



# Investor Presentation

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**Pioneering Differentiated Radiotherapies  
for Patients with Unmet Needs**

April 2026

ATNM: NYSE AMERICAN

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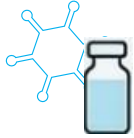
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# Investment Thesis: Differentiated Radio Pipeline with Strong Momentum

*Novel multi-asset portfolio at an inflection point with several key data readouts in 2026 supported by strong balance sheet providing runway into 2028*



## 2 Pan-Tumor Blockbuster Solid Tumor Opportunities

ATNM-400: First-in-class pan-tumor asset supported by compelling preclinical data in mCRPC, NSCLC & Breast Cancer  
Actimab-A (MDSC): Potential to unlock the power of immunotherapy by depleting MDSCs in solid tumors



## De-risked Partner Ready Hematology and Conditioning Franchise with 2 Phase 2/3 Assets

Late-stage (Phase 2/3 ready) Actimab-A and Iomab-B can address the entire AML treatment journey and Iomab-ACT has the potential to be a universal conditioning agent for cell and gene therapies, representing a near-term value opportunity



## Uniquely Positioned Radiotherapies with Novel, First-in-Class Therapies

Poised to fill the innovation gap in the radiopharmaceutical sector, where patients' needs and big pharma's appetite require innovation beyond PSMA, SSTR and FAP



## Robust Capabilities, Platform and Near-Term Catalysts

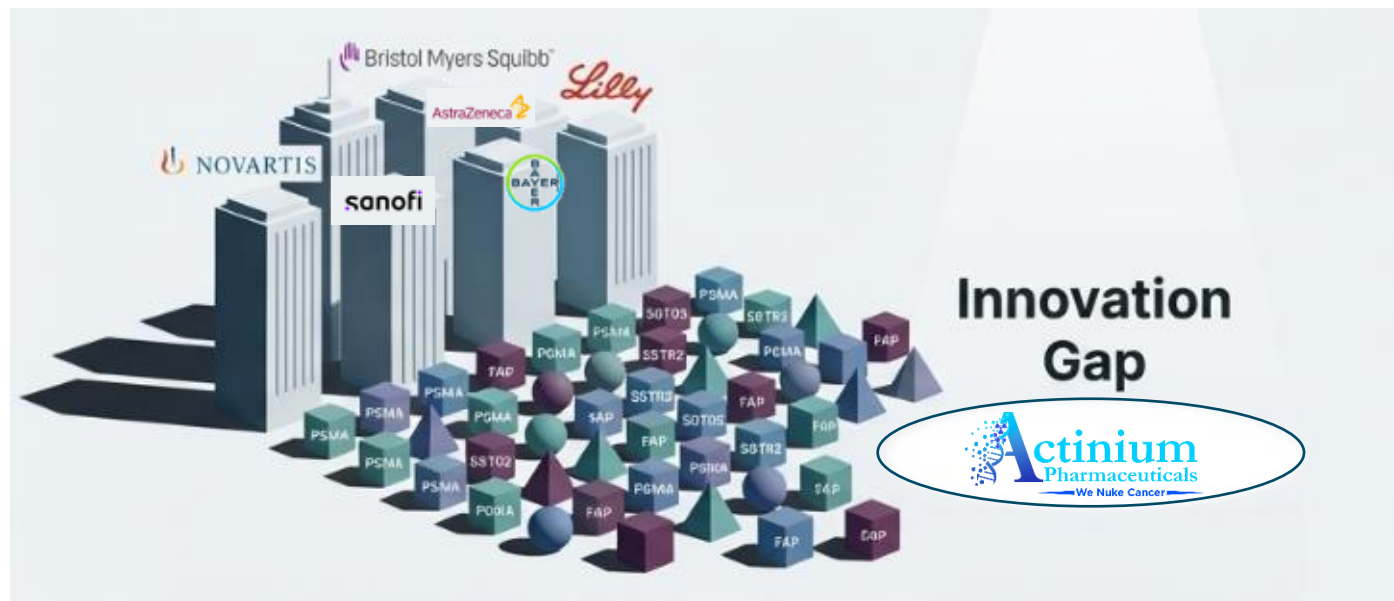
Supported by deep R&D expertise, expanding in-house manufacturing capabilities (operational 2H:26), robust IP approximately 250 patents including Ac-225 production technology and a cascade of significant data readouts expected in 2026



## Strong Financial Position & Clean Capital Structure

Cash runway into 2028 (\$48.0 million in cash and cash equivalents as of December 31, 2025), 31.4 million shares outstanding with no debt, warrants or preferred stock outstanding

# Radiopharma Landscape: A Strategic Void Awaits Differentiated Assets



**6 Big Pharma's**

with an acquired radiotherapy presence

**3 Targets**

Dominate industry pipeline (PSMA, SSTR2 & FAP)

**\$17 Billion**

In high value M&A on late-stage assets and infrastructure

**300,000+**

Sq. feet of underutilized radio mfg. infrastructure




The radio landscape features largely undifferentiated programs by **50+ smaller companies** and **6 Big Pharma players**. A wave of M&A has left them with large infrastructure but sparse pipelines, creating an urgent need for truly novel assets

**\$10+ Billion** in investments focused on novelty & differentiation in 2024 – 2026 YTD

<p><b>\$2.7 B</b> Peptide platform</p>	<p><b>\$1.75 B</b> DLL3 &amp; B7-H3 &amp; platform</p>	<p><b>\$1.1 B</b> Mini-protein platform</p>	<p><b>\$1.1 B</b> GCPR program</p>	<p><b>\$0.65 B</b> PB-212 platform</p>	<p><b>\$1.35 B</b> ACP3 prostate program</p>	<p><b>\$2.1 B</b> Radiopharm platform</p>
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# Actinium is Well Positioned in the Current Radio Landscape

## A Transformative Pipeline Across Three High Value Pillars

Pillar	Program	Differentiation & Indication	Stage of Development			
			Preclinical	Phase 1	Phase 2	Phase 3
<b>Solid Tumors</b>  Growth & Value Driver	<b>ATNM-400 (Undisclosed Target)</b>	First-in-Class Ac-225 Program Targeting mCRPC, NSCLC & Breast Cancer	▶			
	<b>Actimab-A MDSC</b>	Combinations with PD-1 Inhibitors to Overcome Resistance in MDSC-Rich Solid Tumors	▶			
	<b>Undisclosed Targets/Theranostics</b>	Novel Solid Tumor Programs	▶			
<b>Hematology</b>  Value Now/ Partner Ready	<b>Actimab-A + CLAG-M</b>	Mutation Agnostic Backbone Therapy for Fit R/R AML	▶ Seeking collaborator			
	<b>Actimab-A Triplet Combo</b>	Mutation Agnostic Backbone Therapy for Frontline AML	▶			
	<b>Actimab-A Monotherapy</b>	Address Unmet Needs of High-risk HMA refractory MDS	▶			
	<b>Actimab-A Combinations (FLT3, IDH 1/2, Menin)</b>	Novel Combinations for Frontline, R/R & Maintenance – AML/MDS	▶			
<b>Conditioning</b>  Future of Cell & Gene Tx	<b>Iomab-ACT Commercial CAR-T</b>	Universal Conditioning to Improve Patient Access & Outcomes	▶			
	<b>Iomab-ACT BMT / GeneTx</b>	Targeted Non-Chemotherapy Conditioning to Unlock Curative Therapies	▶			
	<b>Iomab-B BMT</b>	Conditioning for Broad Active R/R AML Patient Population	▶ Seeking partner			

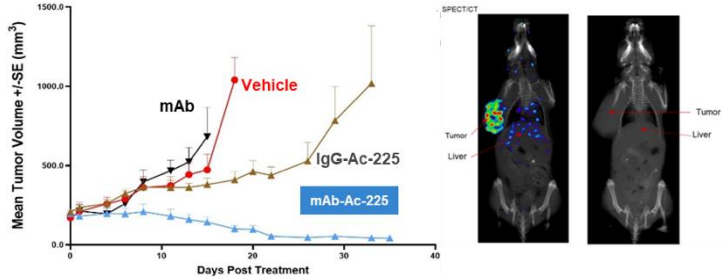
# Innovation Focused R&D & Manufacturing Capabilities To Drive Value

