• Veterans whose records at VHA match identifying data provided to VHA by CMS (submitted by AEs) about individuals who are applying for or are enrolled in private insurance coverage under a qualified health plan through a federally-facilitated health insurance exchange or state-based exchange.

**Categories of Records**

The categories of records used in this matching program are identity records and minimum essential coverage period records, consisting of the following data elements:

- Data provided by CMS to VHA
  - a. first name (required).
  - b. middle name/initial (if provided by applicant).
  - c. surname (applicant’s last name) (required).
  - d. date of birth (required).
  - e. gender (required).
  - f. social security number (SSN) (required).
- g. requested qualified health plan (QHP) coverage effective date (required).
- h. requested QHP coverage end date (required).
  - i. State identification (required).
  - j. transaction ID (required).
- Data provided by VHA to CMS
  - a. SSN (required).
  - b. start/end date(s) of enrollment period(s) (when match occurs).
  - c. a blank date response when a non-match occurs.
  - d. enrollment period(s) is/are defined as the timeframe during which the individual was enrolled in a VHA health care program.

**System(s) of Records**

The records used in this matching program will be disclosed from the following systems of records, as authorized by routine uses published in the system of records notices (SORNs) cited below:

A. **System of Records Maintained by CMS**


**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**Centers for Medicare & Medicaid Services**

**[Document Identifier CMS–10147]**

**Agency Information Collection Activities: Submission for OMB Review; Comment Request**

**AGENCY:** Centers for Medicare & Medicaid Services, Health and Human 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 (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 a second opportunity for public comment on the notice. Interested persons are invited to send comments regarding the burden estimate 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 on the collection(s) of information must be received by the OMB desk officer by April 26, 2021.

**ADDRESSES:** Written comments and recommendations for the proposed information collection should be sent within 30 days of publication of this notice to www.reginfo.gov/public/do/PRAMain. Find this particular information collection by selecting “Currently under 30-day Review—Open for Public Comments” or by using the search function.

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 the following:


**FOR FURTHER INFORMATION CONTACT:**

William Parham at (410) 786–4669.

**SUPPLEMENTARY INFORMATION:**

Under the Paperwork Reduction Act of 1995 (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 (44 U.S.C. 3506(c)(2)(A)) requires federal agencies to publish a 30-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 that summarizes the following proposed collection(s) of information for public comment:

1. **Type of Information Collection Request:** Reinstatement without change of a previously approved collection; **Title of Information Collection:** Medicare Prescription Drug Coverage and Your Rights; **Use:** Section 423.562(a)(3) and an associated regulatory provision at § 423.128(b)(7)(iii) require that Part D plan sponsors’ network pharmacies provide Part D enrollees with a printed copy of our standardized pharmacy notice “Medicare Prescription Drug Coverage and Your Rights” (hereafter, “notice”) if an enrollee’s prescription cannot be filled.

The purpose of this notice is to provide enrollees with information about how to contact their Part D plans to request a coverage determination, including a request for an exception to the Part D plan’s formulary. The notice reminds enrollees about certain rights and protections related to their Medicare prescription drug benefits, including the right to receive a written explanation from the drug plan about why a prescription drug is not covered. Through delivery of this standardized notice, a Part D plan sponsor’s network pharmacies are in the best position to inform enrollees at point of sale about how to contact their Part D plan if the prescription cannot be filled. **Form**
Number: CMS–10147 [OMB control number 0938–0975]; Frequency: Annually; Affected Public: Private Sector, Business or other for-profits; Number of Respondents: 70,000; Number of Responses: 49,681,292; 70,000; Number of Annual Hours: 827,690. (For questions regarding this collection, contact Trevor Rose at (410) 786 7768.)


