



Aclaris Therapeutics Initiates Phase 1b Proof-of-Concept Trial in Atopic Dermatitis (AD) with Its Novel Bispecific Antibody ATI-052

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WAYNE, Pa., Jan. 12, 2026 (GLOBE NEWSWIRE) -- Aclaris Therapeutics, Inc. (NASDAQ: ACRS), a clinical-stage biopharmaceutical company focused on developing novel product candidates for immuno-inflammatory diseases, today announced that it has initiated a placebo-controlled Phase 1b proof-of-concept (POC) trial in atopic dermatitis (AD) for ATI-052, the Company's potential best-in-class investigational bispecific anti-TSLP/IL-4R α antibody.

"We are experiencing strong momentum in the ATI-052 clinical development program including the recent positive Phase 1a interim results which demonstrated a strong safety and tolerability profile, extended pharmacokinetics, and concentration-dependent pharmacodynamics even at the lowest dose," said Dr. Jesse Hall, Chief Medical Officer of Aclaris. "Further to this, we are excited to announce the initiation of the first POC Phase 1b trial in people living with atopic dermatitis. We expect to initiate the second Phase 1b POC trial in asthma this quarter as well. We intend to provide top line results from both trials in the second half of the year."

This randomized (3:1), blinded, placebo-controlled Phase 1b POC trial will evaluate the safety and tolerability of ATI-052 compared to placebo in patients with moderate-to-severe AD. Other endpoints that will be assessed include AD clinical efficacy assessments (Eczema Area and Severity Index (EASI), validated Investigator Global Assessment (IGA) response, and Peak Pruritus Numerical Rating Scale (PP-NRS)), pharmacokinetic parameters, and pharmacodynamic endpoints measured by assays including lesional and non-lesional skin tape strips.

Aclaris also expects to initiate a Phase 1b POC trial in asthma in the first quarter of 2026. Top line results from both POC trials are expected in the second half of 2026.

About ATI-052

ATI-052 is an investigational humanized anti-TSLP and anti-IL-4R α bispecific antibody that simultaneously inhibits thymic stromal lymphopoietin (TSLP) and interleukin-4 receptor (IL-4R α) with high affinity and potency. By targeting TSLP, which sits at the top of the inflammatory cascade, it inhibits a broad range of inflammation; by targeting IL-4R α , it blocks both downstream IL-4 and IL-13, which are key cytokines involved in Th2-mediated inflammation and allergic diseases. ATI-052 exhibits potential best-in-class potency and utilizes the same TSLP antigen-binding fragment (Fab) region as bosakitug (ATI-045), retaining the dissociation kinetics, long residence time, and high potency advantages over comparator antibodies, but is engineered to bind more tightly to the neonatal Fc receptor (FcRn) to extend its half-life. ATI-052 has the potential to treat a variety of atopic, immunologic and respiratory diseases. Aclaris has the exclusive worldwide rights to ATI-052, excluding Greater China.

About Aclaris Therapeutics, Inc.

Aclaris Therapeutics, Inc. is a clinical-stage biopharmaceutical company developing a pipeline of novel product candidates to address the needs of patients with immuno-inflammatory diseases who lack satisfactory treatment options. The company has a multi-stage portfolio of product candidates powered by a robust R&D engine. For additional information, please visit www.aclaristx.com and follow Aclaris on X (formerly Twitter) at @AclarisTx and on [LinkedIn](https://www.linkedin.com/company/aclaris-therapeutics).

Cautionary Note Regarding Forward-Looking Statements

Any statements contained in this press release that do not describe historical facts may constitute forward-looking statements as that term is defined in the Private Securities Litigation Reform Act of 1995. These statements may be identified by words such as "anticipate," "believe," "expect," "intend," "may," "plan," "potential," "will," and similar expressions, and are based on Aclaris' current beliefs and expectations. These forward-looking statements include expectations regarding its development plans for ATI-052, including the timing to initiate its Phase 1b trial of ATI-052 in asthma and the timing to report results from its Phase 1b trials of ATI-052 in AD and asthma, the potential for ATI-052 to be a best-in-class anti-TSLP/IL-4R α bispecific monoclonal antibody, and the therapeutic potential for ATI-052 including in other atopic, immunologic and respiratory diseases. These statements involve risks and uncertainties that could cause actual results to differ materially from those reflected in such statements. Risks and uncertainties that may cause actual results to differ materially include uncertainties inherent in the conduct of clinical trials, potential changes to interim, topline and preliminary data as more subject data become available, Aclaris' reliance on third parties over which it may not always have full control, Aclaris' ability to enter into strategic partnerships on commercially reasonable terms, the uncertainty regarding the macroeconomic environment and other risks and uncertainties that are described in the Risk Factors section of Aclaris' Annual Report on Form 10-K for the year ended December 31, 2024, and other filings Aclaris makes with the U.S. Securities and Exchange Commission from time to time. These documents are available under the "SEC Filings" page of the "Investors" section of Aclaris' website at www.aclaristx.com. Any forward-looking statements speak only as of the date of this press release and are based on information available to Aclaris as of the date of this release, and Aclaris assumes no obligation to, and does not intend to, update any forward-looking statements, whether as a result of new information, future events or otherwise.

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