Use of Multilocus Variable Number of Tandem Repeats Analysis Genotyping to Determine the Role of Asymptomatic Carriers in Clostridium difficile Transmission

Scott R. Curry,1,2 Carlene A. Muto,1,2,3 Jessica L. Schlackman,2 A. William Pasculle,4 Kathleen A. Shutt,1,2 Jane W. Marsh,1,2 and Lee H. Harrison1,2

1Division of Infectious Diseases, Department of Medicine, University of Pittsburgh School of Medicine; 2Infectious Diseases Epidemiology Research Unit, University of Pittsburgh School of Medicine and Graduate School of Public Health; 3Division of Hospital Epidemiology and Infection Control, University of Pittsburgh Medical Center, Presbyterian Campus; and 4Division of Microbiology, Department of Pathology, University of Pittsburgh School of Medicine, Pennsylvania

(See the Editorial Commentary by McDonald on pages 1103–5.)

Background. Previous studies have suggested that asymptomatic carriers of toxigenic Clostridium difficile are a source of hospital-associated (HA) infections. Multilocus variable number of tandem repeats analysis (MLVA) is a highly discriminatory molecular subtyping tool that helps to determine possible transmission sources.

Methods. Clostridium difficile isolates were recovered from perirectal swabs collected for vancomycin-resistant Enterococcus (VRE) surveillance as well as from clinical C. difficile toxin–positive stool samples from July to November 2009 at the University of Pittsburgh Medical Center Presbyterian (UPMC). MLVA was performed to determine the genetic relationships between isolates from asymptomatic carriers and patients with HA C. difficile infection (HA-CDI). Asymptomatic carriage and HA-CDI isolates were considered to be associated if the carriage isolate was collected before the HA-CDI isolate and if the MLVA genotypes had a summed tandem-repeat difference of ≤2.

Results. Of 3006 patients screened, 314 (10.4%) were positive for toxigenic C. difficile, of whom 226 (7.5%) were detected only by VRE surveillance cultures. Of 56 incident cases of CDI classified as HA at UPMC during the study with available isolates, 17 (30%) cases were associated with CDI patients, whereas 16 (29%) cases were associated with carriers. Transmission events from prior bed occupants with CDI (n = 2) or carriers (n = 2) were identified in 4 of 56 cases.

Conclusions. In our hospital with an established infection control program designed to contain transmission from symptomatic CDI patients, asymptomatic carriers appear to have played an important role in transmission. Identification and isolation of carriers may be necessary to further reduce transmission of C. difficile in such settings.

Keywords. Clostridium difficile; screening; transmission; genotyping; MLVA.

Previous studies have established that asymptomatic carriers of Clostridium difficile outnumber symptom-
for prevention of CDI has not been practical until the recent development of molecular tests that allow detection of the *C. difficile* carrier state with a reasonable turnaround time [8].

Although previous studies of CDI transmission based on restriction endonuclease analysis (REA) as the primary molecular epidemiologic approach have shown that asymptomatic carriers have some role in CDI transmission, [1, 9] current infection control guidelines focus on isolation measures for symptomatic patients to assess potential of each to HA-CDI and to determine the potential role of active surveillance for *C. difficile* carriage in a hospital with an established infection control program that has been associated with successful control of HA-CDI [11].

**METHODS**

*Patient Population and Setting*

This study was conducted at the University of Pittsburgh Medical Center Presbyterian–Shadyside (UPMC), a 762-bed (156 intensive care unit [ICU] beds) teaching hospital affiliated with the University of Pittsburgh. Patients undergoing VRE surveillance testing over a 117-day period in July 2009 through November 2009 were also screened for *C. difficile*. VRE surveillance cultures were performed at admission for patients admitted from other healthcare facilities; VRE surveillance was also performed at weekly intervals for all ICU patients and for all other inpatients with length of stay >20 days (>5 days if receiving antibiotics) or who tested positive for MRSA, *C. difficile*, or multidrug-resistant organisms [8]. Patients identified as VRE positive were placed in contact isolation for current and subsequent healthcare and were not rescreened.

*Clostridium difficile* reduction interventions did not change throughout the study, including contact isolation for the duration of hospitalization for patients identified as colonized or infected by *C. difficile* stool toxin testing [11]. Since 2006, the HA-CDI rate at UPMC has never exceeded our target rate of 8 per 10,000 patient-days. The average HA-CDI rate per year (2006–2012) was 5.6 cases per 10,000 patient-days. In addition to previously reported measures, 5000 ppm sodium hypochlorite was routinely used to disinfect all patient care areas. This study was approved by the University of Pittsburgh Institutional Review Board.

