

Azitra Announces First Patient Dosed in Phase 1b Trial of ATR-12 for Netherton Syndrome

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BRANFORD, Conn.—(BUSINESS WIRE)—Aug. 28, 2024—Azitra, Inc. (NYSE American: AZTR), a precision dermatology company, announced that it has initiated dosing the first patient in its Phase 1b clinical trial evaluating ATR-12 for the treatment of Netherton syndrome. The study is designed to enrol approximately 12 adult patients with twice-daily treatment for 14 days, with primary endpoints focusing on safety and tolerability, and secondary and exploratory endpoints assessing efficacy signals and biomarkers. Azitra expects to report interim safety data from the Phase 1b trial in early 2025 with full results accordinated or 10 second half of 2025.

ATR-12 is Azitra's lead candidate, a proprietary strain of Staphylococcus epidermidis engineered to express therapeutic levels of an active subunit of the LEKTI protein to treat Netherton syndrome, a chronic genetic skin disease that can be life threatening,

"Dosing the first patient in our Phase 1b trial of ATR-12 marks a vital milestone for Azitra and for patients suffering from Netherton syndrome," said Francisco Salva, CEO of Azitra. "We are exhilarated to start dosing patients and look forward to executing this trial and delivering clinical data."

Mary Spellman, MD, Azitra's acting CMO, added, "We are thrilled to begin this clinical trial of ATR-12 in patients with Netherton syndrome. These patients often suffer from poor quality of life due to debilitating disease. This first-in-human study will inform future studies for the treatment of Netherton syndrome, including in pediatric patients and for longer treatment durations."

The Phase 1b trial (NCT06137157) is a multicenter, randomized, double-blind, vehicle-controlled study in approximately 12 adult patients with Netherton syndrome. Patients will be treated with 10⁹ CFU / g wice daily with ATR-12 or its vehicle control on the contralateral side of the body twice daily for 14 days. The primary objective is to assess he safety and tolerability of topical ATR-12 application. Secondary objectives include evaluating patients are contral applications and patient global assessments) and skin pharmacokinetics of the LEKTI subunit. Additional exploratory objectives include evaluating pharmacokynamic parameters, biomarkers, and LEKTI response, and cytokine responses.

About ATR-12

ATR-12 (also known as ATR12-351) is an engineered strain of *S. epidermidis* that expresses an active subunit of human lympho-epithelial Kazal-type-related inhibitor (LEKTI) protein, which is missing in patients with Netherton syndrome, a chronic and sometimes fatal disease of the skin estimated to affect approximately one to nine in every 100,000. ATR-12 has been engineered to deliver the missing LEKTI protein when applied topically. Aziltra has initiated a 12-patient, Phase 1b clinical trial, which will assess safety, tolerability, and efficacy endpoints (NCT06137157). Aziltra expects to announce initiat safety data in early 2025.

About Azitra, Inc.

Action. Inc. is an early-stage clinical biopharmacoutical company focused on developing innovative therapies for precision dematology using engineered probine and topical live biotherapeutic products. The Company has built a proprietary polarors that can be screened for unique therapeutic characteristics. The problem is augmented by artificial intelligence and machine learning technology that analyzes, predicts, and helpers go reserve the Company's initial focus. In the development of genetically engineered strains of Supphylococcus epidemiols, which the Company considers to be an optimal therapeutic candidate species for engineer of demandate species for engineer of demandate

Forward-Looking Statements

This press release contains forward-looking statements within the meaning of the Private Securities Litigation Reform Act of 1995, as amended. These statements may be identified by words such as "aims," "anticipates," "believes," "could," "estimates," "expects," "forecasts," "goal," "intends," "may," "plans," "possible," "possible," "possible," "possible," possible," possible and the private of the expects of these words or similar expressions that are intended to be forward-looking statements in this press release that are not statements of the statements and a contract of the expects, "and a contract of the expects," and a contract of the expects of the ex

Any forward-looking statements in this press release are based on current expectations, estimates and projections only as of the date of this release and are subject to a number of risks and uncertainties in this press release are based on current expectations, estimates and projections only as of the date of this release and are subject to a number of risks and uncertainties in the pressure of the product candidates and obtain required approval before commercialization; our product candidates and obtain required approval before commercialization; our product candidates may be inaccurate; we may be inaccurate; we may fail to timely risks additional required funding more efficient competitors or more effective competing treatment may emerge, we may be involved in disputes surrounding the use of our intellectual property crucial to our success; we may not be able to attract and retain key employees and qualified personnel; earlier study results may not be predictive of later stage study outcomes; and we are dependent on thirt-parties to some or all aspects of our product manufacturing, research and preclinical and clinical testing. Additional risks are described in our Form 10-Q filled with the SEC on August 12, 2024. Aztira explicitly disclaims any obligation to update any forward-looking statements except to the extent required by law.

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