



# Tipping the balance in favor of the immune system to fight cancer

Corporate Presentation | April 2024

NASDAQ: CADL

### **Forward Looking Statements**

This Presentation contains forward-looking statements within the meaning of the Private Securities Litigation Reform Act of 1995. All statements other than statements of historical facts contained in this Presentation, including express or implied statements regarding our strategy, future financial condition, future operations, projected costs, prospects, plans, objectives of management and expected market size, are forward-looking statements. In some cases, you can identify forward-looking statements by terminology such as "may," "will," "should," "expect," "intend," "plan," "anticipate," "believe," "estimate," "target," "seek," "predict," "potential," "continue" or the negative of these terms or other comparable terminology. Although we believe that the expectations reflected in these forward-looking statements are reasonable, these statements relate to our strategy, future operations, future financial position, future revenue, projected costs, prospects, plans, objectives of management and expected market size, and involve known and unknown risks, uncertainties and other factors that may cause our actual results, performance or achievements to be materially different from any future results, performance or achievements expressed or implied by these forward-looking statements. Forward-looking statements in this Presentation include, but are not limited to statements about; the initiation, timing, progress, results, and cost of our research and development programs and our current and future preclinical and clinical studies, including statements regarding the timing of initiation and completion of studies or trials and related preparatory work, the period during which the results of the trials will become available, and our research and development programs; the therapeutic benefit of our programs, including the potential for our programs to extend patient survival; our ability to efficiently discover and develop product candidates; our ability to initiate, recruit and enroll patients in and conduct our clinical trials at the pace that we project; our ability to obtain and maintain regulatory approval of our product candidates; our ability to compete with companies currently marketing or engaged in the development of treatments that our product candidates are designed to target; our reliance on third parties to conduct our clinical trials and to manufacture drug substance for use in our clinical trials; the size and growth potential of the markets for our product candidates and our ability to serve those markets; the ability and willingness of our third-party strategic collaborators to continue research and development activities relating to our development candidates and product candidates; our ability to obtain and maintain adequate intellectual property rights; our estimates of our future expenses, revenue, capital requirements or our need for or ability to obtain additional financing; our ability to continue as a going concern, the potential benefits of strategic collaboration agreements, our ability to enter into additional strategic collaborations or arrangements, and our ability to attract collaborators with development, regulatory and commercialization expertise: our financial performance; and developments and projections relating to our competitors or our industry. We caution the recipient not to place considerable reliance on the forwardlooking statements contained in this Presentation. The forward-looking statements in this Presentation speak only as of the date of this document, and we undertake no obligation to update or revise any of these statements. Our business is subject to substantial risks and uncertainties, including those referenced above.

Certain information contained in this Presentation relates to or is based on estimates, projections and other information concerning the Company's industry, its business and the markets for its programs and product candidates and studies, publications, surveys and other data obtained from third-party sources and the Company's own internal estimates and research. While the Company believes these third-party sources to be reliable as of the date of this Presentation, it has not independently verified, and makes no representation as to the adequacy, fairness, accuracy or completeness of, any information obtained from third-party sources. In addition, all of the market data included in this Presentation involves a number of assumptions; there can be no guarantee as to the accuracy or reliability of such assumptions. Finally, while we believe our own internal research is reliable, such research has not been verified by any independent source.

These forward-looking statements are based on the beliefs of our management as well as assumptions made by and information currently available to us. Although we believe the expectations reflected in such forward-looking statements are reasonable, we can give no assurance that such expectations will prove to be correct. If such assumptions do not fully materialize or prove incorrect, the events or circumstances referred to in the forward-looking statements may not occur. We undertake no obligation to update publicly any forward-looking statements for any reason after the date of this presentation to conform these statements to actual results or to changes in our expectations, except as required by law. Accordingly, readers are cautioned not to place undue reliance on these forward-looking statements. Additional risks and uncertainties that could affect our business are included under the caption "Risk Factors" in our most recent Form 10-K filed with the Securities and Exchange Commission on March 28, 2024.



## CAN-2409: Mechanism of action Please visit https://vimeo.com/822135123

1. CAN-2409 locally administered combined with oral prodrug 3. CAN-2409 induces CD8+ cytotoxic T cells Valacyclovir Inflammatory Tumor Dendritic cell B-cell mediators antigens Macrophag Fibroblast 0 0 CAN-2409 Cytotoxic metabolite Thymidine kinase 0000 C enzyme Valacyclovir CAN-2409 0 4. Local immunization yields systemic CD8+ T cell mediated 2. Localized cytolytic mechanism combined response against injected tumor and uninjected metastases with proinflammatory viral particles

CAN-2409 is an investigational product and its mechanism of action in humans has not been definitively established. This depiction of the CAN-2409 mechanism of action and the MoA video linked above are based on preclinical data and observations in clinical studies to date

## CAN-2409: Replication-defective adenoviral gene construct engineered for in situ vaccination against pan-solid tumors

- Proof of concept in patients with prostate cancer, non-small cell lung cancer, pancreatic cancer, and other solid tumors
- > 1,000 patients dosed
- Fast Track Designation in prostate cancer, non-small cell lung cancer, and pancreatic cancer
- Special Protocol Assessment (SPA) in localized prostate cancer

 Monotherapy activity of CAN-2409 in NSCLC patient: Nearly 50% decrease in tumor volume in 3 weeks



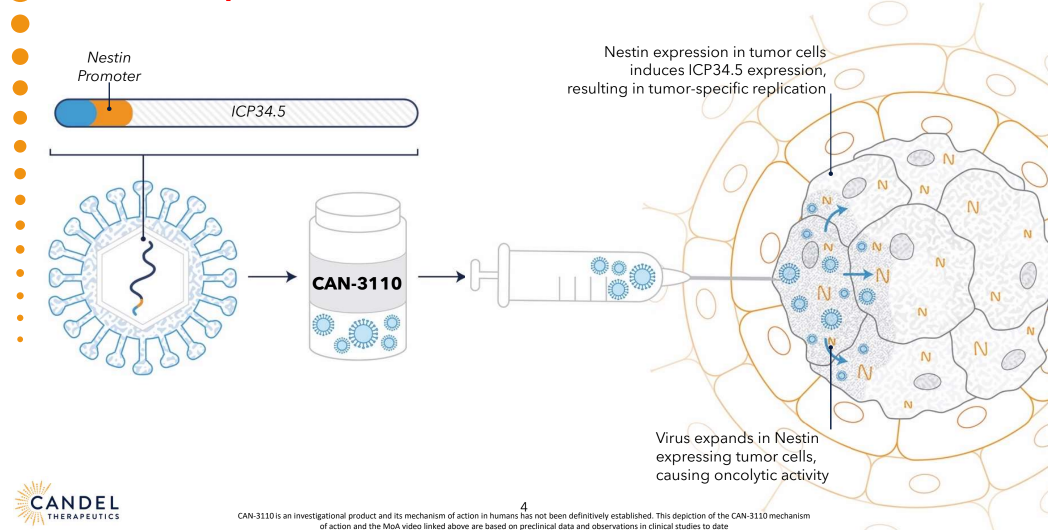
Day 0
Tumor Dimensions: 148 x
40 x 82 mm
(1012 vp dose)



Day 22 Tumor Dimensions: 100 x 34 x 75 mm



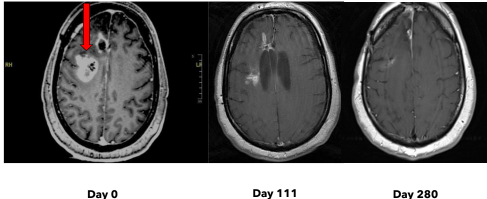
- CAN-3110: Mechanism of action
- Please visit https://vimeo.com/822133681



### CAN-3110: Replication-competent HSV-1 engineered to enhance selective killing of cancer cells while sparing healthy neighboring cells

- o Proof of concept in patients with recurrenthigh grade glioma (mostly glioblastoma)
- > 50 patients dosed
- Data published in Nature\*
- Fast Track Designation in recurrent highgrade glioma
- First cohort of patients treated with multiple injections of CAN-3110
- IND-enabling work in a second indication characterized by Nestin expression being planned

Monotherapy activity of CAN-3110 in recurrent high-grade glioma: Clinical effect on injected tumor and uninjected tumor



