B. Federal Reserve Bank of Kansas City (Dennis Denney, Assistant Vice President) 1 Memorial Drive, Kansas City, Missouri 64198–0001:

1. Lance L. White, as co-trustee of the Lance L. White Revocable Trust and as trustee of the Lance L. White Irrevocable Trust, and Cherlyn D. White, as trustee of the Cherlyn D. White-Conklin Irrevocable Trust and as co-trustee of the Cherlyn White-Conklin Trust, all of Wamego, Kansas; and Monte W. White, individually, and as trustee of the MWW Irrevocable Trust #1, both of Salina, Kansas; as members of the White Family Group, a group acting in concert, to retain voting shares of Wamego Bancshares, Inc., and thereby indirectly retain voting shares of Bank of the Flint Hills, both of Wamego, Kansas.

Additionally, Kara L. White, as co-trustee of the Lance L. White Revocable Trust, Cherlyn White-Conklin Trust, Erich Conklin, as co-trustee, and certain minor children, all of Salina and Wamego, Kansas; to join the White Family Group, and retain voting shares of Wamego Bancshares, Inc., and thereby indirectly retain voting shares of Bank of the Flint Hills.


Michele Taylor Fennell, Deputy Associate Secretary of the Board.

[FR Doc. 2021–03841 Filed 2–24–21; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[Docket No. CDC–2021–0021]

Advisory Committee on Immunization Practices (ACIP)

AGENCY: Centers for Disease Control and Prevention (CDC), Department of Health and Human Services (HHS).

ACTION: Notice of meeting and request for comment.

SUMMARY: In accordance with the Federal Advisory Committee Act, the Centers for Disease Control and Prevention (CDC), announces the following meeting of the Advisory Committee on Immunization Practices (ACIP). This meeting is open to the public. Time will be available for public comment. The meeting will be webcast live via the World Wide Web.

DATES: The meeting will be held on February 28, 2021–March 1, 2021, from 10:00 a.m. to 5:00 p.m. EDT (times subject to change). Written comments must be received on or before March 1, 2021.

ADDRESSES: For more information on ACIP please visit the ACIP website: http://www.cdc.gov/vaccines/acip/index.html.

You may submit comments, identified by Docket No. CDC–2021–0021 by any of the following methods:

• Federal eRulemaking Portal: https://www.regulations.gov. Follow the instructions for submitting comments.

• Mail: Docket No. CDC–2021–0021, c/o Attn: ACIP Meeting, Centers for Disease Control and Prevention, 1600 Clifton Road NE, MS H24–8, Atlanta, GA 30329–4027.

Instructions: All submissions received must include the Agency name and Docket Number. All relevant comments received in conformance with the https://www.regulations.gov suitability policy will be posted without change to https://www.regulations.gov, including any personal information provided. For access to the docket to read background documents or comments received, go to https://www.regulations.gov.

Written public comments submitted 24 hours prior to the ACIP meeting will be provided to ACIP members before the meeting.

FOR FURTHER INFORMATION CONTACT:

Stephanie Thomas, ACIP Committee Management Specialist, Centers for Disease Control and Prevention, National Center for Immunization and Respiratory Diseases, 1600 Clifton Road NE, MS–H24–8, Atlanta, GA 30329–4027; Telephone: 404–639–8367; Email: ACIP@cdc.gov.

SUPPLEMENTARY INFORMATION: In accordance with 41 CFR 102–3.150(b), less than 15 calendar days notice is being given for this meeting due to the exceptional circumstances of the COVID–19 pandemic and rapidly evolving COVID–19 vaccine development and regulatory processes. The Secretary of Health and Human Services has determined that COVID–19 is a Public Health Emergency.

Purpose: The committee is charged with advising the Director, CDC, on the use of immunizing agents. In addition, under 42 U.S.C. 1396s, the committee is mandated to establish and periodically review and, as appropriate, revise the list of vaccines for administration to vaccine-eligible children through the Vaccines for Children (VFC) program, along with schedules regarding dosing interval, dosage, and contraindications to administration of vaccines. Further, under provisions of the Affordable Care Act, section 2713 of the Public Health Service Act, immunization recommendations of the ACIP that have been approved by the Director of the Centers for Disease Control and Prevention and appear on CDC immunization schedules must be covered by applicable health plans.

Matters To Be Considered: The agenda will include discussions on COVID–19 vaccines. A recommendation vote is scheduled. Agenda items are subject to change as priorities dictate. For more information on the meeting agenda visit https://www.cdc.gov/vaccines/acip/meetings/meetings-info.html.

Meeting Information: The meeting will be webcast live via the World Wide Web; for more information on ACIP please visit the ACIP website: http://www.cdc.gov/vaccines/acip/index.html.

Public Participation

Interested persons or organizations are invited to participate by submitting written views, recommendations, and data. Please note that comments received, including attachments and other supporting materials are part of the public record and are subject to public disclosure. Do not include any information in your comment or supporting materials that you consider confidential or inappropriate for public disclosure. If you include your name, contact information, or other information that identifies you in the body of your comments, that information will be on public display. CDC will review all submissions and may choose to redact, or withhold, submissions containing private or proprietary information such as Social Security numbers, medical information, inappropriate language, or duplicate/near duplicate examples of a mass-mail campaign. CDC will carefully consider all comments submitted into the docket. CDC does not accept comment by email.