*Clostridium difficile* Diagnosis and Screening

Laboratory diagnosis of CDI was done using cell culture cytotoxicity assays (stool toxins) ordered by physicians (Diagnostic Hybrids, Athens, Ohio) of formed and unformed stool specimens. Both perirectal swabs collected for VRE surveillance (screening tests) and positive stool toxins were cultured for *C. difficile* using 48-hour broth enrichment and anaerobic culture [8]. Positive stool toxin specimens were cultured for *C. difficile* ±30 days relative to the 117-day study period. Toxigenic *C. difficile* was confirmed by *tcdC* genotyping for both stool toxins and screening tests [12]. The field sensitivity of screening tests for detection of *C. difficile* was calculated as the number of screening tests positive for toxigenic *C. difficile* divided by the total number of screening tests collected patients with positive stool toxins from −1 to 0 days prior to collection the stool toxin specimen. The results of *C. difficile* screening were not released to clinicians.

**Molecular Typing of *C. difficile***

*Clostridium difficile* isolates were typed using MLVA and *tcdC* genotyping [10, 12, 13]. The *tcdC* genotypes in this study were assigned according to the PubMLST database (http://pubmlst.org/cdifficile) and were used to infer ribotype as described by Dingle et al [14]. Allele designations for *tcdC* conform to those previously described with the exception of genotypes 0, A1, and B, which have been renumbered as 19, 20, and 47 respectively [13, 15]. Minimum-spanning-tree analysis of all isolates recovered in the study was performed using BioNumerics software version 6.6 (Applied Maths, Austin, Texas), using a Manhattan coefficient to calculate the summed tandem-repeat difference (STRD) between isolates [10]. Complexes containing ≥2 isolates whose MLVA genotypes generated an STRD of ≤2 were defined as highly related isolates.

**Epidemiologic Definitions and Analysis**

Patients with positive stool toxins collected at UPMC during the study period plus a lag period of 10 days (to account for the potential incubation period of CDI) were epidemiologically classified by UPMC infection control according to modified criteria for CDI surveillance [16]. Hospital-associated CDI was defined by toxin positive stool and CDI symptom onset ≥3 days after admission to UPMC or present on readmission with documented UPMC exposure (without exposures to other healthcare facilities) in the preceding 12 weeks. CDIs occurring within 6 months of any previous CDI were classified as recurrent CDI. Patients with positive stool toxins but without signs of CDI on chart review were classified as carriers by infection control personnel.

Patients with *C. difficile*–positive screening tests underwent electronic record review of stool toxins sent during the study period ±12 weeks, and patients without any stool toxins sent were defined as carriers and were categorized into 3 groups: (1) persistent carriers: patients with at least 2 screening tests positive for toxigenic *C. difficile* collected ≥7 days apart and no stool toxins ordered by clinicians; (2) transient carriers: patients with only 1 screening test positive for toxigenic *C. difficile*.
**Results**

**Incident HA-CDI**

During the screening period, 158 patients had positive stool toxin assays for *C. difficile* ordered by clinicians, of whom 34 (22%) were classified as carriers. Of the remaining 124 symptomatic patients, 53 (43%) were classified as HA-CDI at UPMC, 13 (10%) as community-acquired, and 57 (46%) as acquired at other facilities or relapses.

Of 61 cases of HA-CDI during the screening period plus the 10-day lag period, isolates were obtained from 56 (92%) cases. Of these, 17 cases were unrelated by MLVA to any isolate recovered during the study and 5 were chronologically first within a complex of highly related isolates; these latter were recovered as late as study day 74. Detailed molecular combined with time–place epidemiologic analysis was thus performed for 33 of 61 (54%) HA-CDI cases (Table 1).

Carriers identified by screening tests only were identified as the sources for a total of 16 of 56 (29%) HA-CDI cases in the study period; of these, 9 were classified as non–ward transmissions, 2 as ward transmissions, 2 as environmental transmissions, and 2 as indeterminate. The presumed source for 1 HA-CDI case classified as a non–ward carrier transmission was a patient who was later diagnosed with CDI; this source patient’s screening tests were positive 35 days prior to the collection of the first stool toxin assay, and the patient was not in contact isolation at the time of the presumed transmission event. Four of the carriers identified as source patients were already in contact isolation for infections with other organisms (Table 1).

