



# oncosec™

Powerful, Tumor-agnostic Immunotherapy Treatment

ONCOSEC'S PLATFORM TO ATTACK VISCERAL, CUTANEOUS AND SUBCUTANEOUS TUMORS

NASDAQ: ONCS  
SEPTEMBER 2020

# FORWARD LOOKING STATEMENTS

To the extent statements contained in the following presentations are not descriptions of historical facts regarding OncoSec Medical Incorporated, they should be considered “forward-looking statements,” as described in the Private Securities Litigation Reform Act of 1995, that reflect management’s current beliefs and expectations. You can identify forward-looking statements by words such as “anticipate,” “believe,” “could,” “estimate,” “expect,” “forecast,” “goal,” “hope,” “hypothesis,” “intend,” “may,” “plan,” “potential,” “predict,” “project,” “should,” “strategy,” “will,” “would,” or the negative of those terms, and similar expressions that convey uncertainty of future events or outcomes. Forward-looking statements are not assurances of future performance and include, but are not limited to, statements regarding: (i) the success and timing of our product development activities and clinical trials; (ii) our ability to develop and commercialize our product candidates; (iii) our plans to research, discover, evaluate and develop additional potential product, technology and business candidates and opportunities; (iv) our and our partners’ ability to develop, manufacture and commercialize our product candidates and to improve the manufacturing process; (v) the size and growth potential of the markets for our product candidates, and our ability to serve those markets; (vi) the rate and degree of acceptance of our product candidates; (vii) our ability to attract and

retain key scientific or management personnel; (viii) the anticipated timing of clinical data availability; (ix) the anticipated timing of commercial launch of ImmunoPulse® IL-12; (x) our ability to meet our milestones; (xi) our expectations regarding our ability to obtain and maintain intellectual property protection; (xii) the level of our corporate expenditures; (xiii) the assessment of our technology by potential corporate partners; and (xiv) the impact of capital market conditions on us. Forward-looking statements are subject to known and unknown factors, risks and uncertainties that may cause actual results to differ materially from those expressed or implied by such forward looking statements. These statements are also subject to a number of material risks and uncertainties that are described in OncoSec’s most recent Annual Report on Form 10-K filed with the Securities and Exchange Commission, as updated by its subsequent filings with the Securities and Exchange Commission. Undue reliance should not be placed on forward-looking statements. We undertake no obligation to publicly update any forward-looking statements, except as required by law. OncoSec’s investigational drug and device products have not been approved or cleared by the FDA.

# OncoSec at a Glance

We are a late-stage biotechnology company focused on intratumoral DNA based immunotherapies.

- **Headquarters:** Pennington, NJ, with R&D Laboratories in San Diego, CA
- **Ticker:** NASDAQ: ONCS
- **Market Cap (as of 7/20/20):** ~\$82M
- **Stock Price (as of 7/20/20):** \$3.57



**Technology:** Our lead product candidate, TAVO™, intratumoral DNA plasmid-based IL-12 delivered via electroporation, has demonstrated promising anti-tumor activity with abscopal responses in melanoma and four other cancer types. Anti-tumor activity observed both as a monotherapy and in combination with anti-PD-1 checkpoint inhibitors.



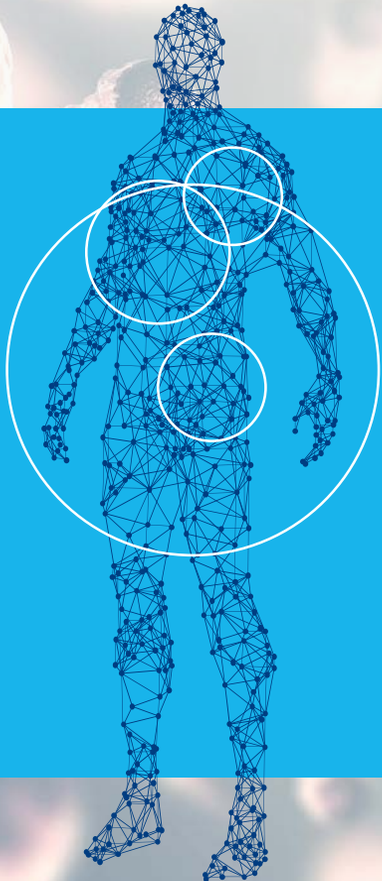
**Clinical Pipeline:** TAVO is being evaluated in combination with Merck's KEYTRUDA® in two ongoing KEYNOTE clinical trials - a pivotal study in anti-PD-1 checkpoint resistant metastatic melanoma and a phase 2 study in metastatic triple negative breast cancer.



