

HEMISPHERX BIOPHARMA INC
Form 10-K
April 01, 2019

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, 2018

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. 000-27072

HEMISPHERX BIOPHARMA, INC.

(Exact name of registrant as specified in its charter)

Edgar Filing: HEMISPHERX BIOPHARMA INC - Form 10-K

Delaware 52-0845822
(State or other jurisdiction of (I.R.S. Employer Identification
incorporation or organization) Number)

2117 SW Highway 484, Ocala FL 34473
(Address of principal executive offices) (Zip Code)

Registrant's telephone number, including area code: (215) 988-0080

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

Common Stock, \$.001 par value

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

(Title of Each Class)

NONE

Indicate by check mark if the registrant is a well-known seasoned issuer, as 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 Section 15(d) of the Act. Yes No

Indicate by check mark whether the registrant (1) has filed all reports 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

Edgar Filing: HEMISPHERX BIOPHARMA INC - Form 10-K

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 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. [X]

Indicate by check mark whether the registrant is a large accelerated filer, an accelerated filer, a non-accelerated filer or a smaller reporting company. See definition of "large accelerated filer," "accelerated filer" and "smaller reporting company" in Rule 12b-2 of the Exchange Act:

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 has elected not to use the extended transition period for complying with any new or revised financial accounting standards pursuant to Section 13(a) of the Exchange Act.

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 Common Stock held by non-affiliates at June 30, 2018, the last business day of the registrant's most recently completed second fiscal quarter was \$14,981,748.

The number of shares of the registrant's Common Stock outstanding as of March 22, 2019 was 62,290,318.

DOCUMENTS INCORPORATED BY REFERENCE: None.

TABLE OF CONTENTS

	Page
<u>PART I</u>	
ITEM 1. <u>Business.</u>	3
ITEM 1A. <u>Risk Factors.</u>	11
ITEM 1B. <u>Unresolved Staff Comments.</u>	23
ITEM 2. <u>Properties.</u>	23
ITEM 3. <u>Legal Proceedings.</u>	23
ITEM 4. <u>Mine Safety Disclosures.</u>	23
<u>PART II</u>	
ITEM 5. <u>Market for the Registrant’s Common Equity, Related Stockholder Matters and Issuer Purchases of Equity Securities.</u>	24
ITEM 6. <u>Selected Financial Data.</u>	24
ITEM 7. <u>Management’s Discussion and Analysis of Financial Condition and Results of Operations.</u>	24
ITEM 7A. <u>Quantitative and Qualitative Disclosures About Market Risk.</u>	30
ITEM 8. <u>Financial Statements and Supplementary Data.</u>	31
ITEM 9. <u>Changes in and Disagreements with Accountants on Accounting and Financial Disclosure.</u>	31
ITEM 9A. <u>Controls and Procedures.</u>	31
ITEM 9B. <u>Other Information.</u>	32
<u>PART III</u>	
ITEM 10. <u>Directors, Executive Officers and Corporate Governance.</u>	32
ITEM 11. <u>Executive Compensation.</u>	36

ITEM 12. Security Ownership of Certain Beneficial Owners and Management and Related Stockholder Matters. 44

ITEM 13. Certain Relationships and Related Transactions and Director Independence. 49

ITEM 14. Principal Accountant Fees and Services. 49

PART IV

ITEM 15. Exhibits and Financial Statement Schedules. 50

PART I

ITEM 1. Business

GENERAL

Hemispherx Biopharma, Inc. and its subsidiaries (collectively, “Hemispherx”, “Company”, “we” or “us”) are an immuno-pharma company headquartered in Ocala, Florida and focused on the research and development of therapeutics to treat multiple types of cancers, as well as immune-deficiency disorders. We have established a strong foundation of laboratory, pre-clinical and clinical data with respect to the development of nucleic acids and natural interferon to enhance the natural antiviral defense system of the human body and to aid the development of therapeutic products for the treatment of certain cancers and chronic diseases.

Hemispherx’s flagship products include Ampligen® (Rintatolimod), a first-in-class drug of large macromolecular RNA (ribonucleic acid) molecules, and Alferon N Injection® (Interferon Alfa-N3). A first-in-class drug also known as a new chemical entity, is a drug that contains an active moiety that has not been approved by the FDA or marketed in the US.

Ampligen® represents an RNA being developed for globally important cancers, viral diseases and disorders of the immune system. Ampligen® has in the clinic demonstrated the potential for standalone efficacy in a number of solid tumors. We have also seen success in increasing survival rates and efficacy in the treatment of animal tumors when Ampligen® is used in combination with checkpoint blockade therapies. This success in the field of immuno-oncology has guided our focus toward the potential use of Ampligen® as a combinational therapy for the treatment of a variety of solid tumor types. There are currently multiple Ampligen® clinical trials — both underway and planned — at major cancer research centers around the country. Ampligen® is also being used as a monotherapy to treat pancreatic cancer patients in an Early Access Program (EAP) approved by the Inspectorate of Healthcare in the Netherlands at Erasmus Medical Center.

Ampligen® is also being evaluated for the treatment of myalgic encephalomyelitis/chronic fatigue syndrome (ME/CFS). Hemispherx is currently sponsoring an expanded access program (EAP) for ME/CFS patients in the U.S. In August 2016, we received approval of our NDA from Administracion Nacional de Medicamentos, Alimentos y Tecnologia Medica (ANMAT) for commercial sale of Ampligen® in the Argentine Republic for the treatment of severe CFS. With regulatory approval in Argentina, Ampligen® is the world’s only approved therapeutic for ME/CFS. We continue to pursue our Ampligen New Drug Application, or NDA, for the treatment of CFS with the Food and Drug Administration, or FDA. Please see “Research And Development (“R&D”); Myalgic Encephalomyelitis/Chronic Fatigue Syndrome (“ME/CFS”)” below.

Alferon N Injection® is approved for a category of STD infection and patients that are intolerant to recombinant interferon in Argentina. Alferon is the only natural-source, multi-species alpha interferon currently approved for sale in the U.S. for the intralesional treatment of refractory (resistant to other treatment) or recurring external Condylomata Acuminata/genital warts (GW) in patients 18 years of age or older. Certain types of human papilloma viruses cause GW. Hemispherx also has approval from ANMAT for the treatment of refractory patients that failed or were intolerant to treatment with recombinant interferon in Argentina. We have developed and, with proper funding, will be seeking FDA Pre-Approval Inspection of a high-volume, high-efficiency, upgraded manufacturing process to allow for the commercial viability of Alferon®.

We operate a 30,000 sq. ft. facility in New Brunswick, NJ with the objective of producing Ampligen® and Alferon®. We are committed to a focused business plan oriented toward finding senior co-development partners with the capital and expertise needed to commercialize the many potential therapeutic aspects of Ampligen® and our FDA-approved drug Alferon® N.

AVAILABLE INFORMATION

We file our annual reports on Form 10-K, quarterly reports on Form 10-Q and current reports on Form 8-K pursuant to Section 13(a) or 15(d) of the Exchange Act electronically with the Securities and Exchange Commission, or SEC. The public may obtain information on the operation of the Public Reference Room by calling the SEC at 1-800-SEC-0330. The SEC maintains an Internet site that contains reports, proxy and information statements, and other information regarding issuers that file electronically with the SEC. The address of that site is <http://www.sec.gov>.

You may obtain a free copy of our annual reports on Form 10-K, quarterly reports on Form 10-Q and current reports on Form 8-K and amendments to those reports on the day of filing with the SEC on our website on the World Wide Web at <http://www.hemispherx.net> under the Investor Relations tab for SEC Filings or by contacting the Investor Relations Department by calling 888-557-6480 or sending an e-mail message to ir@hemispherx.net.

OUR PRODUCTS

Our primary pharmaceutical product platform consists of Ampligen®, first-in-class drug of large macromolecular double-stranded (ds) RNA (ribonucleic acid) molecules and our FDA approved natural alpha-interferon product, Alferon N Injection®.

Ampligen®

Ampligen® is approved for sale in Argentina for severe Chronic Fatigue Syndrome (CFS) and is an experimental drug in the United States currently undergoing clinical development for the treatment of certain cancers and ME/CFS. Over its developmental history, Ampligen® has received various designations, including Orphan Drug Product Designation (FDA and European Medicines Agency (“EMA”)), Treatment protocol (e.g., “Expanded Access” or “Compassionate” use authorization) with Cost Recovery Authorization (FDA) and “promising” clinical outcome recognition based on the evaluation of certain summary clinical reports (“AHRQ” or Agency for Healthcare Research and Quality). Ampligen® represents the first drug in the class of large (macromolecular) dsRNA molecules to apply for NDA review. Based on the results of published, peer reviewed pre-clinical studies and clinical trials, we believe that Ampligen® may have broad-spectrum anti-viral and anti-cancer properties.

We believe that nucleic acid compounds represent a potential new class of pharmaceutical products designed to act at the molecular level for treatment of many human diseases. There are two forms of nucleic acids, deoxyribonucleic acid (“DNA”) and ribonucleic acid (“RNA”). DNA is a group of naturally occurring molecules found in chromosomes, the cell’s genetic machinery. RNA is a group of naturally occurring informational molecules which orchestrate a cell’s behavior which, in turn, regulates the action of groups of cells, including the cells which compromise the body’s immune system. RNA directs the production of proteins and regulates certain cell activities including the activation of an otherwise dormant cellular defense against viruses and tumors. Our drug technology utilizes specifically-configured RNA and is a selective TLR3 agonist that is administered intravenously. Ampligen® has been assigned the generic name rintatolimod by the United States Adopted Names Council (USANC) and has the chemical designation poly(I):poly(C₁₂U).

EAP/clinical trials of Ampligen® that have been conducted or that are ongoing include studies of the potential treatment of patients with renal cell carcinoma, malignant melanoma, non-small cell lung, ovarian, breast, colorectal, urothelial, prostate and pancreatic cancer, CFS, Hepatitis B and HIV.

We have received approval of our NDA from ANMAT for commercial sale of rintatolimod (U.S. tradename: Ampligen®) in the Argentine Republic for the treatment of severe CFS. The product will be marketed by GP Pharm, our commercial partner in Latin America. Commercialization in Argentina will require, among other things, GP

Pharm to establish disease awareness, medical education, creation of an appropriate reimbursement level, design of marketing strategies and completion of manufacturing preparations for launch.

The FDA has authorized an open-label expanded access treatment protocol, (“AMP-511”), allowing patient access to Ampligen® in an open-label safety study under which severely debilitated CFS patients have the opportunity to be on Ampligen® to treat this very serious and chronic condition. The data collected from the AMP-511 protocol through clinical sites provide safety information regarding the use of Ampligen® in patients with CFS. We are establishing an enlarged data base of clinical safety information which we believe will provide further documentation regarding the absence of autoimmune disease associated with Ampligen® treatment. We believe that continued efforts to understand existing data, and to advance the development of new data and information, will ultimately support our future filings for Ampligen® and/or the design of future clinical studies that the FDA requested in a complete response letter. The FDA recently approved the increase reimbursement level from \$200 to \$345 per 200mg vial of Ampligen, due to increased production costs. At this time, we do not plan on passing this adjustment along the patients in this program. As of December 31, 2018, there are 15 patients being treated in this open-label expanded access treatment protocol.

In May 2016, we entered into a five-year agreement with myTomorrows, a Netherlands based company, for the commencement and management of an Early Access Program (“EAP”) in Europe and Turkey (the “Territory”) related to ME/CFS. Pursuant to the agreement, as amended, myTomorrows also will manage all Early Access Programs and Special Access Programs in Europe, Canada and Turkey to treat pancreatic cancer and ME/CFS patients.

In April 2018, we completed data analysis of an intranasal human safety study of Ampligen® plus FluMist® known as AMP-600. The study was previously closed after the US Centers for Disease Control and Prevention (“CDC”) recommended against the use of FluMist®. Intranasal Ampligen® in combination with FluMist® was generally well-tolerated in the study.

In June 2018, Ampligen® was cited as outperforming two other TLR3 agonists, poly IC and natural double stranded RNA, in creating an enhanced tumor microenvironment for checkpoint blockage therapy in the journal of Cancer Research (<http://cancerres.aacrjournals.org/content/early/2018/05/31/0008-5472.CAN-17-3985>). In a head-to-head study in explant culture models, Ampligen® activated the TLR3 pathway and promoted an accumulation of killer T cells but, unlike the other two TLR3 agonists, it did so without causing regulatory T cell (Treg) attraction. These findings were considered important because they indicate that Ampligen® selectively reprograms the tumor microenvironment by inducing the beneficial aspects of tumor inflammation (attracting killer T cells), without amplifying immune suppressive elements such as regulatory T cells. The study was conducted at the University of Pittsburgh and Roswell Park Comprehensive Cancer Center, as a part of the NIH-funded P01 CA132714 and Ovarian Cancer Specialized Program of Research Excellence (SPORE). Based upon these findings Hemispherx and Roswell Park Comprehensive Cancer Center expanded their existing scientific collaboration to advance the clinical development of Ampligen® which has shown promise in preclinical studies when combined with checkpoint inhibitors (CPIs). The parties executed a Memorandum of Understanding (“MOU”) designed to further assess the clinical potential of Ampligen® in treating certain cancers. This phase I/II study will evaluate the potential of Ampligen® to enhance the immune mediated effects of CPIs in patients with advanced solid tumors including bladder, melanoma and renal cell carcinoma.

In 2018, we also reported that we completed production of two commercial-size batches of more than 16,000 vials of Ampligen®, following its “Fill & Finish” at the Contract Manufacturing Organization. These lots passed all required testing for regulatory release for human use and are being used for multiple programs including the product launch in Argentina, for the treatment of ME/CFS, the pancreatic cancer EAP in the Netherlands, ongoing and future clinical studies in oncology, and our ME/CFS EAP in the U.S. and Europe.

Alferon N Injection®

Alferon N Injection® is the registered trademark for our injectable formulation of natural alpha interferon. Alferon® is the only natural-source, multi-species alpha interferon currently approved for sale in the U.S. and Argentina for the intralesional (within lesions) treatment of refractory (resistant to other treatment) or recurring external genital warts in patients 18 years of age or older. Alferon® is also approved in Argentina for the treatment of refractory patients that failed or were intolerant to treatment with recombinant interferons. Certain types of human papilloma viruses (“HPV”) cause genital warts, a sexually transmitted disease (“STD”). According to the CDC, HPV is the most common sexually transmitted infection, with approximately 79 million Americans — most in their late teens and early 20s — infected with HPV. In fact, the CDC states that “HPV is so common that nearly all sexually active men and women get the virus at some point in their lives.” Although they do not usually result in death, genital warts commonly recur, causing significant morbidity and entail substantial health care costs.

Interferons are a group of proteins produced and secreted by cells to combat diseases. Researchers have identified four major classes of human interferon: alpha, beta, gamma and omega. Alferon N Injection® contains a multi-species form of alpha interferon. The world-wide market for injectable alpha interferon-based products has experienced rapid growth and various alpha interferon injectable products are approved for many major medical uses worldwide. Alpha

interferons are manufactured commercially in three ways: by genetic engineering, by cell culture, and from human white blood cells. All three of these types of alpha interferon are or were approved for commercial sale in the U.S. Our natural alpha interferon is produced from human white blood cells.

The potential advantages of natural alpha interferon over recombinant (synthetic) interferon produced and marketed by other pharmaceutical firms may be based upon their respective molecular compositions. Natural alpha interferon is composed of a family of proteins containing many molecular species of interferon. In contrast, commercial recombinant alpha interferon products each contain only a single species. Researchers have reported that the various species of interferons may have differing antiviral activity depending upon the type of virus. Natural alpha interferon presents a broad complement of species, which we believe may account for its higher activity in laboratory studies. Natural alpha interferon is also glycosylated (partially covered with sugar molecules). Such glycosylation is not present on the currently U.S. marketed recombinant alpha interferons. We believe that the absence of glycosylation may be, in part, responsible for the production of interferon-neutralizing antibodies seen in patients treated with recombinant alpha interferon. Although cell culture-derived interferon is also composed of multiple glycosylated alpha interferon species, the types and relative quantity of these species are different from our natural alpha interferon.

Alferon N Injection® [Interferon alfa-n3 (human leukocyte derived)] is a highly purified, natural-source, glycosylated, multi-species alpha interferon product. There are essentially no neutralizing antibodies observed against Alferon N Injection® to date and the product has a relatively low side-effect profile. The recombinant DNA derived alpha interferon formulations have been reported to have decreased effectiveness after one year of treatment, probably due to neutralizing antibody formation.

See “Manufacturing” and “Marketing/Distribution” sections below for more details on the manufacture and marketing/distribution of Alferon N Injection®.

PATENTS AND NON-PATENT EXCLUSIVITY RIGHTS

As of December 31, 2018, we had 57 patents worldwide with 6 additional pending patent applications comprising our intellectual property. Please see “Note 5: Patents, Trademark Rights and Other Intangibles (FASB ASC 350 General Intangibles Other than Goodwill)” under Notes to Consolidated Financial Statements for more information on these patents. We continually review our patents’ rights to determine whether they have continuing value.

In 2016, we received a new Ampligen® composition of matter patent in the US (#9,315,538). In 2015, we were granted a new composition of matter patent (#2340307) by the European Patent Office and we received twenty-eight new patents in various EU countries. In 2014, we were granted a new composition of matter patent in the United States (#8722874) covering Ampligen® formulations.

The Ampligen® U.S. CFS treatment patent (#6130206) expired October 10, 2017 (we believe that the expiration of this patent will have minimal impact on us; see details on U.S. #9315538, U.S. #8722874 and information on the FDA has granted “orphan drug status” to the drug for CFS below). Our U.S. Ampligen® Trademark (#73617687) has been renewed through December 6, 2028. New therapeutic use patent applications are pending. On May 13, 2014, the United States Patent Office issued patent U.S. #8722874 titled “Double-Stranded Ribonucleic Acids with Rugged Physiochemical Structure and Highly Specific Biologic Activity,” with all rights assigned to Hemispherx. The patent claims a novel form of rugged dsRNA. Rugged dsRNA are nucleic acids with a unique composition and physical characteristic identified with high specificity of binding to Toll-Like Receptor 3 (TLR3), thereby conveying an important range of therapeutic opportunities. The newly discovered form of dsRNA has increased bioactivity and binding affinity to the TLR 3 receptor because of its reduced tendency to form branched dsRNA which can inhibit receptor binding. Pharmaceutical formulations containing the newly discovered nucleic acid as active ingredients and methods of treatment with those formulations are also described in the issued patent. Hemispherx believes that the issuance of U.S. Patents #9315538 and #8722874 will help ensure that Hemispherx retains patent protection for novel formulations of Ampligen® products until at least 2029.

In September 2015, the European Patent Office granted the European version of U.S. Patent #9315538, with all rights assigned to Hemispherx.

In addition to our patent rights relating to Ampligen®, the FDA has granted “orphan drug status” to the drug for CFS, HIV/AIDS, renal cell carcinoma and malignant melanoma. Orphan drug status grants us protection against the potential subsequent approval of other sponsors’ versions of the drug for these uses for a period of seven years following FDA approval of Ampligen® for each of these designated uses. The first NDA approval for Ampligen® as a new chemical entity will also qualify for four or five years of non-patent exclusivity during which abbreviated new drug applications seeking approval to market generic versions of the drug cannot be submitted to the FDA. (See “Government Regulation” below.)

In May 2011, a new United States Patent #7943147 was granted for the use of Ampligen® as a vaccine adjuvant for use with seasonal influenza vaccine to induce an enhanced immune response against H5N1 avian influenza.

With respect to Alferon®, the composition is a complex mixture of natural interferon species that is manufactured from human leukocytes obtained from human blood donors. In addition, while it is the current standard by the FDA to treat biological drug products like interferon as “Well Characterized” biologics, a process for which chemical entities can have their identity, purity, impurities, potency, and quality controlled by chemical testing, Alferon®, as a natural interferon, does not lend itself well to such testing. Moreover, FDA continues to require that each lot of Alferon® we produce be tested and released by the FDA before it can be distributed for commercial sales. Because of the complexity of the Alferon® manufacturing process and these additional regulatory requirements, we believe that potential manufacturers of generic, or so-called “bio-similar,” drug products are focused on developing recombinant interferon products, rather than natural interferon products. For these reasons, we believe that not having patent protection should have no or little impact on the Company. Additionally, at the receipt of the FDA certification for the revised Alferon® manufacturing process and techniques in New Brunswick, NJ, it is our intention to file for additional patent protection.

RESEARCH AND DEVELOPMENT (“R&D”)

Our general focus during the past two fiscal years has been on the clinical development of new drug therapies based on natural immune system enhancing technologies for the treatment of immune-based disorders including cancer and CFS.

Cancer

We have been working with the University of Pittsburgh’s chemokine modulation research initiative which includes the use of Ampligen® as a potential adjuvant to modify the tumor microenvironment (TME) with the goal of increasing anti-tumor responses to check point inhibitors (CPI). As part of this collaboration, Hemispherx has supplied Ampligen® (rintatolimod) to the University. The study, under the leadership of Robert P. Edwards, MD, chair of gynecologic services at Magee-Women’s Hospital of the University of Pittsburgh School of Medicine, and Professor of Surgery Pawel Kalinski, M.D., Ph.D., at Roswell Park Comprehensive Cancer Center, Buffalo, N.Y., involved the chemokine modulatory regimen developed by Dr. Kalinski’s group and successfully completed the Phase 1 dose escalation in patients with resectable colorectal cancer. In the 1st quarter of 2017, Dr. Kalinski relocated to Roswell Park Comprehensive Cancer Center (“RPCCC”) in Buffalo, NY and has established a cancer program which will continue to require a supply of Ampligen®.

In October 2018, we signed a clinical trial agreement with Roswell Park Comprehensive Cancer Center to evaluate Ampligen® in combination with checkpoint inhibitors (CPIs). The Phase IIa clinical trial will evaluate the immune-mediated effects of cytokine modulation in combination with CPIs in patients with primary resistance to CPI therapy. The protocol will seek to evaluate the combination of Ampligen® and CPIs in patients with advanced urothelial carcinoma, renal cell carcinoma and melanoma. Ampligen® is our investigational immune-enhancing TLR3 agonist that has demonstrated a robust anti-cancer effect in preclinical models when combined with CPIs. This new agreement expands the extensive prior clinical and preclinical work into the clinical checkpoint blockade arena and offers the opportunity to begin evaluation of this combination therapy in patients with a variety of solid tumors where large numbers of patients do not respond or progress following treatment with standard CPI-based therapy.

Currently, four Ampligen® clinical trials are underway at university cancer centers testing whether tumor microenvironments can be reprogrammed to increase the effectiveness of cancer immunotherapy, including checkpoint inhibitors:

Recurrent Ovarian Cancer - Phase 1 / 2 study of intraperitoneal chemo-immunotherapy in recurrent ovarian cancer at University of Pittsburgh Medical Center. Dr. R. Edwards, PI. Study underway. An interim report from Dr. Edwards' team is expected within thirty days and a summary of same will be disclosed upon receipt. See: <https://clinicaltrials.gov/ct2/show/NCT02432378>

Colorectal Cancer - Phase 2a study of Ampligen as component of chemokine modulatory regimen on colorectal cancer metastatic to liver at Roswell Park Comprehensive Cancer Center. Dr. P. Boland, PI. Study underway. See: <https://clinicaltrials.gov/ct2/show/NCT03403634>

Metastatic Triple Negative Breast Cancer - Open label study of metastatic triple-negative breast cancer using chemokine modulation therapy, including Ampligen and pembrolizumab, at Roswell Park Comprehensive Cancer Center. Dr. M. Opyrchal, PI. Initiation of study is expected in the near future and will be announced forthwith. See: <https://www.clinicaltrials.gov/ct2/show/NCT03599453>

Recurrent Ovarian Cancer – This is a Phase 2 investigator-sponsored trial being conducted in advanced recurrent ovarian cancer at the University of Pittsburgh Medical Center that will evaluate Ampligen in combination with pembrolizumab. Patient enrollment has been initiated in this study designed for 45 subjects. Dr. Robert Edwards, world renowned expert in ovarian cancer is the lead investigator. For more important details see: <https://clinicaltrials.gov/ct2/show/NCT03734692>

In addition, five Ampligen clinical trials are planned for initiation in 2019, subject to funding:

Phase 2 study that will evaluate Ampligen in combination with pembrolizumab in refractory metastatic colorectal carcinoma at Roswell Park Comprehensive Cancer Center. Dr. P. Boland, PI. Study design and budget being developed.

Phase 2 study of advanced urothelial (bladder), melanoma and renal cell carcinoma, resistant to checkpoint blockade, that will evaluate Ampligen in combination with a checkpoint blockade therapy at Roswell Park Comprehensive Cancer Center. Dr. M. Opyrchal, PI. Protocol design currently being finalized. Hemispherx Biopharma signed a clinical trial agreement with Roswell Park Comprehensive Cancer Center to study Ampligen in combination with checkpoint inhibitors in a phase 2a study in urothelial carcinoma, renal cell carcinoma and melanoma. This Phase 2a study will be led by Mateusz Opyrchal, MD, PhD, Assistant Professor of Medicine and Associate Director of the Early Phase Clinical Trial Program at Roswell Park, in collaboration with Dr. Kalinski.

First-line therapy for non-small cell lung cancer with SOC chemotherapy that will evaluate Ampligen in combination with pembrolizumab at University of Nebraska Medical Center. Dr. V. Ernani, PI. Study design and budget being developed.

Phase 2 study in advanced pancreatic cancer using checkpoint blockade plus Ampligen at University of Nebraska Medical Center. Dr. K. Klute, PI. Protocol and budget being developed. Based upon success in the initial animal studies, an additional round of more extensive and comprehensive pre-clinical animal pancreatic cancer studies are being conducted at University of Nebraska to reconfirm results, test additional PC tumor types, examine anti-PD-1 in addition to the prior anti-PD-L1 analysis, then fine tune the focus of the proposed future pancreatic cancer clinical trial and reduce the chances of error in clinical trial design. This information will also be used to formulate proposed future combination therapy clinical activity in the Kingdom of the Netherlands.

Phase 2 study of neoadjuvant conditioning of prostate cancer using Ampligen as a component of chemokine modulation at Roswell Park Comprehensive Cancer Center. Dr. G. Chatta, PI. This protocol is under review by the FDA.

In January 2017, the EAP through our agreement with myTomorrows designed to enable access of Ampligen® to ME/CFS patients had been extended to pancreatic cancer patients beginning in the Netherlands. myTomorrows is our exclusive service provider in Europe and Turkey and will manage all EAP activities relating to the pancreatic cancer extension of the program. In February 2018, the agreement with myTomorrows was extended to cover Canada to treat pancreatic cancer patients, pending government approval.

As of December 31, 2018, 40 pancreatic cancer patients have received treatment with Ampligen® immuno-oncology therapy under the EAP program at Erasmus University in the Netherlands.

Supervised by Prof. Casper van Eijck, MD, a world-renowned specialist in this dread malignancy, and Diba Latifi, MD, the team at Erasmus is making progress. As disclosed recently, the Dutch government has approved and extended the therapeutic program for an additional year. Early progress was reported in a published abstract from Erasmus, and a copy of the abstract can be found at http://ir.hemispherx.net/Events_Presentations. The abstract was part of a larger original report covering a variety of medical topics, which can be found at <https://www.pancreasclub.com/wp-content/uploads/2018/06/Poster-Abstracts.pdf>.

As of today, we are pleased to report that four out of 24 patients with either locally advanced or metastatic disease have survived for more than one year on the Ampligen protocol without additional therapy. Another four patients have survived for more than one year since the start of the Ampligen protocol with palliative chemotherapy. However, in this group of patients 15 died within seven months since start of Ampligen. Of the five resected patients two died on Ampligen, 24 and 27 months after resection. The other three patients are still alive with a mean survival of 26 months after resection and adjuvant Ampligen treatment.

All patients reported improvement in quality of life during treatment. We expect within 60 days a more comprehensive update from the Erasmus team on the immunological response in relation to survival. Hemispherx hopes to work with Dr. Van Eijck, Dr. Latifi, and Erasmus M.C. to initiate a combination therapy program to extend the results seen thus far in the Netherlands by combining Ampligen with checkpoint blockade therapy.

Myalgic Encephalomyelitis/Chronic Fatigue Syndrome (“ME/CFS”)

Myalgic Encephalomyelitis/Chronic Fatigue Syndrome (“ME/CFS”), also known as Chronic Fatigue Immune Dysfunction Syndrome (“CFIDS”) and Chronic Fatigue Syndrome (“CFS”), is a serious and debilitating chronic illness and a major public health problem. ME/CFS is recognized by both the government and private sector as a significant unmet medical need, including the U.S. National Institutes of Health (“NIH”), FDA and the CDC. The CDC states on its website at <https://www.cdc.gov/me-cfs/> that “*Myalgic encephalomyelitis/chronic fatigue syndrome (ME/CFS) is a serious, long-term illness that affects many body systems. People with ME/CFS are often not able to do their usual*

activities. At times, ME/CFS may confine them to bed. People with ME/CFS have severe fatigue and sleep problems. ME/CFS may get worse after people with the illness try to do as much as they want or need to do. This symptom is known as post-exertional malaise (PEM). Other symptoms can include problems with thinking and concentrating, pain, and dizziness.”

Many severe ME/CFS patients become completely disabled or totally bedridden and are afflicted with severe pain and mental confusion even at rest. ME/CFS is characterized by incapacitating fatigue with profound exhaustion and extremely poor stamina, sleep difficulties and problems with concentration and short-term memory. It is also accompanied by flu-like symptoms, pain in the joints and muscles, tender lymph nodes, sore throat and new headaches. A distinctive characteristic of the illness is a worsening of symptoms following physical or mental exertion, which do not subside with rest.

In October 2016, an analysis of a subset of CFS patients from the AMP-516 Phase 3 study was performed and presented at the IACFS/ME annual meeting in Fort Lauderdale, FL. The ITT Population (n=208) was separated into two subsets based primarily on baseline CFS symptom duration (2-8 years (n=75) and <2 years plus >8 years (n=133)). Responder analyses of the ITT Population and both subsets were performed. Responder analyses of rintatolimod vs. placebo patients improving ET duration from baseline by $\geq 25\%$ shows over twice the % of patients with clinical enhancement in ET effect in the rintatolimod cohort compared to placebo for the 2-8 year subset vs. the ITT population. This subset may assist in the design of future clinical studies of Ampligen® in the treatment for ME/CFS patients.

Other Diseases

In Europe, the EMA has approved the Orphan Medicinal Products Designation for rintatolimod (Ampligen®) as a potential treatment of Ebola virus disease and for Alferon® N Injection, also known as interferon alfa-n3, as a potential treatment of MERS.

We concluded our series of collaborations designed to determine the potential effectiveness of Ampligen® and Alferon® N as potential preventative and/or therapeutic treatments for Ebola related disorders. Although we believe that the threat of both MERS and Ebola globally may reemerge in the future, it appears that the spread of these disorders has somewhat diminished. As a result, we have elected to focus our research and development efforts on other areas at this time.

MANUFACTURING

In January 2017, Hemispherx approved a quote and provided a purchase order commitment with Jubilant Hollister-Stier LLC (“Jubilant”) pursuant to which Jubilant will manufacture commercial size batches of Ampligen®. Additional orders will be placed upon approved quotes and purchase orders provided by Hemispherx to Jubilant. Jubilant was approved by the FDA as a manufacturer of Ampligen by the successful completion of a previous preapproval inspection by the agency. The National Administration of Drugs, Food and Medical Devices (A.N.M.A.T) in Argentina has approved Ampligen for commercial distribution for the treatment of Chronic Fatigue Syndrome (CFS). Shipment of the drug product to Argentina was initiated in 2018 to complete the release testing by A.N.M.A.T. needed for commercial distribution.

Since the commencement of the 2017 commitment between Jubilant and Hemispherx, two lots of Ampligen consisting of more than 16,000 units have been manufactured and released. The first lot was designated for human use in the US in the cost recovery CFS program and for expanded oncology clinical trials. The second lot has been designated for these programs in addition to commercial distribution in Argentina for the treatment of CFS. Additional lots of Ampligen are being planned for manufacture at Jubilant. The production of additional polymer (Ampligen intermediates) at our New Brunswick facility is required to produce the additional lots of Ampligen. Polymer manufacture is on schedule to provide the intermediates needed for the Ampligen manufacture planned at Jubilant

Alferon® is approved by the FDA for commercial sales in the US for the treatment of genital warts. It is also approved by A.N.M.A.T in Argentina for commercial sales for the treatment of genital warts and in patients refractory to treatment with recombinant interferons. While the Hemispherx facility in New Brunswick is approved by the FDA under the Biologic License Application (BLA) for Alferon®, this status will need to be reaffirmed by an FDA pre-approval inspection which will not occur until new batches of commercial filled and finished product are

produced and released by the FDA. Currently, the manufacturing process is on hold and there is no definitive timetable to have the facility back online until additional funding is obtained.

Licensing/Collaborations/Joint Ventures

To maximize the availability of Ampligen® to patients on a worldwide basis, we have embarked on a strategy to license the product and/or to collaborate and/or create a joint venture with companies that have the demonstrated capabilities and commitment to successfully gain approval and commercialize Ampligen® in their respective territories of the world. Ideal partners would have the following characteristics: well established global and regional experience and coverage, robust commercial infrastructure, strong track record of successful development and registration of in-licensed products, as well as a therapeutic area fit (ME/CFS, immuno-oncology, etc.).

MARKETING/DISTRIBUTION

In May 2016, we entered into a five-year exclusive Renewed Sales, Marketing, Distribution and Supply Agreement (the “Agreement”) with GP Pharm. Under this Agreement, GP Pharm was responsible for gaining regulatory approval in Argentina for Ampligen® to treat severe CFS in Argentina and for commercializing Ampligen® for this indication in Argentina. We granted GP Pharm the right to expand rights to sell this experimental therapeutic into other Latin America countries based upon GP Pharm achieving certain performance milestones. We also granted GP Pharm an option to market Alferon N Injection® in Argentina and other Latin America countries.

In January 2017, the ANMAT granted a five-year extension to a previous approval to sell and distribute Alferon N Injection® (under the brand name “Naturaferon”) in Argentina. This extends the approval until 2022. In February 2013, we received the ANMAT approval for the treatment of refractory patients that failed or were intolerant to treatment with recombinant interferon, with Naturaferon® in Argentina.

In May 2016, we entered into a five-year agreement (the “Impatients Agreement”) with Impatients, N.V. (“myTomorrows”), a Netherlands based company, for the commencement and management of an EAP in Europe and Turkey (the “Territory”) related to ME/CFS. Pursuant to the agreement, myTomorrows, as our exclusive service provider and distributor in the Territory, is performing EAP activities. These activities will be directed to (a) the education of physicians and patients regarding the possibility of early access to innovative medical treatments not yet the subject of a Marketing Authorization (regulatory approval) through named-patient use, compassionate use, expanded access and hospital exemption, (b) patient and physician outreach related to a patient-physician platform, (c) the securing of Early Access Approvals (exemptions and/or waivers required by regulatory authorities for medical treatments prior to Marketing Authorization) for the use of such treatments, (d) the distribution and sale of such treatments pursuant to such Early Access Approvals, (e) pharmacovigilance (drug safety) activities and/or (f) the collection of data such as patient-reported outcomes, doctor-reported experiences and registry data. We are supporting these efforts and supplying Ampligen® to myTomorrows at a predetermined transfer price. In the event that we receive Marketing Authorization in any country in the Territory, we will pay myTomorrows a royalty on products sold. Pursuant to the Impatients Agreement, the royalty would be a percentage of Net Sales (as defined in the Impatients Agreement) of Ampligen® sold in the Territory where Marketing Authorization was obtained, and the maximum royalty would be a percentage of Net Sales. The formula to determine the percentage of Net Sales will be based on the number of patients that are entered into the EAP. The Company believes that disclosure of the exact maximum royalty rate and royalty termination date could cause competitive harm. However, to assist the public in gauging these terms, the actual maximum royalty rate is somewhere between 2% and 10% and the royalty termination date is somewhere between five and fifteen years from the First Commercial Sale of a product within a specific country. The parties established a Joint Steering Committee comprised of representatives of both parties to oversee the EAP. No assurance can be given that activities under the EAP will result in Marketing Authorization or the sale of substantial amounts of Ampligen® in the Territory.