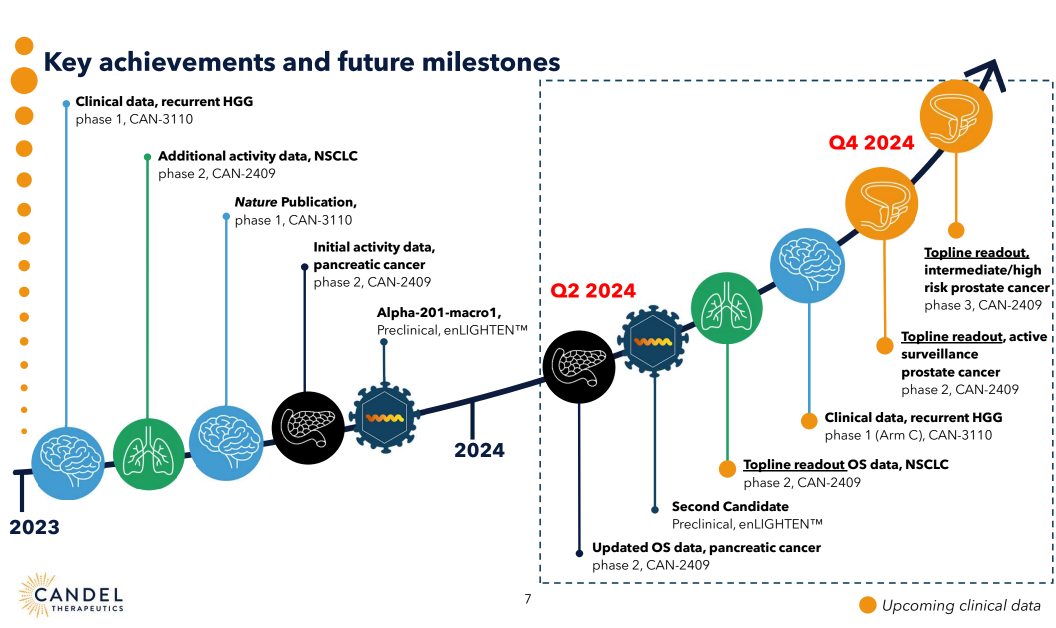
Patient back to work



## Pipeline focused on value creation

PROGRAM	INDICATION	PRECLINICAL	PHASE 1	PHASE 2	PHASE 3
Adenovirus Platform	ו				
<b>CAN-2409</b> Prostate Cancer	Localized, Intermediate/High Risk, Fast Track Designation (FDA), Special Protocol Assessment (FDA)				
	Active Surveillance				
CAN-2409 Lung Cancer	NSCLC + PD-1/PD-(L)1, Fast Track Designation (FDA)				
CAN-2409 Pancreatic Cancer	Borderline Resectable Pancreatic Adenocarcinoma, Fast Track Designation (FDA)				
HSV Platform					
<b>CAN-3110</b> Brain Cancer	Recurrent High-Grade Glioma, Fast Track Designation (FDA)				+ + +
enLIGHTEN™ Discovery Programs	Solid Tumors				





## Milestones for CAN-2409 this year (2024)

Study	Population	Potential success scenario			
Phase 2 randomized clinical trial in borderline resectable pancreatic cancer	US + EU5: ~13,000 patients/year	mOS 28.8 months vs. 12.5 months in control group  Disclosed in April 2024  ACHIEVED			
Phase 2 clinical trial in NSCLC with inadequate response to ICI	US + EU5: ~74,000 patients/yr	mOS > 10-13 months in per protocol population (Q2 2024)			
Phase 2b randomized clinical trial in localized low/intermediate risk prostate cancer (active surveillance population)	US + EU5: ~98,000 patients/yr	Significant improvement in progression-free survival (Q4 2024)			
Phase 3 randomized clinical trial in localized intermediate/high risk prostate cancer	US + EU5: ~110,000 patients/yr	Significant improvement in disease-free survival (Q4 2024)			



### Leadership team with decades of experience in oncology, immunology and drug development



Paul Peter Tak, MD, PhD, FMedSci

President & Chief Executive Officer









Charles Schoch, MBA, MSA, CPA

Interim Chief Financial Officer







Francesca Barone, MD, PhD

Chief Scientific Officer









**Garrett Nichols, MD, MS** 

Chief Medical Officer











Seshu Tyagarajan, PhD, RAC

Chief Technical and Development Officer











Susan Stewart, JD

Chief Regulatory Officer









### Research Advisory Board of premier thought leaders



James Allison, Ph.D. Chair of the Department of Immunology MD Anderson Cancer Center Director of the Parker Institute for Cancer Research 2018 Nobel Recipient





President and CEO Emeritus Dana-Farber Cancer Institute



Henry Brem, M.D. Director, Department of Neurosurgery Professor of Neurosurgery Johns Hopkins University



Roy Herbst, M.D., Ph.D. Chief of Medical Oncology Yale Cancer Center



Philip Kantoff, M.D. Former Chair, Department of Medicine Memorial Sloan Kettering Cancer Center



Gary Nabel, M.D., Ph.D. Chief Innovation Officer of OPKO and President/CEO of ModeX Therapeutics Former CSO Sanofi



Padmanee Sharma, M.D., Ph.D. Professor of Genitourinary Medical Oncology and Immunology MD Anderson Cancer Center



### **Candel at a glance**



- o CAN-2409: Off-the-shelf pan-solid tumor therapy, individualized anti-cancer immune response
  - Proof of concept in patients across multiple solid tumors
  - Fast Track Designation in NSCLC, pancreatic cancer, and prostate cancer
  - "Pipeline in a product" strategy advancing multiple programs in several large indications
  - Upcoming catalysts:
    - Topline phase 2 OS data in NSCLC (Q2 2024)
    - Topline phase 2b (Active Surveillance) and phase 3 (Intermediate/High Risk) prostate cancer clinical data (Q4 2024)



#### o CAN-3110: Oncolytic HSV-1 designed for tumor-specific replication

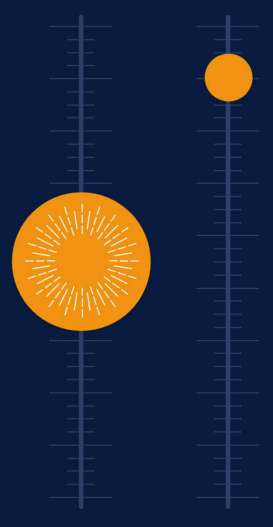
- · Proof of concept in patients with recurrent high-grade glioma published in Nature
- Fast Track Designation
- Opportunity for creation of "pipeline in a product" by expansion into indications beyond brain cancers
- Clinical and immunological biomarker data, evaluating repeat dosing regimen of CAN-3110 (H2 2024)



#### Corporate Highlights

- · Very experienced Executive Team and strong scientific support from high-profile Research Advisory Board
- · Cash and cash equivalents of \$35.4 million as of December 31, 2023; expected runway into Q4 2024
- IP protection: CAN-2409 (2034, method of use); CAN-3110 (2036, composition of matter); 12 years data exclusivity
- Low-cost manufacturing





## **CAN-2409**

• • • • • • • • • • •

Off-the-shelf therapy, individualized cancer response

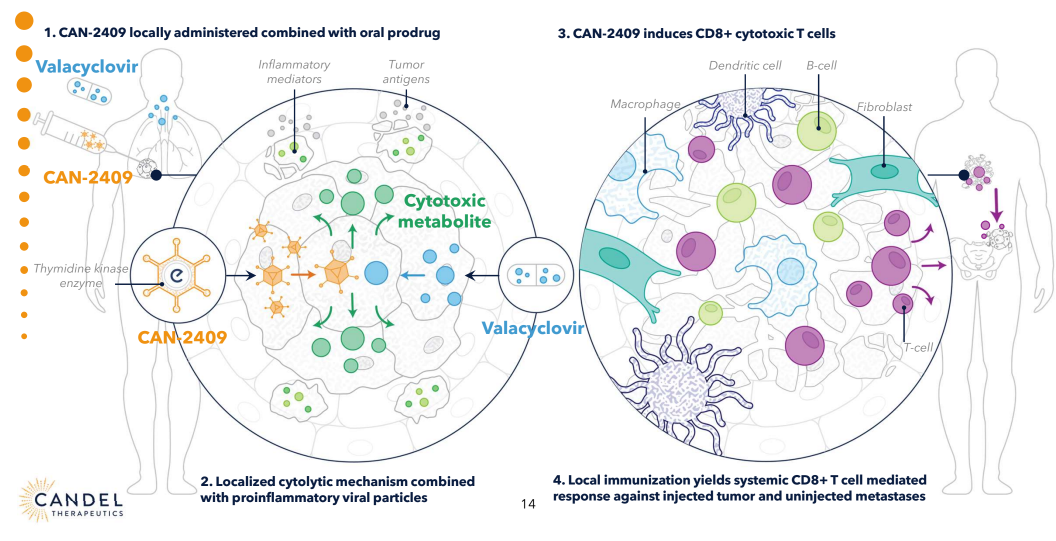


### **CAN-2409: Development program**

"Pipeline in a Product" approach advancing multiple programs in several large indications

Candidate	Indication	<b>Description</b>	Current Phase			Timing of Next	
			Phase 1	Phase 2	Phase 3	Milestone	
CAN-2409	Localized Prostate Cancer Intermediate / High Risk	<ul> <li>Fast-track status</li> <li>711 patients</li> <li>2:1 Randomization</li> <li>Primary Endpoint:</li> <li>Disease-free survival</li> </ul>			-	Q4:2024	
CAN-2409	Localized Prostate Cancer Active Surveillance	<ul> <li>187 patients</li> <li>2:1 Randomization</li> <li>Primary Endpoint:     Progression-free     survival</li> </ul>				Q4:2024	
CAN-2409 +PD-1/PD-(L)1	Non-Small Cell Lung Cancer	<ul> <li>Fast-track status</li> <li>80 patients</li> <li>Primary Endpoint:         Response by RECIST         criteria and disease         control rate</li> </ul>				Q2:2024	
CAN-2409	Borderline Resectable Pancreatic Adenocarcinoma	<ul> <li>Fast-track status</li> <li>13 patients</li> <li>1:1 Randomization</li> <li>Primary Endpoint:</li> <li>Safety and survival rate at 24 mos</li> </ul>				Q2:2024	
INDEL IERAPEUTICS		13	3				