**Partnerships:** We have a track record of establishing, operating and evolving high-performance partnerships globally. Ongoing collaboration partners include Merck, CGP, Sirtex, among others.



# The Promise of Immunotherapy Has Yet to Be Fully Realized



## THE PROMISE

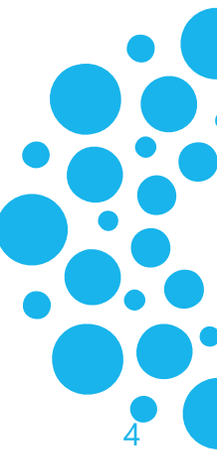
Instead of cytotoxic agents, use the body's own immune system against tumors

Immunotherapy in many forms — like checkpoint therapy — have had **unprecedented success** in halting or shrinking cancer

## How Checkpoint Inhibitors Work

- 1 Molecular switches known as checkpoints normally prevent T-cells from attacking healthy tissue
- 2 When these checkpoints, such as PD-1 and PD-L1, are hijacked by cancer cells, the immune system's T-cell response is switched off, allowing tumors to grow
- 3 Checkpoint inhibitors flip the switch back on, freeing the immune response so that T-cells are activated and destroy the cancer cells

Yet, there are still too many patients who are not benefiting from these therapies



# Checkpoint Non-response in 60-90% of Cases

## TUMOR TYPE

% OF CHECKPOINT  
NON-RESPONDERS

MELANOMA

~60-80%

TRIPLE NEGATIVE BREAST

~95%

HEAD AND NECK

~68-86%

CERVICAL

~86%

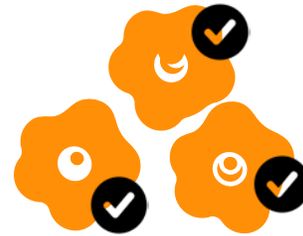
SUBCUTANEOUS  
T-CELL LYMPHOMA

~57%

**KEYTRUDA®**

Powerful drugs (like KEYTRUDA®) have been highly successful for some patients, but not the majority

90% of cancers are solid tumors. Of these:

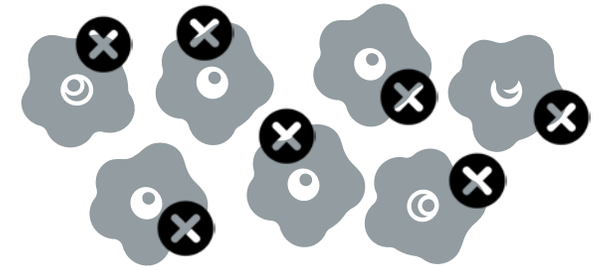


~30%

### Hot Tumors

Have T-cells and cancer fighters

Respond to checkpoint therapies



~70%

### Cold Tumors

Have immunosuppressive cells

Have few or no T-cells

Do not respond to checkpoint therapies

***There is an industry effort underway to improve response rates through new therapies or additional therapies***

# TAVO™ is Capable of Reversing Resistance to Checkpoint Therapies



TAVO (plasmid-based interleukin-12) is administered locally at the tumor site using OncoSec's electroporation gene delivery system. TAVO is designed to induce local expression of IL-12, turning "cold" tumors "hot" and **enabling checkpoint therapies to be effective.**



## Well Tolerated

TAVO leverages IL-12, a naturally occurring chemical in the body; intratumoral approach avoids systemic toxicity



## Cold to Hot

Clinical data shows TAVO induces local expression of IL-12, converting immunologically suppressed "cold" tumors into T-cell inflamed "hot" tumors



## Intratumoral Approach with Abscopal Effect

Clinical data in five tumor types showing evidence of anti-tumor activity with whole body (abscopal) effect



## Sustainable

Highly scalable with low manufacturing costs, potentially offering an innovative treatment option well below the cost of other biologic drug therapies

# Benefits of Electroporation (EP) Gene Delivery System

**A non-invasive,  
non-chemical,  
non-toxic method  
that is easy  
to perform**



## Rapid Transfection

More rapid than traditional chemical or biologic cell transfection techniques



## Surface & Visceral Lesions

Beyond cutaneous and subcutaneous; tumors can be accessed with an endoscope, bronchoscope, catheter, or trocar



## Versatile

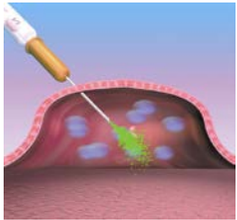
Wide array of molecules can be transfected, and can be applied to a broad selection of cell types



