involved in the response. The number of samples tested during a response to a possible event could range from 10,000 to more than 500,000 samples depending on the length and breadth of the event. Since there is potentially a large range in the number of samples for a surge event, CDC estimates the annualized burden for this event will be 2,250,000 hours or 625 responses per respondent.

There is no cost to the respondents other than their time. The total estimated annualized burden is 2,382,300 hours.

### ESTIMATED ANNUALIZED BURDEN HOURS

<table>
<thead>
<tr>
<th>Type of respondents</th>
<th>Form name</th>
<th>Number of respondents</th>
<th>Number of responses per respondent</th>
<th>Avg. burden per response (in hrs)</th>
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</thead>
<tbody>
<tr>
<td>Public Health Laboratories</td>
<td>Biennial Requalification</td>
<td>150</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Public Health Laboratories</td>
<td>General Surveillance Testing Results</td>
<td>150</td>
<td>25</td>
<td>24</td>
</tr>
<tr>
<td>Public Health Laboratories</td>
<td>Proficiency Testing/Validation Testing Results</td>
<td>150</td>
<td>5</td>
<td>56</td>
</tr>
<tr>
<td>Public Health Laboratories</td>
<td>Surge Event Testing Results</td>
<td>150</td>
<td>625</td>
<td>24</td>
</tr>
</tbody>
</table>


Ron A. Otten,
Director, Office of Scientific Integrity (OSI),
Office of the Associate Director for Science (OADS), Office of the Director, Centers for Disease Control and Prevention.

[FR Doc. 2013–04900 Filed 3–1–13; 8:45 am]

### DEPARTMENT OF HEALTH AND HUMAN SERVICES

#### Administration for Children and Families

**Agency Recordkeeping/Reporting Requirements Under Emergency Review by the Office of Management and Budget (OMB)**

**Title:** TANF Quarterly Financial Report, ACF–196.

**OMB No.:** 0970–0247.

**Description:** This information collection is authorized under Section 411(a)(3) of the Social Security Act. This request is for revision of approval to use the Administration for Children and Families’ (ACF) 196 form for periodic financial reporting under the Temporary Assistance for Needy Families (TANF) program. States participating in the TANF program are required by statute to report financial data on a quarterly basis. This form meets the legal standard and provides essential data on the use of Federal funds. Failure to collect the data would seriously compromise ACF’s ability to monitor program expenditures, estimate funding needs, and to prepare budget submissions required by Congress. Financial reporting under the TANF program is governed by 45 CFR part 265. This renewal restores columns for reporting Emergency Contingency Fund and Supplemental Grant expenditures.

**Respondents:** TANF Agencies.

### ANNUAL BURDEN ESTIMATES

<table>
<thead>
<tr>
<th>Instrument</th>
<th>Number of respondents</th>
<th>Number of responses per respondent</th>
<th>Average burden hours per response</th>
<th>Total burden hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACF–196</td>
<td>51</td>
<td>4</td>
<td>10</td>
<td>2040</td>
</tr>
</tbody>
</table>

Estimated Total Annual Burden Hours: 2040.

### Additional Information

ACF is requesting that OMB grant a 180 day approval for this information collection under procedures for emergency processing by March 15, 2013. A copy of this information collection, with applicable supporting documentation, may be obtained by emailing the Administration for Children and Families, Reports Clearance Officer: rsargs@acf.hhs.gov.

Comments and questions about the information collection described above should be directed to the Office of Information and Regulatory Affairs, Attn: OMB Desk Officer for ACF, Office of Management and Budget, Paperwork Reduction Project; 725 17th Street NW., Washington, DC 20503; FAX: (202) 395–7285; email: oira_submission@omb.eop.gov.

Robert Sargsis,
Reports Clearance Officer.

[FR Doc. 2013–04826 Filed 3–1–13; 8:45 am]

### DEPARTMENT OF HEALTH AND HUMAN SERVICES

#### Food and Drug Administration

**[Docket No. FDA–2012–P–1071]**

**Determination That GEREF (Sermorelin Acetate) Injection, 0.5 Milligrams Base/Vial and 1.0 Milligrams Base/Vial, and GEREF (Sermorelin Acetate) Injection, 0.05 Milligrams Base/Amp, Were Not Withdrawn From Sale for Reasons of Safety or Effectiveness**

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

**ACTION:** Notice.

**SUMMARY:** The Food and Drug Administration (FDA) has determined that GEREF (Sermorelin Acetate)
injection, 0.5 milligrams (mg) base/vial and 1.0 mg base/vial, and GEREF (Sermorelin Acetate) injection, 0.05 mg base/amp, were not withdrawn from sale for reasons of safety or effectiveness. This determination will allow FDA to approve abbreviated new drug applications (ANDAs) for GEREF (Sermorelin Acetate) injection, 0.5 mg base/vial and 1.0 mg base/vial, and GEREF (Sermorelin Acetate) injection, 0.05 mg base/amp, if all other legal and regulatory requirements are met.

FOR FURTHER INFORMATION CONTACT: Kathy Schreier, Center for Drug Evaluation and Research, Food and Drug Administration, 10903 New Hampshire Ave, Bldg. 51, Rm. 6252, Silver Spring, MD 20993–0002, 301–796–3432.

SUPPLEMENTARY INFORMATION: In 1984, Congress enacted the Drug Price Competition and Patent Term Restoration Act of 1984 (Public Law 98–417) (the 1984 amendments), which authorized the approval of duplicate versions of drug products under an ANDA procedure. ANDA applicants must, with certain exceptions, show that the drug for which they are seeking approval contains the same active ingredient in the same strength and dosage form as the “listed drug,” which is a version of the drug that was previously approved. ANDA applicants do not have to repeat the extensive clinical testing otherwise necessary to gain approval of a new drug application (NDA). The only clinical data required in an ANDA are data to show that the drug that is the subject of the ANDA is bioequivalent to the listed drug.

