maturities greater than three months from the date of purchase that have historically included commercial paper, government-backed securities, treasury securities, corporate notes and agency securities. Classification of marketable securities between current and non-current is dependent upon the maturity date at the balance sheet date taking into consideration the Company’s ability and intent to hold the investment to maturity.

Interest and dividend income are recorded when earned and included in investment income in the consolidated statements of operations. Premiums and discounts, if any, on marketable securities are amortized or accreted to maturity and included in investment income in the consolidated statements of operations. The specific identification method is used in computing realized gains and losses on the sale of the Company’s securities.

The Company classifies its marketable securities with readily determinable fair values as "available-for-sale." Investments in securities that are classified as available-for-sale are measured at fair market value in the consolidated balance sheets, and unrealized gains and losses on marketable securities are reported as a separate component of stockholders’ equity (deficit) until realized. Marketable securities are evaluated periodically to determine whether a decline in value is "other-than-temporary." The term "other-than-temporary" is not intended to indicate a permanent decline in value. Rather, it means that the prospects for a near-term recovery of value are not necessarily unfavorable, or that there is a lack of evidence to support fair values equal to, or greater than, the carrying value of the security. Management reviews criteria, such as the magnitude and duration of the decline, as well as the Company’s ability to hold the securities, including whether the Company will be required to sell a security prior to recovery of its amortized cost basis, the investment issuer’s financial condition and business outlook to predict whether the loss in value is other-than-temporary. If a decline in value is determined to be other-than-temporary, the value of the security is reduced, and the impairment is recorded as other income (expense) in the consolidated statements of operations.

Concentration of Credit Risk

Financial instruments expose the Company to concentration of credit risk and consist primarily of cash and cash equivalents and marketable securities. The Company’s investment policy limits investments to certain types of instruments, including asset-backed securities, high-grade corporate debt securities and money market funds, places restrictions on maturities and concentrations in certain industries and requires the Company to maintain a certain level of liquidity. At times, the Company maintains cash balances in financial institutions, which may exceed federally insured limits. The Company has not experienced any losses relating to such accounts and believes it is not exposed to a significant credit risk on its cash and cash equivalents.

Fair Value Measurements

The Company applies Accounting Standards Codification ("ASC") Topic 820, Fair Value Measurements and Disclosures ("ASC 820"), for financial and non-financial assets and liabilities.

ASC 820 discusses valuation techniques, such as the market approach (comparable market prices), the income approach (present value of future income or cash flow) and the cost approach (cost to replace the service capacity of an asset or replacement cost). The statement utilizes a fair value hierarchy that prioritizes the inputs to valuation techniques used to measure fair value into three broad levels. The following is a brief description of those three levels:

- Level 1: Observable inputs such as quoted prices (unadjusted) in active markets for identical assets or liabilities.
- Level 2: Inputs other than quoted prices that are observable for the asset or liability, either directly or indirectly. These include quoted prices for similar assets or liabilities in active markets and quoted prices for identical or similar assets or liabilities in markets that are not active.
- Level 3: Unobservable inputs that reflect the reporting entity’s own assumptions.

Restricted Cash

The Company’s current and non-current restricted cash includes payments received under the Coalition for Epidemic Preparedness Innovations ("CEPI") funding agreements (see Note 8), payments received under the Bill & Melinda Gates Foundation ("BMGF") grant agreements (see Note 8), escrow funds paid in connection with the acquisition of Novavax CZ (see Note 6), escrow funds received in connection with a sale of assets transaction in 2019, and cash collateral accounts under letters of credit that serve as security deposits for certain facility leases. The Company will utilize the CEPI and BMGF funds as it incurs expenses for services performed under these agreements.

As of December 31, 2020, the restricted cash balances (both current and non-current) consisted of $1.5 million for payments received from BMGF, $92.4 million of payments under the CEPI funding agreements, and $1.5 million of security deposits. As of December 31, 2019, the restricted cash balances (both current and non-current) consisted of $1.4 million for...
payments received from BMGF, $1.5 million held in escrow received in connection with the sale of assets transaction and $0.4 million of security deposits.

The following table provides a reconciliation of cash, cash equivalents and restricted cash reported within the consolidated balance sheets that sum to the total of the same such amounts shown in the statement of cash flows at December 31 (in thousands):

<table>
<thead>
<tr>
<th></th>
<th>2020</th>
<th>2019</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cash and cash equivalents</td>
<td>$ 553,398</td>
<td>$ 78,823</td>
</tr>
<tr>
<td>Restricted cash current</td>
<td>93,880</td>
<td>2,947</td>
</tr>
<tr>
<td>Restricted cash non-current</td>
<td>1,460</td>
<td>410</td>
</tr>
<tr>
<td>Cash, cash equivalents and restricted cash</td>
<td>$ 648,738</td>
<td>$ 82,180</td>
</tr>
</tbody>
</table>

**Property and Equipment**

Property and equipment are stated at cost and are depreciated using the straight-line method over the estimated useful lives of the assets, generally three to twenty-five years. Amortization of leasehold improvements is computed using the straight-line method over the shorter of the estimated useful lives of the improvements or the remaining term of the lease. Repairs and maintenance costs are expensed as incurred.

**Lease Accounting**

The Company determines at the inception or modification of a contract if an arrangement is, or contains, a lease, which exists when the contract conveys the right to control the use of identified property or equipment for a period of time in exchange for consideration. In determining if a contract contains a lease, the Company evaluates whether the contract, either explicitly or implicitly, is for the use of an identified asset and the Company has the right to direct the use of, and obtain substantially all of the benefit from, the identified asset. Depending on the contract, the lease commencement date, defined as the date on which the lessor makes the underlying asset available for use by the lessee and is the date on which the Company is required to accrue lease expenses, may be different than the inception date of the contract. The Company evaluates changes to the terms and conditions of a lease contract to determine if they result in a new lease or a modification of an existing lease. For lease modifications, the Company reassesses the lease classification at the effective date of the modification. Leases are classified as either operating or finance leases based on the economic substance of the agreement.

The Company enters into non-cancelable lease agreements for facilities and certain equipment. Further, the Company enters into manufacturing supply agreements with contract manufacturing organizations and contract development and manufacturing organizations to manufacture its vaccine candidates. Certain of these manufacturing supply agreements include the use of identified manufacturing facilities and equipment that are controlled by the Company and, if the Company receives substantially all of the output of the underlying assets, qualify as an embedded lease. Manufacturing supply agreements that contain a lease are treated as lease arrangements in their entirety.

For leases that have a lease term of more than 12 months at the lease commencement date, the Company recognizes lease liabilities, which represent the Company’s obligation to make lease payments arising from the lease, and corresponding right-of-use (“ROU”) assets, which represent the right to use an underlying asset for the lease term, based on the present value of the fixed future payments over the lease term. The Company calculates the present value of future payments using the discount rate implicit in the lease, if available, or the Company’s incremental borrowing rate. For all leases that have a lease term of 12 months or less at the commencement date (referred to as “short-term” leases), the Company has elected to apply the practical expedient in ASC Topic 842, *Leases* (“ASC 842”), to not recognize a lease liability or ROU asset but instead, recognize lease payments as an expense on a straight-line basis over the lease term and variable lease payments that do not depend on an index or rate, as an expense in the period in which the variable lease costs are incurred based on performance or usage in accordance with contractual agreements. In determining the lease period, the Company evaluates facts and circumstances that could affect the period over which it is reasonably certain to use the underlying asset while taking into consideration the non-cancelable period over which it has the right to use the underlying asset and any option period to extend or terminate the lease if it is reasonably certain to exercise the option. The Company re-evaluates short-term leases that are modified and if they no longer meet the requirements to be treated as short-term leases, recognizes and measures the lease liability and ROU asset as if the date of the modification is the lease commencement date.

For operating leases, the Company recognizes lease expense related to fixed payments on a straight-line basis over the lease term and lease expense related to variable payments as incurred based on performance or usage in accordance with the contractual agreements. For finance leases, the Company recognizes the amortization of the ROU asset over the shorter of the lease term or useful life of the underlying asset. The Company expenses ROU assets acquired for research and development activities under ASC Topic 730, *Research and Development*, if they do not have an alternative future use, in research and development projects or otherwise.

The Company uses significant assumptions and judgment in evaluating its lease contracts and other agreements under ASC 842, including the determination of whether an agreement is or contains a lease, whether a change in the terms and conditions of a lease contract represent a new or modified lease, whether a lease represents an operating or finance lease, the discount rate used to determine the present value of lease obligations and the term of a lease embedded in its manufacturing supply agreements.

**Revenue**

The Company performs research and development under government funding, grant, license and clinical development agreements. The revenue primarily consists of funding under U.S. government contracts and other arrangements to advance the clinical development and manufacturing of NVX-CoV2373. The Company’s U.S. government contracts are with the U.S. Department of Defense (the “DoD”) and its participation in formerly known as Operation Warp Speed (“OWS”) (see Note 8). Other funding arrangements primarily include a grant and forgivable loan funding from CEPI (see Note 8).