## Non-Invasive

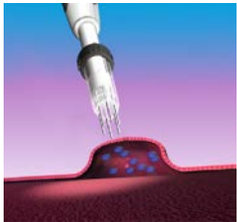
Electroporation gene delivery is noninvasive, nonchemical, nontoxic method of cell transfection, applicable to a wide array of immunologically relevant molecules

# Seamless Delivery of Plasmid IL-12 + Energy



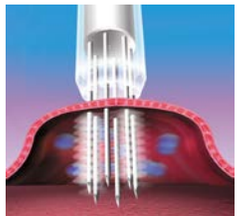
## Step 1: TAVO™ Injection

Multiple copies of IL-12 coded DNA plasmids to produce immune modulatory proteins are injected directly into the tumor using a conventional needle and syringe.



## Step 2: Applicator Insertion

The applicator's tip needle array is inserted into the tumor, up to a depth of 15mm.



## Step 3: Electroporation

Electrical pulses, activated by a foot switch administered between hexagonal needle electrodes increases the permeability of cell membranes, facilitating uptake ("transfection") of IL-12 coded DNA into cells.



## Genpulse™ Generator

Fixed electrical field intensity. Momentary electrical pulses (100  $\mu$ sec duration and 300 millisecond interval). Pulses activated by foot switch. 16 lbs. 12.5" w x 5.5" h x 13" d

## Sub / Cutaneous Applicator

Handle with electrode needed array disposable tip. Applicator 0.5 or 1.0 cm in diameter. Needle array hexagonal. Adjustable needles 1-15 mm.

**Entire procedure takes approximately 30 minutes**

5

CLINICAL/  
IMMUNOLOGICAL  
SIGNAL IN FIVE  
TUMOR TYPES

# TAVO: A Tumor Agnostic Platform with Single Agent Activity

## TAVO + PEMBRO COMBINATION SPONSORED TRIALS

**Metastatic Melanoma**  
U.S. Orphan  
Fast Track  
EU ATMP

**Metastatic TNBC**  
Accelerated Approval  
Opportunity

**KEYNOTE-695**  
TAVO + PEMBRO  
Second Line CPI  
refractory  
N=100

**KEYNOTE-890**  
TAVO + PEMBRO + Chemo  
First Line  
N=65

## TAVO SINGLE AGENT INVESTIGATOR-SPONSORED TRIALS

**Metastatic  
Melanoma**

**Metastatic  
Head & Neck**

**Metastatic  
Merkel  
Cell  
Carcinoma**

**Metastatic  
Cutaneous  
T-cell  
Lymphoma**

# Clinical Pipeline is Well Diversified with Multiple Growth Opportunities

**TAVO™**

REGIMEN	TRIAL	INDICATION	N	PARTNER	PHASE 1	PHASE 2	PIVOTAL
TAVO + pembrolizumab	<b>KEYNOTE-695</b>	Checkpoint Resistant Metastatic Melanoma	~100	 <b>MERCK</b>	→	→	→
TAVO + pembrolizumab	<b>KEYNOTE-890</b>	Triple Negative Breast Cancer (TNBC)	~65	 <b>MERCK</b>	→	→	→
TAVO + epacadostat + pembrolizumab	OMS-131 (Investigator Sponsored Study)	Squamous cell carcinoma head and neck (SCCHN) cancer	~34		→	→	→
TAVO + nivolumab	OMS-104 (Investigator Sponsored Study)	Neoadjuvant for operable locally/regionally advanced melanoma	~33		→	→	→

# Recent Peer Reviewed Publications

\*Click thumbnails to access full articles

## TAVO SINGLE AGENT ABCOPAL EFFECT

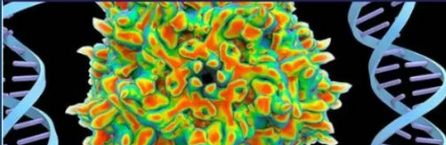
**Drug Evaluation**  
For reprint orders, please contact: [reprints@futuremedicine.com](mailto:reprints@futuremedicine.com)

**Immunotherapy**



Melanoma treatment with intratumoral electroporation of tavokinogene telseplasmid (pIL-12, tavokinogene telseplasmid)

**Gene Therapy**



Nature Gene Therapy  
Characterization of abscopal effects of intratumoral electroporation-mediated IL-12 gene therapy.