Written Public Comment: Written comments must be received on or before March 1, 2021. Oral Public Comment: This meeting will include time for members of the public to make an oral comment. Oral public comment will occur before any scheduled votes including all votes relevant to the ACIP’s Affordable Care Act and Vaccines for Children Program roles. Priority will be given to individuals who submit a request to make an oral public comment before the meeting according to the procedures below.

Procedure for Oral Public Comment: All persons interested in making an oral public comment at the February 28, 2021–March 1, 2021 ACIP meeting must submit a request at http://www.cdc.gov/vaccines/acip/meetings/ no later than 11:59 p.m., EDT, February 25, 2021 according to the instructions provided.
If the number of persons requesting to speak is greater than can be reasonably accommodated during the scheduled time, CDC will conduct a lottery to determine the speakers for the scheduled public comment session. CDC staff will notify individuals regarding their request to speak by email by February 26, 2021. To accommodate the significant interest in participation in the oral public comment session of ACIP meetings, each speaker will be limited to 3 minutes, and each speaker may only speak once per meeting.

The Director, Strategic Business Initiatives Unit, Office of the Chief Operating Officer, Centers for Disease Control and Prevention, has been delegated the authority to sign Federal Register notices pertaining to announcements of meetings and other committee management activities, for both the Centers for Disease Control and Prevention and the Agency for Toxic Substances and Disease Registry.

Kalwant Smagh,
Director, Strategic Business Initiatives Unit, Office of the Chief Operating Officer, Centers for Disease Control and Prevention.

[FR Doc. 2021–03959 Filed 2–23–21; 11:15 am]
BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[60Day–21–21DC; Docket No. CDC–2021–0012]

Proposed Data Collection Submitted for Public Comment and Recommendations

AGENCY: Centers for Disease Control and Prevention (CDC), Department of Health and Human Services (HHS).

ACTION: Notice with comment period.

SUMMARY: The Centers for Disease Control and Prevention (CDC), as part of its continuing effort to reduce public burden and maximize the utility of government information, invites the general public and other Federal agencies the opportunity to comment on a proposed and/or continuing information collection, as required by the Paperwork Reduction Act of 1995. This notice invites comment on a proposed information collection project titled 'National Syringe Services Program (SSP) Evaluation', which proposes to: (1) Assess and monitor SSP operational characteristics and services, client characteristics and drug use patterns, client satisfaction, funding resources, community relations, and key operational and programmatic successes and challenges and (2) support timely analysis and dissemination of national program evaluation survey findings.

DATES: CDC must receive written comments on or before April 26, 2021.

ADDRESSES: You may submit comments, identified by Docket No. CDC–2021–0012 by any of the following methods:

• Federal eRulemaking Portal: Regulations.gov. Follow the instructions for submitting comments.

• Mail: Jeffrey M. Zirger, Information Collection Review Office, Centers for Disease Control and Prevention, 1600 Clifton Road NE, MS–D74, Atlanta, Georgia 30329.

Instructions: All submissions received must include the agency name and Docket Number. CDC will post, without change, all relevant comments to Regulations.gov.

Please note: Submit all comments through the Federal eRulemaking portal (regulations.gov) or by U.S. mail to the address listed above.

FOR FURTHER INFORMATION CONTACT: To request more information on the proposed project or to obtain a copy of the information collection plan and instruments, contact Jeffrey M. Zirger, Information Collection Review Office, Centers for Disease Control and Prevention, 1600 Clifton Road NE, MS–D74, Atlanta, Georgia 30329; phone: 404–639–7570; Email: omb@cdc.gov.

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. In addition, the PRA also requires Federal agencies to provide a 60-day notice in the Federal Register concerning each proposed collection of information, including each new proposed collection, each proposed extension of existing collection of information, and each reinstatement of previously approved information collection before submitting the collection to the OMB for approval. To comply with this requirement, we are publishing this notice of a proposed data collection as described below.

The OMB is particularly interested in comments that will help:

1. Evaluate whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information will have practical utility;

2. Evaluate whether the agency's estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used;

3. Enhance the quality, utility, and clarity of the information to be collected; and

4. Minimize the burden of the collection of information on those who are to respond, including through the use of appropriate automated, electronic, mechanical, or other technological collection techniques or other forms of information technology, e.g., permitting electronic submissions of responses.

5. Assess information collection costs.

Proposed Project


Background and Brief Description

The primary purpose of the National Syringe Services Program (SSP) Evaluation is to strengthen and improve the capacity of SSPs to conduct regular monitoring and evaluation to ensure that comprehensive prevention services are provided to meet the needs of people who inject drugs (PWID) and reduce infectious disease and other harms related to intravenous drug use (IDU). The project will invite the participation of all SSPs that are listed in a publicly available directory of all known SSPs in the United States maintained by the North American Syringe Exchange Network (NASEN; https://nasan.org). SSPs will be sent a letter of invitation to participate in a 35-minute program survey. Participating programs will have the option of completing the survey via different modalities to enhance feasibility and comfort in completing the survey, for example via the Research Electronic Data Capture (REDCap) or a similarly secure web-based application. Other modalities for survey administration will include a coordinated telephone or videoconferencing interview. SSPs will be sent reminder letters for an approximately three-month data collection period. SSPs that do not respond to prior reminders will be sent one final reminder, and if the SSP still does not want to participate, one (optional) question on why the SSP did not complete the survey will be offered.

The survey will include questions on operational characteristics and services, client characteristics and drug use patterns, client satisfaction, funding resources, community relations, and key operational successes and challenges. Approximately 400 SSPs will be able to