in reducing selected risk factors or enhancing protective factors; and (2) document the process of service delivery and program implementation. Data will be collected from both program participants and comparison group youth at four points in time over a 4-year period. The annual burden estimates are as follows:

	Number of re- spondents	Number of responses per respondent	Average burden/re- sponse
Demonstration Project Staff	245	.75	1.1 hours.
	11,000	1.0	1.0 hour.

Dated: July 20, 1995.

Richard Kopanda,

Acting Executive Officer, SAMHSA.
[FR Doc. 95–18324 Filed 7–25–95; 8:45 am]
BILLING CODE 4162–20–P

Food and Drug Administration

Muscle Monitoring Devices; Decision Not to Rely on Dental Products Panel Recommendations

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing that it does not intend to rely on recommendations arising out of the October 13 and 14, 1994, meeting of the Dental Products Panel of the Medical **Devices Advisory Committee** concerning the classification of muscle monitoring devices. It is FDA's view that the October 1994 meeting was flawed and should not be the basis for decisions made about the use of these devices. FDA plans to fully and comprehensively consider the classification of muscle monitoring devices at a future meeting of the Dental Products Panel of the Medical Devices Advisory Committee.

FOR FURTHER INFORMATION CONTACT: Nancy J. Pluhowski, Center for Devices and Radiological Health (HFZ–400), Food and Drug Administration, 9200 Corporate Blvd., Rockville, MD 20850, 301–594–2022.

SUPPLEMENTARY INFORMATION: In the Federal Register of September 19, 1994 (59 FR 47880 at 47881), FDA announced that a meeting of the Dental Products Panel of the Medical Devices Advisory Committee would be held on October 13 and 14, 1994, to consider the classification of muscle monitoring devices. Because of substantive and procedural issues in connection with the October 1994 meeting, including the scope of products included and concerns that all interested parties may not have received adequate notice of the devices to be discussed at the meeting, FDA does not intend to rely on the

Panel's discussion or recommendations for the use or classification of these devices. In addition, it is FDA's view that the October 1994 meeting should not be the basis for decisions about the use of these devices. FDA plans a full and comprehensive consideration of muscle monitoring devices at a future meeting of the Dental Products Panel.

The Dental Products Panel of the Medical Devices Advisory Committee meeting tentatively scheduled for August 8, 9, and 10, 1995, which was announced in the **Federal Register** on March 9, 1995 (60 FR 12960 at 12962), will not include discussion of muscle monitoring devices. FDA will announce future meetings of the Dental Products Panel of the Medical Devices Advisory Committee in the **Federal Register** at least 15 days in advance of the upcoming meetings.

Dated: July 21, 1995.

William B. Schultz,

Deputy Commissioner for Policy.
[FR Doc. 95–18450 Filed 7–24–95; 11:46 am]
BILLING CODE 4160–01–F

DEPARTMENT OF HOUSING AND URBAN DEVELOPMENT

Office of the Assistant Secretary for Fair Housing and Equal Opportunity

[Docket Nos. FR-3622-N-07 and FR-3878-N-03]

Announcement of Funding Awards Fair Housing Initiatives Program FY 1994

AGENCY: Office of the Assistant Secretary for Fair Housing and Equal Opportunity, HUD.

ACTION: Announcement of funding awards.

SUMMARY: In accordance with section 102(a)(4)(C) of the Department of Housing and Urban Development Reform Act of 1989, this document notifies the public of FY 1994 funding awards made under the Fair Housing Initiatives Program (FHIP). The purpose of this document is to announce the names and addresses of the award winners and the amount of the awards

to be used to strengthen the Department's enforcement of the Fair Housing Act and to further fair housing. FOR FURTHER INFORMATION CONTACT: Maxine Cunningham, Director, Office of Fair Housing Initiatives and Voluntary Programs, Room 5234, 451 Seventh Street, S.W., Washington, D.C. 20410-2000. Telephone number (202) 708-0800. A telecommunications device (TDD) for hearing and speech impaired persons is available at (202) 708–3216. (These are not toll-free numbers.) SUPPLEMENTARY INFORMATION: Title VIII of the Civil Rights Act of 1968, as amended, 42 U.S.C. 3601-19 (The Fair Housing Act), charges the Secretary of Housing and Urban Development with responsibility to accept and investigate complaints alleging discrimination based on race, color, religion, sex, handicap, familial status or national origin in the sale, rental, or financing of most housing. In addition, the Fair Housing Act directs the Secretary to coordinate with State and local agencies administering fair housing laws and to cooperate with and render technical assistance to public or private entities carrying out programs to prevent and eliminate discriminatory housing practices.

Section 561 of the Housing and Community Development Act of 1987, 42 U.S.C. 3616 note, established the FHIP to strengthen the Department's enforcement of the Fair Housing Act and to further fair housing. This program assists projects and activities designed to enhance compliance with the Fair Housing Act and substantially equivalent State and local fair housing laws. Implementing regulations are found at 24 CFR Part 125.

The FHIP has four funding categories: The Administrative Enforcement Initiative, the Education and Outreach Initiative, the Private Enforcement Initiative, and the Fair Housing Organizations Initiative.

In the FY 1995 FHIP Notice of Funding Availability (NOFA) published in the **Federal Register** on April 11, 1995 (60 FR 18444), the Department announced the availability of up to \$1,457,446 for funding of FY 1994 awards. This Notice announces the