in the ESRD program are required by
Pub. L. 95–292 to supply data to this
system. Form Number: CMS–2746
(OMB control number: 0938–0448);
Frequency: Yearly; Affected Public:
Private Sector (Business or other for-
profits, Not-For-Profit Institutions);
Number of Respondents: 7,311; Total
Annual Hours: 92,023; Total
Annual Hours: 46,011.50. (For policy
questions regarding this collection
contact Gequinicia Polk at 410–786–
2305.)

3. Type of Information Collection
Request: Reinstatement of previously
approved collection; Title of
Information Collection: End Stage Renal
Disease Medical Evidence Report
Medicare Entitlement and/or Patient
Registration; Use: The primary purpose
of this form is to have a physician
medically determine that a patient has
end stage renal disease for purposes of
filing for Medicare benefits. The End
Stage Renal Disease (ESRD) Medical
Evidence (CMS–2728) is completed for
all ESRD patients either by the first
treatment facility or by a Medicare-
approved ESRD facility when it is
determined by a physician that the
patient’s condition has reached that
stage of renal impairment that a regular
course of kidney dialysis or a kidney
transplant is necessary to maintain life.
The data reported on the CMS–2728 is
to monitor and assess the quality and
type of care provided to end stage renal
disease beneficiaries. Collection of these
data are also necessary for the
maintenance of a single, nationwide
kidney disease registry for dialysis,
transplant, and prospective transplant
patients. Form Number: CMS–2728
(OMB control number: 0938–0046);
Frequency: Yearly; Affected Public:
Private Sector (Business or other for-
profits, Not-For-Profit Institutions);
Number of Respondents: 7,311; Total
Annual Responses: 138,000; Total
Annual Hours: 103,500. (For policy
questions regarding this collection
contact Gequinicia Polk at 410–786–
2305.)

4. Type of Information Collection
Request: Revision of a currently
approved collection; Title of
Information Collection: Hospital
Notices: IM/DND; Use: The purpose of
the IM is to inform beneficiaries and
enrollees of their rights as hospital
inpatients and how to request a
discharge appeal by a Quality
Improvement Organization (QIO) and
how to file a request. For all Medicare
beneficiaries, hospitals must deliver
valid, written notice of a beneficiary’s
rights as a hospital inpatient, including
dischARGE appeal rights. The hospital
must use a standardized notice, as
specified by CMS. This is satisfied by
IM delivery.
Consistent with 42 CFR 405.1205 for
Original Medicare and 422.620 for
Medicare health plans, hospitals must
provide the initial IM within 2 calendar
days of admission. A follow-up copy of
the signed IM is given no more than 2
calendar days before discharge. The
follow-up copy is not required if the
first IM is provided within 2 calendar
days of discharge. In accordance with 42
CFR 405.1206 for Original Medicare and
422.622 for Medicare health plans, if a
beneficiary/enrollee appeals the
discharge decision, the beneficiary/
enrollee and the QIO must receive a
detailed explanation of the reasons
services should end. This detailed
explanation is provided to the
beneficiary/enrollee using the DND, the
second notice included in this renewal
package. Form Number: CMS–10065/
10066 (OMB control number: 0938–
1019); Frequency: Yearly; Affected
Public: Private Sector (Business or other
for-profits, Not-For-Profit Institutions);
Number of Respondents: 6,123; Total
Annual Responses: 17,742,803; Total
Annual Hours: 2,990,720. (For policy
questions regarding this collection
contact Janet Miller at 410–786–1799.)

William N. Parham, III,
Director, Paperwork Reduction Staff, Office
of Strategic Operations and Regulatory
Affairs.
[FR Doc. 2019–03015 Filed 2–21–19; 8:45 am]
BILLING CODE 4120–01–P

DEPARTMENT OF HEALTH AND
HUMAN SERVICES

Centers for Medicare & Medicaid Services
[Document Identifier: CMS–643]
Agency Information Collection Activities: Proposed Collection; Comment Request
AGENCY: Centers for Medicare & Medicaid Services, HHS.
ACTION: Notice.
SUMMARY: The Centers for Medicare & Medicaid Services (CMS) is announcing an opportunity for the public to comment on CMS’ intention to collect information from the public. Under the Paperwork Reduction Act of 1995 (the PRA), federal agencies are required to publish notice in the Federal Register concerning each proposed collection of information (including each proposed extension or reinstatement of an existing collection of information) and to allow 60 days for public comment on the proposed action. Interested persons are invited to send comments regarding our burden estimates or any other aspect of this collection of information, including the necessity and utility of the proposed information collection for the proper performance of the agency’s functions, the accuracy of the estimated burden, ways to enhance the quality, utility, and clarity of the information to be collected, and the use of automated collection techniques or other forms of information technology to minimize the information collection burden.
DATES: Comments must be received by April 23, 2019.
ADDRESSES: When commenting, please reference the document identifier or OMB control number. To be assured consideration, comments and recommendations must be submitted in any one of the following ways:
1. Electronically. You may send 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) that are 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.
To obtain copies of a supporting statement and any related forms for the proposed collection(s) summarized in this notice, you may make your request using one of following:
2. Email your request, including your address, phone number, OMB number, and CMS document identifier, to Paperwork@cms.hhs.gov.
3. Call the Reports Clearance Office at (410) 786–1326.
FOR FURTHER INFORMATION CONTACT: William N. Parham at (410) 786–4699.
SUPPLEMENTARY INFORMATION:
Contents
This notice sets out a summary of the use and burden associated with the following information collections. More detailed information can be found in each collection’s supporting statement and associated materials (see ADDRESSES).
CMS–643 Hospice Survey and Deficiencies Report Form and Supporting Regulations

