

Actinium Pharmaceuticals Presents New Preclinical Data Demonstrating Potent Anti-Tumor Activity of ATNM-400 Across Multiple Breast Cancer Subtypes Including Hormone Receptor-Positive, Triple-Negative, and Tamoxifen- and HER2 Therapy-Resistant Breast Cancer Models at SABCS 2025

- ATNM-400's target antigen is overexpressed in breast cancer and its expression is further increased in breast cancer cells resistant to standard of care endocrine therapy, tamoxifen and the HER2 therapy trastuzumab
- The Actinium-225 alpha-emitter payload of ATNM-400 induced irreversible doublestrand DNA breaks and has the potential to produce potent localized tumor killing with reduced off-target lung toxicity that limits the use of antibody drug conjugates
- Data supports the continued development of ATNM-400 as a monotherapy or combination therapy to address multiple breast cancer subtypes with limited treatment options

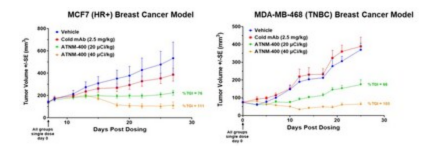
NEW YORK, Dec. 12, 2025 /PRNewswire/ -- Actinium Pharmaceuticals, Inc. (NYSE American: ATNM), a leader in the development of differentiated targeted radiotherapies, today announced the presentation of new preclinical data for ATNM-400, its first-in-class Actinium-225 (Ac-225) based antibody radioconjugate, at the 2025 San Antonio Breast Cancer Symposium (SABCS). The data highlight robust anti-tumor activity and a favorable tolerability profile across a broad panel of preclinical breast cancer models, including hormone receptor-positive (HR+), triple-negative breast cancer (TNBC), and tamoxifen- and trastuzumab-resistant tumors. Breast cancer is the most diagnosed cancer among women in the United States with approximately 316,950 women expected to be diagnosed with the disease in 2025 according to the National Cancer institute. It is estimated that approximately 200,000 women are living with metastatic breast cancer in 2025, which is expected to grow to 250,000 in 2030. Of those diagnosed, hormone receptor-positive, HER2-negative (HR+/HER2-) breast cancer accounts for 70-75% of breast cancer, representing the largest subtype. ATNM-400 targets a validated cancer antigen that is overexpressed in breast cancer as well as multiple other solid tumor indications including prostate cancer and non-small cell lung cancer (NSCLC). Actinium is developing ATNM-400

as a potential pan-tumor targeted radiotherapy.

Highlights from the SABCS 2025 Poster Presentation

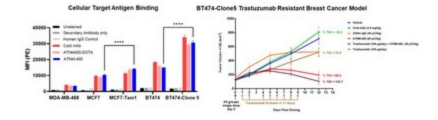
The poster, titled "Anti-Tumor Activity of ATNM-400, a First-in-Class Actinium-225 Antibody Radioconjugate, in Hormone-Positive, Triple-Negative, Tamoxifen-Resistant and Trastuzumab-Resistant Breast Cancer Models," showcases the following key findings:

Potent Efficacy Across Breast Cancer Subtypes: ATNM-400 demonstrated significant tumor-growth inhibition (TGI) in HR+ (MCF7) and TNBC (MDA-MB-468) in vivo models, with all treatment regimens well tolerated and no significant changes in body weight observed.



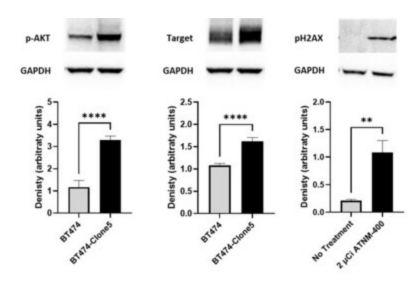
Potent Activity in Standard-of-care (SOC) Treatment-Resistant Breast Cancer Models

- Trastuzumab-resistant BT474-Clone5 breast cancer cells or Tamoxifen-resistant MCF7-Tam1 breast cancer cells exhibited increased target expression, resulting in enhanced in vitro cytotoxicity with ATNM-400.
- Combining ATNM-400 with either trastuzumab or tamoxifen resulted in greater cytotoxicity versus monotherapy and produced in vivo tumor regression in the trastuzumab-resistant model.



