International Overdose Awareness Day — August 31, 2019

August 31, 2019, is International Overdose Awareness Day, a global event that aims to raise awareness that overdose death is preventable and to reduce the stigma associated with drug-related death. Goals also include providing information about risk for overdose and community services and preventing drug-related harm through evidence-based policy and practice (https://www.overdoseday.com).

The opioid overdose epidemic, which killed 47,600 U.S. persons in 2017,* substantially expanded in 2013 driven by rapid increases in overdose deaths involving synthetic opioids (excluding methadone), particularly illicitly manufactured fentanyl.† Cocaine and methamphetamine overdose deaths co-involving synthetic opioids also rapidly increased during this period (1).

A report in this issue of MMWR documented decreases in opioid-involved overdose deaths in 25 states from July–December 2017 to January–June 2018, especially those involving fentanyl analogs and prescription opioids. Overdose deaths involving illicitly manufactured fentanyl (including those co-occurring with illicit opioids and stimulants) increased (2). Improved identification of persons at high risk for overdoses involving illicitly manufactured fentanyl and linkage to risk-reduction services and evidence-based treatment are critical to reducing opioid deaths. Further information on CDC’s state efforts and overdose data is available at https://www.cdc.gov/drugoverdose/index.html.

Changes in Opioid-Involved Overdose Deaths by Opioid Type and Presence of Benzodiazepines, Cocaine, and Methamphetamine — 25 States, July–December 2017 to January–June 2018

R. Matt Gladden, PhD1; Julie O’Donnell, PhD1; Christine L. Mattson, PhD1; Puja Seth, PhD1

From 2013 to 2017, the number of opioid-involved overdose deaths (opioid deaths) in the United States increased 90%, from 25,052 to 47,600.* This increase was primarily driven by substantial increases in deaths involving illicitly manufactured fentanyl (IMF) or fentanyl analogs† mixed with heroin, sold as heroin, or pressed into counterfeit prescription pills (I–3). Methamphetamine-involved and cocaine-involved deaths that

† Fentanyl is a synthetic opioid 50–100 times more potent than morphine and is approved for treatment of severe (typically advanced cancer) pain. Illicitly manufactured fentanyl is manufactured illegally and sold through illegal drug markets for its heroin-like effect. Fentanyl analogs, also known as fentanyl-related substances, are synthetic opioids that are similar in chemical structure to fentanyl but modified to generate distinct substances. Fentanyl analogs vary in potency, with some more potent than fentanyl and others with potency similar to or less than fentanyl. https://www.cdc.gov/drugoverdose/opioids/fentanyl.html; https://www.deadiversion.usdoj.gov/drug_chem_info/hrs.pdf.

References

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co-involved opioids also substantially increased from 2016 to 2017 (4). Provisional 2018\(^\text{††}\) estimates of the number of opioid deaths suggest a small decrease from 2017. Investigating the extent to which decreases occurred broadly or were limited to a subset of opioid types (e.g., prescription opioids versus IMF) and drug combinations (e.g., IMF co-involving cocaine) can assist in targeting of intervention efforts. This report describes opioid deaths during January–June 2018 and changes from July–December 2017 in 25\(^\text{§}\) of 32 states and the District of Columbia participating in CDC’s State Unintentional Drug Overdose Reporting System (SUDORS).** Opioid deaths were analyzed by involvement (opioid determined by medical examiner or coroner to contribute to overdose death) of prescription or illicit opioids, as well as by the presence (detection of the drug in decedent) of co-occurring nonopioid drugs (cocaine, methamphetamine, and benzodiazepines). Three key findings emerged regarding changes in opioid deaths from July–December 2017 to January–June 2018. First, overall opioid deaths decreased 4.6%. Second, decreases occurred in prescription opioid deaths without co-involved illicit opioids and deaths involving non-IMF illicit synthetic opioids (fentanyl analogs and U-series drugs) (5). Third, IMF deaths, especially those with multiple illicit opioids and common nonopioids, increased. Consequently, IMF was involved in approximately two-thirds of opioid deaths during January–June 2018. Notably, during January–June 2018, 62.6% of all opioid deaths co-occurred with at least one common nonopioid drug. To maintain and accelerate reductions in opioid deaths, efforts to prevent IMF-involved deaths and address polysubstance misuse with opioids must be enhanced. Key interventions include broadening outreach to groups at high risk for IMF or fentanyl analog exposure and overdose. Improving linkage to and engagement in risk-reduction services and evidence-based treatment for persons with opioid and other substance use disorders with attention to polysubstance use or misuse is also needed.

\(\text{††} \text{ CDC’s State Unintentional Drug Overdose Reporting System (SUDORS).} \)

\(\text{‡‡} \text{ IMF, fentanyl analogs, heroin, and illicitly manufactured U-series drugs.} \)

\(\text{\textsuperscript{§} Alaska, Connecticut, Delaware, Florida, Georgia, Illinois, Kentucky, Maine, Massachusetts, Minnesota, Missouri, Nevada, New Jersey, New Mexico, North Carolina, Ohio, Oklahoma, Pennsylvania, Rhode Island, Tennessee, Utah, Vermont, Virginia, Washington, and Wisconsin.} \)

\(\text{\textsuperscript{25} The SUDORS captures detailed information on toxicology, death scene investigations, overdose death) of prescription or illicit opioids,} \)

\(\text{\textsuperscript{2017} Broadly or were limited to a subset of opioid types (e.g., prescription opioids versus IMF) and drug} \)

\(\text{\textsuperscript{2018} Provisional 2018} \)

\(\text{\textsuperscript{4} Third, IMF deaths, especially those with multiple illicit opioids and common nonopioids, increased.} \)

\(\text{\textsuperscript{5} Third, IMF deaths, especially those with multiple illicit opioids and common nonopioids, increased.} \)

\(\text{\textsuperscript{6} Notably, during January–June 2018, 62.6% of all opioid deaths co-occurred with at least one common nonopioid drug.} \)

\(\text{\textsuperscript{7} Key interventions include broadening outreach to groups at high risk for IMF or fentanyl analog exposure and overdose.} \)