One carrier detected by positive stool toxin testing was implicated as the source of a ward transmission (Table 1). No other HA-CDI cases could be traced to a source that was determined to be colonized by stool toxin testing.

Transmission events from CDI patients comprised 17 of 56 (30%) cases; of these, 9 were classified as non–ward transmissions, 7 as ward CDI transmissions, and 2 as environmental transmissions (Table 1). The ward transmission events occurred in 7 different units.

**Screening Cultures**

During the study period, 422 of 4979 (8.5%) screening tests performed on 3006 patients were positive for toxigenic *C. difficile*. Of 3006 patients screened, 314 (10.4%) and 506 (16.8%) were positive for toxigenic *C. difficile* and VRE on at least 1 specimen, respectively. Of the 3006 patients, 1957 (65.1%), 598 (19.9%), 29 (7.6%), 104 (3.5%), 53 (1.8%), and 65 (2.2%) patients had a total of 1, 2, 3, 4, 5, and >5 screening tests performed, respectively. Of 314 patients positive for toxigenic *C. difficile*, 120 (38.2%) patients were also positive for VRE at least once, whereas 194 patients had no positive VRE tests. Positive screening for VRE during the study was associated with increased risk of being positive for toxigenic *C. difficile* on at least 1 specimen (odds ratio [OR], 3.7; 95% confidence interval [CI], 2.9–4.8). Of the 120 patients positive for both *C. difficile* and VRE, *C. difficile* was identified first in 17 of 120 (14%) patients, on the same day in 95 of 120 patients (80%), and afterward in 6 of 120 (6%) patients.

During the 117-day screening period, 417 *C. difficile* isolates were obtained from the positive stool toxins and/or screening cultures of 384 patients. Of these patients, 226 of 384 (58.9%) were detected using screening cultures only. Of these patients, 131 of 226 (58.0%), 58 of 226 (25.7%), 27 of 226 (11.9%), and 10 of 226 (4.4%) were classified as indeterminate-length,
discrepant, persistent, and transient carriers, respectively. During the screening period, 3006 of 12,054 (24.9%) admissions were tested for VRE.

Environmental Cultures

Five of 6 patients who met the criteria for environmental culturing had recovery of *C. difficile* from at least 1 site. The
patient with no recovery of *C. difficile* was classified as a transient carrier and had been moved to a different room by the date of room sampling. All 5 patients with room sites positive for *C. difficile* had isolates highly related by MLVA to isolates from 1 or more screening cultures (Table 2). Additionally, 3 environmental isolates from 2 patients were unrelated (STRD >10) to any of their screening culture isolates. The most frequently contaminated site was the patient call bell.

**Molecular Epidemiology**

MLVA genotyping of all 739 toxigenic *C. difficile* isolates in this study yielded 524 unique genotypes from 479 patients and 13 environmental samples (Figure 1). Two hundred thirty of 739 isolates formed 78 complexes of highly related isolates (STRD ≤2). In 5 of 18 complexes, genotypes contributed by carriers were essential to formation of the complex (Figure 2).

The 17 isolates from HA-CDI cases that were unrelated to other isolates comprised *tcdC* genotypes 1 (n = 6), 19 (n = 5), 14 (n = 2), 10 (n = 2), 9 (n = 1), and 3 (n = 1). Epidemic strain isolates (*tcdC* genotype 1, inferred ribotype 027) comprised 302 of 739 (40.9%) isolates in this study; these were discriminated by MLVA into 170 genotypes, 26 of which formed complexes (Table 1; Figure 2). Epidemic strain isolates were found in 62 of 225 (28%) carriers and in 105 of 194 (54%) of patients with CDI (OR, 3.1; 95% CI, 2.1–4.7; *P* < .001). No other significant differences were seen in strain prevalence by any other *tcdC* genotype over different epidemiological classes. Detailed typing data for all patient isolates in this study are found in Supplementary Table 1.

