

## PharmaCyte Biotech Successfully Completes All Release Tests on Clinical Trial Product

LAGUNA HILLS, Calif.--(BUSINESS WIRE)-- PharmaCyte Biotech, Inc. (OTCQB: PMCB), a biotechnology company focused on developing cellular therapies for cancer and diabetes using its signature live-cell encapsulation technology, Cell-in-a-Box<sup>®</sup>, announced today that both batches of the company's manufactured clinical trial product have undergone and passed all 10 of the necessary "release tests" required by the U.S. Food and Drug Administration (FDA).

PharmaCyte's partner, Austrianova Singapore (Austrianova), is now completing and signing the paperwork to issue PharmaCyte a Certificate of Analysis for each batch of the company's clinical trial product. Austrianova will also be turning over all of the completed batch records to PharmaCyte for both production runs that tested the company's clinical trial product for the "functionality" of the encapsulated cells and the "safety" of the clinical trial product. These tests will provide essential data for PharmaCyte to complete its Investigational New Drug application (IND) to the FDA to request approval for our planned clinical trial in locally advanced, inoperable pancreatic cancer (LAPC).

PharmaCyte's Chief Executive Officer, Kenneth L. Waggoner, said, "After completing the production of our clinical trial product, it is extremely good news to learn that our capsules and the live cells inside are functioning and safe to put inside patients in our upcoming planned clinical trial in locally advanced, inoperable pancreatic cancer. There were 20 tests in total over both batches, and our clinical trial product passed all of them. This proves that our years of hard work and working through each challenging hurdle was well worth it. We now eagerly await the two Certificates of Analysis and the records from both batches from Austrianova so that we can begin to compile and enter that data into our IND application."

To learn more about PharmaCyte's pancreatic cancer treatment and how it works inside the body to treat locally advanced inoperable pancreatic cancer, we encourage you to watch the company's documentary video complete with medical animations at: <a href="https://www.PharmaCyte.com/Cancer">https://www.PharmaCyte.com/Cancer</a>

## **About PharmaCyte Biotech**

PharmaCyte Biotech, Inc. (PharmaCyte) is a biotechnology company developing cellular therapies for cancer and diabetes based upon a proprietary cellulose-based live cell encapsulation technology known as "Cell-in-a-Box®." This technology will be used as a platform upon which therapies for several types of cancer and diabetes are being developed.

PharmaCyte's therapy for cancer involves encapsulating genetically engineered human cells that convert an inactive chemotherapy drug into its active or "cancer-killing" form. For

pancreatic cancer, these encapsulated cells are implanted in the blood supply to the patient's tumor as close as possible to the site of the tumor. Once implanted, a chemotherapy drug that is normally activated in the liver (ifosfamide) is given intravenously at one-third the normal dose. The ifosfamide is carried by the circulatory system to where the encapsulated cells have been implanted. When the ifosfamide flows through pores in the capsules, the live cells inside act as a "bio-artificial liver" and activate the chemotherapy drug at the site of the cancer. This "targeted chemotherapy" has proven effective and safe to use in past clinical trials and results in little to no treatment related side effects.

PharmaCyte's therapy for Type 1 diabetes and insulin-dependent Type 2 diabetes involves encapsulating a human cell line that has been genetically engineered to produce and release insulin in response to the levels of blood sugar in the human body. PharmaCyte is developing the use of genetically modified liver cells and stem cells, as well as beta islet cells, to treat diabetes. The encapsulation will be done using the Cell-in-a-Box<sup>®</sup> technology. Once the encapsulated cells are implanted in a diabetic patient, they will function as a "bio-artificial pancreas" for purposes of insulin production.

## Safe Harbor

This press release may contain forward-looking statements within the meaning of the Private Securities Litigation Reform Act of 1995 that express the current beliefs and expectations of the management of PharmaCyte, including statements regarding the timing and commencement of our first Phase 2b clinical trial. Any statements contained herein that do not describe historical facts are forward-looking statements that are subject to risks and uncertainties that could cause actual results, performance and achievements to differ materially from those discussed in such forward-looking statements. Factors that could affect our actual results are included in the periodic reports on Form 10-K and Form 10-Q that we file with the U.S. Securities and Exchange Commission. These forward-looking statements are made only as of the date hereof, and we undertake no obligation to update or revise the forward-looking statements, except as otherwise required by law, whether as a result of new information, future events or otherwise

More information about PharmaCyte can be found at <a href="www.PharmaCyte.com">www.PharmaCyte.com</a>. Information may also be obtained by contacting PharmaCyte's Investor Relations Department.

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