

Corporate Overview June 3rd, 2024

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[®] 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[®] 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 our research and the research of our collaborators may not be successful, (ii) the risk that promising observations in preclinical studies do not ensure that later studies and development will be successful, (iii) the risk that we may not be successful in licensing potential candidates or in completing potential partnerships and collaborations, (iv) 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, (v) the risk that no drug product based on our proprietary RADR[®] AI platform has received FDA marketing approval or otherwise been incorporated into a commercial product, and (vi) those other factors set forth in the Risk Factors section in our Annual Report on Form 10-K for the year ended December 31, 2023, filed with the Securities and Exchange Commission on March 18, 2024. You may access our Annual Report on Form 10-K for the year ended December 31, 2023 under the investor SEC filings tab of our website at www.lanternpharma.com or on the SEC's website at 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 Al platform, RADR[®], is transforming the cost, pace, and timeline of cancer drug discovery and development

12

Lead drug candidates* powered by Al Clinical stage lead drug candidates*

5

100+

Issued patents & pending applications

\$38.4M**

Cash/cash eq./ marketable securities

2.5 years

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

\$1.5M

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

* Includes drug programs being developed in collaboration ** at 3/31/2024

Only **6%*** of clinical trials using traditional drug discovery approaches succeed

*Clinical Development Success Rates and Contributing Factors 2011–2020, BIO Stats

Current Challenges



Average cost to bring a new cancer drug to market is **\$2.8 billion**



Out of 20,000 trials from 2012-2022, **19,200 trials failed**



Early-Stage development takes **3-5+ Years**, late-stage development takes **6-12+ Years**

Current oncology drug development is being improved by **data-driven**, and **AI-enabled approaches** and technology

Lantern is Transforming Drug Discovery Timelines & Costs with AI

Al 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

RA DR 。



Large Scale/Multi-omics Oncology Data



Proprietary Al platform RADR[®]



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Accelerated timelines; reduced costs and risks

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





Precision Medicine Platform

Response Algorithm for **D**rug Positioning & **R**escue

A proprietary integrated data analytics, experimental biology, oncologyfocused, machine-learning-based platform focused on drug development



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**

Discover mechanism of action of any compound or drug



m1

Identify/prioritize a compound's disease indications or subtypes

m3

Determine optimal drug combos to improve therapeutic potential

Generate ML-driven m4 biomarker signatures for patient selection



Characterize specialized attributes of a molecule such as BBB permeability



Enhance the selection of optimal combination of ADC components



Discover drug combos for checkpoint inhibitors to improve therapeutic index

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



According to Deep Pharma Intelligence (May 04, 2022)



RADR®'s AI Framework

RADR®'s AI framework develops actionable insights using billions of datapoints



RADR[®] Case Study – Actuate Therapeutics

Advanced RADR[®] machine learning models predict clinical trial patient responses at 88% accuracy





Lantern is accelerating the development of Actuate Therapeutic's drug candidate, Elraglusib^{*} (9-ING-41), using AI insights produced by RADR[®]

- Predicted patient response with greater than 88% accuracy
- Identified metastatic melanoma patients resistant to PD-1 therapies may benefit from Elraglusib
- Insights and new data including RNA, ctDNA, and protein biomarkers are informing design of an upcoming Phase 2 clinical trial
- Lantern received equity in Actuate as part of the collaboration





*Elraglusib is a widely researched GSK-3β inhibitor. Currently, Elraglusib is in multiple active Phase I/II clinical trials as a monotherapy and in combination with other agents (<u>NCT03678883)</u>



Collaborations

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



Biopharma Collaborations







OREGON

THERAPEUTICS

Lantern's Diverse & Unique Al Driven Pipeline of Drug Programs

Lantern has 12 disclosed and collaborative lead drug programs including the Phase 2 Harmonic[™] trial

Lantern Pharma (NASDAQ: LTRN)				🚫 Lantern Pharma			
Lead Candidate	Indication	Discovery	Preclinical	Phase I	Phase II	Orphan Designation	Rare Pediatric Disease
LP-300	Non-Small Cell Lung Cancer for Never Smokers				Marmonic		
LP-184	Recurrent Advanced Solid Tumors (Pancreatic, TNBC, Bladder, & Other Solid Tumors)					* for Pancreatic & HGG	
LP-284	Recurrent Non-Hodgkin's Lymphomas (Mantle cell, Double-hit lymphomas, & HGBL)					for Mantle Cell & HGBL	
ADC	Select Solid Tumors						

