ABSTRACT

Background: In response to Clostridioides (Clostridium) difficile infections (CDI), infection prevention and control practices in hospital settings tend to focus on symptomatic patients, potentially neglecting other sources of C. difficile. The purpose of the study was to identify epidemiological connections between C. difficile positive patients to explore the possibility of transmission occurring. This would allow an assessment of IPAC practices to ensure resources were being optimized and targeted to the most appropriate strategies to prevent transmission.

Methods: C. difficile was isolated and characterized from 125 patient stool specimens. Isolates were subjected to toxin profiling and ribotyping. Patient locations in the hospital were mapped and epidemiological connections between patients with the same C. difficile ribotype were assessed.

Results: A total of 47 distinct ribotypes were identified, with the most common being ribotype 027/NAP1. Of the 41 cases identified as hospital-associated, only four (9.8%) of the cases could be epidemiologically linked to another patient with known CDI.

Conclusions: A small minority of hospital-associated infections were found to have an epidemiological link to another known case of CDI suggesting transmission from known cases is rare. This suggested that current IPAC practices were effective in preventing transmission from symptomatic patients, but other sources of C. difficile are potentially unrecognized.

KEYWORDS

C. difficile; Ribotyping; Transmission; Hospital

INTRODUCTION

Clostridioides (Clostridium) difficile is the leading cause of antibiotic-associated diarrhea and a common cause of healthcare-associated infections [1]. Infection prevention and control (IPAC) practices, such as the use of personal protective equipment and the use of sporicides for environmental disinfection, tend to target symptomatic patients with C. difficile infection (CDI) because they have been identified as important sources of C. difficile and pose a potential risk for transmission to other patients [2,3]. However, transmission to hospitalized patients may occur from a variety of sources in addition to symptomatic patients with CDI. Asymptomatically colonized patients, environmental sources such as contaminated surfaces or equipment, or the healthcare workers providing care may also be a source for transmission, or alternatively, the patients may be colonized prior to admission [4, 5, 6]. Despite transmission potentially occurring from a variety of sources and enhanced IPAC practices often only targeting symptomatic patients and the rooms they occupy, other potential sources are often left unaddressed.

Asymptomatically colonized patients are typically not recognized since testing formed stool for C. difficile is not recommended [7] and the contribution of asymptomatically colonized patients to the transmission of C. difficile remains unclear. Colonization rates in hospitalized patients have been reported to be low, between four to 13% [8, 9, 10], and isolating colonized patients has led to a reduction in hospital-associated cases of CDI [11]. Colonized patients are themselves at a greater risk of developing CDI than non-colonized patients [5].

Environmental sources such as shared patient equipment, computers in clinical care areas, medication carts and even laundered linens have been shown to be contaminated with spores and may also play a role in transmission [12, 13]. C. difficile spores have also been found in areas not involved in direct patient care, such as physician and nurse work areas [12]. Aithinne et al. (2018) [14] demonstrated that C. difficile can even persist in toilet bowl water after multiple flushes and can be aerosolized during flushing, potentially allowing spores...
to settle and persist on surfaces patients may have contact with. As such, many possible sources of exposure exist in hospital settings.

In recent years, investigations into *C. difficile* in hospital environments have suggested that, under non-outbreak conditions, transmission onward from symptomatic patients isn’t as common as previously thought. Eyre et al. (2013b) [15] found only 35% of *C. difficile* cases were genetically related to a previous case, despite apparent epidemiological links. Walker et al. (2012) [16] also found that the majority of cases (66%) could not be linked to other known cases. These suggest that there might be over-diagnosis of hospital-associated transmission and outbreaks in situations where unrelated strains are causing disease in an epidemiologically similar fashion. Identifying the sources of *C. difficile* and how transmission occurs is complicated and remains poorly understood.

The objectives of this study were to isolate and characterize *C. difficile* from patients diagnosed with CDI at a healthcare facility over a two-year study period and to identify any epidemiological connections between patients that could support the possibility of transmission occurring. This would allow an assessment of IPC practices to ensure resources are being optimized and targeted to the most appropriate strategies to prevent transmission.

**METHODS**

**Specimen collection**

The study took place in a 182-bed community hospital located in Southern Ontario, Canada. The hospital provides care for inpatients in medicine, surgery, obstetrics, paediatrics, step-down and intensive care units. *C. difficile* positive specimens from all inpatient and emergency department patients were included for study. All specimens came from patients who met the case definition for CDI [7]. All patients exhibited diarrhea defined as three or more watery or loose stools in a 24-hour period and tested positive for *C. difficile* by a polymerase chain reaction (PCR) based diagnostic test following the manufacturer’s instructions (GenXpert® *C. difficile*/Epi Assay, Cepheid, Sunnyvale, CA) in the hospital microbiology laboratory. All specimens testing positive from June 2014 – June 2016 were included. The specimens were stored at -80°C until at least 10 specimens were available for sending to a microbiology research laboratory at the University of Guelph. This study was approved by the research ethics board at the participating hospital and the University of Guelph.