\(\text{\textsuperscript{8} Improving linkage to and engagement in risk-reduction services and evidence-based treatment for persons with opioid and other substance use disorders with attention to polysubstance use or misuse is also needed.} \)
Numbers of opioid deaths of unintentional and undetermined intent occurring during January–June 2018 and changes from July–December 2017 were analyzed for 25 of the 32 states and the District of Columbia that participate in SUDORS (data for these periods were the most recent and complete). The states abstract death certificate and medical examiner and coroner report data, including death scene investigation and toxicology findings. States list drugs detected in (i.e., contributing to) the opioid death as determined by medical examiners and coroners and all drugs detected (present or co-occurring) by toxicologic tests. Fentanyl and morphine deaths were classified as prescription opioid deaths or illicit opioid deaths based on scene evidence and toxicology findings. Changes in the number of opioid deaths from July–December 2017 to January–June 2018 were analyzed by five opioid types: 1) prescription, 2) IMF, 3) fentanyl analog, 4) heroin, and 5) U-series. Because the frequency and changes in opioid deaths might vary by co-involvement with IMF or other illicit opioids, opioid deaths were also grouped into the following eight mutually exclusive categories: 1) IMF with no other illicit opioids involved; 2) IMF co-involving heroin; 3) IMF co-involving fentanyl analogs; 4) co-involved IMF, heroin, and fentanyl analogs; 5) heroin with no other illicit opioids involved; 6) fentanyl analogs with no other illicit opioids involved; 7) prescription opioids with no illicit opioids involved; and 8) all other opioid combinations. Finally, deaths were analyzed by nonopioids (cocaine, methamphetamine, and benzodiazepines) that are commonly present and involved in opioid deaths. Tracking the presence of commonly occurring nonopioids is important to inform public health action and has implications for treatment approaches. Some opioid deaths were grouped into one or more of the five opioid type categories and nonopioid drug combinations because multiple opioids and nonopioids might be involved in a single death (e.g., an opioid death involving IMF, heroin, cocaine, and benzodiazepine). Changes in numbers of opioid deaths over the analysis period were tested using z-tests or nonoverlapping confidence intervals if the number of deaths was <100. SAS statistical software (version 9.4; SAS Institute, Inc.) was used for all analyses; p-values <0.05 were considered statistically significant.

During January–June 2018, among 13,631 opioid deaths in the 25 states, data on specific opioids involved were available for 13,415 (98.4%). IMF was co-involved in 68.0% of 5,281 heroin deaths and most (82.1%) of 2,678 fentanyl analog deaths (Table 1). In addition, 1,562 (40.5%) of 3,853 prescription opioid deaths co-involved illicit opioids. Opioids commonly involved in opioid deaths were IMF (67.9%), heroin (39.4%), prescription opioids (28.7%), and fentanyl analogs (20.0%) (Table 2). Among categories of deaths involving IMF, those with no other illicit opioids involved, those co-involved with heroin, those co-involved with fentanyl analogs, and those co-involved with heroin and fentanyl analogs accounted for

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§§§ A death was included in the analysis if 1) the fatal injury occurred within, and was reported by, one of the 25 SUDORS states, and 2) the death was classified as an overdose death involving opioids either through review of the medical examiner/coroner report or the death certificate had International Classification of Diseases, Tenth Revision underlying cause-of-death codes X40–44 (unintentional) or Y10–Y14 (undetermined intent) and multiple cause-of-death codes T40.0, T40.1, T40.2, T40.3, T40.4, or T40.6. Data for this report were downloaded on June 26, 2019, and might differ from earlier or future reports because states continually update death data and investigations of suspected drug overdose deaths might involve lengthy investigations.

¶¶¶ All substances detected were categorized as contributing to the death when deaths were classified by the medical examiner/coroner as a drug overdose and toxicology results were available, but no information was available on the specific drugs contributing to death.

††† Fentanyl was classified on the basis of toxicology, scene, and witness evidence that indicated the likely source as either prescription (e.g., scene evidence of fentanyl patches at the overdose location) or illicit (e.g., toxicology evidence of a fentanyl analog, or scene or witness evidence of illicit drug use including injection or snorting). If evidence of prescription or illicit use was not available, fentanyl was categorized as IMF because the vast majority of fentanyl overdose deaths involve IMF. Morphine in the absence of 6-acetylmorphine (a metabolite of heroin) was classified as likely prescription morphine (scene or witness evidence of prescription morphine use) or as likely heroin (toxicology evidence of heroin impurities or other illicit drug detected or scene or witness evidence that indicated injection, illicit drug use, or a history of heroin use). If evidence of prescription or illicit use was not available, morphine was categorized as prescription because the investigation did not obtain scene evidence of heroin use or detect 6-acetylmorphine. If morphine was detected along with 6-acetylmorphine, it was classified as heroin.

§§§§ Substances coded as prescription opioids were oxycodone, oxymorphone, hydrocodone, hydromorphone, tramadol, buprenorphine, methadone, prescription fentanyl, morphine, codeine, meperidine, tapentadol, dextorphan, levorphanol, propoxyphene, noscapine, and pentazocine. Also included as prescription drugs were brand names (e.g., Opana) and metabolites (e.g., normtramadol) of these substances and combinations of these substances and nonopioids (e.g., acetylmophonen-oxycodone). Substances coded as illicit opioids were IMF, heroin, fentanyl analogs, and other illicit synthetic opioids, such as U-series drugs (e.g., U-47700), AH-7921, and MT-45. This analysis does not distinguish between prescription drugs prescribed to the decedent and those that were diverted. Data on AH-7921 and MT-45 were not included because they were involved in fewer than five deaths.

†††† Only commonly occurring (present in >10% of opioid deaths) and contributing (involved in >50% of opioid deaths in which present) nonopioids were included. Cutoff for nonopioid inclusion was determined by a review of the data.

To verify that changes in population size did not account for observed changes, sensitivity analyses were conducted in which crude population death rates per 100,000 residents for July–December 2017 were compared with death rates for January–June 2018 using z-tests or nonoverlapping confidence intervals if the number of deaths was <100. U.S. Department of Commerce’s Bureau of Economic Analysis data were used to estimate populations for the 25 states in the middle of the last quarter of 2017 and the middle of the second quarter of 2018 (https://apps.bea.gov/ITable/index.cfm/). For states collecting data on opioid deaths in a subset of counties, the estimates of state population were used and should have introduced minimal bias because of the >75% coverage of opioid deaths in these states and minimal population changes over a 6-month period.
TABLE 1. Number and percentage of opioid overdose deaths that co-involved another opioid, by opioid type (illicitly manufactured fentanyl [IMF],† fentanyl analogs,‡ heroin, and prescription opioids§) — 25 states,§ State Unintentional Drug Overdose Reporting System (SUDORS), January–June 2018

<table>
<thead>
<tr>
<th>Opioid type involved in opioid death**</th>
<th>No. of deaths</th>
<th>No. of deaths with co-involved opioid types (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Any suspected IMF</td>
<td>9,105</td>
<td>2,199 (24.2)</td>
</tr>
<tr>
<td>Any fentanyl analog</td>
<td>2,678</td>
<td>2,199 (82.1)</td>
</tr>
<tr>
<td>Any suspected heroin</td>
<td>5,281</td>
<td>1,172 (22.2)</td>
</tr>
<tr>
<td>Any prescription opioid</td>
<td>3,853</td>
<td>1,250 (32.4)</td>
</tr>
</tbody>
</table>

** Among fentanyl-involved deaths, 87.2%, 11.2%, and 1.6% were suspected to involve IMF, had insufficient data to classify the fentanyl death as IMF or prescription fentanyl, and were suspected to involve prescription fentanyl, respectively. Because the majority of identified cases involved IMF, and characteristics of unclassified fentanyl deaths were more similar to IMF-involved deaths than to prescription fentanyl-involved deaths, unclassified fentanyl deaths were categorized as suspected IMF-involved.