In January 2017, the EAP through our agreement with myTomorrows designed to enable access of Ampligen® to ME/CFS patients has been extended to pancreatic cancer patients beginning in the Netherlands. myTomorrows is our exclusive service provider in the Territory and will manage all EAP activities relating to the pancreatic cancer extension of the program.

In February 2018, we signed an amendment to the EAP with myTomorrows. This amendment extended the territory to cover Canada to treat pancreatic cancer patients, pending government approval.

In March 2018, we signed an amendment to the EAP with myTomorrows, pursuant to which myTomorrows will be our exclusive service provider for special access activities in Canada for the supply of Ampligen® for the treatment of ME/CFS.

In August 2017, we extended our agreement with Asembia, formerly Armada Healthcare, LLC, to undertake the marketing, education and sales of Alferon N Injection® throughout the United States.

COMPETITION

The major pharmaceutical competitors for Ampligen include Pfizer, GlaxoSmithKline, Merck & Co., Novartis and AstraZeneca. Biotech competitors include Baxter International, Fletcher/CSI, AVANT Immunotherapeutics, AVI BioPharma and Genta. When we recommence sales of Alferon N Injection®, it will compete with Intron® A, an injectable from Merck & Co.

GOVERNMENT REGULATION

Regulation by governmental authorities in the U.S. and foreign countries is and will be a significant factor in the manufacture and marketing of Alferon® products and our ongoing research and product development activities. Ampligen® and other products developed from the ongoing research and product development activities will require regulatory clearances prior to commercialization. In particular, new drug products for humans are subject to rigorous pre-clinical and clinical testing as a condition for clearance by the FDA and by similar authorities in foreign countries. The process of seeking these approvals, and the ongoing process of compliance with applicable statutes and regulations, has and will continue to require the expenditure of substantial resources. Any failure by us or our collaborators or licensees to obtain, or any delay in obtaining, regulatory approvals could materially adversely affect the marketing of any products developed by us and our ability to receive product or royalty revenue. We have received Orphan Drug designation for certain therapeutic indications, which we believe might under certain conditions help to accelerate the process of drug development and commercialization. Alferon N Injection® is only approved for use in intralesional treatment of refractory or recurring external genital warts in patients 18 years of age or older. Use of Alferon N Injection® for other applications requires regulatory approval.

We are subject to various federal, state and local laws, regulations and recommendations relating to such matters as safe working conditions, laboratory and manufacturing practices, the experimental use of animals and the use of and disposal of hazardous or potentially hazardous substances, including infectious disease agents, used in connection with our research work. Prior to our construction phase, our laboratory and production facility in New Brunswick, New Jersey was approved for the manufacture of Alferon N Injection®. While our facility had been granted approval of its BLA by the FDA for the manufacture of Alferon®, this status will need to be reaffirmed as we have completed the facility's enhancements and believe, with adequate funding, it will again be able to obtain FDA approval.

For more information about the current status of Alferon N Injection® and Ampligen® please see “Our Products” above.

HUMAN RESOURCES

As of February 1, 2019, we had personnel consisting of 31 full-time employees and two part-time employees. Seventeen of the combined personnel are engaged in our research, development, clinical, and manufacturing effort with 16 performing regulatory, general administration, data processing, including bio-statistics, financial and investor relations functions. We have no union employees.

While we have been successful in attracting skilled and experienced scientific personnel, there can be no assurance that we will be able to attract or retain the necessary qualified employees and/or consultants in the future.

ITEM 1A: Risk Factors

The following cautionary statements identify important factors that could cause our actual results to differ materially from those projected in the forward-looking statements made in this Form 10-K. Please see “Special Note Regarding Forward Looking Statements” below.

Risks Associated with Our General Business

No assurance of successful product development and finding co-development partners.

We are committed to a focused business plan oriented toward finding co-development partners with the necessary capital and expertise required to commercialize the many therapeutic aspects of our experimental drugs and our FDA approved drug Alferon® N. If we are unable to find a suitable co-development partner to assist in the product development and commercialization of our experimental drugs and our FDA approved drug Alferon® N, we may be unable to continue or complete our development and commercialization of our products. In addition, there can be no assurance that such co-development partnerships would be on acceptable terms, or that such partnerships, will be acceptable from a profitability standpoint.

We will require additional financing which may not be available.

The development of our products requires the commitment of substantial resources to conduct the time consuming research, preclinical development, and clinical trials that are necessary to bring pharmaceutical products to market. As of December 31, 2018, we had approximately \$1,825,000 in cash, cash equivalents and marketable securities (inclusive of approximately \$1,526,000 in Marketable Securities). However, if we are unable to commercialize and sell Ampligen® and/or recommence material sales of Alferon N Injection®, our operations, financial position and liquidity may be adversely impacted.

Given the challenging economic conditions, we continue to review every aspect of our operations for cost and spending reductions to assure our long-term financial stability while maintaining the resources necessary to achieve our primary objectives of obtaining FDA approval of Ampligen® along with the manufacturing, marketing and distribution of our products, including Alferon N Injection®. Due to the high cost estimates to bring the facility back online, we will need additional funds to finance the revalidation process in our facility to initiate commercial manufacturing, thereby readying ourselves for an FDA Pre-Approval Inspection. We also will need additional capital to eventually commercialize and sell Ampligen® and/or recommence and increase sales of Alferon N Injection® or our other products. We anticipate considering multiple options in an attempt to secure funding, including but not limited to such methods as the sales of additional equity, licensing agreements, partnering with other organizations, debt financing or other sources of capital. If we are unable to obtain additional funding, through an Equity Distribution Agreement (“EDA”) or other sales of securities and/or otherwise, our ability to develop our products, commercially produce inventory or continue our operations may be materially adversely affected.

We may continue to incur substantial losses and our future profitability is uncertain.

As of December 31, 2018, our accumulated deficit was approximately \$318,573,000. As with many biotechnology companies we have not yet generated significant revenues from our products and may incur substantial and increased losses in the future. We cannot assure that we will ever achieve significant revenues from product sales or become profitable. We require, and will continue to require, the commitment of substantial resources to develop our products. We cannot assure that our product development efforts will be successfully completed or that required regulatory approvals will be obtained or that any products will be manufactured and marketed successfully, or be profitable.

Our drug and related technologies are investigational and subject to regulatory approval. If we are unable to obtain regulatory approval in a timely manner, or at all, our operations will be materially harmed and our stock adversely affected.

All of our drugs and associated technologies, other than Alferon N Injection®, are investigational in the U.S. and must receive prior regulatory approval by appropriate regulatory authorities for commercial distribution and sale and are currently legally available only through clinical trials in the U.S. with specified disorders. At present, Alferon N Injection® is approved for the intralesional treatment of refractory or recurring external genital warts in patients 18 years of age or older. Use of Alferon N Injection® for other indications will require regulatory approval in the U.S. and abroad.

Our products, including Ampligen®, are subject to extensive regulation by numerous governmental authorities in the U.S. and other countries, including, but not limited to, the FDA in the U.S., the Health Protection Branch (“HPB”) of Canada, the Agency for the European Medicines Agency (“EMA”) in Europe and the Administracion Nacional de Medicamentos, Alimentos y Tecnologia Medica (“ANMAT”) in Argentina. Obtaining regulatory approvals is a rigorous and lengthy process and requires the expenditure of substantial resources. In order to obtain final regulatory approval of a new drug, we must demonstrate to the satisfaction of the regulatory agency that the product is safe and effective for its intended uses and that we are capable of manufacturing the product to the applicable regulatory standards. We require regulatory approval in order to market Ampligen® or any other proposed product and receive product revenues or royalties. We cannot assure you that Ampligen® will ultimately be demonstrated to be safe and efficacious. While Ampligen® is authorized for use in clinical trials in the U.S., we cannot assure you that additional clinical trial approvals will be authorized in the United States or in other countries, in a timely fashion or at all, or that we will complete these clinical trials. In addition, although Ampligen® has been authorized by the FDA for treatment use under certain conditions, including provision for cost recovery, there can be no assurance that such authorization will continue in effect.

While we received approval of our Argentinian NDA from ANMAT for commercial sale of rintatolimod (U.S. tradename: Ampligen®) in the Argentine Republic for the treatment of severe ME/CFS, ANMAT approval is only an initial, but important, step in the overall successful commercialization of our product. There are a number of actions that must occur before we would be able to commence commercial sales in Argentina.

The FDA’s regulatory review and approval process is extensive, lengthy, expensive and inherently uncertain. To receive approval for a product candidate, we must, among other things, demonstrate to the FDA’s satisfaction with substantial evidence from well-controlled pre-clinical and clinical trials that the product candidate is both safe and effective for each indication for which approval is sought. Before we can sell Ampligen® for any use, or promote Alferon® for any use other than as Alferon N Injection® for treatment of refractory or recurring genital warts, we will need to file the appropriate NDA with the FDA in the U.S. and the appropriate regulatory agency outside of the U.S. where we intend to market and sell such products. At present the only NDA we have filed with the FDA is the NDA for the use of Ampligen® to treat CFS. The FDA issued a Complete Response Letter (“CRL”) in February 2013 for this NDA and provided recommendations to address certain outstanding issues before they could approve Ampligen for

Commercial Sales. The Agency stated that the submitted data do not provide substantial evidence of efficacy of Ampligen® for the treatment of CFS and that the data do not provide sufficient information to determine whether the product is safe for use in CFS due to the limited size of the safety database and multiple discrepancies within the submitted data. The FDA indicated that we needed to conduct additional work. Therefore, ultimate FDA approval, if any, may be delayed indefinitely and may require us to expend more resources than we have available. It is also possible that additional studies, if performed and completed, may not be successful or considered sufficient by the FDA for approval or even to make our applications approvable. If any of these outcomes occur, we may be forced to abandon one or more of our future applications for approval, which might significantly harm our business and prospects. As a result, we cannot predict if or when we might receive regulatory approval for the use of Ampligen® to treat CFS or for the use of any other products. Even if regulatory approval from the FDA is received for the use of Ampligen® to treat CFS or eventually, for the use of any other product, any approvals that we obtain could contain significant limitations in the form of narrow indications, patient populations, warnings, precautions or contra-indications or other conditions of use, or the requirement that we implement a risk evaluation and mitigation strategy. In such an event, our ability to generate revenues from such products could be greatly reduced and our business could be harmed.

If we are unable to gain necessary FDA approvals related to Ampligen® and Alferon® on a timely basis, or we are unable to generate the additional data, successfully complete inspections or obtain approvals as required by the FDA on a timely manner, or at all, or determine that any of our clinical studies are not cost/justified to undertake or if, for that or any other reason, Ampligen®, Alferon® or one of our other products or production processes do not receive necessary regulatory approval in the U.S. or elsewhere, our operations most likely will be materially and/or adversely affected.

Generally, obtaining approval of a NDA by the FDA, or a comparable foreign regulatory authority, is inherently uncertain. Even after completing clinical trials and other studies, a product candidate could fail to receive regulatory approval for many reasons, including the following:

- not be able to demonstrate to the satisfaction of the FDA that our product candidate is safe and effective for any indication;
- the FDA may disagree with the design or implementation of our clinical trials or other studies;
- the results of the clinical trials or other studies may not demonstrate that a product candidate's clinical and other benefits outweigh its safety risks;
- the FDA may disagree with our interpretation of data from clinical trials or other studies;
- the data collected from clinical trials and other studies of a product candidate may not be sufficient to support the submission of a NDA;
- the approval policies or regulations of the FDA may significantly change in a manner rendering our clinical and other study data insufficient for approval; and
- the FDA may not approve the proposed manufacturing processes and facilities for a product candidate.

We may be subject to product liability claims from the use of Ampligen®, Alferon N Injection®, or other of our products which could negatively affect our future operations. We have limited product liability and clinical trial insurance.

We maintain a limited amount of Products Liability and Clinical Trial insurance coverage world-wide for Ampligen® and Alferon® due to the minimal amount of historical loss claims regarding these products in the marketplace. Any claims against our products, Ampligen® and Alferon N Injection®, could have a materially adverse effect on our business and financial condition.

We face an inherent business risk of exposure to product liability claims in the event that the use of Ampligen®, Alferon N Injection® or other of our products results in adverse effects. This liability might result from claims made directly by patients, hospitals, clinics or other consumers, or by pharmaceutical companies or others manufacturing these products on our behalf. Our future operations may be negatively affected from the litigation costs, settlement expenses and lost product sales inherent to these claims. While we will continue to attempt to take appropriate precautions, we cannot assure that we will avoid significant product liability exposure.

Uncertainty of health care reimbursement for our products.

Our ability to successfully commercialize our products will depend, in part, on the extent to which reimbursement for the cost of such products and related treatment will be available from government health administration authorities, private health coverage insurers and other organizations. Significant uncertainty exists as to the reimbursement status of newly approved health care products, and from time to time legislation is proposed, which, if adopted, could further restrict the prices charged by and/or amounts reimbursable to manufacturers of pharmaceutical products. We cannot predict what, if any, legislation will ultimately be adopted or the impact of such legislation on us. There can be no assurance that third party insurance companies will allow us to charge and receive payments for products sufficient to realize an appropriate return on our investment in product development.

There are risks of liabilities associated with handling and disposing of hazardous materials.

Our business involves the controlled use of hazardous materials, carcinogenic chemicals, and flammable solvents. Although we believe that our safety procedures for handling and disposing of such materials comply in all material respects with the standards prescribed by applicable regulations, the risk of accidental contamination or injury from these materials cannot be completely eliminated. In the event of such an accident or the failure to comply with applicable regulations, we could be held liable for any damages that result. However, we have obtained insurance coverage to mitigate any potential significant loss in this area.

We rely upon information technology and any failure, inadequacy, interruption or security lapse of that technology, including any cyber security incidents, could harm our ability to operate our business effectively.

Despite the implementation of security measures, our internal computer systems and those of third parties with which we contract are vulnerable to damage from cyber-attacks, computer viruses, unauthorized access, natural disasters, terrorism, war and telecommunication and electrical failures. System failures, accidents or security breaches could cause interruptions in our operations, and could result in a material disruption of our business operations, in addition to possibly requiring substantial expenditures of resources to remedy. The loss of clinical trial data could result in delays in our regulatory approval efforts and significantly increase our costs to recover or reproduce the data. To the extent that any disruption or security breach were to result in a loss of, or damage to, our data or applications, or inappropriate public disclosure of confidential or proprietary information, we could incur liability and our product development and commercialization efforts could be delayed.

The loss of services of key personnel could hurt our chances for success.

Our success is dependent on the continued efforts of our staff, especially certain doctors and researchers. The loss of the services of personnel key to our operations could have a material adverse effect on our operations and chances for success. The loss of key personnel or the failure to recruit additional personnel as needed could have a materially adverse effect on our ability to achieve our objectives.

Risks Associated with Our Products

In addition to the risks disclosed above, the development of Ampligen® is subject to a number of significant risks. Ampligen® may be found to be ineffective or to have adverse side effects, fail to receive necessary regulatory clearances, be difficult to manufacture on a commercial scale, be uneconomical to market or be precluded from commercialization by proprietary right of third parties. Our investigational products are in various stages of clinical and pre-clinical development and require further clinical studies and appropriate regulatory approval processes before any such products can be marketed. We do not know when, if ever, Ampligen® or our other products will be generally available for commercial sale for any indication. Generally, only a small percentage of potential therapeutic products are eventually approved by the FDA for commercial sale.

To the extent that we are required by the FDA, pursuant to the Ampligen® NDA, to conduct additional studies and take additional actions, approval of any applications that we submit may be delayed by several years, or may require us to expend more resources than we have available. It is also possible that additional studies, if performed and completed, may not be successful or considered sufficient by the FDA for approval or even to make our applications approvable. If any of these outcomes occur, we may be forced to abandon one or more of our future applications for approval, which might significantly harm our business and prospects. As a result, we cannot predict when or whether regulatory approval will be obtained for any product candidate we develop.

Ampligen®. We believe that Ampligen® has been generally well tolerated with a low incidence of clinical toxicity, particularly given the severely debilitating or life threatening diseases that have been treated. A mild flushing reaction has been observed in approximately 15-20% of patients treated in our various studies. This reaction is occasionally accompanied by a rapid heartbeat, a tightness of the chest, urticaria (swelling of the skin), anxiety, shortness of breath, subjective reports of “feeling hot”, sweating and nausea. The reaction is usually infusion-rate related and can generally be controlled by reducing the rate of infusion. Other adverse side effects include liver enzyme level elevations, diarrhea, itching, asthma, low blood pressure, photophobia, rash, visual disturbances, slow or irregular heart rate, decreases in platelets and white blood cell counts, anemia, dizziness, confusion, elevation of kidney function tests, occasional temporary hair loss and various flu-like symptoms, including fever, chills, fatigue, muscular aches, joint pains, headaches, nausea and vomiting. These flu-like side effects typically subside within several months.

If approved, one or more of the potential side effects of the drug might deter usage of Ampligen® in certain clinical situations and, therefore, could adversely affect potential revenues and physician/patient acceptability of our product.

Alferon N Injection®. Although Alferon N Injection® is approved for marketing in the United States for intralesional treatment of refractory or recurring external genital warts in patients 18 years of age or older, to date it has not been approved for other indications.

Possible side effects from the use of Ampligen® or Alferon N Injection® could adversely affect potential revenues and physician/patient acceptability of our product.

Ampligen®. We believe that Ampligen® has been generally well tolerated with a low incidence of clinical toxicity, particularly given the severely debilitating or life threatening diseases that have been treated. A mild flushing reaction has been observed in approximately 15-20% of patients treated in our various studies. This reaction is occasionally accompanied by a rapid heartbeat, a tightness of the chest, urticaria (swelling of the skin), anxiety, shortness of breath, subjective reports of “feeling hot”, sweating and nausea. The reaction is usually infusion-rate related and can generally be controlled by reducing the rate of infusion. Other adverse side effects include liver enzyme level elevations, diarrhea, itching, asthma, low blood pressure, photophobia, rash, visual disturbances, slow or irregular heart rate, decreases in platelets and white blood cell counts, anemia, dizziness, confusion, elevation of kidney function tests, occasional temporary hair loss and various flu-like symptoms, including fever, chills, fatigue, muscular aches, joint pains, headaches, nausea and vomiting. These flu-like side effects typically subside within several months.

The FDA in its February 1, 2013 CRL, provided recommendations to address certain outstanding issues before they could approve Ampligen for Commercial Sales. The Agency stated that the submitted data do not provide sufficient information to determine whether the product is safe for use in CFS due to the limited size of the safety database and multiple discrepancies within the submitted data.

If approved, one or more of the potential side effects of the drug might deter usage of Ampligen® in certain clinical situations and therefore, could adversely affect potential revenues and physician/patient acceptability of our product.

Alferon N Injection®. At present, Alferon N Injection® is approved for the intralesional (within the lesion) treatment of refractory or recurring external genital warts in adults. In clinical trials conducted for the treatment of genital warts with Alferon N Injection®, patients did not experience serious side effects; however, there can be no assurance that unexpected or unacceptable side effects will not be found in the future for this use or other potential uses of Alferon N Injection® which could threaten or limit such product’s usefulness.

Risks Associated with Our Intellectual Property

We may not be profitable unless we can protect our patents and/or receive approval for additional pending patents.

We need to preserve and acquire enforceable patents covering the use of Ampligen® for a particular disease in order to obtain exclusive rights for the commercial sale of Ampligen® for such disease. We obtained all rights to Alferon N Injection®, and we plan to preserve and acquire enforceable patents covering its use for existing and potentially new diseases once we have had a successful FDA Pre Approval Inspection. Our success depends, in large part, on our ability to preserve and obtain patent protection for our products and to obtain and preserve our trade secrets and expertise. Certain of our know-how and technology is not patentable, particularly the procedures for the manufacture of our experimental drug, Ampligen®. We also have been issued a patent which affords protection on the use of Ampligen® in patients with Chronic Fatigue Syndrome. We have not yet been issued any patents in the United States for the use of Ampligen® as a sole treatment for any of the cancers which we have sought to target. For more information on Patents, please see PART I, Item 1 – “Business; Patents”.

We cannot assure that our competitors will not seek and obtain patents regarding the use of similar products in combination with various other agents, for a particular target indication prior to our doing so. If we cannot protect our patents covering the use of our products for a particular disease, or obtain additional patents, we may not be able to successfully market our products.

The patent position of biotechnology and pharmaceutical firms is highly uncertain and involves complex legal and factual questions.

To date, no consistent policy has emerged regarding the breadth of protection afforded by pharmaceutical and biotechnology patents. There can be no assurance that new patent applications relating to our products, process or technology will result in patents being issued or that, if issued, such patents will afford meaningful protection against competitors with similar technology. It is generally anticipated that there may be significant litigation in the industry regarding patent and intellectual property rights. Such litigation could require substantial resources from us and we may not have the financial resources necessary to enforce the patent rights that we hold. No assurance can be made that our patents will provide competitive advantages for our products, process and technology or will not be successfully challenged by competitors. No assurance can be given that patents do not exist or could not be filed which would have a materially adverse effect on our ability to develop or market our products or to obtain or maintain any competitive position that we may achieve with respect to our products. Our patents also may not prevent others from developing competitive products or processes using related technology.

There can be no assurance that we will be able to obtain necessary licenses if we cannot enforce patent rights we may hold. In addition, the failure of third parties from whom we currently license certain proprietary information or from whom we may be required to obtain such licenses in the future, to adequately enforce their rights to such proprietary information, could adversely affect the value of such licenses to us.

If we cannot enforce the patent rights we currently hold we may be required to obtain licenses from others to develop, manufacture or market our products. There can be no assurance that we would be able to obtain any such licenses on commercially reasonable terms, if at all. We currently license certain proprietary information from third parties, some of which may have been developed with government grants under circumstances where the government maintained certain rights with respect to the proprietary information developed. No assurances can be given that such third parties will adequately enforce any rights they may have or that the rights, if any, retained by the government will not adversely affect the value of our license.

There is no guarantee that our trade secrets will not be disclosed or known by our competitors.

To protect our rights, we require all employees and certain consultants to enter into confidentiality agreements with us. There can be no assurance that these agreements will not be breached, that we would have adequate and enforceable remedies for any breach, or that any trade secrets of ours will not otherwise become known or be independently developed by competitors.

Risks Associated with Our R&D

Due to the inherent uncertainty involved in the design and conduct of clinical trials and the applicable regulatory requirements, including the factors discussed above in “Our Products”, we cannot predict what additional studies and/or additional testing or information may be required by the FDA. Accordingly, we are unable to estimate the nature, timing, costs and necessary efforts to complete these projects nor the anticipated completion dates. In addition, we have no basis for estimating when material net cash inflows may commence. We have yet to generate significant revenues from the sale of these developmental products. As of December 31, 2018, we had approximately \$1,825,000 in Cash, Cash Equivalents and Marketable Securities, (inclusive of approximately \$1,526,000 in Marketable Securities). Please see “*We will require additional financing which may not be available*” above.

Risks Associated with Our Manufacturing

Our Alferon N Injection® Commercial Sales were halted due to lack of finished goods inventory. If we are unable to gain the necessary FDA approvals related to Alferon®, our operations most likely will be materially and/or adversely affected.

While our facility is FDA approved under the BLA by the FDA for Alferon®, this status will need to be reaffirmed upon the completion of the facility's upgrades for Alferon®. We cannot provide any guarantee that the facility will necessarily pass a FDA pre-approval inspection for Ampligen® or Alferon® manufacture, which are conducted in separately dedicated areas within the overall New Brunswick manufacturing complex.

If we are unable to gain the necessary FDA approvals related to the manufacturing process and/or final product of new Alferon® inventory, our operations most likely will be materially and/or adversely affected. For more information on Alferon N Injection® regarding potential commercial sales, please see PART I, Item 1 - "Business; Manufacturing".

There are no long-term agreements with suppliers of required materials and services for Ampligen® and there are a limited number of raw material suppliers. If we are unable to obtain the required raw materials and/or services, we may not be able to manufacture Ampligen®.

A number of essential raw materials are used in the production of Ampligen® as well as packaging materials utilized in the fill and finish process. We do not have, but continue to work towards having long-term agreements for the supply of such materials, when possible. There can be no assurance we can enter into long-term supply agreements covering essential materials on commercially reasonable terms, if at all.

There are a limited number of suppliers in the United States and abroad available to provide the raw and packaging materials/reagents for use in manufacturing Ampligen® and Alferon®. At present, we do not have any agreements with third parties for the supply of any of these materials or we are relying on a limited source of reagent suppliers necessary for the manufacture of Alferon®. In January 2017, we approved a quote and provided a purchase order with Jubilant Hollister-Stier LLC ("Jubilant") pursuant to which Jubilant manufactured batches of Ampligen® for us. We anticipate that additional orders will be placed upon approved quotes and purchase orders provided by Hemispherx to Jubilant. If we are unable to place adequate acceptable purchase orders with Jubilant in the future at acceptable prices upon acceptable terms, we will need to find another manufacturer. If we need to find another contract manufacturer to produce Ampligen, it would create a significant delay and expense to get the manufacturer up and running. The costs and availability of products and materials we would need for the production of Ampligen® are subject to fluctuation depending on a variety of factors beyond our control, including competitive factors, changes in technology, ownership of intellectual property, FDA and other governmental regulations. There can be no assurance that we will be able to

obtain such products and materials on terms acceptable to us or at all.

We have established relevant manufacturing operations within our New Brunswick, New Jersey facility for the production of Ampligen® polymers from raw materials in order to obtain a more consistent manufacturing basis in the quantities necessary for clinical testing.

Currently, the Alferon manufacturing process is on hold until additional funding is attained; there is no definitive timetable to have the facility back online. If we are unable to gain the necessary funding and FDA approvals related to the manufacturing process and/or final product of new Alferon® inventory, our operations most likely will be materially and/or adversely affected. In light of these contingencies, there can be no assurances that the approved Alferon N Injection® product will be returned to production on a timely basis, if at all, or that if and when it is again made commercially available, it will return to prior sales levels.

If we are unable to obtain or manufacture the required materials/reagents, and/or procure services needed in the final steps in the manufacturing process, we may be unable to manufacture Ampligen®. The costs and availability of products and materials we need for the production of Ampligen® are subject to fluctuation depending on a variety of factors beyond our control, including competitive factors, changes in technology, ownership of intellectual property, FDA and other governmental regulations. There can be no assurance that we will be able to obtain such products and materials on terms acceptable to us or at all. For more information on Ampligen® manufacturing, please see PART I, Item 1 - “Business; Our Products; Manufacturing” above.

There are a limited number of organizations in the United States available to provide the final manufacturing steps of formulation, fill, finish and packing sets for Alferon N Injection® and Ampligen®.

There are a limited number of organizations in the United States available to provide the final steps in the manufacturing for Alferon N Injection® and Ampligen®. To formulate, fill, finish and package our products (“fill and finish”), we require a FDA approved third party CMO.

In January 2017, we approved a quote and provided a purchase order with Jubilant Hollister-Stier LLC (“Jubilant”) pursuant to which Jubilant manufactured batches of Ampligen® for us. We anticipate that additional orders will be placed upon approved quotes and purchase orders provided by Hemispherx to Jubilant. If we are unable to place adequate acceptable purchase orders with Jubilant in the future at acceptable prices upon acceptable terms our business would be materially and adversely affected. Please see the prior risk factor.

Should there be an unanticipated delay in receiving new product or should we experience an unexpected demand for Ampligen®, our ability to supply Ampligen® most likely will be adversely affected. If we are unable to procure services needed in the final steps in the manufacturing process, we may be unable to manufacture Alferon N Injection® and/or Ampligen®. The costs and availability of products and materials we need for the production of Ampligen® and the commercial production of Alferon N Injection® and other products which we may commercially produce are subject to fluctuation depending on a variety of factors beyond our control, including competitive factors, changes in technology, and FDA and other governmental regulations and there can be no assurance that we will be able to obtain such products and materials on terms acceptable to us or at all. For more information on Ampligen® and Alferon N Injection® manufacturing, please see PART I, Item 1 - “Business; Our Products; Manufacturing” above.

There is no assurance that upon successful manufacture of a drug on a limited scale basis for investigational use will lead to a successful transition to commercial, large-scale production.

Changes in methods of manufacturing, including commercial scale-up, may affect the chemical structure of Ampligen® and other RNA drugs, as well as their safety and efficacy. The transition from limited production of pre-clinical and clinical research quantities to production of commercial quantities of our products will involve distinct management and technical challenges and may require additional management, technical personnel and capital to the extent such manufacturing is not handled by third parties. While we believe that we could successfully upgrade our production capability at our New Brunswick, NJ facility in a commercial scale-up of Ampligen®, there can be no assurance that our manufacturing will be successful or that any given product will be determined to be safe and effective, or capable of being manufactured under applicable quality standards, economically, and in commercial quantities, or successfully marketed.

We have limited manufacturing experience for Ampligen® and Alferon®. We may not be profitable unless we can produce Ampligen®, Alferon® or other products in commercial quantities at costs acceptable to us.

Ampligen® has been produced to date in limited quantities for use in our clinical trials and Early Access Programs. To be successful, our products must be manufactured in commercial quantities in compliance with regulatory requirements and at acceptable costs. We believe, but cannot assure, that our enhancements to our manufacturing facilities will be adequate for our future needs for the production of our proposed products for large-scale commercialization. We intend to ramp up our existing facility and/or utilize third party facilities if and when the need arises or, if we are unable to do so, to build or acquire commercial-scale manufacturing facilities. We will need to comply with regulatory requirements for such facilities, including those of the FDA pertaining to cGMP requirements or maintaining our BLA status. There can be no assurance that such facilities can be used, built, or acquired on commercially acceptable terms, or that such facilities, if used, built, or acquired, will be adequate for the production of our proposed products for large-scale commercialization or our long-term needs.

We have never produced Ampligen®, Alferon® or any other products in large commercial quantities. We must manufacture our products in compliance with regulatory requirements in large commercial quantities and at acceptable costs in order for us to be profitable. We intend to utilize third party manufacturers and/or facilities if and when the need arises or, if we are unable to do so, to build or acquire commercial-scale manufacturing facilities. If we cannot manufacture commercial quantities of Ampligen® and/or Alferon®, or continue to maintain third party agreements for its manufacture at costs acceptable to us, our operations will be significantly affected. If and when the Ampligen® NDA is approved, we may need to find an additional vendor to manufacture the product for commercial sales. Also, each production lot of Alferon N Injection® is subject to FDA review and approval prior to releasing the lots to be sold. This review and approval process could take considerable time, which would delay our having product in inventory to sell, nor can we provide any assurance as to the receipt of FDA approval of our finished inventory product. There can be no assurances that the Ampligen® and/or Alferon® can be commercially produced at costs acceptable to us.

Risks Associated with Our Licensing/Collaborations/Joint Ventures

If we are unable to achieve licensing, collaboration and/or joint ventures, our marketing strategy for Ampligen will be part of the differing health care systems around the world along with the different marketing and distribution systems that are used to supply pharmaceutical products to those systems.

We have received approval of our NDA from ANMAT for commercial sale of rintatolimod (U.S. tradename: Ampligen®) in the Argentine Republic for the treatment of severe CFS. The product will be marketed by GP Pharm, our commercial partner in Latin America. Commercialization in Argentina will require, among other things, GP Pharm to establish disease awareness, medical education, creation of an appropriate reimbursement level, design of marketing strategies and completion of manufacturing preparations for launch.

Risks Associated with Our Marketing and Distribution

We have limited marketing and sales capability. If we are unable to obtain additional distributors and our current and future distributors do not market our products successfully, we may not generate significant revenues or become profitable.

We have limited marketing and sales capability. We are dependent upon existing and, possibly future, marketing agreements and third party distribution agreements for our products in order to generate significant revenues and become profitable. As a result, any revenues received by us will be dependent in large part on the efforts of third parties, and there is no assurance that these efforts will be successful.

Our commercialization strategy for Ampligen®, if and when it is approved for marketing and sale by the FDA, may include licensing/co-marketing agreements utilizing the resources and capacities of a strategic partner(s). We continue to seek a world-wide marketing partner with the goal of having a relationship in place before approval is obtained. In parallel to partnering discussions, appropriate pre-marketing activities will be undertaken. It is our current intention to control manufacturing of Ampligen® on a world-wide basis.

Our commercialization strategy for Alferon N Injection® may include the utilization of internal functions and/or licensing/co-marketing agreements that would utilize the resources and capacities of one or more strategic partners. Accordingly, we have engaged Asembia, formerly Armada Healthcare, LLC, to undertake the marketing, education and sales of Alferon N Injection® throughout the United States along with GP Pharm for both Ampligen® and Alferon® in Argentina along with other South American countries.

We cannot assure that our U.S. or foreign marketing strategy will be successful or that we will be able to establish future marketing or third party distribution agreements on terms acceptable to us, or that the cost of establishing these arrangements will not exceed any product revenues. Our inability to establish viable marketing and sales capabilities would most likely have a materially adverse effect on us. There can be no assurances that the approved Alferon N Injection® product will be returned to prior sales levels.

Risks Associated with Our Competition

Rapid technological change may render our products obsolete or non-competitive.

The pharmaceutical and biotechnology industries are subject to rapid and substantial technological change. Technological competition from pharmaceutical and biotechnology companies, universities, governmental entities and others diversifying into the field is intense and is expected to increase. Most of these entities have significantly greater research and development capabilities than us, as well as substantial marketing, financial and managerial resources, and represent significant competition for us. There can be no assurance that developments by others will not render our products or technologies obsolete or noncompetitive or that we will be able to keep pace with technological developments.

Our products may be subject to substantial competition.

Ampligen®. Competitors may be developing technologies that are, or in the future may be, the basis for competitive products. Some of these potential products may have an entirely different approach or means of accomplishing similar therapeutic effects to products being developed by us. These competing products may be more effective and less costly than our products. In addition, conventional drug therapy, surgery and other more familiar treatments may offer competition to our products. Furthermore, many of our competitors have significantly greater experience than we do in preclinical testing and human clinical trials of pharmaceutical products and in obtaining Food and Drug Administration (FDA), The Health Protection Branch of the Canada Department of National Health and Welfare (HPB) and other regulatory approvals of products. Accordingly, our competitors may succeed in obtaining FDA, HPB or other regulatory product approvals more rapidly than us. There are no drugs approved for commercial sale with respect to treating CFS in the United States. The dominant competitors with drugs to treat disease indications which we plan to address include Pfizer, GlaxoSmithKline, Merck & Co., Novartis and AstraZeneca. Biotech competitors include Baxter International, Fletcher/CSI, AVANT Immunotherapeutics, AVI BioPharma and Genta. These potential competitors are among the largest pharmaceutical companies in the world, are well known to the public and the medical community, and have substantially greater financial resources, product development, and manufacturing and marketing capabilities than we have. Although we believe our principal advantage is the unique mechanism of action of Ampligen® on the immune system, we cannot assure that we will be able to compete.

Alferon N Injection®. Our competitors are among the largest pharmaceutical companies in the world, are well known to the public and the medical community, and have substantially greater financial resources, product development, and manufacturing and marketing capabilities than we have. Alferon N Injection® currently competes with Merck's injectable recombinant alpha interferon product (Intron® A) for the treatment of genital warts. In addition, other pharmaceutical firms offer self-administered topical cream, for the treatment of external genital and perianal warts such as Graceway Pharmaceuticals (Aldara®), Perrigo Company (Imiquimod Cream - Generic Equivalent to Aldara®), Watson Pharma (Condylox®) and MediGene (Veregen®). Alferon N Injection® also competes with surgical, chemical, and other methods of treating genital warts. We cannot assess the impact products developed by our competitors, or advances in other methods of the treatment of genital warts, will have on the commercial viability of Alferon N Injection®. If and when we obtain additional approvals of uses of this product, we expect to compete primarily on the basis of product performance. Our competitors have developed or may develop products (containing either alpha or beta interferon or other therapeutic compounds) or other treatment modalities for those uses. There can be no assurance that, if we are able to obtain regulatory approval of Alferon N Injection® for the treatment of new indications, we will be able to achieve any significant penetration into those markets. In addition, because certain competitive products are not dependent on a source of human blood cells, such products may be able to be produced in greater volume and at a lower cost than Alferon N Injection®. Currently, our wholesale price on a per unit basis of Alferon N Injection® is higher than that of the competitive recombinant alpha and beta interferon products. Please see risk factor "We may not be profitable unless we can protect our patents and/or receive approval for additional pending patents" above for additional information.