The 1984 amendments include what is now section 505(j)(7) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 355(j)(7)), which requires FDA to publish a list of all approved drugs. FDA publishes this list as part of the “Approved Drug Products With Therapeutic Equivalence Evaluations,” which is known generally as the “Orange Book.” Under FDA regulations, drugs are removed from the list if the Agency withdraws or suspends approval of the drug’s NDA or ANDA for reasons of safety or effectiveness or if FDA determines that the listed drug was withdrawn from sale for reasons of safety or effectiveness (21 CFR 314.162).

A person may petition the Agency to determine, or the Agency may determine on its own initiative, whether a listed drug was withdrawn from sale for reasons of safety or effectiveness. This determination may be made at any time after the drug has been withdrawn from sale, but must be made prior to approving an ANDA that refers to the listed drug § 314.161 (21 CFR 314.161). FDA may not approve an ANDA that does not refer to a listed drug.

GEREF (Sermorelin Acetate) injection, 0.5 mg base/vial and 1.0 mg base/vial, is the subject of NDA 20–443, held by EMD Serono, and initially approved on September 26, 1997; and GEREF (Sermorelin Acetate) injection, 0.05 mg base/amp, is the subject of NDA 19–863, held by EMD Serono, and initially approved on December 28, 1990. GEREF (Sermorelin Acetate) injection, 0.5 mg base/vial and 1.0 mg base/vial, is indicated for the treatment of idiopathic growth hormone deficiency (GHD) in children with growth failure, and GEREF (Sermorelin Acetate) injection, 0.05 mg base/amp, is indicated for evaluating the ability of the somatotroph of the pituitary gland to secrete growth hormone.

In a letter dated December 2, 2008, EMD Serono notified FDA that GEREF (Sermorelin Acetate) injection, 0.5 mg base/vial and 1.0 mg base/vial, was being discontinued and requested withdrawal of NDA 20–443; and FDA moved that drug product to the “Discontinued Drug Product List” section of the Orange Book. In a letter dated July 11, 2008, EMD Serono also notified FDA that GEREF (Sermorelin Acetate) injection, 0.05 mg base/amp, was being discontinued, and FDA moved the drug product to the “Discontinued Drug Product List” section of the Orange Book as well. In addition, in a letter dated December 12, 2008, EMD Serono requested withdrawal of NDA 19–863 for GEREF (Sermorelin Acetate) injection, 0.05 mg base/amp. In the Federal Register of May 19, 2009 (74 FR 23407), FDA announced that it was withdrawing approval of NDA 19–863 and NDA 20–443, effective June 18, 2009.

Alvin J. Lorman submitted a citizen petition dated October 12, 2012 (Docket No. FDA–2012–P–1071), under 21 CFR 10.30, requesting that the Agency determine whether GEREF (Sermorelin Acetate) injection, 0.5 mg base/vial and 1.0 mg base/vial, was withdrawn from the market for reasons of safety and efficacy. Although the citizen petition did not request that we determine whether GEREF (Sermorelin Acetate) injection, 0.05 mg base/amp, approved under NDA 19–863, was withdrawn for safety or efficacy, that product has also been discontinued. On our own initiative, we have also determined whether GEREF (Sermorelin Acetate) injection, 0.05 mg base/amp, was withdrawn for safety or effectiveness reasons.

After considering the citizen petition and reviewing Agency records and based on the information we have at this time, FDA has determined under § 314.161 that GEREF (Sermorelin Acetate) injection, 0.5 mg base/vial and 1.0 mg base/vial, and GEREF (Sermorelin Acetate) injection, 0.05 mg base/amp, were not withdrawn for reasons of safety or effectiveness. The petitioner has identified no data or other information suggesting that GEREF (Sermorelin Acetate) injection, 0.5 mg base/vial and 1.0 mg base/vial, and GEREF (Sermorelin Acetate) injection, 0.05 mg base/amp, were withdrawn for reasons of safety or effectiveness. We have carefully reviewed our files for records concerning the withdrawal of GEREF (Sermorelin Acetate) injection, 0.5 mg base/vial and 1.0 mg base/vial, and GEREF (Sermorelin Acetate) injection, 0.05 mg base/amp, from sale. We have also independently evaluated relevant literature and data for possible postmarketing adverse events for both GEREF products. We have reviewed the available evidence and determined that both GEREF products were not withdrawn from sale for reasons of safety or effectiveness.

Accordingly, the Agency will continue to list GEREF (Sermorelin Acetate) injection, 0.5 mg base/vial and 1.0 mg base/vial, and GEREF (Sermorelin Acetate) injection, 0.05 mg base/amp, in the “Discontinued Drug Product List” section of the Orange Book. The “Discontinued Drug Product List” delineates, among other items, drug products that have been discontinued from marketing for reasons other than safety or effectiveness. ANDAs that refer to GEREF (Sermorelin Acetate) injection, 0.5 mg base/vial and 1.0 mg base/vial, and GEREF (Sermorelin Acetate) injection, 0.05 mg base/amp, may be approved by the Agency as long as they meet all other legal and regulatory requirements for the approval of ANDAs. If FDA determines that labeling for this drug product should be revised to meet current standards, the Agency will advise ANDA applicants to submit such labeling.


Leslie Kux,
Assistant Commissioner for Policy.

[FR Doc. 2013–04827 Filed 3–1–13; 8:45 am]

BILLING CODE 4160–01–P