DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Prospective Grant of Exclusive License: Development of T Cell Receptors and Chimeric Antigen Receptors into Therapeutics for Adoptive Transfer in Humans To Treat Cancer

AGENCY: National Institutes of Health, Public Health Service, HHS.

ACTION: Notice.


Other than license applications submitted as objections to this Notice of Intent to Grant an Exclusive License, no further license applications will be considered for the exclusive field of use set forth below if Kite Pharma, Inc. is granted an exclusive license pursuant to this Notice of Intent to Grant an Exclusive License. The prospective exclusive license territory may be worldwide and the field of use may be limited to the treatment of cancers, which may include brain cancer, breast cancer, colorectal cancer, esophageal cancer, gastric cancer, head and neck cancer, liver cancer, lung cancer, melanoma, multiple myeloma, ovarian cancer, prostate cancer, sarcoma, and urothelial cancer, as claimed in the Licensed Patent Rights.

DATES: Only written comments and/or applications for a license which are received by the NIH Office of Technology Transfer on or before February 23, 2012 will be considered, in addition to the current non-exclusive applications under consideration, for the prospective license territory and field of use to be granted under the contemplated exclusive patent license.

ADDRESSES: Requests for copies of the patent application, inquiries, comments, and other materials relating to the contemplated exclusive license should be directed to: Samuel E. Bish, Ph.D., Licensing and Patenting Manager, Office of Technology Transfer, National Institutes of Health, 6011 Executive Boulevard, Suite 325, Rockville, MD 20852–3804; Telephone: (301) 435–5282; Facsimile: (301) 402–0220; Email: bishse@mail.nih.gov.

SUPPLEMENTARY INFORMATION: The technologies describe T cells engineered to express MAGE–A3, MAGE–A12, or SSX–2 T cell receptors (TCRs) or EGFRvIII chimeric antigen (CARs) and methods of using these engineered T cells to treat and/or prevent cancer. These technologies include the TCR and CAR amino acid sequences, the nucleic acid sequences that encode these compositions, vectors to express the TCRs and CARs, host cells and populations of host cells, such as T cells, that express the compositions, antibodies to the TCRs and CARs, pharmaceutical compositions, and associated methods of detecting, preventing, and treating diseases, such as cancer, with these TCRs and CARs. TCRs and CARs are proteins that recognize antigens, such as cancer antigens, and activate the cells expressing these compositions to destroy the antigen-expressing cell. TCRs consist of two domains, one variable domain that recognizes the antigen and one constant region that helps the TCR anchor to the membrane and transmit recognition signals by interacting with other proteins. CARs are hybrid proteins consisting of a portion of an antibody that recognizes an antigen fused to protein domains that signal to activate the CAR-expressing cell. Therapies utilizing these technologies involve isolating a cancer patient’s own T cells to be engineered with the TCR and/or CAR that recognize the tumor antigen(s) expressed on that specific patient’s cancer cell.

Afterwards, the engineered T cells from the patient are adoptively transferred back into the patient to mediate tumor regression. Personalized adoptive cell transfer therapies developed from these technologies could yield innovative therapeutics for any cancers that express the antigens recognized by these TCRs and CARs.

The prospective exclusive license, subject to current non-exclusive license applications under consideration and any further license applications received as objections to this Notice of Intent to Grant an Exclusive License, will be royalty bearing and will comply with the terms and conditions of 35 U.S.C. 209 and 37 CFR Part 404.7. The prospective exclusive license may be granted unless within thirty (30) days from the date of this published notice, the NIH receives written evidence and argument that establishes that the grant of the license would not be consistent with the requirements of 35 U.S.C. 209 and 37 CFR 404.7.

Any additional applications for a license in the field of use filed in response to this notice will be treated as objections to the grant of the contemplated exclusive license. Comments and objections submitted to this notice will not be made available for public inspection and, to the extent permitted by law, will not be released under the Freedom of Information Act, 5 U.S.C. 552.


Richard U. Rodriguez,
Director, Division of Technology Development and Transfer, Office of Technology Transfer, National Institutes of Health.

[FR Doc. 2012–1383 Filed 1–23–12; 8:45 am]

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DEPARTMENT OF HOMELAND SECURITY

Federal Emergency Management Agency

[Docket ID FEMA–2011–0029; OMB No. 1660–0095]

Agency Information Collection Activities: Submission for OMB Review; Comment Request, National Flood Insurance Claims Appeals Process

AGENCY: Federal Emergency Management Agency, DHS.

ACTION: Notice.

SUMMARY: The Federal Emergency Management Agency (FEMA) has submitted the following information collection to the Office of Management