



Forward-Looking Statements

This presentation includes statements that are, or may be deemed, "forward-looking statements," within the meaning of Section 27A of the Securities Act of 1933, as amended. All statements, other than statements of historical facts, included in this presentation regarding our strategy, future operations, future financial position, future revenue, projected costs, prospects, plans and objectives of management are forward-looking statements. In some cases, you can identify forward-looking statements by terms such as "may," "might," "will," "objective," "intend," "should," "could," "can," "would," "expect," "believe," "anticipate," "project," "target," "design," "estimate," "predict," "opportunity," "proposition," "strategy," "potential," "plan" or the negative of these terms and similar expressions intended to identify forward-looking statements.

You should not place undue reliance on these forward-looking statements. Forward-looking statements include, but are not limited to, statements about: the timing and success of preclinical studies and clinical trials; the ability to obtain and maintain regulatory approval of our product candidates; the scope, progress, expansion and costs of developing and commercializing our product candidates; our expectations regarding the amount and timing of our expenses and revenue; the sufficiency of our cash resources, plans for the use of our cash resources and needs for additional financing; our ability to adequately manufacture our product candidates; our ability to obtain and maintain intellectual property protection for our product candidates; our expectations regarding competition; the size and growth of the potential markets for our product candidates and the ability to serve those markets; the rate and degree of market acceptance of any of our product candidates; our anticipated growth strategies; the anticipated trends and challenges in our business and the market in which we operate; our ability to establish and maintain development partnerships; our ability to attract or retain key personnel; our expectations regarding federal, state and foreign regulatory requirements; regulatory developments in the United States and foreign countries and other factors that are described in the "Risk Factors" and "Management's Discussion and Analysis of Financial Condition and Results of Operations" sections of our Annual Report on Form 10-K for the year ended December 31, 2019, which has been filed with the Securities and Exchange Commission (SEC) and is available on the SEC's website at <u>www.sec.gov</u>.

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Powering a New Decade of DNA Medicines

Precisely Designed Plasmids Delivered Through Proprietary Smart Device

Safe and Robust Immune Responses in More Than 2,000 Patients

In Vivo Immune Responses for "Off-the-Shelf" Speed, Efficiency

Extensive Patent Portfolio Protecting Technology Platform FIRST DNA Medicine in Phase 3 Clinical Trials (VGX-3100) for Precancerous Cervical Dysplasia

> FIRST to Show Clearance of High-Risk HPV 16/18 in Phase 2b Trial (VGX-3100)

FIRST to Show Complete Response in Phase 1 w/2 PD-1s for Head and Neck Cancer (MEDI0457)

FIRST dMAb[™] Plasmid in Phase 1 for Zika (INO-A002)



INOVIO Vision to Build the Leading DNA Medicine Company

Founding Vision

- Create precisely designed plasmids that target antigens to address urgent medical needs
- Develop proprietary device to deliver plasmid safely in vivo directly into the cell to produce robust immune response
- Build scientific, medical, and commercial team and outstanding partnerships to drive value

Near-Term Execution

- Rapidly bring to market precisely designed DNA medicines to potentially treat and prevent diseases associated with HPV, cancer, and infectious diseases
- Maximize value of lead candidates worldwide

Long-Term Strategy

- Create new market in safe, effective
 DNA medicines
- Aggressively seek partners to ensure DNA medicines reach patients in need
- Be capital efficient
 - \$89.5M cash/investments as of last reported earnings (12/31/19)
 - \$208.2M net proceeds from new financing in 1Q20



Vision Built on INOVIO Proprietary Technology

OPTIMIZED PLASMID DESIGN AND DELIVERY TECHNOLOGY

PRECISELY DESIGNED PLASMIDS (SynCon®)



PROPRIETARY SMART DEVICE (CELLECTRA®)

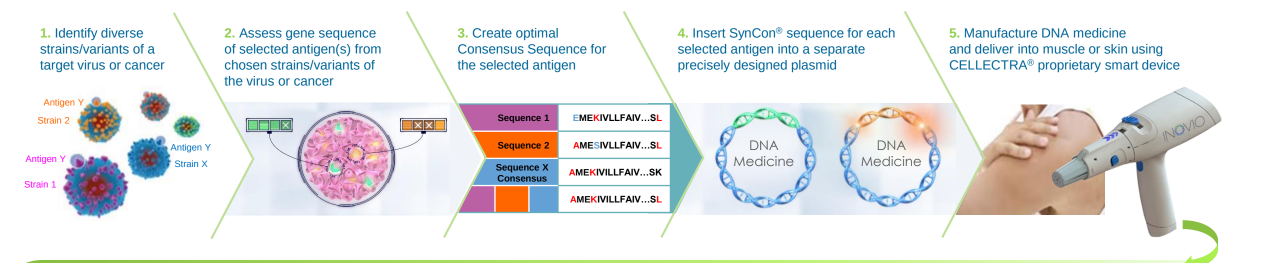




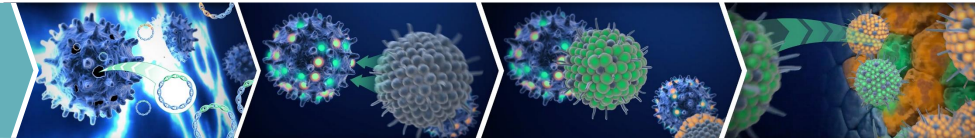


INOVIO Technology – Powering Potent Antigen Specific Immune Responses

INOVIO DNA medicines power a patient's immune system to generate functional antibodies and killer T cells *in vivo* to fight cancer and infectious disease



6. Protective antibodies and killer T cells produced by immune system against diverse strains of a virus or cancer





CELLECTRA® 5PSP – INOVIO's First Commercial Smart Device

CELLECTRA® 5PSP

- World's first commercial smart device for DNA medicine – CE Marking in Europe
- Proprietary smart device currently used in Phase 3 trials
- Simplified interaction and automated injection using prefilled cartridge
- Disposable single use array which includes used drug cartridge
- Touch screen interface, automated sensors and trigger start
- Records data file for post-treatment review
- Data files can be downloaded from system and uploaded to web-based interface
- Several rounds of Usability Testing that refined development











INOVIO's Technology Advantages

Clinical Efficacy

- Demonstrated clinical efficacy in Phase 2b study
- Lead candidate VGX-3100 in Phase 3 evaluation for cervical dysplasia

Safety

- Favorable safety profile tested in over 2,000 patients in over 6,000 administrations
- Carries no potential toxicity from plasmid vector

Versatility and Boosting

- Targets virtually any antigenic sequence; combining multi-antigens into single vial
- Initiated first-in-human study of optimized dMAb[™] plasmid
- No anti-vector response allows for effective boosting

Rapid and Scalable Manufacturing

- "Off-the-shelf" product; no frozen storage issues (room temp storage >1 yr.)
- Rapid development from concept to human in <3 months (COVID-19 vaccine)
- Relatively inexpensive to manufacture; produce large quantities



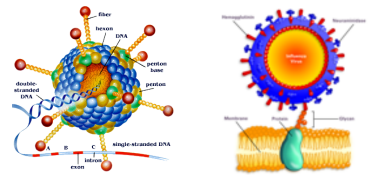
Limitations of Other Approaches

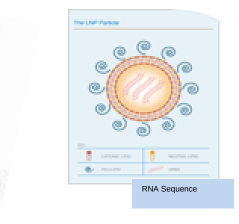
Viral Vectors – Receptor/cell target based mediated entry

- Systemic delivery/local injection
- Preexisting or induced immunity is an issue
- Biologic variability of take
- Immune bias tuned by vector
- Hard to re-administer/tissue tropism limits and positives