**Case classification**

Hospital-associated infections (HAIs) were defined as cases of CDI in a patient with diarrhea having an onset of symptoms at least 72 hours after admission or within four weeks of a previous hospitalization, when they had not had a *C. difficile* infection in the previous eight weeks. Infections attributed to another healthcare (other HAIs) facility were defined as cases of CDI in a patient with the onset of symptoms less than 72 hours into admission and having been exposed to another healthcare facility within the last four weeks. Infections classified as community-acquired (CAs) or indeterminate were defined as cases of CDI in a patient with the onset of symptoms less than 72 hours after admission and no known exposure to a healthcare facility within the previous four weeks, or where the source of infection could not be determined, respectively. Cases were classified as a relapse if symptoms recurred within eight weeks of the previous infection.

**C. difficile culture**

Stools specimens were refrigerated at 4°C and cultured within 24 hours of receipt from the hospital. Approximately 200 mg of the stool specimen was immersed in 9 ml of *C. difficile* moxalactam norfloxacin broth with 0.1% sodium taurocholate and incubated anaerobically at 37°C for seven days. Cultures were alcohol shocked (at a 1:1 ratio with anhydrous ethanol) for one hour for spore selection, centrifuged and plated onto CDMN plates (Oxoid, Nepean, Canada). The plates were incubated anaerobically at 37°C for 48 hours. *C. difficile* was initially identified by colony morphology, characteristic odour, and a positive l-proline aminopeptidase activity test (Pro Disc, Key Scientific Products, Stamford, TX, USA).

**DNA extraction**

DNA was extracted using a commercial kit following the manufacturer’s instructions (Instagene Matrix, Biorad, Richmond, CA). Briefly, a 10 μl loopful of the culture was suspended in 1 ml sterile water in a 1.5 ml microcentrifuge tube, centrifuged at 12 000 x g for 60 seconds and the supernatant was discarded. The pellets were re-suspended in 200 μl of Instagene Matrix, vortexed briefly and incubated at 56°C for 30 minutes. The tubes were then incubated at 100°C for eight minutes, centrifuged at 12 000 x g for two minutes and the supernatants were removed to a fresh tube and stored at -20°C for future use.

**Detection of C. difficile toxins**

PCR to detect the toxin A gene (*tcdA*), the toxin B gene (*tcdB*), and the binary toxin genes (*cdtA* and *cdtB*) was performed as previously described [17]. Amplification reactions were performed using a Mastercycler® pro S Thermal Cycler (Eppendorf, Mississauga, Ontario) and the following cycling parameters: 10 minutes at 94°C, 35 cycles of 50 seconds at 94°C, 40 seconds at 54°C, 50 seconds at 72°C, and a final extension of three minutes at 72°C. The 25 μl reaction contained 2 μl DNA, 1.0 μl of 25 mM MgCl2, 12.5 μl KAPA2G™ Fast HotStart Ready Mix (Kapa Biosystems, Boston, MA), and primers as previously described [17]. PCR products were resolved on a 1.5% agarose gel and visualized using GelRed DNA stain (Biotium, Hayward, CA).

**Capillary ribotyping**

Briefly, 200 ng of purified DNA is used with 0.2 μM of the 16S (5’-GTCCGGCTGATCCACCTCCTC-3’) and 23S (5’-CCCTGACCCCTTAAACTTGACC-3’) primers.
The 16S primer was labelled at the 5’ end with a fluorescent label. Amplification reactions were performed using a Mastercycler® pro S Thermal Cycler (Eppendorf, Missisauga, Ontario) with the following cycling parameters: 15 minutes at 95°C, 24 cycles of 60 seconds at 95°C, 60 seconds at 57°C, 60 seconds at 72°C, and a final extension of 30 minutes at 72°C. The 25 μl reaction contained 2 μl DNA, 0.5 μl of 10 μM primers, 12.5 μl KAPA2G™ Fast HotStart Ready Mix (Kapa Biosystems, Boston, MA).

PCR products were analyzed using the Webribo server (https://webribo.ages.at). Ribotypes identified as international ribotypes based on comparison to reference strains were assigned the appropriate numerical designation and an internal laboratory number was assigned for all other isolates.