RADR [®] Colla	borations	RA Precision Medicine Platform
Elraglusib owned by - Actuate Thera.	Multiple Solid Tumors	Collaboration partner
TTC-352 owned by- TTC Oncology	ER+ Breast Cancers	Collaboration partner
XCE853 owned by – Oregon Thera.	Protein Disulfide Isomerase (PDI) Inhibitor	Collaboration Definition Collaboration Colla
ADC	Cryptophycin Conjugate for Solid Tumors	Collaboration UNIVERSITÄT partner BIELEFELD



Starlight's pipeline is focused on multiple CNS indications in both adult and pediatric patients



Starlight Therapeutics

ADULT CNS CANCERS

Lead Candidate	Indication	Discovery	Preclinical	Phase I	Phase II	Orphan Designation	Rare Pediatric Disease
	Glioblastoma (GBM)*					•	
STAR-001	Brain Metastases (TNBC)**						
	Brain Metastases (NSCLC)**						

* Multiple GBM patients have been enrolled in the ongoing phase 1a being conducted by Lantern Pharma

**The MTD from the ongoing Phase 1A LP-184 clinical trial is expected to support the later expansion to brain metastases

PEDIATRIC CNS CANCERS

STAR-001	Atypical Teratoid Rhabdoid Tumors (ATRT)	Pediatric CNS	•	•
	Diffuse Midline Glioma (DMG)	indications will enter clinical trials after the adult		
	High-Grade Hemispheric Glioma	trials begin		



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



Lead Indication	Indication Relapsed NSCLC for Never Smokers	
Clinical Status	Phase 2 (multiple patients dosed)	
Market Potential*	\$1.3 billion (USD)	
Indication Size*	20,000-40,000 Cases	
Target/ MOA	Tyrosine Kinases & Cell Redox Enzymes	
Molecule Type	Disulfide Small Molecule	
Combination	With carboplatin and pemetrexed	
IP Estate	Claims extending to at least 2032	
	*Estimated Annual LIS	



Disease Overview – NSCLC in Never Smokers – LP-300

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

Global Annual Market Potential: \$2.5+ Billion



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



initiation 6

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

20,000-40,000

never smokers will be diagnosed with NSCLC each year In the US *Cancer.gov*

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

*Subpopulations receiving paclitaxel/cisplatin



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NASDAQ: LTRN

Mechanism of Action – LP-300

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

- 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)

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

*after progressing from TKI

Patients will receive standard of

Primary Outcomes: Overall and progression free survival

care (pemetrexed and carboplatin)

Accelerating recruitment efforts for a growing indication with limited treatment options



- Initial patients dosed in first half of 2023
- Multiple additional patients and sites across the US anticipated to be enrolled during 2024
- Received regulatory approval to expand the trial in both Japan and Taiwan in Q1 2024



LP-184 for the Treatment of Advanced Solid Tumors



Lead Indications	DDR deficient solid tumors including Pancreatic cancer, Bladder cancer, and TNBC
Clinical StatusPhase 1a (multiple patients dosed)	
Market Potential*	\$14+ Billion
Indication Size*	170,000 + Cases, Estimated 400,000 + Cases Global
Target/ MOA	Double-stranded DNA breaks; alkylates DNA in the 3' of Adenine
Molecule Type	Acylfulvene Class
Combination Potential	Checkpoint inhibitors, PARP inhibitors, Spironolactone, Chemotherapy and Radiation Therapy
IP Estate	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: \$14+ Billion

(DDR Deficient Solid Tumors)



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
- Current treatment options include ; surgery, chemotherapy, radiation therapy, targeted therapy, and immunotherapy

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 "Achiles Heel" for drugs leveraging synthetic lethality

Mechanism of Action – LP-184

LP-184 has a unique mechanism of action – leveraging synthetic lethality



Al insights generated by RADR[®] – LP-184

LP-184's MoA was predicted by RADR® and validated with In vitro and In vivo studies



- **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[®] 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[®] insights and that LP-184 was highly potent in cells with PTGR1

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RA

LP-184 treatment results in complete regression in DDR deficient pancreatic cancer 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



TNBC PDX Tumor Models



🔪 Lantern Pharma

LP-184 shows significantly improved survival and tumor shrinkage in GBM xenografts



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Cancer models with common DNA damage response deficiencies are highly sensitive to LP-184 treatment



LP-184 in **NERD** Cancers

- LP-184 shows exquisite potency in cancers with deficiencies in Nucleotide Excision Repair (NERD) pathways
- There are currently **no approved therapies** for NERD cancers

