



## Corporate Overview

March 2026

NASDAQ: LTRN

# Forward Looking Statements

This presentation contains forward-looking statements within the meaning of Section 27A of the Securities Act of 1933, as amended, and Section 21E of the Securities Exchange Act of 1934, as amended. These forward-looking statements include, among other things, statements relating to: future events or our future financial performance; the potential advantages of our RADR<sup>®</sup> platform in identifying drug candidates and patient populations that are likely to respond to a drug candidate; our strategic plans to advance the development of our drug candidates and antibody drug conjugate (ADC) development program; estimates regarding the development timing for our drug candidates and ADC development program; expectations and estimates regarding clinical trial timing and patient enrollment; our research and development efforts of our internal drug discovery programs and the utilization of our RADR<sup>®</sup> platform to streamline the drug development process; our intention to leverage artificial intelligence, machine learning and genomic data to streamline and transform the pace, risk and cost of oncology drug discovery and development and to identify patient populations that would likely respond to a drug candidate; estimates regarding patient populations, potential markets and potential market sizes; sales estimates for our drug candidates and our plans to discover and develop drug candidates and to maximize their commercial potential by advancing such drug candidates ourselves or in collaboration with others. Any statements that are not statements of historical fact (including, without limitation, statements that use words such as "anticipate," "believe," "contemplate," "could," "estimate," "expect," "intend," "seek," "may," "might," "plan," "potential," "predict," "project," "target," "model," "objective," "aim," "upcoming," "should," "will," "would," or the negative of these words or other similar expressions) should be considered forward-looking statements. There are a number of important factors that could cause our actual results to differ materially from those indicated by the forward-looking statements, such as (i) the risk that we may not be able to secure sufficient future funding when needed and as required to advance and support our existing and planned clinical trials and operations, (ii) the risk that observations in preclinical studies and early or preliminary observations in clinical studies do not ensure that later observations, studies and development will be consistent or successful, (iii) the risk that our research and the research of our collaborators may not be successful, (iv) the risk that we may not be successful in licensing potential candidates or in completing potential partnerships and collaborations, (v) the risk that none of our product candidates has received FDA marketing approval, and we may not be able to successfully initiate, conduct, or conclude clinical testing for or obtain marketing approval for our product candidates, (vi) the risk that no drug product based on our proprietary RADR<sup>®</sup> AI platform has received FDA marketing approval or otherwise been incorporated into a commercial product, and (vii) those other factors set forth in the Risk Factors section in our Annual Report on Form 10-K for the year ended December 31, 2025, filed with the Securities and Exchange Commission on March 30, 2026. You may access our Annual Report on Form 10-K for the year ended December 31, 2025 under the investor SEC filings tab of our website at [www.lanternpharma.com](http://www.lanternpharma.com) or on the SEC's website at [www.sec.gov](http://www.sec.gov). Given these risks and uncertainties, we can give no assurances that our forward-looking statements will prove to be accurate, or that any other results or events projected or contemplated by our forward-looking statements will in fact occur, and we caution investors not to place undue reliance on these statements. All forward-looking statements in this presentation represent our judgment as of the date hereof, and, except as otherwise required by law, we disclaim any obligation to update any forward-looking statements to conform the statement to actual results or changes in our expectations.

Lantern's AI platform, RADR<sup>®</sup>, is transforming the **cost, pace, and timeline** of cancer drug discovery and development

**12**

Lead drug programs\*  
powered by AI

**5**

Clinical stage lead  
drug candidates\*

**100+**

Issued patents &  
pending applications

**\$100M**

Approximate total capital  
raised since 2019

**2.5 years**

Avg. time for new  
LTRN programs  
to Ph. 1 Trial

**\$2M**

Avg. cost for new  
LTRN programs  
to Ph. 1 Trial

*\* Includes drug programs being  
developed in collaboration*

# Lantern is Transforming Drug Discovery Timelines & Costs with AI

**AI insights and biomarkers** can increase the odds of clinical trial success by **12X\***

(\*Parker et al., 2021)

**RADR®** can **predict and stratify real-world patients** for clinical trials with **88% accuracy**



Lantern can **compress the timeline** of early-stage drug development by **70%** and **reduce the cost** by **80%**

Lantern has launched **10 new programs in 2 years**, and has active ongoing Ph.1 and Ph.2 clinical trials

## LANTERN'S DRUG DEVELOPMENT MODEL AND OBJECTIVES



Large Scale/Multi-omics  
Oncology Data

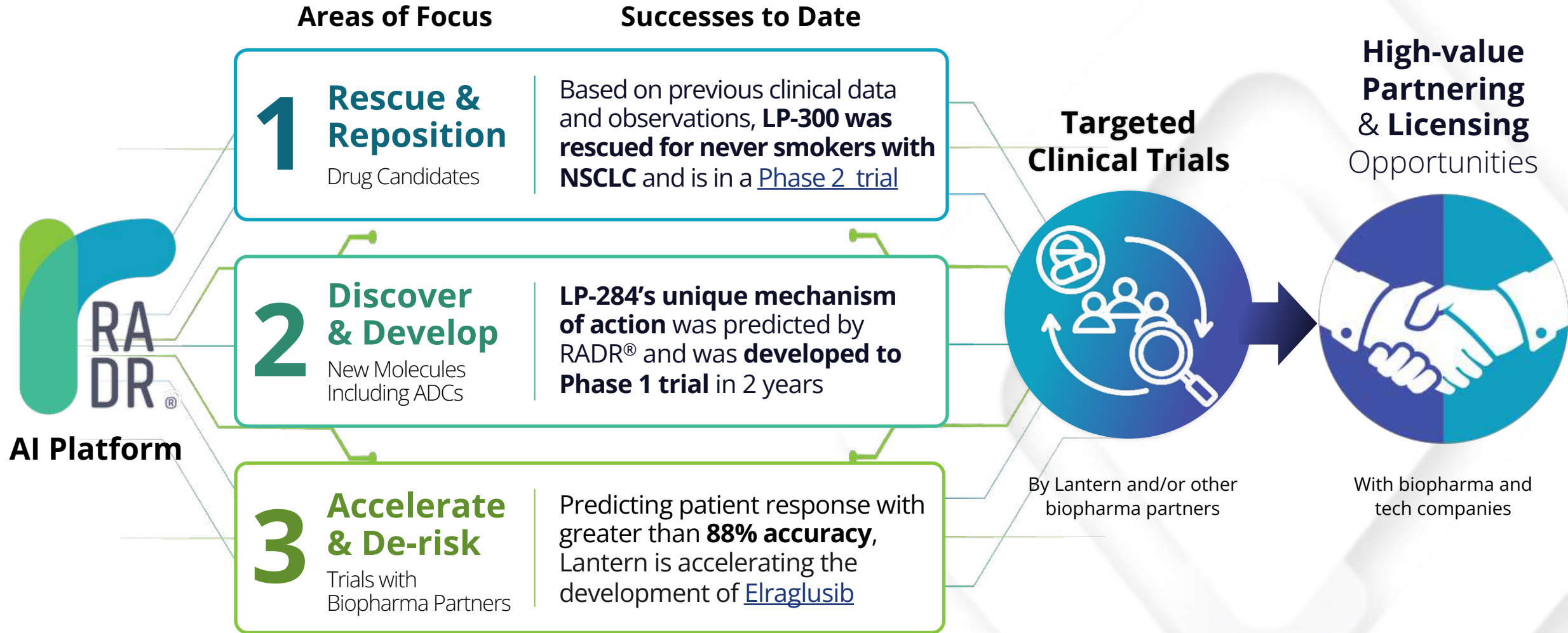


Proprietary AI  
platform RADR®



Accelerated  
timelines; reduced  
costs and risks

# Lantern's AI-Driven Business Model has Multiple Routes Towards Success



# Lantern's Diverse & Unique AI Driven Pipeline of Drug Programs

Lantern has 10 disclosed drug programs including the Phase 2 Harmonic™ trial

## Lantern Pharma (NASDAQ: LTRN)



Program	Indication	Discovery	Preclinical	Phase 1a	Phase 1b	Phase II	Orphan Designation	Rare Pediatric Disease	Fast Track	
<b>LP-300</b>	Non-Small Cell Lung Cancer for Never Smokers						harmonic			
<b>LP-184</b>	Monotherapy & Combination w/ Olaparib for TNBC									●
	Combination w/ Immune Checkpoint Inhibitors for NSCLC									
	Advanced Recurrent PTGR1-Positive Bladder Cancer						Investigator-led trial by Dr. Helle Pappot in Denmark			
<b>LP-284</b>	Recurrent Non-Hodgkin's Lymphomas ( <i>Mantle cell, Double-hit lymphomas</i> ) and Adult Soft Tissue Sarcomas							●		
<b>ADC</b>	Select Solid Tumors									





## Starlight Therapeutics (Wholly Owned Subsidiary)



<b>STAR-001</b> <i>(LP-184 for CNS and Brain Cancers Only)</i>	First Recurrent Glioblastoma in adults							●		●
	Newly Diagnosed MGMT Unmethylated Glioblastoma (investigator led trial)							●		●
	Phase 1a monotherapy including ATRT, DIPG and Medulloblastoma						FDA IND Cleared	●	●	
	Phase 1b combination select pediatric CNS cancers							●	●	

# RADR® AI-Driven Strategic Collaborations

Collaborating with top-tier oncology innovators to unlock data-driven therapeutic breakthroughs

Collaborator	Program	Indication	Stage
	Elraglusib (9-ING-41)	Multiple Solid Tumors	Phase 2 Completed
	XCE853	Protein Disulfide Isomerase (PDI) Inhibitor	Preclinical
	TTC-352	ER+ Breast Cancers	Phase 1
	ADC	Cryptophycin Conjugate for Solid Tumors	Preclinical



Precision  
Medicine  
Platform

A proprietary integrated experimental biology, oncology-focused, machine-learning-based drug development platform

**200+** Billion\*



Data points from oncology focused real-world patient and clinical data and preclinical studies

**80%+**

Prediction  
Success

**130K+**

Patient  
Records

**200+**

Advanced ML  
Algorithms

**8,163+**

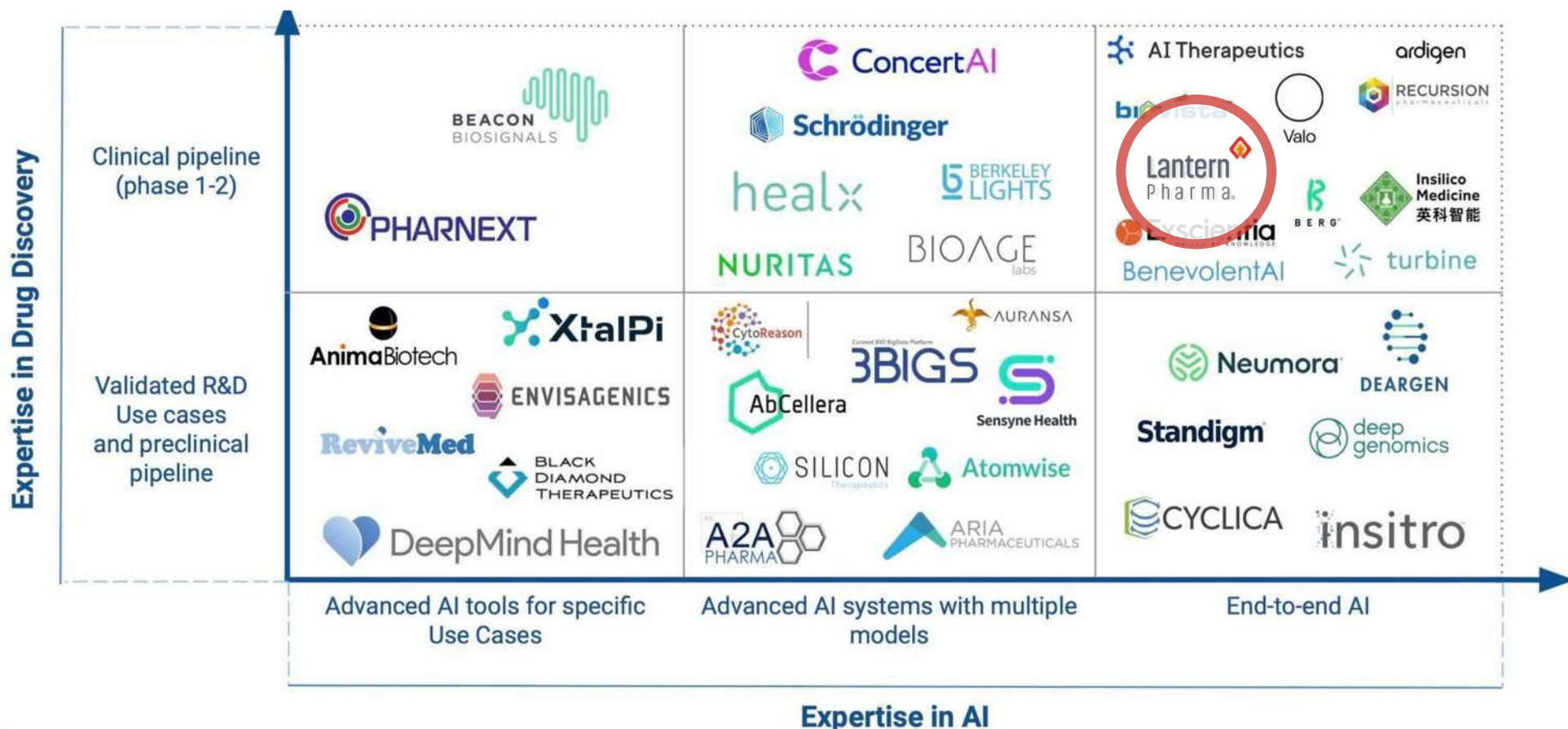
Data Sets

## AI-Powered RADR® Modules for Oncology Drug Discovery and Development

- m1** Discover mechanism of action
- m2** Identify/prioritize disease indications or subtypes
- m3** Determine optimal drug combinations
- m4** Generate ML-driven biomarker signatures
- m5** Characterize specialized attributes of a molecule
- m6** Understand potential binding site interactions
- m7** Discover combinations with checkpoint inhibitors
- m8** ADC design and optimization

# Lantern Pharma is a Top 10 End-to-End AI Drug Discovery Company

Comparison of Top-40 Leading AI for Drug Discovery Companies Expertise in Drug Discovery R&D



According to Deep Pharma Intelligence

# PredictBBB - Predict any molecule's brain permeability

Open-access AI for predicting and optimizing CNS drug's brain permeability

## The Challenge

Less than **6% of molecules** cross the Blood Brain Barrier (BBB) - one of pharmaceutical development's most persistent challenges

## The Solution

**PredictBBB™** uses AI to predict whether a molecule can cross the BBB - instantly

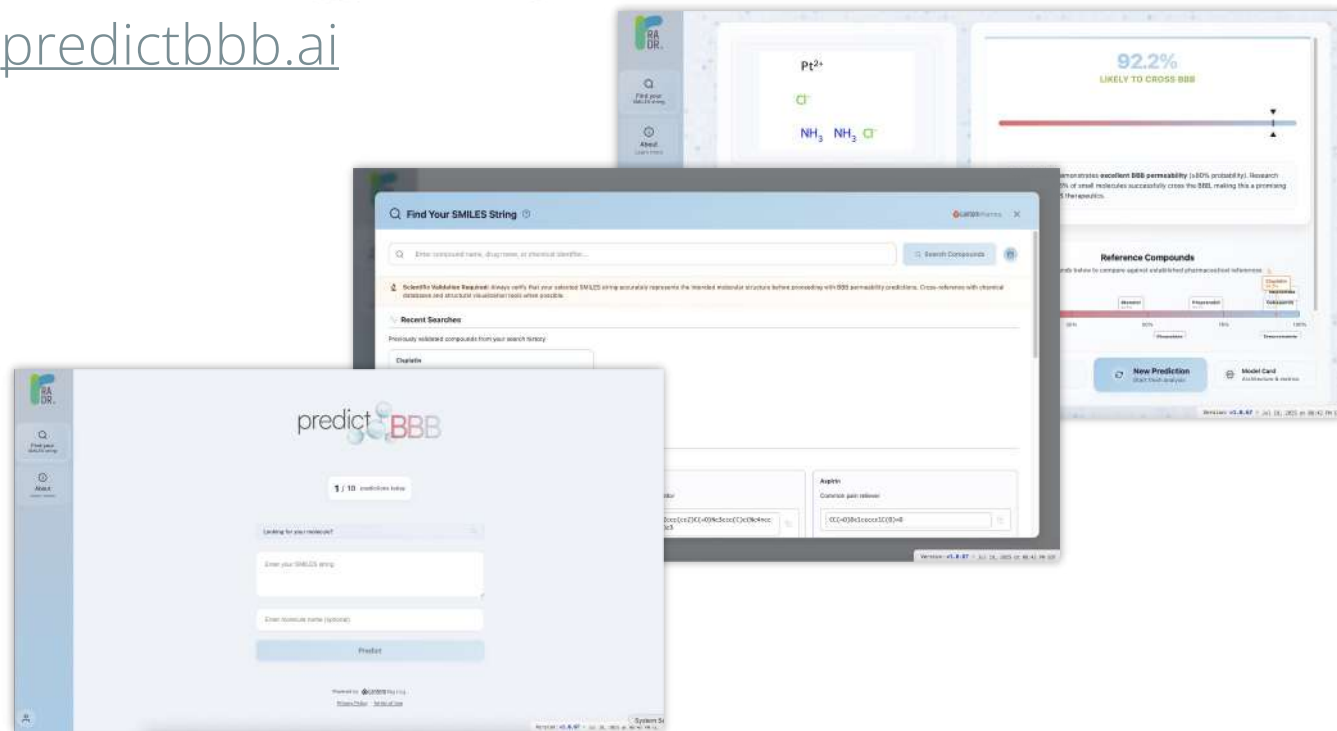
- Built on Lantern's RADR® AI platform
- Converts molecular structure into thousands of predictive features
- Evaluates permeability using ensemble machine learning models

