Contact Lens Health Week — August 24–28, 2015

August 24–28, 2015, marks the second annual Contact Lens Health Week. In collaboration with partners from clinical, public health, industry, and regulatory sectors, CDC is promoting healthy contact lens wear and care practices to reduce the risk for eye infections and complications associated with poor contact lens hygiene.

Research following outbreaks of rare but serious eye infections in the United States showed that these types of infections occur most often in contact lens wearers who do not take proper care of their contact lenses and cases. This finding signaled that action needed to be taken to promote safer contact lens wear and care.

A report in this issue of MMWR provides an updated population-based estimate of the number of contact lens wearers in the United States. The report finds that there are 40.9 million contact lens wearers aged ≥18 years. It also includes results of a survey that found more than 99% of contact lens wearers report at least one contact lens hygiene habit that could put them at risk for an eye infection, with the majority of respondents reporting behaviors that can raise the risk for eye infection. Nearly one third of contact lens wearers reported ever experiencing a contact lens-related red or painful eye that required a doctor’s visit.

Contact lens wearers represent a significant proportion of the U.S. population, and their contact lens hygiene habits put them at risk for painful, costly eye infections that could lead to vision problems. This year’s observance targets teenage contact lens wearers, who have been associated with lower contact lens compliance and higher risk for serious eye infections. Proper contact lens hygiene habits, supplies, and regular visits to the eye doctor are all essential to keeping contact lens wearers’ eyes healthy. Additional information on Contact Lens Health Week and the proper wear and care of contact lenses is available at http://www.cdc.gov/contactlenses.
Panel members are recruited using address-based probability sampling methods and are provided with internet access and a computer if needed. ConsumerStyles survey participants receive entry into a monthly sweepstakes with a prize usually worth <$500. Statistical weighting was used to make the panel representative of the U.S. population on age, sex, race/ethnicity, education level, household income, household size, census region, metropolitan status, and internet access before joining the panel. Respondents were asked demographic questions and what type of contact lenses they wore.

To describe the prevalence of contact lens hygiene-related risk behaviors, an adapted version of the Contact Lens Risk Survey, a previously validated survey, was administered to a convenience sample of online, contact lens-wearing panelists to describe the prevalence of usual contact lens hygiene-related risk behaviors. Participants were members of market research firm Schlesinger Associates’ research panel and wore contact lenses. Panel members are recruited in-person or via internet advertising, email campaigns, or telephone calls. Questions about usual contact lens-related behaviors included the following responses regarding the usual frequency of the behavior: always, fairly often, sometimes, infrequently, or never. For this report, questions with these responses were coded as “ever” if the response was not “never.”


Using the population-based survey, an estimated 40.9 million persons in the United States aged ≥18 years wear contact lenses (16.7% of U.S. adults); 93.0% of contact lens wearers reported wearing soft contact lenses (lenses made of soft, flexible plastics that allow oxygen to pass through to the cornea). Overall, contact lens wearers were younger, female, more educated, and of white, non-Hispanic race/ethnicity when compared with non-contact lens wearers (Table 1). No significant geographic differences between contact lens wearers and non-contact lens wearers were found. Among subtypes of contact lens wearers, rigid contact lens (lenses made of more durable materials resistant to deposit buildup) wearers did not differ significantly in age from non-contact lens wearers, although wearers of soft, daily disposable (lenses worn once and discarded) and overnight contact lens (lenses prescribed for wear while sleeping) were significantly younger.

Approximately 1,000 contact lens wearers completed the Contact Lens Risk Survey. Respondents were mostly female (82%) and aged ≥40 years (62%). Approximately 99% of respondents reported at least one contact lens hygiene behavior previously associated with an increased risk for eye infection or inflammation (Table 2). Half or more of wearers reported ever sleeping overnight in contact lenses (50.2%), ever napping in contact lenses (87.1%), ever topping off disinfecting solution (adding new solution to existing solution in the contact lens case instead of emptying and cleaning the case before adding new solution, 55.1%), extending the recommended replacement frequency of lenses (49.9%) or cases (82.3%), and ever showering (84.9%) or swimming (61.0%) in contact lenses. Approximately one third (35.5%) of contact lens wearers reported ever rinsing their lenses in tap water and 16.8% reported ever storing their lenses in tap water. Almost all rigid wearers (91.3%) reported ever rinsing their lenses in tap water and 33.3% reported ever storing their lenses in water. Nearly one third of all wearers reported ever having experienced a contact lens-related red or painful eye that required a doctor’s visit.

Discussion

An estimated one in six adults in the United States wears contact lenses, and one third of them report at least one health care visit for a red or painful eye while wearing lenses. Approximately 99% of contact lens wearers reported at least one risk behavior ever for eye infections or inflammation. Of particular concern, contact lens wearers of all types frequently reported exposure of their contact lenses to water, including storing or rinsing their lenses in tap water and showering or swimming while wearing lenses. Exposure of lenses to water raises the risk for infection because microorganisms living in water can be transferred to the eye. Even household tap water, although treated to be safe for drinking, is not sterile and contains microorganisms that can contaminate lens cases and contact lenses and cause eye infections.

Sleeping in contact lenses was a frequently reported behavior. Although many soft and some rigid contact lenses have U.S. Food and Drug Administration-approved indications for overnight wear, sleeping in any type of contact lens increases risk for eye infection, although the precise mechanism is not known (4). Noncompliance with recommended lens and case replacement schedules was also commonly reported. Infrequent replacement of contact lens cases has been linked to serious eye infections (5). Additionally, contact lens wearers who do not follow recommended contact lens replacement schedules have more complications and eye discomfort (6). These behaviors raise the risk for eye infections because repeated handling of the lens and case provides opportunities for introduction of microorganisms, while the moist surface of the lens and case provide an environment conducive to microbial growth. This risk is compounded if wearers top off solution in the case, as a majority of surveyed contact lens wearers reported having done at least once. Topping off also decreases the effectiveness of contact lens disinfection (7).

Daily disposable contact lens wearers might have a lower risk for infection if contact lenses are disposed of daily as recommended. Although 40% of daily disposable contact lens wearers did not use a case, thereby avoiding potential contamination associated with the case, a large proportion of daily disposable contact lens wearers did use a case and did so improperly, using tap water to store their lenses.

The number of contact lens wearers in the United States presented here is higher than previous estimates. Another study estimated 38 million contact lens wearers, although the data collection methods were not described (8). A more recent study used data from the National Health and Nutrition Examination Survey (NHANES) and estimated that 18.6 million persons aged ≥12 years wore contact lenses (9). However, the NHANES protocol used a more restrictive contact lens wearer definition and might have underestimated the total number of contact lens wearers in the United States. The demographic patterns observed in the population used for the estimate reported here were similar to the NHANES population; however, the estimate reported here, based on self-reported contact lens use, is a more inclusive estimate. Contact lens wearers are younger...
TABLE 1. Demographic characteristics of wearers and non-wearers of contact lenses, by type of contact lens — United States, 2014*