*Demonstrated ability to generate novel and highly differentiated radiotherapy programs as evidenced by our highly innovative pipeline*

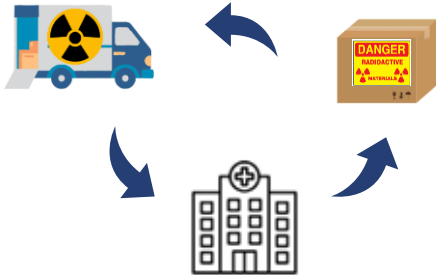
**Strong In-house R&D to be Complemented by RLT Manufacturing 2H:26**



**Demonstrated Leading-Edge Preclinical Radiochemistry & Translational Biology**

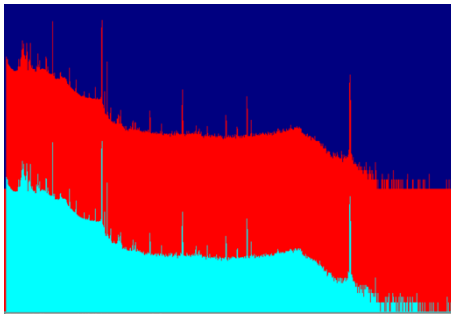
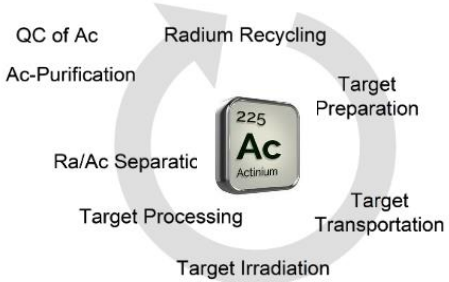


**End-to-end Supply Chain Established at ~50 leading Hospitals & Redundant Isotope Supply**



**Proprietary Cyclotron Manufacturing Technology Enables Commercial-Scale, Low-Cost AC-225 Production**

**Radiochemical Purity Equivalent to Gold-Standard Thorium Method & No Long-Lived Contaminants**





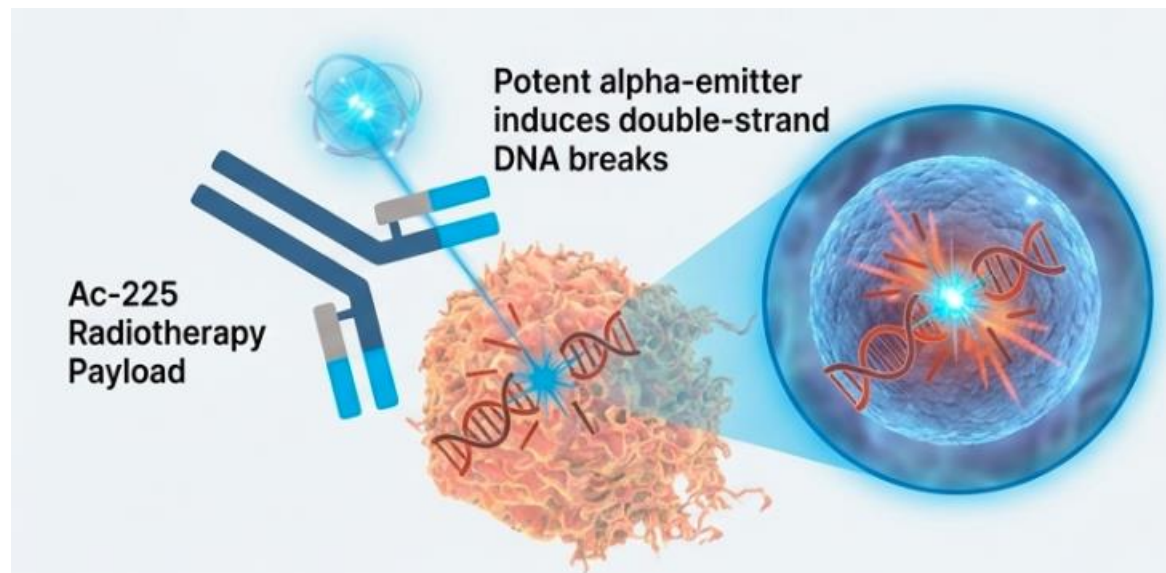
# Solid Tumors: ATNM-400 & Actimab-A

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**ATNM-400: First-in-class, multi-indication Ac-225 targeted radiotherapy supported by robust preclinical data in mCRPC, NSCLC & Breast Cancer**

**Actimab-A: Potential to synergize with PD-1 and other immune checkpoint inhibitors in solid tumors by depleting CD33+ MDSCs**

# ATNM-400: Differentiated Radiotherapy Targeting 3 Blockbuster Indications



- ✓ Target implicated in disease biology & treatment resistance
- ✓ Pan-tumor potential: target expressed across multiple solid tumor indications
- ✓ Theranostic approach with Zr-89 for imaging and Ac-225 for therapy



## Prostate Cancer (~300K U.S. patients)

- Non-PSMA target differentiates from Pluvicto, addresses non-responders/relapses and shows synergy with ARPIs for earlier line therapy



## Lung Cancer (~200K U.S. patients)

- Target is highly expressed in EGFR NSCLC and increases in resistant disease, outperformed 1<sup>st</sup>, 2<sup>nd</sup> and 3<sup>rd</sup> line EGFR therapies and synergistic with 1<sup>st</sup> line TAGRISSO

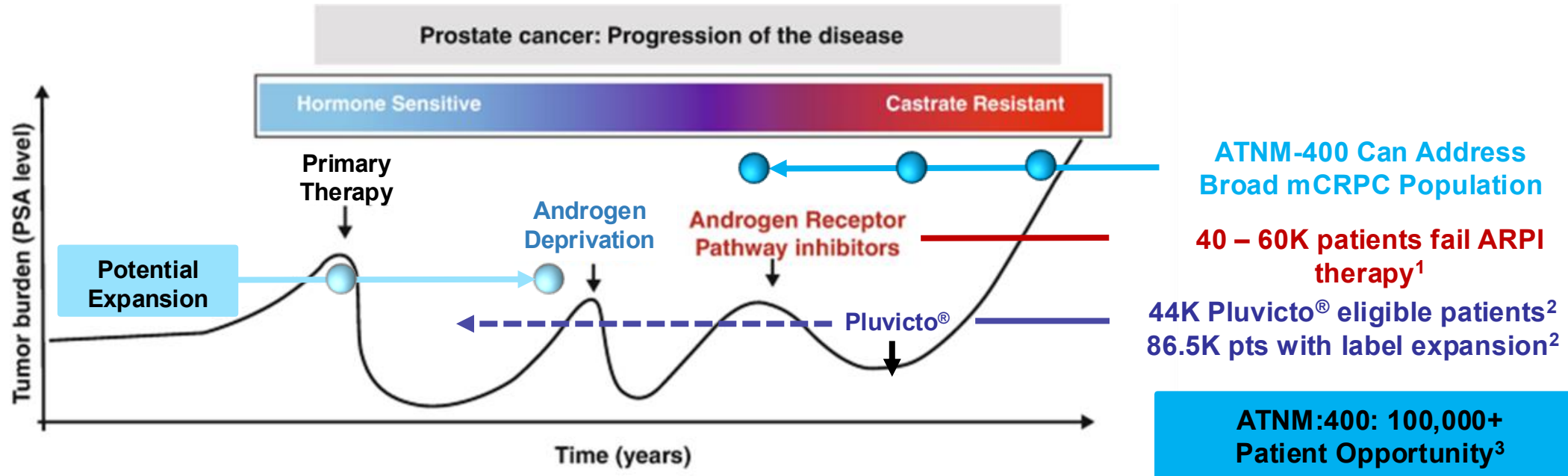


## Breast Cancer (~300K U.S. patients)

- Robust Efficacy in HR+, TNBC and HER2-resistant models with the potential to avoid toxicities like ILD seen with ADCs

