

ACTION: Notice.

SUMMARY: The Postal Service gives notice of filing a request with the Postal Regulatory Commission to add a domestic shipping services contract to the list of Negotiated Service Agreements in the Mail Classification Schedule's Competitive Products List.

DATES: *Date of required notice:* December 30, 2022.

FOR FURTHER INFORMATION CONTACT: Sean Robinson, 202-268-8405.

SUPPLEMENTARY INFORMATION: The United States Postal Service® hereby gives notice that, pursuant to 39 U.S.C. 3642 and 3632(b)(3), on December 22, 2022, it filed with the Postal Regulatory Commission a *USPS Request to Add Priority Mail Express, Priority Mail, First-Class Package Service, and Parcel Select Service Contract 105 to Competitive Product List*. Documents are available at www.prc.gov, Docket Nos. MC2023-96, CP2023-97.

Sarah Sullivan,

Attorney, Ethics & Legal Compliance.

[FR Doc. 2022-28413 Filed 12-29-22; 8:45 am]

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POSTAL SERVICE

Product Change—Priority Mail Express, Priority Mail, & First-Class Package Service Negotiated Service Agreement

AGENCY: Postal Service™.

ACTION: Notice.

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DATES: *Date of required notice:* December 30, 2022.

FOR FURTHER INFORMATION CONTACT: Sean Robinson, 202-268-8405.

SUPPLEMENTARY INFORMATION: The United States Postal Service® hereby gives notice that, pursuant to 39 U.S.C. 3642 and 3632(b)(3), on December 19, 2022, it filed with the Postal Regulatory Commission a *USPS Request to Add Priority Mail Express, Priority Mail, & First-Class Package Service Contract 80 to Competitive Product List*. Documents are available at www.prc.gov, Docket Nos. MC2023-94, CP2023-95.

Sarah Sullivan,

Attorney, Ethics & Legal Compliance.

[FR Doc. 2022-28417 Filed 12-29-22; 8:45 am]

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SOCIAL SECURITY ADMINISTRATION

[Docket No: SSA-2022-0067]

Agency Information Collection Activities: Proposed Request and Comment Request

The Social Security Administration (SSA) publishes a list of information collection packages requiring clearance by the Office of Management and Budget (OMB) in compliance with Public Law 104-13, the Paperwork Reduction Act of 1995, effective October 1, 1995. This notice includes revisions of OMB-approved information collections and one new collection.

SSA is soliciting comments on the accuracy of the agency's burden estimate; the need for the information; its practical utility; ways to enhance its quality, utility, and clarity; and ways to minimize burden on respondents, including the use of automated collection techniques or other forms of information technology. Mail, email, or fax your comments and recommendations on the information collection(s) to the OMB Desk Officer and SSA Reports Clearance Officer at the following addresses or fax numbers.

(OMB) Office of Management and Budget, Attn: Desk Officer for SSA, Comments: <https://www.reginfo.gov/public/do/PRAMain>. Submit your comments online referencing Docket ID Number [SSA-2022-0067].

(SSA) Social Security Administration, OLCA, Attn: Reports Clearance Director, 3100 West High Rise, 6401 Security Blvd., Baltimore, MD 21235, Fax: 410-966-2830, Email address:

OR.Reports.Clearance@ssa.gov.

Or you may submit your comments online through <https://www.reginfo.gov/public/do/PRAMain>, referencing Docket ID Number [SSA-2022-0067].

I. The information collections below are pending at SSA. SSA will submit them to OMB within 60 days from the date of this notice. To be sure we consider your comments, we must receive them no later than February 28, 2023. Individuals can obtain copies of the collection instruments by writing to the above email address.

1. Vocational Resource Facilitator Demonstration—0960-NEW. SSA is undertaking the Vocational Resource Facilitator Demonstration (VRFD) under the Interventional Cooperative Agreement Program (ICAP). ICAP allows SSA to partner with various non-federal groups and organizations to advance interventional research connected to the Supplemental Security Income (SSI) and Social Security Disability Insurance (SSDI) programs. VRFD will test the Vocational Resource Facilitator (VRF)

intervention, which helps newly injured spinal cord injury or disease (SCI) or brain injury (BI) patients pursue their employment goals. The VRFD will provide empirical evidence on the impact of the intervention on patients in several critical areas: (1) employment and earnings; (2) SSI and SSDI benefit receipt; and (3) satisfaction and well-being. A rigorous evaluation of VRFD is critical to help SSA and other interested parties assess promising options to improve employment-related outcomes and decrease benefit receipt. The VRFD evaluation uses a randomized control experimental design that includes one treatment group and one control group. Control group members will receive a referral for services to the Division of Vocational Rehabilitation Services (DVRS), New Jersey's state Vocational Rehabilitation agency. The treatment group will receive a referral to DVRS and employment services from a resource facilitator (RF). RFs are fully integrated members of clinical teams who engage with injured workers during inpatient rehabilitation about return to work. The central research questions include:

- Was the intervention implemented as planned?
- What are key considerations for scaling up or adopting the VRF model at other facilities?
- What were the impacts of VRF on outcomes of interest?
- Did treatment group members earn or work more than control group members?
- Were treatment group members relatively less likely to apply to or receive SSI or SSDI benefits?
- Did treatment group members experience greater satisfaction and well-being than control group members?
- What were the benefits and costs of the demonstration across key groups?

The proposed public survey data collections will support three components of the planned implementation, impact, and benefit-cost analyses. The data collection efforts will provide information that is not available in SSA program records about the characteristics and outcomes of VRFD participants in the treatment and control groups. Respondents are newly injured SCI and BI patients, who will provide written consent before agreeing to participate in the study and be randomly assigned to one of the study groups.

Type of Request: Request for a new information collection.