**Clostridium difficile** Infection: Update on Diagnosis, Epidemiology, and Treatment Strategies

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Abstract

**Background:** **Clostridium difficile** infection (CDI) has increased in incidence and severity over the past quarter century, and is now considered a major cause of healthcare-associated infections.

**Methods:** Review of the pertinent English-language medical literature.

**Results:** There has been a substantial change in the management of CDI. The emergence of the NAP1/BI/O27 strain in the early to mid-2000s has been associated with more severe forms of CDI. The pathophysiology, epidemiology, clinical manifestations and diagnosis, as well as new strategies for medical and surgical management are discussed in this review.

**Conclusions:** **Clostridium difficile** infection can range from benign diarrhea to severe disease associated with substantial morbidity and mortality. Treatment modalities vary based on disease severity and timing of onset. The mainstay of medical treatment remains metronidazole and oral/rectal vancomycin. New management strategies are evolving, including adjunctive treatments such as monoclonal antibodies, vaccination, and fecal transplant. In patients with severe disease or clinical deterioration, early surgical consultation for total colectomy or loop ileostomy may be life-saving. Infection control measures are vital to mitigating the spread of CDI.

During the past 20 years, the incidence and severity of **Clostridium difficile** infections have increased dramatically, especially with the emergence of the NAP1/BI/O27 strain in the early-to-mid-2000s. **Clostridium difficile** is now considered one of the most important causes of healthcare-associated infections, and community-acquired **C. difficile** disease also has emerged as an important entity. Both medical and surgical management of **C. difficile** colitis have changed over the past decade. This review provides an update regarding pathophysiology, epidemiology, clinical manifestations and diagnosis, current medical and surgical management, and opportunities for advancement in treatment modalities for **C. difficile** infection (CDI).

**Pathophysiology**

**Clostridium** was first cultured by Hall and O’Toole in 1935 from the feces of healthy newborns [1]. It was named “Bacillus difficilis” due to the difficulty in culturing this slow-growing, gram-positive bacillus from an anaerobic medium. While of academic interest, the **C. difficile** bacterium was not believed to be important clinically significant until the late 1970s, when its pathogenic role in antibiotic-associated diarrhea and pseudomembranous colitis was described [2]. We now know that **C. difficile** is a rod-shaped sporulating, anaerobic gram-positive bacillus whose potent toxins can cause symptoms ranging from mild diarrhea to life-threatening pseudomembranous colitis and toxic megacolon. Its virulence is related to the production of endospores and endotoxins A and B.

The life cycle of the **C. difficile** bacterium is shown in Figure 1 [3]. Transmission occurs via ingestion of spores and vegetative cells. Most of the vegetative cells are killed by gastric acid. The spores, on the other hand, can survive the acidic gastric environment and will germinate in the small bowel upon exposure to bile acids. These newly flagellate bacteria then migrate to the colon where they can multiply, especially in the setting of antibiotic-induced changes to the composition of colonic flora. The bacteria adhere to the gut mucosa and produce toxins A and B, which leads to a massive inflammatory response cascade, epithelial cell junction opening and apoptosis, and degradation...
of connective tissue by hydrolytic enzymes. The end result is the development of pseudomembranous colitis and watery diarrhea.

*Clostridium difficile* can be part of the normal colon flora colonizing the colon of an asymptomatic carrier. However, when there is an alteration to the composition of normal colonic flora, it can lead to organism overgrowth and toxin elaboration. Kyne et al. showed that as many as 31% of patients who received antibiotics were colonized with *C. difficile*, and 56% of these patients subsequently developed symptomatic *C. difficile* disease [4].