# Blockbuster Market Opportunity for ATNM-400 in Prostate Cancer

*Multiple near-term opportunities in non-responders and progressors with future expansion to earlier lines of treatment*

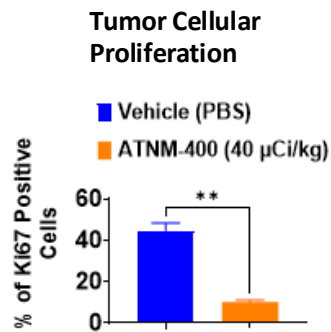
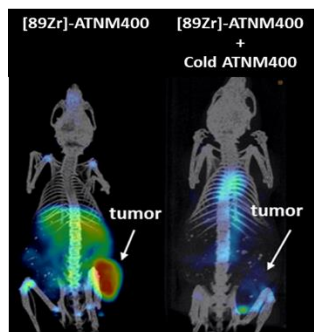


**ATNM-400 Addresses Blockbuster Segments of MCRPC : ARPIs: \$10+ billion | Pluvicto®: \$2.0 billion<sup>4</sup>**

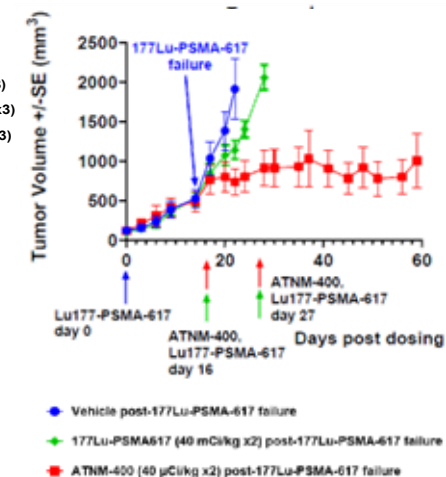
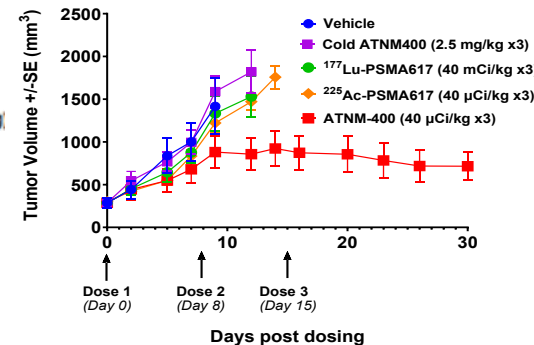
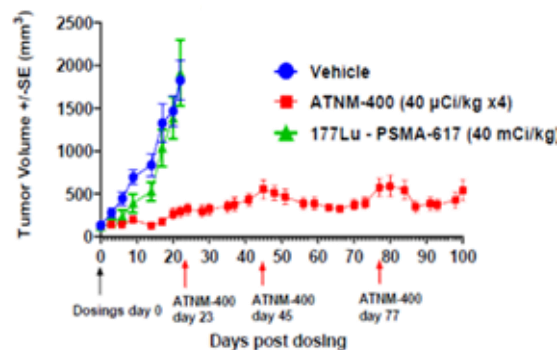
# Transformative Therapeutic Potential of ATNM-400 for Prostate Cancer

*Robust in vivo data show ATNM-400 achieves specific tumor uptake, decreased tumor cell proliferation, and superior efficacy versus 177Lu-PSMA-617, active agent in blockbuster radiotherapy Pluvicto® and next-generation 225Ac-PSMA-617*

## Specific Tumor Uptake & Decreased Cell Proliferation



## Greater Efficacy Than 177Lu-PSMA-617 and 225Ac-PSMA-617 in 22Rv1 PSMA-Low Prostate Cancer Model; Overcomes Resistance



Target Considerations	ATNM-400	PSMA
Implicated in prostate cancer cell survival	✓	✗
Linked to rapid disease progression	✓	✗
Drives shorter time to castration resistance	✓	✗
Expressed in multiple solid tumors	✓	✗

- 30% mCRPC patients have low or no PSMA expression
- Up to 70% patients do not respond
- Nearly all patients progress on Pluvicto® in <12-mos

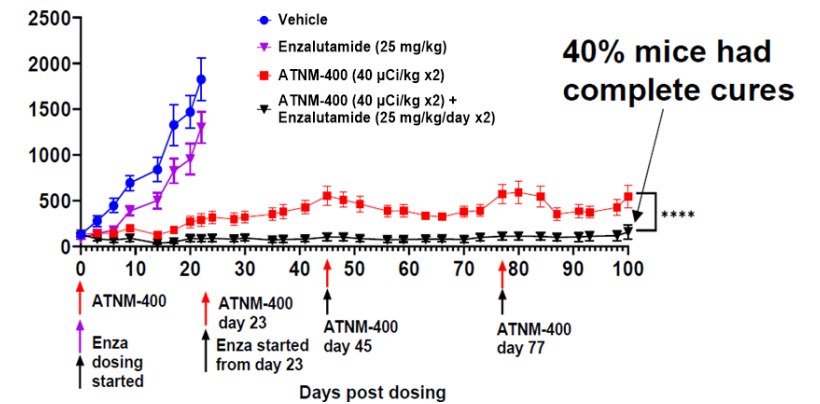
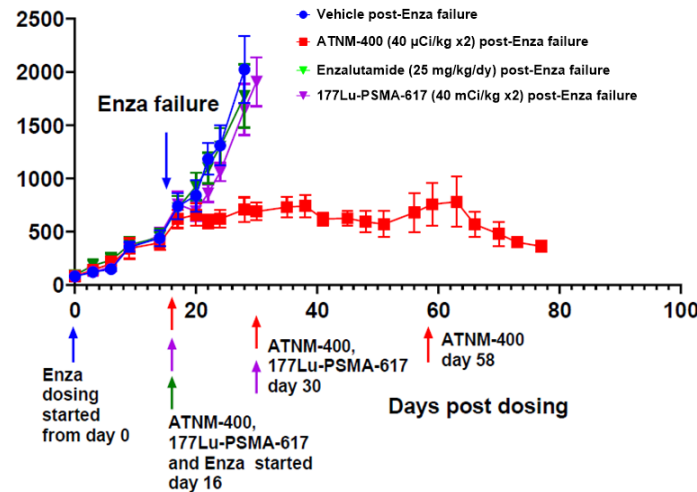
# Greater Efficacy and Synergy with ARPIs Supports Large Market

*Mechanistic rationale and data supports potential of ATNM-400 in earlier line mCRPC as a monotherapy or in combination*

## Superior to Enzalutamide and 177Lu-PSMA-617 in ARPI-Resistant Prostate Cancer and Strong Combination Activity with Durable Efficacy

### Mechanistic Synergy Supports Positive Results

- Enzalutamide resistance increases ATNM-400 target expression in prostate cancer models and mCRPC patients
- This provides a mechanistic rationale for the strong combination activity observed with ATNM-400 and ARPI enzalutamide

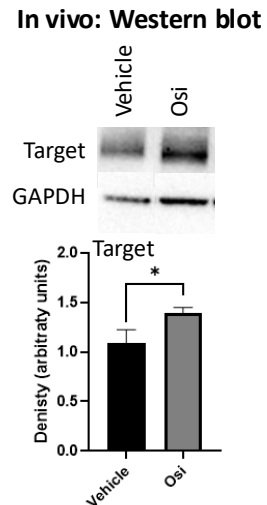
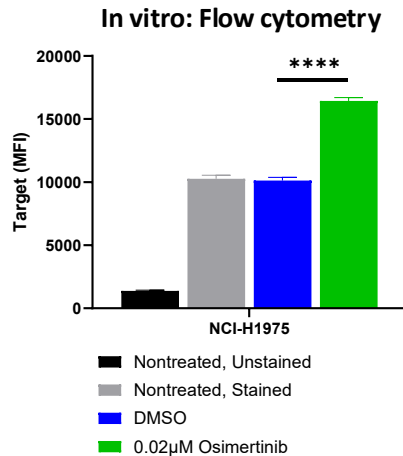


