



January 6, 2017

Curis Reports Inducement Grants Under NASDAQ Listing Rule 5635(c)(4)

LEXINGTON, Mass., Jan. 06, 2017 (GLOBE NEWSWIRE) -- Curis, Inc. (NASDAQ:CRIS), a biotechnology company focused on the development and commercialization of innovative and effective drug candidates for the treatment of human cancers, today announced that on January 1, 2017, the independent Compensation Committee of the Board of Directors of Curis approved the grant of inducement stock options to purchase a total of 185,000 shares of Curis common stock to eight new employees, with a grant date of January 1, 2017 (the "Q1 2017 Inducement Grants").

Each of the Q1 2017 Inducement Grants has an exercise price per share equal to the closing price of the Company's common stock on December 30, 2016, the last trading date prior to the date of grant. Each stock option has a 10 year term and vests over four years, with 25% of the original number of shares underlying the award vesting on the first anniversary of the employee's date of hire and an additional 6.25% of the original number of shares underlying the award vesting on each successive three-month period thereafter, subject to the new employee's continued service with the Company through the respective vesting dates. Each stock option was granted as an inducement equity award outside of the Company's Amended and Restated 2010 Stock Incentive Plan, as amended, and was made as an inducement material to such employee's acceptance of employment with the Company.

About Curis, Inc.

Curis is a biotechnology company focused on the development and commercialization of innovative and effective drug candidates for the treatment of human cancers, including its lead development candidate, CUDC-907 that is being investigated in clinical studies in patients with lymphomas and solid tumors. Curis is also engaged in a broad collaboration with Aurigene in the areas of immuno-oncology and precision oncology. As part of this collaboration, Curis has exclusive licenses to oral small molecule antagonists of the PD-1 and VISTA pathways, including PD-L1/VISTA antagonist CA-170, and oral small molecule antagonists of the PD-1 and TIM-3 pathways, including PD-L1/TIM-3 antagonist CA-327, as well as to molecules designed to inhibit the IRAK4 kinase, including CA-4948. CA-170 is currently undergoing testing in a Phase 1 trial in patients with advanced solid tumors and lymphomas. Curis is also party to a collaboration with Genentech, a member of the Roche Group, under which Genentech and Roche are commercializing Erivedge® for the treatment of advanced basal cell carcinoma, and are further developing Erivedge in other diseases including idiopathic pulmonary fibrosis and myelofibrosis. For more information, visit Curis's website at www.curis.com.

Cautionary Note Regarding Forward-Looking Statements:

This press release contains forward-looking statements within the meaning of the Private Securities Litigation Reform Act of 1995, including without limitation statements regarding the potential advantages and benefits of small molecule checkpoint inhibitors and Curis's plans and expectations for the collaboration with Aurigene, including its plans to discover and develop multiple first-in-class oral, small molecule checkpoint inhibitors for the treatment of patients with cancer. Forward-looking statements may contain the words "believes," "expects," "anticipates," "plans," "seeks," "estimates," "assumes," "will," "may," "could" or similar expressions. These forward-looking statements are not guarantees of future performance and involve risks, uncertainties, assumptions and other important factors that may cause actual results to be materially different from those indicated by such forward-looking statements. For example, Curis may experience adverse results, delays and/or failures in its drug development programs and may not be able to successfully advance the development of its drug candidates in the time frames it projects, if at all. Curis's drug candidates may cause unexpected toxicities, fail to demonstrate sufficient safety and efficacy in clinical studies and/or may never achieve the requisite regulatory approvals needed for commercialization. Favorable results seen in preclinical studies and early clinical trials of Curis's drug candidates may not be replicated in later trials. There can be no guarantee that the collaboration agreement with Aurigene will continue for its full term, that Curis or Aurigene will each maintain the financial and other resources necessary to continue financing its portion of the research, development and commercialization costs, that the parties will successfully discover, develop or commercialize drug candidates under the collaboration, or that Curis receive full or partial benefit of payments waived by Aurigene. Curis Royalty may not receive sufficient levels of royalty revenue from sales of Erivedge to satisfy its royalty-collateralized debt obligation or may otherwise lose its rights to Erivedge royalties and royalty-related payments as a result of a foreclosure of the loan. Curis will require substantial additional capital to fund its business and such capital may not be available on reasonable terms, or at all. Curis also faces risks relating to: potential adverse decisions made by the FDA and other regulatory authorities, investigational review boards, and publication review bodies; competition; its ability to obtain or maintain necessary patent protection; unstable market and economic conditions; unplanned expenses; and other important risks relating to its business, operations, financial condition and future prospects that are discussed in its most

recent Form 10-K and Form 10-Q and other filings that it periodically makes with the Securities and Exchange Commission.

In addition, any forward-looking statements represent the views of Curis only as of today and should not be relied upon as representing Curis's views as of any subsequent date. Curis disclaims any intention or obligation to update any of the forward-looking statements after the date of this press release whether as a result of new information, future events or otherwise, except as may be required by law.

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