The *C. difficile* bacterium produces two distinct toxins. Toxin A is an enterotoxin that causes local tissue damage and fluid secretion along with the inflammatory response. This is the less potent of the two toxins produced. Toxin B activates the release of cytotoxins from monocytes, leading to the destruction of the cytoskeleton. Toxin B is approximately 1,000 times more potent than toxin A. In some strains of

FIG. 1. Lifecycle of the *Clostridium difficile* bacterium. From Poutanen and Simor [3].
Table 1. Classification of Clostridium difficile Infection (CDI) Based on Exposure Setting and Timing

<table>
<thead>
<tr>
<th>Disease classification</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>HCF onset, HCF-associated CDI</td>
<td>Symptom onset ≥48 h after admission and before discharge from a HCF</td>
</tr>
<tr>
<td>Community onset, HCF-associated CDI</td>
<td>Symptom onset in the community or within 48 h of admission to a HCF, provided symptom onset was ≤4 weeks after the last discharge from a HCF.</td>
</tr>
<tr>
<td>Community-associated CDI</td>
<td>Symptom onset in the community or within 48 h of admission to an HCF, provided symptom onset was &gt;12 weeks after the last discharge from a HCF.</td>
</tr>
<tr>
<td>Indeterminate CDI</td>
<td>Symptom onset does not fit any of the above criteria for an exposure setting</td>
</tr>
<tr>
<td>Unknown CDI</td>
<td>Exposure setting cannot be determined because of lack of available data</td>
</tr>
</tbody>
</table>

From: McDonald et al. [4].
HCF=healthcare facility.

C. difficile, the tcd-A and tcd-B genes also produce a binary toxin, the clinical implications of which are as yet uncertain.

Classification and Definitions

Clostridium difficile infection is associated commonly with diarrhea, defined as ≥3 stools in less than 24 h. However, the lack of diarrhea does not preclude a CDI, as approximately 1% of patients may present with ileus and colonic distention rather than diarrhea. Symptoms may vary from mild diarrhea treated on an outpatient basis to fulminant severe colitis requiring multimodal therapy, intensive care unit (ICU) support, and possible diverting ileostomy or total abdominal colectomy. Extent and duration of CDI treatment is based on clinical severity.

Most patients with CDI will have a recent history of antimicrobial or antineoplastic treatment. However, there are increasing reports of cases from the community that have not been preceded by antimicrobial use. For epidemiologic surveillance, distinction is made between community-onset, community-associated, healthcare facility-onset and healthcare facility–associated CDI. Cases where symptom onset or exposure setting do not fit criteria or cannot be determined are classified as “indeterminate” or “unknown,” respectively. Detailed criteria for classification are listed in Table 1 and Figure 2 [5].

Epidemiology

Prior to the 2000s, the rate of CDI did not vary much from year to year. However, in the past decade and a half, the total number of cases reported has increased dramatically by approximately 25% per year. During the mid- and late-1990s, the reported incidence of CDI in acute care hospitals in the United States was stable at 30 to 40 cases per 100,000 population, with an increase to 50/100,000 cases in 2001 and 84/100,000 cases in 2005—nearly three times the baseline rate of 31/100,000 in 1996 [6]. The incidence and severity of CDI has been on the increase over the last 10 to 20 years, and it is described increasingly outside healthcare settings and in populations previously believed to be at low risk [7].

There also has been a substantial increase in the morbidity, mortality and economic burden associated with CDI in the last several years [8,9]. Increasing rates of severe and fatal infections are of great concern. Clostridium difficile infection was reported as the primary cause of death in England for 499 patients in 1999; this number increased to 2,998 in 2005 and 3,933 in 2006 [10]. The 2003 outbreak in Quebec, Canada was notable, with a reported four-fold increase in the incidence of CDI from 22.2 and 25.2 per 100,000 population (in 1991 and 2002, respectively) to 92.2 per 100,000 in 2003 [11]. This outbreak was associated with major mortality, with C. difficile as the attributable cause of death in 6.9% of cases and as a contributing factor in an additional 7.5% of cases [12].

A recent multicenter study documented that CDI is a common nosocomial infection in mechanically-ventilated ICU patients (n=386; 6.6% of a total cohort of 5,852 patients), and that septic shock occurred in 34.7% of patients. For C. difficile patients, the mortality rate was high (25.1%) and prolonged hospital (23 d vs. 15 d; p<0.001) and ICU (12 d vs. 8 d; p<0.001) length of stay (LOS) were noted, with increased need for skilled nursing care or rehabilitation following hospital discharge [13].

A major contributor to this increase was the emergence of a new strain of C. difficile, the toxigenic strain III NAP1/BI/O27. Between 2002 and 2006, this particular strain gained

![Timeline for surveillance definitions of Clostridium difficile–associated infection (CDI) exposures. A case patient who had symptom onset during the window of hospitalization marked by an asterisk (*) would be classified as having community-onset, healthcare facility–associated disease (CO-HCFA) if the patient had been discharged from a healthcare facility within the previous four weeks; would be classified as having indeterminate disease if the patient had been discharged from a healthcare facility within the previous four to 12 weeks; or would be classified as having community-associated CDI (CA-CDI) if the patient had not been discharged from a healthcare facility in the previous 12 weeks. HO-HCFA=healthcare facility-onset, healthcare facility–associated CDI. From Conen et al. [15].](image-url)
C. DIFFICILE INFECTION UPDATE

public notoriety as the cause of a deadly new epidemic; as of 2010 this strain has spread to 40 U.S. states, seven Canadian provinces, as well as England, continental Europe, and Asia [14,15]. The NAP1 strain is associated with the use of fluoroquinolones, and contains an 18-pair deletion as well as a novel single-base deletion in the tcd-C gene, which results in overproduction of toxins A and B, as well as the emergence of a binary toxin whose clinical significance has not been elucidated yet [16,17]. Clinically, patients manifest more severe cases of CDI, with more presentations of toxic megacolon than diarrhea, profound leukemoid reactions, severe hypoalbuminemia, septic shock, and increased mortality. A recent analysis identified the key genetic changes linked to the rapid transcontinental dissemination of this epidemic C. difficile and its spread throughout the global healthcare system [18].

Epidemiologic data confirm that CDI also is increasing in surgical patients in the United States, with a 34% increase over a 5-y period (1999–2003). In particular, CDI was most prevalent after emergency surgery and among patients having intestinal tract resections. Multivariable analysis demonstrated that C. difficile was an independent predictor of LOS, which increased by 16.0 d, (95% confidence interval [CI] 15.6–16.4; p < 0.0001) in the presence of infection. Total charges increased by $77,483 (95% CI $75,174–$79,793; p < 0.0001), and there was a 3.4-fold increase in the mortality rate (95% CI 3.02–3.77; p < 0.0001), compared with patients who did not acquire C. difficile [19]. The 2010 clinical practice guidelines for C. difficile infection in adults—published by the Society for Healthcare Epidemiology of America (SHEA) and the Infectious Disease Society of America (IDSA)—make the following evidence-based recommendations with regard to epidemiologic surveillance for CDI: To increase comparability between clinical settings, use available standardized case definitions for surveillance of 1) healthcare facility (HCF)-onset, HCF-associated CDI, 2) community-onset, HCF-associated CDI, and 3) community-associated CDI (Fig. 2; strength of evidence and recommendation B-III); at a minimum, conduct surveillance for HCF-onset, HCF-associated CDI in all inpatient healthcare facilities, to detect outbreaks and monitor patient safety (B-III); express the rate of healthcare-associated CDI as the number of cases per 10,000 patient-days (B-III); and if CDI rates are high, compared with those at other facilities, or if an outbreak is noted, stratify rates by patient location in order to target control measures (B-III).

Risk Factors

Classically, any exposure to antibiotics (particularly clindamycin, ampicillin or amoxicillin, cephalosporins and fluoroquinolones) was a major risk factor for the development of CDI. However, there has been emergence of community-acquired CDI where antimicrobial exposure does not precede clinical infection. Age older than 65 y alone confers a 16.3-fold risk for development of CDI. Increased length of hospital stay, manipulation of the gastrointestinal tract (e.g., surgery), immunosuppressed state (e.g., chemotherapy, HIV + patients, and medical comorbidities leading to immunosuppression) also make the patient more susceptible to developing CDI. There also has been much debate regarding the role of acid-suppressive medications such as histamine-2 receptor antagonists and proton pump inhibitors in the development of CDI [20–22].

Clinical Manifestations

The spectrum of CDI ranges from the asymptomatic carrier (found in 7–26% of adults in acute care facilities) to mild/moderate diarrhea to fulminant colitis and pseudomembranous colitis resulting in multiple organ dysfunction syndrome and death. Infection can manifest as multiple loose stools with passage of mucus or occult in milder cases of CDI; frank melena or hematochezia are rare. With more severe forms of CDI, patients may develop unexplained profound leukocytosis, colonic ileus, or toxic megacolon seen on abdominal imaging with minimal to no diarrhea. The NAP1/BI/027 strain of C. difficile has been associated with these more severe clinical manifestations of CDI in major outbreaks around the world [17].