Mechanistic Evidence of Irreversible DNA Damage

- Activation of phosphorylation of AKT (pAKT) was observed in trastuzumab resistant BT474-Clone5 breast cancer cells, as well as a significant increase in the total level of the ATNM-400 target antigen in the in vivo trastuzumab-resistant breast cancer model.
- ATNM-400 treatment of these trastuzumab-resistant breast cancer cells caused significant increase in phosphorylated H2AZ (pH2AX), consistent with alpha-particle driven double-strand DNA damage.



Favorable Biodistribution: Sustained tumor uptake in a breast cancer model through 144 hours and rapid clearance from normal organs supports a potentially differentiated safety profile.

Pan-Tumor Potential: These results, together with previously published ATNM-400 data in prostate and lung cancer, reinforce the program's broad applicability across solid tumors.

Dr. Aditya Bardia, Professor of Medicine and Program Director, Breast Medical Oncology at the Geffen School of Medicine, University of California Los Angeles, was an author on the ATNM-400 SABCS poster.

"Patients whose tumors progress after endocrine therapy or HER2-targeted therapies face limited and often toxic treatment options. The data presented at SABCS show that ATNM-400 generates potent, targeted DNA damage even in highly resistant breast cancer models, while maintaining a favorable tolerability profile. This therapeutic approach has the potential to address some of the most pressing unmet needs in breast cancer care, particularly for those who have exhausted established modalities", said Dr. Bardia.

Breast cancer remains a heterogeneous disease with significant unmet need, particularly for patients who relapse following endocrine or HER2-directed therapies—an issue impacting 20–30% of HR+ patients and many with HER2-positive disease. In this setting, tamoxifen and trastuzumab (Herceptin[®], Roche and biosimilars) generated sales of approximately \$4.0 billion in 2024. ATNM-400 also demonstrated potency in triple negative breast cancer (TNBC), which accounts for up to 15% of all breast cancer cases and is associated with poor patient outcomes. ATNM-400 was designed to deliver highly potent Actinium-225 alphaparticle radiation, enabling precise tumor killing with limited off-target toxicity due to the ultrashort path length of alpha emissions. This has the potential to address the limitations of antibody drug conjugates, which have shown to cause interstitial lung disease (ILD), a toxicity that restricts their use.

Sandesh Seth, Chairman and CEO of Actinium Pharmaceuticals, added, "ATNM-400 represents an important component of Actinium's strategy to develop next-generation radiotherapeutics that can meaningfully expand treatment options for patients with hard-to-treat solid tumors. The breadth of activity seen in both endocrine-resistant and HER2-resistant breast cancer reinforces ATNM-400's potential to become a differentiated, first-in-

class therapy. Beyond breast cancer, we have presented compelling ATNM-400 data in prostate cancer and lung cancer that demonstrate its pan-tumor potential. These data further support our excitement for ATNM-400 and our plans to accelerate development of ATNM-400 as we advance toward clinical readiness."

The ATNM-400 SABCS poster presentation can be access on Actinium's investor relations website <u>HERE</u>.

