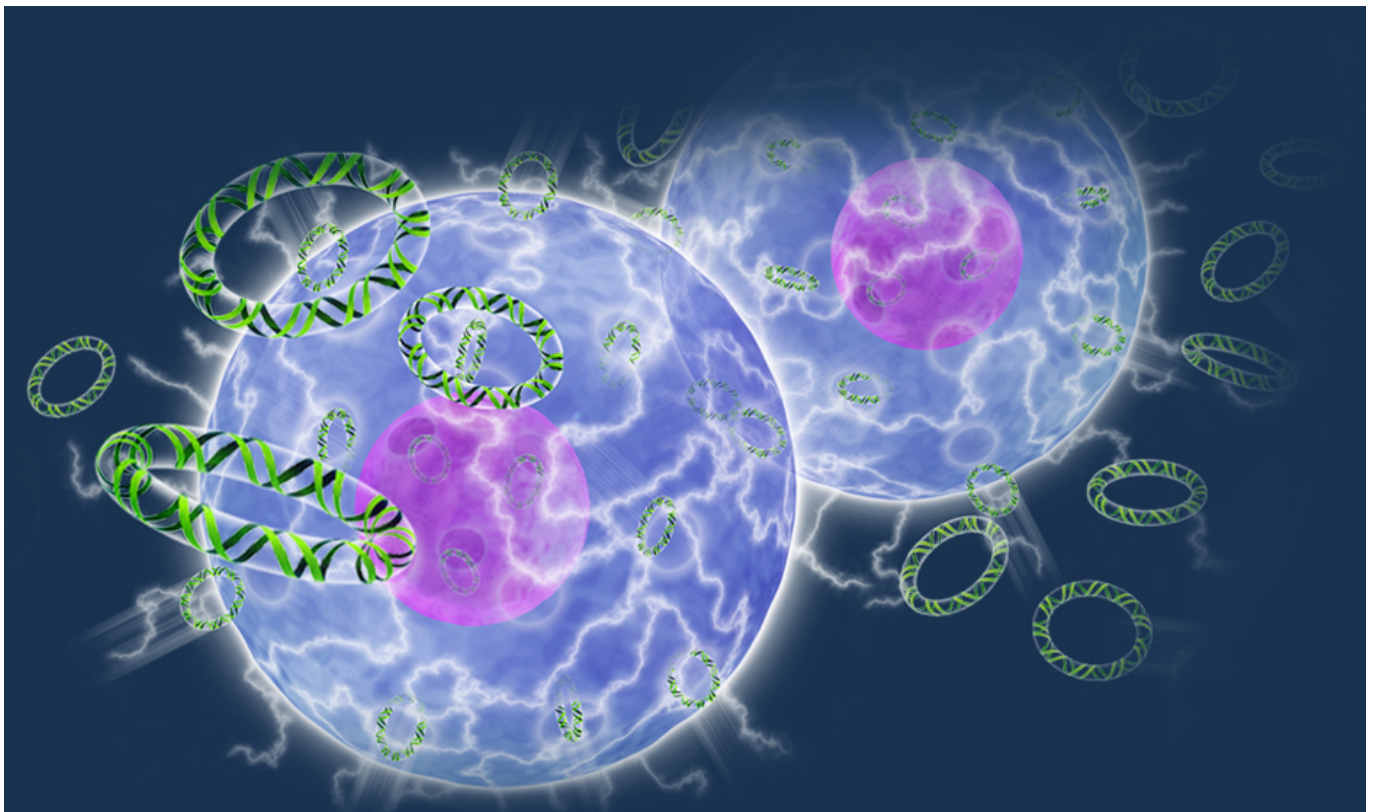


Inovio Shrink Prostate Tumours and Protect Against Lethal Antibiotic-Resistant Bacterial Infection

Inovio Pharmaceuticals, Inc. {NASDAQ: INO} today announced that the development of its DNA-based monoclonal antibody program received a boost from two peer-reviewed scientific papers that demonstrate their impact on prostate tumours, and in preventing infection from a pneumonia-causing bacteria in preclinical studies.



Inovio dDNA® Constructs Shield Prostate Tumors and Protect Against Lethal Antibiotic-Resistant Bacterial Infection in Preclinical Studies

Inovio's dDNA-based Monoclonal Antibody Program
Summary of Scientific Achievements
PUNNETT, RICHMOND, PA – September 21, 2017 – Inovio Pharmaceuticals, Inc. (NASDAQ: INOV) today announced that the development of its dDNA-based monoclonal antibody program resulted in a breakthrough from two years' research. Recent scientific papers that demonstrate their impact on prostate tumors and on preventing infection from a pneumonia-causing bacteria in preclinical studies.