## The Impact

- **94% prediction accuracy** on benchmark datasets
- Screens compounds **in seconds** → scalable to **100K+ molecules**
- Enables earlier, smarter selection of CNS-ready drug candidates



[predictbbb.ai](http://predictbbb.ai)



## From Prediction to Understanding

PredictBBB™ now extends beyond binary BBB prediction to provide deep molecular analysis across key dimensions of drug design

### Multi-Dimensional Molecular Intelligence



#### Overview

Integrates structure, physicochemical properties, and AI prediction into a single interpretable view- delivering probability scores, key drivers, and decision-ready insights at a glance



#### Drug-likeness Analysis

Evaluate developability using established rules (Lipinski, Veber, Ghose) with CNS-specific context



#### Structural Analysis

Assess molecular composition, 3D shape, and structural features influencing permeability and stability



#### Surface Area Profiling

Map charge, lipophilicity, polarizability, and electronic distribution across the molecule



#### Topology Analysis

Quantify molecular connectivity, branching, and complexity through advanced graph-based descriptors

# withZeta.ai – The Multi-Agentive Co-Scientist & AI System For Rare Cancer Drug Development and Research



**withZeta** addresses the fundamental challenge in rare cancer research and drug development where critical insights are scattered across disconnected data sources. Our platform integrates curated databases and external sources into an agential LLM architecture, leveraging recursive reasoning loops to transform fragmented biomedical knowledge into an interconnected investigation platform.

## Core Capabilities

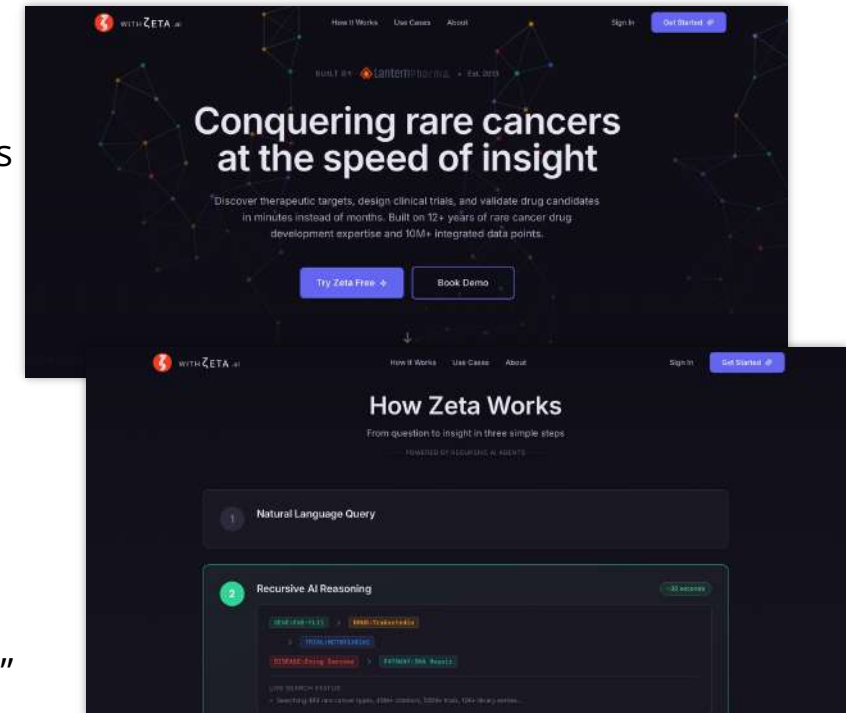
- Curated rare cancer databases and ontology
- Integrated 500k+ clinical trials, 250k+ publications, 1.2M knowledge objects
- Real-time bioinformatics and chemo informatics toolkits
- Links to RADR® predictive modules (e.g., PredictBBB.ai)

## Industry & Business Value

- Faster timelines: weeks → minutes for insights
- Smarter decisions: enhanced oncology guidance
- Novel discovery: identify new drug connections
- Improved outcomes: faster access to treatments
- Efficiency: major cost and time savings

## Strategic Impact

- Unified AI interface for complex, scattered data
- Accelerates novel therapy discovery and trial design
- Shortens drug development by months or more
- Positions Lantern as the “Perplexity for cancer research”



# Collaborations

Strategic collaborations that are providing unique real-world insights and accelerating timelines

## World-Class Academic and Research Institutions



## Biopharma Collaborations



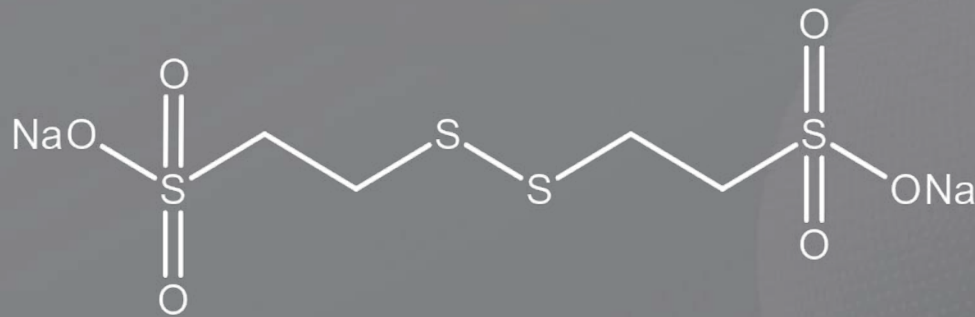
# Twelve FDA Designations Demonstrate our Data-driven, AI-enabled Approach to Transform Drug Development & Strengthen Commercial Value



12 designations

Designation	Candidate	Indication	Date
Fast Track Designation	LP-184	Glioblastoma	Sep. 2024
	LP-184	Triple Negative Breast Cancer	Dec. 2024
Orphan Drug Designation	LP-184	Pancreatic Cancer	Aug. 2021
	LP-184	Glioblastoma	Aug. 2021
	LP-184	Malignant Glioma	Aug. 2021
	LP-284	Mantle Cell Lymphoma	Jan. 2023
	LP-284	High Grade B-Cell Lymphoma	Nov. 2023
	LP-284	Soft Tissue Sarcomas	Jan. 2026
Orphan Drug and Rare Pediatric Disease Designation	LP-184	ATRT	Jan. 2022
	LP-184	Malignant Rhabdoid Tumors	Sep. 2024
	LP-184	Rhabdomyosarcoma	Sep. 2024
	LP-184	Hepatoblastoma	Sep. 2024

# LP-300 for the Treatment of Non-Small Cell Lung Cancer (NSCLC) in Never Smokers



<b>Lead Indication</b>	Relapsed NSCLC for Never Smokers
<b>Clinical Status</b>	Phase 2 (multiple patients dosed globally, Japan enrollment complete)
<b>Market Potential*</b>	\$4+ billion
<b>Indication Size*</b>	150,000 + Cases
<b>Target/ MOA</b>	Tyrosine Kinases & Cell Redox Enzymes
<b>Molecule Type</b>	Disulfide Small Molecule
<b>Combination</b>	With Carboplatin and Pemetrexed
<b>IP Estate</b>	Claims extending to at least 2032

*\*Estimated Annual Global*

# Disease Overview – NSCLC in Never Smokers – LP-300

NSCLC in never smokers is one of the largest unaddressed cancer populations

**Global Annual Market Potential: \$ 4+ Billion**

Lung cancer is the **#1** cause of death among cancer patients in the US

**1** in 6

lung cancer deaths will occur in patients that are never smokers with NSCLC

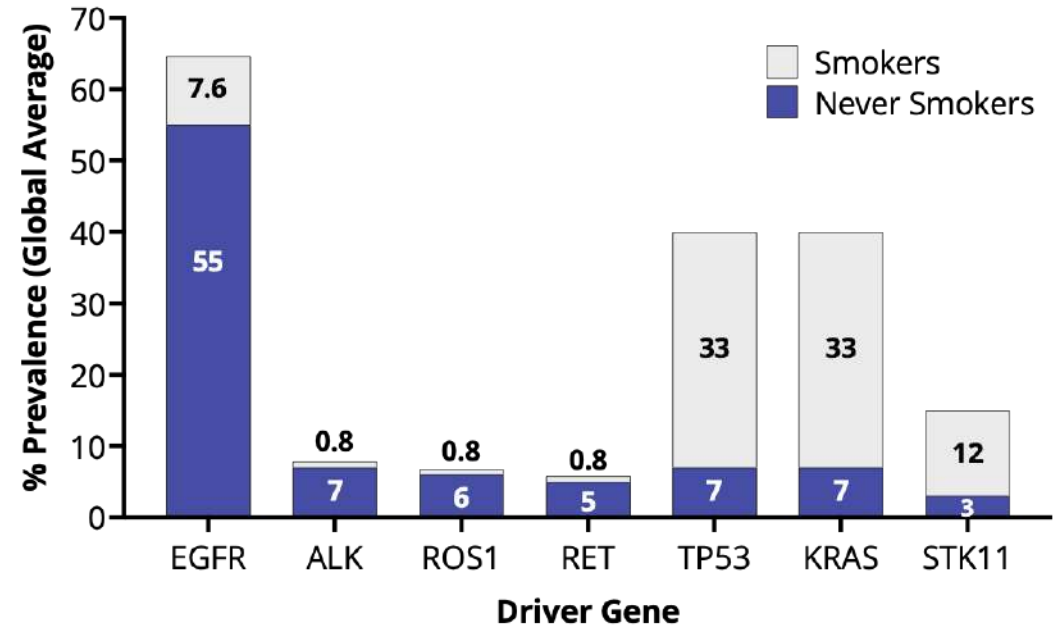
**150,000~175,000**

never smokers will be diagnosed with **NSCLC** Globally  
*Cancer.gov*

## NSCLC in Never Smokers is a Different Disease

Lung Cancer in never smokers has **higher percentage of genetic mutations in Tyrosine Kinases (TK)**, a family of cancer-promoting genes, such as EGFR, ALK, ROS and MET

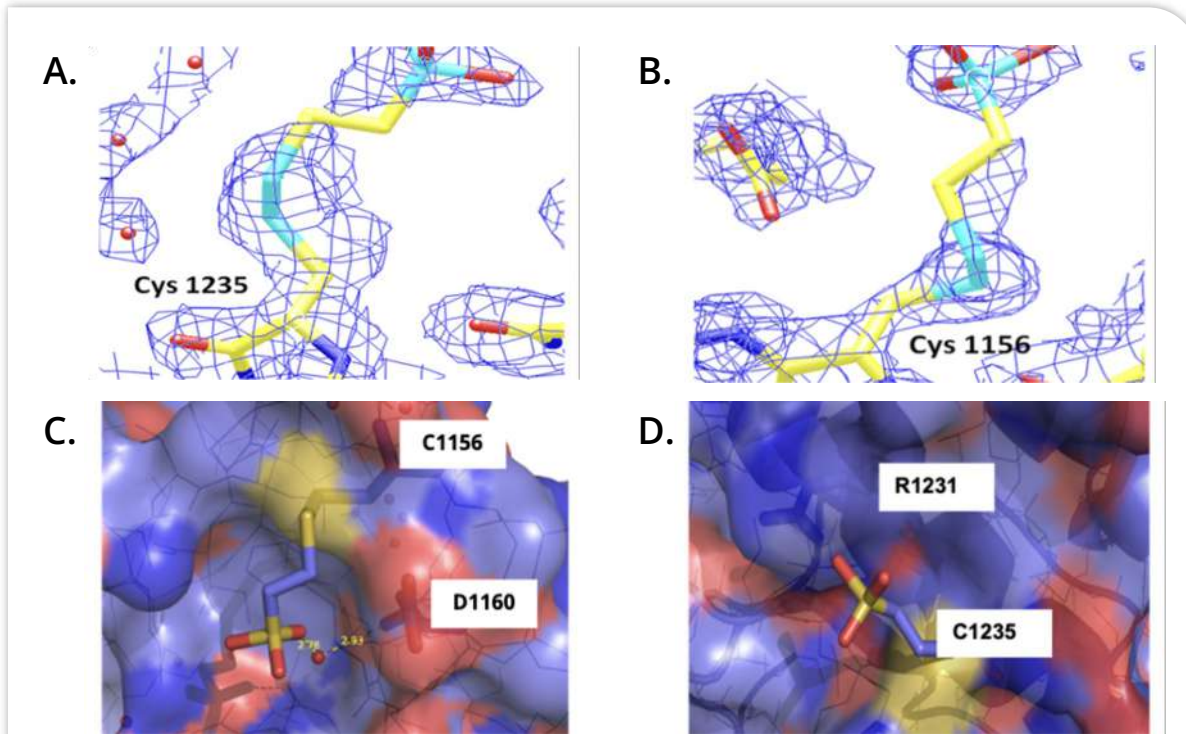
**Mutation Frequency by Smoker Status**



# Mechanism of Action – LP-300

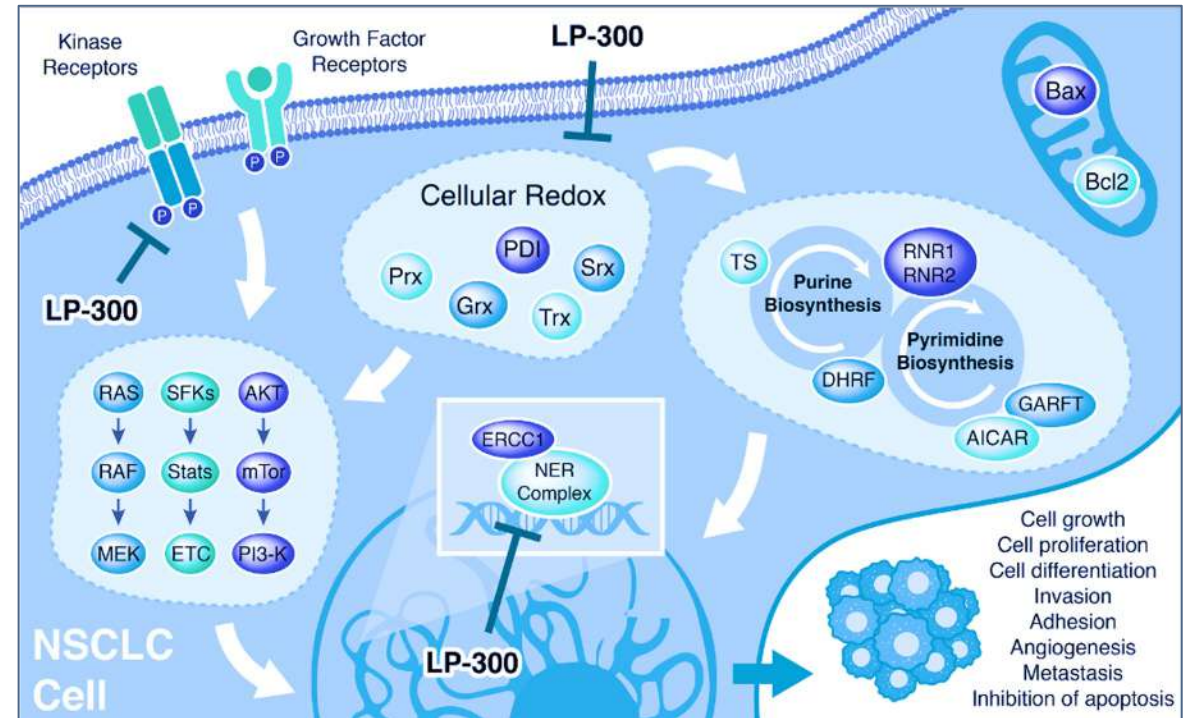
LP-300's multimodal MoA resensitizes NSCLC to chemo in the never smoker population