Other companies may succeed in developing products earlier than we do, obtaining approvals for such products from the FDA more rapidly than we do, or developing products that are more effective than those we may develop. While we will attempt to expand our technological capabilities in order to remain competitive, there can be no assurance that research and development by others or other medical advances will not render our technology or products obsolete or non-competitive or result in treatments or cures superior to any therapy we develop.

Risks Associated with an Investment in Our Common Stock:

The market price of our stock may be adversely affected by market volatility.

The market price of our common stock has been and is likely to be volatile. This is especially true given the current significant instability in the financial markets. In addition to general economic, political and market conditions, the price and trading volume of our stock could fluctuate widely in response to many factors, including:

announcements of the results of clinical trials by us or our competitors;

announcements of availability or projections of our products for commercial sale;

announcements of legal actions against us and/or settlements or verdicts adverse to us;

adverse reactions to products;

governmental approvals, delays in expected governmental approvals or withdrawals of any prior governmental approvals or public or regulatory agency comments regarding the safety or effectiveness of our products, or the adequacy of the procedures, facilities or controls employed in the manufacture of our products;

changes in U.S. or foreign regulatory policy during the period of product development;

developments in patent or other proprietary rights, including any third party challenges of our intellectual property rights;

announcements of technological innovations by us or our competitors;

announcements of new products or new contracts by us or our competitors;

actual or anticipated variations in our operating results due to the level of development expenses and other factors;

changes in financial estimates by securities analysts and whether our earnings meet or exceed the estimates;

conditions and trends in the pharmaceutical and other industries;

new accounting standards;

overall investment market fluctuation;

restatement of prior financial results;

notice of NYSE American non-compliance with requirements; and

occurrence of any of the risks described in these “Risk Factors”.

Our common stock is listed for quotation on the NYSE American. For the year ended December 31, 2018, the trading price of our common stock has ranged from \$0.18 to \$0.65 per share. We expect the price of our common stock to remain volatile. The average daily trading volume of our common stock varies significantly.

Our stock price may be adversely affected if a significant amount of shares is sold in the public market.

We may issue shares to be used to meet our capital requirements or use shares to compensate employees, consultants and/or Directors. In this regard, we completed a rights offering to our stockholders and certain option and warrant holders in March 2019, pursuant to which we issued Preferred stock convertible into an aggregate of 26,560,000 shares of common stock and warrants exercisable for up to an additional 26,560,000 shares of common stock. All of these shares of common stock have been registered for public sale. In addition, we have registered securities for public sale pursuant to a universal shelf registration statement and we had been selling shares under this shelf registration statement. Since December 5, 2017, we have sold an aggregate of 2,970,273 shares under our equity distribution agreements with Maxim Group LLC. In September 2016, we sold 3,333,334 shares of our common stock and issued warrants to purchase 2,500,000 shares of common stock. The warrants were exercised in June and July 2017. In February 2017 we sold 1,818,185 shares of our common stock and issued warrants. In February 2017, these warrants were exchanged for warrants to purchase an aggregate of 5,300,000 shares of common stock at an exercise price of \$0.45 per share, most exercisable commencing December 1, 2017. We have registered the shares issuable upon exercise of these warrants for public sale and, should the market price of our common stock exceed the exercise price of these warrants, some or all of these warrants may be exercised. There were 2,800,000 warrants with an expiration date of March 1, 2018 and an exercise price on \$0.45. These warrants were exercised in January and February 2018. We realized proceeds of \$1,260,000 from these exercises.

We are unable to estimate the amount, timing or nature of future sales of outstanding common stock or instruments convertible into or exercisable for our common stock. Sales of substantial amounts of our common stock in the public market, including additional sale of securities pursuant to our equity distribution agreements with Maxim Group LLC or otherwise under the universal shelf registration statement or upon exercise of outstanding options and warrants, could cause the market price for our common stock to decrease. Furthermore, a decline in the price of our common stock would likely impede our ability to raise capital through the issuance of additional shares of common stock or other equity securities. Please see Item 7- "Management's Discussion and Analysis of Financial Condition and Result of Operations; Liquidity and Capital Resources" in PART II.

The trading price of our common stock has decreased significantly and, as a result, the NYSE American has informed us that we are not in compliance with the standards for continued listing on the NYSE American. If we are unable to raise the trading price, the market for our common stock most likely will be adversely affected.

The price per share of our common stock has closed at or below \$0.20 since February 26, 2019 and most recently closed on March 26, 2019 at \$0.16, with a 30 day average of \$0.19. On March 26, 2016, we received written notice

from the NYSE American LLC (the “NYSE American”) that we are not in compliance with the continued listing standards set forth in Section 1003(f)(v) of the NYSE American Company Guide because our common stock has been selling for a low price per share for a substantial period of time. The NYSE American has determined that the continued listing of our common stock is predicated on us effecting a reverse stock split of our common stock or otherwise demonstrating sustained price improvement within a reasonable period of time. We have until September 26, 2019 to demonstrate compliance. Please see PART II, Item 9B – “Other Information” for a discussion of our plans to regain compliance. No assurance can be given that our plans will prove successful.

If we are unable to sufficiently raise the trading price of our common stock, we risk delisting on the NYSE American. Should our stock be so delisted, stockholders’ ability to sell their shares in the open market most likely will be adversely affected even if the shares are then quoted for trading on an interdealer quotation system such as the OTCBB or OTC Markets.

Provisions of our Certificate of Incorporation and Delaware law could defer a change of our Management which could discourage or delay offers to acquire us.

Provisions of our Certificate of Incorporation and Delaware law may make it more difficult for someone to acquire control of us or for our stockholders to remove existing management, and might discourage a third party from offering to acquire us, even if a change in control or in Management would be beneficial to our stockholders. For example, our Certificate of Incorporation allows us to issue shares of preferred stock without any vote or further action by our stockholders. Our Board of Directors has the authority to fix and determine the relative rights and preferences of preferred stock. Our Board of Directors also has the authority to issue preferred stock without further stockholder approval. As a result, our Board of Directors could authorize the issuance of a series of preferred stock that would grant to holders the preferred right to our assets upon liquidation, the right to receive dividend payments before dividends are distributed to the holders of common stock and the right to the redemption of the shares, together with a premium, prior to the redemption of our common stock. On November 14, 2017, at the direction of the Board, we amended and restated the Rights Agreement between the Company and, American Stock Transfer & Trust Company, LLC, its current Rights Agent. Pursuant to the original Rights Agreement, our Board of Directors declared a dividend distribution of one Right for each outstanding share of Common Stock to stockholders of record at the close of business on November 29, 2002. Each Right entitles the registered holder to purchase from the Company a unit consisting of one one-hundredth of a share (a “Unit”) of Series A Junior Participating Preferred Stock, par value \$0.01 per share at a Purchase Price of \$21.00 per Unit, subject to adjustment.

Special Note Regarding Forward Looking Statements

Certain statements in this Report contain forward-looking statements within the meaning of Section 27A of the Securities Act and Section 21E of the Securities Exchange Act of 1934, as amended, which we refer to as the Exchange Act. These statements are based on our management's current beliefs, expectations and assumptions about future events, conditions and results and on information currently available to us. Discussions containing these forward-looking statements may be found, among other places, in this "Risk Factors" section; Item 1. "Business", Part I; Item 3. "Legal Proceedings" and Part II; Item 7. "Management's Discussion and Analysis of Financial Condition and Results of Operations".

All statements, other than statements of historical fact, included or incorporated herein regarding our strategy, future operations, financial position, future revenues, projected costs, plans, prospects and objectives are forward-looking statements. Words such as "expect," "anticipate," "intend," "plan," "believe," "seek," "estimate," "think," "may," "could," "will," "should," "continue," "potential," "likely," "opportunity" and similar expressions or variations of such words are intended to identify forward-looking statements but are not the exclusive means of identifying forward-looking statements.

Among the factors that could cause actual results to differ materially from those indicated in the forward-looking statements are risks and uncertainties inherent in our business including, without limitation: our ability to adequately fund our projects as we will need additional funding to proceed with our objectives, the potential therapeutic effect of our products, the possibility of obtaining regulatory approval, our ability to find senior co-development partners with the capital and expertise needed to commercialize our products and to enter into arrangements with them on commercially reasonable terms, our ability to manufacture and sell any products, our ability to enter into arrangements with third party vendors, market acceptance of our products, our ability to earn a profit from sales or licenses of any drugs, our ability to discover new drugs in the future, changing market conditions, changes in laws and regulations affecting our industry, and issues related to our New Brunswick, New Jersey facility. In February 2013, we received a Complete Response Letter from the Food and Drug Administration, or FDA, for our Ampligen New Drug Application, or NDA, for the treatment of CFS. The FDA communicated that we should conduct at least one additional clinical trial, complete various nonclinical studies and perform a number of data analysis. Accordingly, the remaining steps to potentially gain FDA approval of the Ampligen NDA, the final results of these and other ongoing activities could vary materially from our expectations and could adversely affect the chances for approval of the Ampligen NDA. These activities and the ultimate outcomes are subject to a variety of risks and uncertainties, including but not limited to risks that (i) the FDA may ask for additional data, information or studies to be completed or provided; and (ii) the FDA may require additional work related to the commercial manufacturing process to be completed or may, in the course of the inspection of manufacturing facilities, identify issues to be resolved.

In August 2016, we received approval of our NDA from Administracion Nacional de Medicamentos, Alimentos y Tecnologia Medica, or ANMAT, for commercial sale of rintatolimod (U.S. tradename: Ampligen®) in the Argentine Republic for the treatment of severe CFS. The product will be marketed by GP Pharm, our commercial partner in Latin America. We believe, but cannot assure, that this approval provides a platform for potential sales in certain countries within the European Union under regulations that support cross-border pharmaceutical sales of licensed

drugs. In Europe, approval in a country with a stringent regulatory process in place, such as Argentina, should add further validation for the product as the Early Access Program, or EAP, as discussed below and underway in Europe in pancreatic cancer. ANMAT approval is only an initial, but important, step in the overall successful commercialization of our product. There are a number of actions that must occur before we could be able to commence commercial sales in Argentina. Commercialization in Argentina will require, among other things, an appropriate reimbursement level, appropriate marketing strategies, completion of manufacturing preparations for launch. Approval of rintatolimod for severe CFS in the Argentine Republic does not in any way suggest that the Ampligen NDA in the United States or any comparable application filed in the European Union or elsewhere will obtain commercial approval.

In May 2016, we entered into a five-year agreement with myTomorrows, a Netherlands based company, for the commencement and management of an EAP in Europe and Turkey related to CFS. Pursuant to the agreement, myTomorrows, as our exclusive service provider and distributor in this territory, is performing EAP activities. In January 2017, the EAP was extended to pancreatic cancer patients beginning in the Netherlands. In February 2018, we signed an amendment to extend the territory to cover Canada to treat pancreatic cancer patients, pending government approval. In March 2018, we signed an amendment to which myTomorrows will be our exclusive service provider for special access activities in Canada for the supply of Ampligen for the treatment of CFS. No assurance can be given that we can sufficiently supply product should we experience an unexpected demand for Ampligen in our clinical studies, the commercial launch in Argentina or pursuant to the EAPs. No assurance can be given that Ampligen will prove effective in the treatment of pancreatic cancer.

Currently, two Ampligen clinical trials are underway with a number of subjects enrolled at university cancer centers testing whether tumor microenvironments can be reprogrammed to increase the effectiveness of cancer immunotherapy, including checkpoint blockade. One is at Roswell Park Comprehensive Cancer Center and the other is at the University of Pittsburgh Medical Center. Two additional studies have been approved for enrollment and subjects are being screened for enrollment recruited at Roswell Park Comprehensive Cancer Center and the University of Pittsburgh Medical Center using Ampligen in conjunction with pembrolizumab. No assurance can be given as to the results of these underway trials. Four additional cancer trials in collaboration with University Medical/Cancer Research Centers using Ampligen plus checkpoint blockade are in various pre-enrollment stages. No assurance can be given as to whether some or all of the planned additional oncology clinical trials will occur and they are subject to many factors including lack of regulatory approval(s), lack of study drug, or a change in priorities at the sponsoring Universities or Cancer Centers. Even if these additional clinical trials are initiated, we cannot assure that these clinical studies or the two studies underway will be successful or yield any useful data.

Our overall objectives include plans to continue seeking approval for commercialization of Ampligen in the United States and abroad as well as seeking to broaden commercial therapeutic indications for Alferon N Injection presently approved in the United States and Argentina. We continue to pursue senior co-development partners with the capital and expertise needed to commercialize our products and to enter into arrangements with them on commercially reasonable terms. Our ability to commercialize our products, widen commercial therapeutic indications of Alferon N Injection and/or capitalize on our collaborations with research laboratories to examine our products are subject to a number of significant risks and uncertainties including, but not limited to our ability to enter into more definitive agreements with some of the research laboratories and others that we are collaborating with, to fund and conduct additional testing and studies, whether or not such testing is successful or requires additional testing and meets the requirements of the FDA and comparable foreign regulatory agencies. We do not know when, if ever, our products will be generally available for commercial sale for any indication.

We outsource certain components of our manufacturing, quality control, marketing and distribution while maintaining control over the entire process through our quality assurance and regulatory groups. We cannot provide any guarantee that the facility or our contract manufacturer will necessarily pass an FDA pre-approval inspection for Alferon manufacture.

The production of new Alferon Active Pharmaceutical Ingredient, or API, inventory will begin once the validation phase is complete. While the facility has already been approved by the FDA under the Biological License Application, or BLA, for Alferon, this status will need to be reaffirmed by a successful Pre-Approval Inspection by the FDA prior to commercial sale of newly produced inventory product. If and when the Company obtains a reaffirmation of FDA BLA status and has begun production of new Alferon API, it will need FDA approval as to the quality and stability of the final product before commercial sales can resume. We will need additional funds to finance the revalidation process in our facility to initiate commercial manufacturing, thereby readying ourselves for an FDA Pre-Approval Inspection. If we are unable to gain the necessary FDA approvals related to the manufacturing process and/or final product of new Alferon inventory, our operations most likely will be materially and/or adversely affected. In light of these contingencies, there can be no assurances that the approved Alferon N Injection product will be returned to production on a timely basis, if at all, or that if and when it is again made commercially available, it will return to prior sales levels. In addition, we are currently readying the New Brunswick facility to start manufacturing polymers

used for the production of Ampligen to satisfy our future needs. While we anticipate that we will be able to commence manufacturing polymers at the New Brunswick facility, we may need additional funding to continue manufacturing. There cannot be any guarantee that we will obtain adequate funds to sustain manufacturing at the New Brunswick facility or that the facility will be able to manufacture sufficient lots for the commercial launch of Ampligen.

We believe, and are investigating, Ampligen's potential role in enhancing the activity of influenza vaccines. While certain studies involving rodents, non-human primates (monkeys) and healthy human subjects indicate that Ampligen may enhance the activity of influenza vaccines by conferring increased cross-reactivity or cross-protection, further studies will be required and no assurance can be given that Ampligen will assist in the development of a universal vaccine for influenza or other viruses.

Because forward-looking statements are inherently subject to risks and uncertainties, some of which cannot be predicted or quantified and some of which are beyond our control, you should not rely on these forward-looking statements as predictions of future events. The events and circumstances reflected in our forward-looking statements may not be achieved or occur and actual results could differ materially from those projected in the forward-looking statements. Moreover, we operate in an evolving environment. New risk factors and uncertainties may emerge from time to time, and it is not possible for management to predict all risk factors and uncertainties. Except as required by applicable law, we do not plan to publicly update or revise any forward-looking statements contained herein, whether as a result of any new information, future events, changed circumstances or otherwise.

This Report also refer to estimates and other statistical data made by independent parties and by us relating to market size and growth and other data about our industry. This data involves a number of assumptions and limitations, and you are cautioned not to give undue weight to such estimates. In addition, projections, assumptions and estimates of our future performance and the future performance of the markets in which we operate are necessarily subject to a high degree of uncertainty and risk.

ITEM 1B. Unresolved Staff Comments.

None.

ITEM 2. Properties.

Our principal executive office is located at 2117 SW Highway 484, Ocala FL 34473 and our finance and human resource office is located at 600 Main Street, Suite 2, Riverton, NJ 08077. We currently lease our principal executive office for \$3,329 per month and our accounting and human resource office for about \$1,100 per month.

In May 2017, we entered into a mortgage and note payable agreement with a bridge funding company to obtain a two-year funding line of up to \$4,000,000 secured by our assets and property located at 783 Jersey Ave., New Brunswick, New Jersey. As of March 16, 2018, this note was paid off in full. See Note 14 - Note Payable below for a more complete description of the terms of the note payable.

On March 16, 2018, we sold our property located at 783 Jersey Ave., New Brunswick, NJ. This property houses our development and production facilities. The purchase price was \$4,080,000 and purchaser received 3,225,806 warrants to purchase common stock. We believe that the sale and lease-back of this building will not have a material impact on our business operations. Simultaneously with the closing of the sale, the purchaser leased the facility back to us. The lease runs for 10 years, with two five year extensions. The initial annual base rent is \$408,000 and will continue for the first and second year. In the third and fourth it will escalate at the rate of 2.5% per year. For all subsequent years it will escalate at the rate of 3% per year. We also will be responsible for additional rent consisting of taxes and certain insurance expenses of the purchaser. The lease contains a repurchase option pursuant to which we can repurchase the facility within the initial 10 year lease period. The purchase price would be based on a multiple of the sale price of \$4,080,000. The multiple would be 1.05 plus .0025N where N represents the number of months between lease commencement and closing of repurchase.

In February 2018, the Company sold the unencumbered, unutilized, and wholly owned property located at 5 Jules Lane, New Brunswick, New Jersey to Acellories, NJ LLC, a New Jersey limited liability company, pursuant to a sale agreement dated September, 11, 2017. The sale price was \$1,050,000.

ITEM 3. Legal Proceedings.

Hemispherx commenced an action against BioLife in December of 2017 for Breach of Contract. The amount of damages we are seeking in this matter are yet to be determined. Damages are not covered by insurance. BioLife, the

defendant, has filed its Answer, Affirmative Defenses and a Counterclaim in the amount of \$96,676.39 representing the Invoices withheld after BioLife indicated that they were not intending to fulfill the balance of the contract. Hemispherx has denied the allegations of the counterclaim. We recently attempted mediation and were, to date, unable to resolve the matter. A discovery schedule has been issued by the Court and we are now in the initial stages of discovery. Although it cannot be determined, we believe there is little chance for an unfavorable outcome in this matter.

Hemispherx recently engaged in mediation with its insurance carrier Travelers over a Business Interruption Loss due to a flood in our New Brunswick manufacturing facility in January of 2016. The carrier to date has covered us for repairs to the facility but there still remains the unresolved issue of the amount of the claim for our Business Interruption Loss for which we have coverage under the policy. The Business Interruption Loss damages sustained after calculations by an independent Forensic Accountant (which was paid for by Travelers under the policy) exceed \$4.5 million, which is the limit allowed under the policy. We have to date been unable to settle through mediation. Therefore, Hemispherx filed and served a complaint in Philadelphia Court of Common Pleas against Travelers at the end of March 2019 seeking the policy limits and additional damages.

ITEM 4. Mine Safety Disclosures.

Not Applicable.

PART II

ITEM 5. Market for Registrant’s Common Equity, Related Stockholder Matters and Issuer Purchases of Equity Securities.

Our common stock is listed and traded on the NYSE American under the symbol HEB.

As of March 21, 2019, there were approximately 166 holders of record of our Common Stock. This number was determined from records maintained by our transfer agent and does not include beneficial owners of our securities whose securities are held in the names of various dealers and/or clearing agencies.

We have not paid any cash dividends on our Common Stock in recent years. It is management’s intention not to declare or pay dividends on our Common Stock, but to retain earnings, if any, for the operation and expansion of our business.

ITEM 6. Selected Financial Data.

Not Applicable.

ITEM 7. Management’s Discussion and Analysis of Financial Condition and Results of Operations.

The following discussion and analysis is related to our financial condition and results of operations for the two years ended December 31, 2018. This information should be read in conjunction with our consolidated financial statements and related notes thereto beginning on F-1 of this Form 10-K. Please also see “Special Note Regarding Forward Looking Statements” in ITEM 1A. Risk Factors.

Fair Value

We have issued warrants (the “Warrants”) in August 2016, February 2017, June 2017, August 2017, and August 2018 that are single compound derivatives containing both an embedded right to obtain stock upon exercise (a “Call”) and a series of embedded rights to settle the Warrants for cash upon the occurrence of certain events (each, a “Put”).

Generally, the Put provisions allow the Warrant Holders liquidity protection; the right to receive cash in certain situations where the Holders would not have a means of readily selling the shares issuable upon exercise of the Warrants (e.g., where there would no longer be a significant public market for our common stock). However, because the contractual formula used to determine the cash settlement value of the embedded Put requires use of certain assumptions, the cash settlement value of the embedded Put can differ from the fair value of the unexercised embedded Call option at the time the embedded Put option is exercised.

We recompute the fair value of the Warrants at the end of each quarterly reporting period. Such value computation includes subjective input assumptions that are consistently applied each period. If we were to alter our assumptions or the numbers input based on such assumptions, the resulting fair value could be materially different.

On September 28, 2018, We entered into a \$3,170,000 10% Secured Convertible Promissory Note (the “Note”) with Iliad Research and Trading, L.P. (the “Holder”), which was issued to the Holder in conjunction with 500,000 shares of Common Stock (the “Origination Shares”). We collected \$3,000,000 in cash from the Holder during September 2018 and the remainder \$170,000 was retained by the Holder for the Holder’s legal fees of \$20,000 for the issuance of the Note and the Original Issue Discount of \$150,000. We incurred \$210,000 in third-party fees directly attributed to the issuance of the Note.

We determined the Note should be recorded at fair value with subsequent changes in fair value recorded in earnings. This conclusion is based on the redemption conversion feature, which allows the Holder to trigger the redemption of the Note for cash or conversion of the Note for common shares prior to its maturity date at a price of the lesser of \$0.30 per share or the Market Price as defined within the Note. The choice of cash redemption or conversion of the Note for common shares is at our option. This feature may require that we issue a variable number of common shares to settle the Note which was determined to have a predominantly fixed monetary value at inception. In connection with the Note, we recorded a loss in the Company’s Consolidated Statements of Comprehensive Income (Loss) equal to \$582,000 for the year ended December 31, 2018.

On March 13, 2019, the Convertible Note was extended from September 28, 2019 to September 28, 2020. In addition, the conversion and redemption rates were revised to a rate to be mutually agreed to us and the Lender. See note 20-Subsequent Events.

RESULTS OF OPERATIONS

Year ended December 31, 2018 versus year ended December 31, 2017

Our net loss was approximately \$9,813,000 and \$8,259,000 for the years ended December 31, 2018 and 2017, respectively, representing an increase in loss of approximately \$1,554,000 or 19% when compared to the same period

in 2017. This increase in loss for the year ended December 31, 2018 was primarily due to the following:

- 1) a decrease in revenues of \$70,000 or 16%;
- 2) an increase in interest and finance costs of \$363,000;
- 3) the quarterly revaluation of certain redeemable warrants resulted in a non-cash gain of \$1,165,000 in the year ended December 31, 2018 compared to a gain of \$2,417,000 in the year ended December 31, 2017, a decrease of \$1,252,000;
- 4) the fair value adjustment for the convertible note resulted in a loss of \$582,000 in the year ended December 31, 2018, which did not occur in 2017;
- 5) an increase in research and development expense of \$680,000 or 17%; offset by
- 6) a decrease in production costs of \$299,000;
- 7) a decrease in general and administrative expenses of \$371,000
- 8) a gain resulting from a settlement of litigation with a vendor of \$474,000;
- 9) a gain from the sale of the underutilized building in New Brunswick of \$223,000; and
- 10) a decrease in legal fees due to a favorable settlement of legal fees of \$342,000.

Net loss per share was \$(0.22) and \$(0.29) for the years ended December 31, 2018 and 2017, respectively. The weighted average number of shares of our common stock outstanding as of December 31, 2018 was 44,189,217 as compared to 28,676,076 as of December 31, 2017.

Revenues

Revenues from our Ampligen® Cost Recovery Program were \$367,000 and \$437,000 for the years ended December 31, 2018 and 2017, respectively. The decrease in revenues of \$70,000, a decrease of 16%, between periods was primarily due to the unavailability of Ampligen for our EAP through our agreement with MyTomorrows designed to enable access of Ampligen to pancreatic cancer patients in the Netherlands.

For the years ended December 31, 2018 and 2017, we had no Alferon N Injection® Finished Good product to commercially sell and all revenue was generated from the EAP and our FDA approved open-label treatment protocol, (“AMP 511”), that allows patient access to Ampligen® for treatment in an open-label safety study.

Production Costs

Production costs were approximately \$884,000 and \$1,183,000, respectively, for the years ended December 31, 2018 and 2017, representing a decrease of \$299,000 in production costs in the current period. These costs primarily represent stability testing and pre-production expenses related to Alferon®. The reduction in costs was due to a write-off of \$210,000 for expired Alferon fill and finish costs in the prior year and a decrease in other Alferon production costs of \$89,000.

Research and Development Costs

Overall Research and Development (“R&D”) costs for the year ended December 31, 2018 were approximately \$4,778,000 as compared to \$4,098,000 for the same period a year ago, reflecting an increase of approximately \$680,000 or 17%. The primary reason for the increase in research and development costs was due to an increase of \$653,000 for the completion of the manufacture of 8,484 vials and 7,907 vials in June and September 2018, respectively, an increase of \$776,000 for the production of polymers offset by a reduction of U.S. clinical costs of \$677,000 as a result of reduction in amounts due to clinical investigators resulting from renegotiated terms with the investigators.

General and Administrative Expenses

General and Administrative (“G&A”) expenses for the years ended December 31, 2018 and 2017, were approximately \$6,201,000 and \$6,572,000, respectively, reflecting a decrease of approximately \$371,000 or 6%. The decrease in G&A expenses during the current period was mainly due to a favorable settlement of legal fees of \$342,000.

Interest Expense and Finance Costs

Interest and finance costs for the year ended December 31, 2018 was \$502,000 compared to \$139,000 in the prior year, an increase of \$363,000 or 261%. The increase is mainly due to a note/mortgage payable incurred in May 2017 which was paid off in March 2018 with the resulting write off of the balance of the unamortized mortgage settlement costs in addition to the interest expense on the mortgage; plus the interest settlement costs on the Finance Obligation from the sale leaseback of the main New Brunswick building and finance costs and interest related to the convertible note from September 2018. None of these were incurred in the years ended December 31, 2017.

Interest and Other Income

Interest and other income for the years ended December 31, 2018 and 2017 was approximately \$46,000 and \$88,000, respectively, representing a decrease of approximately \$42,000 or 48%. The primary cause for the decrease in investment income during the current period was primarily due to lower balances available to invest in the current period as compared to the prior period.

Redeemable Warrants

The quarterly revaluation of certain redeemable warrants resulted in a non-cash adjustment to the redeemable warrants liability for the year ended December 31, 2018 amounted to a gain of approximately \$1,165,000, compared to a gain of \$2,417,000 for the year ended December 31, 2017, which represents a decrease of \$1,252,000 or 52% (see “Financial Statements: Note 18: Fair Value” for the various factors considered in the valuation of redeemable warrants).

Sale of New Jersey Tax Net Operating Loss

In December 2017, the Company effectively sold \$8,000,000 New Jersey state net operating loss for approximately \$622,000 and sold research credits for \$169,000. In December 2018, the Company effectively sold \$10,000,000 New Jersey state net operating loss for approximately \$859,000. The money was received in January 2019.

Convertible Note Payable

The quarterly valuation of the convertible note payable resulted in a non-cash loss of \$582,000 in 2018 which did not occur in 2017.

Other Transactions

During the year ended December 31, 2018 there was a gain of \$474,000 resulting from the settlement of litigation with Nitto Avecia Pharma Services, Inc. ("NAPS").

There was also a gain of \$223,000 resulting from the sale of the second building in New Brunswick, New Jersey.

Liquidity and Capital Resources

In 2018, we sold an under-utilized warehouse at 5 Jules Lane for \$1,050,000 and we sold our manufacturing facility for \$4,080,000 while simultaneously entering into a favorable long term lease with an option to repurchase the facility. In 2018, we also realized \$1,260,000 through the exercising of outstanding warrants.

In March 2019, we completed a rights offering to our stockholders and certain option and warrant holders, pursuant to which we issued Preferred stock convertible into an aggregate of 26,560,000 shares of common stock and warrants exercisable for up to an additional 26,560,000 shares of common stock. We netted approximately \$4.69 million from the sale of securities in the rights offering.

In December 2017, we reactivated the Equity Distribution Agreement (“EDA”) and, through strategic management, have raised \$1,039,000. On March 24, 2018, we sold common stock netting us an additional \$475,000. On April 24, 2018 we sold common stock netting us an additional \$2,344,000.

Cash used in operating activities for the year ended December 31, 2018 was approximately \$10,640,000 compared to approximately \$7,941,000 for the same period in 2017, an increase of \$2,699,000 or 34%. The primary reasons for this increase in cash used in operations in 2018 was the receipt of \$791,000 in funds in 2017 from the sale of our New Jersey state net operating loss carryforwards. In 2018, we did not receive the funds from the sale of our New Jersey net operating loss carryforwards until January 2019. In 2018, we also expended about \$1,500,000 toward the manufacturing of additional polymers for Ampligen for commercial launch in Argentina in May or June on 2019.

Cash provided by investing activities for the year ended December 31, 2018 was approximately \$92,000 compared to cash provided by investing activities of approximately \$2,730,000 for the same period in 2017, representing a decrease of \$2,638,000. The primary reason for the decrease was the sale of marketable securities of approximately \$831,000 during the current period compared to \$2,799,000 the year ended December 31, 2017.

Cash provided by financing activities for the year ended December 31, 2018 was approximately \$9,435,000 compared to approximately \$4,215,000 for the same period in 2017, an increase of \$5,220,000. The primary reasons for this increase was that we received net proceeds of \$3,377,000 from the sale leaseback of our manufacturing facility and \$5,070,000 in 2018 from the sale of shares compared to \$2,417,000 from the sale of shares in 2017.

As of December 31, 2018, we had approximately \$1,825,000 in cash, cash equivalents and marketable securities, inclusive of approximately \$1,526,000 in Marketable Securities, representing a decrease of approximately \$282,000 from December 31, 2017.

If we are unable to commercialize and sell Ampligen and/or recommence material sales of Alferon N Injection, our operations, financial position and liquidity may be adversely impacted, and additional financing may be required. In this regard, due to the high cost estimates to bring the facility back online, we will need additional funds to finance the revalidation process in our facility and to initiate commercial manufacturing, thereby readying ourselves for an FDA Pre-Approval Inspection and to commercialize our products. However, there is no assurance that such financing will be available.

In an effort to conserve cash, effective with the semi-monthly period ended April 30, 2017, all of the members of the Company's Board of Directors agreed to accept 100% of their directors' fees in the form of options to purchase Company Common Stock. This program was terminated as of August 31, 2017. As of September 1, 2017, the directors agreed to defer 100% of their fees until cash is available. On February 13, 2018, 226,023 options were issued to each of the two independent directors with an exercise price of \$0.37 for a period of 10 years with a vesting period of 3 years. In addition, commencing with the semi-monthly period ended June 15, 2017, certain officers of the Company, and certain other employees of the Company, agreed to accept 20% of their salary in options to purchase Company Common Stock. This program was also terminated as of August 31, 2017. In this regard, options to purchase 214,866 shares of Company common stock were issued with exercise prices ranging from \$0.36 to \$0.67, a holding period of 10 years and vesting over three years.

As part of the cash conservation program adopted on August 28, 2017, starting with the month of September 2017, the directors agreed to defer 100% of their fees until cash is available. In consideration of this deferral, 226,023 options were issued to each of the two independent directors in February 2018 with an exercise price of \$0.37; 152,053 options were issued to each of the two independent directors in May 2018 with an exercise price of \$0.30, and 98,098 options were issued in July 2018 with an exercise price of \$0.31. All of the foregoing options and the options discussed below are exercisable for a period of 10 years with a vesting period of three years. This program was suspended as of July 15, 2018 and all remaining deferred fees were paid in July 2018. This Program was reactivated as of August 16, 2018 with the understanding that options would not be issued on the deferred amounts until the 2018 Equity Incentive Plan was approved by the stockholders. The 2018 Equity Incentive Plan was approved by the stockholders and the securities issuable thereunder were registered with the SEC and, on October 17, 2018, 172,786 options were issued to each of the two independent directors with an exercise price on \$0.22 for a period of ten years with a vesting period of one year. On January 28, 2019, 207,343 options were issued to each of the two independent directors with an exercise price of \$0.22 for a period of ten years with a vesting period of one year. Also on January 28, 2019, 50,000 options were issued to each of the two independent directors with an exercise price of \$0.22 for a period of ten years with a vesting period of one year for chairing the committees in 2018.

Also as part of the cash conservation program adopted on August 28, 2017, starting with the month of September 2017, certain officers agreed to defer 40% of their salaries until cash is available. In consideration of this deferral, 884,459 options were issued to these officers in February 2018 with an exercise price of \$0.37; 599,168 options were issued to these officers in May 2018 with an exercise price of \$0.30, and 389,249 options were issued to these officers in July 2018 with an exercise price of \$0.31. This program was suspended as of July 15, 2018 and all remaining deferred salaries were paid on July 2018. This Program was reactivated as of August 16, 2018 for 50% of their salaries with the understanding that options would not be issued on the deferred amounts until the 2018 Equity Incentive Plan was approved by the shareholders and the plan registered with the SEC. The 2018 Equity Incentive Plan has been approved by the shareholders and registered with the SEC and on October 17, 2018, 808,712 options were issued to these officers with an exercise price on \$0.22 for a period of ten years with a vesting period of one year. On January 28, 2019, 1,213,069 options were issued to these officers with an exercise price of \$0.22 for a period of ten years with a vesting period of one year.

Also as part of the cash conservation program adopted on August 28, 2017, all employees agreed to be paid 50% of their salaries in the form of unrestricted common stock of the Company. Starting with the month of September 2017,

the salaries of all the employees of the Company were paid 50% in the form of unrestricted common stock of the Company. The total number of shares issued as of June 30, 2018 to the employees under this program was 2,116,881 shares at stock prices ranging from \$0.31 to \$0.55 per share. This program was suspended by the Board of Directors on June 30, 2018.

On March 24, 2018, the Company sold 1,250,000 shares of common stock under its S-3 shelf registration. The Company realized net proceeds of \$475,000 from this stock offering and paid \$25,000 in placement agent fees.

On April 20, 2018, the Company entered into Securities Purchase Agreements (the "Purchase Agreements") with certain investors (the "Investors") for the sale by the Company of an aggregate of 6,600,000 shares (the "Common Shares") of the Company's common stock, par value \$0.001 per share (the "Common Stock"), at a purchase price of \$0.39 per share. Concurrently with the sale of the Common Shares, pursuant to the Purchase Agreements the Company also sold 6,600,000 warrants, 50% of which are Class A Warrants and 50% of which are Class B Warrants (collectively, the "Warrants"). The Company received gross proceeds from the sale of the Warrants solely to the extent such Warrants are exercised for cash. Both classes of Warrants will not be exercisable until six months after issuance and will have an exercise price of \$0.39 per share, subject to adjustments as provided under the terms of the Warrants. The Class A Warrants and Class B Warrants will expire, respectively, two and five years after the date on which they are first exercisable. The closing of the sales of these securities under the Purchase Agreements took place on April 24, 2018. The Company received net proceeds from the transactions of \$2,343,820 after deducting certain fees due to the placement agent and the Company's transaction expenses.

On November 27, 2017, we reactivated the EDA. During the year ended December 31, 2018, we sold an aggregate of 2,176,392 shares under the EDA for proceeds of \$827,000 net of \$25,000 in commissions. Pursuant to a prospectus supplement dated February 7, 2018, we were able to sell up to 6,549,157 of our common stock (inclusive of shares already sold under the prospectus supplement) under the EDA. From January 1, 2019 through March 25, 2019 we sold an aggregate of 115,606 shares under the EDA for proceeds of \$26,000 net of \$1,000 in commissions. The actual number of shares that we can sell and the proceeds to be received therefrom are dependent upon the market price of our common stock.

