subsequent 60 day notices for later components of the evaluation. The first phase includes data collection for a cross-site implementation study and site-specific impact evaluations in two PIT grantee sites (Kansas; Washoe County, Nevada) that will begin implementing interventions during the second year of the PII grant period. The second phase includes a cross-site cost evaluation and site-specific impact evaluations in four PII grantee sites

expected to implement interventions in the third year of the PII grant period.

Data for the cross-site implementation study will be collected through: (1) Interviews with grantee staff and key informants conducted by telephone and during site visits; (2) web-based instruments completed by grantee staff and key informants; and (3) retrieval and submission of aggregate data from grantee data systems. Data for the Kansas impact evaluation will be collected through (1) family

assessments; (2) caseworkers' clinical assessments of children and families; and (3) caseworker discussions. Data for the Washoe County impact evaluation will be collected through family assessments.

Respondents: Families (parents, or permanent or foster caregivers; children), caseworkers, supervisors, service providers, and key informants such as grantee project directors, data managers, and representatives of partner agencies.

ANNUAL BURDEN ESTIMATES

Instrument	Annual number of re- spondents	Number of responses per respondent	Average burden hours per response	Total annual burden hours
CROSS-SITE IMPLEMENTATION STUDY:				
Survey of Organization/System Readiness	60	1	0.3	18
Implementation Drivers Web Survey	150	2	0.8	240
Grantee Case Study Protocol	30	4	2.0	240
Fidelity Data (Implementation Quotient Tracker)	2	8	1.5	24
Cross-Site Estimated Total	_	_	_	522
KANSAS:				
Caregiver Initial Information Form	300	1	0.1	30
Family Assessment Battery	300	3	1.5	1350
CAFAS/PECFAS	4	150	1.0	600
Caseworker Discussions for NCFAS–G&R Completion	4	150	0.5	300
Kansas Estimated Total	_	_	_	2280
WASHOE COUNTY:				
Family Assessment Battery	175	2	1.5	525
Washoe Estimated Total	_	_	_	525

Estimated Total Annual Burden Hours: 3327.

Additional Information: Copies of the proposed collection may be obtained by writing to the Administration for Children and Families, Office of Planning, Research and Evaluation, 370 L'Enfant Promenade SW., Washington, DC 20447, Attn: OPRE Reports Clearance Officer. All requests should be identified by the title of the information collection. Email address: OPREinfocollection@acf.hhs.gov.

OMB Comment: OMB is required to make a decision concerning the collection of information between 30 and 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, Email: OIRA_SUBMISSION@OMB.EOP.GOV, Attn: Desk Officer for the

Administration, for Children and Families.

Steven M. Hanmer,

OPRE Reports Clearance Officer. [FR Doc. 2012–10848 Filed 5–7–12; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration [Docket No. FDA-2012-N-0001]

Gastrointestinal Drugs Advisory Committee; Cancellation

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The meeting of the Gastrointestinal Drugs Advisory Committee scheduled for May 31, 2012, is canceled. This meeting was announced in the Federal Register of March 23, 2012 (77 FR 17078). The meeting is being canceled because the Agency no longer needs to discuss the issues that were originally under consideration in the review of the application. The sponsor of the new drug application (NDA) submitted new

information which negated the necessity for the planned meeting. The Agency intends to continue evaluating NDA 200–436 and, as needed, may schedule an Advisory Committee meeting in the future.

FOR FURTHER INFORMATION CONTACT:

Minh Doan, Center for Drug Evaluation and Research, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 31, rm. 2417, Silver Spring, MD 20993–0002, 301–796–9001, Fax: 301–847–8533, email: GIDAC@fda.hhs.gov, or FDA Advisory Committee Information Line, 1–800–741–8138 (301–443–0572 in the Washington, DC area), to find out further information regarding FDA advisory committee information or visit our Web site at http://www.fda.gov/AdvisoryCommittees/default.htm.

Dated: May 2, 2012.

Jill Hartzler Warner,

Acting Associate Commissioner for Special Medical Programs.

[FR Doc. 2012-10990 Filed 5-7-12; 8:45 am]

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