Diagnosis

Laboratory studies

The diagnostic tests of choice are either a stool sample positive for C. difficile or its toxins, or colonoscopic/histopathologic findings of pseudomembranous colitis. According to the 2010 SHEA/IDSA clinical practice guidelines, the gold standard for diagnosis of CDI is with toxigenic stool culture; however, this is not practical for routine diagnosis and has been recommended for epidemiologic studies only. Stool samples should only be collected from symptomatic patients with diarrhea unless colonic ileus is suspected. The recommended route for testing is the use of a two-step algorithm,

### Table 2. Diagnostic Testing for Clostridium difficile

<table>
<thead>
<tr>
<th>Test</th>
<th>Sensitivity</th>
<th>Specificity</th>
<th>Availability</th>
<th>Expense</th>
<th>Utilization</th>
</tr>
</thead>
<tbody>
<tr>
<td>C. difficile culture</td>
<td>Low</td>
<td>Moderate</td>
<td>Limited</td>
<td>$5–10</td>
<td>No diagnostic use; only toxigenic organisms cause disease</td>
</tr>
<tr>
<td>Toxigenic culture</td>
<td>High</td>
<td>High</td>
<td>Limited</td>
<td>$10–30</td>
<td>Reference method; epidemiologic tool</td>
</tr>
<tr>
<td>CCNA</td>
<td>High</td>
<td>High</td>
<td>Limited</td>
<td>$15–25</td>
<td>Limited diagnostic use</td>
</tr>
<tr>
<td>GDH</td>
<td>High</td>
<td>Low</td>
<td>Wide</td>
<td>$5–15</td>
<td>Diagnostically as a screening test; must be confirmed</td>
</tr>
<tr>
<td>Toxin EIA tests</td>
<td>Low</td>
<td>High</td>
<td>Wide</td>
<td>$5–15</td>
<td>Must detect toxins A+B; inferior sensitivity</td>
</tr>
<tr>
<td>NAATs</td>
<td>High</td>
<td>High</td>
<td>Wide</td>
<td>$20–50</td>
<td>Use only in acute disease; false-positives of concern</td>
</tr>
</tbody>
</table>

 From Surawicz et al. [25]. (Reprinted by permission from Macmillan Publishers Ltd.).

*Cost of goods; does not reflect laboratory changes.

CCNA = C. difficile cytotoxin neutralization assay; GDH = glutamate dehydrogenase; EIA = enzyme immunoassay; NAATs = nucleic acid amplification tests.
which employs a glutamine dehydrogenase (GDH) assay for the C. difficile antigen, followed by enzyme immunoassay (EIA) toxin testing. If the antigen is positive and toxin is negative, some laboratories will send the specimen for polymerase chain reaction (PCR) testing as a final confirmatory step. Nucleic acid amplification tests (NAATs) such as PCR are superior to EIA in terms of sensitivity and specificity (Table 2) but are much more costly, and false positives may result in inappropriate treatment. It is important to note that when results are negative, repeat testing is of limited value [23–25]. Furthermore, testing for cure is not recommended.

**Adjunctive studies**

In cases where index of suspicion is high in the absence of a diarrheal specimen for laboratory testing, endoscopic evaluation of the colon may be helpful. The classic endoscopic description is of normal-to-patchy, nonspecific colitis with inflamed mucosa studded with adherent raised white and yellow plaques (Fig. 3, Panel A), but nodular C. difficile colitis has also been reported (Fig. 3, Panel B) [26,27]. It is important to note that the false-negative rate for endoscopy may be as high as 10–25%. Also, in patients suspected of having fulminant colitis and toxic megacolon, there is an increased risk for perforation during endoscopic evaluation. Radiologic evaluation may be useful adjuncts in evaluating the critically ill patient for possible CDI. Plain abdominal x-rays may show dilated colon consistent with toxic megacolon. Computed tomography (CT) may show “thumbprinting” of the mucosa indicative of colonic mucosal edema, wall thickening, abnormal bowel wall attenuation (target sign), nodular fold thickening, the “accordion” sign (alternating edematous haustral folds separated by transverse mucosal ridges filled with enteral contrast material), fat stranding, or in severe cases, ascites or perforation. Overall computed tomography (CT) sensitivity is 52–85%, with specificity of 48–93% [28,29]. Some institutions advocate more rigid adherence to CT scan diagnostic criteria for C. difficile colitis of colon wall thickening of greater than 4 mm combined with any one or more findings of pericolonic stranding, colon wall nodularity, the “accordion” sign, or otherwise unexplained ascites, with a reported sensitivity of 70% and specificity of 93% [30].