In February 2017, we entered into Securities Purchase Agreements (each, a “Purchase Agreement”) with certain investors for the sale by us of 1,818,185 shares of our common stock at a purchase price of \$0.55 per share. Concurrently with the sale of the common stock, pursuant to the Purchase Agreement, we also sold warrants to purchase 1,363,639 shares of common stock for aggregate net proceeds of approximately \$875,000. We also issued placement agent warrants for the purchase of an aggregate of 90,910 shares of our common stock.

In May 2017, we entered into a mortgage and note payable agreement with a bridge funding company to obtain a two-year funding line of up to \$4,000,000 secured by our assets and property located at 783 Jersey Ave., New Brunswick, New Jersey. We paid interest on this note at a fixed rate of 12% per annum. We were permitted to prepay the line without penalty commencing after six months. The balance on this note was \$1,835,000 as of December 31, 2017; however, it was paid off on March 16, 2018 in conjunction with the sale of 783 Jersey Ave.

On March 16, 2018, we sold our property located at 783 Jersey Ave, New Brunswick, NJ for \$4,080,000 and the purchasers received 3,225,806 warrants to purchase common stock. Simultaneously therewith, we leased the facility back. See PART I, Item 2 - “Properties.”

In February 2018, the Company sold the unencumbered, unutilized, and wholly owned property located at 5 Jules Lane, New Brunswick, New Jersey to Acellories, NJ LLC, a New Jersey limited liability company, pursuant to a sale agreement dated September, 11, 2017. The sale price was \$1,050,000.

On June 1, 2017, pursuant to an offer (the “Exchange Transaction”) to the holders of warrants issued to investors in September 2016 (the “2016 Warrants”), the exercise price of the 2016 warrants was changed to \$0.50. As a result the warrant holders exercised 2016 Warrants and purchased 2,370,000 shares of Company common stock. The Company realized net proceeds of \$1,055,000 from this exercise. As part of the Exchange Transaction, the Company issued 2,370,000 series A warrants with an exercise price of \$0.60 per share, an initial exercise date of December 1, 2017 and expiring March 6, 2022, and 7,584,000 series B warrants with an exercise price of \$0.60, an initial exercise date of December 1, 2017 per share and expiring March 1, 2018. These warrants were exercised in January and February 2018 for proceeds of \$1,260,000. In addition, in July 2017, the warrant holders exercised the remaining 130,000 2016 Warrants and purchased 130,000 shares of common stock. The Company realized net proceeds of \$65,000 from this exercise. In conjunction with the foregoing the Company issued 130,000 series A warrants with an exercise price of

\$0.60 per share and an initial exercise date of January 10, 2018 and expiring March 6, 2022, and 416,000 series B warrants with an exercise price of \$0.60 and an initial exercise date of January 10, 2018.

On August 23, 2017, the Holders of the series A warrants and series B warrants exchanged all of their series A warrants and series B warrants for new warrants (respectively, the “Series A Exchange Warrants” and the “Series B Exchange Warrants” and, collectively, the “Exchange Warrants”) identical to the series A warrants and series B warrants except as follows: the exercise price of both Exchange Warrants is \$0.45 per share, subject to adjustment therein, and the number of Series B Exchange Warrants issued was proportionately reduced so that all Exchange Warrants in the Exchange Transaction do not exceed 19.9% of the number of the Company’s issued and outstanding shares of Common Stock as of May 31, 2017, the date of the Exchange Transaction offer letters. The issuance of the Exchange Warrants by the Company and the shares of Common Stock issuable upon exercise of the Exchange Warrants is exempt from registration pursuant to Sections 3(a)(9) and 4(a)(2) of the Securities Act of 1933, as amended (the “Securities Act”). There were 2,800,000 warrants with an expiration date of March 1, 2018 and an exercise price on \$0.45. These warrants were exercised in January and February 2018. The Company realized proceeds of \$1,260,000 from these exercises.

There can be no assurances that, if needed, we will be able to raise adequate funds from these or other sources or enter into licensing, partnering or other arrangements to advance our business goals. Our inability to raise such funds or enter into such arrangements, if needed, could have a material adverse effect on our ability to develop our products. Also, we have the ability to curtail discretionary spending, including some research and development activities, if required to conserve cash. Because of our long-term capital requirements, we may seek to access the public equity market whenever conditions are favorable, even if we do not have an immediate need for additional capital at that time. We are unable to estimate the amount, timing or nature of future sales of outstanding common stock or instruments convertible into or exercisable for our common stock. Any additional funding may result in significant dilution and could involve the issuance of securities with rights, which are senior to those of existing stockholders. We may also need additional funding earlier than anticipated, and our cash requirements, in general, may vary materially from those now planned, for reasons including, but not limited to, changes in our research and development programs, clinical trials, acquisitions of intellectual property or assets, enhancements to the manufacturing process, competitive and technological advances, the regulatory processes including the commercializing of Ampligen® products or new utilization of Alferon® products. See Part I, Item 1A - “Risk Factors; *We will require additional financing which may not be available*”.

The proceeds from our financing have been used to fund infrastructure growth including manufacturing, regulatory compliance and market development along with our efforts regarding the Ampligen® NDA and preparedness for the FDA pre-approval inspections of the New Brunswick manufacturing facility. There can be no assurances that, if needed, we will raise adequate funds from these or other sources, which may have a material adverse effect on our ability to develop our products. Also, we have the ability to curtail discretionary spending, including some research and development activities, if required to conserve cash.

Certain Relationships and Related Transactions

Refer to PART III, ITEM 13 - “Certain Relationships and Related Transactions, and Director Independence.”

New Accounting Pronouncements

Refer to “Note 2(i) – Recent Accounting Standards and Pronouncements” under Notes to Consolidated Financial Statements.

Disclosure about Off-Balance Sheet Arrangements

None.

Critical Accounting Policies

Financial Reporting Release No. 60 requires all companies to include a discussion of critical accounting policies or methods used in the preparation of financial statements. Our significant accounting policies are described in the Notes to Consolidated Financial Statements. The significant accounting policies that we believe are most critical to aid in fully understanding our reported financial results are the following:

Revenue

The Company has elected to apply the Full Retrospective Application to implement the new revenue recognition standard ASC 606. The Company, based on the nature of its Ampligen® sales under its cost recovery programs, determined that there were no material differences between the new accounting standard and legacy GAAP and that difficulties will not arise for any “open” contract issues with its customers during the transition period. The Company also determined that the adoption of this standard will have little or no impact to the Company’s opening balance of retained earnings.

Revenue from the sale of Ampligen® under cost recovery clinical treatment protocols approved by the FDA is recognized when the treatment is provided to the patient.

Revenues from the sale of product are recognized when the product is delivered, as title is then transferred to the customer. We have no other obligation associated with our products once shipment has been accepted by the customer

Inventories

We use the lower of first-in, first-out (“FIFO”) cost and net realizable value method of accounting for inventory.

Patents and Trademarks

Patents and trademarks are stated at cost (primarily legal fees) and are amortized using the straight-line method over the estimated useful life of 17 years. We review our patents and trademark rights periodically to determine whether they have continuing value. Such review includes an analysis of the patent and trademark’s ultimate revenue and profitability potential. In addition, Management’s review addresses whether each patent continues to fit into our strategic business plans.

Long-Lived Assets

We assess long-lived assets for impairment when events or changes in circumstances indicate that the carrying value of the assets or the asset grouping may not be recoverable. Factors that we consider in deciding when to perform an impairment review include significant under-performance of a business or product line in relation to expectations, significant negative industry or economic trends, and significant changes or planned changes in its use of the assets. We measure the recoverability of assets that it will continue to use in its operations by comparing the carrying value of the asset grouping to our estimate of the related total future undiscounted net cash flows. If an asset grouping’s carrying value is not recoverable through the related undiscounted cash flows, the asset grouping is considered to be impaired.

We measure the impairment by comparing the difference between the asset grouping's carrying value and its fair value. Long-lived assets are considered a non-financial asset and are recorded at fair value only if an impairment charge is recognized. Impairments are determined for groups of assets related to the lowest level of identifiable independent cash flows. We make subjective judgments in determining the independent cash flows that can be related to specific asset groupings. In addition, as we review our manufacturing process and other manufacturing planning decisions, we must make subjective judgments regarding the remaining useful lives of assets. When we determine that the useful lives of assets are shorter than originally estimated, we accelerate the rate of depreciation over the assets' new, shorter useful lives.

Stock-Based Compensation

Under FASB ASC 718-Compensation-Stock Compensation ("ASC 718") share-based compensation cost is measured at the grant date, based on the estimated fair value of the award, and is recognized as expense over the requisite service period. Under this method, compensation cost is recognized for all share-based payments granted, modified or settled after the date of adoption, as well as for any unvested awards that were granted prior to the date of adoption.

The fair value of each option award is estimated on the date of grant using a Black-Scholes-Merton pricing option valuation model. Expected volatility is based on the historical volatility of the price of our common stock. The risk-free interest rate is based on U.S. Treasury issues with a term equal to the expected life of the option. We use historical data to estimate expected dividend yield, expected life, which represents the period of time the options are expected to be outstanding until they are exercised, and forfeiture rates.

Redeemable Warrants

We utilize the guidance contained in ASC 480 in the determination of whether to record warrants and options as Equity and/or Liability. If the guidance of ASC 480 is deemed inconclusive, we continue our analysis utilizing ASC 815

Our method of recording the related value attempts to be consistent with the standards as defined by the Financial Accounting Standards Board utilizing the concept of "Fair Value" from ASC 820-10-55-1 that states that any fair value measurement requires that the reporting entity, to determine the valuation technique(s) appropriate for the measurement, consider the availability of data with which to develop inputs that represent the assumptions that market participants would use in pricing the asset or liability and the level in the fair value hierarchy within which the inputs fall.

We recomputed the value of the redeemable warrants at the end of each quarterly period. We use the Monte Carlo Simulation approach which includes subjective input assumptions that are consistently applied each quarter. If we were to alter our assumptions or the numbers input based on such assumptions, the resulting fair value could be materially different. As discussed in greater detail in “Fair Value” at the beginning of this ITEM 7, the significant assumptions using this model are: (i) Risk-Free Interest Rate; (ii) Expected Holding Period; (iii) Expected Volatility; (iv) Expected Dividend Yield; (v) Expected Probability of a Fundamental Transaction; (vi) Expected Timing of Announcement of a Fundamental Transaction; (vii) Expected 100 Day Volatility at Announcement of a Fundamental Transaction; (viii) Expected Risk-Free Interest Rate at Announcement of a Fundamental Transaction; and (ix) Expected Time Between Announcement and Consummation of a Fundamental Transaction.

Convertible Note Payable

In September 2018, we entered into a \$3,170,000 10% Secured Convertible Promissory Note with Iliad Research and Trading, L.P. (the “Note”). We determined the Note should be recorded at fair value with subsequent changes in fair value recorded in earnings. This conclusion is based on the redemption conversion feature, which allows the Holder to trigger the redemption of the Note for cash or conversion of the Note for common shares prior to its maturity date at a price of the lesser of \$0.30 per share or the Market Price as defined within the Note. The choice of cash redemption or conversion of the Note for common shares is at our option. This feature may require that we issue a variable number of common shares to settle the Note which was determined to have a predominantly fixed monetary value at inception. In connection with the Note, we recorded a loss in our Consolidated Statements of Comprehensive Income (Loss) equal to \$582,000 for the year ended December 31, 2018. For more detail about the Note, please see the disclosure in “Fair Value” above.

Concentration of Credit Risk

Our policy is to limit the amount of credit exposure to any one financial institution and place investments with financial institutions evaluated as being credit worthy, or in short-term money markets, which are exposed to minimal interest rate and credit risks. We have had bank deposits and overnight repurchase agreements that exceed federally insured limits.

Concentration of credit risk, with respect to receivables, is limited through our credit evaluation process. We do not require collateral on our receivables. Our receivables historically consisted principally of amounts due from wholesale drug companies.

ITEM 7A. Quantitative and Qualitative Disclosures About Market Risk.

Not Applicable.

ITEM 8. Financial Statements and Supplementary Data.

The consolidated balance sheets as of December 31, 2018 and 2017, and our consolidated statements of comprehensive loss, changes in stockholders' equity, and cash flows for each of the years in the two-year period ended December 31, 2018, together with the reports of Morrison, Brown, Argiz & Farra, LLC, our current independent registered public accountants, and RSM US LLP, our prior independent registered public accountants, are included at the end of this report. Reference is made to the "Index to Financial Statements and Financial Statement Schedule" on page F-1.

ITEM 9. Changes in and Disagreements with Accountants on Accounting and Financial Disclosures.

On April 2, 2018, we released RSM US LLP ("RSM") as our independent registered public accounting firm. The decision to rotate the independent registered public accounting firms was approved by the Audit Committee of our Board of Directors (the "Audit Committee"). The Audit Committee determined to transition to another accounting firm for best practices as RSM had been the Company's auditors for 12 years. In this regard, on April 4, 2018, the Audit Committee entered into an agreement with Morrison, Brown, Argiz & Farra, LLC ("MBAF") to serve as our independent registered public accounting firm. We did not consult with MBAF regarding any of the matters or events set forth in Item 304(a)(2)(ii) of Regulation S-K.

During the fiscal years ended December 31, 2017 and 2018, there were (i) no disagreements between us and RSM on any matter of accounting principles or practices, financial statement disclosure, or auditing scope or procedures, which disagreement, if not resolved to the satisfaction of RSM, would have caused RSM to make reference thereto in their reports on the financial statements for the year ended December 31, 2017, and (ii) no "reportable events" as that term is defined in Item 304(a)(1)(v) of Regulation S-K, except as follows. As noted in our quarterly reports on Form 10-Q for the second and third quarters of 2017, we, in carrying out an evaluation of the effectiveness of our disclosure controls and procedures, determined that, at the end of these quarters, there was a material weakness. The material weakness related to the completeness and accuracy of the recording of the exercise of certain redeemable warrants. The Audit Committee discussed this material weakness with RSM during the second and third quarters of 2017. We authorized RSM to respond fully to the inquiries of MBAF regarding the previously reported material weakness. We believe that we have corrected this issue and no such material weakness was found during the fourth quarter of 2017.

The reports of RSM for the fiscal years ended December 31, 2017 contained no adverse opinion or disclaimer of opinion and were not qualified or modified as to uncertainty, audit scope or accounting principles.

We provided RSM and MBAF with a copy of the Current Report on Form 8-K related to the change in accounting firms prior to its filing with the Commission and RSM furnish us with a letter addressed to the SEC stating that it agreed with the above statements.

ITEM 9A. Controls and Procedures.

Effectiveness of Control Procedures

As of December 31, 2018, the end of the period covered by this report, we carried out an evaluation under the supervision and with the participation of our Management, including our Chief Executive Officer and our Chief Financial Officer, of the effectiveness of the design and operation of our disclosure controls and procedures as defined in Rules 13a-15(e) and 15d-15(e) promulgated under the Exchange Act. Our disclosure controls and procedures are intended to ensure that the information we are required to disclose in the reports that we file or submit under the Securities Exchange Act is (i) recorded, processed, summarized and reported within the time periods specified in the Securities Exchange Commission's rules and forms and (ii) accumulated and communicated to our management, including the Chief Executive Officer and Chief Financial Officer, as the principal executive and financial officers, respectively, to allow final decisions regarding required disclosures. Based on that evaluation, our Chief Executive Officer and Chief Financial Officer concluded that the controls and procedures were effective as of December 31, 2018 to ensure that material information was accumulated and communicated to our management, including our Chief Executive Officer and Chief Financial Officer, as appropriate to allow timely decisions regarding required disclosure. Our management has concluded that the financial statements included in this Form 10-K present fairly, in all material respects our financial position, results of operations and cash flows for the periods presented in conformity with accounting principles generally accepted in the United States of America.

Changes in Internal Control over Financial Reporting

We made no changes in our internal control over financial reporting during the last fiscal quarter that have materially affected, or are reasonably likely to materially affect, our internal control over financial reporting (as defined in Rules 13a-15(f) and 15d-15(f) under the Exchange Act).

Management's Report on Internal Control over Financial Reporting

Our Management is responsible for establishing and maintaining adequate internal control over financial reporting as such term is defined in Rules 13a-15(f) or 15d-15(f), under the Exchange Act. Internal control over financial reporting is a process designed by, or under the supervision of, our principal executive and principal financial officers and affected by our Board of Directors, Management and other personnel, and 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 (i) pertain to the maintenance of records that, in reasonable detail, accurately and fairly reflect the transactions and dispositions of our assets; (ii) 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 (iii) 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 its 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 risk that controls may become inadequate because of changes in conditions, or that the degree of compliance with the policies or procedures may deteriorate.

Management has assessed the effectiveness of our internal control over financial reporting as of December 31, 2018. In making this assessment, Management used the criteria set forth in the framework in 2013 established by the Committee of Sponsoring Organizations of the Treadway Commission Internal Control—Integrated Framework, (COSO). Based on this assessment, Management has not identified any material weaknesses as of December 31, 2018. A material weakness is a control deficiency, or combination of control deficiencies, that results in more than a remote likelihood that a material misstatement of the annual or interim financial statements will not be prevented or detected.

Management has concluded that we did maintain effective internal control over financial reporting as of December 31, 2018, based on the criteria set forth in "Internal Control—Integrated Framework" issued by the COSO.

ITEM 9B. Other Information.

The price per share of our common stock has closed at or below \$0.20 since February 26, 2019 and most recently closed on March 26, 2019 at \$0.16, with a 30-day average of \$0.19. On March 26, 2016, we received written notice (the “Notice”) from the NYSE American LLC (the “NYSE American”) that we are not in compliance with the continued listing standards set forth in Section 1003(f)(v) of the NYSE American Company Guide because our common stock has been selling for a low price per share for a substantial period of time. The NYSE American has determined that the continued listing of our common stock is predicated on us effecting a reverse stock split of our common stock or otherwise demonstrating sustained price improvement within a reasonable period of time. We have until September 26, 2019 to demonstrate compliance.

Prior to our receipt of the Notice, and based upon our significant advances in oncology, we were contemplating seeking stockholder approval at our next annual meeting for a reverse stock split to raise the market price of our common stock to a range where it could allow a broader variety of institutions to invest in the common stock. Currently many funds are prohibited from buying stocks with a price below a certain threshold price. Should we have success in the current clinical trials, potentially increasing the trading volume and liquidity of the common stock confers major benefits. We believe that such a reverse stock split could help increase analyst and broker interest in the common stock, as their policies can discourage them from following or recommending companies with lower stock prices. Because of the trading volatility often associated with lower-priced stock, many brokerage houses and institutional investors have adopted internal policies and practices that either prohibit or discourage them from investing in such stocks or recommending them to their customers. Some of those policies and practices may also function to make the processing of trades in lower-priced stocks economically unattractive to brokers.

The Company now plans on holding a special meeting of its stockholders to approve a reverse stock split which it anticipates holding in late May or early June 2019.

As discussed above in PART I. Item 1 – “Business”, there are a number of existing and planned business activities that, we believe, should increase stockholder value and the market price of our common stock. However, we cannot assure that ongoing activities will continue to be positive or that such activities will increase stockholder value or the market price of our common stock. Nor can we assure that stockholders will approve the reverse split.

There is no immediate impact on the listing of our common stock, which will continue to trade on the NYSE American, subject to our compliance with other listing standards.

PART III

ITEM 10. Directors and Executive Officers and Corporate Governance.

The following sets forth biographical information about each of our Directors and Executive Officers as of the date of this report:

Name	Age	Position
Thomas K. Equels, Esq	66	Chief Executive Officer, President, and Director
Peter W. Rodino III	67	Executive Director Gov't Relations, General Counsel, & Secretary
William M. Mitchell, M.D., Ph.D.	84	Chairman of the Board and Director
Stewart L. Appelrouth	65	Director
Adam Pascale	71	Chief Financial Officer
David R. Strayer, M.D.	73	Chief Scientific Officer and Medical Director
Wayne S. Springate	47	Senior Vice President of Operations

Each Director has been elected to serve until the next annual meeting of stockholders, or until his earlier resignation, removal from office, death or incapacity. Each Executive Officer serves at the discretion of the Board of Directors, subject to rights, if any, under contracts of employment.

We believe our Board Members represent a desirable diversity of backgrounds, skills, education and experiences, and they all share the personal attributes of dedication to be effective directors. In recommending Board candidates, Corporate Governance and Nomination Committee considers a candidate's: (1) general understanding of elements relevant to the success of a publicly traded company in the current business environment; (2) understanding of our business; and (3) diversity in educational and professional background. The Committee also gives consideration to a candidate's judgment, competence, dedication and anticipated participation in Board activities along with experience, geographic location and special talents or personal attributes. The following are qualifications, experience and skills for Board members which are important to Hemispherx' business and its future:

Leadership Experience: We seek directors who have demonstrated strong leadership qualities. Such leaders bring diverse perspectives and broad business insight to our Company. The relevant leadership experience that we seek includes a past or current leadership role in a large or entrepreneurial company, a senior faculty position at a prominent educational institution or a past elected or appointed senior government position.

Industry or Academic Experience: We seek directors who have relevant industry experience, both with respect to the disease areas where we are developing new therapies as well as with the economic and competitive dynamics of pharmaceutical markets, including those in which our drugs will be prescribed.

Scientific, Legal or Regulatory Experience: Given the highly technical and specialized nature of biotechnology, we desire that certain of our directors have advanced degrees, as well as drug development experience. Since we are subject to substantial regulatory oversight, both here and abroad by the FDA and other agencies, we also desire directors who have legal or regulatory experience.

Finance Experience: We believe that our directors should possess an understanding of finance and related reporting processes, particularly given the complex budgets and long timelines associated with drug development programs.

THOMAS K. EQUELS, Esq., has been a Director and serves as our Executive Vice Chairman (since 2008), Chief Executive Officer (since 2016) and President (since 2015). Mr. Equels is the owner of and former President and Managing Director of the Equels Law Firm headquartered in Miami, Florida that focuses on litigation. For over a quarter century, Mr. Equels has represented national and state governments as well as companies in the banking, insurance, aviation, pharmaceutical and construction industries. Mr. Equels received his Juris Doctor degree with high honors from Florida State University. He is a summa cum laude graduate of Troy University and also obtained his

Masters' Degree from Troy. He is a member of the Florida Bar Association and the American Bar Association.

THOMAS K. EQUELS, Esq. - Director Qualifications:

Leadership Experience – Owner and former President, Managing Director of Equels Law Firm;

Industry Experience –legal counsel to Hemispherx; and

Scientific, Legal or Regulatory Experience - Law degree with over 25 years as a practicing attorney specializing in litigation.

WILLIAM M. MITCHELL, M.D., Ph.D., has been a Director since July 1998. On February 17, 2016, Dr. Mitchell was appointed as Chairman of the Board upon Dr. Carter's termination. Dr. Mitchell is a Professor of Pathology at Vanderbilt University School of Medicine and is a board certified physician. Dr. Mitchell earned a M.D. from Vanderbilt and a Ph.D. from Johns Hopkins University, where he served as House Officer in Internal Medicine, followed by a Fellowship at its School of Medicine. Dr. Mitchell has published over 200 papers, reviews and abstracts that relate to viruses, anti-viral drugs, immune responses to HIV infection, and other biomedical topics. Dr. Mitchell has worked for and with many professional societies that have included the American Society of Investigative Pathology, the International Society for Antiviral Research, the American Society of Clinical Oncology, the American Society of Biochemistry and Molecular Biology and the American Society of Microbiology. Dr. Mitchell is a member of the American Medical Association. He has served on numerous government review committees, among them the National Institutes of Health, AIDS and Related Research Review Group. Dr. Mitchell previously served as one of our Directors from 1987 to 1989.

WILLIAM M. MITCHELL, M.D., Ph.D. - Director Qualifications:

Leadership Experience – Professor at Vanderbilt University School of Medicine. He is a member of the Board of Directors for Chronix Biomedical and is Chairman of its Medical Advisory Board. Additionally, he has served on multiple governmental review committees of the National Institutes of Health, Centers for Disease Control and Prevention and for the European Union, including key roles as Chairman;

Academic and Industry Experience – Well published medical researcher with extensive investigative experience on virus and immunology issues relevant to the scientific business of Hemispherx along with being a Director of an entrepreneurial diagnostic company (Chronix Biomedical) that is involved in next generation DNA sequencing for medical diagnostics; and

Scientific, Legal or Regulatory Experience - M.D., Ph.D. and professor at a top ranked school of medicine, and inventor of record on numerous U.S. and international patents who is experienced in regulatory affairs through filings with the FDA.

STEWART L. APPELROUTH, CPA was appointed as a director and head of the Audit Committee in August 2016 and is a certified public accountant and partner at Appelrouth Farah & Co., P.A., Certified Public Accountants and Advisors. Mr. Appelrouth is also a certified forensic accountant and possesses 40 years of experience in Accounting and Consulting. He is a member of or has affiliations with the AICPA, American College of Forensic Examiners, Association of Certified Fraud Examiners, Florida Bar Grievance Committee, Florida Institute of Certified Public Accountants and InfraGard Member, a national information sharing program between the Federal Bureau of Investigation and the private sector.

Mr. Appelrouth graduated from Florida State University in 1975 and received his Master's Degree in Finance from Florida International University in 1980. The Board has determined Mr. Appelrouth to be an Independent Director as required under Section 803(2) of the NYSE: American Company Guide and Rule 10A-3 under the Exchange Act.

STEWART L. APPELROUTH - Director Qualifications:

Leadership Experience –has served in leadership positions on numerous Boards and other organizations;

Industry Experience – Partner at certified public accounting and advisory firm; Certified Public Accountant and Certified Fraud Examiner;

Regulatory Experience – FINRA Arbitrator.

Financial Expert – over 40 years of accounting and audit experience.

ADAM PASCALE was promoted to Chief Financial Officer on February 22, 2016. He is also the Company's Chief Accounting Officer. Mr. Pascale has been employed with the Company for 23 years, with more than two decades of public accounting experience and prior public company experience. He earned a Bachelor of Arts degree in Accounting and Finance from Rutgers University. Mr. Pascale served for several years as a CPA prior to joining Hemispherx, and is a member of both the American and the Pennsylvania Institutes of Certified Public Accountants.

DAVID R. STRAYER, M.D. has acted as our Medical Director since 1986. On February 19, 2016, Dr. Strayer was appointed as Chief Scientific Officer upon Dr. Carter's termination. He has served as Professor of Medicine at the Medical College of Pennsylvania and Hahnemann University. Dr. Strayer is Board Certified in Medical Oncology and Internal Medicine with research interests in the fields of cancer and immune system disorders. He has served as principal investigator in studies funded by the Leukemia Society of America, the American Cancer Society, and the National Institutes of Health. Dr. Strayer attended the School of Medicine at the University of California at Los Angeles where he received his M.D. in 1972.

PETER W. RODINO III was appointed a Director in July 2013. On September 30, 2016, Mr. Rodino resigned as a member of our Board to permit him to serve us in a new capacity. In this regard, effective October 1, 2016, we retained Mr. Rodino as our Executive Director for Governmental Relations, and as our General Counsel. In that capacity, Mr. Rodino handles all government affairs and litigation matters on a going forward basis. Mr. Rodino was also appointed Secretary of the Company in November 2016. Through September 30, 2016, Mr. Rodino served as Lead Director and Chairman and Financial Expert of the Audit Committee, a member of the Compensation Committee and a member of the Governance and Nomination Committee of the Board of Directors. Mr. Rodino has broad legal, financial, and executive experience. In addition to being President of Rodino Consulting LLC and managing partner at several law firms during his many years as a practicing attorney, he served as Chairman and CEO of Crossroads Health Plan, the first major Health Maintenance Organization in New Jersey. He also has had experience as an investment executive in the securities industry and acted as trustee in numerous Chapter 11 complex corporate reorganizations. Previously, as founder and president of Rodino Consulting, Mr. Rodino provided business and government relations consulting services to smaller companies with a focus on helping them develop business plans, implement marketing strategies and acquire investment capital. Mr. Rodino holds a B.S. in Business Administration from Georgetown University and a J.D. degree from Seton Hall University.

WAYNE S. SPRINGATE was promoted to Senior Vice President of Operations on May 1, 2011. Mr. Springate joined Hemispherx in 2002 as Vice President of Business Development when Hemispherx acquired Alferon N Injection® and its New Brunswick, NJ manufacturing facilities. He led the consolidation of our Rockville facility to our New Brunswick location as well as coordinated the relocation of manufacturing polymers from South Africa to our production facility in New Brunswick. Previously, Mr. Springate served as President for World Fashion Concepts in New York and oversaw operations at several locations throughout the United States and overseas. Mr. Springate assists the CEO in details of operations on a daily basis and is involved in all aspects of manufacturing, warehouse management, distribution and logistics.

Compliance with Section 16(a) of the Exchange Act

Section 16(a) of the Exchange Act requires our Officers and Directors, and persons who own more than ten percent of a registered class of equity securities, to file reports with the Securities and Exchange Commission reflecting their initial position of ownership on Form 3 and changes in ownership on Form 4 or Form 5. Based solely on a review of the copies of such Forms received by us, we found that, during the fiscal year ended December 31, 2018, all of our Officers and Directors had complied with all applicable Section 16(a) filing requirements on a timely basis with regard to transactions occurring in 2018.

Audit Committee and Audit Committee Expert

The Audit Committee of our Board of Directors consists of William Mitchell, M.D. and Stewart L. Appelrouth. Dr. Mitchell and Mr. Appelrouth are determined by the Board of Directors to be Independent Directors as required under Section 803(2) of the NYSE: American Company Guide and Rule 10A-3 under the Exchange Act. The Board has determined that Mr. Appelrouth qualifies as an “audit committee financial expert” as that term is defined by Section 803B(2) of the NYSE: American Company Guide and the rules and regulations of the SEC.

We believe Dr. Mitchell and Mr. Appelrouth to be independent of management and free of any relationship that would interfere with their exercise of independent judgment as members of this Committee. The principal functions of the Audit Committee are to (i) assist the Board in fulfilling its oversight responsibility relating to the annual independent audit of our consolidated financial statements and internal control over financial reporting, the engagement of the independent registered public accounting firm and the evaluation of the independent registered public accounting firm’s qualifications, independence and performance; (ii) prepare the reports or statements as may be required by NYSE American or the securities laws; (iii) assist the Board in fulfilling its oversight responsibility relating to the integrity of our financial statements and financial reporting process and our system of internal accounting and financial controls; (iv) discuss the financial statements and reports with management, including any significant adjustments, management judgments and estimates, new accounting policies and disagreements with management; and (v) review disclosures by our independent registered public accounting firm concerning relationships with us and the performance of our independent accountants.

This Audit Committee formally met six times in 2018 with all committee members in attendance. Our General Counsel and Chief Financial Officer support the Audit Committee in its work. The full text of the Audit Committee's Charter, as approved by the Board, is available on our website: www.hemispherx.net in the "Investor Relations" tab under "Corporate Governance".

Scientific Advisory Board

The SAB was established to leverage its member's scientific and pharmaceutical expertise and advice to advance our drug development programs by providing guidance on steering us forward and capitalizing on business opportunities as well as interactions with the FDA. It is responsible for: (i) reviewing all submissions made by us to the FDA and other regulators to ensure that the submissions fully, accurately, and timely describe the status of any clinical trials, tests, or other studies or analyses of drug safety and efficacy undertaken by us, and any agreements, protocols, or guidance provided by relevant regulatory agencies; and (ii) monitoring and supervising our relationship with the FDA. The SAB shall have free and open access to our scientific and executive personnel, including the Chief Scientific Officer and the members of our Board of Directors. The SAB is comprised of William Mitchell, M.D., Chairman, and Ronald Brus, M.D., W. Neal Burnette, M.D., and Christopher Nicodemus, M.D., all of whom are members. The SAB reports to the independent directors of the Company and closely interacts with the Disclosure Controls Committee. The Scientific Advisory Board met two times in 2018.

Disclosure Controls Committee

The DCC reports to the Audit Committee and is responsible for procedures and guidelines on managing disclosure information. The purpose of the DCC is to make certain that information required to be publicly disclosed is properly accumulated, recorded, summarized and communicated to the Board and management. This process is intended to allow for timely decisions regarding communications and disclosures and to help ensure that we comply with related SEC rules and regulations. Wayne Springate, Senior Vice President of Operations is the DCC's Investor Relations Coordinator and Chairperson. The other members of the DCC are Peter Rodino, our General Counsel, William Mitchell, one of our Independent Directors, Dr. David Strayer, our Chief Scientific Officer, Adam Pascale, our Chief Financial and Accounting Officer, and Ann Marie Coverly, Director of HR and Administration serving as the Deputy Investor Relations Coordinator. The full text of the DCC's Charter, as approved by the Board, is available on our website: www.hemispherx.net in the "Investor Relations" tab under "Corporate Governance". The DCC actively met on numerous occasions in 2018.

Executive Committee

In February 2016, our Board formed the Executive Committee. The Executive Committee reports to the Board and its purpose is to aid the Board in handling matters which, in the opinion of the Chairman of the Board, should not be postponed until the next scheduled meeting of the Board. Mr. Equels, our Chief Executive Officer is the chairman of the Committee, along with two of our independent directors, Mr. Appelrouth and Dr. Mitchell. The full text of the Executive Committee Charter, as approved by the Board, is available on our website: www.hemispherx.net in the "Investor Relations" tab under "Corporate Governance". The Committee did not meet in 2018.

Code of Ethics

Our Board of Directors adopted a revision to the 2003 Code of Ethics and business conduct for officers, directors, employees, agents and consultants. The principal amendments included broadening the Code's application to our agents and consultants, adoption of a regulatory compliance policy and adoption of a policy for protection and use of Company computer technology for business purposes only. On an annual basis, this Code is reviewed and signed by each Officer, Director, employee and strategic consultant with none of the amendments constituting a waiver of provision of the Code of Ethics on behalf of our Chief Executive Officer, Chief Financial Officer, or persons performing similar functions.

You may obtain a copy of this Code by visiting our web site at www.hemispherx.net (Investor Relations / Corporate Governance) or by written request to our office at 2117 SW Highway 484, Ocala, FL 34473.

ITEM 11. Executive Compensation.

COMPENSATION DISCUSSION AND ANALYSIS

This discussion and analysis describes our executive compensation philosophy, process, plans and practices as they relate to our “Named Executive Officers” (“NEO”) listed below and gives the context for understanding and evaluating the more specific compensation information contained in the narratives, tables and related disclosures that follow. For the purposes of discussion and analysis, the following NEOs are included in the narratives, tables and related disclosures that follow:

Thomas K. Equels, Chief Executive Officer (“CEO”) and President;

Adam Pascale, Chief Financial Officer (“CFO”); and

Peter Rodino, General Counsel and Company Secretary (“CS”)

Governance of Compensation Committee

The Compensation Committee consists of the following two directors, each of whom is “independent” under applicable NYSE American rules, a “Non-Employee Director” as defined in Rule 16b-3 under the Exchange Act, and an “Outside Director” as defined under the U.S. Treasury regulations promulgated under Section 162(m) of the Internal Revenue Code of 1986, as amended (the “Internal Revenue Code”): Dr. William Mitchell, M.D. (Chair) and Stewart L. Appelrouth. The Compensation Committee makes recommendations concerning salaries and compensation for senior management and other highly paid professionals or consultants to Hemispherx. The full text of the Compensation Committee’s Charter, as approved by the Board, is available on our website: www.hemispherx.net in the “Investor Relations” tab under “Corporate Governance”.

This Committee formally met two times in 2018 and all committee members were in attendance for the meetings with the exception of one meeting. Our General Counsel, Chief Financial Officer and Director of Human Resources support the Compensation Committee in its work.

Results of Stockholder Advisory Vote on Executive Compensation

At the September 12, 2018 Annual Meeting of Stockholders, the Stockholders approved the annual, non-binding advisory vote on Executive Compensation.