RNA – LNP/nanoparticle delivery dependent

- Systemic delivery, localized expression (liver>lung or spleen)
- Process for manufacture and release work in progress
- Formulations + RNA follow tissue targeting of the particles/cold chain required, include focus on IV route
- DLT observed, low CTL induced, inflammatory
- High cost of goods







INOVIO DNA Medicines Pipeline

PRODUCT	INDICATION	ANTIGEN	PRECLINICAL	PHASE 1	PHASE 2	PHASE 3	PARTNER/COLLABORATOR/FUNDE
HPV-TARGETED							
	Cervical HSIL	HPV 16 E6, E7/ HPV 18 E6, E7					_
VGX-3100	Vulvar HSIL						Apollobio
	Anal HSIL						(China only)
INO-3107	Recurrent Respiratory Papillomatosis (RRP)	HPV 6 E6, E7/ HPV 11 E6, E7					
		,					
	Head & Neck Cancer	HPV 16 E6, E7/					
MEDI0457	Cervical, Anal, Penile,	HPV 18 E6, E7					AstraZeneca
	Vulvar Cancers						

IMMUNO-ONCOLOGY (NON HPV-ASSOCIATED)

	Glioblastoma			
INO-5401	Multiforme (GBM)	WT1, PSMA, hTERT		REGENERON
INO-5151	Prostate Cancer	PSA, PSMA		CANCER RESEARCH INSTITUTE
				INSTITUTE

INTERNALLY FUNDED

EXTERNALLY FUNDED



INOVIO DNA Medicines Pipeline (Continued)

PRODUCT	INDICATION	ANTIGEN	PRECLINICAL	PHASE 1	PHASE 2	PHASE 3	PARTNER/COLLABORATOR/FUNDER
INFECTIOUS DISEAS	ES (NON HPV-ASSOCIATED)						
PENNVAX-GP	HIV	Gag, pol, env					NIAID HIV VACCINE
INO-4201	Ebola	Glycoprotein					DARPA
INO-4700 (GLS-5300)	MERS	Spike					
INO-4600 (GLS-5700)	Zika	Glycoprotein					GENE
INO-4500	Lassa Fever	Glycoprotein					CEPI
INO-4800	COVID-19 (Coronavirus)	Spike					CEPI () BILL&MELINDA GATES foundation

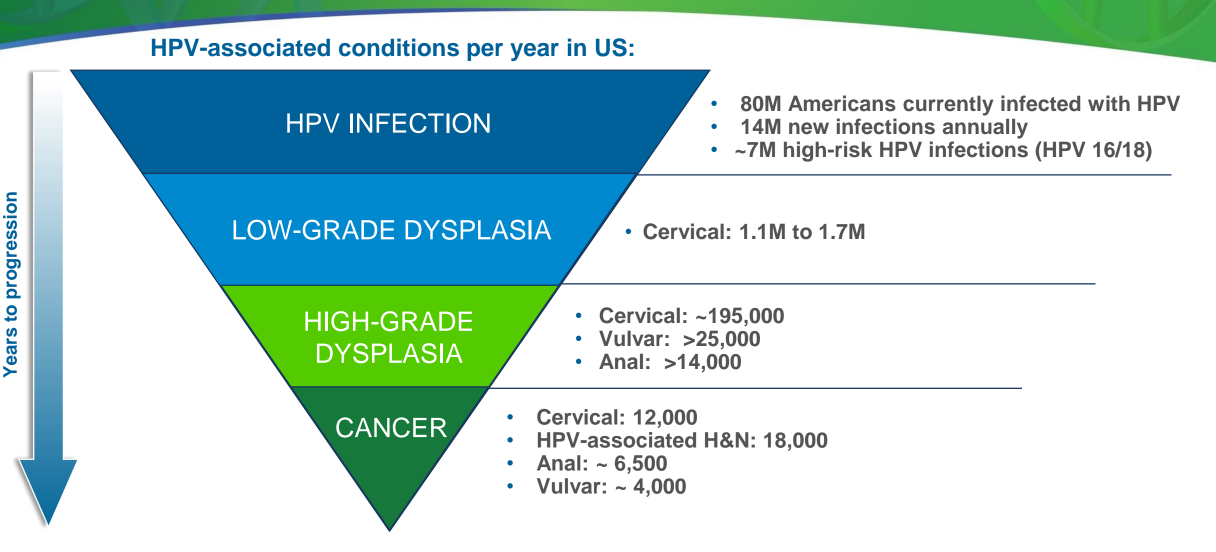
dMAb[™] (DNA-ENCODED MONOCLONAL ANTIBODIES)

INO-A002	Zika	Glycoprotein				BILL& MELINDA GATES foundation
			INTER	NALLY FUND	ED	EXTERNALLY FUNDED



HPV-Related Programs

HPV-Associated Diseases Market Overview

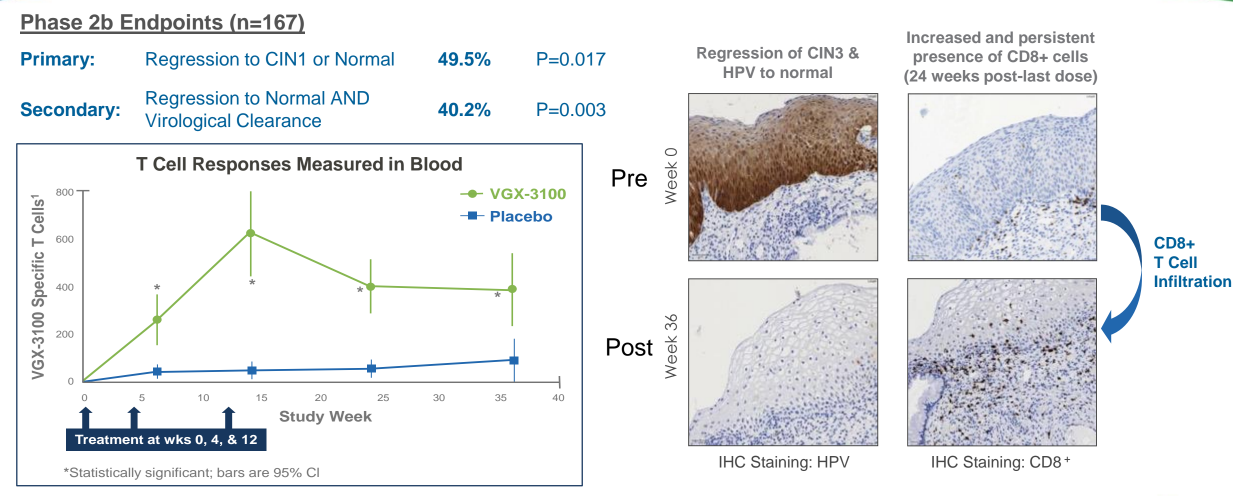


Sources: US CDC (2018) HPV and Cancer, available at: https://www.cdc.gov/cancer/hpv/statistics/cases.htm (accessed July 22, 2019); Saraiya M, Unger ER, Thompson TD, Lynch CF, Hernandez BY, Lyu CW, Steinau M, Watson M, Wilkinson EJ, Hopenhayn C, Copeland G, Cozen W, Peters ES, Huang Y, Saber MS, Altekruse S, Goodman MT; HPV Typing of Cancers Workgroup. US assessment of HPV types in cancers: implications for current and 9-valent HPV vaccines. J Natl Cancer Inst. 2015 Apr 29;107(6):djv086; Inovio Pharmaceuticals, internal estimates from published data (2015-16, 2017-18); US CDC, personal communication (2015); NCI SEER Cancer Stat Facts: Cervix Uteri, Vulvar, and Anal Cancers – https://seer.cancer.gov/statfacts (accessed 2017-18); 'User Initial Visits to Physicians' Offices, United States, 1966-2014. Fig. 47; Schiffman M, Solomon D. Findings to date from the ASCUS-LSIL Triage Study (ALTS). Arch Pathol Lab Med. 2003 Aug;127(8):946-9; US CDC. Genital HPV Infection – Fact Sheet.



