the following issues. The CMS–855A enrollment form currently captures ownership/managerial information on providers. The data required under sections 6401 and 6001, however, is more specific than that currently obtained on the CMS–855A. CMS will therefore create four attachments to the CMS–855A—two for SNFs and the other two for physician-owned hospitals—to secure this information. In addition to the application changes triggered by ACA, CMS is making other revisions to the forms as well.

This information collection request has been revised since the 60-day Federal Register notice published on March 22, 2011 (76 FR 13415). The group/clinic and individual burden has decreased due to the removal of a previously proposed supplier attachment. However, the overall burden hour estimate has increased slightly due to additional role-specific ownership and managerial control data collection for institutional providers. Form Number: CMS-855(A, B, I, R) (OMB#: 0938-0685); Frequency: Yearly; Affected Public: Private Sector; Business or other for-profit and not-for-profit institutions; Number of Respondents: 440,450; Total Annual Responses: 440,450; Total Annual Hours: 856,395. (For policy questions regarding this collection contact Kim McPhillips at 410-786-5374. For all other issues call 410-786-1326.)

To be assured consideration, comments and recommendations for the proposed information collections must be received by the OMB desk officer at the address below, no later than 5 p.m. on June 20, 2011. OMB, Office of Information and Regulatory Affairs, Attention: CMS Desk Officer, Fax Number: (202) 395–6974, E-mail: OIRA submission@omb.eop.gov.

Dated: May 17, 2011.

Martique Jones,

Director, Regulations Development Group, Division-B, Office of Strategic Operations and Regulatory Affairs.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

[Document Identifier: CMS-R-235]

Agency Information Collection Activities: Proposed Collection; Comment Request

AGENCY: Centers for Medicare & Medicaid Services, HHS.

In compliance with the requirement of section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995, the Centers for Medicare & Medicaid Services (CMS) is publishing the following summary of proposed collections for public comment. Interested persons are invited to send comments regarding this burden estimate or any other aspect of this collection of information, including any of the following subjects: (1) The necessity and utility of the proposed information collection for the proper performance of the agency's functions; (2) the accuracy of the estimated burden; (3) ways to enhance the quality, utility, and clarity of the information to be collected; and (4) the use of automated collection techniques or other forms of information technology to minimize the information collection hurden

1. Type of Information Collection Request: Revision of a currently approved collection; Title of Information Collection: Data Use Agreement (DUA) for Data Acquired from the Centers for Medicare & Medicaid Services (CMS); Use: The Privacy Act of 1976, § 552a requires the Centers for Medicare & Medicaid Services (CMS) to track all disclosures of the agency's Personally Identifiable Information (PII) and the exceptions for these data releases. CMS is also required by the Health Insurance Portability and Accountability Act (HIPAA) of 1996 and the Federal Information Security Management Act (FISMA) of 2002 to properly protect all PII data maintained by the agency. When entities request CMS PII data, they enter into a Data Use Agreement (DUA) with CMS. The DUA stipulates that the recipient of CMS PII data must properly protect the data according to FISMA and also provide for its appropriate destruction at the completion of the project/study or the expiration date of the DUA. The DUA form enables the data recipient and CMS to document the request and approval for release of CMS PII data. The form requires the submitter to provide the Requestor's organization; project/study name; CMS contract number (if applicable); data descriptions and the years of the data; retention date; attachments to the agreement; name, title, contact information to include address, city, state, zip code, phone, email, signature and date signed by the requester and custodian; disclosure provision; name of Federal Agency sponsor; Federal Representative name, title, contact information, signature, date; CMS representative name, title, contact information, signature and date;

and concurrence/non-concurrence signatures and dates from 3 CMS System Manager or Business Owners. While the data elements collected are not subject to change, the individualized clauses that are incorporated into any specific DUA are subject to change based on a specific case or situation such as disclosures to states, oversight agencies or DUAs for disproportionate share hospital (DSH) data requests as well as updates to DUAs with additional data descriptions, changes to the requestor or adding custodians to current DUAs. Form Number: CMS-R-235 (OCN: 0938-0734) Frequency: Once; Affected Public: Private Sector—Business or other Forprofits and Not-for-profit Institutions; Number of Respondents: 2,200; Number of Responses: 2,200; Total Annual *Hours:* 916. (For policy questions regarding this collection, contact Sharon Kavanagh at 410-786-5441. For all other issues call (410) 786-1326.)

To obtain copies of the supporting statement and any related forms for the proposed paperwork collections referenced above, access CMS' Web site at http://www.cms.gov/Paperwork ReductionActof1995/PRAL/ list.asp#TopOfPage or e-mail your request, including your address, phone number, OMB number, and CMS document identifier, to Paperwork@cms.hhs.gov, or call the Reports Clearance Office at 410–786–1326.

In commenting on the proposed information collections please reference the document identifier or OMB control number. To be assured consideration, comments and recommendations must be submitted in one of the following ways by July 19, 2011:

1. Electronically. You may submit your comments electronically to http://www.regulations.gov. Follow the instructions for "Comment or Submission" or "More Search Options" to find the information collection document(s) accepting comments.

2. By regular mail. You may mail written comments to the following address: CMS, Office of Strategic Operations and Regulatory Affairs, Division of Regulations Development, Attention: Document Identifier/OMB Control Number, Room C4–26–05, 7500 Security Boulevard, Baltimore, Maryland 21244–1850.

Dated: May 17, 2011.

Martique Jones,

Director, Regulations Development Group, Division B, Office of Strategic Operations and Regulatory Affairs.

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