### CAN-2409: Systemic immunotherapy delivered intratumorally



### **CAN-2409: Prostate cancer opportunity**

## Incidence of localized prostate cancer in the US by risk level



- Prostate cancer is the second leading cause of cancer death among men in the US
- Global annual incidence projected to rise from 1.4 million in 2020 to 2.9 million by 2040 due to aging populations and increase screening^
- No new treatments approved for newly diagnosed, localized prostate cancer during the last > 30 years
- Significant opportunity for new treatment for both the active surveillance and intermediate/high risk populations with a favorable tolerability profile and potential to reduce progression and/or recurrence
- Prostate cancer therapy market globally was estimated at \$13B in 2022 and is expected to grow to \$21B by 2028\*

#### Target label for CAN-2409#

- Indicated in newly diagnosed localized prostate cancer in combination with radiotherapy +/- short-term ADT, in patients with intermediate- to high-risk disease
- Indicated in newly diagnosed localized prostate cancer in patients with NCCN-defined low-risk disease, or patients with intermediate-risk disease undergoing active surveillance



^Source: James ND et al. Lancet 2024 Apr 4:S0140-6736(24)00651-2 #Consistent with market research combined with interviews with 22 KOLs (12 US; 10 EU) and 16 US payors. May 2023

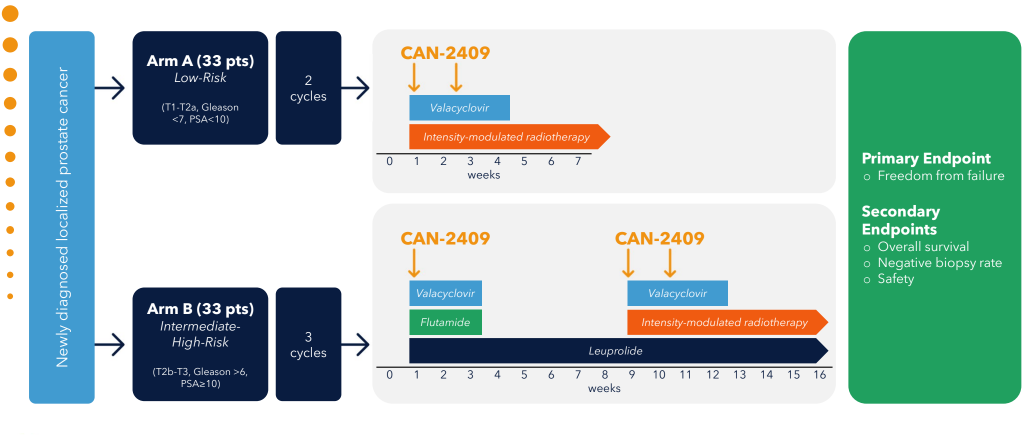
\*Source: EvaluatePharma, accessed May 2023

### **CAN-2409: Prostate cancer opportunity in active surveillance population**

- Active surveillance population (US) ~40,000 pts / year
- There is a ~30% risk of reporting cancer-specific anxiety in this population
- o 30-50% of patients will still undergo radical therapy due to disease progression within 5 yrs
  - Patients will then face side effects and complications of radical therapy, including urinary incontinence and erectile dysfunction
  - Next, one third of these patients will still have recurrence after radical therapy
- Lifetime cost of active surveillance estimated at ~\$21,000 without providing therapeutic treatment
- o Opportunity for development of a well-tolerated therapy that stops progression of the disease



## Completed phase 2a clinical trial of CAN-2409 combined with radiotherapy +/- androgen deprivation therapy

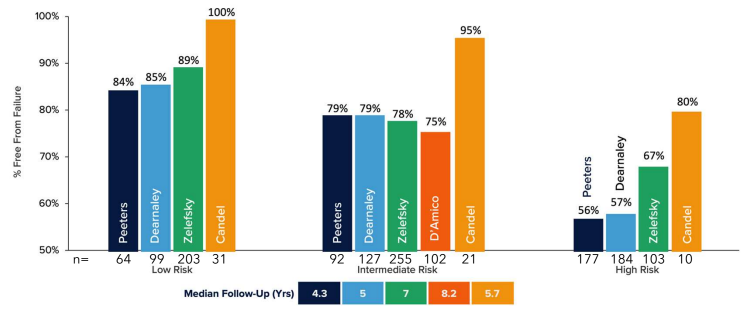




## Completed phase 2a trial demonstrated consistently improved freedom from failure rates in newly diagnosed, localized prostate cancer

- o Pathological complete response (pCR) observed in 93% of the biopsies available at 2yrs (37%-73% in control populations)
- Low risk patients achieved PSA < 2ng/ml in 77% of CAN-2409 treated patients versus 58% in control populations
- Significant increase in activated CD8+ T cells compared to baseline (p=0.0003)

Clinical evidence supports ongoing phase 3 clinical trial of CAN-2409 in combination with SoC as first line treatment

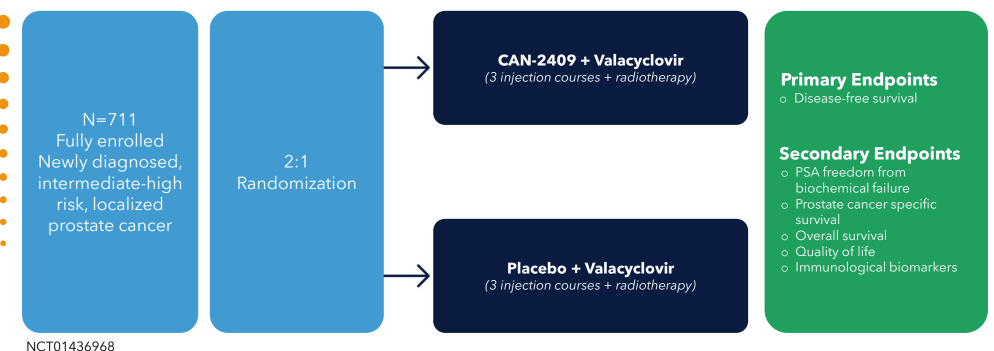


Failure was measured from the start date of treatment until the date of treatment failure defined as clinical failure or biochemical failure, whichever first, according to the Phoenix ASTRO consensus (Nadir +2)



### Fully accrued phase 3 clinical trial of CAN-2409 in patients with prostate cancer - Newly diagnosed, intermediate/high risk population

Pls: Dr. T. DeWeese (JHU) and Dr. P. Scardino (MSKCC)

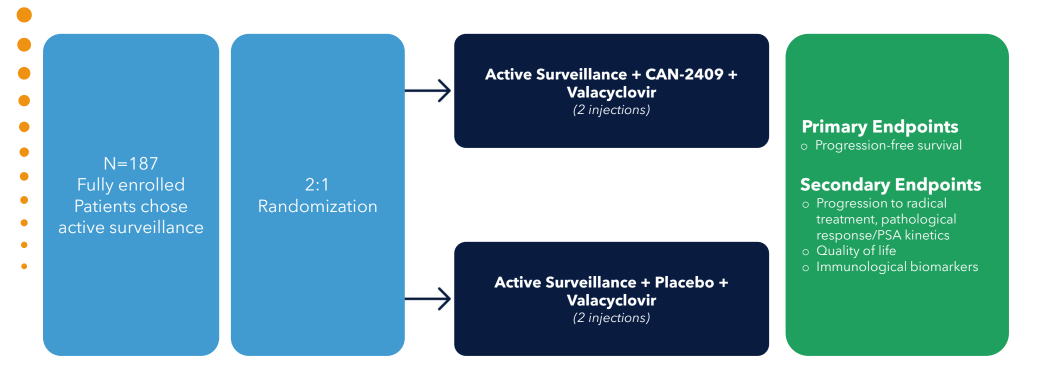


**Conducted under agreement with FDA under Special Protocol Assessment** 



## Fully accrued phase 2b clinical trial of CAN-2409 in patients with prostate cancer – Active surveillance population

PI: Dr. S. Eggener (UChicago)





## Ongoing phase 2b clinical trial: CAN-2409 is generally well-tolerated Monotherapy – Active surveillance population

~ 33% patients experienced flu-like symptoms

< 1% infections requiring hospitalization

Study is still blinded 187 patients treated 362 injections performed

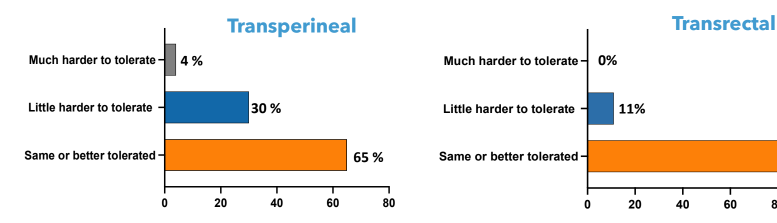
Most samman DT (> -E9/)		n=187			
Most common PT (>=5%)	1	2	3	4	Total (%)
Flu-like symptoms	40 (21)	20 (11)	1 (1)	0	61 (33)
Chills	39 (21)	13 (7)	1 (1)	0	53 (28)
Fever	39 (21)	9 (5)	1 (1)	0	49 (26)
Fatigue	27 (14)	10 (5)	1 (1)	0	38 (20)
Elevated AST/ALT	28 (15)	3 (2)	1 (1)	0	32 (17)
Elevated Creatinine	23 (12)	5 (3)	1 (1)	2 (1)	31 (17)
Headache	20 (11)	5 (3)	0	0	25 (13)
Urinary tract infection	1 (1)	15 (8)	2 (1)	0	18 (10)
Nausea	12 (6)	4 (2)	0	0	16 (9)
Low Hemoglobin	15 (8)	0	0	0	15 (8)
Diarrhea	10 (5)	3 (2)	0	0	13 (7)
Malaise	10 (5)	2 (1)	0	0	12 (6)
Hematuria	12 (6)	0	0	0	12 (6)
Urinary frequency	9 (5)	2 (1)	0	0	11 (6)
Urinary tract pain	6 (3)	3 (2)	0	0	9 (5)
Urinary urgency	7 (4)	2 (1)	0	0	9 (5)
Elevated Alkaline Phosphatase	8 (4)	1 (1)	0	0	9 (5)
Elevated Bilirubin	7 (4)	3 (2)	0	0	10 (5)



## Ongoing phase 3 clinical trial: Most patients tolerate intraprostatic injection same or better than prostate biopsy

Patient questionnaire substudy n=32

In total > 2000 intraprostatic injections
(40% transperineal; 56% transrectal; 4% not reported)
"How did you tolerate the study procedure as compared to a prostate biopsy?"





