

Active duty service members are authorized civilian medical care under 10 U.S.C. 1074(c), and may be referred for unproven therapy when controlled in a formal clinical research trial. The trial must be operating under the structure of an institutional review board process, which conforms to the requirements of DoD Directive 3216.2, Protection of Human Subjects and Adherence of Ethical Standards in DoD Supported Research, the requirements of 32 CFR part 219, Protection of Human Subjects, as well as Service specific human experimentation regulations.

DoD has the authority to waive the statutory limitation for all other DoD beneficiaries that health care services must be medically necessary, appropriate, and proven care, as long as these services are provided within the context of an interagency agreement with the National Institutes of Health (NIH) for beneficiary participation in NIH-sponsored or approved clinical trials. The Secretary of Defense must also determine that such waiver will promote access by covered beneficiaries to promising new treatments and contribute to the development of such treatments.

B. Caseload, Costs

Each year approximately 60,000 TRICARE births occur at the Military Treatment Facilities (MTFs). Approximately 40,000 TRICARE births occur in civilian hospitals. According to the Center of Disease Control, in 2001 there were 20.09 cases of spina bifida per 100,000 births. Based on various studies, we estimate that 95 percent of these reported cases are related to myelomeningocele. We then interpret that approximately 19 cases would occur annually in TRICARE. We expect six to sixteen TRICARE members each year would have a fetus with a prenatal diagnosis of spina bifida that would be eligible for the NICHD clinical trial and would agree to participate.

Treatment protocol costs are estimated between \$300,000 and \$1.3 million over Fiscal Years 2003 through 2006 for TRICARE participation in the NICHD clinical trial of myelomeningocele fetal repair.

C. Operation of the Demonstration

The National Institute of Child Health and Human Development (NICHD) will fund an unblinded randomized controlled clinical trial conducted by three participating centers. The NICHD will provide administrative support, all NICHD enrollments, and study monitoring activities. DoD will provide a Project Officer who will coordinate DoD activities.

The DoD will develop initiatives to educate military healthcare providers and civilian TRICARE network providers about this initiative and the processes that are available for referral and pre-authorization of individuals with affected fetuses. DoD will require pre-authorization for any clinical services necessary or resultant from participation in an NICHD sponsored clinical trial before reimbursement by TRICARE. A pre-authorization for enrollment in the trial will suffice to cover each incidental expense or claim related to participation in the clinical trial extending through the duration of the clinical trial. The pre-authorization process will include verification with the NICHD that the patient has been enrolled in the study.

The TRICARE contractor(s) would not be involved in clinical issues or in directing patients to a particular institution.

D. Requirements for Participation

Active duty members, former members, and their dependents eligible for TRICARE who meet the clinical trial protocol would be eligible to participate in the demonstration. NICHD anticipates a total of two hundred patients whose fetuses have been diagnosed with myelomeningocele at 16 to 25 weeks' gestation who are over the age of 18 years would be enrolled and referred to the Data and Study Coordinating Center (DSCC) at George Washington University in Rockville, Maryland, to undergo an initial evaluation. Those individuals who remain eligible and interested would be assigned by the DSCC to one of the three centers (Vanderbilt University medical Center in Nashville, the University of California at San Francisco, and Children's Hospital of Philadelphia) where final evaluation and screening will be performed. Patient selection to the three Management of Myelomeningocele Study (MOMS) Centers would be based on convenience to the individual as well as the need to divide evenly the participants among the three centers.

E. Costs

Patients who choose to participate in the clinical trial will have no additional costs for prenatal care beyond what is normally paid by the beneficiary. If TRICARE beneficiaries have other health insurance, the other health insurance is required to pay first before TRICARE to the extent the health care is a benefit under the other plan as stated under 10 U.S.C. 1079(j)(1). If patients are in a prenatal surgery group, the travel, meal and lodging costs for the

patient and a relative or friend will be covered by NICHD grant support or the grantee institution until delivery and after delivery, until the patient and baby go home.