**Sensitivity of Screening for *C. difficile* and Patterns of Carriage**

The field sensitivity of screening cultures for *C. difficile* was 21 of 24 (88%) and 12 of 13 (92%) collected at intervals of −1 and

<table>
<thead>
<tr>
<th>Patient</th>
<th>Source/Site Cultured</th>
<th>Interval, d</th>
<th><em>tcdC</em> Genotype</th>
<th>MLVA Locus (CDR)</th>
<th>MLVA Cluster</th>
<th>Epidemiologic Classification</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Screening culture</td>
<td>0</td>
<td>9</td>
<td>27 6 12 5 18 2</td>
<td>NA</td>
<td>u Persistent carrier</td>
</tr>
<tr>
<td>1</td>
<td>Call bell #1</td>
<td>9</td>
<td>9</td>
<td>41 6 12 5 12 3</td>
<td>21</td>
<td>w Environmental</td>
</tr>
<tr>
<td>1</td>
<td>Call bell #2</td>
<td>9</td>
<td>9</td>
<td>27 6 12 5 18 2</td>
<td>0</td>
<td>u Environmental</td>
</tr>
<tr>
<td>1</td>
<td>Computer keyboard #2</td>
<td>9</td>
<td>9</td>
<td>42 6 12 5 12 3</td>
<td>22</td>
<td>w Environmental</td>
</tr>
<tr>
<td>1</td>
<td>Screening culture</td>
<td>19</td>
<td>9</td>
<td>28 6 12 5 18 2</td>
<td>1</td>
<td>u Persistent carrier</td>
</tr>
<tr>
<td>2</td>
<td>Screening culture</td>
<td>0</td>
<td>9</td>
<td>49 6 17 5 15 3</td>
<td>NA</td>
<td>x Discrepant</td>
</tr>
<tr>
<td>2</td>
<td>Screening culture</td>
<td>15</td>
<td>9</td>
<td>48 6 17 5 15 3</td>
<td>1</td>
<td>x Discrepant</td>
</tr>
<tr>
<td>2</td>
<td>Screening culture</td>
<td>22</td>
<td>9</td>
<td>48 6 17 5 15 3</td>
<td>1</td>
<td>x Discrepant</td>
</tr>
<tr>
<td>2</td>
<td>Bedside table</td>
<td>29</td>
<td>9</td>
<td>49 6 17 5 15 3</td>
<td>0</td>
<td>x Environmental</td>
</tr>
<tr>
<td>2</td>
<td>Toilet seat</td>
<td>29</td>
<td>9</td>
<td>40 6 8 4 13 11</td>
<td>29</td>
<td>None Environmental</td>
</tr>
<tr>
<td>3</td>
<td>Screening culture</td>
<td>0</td>
<td>1</td>
<td>22 3 8 9 12 9</td>
<td>NA</td>
<td>C Persistent carrier</td>
</tr>
<tr>
<td>3</td>
<td>Screening culture</td>
<td>12</td>
<td>1</td>
<td>24 3 8 9 13 9</td>
<td>3</td>
<td>C Persistent carrier</td>
</tr>
<tr>
<td>3</td>
<td>Call bell</td>
<td>30</td>
<td>1</td>
<td>24 3 8 9 13 9</td>
<td>3</td>
<td>C Environmental</td>
</tr>
<tr>
<td>4</td>
<td>Screening culture</td>
<td>0</td>
<td>5</td>
<td>37 6 6 4 10 18</td>
<td>NA</td>
<td>None Persistent carrier</td>
</tr>
<tr>
<td>4</td>
<td>Call bell</td>
<td>16</td>
<td>19</td>
<td>16 6 9 2 16 13</td>
<td>37</td>
<td>V Environmental</td>
</tr>
<tr>
<td>4</td>
<td>Screening culture</td>
<td>19</td>
<td>19</td>
<td>16 6 9 2 16 13</td>
<td>0</td>
<td>V Persistent carrier</td>
</tr>
<tr>
<td>5</td>
<td>Screening culture</td>
<td>0</td>
<td>1</td>
<td>21 3 16 9 12 4</td>
<td>NA</td>
<td>N Discrepant</td>
</tr>
<tr>
<td>5</td>
<td>Screening culture</td>
<td>17</td>
<td>1</td>
<td>20 3 16 9 12 4</td>
<td>2</td>
<td>N Discrepant</td>
</tr>
<tr>
<td>5</td>
<td>Screening culture</td>
<td>17</td>
<td>1</td>
<td>20 3 16 9 12 4</td>
<td>1</td>
<td>N Discrepant</td>
</tr>
<tr>
<td>5</td>
<td>Bed rail</td>
<td>21</td>
<td>1</td>
<td>20 3 16 9 12 4</td>
<td>1</td>
<td>N Environmental</td>
</tr>
<tr>
<td>5</td>
<td>Bedside table</td>
<td>21</td>
<td>1</td>
<td>21 3 16 9 12 4</td>
<td>0</td>
<td>N Environmental</td>
</tr>
<tr>
<td>5</td>
<td>Telephone</td>
<td>21</td>
<td>1</td>
<td>20 3 16 9 12 4</td>
<td>1</td>
<td>N Environmental</td>
</tr>
<tr>
<td>5</td>
<td>Toilet seat</td>
<td>21</td>
<td>1</td>
<td>20 3 16 9 12 4</td>
<td>1</td>
<td>N Environmental</td>
</tr>
<tr>
<td>5</td>
<td>Screening culture</td>
<td>24</td>
<td>1</td>
<td>20 3 16 9 12 4</td>
<td>1</td>
<td>N Discrepant</td>
</tr>
</tbody>
</table>