13

Published VGX-3100 Phase 2b Study Achieves All Primary and Secondary Endpoints





VGX-3100 Phase 3 Program: HPV-Associated Cervical HSIL/ Precancerous Dysplasia

TRIAL: VGX-3100

- Targets HPV 16/18 subtypes; E6/E7 oncogenes
- Treats high-grade squamous intraepithelial lesions (HSIL)

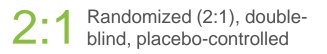


Phase 3 consists of 2 studies in parallel:

REVEAL1 (primary) n=198 – Enrollment Closed Study follow-up through week 88 (as in P2b) Topline efficacy data expected by 4Q 2020 **REVEAL2 (confirmatory) n=198 – Now Enrolling** Study follow-up through week 40

FIRST treatment for HPV infection of the cervix

FIRST non-invasive treatment for cervical pre-cancer Primary endpoint: Regression of HSIL (CIN2/3) AND clearance of HPV 16/18 in the cervix





Dosing: month 0, 1, 3 (as in P2b)



Primary endpoint measured at month 9 (as in P2b)



VGX-3100 Phase 2 Studies in HPV-Associated Vulvar and Anal HSIL/Precancerous Dysplasias

TRIALS: VGX-3100

- Target HPV 16/18 subtypes; E6/E7 oncogenes
- Treat high-grade squamous intraepithelial lesions (HSIL)

Precancerous Vulvar Dysplasia:



Phase 2 open-label study



33 women enrolled Interim data reported for 10

Interim findings (6 months after start of treatment) Decrease in lesion area: 80% of patients Resolution of vulvar dysplasia: 20% of patients

Non-detectability of HPV 16/18: 20% of patients

Precancerous Anal Dysplasia:



Phase 2 open-label study

Interim findings (6 months after start of treatment) Clearance of lesions: 50% of patients ∯ x23

23 patients enrolled Interim data reported for 20

Decrease in number of lesions: 75% of patients



INOVIO and QIAGEN Developing Biomarker to Optimize Patient Selection





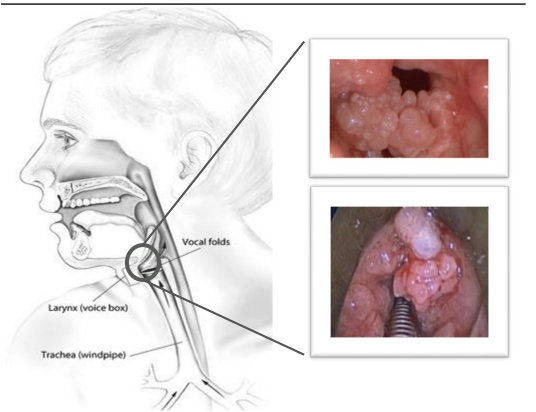
In 2Q 2019, INOVIO entered into collaboration with QIAGEN to co-develop a liquid biopsy-based pretreatment commercial test kit to guide patient selection for VGX-3100:

- Aimed to produce an accurate test that would increase absolute efficacy of VGX-3100 among HPV-infected women who have progressed to Cervical HSIL (pre-cancer)
- Commercialization of a CDx test concurrently with VGX-3100 could enhance market adoption of this first-in-class DNA medicine



Recurrent Respiratory Papillomatosis (RRP) Caused by HPV 6 and 11

Areas affected by Recurrent Respiratory Papillomatosis (RRP)



- Rare, orphan disease with ~15,000 total active cases within the U.S., where virtually all of those require surgical procedures
 - ~6,000 new cases per yr. in the U.S.
- HPV-associated disease; caused by HPV 6 and 11
- Growths can lead to life-threatening airway obstructions
- SoC is lifelong surgery (repeated/multiple times a yr)
 - Currently, disease is incurable and can only be treated by surgery to remove tumors, which temporarily restores the airway
- RRP may occur in adults as well as in children who are thought to have contracted the virus during childbirth



Sources: RRP Foundation; Venkatesan et al. Otolaryngol Clin North Am 2013; Ivancic et al. Laryngoscope Investigative Otolaryngol; www.nidcd.nih.gov/health/recurrent-respiratory-papillomatosis; Derkay et al *Arch Otolaryngol Head Neck Surg* (1995); Armstrong et al *Arch Otolarygol Head Neck Surg* (1995); Marsico et al STDs (2014).

INO-3106 Pilot Study in RRP – Completed

TRIAL: INO-3106 (for HPV 6-caused RRP)



Phase 1 pilot, single-site, clinical study



Enrolled 2 adult patients with RRP, HPV 6+



4 doses of vaccine, 3 weeks apart on Day 0, Weeks 3, 6, 9



CELLECTRA-delivered INO-3106 (only for HPV 6) plasmid encoded antigens

Two RRP patients had prior surgeries every 6 months After receiving 4 doses, 1 patient has gone >915 days without surgery, and the second went 584 days without surgery Planning potential registrational study of INO-3107 (for both HPV 6 and 11) by 1H 2020



Published in Vaccines (MDPI); entitled "Immune Therapy Targeting E6/E7 Oncogenes of Human Papillomavirus Type 6 (HPV-6) Reduces or Eliminates the Need for Surgical Intervention in the Treatment of HPV-6 Associated Recurrent Respiratory Papillomatosis"; January 23, 2020.

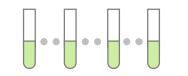
INO-3107 Phase 1/2 Study in RRP – IND Accepted

TRIAL: INO-3107 (for HPV 6 and/or 11-caused RRP)



Phase 1/2 openlabel, multicenter clinical study **₽ x63**

Target enrollment



4 doses of vaccine, 3 weeks apart on Day 0, Weeks 3, 6, 9

CELLECTRA-delivered INO-3107 plasmid encoded antigens

Enrollment criteria: Subjects who have required at least two surgical interventions per year for the past three years for the removal of associated papilloma(s)

Primary endpoint: A doubling or more in the time between surgical interventions following the first dose of INO-3107 relative to the frequency prior to study therapy



MEDI0457 for HPV-Related Cancers in Partnership with AstraZeneca



- **MEDI0457** (formerly INO-3112) = VGX-3100 + INO-9012 (IL-12 plasmid)
- In 2015, AstraZeneca acquired exclusive rights to MEDI0457
 - \$27.5M upfront
 - ~\$250M in potential development and commercial milestones
 - Double-digit tiered royalties on MEDI0457 sales
- AstraZeneca is evaluating MEDI0457 in combination with its PD-L1 checkpoint inhibitor, durvalumab, in HPV-associated cancers



MEDI0457 Potential to Treat Head and Neck Cancer Demonstrated in Phase 1 Trial



SCA

Primary: Safety and tolerability of DNA based immunotherapy
Secondary: Cellular and humoral immune responses
Exploratory: Anti-tumor