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Non-wearers (n = 3,528)</th>
<th>Daily disposables (n = 82)</th>
<th>Planned replacement, soft (n = 461)</th>
<th>Overnight, soft (n = 55)</th>
<th>Rigid (n = 46)</th>
<th>Other† (n = 65)</th>
<th>All contact lens wearers (n = 709)</th>
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<tbody>
<tr>
<td></td>
<td>(%) (95% CI)</td>
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<tr>
<td>Age group (yrs)</td>
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<tr>
<td>18–29</td>
<td>(11.1) (9.6–12.6)</td>
<td>(19.5) (17.3–21.6)</td>
<td>(20.7) (19.4–22.1)</td>
<td>(16.5) (15.2–17.9)</td>
<td>(11.9) (10.5–13.3)</td>
<td>(11.9) (10.5–13.3)</td>
<td>(11.9) (10.5–13.3)</td>
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<tr>
<td>30–39</td>
<td>(15.4) (13.8–16.9)</td>
<td>(22.5) (20.5–24.6)</td>
<td>(26.1) (24.7–27.6)</td>
<td>(37.0) (35.7–38.3)</td>
<td>(19.9) (18.6–21.2)</td>
<td>(19.9) (18.6–21.2)</td>
<td>(19.9) (18.6–21.2)</td>
</tr>
<tr>
<td>40–49</td>
<td>(15.7) (14.4–17.0)</td>
<td>(22.9) (21.5–24.3)</td>
<td>(19.5) (18.2–20.8)</td>
<td>(11.1) (10.0–12.2)</td>
<td>(6.3) (5.0–7.7)</td>
<td>(6.3) (5.0–7.7)</td>
<td>(6.3) (5.0–7.7)</td>
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<tr>
<td>60–69</td>
<td>(18.2) (16.8–19.5)</td>
<td>(6.4) (5.1–7.8)</td>
<td>(6.0) (4.8–7.3)</td>
<td>(7.4) (5.9–9.0)</td>
<td>(7.4) (5.9–9.0)</td>
<td>(7.4) (5.9–9.0)</td>
<td>(7.4) (5.9–9.0)</td>
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<tr>
<td>≥70</td>
<td>(12.0) (10.9–13.1)</td>
<td>(0.7) (0.0–2.1)</td>
<td>(1.4) (0.4–2.4)</td>
<td>NA (0.0–1.1)</td>
<td>(3.4) (1.1–6.0)</td>
<td>(3.4) (1.1–6.0)</td>
<td>(3.4) (1.1–6.0)</td>
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p-value

|               | <0.001 §         | <0.001 §         | <0.001 §         | <0.001 §         | <0.001 §         | <0.001 §         | <0.001 §         |

Sex

<table>
<thead>
<tr>
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<th>Female (50.2) (48.2–52.1)</th>
<th>Male (49.8) (47.9–51.8)</th>
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<tr>
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Education

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<tr>
<th></th>
<th>Less than high school (12.7)</th>
<th>High school (31.5)</th>
<th>Some college (29.1)</th>
<th>Bachelor's or higher (26.7)</th>
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<td>&lt;0.001 §</td>
<td>&lt;0.001 §</td>
<td>&lt;0.001 §</td>
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Race/Ethnicity

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<thead>
<tr>
<th></th>
<th>White, non-Hispanic (66.4)</th>
<th>Black, non-Hispanic (11.9)</th>
<th>Hispanic (14.8)</th>
<th>Other, or ≥2 races (6.8)</th>
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</thead>
<tbody>
<tr>
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<td>&lt;0.001 §</td>
<td>&lt;0.001 §</td>
<td>&lt;0.001 §</td>
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</table>

Metropolitan living area

<table>
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<tr>
<th></th>
<th>Metro (83.7)</th>
<th>Nonmetro (16.3)</th>
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</thead>
<tbody>
<tr>
<td>p-value</td>
<td>0.81</td>
<td>0.05</td>
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Region

<table>
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<tr>
<th></th>
<th>Northeast (18.1)</th>
<th>Midwest (21.1)</th>
<th>South (37.2)</th>
<th>West (23.6)</th>
</tr>
</thead>
<tbody>
<tr>
<td>p-value</td>
<td>0.05</td>
<td>0.05</td>
<td>0.05</td>
<td>0.05</td>
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</table>

Abbreviations: CI = confidence interval; NA = not available (insufficient sample size).

* Based on responses to Porter Novelli 2014 summer ConsumerStyles survey with questions on contact lens use and wearer/non-wearer demographics as of summer 2014.
† Other = Contact lens wearers that said they wore another type of contact lens not captured by the survey choices.
§ Significantly different from non-wearers at the 95% confidence level.

on average than non-contact lens wearers. Teens and college age persons (those aged 15–25 years) have been associated with lower contact lens compliance and with higher risk for corneal inflammatory events, a category of eye problems that includes serious eye infections (10).

The findings in this report are subject to at least two limitations. First, the estimated number of contact lens wearers in the United States reported here does not include those aged <18 years. Since younger age is a predictor of more frequent complications, the current estimate does not include some contact lens wearers who might be most at risk for complications. Second, the Contact Lens Risk Survey used a convenience sample and respondents were more likely to be older and female than the general contact lens-wearing population. Because risk factors have been shown to vary by age, the survey might have underestimated the prevalence of contact lens risk behaviors.
TABLE 2. Prevalence of risk behaviors for eye infections* among contact lens wearers, stratified by type of contact lens — United States, 2014

<table>
<thead>
<tr>
<th>Risk factor/Behavior</th>
<th>Daily disposable (n = 154)</th>
<th>Planned replacement, soft (n = 730)</th>
<th>Overnight, soft† (n = 182)</th>
<th>Rigid (n = 85)</th>
<th>Overall (n = 1,141)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sleeping overnight in contact lens (ever)§</td>
<td>(48.7)</td>
<td>(45.1)</td>
<td>(88.6)</td>
<td>(17.3)</td>
<td>(50.2)</td>
</tr>
<tr>
<td>Napping in contact lens (ever)</td>
<td>(85.1)</td>
<td>(86.9)</td>
<td>(96.4)</td>
<td>(74.1)</td>
<td>(87.1)</td>
</tr>
<tr>
<td>Topping off solution (ever)</td>
<td>(72.0)</td>
<td>(51.3)</td>
<td>(59.3)</td>
<td>(60.5)</td>
<td>(55.1)</td>
</tr>
<tr>
<td>Replacing lenses at interval longer than recommended or when problem</td>
<td>(39.0)</td>
<td>(48.5)</td>
<td>(47.4)</td>
<td>NA‡</td>
<td>(49.9)</td>
</tr>
<tr>
<td>Not using contact lens case</td>
<td>(39.6)</td>
<td>(1.9)</td>
<td>(13.4)</td>
<td>(0.0)</td>
<td>(8.9)</td>
</tr>
<tr>
<td>Replacing contact lens case at interval longer than recommended</td>
<td>(83.9)**</td>
<td>(81.1)</td>
<td>(82.0)</td>
<td>(91.4)</td>
<td>(82.3)</td>
</tr>
<tr>
<td>Storing lenses in tap water (ever)</td>
<td>(28.0)**</td>
<td>(12.4)</td>
<td>(20.9)</td>
<td>(33.3)</td>
<td>(16.8)</td>
</tr>
<tr>
<td>Rinsing lenses in tap water (ever)</td>
<td>(40.3)</td>
<td>(27.2)</td>
<td>(38.3)</td>
<td>(91.4)</td>
<td>(35.5)</td>
</tr>
<tr>
<td>Showering in contact lens (ever)</td>
<td>(85.1)</td>
<td>(84.6)</td>
<td>(94.6)</td>
<td>(67.5)</td>
<td>(84.9)</td>
</tr>
<tr>
<td>Swimming in contact lens (ever)</td>
<td>(59.1)</td>
<td>(61.7)</td>
<td>(64.9)</td>
<td>(50.6)</td>
<td>(61.0)</td>
</tr>
<tr>
<td>Infrequently or never washing hands before inserting lenses</td>
<td>(1.3)</td>
<td>(4.8)</td>
<td>(2.4)</td>
<td>(2.5)</td>
<td>(3.7)</td>
</tr>
<tr>
<td>Infrequently or never washing hands before removing lenses</td>
<td>(19.5)</td>
<td>(12.5)</td>
<td>(9.0)</td>
<td>(17.3)</td>
<td>(13.3)</td>
</tr>
<tr>
<td>Where lenses were purchased</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Provider office</td>
<td>(66.9)</td>
<td>(64.7)</td>
<td>(67.5)</td>
<td>(84.0)</td>
<td>(66.9)</td>
</tr>
<tr>
<td>Retail store without eye exam</td>
<td>(8.4)</td>
<td>(11.8)</td>
<td>(7.5)</td>
<td>(8.6)</td>
<td>(10.4)</td>
</tr>
<tr>
<td>Internet</td>
<td>(23.4)</td>
<td>(21.3)</td>
<td>(24.4)</td>
<td>(4.9)</td>
<td>(20.8)</td>
</tr>
<tr>
<td>Had a red/painful eye while wearing contact lens that required a doctor’s visit (ever)</td>
<td>(29.2)</td>
<td>(29.3)</td>
<td>(35.3)</td>
<td>(28.9)</td>
<td>(30.2)</td>
</tr>
</tbody>
</table>

* Based on responses to Contact Lens Risk Survey, reflecting usual behaviors as assessed in December 2014.
† Overnight contact lens wearers replied “yes” to “Are your contact lenses recommended by your eye doctor for overnight wear?”
§ Ever indicates the combined results of those who answered question “always,” “fairly often,” “sometimes,” or “infrequently” (i.e., questions with these responses were coded as “ever” if the response was not “never”).
‡ NA = 100% of rigid wearers reported replacing their lenses when they had a problem, which is compliant with recommendations for rigid lenses.
** Case replacement and storage in tap water questions were only asked if respondent reported using a contact lens case; 39.6% of daily disposable wearers did not use a case. Thus, the reported percentages are the proportion of the 60.4% (n = 93) of daily disposable users that reported using a case.