September 30, 2019

Published OnlineFirst December 18, 2019; DOI: 10.1158/2326-6066.CIR-19-0359

CANCER IMMUNOLOGY RESEARCH | RESEARCH ARTICLE

**Intratumoral Plasmid IL12 Electroporation Therapy in Patients with Advanced Melanoma Induces Systemic and Intratumoral T-cell Responses** 



Clinical Cancer Research  
Intratumoral Delivery of Plasmid IL12 Via Electroporation Leads to Regression of Injected and Noninjected Tumors in Merkel Cell Carcinoma.

February 1, 2020



**ESMO** GOOD SCIENCE BETTER MEDICINE BEST PRACTICE

**ORIGINAL ARTICLE**

**Intratumoral delivery of tavokinogene telseplasmid yields systemic immune responses in metastatic melanoma patients**

## TAVO + PEMBRO IN ANTICIPATED ANTI-PD-1 CHECKPOINT FAILURES

**JCI** The Journal of Clinical Investigation

**Tumor immune profiling predicts response to anti-PD-1 therapy in human melanoma**

JOURNAL OF CLINICAL ONCOLOGY

ORIGINAL REPORT

Programmed Death-Ligand 1 Expression and Response to the Anti-Programmed Death 1 Antibody Pembrolizumab in Melanoma

**JCI insight** CLINICAL MEDICINE

**Partially exhausted tumor-infiltrating lymphocytes predict response to combination immunotherapy**

Published OnlineFirst May 6, 2020; DOI: 10.1158/1078-0432.CCR-19-2217

CLINICAL CANCER RESEARCH | CLINICAL TRIALS: IMMUNOTHERAPY

**Phase II Trial of IL-12 Plasmid Transfection and PD-1 Blockade in Immunologically Quiescent Melanoma**

# FDA Fast Track

## KEYNOTE-695

Ongoing pivotal trial in PD-1 checkpoint resistant metastatic melanoma provides a pathway towards accelerated approval

### GRANTED FAST TRACK AND ORPHAN DRUG DESIGNATIONS



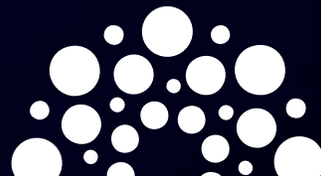
Fast-track makes TAVO™ eligible for accelerated approval program



Program for patients with no FDA approved treatment options



Must meet RECIST (tumor shrinkage) criteria – demonstrating that TAVO works



## KEYNOTE695

Up To  
30 SitesNumber Of  
PATIENTS  
TO BE TREATED

100

## DEFINITIVE CLINICAL PROGRESSION ON FULL COURSE ANTI-PD-1

Stage 3/4  
MelanomaReceived at  
least 12 weeks  
of anti-PD-1  
therapyProgressive  
disease  
according to  
RECIST v1.1Documented  
disease  
progression  
<12 weeks of  
last doseNo intervening  
therapies permitted  
between checkpoint  
failure and trial  
enrollmentEligible  
patient/definitive  
by RECIST V1.1  
anti-PD-1  
refractory patient

## PRIMARY OUTCOME

ORR (by BICR) based on RECIST v1.1

## SECONDARY OUTCOMES

ORR, DOR, PFS and OS

# PRELIMINARY DATA Metastatic Melanoma

TAVO™ + KEYTRUDA® (pembrolizumab) for Checkpoint Refractory Metastatic/Recurrent Melanoma

KEYNOTE-695 STUDY EVALUATING  
100 PATIENTS - **ONGOING**

4 PRs and 1 CR out of 21 patients evaluated by RECIST v1.1 as of 12/15/18

Durable responses observed

Responders are patients with bulky disease

Responders demonstrating regression of distant visceral lesions



**BASELINE**



**12 WEEKS**



**24 WEEKS**

Patient no longer treated with TAVO as there are no accessible lesions

Patient continues maintenance pembrolizumab

Notes: PR = partial response ;  
CR = complete response

# US Market Opportunity

**90%** of all cancer cases are solid tumors

**1.6M** new cases of solid tumors in the US

## Focusing First on Metastatic Melanoma in the United States



91,000 diagnosis each year



9,000 deaths each year



Incidence of melanoma on the rise (1.4% yearly for a decade)



US melanoma market projected to almost double from \$2B in less than 10 years

**15,000**

Patients receiving PD-1 inhibitors

**9,000**

PD-1 refractory patients

**2,700**

PD-1 refractory with accessible lesions

# Commercialization by 2022 Targeted



## Potential US Regulatory Timeline

2020

Pre-BLA Meeting with FDA

2022

FDA Accelerated Approval of TAVO for Metastatic Melanoma

2021

Submission of BLA for Accelerated Approval

## KEYNOTE 695

*TAVO is US Orphan Designated and Keynote-695 is fast tracked. OncoSec will seek accelerated approval*