## 1. LP-300 Directly Engages with TKI Receptors via Cysteine Modification



**A-B.** LP-300 adduct at **Cys1235** **Cys1156** **C.** Molecular surface of ALK with the LP-300-derived adduct at **Cys1156** (*yellow highlight*) **D.** Binding site of the LP-300-derived adduct at **Cys 1235** (*yellow highlight*)

## 2. LP-300 Modulates Cellular Redox in Key Signaling Pathways in NSCLC



- Restoring apoptosis sensitivity
- Oxidative stress modulation
- Anti-angiogenesis
- Reduced DNA synthesis and gene expression
- Reduce glutathione/thioredoxin mediated tumor resistance to therapy
- Nephrotoxicity protection against chemotherapy

# Clinical Trial – The Harmonic™ Phase 2 Trial for LP-300

Accelerating recruitment efforts for a growing indication with limited treatment options



[NCT05456256](https://clinicaltrials.gov/ct2/show/study/NCT05456256)



Non-Small Cell Lung Cancer



Never Smokers

90

Patients



Two arm, Open-label, Randomized Trial



Multi-Site in US & Asia

## Trial Highlights

- Completed Japanese patient cohort enrollment ahead of schedule at multiple clinical sites including the National Cancer Center in Tokyo
- Patient showed durable complete response with survival continuing for nearly **two years**
- Preliminary patient data and clinical readouts showed an **86% clinical benefit rate**

**Primary Outcomes:** Overall and progression free survival

Announced preliminary patient data showing an 86% clinical benefit rate - Scan the QR code for the full initial result release



**Multi-national Phase 2 Trial with 8 sites in the US, 5 sites in Japan, and 5 sites in Taiwan**



## KEY PATIENT CHARACTERISTICS

- ✓ Patients who are never smokers with lung cancer and histopathological evidence of stage III or IV primary lung adenocarcinoma
- ✓ Molecular alterations, including EGFR, MET exon 14 skipping, ROS1, BRAF, ALK, and NTRK fusions
- ✓ Relapsed after one or more lines of therapy with tyrosine kinase inhibitors

## STUDY ENDPOINTS

- ✓ Primary: Progression-free survival (PFS) and overall survival (OS)
- ✓ Secondary: Objective response rate (ORR), duration of response (DOR), and clinical benefit rate (CBR)

Tumor Response	LP-300+ Carboplatin + Pemetrexed
Partial Response	3/7 (43%)
Stable Disease	3/7 (43%)
Progressive Disease (clinical)	1/7 (14%)
<b>Clinical Benefit Rate (CBR)</b>	<b>6/7 (86%)</b>
<b>Objective Response Rate (ORR)</b>	<b>3/7 (43%)</b>

*All patient data as of July 25, 2024*

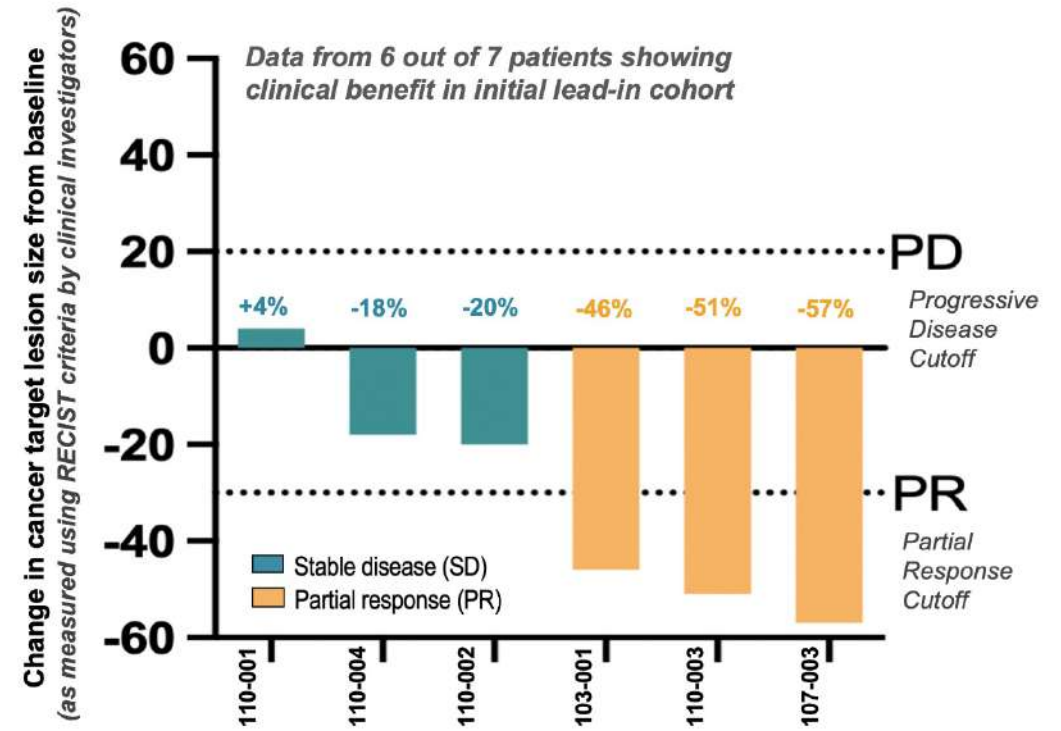
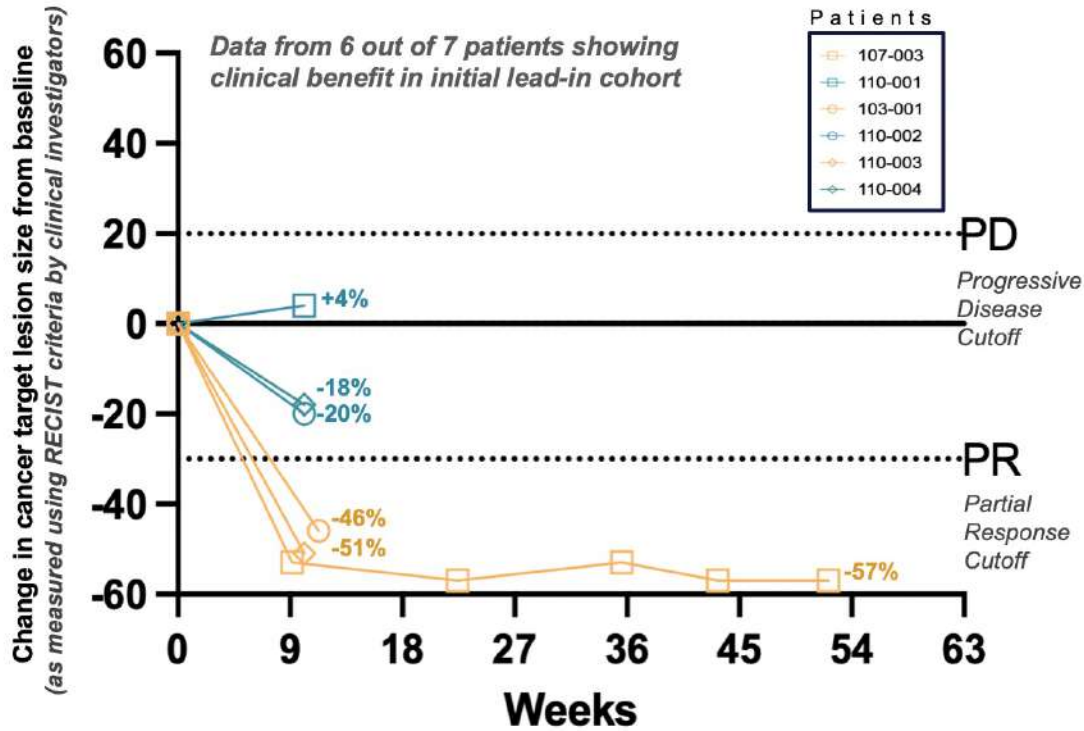
## Patient Highlights from Initial Cohort

- 7 patients enrolled from different geographies
- Sites included were in CA, VA, TX
- 3 Female and 4 Male
- Average age of 62
- Median prior lines of therapy: 2 (1 to 4)
- Recent historical trials in similar patient groups receiving the chemo doublet have had an ORR of 26% to 36% with a PFS of 5.1 months

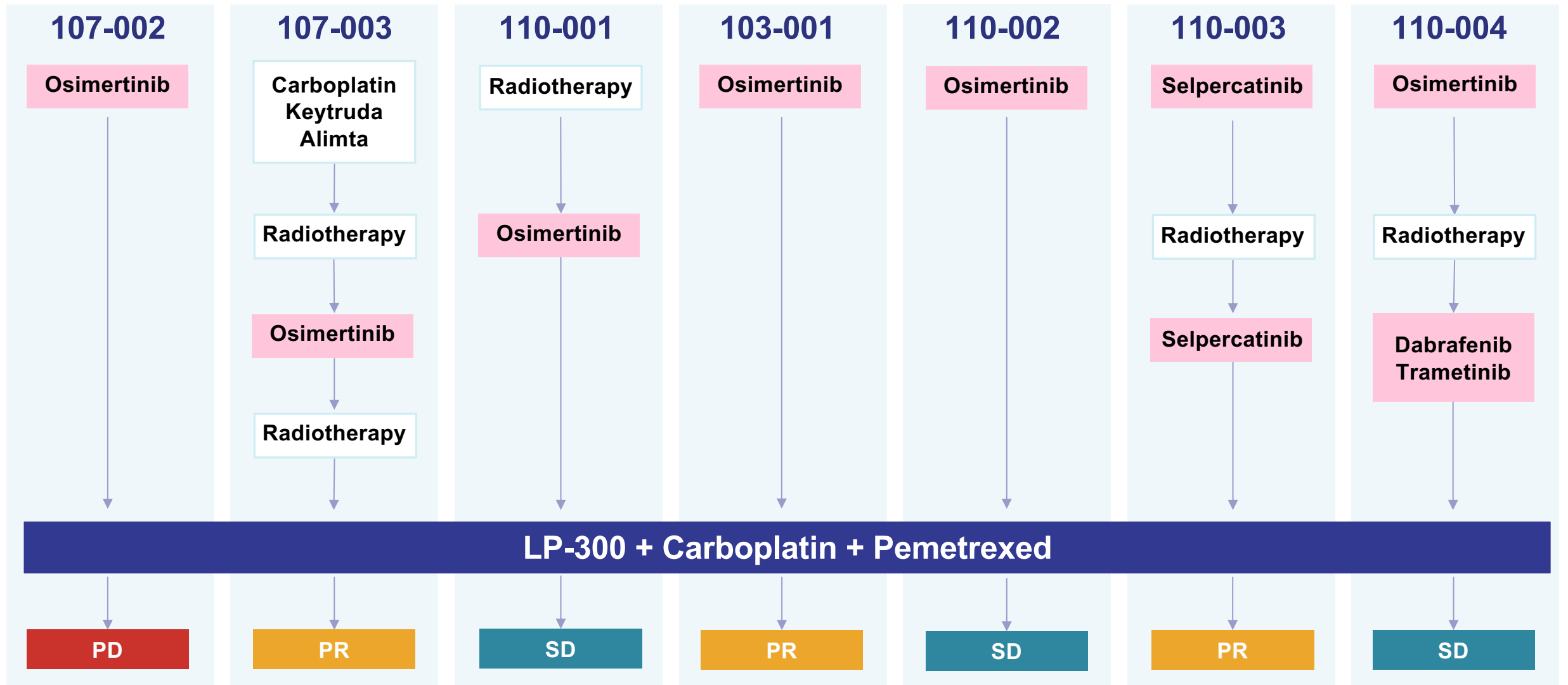
Initial patient responses in the Harmonic™ trial include an **86% disease control rate** in the cohort of lead-in patients and a **43% objective response rate (ORR)** including one patient maintaining a **50+% reduction in tumor size** over 14 months