Under the PRA (44 U.S.C. 3501–3520), federal agencies must obtain approval from the Office of Management and Budget (OMB) for each collection of information they conduct or sponsor. The term “collection of information” is defined in 44 U.S.C. 3502(3) and 5 CFR 1320.3(c) and includes agency requests or requirements that members of the public submit reports, keep records, or provide information to a third party. Section 3506(c)(2)(A) of the PRA requires federal agencies to publish a 60-day notice in the Federal Register concerning each proposed collection of information, including each proposed extension or reinstatement of an existing collection of information, before submitting the collection to OMB for approval. To comply with this requirement, CMS is publishing this notice.

Information Collection

1. Type of Information Collection Request: Extension without change of a currently approved collection; Title of Information Collection: Hospice Survey and Deficiencies Report Form and Supporting Regulations; Use: We use the information collected as the basis for certification decisions for hospices that wish to obtain or retain participation in the Medicare and Medicaid programs. The information is used by CMS regional offices, which have delegated authority to certify Medicare facilities for participation, and by State Medicaid agencies, which have comparable authority under Medicaid. The information on the Hospice Survey and Deficiencies Report Form is coded for entry into the OSCAR system. The data is analyzed by the CMS regional offices and by the CMS central office components for program evaluation and monitoring purposes. The information is also available to the public upon request. Form Number: CMS–643 (OMB control number: 0938–0379); Frequency: Yearly; Affected Public: State, Local, or Tribal Governments; Number of Respondents: 8,111; Total Annual Responses: 1,603; Total Annual Hours: 1,603. (For policy questions regarding this collection contact Thomas Pryor at 410–786–1132.)


William N. Parham, III,
Director, Paperwork Reduction Staff, Office of Strategic Operations and Regulatory Affairs.

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Submission for OMB Review; ACF’s Generic Clearance for Grant Reviewer Recruitment Forms (OMB #0970–0477)

AGENCY: Office of Planning, Research, and Evaluation; Administration for Children and Families; HHS.

ACTION: Request for public comment.

SUMMARY: The Administration for Children and Families (ACF), Office of Planning, Research, and Evaluation (OPRE) is proposing an extension of a currently approved generic clearance (OMB No. 0970–0477) for Grant Reviewer Recruitment (GRR) forms. The GRR forms will be used to select reviewers who will participate in the grant peer review process for the purpose of selecting successful applications.

DATES: Comments due within 30 days of publication. 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.

ADDRESSES: 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@EOP.EOP.GOV, Attn: Desk Officer for the Administration for Children and Families.

Copies of the proposed collection may be obtained by writing to the Administration for Children and Families, Office of Planning, Research and Evaluation, 330 C Street SW, Washington, DC 20201. Attn: OPRE Reports Clearance Officer. All requests should be identified by the title of the information collection. Email address: OPREinfocollection@acf.hhs.gov.

SUPPLEMENTARY INFORMATION:

Description: Under this generic approval, ACF conducts and proposes to continue to conduct more than one information collection that is very similar, voluntary, low-burden and uncontroversial. The purpose is to select qualified reviewers for the grant peer review process based on professional qualifications using data entered by candidates and the uploaded writing sample and/or curriculum vitae and/or resume. The grant review process is in accordance with the U.S. Department of Health and Human Services’ (DHHS) Grants Policy Directive (GPD) 2.04 “Awarding Grants”, the DHHS Awarding Agency Grants Administration Manual (AAGAM), Chapter 2.04.104C “Objective Review of Grant Applications”, and the Public Health Service (PHS) Act, Sections 799(f) and 806(e).

Respondents: Individuals who may apply to review ACF grant applications.

ANNUAL BURDEN ESTIMATES

<table>
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<tr>
<th>Instrument</th>
<th>Total number of responses</th>
<th>Number of responses per respondent</th>
<th>Average burden hours per response</th>
<th>Annual burden hours</th>
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<td>1</td>
<td>.5</td>
<td>1500</td>
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</tbody>
</table>

Estimated Total Annual Burden Hours: 1500.

Mary B. Jones,
ACF/OPRE Certifying Officer.

[FR Doc. 2019–03079 Filed 2–21–19; 8:45 am]