Aguilar L. 28th Annual Prostate Cancer Foundation Scientific Retreat, October 2021

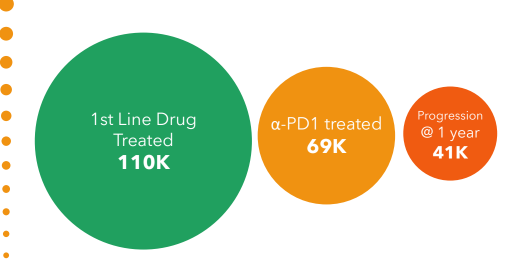
80

89 %

100

### CAN-2409: Non-small cell lung cancer opportunity

#### Incident advanced NSCLC in the US1

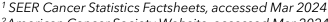


- Lung cancer is the most commonly diagnosed cancer in the US with NSCLC representing over 80% of all lung cancer diagnoses<sup>2</sup>
- Majority of NSCLC patients without actionable mutations are treated with immune checkpoint inhibitors (ICI) as 1<sup>st</sup> line treatment
- After one year of ICI treatment, 60% of anti-PD1 treated patients will have progressive disease<sup>3</sup>
  - In ICI inadequate responders with SoC docetaxel<sup>4</sup>
    - Median overall survival (mOS) <12 months</li>
- Significant opportunity to improve response to ICIs by teaching the immune system how to recognize the cancer cells
- NSCLC cancer therapy global market was estimated at \$32B in 2023 and is expected to grow to \$52B by 2028<sup>5</sup>

Target label for CAN-2409: Indicated as adjunct to standard-of-care immune checkpoint inhibitor treatment (+/- chemotherapy) in stage III / IV NSCLC patients not responding to first-line or later ICI treatment#

Upside to this target population as ~25% of patients treated in the 2L setting receive an anti-PD-1/L1

THERAPEUTICS



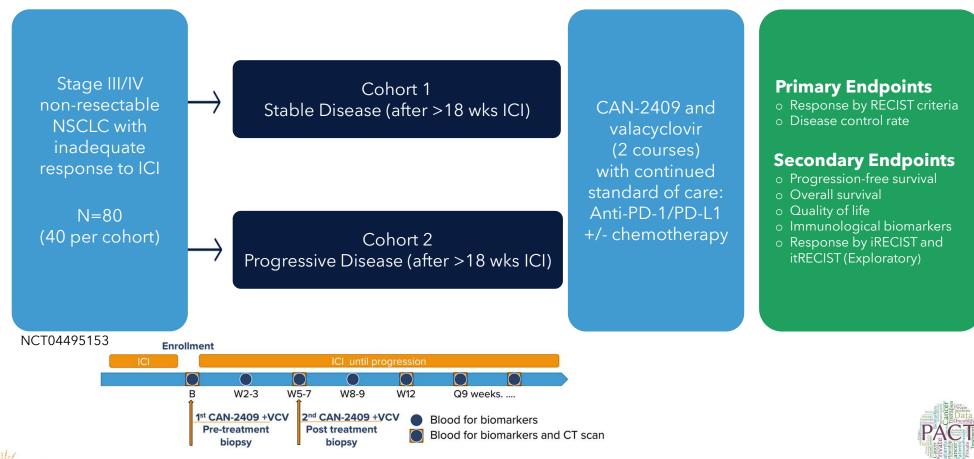
<sup>&</sup>lt;sup>2</sup>American Cancer Society Website, accessed Mar 2024

<sup>&</sup>lt;sup>3</sup> Gandi L et al. NEJM 2018; 378:2078-92

<sup>&</sup>lt;sup>4</sup> Reckamp K et al. J Clin Onc 2022;40:2295-2306

<sup>&</sup>lt;sup>5</sup> EvaluatePharma, accessed May 2023

### Ongoing phase 2 clinical trial of CAN-2409 + continued ICI in stage III/IV **NSCLC** patients with an inadequate response to ICI

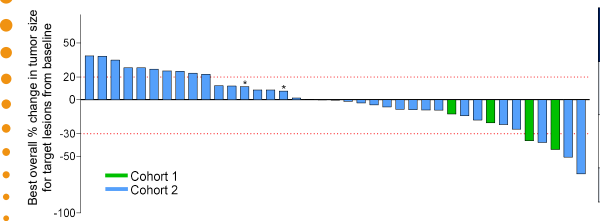




THERAPEUTICS

#### **Evidence that CAN-2409 can control disease**

Most patients entering clinical trial with progressive disease despite immune checkpoint inhibitor treatment achieved disease control after administration of CAN-2409



Coh	N	PR	SD	PD	ORR	DCR	DoR for PR <sup>2</sup>	SD duration <sup>2</sup>
1	4 <sup>1</sup>	2	2	0	50%	N/A	7.7 mo. (2.7+ to 12.8+)	4.9 mo (3.6+ to 6.2)
2	35	3	20	12	9%	66%	6.1 mo (2.8+ to 16.3)	3.9 mo (1.4+ to 14.5)
Total	39	5	22	12	13%	N/A		

 $<sup>^{\</sup>rm 1}$  An additional evaluable patient in Cohort 1 had a pending central read at time of data cutoff

Data shown based on blinded independent central review (BICR) of radiographic data per RECIST 1.1 in evaluable patients

Evaluable patients are those who received 2 courses of CAN-2409 + prodrug (valacyclovir) and completed the 12-week treatment window

\*Disease progression due to a new lesion

PR: partial response, SD: stable disease, PD: progressive disease, ORR: objective response rate, DCR: disease control rate, DoR: duration of response



<sup>&</sup>lt;sup>2</sup> Median (range) for DoR and SD duration

<sup>+</sup> indicates response was ongoing at date of last follow up

## Partial response in extended lung mass with durable post-treatment tumor regression after CAN-2409 treatment (> 2 years, ongoing)

#### **PA-003 (Cohort 1)**

73M, Stage III non-squamous NSCLC diagnosed Jan'20 PD-L1<1%

Initial therapy: pembro + carbo + pemetrexed Feb'20 Maintenance: pembro + pemetrexed from Jun'20 which continued on-trial

OS 24 mo (ongoing as of LFV)

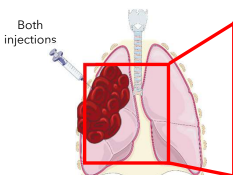
#### **Legend**

LFV: last follow up visit

#### **RECIST target lesions (red)**

LN = lymph node; LA = long axis; SA = short axis

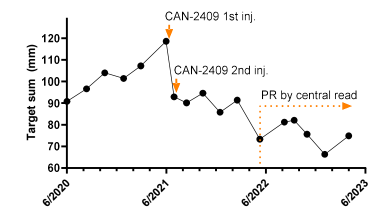
#### Baseline

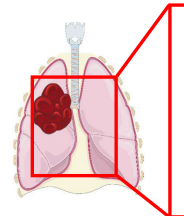




Right middle lobe LA: 118.6 mm Target lesion

Site of both injections







6 Months LA: 85.8 mm Target lesion



24 Months

LA: 76.5 mm Target lesion



### Patient with continued tumor shrinkage after CAN-2409 treatment

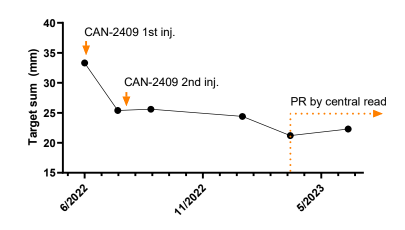
#### **VB-007 (Cohort 1)**

84F, Stage IV non-squamous NSCLC diagnosed Aug'21 PD-L1 < 1%; SMARC4 alteration

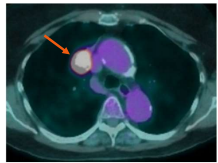
Initial therapy: platinum-based chemotherapy + pembro

Maintenance: pembro which continued on-trial

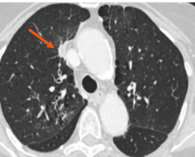
OS 12.1 mo (PR is ongoing as of LFV)



**FDG-PET** 







**Treatment Naïve** 

THERAPEUTICS

**Prior to 1st injection** 

**Post 2nd injection** 

1 year after 1st injection

Scans kindly provided by Wade lams, MD

### Local injection induces systemic anti-tumor activity

**Evidence of abscopal effect, survival > 27 months (ongoing) after CAN-2409 treatment** 

#### **NY-007 (Cohort 2)**

74M, Stage IV non-squamous NSCLC diagnosed Feb'19 PD-L1 < 1%

Initial therapy: cisplatin/etoposide Feb-Jul'19

Maintenance: nivolumab from Sep'19, continued on-

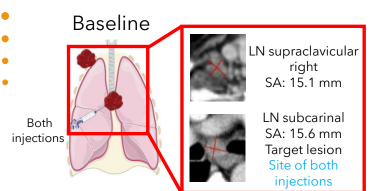
study

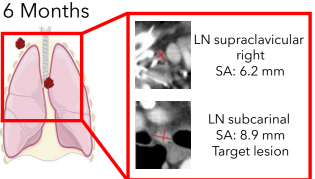
OS 27.9 mo (ongoing as of LFV)

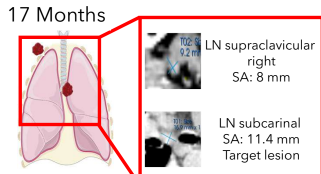
#### Legend

#### **RECIST target lesions (red)**

LN = lymph node; LA = long axis; SA = short axis









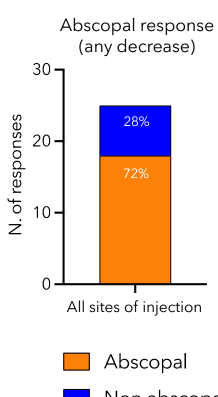
Schematics to show general lesion injection orientation; not to scale 28

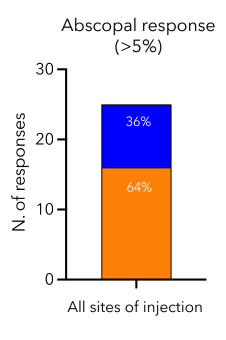
Note: From ongoing phase 2 NSCLC trial as of cutoff date, 1 Aug 2023.

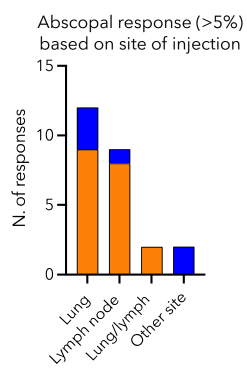
Data on file, September 2023

### Local injection induces systemic anti-tumor activity

Regression of uninjected lesions in about two-thirds of evaluable patients presenting with multiple lesions







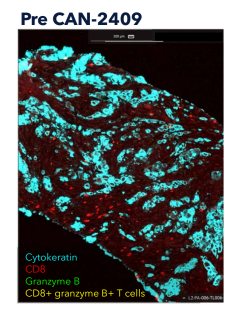
Non abscopal

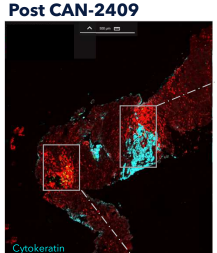
Center and right panel: Decrease of at least 5% observed in at least one noninjected lesion

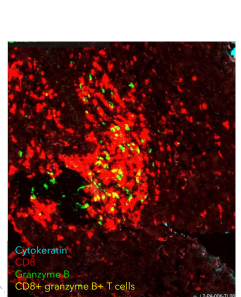


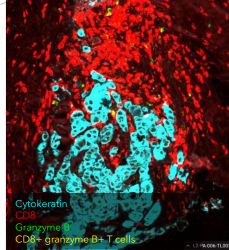
# CAN-2409 induces expansion of CD8+ cytotoxic tumor infiltrating lymphocytes in the tumor microenvironment

**PA006** 

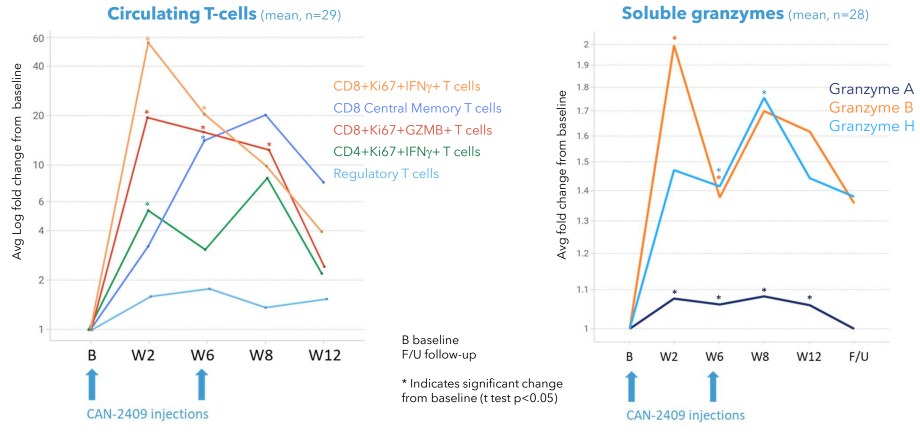








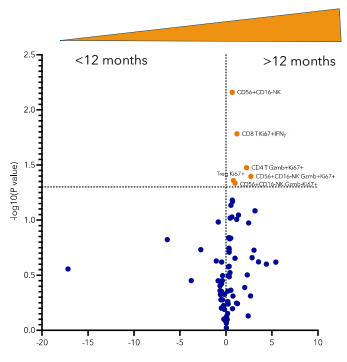
## CAN-2409 significantly increases frequency of circulating cytotoxic T cells and serum levels of soluble granzymes





## Changes in immune cells in peripheral blood after 2<sup>nd</sup> injection of CAN-2409 are associated with subsequent prolonged survival

#### Changes in circulating cells post 2nd injection



Multiparameter flow cytometry Fold changes between 1st and 2nd injection in short (< 12months; n=6) and long (> 12 months; n=11) survivors



## Bronchoscopic delivery of CAN-2409 is a simple extension of existing care for NSCLC patients and is well-tolerated

Majority of lung and thoracic lymph node lesions are accessible for outpatient bronchoscopic injection



- Therapeutic delivery tool based on extensive experience with bronchoscopic biopsy, a routine outpatient procedure (15 min, outpatient setting)
- Transbronchial needle injection (TBNI) presents similar complication rate as biopsy (extremely rare)
- Latest generation of TBNI includes ultrasound-guided transbronchial injection of lymph nodes and robotic bronchoscopy (already in use in LuTK02)



### Systemic immunotherapy delivered intratumorally

- o Intra-tissue injection is a proven strategy for in situ vaccination
  - Optimal benefit/risk by minimizing systemic toxicity
  - Systemic immune response: not all metastases need to be injected
- Durable responses after only 2-3 administrations
- Procedure is well tolerated by patients:
  - Administration to prostate takes 15-20 min in outpatient setting, often better tolerated than prostate biopsy, which
    is a routine procedure in urology
  - Administration to lung tumor takes 15-20 min in outpatient setting via bronchoscopy, which is a routine procedure
    in pulmonary medicine
  - For future indications, any location can be reached using image-guided injection, robotic delivery etc.
  - Procedures with proven benefit/risk, patient tolerability, and cost-effectiveness will be implemented by clinicians
    - Stem cell transplantation, implant radiotherapy, interventional radiology, interventional cardiology



## **Encouraging safety data, clinical activity and immunological changes after CAN-2409 in NSCLC**

#### Initial data suggests 12-month survival is consistent with an increased tail on the maturing survival curve

- Encouraging number of long survivors suggests CAN-2409 may induce a new state of functional immunosurveillance and durable disease control in a subset of patients
- Of the 40 evaluable patients, 15 patients have lived ≥ 12 months; of these, 10 have lived > 18 months, of whom 70% (7/10) were alive as of last follow up. All 4 patients (100%) with OS > 24 months were alive at last follow up, with the longest reaching 31.7 months (data cutoff Aug 1, 2023)
- An additional 18 (out of the 40 evaluable) patients are also alive but have not yet reached 12 months of follow up

#### Negative or low PD-L1 status appears to be associated with long survival in CAN-2409 treated patients

- Many patients treated with CAN-2409 have had long survival (≥ 12 months) despite having disease features generally associated with advanced disease and reduced likelihood to benefit from immune checkpoint inhibitor therapy, such as low or negative PD-L1 expression
- Biomarker data suggests association between immune cell activation and survival
  - Scope of antitumoral immune response broadened through demonstration by CAN-2409 to engage the humoral arm of the immune system
  - Increase observed in effector/cytotoxic T cells and NK cells in peripheral blood after the second CAN-2409 administration

#### Topline overall survival data for Cohort 2 expected in Q2 2024



### CAN-2409: Pancreatic ductal adenocarcinoma opportunity

Incidence of pancreatic ductal adenocarcinoma in the US by risk level<sup>1</sup>



- Limited available treatment options for patients suffering from pancreatic cancer beyond resection when possible and chemotherapy
- Borderline resectable disease: median overall survival <18 months (with neoadjuvant chemoradiation and resection)<sup>2</sup>
- Metastatic disease: median overall survival ~11 months (with FOLFIRINOX)<sup>3</sup>
- Significant opportunity to improve response rates by teaching the immune system how to recognize cancer cells
- Pancreatic cancer therapy global market was estimated at \$800 M in 2022 and is expected to grow to \$3.5 B by 2028<sup>4</sup>