† Fentanyl analog-involved deaths included deaths involving carfentanil, acetylfentanyl, acrylfentanyl, furanylfentanyl, 3-methylfentanyl, butyrylfentanyl, cyclopropyfentanyl, crotonylfentanyl, 4/para-fluorofentanyl, 4/para-fluorobutyrylfentanyl, 4/para-isobutyrylfentanyl, cyclopentylfentanyl, methoxyacetyl fentanyl, isobutyrylfentanyl, furanylfentanyl, methoxybutyrylfentanyl, benzylfentanyl, valeryl fentanyl, alpha-methylfentanyl, tetrahydrofuranylfentanyl, octofentanyl, betahydroxythiofentanyl, alfentanil, sufentanil, methylcarfentanil, methylthiofentanyl, phenylfentanyl, omethyacetyl fentanyl, and isovalerylfentanyl.

§ Included any opioid deaths involving prescription opioids (oxycodone, oxymorphone, hydrocodone, hydromorphone, tramadol, buprenorphine, methadone, morphine, codeine, prescription fentanyl, meperidine, tapentadol, dextrophen, levorphanol, propoxyphene, noscapine, and pentazocine). Other drugs might have been involved or co-occurred with the prescription opioid.

§§ U-series deaths were not reported in the analyses because only 63 deaths involved U-series drugs in the 25 SUDORS states during January–June 2018.

¶ Any illicit opioid other than that listed in drug category. For deaths involving illicit opioids (IMF, fentanyl analogs, and heroin) the illicit drug is excluded from this column. For example, 52.6% of IMF deaths co-involved at least one fentanyl analog, heroin, or U-series drug.

### Table Notes:

- 32.2%, 19.1%, 8.7%, and 7.5% of deaths, respectively. Heroin without other illicit opioids involved accounted for 11.4% of deaths, fentanyl analogs with no other illicit opioids involved for 2.3%, prescription opioids with no illicit opioids involved for 17.1%, and all other opioid combinations for 1.6%. In the Midwest, Northeast, and South U.S. Census regions, deaths involving any IMF were more common than those involving any heroin. In the West, heroin-involved deaths (47.5%) were more common than were IMF-involved deaths (15.8%) (data not shown).

Three principal changes occurred in opioid deaths from July–December 2017 to January–June 2018. First, opioid deaths in the 25 states declined by 4.6% (Table 2). Second, declines occurred in prescription opioid deaths with no co-involved illicit opioids (10.6%) and non-IMF illicit synthetic opioid deaths, including fentanyl analogs (19.0% decline) and U-series drugs (75.1% decline). With the exception of acetylfentanyl, decreases in fentanyl analog deaths occurred broadly across all fentanyl analogs (52.7% decline). Acetylfentanyl deaths co-involved IMF showed a sharp increase (57.5%). Third, IMF deaths increased by 11.1% overall, with increases of 9.5%–33.0% in those co-involving other illicit opioids and 9.4% among those with no other illicit opioids involved. Illicit opioid overdose deaths involving heroin and fentanyl analogs increased when IMF was co-involved, but decreased when IMF and other illicit opioids were not co-involved. Specifically, increases occurred in IMF deaths co-involving heroin (9.5%), fentanyl analogs (11.4%), and both heroin and fentanyl analogs (33.0%). In contrast, substantial declines were observed in heroin deaths with no other illicit opioids involved (16.6% decline) and fentanyl analog deaths with no other illicit opioids involved (67.9% decline). Declines in heroin deaths with no other illicit opioids involved were offset by increases in heroin deaths co-involved IMF, resulting in no significant change in heroin deaths.

The majority of opioid deaths (62.6%) co-occurred with one or more of the following drugs: benzodiazepines, cocaine, and methamphetamine, which were each present in 32.5%, 34.0%, and 12.1% of deaths, respectively. From July–December 2017 to January–June 2018, opioid deaths without benzodiazepines, cocaine, or methamphetamine decreased 8.0%, and opioid deaths co-occurring with benzodiazepines significantly decreased 5.7% (Table 3). Conversely, opioid deaths co-occurring with methamphetamine significantly increased by 14.6%. IMF deaths that co-occurred with benzodiazepines, cocaine, and methamphetamine significantly increased from July–December 2017 to January–June 2018 by 11.3%, 14.0%, and 31.0%, respectively, as IMF deaths without benzodiazepines, cocaine, or methamphetamine increased 6.7%.

**Discussion**

Among 25 states participating in SUDORS, three major changes in opioid deaths from July–December 2017 to January–June 2018 were identified. These included 1) overall decreases in opioid overdose deaths; 2) decreases in both prescription opioid deaths without co-involved illicit opioids and non-IMF illicit synthetic opioids (i.e., fentanyl analogs and U-series drugs) deaths; and 3) increases in IMF deaths, especially those with heroin, fentanyl analogs or nonopioid drugs. Also, at least one nonopioid drug (benzodiazepines,
### Table 2. Change in the number of opioid overdose deaths, by opioid type, eight common opioid drug combinations, and commonly co-occurring nonopioids (cocaine, methamphetamine, and benzodiazepines) — 25 states, State Unintentional Drug Overdose Reporting System (SUDORS), July–December 2017 to January–June 2018

| Characteristic | Opioid deaths with information on involved opioids, Jan–Jun 2018, no. (%) | Change in no. of opioid deaths, Jul–Dec 2017 to Jan–Jun 2018, no. (%) | % Nonopioid drugs commonly present in opioid deaths, Jan–Jun 2018
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Total opioid overdose deaths</td>
<td>13,415 (100)</td>
<td>648 (−4.6)</td>
</tr>
<tr>
<td></td>
<td>% of deaths with contributing nonopioids present</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Opioid drug class or drug involved in opioid deaths</td>
<td>Any prescription opioid†††</td>
<td>3,853 (28.7)</td>
<td>−271 (−6.6)</td>
</tr>
<tr>
<td></td>
<td>Any illicit opioid††</td>
<td>11,124 (82.9)</td>
<td>−376 (−3.3)</td>
</tr>
<tr>
<td></td>
<td>Any suspected IMF†</td>
<td>9,105 (67.9)</td>
<td>−910 (11.1)</td>
</tr>
<tr>
<td></td>
<td>Any suspected heroin</td>
<td>5,281 (39.4)</td>
<td>−83 (−1.5)</td>
</tr>
<tr>
<td></td>
<td>Any fentanyl analog†††</td>
<td>2,678 (20.0)</td>
<td>−627 (−19.0)</td>
</tr>
<tr>
<td></td>
<td>Any U-series†††</td>
<td>63 (0.5)</td>
<td>−190 (−75.1)</td>
</tr>
</tbody>
</table>

Commonly mutually exclusive combinations of opioids involved in opioid deaths

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Opioid combinations co-involving IMF</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>IMF with no other illicit opioids</td>
</tr>
<tr>
<td></td>
<td>IMF with heroin</td>
</tr>
<tr>
<td></td>
<td>IMF with fentanyl analogs</td>
</tr>
<tr>
<td></td>
<td>IMF with heroin and fentanyl analogs</td>
</tr>
<tr>
<td></td>
<td>All other combinations of opioids</td>
</tr>
</tbody>
</table>

Ilicit opioid combinations not co-involving IMF

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Any acetylfentanyl</th>
<th>1,716 (12.8)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Acetylfentanyl with IMF</td>
<td>1,685 (12.6)</td>
</tr>
<tr>
<td></td>
<td>Acetylfentanyl no IMF</td>
<td>31 (0.2)</td>
</tr>
<tr>
<td></td>
<td>All other fentanyl analogs</td>
<td>1,100 (8.2)</td>
</tr>
<tr>
<td></td>
<td>Other fentanyl analogs with IMF</td>
<td>645 (4.8)</td>
</tr>
<tr>
<td></td>
<td>Other fentanyl analogs no IMF</td>
<td>453 (3.4)</td>
</tr>
</tbody>
</table>

Abbreviations: IMF = illicitly manufactured fentanyl; NA = not applicable.