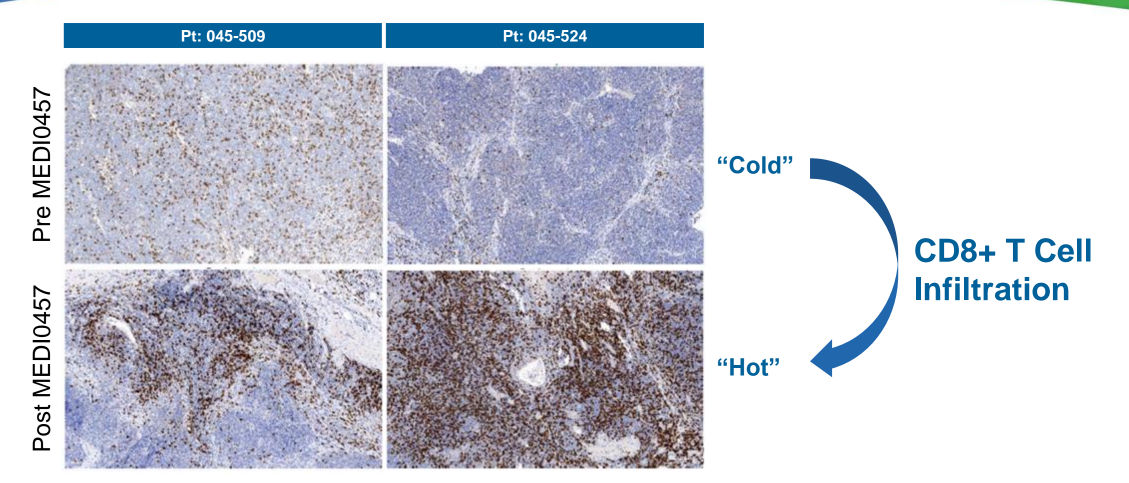
response and progression free survival



MEDI0457: 6 mg of VGX-3100 + 1 mg of INO-9012

In Cohort 1, if time allows, up to 2 treatments can be administrated prior to surgery, but total 4 treatments are scheduled

CD8+ T Cell Infiltration into Tumor Following MEDI0457 Treatment

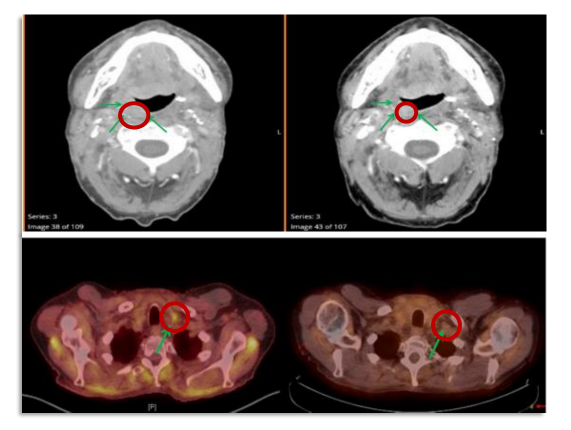


Robust antigen-specific CD8+ killer T cell responses observed in 20/22 - 90.1% – of patients (both tumor tissue and peripheral blood)



Aggarwal et al Clinical Cancer Research 2018

MEDI0457 Phase 1 Study Demonstrates Complete Response



- (Top image) CT neck with IV contrast demonstrating partial response pre- and 6 weeks post-nivolumab.
- (Bottom image) PET scan images pre- and 6 weeks post-nivolumab.

Phase 1 study of MEDI0457 (VGX-3100+IL-12) in 22 HPV+ H&N cancer patients

- Robust antigen-specific CD8+ killer T cell responses observed in 20/22 – 90.1% – of patients (both tumor tissue and peripheral blood)
- 4 progressed over several year period exhibiting recurrence with metastatic disease; treated with PD-1
- 2/4 (50%) show complete response to PD-1 therapy and remained tumor free for 2+ years
- 50% CR rate compares well in metastatic HPV+ H&N:
 - 4% CR rate (8/192) by KEYTRUDA alone
 - 3% CR rate (6/240) by OPDIVO alone
- AstraZeneca conducting Phase 2 studies combining MEDI0457 and durvalumab (PD-L1 inhibitor)



MEDI0457 for HPV-Associated Head & Neck Cancer in Phase 1b/2a in Partnership with AstraZeneca

TRIAL: MEDI0457 (VGX-3100 + IL-12) AstraZeneca



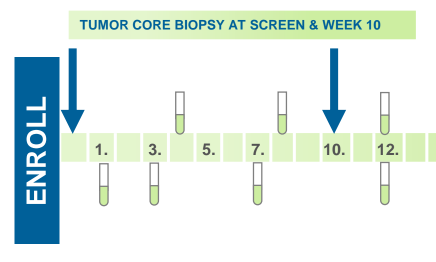
Phase 1b/2a open label study for metastatic HPV+ HNSCC with persistent or recurrent disease after chemotherapy treatment



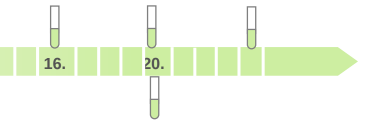
Combination with AstraZeneca's PD-L1 checkpoint inhibitor (durvalumab) Primary Endpoints: Safety, tolerability Secondary Endpoints: Immunogenicity, ORR, PFS, Disease CR, OS

x35

Completed enrollment of 35 subjects in August 2019



Planned Dosing Schedule: durvalumab at Weeks 4, 8 and 12 and then every 4 weeks until disease progression, unacceptable toxicity or withdrawal of consent



Planned Dosing Schedule: MEDI0457: Weeks 1, 3, 7 and 12 and then every 8 weeks until disease progression, unacceptable toxicity or withdrawal of consent



HPV-Related Clinical Program Overview

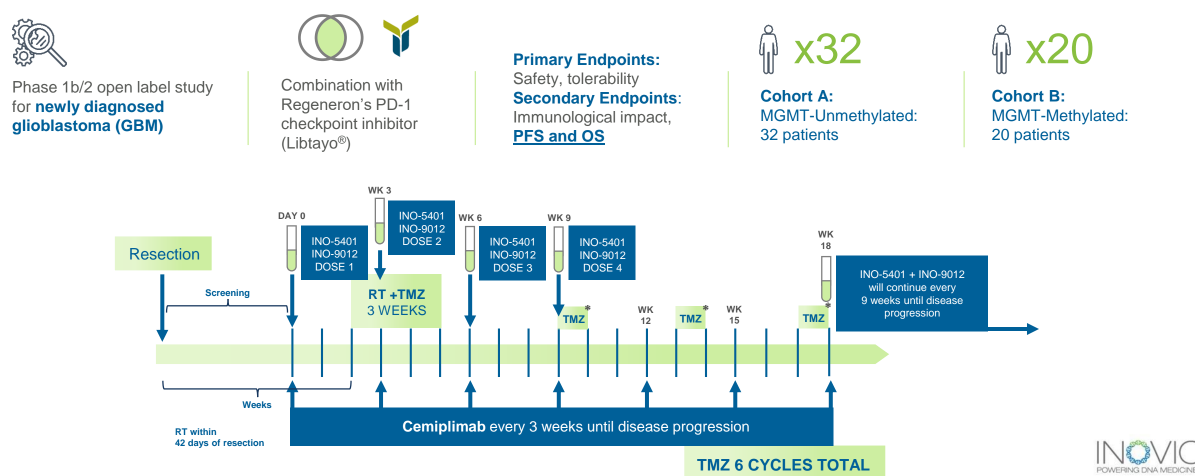
Precancerous Dysplasias (VGX-3100)	 Cervical dysplasia: Phase 2b PoC trial demonstrated a complete response in 43 out of 107 patients in regression of high-grade cervical lesions <i>and</i> elimination of HPV infection Vulvar dysplasia: Open-label Phase 2 trial showed 8 out of 10 women had reduction in lesion area; 2 of 10 had no virus at 6 months (interim) Anal dysplasia: Open-label Phase 2 trial showed clearance of precancerous lesions in 10 out of 20 patients, decrease in lesions for 15 of 20 (interim)
Head & Neck Cancer (MEDI0457)	 Phase 1 trial for HNSCC, 2 out of 4 patients treated with MEDI0457 and 2 different PD-1 checkpoint inhibitors experienced a long-term complete response for >2 years MEDI0457 is licensed by AstraZeneca and currently in a Phase 1b/2a study in combination with durvalumab (PD-L1 checkpoint inhibitor)
RRP (INO-3107)	 Pilot study for Recurrent Respiratory Papillomatosis (RRP) demonstrated a clinical benefit in 2 out of 2 patients by delaying surgery due to lack of tumor recurrence A Phase 1/2 clinical trial for treating RRP with INO-3107, which includes both HPV 6 and HPV 11 antigens, is planned