Up to 60,000 patients fail ARPI therapy and ~35% fail treatment in less than 12-months

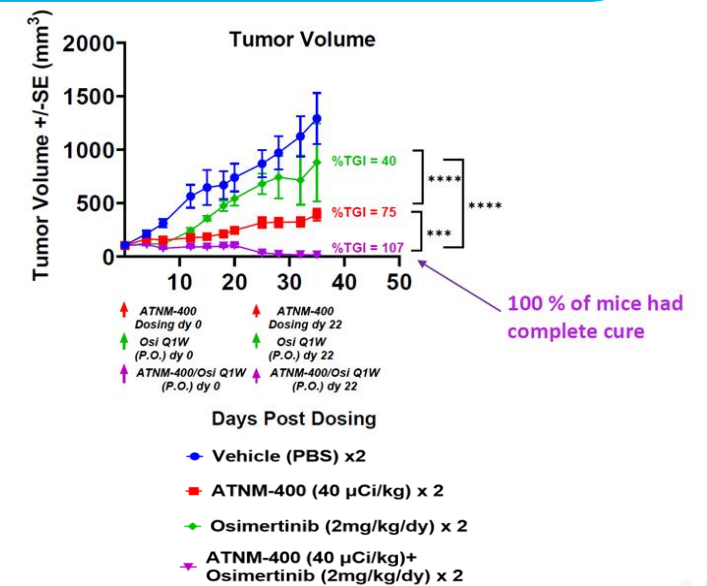
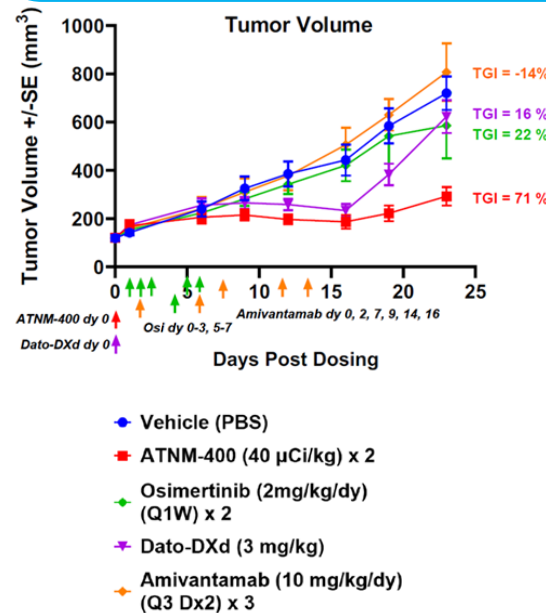
# Significant Market Expansion Opportunity Across NSCLC

ATNM-400 outperformed 1<sup>st</sup>, 2<sup>nd</sup> & 3<sup>rd</sup> line approved therapies for EGFR-mutant NSCLC by 3-5x and had robust synergy in combination with 1<sup>st</sup> line osimertinib that is supported by clinical data with EBRT

Osimertinib treatment increases ATNM-400 target expression both in vitro and in vivo



ATNM-400 had 3-5x greater tumor growth inhibition vs approved EGFR therapies and 100% cures with osimertinib combination



Clinical rationale: EBRT + osimertinib Improved PFS vs. osimertinib (32.2 vs 20 months)

Ac-225 is 4-8× more biologically lethal than EBRT and could drive superior efficacy, reduced toxicity, and access to earlier-line treatment segments

# Compelling Efficacy Across All EGFR-Mutant NSCLC Treatment Settings

**Results Strongly Support ATNM-400's Differentiated Profile in the Highly Competitive NSCLC Space and Potential Across 1<sup>st</sup>, 2<sup>nd</sup> & 3<sup>rd</sup> Line Settings, Alone or In Combination with Successful Therapies Competing for Market Share**

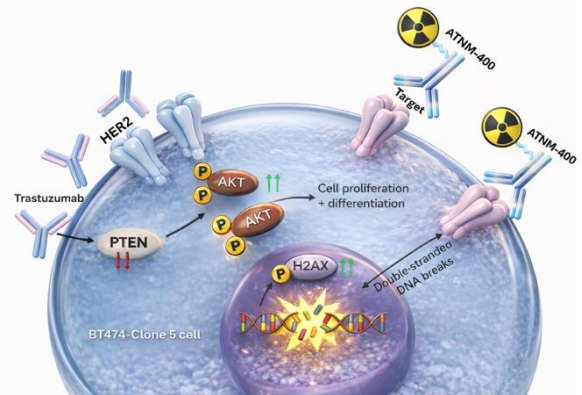
- ATNM-400's target antigen is overexpressed in NSCLC, associated with poor prognosis and linked to treatment resistance
- NSCLC accounts for ~85% of the 2+ million lung cancer cases globally
- AstraZeneca and Johnson & Johnson are competing for market share with TAGRISSO® and RYBREVANT® combinations
- ATNM-400 offers a novel EGFR + radiotherapy combination supported by preclinical and clinical data

	EGFR - 1 <sup>st</sup> Line	EGFR - 2 <sup>nd</sup> Line	EGFR - 3 <sup>rd</sup> Line
<b>ATNM-400<sup>1</sup> Efficacy</b>	✓ 3x Superior TGI ✓ Synergy in combination	✓ 5x Superior TGI	✓ 85% greater TGI
<b>Therapy &amp; Mechanism</b>	TAGRISSO® osimertinib EGFR-TKI	DATROWAY® Dato-DXd Trop-2 ADC	RYBREVANT® amivantamab EGFR-cMET Bispecific
<b>Company</b>	AstraZeneca (AZ)	Daiichi Sankyo/AZ	J&J
<b>Radiotherapy Presence</b>	Yes - Prostate Cancer	Yes - Prostate Cancer	Yes - Prostate Cancer

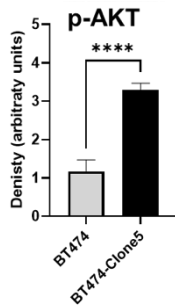
# Broad Potential Demonstrated Across Multiple Breast Cancer Settings

ATNM-400's target antigen is overexpressed in breast cancer including tumors resistant to endocrine therapies such as tamoxifen and HER2 targeted therapies like Herceptin® as well as in TNBC

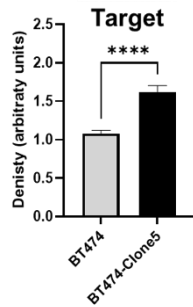
HER2 Therapy Resistance Increases Survival Pathway via p-AKT & ATNM-400 Target Expression



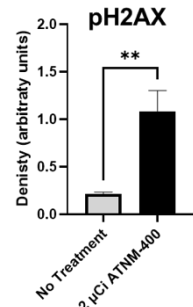
HER2 Resistance Marker



ATNM-400 Target Expression

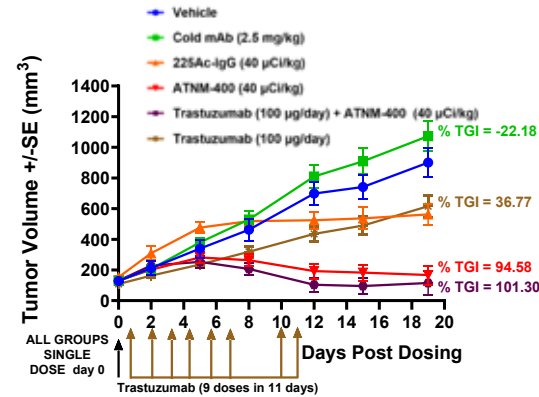


Double-Strand DNA Break Marker

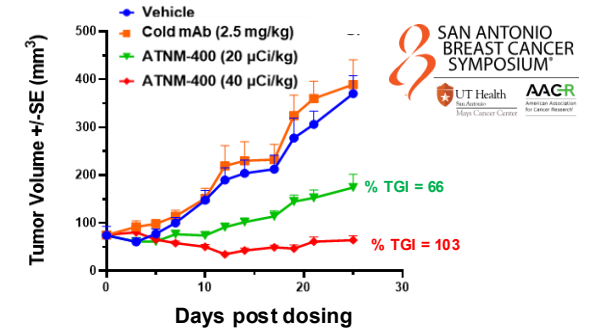


Monotherapy and Combinations Eradicates Trastuzumab-Resistant Tumors and Triple-Negative Breast Cancer (TNBC)

