of FDA’s functions, including whether the information will have practical utility; (2) the accuracy of FDA’s estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used; (3) ways to enhance the quality, utility, and clarity of the information to be collected; and (4) ways to minimize the burden of the collection of information on respondents, including through the use of automated collection techniques, when appropriate, and other forms of information technology.

Medical Device Recall Authority—21 CFR Part 810 (OMB Control Number 0910–0432)—Extension

This collection of information implements section 518(e) of the Federal Food, Drug, and Cosmetic Act (FD&C Act) (21 U.S.C. 360h(e)) and part 810 (21 CFR part 810), medical device recall authority provisions. Section 518(e) of the FD&C Act provides FDA with the authority to issue an order requiring an appropriate person, including manufacturers, importers, distributors, and retailers of a device, if FDA finds that there is reasonable probability that the device intended for human use would cause serious adverse health consequences or death, to: (1) Immediately cease distribution of such device, (2) immediately notify health professionals and device-user facilities of the order, and (3) instruct such professionals and facilities to cease use of such device.

Further, the provisions under section 518(e) of the FD&C Act set out the following three-step procedure for issuance of a mandatory device recall order:

1. If there is a reasonable probability that a device intended for human use would cause serious adverse health consequences or death, FDA may issue a cease distribution and notification order requiring the appropriate person to immediately:
   - Cease distribution of the device,
   - Notify health professionals and device user facilities of the order, and
   - Instruct those professionals and facilities to cease use of the device;

2. FDA will provide the person named in the cease distribution and notification order with the opportunity for an informal hearing on whether the order should be modified, vacated, or amended to require a mandatory recall of the device; and

3. After providing the opportunity for an informal hearing, FDA may issue a mandatory recall order if the Agency determines that such an order is necessary.

The information collected under the recall authority provisions will be used by FDA to do the following: (1) Ensure that all devices entering the market are safe and effective, (2) accurately and immediately detect serious problems with medical devices, and (3) remove dangerous and defective devices from the market.

FDA estimates the burden of this collection of information as follows:

### TABLE 1—ESTIMATED ANNUAL REPORTING BURDEN 1

<table>
<thead>
<tr>
<th>21 CFR Section</th>
<th>Number of respondents</th>
<th>No. of responses per respondent</th>
<th>Total annual responses</th>
<th>Average burden per response</th>
<th>Total hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>810.10(d)</td>
<td>2</td>
<td>1</td>
<td>2</td>
<td>8</td>
<td>16</td>
</tr>
<tr>
<td>810.11(a)</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>8</td>
<td>8</td>
</tr>
<tr>
<td>810.12(a) and (b)</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>8</td>
<td>8</td>
</tr>
<tr>
<td>810.14</td>
<td>2</td>
<td>1</td>
<td>2</td>
<td>16</td>
<td>32</td>
</tr>
<tr>
<td>810.15(a), (b), and (c)</td>
<td>2</td>
<td>1</td>
<td>2</td>
<td>12</td>
<td>24</td>
</tr>
<tr>
<td>810.15(d)</td>
<td>2</td>
<td>1</td>
<td>2</td>
<td>4</td>
<td>8</td>
</tr>
<tr>
<td>810.15(e)</td>
<td>10</td>
<td>1</td>
<td>10</td>
<td>1</td>
<td>10</td>
</tr>
<tr>
<td>810.16(a) and (b)</td>
<td>2</td>
<td>12</td>
<td>24</td>
<td>40</td>
<td>960</td>
</tr>
<tr>
<td>810.17(a)</td>
<td>2</td>
<td>1</td>
<td>2</td>
<td>8</td>
<td>16</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>1,082</td>
</tr>
</tbody>
</table>

1 There are no capital costs or operating and maintenance costs associated with this collection of information.

### TABLE 2—ESTIMATED ANNUAL RECORDKEEPING BURDEN 1

<table>
<thead>
<tr>
<th>21 CFR Section</th>
<th>No. of record-keepers</th>
<th>No. of records per recordkeeper</th>
<th>Total annual records</th>
<th>Average burden per recordkeeping</th>
<th>Total hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>810.15(b)</td>
<td>2</td>
<td>1</td>
<td>1</td>
<td>8</td>
<td>8</td>
</tr>
</tbody>
</table>

1 There are no capital costs or operating and maintenance costs associated with this collection of information.

**Explanation for Burden Estimates**

The burden estimates for tables 1 and 2 of this document are based on FDA’s experience with voluntary recalls under part 810 of the regulations. FDA expects no more than two mandatory recalls per year, as most recalls are done voluntarily. Since the last time this collection of information was submitted to OMB for renewal/approval, there has been one mandatory recall.

Dated: November 9, 2011.

Leslie Kux,

* Acting Assistant Commissioner for Policy.