Patients were either still in the room or sampled before terminal disinfection. All *Clostridium difficile* isolates from patients’ environment and screening cultures are included. Five sites (toilet seat, call bell, telephone, bedrail, and bedside table [computer keyboard instead of telephone for intensive care unit beds]) were sampled per patient. Patient 1 had 2 intensive care unit rooms sampled, as the patient had just been transferred <60 minutes before sampling. Interval is the number of days between the patient’s first screening culture and any subsequent culture. The MLVA complexes conform to Figure 2 and Supplementary Table 1. The STRD listed for each isolate is the STRD between the patient’s first screening culture and the listed isolate. Epidemiological classifications conform to those outlined for *C. difficile* carriers in the Methods section.

Abbreviations: MLVA, multilocus variable number of tandem repeats analysis; NA, not applicable; STRD, summed tandem-repeat distance.
Figure 1. Minimum spanning tree depicting the relationships between 739 isolates in 524 genotypes by multilocus variable number of tandem repeats analysis (MLVA) recovered at the University of Pittsburgh Medical Center (UPMC) July–November 2009. Duplicate isolates from the same patient are included. Single circles represent 1 MLVA genotype; double circles represent complexes of highly related isolates by MLVA (summed tandem-repeat distances of ≤2), each labeled by capital letter designations conforming to those in Table 1. The numbers within each complex/genotype represent the tcdC genotype. Circle/complex size is proportional to the number of isolates included. Complexes labeled by capital letters contain cases of Clostridium difficile infection (CDI) acquired at UPMC according to epidemiologic criteria occurring within the screening period +10 days; complexes with hospital-associated (HA) UPMC case isolates only from the 30 days before and after this period remain unlabeled. Complexes and genotypes are color-coded according to epidemiologic classifications: red = HA-CDI at UPMC, yellow = colonization detected by toxin testing; blue = C. difficile colonization detected on screening tests; white = environmental isolates; green = all other C. difficile infections detected on toxin testing (recurrent CDI, community-acquired CDI, and HA-CDI not acquired at UPMC).
DISCUSSION

This study supports the hypothesis that carriers of *C. difficile* contribute to transmission within hospitals. Approximately a quarter of isolates from HA-CDI cases were highly related to isolates from asymptomatic patients identified from screening tests. This finding suggests that screening and isolating patients with carriage could potentially lead to further reductions in CDI incidence. Some CDI patients had isolates that could be traced by MLVA to carriers or other CDI patients in isolation at the time of screening. These patients may have contaminated the environment prior to isolation initiation or may represent noncompliance with contact precautions. However, it is conceivable that earlier isolation initiation for some of these sources of new cases through screening for *C. difficile* carriage might have prevented spread. These cases would have then required hand hygiene with soap and water in addition to existing contact isolation precautions.

This study supports our practice of extending *C. difficile* isolation precautions for the duration of hospitalization [11]. As >50% of CDI patients screened at >14 days after their first positive toxin were negative by perirectal swab culture, an active surveillance screening program for *C. difficile* may have a role in determining the timing of contact precaution discontinuation.