Percent change in cancer lesion size over time

Percent change in cancer lesion size by patient



All patient data as of July 25, 2024



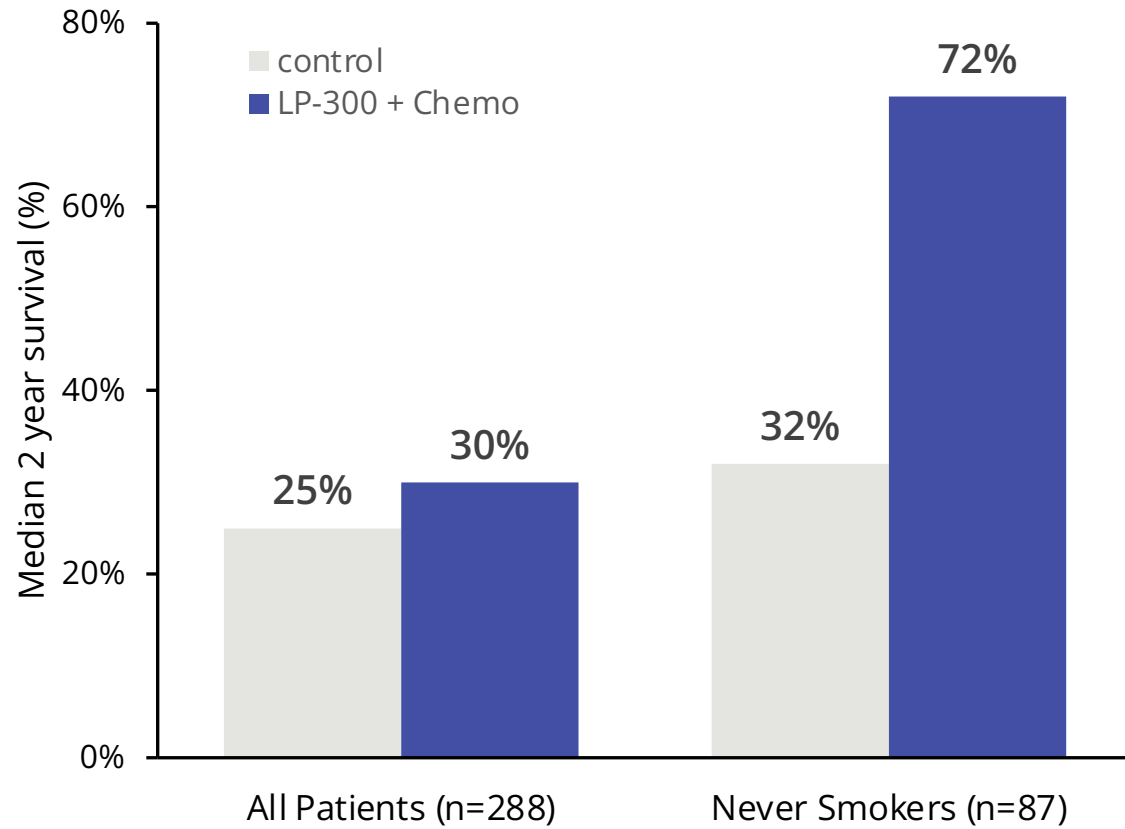
All patient data as of July 25, 2024

TKI PR: Partial Response, SD: Stable Disease, PD: Progressive Disease

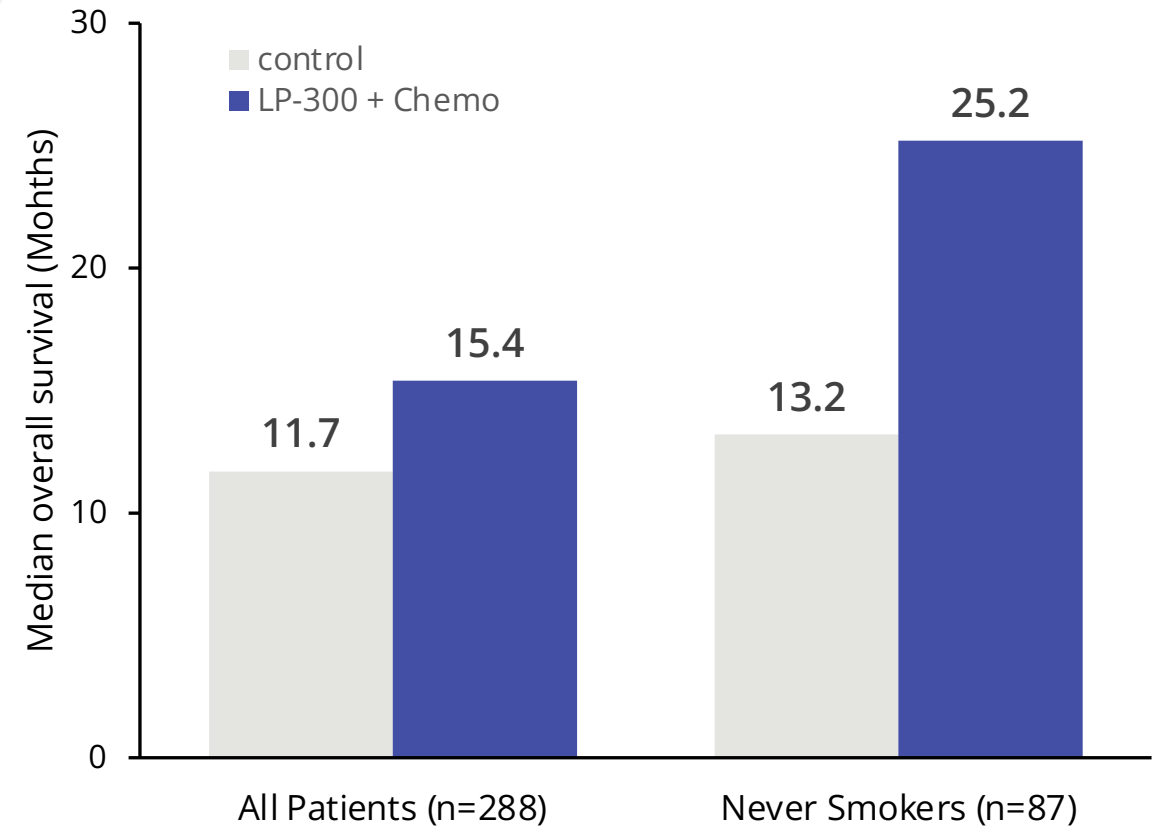
# LP-300 Nearly Doubled Survival Outcomes for Never Smoker Subgroups with NSCLC in Previous Clinical Trial\*

\*Subpopulations receiving paclitaxel/cisplatin

**+ 125% increase** in median 2 year survival



**+ 91% increase** in median overall survival



\*Overall study did not meet clinical efficacy endpoints

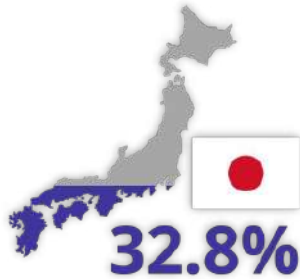
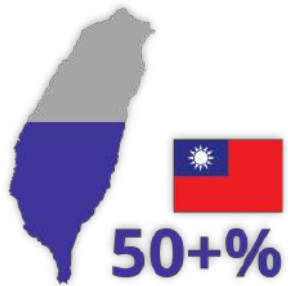
Clinicaltrials.gov ([NCT00966914](https://clinicaltrials.gov/ct2/show/study/NCT00966914))

# Initiated East Asia: Boosting Patient Enrollment in Countries with High Incidences of NSCLC in Never Smokers

**2 in 5**

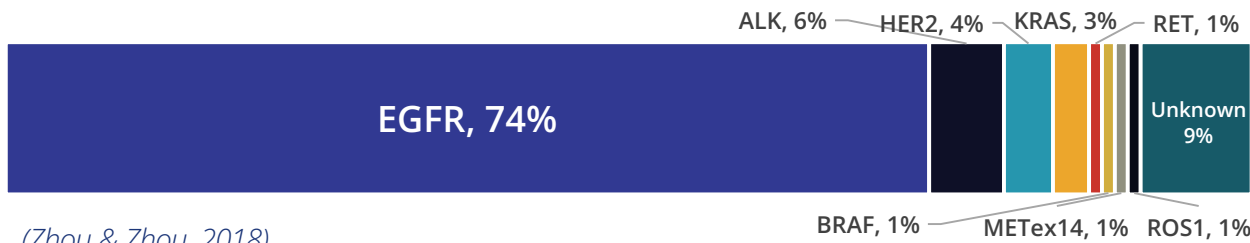
of all lung cancer patients in East Asia are **never smokers\***

*\*Approximately*



% of **never smokers** among lung cancer patients in Taiwan and Japan

Lung cancer in East Asian never-smokers is a **distinct subtype** that can be largely defined by targetable mutations



*(Zhou & Zhou, 2018)*

## Highlights

- Study expansion to Taiwan and Japan with 5 sites in each country
- Enrollment completed in Japan

## Key Opinion Leaders



**Dr. Yasushi Goto**  
National Cancer Center Hospital



**Dr. Chun-Hui Lee**  
National Cheng Kung University Hospital

**Q2-Q3 2024**

Regulatory and Site Submissions

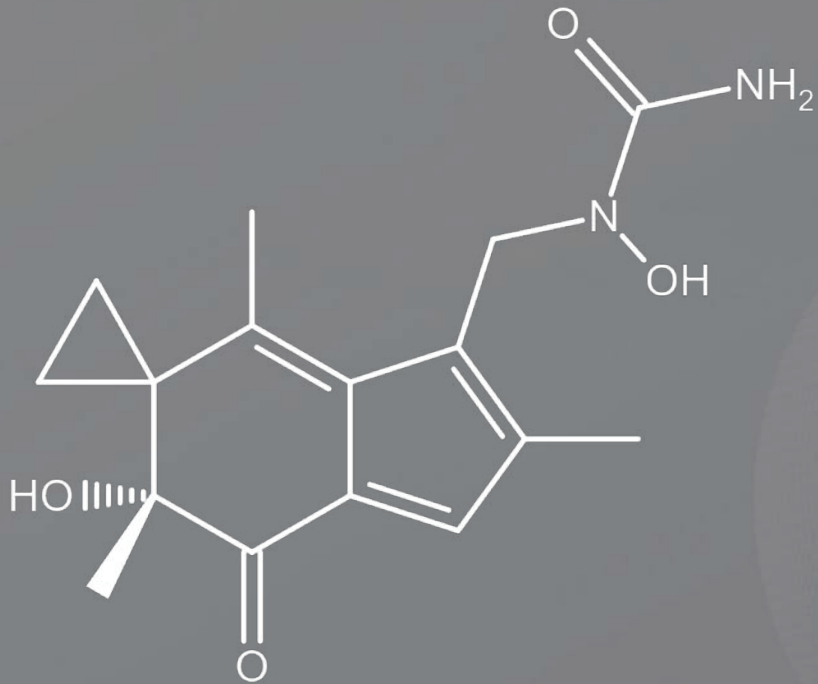
**Q4 2024**

Site Activation and First Patients Dosed

**Q4 2025**

Review of initial patient response in Asia and updates from first US cohort

# LP-184 for the Treatment of Advanced Solid Tumors



<b>Lead Indications</b>	DDR deficient solid tumors including Pancreatic cancer, Bladder cancer, and TNBC
<b>Clinical Status</b>	Phase 1a enrollment completed, Phase 1b/2 planned
<b>Market Potential*</b>	\$10+ Billion
<b>Indication Size*</b>	170,000 + Cases, Estimated 400,000 + Cases Global
<b>Target/ MOA</b>	Double-stranded DNA breaks; alkylates DNA in the 3' of Adenine
<b>Molecule Type</b>	Acylfulvene Class
<b>Combination Potential</b>	Checkpoint inhibitors, PARP inhibitors, Spironolactone, Chemotherapy and Radiation Therapy
<b>IP Estate</b>	10+ patents/pending apps., Claims extending into 2041

\*Estimated Annual USA

# Disease Overview – Advanced Solid Tumors with DDR Deficiencies

LP-184 has Blockbuster Potential Across Multiple Cancers as a Single Agent or in Combination Therapy

## Annual US Market Potential: \$10+ Billion

(DDR Deficient Solid Tumors)

 **1 in 4** people have solid tumors with DDR Deficiencies



Pancreatic Cancer



Triple Negative Breast Cancer



Bladder Cancer



Lung Cancer

## Advanced Solid Tumors

- Advanced solid tumor cancers, having spread beyond the primary site, are often more challenging to treat than earlier stage tumors due to their advanced progression
- Demonstrated preclinical synergy with multiple FDA approved drugs (e.g. PARPi, PD-1, and Spironolactone)
- Many of these indications - *reinforced with AI insights* - have limited or no standard of care, making them ideal and efficient entry points for LP-184 as an approved therapy

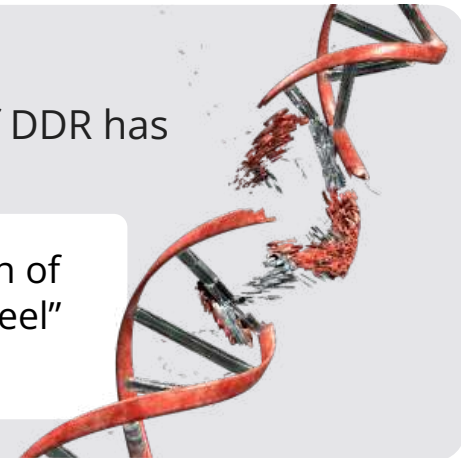
## DNA Damage Response (DDR) Deficiency

DDR is essential for maintaining genomic stability by repairing different types of DNA damage. Inhibition of DDR has been shown to increase the effectiveness of anticancer immunotherapies

**Cancer cells** with high underlying levels of DNA damage are **more dependent on DDR** for survival when compared to normal cells



**DDR Deficiencies** result in the accumulation of DNA damage, which produces an “Achilles Heel” for drugs leveraging synthetic lethality

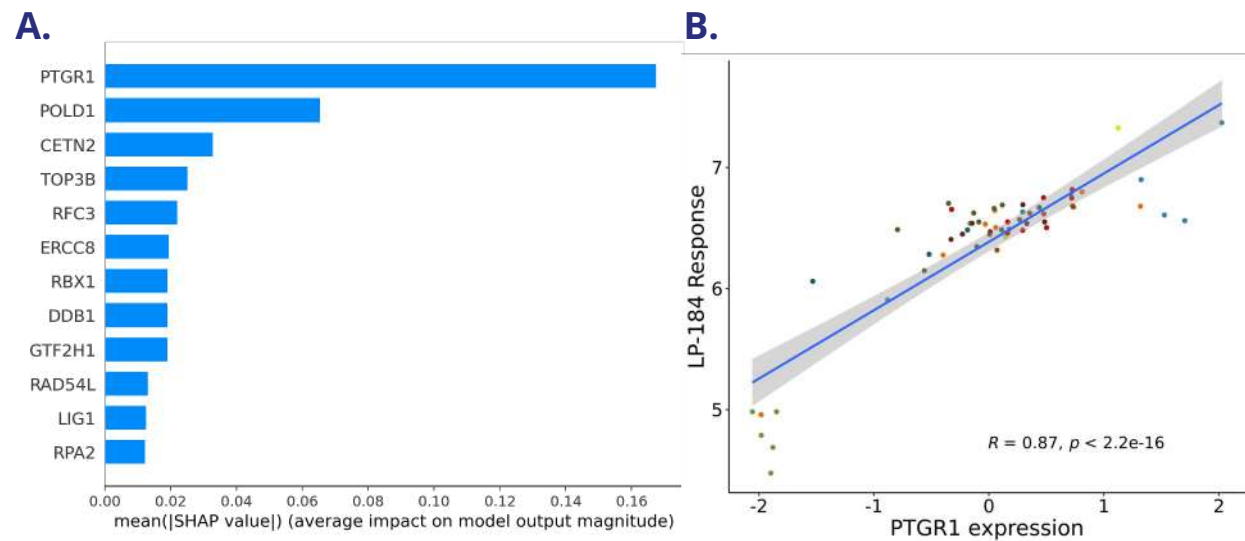


# LP-184's MoA was Predicted by RADR<sup>®</sup> and Validated in Multiple Lab Studies

*In silico*



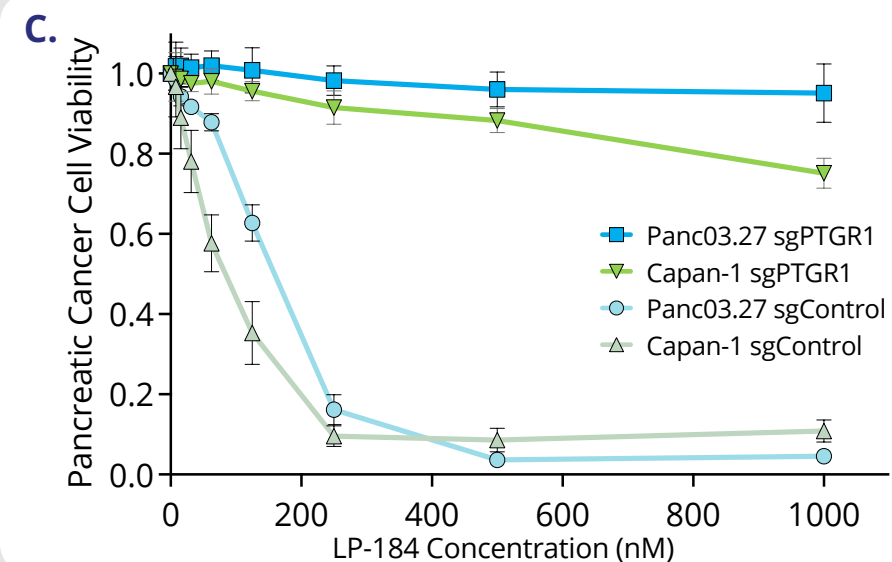
Using RADR<sup>®</sup>, PTGR1 was Identified as a Biomarker that Predicts LP-184 Response



*In vitro*



Validated using CRISPR Experiments



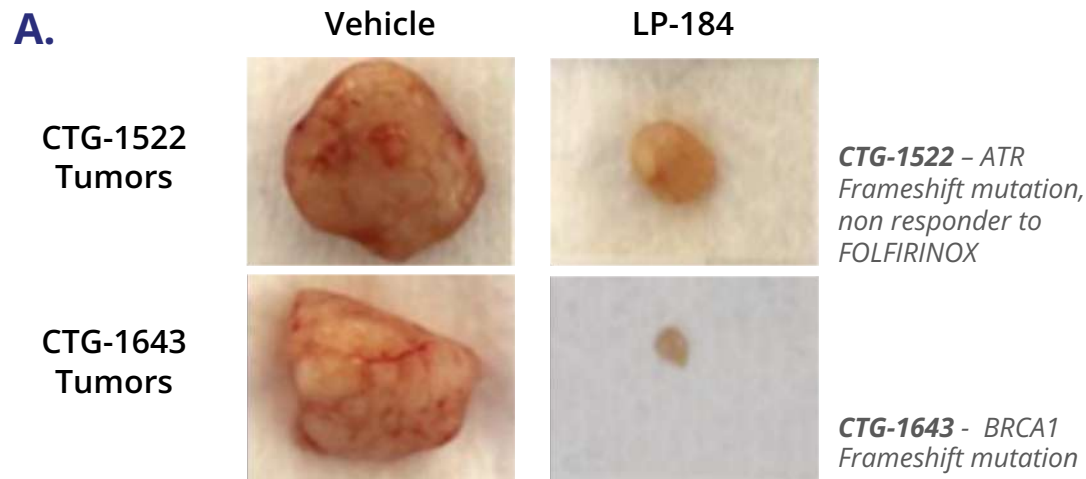
- **Prostaglandin Reductase 1 (PTGR1)** is an oxidoreductase enzyme that is frequently elevated in cancers
- PTGR1 activates LP-184 into its highly potent and cytotoxic form
- RADR<sup>®</sup> insights predicted that LP-184 activity positively correlates with PTGR1 transcript levels in the NCI60 cancer cell line panel

- CRISPR-mediated depletion of PTGR1 expression in a pancreatic cancer cell line is sufficient to **fully diminish LP-184 activity**
- This **confirmed the RADR<sup>®</sup> insights** and that LP-184 was highly potent in cells with PTGR1