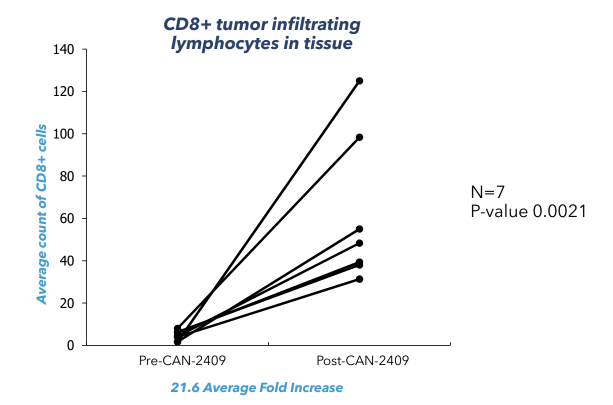
<sup>&</sup>lt;sup>1</sup> Park W et al. JAMA 2021;326:851-862

<sup>&</sup>lt;sup>2</sup> Versteijne E et al. J Clin Onc 2020; 38:1763-1773

<sup>&</sup>lt;sup>3</sup> Conroy T et al. NEJM 2011; 364:1817-1825

<sup>&</sup>lt;sup>4</sup> Source: EvaluatePharma, accessed May 2023

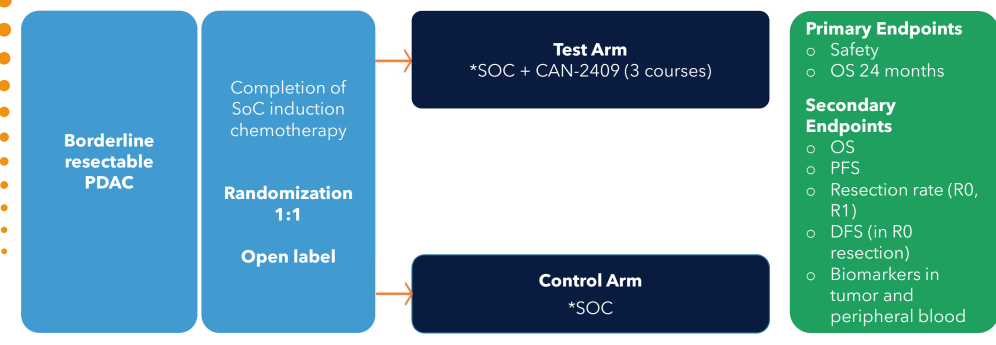
## Completed phase 1 clinical trial of CAN-2409 in pancreatic ductal adenocarcinoma: Infiltration by CD8+ tumor infiltrating lymphocytes





### Randomized phase 2 clinical trial of CAN-2409 in borderline resectable pancreatic ductal adenocarcinoma (PDAC)

Reflecting v5/v6 of protocol (data collected to date reflects this design)

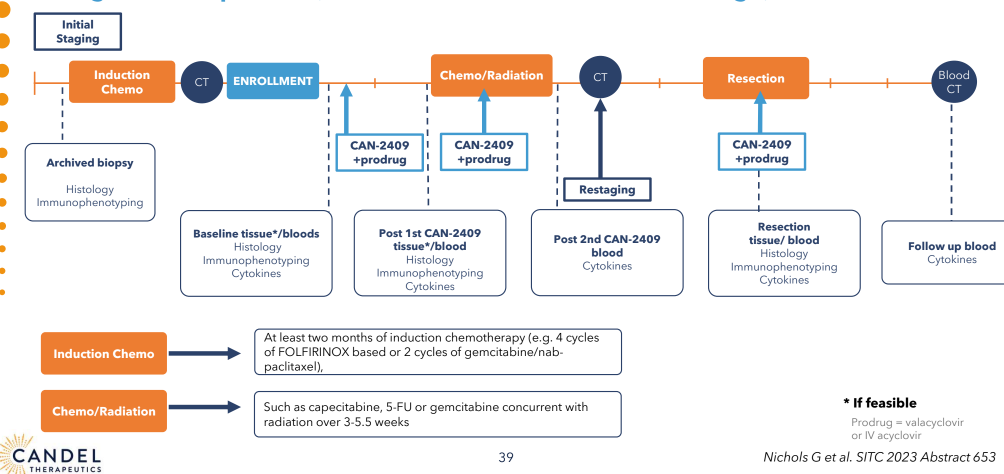


\*SOC= Chemoradiation + Resection

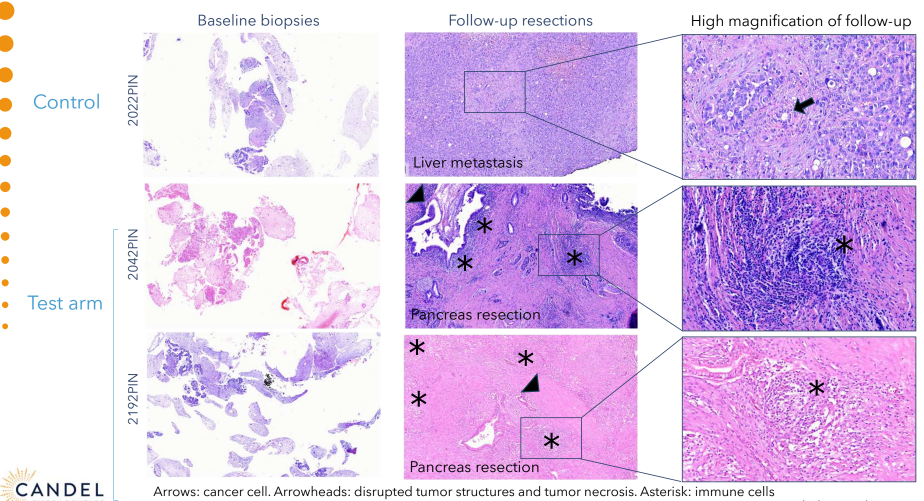


# SOC treatment timeline in non-metastatic PDAC and timing of CAN-2409 injections

Through v5/v6 of protocol (data collected to date reflects this design)



## CAN-2409 induces formation of dense lymphocyte aggregates, disruption of tumor structures, and necrosis in PDAC



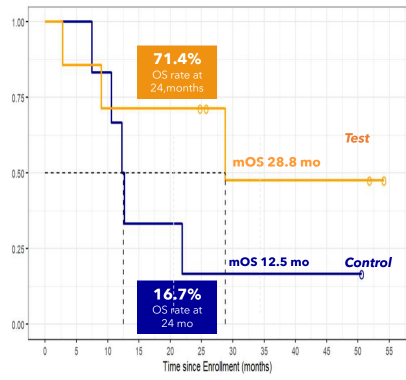
Nichols G et al. SITC 2023 Abstract 653

### Overall survival in borderline resectable PDAC patients

Data as of 3/29/2024

	PCN	Arm	Surgical result	pStage #	Date of last follow-up	OS mo (enrollment)	OS mo (diagnosis)	Alive (A) / Dead (D)
	2022PIN	С	Unresected	IV	6/16/2020	10.6	17.2	D
	2072PIN	С	Unresected	N/A*	11/13/2020	12.7	52.4	D
	2092POS	С	Unresected	N/A*	7/23/2020	7.5	10.3	D
	2052PLB	С	Resected	IIA	10/3/2020	12.3	16.9	D
	2152PLB	С	Resected	IIB	9/25/2022	21.9	26.8	D
	2112PLB	С	Resected	N/A*	3/28/2024	50.6+	54.8+	А
	2102PLB	Т	Unresected	IV	9/7/2020	9.0	13.7	D
•	2162PLB	Т	Unresected	N/A*	6/9/2021	2.8	8.3	D
•	2042PIN	Т	Unresected	IV	2/22/2024	54.2+	61.7+	А
•	2172PIN	Т	Unresected	N/A*	1/14/2024	28.8	34.7	D
•	2082PLB	Т	Resected	IA	2/26/2024	51.9+	57.0+	А
	2182PLB	Т	Resected	IB	3/04/2024	25.8+	32.3+	А
	2192PIN	Т	Resected	IA	3/20/2024	24.8+	30.3+	А

#### Time since enrollment



Censored = alive, still under follow-up

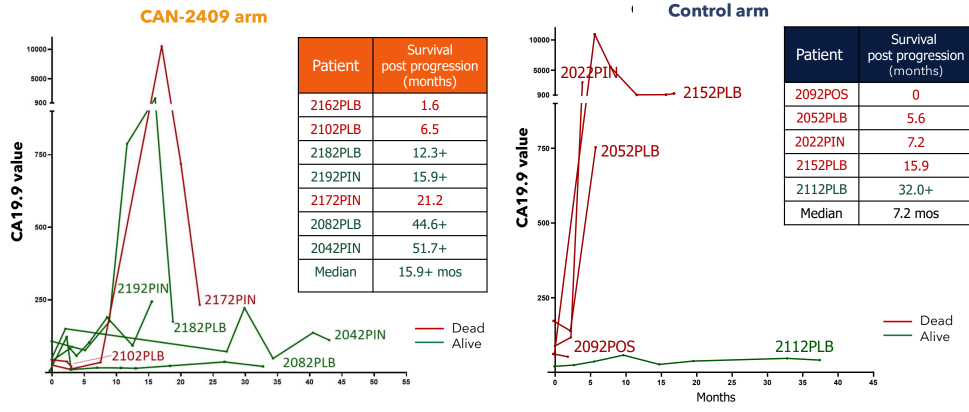
# pathologic tumor stage at resection

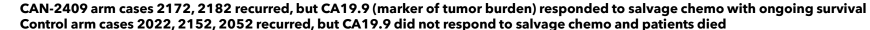


Survival probability

<sup>\*</sup>Refer to slide with details on surgical status

## CA19-9 biomarker response associated with ongoing survival in CAN-2409 arm, but not in control arm, in patients with progressive disease