Trastuzumab-resistant model BT474-Clone5



TNBC model MDA-MB-468

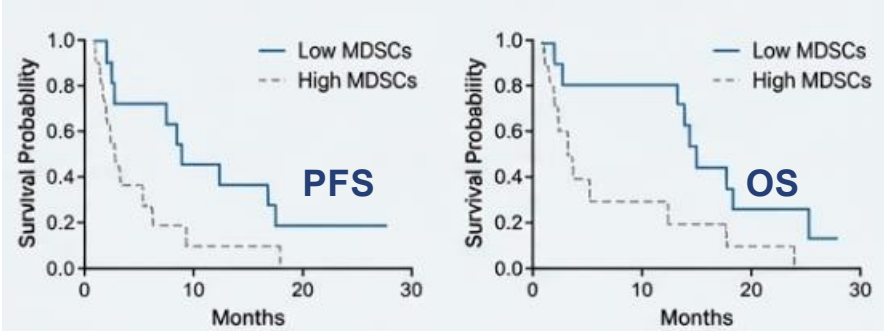


- Robust efficacy and tumor regression as monotherapy and in combination with trastuzumab (Herceptin®) in trastuzumab-resistant and TNBC models
- Herceptin® (Roche) and biosimilars generated sales of over \$5 billion in 2025<sup>1</sup>
- Off-target toxicities like ILD with 2L therapies (HER2 or Trop-2 ADCs) restricts use
- ATNM-400 represents a novel therapeutic approach to address the high unmet needs in HR+, HER2- and TNBC as a monotherapy or in combinations

# Actimab-A: Tap Into the Blockbuster PD-1 Market By Depleting MDSCs

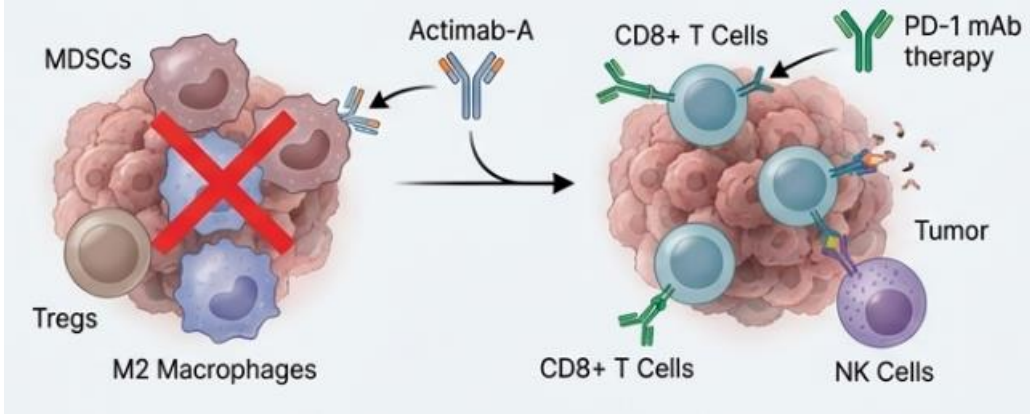
*Actimab-A can be synergistic with PD-1 inhibitors by depleting MDSCs, an immune cell subtype implicated in poor response and outcomes to PD-1 inhibitors, potentially opening and expanding a \$40+ billion market*

### Clinical Data Shows High MDSCs Lead to PD-1 Treatment Failure<sup>1</sup>



Outcomes	Low MDSCs	High MDSCs
PFS	8.39 months	1.94 months
OS	15.15 months	3.03 months

### CD33+ MDSCs are Primed for Depletion with Actimab-A = ↑ T-cell Proliferation



## The Opportunity

### Re-sensitize/Extend Responses

Enhance outcomes in the existing \$40 billion PD-1 approved market (Lung, melanoma, etc.) and potential for new IP

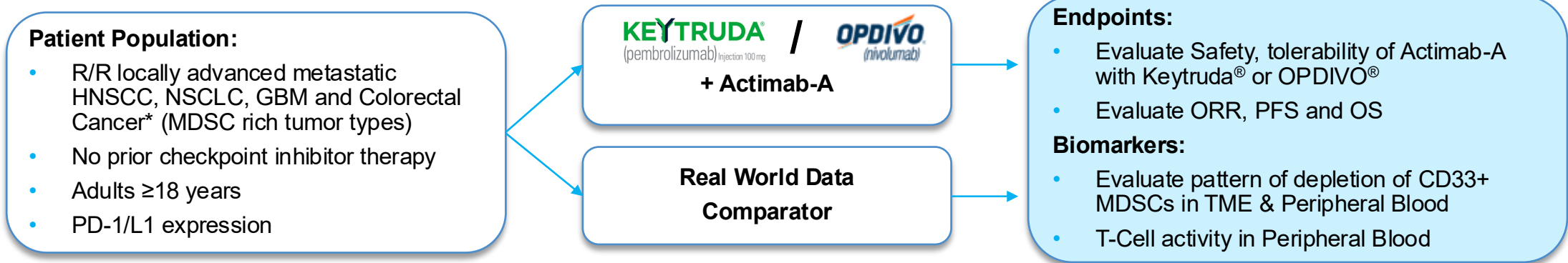
### Sensitize/Expand Indications

Open new markets for tumors with limited PD-1 response (Pancreatic, Prostate, Ovarian, etc.), representing 400,000+ patients annually

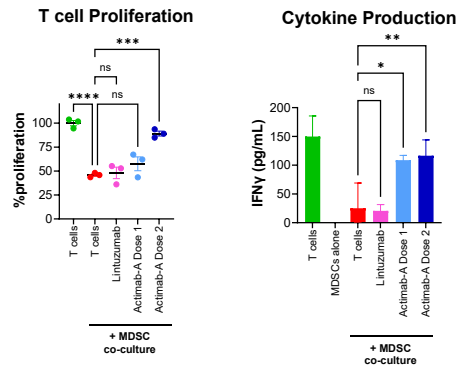
<sup>1</sup> Bronte et al. High Levels of Circulating Monocytic Myeloid-Derived Suppressive-Like Cells Are Associated With the Primary Resistance to Immune Checkpoint Inhibitors in Advanced Non-Small Cell Lung Cancer: An Exploratory Analysis <https://pmc.ncbi.nlm.nih.gov/articles/PMC9043492/>. Frontiers in Immunology. 2022 Apr 13;13:866561

# Actimab-A MDSC PD-1 Basket Trial: Data Expected 2H:2026

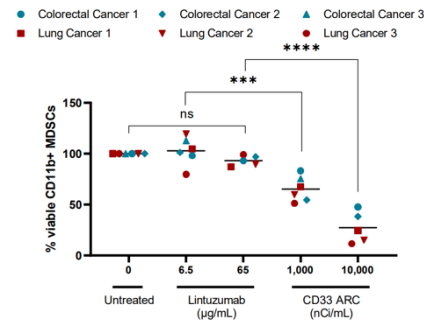
Trial supported by compelling preclinical data and mechanistic synergy will enroll patients across 4 tumor types



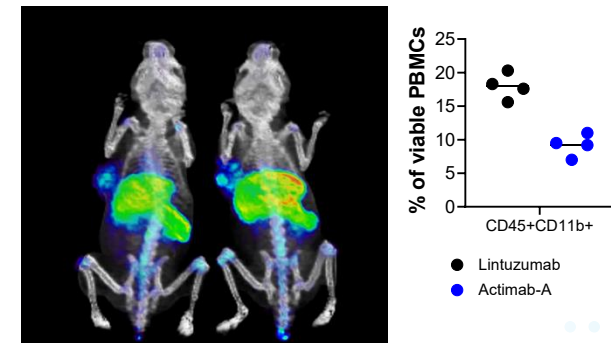
## Actimab-A targeting of MDSCs rescues T cell response *ex vivo*<sup>1</sup>



