

The Office of Population Affairs is reviewing the family planning services delivery improvement research priorities and may revise those priorities. Accordingly, the November 18, 1985 standing announcement is hereby withdrawn.

**FOR FURTHER INFORMATION CONTACT:** Eugenia Eckard, (301) 594-6534.

Dated: April 12, 1999.

**Denese O. Shervington,**  
Deputy Assistant Secretary for Population Affairs.

[FR Doc. 99-10305 Filed 4-23-99; 8:45 am]

BILLING CODE 4160-17-M

**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**Administration for Children and Families**

**Submission for OMB Review; Comment Request**

*Title:* Family Preservation and Family Support (FP/FS) Services Implementation Study—State Level Data Collection.

*OMB:* 0970-0137.

*Description:* The Omnibus Budget Reconciliation Act of 1993 (OBRA 93)

established title IV-B, subpart 2 of the Social Security Act (42 U.S.C. 62-628) to provide funds to states for the development of family preservation and family support programs and services. Subpart 2, Section 435 of OBRA 93 requires the Secretary of HHS to evaluate the effectiveness of programs carried out under the legislation. The Adoption and Safe Families Act of 1997, P.L. 105-89, reauthorized the family preservation and family support programs and services and amended Section 431 [42 U.S.C. 629a] to add two new services: Time-Limited Family Reunification Services and Adoption Promotion and Support Services.

In this second phase of data collection, the five data collection instruments, which were used during the previous phase (1996-1999) will be used. Each instrument is geared toward obtaining information from individuals/agencies who will have a slightly different perspective on the context, planning, and implementation of the legislation. The data collection instruments will seek information on the programs and services funded, the goals of the planning process, populations targeted, reform efforts initiated, the relationship between

family preservation, family support and child welfare, staffing and training, information systems.

Data collection on states' planning and implementation experiences will be accomplished through semi-structured interviews with state officials and other key stakeholders who are knowledgeable about child welfare.

Both qualitative and quantitative analyses will be completed to highlight the process states employ to implement the legislation, coordinate with other funding sources, develop new systems, and improve service delivery systems. Data analyses also will focus on the impact of legislative changes on the state implementation of the program and comparisons of state implementation before and after the legislative reauthorization. Information obtained from data analyses will provide feedback to ACF in the determination of future policy guidance and the scope and nature of technical assistance to be provided to states. The information will also provide direct feedback to states concerning successful implementation strategies.

*Respondents:* State, Local or Tribal Government and Not-for-Profit Institutions.

ANNUAL BURDEN ESTIMATES

Instrument	Number of respondents	Number of responses per respondent	Average burden house per response	Total burden hours
State Level Data Collection .....	150	1	.0849	127.40

*Estimated Total Annual Burden Hours:* 127.40.

*Additional Information:* Copies of the proposed collection may be obtained by writing to the Administration for Children and Families, Office of Information Services, 370 L'Enfant Promenade, S.W., Washington, D.C. 20447, Attn: ACF Reports Clearance Officer.

*OMB Comment:* OMB is required to make a decision concerning the collection of information between 30 to 60 days after publication of this document in the **Federal Register**. Therefore, a comment is best assured of having its full effect if OMB receives it within 30 days of publication. Written comments and recommendations for the proposed information collection should be sent directly to the following: Office of Management and Budget Paperwork Reduction Project, 725 17th Street, N.W., Washington, D.C., 20503, Attn: Ms. Lori Schack.

Dated: April 19, 1999.

**Bob Sargis,**  
Acting Reports Clearance Officer.  
[FR Doc. 99-10304 Filed 4-23-99; 8:45 am]  
BILLING CODE 4184-01-M

**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**Food and Drug Administration**  
[Docket No. 99N-0672]

**Iatric Corp.; Revocation of U.S. License No. 0416**

**AGENCY:** Food and Drug Administration, HHS.  
**ACTION:** Notice.

**SUMMARY:** The Food and Drug Administration (FDA) is announcing the revocation of the establishment license (U.S. License No. 0416) and the product license issued to Iatric Corp. for the manufacture of Allergenic Extracts. In letters to FDA dated June 26 and June

30, 1998, the firm voluntarily requested revocation of its establishment and product licenses. In a letter dated August 28, 1998, FDA informed the firm that the establishment and product licenses were revoked.

**DATES:** The revocation of the establishment license (U.S. License No. 0416) and the product license became effective August 28, 1998.

**FOR FURTHER INFORMATION CONTACT:** Astrid L. Szeto, Center for Biologics Evaluation and Research (HFM-17), Food and Drug Administration, 1401 Rockville Pike, Rockville, MD 20852-1448, 301-827-6210.

**SUPPLEMENTARY INFORMATION:** FDA has revoked the establishment license (U.S. License No. 0416) and the product license for the manufacture of Allergenic Extracts issued to Iatric Corp., 2330 South Industrial Park Dr., Tempe, AZ 85282.

FDA inspected Iatric Corp. from April 7 through April 11, 1997. The inspection of the facility revealed