At contract inception, the Company analyzes the revenue arrangement to determine the appropriate accounting under U.S. GAAP. Currently, the Company’s revenue arrangements represent customer contracts within the scope of ASC 606, *Revenue from Contracts with Customers (Topic 606)* (“ASC 606”) or are subject to the contribution guidance in ASC Topic 958-605, Not-for-Profit Entities – Revenue Recognition (“ASC 958-605”), which applies to business entities that receive contributions within the scope of ASC 958-605. The Company recognizes revenue from arrangements within the scope of ASC 606 following the five-step model: (i) identify the contract(s) with a customer; (ii) identify the performance obligation(s) in the contract; (iii) determine the transaction price; (iv) allocate the transaction price to the performance obligation(s) in the contract; and (v) recognize revenue when (or as) it satisfies a performance obligation. The Company only applies the five-step model to contracts when it is probable that it will collect the consideration it is entitled to in exchange for the goods or services it transfers to its customer. The Company recognizes contribution revenue within the scope of ASC 958-605 when the funder-imposed conditions have been substantially met. Contributions are recorded as deferred revenue until the period in which research and development activities are performed that satisfy the funder-imposed conditions.

Under the U.S. government contracts, the Company is entitled to receive funding of up to $1.8 billion, on a reimbursable-cost or reimbursable-cost-plus-fixed-fee basis, to support certain activities related to the development, manufacture and delivery of NVX-CoV2373 to the U.S. government. The Company analyzed these contracts and determined that they are within the scope of ASC 606. The obligations under each of the contracts are not distinct in the context of the contract as they are highly interdependent or interrelated and, as such, they are accounted for as a single performance obligation. The transaction price under these arrangements is the consideration the Company is expecting to receive and consists of the funded contract amount and the unfunded variable amount to the extent that it is probable that a significant reversal of revenue will not occur. The Company recognizes revenue for these contracts over time as the Company transfers control over the goods and services and satisfies the performance obligation. The Company measures progress toward satisfaction of the performance obligation using an Estimate-at-Completion (“EAC”) process, which is a cost-based input method that reviews and monitors the progress towards the completion of the contract. Under this process, management considers the costs that have been incurred to-date, as well as projections to completion using various inputs and assumptions, including, but not limited to, progress towards completion, labor costs and level of effort, material and subcontractor costs, indirect administrative costs and other...
identified risks. Estimating the total allowable cost at completion of the performance obligation under a contract is subjective and requires the Company to make assumptions about future activity and cost drivers. Changes in these estimates can occur for a variety of reasons and, if significant, may impact the timing of revenue and fee recognition on the Company’s contracts. Allowable contract costs include direct costs incurred on the contract and indirect costs that are applied in the form of rates to the direct costs. Progress billings under the contracts are initially based on provisional indirect billing rates, agreed upon between the Company and the U.S. government. These indirect rates are subject to audit on an annual basis. The Company records the impact of changes in the indirect billing rates in the period when such changes are identified. These changes reflect the difference between actual indirect costs incurred compared to the estimated amounts used to determine the provisional indirect billing rates agreed upon with the U.S. government. The Company recognizes revenue on the U.S. government contracts based on reimbursable allowable contract costs incurred in the period up to the transaction price. For reimbursable-cost-plus-fixed-fee contracts, the Company recognizes the fixed-fee based on the proportion of reimbursable contract costs incurred to total estimated allowable contract costs expected to be incurred on completion of the underlying performance obligation as determined by the EAC process. The Company recognizes changes in estimates related to the EAC process in the period when such changes are made on a cumulative catch-up basis. The Company includes the transaction price comprising both funded and unfunded portions of customer contracts, in this estimate.

The Company’s other funding agreements currently include funding from CEPI of $399.5 million in the form of a grant of $257.0 million (“CEPI Grant Funding”) and one or more forgivable no interest term loans of $142.5 million (“CEPI Forgivable Loan Funding”). The Company’s arrangements, including the CEPI Grant Funding and CEPI Forgivable Loan Funding, are unfunded and entitled to reimbursement for costs that support development related activities of NVX-CoV2373. The CEPI Forgivable Loan Funding is designated for the prepayment of certain manufacturing activities. The Company analyzed these other funding arrangements and determined that they are not within the scope of ASC 606 as they do not provide a direct economic benefit to the grantor. Payments received under the grant funding arrangements are considered conditional contributions under the scope of ASC 958-605 and are recorded as deferred revenue until the period in which such research and development activities are actually performed that satisfy the funder-imposed conditions. Payments received under the CEPI Forgivable Loan Funding agreements are only repayable if the proceeds of sales to one or more third-parties of NVX-CoV2373 cover the Company’s costs of manufacturing such vaccine candidate, not including manufacturing costs funded by CEPI. As the financial risk remains with CEPI, the Company determined that the use of the CEPI Forgivable Loan Funding is outside the scope of ASC 470, Debt. The research and development risk is considered substantive, such that it is not yet probable that the development will be successful. Therefore, the Company has concluded that ASC 730 is considered applicable and most appropriate. Given the Company’s capital stock is the appropriate measurement of fair value, the Company considers factors such as its trading volume, diversity of investors and analyst coverage. If considered necessary, the income approach is used to corroborate the results of the market approach. Goodwill impairment may exist if the carrying value of the reporting unit exceeds its estimated fair value. If the carrying value of the reporting unit exceeds its fair value, step two of the impairment analysis is performed. In step two of the analysis, an impairment loss is recorded equal to the excess of the carrying value of the reporting unit’s goodwill over its implied fair value, should such a circumstance arise.

During 2020, the Company changed its annual goodwill impairment testing date from December 31 to October 1. Management has determined that the change in the testing date does not represent a material change to a method of applying an accounting principle as it does not have a material effect on the Company’s consolidated financial statements in light of the Company’s internal controls and requirements under ASC Topic 350, Intangibles—Goodwill and Other, to assess goodwill impairment upon certain triggering events.

At October 1, 2020 and December 31, 2019, the Company used the market approach to determine if the Company had an impairment of its goodwill. The fair value of the Company’s single reporting unit was substantially higher than its carrying value, resulting in no impairment to goodwill as of October 1, 2020 and December 31, 2019.

**Stock-Based Compensation**

The expected term of stock options and stock appreciation rights, restricted stock awards and purchases under the Company’s Employee Stock Purchase Plan, as amended and restated (the “ESPP”) at fair value. The Company recognizes compensation expense related to such awards on a straight-line basis over the requisite service period (generally the vesting period) of the equity awards, which typically occurs ratably over periods ranging from one year to four years.

The expected term of stock options and stock appreciation rights granted is based on the Company’s historical option exercise experience and post-vesting forfeitures. The expected term for purchases under the ESPP is based on the purchase periods included in the offering. The expected volatility is determined using historical volatilities based on stock prices over a look-back period corresponding to the expected term. The risk-free interest rate is determined using the yield available for zero-coupon U.S. government issues with a remaining term equal to the expected term. The Company has never paid a dividend, and as such, the dividend yield is zero, and the Company does not intend to pay dividends in the foreseeable future.
Restricted stock awards are recorded as compensation expense over the expected vesting period based on the fair value at the award date using the straight-line method of amortization.

See Note 13 for a further discussion on stock-based compensation.

Research and Development Expenses

Research and development expenses include salaries, stock-based compensation, laboratory supplies, consultants and subcontractors, including external contract research organizations ("CROs"), contract management organizations ("CMOs") and contract development and manufacturing organizations ("CDMOS") and other expenses associated with the Company’s process development, manufacturing, clinical, regulatory and quality assurance activities for its clinical development programs. In addition, related indirect costs such as fringe benefits and overhead expenses are also included in research and development expenses.

The Company estimates its research and development expense related to services performed under its contracts with external service providers based on an estimate of the level of service performed in the period. Research and development activities are expensed as incurred.

Accrued Research and Development Expenses

The Company accrues research and development expenses, including clinical trial-related expenses, as the services are performed, which may include estimates of those expenses incurred, but not invoiced. The Company uses information provided by third-party service providers and CROs, CMO’s and CDMO’s invoices and internal estimates to determine the progress of work performed on the Company’s behalf. Assumptions based on clinical trial protocols, contracts and participant enrollment data are also developed to determine and analyze these estimates and accruals.

Income Taxes

The Company accounts for income taxes in accordance with ASC Topic 740, Income Taxes. Under the liability method, deferred income taxes are recognized for the future tax consequences attributable to differences between the financial statement carrying amounts of existing assets and liabilities and their respective tax basis and operating loss carryforwards. Deferred tax assets and liabilities are measured using enacted tax rates expected to apply to taxable income in the year in which those temporary differences are expected to be recovered or settled. The effect of changes in tax rates on deferred tax assets and liabilities is recognized in income in the period such changes are enacted. A valuation allowance is established when necessary to reduce net deferred tax assets to the amount expected to be realized.

Tax benefits associated with uncertain tax positions are recognized in the period in which one of the following conditions is satisfied: (1) the more likely than not recognition threshold is satisfied; (2) the position is ultimately settled through negotiation or litigation; or (3) the statute of limitations for the taxing authority to examine and challenge the position has expired. Tax benefits associated with an uncertain tax position are reversed in the period in which the more likely than not recognition threshold is no longer satisfied.

Interest and penalties related to income tax matters are recorded as income tax expense. At December 31, 2020 and 2019, the Company had no accruals for interest or penalties related to income tax matters.

Net Loss per Share

Net loss per share is computed using the weighted average number of shares of common stock outstanding. At December 31, 2020, 2019 and 2018, the Company had outstanding stock options and unvested restricted stock awards totaling 6,679,629, 4,992,792 and 2,975,481 underlying shares of the Company’s common stock, respectively. At December 31, 2020 and 2019, the Company’s Notes (as defined in Note 11) would have been convertible into approximately 2,385,800 shares of the Company’s common stock assuming a common stock price of $136.20 or higher. These and any other shares due to the Company upon settlement of its capped call transactions are excluded from the computation, as their effect is antidilutive.