About ATNM-400

ATNM-400 is a highly innovative, first-in-class, and multi-indication Actinium-225 (Ac-225) targeted radiotherapy candidate in development for prostate cancer, non-small cell lung cancer (NSCLC) and breast cancer. ATNM-400 is highly differentiated in prostate cancer as it targets a distinct non-PSMA protein strongly implicated in prostate cancer disease biology including progression and treatment resistance. Unlike 177Lu-PSMA-617, the active agent in Pluvicto[®] and the majority of radiotherapies under development, which rely on PSMA targeting, ATNM-400 is designed to maintain efficacy in low-PSMA or high-PSMA resistant disease, a major unmet clinical need as up to 30% of patients do not respond to PSMA radioligand therapies and up to 60% of patients have at least one PSMA-negative tumor lesion. Ac-225 delivers high-linear-energy-transfer alpha particles that induce irreparable double-strand DNA breaks, offering superior potency over beta emitters like Lutetium-177 (177Lu), and has a shorter tissue path length that may reduce off-target toxicity. The receptor specifically targeted by ATNM-400 continues to be expressed at a high level even after androgen receptor inhibitor (ARPI) and ATNM-400 has shown to overcome resistance to the ARPI therapy enzalutamide and work synergistically in combination with enhanced tumor control including complete tumor regression. In NSCLC, ATNM-400 has shown superior efficacy compared to approved first, second- and third-line EGFR therapies including small molecules, antibody drug conjugates and bispecific antibodies that is synergistic with osimertinib, an EGFR tyrosine kinase inhibitor (TKI) that is a standard of care therapy approved for treatment of patients in the frontline setting and is also able to overcome osimertinib resistance. ATNM-400 has also shown potent anti-tumor activity across multiple breast cancer subtypes including hormone receptor-positive, triple-negative, and tamoxifen- and HER2 therapy-resistant breast cancer models.

Prostate cancer is the most commonly diagnosed cancer in men, with ~1.5 million new cases globally and over 313,000 expected in the U.S. in 2025. While early-stage disease is typically managed with surgery, radiation, and ARPI therapy, up to 20% of cases progress to mCRPC - a lethal stage with limited treatment options. Targeted radiotherapy is a growing field in prostate cancer, dominated by PSMA-targeting agents like Pluvicto[®], which had sales of over \$1.3 billion in 2024, yet up to 30% of patients either lack or have no PSMA expression and virtually all patients develop resistance to Pluvicto[®] within 1-year. In the U.S., 40,000–60,000 mCRPC patients annually progress after ARPI therapy with approximately 35% of patients progressing within 1-year. As a class, ARPI therapies had sales of over \$10.0 billion in 2024 including enzalutamide (Xtandi[®]) that led the class with sales of over \$5.9 billion in 2024, highlighting a significant unmet need. Lung cancer is the leading cause of cancer deaths and there are there are over 200,000 new cases expected in the U.S. in 2025 and over 2 million cases globally. NSCLC accounts for approximately 85% of all lung cancer cases. EGFR targeting therapies including front-line osimertinib (TAGRISSO[®], AstraZeneca) an EGFR tyrosine kinase inhibitor (TKI), second-line Dato-DXd

(Datroway[®], AstraZeneca/Daiichi Sankyo) a Trop-2 ADC, and third-line amivantamab (Rybrevant[®], Johnson & Johnson) an EGFR-cMET bispecific had sales of approximately \$7 billion in 2024 with the EGFR TKI Osimertinib (TAGRISSO[®], AstraZeneca) generating sales of \$6.6 billion in 2024. Breast cancer is the most diagnosed cancer among woman in the United States with approximately 316,950 women expected to be diagnosed with the disease in 2025 according to the National Cancer institute. It is estimated that approximately 200,000 women are living with metastatic breast cancer in 2025, which is expected to grow to 250,000 in 2030. Of those diagnosed, hormone receptor-positive, HER2-negative (HR+/HER2-) breast cancer accounts for 70-75% of breast cancer, representing the largest subtype. In this setting, tamoxifen and trastuzumab (Herceptin[®], Roche and biosimilars) generated sales of approximately \$4.0 billion in 2024. ATNM-400 also demonstrated potency in triple negative breast cancer (TNBC), which accounts for up to 15% of all breast cancer cases is associated with poor patient outcomes. Across prostate cancer, NSCLC and breast cancer, ATNM-400 has demonstrated treatment paradigm changing potential in these indications, which have over 800,000 new cases in the U.S. alone.

About Actinium Pharmaceuticals, Inc.