Immuno-Oncology Programs (Non-HPV Associated)

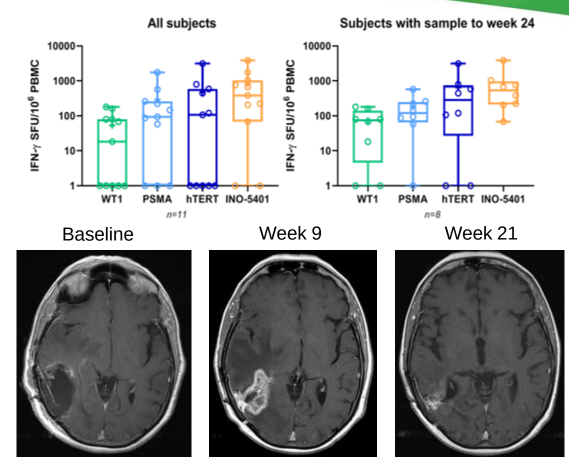
INO-5401 for Newly Diagnosed GBM in Phase 1/2 Study in Collaboration with Regeneron

TRIAL: INO-5401 (encoding tumor-associated antigens: hTERT, WT1, PSMA)



INO-5401 Interim Results: Promising 6-Month Progression-Free Survival Data, 12- and 18-Month Overall Survival Data in 2020

- SITC late-breaking abstract presented November 6-10th 2019
 - MGMT-Unmethylated PFS6 (24/32) <u>75% vs ~40% SOC</u>
 - MGMT-Methylated PFS6 (16/20) <u>80% vs ~60% SOC</u>
 - Supportive safety, tolerability, and immunogenicity data
 - Acceptable safety profile consistent with that of Libtayo and INOVIO's platform technology
 - Majority of patients tested had a T cell immune response to one or more tumor-associated antigens encoded by INO-5401
- The combination of INO-5401 + INO-9012 with cemiplimab, given with RT and TMZ, is promising
- Overall survival results (OS12, OS18) will be available in 2020



Several patients have experienced pseudo-progression, with progression by RANO criteria and radiographic evidence of progression on MRI, without evidence of tumor on repeat biopsy



INO-5151 Phase 2 Prostate Cancer Combination Study

TRIAL: INO-5151 (encoding tumor-associated antigens: PSA, PSMA)



Phase 2 study (PORTER) for metastatic castration-resistant prostate cancer



Three cohort, 45-patient platform study, **INO-5151 in Cohort C**

Cohort C – 15 patients



INO-5151 (DNA immunotherapy)CDX-301 (FLT3 ligand) from Celldex TherapeuticsNivolumab (anti-PD-1) from Bristol-Myers Squibb

PICI/CRI will fund & execute the clinical study







Infectious Disease Programs (Non-HPV Associated)

Positive Clinical Data and Partnering Opportunities

Product	Indication	Data Reported (to date)	Partner/s	Next Milestone
PENNVAX-GP	HIV	 Phase 1: 93% (71 of 76) evaluable vaccinated participants showed a CD4+ or CD8+ cellular immune response to at least one of the vaccine antigens 94% (62 of 66) demonstrated an env specific antibody response 	NIH) NIAID	Interim results from Phase 1/2 HIV trial study 2020 (UCSF; Deeks)
INO-4201	Ebola	 Phase 1: High levels of binding antibodies measured (ELISA) in 95% (170 of 179) of evaluated subjects Published: The Journal of Infectious Diseases, March 2019 	DARPA	Seeking additional grant funding for Phase 2 development
INO-4700 (GLS-5300)	MERS	 Phase 1: High levels of binding antibodies measured (ELISA) in 92% (57 of 62) of evaluated subjects 98% (61 of 62) generated an antibody and/or T cell response against MERS Published: The Lancet Infectious Diseases, July 2019 	전원생명과학(주) GeneOne Life Science	Publish Phase1 data
INO-4600 (GLS-5700)	Zika	 Phase 1: High levels of binding antibodies measured (ELISA) in 100% (39 of 39) of evaluated subjects Published: New England Journal of Medicine, October 2017 	GENE 진원생명과학(주) GeneOne Life Science	Report on Puerto Rico study 2020



INOVIO's COVID-19 DNA Vaccine INO-4800 Development Timeline

DECEMBER 31, 2019	JANUARY 10, 2020	JANUARY 10 - JANUARY 23, 2020	JANUARY 23, 2020	JANUARY 23 - FEBRUARY 29, 2020	MARCH 2020
INOVIO coronavirus experts learn about a novel coronavirus (SARS-CoV-2) which caused an outbreak of respiratory disease in Wuhan, China, now referred to as COVID-19	Chinese researchers share the genetic sequence of the novel coronavirus INOVIO designs DNA vaccine INO-4800 in three hours after receiving the genetic sequence using its proprietary DNA medicines platform technology INO-4800 was designed to precisely match the DNA sequence of the virus	INOVIO coronavirus experts race to manufacture INO-4800 and begin preclinical testing	INOVIO receives a grant of up to \$9 million from the Coalition for Epidemic Preparedness Innova- tions (CEPI) to fund ongoing preclinical and initial clinical development of INO-4800	Preclinical testing continues, with immune responses generated in animal models; initial preclini- cal data submitted to peer-reviewed journal; human clinical trial designs developed and shared with regulators	Ongoing preclinical studies, including challenge studies; initial preclinical data "under consideration" at peer-reviewed journal; human clinical trial designs finalized; 3,000 human trial doses prepared for clinical trials in the U.S., China, and South Korea ; large-scale manufacturing plans developed
MARCH 12, 2020	MARCH 26, 2020	APRIL 6, 2020	SPRING - SUMMER 2020	FALL 2020	END OF 2020
INOVIO announces \$5 million grant from the Bill & Melinda Gates Foundation to continue advancing the development of INO-4800, specifically to accelerate the testing and scale up of INOVIO's proprietary smart device CELLECTRA® 3PSP BILL&MELINDA GATES foundation	INOVIO announces the Department of Defense (DOD) awarded Ology Bioservices an \$11.9 million contract to work with INOVIO on DNA technology transfer to rapidly manufacture INO-4800 for the DOD for upcoming clinical trials	INOVIO announces initiation of Phase 1 human clinical trial in the U.S. following authorization by the U.S. Food and Drug Administration of its Investigative New Drug (IND) application U.S. study to include up to 40 healthy volunteers; study sites are the University of Pennsylvania and a clinic in Kansas City, MO	U.S. Phase 1 study initial immune responses and safety data expected by late summer Human clinical trials expected to begin in China and South Korea;* CEPI provides \$6.9 million to INOVIO and International Vaccine Institute for Phase 1/2 study in South Korea with 160 participants	Upon successful completion of Phase 1 study, Phase 2 trial to begin Human clinical trial results presented/ published	1 million doses of INO-4800 COVID-19 DNA vaccine planned for production*

Leading a Global Public Health Dialogue, Addressing Urgent Health Needs



A deadly new virus. Thousands of people infected. No cure. No vaccine.