Foreign Currency

The accompanying consolidated financial statements are presented in U.S. dollars. The functional currency of Novavax AB, which is located in Sweden, is the local currency (Swedish Krona) and the functional currency of Novavax CZ, which is located in the Czech Republic, is the local currency (Czech Koruna). The translation of assets and liabilities of Novavax AB and Novavax CZ to U.S. dollars are made at the exchange rate in effect at the consolidated balance sheet date, while equity accounts are translated at historical rates. The translation of the statement of operations data is made at the average exchange rate in effect for the period. The translation of operating cash flow data is made at the average exchange rate in effect for the period, and investing and financing cash flow data is translated at the exchange rate in effect at the date of the underlying transaction. Translation gains and losses are recognized as a component of accumulated other comprehensive income (loss) in the accompanying consolidated balance sheets. The foreign currency translation adjustment balance included in accumulated other comprehensive income (loss) was $7.0 million and $(12.5) million at December 31, 2020 and 2019, respectively.

Segment Information

The Company manages its business as one operating segment: the development of recombinant vaccines. The Company does not operate separate lines of business with respect to its vaccine candidates. Accordingly, the Company does not have separately reportable segments as defined by ASC Topic 280, Segment Reporting.

Recent Accounting Pronouncements

Recently Adopted

In January 2017, the FASB issued Accounting Standards Update ("ASU") No. 2017-04, Intangibles—Goodwill and Other (Topic 350) ("ASU 2017-04"), which will simplify the goodwill impairment calculation by eliminating Step 2 from the current goodwill impairment test. The new standard does not change how a goodwill impairment is identified. The Company will continue to perform its quantitative goodwill impairment test by comparing the fair value of its reporting unit to its carrying amount, but if the Company is required to recognize a goodwill impairment charge, under the new standard, the amount of the charge will be calculated by subtracting the reporting unit’s fair value from its carrying amount. Under the current standard, if the Company is required to recognize a goodwill impairment charge, Step 2 requires it to calculate the implied value of goodwill by assigning the fair value of a reporting unit to all of its assets and liabilities as if that reporting unit had been acquired in a business combination and the amount of the charge is calculated by subtracting the reporting unit’s implied fair value of goodwill from the goodwill carrying amount. The standard was effective January 1, 2020 for the Company and will be applied prospectively from the date of adoption. The adoption of ASU 2017-04 did not have a material impact on the Company’s historical financial statements.

Not Yet Adopted

In August 2020, the FASB issued ASU No. 2020-06, Debt—Debt with Conversion and Other Options (Subtopic 470-20) and Derivatives and Hedging—Contracts in Entity’s Own Equity (Subtopic 815-40): Accounting for Convertible Instruments and Contracts in an Entity’s Own Equity ("ASU 2020-06"), which will simplify the accounting for certain financial instruments with characteristics of liabilities and equity, including certain convertible instruments and contracts on an entity’s own equity. Specifically, the new standard will remove the separation models required for convertible debt with cash conversion features and convertible instruments with beneficial conversion features. It will also remove certain settlement conditions that are currently required for equity contracts to qualify for the derivative scope exception and will simplify the diluted earnings per share calculation for convertible instruments. ASU 2020-06 will be effective January 1, 2022 for the Company and may be applied using a full or modified retrospective approach. Early adoption is permitted, but no earlier than January 1, 2021 for the Company. Management has evaluated the impact of adopting ASU 2020-06 and has determined such adoption will not have a material impact on the overall stockholders’ equity (deficit) in the Company’s consolidated financial statements.
Note 3 – Fair Value Measurements

The following table represents the estimated fair value of the Company’s financial assets and liabilities (in thousands):

<table>
<thead>
<tr>
<th>Assets</th>
<th>Fair Value at December 31, 2020</th>
<th>Fair Value at December 31, 2019</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Level 1</td>
<td>Level 2</td>
</tr>
<tr>
<td>Money market funds(1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Government-backed securities(2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Treasury securities(3)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Corporate debt securities(4)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Agency securities</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total cash equivalents and</td>
<td>$96,116</td>
<td>$407,238</td>
</tr>
<tr>
<td>marketable securities</td>
<td>$63,156</td>
<td>$24,250</td>
</tr>
<tr>
<td>Liabilities</td>
<td>$—</td>
<td>$—</td>
</tr>
</tbody>
</table>

(1) Classified as cash and cash equivalents as of December 31, 2020 and 2019, respectively (see Note 2).
(2) Includes $44,250 and $20,000 classified as cash and cash equivalents as of December 31, 2020 and 2019, respectively, on the consolidated balance sheets.
(3) Includes $44,052 classified as cash and cash equivalents as of December 31, 2020 on the consolidated balance sheets.
(4) Includes $246,668 classified as cash and cash equivalents as of December 31, 2020 on the consolidated balance sheets.

Fixed-income investments categorized as Level 2 are valued at the custodian bank by a third-party pricing vendor’s valuation models that use verifiable observable market data, e.g., interest rates and yield curves observable at commonly quoted intervals and credit spreads, bids provided by brokers or dealers or quoted prices of securities with similar characteristics. Pricing of the Company’s Notes (as defined in Note 11) has been estimated using other observable inputs, including the price of the Company’s common stock, implied volatility, interest rates and credit spreads among others.

During the years ended December 31, 2020 and 2019, the Company did not have any transfers between Levels.

The amount in the Company’s consolidated balance sheets for accounts payable and accrued expenses approximates its fair value due to its short-term nature.

Note 4 – Marketable Securities

Marketable securities classified as available-for-sale as of December 31, 2020 and 2019 were comprised of (in thousands):

<table>
<thead>
<tr>
<th></th>
<th>Amortized Cost</th>
<th>Gross Unrealized Gains</th>
<th>Gross Unrealized Losses</th>
<th>Fair Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Treasury securities</td>
<td>$10,038</td>
<td>$—</td>
<td>$(2)</td>
<td>$10,036</td>
</tr>
<tr>
<td>Corporate debt securities</td>
<td>127,003</td>
<td>13</td>
<td>(3)</td>
<td>127,013</td>
</tr>
<tr>
<td>Agency securities</td>
<td>20,599</td>
<td>1</td>
<td>—</td>
<td>20,600</td>
</tr>
<tr>
<td>Total</td>
<td>$157,640</td>
<td>$14</td>
<td>$(5)</td>
<td>$157,649</td>
</tr>
</tbody>
</table>

As of December 31, 2020, investments in marketable securities, including corporate debt securities, were due to mature within one year.

Note 5 – Goodwill and Other Intangible Assets

Goodwill

The changes in the carrying amounts of goodwill for the years ended December 31, 2020 and 2019 were as follows (in thousands):

<table>
<thead>
<tr>
<th>Year Ended December 31,</th>
<th>2020</th>
<th>2019</th>
</tr>
</thead>
<tbody>
<tr>
<td>Beginning balance</td>
<td>$51,154</td>
<td>$51,967</td>
</tr>
<tr>
<td>Goodwill resulting from the acquisition of Novavax CZ</td>
<td>70,662</td>
<td>—</td>
</tr>
<tr>
<td>Currency translation adjustments</td>
<td>13,563</td>
<td>(813)</td>
</tr>
<tr>
<td>Ending balance</td>
<td>$135,379</td>
<td>$51,154</td>
</tr>
</tbody>
</table>

Identifiable Intangible Assets

Purchased intangible assets consisted of the following as of December 31, 2020 and 2019 (in thousands):

<table>
<thead>
<tr>
<th></th>
<th>Gross Carrying Amount</th>
<th>Accumulated Amortization</th>
<th>Intangible Assets, Net</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>December 31, 2020</td>
<td>December 31, 2019</td>
<td>December 31, 2020</td>
</tr>
<tr>
<td></td>
<td>$13,208</td>
<td>$(7,483)</td>
<td>$5,725</td>
</tr>
</tbody>
</table>

Amortization expense for the years ended December 2020, 2019 and 2018 was $0.6 million, $0.7 million and $0.7 million, respectively. Estimated amortization expense for existing intangible assets for each of the five succeeding years ending December 31, 2023, is as follows (in thousands):

<table>
<thead>
<tr>
<th>Year</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>2021</td>
<td>$455</td>
</tr>
<tr>
<td>2022</td>
<td>$455</td>
</tr>
<tr>
<td>2023</td>
<td>$455</td>
</tr>
<tr>
<td>2024</td>
<td>$455</td>
</tr>
<tr>
<td>2025</td>
<td>$455</td>
</tr>
</tbody>
</table>

Note 6 - Acquisition of Novavax CZ

On May 27, 2020 (the “Acquisition Date”), the Company entered into a Share Purchase Agreement (the “Deed”) by and among Novavax AB, the Company’s wholly-owned Swedish subsidiary (the “Buyer”), and De Bilt Holdings B.V., Poonawalla Science Park B.V., and Bilthoven Biologicals B.V. (collectively, the “Sellers”) and, solely as guarantors, each of Serum International B.V. and the Company. Pursuant to the terms and conditions of the Deed, the Buyer acquired all the issued and outstanding shares of Novavax CZ (formerly Praha Vaccines a.s.), a vaccine manufacturing company (the “Acquisition”). The assets of Novavax CZ acquired as part of the Acquisition include a biologics manufacturing facility and associated assets in Bohumil, Czech Republic and will be used by the Company to expand its manufacturing capacity.
The Company has accounted for the Acquisition as a business combination using the acquisition method of accounting, with the Company as the acquirer. The acquisition method requires the Company to record the assets acquired and liabilities assumed at fair value. The amount by which the purchase price exceeds the fair value of net assets acquired is recorded as goodwill. The Company completed the appraisal process necessary to assess the fair values of the assets acquired and liabilities assumed to determine the amount of goodwill to be recognized as of the Acquisition Date. The final determination of the fair value of all assets and liabilities is presented in the table below.