† Only the two most frequently co-occurring types of stimulants (cocaine and methamphetamine) are reported because other types of stimulants such as amphetamines did not meet inclusion criteria.

§ Statistically significantly change from July–December 2017 to January–June 2018 based on z-tests or nonoverlapping confidence intervals if the number of deaths was <100 (p<0.05).

¢ Cocaine, methamphetamine, and benzodiazepines, this row reports a percentage calculated by dividing the number of opioid deaths in which the drug was present and reported as contributing to the opioid death (numerator) by the number of opioid deaths in which the drug was present (i.e., detected by toxicology tests) irrespective of whether it contributed to the opioid death (denominator).

** Percentage of all opioid deaths in which cocaine, methamphetamine, or benzodiazepines contributed to death.

†† An opioid death might involve multiple opioids. Thus, total opioid deaths and change in opioid deaths will be different than the sum of the deaths associated with each opioid type.

† † Other nonopioid drugs might have been involved or co-occurred.

††† Included any opioid death involving IMF, heroin, fentanyl analogs, or U-series drugs. Other drugs might have been involved or co-occurred.

††‡ Among fentanyl-involved deaths, 87.2%, 11.2%, and 1.6% were suspected to involve IMF, had insufficient data to classify the fentanyl death as IMF or prescription fentanyl, and were suspected to involve prescription fentanyl, respectively. Because the majority of identified cases involved IMF, and characteristics of unclassified fentanyl deaths were more similar to IMF-involved deaths than to prescription fentanyl-involved deaths, unclassified fentanyl deaths were categorized as suspected IMF-involved.

†††† Fentanyl analog deaths included deaths involving carfentanil, acetylfentanyl, acrylfentanyl, furanylfentanyl, 3-methylfentanyl, butyrylfentanyl, cyclopropylfentanyl, crotylfentanyl, 4/para-fluoroacrylfentanyl, 4/para-fluoroacetylphenylacetic acid, cyclopropylfentanyl, methoxyacetylphenylacetic acid, isobutylphenylacetic acid, isobutylphenylacetic acid, benzylphenylacetic acid, valerylphenylacetic acid, alpha-methylfentanyl, tetrahydrofurfurylfentanyl, ofentanil, betahydroxythiofentanyl, alfentanil, sufentanil, methylcarfentanil, methyllithiofentanyl, phenylfentanyl, methoxymethyllithiofentanyl, and isovalerylfentanyl. Nonfentanyl-related synthetic opioids with no authorized medical uses. U-series drug deaths include those involving U-47700 and its analogs U-48800 and U-49900. U-47700, a nonfentanyl benzamide compound developed by a pharmaceutical company, is not authorized for medical use in the United States and is currently distributed illicitly for its heroin-like effect. Deaths involving U-50488 and U-51754 were also included in this category, but each was involved in five or fewer deaths.

§§§ Six categories are combinations of the illicit opioids involved in death (IMF, heroin, fentanyl analog, and U-series) that were involved in >200 deaths during January–June 2018. These deaths might co-involve prescription opioids and co-occur with nonopioids. The “prescription opioids with no illicit opioid” category includes only deaths involving prescription opioids with no illicit opioid co-involvement but might co-occur with other nonopioid drugs. The “all other combinations of opioids” category includes opioid deaths that involved opioid drug combinations not listed, primarily opioid deaths involving U-series drugs or heroin deaths co-involving fentanyl analogs.

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person misusing heroin,**** expanding naloxone access,††††† or changing behaviors of persons injecting drugs to reduce the likelihood of an IMF-involved overdose (8).

Evidence suggests that persons using powdered heroin are often unaware of whether IMF or fentanyl analogs are present in illicit products (2,3). Consequently, IMF, heroin, and fentanyl analog combinations in opioid deaths might represent mixed drug products rather than purposeful co-use. IMF deaths without other illicit opioids co-involved and IMF deaths co-involved heroin are the two most frequent drug combinations in opioid deaths and are consistent with combinations found when drug products test positive for fentanyl by law enforcement (2). As IMF supply expanded during January–June, 2018 (9), a large, nationally accredited laboratory reported that the majority of patients east of the Mississippi River who tested positive for heroin also tested positive for fentanyl, and this percentage increased in early 2018.§§§§§ This suggests increased mixing of IMF with powdered heroin and fewer heroin-only products, consistent with the increases in IMF deaths co-involved heroin and decreases in heroin deaths without IMF documented in this report. In Western states, heroin deaths predominated, possibly because of the limited mixing of IMF with powdered heroin and fewer heroin-only products, consistent with the increases in IMF deaths co-involved heroin and decreases in heroin deaths without IMF documented in this report. In Western states, heroin deaths predominated, possibly because of the limited mixing of IMF with powdered heroin.

Although concerning, the 6-month 11.1% increase in IMF deaths in the 25 states is smaller than the approximate doubling of U.S. fentanyl deaths each year during 2014–2016.§§§§ Because IMF is distributed primarily in the powder heroin market,**** slower increases in IMF deaths might reflect successes in one or more objectives: reducing the number of persons who initiate heroin use, increasing treatment access for persons misusing heroin,***** expanding naloxone access,††††† or changing behaviors of persons injecting drugs to reduce the likelihood of an IMF-involved overdose (8).

Evidence suggests that persons using powdered heroin are often unaware of whether IMF or fentanyl analogs are present in illicit products (2,3). Consequently, IMF, heroin, and fentanyl analog combinations in opioid deaths might represent mixed drug products rather than purposeful co-use. IMF deaths without other illicit opioids co-involved and IMF deaths co-involved heroin are the two most frequent drug combinations in opioid deaths and are consistent with combinations found when drug products test positive for fentanyl by law enforcement (2). As IMF supply expanded during January–June, 2018 (9), a large, nationally accredited laboratory reported that the majority of patients east of the Mississippi River who tested positive for heroin also tested positive for fentanyl, and this percentage increased in early 2018.§§§§§ This suggests increased mixing of IMF with powdered heroin and fewer heroin-only products, consistent with the increases in IMF deaths co-involved heroin and decreases in heroin deaths without IMF documented in this report. In Western states, heroin deaths predominated, possibly because of the limited mixing of IMF with powdered heroin and fewer heroin-only products, consistent with the increases in IMF deaths co-involved heroin and decreases in heroin deaths without IMF documented in this report. In Western states, heroin deaths predominated, possibly because of the limited mixing of IMF with powdered heroin.

Although concerning, the 6-month 11.1% increase in IMF deaths in the 25 states is smaller than the approximate doubling of U.S. fentanyl deaths each year during 2014–2016.§§§§ Because IMF is distributed primarily in the powder heroin market,**** slower increases in IMF deaths might reflect successes in one or more objectives: reducing the number of persons who initiate heroin use, increasing treatment access for persons misusing heroin,***** expanding naloxone access,††††† or changing behaviors of persons injecting drugs to reduce the likelihood of an IMF-involved overdose (8).