# LP-184 Treatment Results in Complete Regression in Multiple DDR Deficient PDX Models

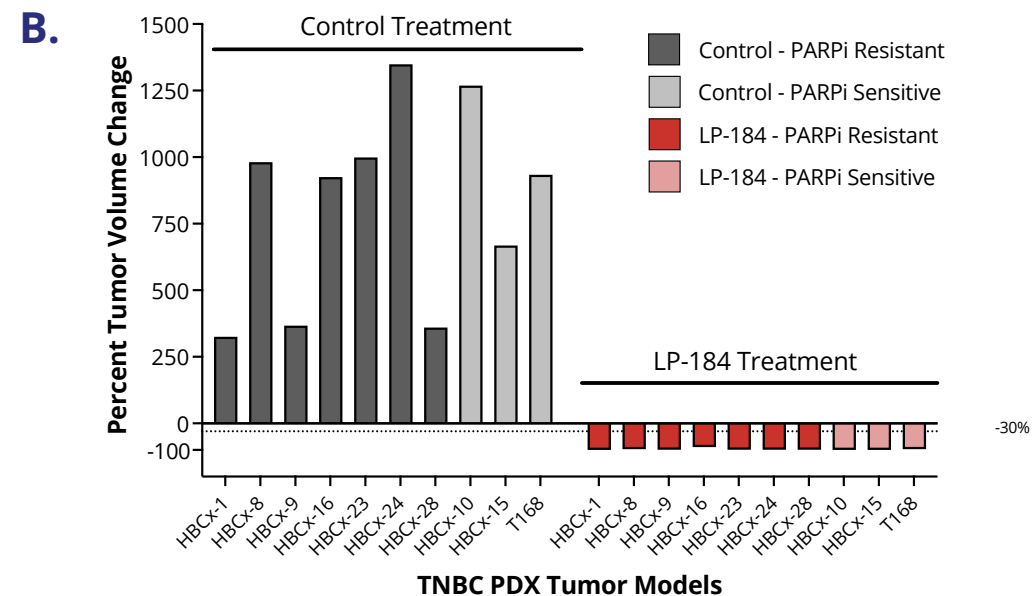
## Pancreatic Cancer

In-vitro PDX pancreatic mouse models treated with LP-184 - CTG-1522 and CTG-1643 models showed a **tumor growth inhibition of >100%**



## Triple Negative Breast Cancer (TNBC)

Across 10 TNBC PDX mouse models (*All 10 TNBC PDX models were HR deficient*) LP-184 treatment resulted in 107-141% tumor growth inhibition



In collab. with 

Poster: 

- LP-184 exhibits nanomolar potency in PTGR1 overexpressing tumors with DDR deficiencies
- Positioned for 2nd and 3rd line treatment, where there is unmet need for novel therapies
- FDA **Orphan Drug Designation** granted in pancreatic and **Fast Track Designation** in TNBC
- Combination therapy potential with SOC agents: Spironolactone, PARP inhibitors, Gemcitabine, Irinotecan, Oxaliplatin, and PD-1

# Clinical Trial – Completed LP-184 Phase 1a Basket Trial

Potential blockbuster molecule with a market of \$10+ billion in annual sales

## First-In-Human Trial for LP-184

[Clinicaltrials.gov \(NCT05933265\)](https://clinicaltrials.gov/ct2/show/study/NCT05933265)



Solid Tumors /  
Brain & CNS Cancers

~60

Patients expected  
to be enrolled

\$10+ Bn

Annual US market potential in  
DDR deficient solid tumors

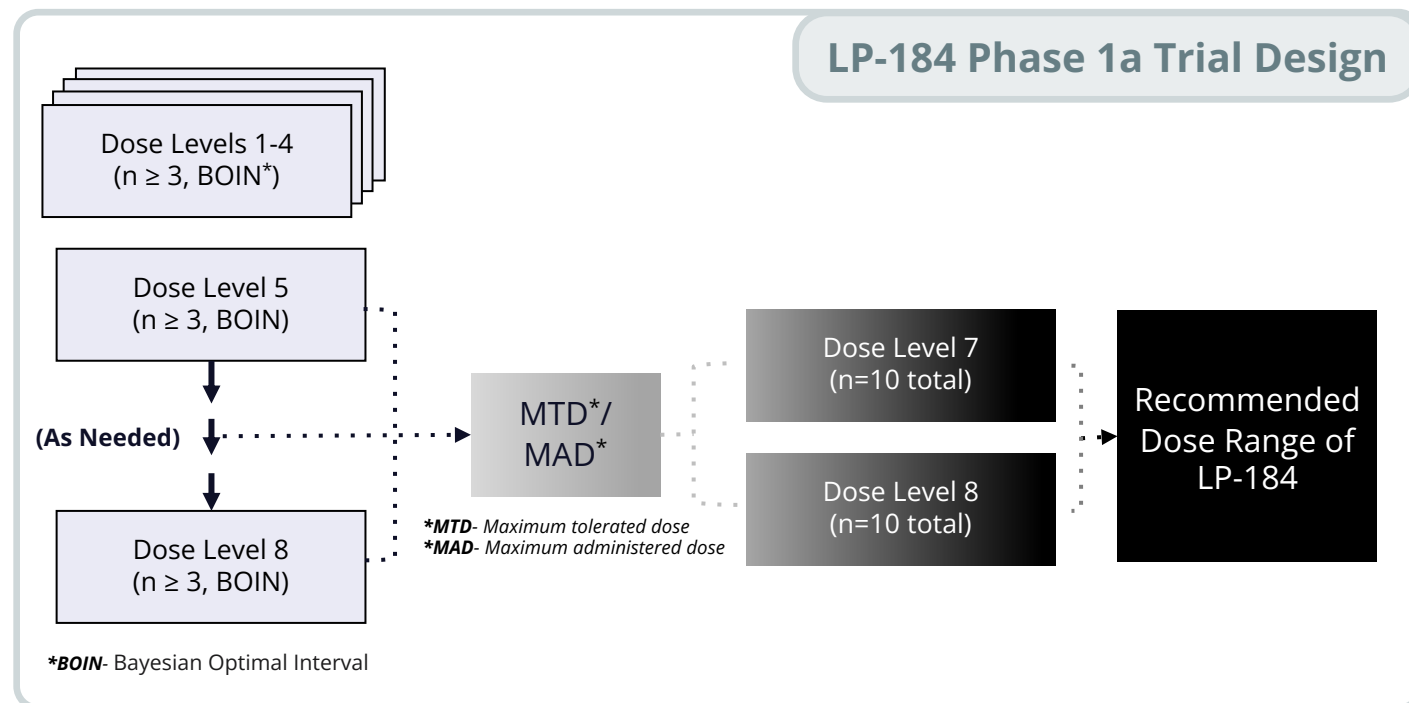


Multi-Site

## Phase 1 Trial Highlights

- Successfully **completed** with all primary endpoints met, demonstrating a favorable safety and pharmacokinetic (PK) profile, and early signs of antitumor activity.
- **Potential future studies: Phase 2 in GBM (through Starlight) and Phase 1b/2 in other solid tumors** to be initiated after determination of MTD
- Enrollment is complete, with several patients continuing treatment due to ongoing clinical benefit.

## LP-184 Phase 1a Trial Design

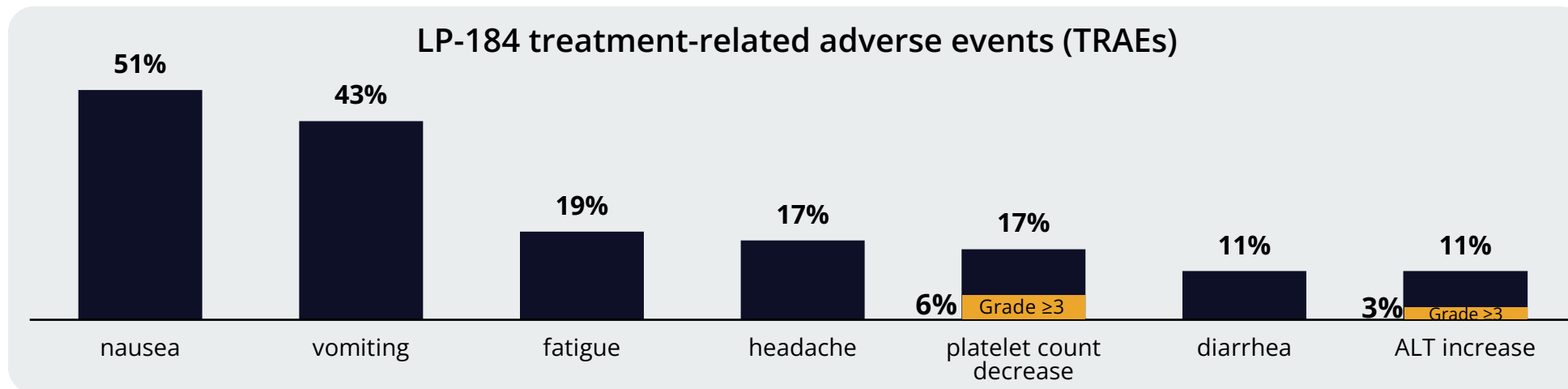


# LP-184 Phase 1a Trial Achieved All Primary Endpoints with Robust Safety Profile and Promising Antitumor Activity in Multiple Advanced Solid Tumors

LP-184 exhibited a robust safety profile, with no dose-limiting toxicities in the majority of cohorts

# 89%





of treatment emergent adverse events (AEs) were Grade 1-2



## TRIAL RESULT HIGHLIGHTS

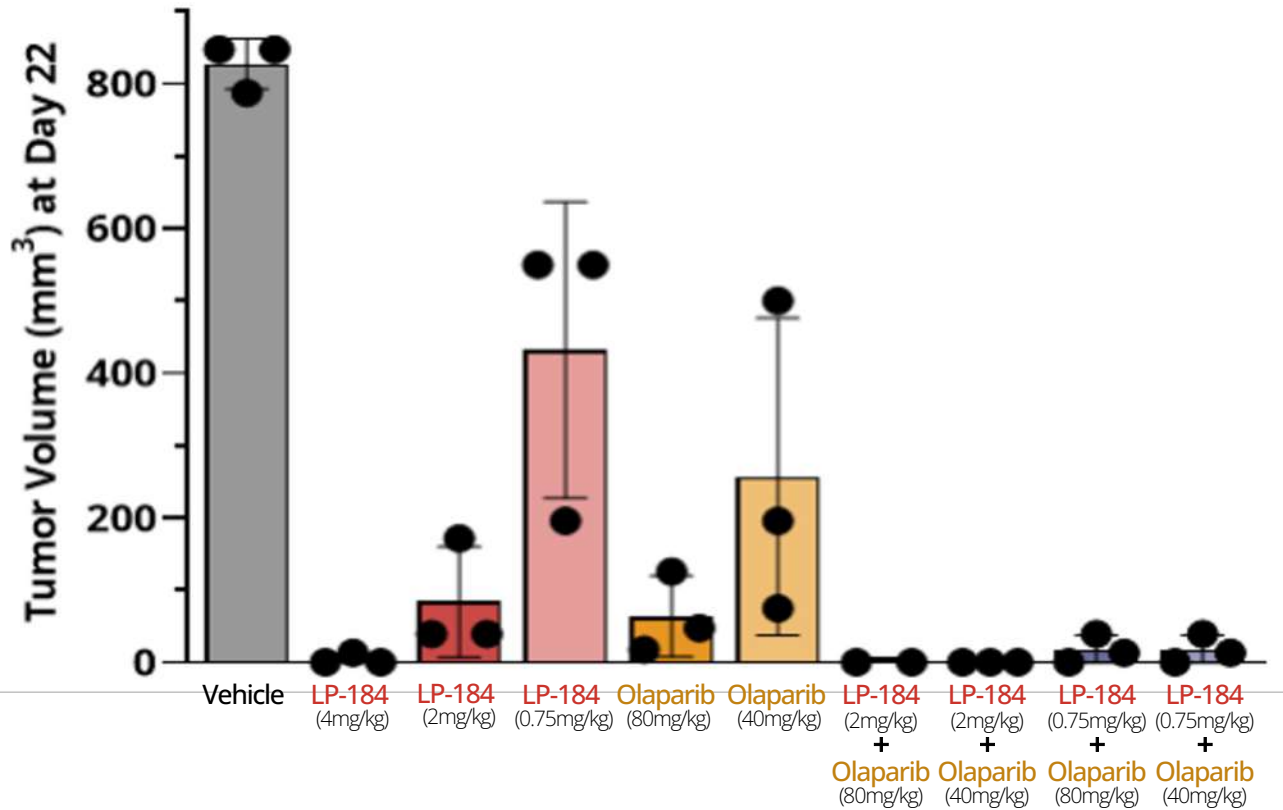
- **Clinical benefit observed in 48%** of evaluable cancer patients at or above the therapeutic dose threshold
- **Durable clinical benefits** were observed in **hard-to-treat tumors** like glioblastoma multiforme (GBM), gastrointestinal stromal tumor (GIST) and thymic carcinoma
- PK data confirmed that therapeutic concentrations were achieved at **dose level 8 (0.25 mg/kg)** and above
- Biomarker insights highlight potential in **DDR-mutated cancers**, with marked tumor reductions in patients with CHK2, ATM, and STK11/KEAP1 alterations
- **Recommended Phase 2 dose (RP2D)** established for targeted Phase 1b/2 trials in triple-negative breast cancer (TNBC), non-small cell lung cancer (NSCLC), and bladder cancer

# Planned Clinical Trials – LP-184 Phase 1b/2 Trials informed by RADR® AI Insights

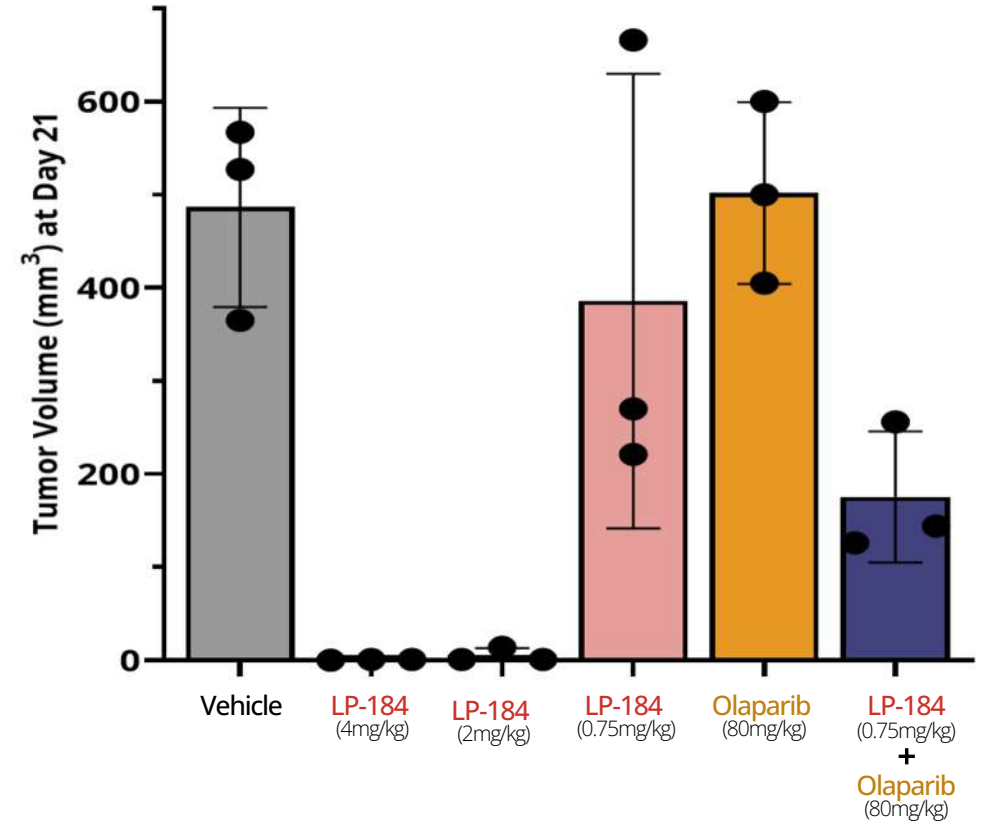
Trial	Indication	Market potential	Trial Size	Trial Highlights
Phase 1b/2 Monotherapy & Combination with <b>Olaparib</b> for <b>TNBC</b>	 Triple Negative Breast Cancer	<b>\$4+Bn</b> Annual US market potential	<b>~60</b> Patients expected to be enrolled	<ul style="list-style-type: none"> <li>Granted FDA Fast Track Designation for monotherapy of LP-184</li> <li><b>Monotherapy Trial:</b> Evaluating optimal dose and early efficacy of LP-184 in advanced TNBC with DNA repair gene mutations.</li> <li><b>Combination Trial:</b> Assessing safety and efficacy of LP-184 + Olaparib in advanced TNBC with BRCA mutations.</li> </ul>
Phase 1b/2 Combination with <b>Immune Checkpoint Inhibitors</b> for <b>NSCLC</b>	 KEAP1 and/or STK11 mutated NSCLC	<b>\$2+Bn</b> Annual US market potential	<b>~34</b> Patients expected to be enrolled	<ul style="list-style-type: none"> <li>Submission for FDA Fast Track Designation in process</li> <li>Open-label study evaluating safety and early efficacy of LP-184 with nivolumab and ipilimumab in advanced NSCLC with KEAP1/STK11 mutations and low PD-L1.</li> </ul>
Phase 1b/2 Investigator Led Trial in Denmark for <b>Bladder Cancer</b>	 Bladder cancer with TC-NER deficiency	<b>\$0.5+Bn</b> Annual Global market potential	<b>~39</b> Patients expected to be enrolled	<ul style="list-style-type: none"> <li>Investigator-sponsored trial (Dr. Helle Pappot, Rigshospitalet University, Denmark)</li> <li>Open-label study evaluating safety and early efficacy of LP-184 in advanced/metastatic urothelial carcinoma with PTGR1 positive and TC-NER/HR deficiency</li> </ul>
Phase 1b/2a Combination with <b>Spirolactone</b> for <b>Glioblastoma</b>	 Recurrent Glioblastoma	<b>\$1+Bn</b> Annual US market potential	<b>~38</b> Patients expected to be enrolled	<ul style="list-style-type: none"> <li>Granted FDA Fast Track Designation and Orphan Drug Designation for monotherapy of LP-184</li> <li>First recurrent Glioblastoma</li> <li>Simon 2-stage design</li> <li>2 arms; IDHm and IDHwt</li> </ul>