Objectives and Philosophy of Executive Compensation

The primary objectives of the Compensation Committee of our Board of Directors with respect to Executive compensation are to attract and retain the most talented and dedicated Executives possible, to tie annual and long-term cash and stock incentives to achievement of measurable performance objectives, and to align Executives' incentives with stockholder value creation. To achieve these objectives, the Compensation Committee expects to implement and maintain compensation plans that tie a substantial portion of Executives' overall compensation to key strategic financial and operational goals such as the establishment and maintenance of key strategic relationships, the development of our products, the identification and advancement of additional products and the performance of our common stock price. The Compensation Committee evaluates individual Executive performance with the goal of setting compensation at levels the Committee believes are comparable with Executives in other companies of similar size and stage of development operating in the biotechnology industry while taking into account our relative performance, our own strategic goals, governmental regulations and the results of Stockholder Advisory Votes regarding executive compensation.

EXECUTIVE COMPENSATION

The following table provides information on the compensation during the fiscal years ended December 31, 2018 and 2017 of our Chief Executive Officer, Chief Financial Officer, and Peter Rodino, General Counsel and Company Secretary constituting the Company's Named Executive Officers, based on the year ended 2018 for each fiscal year.

Summary Compensation Table

Name & Principal Position	Year	Salary / Fees (2)(3)	Bonus	Stock Awards	Option Awards (1)	Non-Equity Incentive Plan Compensation	Change in Pension Value and NQDC	All Other Compensation	Total (3)
---------------------------	------	-------------------------	-------	--------------	-------------------	--	----------------------------------	------------------------	-----------

							Earnings (\$)			
Thomas K. Equels CEO & President (2) (3)	2018	\$751,000	\$18,350(3)	\$500	\$335,731(1)	—	—	\$65,927	(4)	\$1,171,508
	2017	\$712,500	\$22,067(3)	—	\$174,052(1)	—	—	\$78,604	(4)	\$987,223
Adam Pascale CFO	2018	\$251,000	\$—	\$500	\$99,979	—	—	\$40,973	(6)	\$392,452
	2017	\$234,500	\$—	\$—	\$12,501	—	—	\$48,379	(6)	\$295,380
Peter Rodino General Counsel and Secretary	2018	\$351,000	\$—	\$500	\$138,599	—	—	\$36,684	(5)	\$526,783
	2017	\$332,500	\$—	—	\$17,500	—	—	\$48,656	(5)	\$398,656

Notes:

(1) All option awards were valued using the Black-Scholes method.

For Named Executive Officers, who are also Directors that receive compensation for their services as a Director, the Salary/Fees and Option Awards columns include compensation that was received by them for their role as a (2) member of the Board of Directors. As is required by Regulation S-K, Item 402(c), compensation for services as a Director have been reported within the “Summary Compensation Table” (above) for fiscal years of 2018 and 2017 as well as reported separately in the “Compensation of Directors” section (see below) for calendar year 2018.

(3) As stated in Thomas Equels’ employment contract, he is entitled to 5% of Ampligen® sales. In 2018 and 2017, a bonus of 5% of Ampligen® sales for 2018 and 2017 was accrued.

For 2017, salaries for Messrs. Equels, Pascale and Rodino include 40% deferred salaries of \$100,000, \$33,333 and \$46,667, respectively, starting from September 1, 2017.

For 2018, salaries for Messrs. Equels, Pascale and Rodino include 50% deferred salaries of \$140,625, \$46,875 and \$65,625, respectively, starting from August 1, 2018

(4) Mr. Equels’ All Other Compensation consists of:

	2018	2017
Life and Disability Insurance	\$22,677	\$26,837
Healthcare Insurance	25,250	33,767
Car Expenses / Allowance	18,000	18,000
401(k) Matching Funds	—	—
	\$65,927	\$78,604

(5) Mr. Rodino’s All Other Compensation consists of:

Edgar Filing: HEMISPHERX BIOPHARMA INC - Form 10-K

	2018	2017
Life and Disability Insurance	\$2,542	\$2,504
Healthcare Insurance	19,742	31,752
Car Expenses / Allowance	14,400	14,400
401(k) Matching Funds	—	—
	\$36,684	\$48,656

(6)Mr. Pascale's All Other Compensation consists of:

	2018	2017
Life and Disability Insurance	\$2,272	\$2,212
Healthcare Insurance	24,301	31,767
Car Expenses / Allowance	14,400	14,400
401(k) Matching Funds	-	-
	\$40,973	\$48,379

Outstanding Equity Awards at Fiscal Year End

Name	Option Awards					Stock Awards				
	Number of Securities Underlying Unexercised Options (#) Exercisable	Number of Securities Underlying Unexercised Options (#) Unexercisable	Equity Incentive Plan Awards: Number of Securities Underlying Unearned Options (#)	Option Exercise Price (\$)	Option Expiration Date	Number of Shares or Units of Stock That Have Not Vested (#)	Market Value of Shares or Units of Stock That Have Not Vested (\$)	Plan Awards: Number of Shares, Units or Rights That Have Not Vested (#)	Equity Incentive Awards: Payout Value of Unearned Shares, Units or Rights that Have Not Vested (#)	Equity Incentive Awards: Market Value of Unearned Shares, Units or Rights that Have Not Vested (#)
Thomas	25,000	—	—	7.92	6/11/2020	—	—	—	—	—
Equels,	25,000	—	—	4.92	6/24/2021	—	—	—	—	—
President and Chief	8,333	—	—	3.48	6/6/2022	—	—	—	—	—
Executive Officer	25,000	—	—	3.72	6/11/2022	—	—	—	—	—
	25,000	—	—	3.72	6/6/2023	—	—	—	—	—
	12,500	—	—	3.00	8/2/2023	—	—	—	—	—
	25,000	—	—	4.32	6/6/2024	—	—	—	—	—
	25,000	—	—	3.00	6/8/2025	—	—	—	—	—
	25,000	—	—	1.68	6/8/2026	—	—	—	—	—
	300,000	—	—	0.56	6/8/2027	—	—	—	—	—
	14,212	—	—	0.49	6/15/2027	—	—	—	—	—
	14,214	—	—	0.49	6/30/2027	—	—	—	—	—
	—	18,124	—	0.48	7/15/2027	—	—	—	—	—
	—	20,786	—	0.42	7/31/2027	—	—	—	—	—
	—	21,336	—	0.41	8/15/2027	—	—	—	—	—
	—	24,463	—	0.36	8/31/2027	—	—	—	—	—
	—	371,622	—	0.37	2/13/2028	—	—	—	—	—
	—	125,000	—	0.38	4/12/28	—	—	—	—	—
	—	300,000	—	0.30	5/16/28	—	—	—	—	—
	—	250,000	—	0.30	5/16/28	—	—	—	—	—
	—	161,290	—	0.31	7/18/21	—	—	—	—	—
	—	284,091	—	0.22	10/17/28	—	—	—	—	—
	—	1,000	—	0.22	10/17/28	—	—	—	—	—

Edgar Filing: HEMISPHERX BIOPHARMA INC - Form 10-K

	—	426,136	—	0.22	1/28/2029	—	—	—	—
Adam Pascale, Chief Financial Officer	500	—	—	48.36	4/13/2022	—	—	—	—
	4,167	—	—	3.96	7/8/2024	—	—	—	—
	598	—	—	1.56	6/21/2026	—	—	—	—
	12,500	—	—	0.49	6/15/2027	—	—	—	—
	—	4,738	—	0.49	6/30/2027	—	—	—	—
	—	7,738	—	0.48	7/15/2027	—	—	—	—
	—	6,042	—	0.42	7/31/2027	—	—	—	—
	—	6,929	—	0.41	8/15/2027	—	—	—	—
	—	7,112	—	0.36	8/31/2027	—	—	—	—
	—	8,155	—	0.37	2/13/2028	—	—	—	—
	—	123,874	—	0.38	4/12/28	—	—	—	—
	—	75,000	—	0.30	5/16/21	—	—	—	—
	—	83,334	—	0.31	7/18/21	—	—	—	—
	—	53,763	—	0.22	10/17/19	—	—	—	—
	—	94,697	—	0.22	11/14/19	—	—	—	—
	—	1,000	—	0.22	1/28/2029	—	—	—	—
	—	142,046	—	3.00	8/2/2023	—	—	—	—
Peter Rodino, General Counsel and Secretary	12,500	—	—	1.56	6/21/2026	—	—	—	—
	12,500	—	—	0.49	6/15/2027	—	—	—	—
	—	6,632	—	0.49	6/30/2027	—	—	—	—
	—	6,633	—	0.48	7/15/2027	—	—	—	—
	—	8,458	—	0.42	7/31/2027	—	—	—	—
	—	9,700	—	0.41	8/15/2027	—	—	—	—
	—	9,957	—	0.36	8/31/2027	—	—	—	—
	—	11,416	—	0.37	2/13/2028	—	—	—	—
	—	173,423	—	0.38	4/12/2028	—	—	—	—
	—	100,000	—	0.30	5/16/2028	—	—	—	—
	—	116,667	—	0.31	7/18/2028	—	—	—	—
	—	75,269	—	0.22	10/17/2028	—	—	—	—
	—	132,576	—	0.22	11/14/2028	—	—	—	—
	—	1,000	—	0.22	1/28/2029	—	—	—	—
	—	198,864	—	0.22	1/28/2029	—	—	—	—

Payments on Disability

At December 31, 2018, we had an employment agreement with Mr. Equels which entitled him Base Salary and applicable benefits otherwise due and payable through the last day of the month in which disability occurs and for an additional twelve-month period. Each current NEO, including Mr. Pascale and Mr. Rodino, has the same short and long-term disability coverage which is available to all eligible employees. The coverage for short-term disability provides up to six months of full salary continuation up to 60% of weekly pay, less other income, with a \$1,500 weekly maximum limit. The coverage for group long-term disability provides coverage at the exhaustion of short-term disability benefits of full salary continuation up to 60% of monthly pay, less other income, with a \$10,000 monthly maximum limit. The maximum benefit period for the group long-term disability coverage is 60 months for those age 60 and younger at the time of the claim with the coverage period proportionately reduced with the advanced age of the eligible employee to a minimum coverage period of 12 months for those of 69 years old and older as of the date of the claim. For the period June 2010 through 2017 pursuant to his respective employment agreement and payable by us, Mr. Equels is entitled to receive total disability coverage of \$400,000.

Payments on Death

At December 31, 2018, we had an employment agreement with Mr. Equels which entitled him Base Salary and applicable benefits otherwise due and payable through the last day of the month in which death occurs and for an additional twelve-month period. Each NEO, including Mr. Pascale and Mr. Rodino, has coverage of group life insurance, along with accidental death and dismemberment benefits, consistent to the dollar value available to all eligible employees. The benefit is equal to two times current salary or wage with a maximum limit of \$300,000, plus any supplemental life insurance elected and paid for by the NEO. For the period June 2010 and through 2018 pursuant to his respective employment agreements and payable by us, Mr. Equels is entitled to receive total death benefit coverage of \$3,000,000.

Estimated Payments Following Severance — Named Executive Officers (NEO)

At December 31, 2018, we had an employment agreement with Mr. Equels which entitled him to severance benefits on certain types of employment terminations not related to a change in control. Mr. Rodino and Mr. Pascale are not covered by an employment severance agreement and therefore would only receive severance as determined by the Compensation Committee in its discretion.

The dollar amounts below assume that the termination occurred on January 1, 2019. The actual dollar amounts to be paid can only be determined at the time of the NEO's separation from Hemispherx based on their prevailing compensation and employment agreements along with any determination by the Compensation Committee in its

discretion.

39

Name	Event	Cash Severance (\$)	Value of Stock Awards That Will Become Vested (1) (\$)	Continuation of Medical Benefits (\$)	Additional Life Insurance (\$)	Total (\$)
Thomas K. Equels, CEO & President	Involuntary (no cause)	\$ 768,000	\$ 62,320	—	—	\$ 830,320
	Termination (for cause)	—	—	—	—	—
	Death or disability	\$ 768,000	\$ 62,320	—	—	\$ 830,320
	Termination by employee or retirement	\$ 768,000	\$ 62,320	—	—	\$ 830,320
Adam Pascale CFO	Involuntary (no cause)	—	—	—	—	—
	Termination (for cause)	—	—	—	—	—
	Death or disability	—	—	—	—	—
	Termination by employee or retirement	—	—	—	—	—
Peter Rodino General Counsel and Secretary	Involuntary (no cause)	—	—	—	—	—
	Termination (for cause)	—	—	—	—	—
	Death or disability	—	—	—	—	—
	Termination by employee or retirement	—	—	—	—	—

Notes:

- Consists of stock options contractually required per the employee's respective Employment Agreement to be granted during each calendar year of the term under our 2009 Equity Incentive Plan. The stock options have a ten-year term and an exercise price equal to the closing market price of our common stock on the date of grant.
- (1) For the purpose of this schedule, an NYSE American closing price at March 16, 2018 of \$0.30 was used with an estimated exercise price of \$0.30 for Mr. Equels. The value was obtained using the Black-Scholes-Merton pricing model for stock-based compensation in accordance with FASB ASC 718.

Payments on Termination in Connection with a Change in Control Named Executive Officers

At December 31, 2018, we had an employment agreement with Mr. Equels which entitled him to severance benefits on certain types of employment terminations related to a change in control whereby the term of his respective agreement would automatically be extended for three additional years. Mr. Rodino and Mr. Pascale are not covered by employment severance agreement and therefore would only receive severance from a change in control as determined

by the Compensation Committee in its discretion. Any specific benefits for these two NEO would be determined by the Compensation Committee in its discretion.

The dollar amounts in the chart below assume that change in control termination occurred on January 1, 2019, based on the employment agreements that existed at that time. The actual dollar amounts to be paid can only be determined at the time of the NEO's separation from Hemispherx based on their prevailing compensation and employment agreements along with any determination by the Compensation Committee in its discretion.

Estimated Benefits on Termination Following a Change in Control — December 31, 2018

The following table shows potential payments to the NEO if employment terminates following a change in control under contracts, agreements, plans or arrangements at December 31, 2018. The amounts assume a January 1, 2019 termination date regarding base pay and use of the opening price of \$0.18 on the NYSE American for our common stock at that date.

Name	Aggregate Severance Pay (\$)	PVSU Acceleration (2) (\$)	Early Vesting of Restricted Stock (4) (\$)	Early Vesting of Stock and SARs (3) (\$)	Acceleration and Vesting of Supplemental Award (5) (\$)	Welfare Benefits Continuation (\$)	Outplacement Assistance (\$)	Parachute Tax Gross-up Payment (\$)	Total (\$)
Thomas K. Equels	768,000 (1)	—	—	—	\$62,320 (4)	—	—	—	\$830,320
Adam Pascale	—	—	—	—	—	—	—	—	—
Peter Rodino	—	—	—	—	—	—	—	—	—

Notes:

(1) This amount represents the base salary and benefits for remaining term of the NEO's employment agreement plus a three-year extension in the term upon the occurrence of a termination from a change in control. The employment agreement with Mr. Equels had a term through December 31, 2016; however, this was automatically extended for an additional three-year period through December 31, 2019.

(2) This amount represents the payout of all outstanding performance-vesting share units ("PVSU") awarded on a change in control at the target payout level with each award then pro-rated based on the time elapsed for the applicable three-year performance period.

(3) This amount is the intrinsic value [fair market value on January 1, 2018 (\$0.18 per share) minus the per share exercise price of \$0.30 of all unvested stock options for each NEO, including Stock Appreciation Rights ("SAR"). Any option with an exercise price of greater than fair market value was assumed to be cancelled for no consideration and, therefore, had no intrinsic value.

(4) This amount represents the options to be issued annually for the remaining term of the NEO's employment agreement plus a three-year extension in the occurrence of termination from a change in control. For the purpose of this schedule, an NYSE American closing price at March 16, 2018 of \$0.30 was used with an estimated exercise price of \$0.30 for Mr. Equels. The value was obtained using the Black-Scholes-Merton pricing model for stock-based compensation in accordance with FASB ASC 718.

(5) Any purchase rights represented by the Option not then vested shall, upon a change in control, shall become vested.

Definition of "Change in Control" for each agreement, a "Change in Control" is defined generally as any such event that requires a report to the SEC, but includes any of the following:

Any person or entity other than Hemispherx, any of our current Directors or Officers or a Trustee or fiduciary holding our securities, becomes the beneficial owner of more than 50% of the combined voting power of our outstanding securities;

An acquisition, sale, merger or other transaction that results in a change in ownership of more than 50% of the combined voting power of our stock or the sale/transfer of more than 75% of our assets;

A change in the majority of our Board of Directors over a two-year period that is not approved by at least two-thirds of the Directors then in office who were Directors at the beginning of the period; or

Execution of an agreement with Hemispherx, which if consummated, would result in any of the above events.

Definition of "Constructive Termination". A "Constructive Termination" generally includes any of the following actions taken by Hemispherx without the Executive's written consent following a change in control:

Significantly reducing or diminishing the nature or scope of the executive's authority or duties;

Materially reducing the executive's annual salary or incentive compensation opportunities;

Changing the executive's office location so that he must commute more than 50 miles, as compared to his commute as of the date of the agreement;

Failing to provide substantially similar fringe benefits, or substitute benefits that were substantially similar taken as a whole, to the benefits provided as of the date of the agreement; or

Failing to obtain a satisfactory agreement from any successor to Hemispherx to assume and agree to perform the obligations under the agreement.

However, no constructive termination occurs if the executive:

Fails to give us written notice of his intention to claim constructive termination and the basis for that claim at least 10 days in advance of the effective date of the executive's resignation; or

We cure the circumstances giving rise to the constructive termination before the effective date of the executive's resignation.

Available Information

Our Internet website is www.hemispherx.net and you may find our SEC filings in the "Investor Relations" under "SEC Filings". We provide access to our filings with the SEC, free of charge through www.sec.gov, as soon as reasonably practicable after filing with the SEC. Our Internet website and the information contained on that website, or accessible from our website, is not intended to be incorporated into this Annual Report on Form 10-K or any other filings we make with the SEC.

Post-Employment Compensation

We have an agreement with the following NEO who has benefits upon termination as a condition of his respective employment agreement: Thomas K. Equels, our CEO.

The following is a description of post-employment compensation payable to the respective NEO. If a NEO does not have a specific benefit, they will not be mentioned in the subsection. In such event, the NEO does not have any such benefits upon termination unless otherwise required by law.

Termination for Cause

All of our NEOs can be terminated for cause. For Mr. Equels, “Cause” means willful engaging in illegal conduct, gross misconduct or gross violation of the Company’s Code of Ethics and Business Conduct for Officers which is demonstrably and materially injurious to the Company. For purposes of his respective agreement, no act, or failure to act, on employee’s part shall be deemed “willful” unless done intentionally by employee and not in good faith and without reasonable belief that employee’s action or omission was in the best interest of the Company. Notwithstanding the foregoing, employee shall not be deemed to have been terminated for Cause unless and until the Company delivers to the employee a copy of a resolution duly adopted by the affirmative vote of not less than three-quarters of the Directors of the Board at a meeting of the Board called and held for such purpose (after reasonable notice to employee and an opportunity for Employee, together with counsel, to be heard before the Board) finding that, in the good faith opinion of the Board, employee was guilty of conduct set forth above and specifying the particulars thereof in detail. In the event that his employment is terminated for Cause, the Company shall pay him, at the time of such termination, only the compensation and benefits otherwise due and payable to them through the last day of their actual employment by the Company.

Termination without Cause

Mr. Equels is entitled to the compensation and benefits otherwise due and payable to him through the last day of the then current term of their respective agreements. In the event that he is terminated at any time without “Cause” the Company shall pay to him, at the time of such termination, the compensation and benefits otherwise due and payable through the last day of the then current term of their Agreement. However, benefit distributions that are made due to a “separation from service” occurring while he is a Named Executive Officer shall not be made during the first six months following separation from service. Rather, any distribution which would otherwise be paid to him during such period shall be accumulated and paid to him in a lump sum on the first day of the seventh month following the “separation from service”. All subsequent distributions shall be paid in the manner specified.

Death or Disability

Mr. Equels can be terminated for death or disability. For each, “Disability” means the inability to effectively carry out substantially all of his duties under their agreement by reason of any medically determinable physical or mental impairment which can be expected to result in death or which has lasted for a continuous period of not less than 12 months. In the event the employment is terminated due to his death or disability, the Company will pay (or their respective estate as the case may be), at the time of such termination, the Base Salary and applicable benefits otherwise due and payable through the last day of the month in which such termination occurs and for an additional 12 month period.

Termination by Officer and Employee

All NEO employment agreements have the right to terminate their respective agreement upon thirty (30) days or less of prior written notice of termination. In such event, Mr. Equels is specifically entitled to fees due to him through the last day of the month in which such termination occurs and for 12 months thereafter. All other NEOs are entitled to the fees due to them through the last day of the month in which such termination occurs.

Change in Control

As an element of his employment agreement, Mr. Equels is entitled to benefits upon a Change in Control or Constructive Termination that include that any unvested Options immediately vest and the term of his respective employment agreement automatically extend for an additional three years. In the event of a Change in Control, the Company is responsible for the base salary or benefits for remaining term of the NEO's employment agreement plus an automatic three-year extension in the term of the agreement. The existing employment agreement with Mr. Equels had a term through December 31, 2016; however, this employment agreement automatically extended for an additional three-years through December 31, 2019.

Compensation of Directors

Our Compensation, Audit and Corporate Governance and Nomination Committees, consist of Dr. William M. Mitchell, Compensation and Corporate Governance and Nomination Committee Chair, and Stewart L. Appelrouth, Audit Committee Chair, both of whom are independent Board of Director members.

Hemispherx reimburses Directors for travel expenses incurred in connection with attending board, committee, stockholder and special meetings along with other Company business-related expenses. Hemispherx does not provide retirement benefits or other perquisites to non-employee Directors under any current program.

There was no cost of living increase granted in 2017 or 2018. Directors' fees are currently being deferred and will continue to be deferred until cash is available.

All Directors have been granted options to purchase common stock under our Stock Option Plans and/or Warrants to purchase common stock. We believe such compensation and payments are necessary in order for us to attract and

retain qualified outside directors. Options shares for stock compensation were issued under the 2009 and 2018 Equity Incentive Plans.

Director Compensation – 2018

Name and Title of Director	Fees Earned or Paid in Cash (\$)	Stock Award (\$)	Option Awards (\$)	Non-Equity Incentive Plan Compensation (\$)	Change in Pension Value and Nonqualified Deferred Compensation Earnings (\$)	All Other Compensation As Director (\$)	Total (\$)
T. Equels, Executive Vice Chairman	-	(2)	-	-	-	-	-
W. Mitchell, Chairman of the Board (1)	\$114,039	\$500	\$193,411	-	-	-	\$307,950
Stewart L. Appelrouth, Director (1)	\$114,039	\$500	\$193,411	-	-	-	\$307,950

Notes:

- (1) Independent Director of the Company. Beginning August 16, 2018, the independent directors are deferring payment of 100% of their director's fees until cash is available.

Only includes compensation received in the role as member of the Board of Directors and does not include compensation received in the capacity of a Named Executive Officer. As is required by Regulation S-K, Item (2)402(c), compensation as a Director has also been reported within the "Summary Compensation Table" regarding Named Executive Officer Compensation during fiscal years of 2018 and 2017 (see above). Mr. Equels stopped receiving Board fees in March 2016.

ITEM 12. Security Ownership of Certain Beneficial Owners and Management and Related Stockholder Matters.

The following table sets forth as of March 18, 2019, the number and percentage of outstanding shares of common stock beneficially owned by:

Each person, individually or as a group, known to us to be deemed the beneficial owners of five percent or more of our issued and outstanding common stock;

Each of our Directors and the Named Executives Officers; and

All of our officers and directors as a group.

Name and Address of Beneficial Owner	Shares Beneficially Owned		% Of Shares Beneficially Owned	
Thomas K. Equels	3,976,414	(1)(8)	6.4	%
Peter W. Rodino III 17400 Sterling Lake Drive Fort Myers, FL 33967	1,073,968	(2)(8)	1.7	%
William M. Mitchell, M.D. Vanderbilt University Department of Pathology Medical Center North 21st and Garland Nashville, TN 37232	1,750,550	(3)(8)	2.8	%
Stewart L. Appelrouth 999 Ponce de Leon., Suite 625 Coral Cables, FL33134	1,954,652	(7)(8)	3.1	%
Wayne S. Springate 783 Jersey Ave. New Brunswick, NJ 08901	791,026	(4)(8)	1.3	%
David R. Strayer, M.D.	542,235	(5)(8)	.9	%
Adam Pascale	791,341	(6)(8)	1.3	%

Edgar Filing: HEMISPHERX BIOPHARMA INC - Form 10-K

All directors and executive officers as a group (6 persons) 10,880,186 17.5 %

Mr. Equels is Executive Vice Chairman of our Board of Directors, Chief Executive Officer and President who (1) owns 1,123,307 shares of common stock and beneficially owns 2,853,107 shares issuable or issued upon exercise of:

Options	Plan	Date Issued	Exercise Price	Number Of Shares	Expiration Date
		2009 6/11/2010	\$ 7.92	25,000	6/11/2020
		2009 6/24/2011	\$ 4.92	25,000	6/24/2021
		2009 6/5/2012	\$ 3.48	8,333	6/6/2022
		2009 6/11/2012	\$ 3.72	25,000	6/11/2022
		2009 6/6/2013	\$ 3.72	25,000	6/6/2023
		2009 8/2/2013	\$ 3.00	12,500	8/2/2023
		2009 6/6/2014	\$ 4.32	25,000	6/6/2024
		2009 6/6/2015	\$ 3.00	25,000	6/6/2025
		2009 6/8/2016	\$ 1.68	25,000	6/6/2026
		2009 6/8/2017	\$ 0.56	300,000	6/8/2027
		2009 6/15/2017	\$ 0.49	14,212	6/15/2027
		2009 6/30/2017	\$ 0.48	14,214	6/30/2027
		2009 7/15/2017	\$ 0.49	18,124	7/15/2027
		2009 7/31/2017	\$ 0.42	20,786	7/31/2027
		2009 8/15/2017	\$ 0.41	21,336	8/15/2027
		2009 8/31/2017	\$ 0.36	24,463	8/31/2027
		2009 2/13/2018	\$ 0.37	371,622	2/13/2028
		2009 4/12/2018	\$ 0.38	125,000	4/12/2028
		2009 5/16/2018	\$ 0.30	300,000	5/16/2028
		2009 5/16/2018	\$ 0.30	250,000	5/16/2028
		2009 7/18/2018	\$ 0.31	161,290	7/18/2028
		2018 10/17/2018	\$ 0.22	284,091	10/17/2028
		2018 11/14/2028	\$ 0.22	1,000	11/14/2028
		2018 1/28/2019	\$ 0.22	426,136	1/28/2029
Total Options				2,528,107	

Warrants	Date	Exercise Price	Number Of Shares	Expiration Date
	3/4/2019	\$.20	325,000	3/4/2024

(2) Mr. Rodino is our General Counsel, Exec. Dir. of Governmental Relations and Secretary who owns 108,373 shares of common stock and beneficially owns 965,595 shares issuable or issued upon exercise of:

Options	Plan	Date Issued	Exercise Price	Number Of Shares	Expiration Date
	2009	8/2/2013	\$ 3.00	12,500	8/2/2023
	2009	6/21/2016	\$ 1.56	12,500	6/21/2026
	2009	6/15/2017	\$ 0.49	6,632	6/15/2027
	2009	6/30/2027	\$ 0.49	6,633	6/30/2027
	2009	7/15/2027	\$ 0.48	8,458	7/15/2027
	2009	7/31/2027	\$ 0.42	9,700	7/31/2027
	2009	8/15/2027	\$ 0.41	9,957	8/15/2027
	2009	8/31/2027	\$ 0.36	11,416	8/31/2027
	2009	2/13/2018	\$ 0.37	173,423	2/13/2028
	2009	4/12/2018	\$ 0.38	100,000	4/12/2028
	2009	5/16/2018	\$ 0.30	116,667	5/16/2028
	2009	7/18/2018	\$ 0.31	75,269	7/18/2028
	2018	10/17/2018	\$ 0.22	132,576	10/17/2028
	2018	11/14/2018	\$ 0.22	1,000	11/14/2028
	2018	1/28/2019	\$ 0.22	198,864	1/28/2029
Total Options				875,595	

Warrants	Date	Exercise Price	Number Of Shares	Expiration Date
	3/4/2019	\$.20	90,000	3/4/2024

(3) Dr. Mitchell is our Chairman of the Board who owns 211,197 shares of common stock and beneficially owns 1,522,714 shares issuable upon exercise of:

Options	Plan	Date Issued	Exercise Price	Number Of Shares	Expiration Date
	2009	6/5/2012	\$ 3.48	8,333	6/6/2022
	2009	8/2/2013	\$ 3.00	12,500	8/2/2023
	2009	9/9/2014	\$ 31.20	4,167	9/9/2024
	2009	9/9/2014	\$ 1.56	12,500	9/9/2024
	2009	4/30/2017	\$ 0.67	12,644	4/30/2027
	2009	5/15/2017	\$ 0.64	13,288	5/15/2027
	2009	5/31/2017	\$ 0.59	14,361	5/31/2027
	2009	6/15/2017	\$ 0.49	17,287	6/15/2027
	2009	6/30/2017	\$ 0.49	17,290	6/30/2027
	2009	7/15/2017	\$ 0.48	22,047	7/15/2027
	2009	7/31/2017	\$ 0.42	25,284	7/31/2027
	2009	8/15/2017	\$ 0.41	25,953	8/15/2027
	2009	8/31/2017	\$ 0.36	29,757	8/31/2027
	2009	2/13/2018	\$ 0.37	226,023	2/13/2028
	2009	4/12/2018	\$ 0.38	100,000	4/12/2028
	2009	5/16/2018	\$ 0.30	100,000	5/16/2028
	2009	5/16/2018	\$ 0.30	152,053	5/16/2028
	2009	7/18/2018	\$ 0.31	98,098	7/18/2028
	2018	10/17/2018	\$ 0.22	172,786	10/17/2028
	2018	11/14/2018	\$ 0.22	1,000	11/14/2028
	2018	1/28/2019	\$ 0.22	207,343	1/28/2029
	2018	1/28/2019	\$ 0.22	50,000	1/28/2029
Total Options				1,322,714	

Dr. Mitchell beneficially owns 16,639 shares of common stock of which 8,318 shares are held by Shirley Mitchell (Spouse), 4,098 shares are held by the Aesclepius Irrevocable Trust (Shirley Mitchell Trustee), and 4,223 shares are held by the Aesclepius Irrevocable Trust II (William Mitchell Trustee).

Warrants	Date	Exercise Price	Number Of Shares	Expiration Date
	3/4/2019	\$.20	200,000	3/4/2024

(4) Mr. Springate is our Senior Vice President of Operations and owns 117,273 shares of common stock and beneficially owns 673,753 shares issuable upon exercise of:

Edgar Filing: HEMISPHERX BIOPHARMA INC - Form 10-K

Options	Plan	Date	Exercise	Number	Expiration
		Issued	Price	Of Shares	Date
	2009	5/31/2011	\$ 6.60	7,500	5/31/2021
	2009	6/5/2012	\$ 3.48	4,167	6/5/2022
	2009	5/9/2013	\$ 2.88	4,167	5/9/2023
	2009	6/6/2014	\$ 4.32	4,167	6/6/2024
	2009	12/8/2014	\$ 22.80	151	12/8/2024
	2009	6/21/2016	\$ 1.56	12,500	6/21/2026
	2009	6/15/2017	\$ 0.49	4,264	6/15/2027
	2009	6/30/2017	\$ 0.49	4,264	6/30/2027
	2009	7/15/2017	\$ 0.48	5,438	7/15/2027
	2009	7/31/2017	\$ 0.42	6,236	7/31/2027
	2009	8/15/2027	\$ 0.41	6,401	8/15/2027
	2009	8/31/2017	\$ 0.36	7,339	8/31/2027
	2009	2/13/2018	\$ 0.37	111,486	2/13/2028
	2009	4/12/2018	\$ 0.38	75,000	4/12/2028
	2009	5/16/2018	\$ 0.30	79,167	5/16/2028
	2009	7/18/2018	\$ 0.31	53,763	7/18/2028
	2018	10/17/2018	\$ 0.22	94,697	10/17/2028
	2018	11/14/2018	\$ 0.22	1,000	11/14/2028
	2018	1/28/2029	\$ 0.22	142,046	1/28/2029
Total Options				623,753	

Warrants	Date	Exercise	Number	Expiration
		Price	Of Shares	Date
	3/4/2019	\$.20	50,000	3/4/2024

(5) Dr. Strayer is our Chief Scientific Officer and Chief Medical Director who has ownership of 94,304 shares of common stock and beneficially owns 447,931 shares issuable upon exercise of:

Options	Plan	Date Issued	Exercise Price	Number Of Shares	Expiration Date
	2009	4/13/2012	\$ 48.36	833	4/13/2022
	2009	12/8/2014	\$ 22.80	833	12/8/2024
	2009	6/21/2016	\$ 1.56	12,500	12/8/2024
	2009	4/12/2018	\$ 0.38	75,000	4/12/2028
	2018	10/17/2018	\$ 0.22	123,106	10/17/2028
	2018	11/14/2018	\$ 0.22	1,000	11/14/2028
	2018	1/28/2019	\$ 0.22	184,659	1/28/2029
Total Options				397,931	

Warrants	Date	Exercise Price	Number Of Shares	Expiration Date
	3/4/2019	\$.20	50,000	3/4/2024

(6) Mr. Pascale is our Chief Financial Officer who has ownership of 112,148 shares of common stock and beneficially owns 679,193 shares issuable upon exercise of:

Options	Plan	Date Issued	Exercise Price	Number Of Shares	Expiration Date
	2009	4/13/2012	\$ 48.36	500	4/13/2022
	2009	7/8/2014	\$ 3.96	4,167	7/8/2024
	2009	12/8/2014	\$ 22.80	598	12/7/2024
	2009	6/21/2016	\$ 1.56	12,500	6/21/2026
	2009	6/15/2017	\$ 0.49	4,738	6/15/2027
	2009	6/30/2027	\$ 0.49	4,738	6/30/2027
	2009	7/15/2027	\$ 0.48	6,042	7/15/2027
	2009	7/31/2017	\$ 0.42	6,929	7/31/2027
	2009	8/15/2017	\$ 0.41	7,112	8/15/2027
	2009	8/31/2027	\$ 0.36	8,155	8/31/2027
	2009	2/13/2018	\$ 0.37	123,874	2/13/2028
	2009	4/12/2018	\$ 0.38	75,000	4/12/2028
	2009	5/16/2018	\$ 0.30	83,334	5/16/2028
	2009	7/18/2018	\$ 0.31	53,763	7/18/2028
	2018	10/17/2018	\$ 0.22	94,697	10/17/2028
	2018	11/14/2018	\$ 0.22	1,000	11/14/2028
	2018	1/28/2019	\$ 0.22	142,046	1/28/2029

Total Options 629,193

Warrants	Date	Exercise Price	Number Of Shares	Expiration Date
	3/4/2019	\$.20	50,000	3/4/2024

(7) Mr. Appelrouth is a Director who owns 469,488 shares, and beneficially owns 1,485,164 shares issuable upon exercise of.

Options	Plan	Date Issued	Exercise Price	Number Of Shares	Expiration Date
	2009	4/30/2017	\$ 0.67	12,644	4/30/2027
	2009	5/15/2017	\$ 0.67	13,238	5/15/2027
	2009	5/31/2017	\$ 0.67	14,361	5/31/2027
	2009	6/15/2017	\$ 0.49	17,287	6/15/2027
	2009	6/30/2017	\$ 0.49	17,290	6/30/2027
	2009	7/15/2017	\$ 0.48	22,047	7/15/2027
	2009	7/31/2017	\$ 0.42	25,284	7/31/2027
	2009	8/15/2027	\$ 0.41	25,953	8/15/2027
	2009	8/31/2017	\$ 0.36	29,757	8/31/2027
	2009	2/13/2018	\$ 0.37	226,023	2/13/2028
	2009	4/18/2018	\$ 0.38	100,000	4/12/2028
	2009	5/16/2018	\$ 0.30	100,000	5/16/2028
	2009	5/16/2018	\$ 0.30	152,053	5/16/2028
	2009	7/18/2018	\$ 0.31	98,098	7/18/2028
	2018	10/17/2018	\$ 0.22	172,786	10/17/2028
	2018	11/14/2018	\$ 0.22	1,000	11/14/2028
	2018	1/28/2029	\$ 0.22	207,343	1/28/2029
	2018	1/28/2029	\$ 0.22	50,000	1/28/2029
Total Options				1,285,164	

Warrants	Date	Exercise Price	Number Of Shares	Expiration Date
	3/4/2019	\$.20	200,000	3/4/2024

(8) Includes shares of Common Stock that may be issued upon conversion of Preferred Stock issued in our March 2019 Rights Offering.