Summary

What is already known on this topic?

Contact lenses are a safe and effective form of vision correction for the millions of Americans who require it, if worn and cared for as directed. Poor contact lens hygiene behaviors such as exposing contact lenses to water and topping off storage cases with disinfection solution put contact lens wearers at risk for eye infections.

What is added by this report?

In 2014, there were an estimated 40.9 million contact lens wearers aged ≥18 years in the United States. Approximately 99% of contact lens wearers completing the Contact Lens Risk Survey in 2014 reported at least one contact lens hygiene behavior ever that could put them at risk for an eye infection. One third of contact lens wearers reported ever experiencing a red or painful eye that required a doctor’s visit.

What are the implications for public health practice?

Prevention efforts could include vigorous health promotion activities that encourage contact lens wearers to improve their hygiene behaviors, such as keeping all water away from contact lenses, discarding used disinfecting solution from the case and cleaning with fresh solution each day, and replacing their contact lens case every 3 months.

disease and increased surveillance capacity for microbial keratitis are needed. Prevention efforts could include vigorous health promotion activities that encourage contact lens wearers to improve their hygiene behaviors, such as keeping all water away from contact lenses, discarding used disinfecting solution from the case and cleaning with fresh solution each day, and replacing their contact lens case every 3 months (Box).

References


**BOX. Wear and care recommendations to reduce the risk for contact lens-associated complications**

**Contact lens habits and hygiene**
- Never sleep in contact lenses unless advised to do so by an eye care provider.
- Keep all water away from contact lenses. Avoid showering while wearing contact lenses, remove them before using a hot tub or swimming, and never rinse or store contact lenses in water.

**Contact lenses and supplies**
- Replace contact lenses as often as recommended by an eye care provider.
- Discard used solution from the contact lens case and clean it with fresh solution, never water, every day. Store contact lens case upside down with the caps off after each use.
- Replace the contact lens case at least once every 3 months.

**Eye care provider involvement**
- Visit an eye care provider as often as recommended by your primary health care provider.
- Remove contact lenses immediately and call an eye care provider if you are experiencing eye pain, discomfort, redness, or blurred vision.

**Be prepared**
- Carry a backup pair of glasses with a current prescription in case contact lenses need to be removed.

Additional information about healthy contact lens wear and care is available at [http://www.cdc.gov/contactlenses](http://www.cdc.gov/contactlenses) and [http://www.cdc.gov/contactlenses/show-me-the-science.html](http://www.cdc.gov/contactlenses/show-me-the-science.html).

† These recommendations were developed through solicitation of expert consensus opinion and scientific literature review by CDC in collaboration with a workgroup that included members from the U.S. Food and Drug Administration, the American Academy of Ophthalmology, the American Academy of Optometry, the American Optometric Association, the Contact Lens Association of Ophthalmologists, the Contact Lens Society of America, and the National Academy of Opticianry. The rationale and publications used to support these recommendations can be found on CDC's Healthy Contact Lens “Show Me the Science” web page, available at [http://www.cdc.gov/contactlenses/show-me-the-science.html](http://www.cdc.gov/contactlenses/show-me-the-science.html).
Each year in the United States, approximately two million persons become infected with antibiotic-resistant bacteria, at least 23,000 persons die as a direct result of these infections, and many more die from conditions complicated by a resistant infection (1). Antibiotic-resistant infections contribute to poor health outcomes, higher health care costs, and use of more toxic treatments (2). Although emerging resistance mechanisms are being identified and resistant infections are on the rise, new antibiotic development has slowed considerably (2).

Inappropriate antibiotic prescribing is an important and modifiable contributor to antibiotic resistance and is a problem in all health care settings (1). Inappropriate antibiotic use contributes to excess health care costs, promotes antibiotic resistance, and contributes to preventable adverse drug reactions. Antibiotics cause approximately 142,000 adult emergency department visits annually for adverse drug reactions; almost four out of five of these visits are for allergic reactions (3). Antibiotics also contribute to both health care- and community-associated Clostridium difficile infections, which are associated with considerable costs to patients and the health care system (1,4). In 2009, approximately $10.7 billion was spent on antibiotic therapy in the United States, including $6.5 billion, $3.6 billion, and $526.7 million in the outpatient, inpatient acute, and long-term care settings, respectively (5). The cost of antibiotic resistance to the U.S. economy is estimated $20 billion annually in excess direct health care costs, with an additional $35 billion in lost productivity (1).

Antibiotic prescribing must be tracked to understand and improve antibiotic use. Several data sources and surveillance systems have been employed to examine antibiotic prescribing in hospitals and the community. These include the National Ambulatory Medical Care Survey, the National Hospital Ambulatory Medical Care Survey, the National Healthcare Safety Network, claims data from health plans and insurance companies, and data from private vendors (6). An accurate assessment of antibiotic prescribing, regardless of clinical setting, is important to identify opportunities to improve prescribing and maintain provider accountability.

An estimated half of antibiotic prescriptions given during pediatric ambulatory care visits are inappropriate, and over one quarter of adult prescriptions are for conditions for which antibiotics are rarely indicated (6,7). Health care providers prescribed 262.5 million courses of antibiotics in 2011 (842 prescriptions per 1000 persons), and prescriptions per 1,000 persons vary markedly according to geography (8). The highest prescribing states in 2011, Kentucky and West Virginia, had a rate more than twice that of the lowest prescribing state (Alaska). Why such variability exists is unclear, but this variability is unlikely to be explained by differences in population distribution and extent of infectious diseases.

Inappropriate antibiotic use is not limited to the outpatient setting. A recent evaluation of prescribing for inpatients in two specific scenarios (urinary tract infections in patients without indwelling catheters and treatment with intravenous vancomycin) identified that antibiotic use could have been improved in 37% of cases (9). Frequency of antibiotic prescribing among inpatients varies considerably among hospitals. A recent study of 19 hospitals that had completed data validation and submitted antibiotic use data from one or more patient care settings, found threefold differences in usage rates among 26 medical/surgical wards (9).

Visits for acute respiratory tract infections lead to more inappropriate antibiotic prescribing than visits for any other group of diagnoses. For example, antibiotic treatment for acute uncomplicated bronchitis is not recommended, and despite decades-long, widespread efforts to curb antibiotic prescribing, in 2010, 71% of all outpatient visits for this condition resulted in an antibiotic prescription (10). Similarly, overprescribing for pharyngitis is common. Only 5%–10% of pharyngitis cases among adults are caused by group A Streptococcus, for which antibiotic treatment is recommended, yet antibiotics are prescribed for approximately 60% of ambulatory care visits for adult pharyngitis (7). Outpatient antibiotic prescribing for children with acute respiratory tract infections has been decreasing since the mid- to late-1990s, but the rate of decline has slowed and might have reached a plateau (11). Several factors have been hypothesized to have contributed to this decrease, including the increased use of pneumococcal conjugate and influenza vaccines, national education campaigns to promote appropriate antibiotic use, and increasing concern among both the general public and health care professionals about antibiotic resistance.