**Analysis**
The patients who tested positive for *C. difficile* during their admission were mapped in the hospital by physical location (unit) from the day of admission until discharge. The patients were mapped to three medicine units, two surgical units, an intensive care unit, a step-down unit, a family birthing unit and the emergency department. An epidemiological link that would support the possibility that transmission had occurred was defined as one patient who goes on to develop CDI after spending a minimum of 12 hours on the same unit as another patient who had tested positive for *C. difficile* within the previous eight weeks, with both patients infected with the same ribotype. Basic patient demographic data, reason for hospital visit/admission and recent antibiotic usage were recorded.

**RESULTS**
During the 25-month study period, 125 specimens were isolated from 119 patients diagnosed with CDI and *C. difficile* was isolated from all samples. The classification of infections and summary of toxin profiles are summarized in Table 1.

All 125 isolates were successfully ribotyped and classified into 47 distinct ribotypes, with two subtypes of ribotype 002 being identified (002 and 002/2) and three subtypes of ribotype 014 being identified (014/0, 014/4 and 014/5). Twenty-two isolates belonged to 19 newly identified ribotypes. The most common ribotypes are summarized in Table 2. The 41 HAIs belonged to 18 distinct ribotypes.

Eight of the 14 cases classified as a relapse had the initial infection diagnosed at another facility therefore the specimens from the initial cases were unobtainable and the ribotype is unknown. Of the remaining six relapse cases, only 4/6 (67%) had the same ribotype identified as the initial infection. Two cases had an initial isolate classified as a different ribotype compared to the relapse episode (Table 3).

<table>
<thead>
<tr>
<th>Classification</th>
<th>Total No. of isolates (%)</th>
<th>Total No. of HAI</th>
<th>Total No. of CA-CDIs</th>
<th>Total No. of other-HAI</th>
<th>Total No. of Relapses</th>
</tr>
</thead>
<tbody>
<tr>
<td>027 (NAP1)</td>
<td>25 (20.0)</td>
<td>9</td>
<td>9</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>020</td>
<td>7 (5.6)</td>
<td>5</td>
<td>2</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>014/0</td>
<td>7 (5.6)</td>
<td>0</td>
<td>3</td>
<td>2</td>
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<tr>
<td>014/5</td>
<td>7 (5.6)</td>
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<tr>
<td>002/2</td>
<td>5 (4.0)</td>
<td>3</td>
<td>0</td>
<td>1</td>
<td>1</td>
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<tr>
<td>012</td>
<td>4 (3.2)</td>
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<td>1</td>
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</tbody>
</table>
Demographic and clinical characteristics, including the indication for previous antibiotic use and the admitting diagnosis, are summarized in Table 4.

For the 41 cases identified as hospital-associated, an epidemiological link to another patient known to have, or have had a recent CDI, was investigated. Only 4/41 (9.8%) of the HAIs could be epidemiologically linked to another patient with CDI when ribotype was considered (Table 5). The remaining 37/41 (90%) cases either shared no time on the same unit as another patient with CDI, or was found to have a different ribotype than a patient on the same unit at the same time.

**DISCUSSION**

In this study, only 9.8% (4/41) of infections classified as hospital-associated had an epidemiological link to another known case of CDI suggesting transmission from known cases is rare. On the onset of diarrhea, a series of infection prevention precautions were initiated. In accordance with best practice [7], contact precautions were initiated, which included the accommodation of patients in private rooms with dedicated toileting facilities and all staff and visitors were required to don gloves and a gown prior to any contact with the patient or the patient environment. A sporicidal disinfectant was used for twice-daily environmental cleaning of the patient room and bathroom until the patient was symptom-free for a specified period of time or discharged. In this study, very few cases developed as a result of onward transmission from known cases and therefore the additional precautions initiated for suspect and confirmed cases of CDI were presumably effective in preventing transmission.

Antimicrobial stewardship may be more effective, compared to additional precautions, in reducing CDIs as this strategy would address patients already colonized, or those becoming colonized by an unidentified source as opposed to strictly relying on targeting prevention strategies to recognized exogenous sources.

Recent antibiotic exposure is a well-known risk factor for CDI [18]. Not surprisingly, the majority of patients (70%) had recognized recent exposure to antibiotics prior to the onset of diarrhea, and the most common classes prescribed were cephalosporins, fluoroquinolones, and penicillins. As noted in Table 4, >1 antibiotic was prescribed to 61.1% of patients indicating the need for closer monitoring of antibiotic use to address potential risk factors.