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prescription pills that contain IMF might increase the risk of overdose in persons who use prescription medications not prescribed to them, especially opioid pain relievers (2). The majority of opioid deaths co-occurred with benzodiazepines, cocaine, or methamphetamine highlighting the need to address polysubstance use in the prevention of overdoses and treatment of opioid misuse. Increases in opioid deaths, especially IMF deaths, co-occurring with methamphetamine are consistent with previous reports (4) and with increases in methamphetamine supply (9) and methamphetamine use among persons seeking treatment for opioid misuse (10). Moreover, IMF deaths co-occurring with benzodiazepines and cocaine increased during January–June 2018 even as overall opioid deaths co-occurring with benzodiazepines and cocaine decreased or did not significantly change, respectively. Increases in IMF deaths co-occurring with cocaine are consistent with previous reports (4) and with high co-use of cocaine among persons injecting heroin and outbreaks linked to rare but increasing numbers of drug products that mix IMF and cocaine (2).

The findings in this report are subject to at least five limitations. First, toxicology testing and classification protocols vary over time and across jurisdictions, which affects whether drugs were detected and classified as contributing to death. Second, misclassification of prescription and illicit substances might occur, but this was minimized by using detailed toxicology results and scene evidence. Third, focus on drugs commonly involved in opioid deaths might obscure emerging drug issues. Fourth, patterns in drugs involved in opioid deaths might vary across states and demographic groups. Finally, findings are limited to the 25 states participating in SUDORS and might not be generalizable to other states. Increases in IMF deaths involving multiple illicit opioids and benzodiazepines, cocaine, and methamphetamine (nonopioids) highlight the need to better understand how the risk of IMF overdose varies by illicit product potency, variation in potency, and form (e.g., powder or counterfeit pill) and a person’s tolerance or polysubstance use patterns. In response, CDC’s Overdose Data to Action funding expands SUDORS from including only opioid-involved deaths to including all drug overdose deaths to better understand increases in IMF and stimulant and drug combination deaths (with and without opioids), as well as identify emerging threats. Key interventions include broadening outreach to groups at high risk for IMF or fentanyl analog exposure and overdose. Improving linkage to and engagement in risk-reduction services and evidence-based treatment for persons with opioid and other substance use disorders with attention to polysubstance use or misuse is also needed.

Acknowledgments

Jurisdictions participating in CDC’s Enhanced State Opioid Overdose Surveillance (ESOOS) program and providing data in the State Unintentional Drug Overdose Reporting System, including state and jurisdictional health departments, vital registrar offices, and coroner and medical examiner offices; the CDC ESOOS team, Division of Unintentional Injury Prevention, National Center for Injury Prevention and Control, CDC; Bruce Goldberger, University of Florida College of Medicine, Gainesville, Florida.

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References


Surveillance of U.S. breastfeeding duration and exclusivity has historically reported estimates among all infants, regardless of whether they had initiated breastfeeding. These surveillance estimates have consistently shown that non-Hispanic black (black) infants are less likely to breastfeed, compared with other racial/ethnic groups. Less is known about disparities in breastfeeding duration when calculated only among infants who had initiated breastfeeding, compared with surveillance estimates based on all infants. CDC analyzed National Immunization Survey-Child (NIS-Child) data for infants born in 2015 to describe breastfeeding duration and exclusivity at ages 3 and 6 months among all black and non-Hispanic white (white) infants, and among only those who had initiated breastfeeding. When calculated among all infants regardless of breastfeeding initiation, breastfeeding differences between black and white infants were 14.7 percentage points (95% confidence interval [CI] = 10.7–18.8) for any breastfeeding at age 3 months and were significantly different for both any and exclusive breastfeeding at both ages 3 and 6 months. Among only infants who had initiated breastfeeding, the magnitude of black-white differences in breastfeeding rates were smaller. This was most notable in rates of any breastfeeding at 3 months, where the percentage point difference between black and white infants was reduced to 1.2 (95% CI = -2.3–4.6) percentage points and was no longer statistically significant. Black-white disparities in breastfeeding duration result, in part, from disparities in initiation. Interventions both to improve breastfeeding initiation and to support continuation among black mothers might help reduce disparities.

Breastfeeding has numerous health benefits for infants and mothers. Breastfed infants have reduced risk for ear, respiratory, and gastrointestinal infections and might be less likely to develop asthma, obesity, and diabetes (1). Mothers who breastfeed have a lower risk for developing type 2 diabetes, hypertension, and breast and ovarian cancers (2). U.S. breastfeeding surveillance has consistently demonstrated that rates of breastfeeding initiation, duration, and exclusivity are 10–20 percentage points lower among black infants, compared with white infants.†

NIS-Child is an ongoing, nationally representative random-digit–dialed telephone survey of U.S. households of children aged 19–35 months. From 2011 to 2017, the NIS-Child used a dual landline and mobile telephone sample frame. Although NIS-Child primarily assesses childhood vaccination coverage, breastfeeding questions were added in 2001 and are the primary data source for U.S. breastfeeding surveillance. Each cross-sectional survey includes children born in 3 different calendar years; for this analysis of infants born in 2015, data from the 2016–2017 surveys were combined, consistent with national surveillance estimates. Landline sample response rates were 55.7% in 2016 and 51.9% in 2017. Mobile telephone sample response rates were 32.1% in 2016 and 25.0% in 2017. Children’s breastfeeding history and race/ethnicity were reported by their parents or guardians.

Breastfeeding initiation rates were calculated for black and white infants born in 2015. Rates of any breastfeeding and exclusive breastfeeding (defined as only breast milk and no solids, water, or other liquids) at ages 3 and 6 months were calculated for black and white infants using two sets of denominators. The first denominator included all infants of the respective racial/ethnic group regardless of breastfeeding initiation. The second denominator included only infants of the respective racial/ethnic group who had initiated breastfeeding. The absolute percentage point difference in each breastfeeding rate between black and white infants was also estimated (hereafter, black-white difference). Estimates were weighted and accounted for the NIS complex sampling design. Data were analyzed using SAS (version 9.4; SAS Institute) and SUDAAN (version 11.0.3; RTI International).

Black women were more likely than were white women to have incomes <100% of the poverty level (49.3% versus 17.8%), to receive Special Supplemental Nutrition Program for Women, Infants, and Children benefits (78.2% versus 34.1%), and to be unmarried (65.5% versus 23.9%); they also had less education and were younger (Table 1). In 2015, 69.4% of black infants initiated breastfeeding, compared with 85.9% of white infants, a difference of 16.5 percentage points (95% CI = 10.7–18.8) for any breastfeeding at age 3 months, where the percentage point difference between black and white infants was 14.7 percentage points (95% CI = 10.7–18.8) for any breastfeeding at age 3 months and were significantly different for both any and exclusive breastfeeding at both ages 3 and 6 months. Among only infants who had initiated breastfeeding, the magnitude of black-white differences in breastfeeding rates were smaller. This was most notable in rates of any breastfeeding at 3 months, where the percentage point difference between black and white infants was reduced to 1.2 (95% CI = -2.3–4.6) percentage points and was no longer statistically significant. Black-white disparities in breastfeeding duration result, in part, from disparities in initiation. Interventions both to improve breastfeeding initiation and to support continuation among black mothers might help reduce disparities.