The following table gives information about our Common Stock that may be issued upon the exercise of options, warrants and rights under all of our equity compensation plans as of December 31, 2018:

Plan Category	Number of Securities to be issued upon exercise of outstanding options, warrants and rights	Weighted-average price of Outstanding options, warrants and rights	Number of securities Remaining available for future issuance under equity compensation plans (excluding securities reflected in column) (a)
	(a)	(b)	(c)
Equity compensation plans approved by security holders:	7,527,945	\$ 0.24	5,170,038
Equity compensation plans not approved by security holders:	14,335,298	\$ 0.49	-
Total	21,863,243	\$ 0.40	5,170,038

ITEM 13. Certain Relationships and Related Transactions, and Director Independence.

Review, Approval or Ratification of Transactions with Related Persons

Our policy is to require that any transaction with a related party required to be reported under applicable SEC rules, other than compensation related matters and waivers of our code of business conduct and ethics, be reviewed and approved or ratified by a majority of independent, disinterested Directors. We have adopted procedures in which the Audit Committee shall conduct an appropriate review of all related party transactions for potential conflict of interest situations on an annual and case-by-case basis with the approval of this Committee required for all such transactions.

We have employment agreements with certain of our executive officers and have granted such Officers and Directors options and warrants to purchase our common stock, as discussed under the headings, Item 11. “Executive Compensation”, and Item 12. “Security Ownership of Certain Beneficial Owners and Management”, as noted above.

ITEM 14. Principal Accountant Fees and Services.

All audit and professional services are approved in advance by the Audit Committee to assure such services do not impair the auditor's independence from us. The total fees by MBAF for 2018 and RSM US LLP ("RSM") for 2017 were \$301,250 and \$331,000 respectively. The following table shows the aggregate fees for professional services rendered during the year ended December 31, 2018 and 2017.

Description of Fees:	Amount (\$)	
	2018	2017
Audit Fees	\$260,000	\$278,000
Audit-Related Fees	41,250	53,000
Tax Fees	—	—
All Other Fees	—	—
Total	\$301,250	\$331,000

Audit Fees

Audit fees include the audit of our annual financial statements and the review of our financial statements included in our quarterly reports and services in connection with statutory and regulatory filings.

Audit-Related Fees

Represents the fees for assurance and related services that were reasonably related to the performance of the audit or review of our financial statements. Audit-related fees include professional services related to the Company's filing of SEC Form S-3 and S-8 (i.e., stock shelf offering procedures).

The Audit Committee has determined that RSM's rendering of these audit-related services and all other fees were compatible with maintaining auditor's independence. The Board of Directors considered RSM to be well qualified to serve as our independent public accountants. The Committee also pre-approved the charges for services performed in 2017 and 2016.

The Audit Committee pre-approves all auditing and accounting services and the terms thereof (which may include providing comfort letters in connection with securities underwriting) and non-audit services (other than non-audit services prohibited under Section 10A(g) of the Exchange Act or the applicable rules of the SEC or the Public Company Accounting Oversight Board) to be provided to us by the independent auditor; provided, however, the pre-approval requirement is waived with respect to the provisions of non-audit services for us if the "de minimus" provisions of Section 10A (i)(1)(B) of the Exchange Act are satisfied. This authority to pre-approve non-audit services

may be delegated to one or more members of the Audit Committee, who shall present all decisions to pre-approve an activity to the full Audit Committee at its first meeting following such decision.

PART IV

ITEM 15. Exhibits and Financial Statement Schedules.

Financial Statements and Schedules - See index to financial statements on page F-1 of this Annual Report. All other schedules called for under regulation S-X are not submitted because they are not applicable or not required, or because the required information is included in the financial statements or notes thereto.

(i) Exhibits - See exhibit index below.

Exhibit No.	Description
3.1	Amended and Restated Certificate of Incorporation of the Company, as amended, along with Certificates of Designations. (2)
3.2	<u>Amendment to Certificate of Incorporation. (3)</u>
3.3	<u>Amendment to Certificate of Incorporation. (4)</u>
3.4	<u>Amended and Restated By-Laws of Registrant. (33)</u>
3.5	<u>Certificate of Designation of Preference, Rights and Limitations of Series B Convertible Preferred Stock. (50)</u>
4.1	Specimen certificate representing our Common Stock. (2)
4.2	<u>Amended and Restated Rights Agreement, dated as of November 14, 2017, between the Company and American Stock Transfer & Trust Company LLC. The Amended and Restated Right Agreement includes the Form of Certificate of Designation, Preferences and Rights of the Series A Junior Participating Preferred Stock, the Form of Rights Certificate and the Summary of the Right to Purchase Preferred Stock. (5)</u>
4.3	<u>Form of Indenture filed with Form S-3 Universal Shelf Registration Statement. (6)</u>
4.4	<u>Form of Warrant pursuant to August 30, 2016 Securities Purchase Agreement. (36)</u>
4.5	<u>Form of Warrant pursuant to February 1, 2017 Securities Purchase Agreement. (38)</u>
4.6	<u>Form of Series A Warrant-June 2017. (41)</u>
4.7	<u>Form of Series B Warrant-June 2017. (41)</u>

- 4.8 Form of New Series A Warrant-August 2017. (40)
- 4.9 Form of New Series B Warrant-August 2017. (40)
- 4.10 Form of Warrant issued to Purchaser of facility. (21)
- 4.11 Form of Series A Warrant- April 2018. (44)
- 4.12 Form of Series B Warrant- April 2018. (44)
- 4.13 September 28, 2018 Secured Convertible Promissory Note from the Company to Iliad Research and Trading, L.P. (48)
- 4.14 Form of Non-Transferable Subscription Rights Certificate. (50)
- 4.15 Form of Warrant Agreement. (50)
- 4.16 Form of Warrant Certificate. (49)

50

- 4.17 March 2019 Amendment to September 28, 2018 Secured Convertible Promissory Note from the Company to Iliad Research and Trading, L.P. (1)
- 10.1 Form of Confidentiality, Invention and Non-Compete Agreement. (2)
- 10.2 Form of Clinical Research Agreement. (2)
- 10.3 Employee Wage or Hours Reduction Program. (7)
- 10.4 Supply Agreement with Hollister-Stier Laboratories LLC dated December 5, 2005. (9)
- 10.5 Amendment to Supply Agreement with Hollister-Stier Laboratories LLC dated February 25, 2010. (10)
- 10.6 Vendor Agreement with Bio Ridge Pharma, LLC dated August 11, 2011. (31).
- 10.7 Vendor Agreement with Armada Healthcare, LLC dated August 11, 2011. (31).
- 10.8 Amended and restated employment agreement with Wayne Springate dated May 1, 2011. (12)
- 10.9 Amended and restated employment agreement with Thomas K. Equels dated December 6, 2011. (15)
- 10.10 Amendment to Supply Agreement with Hollister-Stier Laboratories LLC executed September 9, 2011. (16)
- 10.11 Vendor Agreement extension with Bio Ridge Pharma, LLC dated August 14, 2012. (19)
- 10.12 Vendor Agreement extension with Armada Healthcare, LLC dated August 14, 2012. (19)
- 10.13 Vendor Agreement extension with Armada Healthcare, LLC dated July 19, 2013. (20)
- 10.14 Vendor Agreement extension with Bio Ridge Pharma, LLC dated July 19, 2013. (20)
- 10.15 Vendor Agreement extension with Bio Ridge Pharma, LLC and Armada Healthcare, LLC dated August 8, 2014.(21)
- 10.16 Sales, Marketing, Distribution, and Supply Agreement with Emerge Health Pty Ltd. dated March 9, 2015.(Confidential Treatment granted with respect to portions of the Agreement) (21)
- 10.17 August 4, 2015 Amendment to Equity Distribution Agreement between the registrant and Maxim Group LLC. (22)
- 10.18 Vendor Agreement extension with Armada Healthcare, LLC dated July 29, 2015. (25)
- 10.19 Vendor Agreement extension with Bio Ridge Pharma, LLC dated July 29, 2013. (24)
- 10.20 Early Access Agreement with Impatiens N.V. dated August 3, 2015.(Confidential Treatment granted with respect to portions of the Agreement) (24)
- 10.21

Sales, Marketing, Distribution, and Supply Agreement with Emerge Health Pty Ltd. dated August 6, 2015. (Confidential Treatment granted with respect to portions of the Agreement) (24)

10.22 Addendum to Early Access Agreement with Impatiens N.V. dated October 16, 2015.(Confidential Treatment granted with respect to portions of the Agreement) (25)

10.23 Letter agreement between Dr. Carter and the Company dated September 28, 2015 extending the period for notice of non-renewal to December 1, 2015 within the June 11, 2010 Amended and Restated Engagement Agreement entered into between the Company and Dr. Carter. (25)

10.24 November 23, 2015 William A. Carter Employment Agreement Waiver. (26)

10.25 November 23, 2015 Thomas K. Equels Employment Agreement Waiver. (26)

10.26 Equity Distribution Agreement, dated December 15, 2015 with Chardan Capital Markets, LLC. (27)

51

- 10.27 December 23, 2015 letter to Dr. Carter related to non-renewal of his consulting agreement and continued consulting services. (28)
- 10.28 2016 Senior Executive Deferred Cash Performance Award Plan. (29)
- 10.29 2016 Voluntary Incentive Stock Award Plan. (29)
- 10.30 Amended and Restated 2016 Senior Executive Deferred Cash Performance Award Plan. (30)
- 10.31 Sales, Marketing, Distribution and Supply Agreement (the “Agreement”) with Scientific Products Pharmaceutical Co. LTD dated March 3, 2016 (Confidential Treatment granted with respect to portions of the Agreement). (32)
- 10.32 Agreement between Avrio Biopharmaceuticals (“Avrio”) and the Company dated July 20, 2016 (Confidential Treatment granted with respect to portions of the Agreement). (34)
- 10.33 Licensing Agreement dated April 13, 2016 with Lonza Sales AG (Confidential Treatment granted with respect to portions of the Agreement). (35)
- 10.34 Form of Securities Purchase Agreement entered into on August 30, 2016. (36)
- 10.35 Amended and Restated Early Access Agreement with Impatiens N.V. dated May 20, 2016. (Confidential Treatment granted with respect to portions of the Agreement) (37)
- 10.36 December 13, 2016 Amendment No. 1 to Amended and Restated Early Access Agreement with Impatiens N.V. (21)
- 10.37 June 28, 2017 Amendment No. 2 to Amended and Restated Early Access Agreement with Impatiens N.V. (21)
- 10.38 February 14, 2018 Amendment No. 3 to Amended and Restated Early Access Agreement with Impatiens N.V. (21)
- 10.39 March 26, 2018 Amendment No. 4 to Amended and Restated Early Access Agreement with Impatiens N.V. (21)
- 10.40 Form of Securities Purchase Agreement entered into on February 1, 2017. (39)
- 10.41 August 2017 Form of Employee Pay Reduction Plan. (39)
- 10.42 August 2017 Form of Executive Compensation Deferral Plan. (39)
- 10.43 August 2017 Form of Directors’ Compensation Deferral Plan. (39)
- 10.44 Form of August 2017 Agreement between the Company and the Warrantholders. (40)
- 10.45 Form of June 2017 Agreement between the Company and the Warrantholders. (41)
- 10.46 Mortgage and Security Agreement with SW Partners LLC dated May 12, 2017. (42)

- 10.47 Promissory Note with SW Partners LLC dated May 12, 2017. (42)
- 10.48 September 11, 2017 Purchase and Sale Agreement- 5 Jules Lane. (21)
- 10.49 January 8, 2018 Purchase and Sale Agreement- 783 Jersey Lane. (21)
- 10.50 Lease Agreement for 783 Jersey Lane. (21)
- 10.51 Form of Stock Purchase Agreement entered into on March 21, 2018. (43)
- 10.52 Form of Securities Purchase Agreement entered into on May 24, 2018. (44)

- 10.53 October 9, 2018, Clinical Trial Agreement with Roswell Park Comprehensive Cancer Center. (46)
- 10.54 October 8, 2018, Restated First Amendment to Purchase and Sale Agreement. (46)
- 10.55 October 9, 2018, Restated Bill of Sale for the Restated First Amendment and Sale Agreement. (46)
- 16.1 April 5, 2018 Letter from RSM US LLP. (45)
- 21.1 List of Subsidiaries. (47)
- 23.1* Consent of RSM US LLP.
- 23.2* Consent of Morrison, Brown, Argiz & Farra, LLC.
- 31.1 Certification pursuant to Section 302 of the Sarbanes-Oxley Act of 2002 from the Company's Chief Executive Officer. *
- 31.2 Certification pursuant to Section 302 of the Sarbanes-Oxley Act of 2002 from the Company's Chief Financial Officer. *
- 32.1 Certification pursuant to Section 906 of the Sarbanes-Oxley Act of 2002 from the Company's Chief Executive Officer. *
- 32.2 Certification pursuant to Section 906 of the Sarbanes-Oxley Act of 2002 from the Company's Chief Financial Officer. *

101 The following materials from Hemispherx' Annual Report on Form 10-K for the year ended December 31, 2018, formatted in eXtensible Business Reporting Language ("XBRL"): (i) the Condensed Consolidated Statements of Income; (ii) the Condensed Consolidated Balance Sheets; (iii) the Condensed Consolidated Statements of Cash Flows; and (iv) Notes to Condensed Consolidated Financial Statements.

*Filed herewith.

- (1) Filed with the Securities and Exchange Commission as an exhibit to the Company's Current Report on Form 8-K filed March 15, 2019 and is hereby incorporated by reference.
- (2) Filed with the Securities and Exchange Commission as an exhibit to the Company's Registration Statement on Form S-1 (No. 33-93314) filed November 2, 1995 and is hereby incorporated by reference.
- (3) Filed with the Securities and Exchange Commission as Appendix A to the Company's Definitive Proxy Statement on Schedule 14A filed on September 16, 2011 and is hereby incorporated by reference.
- (4) Filed with the Securities and Exchange Commission as Appendix A to the Company's Definitive Proxy Statement on Schedule 14A filed on June 27, 2016 and is hereby incorporated by reference.

- (5) Filed with the Securities and Exchange Commission on November 14, 2017 as an exhibit to the Company's Registration Statement on Form 8-A12B (No. 0-27072) and is hereby incorporated by reference.
- (6) Filed with the Securities and Exchange Commission as an exhibit to the Company's Form S-3 Registration Statement (No. 333-205228) on June 25, 2015 and is hereby incorporated by reference.
- (7) Filed with the Securities and Exchange Commission as an exhibit to the Company's annual report on Form 10-K (No. 000-27072) for the year ended December 31, 2008 and is hereby incorporated by reference.
- (8) Filed with the Securities and Exchange Commission as an exhibit to the Company's quarterly report on Form 10-Q (No. 000-27072) for the period ended June 30, 2010 and is hereby incorporated by reference.
- (9) Filed with the Securities and Exchange Commission as an exhibit to the Company's annual report on Form 10-K (No. 000-27072) for the year ended December 31, 2005 and is hereby incorporated by reference.

- (10) Filed with the Securities and Exchange Commission as an exhibit to the Company's Annual Report on Form 10-K (No. 000-27072) for the year ended December 31, 2009 and is hereby incorporated by reference.
- (11) Filed with the Securities and Exchange Commission as an exhibit to the Company's Current Report on Form 8-K (No. 000-27072) dated May 28, 2010 and is hereby incorporated by reference.
- (12) Filed with the Securities and Exchange Commission as an exhibit to the Company's quarterly report on Form 10-Q (No. 000-27072) for the period ended March 31, 2011 and is hereby incorporated by reference.
- (13) Filed with the Securities and Exchange Commission as an exhibit to the Company's quarterly report on Form 10-Q (No. 000-27072) for the period ended September 30, 2011 and is hereby incorporated by reference.
- (14) Filed with the Securities and Exchange Commission as an exhibit to the Company's Current Report on Form 8-K (No. 000-27072) filed September 23, 2011 and is hereby incorporated by reference.
- (15) Filed with the Securities and Exchange Commission as an exhibit to the Company's Current Report on Form 8-K (No. 000-27072) filed December 12, 2011 and is hereby incorporated by reference.
- (16) Filed with the Securities and Exchange Commission as an exhibit to the Company's Annual Report on Form 10-K (No. 000-27072) for the year ended December 31, 2011 and is hereby incorporated by reference.
- (17) Filed with the Securities and Exchange Commission as an exhibit to the Company's Current Report on Form 8-K (No. 000-27072) filed August 15, 2012 and is hereby incorporated by reference.
- (18) Filed with the Securities and Exchange Commission as an exhibit to the Company's quarterly report on Form 10-Q (No. 000-27072) for the period ended September 30, 2015 and is hereby incorporated by reference.
- (19) Filed with the Securities and Exchange Commission as an exhibit to the Company's annual report on Form 10-K (No. 000-27072) for the year ended December 31, 2013 and is hereby incorporated by reference.
- (20) Filed with the Securities and Exchange Commission as an exhibit to the Company's annual report on Form 10-K (No. 000-27072) for the year ended December 31, 2014 and is hereby incorporated by reference.
- (21) Filed with the Securities and Exchange Commission as an exhibit to the Company's annual report on Form 10-K (No. 000-27072) for the year ended December 31, 2017 and is hereby incorporated by reference left blank.
- (22) Filed with the Securities and Exchange Commission as an exhibit to the Company's Current Report on Form 8-K (No. 000-27072) filed June 23, 2015 and is hereby incorporated by reference.
- (23) Filed with the Securities and Exchange Commission as an exhibit to the Company's Current Report on Form 8-K (No. 000-27072) filed August 4, 2015 and is hereby incorporated by reference.
- (24) Filed with the Securities and Exchange Commission as an exhibit to the Company's quarterly report on Form 10-Q (No. 000-27072) for the period ended June 30, 2015 and is hereby incorporated by reference.
- (25) Filed with the Securities and Exchange Commission as an exhibit to the Company's quarterly report on Form 10-Q (No. 1-13441) for the period ended September 30, 2015 and is hereby incorporated by reference.

Edgar Filing: HEMISPHERX BIOPHARMA INC - Form 10-K

- (26) Filed with the Securities and Exchange Commission as an exhibit to the Company's Current Report on Form 8-K (No. 000-27072) filed November 23, 2015 and is hereby incorporated by reference.
- (27) Filed with the Securities and Exchange Commission as an exhibit to the Company's Current Report on Form 8-K (No. 000-27072) filed December 15, 2015 and is hereby incorporated by reference.
- (28) Filed with the Securities and Exchange Commission as an exhibit to the Company's Current Report on Form 8-K (No. 000-27072) filed January 14, 2016 and is hereby incorporated by reference.

- (29) Filed with the Securities and Exchange Commission as an exhibit to the Company's Current Report on Form 8-K (No. 000-27072) filed February 4, 2016 and is hereby incorporated by reference.
- (30) Filed with the Securities and Exchange Commission as an exhibit to the Company's Current Report on Form 8-K (No. 000-27072) filed March 1, 2016 and is hereby incorporated by reference.
- (31) Filed with the Securities and Exchange Commission as an exhibit to the Company's amended quarterly report on Form 10-Q/A (No. 000-27072) for the period ended September 30, 2011 and is hereby incorporated by reference.
- (32) Filed with the Securities and Exchange Commission as an exhibit to the Company's quarterly report on Form 10-Q for the period ended March 31, 2016 and is hereby incorporated by reference.
- (33) Filed with the Securities and Exchange Commission as an exhibit to the Company's Current Report on Form 8-K filed June 10, 2016 and is hereby incorporated by reference.
- (34) Filed with the Securities and Exchange Commission as an exhibit to the Company's quarterly report on Form 10-Q for the period ended June 30, 2016 and is hereby incorporated by reference.
- (35) Filed with the Securities and Exchange Commission as an exhibit to the Company's quarterly report on Form 10-Q/A for the period ended March 31, 2016 and is hereby incorporated by reference.
- (36) Filed with the Securities and Exchange Commission as an exhibit to the Company's Current Report on Form 8-K filed September 1, 2016 and is hereby incorporated by reference.
- (37) Filed with the Securities and Exchange Commission as an exhibit to the Company's Current Report on Form 8-K/A filed May 8, 2017 and is hereby incorporated by reference.
- (38) Filed with the Securities and Exchange Commission as an exhibit to the Company's Current Report on Form 8-K filed February 3, 2017 and is hereby incorporated by reference.
- (39) Filed with the Securities and Exchange Commission as an exhibit to the Company's Current Report on Form 8-K filed August 29, 2017 and is hereby incorporated by reference.
- (40) Filed with the Securities and Exchange Commission as an exhibit to the Company's Current Report on Form 8-K filed August 23, 2017 and is hereby incorporated by reference.
- (41) Filed with the Securities and Exchange Commission as an exhibit to the Company's Current Report on Form 8-K filed June 1, 2017 and is hereby incorporated by reference.
- (42) Filed with the Securities and Exchange Commission as an exhibit to the Company's quarterly report on Form 10-Q (No. 000-27072) for the period ended March 31, 2017 and is hereby incorporated by reference.
- (43) Filed with the Securities and Exchange Commission as an exhibit to the Company's Current Report on Form 8-K filed March 22, 2018 and is hereby incorporated by reference.
- (44) Filed with the Securities and Exchange Commission as an exhibit to the Company's Current Report on Form 8-K filed May 20, 2018 and is hereby incorporated by reference.

- (45) Filed with the Securities and Exchange Commission as an exhibit to the Company's Current Report on Form 8-K filed May 6, 2018 and is hereby incorporated by reference.
- (46) Filed with the Securities and Exchange Commission as an exhibit to the Company's quarterly report on Form 10-Q (No. 000-27072) for the period ended September 30, 2018 and is hereby incorporated by reference.
- (47) Filed with the Securities and Exchange Commission as an exhibit to the Company's Registration Statement on Form S-1 (No. 333-226057) filed July 2, 2018 and is hereby incorporated by reference.
- (48) Filed with the Securities and Exchange Commission as an exhibit to the Company's Current Report on Form 8-K filed October 4, 2018 and is hereby incorporated by reference.
- (49) Filed with the Securities and Exchange Commission as an exhibit to the Company's Current Report on Form 8-K filed March 8, 2019 and is hereby incorporated by reference.
- (50) Filed with the Securities and Exchange Commission as an exhibit to the Company's Registration Statement on Form S-1/A (No. 333-229051) filed February 6, 2019 and is hereby incorporated by reference.

(b) Financial Statement Schedules

All schedules have been omitted because either they are not required, are not applicable or the information is otherwise set forth in the financial statements and related notes thereto.

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.

HEMISPHERX
BIOPHARMA, INC.

By: */s/ Thomas K. Equels*
Thomas K. Equels
Chief Executive Officer

April 1, 2019

Pursuant to the requirements of Section 13 or 15(d) of the Securities Exchange of 1934, as amended, this report has been signed below by the following persons on behalf of this Registrant and in the capacities and on the dates indicated.

<i>/s/ Thomas K. Equels</i> Thomas K. Equels	Chief Executive Officer & President, Director of the Board	April 1, 2019
<i>/s/ William Mitchell</i> William Mitchell, M.D., Ph.D.	Chairman of the Board and Director	April 1, 2019
<i>/s/ Stewart L. Appelrouth</i> Stewart L. Appelrouth	Director	April 1, 2019
<i>/s/ Adam Pascale</i> Adam Pascale	Chief Financial Officer	April 1, 2019

HEMISPHERx BIOPHARMA, INC AND SUBSIDIARIES

Index to Consolidated Financial Statements

	Page
<u>Report of Independent Registered Public Accounting Firm</u>	F-2
<u>Report of Independent Registered Public Accounting Firm</u>	F-3
<u>Consolidated Balance Sheets at December 31, 2018 and 2017</u>	F-4
<u>Consolidated Statements of Comprehensive Loss for each of the years in the two-year period ended December 31, 2018</u>	F-5
<u>Consolidated Statements of Changes in Stockholders' Equity for each of the years in the two-year period ended December 31, 2018</u>	F-6
<u>Consolidated Statements of Cash Flows for each of the years in the two-year period ended December 31, 2018</u>	F-7
<u>Notes to Consolidated Financial Statements</u>	F-8

F-1

REPORT OF INDEPENDENT REGISTERED PUBLIC ACCOUNTING FIRM

To the Shareholders and the Board of Directors of Hemispherx Biopharma, Inc.

Opinion on the Financial Statements

We have audited the accompanying balance sheet of Hemispherx Biopharma, Inc. (the “Company”) as of December 31, 2018, the related statements of comprehensive loss, stockholders’ equity and cash flows, for the year ended December 31, 2018, and the related notes (collectively referred to as the “financial statements”). In our opinion, the financial statements present fairly, in all material respects, the financial position of the Company as of December 31, 2018 and the results of its operations and its cash flows for the year ended December 31, 2018, in conformity with U.S. generally accepted accounting principles.

Basis for Opinion

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 audit. We are a public accounting firm registered with the Public Company Accounting Oversight Board (United States) (“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 the financial statements are free of material misstatement, whether due to error or fraud. Our audit 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 audit 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 audit provides a reasonable basis for our opinion.

/s/ Morrison, Brown, Argiz & Farra, LLC

We have served as the Company's auditor since 2018.

Miami, Florida

April 1, 2019

F-2

Report of Independent Registered Public Accounting Firm

To the Stockholders and the Board of Directors of Hemispherx Biopharma, Inc.

Opinion on the Financial Statements

We have audited the accompanying consolidated balance sheet of Hemispherx Biopharma, Inc. and its subsidiaries (the Company) as of December 31, 2017, the related consolidated statements of comprehensive loss, changes in stockholders' equity and cash flows for the year then ended, and the related notes to the consolidated financial statements (collectively, the financial statements). In our opinion, the financial statements present fairly, in all material respects, the financial position of the Company as of December 31, 2017, and the results of its operations and its cash flows for the year ended December 31, 2017, in conformity with accounting principles generally accepted in the United States of America.

Basis for Opinion

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 audit. We are a public accounting firm registered with the Public Company Accounting Oversight Board (United States) (PCAOB) and are required to be independent with respect to the Company in accordance with 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 the financial statements are free of material misstatement, whether due to error or fraud. The Company is not required to have, nor were we engaged to perform, an audit of its internal control over financial reporting. As part of our audit we are required to obtain an understanding of internal control over financial reporting but not for the purpose of expressing an opinion on the effectiveness of the Company's internal control over financial reporting. Accordingly, we express no such opinion.

Our audit 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 audit 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 audit provides a reasonable basis

for our opinion.

/s/ RSM US LLP

We have served as the Company's auditor since 2006.

Blue Bell, Pennsylvania

March 30, 2018

F-3

HEMISPHERx BIOPHARMA, INC. AND SUBSIDIARIES**Consolidated Balance Sheets****December 31, 2018 and 2017**

(in thousands, except for share and per share amounts)

	2018	2017
ASSETS		
Current assets:		
Cash and cash equivalents	\$299	\$1,412
Marketable securities- unrestricted	1,526	695
Funds receivable from sale of New Jersey net operating loss	859	—
Accounts receivable	235	24
Assets held for sale	—	764
Prepaid expenses and other current assets	880	610
Total current assets	3,799	3,505
Property and equipment, net	7,782	8,586
Patent and trademark rights, net	912	858
Other assets	1,352	1,258
Total assets	\$13,845	\$14,207
LIABILITIES AND STOCKHOLDERS' EQUITY		
Current liabilities:		
Accounts payable	\$680	\$741
Accrued expenses	1,005	1,966
Convertible note payable	3,408	—
Current portion of financing obligation	199	—
Total current liabilities	5,292	2,707
Long-term liabilities:		
Note payable	—	1,835
Financing obligation arising from sale leaseback transaction (Note 19)	2,318	—
Redeemable warrants	1,061	962
Commitments and contingencies (Notes 9,11,12,14,15 and 19)		
Stockholders' equity:		
Preferred stock, par value \$0.01 per share, authorized 5,000,000; issued and outstanding; none	—	—
Common stock, par value \$0.001 per share, authorized 350,000,000 shares; issued and outstanding 48,734,712 and 32,884,786, respectively	49	33
Additional paid-in capital	323,701	317,419
Other comprehensive income (loss)	(3)	11
Accumulated deficit	(318,573)	(308,760)
Total stockholders' equity	5,174	8,703

Total liabilities and stockholders' equity	\$13,845	\$14,207
--	----------	----------

See accompanying notes to consolidated financial statements.

F-4

HEMISPHERx BIOPHARMA, INC. AND SUBSIDIARIES**Consolidated Statements of Comprehensive Loss**

(in thousands, except share and per share data)

	Years ended December 31,	
	2018	2017
Revenues:		
Clinical treatment programs - US	\$248	\$102
Clinical treatment programs - Europe	119	335
Total Revenues	367	437
Costs and Expenses:		
Production costs	884	1,183
Research and development	4,778	4,098
General and administrative	6,201	6,572
Total Costs and Expenses	11,863	11,853
Operating loss	(11,496)	(11,416)
Interest and other income (expense)	46	88
Interest expense and finance costs	(502)	(139)
Settlement of litigation	474	-
Fair value of convertible note adjustment	(582)	-
Redeemable warrants valuation adjustment	1,165	2,417
Gain from sale of income tax operating losses	859	791
Gain on sale of building	223	-
Net loss	(9,813)	(8,259)
Other Comprehensive Income (Loss)		
Unrealized gain (loss) on securities	(23)	33
Reclassification adjustments for realized gain (loss) on sales of short-term marketable securities and for impairment losses on investments included in net loss	9	(17)
Net comprehensive loss	\$(9,827)	\$(8,243)
Basic and diluted loss per share	\$(0.22)	\$(0.29)
Weighted average shares outstanding basic and diluted	44,189,217	28,676,076

See accompanying notes to consolidated financial statements.

HEMISPHERx BIOPHARMA, INC. AND SUBSIDIARIES**Consolidated Statements of Changes in Stockholders' Equity**

(in thousands except share data)

	Common Stock Shares	Common Stock .001 Par Value	Additional Paid-in Capital	Other Comprehensive Income (Loss)	Accumulated Deficit	Total Stockholders' Equity	
Balance December 31, 2016	24,202,921	\$ 24	\$ 315,980	\$ (5) \$ (300,501) \$ 15,498	
Shares issued for:							
Shares sold at the market	678,275	1	236	—	—	237	
Common stock issuance	4,646,205	5	2,175	—	—	2,180	
Other issuance	2,241,979	2	896	—	—	898	
Equity-based compensation	1,115,406	1	570	—	—	571	
Redeemable warrants	—	—	(2,050)	—	(2,050)
Deemed dividends	—	—	(388)	—	(388)
Net comprehensive loss	—	—	—	16	(8,259) (8,243)
Balance December 31, 2017	32,884,786	33	317,419	11	(308,760) 8,703	
Shares issued for:							
Shares sold at the market	2,176,392	2	801	—	—	803	
Common stock issuance, net of costs	11,302,372	11	2,771	—	—	2,782	
Convertible note origination shares	500,000	1	83	—	—	84	
Other issuance	831,005	1	329	—	—	330	
Equity-based compensation	1,040,157	1	928	—	—	929	
Redeemable warrants	—	—	221	—	—	221	
Warrants issued for building sales leaseback	—	—	1,149	—	—	1,149	
Net comprehensive loss	—	—	—	(14) (9,813) (9,827)
Balance December 31, 2018	48,734,712	\$ 49	\$ 323,701	\$ (3) \$ (318,573) \$ 5,174	

See accompanying notes to consolidated financial statements

HEMISPHERx BIOPHARMA, INC. AND SUBSIDIARIES**Consolidated Statements of Cash Flows****(in thousands)**

	Years ended December 31, 2018	2017
Cash flows from operating activities:		
Net loss	\$ (9,813)	\$ (8,259)
Adjustments to reconcile net loss to net cash used in operating activities:		
Depreciation of property and equipment	856	948
Fair value of convertible note adjustment	582	—
Amortization of debt issue costs	263	37
Amortization and abandonment of patent and trademark rights	22	63
Redeemable warrants valuation adjustment	(1,165)	(2,417)
Equity-based compensation	929	571
Realized gain (loss) on securities	(14)	(17)
Gain on sale of building	(223)	—
Changes in assets and liabilities:		
Prepaid expenses and other assets	(251)	(13)
Accounts receivable	(1,070)	(24)
Accounts payable	166	566
Accrued expenses	(922)	604
Net cash used in operating activities	(10,640)	(7,941)
Cash flows from investing activities:		

Edgar Filing: HEMISPHERX BIOPHARMA INC - Form 10-K

Purchases of property and equipment	(51)	(20)
Additions to patent and trademark rights	(76)	(49)
Proceeds from sale of building	1,050		—	
Sales and maturities of short-term marketable securities	(831)	2,799	
Net cash provided by investing activities	92		\$ 2,730	
Cash flows from financing activities:				
Proceeds from sale of common stock, net of issuance costs	5,070		2,417	
Proceeds for lease financing obligation	4,080		—	
Financing obligation payment	(264)	—	
Debt issuance costs	(439)	(102)
Proceeds from note payable	3,059		1,900	
Payoff of mortgage note payable	(1,957)	—	
Security deposits paid	(114)	—	
Net cash provided by financing activities	9,435		4,215	
Net decrease in cash and cash equivalents	(1,113)	(996)
Cash and cash equivalents at beginning of year	1,412		2,408	
Cash and cash equivalents at end of year	\$ 299		\$ 1,412	
Supplemental disclosures of non-cash investing and financing cash flow information:				
Issuance of common stock for accounts payable and accrued expenses	\$ 330		\$ 898	
Unrealized (loss) gain on marketable securities	\$ (14)	\$ 16	
Fair value of redeemable warrants granted	\$ 1,265		\$ 2,050	

Supplemental
disclosure of cash
flow information:

Cash paid for interest expense	\$	328	\$	101
--------------------------------	----	-----	----	-----

See accompanying notes to consolidated financial statements.

F-7

HEMISPHERx BIOPHARMA, INC. AND SUBSIDIARIES

NOTES TO CONSOLIDATED FINANCIAL STATEMENTS

(1) Business

Hemispherx Biopharma, Inc. and its subsidiaries (collectively, “Hemispherx”, “Company”, “we” or “us”) are an immuno-pharma company headquartered in Ocala, Florida, focused on the research and development of therapeutics to treat multiple types of cancers, as well as immune-deficiency disorders. We have established a strong foundation of laboratory, pre-clinical and clinical data with respect to the development of nucleic acids and natural interferon to enhance the natural antiviral defense system of the human body and to aid the development of therapeutic products for the treatment of certain cancers and chronic diseases.

Hemispherx’s flagship products include Ampligen® (Rintatolimod), a first-in-class drug of large macromolecular RNA (ribonucleic acid) molecules, and Alferon N Injection® (Interferon Alfa-N3).

Ampligen® represents an RNA being developed for globally important cancers, viral diseases and disorders of the immune system. Ampligen® has in the clinic demonstrated the potential for standalone efficacy in a number of solid tumors. We have also seen success in increasing survival rates and efficacy in the treatment of animal tumors when Ampligen® is used in combination with checkpoint blockade therapies. This success in the field of immuno-oncology has guided the company’s focus toward the potential use of Ampligen® as a combinational therapy for the treatment of a variety of solid tumor types. There are currently multiple Ampligen® clinical trials — both underway and planned — at major cancer research centers around the country.

Ampligen® is also being evaluated for the treatment of myalgic encephalomyelitis/chronic fatigue syndrome (ME/CFS). With regulatory approval in Argentina, Ampligen® is the world’s only approved therapeutic for ME/CFS. Hemispherx is sponsoring multiple expanded access programs (EAP) for ME/CFS patients worldwide. In August 2016, we received approval of our NDA from Administracion Nacional de Medicamentos, Alimentos y Tecnologia Medica (ANMAT) for commercial sale of Ampligen® in the Argentine Republic for the treatment of severe CFS.