The fair value of the assets acquired, and liabilities assumed were determined using market and cost valuation methodologies. The fair value measurements were based on significant unobservable inputs that were developed by the Company using publicly available information, market participant assumptions, and cost and development assumptions. Because of the use of significant unobservable inputs, the fair value measurements represent a Level 3 measurement as defined in ASC 820. The market approach is a valuation technique that uses prices and other relevant information generated by market transactions involving identical or comparable assets, liabilities, or a group of assets or liabilities. The cost approach estimates value by determining the current cost of replacing an asset with another of equivalent utility. The cost to replace a given asset reflects the estimated reproduction or replacement cost for the property, less an allowance for loss in value due to depreciation.

The cost approach was the primary approach used to value fixed assets, including the real property. Fixed assets are depreciated on a straight-line basis over their expected remaining useful lives, ranging from four years to 25 years.

The Company recorded $70.7 million in goodwill related to the Acquisition representing the Purchase Price that was in excess of the fair value of the assets acquired and liabilities assumed. The goodwill generated from the Acquisition is not expected to be deductible for U.S. federal income tax purposes. The goodwill recognized is attributable to intangible assets that do not exceed the fair value of the assets acquired and liabilities assumed (in thousands):
At December 31, 2020, the facility leases, excluding the 700QO lease, have expirations that range from approximately three to six years, some of which include options to extend the leases or terminate the leases early. Options to extend the leases or terminate the leases early are only included in the lease term when it is reasonably certain that the option will be exercised. The facility leases contain provisions for future rent increases and obligate the Company to pay building operating costs. The Company records operating lease expense for each of its operating leases on a straight-line basis from lease commencement date through the end of the lease term.

Supplemental balance sheet information related to leases as of December 31, 2020 was as follows (in thousands, except weighted-average remaining lease term and discount rate):

<table>
<thead>
<tr>
<th>Lease Assets and Liabilities</th>
<th>Classification</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>Assets:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ROU assets, operating, net</td>
<td>Other non-current assets</td>
<td>$7,794</td>
</tr>
<tr>
<td>Liabilities:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Current portion of operating lease liabilities</td>
<td>Other current liabilities</td>
<td>$3,782</td>
</tr>
<tr>
<td>Current portion of finance lease liabilities</td>
<td>Current portion of finance lease liabilities</td>
<td>105,862</td>
</tr>
<tr>
<td>Total current lease liabilities</td>
<td></td>
<td>$109,644</td>
</tr>
<tr>
<td>Non-current portion of operating lease liabilities</td>
<td>Other non-current liabilities</td>
<td>$10,122</td>
</tr>
<tr>
<td>Non-current portion of finance lease liabilities</td>
<td>Non-current finance liabilities</td>
<td>40,083</td>
</tr>
<tr>
<td>Total non-current lease liabilities</td>
<td></td>
<td>$50,205</td>
</tr>
</tbody>
</table>

Weighted-average remaining lease term (years):
- Operating leases: 4.5
- Finance leases: 4.7

Weighted-average discount rate:
- Operating leases: 13.8%
- Finance leases: 6.4%

Lease expense for the operating and short-term leases for the year ended December 31 was as follows (in thousands):

<table>
<thead>
<tr>
<th>Year</th>
<th>Operating lease expense</th>
<th>Short-term lease expense</th>
<th>ROU assets expense</th>
<th>Interest expense</th>
<th>Total finance lease expense</th>
</tr>
</thead>
<tbody>
<tr>
<td>2020</td>
<td>$2,462</td>
<td>66,805</td>
<td>242,009</td>
<td>3,097</td>
<td>$245,106</td>
</tr>
</tbody>
</table>

Supplemental cash flow information related to leases for the year ended December 31, 2020 was as follows (in thousands):

<table>
<thead>
<tr>
<th>Amount</th>
<th>Operating cash flows used in operating leases</th>
<th>Operating cash flows used in finance leases</th>
<th>Financing cash flows used in finance leases</th>
</tr>
</thead>
<tbody>
<tr>
<td>$63,634</td>
<td>3,097</td>
<td>96,065</td>
<td></td>
</tr>
</tbody>
</table>

ROU assets obtained in exchange for operating lease obligations: $5,590
ROU assets obtained in exchange for finance lease obligations: 242,009

As of December 31, 2020, maturities of lease liabilities were as follows (in thousands):

<table>
<thead>
<tr>
<th>Year</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>2021</td>
<td>$118,017</td>
</tr>
<tr>
<td>2022</td>
<td>44,692</td>
</tr>
<tr>
<td>2023</td>
<td>3,385</td>
</tr>
<tr>
<td>2024</td>
<td>2,393</td>
</tr>
<tr>
<td>2025</td>
<td>2,369</td>
</tr>
<tr>
<td>Thereafter</td>
<td>2,079</td>
</tr>
<tr>
<td>Total minimum lease payments</td>
<td>172,935</td>
</tr>
<tr>
<td>Less: imputed interest</td>
<td>(13,086)</td>
</tr>
<tr>
<td>Total lease liabilities</td>
<td>$159,849</td>
</tr>
</tbody>
</table>

Note 8 – U.S. Government Contracts, Grants and Other Revenue Arrangements

U.S. Government Contracts

Operation Warp Speed

In July 2020, the Company entered into a Project Agreement (the “Project Agreement”) with Advanced Technology International, Inc. (“ATI”), the Consortium Management Firm acting on behalf of the Medical CBRN Defense Consortium in connection with OWs. OWS is a partnership among components of the U.S. Department of Health and Human Services and the U.S. Department of Defense working to accelerate the development, manufacturing and distribution of COVID-19 vaccines, therapeutics and diagnostics. The Project Agreement, which was last amended in December 2020, relates to the Base Agreement the Company entered into with ATI in June 2020 (the “Base Agreement,” together with the Project Agreement, the “OWS Agreement”). Under the OWS Agreement, the Company is entitled to receive funding of up to $1.7 billion to support certain activities related to the development of NVX-CoV2373 and the manufacture and delivery of the vaccine candidate to the U.S. government. Pursuant to the OWS Agreement, the Company is authorized to make expenditures or incur obligations of up to $1.6 billion.

The OWS Agreement requires the Company to conduct certain clinical, regulatory and other activities, including a pivotal Phase 3 clinical trial to determine the safety and efficacy of NVX-CoV2373, and to manufacture and deliver to the U.S. government 100 million doses of the vaccine candidate. Funding under the OWS Agreement is payable to the Company for various development, clinical trial, manufacturing, regulatory and other activities. The OWS Agreement contains terms and conditions that are customary for U.S. government agreements of this nature, including provisions giving the U.S. government the right to terminate the Base Agreement and/or the Project Agreement based on a reasonable determination that the funded project will not produce beneficial results commensurate with the expenditure of resources and that termination would be in the
U.S. government’s interest. If the Project Agreement is terminated prior to completion, the Company is entitled to be paid for work performed and costs or obligations incurred prior to termination and consistent with the terms of the OWS Agreement. The performance period under the Project Agreement extends from July 2020 through December 2021, subject to early termination by the U.S. government or extension by mutual agreement of the parties. In 2020, the Company recognized revenue under the OWS Agreement of $204.7 million.

U.S. Department of Defense

In June 2020, the Company entered into a letter contract that was last amended in January 2021 (the “DoD Contract”) with the DoD Joint Program Executive Office for Chemical, Biological, Radiological and Nuclear Defense (“JPEO-CBRND-EB”), under which JPEO-CBRND-EB agreed to provide funding of up to $45.7 million to the Company to support the manufacture of NVX-CoV2373. Under the DoD Contract, the Company is authorized to make expenditures or incur obligations up to the full amount of the funding.

Under the DoD Contract, the Company is expected to deliver 10 million doses of NVX-CoV2373 to the DoD. The 10 million doses of NVX-CoV2373 may be used in Phase 2/3 clinical trials or under an EUA, if approved by the U.S. Food and Drug Administration (“FDA”). Pursuant to the DoD Contract, if NVX-CoV2373 is approved by the FDA, the DoD is entitled to most-favored customer status for a period of five years from the award of the DoD Contract, meaning that the Company cannot give any comparable commercial client in the United States more favorable pricing than the DoD under similar transactional circumstances. In 2020, the Company recognized revenue from the DoD Contract of $12.5 million.

Grants and Other Revenue Arrangements

Coalition for Epidemic Preparedness Innovations

In May 2020, the Company entered into a restated funding agreement which was amended in November 2020 (the “CEPI Funding Agreement”) with CEPI, under which CEPI agreed to provide funding of up to $399.5 million to the Company to support the development of NVX-CoV2373. The CEPI Funding Agreement provides up to $257.0 million in Grant Funding and up to $142.5 million in Forgivable Loan Funding, which loans are in the form of one or more forgivable no interest term loans in order to prepay certain manufacturing activities and are not subject to restrictive or financial covenants. The Company is only required to repay any CEPI Forgivable Loan Funding under certain circumstances to the extent it sells doses of NVX-CoV2373, produced with the funds provided and included in such loans(s), to a third-party.

