for cruise ship operators to test the efficacy of their health and safety
protocols in U.S. waters. In lieu of conducting a simulated voyage, a cruise
ship operator’s responsible officials, at their discretion, may sign and submit to
CDC an attestation under 18 U.S.C. 1001 that 98 percent of crew are fully
vaccinated and submit to CDC a clear and specific vaccination plan and
timeline to limit cruise ship sailings to 95 percent of passengers who have been
verified by the cruise ship operator as fully vaccinated prior to sailing.

CDC’s oversight and inspection of cruise ships during simulated and
restricted passenger voyages will be based on the Operations Manual. The
findings and/or observations of these inspections will be shared with the
cruise ship operator. Cruise ship operators are expected to align their
health and safety protocols with any CDC findings and observations. Such
findings and observations must also be incorporated into the cruise ship
operator’s simulated voyage after-action report or as a condition of applying for
and retaining permission to conduct restricted passenger voyages. Based on
these inspections, CDC may also issue additional recommendations to the
cruise ship operator that the operator should consider for adoption into their
health and safety protocols as best practices.

The Technical Instructions document for Cruise Ship Operator’s Agreement
with Port and Local Health Authorities under CDC’s Framework for Conditional
Sailing Order is found at https://
www.cdc.gov/quarantine/cruise/
instructions-local-agreements.html.

The Technical Instructions document for Simulated Voyages by Cruise Ship
Operators under CDC’s Framework for Conditional Sailing Order is found at
https://www.cdc.gov/quarantine/cruise/
ti-simulated-voyages-cso.html.

The COVID–19 Operations Manual for
Simulated and Restricted Voyages under the
Framework for Conditional Sailing Order is found at https://www.cdc.gov/
quarantine/cruise/covid19-operations-
manual-cso.html.

Authority

The Technical Instructions and
Operations Manual are issued pursuant to the Framework for Conditional
Sailing which was issued under the authority of Sections 361 and 365 of the
Public Health Service Act (42 U.S.C. 264, 268) and 42 CFR 70.2, 71.31(b),
71.32(b).

Sherri Berger,
Acting Chief of Staff, Centers for Disease
Control and Prevention.

DEPARTMENT OF HEALTH AND HUMAN SERVICES
Centers for Disease Control and
Prevention
[Docket No. CDC–2021–0049]
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. The meeting will be webcast live
via the World Wide Web.
DATES: The meeting will be held on May
12, 2021, from 11:00 a.m. to 5:00 p.m.,
EDT (dates and times subject to change,
see the ACIP website for updates: http://
www.cdc.gov/vaccines/acip/
index.html). The public may submit
comments from May 10, 2021 through
May 12, 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–2021–0049 by any of
the following methods:
• Federal eRulemaking Portal:
https://www.regulations.gov. Follow the
instructions for submitting comments.
• Mail: Centers for Disease Control
and Prevention, 1600 Clifton Road NE,
MS–H24–8, Atlanta, GA 30329–4027.
Attn: May 12, 2021 ACIP Meeting.
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
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 102–3.15(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. A notice
of this ACIP meeting has also been
posted on CDC’s ACIP website at:
http://
In addition, CDC has sent notice of
this ACIP meeting by email to those who
subscribe to receive email updates about
ACIP.
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, including use of the Pfizer-
BioNTech COVID–19 vaccine under the
Food and Drug Administration’s (FDA)
expanded Emergency Use Authorization
(EUA) for adolescents 12–15 years of
age. A recommendation vote(s) 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://
Public Participation
Interested persons or organizations are
invited to participate by submitting
written views, recommendations,
data. Please note that comments received, including attachments and other supporting materials, are part of the public record and are subject to public disclosure. Comments will be posted on https://www.regulations.gov. Therefore, 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.

Written Public Comment: Written comments must be received on or before May 12, 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 May 12, 2021 ACIP meeting must submit a request at http://www.cdc.gov/vaccines/acip/meetings/ no later than 11:59 p.m., EDT, May 10, 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 12:00 p.m., EDT, May 11, 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 Smaghi,
Director, Strategic Business Initiatives Unit, Office of the Chief Operating Officer, Centers for Disease Control and Prevention.

[PR Doc. 2021–09893 Filed 5–6–21; 11:15 am]

BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

[CFDA Number: 93.676]

Announcement of Intent To Issue Multiple Single-Source Awards To Provide Residential (Shelter) and Transitional Foster Care Services, and for Finger Print Services for Unaccompanied Children

AGENCY: Office of Refugee Resettlement (ORR), Administration for Children and Families (ACF), Department of Health and Human Services (HHS).

ACTION: Notice of issuance of multiple single-source awards to seven recipients.

SUMMARY: ACF, ORR announces the issuance of multiple Single-Source Awards to seven recipients in the amount of $65,366,800. ORR has been identifying additional permanent capacity to provide shelter for recent increases in apprehensions of Unaccompanied Children (UC) at the Southwest Border. The addition of permanent capacity is a prudent step to ensure that ORR is able to meet its responsibility, by law, to provide shelter and appropriate services for UC referred to its care by the Department of Homeland Security.

DATES: The proposed period of performance is May 1, 2021 to May 1, 2022.


SUPPLEMENTAL INFORMATION: ORR is continuously monitoring its capacity to shelter the UC referred to HHS, as well as the information received from interagency partners, to inform any future decisions or actions. ORR has specific requirements for the provision of services. Award recipients must have the infrastructure, licensing, experience, and appropriate level of trained staff to meet those requirements.

ORR announces the intent to award the following single-source awards:

<table>
<thead>
<tr>
<th>Recipient</th>
<th>Award amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>Child Crisis, Mesa, AZ</td>
<td>$5,780,118</td>
</tr>
<tr>
<td>Catholic Guardian Services, New York, NY</td>
<td>5,183,433</td>
</tr>
<tr>
<td>Center for Family Services, Camden, NJ</td>
<td>1,665,980</td>
</tr>
<tr>
<td>LIRS—Shelter/TFC, Multiple Locations</td>
<td>27,767,725</td>
</tr>
<tr>
<td>Baptiste Group, Memphis, TN</td>
<td>14,135,642</td>
</tr>
<tr>
<td>Bethany Christian Service, Multiple Locations</td>
<td>7,018,576</td>
</tr>
<tr>
<td>LIRS—Safe Release Expansion, Multiple Locations</td>
<td>3,815,326</td>
</tr>
</tbody>
</table>

Statutory Authority: This program is authorized by—

(A) Section 462 of the Homeland Security Act of 2002, which in March 2003, transferred responsibility for the care and custody of UAC from the Commissioner of the former Immigration and Naturalization Service to the Director of ORR within HHS.

(B) The Flores Settlement Agreement, Case No. CV85–4544–RJK (C.D. Cal. 1996), as well as the Willberforce Trafficking Victims Protection Reauthorization Act of 2008 (Pub. L. 110–457), which authorizes post release services under certain conditions to eligible children. All programs must comply with the Flores Settlement Agreement, Case No. CV85–4544–RJK (C.D. Cal. 1996); pertinent regulations; and ORR policies and procedures.

Elizabeth Leo,
Senior Grants Policy Specialist, Office of Grants Policy, Office of Administration.
[PR Doc. 2021–09758 Filed 5–4–21; 4:15 pm]

BILLING CODE 4184–45–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration


Agency Information Collection Activities; Announcement of Office of Management and Budget Approvals

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is publishing a list of information collections that have