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Among all infants, black infants had a significantly lower rate of any breastfeeding at age 3 months (58.0%) than did white infants.

‡https://www.cdc.gov/breastfeeding/data/nis_data/methods.html.
TABLE 1. Demographic characteristics of non-Hispanic white and non-Hispanic black infants born in 2015 included in national prevalence estimates of breastfeeding initiation and duration at ages 3 and 6 months — National Immunization Survey-Child, United States, 2016–2017*

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Non-Hispanic white (n = 9,907)</th>
<th>Non-Hispanic black (n = 1,607)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No.</td>
<td>% (95% CI)†</td>
</tr>
<tr>
<td>% of poverty level§</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;100</td>
<td>1,312</td>
<td>17.8 (16.5–19.1)</td>
</tr>
<tr>
<td>100–199</td>
<td>1,703</td>
<td>18.7 (17.4–20.0)</td>
</tr>
<tr>
<td>200–399</td>
<td>2,909</td>
<td>27.9 (26.5–29.3)</td>
</tr>
<tr>
<td>400–599</td>
<td>1,967</td>
<td>17.7 (16.5–19.0)</td>
</tr>
<tr>
<td>≥600</td>
<td>2,016</td>
<td>17.9 (16.6–19.3)</td>
</tr>
<tr>
<td>Recipient of WIC§</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>2,723</td>
<td>34.1 (32.5–35.8)</td>
</tr>
<tr>
<td>No, but eligible</td>
<td>836</td>
<td>9.0 (8.1–9.8)</td>
</tr>
<tr>
<td>Ineligible</td>
<td>6,298</td>
<td>56.9 (55.2–58.6)</td>
</tr>
<tr>
<td>Mother’s education</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Less than high school diploma or GED</td>
<td>460</td>
<td>7.4 (6.3–8.4)</td>
</tr>
<tr>
<td>High school diploma or GED</td>
<td>1,394</td>
<td>20.2 (18.8–21.6)</td>
</tr>
<tr>
<td>Some college</td>
<td>2,435</td>
<td>23.4 (22.0–24.8)</td>
</tr>
<tr>
<td>College graduate</td>
<td>5,618</td>
<td>49.1 (47.4–50.7)</td>
</tr>
<tr>
<td>Mother’s age group (yrs)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;20</td>
<td>70</td>
<td>1.1 (0.7–1.5)</td>
</tr>
<tr>
<td>20–29</td>
<td>2,943</td>
<td>34.4 (32.8–36.1)</td>
</tr>
<tr>
<td>≥30</td>
<td>6,894</td>
<td>64.5 (62.8–66.1)</td>
</tr>
<tr>
<td>Mother’s marital status</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Married</td>
<td>8,097</td>
<td>76.1 (74.6–77.7)</td>
</tr>
<tr>
<td>Unmarried</td>
<td>1,810</td>
<td>23.9 (22.3–25.4)</td>
</tr>
</tbody>
</table>

Abbreviations: GED = general educational development certificate; WIC = Special Supplemental Nutrition Program for Women, Infants, and Children.

† Statistics in this table are based on participants who responded to questions about any breastfeeding at ages 3 and 6 months (N = 11,514). Sample sizes are slightly smaller for participants who also responded to questions about exclusive breastfeeding at ages 3 and 6 months.
§ Ratio of self-reported family income to the poverty threshold value defined by the U.S. Census Bureau.
¶ Sample sizes for the proportions of participants receiving WIC are slightly smaller due to missing data on WIC status.

TABLE 2. Breastfeeding initiation and duration at ages 3 and 6 months* among non-Hispanic black and non-Hispanic white infants born in 2015 — National Immunization Survey-Child, United States, 2016–2017†

<table>
<thead>
<tr>
<th>Breastfeeding indicator</th>
<th>All infants</th>
<th>Infants who had initiated breastfeeding</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Non-Hispanic white</td>
<td>Non-Hispanic black</td>
</tr>
<tr>
<td></td>
<td>No.</td>
<td>% (95% CI)†</td>
</tr>
<tr>
<td>Initiated breastfeeding</td>
<td>9,907</td>
<td>85.9 (84.7 to 87.1)</td>
</tr>
<tr>
<td>Any breastfeeding at age 3 mos</td>
<td>9,907</td>
<td>72.7 (71.2 to 74.2)</td>
</tr>
<tr>
<td>Exclusive breastfeeding through age 3 mos</td>
<td>9,537</td>
<td>53.0 (51.4 to 54.7)</td>
</tr>
<tr>
<td>Any breastfeeding at age 6 mos</td>
<td>9,907</td>
<td>62.0 (60.4 to 63.6)</td>
</tr>
<tr>
<td>Exclusive breastfeeding through age 6 mos</td>
<td>9,537</td>
<td>29.5 (28.0 to 31.1)</td>
</tr>
</tbody>
</table>

Abbreviations: CI = confidence interval; N/A = not applicable.
* Breastfeeding initiation was determined according to participant’s response to the question “Was [child] ever breastfed or fed breast milk?” Breastfeeding duration was determined according to participant’s response to the question “How old was [child’s name] when he/she was first fed breast milk?” Exclusive breastfeeding was defined as only breast milk (no solids, no water, and no other liquids). To assess the duration of exclusive breastfeeding, participants were asked two questions about age: 1) “How old was [child’s name] when he/she was first fed formula?” and 2) “How old was [child’s name] when he/she was first fed anything other than breast milk or formula?” (This includes juice, cow’s milk, sugar water, baby food, or anything else that [child] might have been given, even water).
§ Differences in breastfeeding rates between non-Hispanic black and non-Hispanic white infants.
infants (72.7%); at age 6 months, the rates were 44.7% among black infants and 62.0% among white infants (p<0.05). Rates for exclusive breastfeeding at age 3 months were 36.0% among black infants and 53.0% among white infants; at age 6 months, the rates were 17.2% among black infants and 29.5% among white infants (p<0.05) (Table 2). At age 3 months, black-white differences were 14.7 percentage points for any breastfeeding (95% CI = 10.7–18.8) and 17.0 percentage points for exclusive breastfeeding (95% CI = 12.9–21.2). At age 6 months, black-white differences were 17.3 percentage points for any breastfeeding (95% CI = 13.1–21.4) and 12.4 percentage points for exclusive breastfeeding (95% CI = 8.9–15.8) (Table 2).

Among only infants who had initiated breastfeeding, the magnitude of black-white differences in any and exclusive breastfeeding rates were smaller (Table 2). This was most notable in rates of any breastfeeding at 3 months, where the percentage point difference between black and white infants was reduced from 14.7 (95% CI = 10.7–18.8) to 1.2 (95% CI = -2.3–4.6) percentage points; this difference was no longer statistically significant. The black-white difference in exclusive breastfeeding at age 3 months was reduced from 17.0 percentage points (95% CI = 12.9–21.2) to 9.9 percentage points (95% CI = 5.0–14.7), in any breastfeeding at 6 months from 17.3 percentage points (95% CI = 13.1–21.4) to 7.8 percentage points (95% CI = 3.3–12.3), and in exclusive breastfeeding at age 6 months from 12.4 percentage points (8.9–15.8) to 9.7 percentage points (95% CI = 5.1–14.2).