## Potential EU Regulatory Timeline

*Awarded EU small-medium enterprise (SME) and ATMP designations by EMA's committee on advanced therapeutics (CAT) in 2020*

2020

Obtain CE Mark for to-be-marketed GenPulse Device

2021

Meetings with EU Rapporteurs

2023

EMA Conditional Approval of TAVO for Metastatic Melanoma

EARLY 2019

Obtained Advanced Therapy Medicinal Product (ATMP) Designation

2022

File MAA for Conditional Approval in EU and File Device Application in EU

# Expanding KEYNOTE-890 into First-Line mTNBC

## PHASE 2

Multi-cohort, open-label study of TAVO in combination with KEYTRUDA with or without chemotherapy in approximately 65 patients with metastatic TNBC

## COHORT 1

TAVO and KEYTRUDA in patients with heavily pre-treated mTNBC

**N=25**

CLOSED ENROLLMENT

## COHORT 2

TAVO and KEYTRUDA plus Chemotherapy in patients with mTNBC (first-line setting)

**N=40**

NOW ENROLLING

Primary Endpoint: ORR by blinded independent central review (BICR) based on RECIST v1.1

If KEYNOTE 890 Cohort 2 is successful, plan to expand into pivotal study

TAVO could be eligible for FDA accelerated approval

## KEY ADVANTAGES OF MOVING TO FIRST-LINE

High unmet medical need

Opportunity to improve PFS and OS

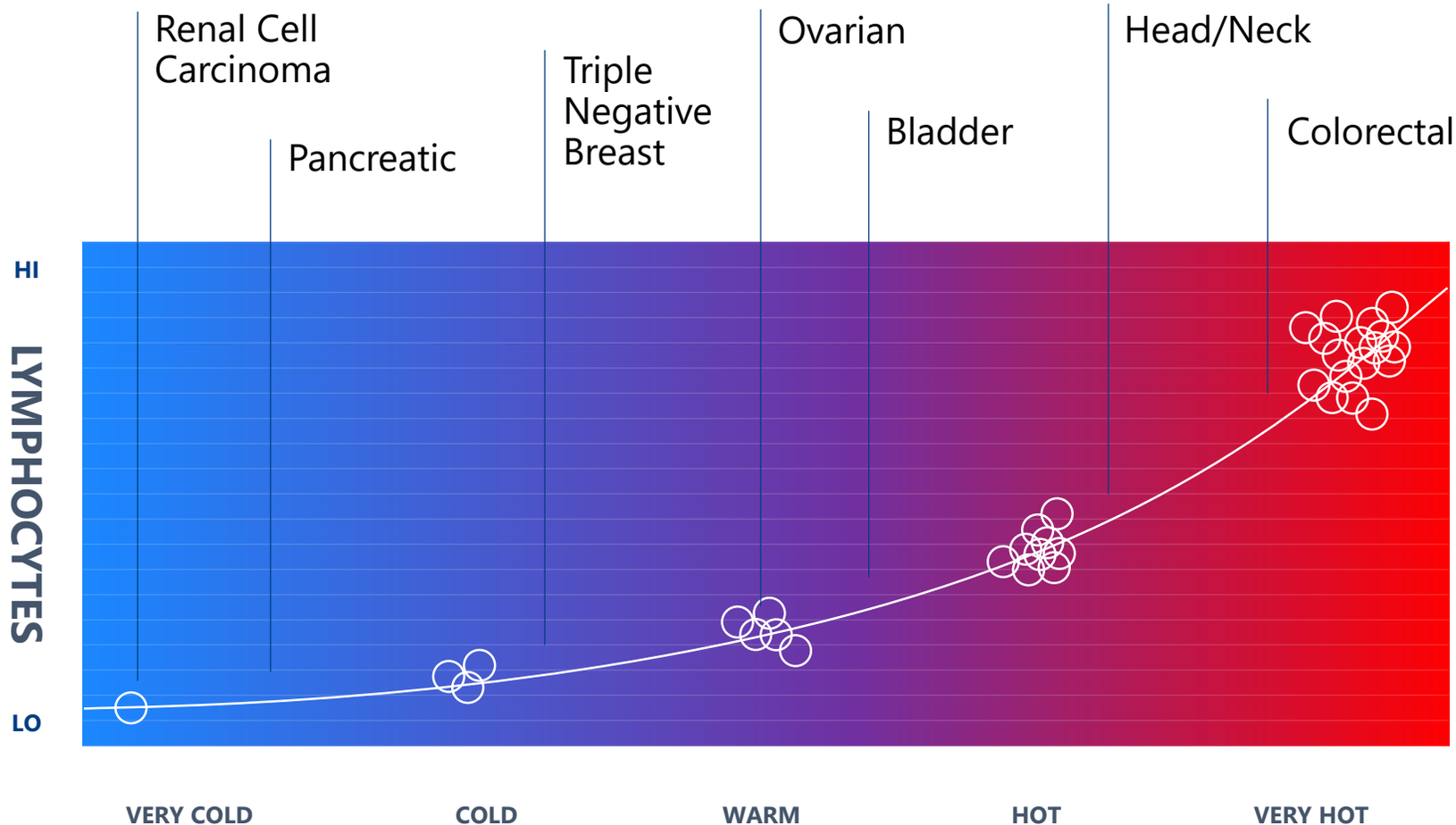
Cohort 2 designed to expand into pivotal study