Under the terms of the CEPI Funding Agreement, among other things, the Company and CEPI agreed on the importance of global equitable access to any vaccines produced pursuant to the CEPI Funding Agreement. Any such vaccines, if approved, are expected to be procured and allocated through global mechanisms under discussion as part of the Access to COVID-19 Tools (ACT) Accelerator, an international initiative launched by the World Health Organization (“WHO”), Gavi the Vaccine Alliance, CEPI and other global non-governmental organizations and governmental leaders in 2020.

The scope and continuation of the CEPI Funding Agreement may be amended depending on ongoing developments of the COVID-19 outbreak and the success of NVX-CoV2373 relative to other third-party COVID-19 vaccine candidates or treatments. If the WHO, CEPI or a regulatory authority having jurisdiction over a clinical trial of NVX-CoV2373 determines that a third-party product candidate has substantially greater potential than a Company’s product candidate, the Company must cease its clinical trial in the relevant region and will be reimbursed for any costs incurred as a result thereof. In addition, CEPI has the right to unilaterally terminate the CEPI Funding Agreement if CEPI reasonably determines that (i) there are material safety, regulatory or ethical issues with the development of NVX-CoV2373, (ii) NVX-CoV2373 development should be limited in scope or terminated, (iii) the Company becomes unable to discharge its obligations under the agreement, (iv) the Company fails to meet certain milestones, or (v) the Company commits fraud or a financial irregularity.

Payments received in advance that are related to future performance are deferred and recognized as revenue when the research and development activities are performed. Cash payments received under the CEPI Funding Agreement are restricted as to their use until expenditures contemplated in the funding agreements are incurred. In 2020, the Company recognized revenue of $222.8 million under the CEPI Funding Agreement.

Bill & Melinda Gates Foundation

In support of the Company’s development of ResVax™, in September 2015, the Company entered into the grant agreement with BMGF (the “BMGF Grant Agreement”), under which it was awarded a grant totaling up to $89.1 million (the “Grant”). The Grant supports ResVax development activities, including the Company’s global Phase 3 clinical trial in pregnant women in their third trimester and other regulatory efforts. Unless terminated earlier by BMGF, the BMGF Grant Agreement will continue in effect until the end of 2021. The Company concurrently entered into a Global Access Commitments Agreement (“GACA”) with BMGF as a part of the BMGF Grant Agreement. Under the terms of the GACA, among other things, the Company agreed to make a certain amount of ResVax available and accessible at affordable pricing to people in certain low- and middle-income countries. Unless terminated earlier by BMGF, the GACA will continue in effect until the later of 15 years from its effective date, or 10 years after the first sale of a product under defined circumstances. The term of the GACA may be extended in certain circumstances, by a period of up to five additional years.

In July 2020, the Company entered into a grant agreement with BMGF (the “BMGF SA Grant Agreement”) under which it was awarded a grant of $15.0 million to support a Phase 2b clinical trial in the Republic of South Africa to evaluate the safety, immunogenicity, and potential efficacy of NVX-CoV2373.

Payments received in advance that are related to future performance are deferred and recognized as revenue when the research and development activities are performed. Cash payments received under the BMGF Grant Agreement and the BMGF SA Grant Agreement are restricted as to their use until expenditures contemplated in the agreements are incurred. In 2020, the Company recognized revenue from the BMGF Grant Agreement of $0.4 million and has recognized approximately $82 million in revenue since the inception of the agreement. In 2020, the Company recognized revenue from the BMGF SA Grant Agreement of $12.4 million.

Serum Institute of India Private Limited

In July 2020, the Company entered into a supply and license agreement with Serum Institute of India Private Limited ("SIPL"), as amended by the parties in September 2020, under which the Company granted exclusive and non-exclusive licenses to SIPL for the development, co-formulation, filling and finishing, registration and commercialization by SIPL of NVX-CoV2373. SIPL has agreed to purchase Matrix-M adjuvant from the Company and the Company has granted SIPL a non-exclusive license to manufacture the antigen drug substance component of NVX-CoV2373 in SIPL’s licensed territory solely for use in the manufacture of NVX-CoV2373 under the terms of the agreement. The parties will equally split the revenue from sale of NVX-CoV2373 by SIPL in its licensed territory, net of agreed costs. The Company granted to SIPL (i) an exclusive license in India during the agreement, and (ii) a non-exclusive license (a) during the “Pandemic Period” (as declared by the World Health Organization), in all countries other than specified countries designated by the World Bank as upper-middle or high-income countries, with respect to which the Company retains rights, and (b) after the Pandemic Period, in only those countries designated as low or middle-income by the World Bank. Following the Pandemic Period, the Company may notify SIPL of any bona fide opportunities for the Company to license NVX-CoV2373 to a third-party in such low and middle-income countries and SIPL would have an opportunity to match or improve such third-party terms, failing which, the Company would have the discretion to remove one or more non-exclusive countries from SIPL’s license.

Takeda Pharmaceutical Company Limited

In August 2020, the Company announced a collaboration agreement with Takeda Pharmaceutical Company Limited ("Takeda") for the exclusive development, manufacturing and commercialization of NVX-CoV2373 in Japan. Takeda will receive funding from the Government of Japan’s Ministry of Health, Labour and Welfare to support the technology transfer, establishment of infrastructure and scale-up of manufacturing. The collaboration agreement was finalized in February 2021. The Company will be entitled to receive payments based on the achievement of certain development and commercial milestones, as well as a portion
of net profits from the sale of the vaccine. In 2020, the Company recognized other revenue as a result of achieving a development milestone from the Takeda arrangement of $20.0 million.

Vaccine Supply Advance Purchase Agreements

In October 2020, the Company entered into a SARS-CoV-2 vaccine supply agreement with The Secretary of State for Business, Energy and Industrial Strategy, acting on behalf of the government of the UK, the purchase of up to 60 million doses of NVX-CoV2373, plus such additional orders as the Authority may make from time to time. The Company agreed to continue to conduct a UK-based Phase 3 clinical trial of NVX-CoV2373 to assess the efficacy of NVX-CoV2373 in the UK population, establish a dedicated supply chain for NVX-CoV2373 in the UK and seek regulatory approval for NVX-CoV2373 in the UK.

In December 2020, the Company finalized the advance purchase agreement with the Australian Federal Government to supply 51 million doses of NVX-CoV2373. The Company will work with Australia’s regulatory agency, the Therapeutics Goods Administration (“TGA”), to obtain product approvals upon demonstrating efficacy in clinical studies. As part of the agreement, Australia will have the option to purchase up to an additional 10 million doses. Further, in December 2020, the Company finalized an advance purchase agreement with the government of New Zealand for the purchase of 10.7 million doses of NVX-CoV2373.

Under the terms of the Company’s advance purchase agreements, government counterparties make upfront payments and have certain termination rights, or rights to reduce or cancel orders, if regulatory approval for the vaccine is not received or if supply is materially interrupted, delayed or deferred. The Company expects to record such upfront payments as deferred revenue and have certain termination rights, or rights to reduce or cancel orders, if regulatory approval for the vaccine is not received or if supply is materially interrupted, delayed or deferred. The Company expects to record such upfront payments as deferred revenue and anticipates recognizing revenue when the vaccine is delivered to its customers.

Note 9 – Preferred Stock

In June 2020, the Company entered into a redeemable Series A Convertible Preferred Stock Subscription Agreement, pursuant to which the Company agreed to issue and sell in a private placement 438,885 shares of its newly designated redeemable Series A Convertible Preferred Stock, par value $0.01 per share (“Preferred Stock”), at a purchase price of $455.70 per share, for total gross proceeds of $200.0 million. During the fourth quarter of 2020, all outstanding shares of Preferred Stock were converted and the Company issued 4,388,850 shares of common stock, par value $0.01 per share and reclassified $199.8 million from preferred stock to additional paid-in capital. The Company recognized a benefit conversion feature of approximately $24.1 million at the time of issuance of the Preferred Stock that was recorded in additional paid-in capital and accumulated deficit as the Preferred Stock issuance was contingent upon recognition and convertibility at any time at the option of the holder.

Note 10 – Other Financial Information

Prepaid Expenses and Other Current Assets

Prepaid expenses and other current assets consist of the following at December 31 (in thousands):

<table>
<thead>
<tr>
<th></th>
<th>2020</th>
<th>2019</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prepaid expenses</td>
<td>$171,602</td>
<td>$3,601</td>
</tr>
<tr>
<td>Other current assets</td>
<td>9,662</td>
<td>4,376</td>
</tr>
<tr>
<td>Prepaid expenses and other current assets</td>
<td>$181,264</td>
<td>$7,977</td>
</tr>
</tbody>
</table>

Property and Equipment, net

Property and equipment is comprised of the following at December 31 (in thousands):

<table>
<thead>
<tr>
<th></th>
<th>2020</th>
<th>2019</th>
</tr>
</thead>
<tbody>
<tr>
<td>Land and buildings</td>
<td>$79,096</td>
<td>—</td>
</tr>
<tr>
<td>Machinery and equipment</td>
<td>31,609</td>
<td>9,946</td>
</tr>
<tr>
<td>Leasohold improvements</td>
<td>9,684</td>
<td>9,088</td>
</tr>
<tr>
<td>Computer hardware</td>
<td>6,126</td>
<td>4,987</td>
</tr>
<tr>
<td>Construction in progress</td>
<td>71,232</td>
<td>448</td>
</tr>
<tr>
<td>Less — accumulated depreciation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Property and equipment, net</td>
<td>$179,954</td>
<td>$11,445</td>
</tr>
</tbody>
</table>

Depreciation expense was approximately $4.3 million, $5.1 million and $7.4 million for the years ended December 31, 2020, 2019 and 2018, respectively.