We've been here many limes before.

In the past five years alone, the world has faced outbreaks of Ebola. Zika, another

SAN DIEGO BUSINESS JOURNAL

MEDICINE: INVESTOR INTEREST SWIRLS AROUND COMPANIES Biotechs in San Diego Join Race to Combat Coronavirus By Jared Whitlack

Thursday, February 6, 2020



Inovio Pharmaceuticals CEO Joseph Kim at the company's San Diego office. Inovio is working on a vaccine for the new coronavirus. Photo by Jamie Scott Lytle

Chinese scientists on Jan. 10 shared online the genetic sequence of a deadly strain of coronavirus. The next day, Inovio Pharmaceuticals whipped up a potential vaccine.

The New York Times

Apr. 7, 2020

A Pharmaceuticals Company is Testing a Possible Vaccine – The Second To Start Human Trials.



Dr. J. Joseph Kim, the chief executive of Inovio Pharmaceuticals, at the White House last month. Inovio's product is the second vaccine candidate to start early human trials in the United States. Andrew Harnik/Associated Press

Inovio Pharmaceuticals announced Monday that it will begin a small safety test of a potential coronavirus vaccine in adults in Philadelphia and Kansas City, Missouri.

Inovio's product is the second vaccine candidate to start early human trials in the United States. Researchers began testing a vaccine candidate developed by the biotech company Moderna in Seattle in mid-March.

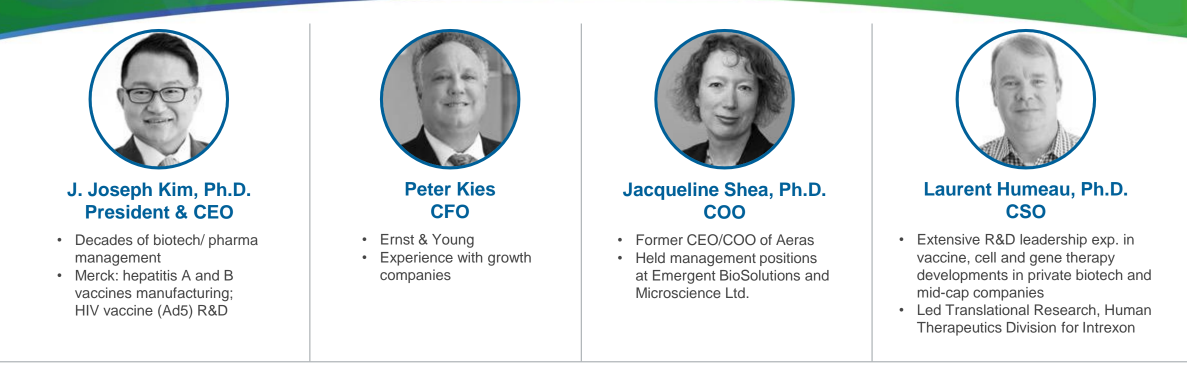


INOVIO Global and Local Coverage Drives Leadership Position



Management & Financials

Experienced Executive Team and Board of Directors



Board of Directors Simon X. Benito Chairman of the Board, Former SVP, Merck Vaccine Division

Angel Cabrera, Ph.D. President, George Mason University

J. Joseph Kim, Ph.D. President & CEO, INOVIO Pharmaceuticals

Ann. C. Miller, M.D. Former Head of Sanofi Oncology Global Marketing Jay Shepard Former President & CEO, Aravive

David B. Weiner, Ph.D. Executive VP, Director, Vaccine Center, The Wistar Institute Wendy L. Yarno, Ph.D.,

Former Executive VP and Chief Marketing Officer, Merck

Lota S. Zoth Former CFO, MedImmune



Strong Balance Sheet to Support Critical Milestones

\$89.5M Cash and short-term investments

As of December 31, 2019

\$208.2M Net proceeds from new financing in 1Q20

144.5M Common stock shares outstanding

As of March 12, 2020

Milestones

VGX-3100

- 1Q20: Report interim data from Phase 2 VIN/AIN clinical trials
- 4Q20: REVEAL 1 Phase 3 top-line efficacy & safety data

MEDI0457

- 2H20: Potential presentation from AZ on MEDI0457 Phase 2 study in HNSCCINO-3107
- ✓ 1H20: Initiate Phase 1/2 trial of INO-3107 for RRP (HPV6 and 11)

INO-5401

- □ 2Q20: OS12 data from Phase 1/2 GBM clinical trial (INO-5401 plus Libtayo[®])
- □ 4Q20: OS18 data from Phase 1/2 GBM clinical trial (INO-5401 plus Libtayo[®])

Platform Development

- ✓ April 2020: Initiate Phase 1 trial of INO-4800 for COVID-19
- 2020: Advance INO-4700 against MERS into Phase 2 field study in Middle East & Africa (CEPI-funded)
- 2020: Interim Phase 1 results from first-in-human trial of dMAb[™] plasmid candidate INO-A002 (for preventing or treating Zika virus infection)



NASDAQ:INO

INOVIO DNA Medicine Value Proposition

NASDAQ:INO

Validated Proprietary Technology

- Platform has demonstrated Phase 2b clinical efficacy of lead asset VGX-3100
- Well-protected with over 1,000 issued and pending patents
- Over 2,000 patients safety data and demonstration of high levels of T cell and antibody immune responses
- Over \$180M in non-dilutive funding since 2009
- **Partnerships** with major pharma and organizations:



12-Month Pipeline Catalysts

- Phase 3 for lead asset VGX-3100 treating high-grade cervical dysplasia
 - Efficacy data from REVEAL 1 in 4Q20
- Phase 2 checkpoint-combination programs in glioblastoma and HPV-associated cancers
 - **INO-5401** GBM OS12 and OS18 data in 2020
 - **MEDI0457** potential pres. from AZ on HNSCC data in 2H20
- Phase 1 trial for INO-4800 DNA vaccine candidate
 against COVID-19
 - Initial immune response/safety data Q3; upon successful completion, begin Phase 2
 - Data published from preclinical studies, including challenge studies





POWERING DNA MEDICINES[™]



Appendix

INOVIO DNA Medicines Will Meet Urgent Health Needs Worldwide

HPV-Related Diseases

- Nearly 80M Americans are currently infected with HPV; ~14M become infected each year
- ~35k Americans get an HPV-attributable cancer per year, including head and neck and cervical, anal, penile and vulvar cancers
- ~23% of Americans age18-59 have genital infections with ≥1 high-risk HPV genotype (e.g., HPV 16, HPV 18), which can lead to cervical, anal, head and neck, and other cancers; no current medicine to destroy/clear the virus
 - ~4% of Americans age 18-69 have oral infection with ≥1 high-risk HPV genotype
- Other HPV genotypes (6/11) can cause debilitating conditions such as Recurrent Respiratory Papillomatosis (RRP), rare and potentially life-threatening in children and adults; only current treatment is multiple, lifelong surgeries

HealthDay

HPV Blamed for Rising Rates of Anal Cancer

OURNAL of the







Anal cancer rates and deaths are climbing in the US, study says



The Washington Post Che Washington Post

A toddler's dwindling voice was chalked up to acid reflux. Her problem was far more serious.