Larger overall patient population – greater commercial opportunity vs. last-line setting

TAVO has shown higher efficacy in less refractory patients in melanoma

## FUTURE OPPORTUNITIES

# TAVO™ is Tumor Agnostic



**Giving life** to the promise of immunotherapies across cancer types

**Few cancers** are highly infiltrated or "HOT" - most fall on a spectrum from warm (cold-acting) to cold

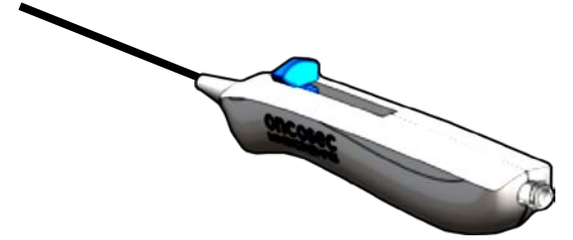
**TAVO is widely applicable**, we plan to expand our studies to include a wider range of tumors that do not respond well to checkpoints, including traditionally "cold"

**Pioneering technology  
designed to target and treat  
tumors located deep inside  
the body**

## INTRODUCING THE **VLA: Visceral Lesion Applicator**



Flexible catheter-  
based applicator



Rigid trocar-based  
applicator



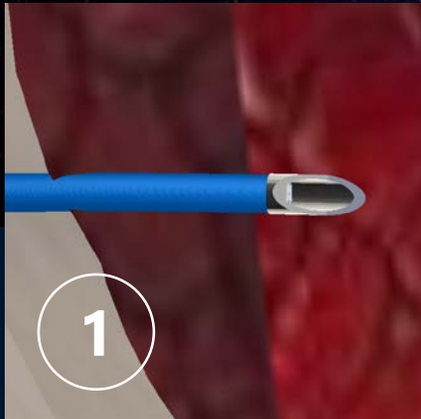
Lower voltage  
Apollo generator to be  
used with VLA

# VLA Highlights

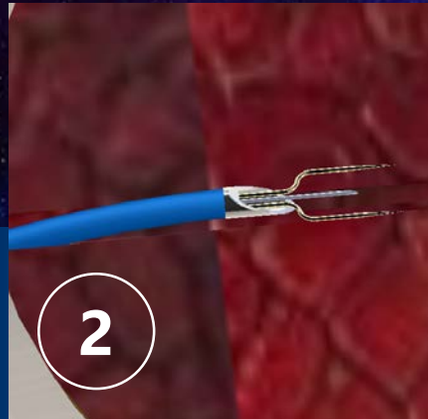
- 1 Rigid and catheter models:  
Used with bronchoscope,  
cystoscope, and trocar
- 2 Prototype successfully tested in  
large animal model: bone,  
pancreas, liver, lung, & bladder



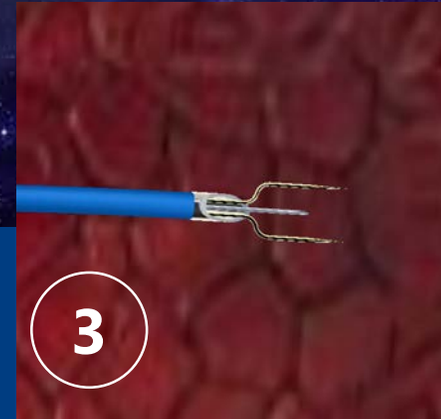
**ENTERS THE ORGAN**



**NEEDLE/ELECTRODES EXPANDED  
DRUG DELIVERY**



**ELECTROPORATION  
OCCURS**





# CORVax12 Vaccine for COVID-19



CORVax12 is a DNA vaccine which combines OncoSec's TAVO / IL-12 immunotherapy with NIH's "spike" protein developed by NIAID as a vaccine against the 2019 novel coronavirus