Alferon N Injection® is approved for a category of STD infection and patients that are intolerant to recombinant interferon in Argentina. Alferon is the only natural-source, multi-species alpha interferon currently approved for sale in the U.S. for the intralesional treatment of refractory (resistant to other treatment) or recurring external Condylomata Acuminata/genital warts in patients 18 years of age or older. Certain types of human papilloma viruses cause genital warts. Hemispherx also has approval from the ANMAT for the treatment of refractory patients that failed or were intolerant to treatment with recombinant interferon in Argentina. The company has developed and will be seeking FDA Pre-Approval Inspection of a high-volume, high-efficiency, upgraded manufacturing process to allow for the

commercial viability of Alferon®.

The Company operates a 30,000 sq. ft. facility in New Brunswick, NJ with the objective of producing Ampligen® and Alferon®. It is committed to a focused business plan oriented toward finding senior co-development partners with the capital and expertise needed to commercialize the many potential therapeutic aspects of Ampligen® and our FDA-approved drug Alferon® .

The Company has incurred numerous years of substantial operating losses as it pursued its clinical and pre-clinical development activities and appropriate regulatory approval processes before any such products can be sold and marketed. As of December 31, 2018, its accumulated deficit was approximately \$318,573,000. The Company has not yet generated significant revenues from our products and may incur substantial losses in the future. The Company evaluated these conditions and events that may raise substantial doubt about the Company's ability to continue as a going concern; however, the Company believes that it has alleviated the substantial doubt by implementing certain actions. The Company reexamined its fundamental priorities in terms of direction, corporate culture and its ability to fund operations. As a result, there were significant changes at the Company including the Company restructuring its executive management team, initiating the pursuit of international sales of clinical grade materials, and implementing a cost saving program which assisted the Company in gained efficiencies and eliminated redundancies within its workforce.

In 2018, the Company sold its property located at 783 Jersey Lane, New Brunswick, NJ. This property houses its development and production facilities. The purchase price was \$4,080,000 and purchaser received 3,225,806 warrants to purchase common stock. Simultaneously with the closing of the sale, the purchaser leased the facility back to the Company. The lease runs for 10 years, with two five year extensions. The initial annual base rent is \$408,000 and will continue for the first and second year. In the third and fourth it will escalate at the rate of 2.5% per year. For all subsequent years it will escalate at the rate of 3% per year. The Company also will be responsible for additional rent consisting of taxes and certain insurance expenses of the purchaser. The lease contains a repurchase option pursuant to which the Company can repurchase the facility within the initial 10 year lease period. The purchase price would \$4,080,000 times a multiple. The multiple would be 1.05 plus .0025N where N represents the number of months between lease commencement and closing of repurchase. The Company sold the building located adjacent to its manufacturing facility located at 5 Jules Lane, New Brunswick, New Jersey to an unaffiliated party. The purchase price was \$1,050,000 and the Company netted \$963,254 in cash.

The consolidated financial statements include the financial statements of Hemispherx Biopharma, Inc. and its wholly-owned subsidiaries, which are incorporated in Delaware and are dormant. The Company's foreign subsidiary, Hemispherx Biopharma Europe N.V./S.A., was established in Belgium in 1998. All significant intercompany balances and transactions have been eliminated in consolidation.

(2) Summary of Significant Accounting Policies**(a) Cash and Cash Equivalents**

Cash and Cash Equivalents consist of cash and money market accounts and total \$299,000 and \$1,412,000 at December 31, 2018 and 2017, respectively.

(b) Marketable Securities

The Company's securities are classified as available for sale and are stated at fair value. Unrealized gains and losses on securities available for sale are excluded from results of operations and are reported as other comprehensive income (loss) on the Statements of Comprehensive Loss, net of taxes. Securities classified as available for sale include securities that may be sold in response to changes in interest rates, changes in prepayment risks or for portfolio management purposes. The cost of securities sold is determined on a specific identification basis. Gains and losses on sales of securities are recognized in the statements of comprehensive loss on the date of sale.

(c) Property and Equipment

	(in thousands)	
	December 31,	
	2018	2017
Land, buildings and improvements	\$10,547	\$10,547
Furniture, fixtures, and equipment	5,045	5,625
Total property and equipment	15,592	16,172
Less: accumulated depreciation and amortization	(7,810)	(7,586)
Property and equipment, net	\$7,782	\$8,586

Property and equipment are recorded at cost. Depreciation and amortization are computed using the straight-line method over the estimated useful lives of the respective assets, ranging from three to thirty-nine years.

As stated above, the Company sold the buildings located at 5 Jules Lane, New Brunswick, NJ and the building located at 783 Jersey Lane, New Brunswick, NJ.

(d) Patent and Trademark Rights

Patents and trademarks are stated at cost (primarily legal fees) and are amortized using the straight line method over the established useful life of 17 years. The Company reviews its patents and trademark rights periodically to determine whether they have continuing value or their value has become impaired. Such review includes an analysis of the patent and trademark's ultimate revenue and profitability potential. Management's review addresses whether each patent continues to fit into the Company's strategic business plans.

(e) Revenue

The Company has elected to apply the Full Retrospective Application to implement the new revenue recognition standard ASC 606. The Company, based on the nature of its Ampligen sales under its cost recovery programs, determined that there were no material differences between the new accounting standard and legacy GAAP and that difficulties did not arise for any "open" contract issues with its customers during the transition period. The Company also determined that the adoption of this standard had little or no impact to the Company's opening balance of retained earnings.

Revenue from the sale of Ampligen® under a cost recovery, open-label treatment protocols approved by the FDA is recognized when the treatment is provided to the patient.

Revenues from the sale of Alferon N Injection® are recognized when the product is shipped and title is transferred to the customer. The Company has no other obligation associated with its products once shipment has been shipped to the customer.

(f) Accounting for Income Taxes

Deferred income tax assets and liabilities are determined based on differences between the financial statement reporting and tax bases of assets and liabilities and are measured using the enacted tax rates and laws in effect when the differences are expected to reverse. The measurement of deferred income tax assets is reduced, if necessary, by a valuation allowance for any tax benefits which are not expected to be realized. The effect on deferred income tax assets and liabilities of a change in tax rates is recognized in the period that such tax rate changes are enacted.

The Company applies the provisions of Financial Accounting Standards Board (“FASB”) Accounting Standards Codification (“ASC”) 740-10 Uncertainty in Income Taxes. There has been no material change to the Company’s tax position as they have not paid any corporate income taxes due to operating losses. All tax benefits will likely not be recognized due to the substantial net operating loss carryforwards which will most likely not be realized prior to expiration. With no tax due for the foreseeable future, the Company has determined that a policy to determine the accounting for interest or penalties related to the payment of tax is not necessary at this time. The 2017 Tax Act, which was signed into law on December 22, 2017, has resulted in significant changes to the U.S. corporate income tax system. These changes include a federal statutory rate reduction from 35% to 21%, the elimination or reduction of certain domestic deductions and credits and limitations on the deductibility of interest expense and executive compensation. The Company determined that there was little or no material impact on its consolidated financial statements resulting from the 2017 Tax Act.

(g) Comprehensive loss

Comprehensive loss consists of net loss, net unrealized gains (losses) on securities and is presented in the consolidated statements of comprehensive loss.

(h) Use of Estimates

The preparation of financial statements in conformity with accounting principles generally accepted in the United States of America 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 financial statements and the reported amounts of revenues and expenses for the reporting period. Actual results could differ from those estimates. Accounts requiring the use of significant estimates include valuation allowances for inventory, determination of other-than-temporary impairment on securities, valuation of deferred taxes, patent and trademark valuations, stock-based compensation calculations, building valuation, fair value of warrants, convertible note payable and contingency accruals.

(i) Recent Accounting Standards and Pronouncements

In May 2014, the Financial Accounting Standards Board (“FASB”) issued Accounting Standards Update No. 2014-09 (ASU 2014-09), *Revenue from Contracts with Customers*. ASU 2014-09 eliminated transaction- and industry-specific revenue recognition guidance under current U.S. GAAP and replace it with a principle based approach for determining revenue recognition. ASU 2014-09 will require that companies recognize revenue based on the value of transferred goods or services as they occur in the contract. ASU 2014-09 also will require additional disclosure about the nature, amount, timing and uncertainty of revenue and cash flows arising from customer contracts, including significant judgments and changes in judgments and assets recognized from costs incurred to obtain or fulfill a contract. ASU 2014-09 is effective for reporting periods beginning after December 15, 2017, and early adoption is not permitted. Entities can transition to the standard either retrospectively or as a cumulative-effect adjustment as of the date of adoption. As of December 31, 2017, we have not identified any accounting changes that would materially impact the amount of reported revenues with respect to our product revenues. The Company applied the Full Retrospective Application to implement the new revenue recognition standard ASC 606. The Company, based on the nature of its Ampligen® sales under its cost recovery programs, determined that there were no material differences between the new accounting standard and legacy GAAP and that difficulties did not arise for any “open” contract issues with its customers during the transition period. The Company also determined that the adoption of this standard had little or no impact to the Company’s opening balance of retained earnings.

In January 2016, the (“FASB”) has issued Accounting Standards Update (ASU) No. 2016-01, *Financial Instruments – Overall (Subtopic 825-10): Recognition and Measurement of Financial Assets and Financial Liabilities*. The new guidance is intended to improve the recognition and measurement of financial instruments. The new guidance is effective for public companies for fiscal years beginning after December 15, 2017, including interim periods within those fiscal years. The new guidance permits early adoption of the own credit provision. The Company believes that the adoption of the guidance had no material impact on the Company’s financial statement presentation or disclosures.

In February 2016, the FASB issued ASU 2016-02 - *Leases*, which amends the existing accounting standards for lease accounting, including requiring lessees to recognize most leases on their balance sheets and making targeted changes to lessor accounting. ASU 2016-02 will be effective for annual reporting periods beginning after December 15, 2018, and early adoption of is permitted as of the standard's issuance date. ASU 2016-02 allows a modified retrospective transition approach for all leases existing at, or entered into after, the date of initial application, with an option to use certain transition relief. The Company is currently evaluating the effects the adoption of this guidance will have on the consolidated financial statements. See note 12 leases.

In August 2016, the FASB issued ASU 2016-15 - *Statement of Cash Flows (Topic 230): Classification of Certain Cash Receipts and Cash Payments* (a consensus of the Emerging Issues Task Force). The new guidance is intended to address the diversity in practice in how certain cash receipts and cash payments are presented and classified in the statement of cash flows under Topic 230, *Statement of Cash Flows*, and other Topics. The guidance addresses eight specific cash flow issues with the objective of reducing the existing diversity in practice. The amendments apply to all entities, including both business entities and not-for-profit entities that are required to present a statement of cash flows under Topic 230. The amendments are effective for public business entities for fiscal years beginning after December 15, 2017, and interim periods within those fiscal years. Early adoption is permitted, including adoption in an interim period. If an entity early adopts the amendments in an interim period, any adjustments should be reflected as of the beginning of the fiscal year that includes that interim period. An entity that elects early adoption must adopt all of the amendments in the same period. The amendments in this Update should be applied using a retrospective transition method to each period presented. The Company believes that the adoption of the guidance did not have a material impact on the Company's financial statement presentation or disclosures.

In 2018, the FASB also issued Accounting Standards Updates ("ASU") 2018-01 through 2018-20. These updates did not have a significant impact on the financial statements.

(j) Stock-Based Compensation

The Company accounts for its stock-based compensation awards in accordance with FASB ASC Topic 718, "Compensation – Stock Compensation", which requires recognition of compensation expense related to stock-based compensation awards over the period during which an employee is required to provide service for the award. Compensation expense is equal to the fair value of the award at the date of grant, net of estimated forfeitures.

(k) Accounts Receivable

Concentration of credit risk, with respect to accounts receivable, is limited due to the Company's credit evaluation process. The Company does not require collateral on its receivables. The Company's receivables were \$235,000 and

\$24,000 as of December 31, 2018 and 2017, respectively.

(l) Common Stock Per Share Calculation

Basic and diluted net loss per share is computed using the weighted average number of shares of common stock outstanding during the period. Equivalent common shares, consisting of 22,054,910, and 9,373,286 of stock options and warrants, are excluded from the calculation of diluted net loss per share for the years ended December 31, 2018 and 2017, respectively, since their effect is antidilutive due to the net loss of the Company.

(m) Long-Lived Assets

The Company assesses long-lived assets for impairment when events or changes in circumstances indicate that the carrying value of the assets or the asset grouping may not be recoverable. Factors that the Company considers in deciding when to perform an impairment review include significant under-performance of a business or product line in relation to expectations, significant negative industry or economic trends, and significant changes or planned changes in its use of the assets. The Company measures the recoverability of assets that it will continue to use in its operations by comparing the carrying value of the asset grouping to our estimate of the related total future undiscounted net cash flows. If an asset grouping's carrying value is not recoverable through the related undiscounted cash flows, the asset grouping is considered to be impaired.

The Company measures the impairment by comparing the difference between the asset grouping's carrying value and its fair value. Long-lived assets are considered a non-financial asset and are recorded at fair value only if an impairment charge is recognized. Impairments are determined for groups of assets related to the lowest level of identifiable independent cash flows. The Company makes subjective judgments in determining the independent cash flows that can be related to specific asset groupings. In addition, as the Company reviews its manufacturing process and other manufacturing planning decisions, the Company must make subjective judgments regarding the remaining useful lives of assets. When the Company determines that the useful lives of assets are shorter than the Company had originally estimated, it accelerates the rate of depreciation over the assets' new, shorter useful lives.

(3) Inventories

The Company uses the lower of first-in, first-out (“FIFO”) cost or net realizable value method of accounting for inventory.

Commercial sales of Alferon in the U.S. will not resume until new batches of commercial filled and finished product are produced and released by the Food and Drug Administration (“FDA”). While the facility is approved by the FDA under the Biologics License Application (“BLA”) for Alferon, this status will need to be reaffirmed by an FDA pre-approval inspection. The Company also will need the FDA’s approval to release commercial product once it has submitted satisfactory stability and quality release data. Currently, the manufacturing process is on hold and there is no definitive timetable to have the facility back online. The Company estimates it will need approximately \$10,000,000 to commence the manufacturing process. Due to the Company extending the timeline of Alferon production to an excess of one year, the Company reclassified Alferon work in process inventory of \$1,095,000 to other assets within our balance sheet as of December 31, 2018 and 2017 and due to the high cost estimates to bring the facility back online. The above estimated cost includes additional funds needed for the revalidation process in the Company’s facility to initiate commercial manufacturing, thereby readying itself for an FDA Pre-Approval Inspection. If the Company is unable to gain the necessary FDA approvals related to the manufacturing process and/or final product of new Alferon inventory, its operations most likely will be materially and/or adversely affected. In light of these contingencies, there can be no assurances that the approved Alferon N Injection product will be returned to production on a timely basis, if at all, or that if and when it is again made commercially available, it will return to prior sales levels.

The Alferon work in process is currently compliant with our internal protocols, is stored in a controlled state, and the Company regularly monitors the stability of the product. All of these factors contribute to the potential sale of the Alferon work in process, after validation lots have been produced and including a successful pre-approval inspection.

(4) Marketable Securities

Marketable securities consist of mutual funds. For the twelve months ended December 31, 2018 and 2017, it was determined that none of the marketable securities had other-than-temporary impairments. At December 31, 2018 and 2017, all securities were classified as available for sale investments and were measured as Level 1 instruments of the fair value measurements standard (see Note 18: Fair Value).

Securities classified as available for sale consisted of:

December 31, 2018

(in thousands)

Securities	Amortized Cost	Gross Unrealized Gains	Gross Unrealized Losses	Fair Value	Short-Term Investments	Long Term Investments
Mutual Funds	\$ 1,529	\$ —	\$ (3)	\$1,526	\$ 1,526	\$ —
Totals	\$ 1,529	\$ —	\$ (3)	\$1,526	\$ 1,526	\$ —

December 31, 2017

(in thousands)

Securities	Amortized Cost	Gross Unrealized Gains	Gross Unrealized Losses	Fair Value	Short-Term Investments	Long Term Investments
Mutual Funds	\$ 684	\$ 11	\$ —	\$ 695	\$ 695	\$ —
Totals	\$ 684	\$ 11	\$ —	\$ 695	\$ 695	\$ —

F-12

Unrealized losses on investments

Investments with continuous unrealized losses for less than 12 months and 12 months or greater and their related fair values were as follows:

There were no investments with continuous unrealized losses for less than 12 months and 12 months or greater at December 31, 2018 and 2017.

(5) Patents, Trademark Rights and Other Intangibles (FASB ASC 350-30 General Intangibles Other than Goodwill)

During the years ended December 31, 2018 and 2017, the Company decided not to pursue certain patents in various countries for strategic reasons and recorded abandonment charges of \$41,000, and \$7,000, respectively, which are included in research and development. Amortization expense was \$ 64,000 and \$56,000 in 2018 and 2017, respectively. The total cost of the patents was \$1,183,000 and \$1,107,000 as of December 31, 2018 and 2017, respectively. The accumulated amortization as of December 31, 2018 and 2017 is \$272,000 and \$249,000, respectively. For the year ended December 31, 2018 and 2017, additions to patents costs and licensing fees were \$118,000 and \$49,000, respectively.

Amortization of patents and trademarks for each of the next five years is as follows: 2019 - \$64,000; 2020 - \$64,000; 2021 - \$64,000; 2022 - \$64,000 and 2023 - \$64,000. No amortization expense is recognized related to patents that are pending.

(6) Accrued Expenses

Accrued expenses at December 31, 2018 and 2017 consist of the following:

	(in thousands)	
	December 31,	
	2018	2017
Compensation	\$613	\$569
Professional fees	83	506

Clinical trial expenses	7	310
Other expenses	302	581
	\$1,005	\$1,966

F-13

(7) Stockholders' Equity

(a) Preferred Stock

The Company is authorized to issue 5,000,000 shares of \$0.01 par value preferred stock with such designations, rights and preferences as may be determined by the Board of Directors. There were no Preferred Shares issued and outstanding at December 31, 2018 and 2017.

(b) Common Stock

The Company has authorized shares of 350,000,000 with specific limitations and restrictions on the usage of 8,000,000 of the 350,000,000 authorized shares.

In August 2016, the Company effected a 12 to 1 reverse stock split of the outstanding shares, in order to become compliant with the NYSE regulations. This did not affect the number of authorized shares.

On September 6, 2016, we entered into a Securities Purchase Agreement (the "September Purchase Agreement") with certain investors for the sale by us of 3,333,334 shares of our common stock registered under our S-3 shelf registration statement on at a purchase price of \$1.50 per share. Concurrently with the sale of the common stock, pursuant to the September Purchase Agreement, we also sold unregistered warrants to purchase 2,500,000 shares of common stock for aggregate gross proceeds of \$5,000,000. Subject to certain ownership limitations, the warrants are initially exercisable six-month after issuance at an exercise price equal to \$2.00 per share of common stock, subject to adjustments as provided under the terms of the warrants. The warrants are exercisable for five years from the initial exercise date. Pursuant to an engagement agreement, we paid our placement agent an aggregate fee equal to 7% of the gross proceeds received by us from the sale of the securities in the offering and granted to our placement agent or its designees warrants to purchase up to 5% of the aggregate number of shares sold in the transactions amounting to 166,667 unregistered warrants. The placement agent warrants have substantially the same terms as the investor warrants, except that the placement agent warrants will expire on September 1, 2021 and have an exercise price equal to \$1.875 per share of common stock.

On February 1, 2017, we entered into Securities Purchase Agreements (each, a "February Purchase Agreement") with certain investors for the sale by us of 1,818,185 shares of our common stock at a purchase price of \$0.55 per share. Concurrently with the sale of the common stock, pursuant to the February Purchase Agreement, we also sold unregistered warrants to purchase 1,363,639 shares of common stock for aggregate gross proceeds of approximately

\$1,000,000. The warrants have an exercise price of \$0.75 per share, are exercisable six months after issuance, and will expire five years from the initial exercise date. Pursuant to an engagement agreement, we paid our placement agent an aggregate fee equal to 7% of the gross proceeds received by us from the sale of the securities in the offering and granted to our placement agent or its designees warrants to purchase up to 5% of the aggregate number of shares sold in the transactions amounting to 90,910 unregistered warrants. The placement agent warrants have substantially the same terms as the investor warrants, except that the placement agent warrants will expire on February 1, 2022 and have an exercise price equal to \$0.6875 per share of common stock. The Company subsequently registered the shares issuable upon exercise of the warrants on form S-1.

The Board of Directors approved up to \$500,000 for all directors, officers and employees to buy company shares from the Company at the market price. As of November 5, 2018, the Company issued 980,392 shares of its common stock at prices between \$0.20 and \$0.69 per share directly to executives and employees, for \$373,852 in a series of private transactions pursuant to stock purchase agreements.

On June 1, 2017, the exercise price of Warrants issued in September 2016 was changed to \$0.50. As a result, the warrant holders exercised these Warrants and purchased 2,370,000 shares of Company common stock. The Company realized net proceeds of \$1,055,000 from this exercise. In conjunction with the foregoing, the Company also issued 2,370,000 series A warrants with an exercise price of \$0.60 per share, an initial exercise date of December 1, 2017 and expiring March 6, 2022 (the "Series A Warrants") and 7,584,000 series B warrants with exercise price of \$0.60, an initial exercise date December 1, 2017 per share and expiring March 1, 2018. The foregoing transactions are hereinafter referred to as the "Exchange Transaction". In addition, on July 10, 2017, the warrant holders exercised the remaining 130,000 warrants issued in September 2016 and purchased 130,000 shares of common stock. The Company realized net proceeds of \$65,000 from this exercise. In conjunction with the foregoing the Company issued 130,000 Series A Warrants and 416,000 Series B Warrants (with an exercise price of \$0.60 and an initial exercise date January 10, 2018 on the three-month anniversary of the of the initial exercise date).

Pursuant to an engagement agreement, the Company paid its placement agent an aggregate fee equal to 7% and 10.5%, respectively, of the gross proceeds received by the Company from the sale of the securities in the offerings and granted to its placement agent or its designees warrants to purchase up to 5% of the aggregate number of shares sold in the transactions amounting to 166,667 and 107,759, respectively, unregistered warrants. The placement agent warrants have substantially the same terms as the investor warrants, except that the 166,667 placement agent warrants issued in September 2017 will expire September 1, 2021 and have an exercise price equal to \$1.875 per share of common stock and the 107,759 placement agent warrants issued in June 2017 will expire June 1, 2022 and have an exercise price of \$0.625.

On August 23, 2017, the Holders of the Series A Warrants and Series B Warrants exchanged all of their Warrants for new warrants (respectively, the “Series A Exchange Warrants” and the “Series B Exchange Warrants” and, collectively, the “Exchange Warrants”) identical to the Warrants except as follows: The exercise price of both Exchange Warrants is \$0.45 per share, subject to adjustment therein, and the number of Series B Exchange Warrants issued was proportionately reduced to an aggregate of 2,800,000 warrants so that all Exchange Warrants in the Exchange Transaction do not exceed 19.9% of the number of the Company’s issued and outstanding shares of Common Stock as of May 31, 2017, the date of the Exchange Transaction offer letters. The issuance of the Exchange Warrants by the Company and the shares of Common Stock issuable upon exercise of the Exchange Warrants is exempt from registration pursuant to Sections 3(a)(9) and 4(a)(2) of the Securities Act of 1933, as amended (the “Securities Act”). The 2,800,000 warrants with an expiration date of March 1, 2018 and an exercise price on \$0.45 were exercised in January and February 2018. The Company realized proceeds of \$1,260,000 from these exercises.

On November 27, 2017, the Company reactivated the EDA. During the year ended December 31, 2018, the Company sold an aggregate of 2,176,392 shares under the EDA for proceeds of \$827,000 net of \$25,000 in commissions. Pursuant to a prospectus supplement dated February 7, 2018, the Company was able to sell up to 6,549,157 of its common stock (inclusive of shares already sold under the prospectus supplement) under the EDA. The actual number of shares, that the Company can sell, and the proceeds to be received there from are dependent upon the market price of its common stock.

Effective with the semi-monthly period ended April 30, 2017, all of the members of the Company’s Board of Directors agreed to accept 100% of their directors’ fees in the form of options to purchase Company Common Stock. This program was terminated as of August 31, 2017. In this regard, options to purchase 226,023 shares of Company common stock were issued with exercise prices ranging from \$0.36 to \$0.67, a holding period of 10 years and vesting over three years. In addition, commencing with the semi-monthly period ended June 15, 2017, certain officers of the Company and certain other employees of the Company, agreed to accept 20% of their salary in options to purchase Company Common Stock. This program was also terminated as of August 31, 2017. In this regard, options to purchase 214,866 shares of Company common stock were issued with exercise prices ranging from \$0.36 to \$0.67, a holding period of 10 years and vesting over three years.

As part of the cash conservation program adopted on August 28, 2017, starting with the month of September 2017, the directors agreed to defer 100% of their fees until cash is available. In consideration of this deferral, 226,023 options

were issued to each of the two independent directors in February 2018 with an exercise price of \$0.37; 152,053 options were issued to each of the two independent directors in May 2018 with an exercise price of \$0.30, and 98,098 options were issued in July 2018 with an exercise price of \$0.31. All of the foregoing options and the options discussed below are exercisable for a period of 10 years with a vesting period of three years. This program was suspended as of July 15, 2018 and all remaining deferred fees were paid in July 2018. This Program was reactivated as of August 16, 2018 with the understanding that options would not be issued on the deferred amounts until the 2018 Equity Incentive Plan was approved by the stockholders and the securities issuable thereunder were registered with the SEC. The 2018 Equity Incentive Plan was approved by the stockholders and the securities issuable thereunder were registered with the SEC and, on October 17, 2018, 172,786 options were issued to each of the two independent directors with an exercise price on \$0.22 for a period of ten years with a vesting period of one year.

Also as part of the cash conservation program adopted on August 28, 2017, starting with the month of September 2017, certain officers agreed to defer 40% of their salaries until cash is available. In consideration of this deferral, 884,459 options were issued to these officers in February 2018 with an exercise price of \$0.37; 599,168 options were issued to these officers in May 2018 with an exercise price of \$0.30, and 389,249 options were issued to these officers in July 2018 with an exercise price of \$0.31. This program was suspended as of July 15, 2018 and all remaining deferred salaries were paid on July 2018. This Program was reactivated as of August 16, 2018 for 50% of their salaries with the understanding that options would not be issued on the deferred amounts until the 2018 Equity Incentive Plan was approved by the shareholders and the plan registered with the SEC. The 2018 Equity Incentive Plan has been approved by the shareholders and registered with the SEC and on October 17, 2018, 808,712 options were issued to these officers with an exercise price on \$0.22 for a period of ten years with a vesting period of one year

Also as part of the cash conservation program adopted on August 28, 2017, all employees agreed to be paid 50% of their salaries in the form of unrestricted common stock of the Company. Starting with the month of September 2017, the salaries of all the employees of the Company were paid 50% in the form of unrestricted common stock of the Company. The total number of shares issued as of June 30, 2018 to the employees under this program was 2,116,881 shares at stock prices ranging from \$0.31 to \$0.55 per share. This program was suspended by the Board of Directors on June 30, 2018.

On March 24, 2018, the Company sold 1,250,000 shares of common stock under its S-3 shelf registration. The Company realized net proceeds of \$475,000 from this stock offering and paid \$25,000 in placement agent fees.

On April 20, 2018, the Company entered into Securities Purchase Agreements (the “Purchase Agreements”) with certain investors (the “Investors”) for the sale by the Company of an aggregate of 6,600,000 shares (the “Common Shares”) of the Company’s common stock, par value \$0.001 per share (the “Common Stock”), at a purchase price of \$0.39 per share. Concurrently with the sale of the Common Shares, pursuant to the Purchase Agreements the Company also sold 6,600,000 warrants, 50% of which are Class A Warrants and 50% of which are Class B Warrants (collectively, the “Warrants”). The Company received gross proceeds from the sale of the Warrants solely to the extent such Warrants are exercised for cash. Both classes of Warrants will not be exercisable until six months after issuance and will have an exercise price of \$0.39 per share, subject to adjustments as provided under the terms of the Warrants. The Class A Warrants and Class B Warrants will expire, respectively, two and five years after the date on which they are first exercisable. The closing of the sales of these securities under the Purchase Agreements took place on April 24, 2018. The Company received net proceeds from the transactions of \$2,343,820 after deducting certain fees due to the placement agent and the Company’s transaction expenses.

The 2009 Equity Incentive Plan, effective June 24, 2009, as amended, authorizes the grant of non-qualified and incentive stock options, stock appreciation rights, restricted stock and other stock awards. A maximum of 22,000,000 shares of common stock is reserved for potential issuance pursuant to awards under the 2009 Equity Incentive Plan. Unless sooner terminated, the 2009 Equity Incentive Plan will continue in effect for a period of 10 years from its effective date. During 2018, there were 4,675,221 options granted by the Company under this Plan.

The 2018 Equity Incentive Plan, effective September 12, 2018, authorizes the grant of (i) Incentive Stock Options, (ii) Nonstatutory Stock Options, (iii) Stock Appreciation Rights, (iv) Restricted Stock Awards, (v) Restricted Stock Unit Awards, (vi) Performance Stock Awards, (vii) Performance Cash Awards, and (viii) Other Stock Awards. Initially, a maximum of 7,000,000 shares of common stock is reserved for potential issuance pursuant to awards under the 2018 Equity Incentive Plan. Unless sooner terminated, the 2018 Equity Incentive Plan will continue in effect for a period of 10 years from its effective date. On October 17, 2018, the Board of Directors issued 1,154,284 options to the officers and directors at the exercise price of \$0.22 expiring in 10 years, and on November 14, 2018, the Board of Directors issued 1,000 options to each employee, officer and director at the exercise price of \$0.22 expiring in ten years.

As of December 31, 2018 and 2017, there were 48,734,712 and 32,884,786 shares outstanding, respectively.

(c) Equity Financings

See (b) above

(d) Common Stock Options and Warrants

(i) Stock Options

The Equity Incentive Plan of 2009, effective June 24, 2009, as amended, authorizes the grant of non-qualified and incentive stock options, stock appreciation rights, restricted stock and other stock awards. A maximum of 22,000,000 shares of common stock is reserved for potential issuance pursuant to awards under the Equity Incentive Plan of 2009. Unless sooner terminated, the Equity Incentive Plan of 2009 will continue in effect for a period of 10 years from its effective date.

The 2018 Equity Incentive Plan, effective September 12, 2018, authorizes the grant of (i) Incentive Stock Options, (ii) Nonstatutory Stock Options, (iii) Stock Appreciation Rights, (iv) Restricted Stock Awards, (v) Restricted Stock Unit Awards, (vi) Performance Stock Awards, (vii) Performance Cash Awards, and (viii) Other Stock Awards. Initially, a maximum of 7,000,000 shares of common stock is reserved for potential issuance pursuant to awards under the 2018 Equity Incentive Plan. Unless sooner terminated, the 2018 Equity Incentive Plan will continue in effect for a period of 10 years from its effective date. On October 17, 2018, the Board of Directors issued 1,154,284 options to the officers and directors at the exercise price of \$0.22 expiring in 10 years, and on November 14, 2018, the Board of Directors issued 1,000 options to each employee, officer and director at the exercise price of \$0.22 expiring in ten years.

The Equity Incentive Plans of 2009 and 2018 are administered by the Board of Directors. The Plans provide for awards to be made to such Officers, other key employees, non-employee Directors, consultants and advisors of the Company and its subsidiaries as the Board may select.

Stock options awarded under the Plans may be exercisable at such times (not later than 10 years after the date of grant) and at such exercise prices (not less than fair market value at the date of grant) as the Board may determine. The Board may provide for options to become immediately exercisable upon a “change in control”, which is defined in the Plans to occur upon any of the following events: (a) the acquisition by any person or group, as beneficial owner, of 20% or more of the outstanding shares or the voting power of the outstanding securities of the Company; (b) either a majority of the Directors of the Company at the annual stockholders meeting has been nominated other than by or at the direction of the incumbent Directors of the Board, or the incumbent Directors cease to constitute a majority of the Company’s Board; (c) the Company’s stockholders approve a merger or other business combination pursuant to which the outstanding common stock of the Company no longer represents more than 50% of the combined entity after the transaction; (d) the Company’s stockholders approve a plan of complete liquidation or an agreement for the sale or disposition of all or substantially all of the Company’s assets; or (e) any other event or circumstance determined by the Company’s Board to affect control of the Company and designated by resolution of the Board as a change in control.

The fair value of each option and equity warrant award is estimated on the date of grant using a Black-Scholes-Merton pricing option valuation model. Expected volatility is based on the historical volatility of the price of the Company’s stock. The risk-free interest rate is based on U.S. Treasury issues with a term equal to the expected life of the option and equity warrant. The Company uses historical data to estimate expected dividend yield, life and forfeiture rates. The expected life of the options and equity warrants was estimated based on historical option and equity warrant holders’ behavior and represents the period of time that options and equity warrants are expected to be outstanding. The fair values of the options and equity warrants granted were estimated based on the following weighted average assumptions:

	Year Ended December 31,	
	2018	2017
Risk-free interest rate	2.6%-3.0%	1.72%-1.89%
Expected dividend yield	-	-
Expected life	5 years	1.25-5 years
Expected volatility	85.68%-86.89%	91.60%-144.15%
Weighted average grant date fair value for options and equity warrants issued	\$0.23 per option for 5,864,505 options	\$0.35 per option/warrant for 1,340,517 options/equity warrants

The exercise price of all stock options and equity warrants granted was equal to or greater than the fair market value of the underlying common stock on the date of the grant.

Information regarding the options approved by the Board of Directors under Equity Plan of 2009 is summarized below. The plan expires June 24, 2019:

	2017		Weighted Average Exercise Price	2018		Weighted Average Exercise Price
	Shares	Option Price		Shares	Option Price	
Outstanding, beginning of year	695,061	1.56-48.36	4.70	1,877,295	0.33-48.36	1.92
Granted	1,190,567	0.33-0.67	0.29	4,675,221	0.30-0.38	0.34
Forfeited	(8,333)	1.56	1.56	(213,855)	8.16-\$48.36	28.26
Exercised	—	—	—	—	—	—
Outstanding, end of year	1,877,295	0.33-48.36	1.92	6,338,661	0.30-48.36	0.36
Exercisable, end of year	1,046,487	0.33-48.36	0.42	2,376,353	0.30-48.36	
Weighted average remaining contractual life (years)	2-10 years			1-10 years		
Available for future grants	4,139,454			159,322		

Information regarding the options approved by the Board of Directors under the Equity Plan of 2018 is summarized below:

	2018		
	Shares	Option Price	Weighted Average Exercise Price
Outstanding, beginning of year	—	—	—
Granted	1,189,284	0.22	0.22
Forfeited	—	—	—
Exercised	—	—	—
Outstanding, end of year	1,189,284	0.22	0.22
Exercisable, end of year	294,404	0.22	0.22
Weighted average remaining contractual life (years)	10 years		
Available for future grants	5,810,716		

Stock option activity during the years ended December 31, 2017 and 2018 is as follows:

Stock option activity for employees

	Number of Options	Weighted Average Exercise Price	Weighted Average Remaining Contracted Term (Years)	Aggregate Intrinsic Value
Outstanding December 31, 2016	836,256	\$ 16.82	4.47	—
Granted	584,794	0.50	—	—
Forfeited	(217,132)	33.35	—	—
Outstanding December 31, 2017	1,203,918	\$ 5.91	6.89	—
Granted	4,164,585	0.32	—	—
Forfeited	(257,917)	18.01	—	—
Outstanding December 31, 2018	5,110,586	\$ 0.75	9.01	—
Vested and expected to vest at December 31, 2018	5,110,586	\$ 0.75	9.01	—
Exercisable at December 31, 2018	1,804,927	\$ 1.04	8.09	—

The weighted-average grant-date fair value of employee options granted during the year 2018 was \$958,000 for 4,164,585 options at \$0.23 per option and during year 2017 was \$230,000 for 584,794 options at \$0.39 per option.

Unvested stock option activity for employees:

	Number of Options	Weighted Average Exercise Price	Average Remaining Contracted Term (Years)	Aggregate Intrinsic Value
Unvested December 31, 2016	90,625	\$ 1.72	9.33	—
Granted	584,794	0.50	—	—
Vested	(309,271)	0.88	—	—
Forfeited	—	—	—	—
Unvested December 31, 2017	366,148	\$ 0.48	9.62	—
Granted	4,164,585	0.32	—	—
Vested	(1,225,074)	0.36	—	—
Forfeited	—	—	—	—
Unvested December 31, 2018	3,305,659	\$ 0.32	9.31	—

The weighted-average grant-date fair value of employee unvested stock options granted during the year 2018 was \$958,000 for 4,164,585 at \$0.23 per option and during the year 2017 was \$230,000 for 584,794 options at \$0.39 per option,.

