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Scancell Holdings Plc
(‘Scancell Holdings’ or the ‘Company’)

Clinical Trial Application for SCIB1 melanoma vaccine

Scancell Holdings Plc, (PLUS:SCLP), the developer of therapeutic cancer vaccines, is pleased to announce that its proposal to conduct a Phase I clinical trial on SCIB1, its DNA ImmunoBody® vaccine being developed for the treatment of melanoma, was submitted to the Gene Therapy Advisory Committee (‘GTAC’) on 29 December 2009. In addition, Scancell and its partner Ichor Medical Systems (‘Ichor’) submitted parallel applications to the Medicines Division and to the Devices Division of the Medicines and Healthcare products Regulatory Agency (‘MHRA’) on 29 January 2010 requesting approvals for the clinical trial of SCIB1 and for the use of Ichor’s TriGrid™ electroporation delivery device to administer SCIB1 to participating patients, respectively.

SCIB1 is a novel DNA ImmunoBody® vaccine being developed using Scancell’s patented ImmunoBody® technology for the treatment of melanoma. ImmunoBody® vaccines generate the high-avidity T-cells that kill cancer cells, which may overcome the current limitations of most cancer vaccines. *In vivo* electroporation is widely regarded as an effective method of enhancing the potency of DNA vaccines by up to 100 -fold compared to conventional methods of delivery. Scancell is confident that TriGrid™ will provide the most effective delivery system for its SCIB1 melanoma vaccine as it enters clinical trials.

Advanced melanoma currently has a very poor prognosis with late stage (stage IV) disease having a median survival of approximately six months. According to the World Health Organisation, 132,000 melanoma skin cancers occur globally each year and the incidence is increasing, especially in the United States, Europe and Australia.

David Evans, Chairman of Scancell, commented:

“Scancell’s CTA submission and proposal to GTAC to conduct the Phase I clinical trial for our first therapeutic cancer vaccine marks a significant step for the Company and we look forward to reporting on its progress. We are confident that, subject to regulatory and ethical approvals, the Phase I clinical trial for SCIB1 will be on target to commence in Q2 2010.”

A copy of this announcement is available for download on the Company’s website at <http://www.scancell.co.uk/>

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About Scancell

Scancell is developing novel therapeutic vaccines for the treatment of cancer and infectious diseases based on its groundbreaking ImmunoBody® technology platform. Scancell's first cancer vaccine, SCIB1, is being developed for the treatment of melanoma and will enter clinical trials in early 2010.

Treating cancer by vaccination allows small non-toxic doses of a vaccine to be administered to a patient, stimulating an immune response. Effective cancer vaccine need to target dendritic cells to stimulate both parts of the cellular immune system; the helper cell system where inflammation is stimulated at the tumour site; and the cytotoxic T-lymphocyte or CTL response where immune system cells are primed to recognise and kill specific cells.

A limitation of many cancer vaccines currently in development is that they cannot specifically target dendritic cells *in vivo*. Several groups have demonstrated successful vaccination by growing dendritic cells *ex vivo*, pulsing them with tumour antigens and re-infusing them. However, this procedure is patient specific, time consuming and expensive. Scancell has developed its breakthrough patent protected ImmunoBody® technology to overcome these limitations.

An ImmunoBody® is a DNA vaccine encoding a human antibody or fusion protein engineered to express helper cell and CTL epitopes from tumour antigens over-expressed by cancer cells. Antibodies are ideal vectors for carrying T cell epitopes from tumour antigens as they can effectively target dendritic cells via their Fc receptors, allowing efficient stimulation of high avidity and high frequency helper and CTL responses.

The ImmunoBody® technology can be adapted to provide the basis for treating any tumour type and may also be of potential utility in the development of vaccines against hepatitis, HIV and other chronic infectious diseases.

About Ichor

Ichor Medical Systems' TriGrid™ Delivery System is the first integrated and fully automated system for electroporation-mediated DNA administration. Ichor, a privately-held biotech company based in San Diego, CA, is collaborating with partners on three continents in a wide range of studies to test the TriGrid™ as an enabling platform for delivery of DNA drugs and vaccines to treat diseases such as pandemic flu, hepatitis, HIV, melanoma, multiple sclerosis, and others. The TriGrid™ is also being tested by the U.S. military as an efficient means of delivering anti-bioterrorism agents.