Accrued Expenses

Accrued expenses consist of the following at December 31 (in thousands):

<table>
<thead>
<tr>
<th></th>
<th>2020</th>
<th>2019</th>
</tr>
</thead>
<tbody>
<tr>
<td>Employee benefits and compensation</td>
<td>$20,752</td>
<td>$7,504</td>
</tr>
<tr>
<td>Research and development accruals</td>
<td>99,994</td>
<td>6,175</td>
</tr>
<tr>
<td>Other accrued expenses</td>
<td>16,644</td>
<td>1,188</td>
</tr>
<tr>
<td>Accrued expenses</td>
<td>$137,300</td>
<td>$14,867</td>
</tr>
</tbody>
</table>

Purchase Commitments

During 2020, the Company entered into agreements in the normal course of business with CMOs and CDMOs supplying the Company with production capabilities, and with vendors for preclinical studies, clinical trials and other goods or services. A number of these arrangements are within the scope of lease accounting (see Note 7). Certain agreements provide for termination rights subject to termination fees. Under such agreements, the Company is contractually obligated to make payments to vendors, mainly to reimburse them for their estimated unrecoverable expenses. The exact amount of such obligations are dependent on the timing of termination, and the terms of the relevant agreement, and cannot be reasonably estimated. As of December 31, 2020, these agreements are active ongoing arrangements, and the Company expects to receive value from these arrangements in the future.

As of December 31, 2020, the Company had approximately $117 million of such non-cancelable purchase commitments with a remaining term of more than one year.

Prepaid expenses and other current assets consist of the following at December 31 (in thousands):

<table>
<thead>
<tr>
<th></th>
<th>2020</th>
<th>2019</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prepaid expenses</td>
<td>$171,602</td>
<td>$3,601</td>
</tr>
<tr>
<td>Other current assets</td>
<td>9,662</td>
<td>4,376</td>
</tr>
<tr>
<td>Prepaid expenses and other current assets</td>
<td>$181,264</td>
<td>$7,977</td>
</tr>
</tbody>
</table>
Note 11—Long-Term Debt

Convertible Notes

In 2016, the Company issued $325 million aggregate principal amount of convertible senior unsecured notes that will mature on February 1, 2023 (the “Notes”). The Notes are senior secured debt obligations and were issued at par. The Notes were issued pursuant to an indenture dated January 29, 2016 (the “Indenture”), between the Company and the trustee. The Company received $315.0 million in net proceeds from the offering after deducting underwriting fees and offering expenses. The Notes bear cash interest at a rate of 3.75%, payable on February 1 and August 1 of each year, beginning on August 1, 2016. The Notes are not redeemable prior to maturity and are convertible into shares of the Company’s common stock. As a result of the Company’s one-for-twenty reverse stock split (see Note 13) and pursuant to Section 14.04(a) of the Indenture, the Notes are initially convertible into approximately 2,385,800 shares of the Company’s common stock based on the initial conversion rate of 7.3411 shares of the Company’s common stock per $1,000 principal amount of the Notes. This represents an initial conversion price of approximately $136.20 per share of the Company’s common stock, representing an approximate 22.2% conversion premium based on the last reported sale price of the Company’s common stock of $111.20 per share on January 25, 2016. In addition, the holders of the Notes may require the Company to repurchase the Notes at par value plus accrued and unpaid interest following the occurrence of a Fundamental Change (as described in the Indenture). If a holder of the Notes converts upon a Make-Whole Adjustment Event (as described in the Indenture), they may be eligible to receive a make-whole premium through an increase to the conversion rate up to a maximum of 8.9928 shares per $1,000 principal amount of Notes (subject to other adjustments as described in the Indenture).

The Notes are accounted for in accordance with ASC 470-20, Debt with Conversion and Other Options (“ASC 470-20”) and ASC 815-40, Contracts in Entity’s Own Equity (“ASC 815-40”). Under ASC 815-40, to qualify for equity classification (or non-derivative, if embedded) the instrument (or embedded feature) must be both (1) indexed to the issuer’s stock and (2) meet the requirements of the equity classification guidance. Based upon the Company’s analysis, it was determined the Notes do contain embedded features indexed to its own stock, but do not meet the requirements for bifurcation, and therefore do not need to be separately accounted for as an equity component. Since the embedded conversion feature meets the equity scope exception from derivative accounting, and also since the embedded conversion option does not need to be separately accounted for as an equity component under ASC 470-20, the proceeds received from the issuance of the convertible debt were recorded as a liability on the consolidated balance sheets.

In connection with the issuance of the Notes, the Company also paid $38.5 million, including expenses, to enter into privately negotiated capped call transactions with certain financial institutions (the “capped call transactions”). The capped call transactions are generally expected to reduce the potential dilution upon conversion of the Notes in the event that the market price per share of the Company’s common stock, as measured under the terms of the capped call transactions, is greater than the strike price of the capped call transactions, which initially corresponds to the conversion price of the Notes, and is subject to anti-dilution adjustments generally similar to those applicable to the conversion rate of the Notes. The cap price of the capped call transactions will be classified as an equity instrument.

The Company incurred approximately $10.0 million of debt issuance costs in 2016 relating to the issuance of the Notes, which were recorded as a reduction to the Notes on the consolidated balance sheet. The $10.0 million of debt issuance costs is being amortized and recognized as additional interest expense over the seven-year contractual term of the Notes on a straight-line basis, which approximates the effective interest rate method. The Company also incurred $0.9 million of expenses related to the capped call transactions, which were recorded as a reduction to additional paid-in-capital.

Note 12—Stockholders’ Equity

In 2020, the Company entered into various At Market Issuance Sales Agreements, which allows it to issue and sell up to $1.0 billion in gross proceeds of its common stock. During 2020, the Company sold 25.2 million shares of common stock under these Sales Agreements resulting in $835.6 million in net proceeds (this amount excludes $3.2 million received in the first quarter of 2021 for shares traded in late December 2020) and 7.2 million shares of common stock resulting in $38.5 million in net proceeds from the remaining portion of its At Market Issuance Sales Agreement entered into prior to 2020. From January 1, 2020 through January 20, 2021, the Company sold 0.9 million shares of common stock from its At Market Issuance Sales Agreement entered into in November 2020 (“November 2020 Sales Agreement”) resulting in $113.0 million in net proceeds, leaving $27.2 million remaining under the agreement. The Company terminated the November 2020 Sales Agreement by mutual agreement upon entering into the January 2021 Sales Agreement.

In 2019, the Company sold 13.0 million shares of common stock resulting in $97.4 million in net proceeds (this amount excludes $0.5 million received in the first quarter of 2020 for shares traded in late December 2019) under its various At Market Issuance Sales Agreements.

On May 8, 2019, the Company’s stockholders of record as of March 25, 2019 approved a one-for-twenty reverse stock split of the Company’s outstanding common stock, which was effected on May 10, 2019. The number of authorized shares of common stock and preferred stock of the Company was not affected and remains at 600,000,000 and 2,000,000, respectively, but the number of shares of common stock outstanding as of May 10, 2019 was reduced from 469,453,883 to 23,472,574. The aggregate par value of the issued common stock was reduced by reclassifying a portion of the par value amount of the outstanding common shares from Common Stock to Additional paid-in-capital for all periods presented. In addition, all per share and share amounts, including stock options and restricted stock awards, have been retroactively restated in the accompanying consolidated financial statements and notes thereto for all periods presented to reflect the reverse stock split.

In 2018, the Company sold 1.2 million shares of common stock resulting in $46.2 million in net proceeds under its various At Market Issuance Sales Agreements and completed a public offering of 1.7 million shares of its common stock, including 0.2 million shares of common stock that were issued upon the exercise in full of the option to purchase additional shares granted to the underwriters, at a price of $33.00 per share resulting in net proceeds, net of offering costs of $3.6 million, of approximately $54 million.
Note 13 – Stock-Based Compensation

Stock Options

The 2015 Stock Incentive Plan, as amended (“2015 Plan”), was approved at the Company’s annual meeting of stockholders in June 2015. Under the 2015 Plan, equity awards may be granted to officers, directors, employees and consultants of and advisers to the Company and any present or future subsidiary.

The 2015 Plan authorizes the issuance of up to 10,900,000 shares of common stock under equity awards granted under the 2015 Plan, which includes an increase of 7,100,000 shares approved for issuance under the 2015 Plan at the Company’s 2020 annual meeting of stockholders. All such shares authorized for issuance under the 2015 Plan have been reserved. The 2015 Plan will expire on March 4, 2025.

The Amended and Restated 2005 Stock Incentive Plan (“2005 Plan”) expired in February 2015 and no new awards may be made under such plan, although awards will continue to be outstanding in accordance with their terms.

The 2015 Plan permits and the 2005 Plan permitted the grant of stock options (including incentive stock options), restricted stock, stock appreciation rights and restricted stock units. In addition, under the 2015 Plan, unvested stock units and performance awards may be granted. Stock options and stock appreciation rights generally have a maximum term of 10 years and may be or were granted with an exercise price that is no less than 100% of the fair market value of the Company’s common stock at the time of grant. Grants of stock options are generally subject to vesting over periods ranging from one to four years.