Woman With Raspy Voice Has Had More Than 300 Surgeries to Treat Rare Vocal Cord Disease

HEALTH: 2:35 PM PDT, April 11, 2019. JOHANNA LI



Try 1 month for SI

INOVIO DNA Medicines Will Meet Urgent Health Needs Worldwide (continued)

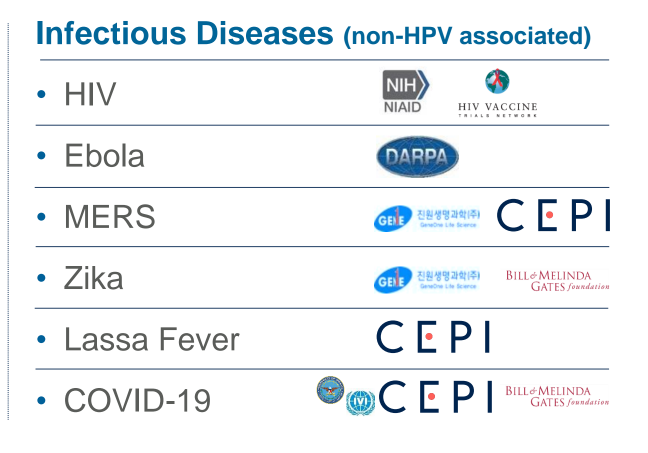
Cancer (non-HPV associated)

>11,000 people in U.S. get glioblastoma (GBM, rare and most aggressive form of brain cancer) each year; 23,000 people in U.S. have GBM

REGENERON

 ~3.1M men in U.S. have prostate cancer, the most common cancer among men except for skin cancer

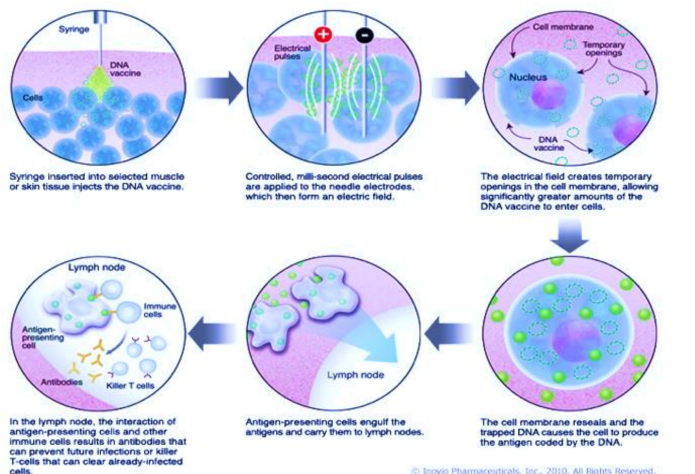






INOVIO's Technology Delivering Precisely Designed Plasmids with Proprietary Smart Devices

INOVIO's DNA medicine powers a patient's immune system to generate functional antibodies and killer T cells





Innovation in the Delivery of SynCon[®] DNA Medicine

CELLECTRA®-3P

• 3mm electrodes

minimally invasive

Intradermal –

In clinical use

CELLECTRA®-5PSP

- Intramuscular
- 13, 19, 25mm electrodes
- In clinical use



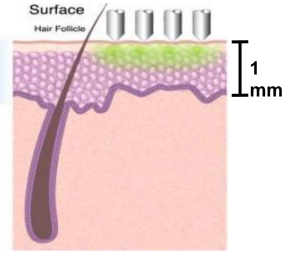
Skin Skeletal Muscle

Full Thickness

Surface EP (SEP)

- Surface
- Noninvasive
- 4x4 electrode array
- Specifically targets epidermis
- In late-stage preclinical development







CELLECTRA® Platform

CELLECTRA-5PSP Intramuscular EP



CELLECTRA-3P Intradermal EP



CELLECTRA-3P technology in a hand-held portable device

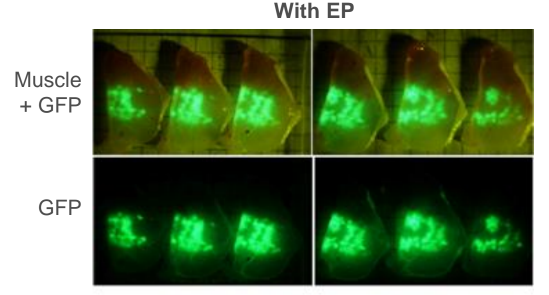


CELLECTRA® 2000 EP Technology – Track record of success in the clinic

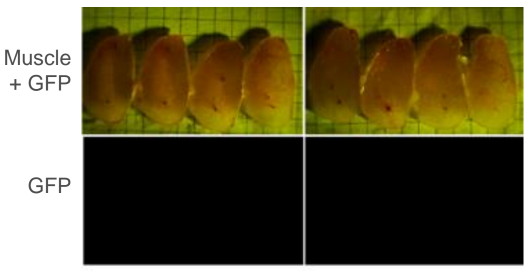
- >2000 human subjects and >6000 doses
- CELLECTRA® 5PSP device developed to support Phase 3 and commercial launch
- Phase 2 efficacy data combining DNA vaccine and EP
- Global Regulatory approval for studies in 6 continents (including Central & Sub-Saharan Africa); both devices CE marked in Europe



Precise Design + Intracellular Delivery = Improved Immune Responses



Display of GFP (green fluorescent protein) gene expression after CELLECTRA[®] delivery into rabbit muscle No EP





CTLA4 or PD1 + DNA Vaccine Improves Tumor Control & Survival in Challenge Model

Checkpoint Inhibitor Therapies Combined with INOVIO DNA Medicine

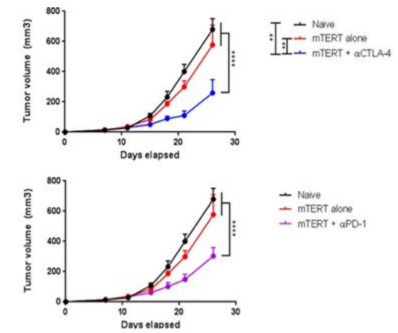
- Potential to improve response rates, without adding toxicity
- Tumor infiltration of antigen-specific, functional CD8+ T cells may prime patients for treatment with checkpoint inhibitors and increase response rates
- Combination studies initiated
 - MEDI0457 with AstraZeneca PDL-1
 - INO-5401 with Regeneron PD-1
 - **INO-5151** with BMS PD-1 + Celldex FTL3L (PICI Study)

Molecular Therapy Original Article



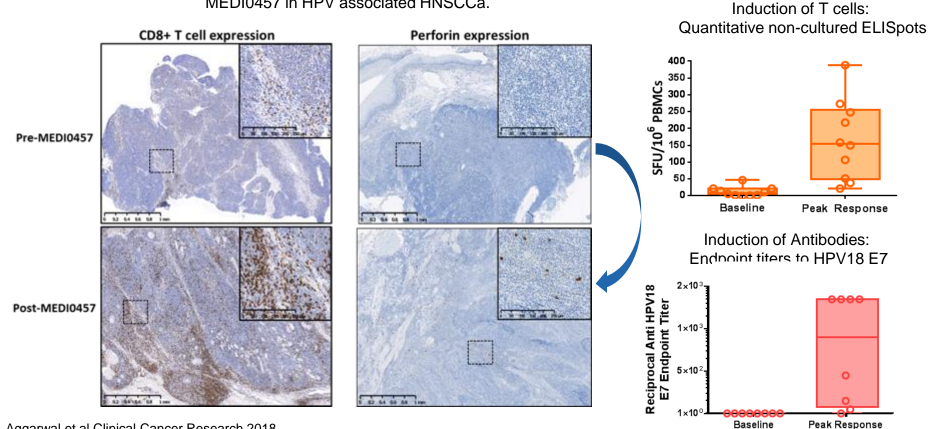
Synergy of Immune Checkpoint Blockade with a Novel Synthetic Consensus DNA Vaccine Targeting TERT

Elizabeth K. Duperret,¹ Megan C. Wise,^{2,3} Aspen Trautz,¹ Daniel O. Villarreal,³ Bernadette Ferraro,² Jewell Walters,² Jian Yan,² Amir Khan,² Emma Masteller,² Laurent Humeau,² and David B. Weiner¹





MEDI0457 (HPV16/18) Induces Robust Anti-Tumor Immunity in Head and Neck Cancer Phase 1 study of MED10457 (INO-3112) in 22 HPV+ HNSCC Patients



Strong invasion by CD8 T cells into tumors following immunization with MEDI0457 in HPV associated HNSCCa.