LP-184 in HRD Cancers

PDX Cancer model	IC50 (nM)	HRD Mutations	
NSCLC	31	ATM	
Prostate	31	PMS2	
Pancreatic	45	ATR, BRIP1, PARP1	
NSCLC	54	CHEK1, FANCA, NBN, RAD50	
Prostate	54	BRCA2, ATM, FANCA, FANCI, FANCM	
Prostate	54	BRCA2, CDK12, FANCI, RAD54L,	
NSCLC	57	ATM, FANCD2, NBN	
Pancreatic	57	BRCA1, BRIP1,	
Prostate	92	ATM, ATR, PALB2,	
Pancreatic	110	BRCA2, ATM, BLM, FANCA	
Pancreatic	270	BRCA2, CDK12, PALB2	
Pancreatic	2,900	ATM, BRCA1, BRCA2	

- PDX-derived cell lines with mutations in key HR and NER genes are highly sensitive to LP-184
- Only 1 model was not highly sensitive to LP-184 (highlighted in blue)

Clinical Trial – LP-184 Phase 1 Basket Trial

Launched Phase 1 basket trial for a blockbuster molecule with a market potential of \$10+ billion in annual sales



(through Starlight) and Phase 1b/2 in other solid tumors to be initiated after determination of MTD

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 $(n \ge 3, BOIN)$

*BOIN- Bayesian Optimal Interval

Preclinical Data on Combination Therapy – LP-184

In-vivo LP-184 has synergy with several SOC agents including spironolactone, radiation therapy, and others



LP-184 + Radiation in the Panc03.27 CDX Model



- **Spironolactone** is an FDA approved agent that can impair DNA damage repair pathways in tumor cells
- **Combination** of LP-184 or LP-284 with Spironolactone:
 - 1) Enhances potency
 - 2) Decreases expected dose needed for treatment
 - 3) Exploits MoA of both drugs

- Terminal tumor volumes from the RT + LP-184 treatment group are significantly (**p < 0.01) smaller than treated with RT or LP-184 alone
- Mean tumor volumes of RT + LP184 were ~1.8 fold lower than tumors treated with LP-184 alone

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



Lead Indications	Mantle Cell, Double Hit Lymphomas, DDR Deficient Non- Hodgkin's Lymphomas
Clinical Status	Phase 1 (multiple patients dosed)
Market Potential*	\$3.75 - 4 Billion
Indication Size*	375,000+
Target/ MOA	Synthetic Lethality
Molecule Type	Acylfulvene Class
Designations	Orphan Drug - Mantle Cell Lymphoma
Combination Potential	Rituximab and Spironolactone
IP Estate	Claims extending into 2039

*Estimated Annual Global



Disease Overview – B-cell Non-Hodgkin's Lymphomas

Superior responses to LP-284 are observed in several B-cell lymphomas

Annual Global Market Potential: \$ 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

the US

leading cause of cancer in the US

of all cancers are NHL in the US



Mantle Cell Lymphoma

(MCL)

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

• Small-medium size cancer cells in the lymph nodes, spleen, bone marrow, blood, and gastrointestinal system

• Often occurs in neck, armpit, groins and can spread to central nervous system

A rare, aggressive type of B-cell NHL distinguished by overexpression of CCND1

Rarely curable with current standard-of-care treatments and poor prognosis

• No standard treatment approach and poor prognosis.

Superior responses to LP-284 are observed in several NHLs including those resistant to SOC agents



LP-284 reduced the volume of tumors resistant to Ibrutinib and Bortezomib



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

Clinical Trial – LP-284 Phase 1 Trial

Ph. 1 trial launched in Q4 2023 for recurrent NHLs with scarce therapeutic options



Preclinical Data on Combination Therapy – LP-284

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 FDAapproved 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:



Advanced the development, synthesis, and preclinical proof-of-concept of a novel, highly potent, cryptophycin-based ADC

Moderate HER2 Expression

ADC Collaboration Update

UNIVERSITÄT BIELEFELD

Collaboration Led by Professor Norbert Sewald, Ph.D.