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of symptoms of CDI. The two most frequent indications for antibiotic therapy were pneumonia and urinary tract infections (UTIs). CDI following pneumonia and UTIs are common and have been shown to have a higher in-hospital mortality rate and to result in a longer length of stay [19]. Antimicrobial stewardship practices targeting pneumonia and UTI cases may reduce overall CDI rates [20, 21].

The sources of *C. difficile* in the majority of the infections classified as hospital-associated in this study are unknown. They may include other asymptptomatically colonized patients or staff, an environmental source, or the patients who develop infections may be colonized prior to admission. Patients may also have had contact with other parts of the hospital not identified in this study, highlighting a potential limitation. The possibility of acquisition of *C. difficile* from an unrecognized location outside of the units the patients were admitted to (e.g., operating rooms, diagnostic imaging rooms) cannot be ruled out. *C. difficile* positive patients may have contaminated those locations creating an unrecognized opportunity for transmission.

The role of asymptomatically colonized patients in transmission remains unclear. Eyre et al. (2013a) [4] demonstrated that onward transmission from asymptomatically colonized patients to cases of CDI was likely very rare and Kong et al. 2019 [22] confirmed this, finding only 6% of new CDIs could be linked solely to colonized patients. Curry et al. (2013) [23] reported a higher rate finding 29% of new CDI cases in their study could be linked to asymptomatically colonized patients. Differences in molecular techniques used and baseline CDI rates in the respective patient populations may account for these differences. Overall, neither infected patients nor asymptomatically colonized patients may be the most common source of incident CDIs in hospitals.

The most common ribotype identified in this study was ribotype 027 (NAP 1) (Table 2) which has previously been reported to be a common type identified in healthcare settings, although it is becoming less frequent [24, 25]. This study found a broader variety of ribotypes than previously reported in similar studies. The 125 isolates were classified into 47 distinct ribotypes while Aptekor et al. (2017) [24] classified 108 isolates from 15 different hospitals into only eight ribotypes and found 7/108 (6.5%) to be non-typable. Furuya-Kanamori et al. (2016) [26] found greater variety, classifying 324 isolates into over 90 ribotypes, but also included isolates from colonized patients. These studies were conducted in different countries in varying patient populations and used different ribotyping techniques compared to this study, which may contribute to the varying results. In Canada, in a non-outbreak setting, 46 isolates from a hospital were classified into 10 different ribotypes by Labbé et al. (2008) [27] and Martin et al., (2008) [28] classified over 1,000 isolates provided by 21 diagnostic laboratories into only 39 distinct ribotypes. The reason why such a heterogeneous population of *C. difficile* was identified in this study isn’t clear, but may indicate that the exposures to *C. difficile* may be from a broad range of sources in this patient population.

Another limitation of this study is that only one molecular typing technique (ribotyping) was performed to differentiate strains. All of the cases with epidemiological links to other cases were among the top five most common ribotypes identified in this study. A more discriminatory typing technique, such as whole genome sequencing, may have further differentiated these strains. If anything, typing using higher resolution methods would reduce the number of likely HAIs even further, if isolates of the same ribotype were identified as distinct.

The identification of patients colonized on admission or throughout the duration of stay was not investigated during this study, but is an opportunity for future research. In the future, as suggested by O’Hagan and McDonald (2018) [29], sampling of healthcare workers’ hands and the collection of environmental samples could provide estimates of transmissions, which may be prevented by a focus on hand hygiene or improved environmental cleaning practices.

Two cases identified as relapses had different ribotypes isolated from the specimen compared to the initial isolates. The patients may have been infected with more than one ribotype of *C. difficile* highlighting another limitation of this study. Only one colony was characterized per specimen and additional strains may have been present and were not identified. Alternatively, the patients may have become infected with a new strain of *C. difficile* within the timeframe to classify the case as a relapse. Given that none of these patients had an epidemiological link to another patient with CDI with the same strain during the time period between the initial and relapse infections, if infection with a new strain had occurred, the source is unknown. Further investigation into the sources of *C. difficile* in cases classified as relapses is warranted.

Without the ribotyping data, 75% (31/41) of the hospital-associated cases of CDI would have an epidemiological link based only on time and location within the hospital. This could result in efforts to improve or enhance the IPAC practices focusing on those symptomatic patients potentially wasting IPAC resources better directed at other potential sources of transmission.

This study identified a heterogeneous population of *C. difficile* in this patient population with ribotype 027 (NAP1) identified as the most common ribotype. The majority of new CDIs in these patients could not be epidemiologically linked to other patients with active CDIs. Other sources of *C. difficile* should be investigated and identified to ensure that the IPAC practices being implemented are the most strategic and effective.

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