This is another in a series of occasional MMWR reports titled CDC Grand Rounds. These reports are based on grand rounds presentations at CDC on high-profile issues in public health science, practice, and policy. Information about CDC Grand Rounds is available at http://www.cdc.gov/cdcgrandrounds.
In addition to the problem of overuse, antibiotic selection is often inappropriate. Prescribers often choose second- or third-line antibiotics, which are typically broad-spectrum drugs, despite established clinical practice guidelines recommending more targeted agents. Overuse of broad-spectrum antibiotics (e.g., second- or third-generation cephalosporins, fluoroquinolones) is especially problematic because of their potential for increased selection of resistant bacterial populations and their role in treating serious infections. Among U.S. ambulatory care visits during 2007–2009, broad-spectrum antibiotics accounted for 74% of antibiotics prescribed to patients during visits for respiratory conditions (7). Among hospitalized patients, 56% received an antibiotic during their stay and 30% received at least 1 dose of a broad-spectrum antibiotic (9).

**Improving Prescribing and Antibiotic Stewardship**

The goal of antibiotic stewardship is to maximize the benefit of antibiotic therapy while minimizing harms to both the individual person and the community. Modest reductions in antibiotic prescribing can make a substantial impact. One study predicted that a 10% decrease in outpatient antibiotic prescribing rates would lead to a 16% decrease in *C. difficile* infection incidence in the community (7). Likewise, reducing exposure of hospitalized patients to broad-spectrum antibiotics by 30% can result in an estimated 26% reduction in inpatient *C. difficile* infections (9).

To reduce inappropriate prescribing, recent guidelines for common outpatient infections emphasize stringent case definitions and clinical observation for mild cases. For example, children aged ≥24 months with unilateral acute otitis media and mild symptoms are less likely to benefit from antibiotics, and are good candidates for close observation with shared decision-making that involves clinicians and caregivers. A mechanism for follow-up in 48–72 hours in such cases is recommended (8).

Several interventions have been shown to improve antibiotic prescribing. Audit and feedback involves tracking individual provider prescribing behaviors and giving feedback on their performance relative to peers or established benchmarks. Academic detailing is a method that adapts some strategies developed by pharmaceutical companies to influence prescribing behaviors that involves active, tailored, and personalized education to promote desired behaviors. Clinical decision support can be integrated with electronic health records to promote appropriate prescribing practices for common infections. Effective ambulatory care interventions have been summarized previously (13) and may be adapted to different settings. Although no single intervention can improve all prescribing behaviors in a given outpatient setting, multifaceted interventions involving active provider education appear to have the greatest benefit. Evidence increasingly supports the reduction of unnecessary antibiotic use through delayed prescribing strategies, where patients are given an antibiotic prescription to be filled within a specified timeframe if symptoms do not improve (8).

Measures promoting appropriate antibiotic prescribing in inpatient settings are primarily implemented through antimicrobial stewardship programs, which CDC recommends for all hospitals in the United States (http://www.cdc.gov/getsmart/healthcare/implementation/core-elements.html) (9). In a recent review of hospital interventions to improve antibiotic prescribing (14), both restrictive interventions (e.g., required approval from an infectious disease specialist to order certain antibiotics) and persuasive interventions (e.g., audit and feedback on prescribing behaviors or provider education) appeared to be equally effective after approximately 6 months. Interventions intended to reduce excess antibiotic prescribing have also been associated with reductions in *C. difficile* infection, and a meta-analysis of clinical outcomes found no significant increases in mortality caused by reductions in antibiotic prescribing when intervention groups were compared with controls (risk for mortality 0.92; 95% confidence interval = 0.81–1.06).

Educational campaigns aim to decrease inappropriate antibiotic prescribing by promoting judicious prescribing among providers and by increasing general public and provider knowledge about antibiotic resistance. Strategies to further employ appropriate antibiotic use messages include distribution of public health messages via pharmacies, child daycare centers, and workplaces. The CDC “Get Smart: Know When Antibiotics Work” and “Get Smart for Healthcare” campaigns (http://www.cdc.gov/getsmart/) inform consumers and providers about antibiotic use and resistance, promote adherence to clinical practice guidelines, and support state- and local-level appropriate antibiotic use programs.

**Challenges, Success Factors, and Directions for the Future**

Although guidelines exist for diagnosis and treatment of common infections, diagnostic uncertainty remains a challenge. Health care providers are frequently influenced by psychosocial factors which drive prescribing decisions, including concerns for both patient satisfaction with a clinical visit and potential negative consequences because of missed diagnoses (15). Providers are also concerned about losing dissatisfied patients to other providers who might be more likely to prescribe antibiotics. Patients who are aware of the potential risks for antibiotic overuse might still express a preference for antibiotic treatment because of perceived benefits. Antibiotic
stewardship interventions and educational efforts aimed at addressing both diagnostic uncertainty and patient expectations will remain important.

Interventions to improve antibiotic prescribing have proven effective in the short-term and within specific settings. It remains less clear which interventions are sustainable and scalable. For this reason, strong stakeholder partnerships and buy-in at the personal, clinic, and health care system levels are fundamental to improving antibiotic prescribing. CDC is working with federal partners, including the Centers for Medicare and Medicaid Services, the U.S. Food and Drug Administration, and the Veterans Health Administration to improve prescribing. CDC partnerships with nonfederal stakeholders, such as vendors of antibiotic prescribing data, state health departments, and professional medical societies are also important.

In March 2015, The National Action Plan for Combating Antibiotic-Resistant Bacteria was released, outlining key actions to combat antibiotic resistance in the United States (https://www.whitehouse.gov). These actions include preventing the development and spread of resistant infections, increasing surveillance efforts, developing new drugs and diagnostic tests, and promoting international collaboration to prevent and control antibiotic resistance. In the United States, changes in health care delivery and increased implementation of quality measures provide opportunities to integrate antibiotic stewardship practices. Tracking antibiotic prescribing, regardless of clinical setting, is important in identifying opportunities to improve prescribing and maintain provider accountability. Priority should be placed on reducing prescribing for diagnoses for which inappropriate antibiotic prescribing is common (e.g., acute bronchitis) and on U.S. regions with higher antibiotic prescription rates. Reducing inappropriate antibiotic use and addressing the threat of antibiotic resistance is critical to improve health care quality and to safeguard patient safety across all health care settings.

References


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Exposure to hydrofluoric acid (HF) causes corrosive chemical burns and potentially fatal systemic toxicity. Car and truck wash cleaning products, rust removers, and aluminum brighteners often contain HF because it is efficient in breaking down roadway matter. The death of a truck wash worker from ingestion of an HF-based wash product and 48 occupational HF burn cases associated with car and truck washing in Washington State during 2001–2013 are summarized in this report. Among seven hospitalized workers, two required surgery, and all but one worker returned to the job. Among 48 injured workers, job titles were primarily auto detailer, car wash worker, truck wash worker, and truck driver. Because HF exposure can result in potentially severe health outcomes, efforts to identify less hazardous alternatives to HF-based industrial wash products are warranted.

HF (Chemical Abstracts Service [CAS] no. 7664-39-3) can produce serious health effects through any exposure route. Exposure of HF solution to the eye can cause irritation as well as potentially permanent ocular damage. Tissue damage from skin contact occurs by two mechanisms. Free hydrogen ions can cause a corrosive burn, and free fluoride ions can cause local cellular destruction and penetrate the skin, causing muscle and bone necrosis. HF is insidiously toxic at the low concentrations (<20%) used in vehicle washing, because no overt corrosive skin burn is present at these concentrations and no initial pain alerts the worker to the exposure (1–3). Numbness, induced by the nerve damage resulting from fluoride ion penetration, leaves the injured worker unaware of the underlying necrosis that can progress for up to 24 hours after exposure (1,2).

Systemically, fluoride toxicity by any route of exposure can cause fatal cardiac arrhythmias precipitated by hypocalcemia and hyperkalemia. Topical application and subcutaneous administration of calcium or magnesium compounds can be used to quench fluoride ions and preempt tissue damage.