# LP-184 + Olaparib Combination Achieves 3-14x Greater Tumor Regression Compared To Olaparib Alone In TNBC PDX Models

Tumor regression is achieved using 5x lower doses of LP-184 in combination as compared to doses used as monotherapy



Tumor Volume in HBCx-10 PARPi sensitive TNBC PDX Model Treated with LP-184 (days 1, 8), Olaparib (daily), or Combination



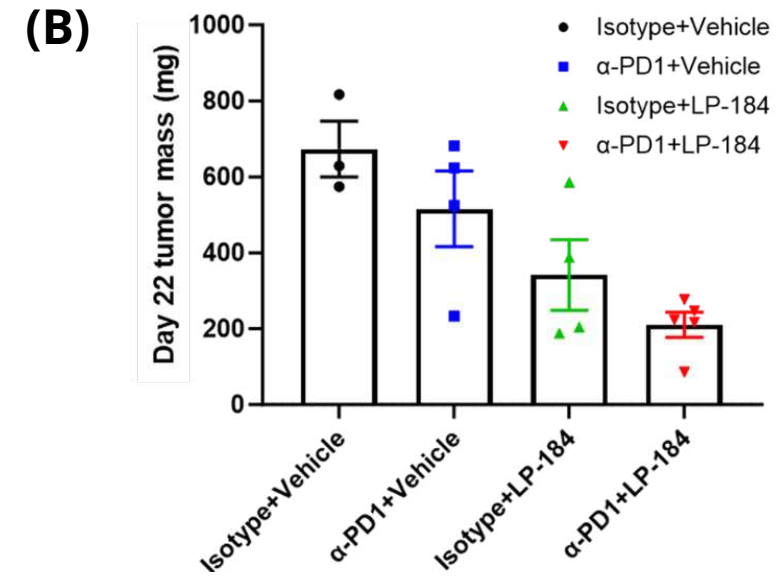
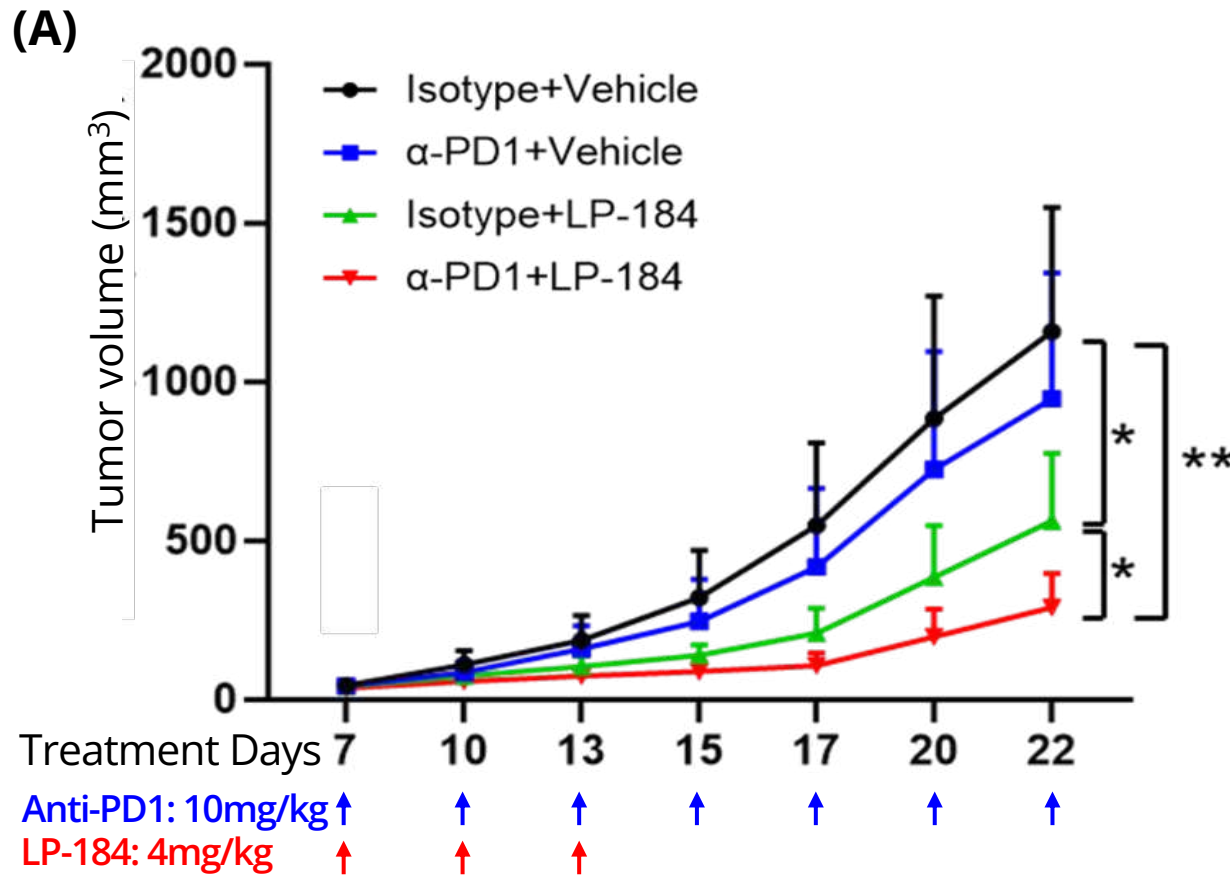
Tumor Volume in HBCx-28 PARPi resistant TNBC PDX Model Treated with LP-184 (days 1, 4, 8, 11), Olaparib (daily), or Combination

Kulkarni, A. et al., Cancer Research Communications, 2024

# LP-184 + Anti-PD1 Combination Significantly Inhibits Tumor Growth And Delays Progression In T11 Mouse TNBC Model

## T11 mouse TNBC tumors treated with LP-184 and anti-PD1 antibody

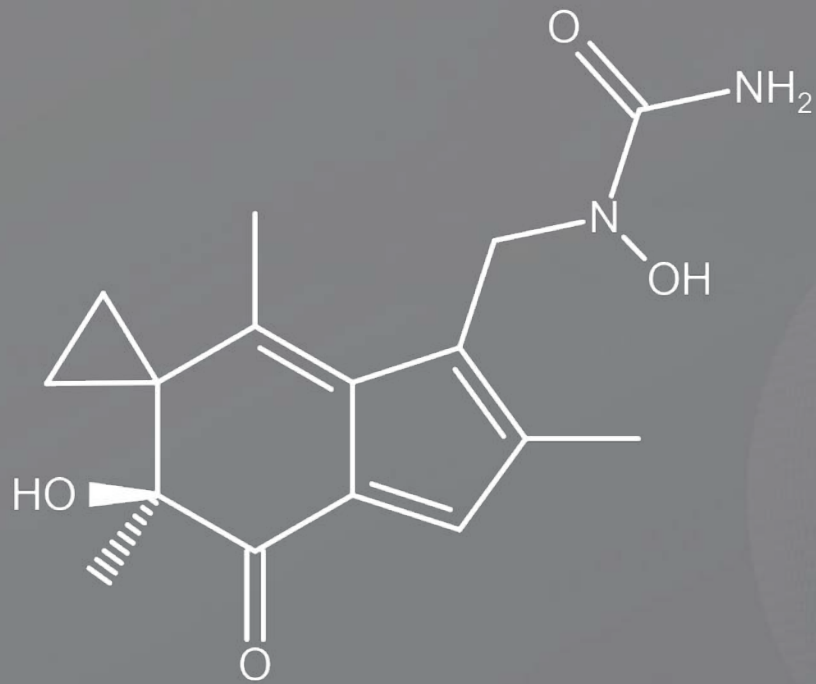
LP-184 Demonstrates Anti-Tumor Efficacy in Mouse TNBC Models and Potential to Sensitize Tumors Non-Responsive to Anti-PD1 Therapy



Treatment arm	Day 22 TGI
■ Anti-PD1 (10mg/kg)	17%
▲ LP-184 (4mg/kg)	51%
▼ LP-184 + anti-PD1	<b>72%</b>

In collaboration with Dr. Shiaw-Yih Lin, MD Anderson Cancer Center

# LP-284 for the Treatment of B-cell Non-Hodgkin's Lymphomas (NHL)



<b>Lead Indications</b>	Mantle Cell, Double Hit Lymphomas, DDR Deficient Non-Hodgkin's Lymphomas
<b>Clinical Status</b>	Phase 1 (Complete response in heavily pre-treated lymphoma patient)
<b>Market Potential*</b>	\$3.75 - 4 Billion
<b>Indication Size*</b>	375,000+
<b>Target/ MOA</b>	Synthetic Lethality
<b>Molecule Type</b>	Acylfulvene Class
<b>Designations</b>	Orphan Drug - Mantle Cell Lymphoma, High-grade B Cell Lymphoma, Soft Tissue Sarcomas
<b>Combination Potential</b>	Rituximab and Spironolactone
<b>IP Estate</b>	Claims extending into 2039

*\*Estimated Annual Global*

# Disease Overview – B-cell Non-Hodgkin’s Lymphomas

Superior responses to LP-284 are observed preclinically

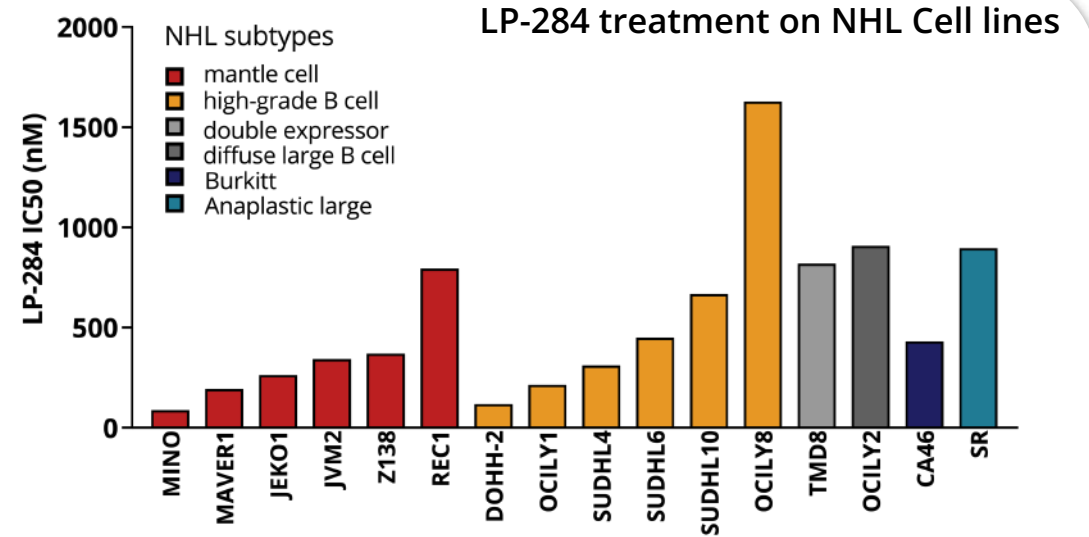
**Annual Global Market Potential: \$ 3-4 Billion**

(NHL)

## B-cell Non-Hodgkin’s Lymphomas

- NHL is a cancer of the lymphatic system and occurs when normal B-cells, T-cells, or Natural Killer (NK)-cells grow out of control
- There are over 30 subtypes of NHL including mantle cell lymphoma (MCL), high-grade b-cell lymphoma(HGBL), and diffuse large B-cell lymphoma

**7th** leading cause of cancer in the US      **4%** of all cancers are NHL in the US



### Mantle Cell Lymphoma

(MCL)

- A rare, aggressive type of B-cell NHL distinguished by overexpression of CCND1
- Small-medium size cancer cells in the lymph nodes, spleen, bone marrow, blood, and gastrointestinal system
- Rarely curable with current standard-of-care treatments and poor prognosis

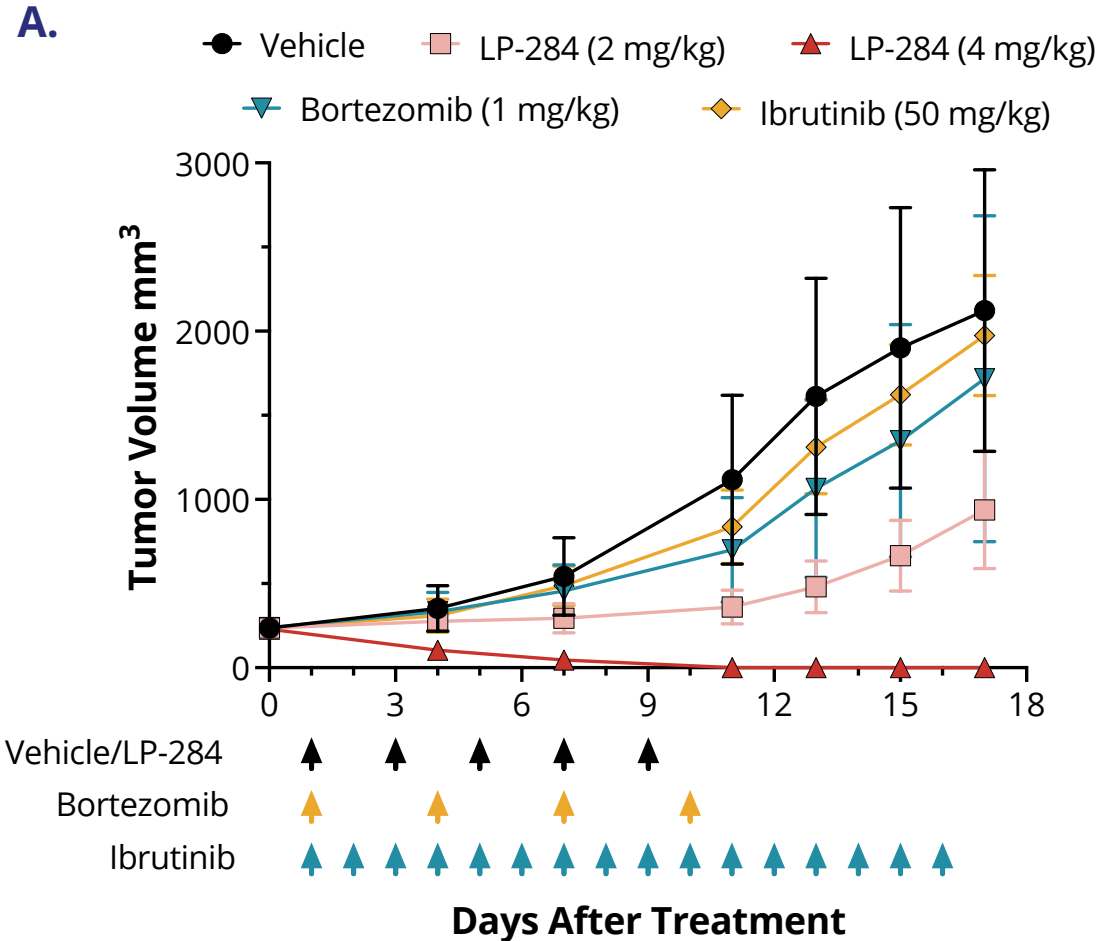
### High-Grade B-Cell Lymphoma

(HGBL)