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

DEPARTMENT OF HEALTH AND HUMAN SERVICES
Food and Drug Administration

Request for Nominations for Individuals and Consumer Organizations for Advisory Committees

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA or Agency) is requesting that any consumer organizations interested in participating in the selection of voting and/or nonvoting consumer representatives to serve on its advisory committees or panels notify FDA in writing. FDA is also requesting nominations for voting and/or nonvoting consumer representatives to serve on advisory committees and/or panels for which vacancies currently exist or are expected to occur in the near future. Nominees recommended to serve as a voting or nonvoting consumer representative may be self-nominated or may be nominated by a consumer organization. FDA seeks to include the views of women and men, members of all racial and ethnic groups, and individuals with and without disabilities on its advisory committees and, therefore, encourages nominations of appropriately qualified candidates from these groups.

DATES: Any consumer organization interested in participating in the selection of an appropriate voting or nonvoting member to represent consumer interests on an FDA advisory committee or panel may send a letter or email stating that interest to FDA (see ADDRESSES) by April 26, 2021, for vacancies listed in this notice. Concurrently, nomination materials for prospective candidates should be sent to FDA (see ADDRESSES) by April 26, 2021. Nominations will be accepted for current vacancies and for those that will or may occur through December 31, 2021.

ADDRESSES: All statements of interest from consumer organizations interested in participating in the selection process should be submitted electronically to ACOMSSubmissions@fda.hhs.gov or by mail to Advisory Committee Oversight and Management Staff, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 32, Rm. 5122, Silver Spring, MD 20993–0002.

Consumer representative nominations should be submitted electronically by logging into the FDA Advisory Committee Membership Nomination Portal: https://www.accessdata.fda.gov/scripts/FACTRSPortal/FACTRIS/index.cfm, or by mail to Advisory Committee Oversight and Management Staff, 10903 New Hampshire Ave., Bldg. 32, Rm. 5122, Silver Spring, MD 20993–0002. Additional information about becoming a member of an FDA advisory committee can also be obtained by visiting FDA’s website at https://www.fda.gov/AdvisoryCommittees/default.htm.

FOR FURTHER INFORMATION CONTACT: For questions relating to participation in the selection process: Kimberly Hamilton, Advisory Committee Oversight and Management Staff, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 32, Rm. 5122, Silver Spring, MD 20993–0002, 301–796–8220, kimberly.hamilton@fda.hhs.gov.

For questions relating to specific advisory committees or panels, contact the appropriate Contact Person listed in table 1.

<table>
<thead>
<tr>
<th>Contact person</th>
<th>Committee/panel</th>
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<tbody>
<tr>
<td>Rakesh Raghuwanshi, Office of the Chief Scientist, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 1, Rm. 3309, Silver Spring, MD 20993–0002, 301–796–4769, <a href="mailto:Rakesh.Raghuwanshi@fda.hhs.gov">Rakesh.Raghuwanshi@fda.hhs.gov</a>.</td>
<td>FDA Science Board Advisory Committee.</td>
</tr>
<tr>
<td>Kathleen Hayes, Center for Biologics Evaluation and Research, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 71, Rm. 5122, Silver Spring, MD 20993–0002, 301–796–7664, <a href="mailto:Kathleen.Hayes@fda.hhs.gov">Kathleen.Hayes@fda.hhs.gov</a>.</td>
<td>Vaccines and Related Biological Products Advisory Committee.</td>
</tr>
<tr>
<td>LaToya Bonner, Center for Drugs Evaluation and Research, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 31, Rm. 2428, Silver Spring, MD 20992–0002, 301–796–2855, <a href="mailto:Latoya.Bonner@fda.hhs.gov">Latoya.Bonner@fda.hhs.gov</a>.</td>
<td>Dermatologic and Ophthalmic Drugs Advisory Committee.</td>
</tr>
<tr>
<td>Yvette Waples, Center for Drugs Evaluation and Research, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 31, Rm. 2510, Silver Spring, MD 20993–0002, 301–796–9034, <a href="mailto:Yvette.Waples@fda.hhs.gov">Yvette.Waples@fda.hhs.gov</a>.</td>
<td>Gastrointestinal Drugs Advisory Committee, Pharmaceutical Science and Clinical Pharmacology Advisory Committee, Psychopharmacologic Drugs Advisory Committee.</td>
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