Injuries in Washington State during 2001–2013 that met the case definition for exposure to HF among workers engaged in car or truck washing, including auto detailing, were identified through Washington's State Fund workers’ compensation data system (4). Washington's law mandates workers’ compensation insurance coverage for all employers, with 97.7% of employers and approximately two thirds of the state workforce insured through the Washington State Fund. Potential nonhospitalized burn patients were identified using the following Occupational Injury and Illness Classification System injury nature codes assigned to workers’ compensation claims: 050 (burns unspecified), 051 (chemical burns), 058 (multiple types of burns), and 059 (burns not elsewhere classified) (5). Among potential cases in both hospitalized and nonhospitalized workers, HF exposure (versus exposure to other or unspecified acids) during car or truck washing was confirmed through review of employer, worker, and/or physician narrative statements in the workers’ compensation medical record. Exposure information, including product Safety Data Sheets, were obtained from WA-DOSH inspection records or the medical record. Time-loss payments begin when work is missed on the fourth calendar day after the date of injury.

In 2012, a truck wash worker aged 38 years died after ingestion of a HF-based truck wash solution.* The victim placed a call to 911 emergency medical services; his 5-hour emergency department course was consistent with previous case reports of HF ingestion, including recurrent ventricular dysrhythmias (6). The product ingested was Fast Bright (NW Chemical, LLC) containing HF at <12% and sulfuric acid at <20% concentrations, with a pH of 1.5–1.6. The product is diluted before use on trucks, and the employer reported a dilution ratio resulting in a solution concentration of 0.65% HF. Both the concentrated and diluted solutions were present in the workplace, and it is not known which was ingested.

Workers’ compensation data from 2001–2013 were reviewed, and 48 HF chemical burn cases were identified. The median age of injured workers was 29 years (range = 15–62 years), three were female, and burn depth included superficial (first-degree), partial-thickness (second-degree), and full-thickness (third-degree) from exposure to products that ranged from 0.5% to 20% HF. HF concentration might have a greater effect on burn severity than the affected total body surface area burned. Eight workers (17%) received a median of 21 days (range = 2–40 days) in time-loss compensation.

* Whether this ingestion was intentional, inadvertent, or attempted self-harm is unknown.
Medical and contextual case details are summarized for the seven hospitalized workers (Table 1). Two required operative intervention, including burn debridement (case 1), split thickness skin graft (case 1), and escharotomy (case 3). Five injuries involved the fingers and hands. At the time of injury, workers wore improper gloves (e.g., cotton gloves) (case 2) or compromised gloves (with holes) (case 3). Two workers (cases 4 and 7) wore no gloves, one of whom manually washed a truck with an HF saturated washing mitt. One worker (case 6) had chemically resistant gloves and a face shield, but while scrubbing carwash walls overhead, the solution dripped down the brush handle and onto the worker’s arm and body. Delay in recognizing the exposure and in seeking medical attention occurred among nearly all hospitalized workers. Although immediate calcium gluconate administration can minimize the local and potential systemic effects of HF, no injured worker received calcium gluconate at their workplace. (Although the federal Occupational Safety and Health Administration (OSHA) and WA-DOSH require employers to provide a safe workplace, no regulation specifies that calcium gluconate be kept at the worksite.) With the exception of one worker (aged 15 years), all hospitalized workers returned to work; two (cases 1 and 7) received time-loss compensation, and two (cases 1 and 3) received permanent partial disability awards.

As a case example, one worker (case 1) splashed his left leg while transferring a cleaning solution of HF and sulfuric acid between containers. He did not irrigate the area and continued to work for approximately 1.5 hours with soaked pants and shoe until he developed an uncomfortable burning sensation. Upon evaluation, the patient was reported to have a quarter-sized brown necrotic area on the anterior left ankle and burn to the anterior left lower leg. Emergency medical technicians irrigated the area with calcium gluconate and transported him to a burn unit, where he received a calcium gluconate injection. He sustained a small area of full-thickness skin loss requiring excision and debridement with a skin graft. The worker received outpatient burn therapy and returned to part-time work 6 weeks after the injury. A foot paresthesia developed, and the worker received a permanent partial disability payment.

Body regions involved in the 41 nonhospitalized burn patients were upper extremity (16 patients, including hands and fingers [14]), head (14 patients, including eyes [14]), lower extremity (seven), multiple body regions (three), and trunk (one).

The exposed population includes workers in 16 industries (Table 2), with nearly half (n=24) occurring in car washes (North American Industry Classification System [NAICS] no. 811192), which includes truck, van and trailer washing as well as auto detailing (7). HF burn injury also commonly occurred in new car dealers (NAICS no. 441110) (n=seven). Truck drivers (n=five) are at risk; three of the seven hospitalized cases were in truck drivers.

Workers apply HF-based solutions to vehicles with handheld sprayers, pressurized metered sprayers, and open wash buckets. In addition to ready-to-use products, car and truck washes dilute concentrated HF-based products with water onsite to create the ‘use dilution’ solution, and exposure can occur during dilution and product transfer. Eight products were named in association with the 17 HF burn patients (Table 3). HF-based products often include additional chemicals that can burn, including sulfuric acid and phosphoric acid. Two products contained ammonium bifluoride (NH₄HF₂, CAS no. 1341-49-7), a chemical that dissociates into HF when dissolved in water and therefore has similar toxicity.

### Discussion
During 2001–2013, one fatal HF ingestion and 48 chemical burns from exposure to HF associated with car and truck washing were reported in Washington State. Although an estimated 134,000 workers are employed in the car wash industry (NAICS no. 811192) in the United States (8), few case reports of HF exposure in car and truck wash workers have been published. In a study that examined nine fatal unintentional occupational HF poisonings investigated by OSHA, none

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**TABLE 1. Summary of cases of hydrofluoric acid exposure occurring during commercial car and truck washing — Washington, 2001–2013**

<table>
<thead>
<tr>
<th>Date of incident</th>
<th>Age*</th>
<th>Assigned task</th>
<th>Burn location</th>
<th>Burn classification (degree)†</th>
<th>Time loss (days)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dec 2012</td>
<td>38</td>
<td>Wash truck</td>
<td>Systemic ingestion</td>
<td>—</td>
<td>Patient died</td>
</tr>
<tr>
<td>Feb 2001</td>
<td>23</td>
<td>Transfer solution</td>
<td>Left ankle, leg</td>
<td>3rd</td>
<td>40</td>
</tr>
<tr>
<td>Dec 2002</td>
<td>62</td>
<td>Wash trailer</td>
<td>Bilateral hands</td>
<td>2nd</td>
<td>0</td>
</tr>
<tr>
<td>Sep 2003</td>
<td>45</td>
<td>Wash truck</td>
<td>Right fingers (4 and 5)</td>
<td>3rd</td>
<td>0</td>
</tr>
<tr>
<td>Aug 2006</td>
<td>53</td>
<td>Wash wheels</td>
<td>Bilateral hands</td>
<td>Not reported</td>
<td>0</td>
</tr>
<tr>
<td>Jan 2007</td>
<td>15</td>
<td>Clean aluminum truck surfaces</td>
<td>Right thigh</td>
<td>3rd</td>
<td>0</td>
</tr>
<tr>
<td>May 2012</td>
<td>21</td>
<td>Wash walls and ceiling</td>
<td>Hands, legs, abdomen</td>
<td>1st</td>
<td>0</td>
</tr>
<tr>
<td>Mar 2013</td>
<td>32</td>
<td>Clean truck</td>
<td>Right thumb</td>
<td>2nd</td>
<td>16</td>
</tr>
</tbody>
</table>

* The fatality and all cases requiring hospitalization occurred in male workers.
† As reported by the physician in the medical record.
were found to be associated with car or truck washing (9). An Oregon-OSHA hazard alert on HF exposure describes two car wash workers with HF burns, one of whom sustained a finger amputation (10). The broad distribution of HF burns associated with vehicle washing but occurring outside of the car wash industry suggests a large population of at-risk workers. Less hazardous alternatives to HF-based wash products are available, and product substitution could have averted the HF burn injuries described in this report (3). When HF-based products are used, workplaces must use engineering and administrative controls to limit exposure. Product Safety Data Sheets reflect the hazardous nature of the product, and employers are faced with the challenge of managing exposure through worker training and use of personal protective equipment (PPE). However, appropriate PPE does not ensure protection; approximately nine of the cases described in this report involved failure of PPE, when product dripped inside rubber boots or gloves, permeated torn resistant gloves, or was sprayed up under safety glasses. Additionally, injury prevention efforts should include education and training with chemical manufacturers and distributors of HF-based products as well as the end users. Among the six identified products, one (made by Zep, Inc.) was produced internationally, and the rest were manufactured and distributed locally.