Actinium is a pioneer in the development of targeted radiotherapies intended to meaningfully improve patient outcomes. ATNM-400, Actinium's lead product candidate, is a novel, first-inclass, and multi-indication Actinium-225 (Ac-225) in development for prostate cancer, nonsmall cell lung cancer (NSCLC) and breast cancer. The antigen specifically targeted by ATNM-400 is highly expressed in metastatic castration-resistant prostate cancer (mCRPC), contributes directly to disease progression, poorer survival outcomes, and continues to be expressed at a high level even after androgen receptor inhibitor (ARPI) and Pluvicto® treatment. ATNM-400 is supported by preclinical data demonstrating tumorspecific uptake, higher efficacy than androgen receptor inhibitor enzalutamide (Xtandi®) and 177Lu-PSMA-617 radiotherapy, the active agent in Pluvicto®, durable tumor control and potent efficacy in prostate cancer models resistant to both enzalutamide and 177Lu-PSMA-617. In addition, ATNM-400 has demonstrated synergy with enzalutamide. In NSCLC, ATNM-400 showed superior efficacy to EGFR targeting therapies including osimertinib (TARGRISSO®, AstraZeneca), Dato-DXd (DATROWAY®, AstraZeneca/Daiichi Sankyo) and amivantamab (RYBREVANT®, J&J) with synergistic activity in combination with osimertinib. In breast cancer, ATNM-400 showed potent anti-tumor activity across multiple breast cancer subtypes including hormone receptor-positive, triple-negative, and tamoxifen- and HER2 therapy-resistant breast cancer models. The data generated to date with ATNM-400 supports its potential across treatment settings to be used either as a monotherapy, or in combination or sequenced with other therapies. Actinium's most advanced product candidate in development is Actimab-A, a CD33 targeting therapeutic, that is a potential backbone therapy for acute myeloid leukemia (AML) and other myeloid malignancies leveraging the mutation agnostic alpha-emitter radioisotope payload Actinium-225 (Ac-225). Actimab-A has demonstrated potential activity in relapsed and refractory acute myeloid leukemia (r/r AML) patients in combination with the chemotherapy CLAG-M including high rates of Complete Remissions (CR) and measurable residual disease (MRD) negativity leading to improved survival outcomes and is being advanced to a pivotal Phase 2/3 trial. In addition, Actinium is engaged with the National Cancer Institute (NCI) under a Cooperative Research and Development Agreement (CRADA) for development of Actimab-A in AML and other myeloid malignancies. The first clinical trial under the CRADA will evaluate the triplet

combination comprised of Actimab-A, Venetoclax (Abbvie/Roche) an oral Bcl-2 inhibitor and ASTX-727 (Taiho Oncology, an Otsuka holdings company) a novel oral hypomethylating agent (HMA) in frontline acute myeloid leukemia (AML) patients. Additionally, Actinium is developing Actimab-A as a potential pan tumor therapy in combination with PD-1 checkpoint inhibitors including KEYTRUDA® and OPDIVO® by depleting myeloid derived suppressor cells (MDSCs), which represents a potential multi-billion-dollar addressable market. Iomab-ACT, Actinium's next generation conditioning candidate, is being developed with the goal of improving patient access and outcomes for potentially curative cell and gene therapies. Iomab-B is an induction and conditioning agent prior to bone marrow transplant in patients with r/r AML, which Actinium is seeking a potential strategic partner for the U.S. In addition, the company's R&D efforts are primarily focused on advancing several preclinical programs for solid tumor indications. Actinium holds approximately 250 patents and patent applications including several patents related to the manufacture of the isotope Ac-225 in a cyclotron.

For more information, please visit: https://www.actiniumpharma.com/

Forward-Looking Statements

This press release may contain projections or other "forward-looking statements" within the meaning of the "safe-harbor" provisions of the private securities litigation reform act of 1995 regarding future events or the future financial performance of the Company which the Company undertakes no obligation to update. These statements are based on management's current expectations and are subject to risks and uncertainties that may cause actual results to differ materially from the anticipated or estimated future results, including the risks and uncertainties associated with preliminary study results varying from final results, estimates of potential markets for drugs under development, clinical trials, actions by the FDA and other governmental agencies, regulatory clearances, responses to regulatory matters, the market demand for and acceptance of Actinium's products and services, performance of clinical research organizations and other risks detailed from time to time in Actinium's filings with the Securities and Exchange Commission (the "SEC"), including without limitation its most recent annual report on form 10-K, subsequent quarterly reports on Forms 10-Q and Forms 8-K, each as amended and supplemented from time to time.

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