**Discussion**

Surveillance of U.S. breastfeeding duration and exclusivity, including monitoring for Healthy People 2020 objectives, reports estimates among all infants, regardless of whether they had initiated breastfeeding. The findings in this report demonstrate that differences between black and white infants in any and exclusive breastfeeding at ages 3 and 6 months are caused, in part, by racial/ethnic differences in breastfeeding initiation. Interventions to improve breastfeeding initiation and support continuation among black mothers might be important to closing the black-white gap in duration.

Black mothers disproportionately experience a number of barriers to breastfeeding, including lack of knowledge about breastfeeding; lack of peer, family, and social support; insufficient education and support from health care settings; and concerns about navigating breastfeeding and employment (3). Subjective norms, or perceptions of approval from others who are important to the person (e.g., family members), are important drivers of breastfeeding behaviors, particularly among black women (3). Increasing interpersonal support for breastfeeding might help increase breastfeeding initiation and duration among black women, who might lack breastfeeding role models in their social networks and be more likely to face negative perceptions of breastfeeding among their peers and communities (3,4). For example, peer counseling might increase breastfeeding initiation and duration among black mothers (3).

In the United States, the rate of implementation of evidence-based maternity care practices supportive of breastfeeding is lower among maternity care facilities in neighborhoods with larger black populations (5). Hospitals’ use of such practices, which include helping women initiate breastfeeding within the first hour of birth and not providing breastfeeding infants with infant formula without a medical indication, increases rates of breastfeeding initiation, duration, and exclusivity (6). A recent analysis indicated that making improvements in these practices among maternity care facilities in four southern states reduced black-white disparities in breastfeeding initiation (7).

Returning to work is another major barrier to breastfeeding initiation and continuation, particularly for black women (3). A woman’s plans for returning to work are associated with her intention to breastfeed; specifically, women planning to return to work before 12 weeks postpartum, planning to work full-time, or both were less likely to intend to exclusively breastfeed, compared with women planning to return to work after 12 weeks postpartum, planning to work part-time, or both (8). Black women, especially those with a low income, return to work earlier than do women in other racial/ethnic groups and are more likely to experience challenges to breastfeeding or expressing milk, including inflexible work hours (9). Policies that enable taking paid leave after giving birth, flexible work schedules, and support for breastfeeding or expressing milk at work might help improve breastfeeding intention, initiation, and duration.**

The findings in this report are subject to at least three limitations. First, response rates averaged 53.8% for the landline sample and 28.6% for the mobile telephone sample; further, households without a telephone are not represented. The possibility exists that selection bias occurs even after adjusting weights for nonresponse and noncoverage. Second, maternal reports of breastfeeding behaviors could be subject to recall bias because mothers reported these behaviors when their children were aged 19–35 months and to social desirability bias because of a desire to provide socially acceptable responses. However, maternal recall of breastfeeding behavior has been found to be valid and reliable, especially when recalled within 3 years (10). Finally, although this report focuses only on black-white

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Summary
What is already known on this topic?
Rates of breastfeeding duration and exclusivity, calculated for all infants regardless of whether they had initiated breastfeeding, are lower among black infants than among white infants.

What is added by this report?
Among infants who had initiated breastfeeding, differences between black infants and white infants in any and exclusive breastfeeding at ages 3 and 6 months were smaller but still present.

What are the implications for public health practice?
Increasing rates of breastfeeding initiation and supporting continuation of breastfeeding among black women might help reduce disparities in breastfeeding duration. Strategies might include improving peer and family support, access to evidence-based maternity care, and employment support.

breastfeeding differences, lower rates of breastfeeding duration and exclusivity among Hispanic infants, compared with non-Hispanic white infants, have been documented (3). However, because Hispanic and white infants have similar rates of breastfeeding initiation, the methods applied in this report did not affect estimates of breastfeeding duration and exclusivity.

Breastfeeding provides optimal nutrition to infants and provides health benefits for both infants and mothers, and CDC works to increase breastfeeding rates among all mothers in the United States. In order to address disparities in breastfeeding duration, continued efforts are needed to increase rates of breastfeeding initiation and support continuation of breastfeeding among black women. Closing the black-white gap in breastfeeding duration might require efforts of multiple groups. Families, hospitals, and employers can help black women initiate and continue breastfeeding, thereby providing their infants with optimal nutrition.

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All authors have completed and submitted the International Committee of Medical Journal Editors form for disclosure of potential conflicts of interest. No potential conflicts of interest were disclosed.

References
**Notes from the Field**

### Mumps in Detention Facilities that House Detained Migrants — United States, September 2018–August 2019

Jessica Leung, MPH\(^1\); Diana Elson, DrPH\(^2\); Kelsey Sanders, MPH\(^3\); Mona Marin, MD\(^1\); Greg Leos, MPH\(^3\); Brandy Cloud, DNP\(^2\); Rebecca J. McNall, PhD\(^1\); Carole J. Hickman, PhD\(^1\); Mariel Marlow, PhD\(^1\)

On October 12, 2018, five confirmed cases of mumps among migrants who had been transferred between two detention facilities were reported by the facilities to the Texas Department of State Health Services (TDSHS). By December 11, eight Texas detention facilities and six facilities in five other states had reported 67 mumps cases to U.S. Immigration and Customs Enforcement (ICE) Health Service Corps (IHSC) or local health departments. On December 12, TDSHS contacted CDC to discuss mumps control in detention facilities and facilitate communication with IHSC. During January 4–17, 2019, six more state health departments reported new cases in detention facilities, which prompted CDC and IHSC to launch a coordinated national outbreak response.

During September 1, 2018–August 22, 2019, a total of 898 confirmed and probable mumps cases (1) in adult migrants detained in 57 facilities (18% of 315 U.S. facilities that house ICE detainees*) were reported in 19 states (Figure); an additional 33 cases occurred among staff members. Private companies operated 34 facilities, 19 were county jails that house detained migrants, and four were ICE-operated. Forty-four percent (394) of cases were reported from facilities that house ICE detainees in Texas. Median patient age was 25 years (range = 17–67); 846 (94%) were male. Based on detainee custody status during their incubation period (12–25 days before symptom onset), most (758, 84%) patients were exposed while in custody of ICE or another U.S. agency\(^1\); 43 (5%) were exposed before apprehension; and the custody status at the time of exposure of 97 (11%) was unknown. Among those with data on complications, 79 (15%) of 527 male patients reported orchitis, and at least 13 patients were hospitalized. More than half (576, 64%) of cases were confirmed by quantitative reverse transcription–polymerase chain reaction testing or viral culture testing at CDC, state public health laboratories, Association of Public Health Laboratories–CDC Vaccine Preventable Disease Reference Centers, or commercial laboratories. Sequencing of isolates from 70 patients identified genotype G, the most common mumps genotype detected in the United States since 2006 (2). IHSC provided >25,000 doses of measles-mumps-rubella (MMR) vaccine in response to mumps in 56 facilities.