Dr. J. Joseph Kim, Inovio's President and CEO, said, "Many of the top-selling drugs on the market today are monoclonal antibodies, and our dDNA products are the first that can easily be genetically designed and in cost production. In fact, we are advancing our first dDNA product, an engineered bacteria product funded by dDNA – into human testing next year, paving the path for commercializing these new products to treat cancer and infection. These two newly published studies further support that Inovio's patent dDNA platform could be expanded to target cancer and bacterial diseases along with viral infectious diseases. Our dDNA program represents a new application of our patent dDNA technology platform to develop valuable new treatments for cancer and infection."

An article in the journal *Cancer Immunology, Immunotherapy* described how the dDNA constructs against prostate specific membrane antigen (PSMA) produced monoclonal antibodies that attack prostate tumors in a preclinical animal model. The article is entitled, "Novel cancer immunotherapy with dDNA encoded anti-prostate-specific membrane antigen monoclonal antibody," by Inovio researchers and collaborators.

This research publication is significant because it is the first to report on the use of Inovio dDNA technology to develop novel monoclonal antibody-based therapies against cancer targets. This new dDNA construct encodes an engineered synthetic DNA to encode therapeutic monoclonal antibody targeted to bind PSMA. When delivered directly into the body, the genetic instructions provided from the dDNA construct enable the patient's own cells to become the factory which manufactures the therapeutic monoclonal antibody products. The PSMA dDNA antibody was expressed in high levels directly in mice and successfully targeted human prostate cancer cells grown in transgenic mice. Inovio has previously published research papers demonstrating its dDNA platform's ability to treat multiple viral targets such as flu, dengue, chikungunya, and HIV.

The anti-cancer dDNA products work by binding the antigens on the cancer cells and killing them by an antibody dependent cytotoxicity (ADCC) mechanism in conjunction with natural killer (NK) cells. These results indicate the potential clinical utility of the dDNA therapeutic strategy against prostate cancer as well as other cancers. The data is especially interesting since the anti-cancer dDNA product utilizes an NK cell-killing mechanism which is distinct from the killer T cell mechanism generated by Inovio's cancer vaccine products like DND-510 and DND-5401.

In another first, Inovio also published results showing its dDNA constructs targeting antibiotic-resistant bacteria protected mice when challenged with a lethal dose of drug-resistant pneumonia – a pneumonia-causing bacteria. Pneumonia infections usually occur in people in the hospital or with weakened immune systems. The CDC reports each year that more than two million American acquire antibiotic-resistant infections that are becoming increasingly more difficult and costly to treat. The bacteria can cause infections of the blood, pneumonia, and infections following surgery that can lead to severe illness and death.

To design an effective treatment, Inovio engineered dDNA constructs encoding the pneumonia antigen. The constructs were protection against lethal pneumonia, prevented severe lung pathology and even exhibited enhanced protective activity when combined with antibiotics. This paper is also significant for Inovio to demonstrate that dDNA constructs could be successfully designed to generate isotype-specific monoclonal antibodies. Isotype-specific monoclonal antibodies are engineered against specific antigens which could simultaneously bind two different antigens, and this continues to be very difficult to achieve using traditional protein-based monoclonal technology. These results further support the sophisticated capabilities built into Inovio dDNA/RN technology. Results from this study were outlined in Nature Communications, in an article entitled, "One delivery of nose- and engineered biologic monoclonal antibodies protect against multiple resistant *Pseudomonas aeruginosa*," authored by Inovio and its collaborators at The Walter Institute and Hoffmann, Hoffmann's clinical biologist research and development arm.

Infused with over 400 million in R&D support from the agencies like DARPA, NIH, and the Gates Foundation, Inovio dDNA products could extend the medical benefits that monoclonal antibodies have already achieved, and even potentially address diseases that conventional monoclonal antibodies cannot. For instance in immunomodulation, Inovio is already developing its dDNA as well as other Checkpoint inhibitors in its cancer dDNA portfolio.

Even though conventional monoclonal antibodies represent one of the most successful segment of biotechnology market, accounting for over \$50 billion in sales today, they are still costly and time consuming to develop, produce and study. They are manufactured outside the body in bioreactors, typically requiring costly large-scale manufacturing facility development and laborious production. Inovio's disruptive dDNA technology has the potential to overcome these limitations by virtue of their simplified design, rapidity of development, product stability, ease of manufacturing and degradable, and cost effectiveness, thereby providing potential, no answer for treating a range of diseases.

