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**ADDRESSES:** Licensing information and a copy of the U.S. patent application referenced below may be obtained by contacting Girish C. Barua, Ph.D., Technology Licensing Specialist, Office of Technology Transfer, National Institutes of Health, 6011 Executive Boulevard, Suite 325, Rockville, Maryland 20852-3804 (telephone 301/496-7735 ext 263; fax 301/402-0220). A signed Confidential Disclosure Agreement will be required to receive a copy of the patent application.

**Antibacterial Therapy With Bacteriophage Genotypically Modified To Delay Inactivation by the Host Defense System**

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Serial No. 08/222,956

The present invention is directed to bacteriophage therapy, using methods that enable the bacteriophage to delay inactivation by any and all parts of the host defense system (HDS) against foreign objects that would tend to reduce the numbers of bacteriophage and/or the efficiency of those phage at killing the host bacteria in an infection. Disclosed is a method of producing bacteriophage modified for anti-HDS purposes, one method being selection by serial passaging, and the other method being genetic engineering of a bacteriophage, so that the modified bacteriophage will remain active in the body for longer periods of time than the wild-type phage. [portfolio: Infectious Diseases—Therapeutics]

Dated: October 4, 1995.

Barbara M. McGarey,  
*Deputy Director, Office of Technology Transfer.*

[FR Doc. 95-25730 Filed 10-16-95; 8:45 am]

**BILLING CODE 4140-01-P**

### Notice of Meeting

Notice is hereby given of the meeting of the NIH AIDS Research Program Evaluation Working Group Area Review Panel on Clinical Trials on November 13-14, 1995 at the Chevy Chase Holiday Inn, 5520 Wisconsin Avenue, Chevy Chase, Maryland. The meeting will be

open to the public from 8:00 a.m. to 12:00 p.m. on November 13, and the closed portions will be from 1:00 p.m. to 5:30 p.m. on November 13, and 8:30 a.m. to 3:00 p.m. on November 14.

The NIH Revitalization Act of 1993 authorizes the Office of AIDS Research (OAR) to evaluate the AIDS research activities of NIH. The NIH AIDS Research Program Evaluation Working Group was established by the OAR to carry out this major evaluation initiative, reviewing and assessing each of the components of the NIH AIDS research endeavor to determine whether those components are appropriately designed and coordinated to answer the critical scientific questions to lead to better treatments, preventions, and a cure for AIDS. Six Area Review Panels were also established to address the following research areas: Natural History and Epidemiology; Etiology and Pathogenesis; Clinical Trials; Drug Discovery; Vaccines; and Behavioral and Social Sciences Research.

The purpose of the meeting is to seek input from individuals and organizations interested in the evaluation of AIDS research in the areas of therapeutics research as it pertains to clinical trials. Examples of areas under consideration by the panel include the effectiveness, efficiency, scientific productivity and clinical impact of NIH clinical trials programs in adults and children for HIV and its sequelae. This includes trials aimed at improving outcomes by limiting HIV replication and enhancing immune function as well as studies aimed at preventing and treating the complications of advanced HIV disease. The NIH AIDS Research Program Evaluation Working Group will develop recommendations to be made to the Office of AIDS Research Advisory Council that address the overall NIH AIDS research initiatives, both intramural and extramural, and identify long-range goals in the relevant areas of science. These recommendations will provide the framework for future planning and budget development of the NIH AIDS research program.

There will be a closed session from 1:00 p.m. to 5:30 p.m. on November 13 and 8:30 a.m. to 3:00 p.m. on November 14, to update the Panel members on privileged information on institute and center grant and contract portfolios.

The open session from 8:00 a.m. to 12:00 p.m. will begin with a brief overview of panel activities by members of the panel. The remainder of the meeting will be devoted to presentations from individuals and organizations. The session is open to the public; however, attendance may be limited by seat availability.

Comments should be confined to statements related to the current status of NIH AIDS research in the areas of therapeutic clinical trials and recommendations for consideration by the panel in assessing and reviewing the relevant research in these areas.

Only one representative of an organization may present oral comments. Each speaker will be permitted 5 minutes for their presentation. Interested individuals and representatives of organizations must submit a letter of intent to present comments and three (3) typewritten copies of the presentation, along with a brief description of the organization represented, to the attention of Dr. Judith Feinberg, Office of AIDS Research, NIH, 31 Center Drive, MSC 2340, Building 31, Room 5C08, Bethesda, MD 20892-2340, (301) 496-0358, FAX: (301) 402-8638. Letters of intent and copies of presentations must be received no later than 5:00 p.m. on Monday, October 30.

Any person attending the meeting who does not request an opportunity to speak in advance of the meeting will be allowed to make a brief oral presentation at the conclusion of the meeting, if time permits, and at the discretion of the Chairperson.

Individuals wishing to provide only written statements should send three (3) typewritten copies of their comments, including a brief description of their organization, to the above address no later than 5 p.m. on October 30. Statements submitted after that date will be accepted. They may not, however, be made available to the Area Review Panel prior to the meeting, though they will be provided subsequently as written testimony.

Individuals who plan to attend and need special assistance, such as sign language interpretation or other reasonable accommodations, should contact Dr. Feinberg in advance of the meeting.

Dated: October 11, 1995.

Susan K. Feldman,

*Committee Management Officer, NIH.*

[FR Doc. 95-25729 Filed 10-16-95; 8:45 am]

**BILLING CODE 4140-01-M**

### National Center for Research Resources; Notice of Closed Meeting

Pursuant to Section 10(d) of the Federal Advisory Committee Act, as amended (5 U.S.C. Appendix 2), notice is hereby given of the following National Center for Research Resources Special Emphasis Panel (SEP) meeting: