costs other than the amount of time required to respond to the survey.

#### ESTIMATED ANNUALIZED BURDEN HOURS

Data collection instrument	Number of respondents	Responses/ respondent	Average bur- den per re- sponse (in hrs)	Average an- nual burden hours
Satisfaction survey (callers)	25,000	1	3/60	1,250
Satisfaction survey (e-mail inquiries)	330	1	3/60	17
Follow up survey	3,125	1	7/60	365
Key informant survey	100	1	7/60	12
Postcard survey for bulk mailing	950	1	1/60	16
Postcard survey for individual publications	2,100	1	1/60	35
Web survey for e-mail publication orders	1,000	1	1/60	17
Web survey for internet publications	950	1	1/60	16
Special event/Outreach survey—General Public	25,600	1	5/60	2,133
Special event/Outreach survey—Professionals	10,400	1	5/60	867
Emergency response survey—Level 1 emergency—General Public	31,151	1	5/60	2596
Emergency response survey—Level 1 emergency—Professionals	7,459	1	5/60	622
Emergency response survey—Level 2 emergency—General Public	57,579	1	5/60	4798
Emergency response survey—Level 2 emergency—Professionals	51,821	1	5/60	4318
Emergency response survey—Level 3 emergency—General Public	351,863	1	5/60	29,322
Emergency response survey—Level 3 emergency—Professional	316,678	1	5/60	26,390
Emergency response survey—Level 4 emergency—General Public	645,630	1	5/60	53,803
Emergency response survey—Level 4 emergency—Professional	596,504	1	5/60	49,709
Total Burden Hours				176,286

Dated: February 6, 2007.

#### Joan F. Karr,

Acting Reports Clearance Officer, Centers for Disease Control and Prevention.

[FR Doc. E7–2637 Filed 2–14–07; 8:45 am] BILLING CODE 4163–18–P

## DEPARTMENT OF HEALTH AND HUMAN SERVICES

## **Food and Drug Administration**

[Docket No. 2006N-0430]

Agency Information Collection Activities; Submission for Office of Management and Budget Review; Comment Request; General Licensing Provisions: Biologics License Application, Changes to an Approved Application, Labeling, Revocation and Suspension, Postmarketing Studies Status Reports, and Forms FDA 356h and 2567; Correction

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

**ACTION:** Notice.

SUMMARY: The Food and Drug Administration (FDA) is correcting a notice that appeared in the Federal Register of February 2, 2007 (72 FR 5057). The document announced that an opportunity for public comment on a proposed collection of information had been submitted to the Office of Management and Budget for review and clearance under the Paperwork Reduction Act of 1995. The notice published with an error in titles referring to an FDA form number in two places in the document. This document corrects those errors.

DATES: February 15, 2007.

#### FOR FURTHER INFORMATION CONTACT:

Jonna Capezzuto, Office of the Chief Information Officer (HFA–250), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301–827– 4659.

**SUPPLEMENTARY INFORMATION:** In the **Federal Register** of Friday, February 2, 2007, the following corrections are made on page 5057:

- 1. In the first column, in the ninth line of the title of the document, the phrase "Forms FDA 456h" is corrected to read "Forms FDA 356h".
- 2. In the second column, in the **SUPPLEMENTARY INFORMATION** section of the document, in the sixth line of the title, the phrase "Forms FDA 456h" is corrected to read "Forms FDA 356h".

Dated: February 8, 2007.

#### Jeffrey Shuren,

Assistant Commissioner for Policy. [FR Doc. E7–2576 Filed 2–14–07; 8:45 am] BILLING CODE 4160–01–8

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

#### **Food and Drug Administration**

[Docket No. 2006N-0436]

Agency Information Collection Activities; Submission for Office of Management and Budget Review; Comment Request; Guidance for Industry on How To Use E-Mail To Submit a Study Protocol

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

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**ACTION:** Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing that a proposed collection of information has been submitted to the Office of Management and Budget (OMB) for review and clearance under the Paperwork Reduction Act of 1995.

**DATES:** Fax written comments on the collection of information by March 19, 2007.

ADDRESSES: To ensure that comments on the information collection are received, OMB recommends that written comments be faxed to the Office of Information and Regulatory Affairs, OMB, Attn: FDA Desk Officer, FAX: 202–395–6974.

### FOR FURTHER INFORMATION CONTACT:

Denver Presley, Jr., Office of the Chief Information Officer (HFA–250), Food and Drug Administration, 5600 Fishers