Aggarwal et al Clinical Cancer Research 2018

Most participants respond immunologically to the vaccine









GBM (Newly-diagnosed) Phase 1/2 Study

Trial Treatment (NCT03491683)

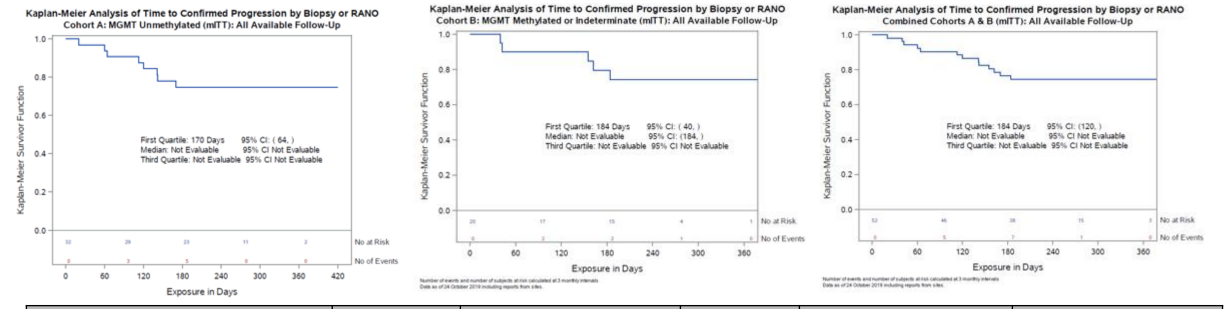
- INO-5401 (3 mg of each <u>WT1, PSMA and hTERT plasmids</u>) combined with 1 mg INO-9012, (total 10 mg of DNA) IM injection followed by EP given <u>every 3 weeks</u> for 4 doses, then <u>every 9 weeks</u>; and
- Cemiplimab (LIBTAYO[®]) (350 mg/dose IV every 3 weeks)

Chemoradiation Treatment

- **Radiotherapy** (RT), given in a hypofractionated schedule (40 Gy over 3 weeks) for all patients post surgery
- Temozolomide (TMZ) concurrent with RT for all patients, and then following RT for 6 cycles in <u>methylated patients only</u>



GBM-001 Progression-Free Survival at Six Months (PFS6)



Cohort	N Subjects	N Event-free Subjects	PFS6 (%)	95% CI Lower Bound	95% CI Upper Bound
Cohort A (MGMT Unmethylated)	32	24	75	56.6	88.5
Cohort B (MGMT Methylated)	20	16	80	56.3	94.3
Both Cohorts Combined	52	40	77	63.2	87.5

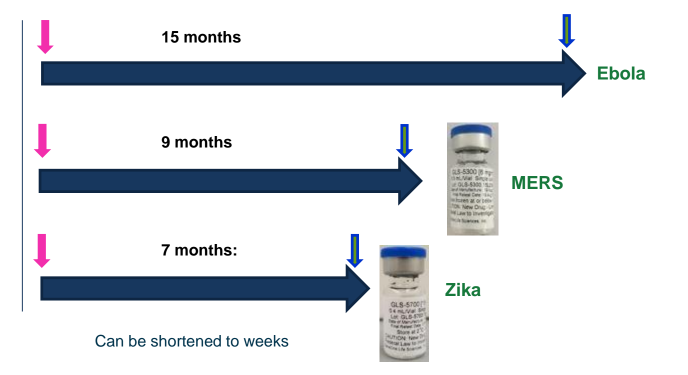
Confirmed PD (RANO) = confirmation by consecutive PD scan \geq 4 weeks from original PD event, or progressed according to biopsy surgery. Subjects who terminated for any reason prior to 6 months other than PD included as confirmed progressive events, including two (2) subjects in Cohort B who came off-study at week three (3), and declined long-term follow-up. Note: subjects with time to events longer than 6 months included; subjects have different time on study durations.



Rapid Clinical Translation of INOVIO Infectious Disease Programs



- Ebola 15 months to clinic: 95% response rate post dose 2 (publication submitted)
- MERS 9 months to clinic: 95% responses post dose 2, 98% overall response rate
- Zika 6.5 months to clinic (including animal preclinical work): 100% response rate-passive transfer protection (Tebas et al NEJM 2017)



Executive Team



J. Joseph Kim, Ph.D., President & CEO

- Decades of biotechnology/pharma management
- Merck: hepatitis A and B vaccines manufacturing; HIV vaccine (Ad5) R&D



Peter Kies CFO

- Ernst & Young
- Experience with growth companies



Jacqueline Shea, Ph.D., COO

- Former CEO of Aeras, the leading not-for-profit organization dedicated to developing new tuberculosis vaccines
- Held management positions at Emergent BioSolutions and Microscience Ltd.



Laurent Humeau, Ph.D., CSO

- Extensive R&D leadership experience in vaccine, cell and gene therapy developments in private biotech and mid-cap companies
- Led Translational Research, Human Therapeutics Division for Intrexon



Board of Directors



Simon X. Benito Chairman, BOD

 Former Senior Vice President, Merck Vaccine Division



Angel Cabrera, Ph.D.President, Georgia Tech



J. Joseph Kim, Ph.D.President & CEO, INOVIO



Ann C. Miller, M.D.
Former Head of Sanofi Oncology Global Marketing



Jay Shepard

 Former President & CEO, Aravive; Former Executive Partner, Sofinnova Ventures



David B. Weiner, Ph.D.

 Executive VP, The Wistar Institute; Director, Vaccine Center



 Wendy Yarno
 Former Chief Marketing Officer, Merck



Lota Zoth, CPA

Former CFO, MedImmune



Scientific Advisory Board



David B. Weiner, Ph.D., Chairman

- "Father of DNA vaccines"
- Executive VP, The Wistar Institute; Director, Vaccine Center



Anthony W. Ford-Hutchinson, Ph.D.

- Former SVP, Vaccines R&D, Merck
- Oversaw development: Singulair[®], Januvia[®], Gardasil[®], Zostavax[®], Proquad[®] and Rotateq[®]



Stanley A. Plotkin, M.D.

- Developed rubella and rabies vaccines
- · Oversaw Sanofi flu vaccine
- Emeritus Professor, Wistar Institute & University of Pennsylvania



Rafi Ahmed, Ph.D.

 Professor, Department of Microbiology and Immunology, Emory University School of Medicine



INOVIO Fully Integrated Capabilities Poised for Rapid Production



Philadelphia Corporate and Operations Site

- Corporate, Clinical, Regulatory, Compliance, Biostatistics, and Data Management functions
- ~80 FTE



San Diego Research Center

- Molecular biology, cell biology, and clinical immune monitoring
- Research-grade DNA manufacture capabilities
- 6,000 sf dedicated BSL-2 research lab (wet lab and cell culture)
- 5,000 sf cGLP labs to process, store, and analyze human clinical trial samples
- Well established QA capability
- ~40 FTE



San Diego Device Engineering and Manufacturing Facility

- Electroporation delivery device and consumable design, engineering, and manufacturing
- Delivery device testing and distribution
- 53,000 sf facility opened in July 2017
- ISO 13485 and MDD certified by TÜV America in San Diego
- ~70 FTE