## Actimab-A is cytotoxic to patient-derived MDSCs *ex vivo*<sup>1</sup>



## Actimab-A homes to tumor-resident MDSCs and depletes MDSCs *in vivo*<sup>1</sup>





# Hematology Portfolio: AML Therapeutics & Targeted Conditioning

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**Actimab-A: Phase 2/3 ready, mutation agnostic Ac-225 AML therapy**

**lomab-B: Phase 2/3 ready BMT targeted conditioning agent**

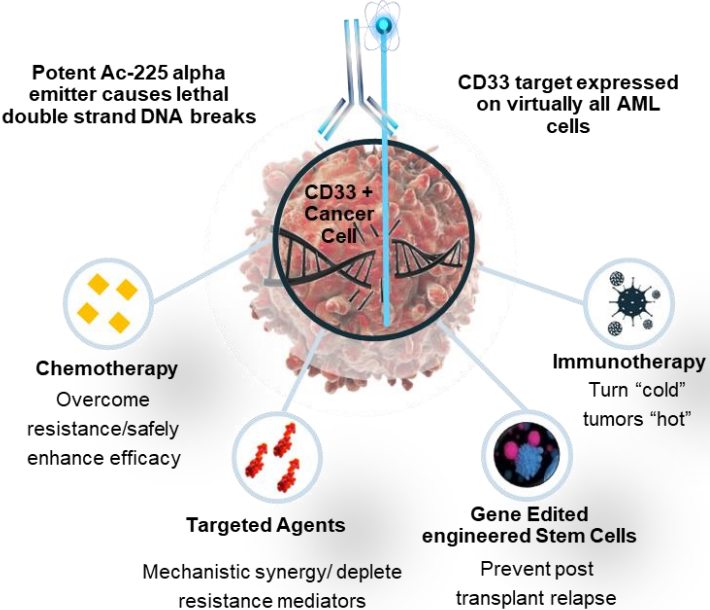
**lomab-ACT: Potential universal conditioning to improve access and outcomes for cell & gene therapies**

# A De-Risked, Late-Stage Hematology Franchise Ready for Partnership

Only CD33 and CD45 targeting clinical stage radiotherapies supported by strong data and Actimab-A NCI CRADA

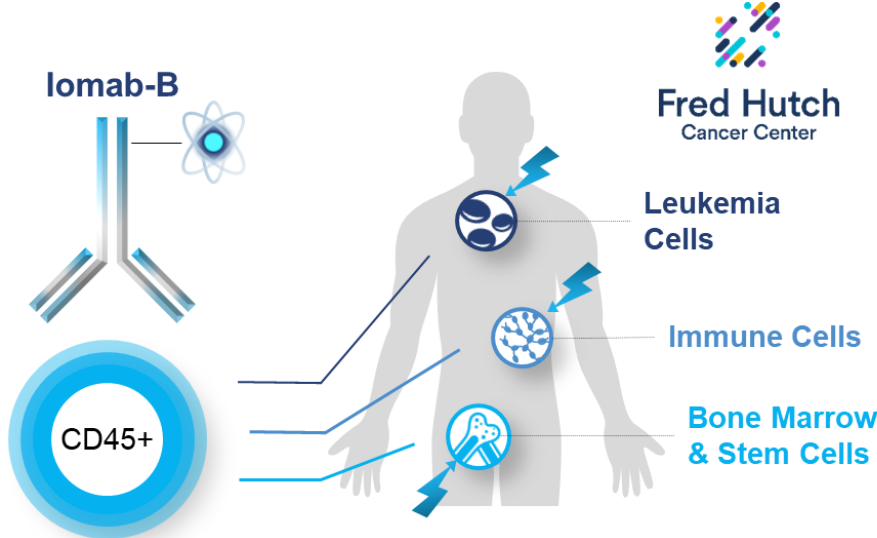
## Actimab-A: AML & MDS Therapeutics

- Mutation agnostic Ac-225 backbone therapy with strong data in high-risk patients
- Aligned with FDA on Phase 2/3 trial for Actimab-A + CLAG-M in R/R AML. **Actively seeking collaborator**
- NCI CRADA supports broad development in cost effective manner



## Iomab-B/ACT: Targeted Conditioning

- CD45 targeted radiotherapy designed to improve access and outcomes to BMT, Cell & Gene Therapies
- Aligned with FDA on Phase 2/3 trial in expanded R/R AML patient population. **Actively seeking partner**
- 3 active trials for Iomab-ACT for cell & gene therapy



# Potential to Transform the AML & MDS Treatment Journey

Late-stage differentiated radio assets have strong synergy with favorable commercial dynamics and can address the unmet needs of 110,000 patients in the U.S. and EU

Iomab-B BMT Conditioning: R/R AML	Phase 2/3
Actimab-A + CLAG-M: R/R AML	Phase 2/3
Actimab-A Triplet Combo: Frontline AML	Phase 1
Actimab-A Monotherapy: High-risk MDS	Phase 1
Actimab-A Combinations (FLT3, IDH 1/2 & Menin inhibitors)	Planned
Actimab-A: AML & MDS Maintenance	Planned

Indication	Therapy		BMT Conditioning	Post-BMT Maintenance
	Front Line	R/R		
AML: 88K pts	●	●	●	●
MDS: 23K pts	Potential Expansion	●	Potential Expansion	Potential Expansion

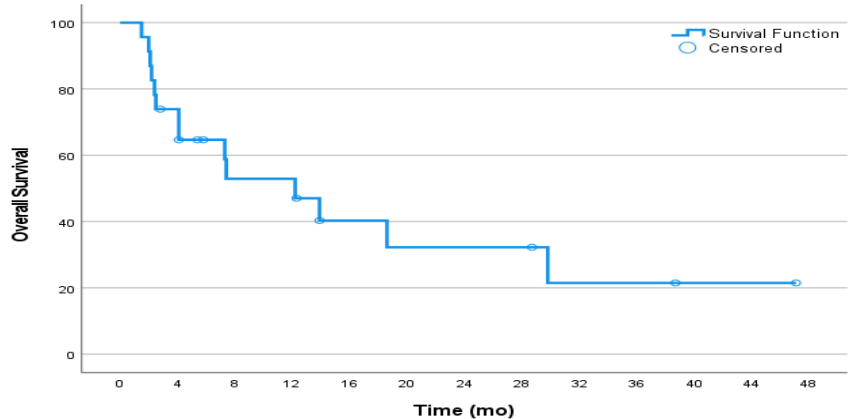
## Key Value Drivers

- ✓ **Clinical Validation:** Over 500 patients treated with Iomab-B
- ✓ **Concentrated Commercial Market:** Patient population largely treated in top 100 quaternary care centers in U.S. & EU
- ✓ **Favorable Commercial Dynamics:** Strong synergy across Actimab-A, Iomab-B and Iomab-ACT
- ✓ **Unmet Medical Need:** Provides opportunity for commercial success with the only radiotherapies in development for these blockbuster markets