TABLE 2. Industry and job titles associated with all hydrofluoric acid burns — Washington, 2001–2013

<table>
<thead>
<tr>
<th>NAICS no.</th>
<th>Industry description</th>
<th>Job title* (no. of workers affected)</th>
<th>No. of cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>811192</td>
<td>Car washes</td>
<td>Auto detailer (5), auto detail manager (1), car washer (5), car wash manager (4), truck washer (7), truck wash manager (1), washer (1)</td>
<td>24</td>
</tr>
<tr>
<td>441110</td>
<td>New car dealers</td>
<td>Auto detailer (6), dealership lot attendant (1)</td>
<td>7</td>
</tr>
<tr>
<td>238990</td>
<td>All other specialty trade contractors</td>
<td>Trucking manager, unknown</td>
<td>2</td>
</tr>
<tr>
<td>327320</td>
<td>Ready mix concrete manufacturing</td>
<td>Truck driver, mixer driver</td>
<td>2</td>
</tr>
<tr>
<td>561790</td>
<td>Other services to buildings and dwellings</td>
<td>Truck washer, cleaner</td>
<td>2</td>
</tr>
<tr>
<td>811310</td>
<td>Commercial and industrial machine and equipment (except auto and electronic) repair and maintenance</td>
<td>Mechanic, truck washer</td>
<td>2</td>
</tr>
<tr>
<td>111219</td>
<td>Other vegetable and melon farming</td>
<td>Unknown</td>
<td>1</td>
</tr>
<tr>
<td>113310</td>
<td>Logging</td>
<td>Truck driver</td>
<td>1</td>
</tr>
<tr>
<td>423830</td>
<td>Industrial machinery and equipment merchant wholesalers</td>
<td>Car washer</td>
<td>1</td>
</tr>
<tr>
<td>484121</td>
<td>General freight trucking, long distance, truckload</td>
<td>Mechanic</td>
<td>1</td>
</tr>
<tr>
<td>484210</td>
<td>Used household and office goods moving</td>
<td>Truck washer</td>
<td>1</td>
</tr>
<tr>
<td>484220</td>
<td>Specialized freight (except used goods) trucking, local</td>
<td>Truck driver</td>
<td>1</td>
</tr>
<tr>
<td>532111</td>
<td>Passenger car rental</td>
<td>Auto detailer</td>
<td>1</td>
</tr>
<tr>
<td>561320</td>
<td>Temporary help services</td>
<td>Mechanic</td>
<td>1</td>
</tr>
<tr>
<td>561431</td>
<td>Private mail centers</td>
<td>Truck driver</td>
<td>1</td>
</tr>
<tr>
<td>611512</td>
<td>Flight training</td>
<td>Truck washer</td>
<td>1</td>
</tr>
</tbody>
</table>

Total no. of cases, including fatality: 49

* Job title as given on the workers’ compensation Report of Accident form (free text).


<table>
<thead>
<tr>
<th>Product</th>
<th>Manufacturer</th>
<th>No. of cases</th>
<th>HF% concentrate*</th>
<th>HF% dilute solution†</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zep-A-Lume</td>
<td>Zep, Inc.</td>
<td>6</td>
<td>5–10</td>
<td>4.2–8.3</td>
</tr>
<tr>
<td>Aluma Brite</td>
<td>—</td>
<td>3</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Aluma-Kleen 1000</td>
<td>Wesmar Co., Inc.</td>
<td>2</td>
<td>10–20§</td>
<td>—</td>
</tr>
<tr>
<td>Fast Bright</td>
<td>NW Chemical, LLC</td>
<td>2</td>
<td>&lt;12</td>
<td>0.65</td>
</tr>
<tr>
<td>A-Wall</td>
<td>CH₂O, Inc.</td>
<td>1</td>
<td>—</td>
<td>0.5</td>
</tr>
<tr>
<td>Lume Brite Aluminum Cleaner and Brightener</td>
<td>—</td>
<td>1</td>
<td>&lt;12</td>
<td>—</td>
</tr>
<tr>
<td>TC-303 Acid Aluminum Truck Brightener</td>
<td>Malco Products, Inc.</td>
<td>1</td>
<td>&lt;5+ &lt;4§</td>
<td>—</td>
</tr>
<tr>
<td>Wheel Bright</td>
<td>Armor Chemical, Co.</td>
<td>1</td>
<td>—</td>
<td>7</td>
</tr>
</tbody>
</table>

* HF% concentrate is that reported on the product’s Safety Data Sheet.
† HF% dilute solution is self-reported by the worker or their employer in the medical record or during inspection by Washington’s Division of Occupational Safety and Health.
§ Product does not contain HF. It contains 10%–20% ammonium bifluoride (Chemical Abstracts Service no. 1341-49-7 [NH₄HF₂]), which dissociates into HF when dissolved in water.
¶ Product contains <5% HF and <4% ammonium bifluoride.

The findings in this report are subject to at least two limitations. First, groups exempted from Washington’s mandatory workers’ compensation law, including self-insured qualified employers, large employers, and sole proprietors, are not represented in the findings. Second, workers who have workers’ compensation coverage but do not file a claim would not be included. Barriers to accessing the workers’ compensation system include a lack of knowledge of the system, language other than English, beliefs about eligibility, and fear of job loss or retribution (10).

Occupational exposure to HF-based wash solutions can result in chemical burns, disability, and death. HF’s potential to cause severe injury combined with the inherent challenge of relying on PPE to protect workers warrants efforts to identify less hazardous alternatives, which would provide the most effective means of prevention.

Summary
What is already known on this topic?
Hydrofluoric acid (HF) causes chemical burns and is a serious systemic poison by all routes of exposure. HF is a chemical component in car and truck wash products, such as rust removers, aluminum brighteners, and wash formulations, because it is inexpensive and highly effective.

What is added by this report?
During 2001–2013, one death and 48 chemical burns from exposure to HF-based products used during car and truck washing, including auto detailing, were reported in Washington. The burns resulted in hospitalization, time lost from work, and disability. Reported diluted-use concentrations were <1% HF, and reported concentrated formulations contained up to 20% HF; both concentrations are hazardous to workers.

What are the implications for public health practice?
Because exposure to HF is toxic and can result in severe health outcomes, efforts to identify less hazardous alternatives to HF-based wash products are warranted. Further characterization of chemical burns from exposure to HF in auto detailers, car and truck wash workers, and truck drivers from other data sources or states would elucidate the magnitude and severity of this occupational health hazard.

Acknowledgments
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References

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Since the 1988 launch of global poliomyelitis eradication efforts, four of the six World Health Organization (WHO) regions have been certified polio-free (1). Nigeria is one of only three countries, along with Afghanistan and Pakistan, where transmission of wild poliovirus (WPV) has never been interrupted. During 2003–2013, northern Nigeria served as a reservoir for WPV reintroduction into 26 previously polio-free countries (2). In 2012, the Nigerian government launched a national polio eradication emergency plan (3) to intensify efforts to interrupt WPV transmission. This report describes polio eradication activities and progress in Nigeria during January 2014–July 2015 and updates previous reports (2–4). No WPV cases have been reported to date in 2015, compared with a total of six cases reported during 2014. Onset of paralysis in the latest reported WPV type 1 (WPV1) case was July 24, 2014. Only one case of circulating vaccine-derived poliovirus type 2 (cVDPV2) has been reported to date in 2015, compared with 20 cVDPV2 cases during the same period in 2014. Pending final laboratory testing of 218 remaining specimens of 16,617 specimens collected since January 2015, Nigeria could be removed from the WHO list of polio-endemic countries in September 2015. Major remaining challenges to the national polio eradication program include sustaining political support and program funding in the absence of active WPV transmission, maintaining high levels of population immunity in hard-to-reach areas, and accessing children in security-compromised areas of the northeastern states.