About Inovio's dDNA-based Monoclonal Antibody Platform
Funded by over \$400 million in R&D support from the last 3 years, Inovio has rapidly advanced this transformative approach in a systemic way. Earlier this year Inovio reported its Influenza dDNA product broadly cross-reactive antibodies that provided complete protection from a lethal challenge with multiple strains from both influenza A and B types. The flu dDNA was combined as earlier with flu and dDNA products for other viral targets including HIV, dengue, and chikungunya. Together with its pre-clinical studies demonstrating the potential of Inovio dDNA platform to cancer and bacterial diseases, these results fully demonstrate the potential of this platform to target multiple medical markets.

The significant advancement seen in Inovio dDNA technology is that the optimized genes for a desired monoclonal antibody is encoded in a dDNA plasmid, which is produced using any cost-effective and highly scalable fermentation techniques. These plasmids are delivered directly into cells of the body using electroporation and the encoded monoclonal antibody is then directly produced by these cells. Previously published studies show that a single administration of a highly optimized dDNA-based monoclonal antibody targeting HIV virus produced a high level of expression of the antibody in the bloodstream of mice. Inovio recently reported data showing that dDNA products against flu, Ebola, chikungunya and dengue protected animals against lethal challenge. Inovio dDNA dDNA product is being developed under a grant from the Defense Advanced Research Projects Agency (DARPA).

About Inovio Pharmaceuticals, Inc.
Inovio is taking immunotherapy to the next level in the fight against cancer and infectious diseases. We are the only immunotherapy company that has reported generating T cells in vivo in high quantity that are fully functional and whose killing capacity correlates with relevant clinical outcomes with a favorable safety profile. With an expanding portfolio of cancer therapies, the company is advancing a growing preclinical and clinical stage cancer pipeline. Partners and collaborators include Moderna, Regeneron, Genentech, Novartis, Novartis, Sanofi, University of Pennsylvania, DARPA, Sanofi Life Sciences, Plasmid Life Sciences, Apollonia Corporation, Brown University, NIH, DARPA, Precision Trial Network, National Cancer Institute, U.S. Military HIV Research Program, and Lowell University.

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This press release contains certain forward-looking statements relating to our business, including our plans to develop electroporation-based drug and gene delivery technologies and dDNA vaccines and dDNA products, our expectations regarding our research and development programs, including the planned initiation and conduct of clinical trials and the availability and timing of data from these trials, and the sufficiency of our capital resources. Actual results or results may differ from the expectations set forth herein as a result of a number of factors, including uncertainties inherent in clinical studies, clinical trials and product development programs, the development of Inovio dDNA products, the availability of funding to support continuing research and studies in an effort to prove safety and efficacy of electroporation technology as a delivery mechanism or develop viable dDNA vaccines, our ability to support our pipeline of systemic active immunotherapy and vaccine products, the ability of our collaborators to attract development and commercial collaboration for products or license any product sales that will enable us to recover from payments and royalties, the ability of our capital resources, the availability of potential availability of alternative therapies or treatments for the conditions targeted by the company or its collaborators, including alternatives that may be more efficacious or cost-effective than any therapy or treatment that the company and its collaborators seek to develop. Issues involving product liability, issues involving patents and whether they or licenses to them will provide the company with meaningful protection from others using the covered technologies, whether such proprietary rights are enforceable or enforceable in foreign or allegedly foreign or rights of others or on intellectual claims or invalidity and whether the company or license or licensees may be required to defend them, the level of corporate expenditures, assessments of the company's technology to potential corporate or other partners or collaborators, capital market conditions, the impact of government healthcare programs and other factors set forth in our Annual Report on Form 10-K for the year ended December 31, 2016, our Form 10-Q for the period ended June 30, 2017, our Form 10-K for the period ended June 30, 2017, and our Form 10-Q for the period ended June 30, 2017. There can be no assurance that any product candidates or Inovio's pipeline will be successfully developed, manufactured or commercialized. The final results of clinical trials will be supportive of regulatory approvals required to market licensed products, or that any of the forward-looking information provided herein will be proven accurate. In addition, the forward-looking statements included in this press release represent Inovio's view as of the date hereof. Inovio anticipates that subsequent events and developments may cause its view to change. However, while Inovio may attempt to update these forward-looking statements at some point in the future, the company specifically disclaims any obligation to do so, except as may be required by law. These forward-looking statements should not be relied upon as representing Inovio's views as of any date subsequent to the date of this release.