Stock Options and Stock Appreciation Rights

The following is a summary of stock options and stock appreciation rights activity under the 2015 Plan and the 2005 Plan for the year ended December 31, 2020:

<table>
<thead>
<tr>
<th></th>
<th>2015 Plan</th>
<th>2005 Plan</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Stock Options</td>
<td>Weighted-Average Exercise Price</td>
</tr>
<tr>
<td>Outstanding at January 1, 2020</td>
<td>3,388,750</td>
<td>$ 35.64</td>
</tr>
<tr>
<td>Granted</td>
<td>3,363,766</td>
<td>38.01</td>
</tr>
<tr>
<td>Exercised</td>
<td>(1,025,025)</td>
<td>31.39</td>
</tr>
<tr>
<td>Canceled</td>
<td>(307,028)</td>
<td>33.51</td>
</tr>
<tr>
<td>Outstanding at December 31, 2020</td>
<td>5,420,463</td>
<td>38.05</td>
</tr>
<tr>
<td>Shares exercisable at December 31, 2020</td>
<td>878,488</td>
<td>71.27</td>
</tr>
<tr>
<td>Shares available for grant at December 31, 2020</td>
<td>2,473,916</td>
<td></td>
</tr>
</tbody>
</table>

In 2019, the Company granted 192,400 stock appreciation rights, with a weighted-average exercise price of $5.95, under the 2015 Plan.

Additionally, in 2019, due to limitations on the equity awards available under the 2015 Plan, the Company granted to certain employees 1,014,240 stock options, with a weighted-average exercise price of $5.95, under the 2015 Plan that were subject to approval of an increase in the number of shares under the 2015 Plan at the Company’s 2020 annual meeting of stockholders. Furthermore, in April 2020, due to limitations on the equity awards available under the 2015 Plan, the Company granted to all of its employees collectively 2,501,600 stock options, with a weighted-average exercise price of $19.08, and 326,050 restricted stock units under the 2015 Plan that include a performance requirement related to its NVX-CoV2373 program that were also subject to approval of an increase in the number of shares under the 2015 Plan at the Company’s 2020 annual meeting of stockholders. Since the proposal to increase the number of shares under the 2015 Plan was approved at the Company’s 2020 annual meeting of stockholders, as discussed in the “Stock Options” section above, the Company began to record stock-based compensation expense for these awards at that time.

The fair value of stock options granted under the 2015 Plan was estimated at the date of grant or the date upon which the 2015 Plan was approved by the Company’s stockholders for stock options discussed above using the Black-Scholes option-pricing model with the following assumptions:

<table>
<thead>
<tr>
<th></th>
<th>2020</th>
<th>2019</th>
<th>2018</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weighted average Black-Scholes fair value of stock options and SARs granted</td>
<td>$80.48</td>
<td>$4.98</td>
<td>$34.80</td>
</tr>
<tr>
<td>Risk-free interest rate</td>
<td>0.2%-1.5%</td>
<td>1.5%-2.6%</td>
<td>2.3%-3.1%</td>
</tr>
<tr>
<td>Dividend yield</td>
<td>—%</td>
<td>—%</td>
<td>—%</td>
</tr>
<tr>
<td>Volatility</td>
<td>116.0%-152.2%</td>
<td>105.4%-134.1%</td>
<td>93.3%-115.6%</td>
</tr>
<tr>
<td>Expected term (in years)</td>
<td>3.9-7.6</td>
<td>3.9-7.5</td>
<td>4.1-7.5</td>
</tr>
</tbody>
</table>

The total aggregate intrinsic value and weighted-average remaining contractual term of stock options and stock appreciation rights outstanding under the 2015 Plan and 2005 Plan as of December 31, 2020 was $427.8 million and 8.5 years, respectively. The total aggregate intrinsic value and weighted-average remaining contractual term of stock options and stock appreciation rights exercisable under the 2015 Plan and 2005 Plan as of December 31, 2020 was $55.9 million and 5.8 years, respectively. The aggregate intrinsic value represents the total intrinsic value (the difference between the Company’s closing stock price on the last trading day of the period and the exercise price, multiplied by the number of in-the-money stock options and stock appreciation rights) that would have been received by the holders had all stock option and stock appreciation rights holders exercised their stock options and stock appreciation rights on December 31, 2020. This amount is subject to change based on changes to the closing price of the Company’s common stock. The aggregate intrinsic value of stock options exercised and vesting of restricted stock awards for 2020, 2019 and 2018 was $187.3 million, $0.5 million and $0.4 million, respectively.

Employee Stock Purchase Plan

The Employee Stock Purchase Plan, as amended (the “ESPP”), was approved at the Company’s annual meeting of stockholders in June 2013. The ESPP currently authorizes an aggregate of 600,000 shares of common stock to be purchased. The ESPP allows employees to purchase shares of common stock of the Company at each purchase date through payroll deductions of up to a maximum of 15% of their compensation, at 85% of the lesser of the market price of the shares at the time of purchase or the market price on the beginning date of an option period (or, if later, the date during the option period when the employee was first eligible to participate). At December 31, 2020, there were 255,596 shares available for issuance under the ESPP.

The ESPP is considered compensatory for financial reporting purposes. As such, the fair value of ESPP shares was estimated at the date of grant using the Black-Scholes option-pricing model with the following assumptions:

<table>
<thead>
<tr>
<th></th>
<th>2020</th>
<th>2019</th>
<th>2018</th>
</tr>
</thead>
<tbody>
<tr>
<td>Range of Black-Scholes fair values of ESPP shares granted</td>
<td>$2.57-$92.67</td>
<td>$2.57-$35.00</td>
<td>$7.20-$70.64</td>
</tr>
<tr>
<td>Risk-free interest rate</td>
<td>0.2%-2.6%</td>
<td>1.2%-2.6%</td>
<td>0.7%-2.2%</td>
</tr>
<tr>
<td>Dividend yield</td>
<td>—%</td>
<td>—%</td>
<td>—%</td>
</tr>
<tr>
<td>Volatility</td>
<td>66.6%-189.7%</td>
<td>52.2%-171.6%</td>
<td>52.2%-203.8%</td>
</tr>
<tr>
<td>Expected term (in years)</td>
<td>0.5-2.0</td>
<td>0.5-2.0</td>
<td>0.5-2.0</td>
</tr>
</tbody>
</table>
Restricted Stock Units

The following is a summary of for the year ended December 31, 2020:

| Outanding and Unvested at January 1, 2020 | 1,102,311 | 5.95 |
| Restricted stock units granted | 837,896 | 94.74 |
| Restricted stock units vested | (840,812) | 9.37 |
| Restricted stock units forfeited | (54,415) | 40.37 |
| Outanding and Unvested at December 31, 2020 | 1,048,980 | 72.59 |

The Company recorded stock-based compensation expense for awards issued under the above mentioned plans in the consolidated statements of operations as follows (in thousands):

<table>
<thead>
<tr>
<th>Year Ended December 31.</th>
<th>2020</th>
<th>2019</th>
<th>2018</th>
</tr>
</thead>
<tbody>
<tr>
<td>Research and development</td>
<td>$55,955</td>
<td>$8,436</td>
<td>$10,575</td>
</tr>
<tr>
<td>General and administrative</td>
<td>72,080</td>
<td>8,612</td>
<td>7,739</td>
</tr>
<tr>
<td>Total stock-based compensation expense</td>
<td>$128,035</td>
<td>$17,048</td>
<td>$18,314</td>
</tr>
</tbody>
</table>

As of December 31, 2020, there was approximately $312 million of total unrecognized compensation expense related to vested options, stock appreciation rights, restricted stock units and the ESPP. This unrecognized non-cash compensation expense is expected to be recognized over a weighted-average period of 1.3 years, and will be allocated between research and development and general and administrative expenses accordingly. This estimate does not include the impact of other possible stock-based awards that may be made during future periods and awards that require approval by the stockholders.

Note 14 – Employee Benefits

The Company maintains a defined contribution 401(k) retirement plan, pursuant to which employees may elect to contribute up to 100% of their compensation on a tax deferred basis up to the maximum amount permitted by the Internal Revenue Code of 1986, as amended.

The Company matches 100% of the first 3% of the participants’ deferral, and 50% on the next 2% of the participants’ deferral, up to a potential 4% Company match. The Company’s matching contributions to the 401(k) plan vest immediately. Under its 401(k) plan, the Company has recorded expense of $0.9 million, $1.0 million and $1.2 million in 2020, 2019 and 2018, respectively.

The Company’s foreign subsidiaries have pension plans under local tax and labor laws and are obligated to make contributions to the plan. Contributions and other expenses related to this plan were $1.0 million, $0.7 million and $0.8 million in 2020, 2019 and 2018, respectively.

Note 15 – Income Taxes

The Company’s loss from operations before income tax expense by jurisdiction for the years ended December 31 are as follows (in thousands):

<table>
<thead>
<tr>
<th>Year Ended December 31</th>
<th>2020</th>
<th>2019</th>
<th>2018</th>
</tr>
</thead>
<tbody>
<tr>
<td>Domestic</td>
<td>(455,253)</td>
<td>(124,189)</td>
<td>(176,290)</td>
</tr>
<tr>
<td>Foreign</td>
<td>36,994</td>
<td>(8,505)</td>
<td>(8,458)</td>
</tr>
<tr>
<td>Total net loss</td>
<td>(418,259)</td>
<td>(132,694)</td>
<td>(184,748)</td>
</tr>
</tbody>
</table>

As a result of current and historical losses, there is no income tax provision for the years ended December 31, 2020, 2019 and 2018.

