

Iovance Biotherapeutics Announces H. Lee Moffitt Cancer Center to Present Lung Cancer Clinical Data at American Association for Cancer Research (AACR) Virtual Annual Meeting 2020

April 13, 2020

Phase 1 Study Combining Tumor Infiltrating Lymphocytes (TIL) and Nivolumab in Non-Small Cell Lung Cancer Funded in Part by Iovance and Stand Up To Cancer

SAN CARLOS, Calif., April 13, 2020 (GLOBE NEWSWIRE) -- Iovance Biotherapeutics, Inc. (NASDAQ: IOVA), a late-stage biotechnology company developing novel T cell-based cancer immunotherapies, today announced that H. Lee Moffitt Cancer Center ("Moffitt") plans to present clinical results from a Phase 1 trial using Moffitt's tumor infiltrating lymphocytes (TIL) in patients with non-small cell lung cancer (NSCLC) at the American Association for Cancer Research (AACR) Virtual Annual Meeting I, to be held April 27-28, 2020. The Phase 1 study is being conducted at Moffitt with support from lovance Biotherapeutics, a Stand Up To Cancer Catalyst® grant, and other partners.

Maria Fardis, Ph.D., MBA, President and Chief Executive Officer of Iovance, stated, "We are very pleased that Moffitt will present clinical data demonstrating the potential for tumor infiltrating lymphocytes, or TIL, in such an unmet medical need indication, non-small cell lung cancer. This data is the basis of our strategy to investigate Iovance TIL in two cohorts of non-small cell lung cancer patients in our IOV-COM-202 'basket' study. We continue to be excited about the broader potential of Iovance TIL in additional tumor types."

 Abstract Title: Durable complete responses to adoptive cell transfer using tumor infiltrating lymphocytes (TIL) in non-small cell lung cancer (NSCLC): a phase I trial.

Authors: Ben Creelan, et al.

Session: VCTPL05 - Adoptive Cell Transfer Therapy

Date and Time: April 28, 2020, 1:00 PM Abstract Number: 20-LB-10617-AACR

Location: AACR Virtual Annual Meeting I at www.aacr.org

The AACR Virtual Annual Meeting I will include more than 30 oral presentations in several clinical trial plenary sessions along with commentaries from expert discussants, as well as clinical trial poster sessions consisting of short videos providing the authors' perspectives. This Virtual Meeting will be available free to everyone, although attendees will be asked to register to participate. For more information please visit the AACR Virtual Annual Meeting I page at www.aacr.org.

About Iovance Biotherapeutics, Inc.

lovance Biotherapeutics aims to improve patient care by making T cell-based immunotherapies broadly accessible for the treatment of patients with solid tumors and blood cancers. Tumor infiltrating lymphocyte (TIL) therapy uses a patient's own immune cells to attack cancer. TIL cells are extracted from a patient's own tumor tissue, expanded through a proprietary process, and infused back into the patient. After infusion, TIL reach tumor tissue, where they attack tumor cells. The company has completed dosing in the pivotal study in patients with metastatic melanoma and is currently conducting a pivotal study in patients with advanced cervical cancer. In addition, the company's TIL therapy is being investigated for the treatment of patients with locally advanced, recurrent or metastatic cancers including head and neck and non-small cell lung cancer. A clinical study to investigate lovance T cell therapy for blood cancers called peripheral blood lymphocyte (PBL) therapy is open to enrollment. For more information, please visit www.iovance.com.

Forward-Looking Statements

Certain matters discussed in this press release are "forward-looking statements" of lovance Biotherapeutics, Inc. (hereinafter referred to as the "Company," "we," "us," or "our"). We may, in some cases, use terms such as "predicts," "believes," "potential," "continue," "estimates," "anticipates," "expects," "plans," "intends," "may," "could," "might," "will," "should" or other words that convey uncertainty of future events or outcomes to identify these forward-looking statements. The forward-looking statements include, but are not limited to, risks and uncertainties relating to the success, timing, projected enrollment, manufacturing and production capabilities, and cost of our ongoing clinical trials and anticipated clinical trials for our current product candidates (including both Company-sponsored and collaborator-sponsored trials in both the U.S. and Europe), such as statements regarding the timing of initiation and completion of these trials: the timing of and our ability to successfully submit, obtain and maintain FDA or other regulatory authority approval of, or other action with respect to, our product candidates, including those product candidates that have been granted breakthrough therapy designation ("BTD") or regenerative medicine advanced therapy designation ("RMAT") by the FDA and new product candidates in both solid tumor and blood cancers; the strength of the Company's product pipeline; the successful implementation of the Company's research and development programs and collaborations; the Company's ability to obtain tax incentives and credits; the guidance provided for the Company's future cash, cash equivalents, and short term investment balances; the success of the Company's manufacturing, license or development agreements; the acceptance by the market of the Company's product candidates, if approved; and other factors, including general economic conditions and regulatory developments, not within the Company's control. The factors discussed herein could cause actual results and developments to be materially different from those expressed in or implied by such statements. Actual results may differ from those set forth in this press release due to the risks and uncertainties inherent in the Company's business, including, without limitation: the preliminary clinical results, which may include efficacy and safety results, from ongoing Phase 2 studies may not be reflected in the final analyses of these trials or subgroups within these trials; a slower rate of enrollment may impact the Company's clinical trial timelines; enrollment may need to be adjusted for the Company's trials and cohorts within those trials based on FDA and other regulatory agency input; the new version of the protocol which further defines the patient population to include more advanced patients in the Company's cervical cancer trial may have an adverse effect on the results reported to date; the data within these trials may

not be supportive of product approval; changes in patient populations may result in changes in preliminary clinical results; the Company's ability or inability to address FDA or other regulatory authority requirements relating to its clinical programs and registrational plans, such requirements including, but not limited to, clinical, safety, manufacturing and control requirements; the Company's interpretation of communications with the FDA may differ from the interpretation of such communications by the FDA; risks related to the Company's ability to maintain and benefit from accelerated FDA review designations, including BTD and RMAT, which may not result in a faster development process or review of the Company's product candidates (and which may later be rescinded by the FDA), and does not assure approval of such product candidates by the FDA or the ability of the Company to obtain FDA approval in time to benefit from commercial opportunities; the ability or inability of the Company to manufacture its therapies using third party manufacturers or its own facility may adversely affect the Company's potential commercial launch; the results of clinical trials with collaborators using different manufacturing processes may not be reflected in the Company's sponsored trials; and additional expenses may decrease our estimated cash balances and increase our estimated capital requirements. A further list and description of the Company's risks, uncertainties and other factors can be found in the Company's most recent Annual Report on Form 10-K and the Company's subsequent filings with the Securities and Exchange Commission. Copies of these filings are available online at www.sec.gov or www.iovance.com. The forward-looking statements are made only as of the date of this press release and the Company undertakes no obligation to publicly update such forward-looking statements to reflect subsequent events or circumstances.

Iovance Biotherapeutics, Inc:

Sara Pellegrino, IRC Vice President, Investor Relations & Public Relations 650-260-7120 ext. 264 Sara.Pellegrino@iovance.com

Solebury Trout:

Annie Chang (investors) 646-378-2972 achang@troutgroup.com

Chad Rubin (investors) 646-378-2947 crubin@troutgroup.com

Rich Allan (media) 646-378-2958 rallan@troutgroup.com

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