**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**Food and Drug Administration**

[FR Doc. 2011–29512 Filed 11–15–11; 8:45 am]

BILLING CODE 4160–01–P

Scott S. Reuben: Debarment Order

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

**ACTION:** Notice.

**SUMMARY:** The Food and Drug Administration (FDA) is issuing an
order under the Federal Food, Drug, and Cosmetic Act permanently debarring Scott S. Reuben, M.D. from providing services in any capacity to a person that has an approved or pending drug product application. FDA bases this order on a finding that Dr. Reuben was convicted of a felony under Federal law for conduct relating to the regulation of a drug product under the Federal Food, Drug, and Cosmetic Act. Dr. Reuben was given notice of the proposed permanent debarment and an opportunity to request a hearing within the timeframe prescribed by regulation. Dr. Reuben failed to respond. Dr. Reuben’s failure to respond constitutes a waiver of his right to a hearing concerning this action.

DATES: This order is effective November 16, 2011.

ADDRESSES: Submit applications for special termination of debarment to the Division of Dockets Management (HFA–305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852.

FOR FURTHER INFORMATION CONTACT: Kenny Shade, Division of Compliance Policy (HFC–230), Office of Enforcement, Office of Regulatory Affairs, Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, (301) 796–4640.

SUPPLEMENTARY INFORMATION:

I. Background

Section 306(a)(2)(B) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 335a(a)(2)(B)) requires debarment of an individual if FDA finds that the individual has been convicted of a felony under Federal law for conduct relating to the regulation of any drug product under the Federal Food, Drug, and Cosmetic Act.

On June 24, 2010, the U.S. District Court for the District of Massachusetts entered judgment against Dr. Reuben for health care fraud in violation of 18 U.S.C. 1347.

The FDA’s finding that debarment is appropriate is based on the felony conviction referenced herein for conduct relating to the regulation of a drug product. The factual basis for this conviction is as follows: Dr. Reuben was a physician licensed by the State of Massachusetts working as an anesthesiologist providing anesthesia services to patients in connection with surgeries, and also treating patients post-surgery in the District of Massachusetts. Dr. Reuben served as the chief of acute pain at a hospital in Western Massachusetts and maintained an office at the hospital for the purpose of conducting research. Dr. Reuben’s interest, from a research perspective, was in post-operative multimodal analgesia therapy. Dr. Reuben made proposals for research funding to pharmaceutical companies that manufactured drugs that he used or proposed to use in multimodal analgesia therapy. Dr. Reuben represented to the companies that, as the principal investigator, he would be performing clinical studies with actual patients to whom he would administer the drug that was the subject of the research grant.

Dr. Reuben entered into contracts to perform research studies funded by the companies from at least as early as 1999. Dr. Reuben purported to perform the research called for by the contracts, and published articles in various medical journals based on the purported results of the research, when in fact those studies had not been performed, and therefore the research results reported in the medical journals were false.

As a result of his convictions, on August 22, 2011, FDA sent Dr. Reuben a notice by certified mail proposing to permanently debar him from providing services in any capacity to a person that has an approved or pending drug product application. The proposal was based on a finding, under section 306(a)(2)(B) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 335a(a)(2)(B)), that Dr. Reuben was convicted of a felony under Federal law for conduct relating to the regulation of a drug product under the Federal Food, Drug, and Cosmetic Act. This conclusion was based on the fact that FDA regulates clinical trials related to drug products such as those described previously as part of the Agency’s regulation of drug products. The proposal also offered Dr. Reuben an opportunity to request a hearing, providing him 30 days from the date of receipt of the letter in which to file the request, and advised him that failure to request a hearing constituted a waiver of the opportunity for a hearing and of any contentions concerning this action. The proposal was received on August 26, 2011. Dr. Reuben failed to respond within the timeframe prescribed by regulation and has, therefore, waived his opportunity for a hearing and has waived any contentions concerning his debarment (21 CFR part 12).

II. Findings and Order

Therefore, the Director, Office of Enforcement, Office of Regulatory Affairs, under section 306(a)(2)(B) of the Federal Food, Drug, and Cosmetic Act, (21 U.S.C. 335a(a)(2)(B)) and 335a(d)(4) of the Federal Food, Drug, and Cosmetic Act, (21 U.S.C. 335a(d)(4)) should be identified with Docket No. FDA–2011–N–0377 and sent to the Division of Dockets Management (see ADDRESSES). All such submissions are to be filed in four copies. The public availability of information in these submissions is governed by 21 CFR 10.20(j).

Publicly available submissions may be seen in the Division of Dockets Management between 9 a.m. and 4 p.m., Monday through Friday.

Dated: November 7, 2011.

Armando Zamora,
Acting Director, Office of Enforcement, Office of Regulatory Affairs.