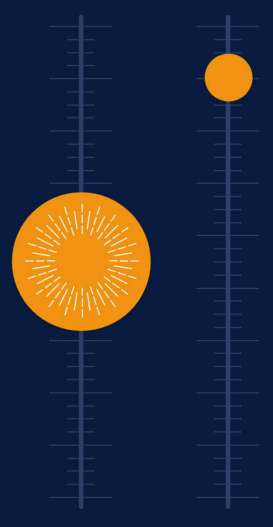




## **Encouraging safety data, clinical activity and immunological changes after CAN-2409 in pancreatic cancer**

- Notable improvements in estimated median overall survival of 28.8 months after experimental treatment with CAN-2409 versus only 12.5 months in control group
- At 24 months, survival rate was 71.4% in CAN-2409 treated patients versus only 16.7% in the control group after chemoradiation. At 36 months, estimated survival was 47.6% in the CAN-2409 group versus 16.7% in the control group
- Multiple injections of CAN-2409 were generally well tolerated, with no dose-limiting toxicities and no cases of pancreatitis
- In patients with progressive disease, there was a CA19-9 and survival response to salvage chemotherapy in the CAN-2409 arm but not in control arm
- CAN-2409 activates the immune response in the pancreatic tumor and peripheral blood



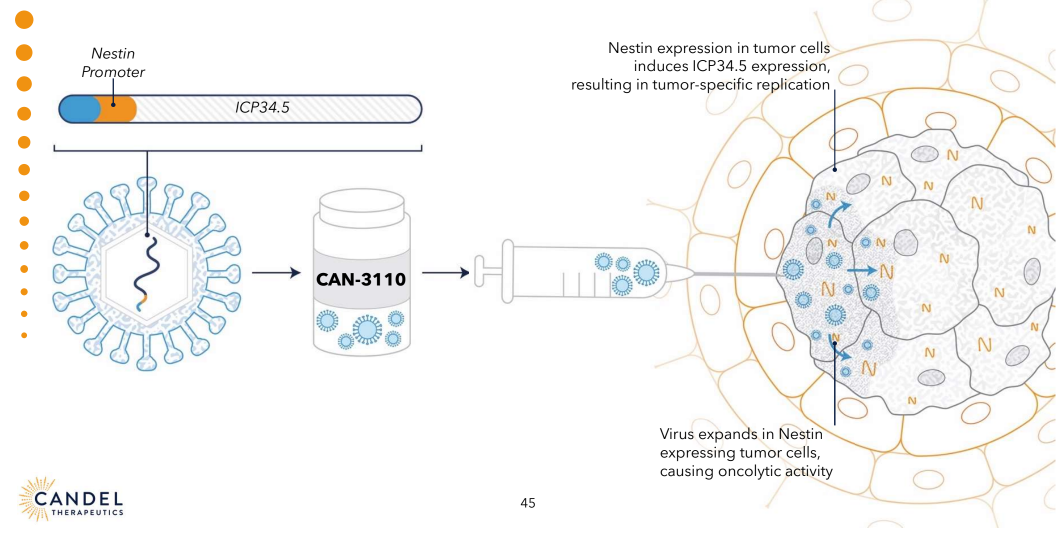


## **CAN-3110**

Oncolytic virus with tumor-specificity



### **CAN-3110:** Replication-competent HSV with tumor-specificity



### **CAN-3110: High-grade glioma opportunity**

#### Prevalence of glioblastoma in the US<sup>1</sup>







- Glioblastoma, the most common form of highgrade glioma, is a rare and often deadly cancer
- Fewer than 10% of patients survive > 5 years past initial diagnosis<sup>2</sup>
- Median overall survival < 6-9 months in recurrent high-grade glioma<sup>3</sup>
- Current standard of care includes surgical resection with few available therapeutic options
- Significant opportunity to improve survival by teaching the immune system how to recognize the cancer cells and turn 'cold tumors' into 'hot tumors'

1 Miller KD et al. CA Cancer J Clin 2021;71:381-406 2 Stupp R et al. Lancet Oncol. 2009;10:459-466 <sup>3</sup> vanLinde MC et al. J Neuro Onc 2017;135:183-192



### Phase 1b clinical trial of CAN-3110 in recurrent high-grade glioma

PI: Dr. E. Antonio Chiocca (Brigham & Women's)

Patients with recurrent high-grade glioma

Lesions ≥ 1.0 cm

rm A

#### **Dose escalation (Cohort I-IX)**

Single stereotactic injection of CAN-3110

3+3 dose escalation  $1 \times 10^6$  to  $1 \times 10^{10}$  PFU in half-log increments 30 patients dosed

#### **Dose expansion (Cohort X)**

1x10<sup>9</sup> PFU 11 patients dosed

\rm B

#### **Pre-Administration of Cytoxan**

3 x 10<sup>8</sup> PFU 6x 10<sup>9</sup> PFU 9 patients dosed



Repeat Dosing (up to 6) +1 x 10<sup>8</sup> PFU x 6 doses +1 x 10<sup>9</sup> PFU x 6 doses 12 patients targeted

#### **Primary Endpoints**

- Safety
- o Determine maximum tolerated dose

#### **Secondary Endpoints**

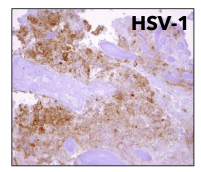
- o Immunological biomarkers
- MRI assessment of disease and progression free survival
- MRI alteration of permeability and flow at injection site



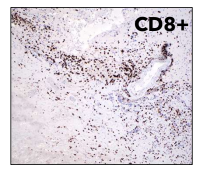
### CAN-3110 treatment in patients with recurrent high-grade glioma

Persistent HSV antigen expression in injected tumors as well as uninjected tumors associated with CD8+ T cell infiltration after single CAN-3110 injection

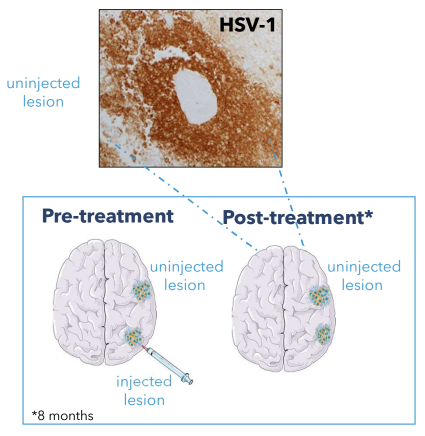
injected lesion



HSV1 antigen 6 weeks after injection of  $1x10^6$  pfu  $1.79 \times 10^6$  copies of viral DNA/mg  $2.97 \times 10^5$  copies of viral RNA transcript (ICP22)/mg



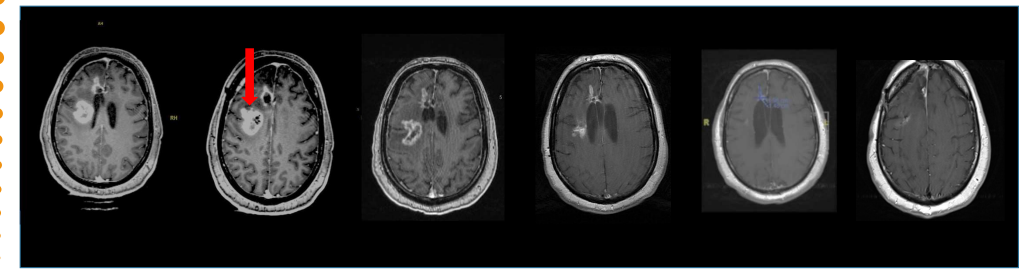
Infiltration by CD8+ cytotoxic T cells (tumor infiltrating lymphocytes)





### Monotherapy activity of CAN-3110 in recurrent high-grade glioma (arm A)

Clinical effect on injected tumor and uninjected tumor



Baseline

Day 0
Black hole within tumor image is injection site 10° PFU dose

Day 56 Reduction in contrast area with no additional treatment Day 111
Patient back to work

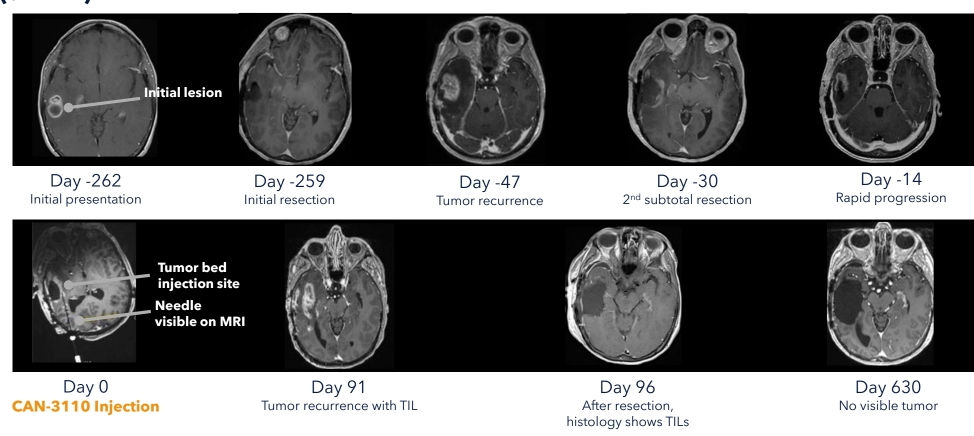
Day 168

Day 280

56 YOM, IDH wild-type, MGMT partially methylated, right frontal mesial lesion initially treated with GTR, chemoradiation. Recurrences at two sites.