Vaccination Activities

Nigeria's routine immunization program includes vaccination with trivalent (types 1, 2, and 3) oral poliovirus vaccine (tOPV) at birth and ages 6, 10, and 14 weeks. In 2014, WHO and the United Nations Children's Fund estimated national 3-dose tOPV coverage (tOPV3)* among children aged 12 months to be 66% (5). In February 2015, inactivated polio vaccine (IPV) was introduced into the routine immunization program and is being rolled out in phases that initially prioritized eleven polio high-risk states† (6), and as of July, had been introduced in 35 of Nigeria's 36 states as well as the Federal Capital Territory. This is part of a global plan to provide immunity to type 2 poliovirus (the most common type of cVDPV) in all OPV-using countries, before a synchronized switch from tOPV to bivalent OPV (bOPV), which contains OPV types 1 and 3 (7).

During January 2014–July 2015, 14 supplemental immunization activities (SIAs)§ were conducted in Nigeria. The majority of the 10 subnational SIAs used bOPV, although some local government areas (LGAs) (equivalent to districts) at increased risk for cVDPV2 emergence used rOPV. Of the four national SIAs conducted during this period, one used tOPV, one used bOPV, and two used bOPV in some states and tOPV in others, depending upon polio risk profiles. During SIAs using both rOPV and IPV in selected high-risk states and LGAs from June 2014 through May 2015, approximately 4.4 million IPV doses were administered in high-risk communities.

A number of strategies were implemented during January 2014–July 2015 to enhance the quality of SIAs and to further engage communities, including continued use of an accountability dashboard tool,¶ directly observed polio vaccination,** health camps, †† and social mobilization by volunteer community mobilizers, religious and traditional leaders, and polio survivors, who continue to assist in reducing noncompliance. Although areas of inaccessibility caused by political insurgency increased in places such as Borno, Yobe, and northern Adamawa states (Figure 1), additional innovative strategies continue to be implemented, including permanent health teams made up of women who deliver OPV to households within their communities, transit-point vaccination, vaccination in camps for internally displaced persons, short-interval SIAs that take advantage of intermittent access to normally inaccessible areas, and vaccination of children attending malnutrition treatment centers.

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* Coverage with the third dose of diphtheria-tetanus-pertussis vaccine is used as a surrogate for routine immunization coverage because reported OPV coverage can include doses given during SIAs.
† Polio high-risk states in northern Nigeria: Bauchi, Borno, Jigawa, Kaduna, Kano, Katsina, Kebbi, Niger, Sokoto, Yobe, and Zamfara.
‡ § ¶ ** ††

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FIGURE 1. Areas inaccessible to vaccination teams, by proportion of inaccessible settlements — Borno and Yobe states, northern Nigeria, January 2014–June 2015
SIA quality is assessed using lot quality assurance sampling (LQAS)\(^9\) surveys to estimate whether OPV coverage in the surveyed area is at or above a threshold of 90%. During January 2014–July 2015, the number of LGAs conducting LQAS surveys in the 11 high-risk states increased from 207 to 226. During the same period, the proportion of LGAs passing at or above the 90% threshold increased from 47% to 75%, the proportion of LGAs at the 80%–89% level decreased from 34% to 22%, and the proportion of LGAs below the 80% level decreased from 18% to 3%.

**Poliovirus Surveillance**

*Acute flaccid paralysis surveillance.* Polio surveillance relies on laboratory-supported acute flaccid paralysis (AFP) case detection and confirmation. Two indicators are used to assess the quality of AFP surveillance: documentation of a nonpolio AFP (NPAFP) rate of two or more cases per 100,000 population aged <15 years (indicating satisfactory sensitivity) and collection of adequate stool specimens from ≥80% of persons with AFP (I). Nigeria’s NPAFP rate for 2014 was 14.8 per 100,000,\(^5\) and 97% of AFP cases had adequate stool specimen collection. For 2015, the annualized NPAFP rate was 13 cases per 100,000, and adequate stool specimens were collected for 99% of AFP cases. All 11 high-risk states exceeded both indicator standards in 2014 and continue to do so in 2015. Efforts have been made to enhance surveillance in insecure areas within Borno and Yobe states by adding reporting sites, increasing the number of community informants, and monitoring the performance of surveillance weekly at the national level. As a result, the 2015 NPAFP rate per 100,000 population <15 years was 17.0 for Borno and 27.7 for Yobe (Figure 2).

*Environmental surveillance.* AFP surveillance is supplemented by environmental surveillance; samples are taken from effluent sewage sites every 2–4 weeks for poliovirus testing. By July 2015, environmental surveillance was being conducted in 38 sites, mostly in northern Nigeria: Borno (four sites), Kaduna (three), Kano (five), Lagos (five), Sokoto (four), the Federal Capital Territory (two), Kebbi (three), Katsina (three), Jigawa (three), Yobe (three), and Adamawa (three). In 2014, WPV1 was detected in one sewage sample collected in May in Kaduna, and cVDPV2 was detected in 54 sewage samples: 14 from Kano (last detected in July 2014); 13 from Borno (June); 12 from Sokoto (August); 11 from Kaduna (October); two from Katsina (October); and one each from Jigawa and Yobe (November). Borno had no further positive environmental samples after mid-2014, following the introduction of IPV and use of tOPV in SIAs in the state. During January–July 2015, cVDPV2 was identified in one sewage sample collected from Kaduna (March).

**Polio Incidence**

*WPV and cVDPV polio cases.* No WPV1 cases have been reported in Nigeria to date in 2015. During 2014, six WPV1 cases were reported, 53 were reported during 2013, and 122 were reported during 2012 (Figure 3). The six WPV1 cases in 2014 were geographically limited to five in Kano and one in Yobe state; onset of paralysis in the last reported WPV1 case was July 24, 2014. The last WPV type 3 case was reported in November 2012. One cVDPV2 case has been reported to date in 2015 in the Federal Capital Territory, with a paralysis onset date of May 16. During 2014, 30 cVDPV2 cases were reported, compared with four cases in 2013. Six polio-compatible**

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\(^9\) A clustered LQAS methodology is used to assess SIA quality by sampling the target population of children at the LGA level and documenting finger markings indicative of OPV receipt. A sample is drawn from six wards (geopolitical subunits) within the LGA, with 10 children in a single settlement selected at random from each sampled ward. This yields a total sample of 60 children per LGA. LGAs are classified into one of four classifications based on the number of unmarked children found: 0–3 high pass (dark green); 4–8 pass (light green); 9–19 unacceptable (yellow); and >19 fail (red). A detailed description of the methodology is available at [http://www.polioeradication.org/Portals/0/Document/Research/OPVDelivery/LQAS.pdf](http://www.polioeradication.org/Portals/0/Document/Research/OPVDelivery/LQAS.pdf).

\(^5\) Calculated using WHO African Region population estimates.

** A case in which both adequate stool specimens were not collected from an AFP case within 2 weeks of the onset of paralysis, for which a panel of experts considers the clinical presentation to be compatible with polio and 1) an acute paralytic illness is reported with polio-compatible residual paralysis at 60 days; 2) death takes place within 60 days; or 3) the case is lost to follow-up. Case definitions are available at [http://www1.paho.org/english/HVP/HVI/hvp_fg_pol.pdf](http://www1.paho.org/english/HVP/HVI/hvp_fg_pol.pdf).

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![FIGURE 2. Cases of nonpolio acute flaccid paralysis reported (N = 435)* — Borno and Yobe states, northeast Nigeria, January–July 2015](image-url)
cases have been reported in 2015 thus far, compared with 21 during the same period in 2014. Overall in 2014, 35 compatible cases were reported.