High HER2 Expression



Key Highlights

- The cryptophycin(Cp) drug-payload and Cp-ADC averaged an 80% cancer cell kill rate
- In a moderate Her2 expression model, the Cp-ADC with a DAR* of 8 (*Tras(C8)-Cp*) was about **10x more potent** than a DAR 8 MMAE**-ADC (*Tras (C8)-MMAE*)
- Cp-ADC showed highly efficient anti-tumor activity in all six cancer cell lines (breast, bladder, colorectal, gastric, pancreatic, and ovarian cancer) with EC-50 values in the picomolar to single-digit nanomolar range
- Additional studies are now being developed to further validate and expand these findings to obtain a deeper understanding of the genomic and biomarker correlates of payload efficacy

*drug to antibody ratio

**Monomethyl auristatin E - potent tubulin inhibitor that is used as the payload for four FDA-approved ADCs

Al-powered module has potential to deliver differentiated and derisked ADCs faster & with significantly reduced costs



Leveraging AI to deliver novel, differentiated ADCs

Ability to select and characterize among potent and super potent payloads with specific and optimized molecular and biochemical characteristics

The ADC module will continue to grow by ingesting and learning from billions of data points each quarter based on both experimental and real-world data

Ability to predict synergy of payloads and antibodies in certain tumors based on both the tumor environment and biological impact

RADR[®] insights generated by understanding the impact of mutations on heterogeneous target expression patterns in cancers can help improve treatment response, enabling personalized targeting of ADCs



Stantight therapeutics

Born from Billions of Datapoints & AI, Starlight has Blockbuster Potential to Provide New Treatment Options for 500,000+ Patients

There are over **120 types of central nervous system (CNS) and brain cancers** and a majority have **no effective treatment options**

- No effective single-agent therapies have been approved for glioblastoma (GBM) in over 18 years
- Effective therapies are needed to improve outcomes for brain metastases patients
- There are no approved therapies for atypical teratoid rhabdoid tumors (ATRT)

5 Year Survival Rates of CNS And Brain Cancers Remain Low Despite Advances in Cancer Therapies



- staright therapeutics
- 500,000+ Potential CNS Patients Globally*
- Multiple Clinical-stage CNS Cancer Indications
- STAR-001 has been Granted FDA Orphan Drug Designation for GBM & ATRT and Rare Pediatric Disease Designation for ATRT
- World Class Collaborators from Johns Hopkins, UT Health San Antonio, and Children's Brain Tumor Network
- 4 US Patents & Patent Applications and 10+ Foreign Pending Patent Applications





STAR-001 has potent anti-tumor efficacy in methylated and unmethylated GBM mouse models

STAR-001 demonstrates **3,000X** higher *in vitro* potency **compared to temozolomide -** independent of MGMT status



STAR-001 KEY TAKEAWAYS

- No effective single-agent therapy has been approved for adult GBM in over 18 years
- STAR-001 was granted an FDA Orphan Drug Designation to treat malignant gliomas including GBM
- STAR-001 has shown effectiveness in both MGMT(+,-) forms of GBM cell lines
- Planning for launch of Phase 1b/2 in second half of 2024

Unique position of STAR-001 compared to current therapeutic options



CURRENTLY APPROVED MAINSTAY GBM THERAPIES

	STAR-001	Temozolomide (TMZ)	Nitrosourea (CCNU)
Molecular weight	304 kD	194 kD	233 kD
Derivation	Mushroom Omphalotus illudens [*]	Synthetic	Synthetic Nitrogen Mustards
Chemistry	Acylfulvene	Imidazotetrazine	Cyclohexylamine + 1-chloro-2-isocyanatoethane
Drug schedule	Intravenous D1, 8 q21d	Oral daily or D1-5 q28d	Oral D1q6 weeks
Mechanism of action	dsDNA breaks @ N ³ adenosine	ssDNA breaks @ O ⁶ & N ⁷ guanine (methyl)	ss & dsDNA breaks @ O ⁶ guanine (chloroethyl)
DNA repair system	TC-NER&HR	MGMT	HR
Tumor/blood concentration ratio	0.2	0.2	0.9
IC_{50} (varies by cell line)	100~ 800 nM	500 μΜ	50 μΜ
Bioactivation	Prodrug, conversion by intracellular PTGR1	Prodrug, spontaneous conversion by hydrolysis to MTIC	Prodrug, spontaneous conversion by hydrolysis
Elimination, half life	Kidney, <30 minutes	Kidney, 2 hours	Kidney, <5 minutes

*Synthetic manufacturing route

IP Portfolio

Intellectual property portfolio builds expanding protections with additional barriers to competition



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Recent Posters/ Publications

Highlighting the strong validation of RADR[®] insights, de-risking the development of Lantern's drug candidates



LP-184, a novel acylfulvene molecule, exhibits anticancer activity against diverse solid tumors with homologous recombination deficiency May 2024



Targeting homologous recombination deficiencies in B-cell non-Hodgkin's lymphomas with the novel anti-tumor small molecule LP-284 Sep 2023