This study also supports the hypothesis that environmental transmission events from prior room occupants with CDI or carriage can occur. Room surfaces were positive with MLVA genotypes closely matching that of the carrier in 5 of 6 patients tested in this study. Despite the use of universal sodium hypochlorite cleaning, environmental transmission may have accounted for 4 of 61 incident HA-CDI cases. This finding is in line with the study of Shaughnessy et al, who reported that occupancy of a hospital room previously containing a CDI patient is an independent risk factor for CDI [18]. In a recent study, housekeeping staff were ineffective in disinfecting rooms contaminated with *C. difficile* spores, remote room contamination from carriers may account for the large number of HA-CDI cases that were unrelated to any isolate in this short study. These data suggest that further comparative effectiveness research in hospital disinfection is needed.

This study suggests that the incubation period for CDI may be >1 week, as 7 symptomatic patients tested 8–28 days prior to their first stool toxin were positive for highly related *C. difficile*. In contrast to the meta-analysis of Shim et al [5], where only 2 patients with *C. difficile* carriage developed CDI in an analysis of 810 patients, 12 patients meeting their definition of colonization developed symptomatic disease in our study. These data suggest that colonization may long precede other factors that precipitate CDI such as antimicrobials.

![Figure 2. Minimum spanning tree depicting the detailed relationships between multilocus variable number of tandem repeats analysis (MLVA) genotypes in complexes M and N (Figure 1). Each circle represents 1 MLVA genotype scaled in size to the number of isolates comprising it. Numbers between genotypes indicate the summed tandem-repeat difference between them. The color coding of MLVA genotypes conforms to that in Figure 1. Numbers within genotypes represent the study day(s) on which each isolate within the genotype was collected. Isolates with capital letters (A–F) indicate isolates from patients with HA *C. difficile* infection (HA-CDI) listed in Table 1; the source isolates for these cases based on epidemiological analysis are labeled by corresponding lower-case letters (a–f). One isolate labeled e/d is both a hospital-associated CDI case isolate as well as a source for a subsequent case. The MLVA genotype indicated by the arrow shows an example of a *C. difficile* genotype recovered by screening tests only that was essential to the formation of a complex.](https://academic.oup.com/cid/article-abstract/57/8/1094/530005/5786194530005?fbclid=IwAR1QKXuEw7yoRst9a4rZqbyrJZ98x3o0nJmRgQxmaoxquEeGWZcRw)
Because our screening cultures used an existing surveillance program for a hospital-acquired organism, we did not screen the entire population at risk for C. difficile carriage; prior ward-based studies have shown that patients without recognized C. difficile risk factors comprise 30%–50% of the population of C. difficile carriers [1, 20]. In addition, many patients in this study who were at high risk for C. difficile carriage were not screened because they acquired VRE. We were unable to track common epidemiologic exposures beyond common inpatient wards such as exposure to diagnostic tests or personnel such as physicians. Such exposures may account for patients identified as non-ward or indeterminate transmissions in this study. Future intervention studies of active surveillance testing for C. difficile may need to include all inpatients to avoid these limitations.

Our study confirms that of Walker et al, who showed that ward-based contact with CDI patients can only account for approximately 25% of CDI transmission events using a less discriminatory typing method than MLVA [21]. In this study, we showed that a substantial balance of transmissions arises from asymptomatic carriers; whether expanding screening to all inpatients would identify more ward transmission events or, conversely, show more non-ward transmissions is a matter of conjecture, but the large number of HA-CDI cases with non-ward transmissions suggests that C. difficile spores may be widely transmitted in time and space within hospitals because of their near-indefinite viability and low infective dose [22, 23].

In summary, there could be added value in performing surveillance testing for C. difficile carriage. A controlled trial is required to evaluate the utility of screening for C. difficile carriage with incident HA-CDI as the main study endpoint.

Supplementary Data

Supplementary materials are available at Clinical Infectious Diseases online (http://cid.oxfordjournals.org). Supplementary materials consist of data provided by the author that are published to benefit the reader. The posted materials are not copyrighted. The contents of all supplementary data are the sole responsibility of the authors. Questions or messages regarding errors should be addressed to the author.

Notes

Acknowledgments. We thank Jamie L. Grey for her maintenance of the infection control database used in this study.

Financial support. This work was supported by the Pennsylvania Department of Health (number 4100047864, all authors) and by a research career award from the National Institute of Allergy and Infectious Diseases (K24 AI52788 to L. H. H.).


All authors have submitted the ICMJE Form for Disclosure of Potential Conflicts of Interest. Conflicts that the editors consider relevant to the content of the manuscript have been disclosed.

References