Since 2015, approximately 150 mumps outbreaks and 16,000 cases have been reported in the United States, typically in close-contact settings such as universities, schools, and athletic events.\(^3\) This is the first report of mumps outbreaks in detention facilities.

MMR vaccination efforts differ among detention facilities; facilities should follow local or state health department recommendations for preventing and responding to mumps (3) and should report cases and follow disease control guidance from their health department. Detainees and staff members at increased risk for mumps should be offered MMR vaccine per existing recommendations for vaccination during outbreaks (4,5). MMR vaccine has not been shown to be effective at preventing disease in persons already infected with mumps; facilities should be aware that cases might occur among detainees exposed before vaccination.

Health departments, CDC, IHSC, and facility health administration can work together to develop appropriate control measures based on local epidemiology and the specific needs of each facility. Identifying and vaccinating close contacts of exposed or symptomatic persons with mumps in detention centers is challenging. IHSC can look up transfer history and facilitate vaccine procurement for detainees in ICE custody upon request from facility health services administrators. CDC is coordinating communication among state and local health departments, IHSC, and other federal partners to mobilize appropriate resources and is providing technical support for implementing appropriate disease control and prevention measures. Effective public health interventions require understanding of facility and custody operations, which often involve frequent transfers of detainees (between facilities and states) and multiple entities with authority for operations and detainee custody.

As of August 22, 2019, mumps outbreaks are ongoing in 15 facilities in seven states, and new introductions into detention facilities through detainees who are transferred or exposed before being taken into custody continue to occur.

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*Personal communication, Dr. Diana Elson, U.S. Immigration and Customs Enforcement. Facility count of 315 detention facilities on August 13, 2019, that housed ICE detainees with an average daily population >0 during Fiscal Year 2019. The number of facilities might change with time.

1 U.S. Customs and Border Protection and U.S. Marshals Service.
FIGURE. Mumps cases among U.S. Immigration and Customs Enforcement (ICE) detainees, by custody status* at time of exposure, by week of onset — United States, September 2018–August 2019 (N = 898)

Abbreviations: CBP = U.S. Customs and Border Protection; USMS = U.S. Marshals Service.
* Based on mumps incubation period of 12–25 days before symptom onset.
† Data collected as of August 22, 2019.

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References

Multistate Outbreak of *Salmonella Agbeni* Associated with Consumption of Raw Cake Mix — Five States, 2018

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In August 2018, two Oregon patients with diagnosed *Salmonella* infection were interviewed using a standard enteric illness questionnaire; both patients reported having eaten raw cake mix. Standardized interview questionnaire data collected from 207 Oregon patients with salmonellosis in 2017 indicated a 5% rate of consumption of raw “cake mix or cornbread mix” (Oregon Health Authority, unpublished data, 2017). The binomial probability that both 2018 patients were exposed to raw cake mix by chance was determined to be 0.003, prompting the Oregon Health Authority (OHA) to collect and test the contents of 43 boxes of unopened cake mix of various brands from six retail locations. OHA sent samples to the Institute for Environmental Health Laboratories in Lake Forest Park, Washington, for pathogen testing. *Salmonella Agbeni* was isolated from an unopened box of white cake mix from manufacturer A, and whole genome sequencing (WGS) data describing the isolate were uploaded to the U.S. National Library of Medicine’s National Center for Biotechnology Information (NCBI) website (https://www.ncbi.nlm.nih.gov/pathogens). OHA used the NCBI database to compare sequence data with the cake mix isolate (PNUSAS056022) and then consulted CDC’s System for Enteric Disease Response, Investigation, and Coordination (SEDRIC), a web-based, outbreak investigation tool designed for collaborative, multistate investigations of enteric disease outbreaks.* On October 19, OHA determined that clinical isolates from four patients from Maryland, Ohio, and Wisconsin, with specimen isolation dates ranging from June to September 2018, were genetically related to the *Salmonella Agbeni* isolate from the unopened box of white cake mix, within four single nucleotide polymorphisms (SNPs).

On October 22, 2018, OHA notified state public health counterparts in the three states of this finding and inquired about raw cake mix exposures among their patients. The Wisconsin patient reported having consumed an entire box of raw white cake mix over several days during the likely exposure period. In addition, WGS analysis indicated that this clinical isolate was closely related genetically (within one SNP) to the isolate cultured from the Oregon white cake mix. On October 25, CDC requested officials in Maryland, Ohio, and Wisconsin to interview patients using a questionnaire with specific questions about baking exposures.

On October 31, the Food and Drug Administration (FDA) initiated an investigation of manufacturer A with regard to the *Salmonella*-positive white cake mix. In addition to the investigation and document collection, FDA collected samples including an ingredient (flour), finished cake mix, and environmental samples. All collected samples tested negative for *Salmonella*. On November 5, a voluntary recall of manufacturer A’s classic white, classic butter golden, signature confetti, and classic yellow cake mixes was announced because they might be contaminated with *Salmonella* bacteria.

On January 14, 2019, CDC declared this outbreak, which totaled seven cases in five states,† to be over (1). This is the first time that OHA used WGS data on the publicly available NCBI website to detect a multistate outbreak associated with a widely distributed consumer product, which resulted in product action. WGS of food and environmental isolates and subsequent analysis on the NCBI and SEDRIC platforms are emerging as useful tools in identifying outbreaks associated with widely distributed products with long shelf lives and low background rates of consumption, such as raw cake mix. Detection of these outbreaks is typically difficult and relies mainly upon epidemiologic evidence from investigation of a larger number of cases (2–4). These efforts also highlight the value of collaboration between public health epidemiologists and laboratorians as well as the use of new technological tools for outbreak detection. During outbreak or cluster investigations, food and environmental samples should be collected as quickly as possible whenever practical, particularly when epidemiologic data suggest an association. WGS, in conjunction with the NCBI website and SEDRIC, can be used to identify genetically related isolates quickly.

* 1 Florida, Maryland, Missouri, Ohio, and Wisconsin.

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References


QuickStats

FROM THE NATIONAL CENTER FOR HEALTH STATISTICS

Percentage* of Currently Employed Adults Who Have Paid Sick Leave, † by Industry§ — National Health Interview Survey, 2009 and 2018¶

<table>
<thead>
<tr>
<th>Industry</th>
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* With 95% confidence intervals shown with error bars.
† Based on responses to a question that asked, "Do you have paid sick leave on this main job or business?"
§ Respondents were asked to identify the business or industry of their main job, and these industries/businesses were then categorized by the North American Industry Classification System (https://www.census.gov/eos/www/naics).
¶ Estimates were based on a sample of the U.S. civilian, noninstitutionalized population aged ≥18 years. Adults not currently employed at the time of interview were not included in the denominators when calculating percentages.

The percentage of all currently employed workers with access to paid sick leave increased from 57.8% in 2009 to 62.4% in 2018. By industry, the percentage increased for workers in construction (32.7% to 43.9%), wholesale & retail trade (48.3% to 53.1%), services (56.7% to 60.8%), and manufacturing (60.7% to 65.5%). In 2018, fewer than half of workers in agriculture, forestry, and fishing and construction industries had access to paid sick leave compared to approximately 90% of workers in public administration.

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