**Medical Management**

**Current standard of care**

The first step in treatment of C. difficile disease is cessation of all antibiotics if possible. Prompt appropriate fluid resuscitation, electrolyte normalization, discontinuation of all antiperistaltic agents (including narcotics) that inhibit the elimination of intestinal luminal toxin, and correction of any other concomitant organ dysfunction remain the initial mainstay of therapy in C. difficile disease. According to the 2010 SHEA/IDSA guidelines, metronidazole is the initial drug of choice for the treatment of mild-to-moderate CDI, with oral vancomycin reserved for more severe cases, and the addition of vancomycin retention enemas in severe or complicated cases of CDIs (Table 3A) [15,31]. Numerous studies have examined the efficacy of metronidazole versus oral vancomycin for the treatment of CDI. Vancomycin is one of only two U.S. Food and Drug Administration (FDA)-approved drug for the treatment of CDIs, and overall has a higher rate of cure, especially in more severe cases of CDI. Oral vancomycin also has a lower treatment failure rate, as there is evidence of some C. difficile strains showing metronidazole resistance in vitro. However, for milder cases of CDI, the difference between the two drugs is marginal (96% for vancomycin and 90% for metronidazole); thus, due to concerns regarding cost and potential emergence of vancomycin-resistant Enterococcus, metronidazole is still the preferred initial agent for these milder infections [17,32–34].

In patients with a first recurrence of CDI, the recommended treatment algorithm is the same as for the index episode; however, for patients with multiple recurrences, oral or rectal vancomycin in a tapered or pulsed regimen is recommended due to concern for neurotoxicity from cumulative metronidazole dosing (Table 3B; Fig. 4; Fig. 5) [14,28,32].

**Alternative medications and adjunctive therapies**

New treatment strategies for CDI are needed, and important data have emerged recently for new anti-C. difficile antibiotics including fidaxomicin, rifampin, rifaximin, rifalazil, nitazoxanide, tigecycline, and ramoplanin [36,37].

**Fidaxomicin**

Fidaxomicin (OPT-80) is a macrocyclic antibiotic with a narrow antimicrobial spectrum. Most of its activity is against C. difficile, most strains of staphylococci, and

**FIG. 3.** Findings of Clostridium difficile colitis. (Courtesy of Dr. Julio Murra-Saca, Hospital Centro de Emergencias [26]; Reprinted with permission from Medscape [27].)
TABLE 3A. GUIDELINES FOR TREATMENT OF INITIAL CLOSTRIDIUM DIFFICILE INFECTION

<table>
<thead>
<tr>
<th>Initial episode</th>
<th>Recommended treatment</th>
</tr>
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<tbody>
<tr>
<td>Mild-moderate CDI</td>
<td>Metronidazole 500 mg PO/IV TID&lt;sup&gt;1&lt;/sup&gt;</td>
</tr>
<tr>
<td>-WBC &lt; 15,000 cells/mm&lt;sup&gt;3&lt;/sup&gt; and -creatinine &lt; 1.5 x baseline</td>
<td>10-14 d</td>
</tr>
<tr>
<td>Severe CDI</td>
<td>Vancomycin 125 mg PO QID (FDA approved)&lt;sup&gt;2&lt;/sup&gt;</td>
</tr>
<tr>
<td>-WBC &gt; 15,000 cells/mm&lt;sup&gt;3&lt;/sup&gt; or -creatinine &gt; 1.5 x baseline</td>
<td>10-14 d</td>
</tr>
<tr>
<td>Severe, complicated CDI</td>
<td>Vancomycin 500 mg PO QID ± IV metronidazole 500 mg IV q8h</td>
</tr>
<tr>
<td>-same as for severe CDI and</td>
<td></td>
</tr>
<tr>
<td>-hypotension, shock, ileus, toxic megacolon, perforation, need for colectomy, or ICU admission</td>
<td></td>
</tr>
</tbody>
</table>

Adapted from Cohen et al. [15].
<sup>1</sup>Neither oral nor parenteral metronidazole reach therapeutic concentrations in normal, nondiarrheal stool.
<sup>2</sup>To perform vancomycin retention enemas, insert an 18-inch Foley catheter with 30 mL balloon into rectum, inflate balloon, instill vancomycin, clamp catheter for 60 min, then deflate balloon and remove catheter with balloon.