F-19

Stock option activity for non-employees during the year:

	Number of Options	Weighted Average Exercise Price	Weighted Average Remaining Contracted Term (Years)	Aggregate Intrinsic Value
Outstanding December 31, 2016	271,500	\$ 10.41	4.66	—
Granted	605,772	0.42	—	—
Exercised	—	—	—	—
Forfeited	(42,396)	19.41	—	—
Outstanding December 31, 2017	834,876	\$ 2.70	6.69	—
Granted	1,699,920	0.31	—	—
Exercised	—	—	—	—
Forfeited	(109,104)	10.33	—	—
Outstanding December 31, 2018	2,425,692	\$ 0.68	8.55	—
Vested and expected to vest at December 31, 2018	2,425,692	\$ 0.68	8.55	—
Exercisable at December 31, 2018	874,164	\$ 1.52	7.78	—

The weighted-average grant-date fair value of non-employee options granted during year 2018 was \$391,000 for 1,699,920 options at \$0.23 per option and during the year 2017 was \$182,000 for 605,772 options at \$0.30 per option.

Unvested stock option activity for non-employees:

	Number of Options	Weighted Average Exercise Price	Weighted Average Remaining Contracted Term (Years)	Aggregate Intrinsic Value
Unvested December 31, 2016	26,389	\$ 1.65	8.61	—
Granted	606,772	0.42	—	—
Vested	(163,335)	0.57	—	—
Forfeited	(4,861)	7.58	—	—
Unvested December 31, 2017	464,965	\$ 0.36	7.84	—
Granted	1,699,920	0.31	—	—
Vested	(613,357)	0.35	—	—
Forfeited	—	—	—	—
Unvested December 31, 2018	1,551,528	\$ 0.31	8.84	—

Stock-based compensation expense was approximately \$929,000 and \$571,000 for the years ended December 31, 2018, and 2017 resulting in an increase in general and administrative expenses and loss per share of \$0.02 and \$0.02, respectively.

As of December 31, 2018 and 2017, there was \$1,273,000 and \$435,000, respectively, of unrecognized stock-based compensation cost related to options granted under the Equity Incentive Plans. Stock-based compensation related to options granted under the Equity Incentive Plans will be recorded over the vesting period which is typically one year or upon reaching agreed upon company and/or individual performance milestones being met which is indefinite.

(ii) Stock Warrants

F-20

Stock warrants are issued as needed by the Board of Directors and have no formal plan.

The fair value of each warrant award is estimated on the date of grant using a Black-Scholes-Merton pricing option valuation model. Expected volatility is based on the historical volatility of the price of the Company's stock. The risk-free interest rate is based on U.S. Treasury issues with a term equal to the expected life of the warrant. The Company uses historical data to estimate expected dividend yield, life and forfeiture rates. The expected life of the warrants was estimated based on historical option holder's behavior and represents the period of time that options are expected to be outstanding. There were 17,512,308 granted in 2017 at \$0.45 to \$0.75 per warrant, and 9,825,808 granted in 2018 at \$0.39 per warrant.

Information regarding warrants outstanding and exercisable into shares of common stock is summarized below:

	2017			2018		
	Shares	Warrant Price	Weighted Average Exercise Price	Shares	Warrant Price	Weighted Average Exercise Price
Outstanding, beginning of year	2,830,516	\$1.08-\$24.00	\$ 2.16	7,334,490	\$0.45-\$10.68	\$ 0.63
Granted	17,512,308	\$0.45-\$0.75	\$ 0.57	9,825,808	\$0.39	\$ 0.39
Forfeited	(10,508,334)	\$0.60-\$24.00	\$ 0.61	(25,000)	\$6.00	\$ 6.00
Exercised	(2,500,000)	\$0.50	\$ 0.50	(2,800,000)	\$0.45	\$ 0.45
Outstanding, end of year	7,334,490	\$0.45-\$10.68	\$ 0.63	14,335,298	\$0.39-\$10.68	\$ 0.49
Exercisable	7,334,490	\$0.45-\$10.68	\$ 0.63	14,335,298	\$0.39-\$10.68	\$ 0.49
Weighted average remaining contractual life	2.7 years			2.3 years		
Years exercisable	2017-2023			2019-2023		

Stock warrants are issued at the discretion of the Board. In 2018 there were 9,825,808 warrants issued at a weighted average exercise price of \$0.39 and in 2017, there were 17,512,308 warrants issued at a weighted average exercise price of \$0.57 per share. 2,800,000 warrants were exercised in 2018 and 2,500,000 warrants were exercised in 2017.

(e) Rights Offering

On November 14, 2017, at the direction of the Board, the Company amended and restated the Rights Agreement between the Company and, American Stock Transfer & Trust Company, LLC, its current Rights Agent (as amended and restated, the "Rights Agreement"). Each Right entitles the registered holder to purchase from the Company a unit consisting of one one-hundredth of a share (a "Unit") of Series A Junior Participating Preferred Stock, par value \$0.01

per share (the “Series A Preferred Stock”) at a Purchase Price of \$21.00 per Unit, subject to adjustment.

(8) Segment and Related Information

The Company operates in one segment, which performs research and development activities related to Ampligen® and other drugs under development, and sales and marketing of Alferon®. The Company’s revenues for the two-year period ended December 31, 2018, were earned in the United States and overseas.

(9) Research, Consulting and Supply Agreements

In 2016, we entered into a five-year agreement (the “Impatients Agreement”) with Impatients, N.V. (“myTomorrows”), a Netherlands based company, for the commencement and management of an EAP in Europe and Turkey (the “Territory”) related to ME/CFS. Pursuant to the agreement, myTomorrows, as our exclusive service provider and distributor in the Territory, is performing EAP activities.

In 2017, we entered into a purchase order with Jubilant pursuant to which Jubilant will manufacture batches of Ampligen® for the Company. Two commercial size batches were filled and finished for human use in 2018. We paid Jubilant \$320,000 in 2017 and \$1,078,000 in 2018 for a total of \$1,398,000 to date for these services.

In 2017, Hemispherx filed a complaint in the Philadelphia County Court of Common Pleas Civil Trial Division against Nitto Avecia Pharma Services, Inc. (“NAPS”), the successor to Avrio Biopharmaceuticals, LLC (“Avrio”), primarily for breach of contract. Pursuant to the applicable agreement, Avrio was to provide fill and finish services of Ampligen®. Hemispherx sought damages due to Avrio’s failures and omissions during the fill and finish process which led to a loss of product. In June 2017, NAPS filed an answer denying liability and counter claiming breach of contract by Hemispherx. In March of 2018, the parties agreed to fully resolve their dispute by agreement for a satisfactory payment to Hemispherx and additional consideration. There was a gain of \$474,000 resulting from the settlement of litigation with Nitto Avecia Pharma Services, Inc

The Company has an agreement with Asembia, formerly Armada Healthcare, LLC to undertake the marketing, education and sales of Alferon N Injection® throughout the United States. This agreement also provides start-up along with ongoing sales and marketing support to the Company. In August 2017, the Company extended this agreement through August 14, 2019 subject to the same terms and conditions. The Company incurred no fees for the years ended December 31, 2018, and 2017, pursuant to the original and amended agreements.

In 2017, we announced that the EAP through our agreement with myTomorrows designed to enable access of Ampligen to ME/CFS patients has been extended to pancreatic cancer patients beginning in the Netherlands. myTomorrows is our exclusive service provider in Europe and Turkey and will manage all EAP activities relating to the pancreatic cancer extension of the program.

In 2018, we signed two amendments to the EAP with myTomorrows. The first extended the territory to cover Canada to treat pancreatic cancer patients, pending government approval and the second to be our exclusive service provider for special access activities in Canada for the supply of Ampligen for the treatment of ME/CFS.

(10) 401(k) Plan

The Company has a defined contribution plan, entitled the Hemispherx Biopharma Employees 401(k) Plan and Trust Agreement (the “401(k) Plan”). Full time employees of the Company are eligible to participate in the 401(k) Plan following one year of employment. Subject to certain limitations imposed by federal tax laws, participants are eligible to contribute up to 15% of their salary (including bonuses and/or commissions) per annum. Participants’ contributions to the 401(k) Plan may be matched by the Company at a rate determined annually by the Board of Directors.

Each participant immediately vests in his or her deferred salary contributions, while Company contributions will vest over one year. A 6% Company matching contribution was established, effective as of January 1, 2010 through December 31, 2015. As of January 1, 2016, the matching has been terminated. For 2018 and 2017, the Company made no contributions towards the 401(k) Plan in these years.

(11) Royalties, License and Employment Agreements

The Company had contractual agreements with Named Executive Officers (“Officers”) in 2018, and 2017. The aggregate annual base compensation for these Officers under their respective contractual agreements for 2018, and 2017 was \$ 1,247,000, and \$1,164,000 respectively. In addition, certain of these Officers were entitled to receive performance bonuses of up to 25% or 20% of their respective annual base salary, at the sole discretion of the Compensation Committee of the Board of Directors. In 2018 and 2017, no Officers’ bonuses were granted.

In 2018, equity was granted as a form of compensation to these Officers:

The Company granted to Thomas K. Equels, Chief Executive Officer, consistent with his employment agreement 300,000 ten year options to purchase common stock with an exercise price of \$0.30 per share which vest in one year and 1,192,003 ten year options with exercise prices of \$0.22 to \$0.38 which vest in one to three years for 40% and 50% salary deferrals; and

The Company granted 430,668 ten year options to purchase common stock with exercise prices of \$0.22 to \$0.38 per share which vest in one to three years to Adam Pascale, Chief Financial Officer for 40% and 50% salary deferrals; and

The Company granted 597,935 ten year options to purchase common stock with exercise prices of \$0.22 to \$0.38 per share which vest in one to three years to Peter Rodino, General Counsel and Company Secretary for 40% and 50% salary deferrals.

In 2017, equity was granted as a form of compensation to these Officers:

The Company granted to Thomas K. Equels, Chief Executive Officer, consistent with his employment agreement 300,000 ten year options to purchase common stock with an exercise price of \$0.56 per share which vest in one year and 113,135 ten year options with exercise prices of \$0.36 to \$0.49 which vest in three years for a 20% reduction in salary; and

The Company granted 37,712 ten year options to purchase common stock with exercise prices of \$0.36 to \$0.49 per share which vest in three years to Adam Pascale, Chief Financial Officer for a 20% reduction in salary; and

During 2017, we granted 52,796 ten year options to purchase common stock with exercise prices of \$0.36 to \$0.49 per share which vest in three years to Peter Rodino, General Counsel and Company Secretary for a 20% reduction in salary.

The Company recorded stock compensation expense of approximately \$145,000 and \$86,000 during the years ended December 31, 2018 and 2017 respectively with regard to these issuances.

(12) Leases

In February 2016, the FASB established Topic 842, Leases, by issuing Accounting Standards Update (ASU) No. 2016-02, which requires lessees to recognize leases on-balance sheet and disclose key information about leasing arrangements. Topic 842 was subsequently amended by ASU No. 2018-01, Land Easement Practical Expedient for Transition to Topic 842; ASU No. 2018-10, Codification Improvements to Topic 842, Leases; ASU No. 2018-11, Targeted Improvements; and ASU No. 2018-20, Narrow-Scope Improvements for Lessors. The new standard establishes a right-of-use model (ROU) that requires a lessee to recognize a ROU asset and lease liability on the balance sheet for all leases with a term longer than 12 months. Leases will be classified as finance or operating, with classification affecting the pattern and classification of expense recognition in the income statement.

The new standard is effective for the Company on January 1, 2019, with early adoption permitted. A modified retrospective transition approach is required, applying the new standard to all leases existing at the date of initial application. An entity may choose to use either (1) its effective date or (2) the beginning of the earliest comparative period presented in the financial statements as its date of initial application. If an entity chooses the second option, the transition requirements for existing leases also apply to leases entered into between the date of initial application and the effective date. The entity must also recast its comparative period financial statements and provide the disclosures required by the new standard for the comparative periods. We expect to adopt the new standard on January 1, 2019 and use the effective date as our date of initial application. Consequently, financial information will not be updated, and the disclosures required under the new standard will not be provided for dates and periods before January 1, 2019.

The new standard provides several optional practical expedients in transition. We expect to elect the 'package of practical expedients', which permits us not to reassess under the new standard our prior conclusions about lease identification, lease classification and initial direct costs. We expect to elect all the new standard's available transition practical expedients other than the use-of hindsight.

The new standard also provides practical expedients for an entity's ongoing accounting. We currently expect to elect the short-term lease recognition exemption for all leases that qualify. This means, for those leases that qualify, we will not recognize ROU assets or lease liabilities, and this includes not recognizing ROU assets or lease liabilities for existing short-term leases of those assets in transition. We also currently expect to elect the practical expedient to not separate lease and non-lease components for leases of office equipment.

We expect that this standard will have a material effect on our financial statements. While we continue to assess all the effects of adoption, we currently believe the most significant effects relate to the recognition of new ROU assets and lease liabilities on our balance sheet for our real estate and equipment operating leases and providing significant new disclosures about our leasing activities. We do not expect a significant change in our leasing activities between now and adoption.

Rent expense charged to operations for the years ended December 31, 2018 and 2017 amounted to approximately \$76,000 and \$253,000 respectively.

(13) Income Taxes (FASB ASC 740 Income Taxes)

The Company applies the provisions of FASB ASC 740-10 Uncertainty in Income Taxes. As a result of the implementation, there has been no material change to the Company's tax position as they have not paid any corporate income taxes due to operating losses. All tax benefits will likely not be recognized due to the substantial net operating loss carryforwards which will most likely not be realized prior to expiration. The 2017 Tax Act, which was signed into law on December 22, 2017, has resulted in significant changes to the U.S. corporate income tax system. These changes include a federal statutory rate reduction from 35% to 21%, the elimination or reduction of certain domestic deductions and credits and limitations on the deductibility of interest expense and executive compensation. The 2017 Tax Act also transitions international taxation from a worldwide system to a modified territorial system and includes base erosion prevention measures on non-U.S. earnings, which has the effect of subjecting certain earnings of our foreign subsidiaries to U.S. taxation as global intangible low taxed income (GILTI). These changes are effective beginning in 2018.

As of December 31, 2018, the Company has approximately \$183,000,000 of Federal net operating loss carryforwards (expiring in the years 2019 through 2037) and \$7,000,000 of Federal net operating loss with no expiration date available to offset future federal taxable income. The Company also has approximately \$34,000,000 of Pennsylvania state net operating loss carryforwards (expiring in the years 2019 through 2033) and approximately \$10,000,000 of New Jersey state net operating loss carryforwards (expiring in 2038) available to offset future state taxable income. In December 2018 the Company effectively sold \$10,000,000 of its New Jersey state net operating loss carryforward for the year 2017 for approximately \$859,000. In December 2017, the Company effectively sold \$8,000,000 of its New Jersey state net operating loss carryforward for the year 2016 for approximately \$622,000, and also sold New Jersey research and development credits for \$169,000.

The utilization of certain state net operating loss carryforwards may be subject to annual limitations. With no tax due for the foreseeable future, the Company has determined that a policy to determine the accounting for interest or penalties related to the payment of tax is not necessary at this time.

Under the Tax Reform Act of 1986, the utilization of a corporation's net operating loss carryforward is limited following a greater than 50% change in ownership. Due to the Company's prior and current equity transactions, the Company's net operating loss carryforwards may be subject to an annual limitation generally determined by multiplying the value of the Company on the date of the ownership change by the federal long-term tax exempt rate. Any unused annual limitation may be carried forward to future years for the balance of the net operating loss carryforward period. The 2017 Tax Act eliminates the three year carryback of net operating losses and allows only the carryforward of losses.

Deferred income taxes reflect the net tax effects of temporary differences between carrying amounts of assets and liabilities for financial reporting purposes and the carrying amounts used for income tax purposes. In assessing the

realizability of deferred tax assets, Management considers whether it is more likely than not that some portion or all of the deferred tax assets will not be realized. The realization of deferred tax assets is dependent upon the generation of future taxable income during the periods in which temporary differences representing net future deductible amounts become deductible. Due to the uncertainty of the Company's ability to realize the benefit of the deferred tax asset, the deferred tax assets are fully offset by a valuation allowance at December 31, 2018 and 2017.

F-24

The components of the net deferred tax assets and liabilities as of December 31, 2018 and 2017 consist of the following:

	(in thousands)	
	December 31,	
	2018	2017
Deferred tax assets:		
Net operating losses	\$39,866	\$38,005
Amortization & depreciation	131	138
Accrued expenses	100	51
Stock compensation	191	120
Total deferred tax assets	40,288	38,314
Deferred tax liabilities:		
Research and development costs	(127)	(189)
Deferred tax assets, net	40,161	38,125
Less: Valuation allowance	(40,161)	(38,125)
Deferred tax assets, net	—	—

(14) Note Payable

In May 2017, the Company entered into a mortgage and note payable agreement with a bridge funding company to obtain a two-year funding line of up to \$4,000,000 secured by the property and assets located at 783 Jersey Avenue, New Brunswick, New Jersey. The Company borrowed \$1,900,000 of the line in monthly advances including accrued interest as of December 31, 2017. The Company was able to request future advances in excess of \$2,000,000 at the lender's discretion and be payable in full upon maturity. The Company paid interest on this note at a fixed rate of 12% per annum for the first 18 months and change to a rate equal to 800 basis points above the prime rate of interest during the remainder of the term; however, the interest rate was not to be less than 12% for the entire term. The note was interest only and payable monthly through the maturity. The Company was permitted to prepay the line without penalty commencing after six months. The balance on the note at December 31, 2017 was \$1,835,000 (\$1,900,000 less unamortized deferred finance costs of \$65,000). The note was paid off on March 16, 2018 in conjunction with the sale leaseback of the Company's above property and assets at an amount of \$1,956,803, which included all accrued interest and fees (See also Note 2(c)– Property and Equipment and Note 19).

(15) Convertible Note Payable

On September 28, 2018, the Company entered into a \$3,170,000 10% Secured Convertible Promissory Note (the "Note") with Iliad Research and Trading, L.P. (the "Holder"), which was issued to the Holder in conjunction with 500,000 shares of Common Stock (the "Origination Shares"). The Company collected \$3,000,000 in cash from the Holder during

September 2018 and the remainder \$170,000 was retained by the Holder for the Holder's legal fees of \$20,000 for the issuance of the Note and the Original Issue Discount of \$150,000. The Company incurred \$210,000 in third-party fees directly attributed to the issuance of the Note. The Company promised to pay the principal amount, together with guaranteed interest at the annual rate of 10%, with principal and accrued interest on the Note due and payable on September 28, 2019 (unless converted under terms and provisions as set forth within the Note). The Note provides the Holder with the right to convert, at any time, all or any part of the outstanding principal and accrued but unpaid interest into shares of the Company's common stock at a conversion price of \$0.30 per share. In addition, beginning on March 28, 2019, the Note also provides the Holder with the right to redeem all or any portion of the Note ("Redemption Amount"). The payments of each Redemption Amount may be made, at the option of the Company, in cash, by converting such Redemption Amount into shares of common stock ("Redemption Conversion Shares"), or a combination thereof. The number of Redemption Conversion Shares equals the portion of the applicable Redemption Amount being converted divided by the lesser of \$0.30 or 80% of the lowest Volume Weighted Average Price ("VWAP") during the ten (10) trading days immediately preceding the applicable measurement date (the "Market Price"). The Purchase Agreement requires the Company to reserve at least 8,900,000 shares of common stock from its authorized and unissued common stock to provide for all issuances of common stock under the Note. However, the Note provides that the aggregate number shares of common stock issued to the Holder under the Note and Purchase Agreement shall not exceed 19.99% of the total number of shares of common stock outstanding as of the closing date unless the Company has obtained stockholder approval of the issuance. The Origination Shares were to be returned to the Company in the event that the Company could provide within 30 days of the closing of the transaction certain requested assets as security for repayment of the Note. The security was not provided so the Origination Shares remained with the Holder.

The Company determined the Note should be recorded at fair value with subsequent changes in fair value recorded in earnings. This conclusion is based on the redemption conversion feature, which allows the Holder to trigger the redemption of the Note for cash or conversion of the Note for common shares prior to its maturity date at a price of the lesser of \$0.30 per share or the Market Price as defined within the Note. The choice of cash redemption or conversion of the Note for common shares is at the option of the Company. This feature may require the Company to issue a variable number of common shares to settle the Note which was determined to have a predominantly fixed monetary value at inception. In connection with the Note, the Company recorded a loss in the Company's Consolidated Statements of Comprehensive Income (Loss) equal to \$582,000 for the year ended December 31, 2018.

Interest expense associated with the Note was approximately \$201,000 for the year ended December 31, 2018, which included approximately \$119,000 associated with the amortization of applicable discounts to the Note.

(16) Certain Relationships and Related Transactions

The Company has employment agreements with certain of their Executive Officers and has granted such officers and directors options and warrants to purchase their common stock. Please see details of these Employment Agreements in Note 11 - Royalties, License and Employment Agreements.

As set forth in Section 3(c)(ii) of his Employment Agreement, Mr. Equels earned \$18,000 and \$22,000 for 5% of the Ampligen® cost recovery sales in 2018 and 2017, respectively.

(17) Concentrations of Credit Risk

Financial instruments, which potentially subject the Company to concentrations of credit risk, consist principally of cash, cash equivalents, investments and accounts receivable. The Company places its cash with high-quality financial institutions and, at times, such amounts in non-interest bearing accounts may be in excess of Federal Deposit Insurance Corporation insurance limits. There were no credit based sales for 2018 and 2017.

(18) Fair Value

The Company is required under GAAP to disclose information about the fair value of all the Company's financial instruments, whether or not these instruments are measured at fair value on the Company's consolidated balance

sheets.

The Company estimates that the fair values of cash and cash equivalents, other assets, accounts payable and accrued expenses approximate their carrying values due to the short-term maturities of these items. The Company also has certain warrants with a cash settlement feature in the unlikely occurrence of a Fundamental Transaction. The fair value of the redeemable warrants (“Warrants”) related to the Company’s August 2016, February 2017, June 2017, August 2017 and April 2018 common stock and warrant issuance, are calculated using a Monte Carlo Simulation. While the Monte Carlo Simulation is one of a number of possible pricing models, the Company has determined it to be industry accepted and fairly presented the fair value of the Warrants. As an additional factor to determine the fair value of the Put’s liability, the occurrence probability of a Fundamental Transaction event was factored into the valuation.

The Company recomputes the fair value of the Warrants at the issuance date and the end of each quarterly reporting period. Such value computation includes subjective input assumptions that are consistently applied each period. If the Company were to alter its assumptions or the numbers input based on such assumptions, the resulting fair value could be materially different.

F-26

The Company utilized the following assumptions to estimate the fair value of the August 2016 Warrants:

	December 31, 2018	December 31, 2017		
Underlying price per share	\$ 0.18	\$ 0.35		
Exercise price per share	\$ 1.88	\$ 1.88		
Risk-free interest rate	2.47	2.05	%	%
Expected holding period	2.67	3.70		
Expected volatility	70	65	%	%
Expected dividend yield	-	-		

The Company utilized the following assumptions to estimate the fair value of the February 2017 Warrants:

	December 31, 2018	December 31, 2017		
Underlying price per share	\$0.18	\$0.35		
Exercise price per share	\$0.69-\$0.75	\$0.69-\$0.75		
Risk-free interest rate	2.47	2.10	%	%
Expected holding period	3.59-3.60	4.1		
Expected volatility	70	65	%	%
Expected dividend yield	-	-		

The Company utilized the following assumptions to estimate the fair value of the June 2017 Warrants:

	December 31, 2018	December, 2017		
Underlying price per share	\$ 0.18	\$ 0.35		
Exercise price per share	\$ 0.63	0.63		
Risk-free interest rate	2.47	2.14	%	%
Expected holding period	3.42	4.4		
Expected volatility	70	65	%	%
Expected dividend yield	-	-		

The Company utilized the following assumptions to estimate the fair value of the August 2017 Warrants:

Edgar Filing: HEMISPHERX BIOPHARMA INC - Form 10-K

	December 31, 2018	December 31, 2017	
Underlying price per share	\$ 0.18	\$0.35	
Exercise price per share	\$ 0.45	\$0.45	
Risk-free interest rate	2.46	% 1.33%-2.11	%
Expected holding period	3.18	0.2-4.2	
Expected volatility	70	% 65	%
Expected dividend yield	-	-	

F-27

The Company utilized the following assumptions to estimate the fair value of the April 2018 Warrants:

	September 30, 2018	April 24, 2018		
Underlying price per share	\$0.18	\$0.34		
Exercise price per share	\$0.39	\$0.39		
Risk-free interest rate	2.51	%	2.56%-2.86	%
Expected holding period	1.82-4.82		2.5-5.5	
Expected volatility	70	%	70	%
Expected dividend yield	-		-	

The significant assumptions using the Monte Carlo Simulation approach for valuation of the Warrants are:

- (i) *Risk-Free Interest Rate.* The risk-free interest rates for the Warrants are based on U.S. Treasury constant maturities for periods commensurate with the remaining expected holding periods of the warrants.
Expected Holding Period. The expected holding period represents the period of time that the Warrants are expected to be outstanding until they are exercised. The Company utilizes the remaining contractual term of the Warrants at each valuation date as the expected holding period.
- (ii) *Expected Volatility.* Expected stock volatility is based on daily observations of the Company's historical stock values for a period commensurate with the remaining expected holding period on the last day of the period for which the computation is made.
- (iii) *Expected Dividend Yield.* Expected dividend yield is based on the Company's anticipated dividend payments over the remaining expected holding period. As the Company has never issued dividends, the expected dividend yield is 0% and this assumption will be continued in future calculations unless the Company changes its dividend policy.
- (iv) *Expected Probability of a Fundamental Transaction.* The possibility of the occurrence of a Fundamental Transaction triggering a Put right is extremely remote. As discussed above, a Put right would only arise if a Fundamental Transaction (1) is an all cash transaction; (2) results in the Company going private; or (3) is a transaction involving a person or entity not traded on a national securities exchange. The Company believes such an occurrence is highly unlikely because:
 - a. The Company only has one product that is FDA approved but is currently not available for commercial sales;
 - b. The Company will have to perform additional clinical trials for FDA approval of its flagship product;
 - c. Industry and market conditions continue to include a global market recession, adding risk to any transaction;
 - d. Available capital for a potential buyer in a cash transaction continues to be limited;
 - e. The nature of a life sciences company is heavily dependent on future funding and high fixed costs, including Research & Development;
 - f. The Company has minimal revenues streams which are insufficient to meet the funding needs for the cost of operations or construction at their manufacturing facility; and
 - g. The Company's Rights Agreement and Executive Agreements make it less attractive to a potential buyer.

With the above factors utilized in analysis of the likelihood of the Put's potential Liability, the Company estimated the range of probabilities related to a Put right being triggered as:

Range of Probability	Probability
Low	0.5 %
Medium	1.0 %
High	5.0 %

The Monte Carlo Simulation has incorporated a 5.0% probability of a Fundamental Transaction to date for the life of the securities.

(vi) *Expected Timing of Announcement of a Fundamental Transaction.* As the Company has no specific expectation of a Fundamental Transaction, for reasons elucidated above, the Company utilized a discrete uniform probability distribution over the Expected Holding Period to model in the potential announcement of a Fundamental Transaction occurring during the Expected Holding Period.

(vii) *Expected 100 Day Volatility at Announcement of a Fundamental Transaction.* An estimate of future volatility is necessary as there is no mechanism for directly measuring future stock price movements. Daily observations of the Company's historical stock values for the 100 days immediately prior to the Warrants' grant dates, with a floor of 100%, were utilized as a proxy for the future volatility.

(viii) *Expected Risk-Free Interest Rate at Announcement of a Fundamental Transaction.* The Company utilized a risk-free interest rate corresponding to the forward U.S. Treasury rate for the period equal to the time between the date forecast for the public announcement of a Fundamental Transaction and the Warrant expiration date for each simulation.

(ix) *Expected Time Between Announcement and Consummation of a Fundamental Transaction.* The expected time between the announcement and the consummation of a Fundamental Transaction is based on the Company's experience with the due diligence process performed by acquirers, and is estimated to be six months. The Monte Carlo Simulation approach incorporates this additional period to reflect the delay Warrant Holders would experience in receiving the proceeds of the Put.

While the assumptions remain consistent from period to period (e.g., utilizing historical stock prices), the numbers input change from period to period (e.g., the actual historical prices input for the relevant period). The carrying amount and estimated fair value of the above Warrants was approximately \$1,161,000 and \$962,000 at December 31, 2018 and 2017, respectively.

The Company applies FASB ASC 820 (formerly Statement No. 157 *Fair Value Measurements*) that defines fair value, establishes a framework for measuring fair value in generally accepted accounting principles, and expands disclosures about fair value measurements. The guidance does not impose any new requirements around which assets and liabilities are to be measured at fair value, and instead applies to asset and liability balances required or permitted to be measured at fair value under existing accounting pronouncements. The Company measures its warrant liability for those warrants with a cash settlement feature at fair value.

FASB ASC 820-10-35-37 (formerly SFAS No. 157) establishes a valuation hierarchy based on the transparency of inputs used in the valuation of an asset or liability. Classification is based on the lowest level of inputs that is significant to the fair value measurement. The valuation hierarchy contains three levels:

Level 1 – Quoted prices are available in active markets for identical assets or liabilities at the reporting date. Generally, this includes debt and equity securities that are traded in an active market.

Level 2 – Observable inputs other than Level 1 prices such as quote prices for similar assets or liabilities; quoted prices in markets that are not active; or other inputs that are observable or can be corroborated by observable market data for substantially the full term of the assets or liabilities. Generally, this includes debt and equity securities that are not traded in an active market.

Level 3 – Unobservable inputs that are supported by little or no market activity and that are significant to the fair value of the assets or liabilities. Level 3 assets and liabilities include financial instruments whose value is determined using pricing models, discounted cash flow methodologies, or other valuation techniques, as well as instruments for which the determination of fair value requires significant management judgment or estimation. As of December 2018, the Company has classified the warrants with cash settlement features and a convertible note payable as Level 3. Management evaluates a variety of inputs and then estimates fair value based on those inputs. As discussed above, the Company utilized the Monte Carlo Simulation Model in valuing the warrants and the convertible note.

Edgar Filing: HEMISPHERX BIOPHARMA INC - Form 10-K

The table below presents the balances of assets and liabilities measured at fair value on a recurring basis by level within the hierarchy as:

	(in thousands)			
	As of December 31, 2018			
	Total	Level 1	Level 2	Level 3
Assets:				
Marketable securities	\$1,526	\$1,526	\$-	\$-
Liabilities:				
Convertible note payable	\$3,408	-	-	\$3,408
Redeemable warrants	\$1,061	-	-	\$1,061

F-29

(in thousands)
As of December 31, 2017

Total	Level 1	Level 2	Level 3
-------	------------	------------	------------

Assets:

Marketable Securities	\$695	\$695	\$-	\$-
-----------------------	-------	-------	-----	-----

Liabilities:

Redeemable warrants	\$962	-	-	\$962
---------------------	-------	---	---	-------

The changes in Level 3 Liabilities measured at fair value on a recurring basis are summarized as follows (in thousands):

Redeemable warrants:

Balance at December 31, 2017	\$962
Warrants exercised and cancelled	(222)
Warrants issued	1,486
Fair value adjustments	(1,165)
Balance at December 31, 2018	\$1,061

Convertible debt:

Balance at December 31, 2017	\$-
Note amount	3,170
Deferred debt discounts	(344)
Fair value adjustments	582
Balance at December 31, 2018	\$3,408

(19) Financing Obligation Arising from Sale Leaseback Transaction

On March 16, 2018, the Company sold land and a building for \$4,080,000 and concurrently entered into an agreement to lease the property back for ten years at \$408,000 per year for two years through March 31, 2020. The lease payments will increase 2.5% per year for the next three years through March 31, 2023 and the lease payments will increase 3% for the remaining five years through March 31, 2028. The sale of the property includes an option to repurchase the property at fair value which does not permanently transfer all the risks and rewards of ownership to the buyer. The option to repurchase the property also would be at a higher price than the sales price and is considered likely based upon the Company's plans going forward. Because the sale of the property includes the option to repurchase the property and includes the above attributes, the transaction was accounted for as a financing transaction whereby the Company debited cash for the amount of cash received and credit financing obligation. The Company will continue to report the property as an asset and the property will continue to be depreciated. The fair value repurchase option is accounted for similar to a share appreciation mortgage. Accordingly, the guidance in ASC 470-30 related to participating mortgage loans would be applied to the liability. If the option expires unused, the sale is

recognized at that time. The gain on the sale would be the excess of the liability (current fair value of the property) over its carrying amount. If the option is exercised, the cash payment by the seller-lessee is to pay off the financing obligation. As part of the sale of this building, warrants were provided to the buyer for the purchase of up to 3,225,806 shares of Company common stock for a period of five years at an exercise price of \$0.3875 per share, 125% of the closing price of the common stock on the NYSE American on the date of execution of the letter of intent for the purchase. The warrants cannot be exercised to the extent that any exercise would result in the purchaser owning in excess of 4.99% of our issued and outstanding shares of common stock.

The Property and equipment in “Note 7 Stockholders’ Equity” above are the property and equipment involved in this transaction. Depreciation on the building will continue until a sale has been recognized.

Future minimum payments required under the Financing Obligation and the balance of the Finance Obligation as of December 31, 2018, are as follows:

During the year:

	(in thousands)
2019	\$ 408
2020	417
2021	427
2022	438
2023	450
Thereafter	2,025
Total of payments	4,165
Less deferred issuance costs	(245)
Less discount on debt instrument	(1,054)
Less imputed interest	(349)
Total balance	2,517
Less current portion	199
Long term portion	\$ 2,318

Interest expense relating to this financing agreement was \$61,000 for the year ended December 31, 2018.

(20) Subsequent Events

A registration statement relating to the rights offering was filed with the U.S. Securities and Exchange Commission (“SEC”). Under the rights offering, Hemispherx distributed to its holders of common stock and to holders of certain options and warrants as of February 14, 2019, at no charge, one non-transferable subscription right for each share of common stock held or deemed held on the record date. Each right entitled the holder to purchase one unit, at a subscription price of \$1,000 per unit, consisting of one share of Series B Convertible Preferred Stock with a face value of \$1,000 (and immediately convertible into common stock at an assumed conversion price of \$0.20) and 5,000 warrants with an assumed exercise price of \$0.20. The warrants will be exercisable for five (5) years after the date of issuance. The funds realized from the offering were approximately \$4.69 million. The \$4.69 million received from the rights offering increased our net equity by this amount on the date the funds were received.

On March 13, 2019, the Company and Iliad Research and Trading, L.P. (the “Lender”) amended the September 28, 2018 Secured Convertible Promissory Note (the “Convertible Note”) issued to the Lender. Pursuant to the amendment, the maturity date of the Convertible Note was extended from September 28, 2019 to September 28, 2020. In addition, the conversion and redemption rates were revised to a rate to be mutually agreed to by the Company and the Lender. In the event that the Company and the Lender are unable to reach agreement on such rate, the Company will be required to pay the redemption or conversion in cash. The amount of the Convertible Note that the Lender is entitled to redeem is limited to a maximum of \$300,000 per calendar month.

The price per share of the Company common stock has closed at or below \$0.20 since February 26, 2019 and most recently closed on March 26, 2019 at \$0.16, with a 30 day average of \$0.19. On March 26, 2016, the Company received written notice from the NYSE American LLC (the “NYSE American”) that the Company is not in compliance with the continued listing standards set forth in Section 1003(f)(v) of the NYSE American Company Guide because the common stock has been selling for a low price per share for a substantial period of time. The NYSE American has determined that the continued listing of the Company’s common stock is predicated on the Company effecting a reverse stock split of the common stock or otherwise demonstrating sustained price improvement within a reasonable period of time. the Company has until September 26, 2019 to demonstrate compliance.

The Company evaluated subsequent events through the date on which these financial statements were issued and determined that no subsequent event, other than the above, constituted a matter that required adjustment to the financial statements.