Phase 1a/1b clinical trial of LP-284, a highly potent TP53 mutation agnostic DNA damaging agent, in patients with refractory or relapsed lymphomas and solid tumors (NCT06132503) April 2024 LP-184, a clinical stage acylfulvene-derived tumor site activated small molecule, inhibits adult and pediatric CNS tumor cell growth Aug 2023



Preclinical Efficacy of LP-184, a Tumor Site Activated Synthetic Lethal Therapeutic, in Glioblastoma

Oct 2023

<text>
 Artificial intelligence platform,
 Strange of the strange

Artificial intelligence platform, RADR[®], aids in the discovery of DNA damaging agent for the ultrarare cancer Atypical Teratoid Rhabdoid Tumors Oct 2022



Financial Highlights And Cap Table

Solid financial position and capital efficiency fuel continued growth anticipated to provide a cash runway into at least Q3 2025

- Approx. **\$38.4 M of cash, cash equivalents and marketable securities** as of March 31, 2024
- Committed to creating enduring growth and value for LTRN shareholders

LANTERN PHARMA INC. (LTRN)				
Exchange	Nasdaq			
52 Week Per Share Price Range (through 6/3/24)	\$2.38 - \$11.99			
Common Shares Outstanding (3/31/24)	10.76M			
Warrants (3/31/24)	81.5K			
Options (Employees, Management and Directors) (3/31/24)	1.08M			
Fully Diluted Shares Outstanding (3/31/24)	11.92M			





Lantern's 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 Theraputics
- VP, Tesaro/GSK
- VP, Pfizer



MARC CHAMBERLAIN M.D.

Chief Medical Officer

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



PETER CARR

Principal Software Architect

PRIOR:

- Sr. Software Engineer, Broad
 Institute Cancer Program
- Sr. Programmer/Analyst, Boston Univ Science & Math Education Center





Investment Highlights

Lantern Pharma (NASDAQ: LTRN)



Proven drug rescue and drug development process and in the clinic with 3 compounds and accelerating additional compounds and combinations to clinical trials...potentially saving tens of millions of dollars and years of development



Growing AI based platform with clear roadmap to 100+ Bn. datapoints focused exquisitely on cancer therapeutic development and companion Dx in a high growth, high demand \$12+ Bn. market



Focused on cancer drug market segments with clear clinical need, understood mechanisms, targeted patient populations that exceed 1 million , and multi-billion USD in annual sales potential



A novel AI-powered ADC platform with the potential to develop and out-license or partner ADC assets in early phases



Several compounds in place with multiple targeted indications, including LP-184 and LP-284 (received Orphan Disease Designations in pancreatic and GBM & Rare Pediatric Disease Designation for ATRT), which can help accelerate development



Proven and growing library of AI & machine-learning methodologies published at ASCO, AACR, and SNO used to generate novel IP & patents and accelerate discovery by potentially years



Experienced and innovative management team w/ 70+ years experience in cancer and a passion to change the cost and outcome for cancer patients by using AI and genomics – *paradigm changing technologies*



Industry leading collaborations with Johns Hopkins, UT Health San Antonio, Fox Chase Cancer Center, and University of Bielefeld

2024 Investment Highlights

est Recent Milestones

Dosed initial patients in the Harmonic[™] clinical trial

- Launched Phase 1A basket trial for LP-184 and multiple patients dosed
- Launched Phase 1A trial for LP-284 and initial patients dosed
- Advanced collaboration with Bielefeld University to develop breakthrough ADCs using AI
- Developed industry leading AI algorithms to predict any compound's ability to cross the BBB
- Expanded RADR[®] AI platform to 60+ billion datapoints
- Received orphan drug designation for LP-284 in High-Grade B-cell Lymphoma

Upcoming Milestones

- Complete Phase 1a clinical trial for LP-184; commence Phase 1b and investigator led trial(s)
- Accelerate enrollment in first-in-human clinical trial for LP-284 in NHL + other cancers
- ♥ Commence enrollment of The Harmonic[™] Trial in targeted sites in Asia
- Progress Starlight Therapeutics towards Phase 1 / 2 adult & pediatric clinical trials
- Expand RADR[®] AI platform to 100+ billion datapoints and develop additional collaborations
- Further ADC preclinical and IND development to support future Phase 1 launch and/or partnership



Develop combination programs for LP-184, LP-284, and LP-300 with existing approved drugs

Lantern Pharma_® NASDAQ: LTRN

IR Contact: IR@lanternpharma.com 1-972-277-1136



- www.lanternpharma.com
- X <u>@LanternPharma</u>
- in linkedin.com/company/lanternpharma