If patients are in a postnatal surgery group, travel back to the center for the patient and a support person will be covered by NICHD grant support or the grantee institution, as well as meals and lodging before and after delivery, until the baby and patient are able to go home. The cost of the study follow up, returning at one year and two and a half years of age will also be covered. Meals and lodging will be covered for those visits as well.

Dated: February 5, 2003.

Patricia L. Toppings,

Alternate OSD Federal Register Liaison Officer, Department of Defense.

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DEPARTMENT OF DEFENSE

Department of the Air Force

HQ USAF Scientific Advisory Board

AGENCY: Department of the Air Force, DoD.

ACTION: Notice of meeting.

SUMMARY: Pursuant to Pub. L. 92-463, notice is hereby given of the forthcoming meeting of the 2003 S&T Review and the Chief of Staff and Secretary of the Air Force. The purpose of the meeting is to allow the SAB leadership to advise the Air Force leadership on the outcome of the 2003 Review. Because classified and contractor-proprietary information will be discussed, this meeting will be closed to the public.

DATES: February 20, 2003.

ADDRESSES: Room 4E869, The Pentagon.

FOR FURTHER INFORMATION CONTACT: Maj John Pernot, Air Force Scientific Advisory Board Secretariat, 1180 Air Force Pentagon, Rm 5D982, Washington DC 20330-1180, (703) 697-4811.

Pamela D. Fitzgerald,

Air Force Federal Register Liaison Officer.

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DEPARTMENT OF DEFENSE

Department of the Air Force

HQ USAF Scientific Advisory Board

AGENCY: Department of the Air Force, DoD.

ACTION: Notice of meeting.

SUMMARY: Pursuant to Pub. L. 92-463, notice is hereby given of the forthcoming meeting of the 2003 S&T Review and the leadership of the Air Force Materiel Command. The purpose of the meeting is to allow the SAB leadership to advise the commander of the AFMC on the outcome of the 2003 Review. Because classified and contractor-proprietary information will be discussed, this meeting will be closed to the public.

DATES: February 18, 2003.

ADDRESSES: Room 4E987, The Pentagon.
FOR FURTHER INFORMATION CONTACT: Maj John Pernot, Air Force Scientific Advisory Board Secretariat, 1180 Air Force Pentagon, Rm 5D982, Washington DC 20330-1180, (703) 697-4811.

Pamela D. Fitzgerald,
Air Force Federal Register Liaison Officer.
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DEPARTMENT OF DEFENSE**Department of the Army; Corps of Engineers**

Intent To Prepare a Joint Environmental Impact Statement/Environmental Impact Report for the Dredged Material Management Plan Feasibility Study, Los Angeles County, CA

AGENCY: Department of the Army, Army Corps of Engineers, DoD.

ACTION: Notice of intent.

SUMMARY: The U.S. Army Corps of Engineers (Corps), the Port of Los Angeles, the City of Long Beach, and the County of Los Angeles propose to evaluate options for managing dredge materials in Los Angeles County.

DATES: A scoping meeting will be held on February 26, 2003, from 6:30 to 8:30 p.m. at the Cesar Chavez Community Center located at 401 Golden Avenue in Long Beach, CA.

FOR FURTHER INFORMATION CONTACT: Questions regarding the scoping process or preparation of the Environmental Impact Statement/Environmental Impact Report (EIS/EIR) may be directed to Mr. Paul Rose, Chief, Environmental Resources Branch, U.S. Army Corps of Engineers, P.O. Box 532711, Los Angeles, CA, 90053-2325, (213) 452-3840.