## Durable response for 2 years after CAN-3110 in recurrent high-grade glioma (arm A)

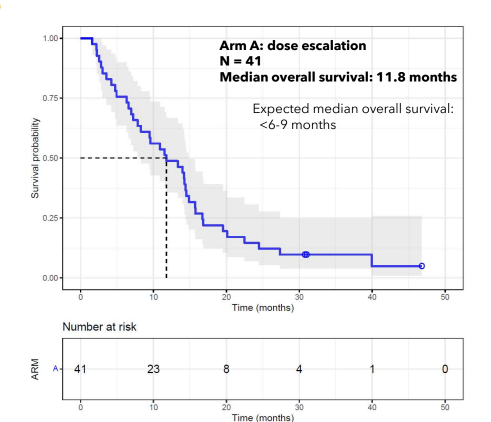


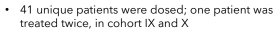
61 YOF, IDH wild-type, MGMT methylated glioblastoma, right temporal lesion initially treated with surgery, chemoradiation and temozolomide CAN-3110 dose: 10<sup>8</sup> PFUs. Patient passed away as passenger in a motor vehicle accident on day 717.

ANDEL

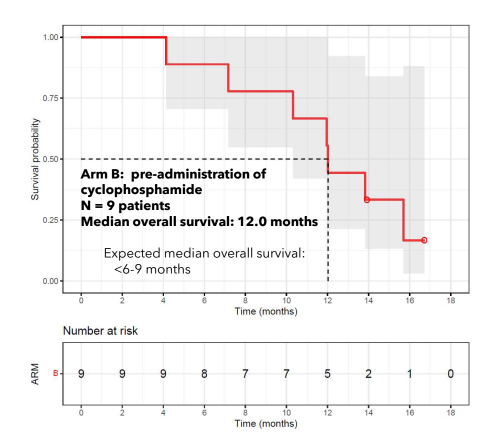
THERAPEUTICS

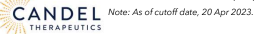
## **Encouraging overall survival in recurrent high-grade glioma after single injection of CAN-3110**



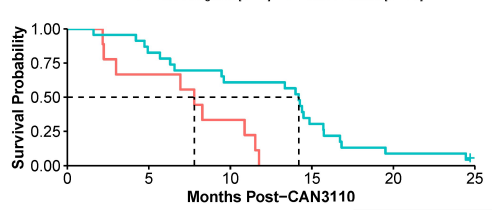


• A total of 50 unique patients





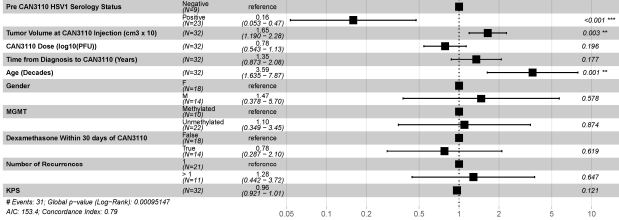
## Prolonged survival after CAN-3110 treatment is associated with HSV1 seropositivity



HSV1 Negative [n = 9] + HSV1 Positive [n = 23]

HSV2 serology status is not associated with survival

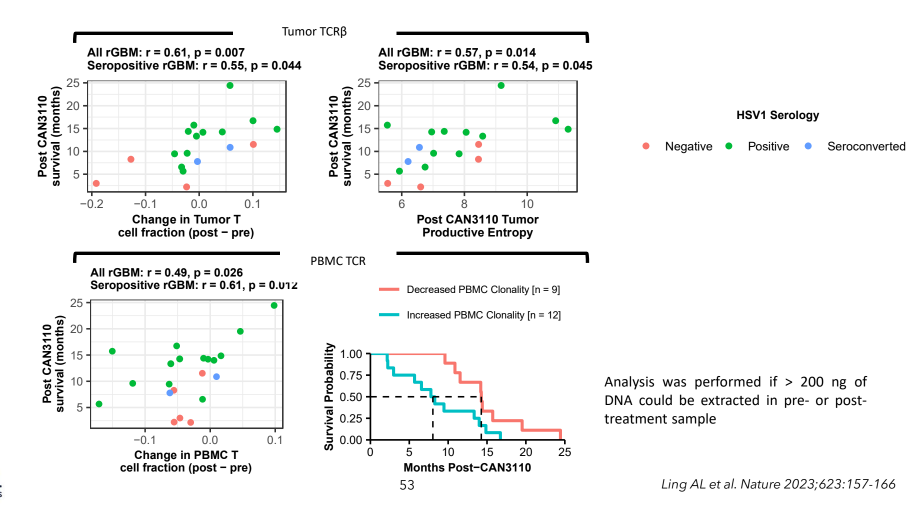
COxPH Hazard Ratios





Ling AL et al. Nature 2023;623:157-166

### Changes in T cell fractions and TCR $\beta$ diversity correlate with survival after CAN-3110 treatment

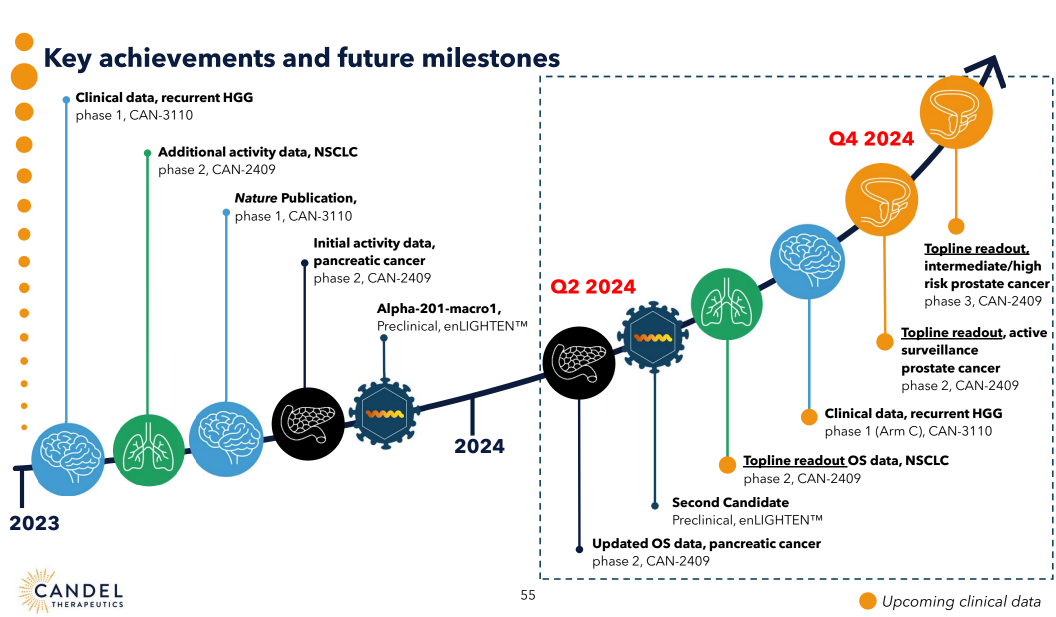




## **Encouraging safety data, clinical activity and immunological changes after CAN-3110 in recurrent high-grade glioma (glioblastoma)**

- Monotherapy treatment with CAN-3110 in rHGG is well tolerated and associated with doubling of expected median overall survival
- Immunological changes in the tumor microenvironment are associated with improved survival and HSV1 seropositivity
- o First six patients have been dosed in cohort C (fully funded by the Break Through Cancer foundation)
- Repeated injections of CAN-3110 (up to six) are safe and well tolerated
- Significant decrease in tumor cells and increased immune cell infiltration after CAN-3110 administration





### Candel at a glance



- o CAN-2409: Off-the-shelf pan-solid tumor therapy, individualized anti-cancer immune response
  - Proof of concept in patients across multiple solid tumors
  - · Fast Track Designation in NSCLC, pancreatic cancer, and prostate cancer
  - · "Pipeline in a product" strategy advancing multiple programs in several large indications
  - Upcoming catalysts:
    - Topline phase 2 OS data in NSCLC (Q2 2024)
    - Topline phase 2b (Active Surveillance) and phase 3 (Intermediate/High Risk) prostate cancer clinical data (Q4 2024)



- o CAN-3110: Oncolytic HSV-1 designed for tumor-specific replication
  - · Proof of concept in patients with recurrent high-grade glioma published in Nature
  - Fast Track Designation
  - Opportunity for creation of "pipeline in a product" by expansion into indications beyond brain cancers
  - Clinical and immunological biomarker data, evaluating repeat dosing regimen of CAN-3110 (H2 2024)



#### Corporate Highlights

- · Very experienced Executive Team and strong scientific support from high-profile Research Advisory Board
- · Cash and cash equivalents of \$35.4 million as of December 31, 2023; expected runway into Q4 2024
- IP protection: CAN-2409 (2034, method of use); CAN-3110 (2036, composition of matter); 12 years data exclusivity
- Low-cost manufacturing