The combination of IL-12 and NIH's spike protein is designed to drive a coordinated vaccine response, capable of eliciting a Th1-biased cellular immune response to drive robust humoral immunity



Providence Cancer Institute to conduct Phase I trial of CORVax12 – Providence submitted Investigator-Initiated IND and plans to initiate trial pending FDA clearance

# Leveraging World Class Strategic Partnerships to Enhance Pipeline Value



Our partnership with CGP gives us the ability to develop and introduce our products to the important Chinese market

Through our partnership with Sirtex, we have access to top-notch commercial talent and resources

We are partnering with Merck on several KEYNOTE clinical programs to potentially address a great unmet need: anti-PD-1 non-responders

Emerge collaboration provides Australian metastatic melanoma patients early access to TAVO™ via the Special Access Scheme

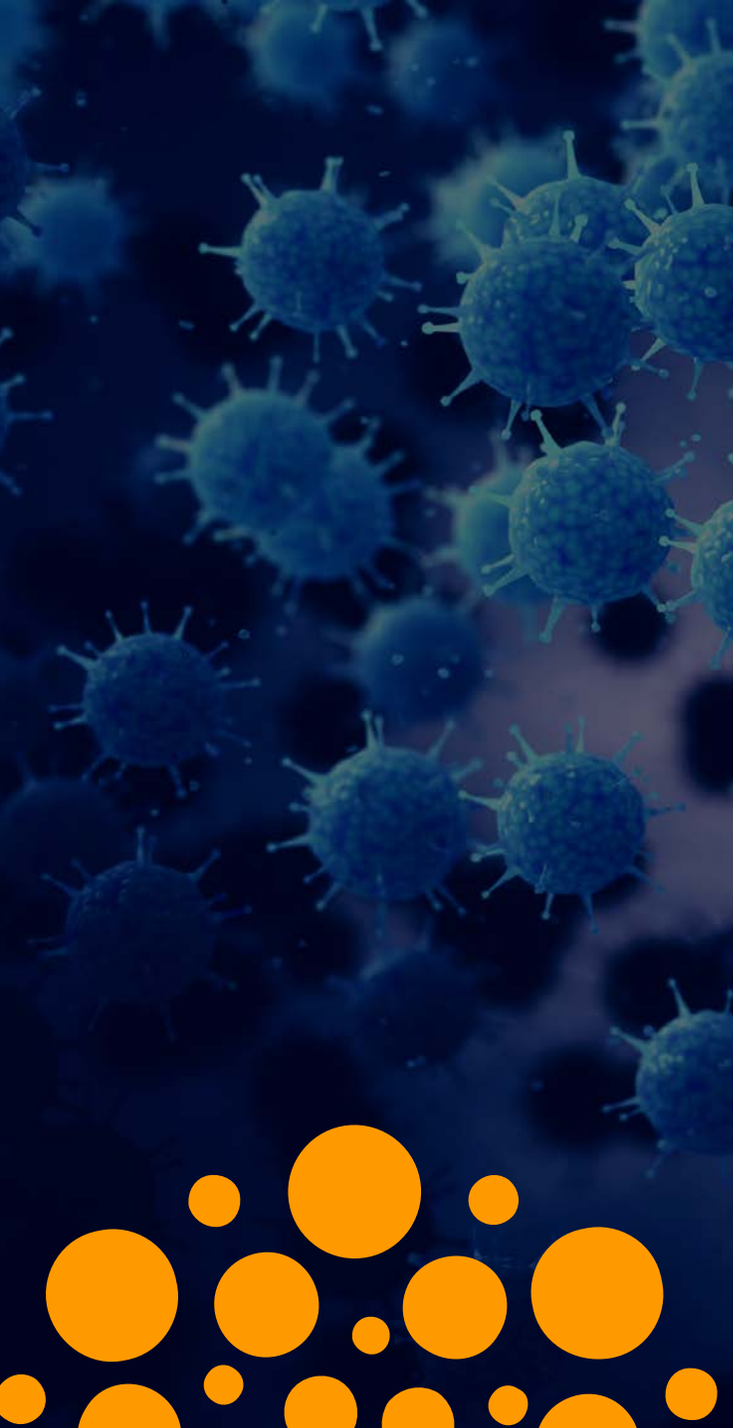
Other TAVO R&D Partnerships Include:



Duke  
UNIVERSITY



ROSWELL  
PARK



# Strong Financial Position to Drive TAVO™ Development Forward

**\$30.1M**

Cash & Equivalent as of  
4/30/20

**27,771,571**

Total Shares Outstanding  
as of 8/31/20

**~\$14M**

Financing Completed  
August 2020

**None**

Debt

# Key Developments in 1H 2020

- ✓ Closed on \$30 million strategic financing and partnership with CGP/Sirtex
- ✓ Presented promising preclinical data on novel visceral lesion applicator
- ✓ Announced partnership with Providence Cancer Institute to investigate OncoSec's CORVax-12 vaccine for COVID-19
- ✓ Appointed renowned oncologist and immunologist – Herbert Kim Lyerly, MD, Duke University – to board of directors
- ✓ TAVO™ granted CAT certification in Europe to support marketing authorization application in melanoma
- ✓ Increased recognition of TAVO data via 4 peer-reviewed publications year-to-date
- ✓ Strengthened IP estate with three new patents covering TAVO and electroporation gene delivery system
- ✓ Initiated a second TAVO / KEYTRUDA® combination study in TNBC designed to expand into a pivotal study

# Anticipated Milestones

- Provide interim data update at SITC 2020 for KEYNOTE-695
- Complete enrollment in pivotal KEYNOTE-695 study
- Prepare to file TAVO™ for accelerated FDA approval in metastatic melanoma
- Initiate TAVO neoadjuvant study in metastatic melanoma
- Complete preclinical assessment and IND prep for VLA, APOLLO™ and SPARK™

# Established Biotech Leaders

## WITH A TRACK RECORD OF SUCCESS

### MANAGEMENT



**Daniel J. O'Connor**  
*President/Director/CEO*



**Kellie Malloy Foerter**  
*Chief Operating Officer*



**Christopher G. Twitty, Ph.D**  
*Chief Scientific Officer*



**Keir Loiacono**  
*General Counsel, Vice President, Corporate Development*



**Robert J. Delaversano, CPA**  
*Principal Accounting Officer And Controller*



**Robert W. Ashworth, Ph.D**  
*Senior Vice President, Regulatory, Quality/CMC*



**Kim Jaffe, Ph.D**  
*Senior Director, Operations*



**John Rodriguez**  
*Vice President, Product Engineering*

### BOARD OF DIRECTORS

**Daniel J. O'Connor, JD**  
*Chief Executive Officer & Director*

**Margaret R. Dalesandro, Ph.D.**  
*Chair*

**Herbert Kim Lyerly, M.D.**  
*Director*

**Chao Zhou**  
*Director*

**Yuhang Zhao, Ph.D., M.B.A.**  
*Director*

**Kevin R. Smith**  
*Director*

**Jim DeMesa, M.D., M.B.A.**  
*Director*

**Robert E. Ward**  
*Director*

**Joon Kim, JD**  
*Director*

### CLINICAL ADVISOR

**Alain Algazi, M.D.**

# OncoSec is Positioned for Success with TAVO™



Positive tumor shrinkage/response data being generated by our lead product candidate, TAVO, across multiple solid tumor types



Well tolerated, natural solution that may increase the efficacy of checkpoint therapies



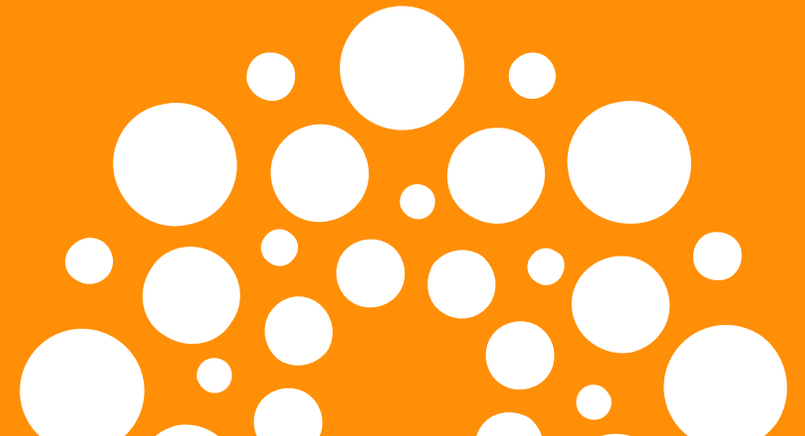
Expanding device development and clinical studies into new tumor types to serve a wider set of patients



Fast track status and partnership with Merck provides opportunity for more robust drug development



Strong financial position with no debt





**Thank You**



**oncosec**<sup>™</sup>  
IMMUNOTHERAPIES

**Keir Loiacono**

HEAD OF CORPORATE DEVELOPMENT

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