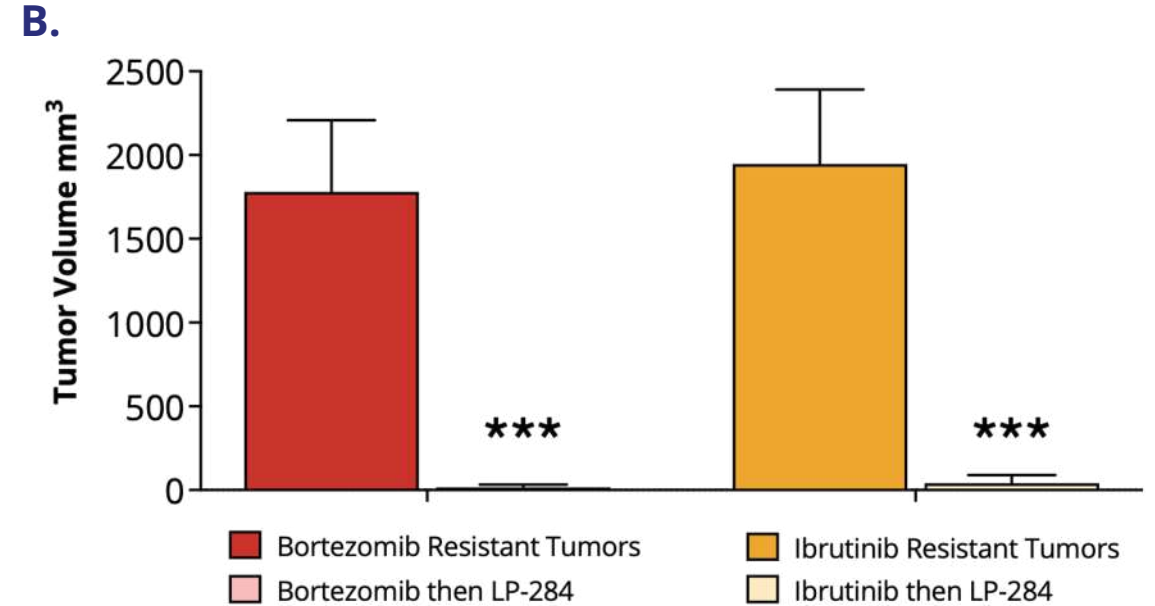
- A rare, aggressive type of B-cell NHL characterized by rearrangements of MYC and BCL2 and/or BCL6 genes
- Often occurs in neck, armpit, groins and can spread to central nervous system
- No standard treatment approach and poor prognosis

# Superior Responses to LP-284 are Observed Preclinically in Several NHLs Including those Resistant to SOC Agents

MCL tumor volumes drastically reduced compared to FDA approved agents in mice models



Tumors resistant to Ibrutinib and Bortezomib has significantly reduced volume



## Nearly all MCL Patients Relapse from SOC Therapies

In cell-derived xenograft MCL models, LP-284 can completely reduce tumors that are resistant to Ibrutinib and Bortezomib

# LP-284 Highlights from NHL Clinical Trial & Potential New Indications

Phase 1a trial for recurrent NHLs with scarce therapeutic options and potential in SLE / Lupus

First-In-Human  
Trial for LP-284



Non-Hodgkin's  
Lymphomas

30-35

Patients expected  
to be enrolled

\$4.0Bn

Estimated global annual  
market potential in NHL

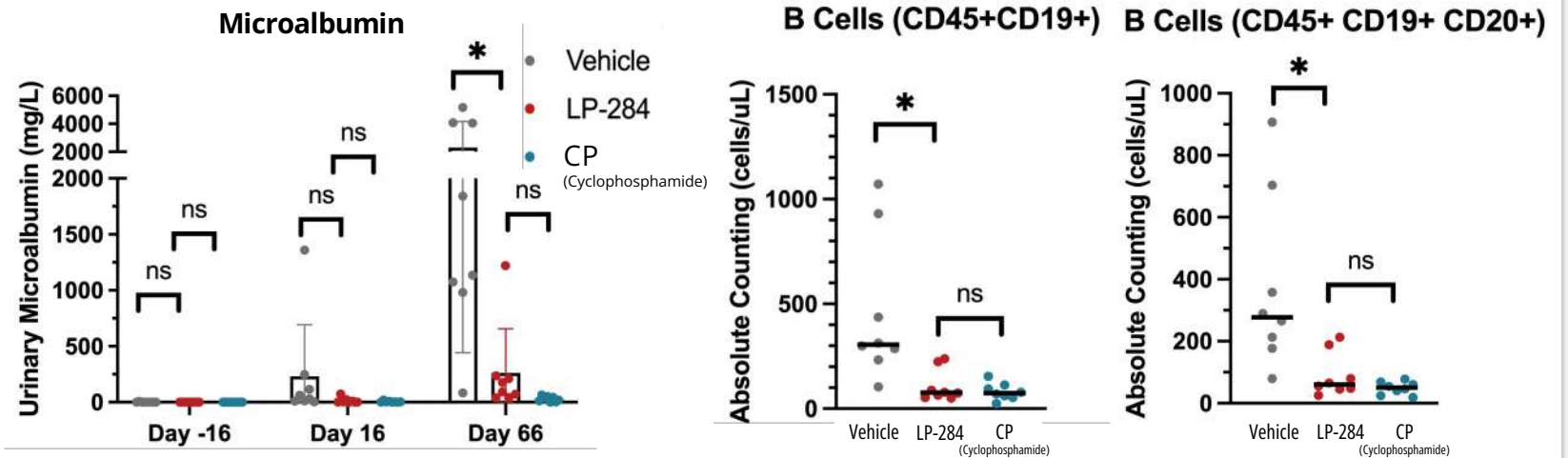


Multi-Site

## Highlights

- Heavily pretreated patient with aggressive Grade 3 B-cell lymphoma (DLBCL) achieved a **complete metabolic response**
- Exploring LP-284 and Rituximab as an alternative to Cyclophosphamide (CP) and Methotrexate in Systemic Lupus Erythematosus (SLE)
- Presented at the Lymphoma Leukemia and Myeloma Congress 2025

## LP-284 + Rituximab: A Potential Next-Generation B-Cell Depleting Therapy for SLE



- LP-284 reduced urinary **microalbumin** ~10x and **B cells** ~4x in an SLE mouse model
- LP-284 + Rituximab combination **further depletes B-lymphoma cells than alone.**

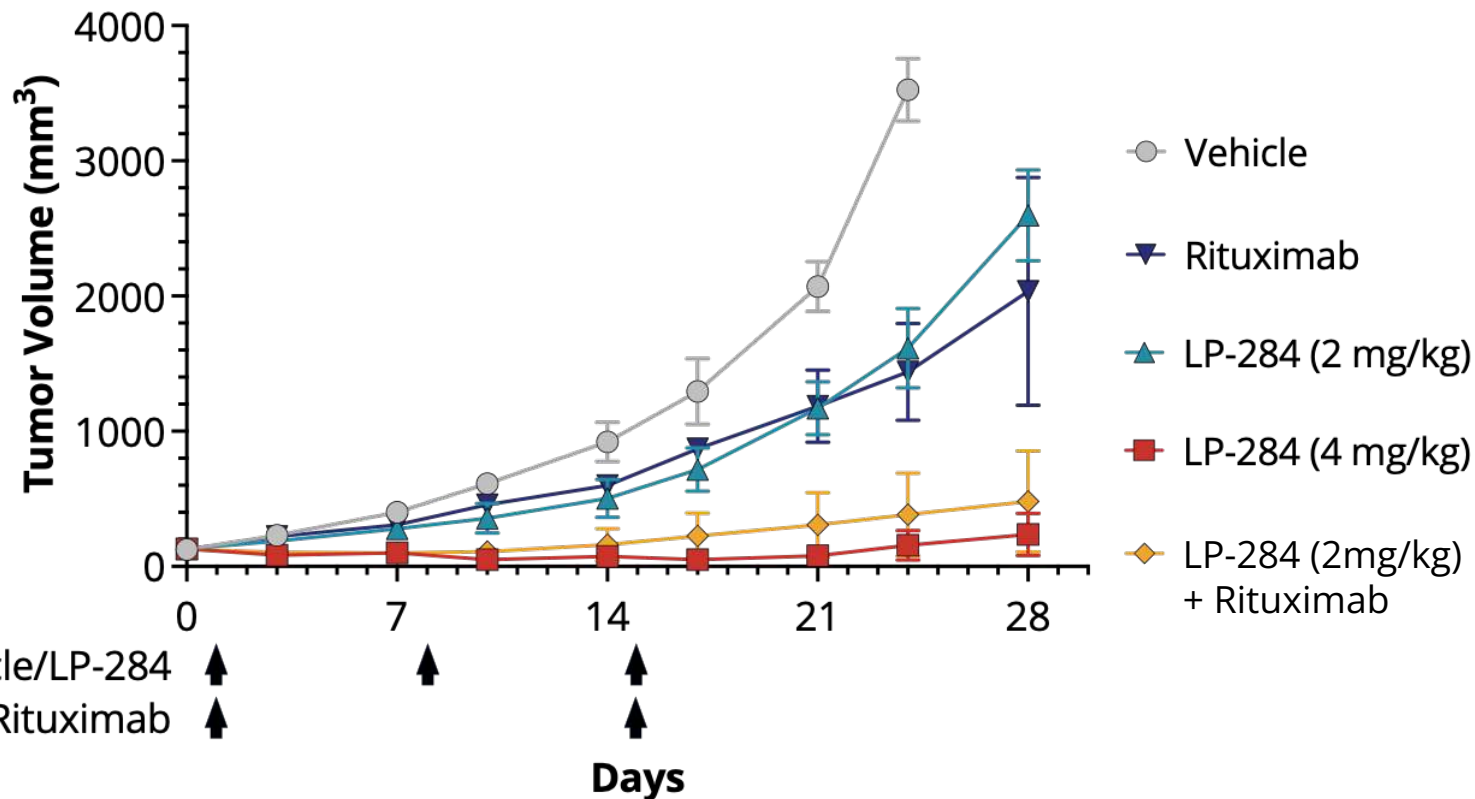
Check out the  
poster now



# LP-284 was Highly Synergistic when Used in Combination with Rituximab in HGBL Xenograft Models

## High Grade B-cell Lymphoma (HGBL) Tumor Volumes in Mice LP-284 – in combination with rituximab

HGBL have universally poor prognosis after chemotherapy, such as EPOCH, Hyper CVAD, and CODOX-M/IVAC - all are given with Rituximab. Novel agents are critically needed for more effective treatments in HGBL



LP-284 treatment led to **near complete tumor growth** inhibition and showed synergistic effects with the FDA-approved agent rituximab

At half of the optimal dose (2mg/kg v. 4mg/kg) **LP-284 when combined with rituximab led to a 63% improvement** in anti-cancer activity (as measured by tumor volumes) versus rituximab alone

- ▼ Rituximab alone = 57% TGI
- ◆ LP-284+ Rituximab = 93% TGI

Results presented at:





Developed from Billions of Datapoints Using AI



\$5-6 Billion Market Potential



Multiple Clinical Stage CNS Cancer Indications



Received Fast Track & Orphan Drug Designation for GBM, Orphan Drug & Rare Pediatric Designation for ATRT



Completed Enrollment for Adult Phase 1a Trial



World Class Collaborators from Johns Hopkins, and UT Health San Antonio

**starlight**  
therapeutics

Scan the QR code for the full Starlight Corporate Overview



# THERE ARE OVER 120 TYPES OF CNS CANCERS AND A MAJORITY HAVE NO CURATIVE TREATMENT OPTIONS

## Starlight's Unique Areas of Focus



**Glioblastoma  
(GBM)**  
13,000/yr in USA

No effective systemic  
therapies have been  
approved for GBM in over  
18 years



**Brain  
Metastases**  
100,000+ /yr

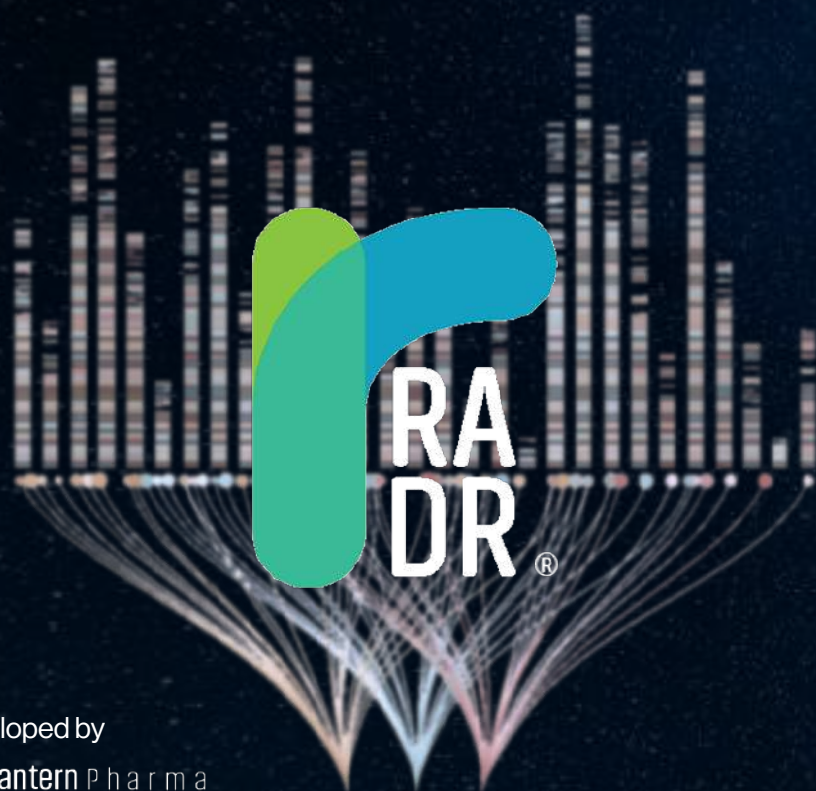
More effective therapies  
are needed to improve  
outcomes for brain  
metastases



**Pediatric  
CNS Cancers**  
4,000/yr in USA

There are no approved  
therapies for atypical  
teratoid rhabdoid tumors  
(ATRT)

# ORIGINATION OF STAR-001: RADR PREDICTIONS POWERED BY AI



## Leading AI Technology developed by Lantern Pharma, RADR®, Helped Identify

- PTGR1 levels correlate with drug response
- Brain penetrant
- GBM has higher levels of PTGR1 relative to normal brain
- Novel alkylation site (at the N<sup>3</sup> adenosine base) causing replication stress and double-strand DNA breaks
- Agnostic to MGMT promoter methylation
- Increased activity with alterations in EGFR and SMARCB1
- Synthetic lethality when co-administered with spironolactone or in tumors deficient in DNA damage repair

Developed by  
 Lantern Pharma

# STAR-001 HAS MULTI-BILLION USD MARKET POTENTIAL IN CNS CANCERS

Annual **5-6B** (USD)