CDI = Clostridium difficile infection; PO = oral; IV = intravenous; TID = 3 times daily; WBC = white blood cells; QID = 4 times daily; FDA = U.S. Food and Drug Administration; q8h = every 8 h; NS = normal saline; q6h = every 6 h; ICU = intensive care unit.

The rates of clinical cure were non-inferior, but fidaxomicin was associated with a significantly lower rate of recurrence of CDI in patients with the NAP1 strain (15% vs. 25% in modified intention-to-treat population; p = 0.005) [38,39]. In a multicenter, multinational, double-blind randomized controlled trial (n = 535) of fidaxomicin versus oral vancomycin, fidaxomicin was shown to have similar rates of clinical cure as oral vancomycin, with improved sustained response and decreased recurrence rates [40]. Fidaxomicin has recently received FDA approval for the treatment of CDI with a recommended treatment dose of 100 mg by mouth (PO) twice daily (BID).

Rifaximin

Rifaximin displays potent activity against C. difficile strains but resistance remains a concern. In a study of C. difficile strains from hospitalized patients, 12 of 163 isolates were resistant to rifaximin [41] and in a single-center Pittsburgh study, 36.8% (173–470) strains displayed reduced susceptibility to rifaximin [42]. In a clinical study of metronidazole monotherapy versus metronidazole/rifaximin combination therapy, no difference in median time to resolution of symptoms or median time to first occurrence of relapse was identified in an interim analysis, and the study was ended early because there was little chance of demonstrating superiority with combination therapy including rifaximin [43].

Rifaximin is currently approved by the FDA for travelers’ diarrhea (200 mg PO three times daily [TID] for 10 d) as first-line therapy in 13 hospitalized patients with CDI documented that all eight patients who completed the study had completed symptom resolution; one patient had recurrent infection during a repeat hospitalization [49]. In a report of 32 patients treated with rifaximin for recurrent C. difficile infection with 1 y follow up, 53% of patients had no relapse. The MIC value of rifaximin seemed to predict the response to rifaximin treatment [50].

A multicenter double-blind randomized controlled trial of rifaximin compared to vancomycin for C. difficile (NCT00269399) was completed; final results are pending. Rifaximin is currently approved by the FDA for travelers’ diarrhea (200 mg PO TID for 3 d) caused by non-invasive enterococci. It is active against clinical isolates of C. difficile, including the NAP1/BI/027 strains. As with oral vancomycin, there is minimal systemic absorption when the drug is given orally, with resultant high fecal concentrations. In vitro and in vivo studies also show limited activity against other components of the normal gut flora. The first published report of fidaxomicin efficacy was a randomized (n = 629) trial comparing fidaxomicin (200 mg twice daily) or vancomycin (125 mg four times daily) orally for 10 d.
strains of *Escherichia coli* and for reduction in the risk of overt hepatic encephalopathy recurrence (550 mg PO BID) in patients 18 y or older but is not approved for the treatment of CDI.

**Nitazoxanide**

Nitazoxanide is synthetic nitrothiazolyl-salicylamide derivative used to treat protozoan and helminthic infections. This is also the only drug that is effective for cryptosporidiosis in patients who have acquired immune deficiency syndrome. Nitazoxanide blocks anaerobic metabolism of eukaryocytes and inhibits *C. difficile* in vitro at low concentrations. Approximately two-thirds of the oral dose is excreted in the stool as the active metabolite tizoxanide. In a prospective, double-blind control trial, a 10-d course of nitazoxanide 500 mg PO BID was as effective as metronidazole in the treatment of CDI [51]. In a second prospective double-blind study of nitazoxanide versus oral vancomycin, the clinical response rate was no different (77% nitazoxanide vs. 74% vancomycin) [52]. An open-label study in 35 CDI patients reported an overall response rate of 66%. In these