# Actimab-A + CLAG-M: Results Support Phase 2/3 Trial in R/R AML

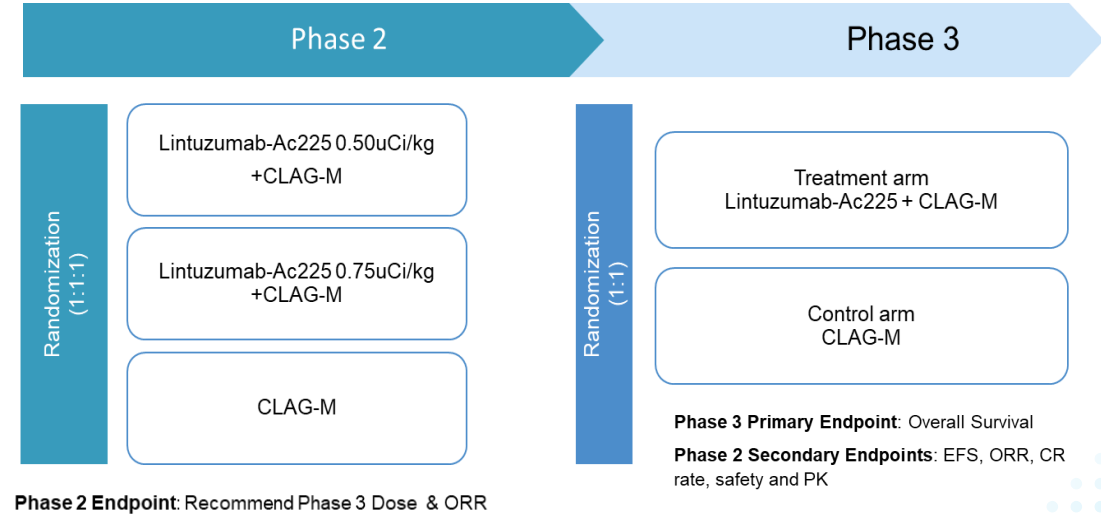
**High Rates of MRD Negativity and Improved Survival Outcomes in High-Risk R/R Patients<sup>1</sup>**



Patient segment	MRD Negativity	Median Overall Survival
1 <sup>st</sup> /2 <sup>nd</sup> Salvage (n=14)	80%	18.4 mo.
TP53mut (n=13)	83%	9.6 mo.
Prior Venetoclax (n=13)	100%	7.3 mo.

**Phase 2/3 Trial to Enroll Patients with 1<sup>st</sup> or 2<sup>nd</sup> Salvage R/R AML, Group Had Best Outcomes in Phase 1b Trial**

- 24-month median overall survival in the 70% of patients who proceeded to a BMT with Actimab-A + CLAG-M
- Results compare highly favorably to <2 mo. – 4 mo. overall survival in TP53+ or prior Venetoclax treated patients<sup>2,3</sup>



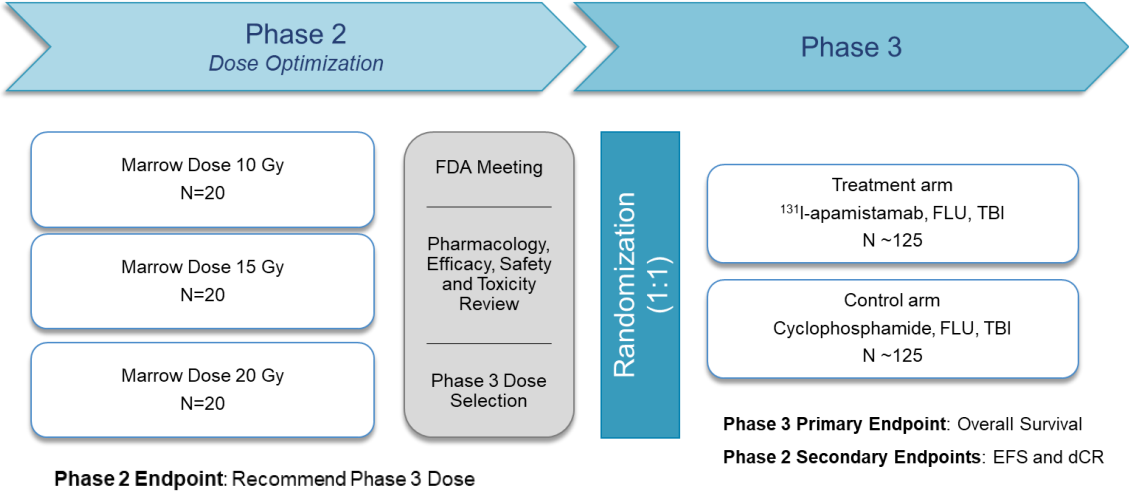
**Aligned with FDA on Phase 2/3 trial, Actively seeking collaborator**



1) Abedin et al. Phase 1 study of lintuzumab-Ac225 combined with CLAG-M salvage therapy in relapsed/refractory acute myeloid leukemia. Leukemia February 2025. 2) Maiti et al. Outcomes of relapsed or refractory acute myeloid leukemia after front-line hypomethylating agent and venetoclax regimens 3) Zucenka, A., et al. Outcomes of relapsed or refractory acute myeloid leukemia patients failing venetoclax-based salvage therapies. Eur J Haematol. 2020; 106: 105– 1133)

# Iomab-B: Phase 2/3 Trial Ready for Initiation and Partnering

**FDA Aligned on Trial in Expanded Patient Population to Include All R/R AML Patients age 18+, Potential Future Market Expansion Supported by Data in 5 Additional Disease Indications**



- Long-lived IP:**
  - Composition of matter patents extend to 2038
- Strong Existing Use Base:**
  - 24 SIERRA sites continue to have strong interest in Iomab-B
- Regulatory Clarity:**
  - Potential to leverage Phase 2 results with SIERRA data
- Market Opportunity:**
  - ~ 150,000 addressable patients across 6 indications via improved BMT conditioning

**Robust and compelling clinical data supports near-term opportunity in r/r AML and indication expansion**

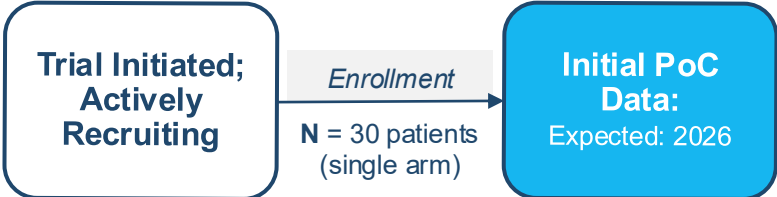
- 400+ patients
- 12 clinical trials
- 6 diseases
- Improved survival and curative outcomes

**Strong market awareness driven by 12 oral presentations at the most prestigious BMT, CAR-T hematology and nuclear medicine conferences**

**SIERRA trial supports Phase 2/3 in expanded population; Actively seeking partner**

# Iomab-ACT Further Expands Heme Portfolio Market Opportunity

## Commercial CAR-T Phase 1B/2 Trial



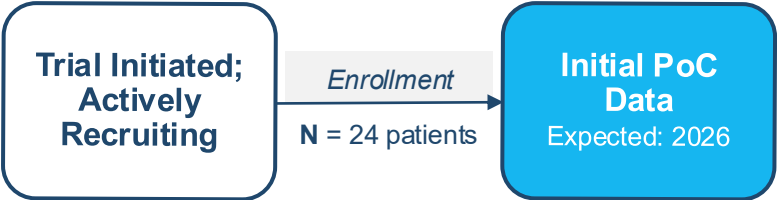
**PRIMARY OBJECTIVES**

Safety, tolerability, and efficacy of Iomab-ACT conditioning with CAR-T

**SECONDARY OBJECTIVES**

Incidences of CRS, ICANS  
In vivo persistence and expansion of CAR-T cells

## BMT & Gene Therapy Conditioning – Sick Cell Disease BMT Phase 1 Trial



**PRIMARY & SECONDARY OBJECTIVES**

Safety assessment by evaluating graft failure, grades III-IV GvHD, or death at 100 days