Estimated Market Potential

Glioblastoma

**1.5-2B\***

Annual US Cases **13K**

Other Gliomas

**1.2B\***

Annual US Cases **22K**

Brain metastases

**3B\***

Annual US Cases **100K**

Pediatric CNS Tumors

**0.1B\***

Annual US Cases **4,000**

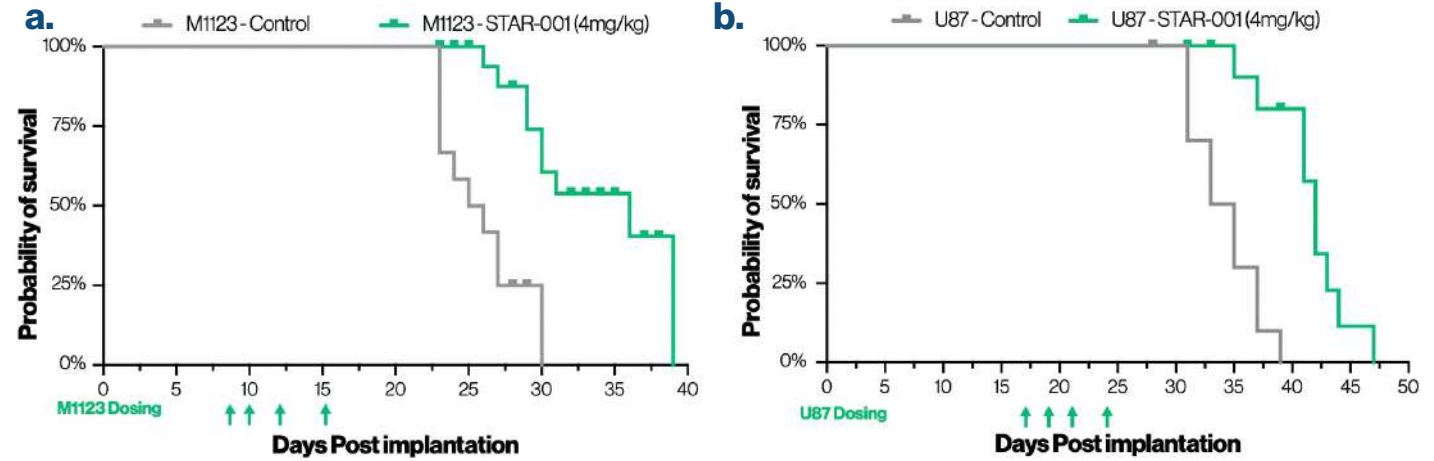
\*Annual Estimated Market Potential

# STAR-001 Treatment Results in Prolongation of Survival in Orthotopic Mouse Models

## KEY TAKEAWAYS

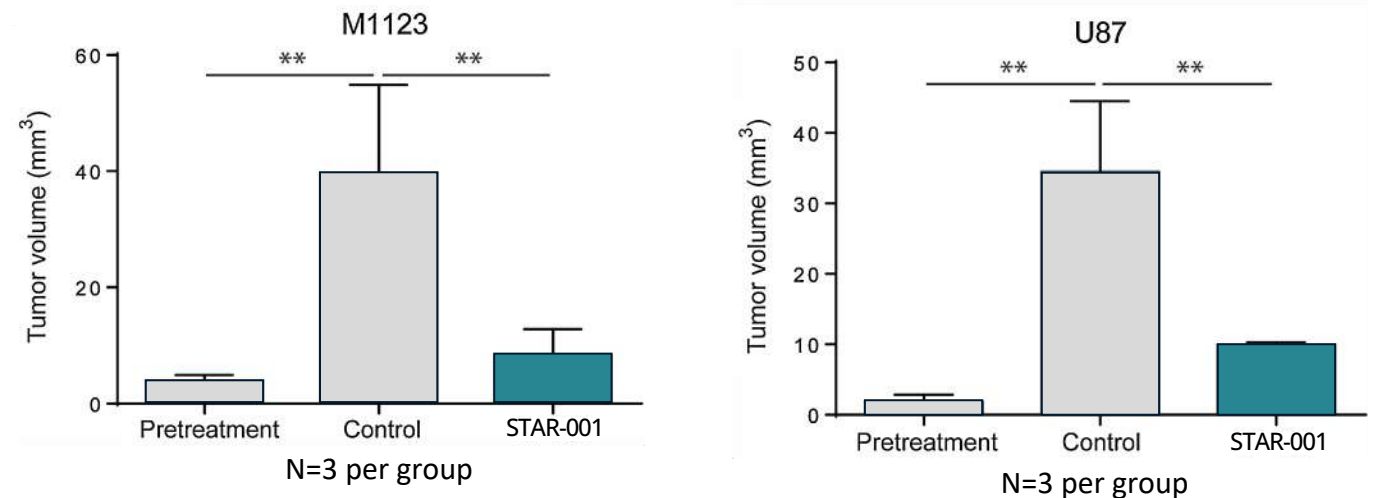
- STAR-001 increased survival of animals by >20%
- STAR-001 reduced M1123 and U87 tumor volume by >75%

### A STAR-001 treatment increased survival of animals :



**a:** Mice with orthotopic M1123 or U87 xenografts received vehicle or STAR-001 (4mg/kg iv) on days indicated by arrow, 10 mice were evaluated for survival. **b:** Tumor volumes, pre-treatment (post implantation day 8 and post treatment-post day 16 implantation for M1123 and for U87 pretreatment (day 16) and post treatment day 25.

### B Tumor volumes before and after STAR-001 treatment



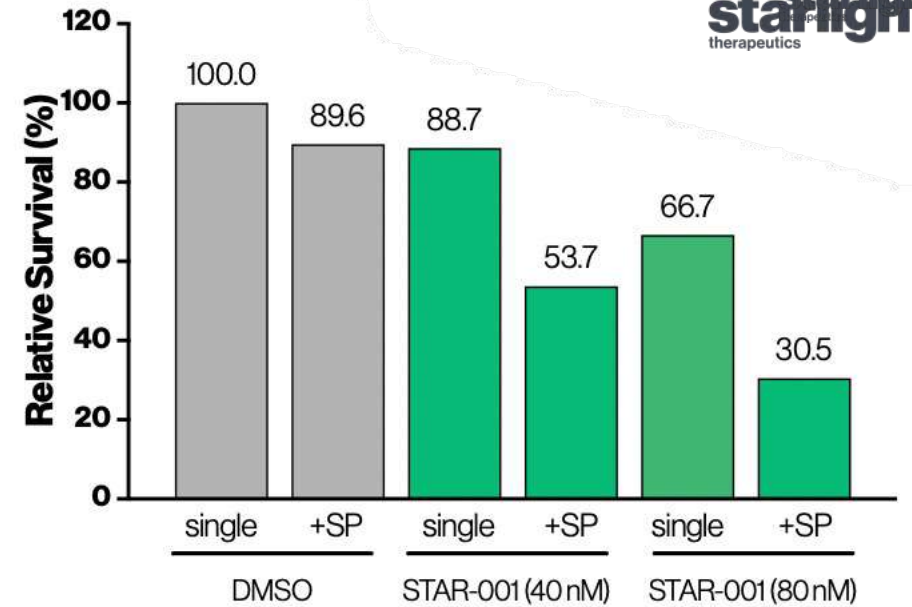
# ATRT *in vivo* Tumors are Exceptionally Sensitive to STAR-001

## KEY TAKEAWAYS

- RADR<sup>®</sup> identified the near universal SMARCB1 mutation and chromatin remodeling deficiency that make ATRT susceptible to STAR-001
- STAR-001 increases survival in ATRT mouse models, decreases tumor volume by >80%
- STAR-001 was granted FDA Rare Pediatric Disease and Orphan Drug Designations to treat ATRT (*Atypical Teratoid Rhabdoid Tumors*)

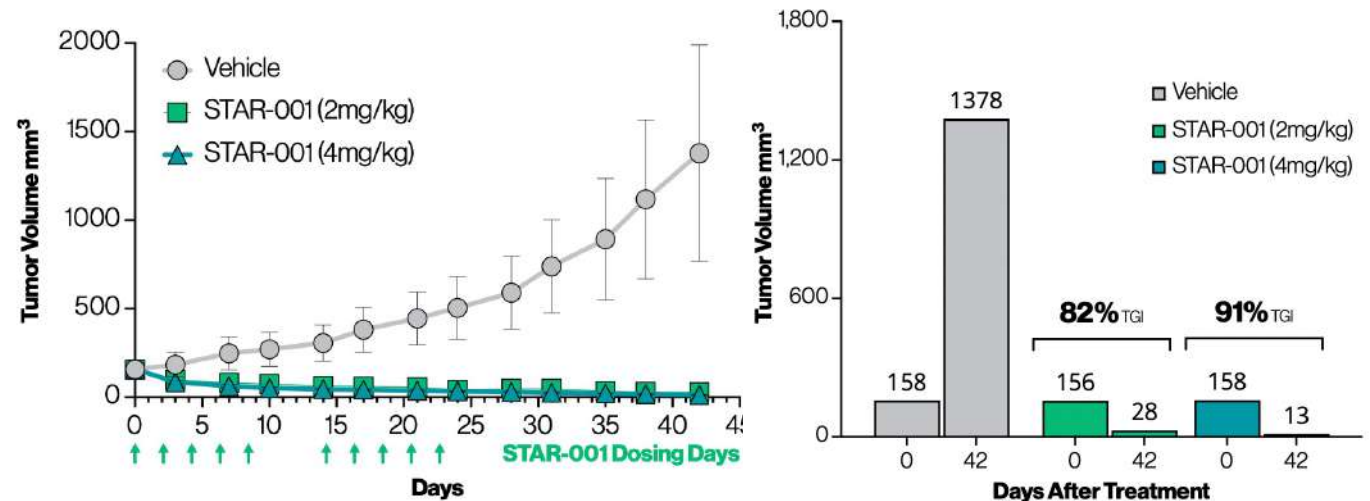
**A**

STAR-001 treatment with Spironolactone in the **ATRT Cell Line CHLA06**







**B**

STAR-001 treatment of **ATRT mouse tumors**



# RESEARCH SUGGEST INCREASING INVESTOR AND INDUSTRY INTEREST IN CNS ONCOLOGY

COMPANY	VALUE	TRANSACTION	DATE	STAGE	COMPOUND	INDICATION
 <p><b>CHIMERIX</b> A Jazz Pharmaceuticals Company</p>	<b>953 M</b>	Acquired by Jazz Pharma	April 21, 2025	PDUFA After Phase II Trial	Dordaviprone	H3K27M Mutant Gliomas
 <p><b>ModifiBio</b></p>	<b>1.3 B</b>	Acquired by Merck	October 23, 2024	Preclinical	MOD246	TMZ-Resistant GBM
 <p><b>Day One</b> BIOPHARMACEUTICALS</p>	<b>461 M</b>	Ex-US rights only sold to Ipsen	July 25, 2024	Phase III Trial	Tovorafenib	RAF-altered PLGG (pediatric low-grade glioma)
 <p><b>agios</b></p>	<b>2 B</b>	Servier acquired Agios CNS oncology	April 1, 2021	Phase II	Vorasidenib	Grade 2 IDH Mutant Gliomas

# IP Portfolio

Intellectual property portfolio builds expanding protections with additional barriers to competition

**100+** Issued Patents & Pending Applications

**5 Families**  
Drug Sensitivity & Response Signatures using Biomarkers

**11 Families**  
Methods of Use

**2 Families**  
Composition of Matter

## RADR



**2041\***

Identifying suitable cancer types and subtypes for a drug candidate



**2043\***

Applying ensemble methods in machine learning and deep learning for drug discovery



**2044\***

Predicting blood-brain barrier permeability

## LP-300



**2041\***

Determining sensitivity to LP-300 based on biomarkers



**2041\***

Treating female (non-smoker) patients with non-small cell lung cancer



Increasing cancer patient survival time using LP-300

## LP-184



**2041**

Treating rhabdoid tumors with LP-184



**2039\***

Treating solid tumor cancers using LP-184 and biomarker



**2041\***

Treating pancreatic cancer using LP-184



**2042\***

Treating cancers with spironolactone and LP-184

## LP-284



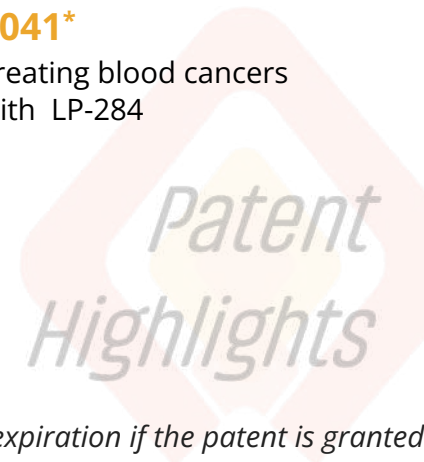
**2040**

Composition of Matter



**2041\***

Treating blood cancers with LP-284



\*Pending patent application. Date referenced indicates estimated year of expiration if the patent is granted.

# Financial Highlights And Cap Table

- Approx. \$10.1M of cash, cash equivalents and marketable securities as of December 31, 2025
- Committed to creating enduring growth and value for LTRN shareholders

LANTERN PHARMA INC. (LTRN)	
Exchange	Nasdaq
52 Week Per Share Price Range (through 3/30/26)	\$1.11 - \$5.74
Common Shares Outstanding (12/31/25)	11.25M
Options (Employees, Management and Directors) (12/31/25)	1.31M
Fully Diluted Shares Outstanding (12/31/25)	12.56M



# Leadership & Board of Directors

## Leadership



### PANNA SHARMA

Chief Executive Officer & President

**PRIOR:** President & CEO, Cancer Genetics (CGIX); CEO & Managing Partner, TSG Partners; Managing Member, Oncospire Genomics (Joint Venture with Mayo Clinic); CSO, iXL Services



### DAVID MARGRAVE

Chief Financial Officer

**PRIOR:** 20+ years of oncology focused management experience; Chairman, Texas Healthcare & Bioscience Institute (current); President & CAO, BioNumerik Pharmaceuticals



### KISHOR BHATIA, Ph.D.

Chief Scientific Officer

**PRIOR:** 40+ years experience in cancer research; Director, Children's cancer Center Riyadh; Director Office of AIDS Malignancy Program, NCI



### REGINALD EWESUEDO, M.D., M.S.c., MBA

VP of Clinical Development

**PRIOR:** VP, Kymera Therapeutics  
VP, Tesaro/GSK  
VP, Pfizer



### MARC CHAMBERLAIN, M.D.

Chief Medical Officer of Starlight

**PRIOR:** Co-director of Neuro-oncology program, UC San Diego; USC; Moffitt Cancer Center; Fred Hutchinson Cancer Center; Medical Director, Cascadian Therapeutics; SeaGen; SystImmune; Pionyr Immunotherapeutics



### SANDRA SINCLAIR, BSBA, MHA/ED, RN

Executive Director, Clinical Operations

**PRIOR:** 30+ years in clinical operations and trial execution  
Alaunos Therapeutics; SAB  
Biotherapeutics; MD Anderson Cancer Center

## Board of Directors

Donald "Jeff" Keyser, J.D., MPH, Ph.D.

*Non-executive Chairman*

David Silberstein, Ph.D.

Vijay Chandru, Ph.D.








Maria Maccacchini, Ph.D.

Panna Sharma








*CEO and President*

# 2026 Investment Highlights

## Recent Milestones

-  Preliminary patient data showing an 86% clinical benefit rate in the initial safety lead-in cohort of the Harmonic™ Phase 2 Trial
-  Reported a durable complete response in a Harmonic™ trial patient, with survival continuing for nearly two years
-  Delivered complete metabolic response after two cycles of LP-284 for in a heavily pre-treated lymphoma patient
-  Received three rare pediatric disease designations for LP-184 in malignant rhabdoid tumors (MRT), rhabdomyosarcoma (RMS), and hepatoblastoma
-  Received orphan drug designation from US FDA for LP-284 in soft tissue sarcomas
-  Expanded RADR® AI platform to 200+ billion datapoints and launched initial modules publicly
-  Expanded the Harmonic™ trial to Taiwan and Japan with 5 sites in each country and completed enrollment in Japan

## Upcoming Milestones & objectives

-  Complete further analysis of Phase 1a LP-184 results; pursue Phase 1b/2 and investigator or grant led opportunities
-  Advance enrollment for LP-284 in NHL + soft tissue sarcoma and initiate partnering discussions
-  Report initial clinical data for Asian cohort in the Harmonic™ Trial and updates on the US patient population
-  Progress and monetize Starlight Therapeutics towards Phase 1/2 adult & pediatric clinical trials
-  Expand RADR® AI & withZeta.ai platforms and develop additional revenue opportunities with AI for drug development
-  Further ADC preclinical and IND development to support future Phase 1 launch and/or partnership
-  Develop and communicate combination programs and trials for Lantern's portfolio with existing FDA approved drugs



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