

March 27, 2018



Cellecstar Granted Seminal U.S. Patent for Phospholipid-Ether Analogs as Cancer-Targeting Drug Vehicles

Covers composition of matter and method of use for proprietary PDCs™ in combination with anti-cancer agents

MADISON, Wis., March 27, 2018 (GLOBE NEWSWIRE) -- Cellecstar Biosciences, Inc. (Nasdaq:CLRB), a clinical-stage biopharmaceutical company focused on the discovery, development and commercialization of targeted treatments for cancer, announces that the U.S. Patent and Trademark Office (USPTO) has granted patent number 9,345,718, titled "Phospholipid-Ether Analogs as Cancer-Targeting Drug Vehicles," related to the company's phospholipid-ether analogs for targeting anticancer therapeutics to tumors and cancer stem cells.

"This seminal patent provides Cellecstar with broad protection for Phospholipid Drug Conjugate™ (PDC™) products created with our phospholipid ether technology, as well as the freedom to operate that is necessary to maximize the value of the platform," stated Jim Caruso, chief executive officer of Cellecstar Biosciences. "The delivery technology forms the backbone of our lead PDC candidate, CLR 131, which continues to advance in clinical trials for a variety of hematologic and solid tumors."

The new patent protects both composition of matter and method of use for PDCs developed with Cellecstar's proprietary phospholipid-ether delivery vehicle conjugated with any known or in-development anti-cancer agent, including the company's pipeline of internal and partnered PDC programs designed for targeted delivery to cancer cells and cancer stem cells.

About Phospholipid Drug Conjugates™

Cellecstar's product candidates are built upon a patented delivery and retention platform that utilizes optimized PDCs to target cancer cells. The PDC platform is used to selectively deliver diverse oncologic payloads to cancerous cells and cancer stem cells, including hematologic cancers and solid tumors. This selective delivery allows a payloads' therapeutic window to be modified, which may maintain or enhance drug potency while reducing the number and severity of adverse events. This platform takes advantage of a metabolic pathway utilized by all tumor cell types in all cell cycle stages. Compared with other targeted delivery platforms, the PDC platform's mechanism of entry does not rely upon specific cell surface epitopes or antigens. In addition, PDCs can be conjugated to molecules in numerous ways, thereby increasing the types of molecules selectively delivered. Cellecstar believes the PDC platform holds potential for the discovery and development of the next generation of cancer-targeting agents.

About Cellecstar Biosciences, Inc.

Cellecstar Biosciences is focused on the discovery, development and commercialization of drugs for the treatment of cancer. The company plans to develop proprietary drugs

independently and through research and development (R&D) collaborations. The core drug development strategy is to leverage our PDC platform to develop therapeutics that specifically target treatment to cancer cells. Through R&D collaborations, the company's strategy is to generate near-term capital, supplement internal resources, gain access to novel molecules or payloads, accelerate product candidate development and broaden our proprietary and partnered product pipelines.

The company's lead PDC therapeutic, CLR 131, is in a Phase 1 clinical study in patients with relapsed or refractory (R/R) multiple myeloma (MM) and a Phase 2 clinical study in R/R MM and a range of B-cell malignancies. In 2018 the company plans to initiate a Phase 1 study with CLR 131 in pediatric solid tumors and lymphoma, and a second Phase 1 study in combination with external beam radiation for head and neck cancer. The company's product pipeline also includes two preclinical PDC chemotherapeutic programs (CLR 1700 and 1900) and partnered assets include PDCs from multiple R&D collaborations.

For more information please visit www.cellectar.com.

Forward-Looking Statement Disclaimer

This news release contains forward-looking statements. You can identify these statements by our use of words such as "may," "expect," "believe," "anticipate," "intend," "could," "estimate," "continue," "plans," or their negatives or cognates. These statements are only estimates and predictions and are subject to known and unknown risks and uncertainties that may cause actual future experience and results to differ materially from the statements made. These statements are based on our current beliefs and expectations as to such future outcomes. Drug discovery and development involve a high degree of risk. Factors that might cause such a material difference include, among others, uncertainties related to the ability to raise additional capital, uncertainties related to the ability to attract and retain partners for our technologies, the identification of lead compounds, the successful preclinical development thereof, the completion of clinical trials, the FDA review process and other government regulation, our pharmaceutical collaborators' ability to successfully develop and commercialize drug candidates, competition from other pharmaceutical companies, product pricing and third-party reimbursement. A complete description of risks and uncertainties related to our business is contained in our periodic reports filed with the Securities and Exchange Commission including our Form 10-K for the year ended December 31, 2017. These forward-looking statements are made only as of the date hereof, and we disclaim any obligation to update any such forward-looking statements.

CONTACT:

LHA Investor Relations
Miriam Weber Miller
212-838-3777
mmiller@lhai.com



Source: Collectar Biosciences, Inc.