Advance to gene therapy conditioning



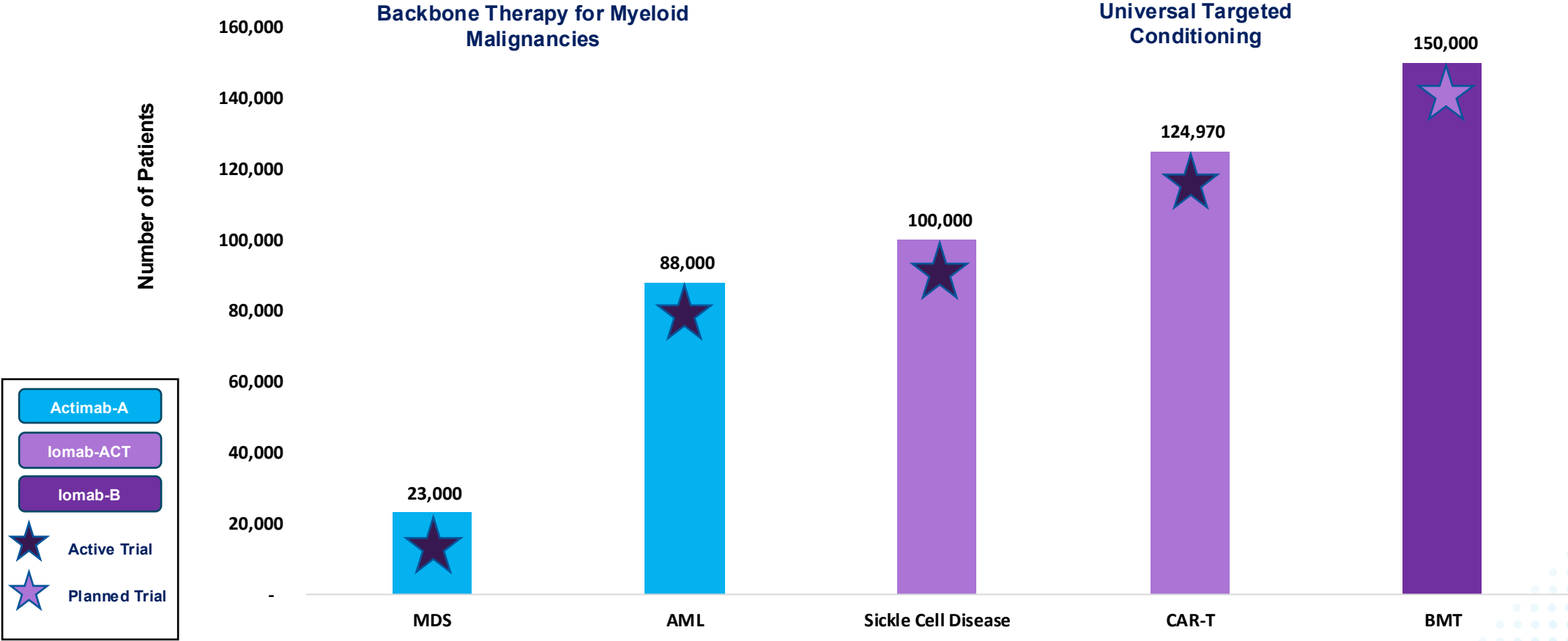
**UTSouthwestern**  
Medical Center  
**Farrukh Awan, MD**  
Professor of Internal Medicine



**COLUMBIA** | COLUMBIA UNIVERSITY  
HERBERT IRVING COMPREHENSIVE  
CANCER CENTER  
**Markus Mapara, MD**  
Professor of Medicine

**Early POC Data Expected 2H:2026 That Can Inform Registration Trials**

# Blockbuster Opportunities for Three First-in-Class Heme Radiotherapies



Indicates number of patients diagnosed with (incidence) or living with the disease (prevalence). Sources: 1. SEER 2. CancerMPact AML Treatment Architecture US 2022 3. CancerMPact AML Treatment Architecture EU5 2022 4. O. Visser et al, Incidence, survival and prevalence of myeloid malignancies in Europe, 2012; RARECARE 5. Competitor data 6. \*High Risk MDS patient population includes select Intermediate Risk patients 7. Leukemia & Lymphoma society Facts & Statistics 8. CDC Sickle Cell Disease Data & Statistics 9. American Cancer Society 10. CIBMTR 2025 Summary Data, leukemia & Lymphoma Society Statistics








# 2026 Outlook & Milestones

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**Transformational year with multiple first-in-class therapies reaching inflection points supporting our vision to be a leading fully-integrated targeted radiotherapy company**



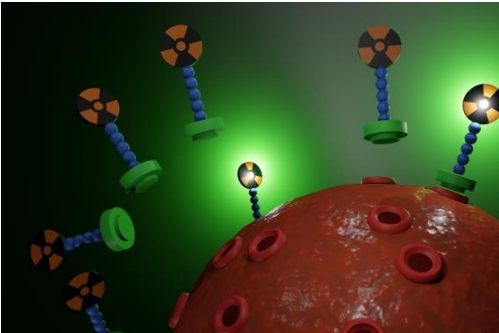
# 2026: A Year of Transformational Milestones and Data Readouts

 <p><b>ATNM-400</b></p>	<ul style="list-style-type: none"> <li>● Mid:26</li> <li>● FY:26</li> </ul>	<ul style="list-style-type: none"> <li>Initial tumor targeting and biodistribution data</li> <li>Present additional preclinical data in mCRPC, NSCLC and Breast Cancer</li> </ul>
 <p><b>Actimab-A (MDSCs)</b></p>	<ul style="list-style-type: none"> <li>● 2H:26</li> <li>● 2H:26</li> </ul>	<ul style="list-style-type: none"> <li>Initial Phase 1 data from MDSC basket trial</li> <li>Multi-tumor Phase 1 data from MDSC basket trial</li> </ul>
 <p><b>Hematology Franchise</b></p>	<ul style="list-style-type: none"> <li>● 2H:26</li> <li>● FY:26</li> <li>● 2H:26</li> </ul>	<ul style="list-style-type: none"> <li>Secure partner, Initiate Actimab-A+CLAG-M Phase 2/3 trial in r/r AML</li> <li>Advance Phase 1 frontline AML trial with Actimab-A triplet</li> <li>Present mutation agnostic data from MSKCC collaboration</li> </ul>
 <p><b>Iomab-B/ACT (Conditioning)</b></p>	<ul style="list-style-type: none"> <li>● 2H:26</li> <li>● 2H:26</li> <li>● 2H:26</li> <li>● FY:26</li> </ul>	<ul style="list-style-type: none"> <li>Initial safety and engraftment data from SCD and commercial CAR-T trials</li> <li>Advance to SCD gene therapy conditioning, explore collaborations</li> <li>Additional clinical data from Iomab-ACT SCD and commercial CAR-T trials</li> <li>Secure partner for Iomab-B Phase 2/3 Trial</li> </ul>
 <p><b>RLT Infrastructure &amp; Supply Chain</b></p>	<ul style="list-style-type: none"> <li>● 2H:26</li> <li>● 2H:26</li> <li>● FY:26</li> </ul>	<ul style="list-style-type: none"> <li>Complete facility buildout</li> <li>Supply first GMP clinical batch</li> <li>Secure partnership to produce lower-cost Ac-225 leveraging proprietary cyclotron technology</li> </ul>

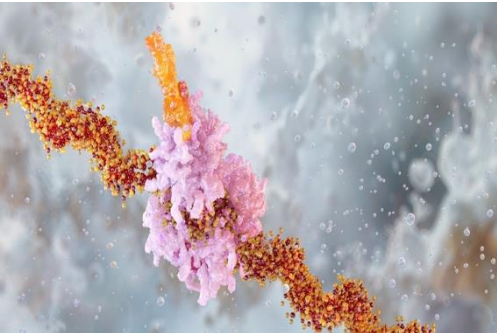
# Actinium is Being Purpose Built to Fill Pharma's Strategic Gaps

*Actinium is leveraging its extensive radiotherapy experience and expanding capabilities to develop novel therapies for high-value oncology indications*

**Radiochemistry Expertise and Know-how**



**Strong Translational Biology Capabilities**



**RLT Manufacturing, Ac-225 Production Technology & Supply Chain Capabilities\***



**Clinical Development & Trial Execution**



***Vision: Be a Leading Fully Integrated Radiopharmaceutical Company Developing Highly Differentiated, First-in-Class Targeted Radiotherapies***



# Thank You

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**Actinium Pharmaceuticals, Inc.**

**ATNM: NYSE AMERICAN**