SUPPLEMENTARY INFORMATION:

1. *Proposed Action:* The Corps estimates that, within the Los Angeles Region, approximately 5 million cubic

meters (m³) of sediments deemed suitable for ocean disposal and 2.5 million m³ of sediments deemed unsuitable for ocean disposal will need to be dredged from the Ports and Harbors of Los Angeles County over the next 10 years. However, there is currently a lack of readily available disposal options for the unsuitable sediments. The U.S. Army Corps of Engineers, L.A. District has determined that there is a federal interest to participate in a detailed feasibility study to develop a regional dredged material management plan.

2. *Alternatives:* Alternatives that may be considered include: no action, upland disposal, aquatic capping, near shore confined disposal facility (California Department of Fish and Game), shallow water habitat creation, chemical stabilization, washing, blending, physical separation, thermal desorption, construction fill, landfill daily cover, reclamation fill, oil well injections, and geotextile encapsulation. These alternatives, and any additional, reasonable, alternatives recommended through the public scoping process, will be evaluated by the Corps of Engineers.

3. *Scoping Process:* The Corps, the Port of Los Angeles, the City of Long Beach, and the County of Los Angeles are preparing a joint EIS/EIR to address potential impacts associated with the proposed project. The Corps is the lead Federal agency for compliance with National Environmental Policy Act (NEPA) for the project, and the County of Los Angeles is the Lead State Agency for compliance with the California Environmental Quality Act (CEQA) for the non-Federal aspects of the project. The draft EIS/EIR (DEIS/EIR) document will incorporate public concerns in the analysis of impacts associated with the proposed action and associated project alternatives. The DEIS/EIR will be sent out for a 45-day public review period, during which time both written and verbal comments will be solicited on the adequacy of the document. The final EIS/EIR (FEIS/EIR) will address the comments received on the DEIS/EIR during public review, and will be furnished to all who commented on the DEIS/EIR, and is made available to anyone that requests a copy during the 30-day public comment period. The final steps involves, for the Federal EIS, preparing a record of decision (ROD) and, for the State EIR, certifying the EIR and adopting a mitigation monitoring and reporting plan. The ROD is a concise summary of the decisions made by the Corps from among the alternatives presented in the FEIS/EIR.

The ROD can be published immediately after the FEIS public

comment period ends. A certified EIR indicates that the environmental document adequately assesses the environmental impacts of the proposed project with respect to CEQA. A formal scoping meeting to solicit public comment and concerns on the proposed action and alternatives will be held on Wednesday, February 26, 2003 (see **DATES**).

Dated: January 31, 2003.

Richard G. Thompson,
Colonel, U.S. Army, District Engineer.

[FR Doc. 03-3587 Filed 2-12-03; 8:45 am]

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DEPARTMENT OF ENERGY

[Docket No. IC03-423-000, FERC Form No. 423]

Federal Energy Regulatory Commission

Commission Collection Activities, Proposed Collection; Comment Request; Extension

February 6, 2003.

AGENCY: Federal Energy Regulatory Commission.

ACTION: Notice.

SUMMARY: In compliance with the requirements of section 3506(c)(2)(a) of the Paperwork Reduction Act of 1995 (Pub. L. No. 104-13), the Federal Energy Regulatory Commission (Commission) is soliciting public comment on the specific aspects of the information collection described below.

DATES: Comments on the collection of information are due by April 9, 2003.

ADDRESSES: Copies of the proposed collection of information can be obtained from Michael Miller, Office of the Executive Director, ED-30, 888 First Street NE., Washington, DC 20426. Comments on the proposed collection of information may be filed either in paper format or electronically. Those parties filing electronically do not need to make a paper filing. For paper filings, the original and 14 copies of such comments should be submitted to the Office of the Secretary, Federal Energy Regulatory Commission, 888 First Street NE., Washington, DC 20426 and should refer to Docket No. 03-423-000.

Documents filed electronically via the Internet must be prepared in WordPerfect, MS Word, Portable Document Format, or ASCII format. To file the document, access the Commission's Web site at <http://www.ferc.gov> and click on "Make an E-filing," and then follow the instructions