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Lee's Pharmaceutical Holdings Limited

李氏大藥廠控股有限公司*

(incorporated in the Cayman Islands with limited liability)

(Stock Code: 950)

VOLUNTARY ANNOUNCEMENT – UPDATE ON THE RESEARCH AND DEVELOPMENT OF AN INVESTIGATIONAL ONCOLOGY PRODUCT

This announcement is made by the board (the “**Board**”) of directors (the “**Directors**”) of Lee’s Pharmaceutical Holdings Limited (the “**Company**” or “**Lee’s Pharm**”, together with its subsidiaries as the “**Group**”) on a voluntary basis.

The Board of the Company is pleased to announce that, on 26 September 2021, China Oncology Focus Limited, a subsidiary of the Group, has obtained approval from the Center for Drug Evaluation (the “**CDE**”) of the China’s National Medical Products Administration (the “**NMPA**”) to submit the new drug application (“**NDA**”) for the Socazolimab (an anti-PD-L1 monoclonal antibody, formerly known as ZKAB001) to treat recurrent or metastatic cervical cancer, which was granted breakthrough therapy designation (“**BTD**”) on 5 February 2021.

Socazolimab is a fully human anti PD-L1 monoclonal antibody targeting tumor PD-L1 protein. It can release the “brake” causing by the tumor cell to the immune system. The clinical study of Socazolimab monotherapy in recurrent and metastatic cervical cancer is led by Doctor Lingying Wu as the principal investigator and consists of two phases, an open labeled 3+3 dose escalation phase followed by a pivotal expansion phase study. A total of 91 patients has participated in the pivotal part of the study and the results indicating that Socazolimab has excellent efficacy profile based on the improved tumor response rate regardless of patient PD-L1 expression level, prolonged duration of response, progression free survival and overall survival.

Socazolimab is an in-licensed product from Sorrento Therapeutics, Inc (“**Sorrento**”) for the People’s Republic of China, Hong Kong, Macau and Taiwan.

* For identification purpose only

ABOUT SOCAZOLIMAB

Socazolimab is a fully human anti-PD-L1 monoclonal antibody identified by Sorrento using its proprietary G-MAB™ library platform. Socazolimab has the following potential advantages over its competitors:

1. Fully human antibody potentially allows it to have minimal immunogenicity; demonstrated by its negative antigen-derived antibody (ADA) generation in humans in studies to date.
2. Potentially lower dose required to achieve efficacy compared to other anti-PD-L1 antibodies.
3. Dual mechanism of action observed with both immune-checkpoint inhibition and antibody-dependent cellular cytotoxicity (ADCC) effect.

The antibody has been tested or is being tested in various cancer indications including recurrent or metastatic cervical cancer, maintenance therapy for high-grade osteosarcoma after adjuvant chemotherapy, locally advanced and metastatic urothelial carcinoma, extensive small cell lung cancer in combination with carboplatin and etoposide, advanced urothelial carcinoma in combination with albumin-bound paclitaxel and esophageal carcinoma.

ABOUT SORRENTO

Sorrento is a clinical stage, antibody-centric, biopharmaceutical company developing new therapies to treat cancers and COVID-19. Sorrento's multimodal, multipronged approach to fighting cancer is made possible by its extensive immuno-oncology platforms, including key assets such as fully human antibodies ("**G-MAB™ library**"), clinical stage immuno-cellular therapies ("**CAR-T**", "**DAR-T™**"), antibody-drug conjugates ("**ADCs**"), and clinical stage oncolytic virus ("**Seprehvir™**"). Sorrento is also developing potential antiviral therapies and vaccines against coronaviruses, including COVI-GUARD™, COVI-AMG™, COVISHIELD™, Gene-MAb™, COVI-MSCTM and COVI-DROPS™; and diagnostic test solutions, including COVI-TRACK™, COVI-STIX™ and COVI-TRACE™.

Sorrento's commitment to life-enhancing therapies for patients is also demonstrated by its effort to advance a first-in-class (TRPV1 agonist) non-opioid pain management small molecule, resiniferatoxin ("**RTX**"), and SP-102 (10 mg, dexamethasone sodium phosphate viscous gel) (SEMDEXA™), a novel, viscous gel formulation of a widely used corticosteroid for epidural injections to treat lumbosacral radicular pain, or sciatica, and to commercialise ZTlido® (lidocaine topical system) 1.8% for the treatment of post-herpetic neuralgia. RTX has completed a phase IB trial for intractable pain associated with cancer and a phase 1B trial in osteoarthritis patients. SEMDEXA is in a pivotal Phase 3 trial for the treatment of lumbosacral radicular pain, or sciatica. ZTlido® was approved by the FDA on 28 February 2018. More information available at www.sorrentotherapeutics.com.

ABOUT COF

COF is a subsidiary of Lee's Pharm and a clinical development stage company focused in oncology with emphasis in immune oncology. COF is currently developing several assets, including Socazolimab (anti-PD-L1 antibody) in pivotal clinical trial stage; Zotiraciclib, an oral multi-kinase inhibitor in Phase I clinical trial for glioblastoma; Gimatécán, a topoisomerase I inhibitor in Phase I clinical trial for ovarian cancer and in Phase Ib/II clinical trial for small cell lung cancer in China; Pexa-vec (oncolytic virus) which is in global Phase Ib clinical trial for renal cell cancer. COF has built a pipeline of 10 assets through internal development and in-licensing. The diversity of its products creates a unique position for the company to use immune oncology as backbone therapy in combination with in-house products and develop potential paradigm-shifting treatment for cancer.

ABOUT LEE'S PHARM

Lee's Pharm is a research-driven and market-oriented biopharmaceutical company with more than 25 years of operation in the pharmaceutical industry in China. The Company is fully integrated with solid infrastructures in drug development, clinical development, regulatory, manufacturing, sales and marketing based in Mainland China with global perspectives. The Company has established extensive partnerships with over 20 international companies and currently markets 25 proprietary, generic and licensed-in pharmaceutical products in Mainland China, Hong Kong, Macau and Taiwan. The Company focuses on several key disease areas such as cardiovascular, woman health, paediatrics, rare diseases, oncology, dermatology and obstetrics, and has more than 40 products under different development stages stemming from both internal research and development as well as from the licensing and development, commercialisation, and manufacturing rights from various United States, European and Japanese companies.

By order of the Board
Lee's Pharmaceutical Holdings Limited
Lee Siu Fong
Chairman

Hong Kong, 27 September 2021

As at the date of this announcement, Ms. Lee Siu Fong (Chairman) and Ms. Leelalertsuphakun Wanee are executive directors of the Company, Dr. Li Xiaoyi is a non-executive director of the Company, Dr. Chan Yau Ching, Bob, Mr. Lam Yat Cheong and Dr. Tsim Wah Keung, Karl are independent non-executive directors of the Company.