A reconciliation of the provision for income tax to the amount computed by applying the U.S. federal statutory tax rate to the Company’s effective tax rate is as follows:

<table>
<thead>
<tr>
<th>Year Ended December 31</th>
<th>2020</th>
<th>2019</th>
<th>2018</th>
</tr>
</thead>
<tbody>
<tr>
<td>Statutory federal tax rate</td>
<td>(21)%</td>
<td>(21)%</td>
<td>(21)%</td>
</tr>
<tr>
<td>State income taxes, net of federal benefit</td>
<td>(3)%</td>
<td>(2)%</td>
<td>(3)%</td>
</tr>
<tr>
<td>Research and development and other tax credits</td>
<td>0%</td>
<td>0%</td>
<td>0%</td>
</tr>
<tr>
<td>Non-deductible expenses</td>
<td>4%</td>
<td>0%</td>
<td>0%</td>
</tr>
<tr>
<td>Non-cash stock-based compensation</td>
<td>(7)%</td>
<td>0%</td>
<td>0%</td>
</tr>
<tr>
<td>Other</td>
<td>1%</td>
<td>1%</td>
<td>1%</td>
</tr>
<tr>
<td>Change in tax rate</td>
<td>(5)%</td>
<td>3%</td>
<td>5%</td>
</tr>
<tr>
<td>Change in valuation allowance</td>
<td>31%</td>
<td>22%</td>
<td>21%</td>
</tr>
<tr>
<td>Income tax provision</td>
<td>0%</td>
<td>0%</td>
<td>0%</td>
</tr>
</tbody>
</table>

As of December 31, 2020, the Company has available federal, state, and foreign net operating losses of $1.3 billion, $756.0 million and $42.7 million, respectively, that may be applied against future taxable income. A significant portion of the federal net operating losses will begin to expire in 2037. A portion of the foreign net operating losses will begin to expire in 2023. The Company also has research tax credits of $35.1 million that begin to expire in 2020. Utilization of the net operating loss carryforwards and credits may be subject to an annual limitation due to ownership changes of the Company. As of December 31, 2020, the Company does not expect such limitation, if any, to impact the use of the net operating losses and business tax credits.

The Company files income tax returns in the U.S. federal jurisdiction and in various states, as well as in Sweden and the Czech Republic. The Company has U.S. tax net operating losses and credit carryforwards that are subject to examination from 2000 through 2020. The returns in Sweden are subject to examination from 2014 through 2020 and the returns for the Czech Republic are subject to examination from 2017 through 2020.
The significant components of the Company’s deferred tax assets and liabilities as of December 31 were as follows (in thousands):

<table>
<thead>
<tr>
<th></th>
<th>2020</th>
<th>2019</th>
</tr>
</thead>
<tbody>
<tr>
<td>Deferred tax assets:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Federal and State net operating loss carryforward</td>
<td>$325,655</td>
<td>$293,736</td>
</tr>
<tr>
<td>Foreign net operating loss carryforward</td>
<td>8,620</td>
<td>13,520</td>
</tr>
<tr>
<td>Research tax credits</td>
<td>35,065</td>
<td>37,066</td>
</tr>
<tr>
<td>Lease liability</td>
<td>39,548</td>
<td>2,164</td>
</tr>
<tr>
<td>Deferred revenue</td>
<td>60,567</td>
<td>973</td>
</tr>
<tr>
<td>Non-cash stock-based compensation</td>
<td>22,577</td>
<td>13,679</td>
</tr>
<tr>
<td>Original discount interest</td>
<td>3,177</td>
<td>4,326</td>
</tr>
<tr>
<td>Other</td>
<td>12,019</td>
<td>2,820</td>
</tr>
<tr>
<td>Total deferred tax assets</td>
<td>507,318</td>
<td>368,284</td>
</tr>
<tr>
<td>Valuation allowance</td>
<td>(504,788)</td>
<td>(365,772)</td>
</tr>
<tr>
<td>Net deferred tax assets</td>
<td>$2,530</td>
<td>$2,512</td>
</tr>
<tr>
<td>Deferred tax liabilities:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ROU assets</td>
<td>(1,253)</td>
<td>(1,033)</td>
</tr>
<tr>
<td>Intangibles</td>
<td>(1,198)</td>
<td>(1,279)</td>
</tr>
<tr>
<td>Other</td>
<td>(79)</td>
<td>(200)</td>
</tr>
<tr>
<td>Total deferred tax liabilities</td>
<td>$ (2,530)</td>
<td>$ (2,512)</td>
</tr>
<tr>
<td>Net deferred tax assets</td>
<td>$ —</td>
<td>$ —</td>
</tr>
</tbody>
</table>

The valuation allowance increased by $139.0 million and $28.3 million for the years ended December 31, 2020 and 2019, respectively, due to increases in deferred tax assets. Realization of net deferred tax assets is dependent on the Company’s ability to generate future taxable income, which is uncertain. Accordingly, a full valuation allowance was recorded against these assets as of December 31, 2020 and 2019 as management believes it is more likely than not that the assets will not be realizable.

The Company recognizes the effect of a tax position when it is more likely than not, based on the technical merits, that the tax position will be sustained upon examination. A reconciliation of the beginning and ending amounts of unrecognized tax benefits in the year ended December 31, 2020, 2019 and 2018 is as follows (in thousands):

<table>
<thead>
<tr>
<th></th>
<th>2020</th>
<th>2019</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unrecognized tax benefits balance at January 1,</td>
<td>$ —</td>
<td>$ —</td>
</tr>
<tr>
<td>Additions for tax positions of current year</td>
<td>1,413</td>
<td>—</td>
</tr>
<tr>
<td>Additions for tax positions of prior years</td>
<td>7,353</td>
<td>—</td>
</tr>
<tr>
<td>Reductions for tax positions of prior year</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Settlements of tax positions of prior years</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Unrecognized tax benefits balance at December 31</td>
<td>$8,766</td>
<td>$ —</td>
</tr>
</tbody>
</table>

The Company's policy is to recognize interest and penalties related to income tax matters in income tax expense. As of December 31, 2020 and 2019, the Company had no accruals for interest or penalties related to income tax matters. The total amount of unrecognized tax benefits that, if recognized, would affect the effective tax rate is $8.8 million.

Note 16 – Related Party Transaction

In June 2020, in advance of David M. Mott joining the Company’s Board of Directors, the Company agreed to sell 32,916 shares of common stock to him at a purchase price of $45.57 per share, reflecting the closing price of the Company’s common stock on the trading date prior to the date the parties’ agreement regarding the sale, for total gross proceeds of $1.5 million. Mr. Mott joined the Company’s Board of Directors later in the same month.

Note 17 – Subsequent Events

In January 2021, the Company entered into an At Market Issuance Sales Agreement (“January 2021 Sales Agreement”), which allows it to issue and sell up to $500 million in gross proceeds of its common stock. From January 22 through February 24, 2021, the Company sold 1.7 million shares of common stock under the January 2021 Sales Agreement resulting in $452.0 million in net proceeds, leaving $42.2 million remaining.

In January and February 2021, the Company finalized multiple advance purchase agreements and one binding Heads of Terms to supply. In total, approximately 75 million doses of NVX-CoV2373 to various government customers. The Company will work with the relevant regulatory agencies to obtain necessary approvals, as necessary.

In February 2021, the Company finalized an expanded collaboration and license agreement with SK biocience to manufacture and commercialize NVX-CoV2373 for sale to the Korean government. Concurrently, SK biocience finalized an advance purchase agreement with the Korean government to supply 40 million doses of NVX-CoV2373 to the Republic of Korea beginning in 2021. The agreement is in addition to the Company’s existing manufacturing arrangement with SK biocience.

In February 2021, the Company entered into a Memorandum of Understanding with Gavi, the Vaccine Alliance (“Gavi”), to provide 1.1 billion cumulative doses of NVX-CoV2373 for the COVAX Facility. The Company will work with Gavi to finalize an advance purchase agreement for vaccine supply and global distribution via the COVAX Facility and its partners. The vaccine doses will be manufactured and distributed globally by the Company and SIIPL.
Corporate Information

Board of Directors

- James F. Young, Ph.D.
  Chairman of the Board of Directors
- Stanley C. Erck
  President and Chief Executive Officer, Director
- Gregg H. Alton, J.D.
  Director
- Richard H. Douglas, Ph.D.
  Director
- Gary C. Evans
  Director

Management Team

- Lyn Caltabiano, Ph.D.
  Senior Vice President, Global Program Management
- Frank Czworka
  Senior Vice President, Global Sales
- Timothy J. Hahn, Ph.D.
  Senior Vice President, CMC, NanoFlu
- Biegie Lee
  Senior Vice President, Chief Information Officer
- Brian Rosen
  Senior Vice President, Commercial Strategy and Public Policy
- Gale E. Smith, Ph.D.
  Senior Vice President, Discovery and Pre-clinical Research, Chief Scientist
- Silvia Taylor
  Senior Vice President, Investor Relations and Corporate Affairs
- Henrietta Ukwu, M.D.
  Senior Vice President, Chief Regulatory and Quality Officer
- Brian Webb
  Senior Vice President, Manufacturing

Annual Meeting

June 17, 2021 at 8:30 a.m. EDT
Live virtual webcast link:
www.virtualshareholdermeeting.com/NVAX2021

Independent Registered Public Accounting Firm

Ernst & Young, LLP
1775 Tysons Boulevard
McLean, VA 22102

Transfer Agent

Computershare, Inc.
250 Royall Street
Canton, MA 02021

Novavax Corporate Headquarters

Novavax, Inc.
21 Firstfield Road
Gaithersburg, MD 20878

Market Information

Novavax is traded on the NASDAQ Global Select Market under “NVAX”.

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21 Firstfield Road
Gaithersburg, MD 20878

Corporate Information

Business Information

Novavax, Inc.
21 Firstfield Road
Gaithersburg, MD 20878

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