**Genomic sequence analysis.** Since 2012, the genetic diversity of WPV in Nigeria has declined. Among eight genetic clusters of poliovirus detected in 2012, four were identified in 2013; among these, two active clusters were found in 2014. Genomic sequence analysis can also be used to identify AFP surveillance gaps not otherwise shown by surveillance performance indicators. In areas with good surveillance, isolates from environmental sampling are usually closely related, having >98.5% nucleotide sequence identity in the coding region of the major capsid protein, VP1. Poliovirus isolates with a nucleotide difference of ≥1.5% in the VP1 coding region indicate undetected chains of transmission. During 2012, 2013, and 2014, VP1 nucleotide differences of ≥1.5% were found in 10 of 103, 10 of 53, and two of six sequenced WPV1 isolates, respectively. During 2014, the proportion of cVDPV2 isolates with a VP1 nucleotide difference of ≥1.5% (7.8%) was similar to that in 2013 (6.8%). The isolate from the single 2015 cVDPV2 case is genetically linked to viruses that were first detected in Kaduna in 2014. For 2015, a VP1 nucleotide difference of ≥1.5% was found in one isolate (of seven sequenced isolates) from an environmental sample taken during March in Kaduna state; it was genetically linked to Nigerian viruses associated with the major cVDPV2 lineage group that first emerged in 2005 (8).

**Discussion**

Since establishing a polio emergency operations center and implementing a national emergency polio eradication action plan supported with global partners in 2012, Nigeria has experienced a progressive decrease in WPV1 cases. The success of strategies implemented to improve SIA quality and increase access to hard-to-reach children is reflected in improved LQAS survey data. Despite a decline in genetic diversity of WPV1 during 2012–2014 and achievement of surveillance performance indicators at the national level, virologic data indicated persistent gaps in AFP surveillance quality even in 2014. Nonetheless, allowing for delays in obtaining results from the remaining 218 laboratory specimens, if no WPV is identified in AFP cases or environmental samples, Nigeria stands poised for imminent removal from the WHO list of polio-endemic countries.

For the African region to be certified polio-free, all countries in the region will have to maintain a zero WPV1 case incidence for ≥36 months with high-quality surveillance. Continued strengthening of surveillance is required, including active case finding and close monitoring of polio-compatible cases, which might indicate missed transmission.

Nigeria is at risk for persistent cVDPV2 transmission because of low routine immunization coverage (9) and predominant use of bOPV in SIAs, which could lead to gaps in immunity to type 2 viruses. Efforts to strengthen routine immunization are ongoing in polio high-risk LGAs with existing polio infrastructure; these include building capacity and increasing accountability for routine immunization service provision at the health facility level. Interrupting cVDPV2 transmission will also require increased use of tOPV in SIAs, boosting immunity to type 2 polioviruses with IPV, and strengthening outbreak response to any newly identified VDPV. Five of the next six planned SIAs will use tOPV.

The national polio program will need to continue to manage the challenges posed by the insecurity in areas of northeastern Nigeria where many children remain inaccessible to vaccination services. Innovative strategies, including use of permanent health teams, transit-point vaccination, short interval SIAs, and vaccination of children who access point of care sites, in
addition to monthly security risk assessments, will be key to achieving consistent coverage in these areas. Nigeria’s polio program, in collaboration with international partners, will need to continue to advocate for its eradication priorities, to ensure sustained support during the post-transmission period and after changes in national political leadership.

Polio program legacy planning in Nigeria has begun. Documentation of lessons learned during the challenging fight to eradicate polio is critical because this knowledge can shape future approaches to global health (10). This process includes evaluation of current programs, planning for post-certification transition of polio assets and further use of polio eradication infrastructure to strengthen routine immunization and other national public health priorities. Continued partner and government support will be essential for creating the polio eradication legacy in Nigeria, and for maintaining a polio-free African region.

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References


Lead Poisoning and Anemia Associated with Use of Ayurvedic Medications Purchased on the Internet — Wisconsin, 2015

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On April 30, 2015, the Wisconsin Division of Public Health (WDPH) was notified by a local health department of an elevated blood lead level (BLL) in a female patient aged 64 years. All Wisconsin laboratories are required to provide BLL testing results performed on any state resident to WDPH, and WDPH and local health departments are statutorily mandated to investigate any single BLL ≥20 µg/dL or BLLs that are persistently ≥15 µg/dL. Review of medical records revealed that the patient had developed progressive fatigue and shortness of breath during a period of multiple weeks that prompted inpatient medical evaluation. Hemoglobin level was 8.3 g/dL (normal range for age and sex of patient = 12.5–15.0 g/dL), and peripheral blood smear showed normochromic, normocytic red blood cells with basophilic stippling. A BLL was obtained and found to be 85.8 µg/dL. Urine toxic metals tests revealed mercury and aluminum levels in the normal range. Combined methylated and inorganic urine arsenic levels were slightly elevated at 53.3 µg/L (normal = <18.9 µg/L). The patient was discharged for outpatient lead chelation therapy with oral meso-2,3-dimercaptosuccinic acid.

WDPH interviewed the patient to determine possible environmental sources of lead. She did not report any home remodeling that involved paint disturbance or plumbing maintenance, symptoms consistent with pica, use of pottery manufactured outside the United States, or ingestion of wild game, which can contain lead shot fragments (1). She reported taking several supplements, including two Ayurvedic (traditional Indian) medications produced in India that she purchased on the Internet: Mahayogaraj Guggulu (MG) (Sri Sri Ayurveda Trust) and Bruhat Vata Chintamani Rasa (BVCR) (Shree Dhootapapeshwar Limited). The patient ingested approximately four tablets of MG and two tablets of BVCR daily during February–April 2015.

The Wisconsin State Laboratory of Hygiene performed metals testing of the patient’s well water using graphite furnace atomic absorption, and of both Ayurvedic medications using inductively coupled plasma optical emission spectroscopy. Well water lead level was 4.3 µg/L (Wisconsin public health standards set acceptable levels at ≤15 µg/L), and arsenic was undetectable. BVCR contained 16.4 mg/kg (0.2%) lead, and MG contained 48,700 mg/kg (4.9%) lead. Both supplements also contained trace amounts of cadmium, chromium, and aluminum, as well as substantial amounts of arsenic (3,830 mg/kg in MG) and thallium (14.7 mg/kg in MG and 17.2 mg/kg in BVCR). On the basis of estimated daily MG and BVCR consumption and the patient’s body weight, the patient’s exposure to arsenic and thallium exceeded thresholds deemed safe for human health, as defined by the U.S. Environmental Protection Agency (2). The patient discontinued Ayurvedic medication use and reported improvement in symptoms after 1 month of chelation therapy.

Lead is a highly toxic substance that has no endogenous physiologic role, and no safe level of exposure has been identified. High levels of exposure can cause anemia, cognitive dysfunction, coma, and death (3). Although strict regulations have substantially reduced environmental contamination in the United States, lead poisoning continues to occur. This case report confirms earlier reported risk for lead poisoning from Ayurvedic medications produced in India (4), and highlights the acute toxicity that can develop from short-term use. Although toxic metals can occur naturally in some Ayurvedic medicines, or result from contamination, metals such as lead are often intentionally added to some preparations because of putative health benefits (e.g., naga bhasma, a lead-based herbal medicine used to treat various conditions). Physicians should be aware of possible toxicity caused by these medications and should consider lead poisoning as a cause of unexplained anemia in patients taking Ayurvedic medication. Although this investigation did not reveal health problems caused by other toxic metals, the elevated levels of arsenic and thallium could have presented health risks if these medications had been consumed for prolonged periods. State and local public health departments should consider outreach to educate the public about potential risks of Ayurvedic medications and consider sales restrictions as permitted by statutory and regulatory authority.

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References

From 2010 to 2014, the percentage of persons aged 19–25 years who had a usual place to go for medical care increased for Hispanics (50.7% to 65.1%) and non-Hispanic blacks (65.4% to 74.3%). In 2010, among persons aged 19–25 years, non-Hispanic blacks (65.4%) were less likely than non-Hispanic whites (73.0%) to have a usual place to go for medical care; however, in 2014, no significant difference between the two groups was found. In 2010 and 2014, Hispanic adults aged 19–25 years were the least likely to have a usual place to go for medical care.


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