**FIG. 4.** Treatment algorithm for management of *Clostridium difficile* infection. From Joseph et al. [31].

**FIG. 5.** Suggested approach to treatment of recurrent *Clostridium difficile* infection (CDI). From Cocanour [35]. IV Ig = intravenous immunoglobulin; MAB = monoclonal antibody.
Monoclonal antibodies

One of the major incongruities in the management of CDI is that antibiotics are the mainstay of treatment for this antibiotic-associated condition. Non-antibiotic treatments such as monoclonal antibodies would be a welcome addition or replacement of current standard therapies.

The first Phase II human multicenter randomized double-blind, placebo-controlled trial of two novel, neutralizing, fully human monoclonal antibodies against *C. difficile* toxins A and B had impressive results [54]. Recurrent infection developed in 7% (seven of 101) of the antibody group compared with 25% (25 of 99) in the placebo group, reflecting a relative reduction of 72%. Interestingly, there was no effect on the duration or severity of the initial episode of infection. The antibodies were not immunogenic and adverse events were not different than placebo. The trial results are consistent with previous studies showing that inadequate circulating antibody levels against *C. difficile* toxins predispose patients to symptomatic and recurrent infection, and with observational data suggesting a benefit associated with passive and active immunization for secondary prevention. Although monoclonal antibodies will not be used for primary treatment, they will likely play a role in patients at risk for recurrent infection, and they may allow a reduction in the duration of standard antimicrobial therapy for acute CDI [55]. Additional Phase III human studies are underway examining the efficacy of monoclonal antibodies against toxins A and B in *C. difficile* disease (Table 4).

Immunoglobulins

Immunoglobulins have been studied retrospectively with some success in patients with severe and recurrent CDI. Since an inability to mount a protective immune response to *C. difficile* and its toxins appears to be responsible for susceptibility to recurrent infection [56,57], this treatment strategy has some merit. More than one-half of all adults have circulating antibodies against *C. difficile* toxins and normal pooled immunoglobulins can neutralize toxins A and B [58]. Intravenous pooled human immunoglobulin products have been used off-label and on an ad hoc basis for passive immunotherapy. However, pharmacokinetic and efficacy data for these products are not available. Variable results have been reported with this treatment strategy [59–63]. No randomized trial data are available.

Fecal transplantation

The use of fecal transplant from healthy donors is another treatment modality under investigation. Stool slurries are given via nasogastric tube or as enemas to CDI patients. A recent study randomized patients (n = 43) with recurrent CDI to receive one of three therapies: An initial vancomycin regimen (500 mg orally four times per day for 4 d), followed by bowel lavage and subsequent infusion of a solution of donor feces through a nasoduodenal tube; a standard vancomycin regimen (500 mg orally four times per day for 14 d); or a standard vancomycin regimen with bowel lavage. The primary end point was the resolution of diarrhea associated with CDI without relapse after 10 wk. The study was stopped early after an interim analysis documented that 81% of the fecal transplantation cohort had resolution after the first infusion, compared with 31% (oral vancomycin therapy alone) and 23% (oral vancomycin with bowel lavage) in the other experimental cohorts [64,65]. This study, although small and unblinded, does confirm the efficacy of fecal transplantation for recurrent CDI.

Active immunization

The *C. difficile* toxoid vaccine was well tolerated and immunogenic in normal healthy subjects [66]. A report of vaccine use in three patients with recurrent infection documented that none had a subsequent relapse [67].

Currently, two vaccine candidates against CDI are being tested clinically. The more advanced program is that of Sanofi-Pasteur (Lyon, France) with the toxoid ACAM-CDIFF™ (NCT00772343 and NCT01230957). The vaccine candidate is composed of a mixture of formalin-inactivated toxins A and B that is given three times intramuscularly. The vaccine has been shown to be safe, well tolerated, and immunogenic in healthy adults [68,69]. Two phase II studies are currently being conducted in prophylactic and therapeutic settings to better define the optimal dose and to tackle the necessity of adsorbing the toxoids on aluminum salt [70]. Because the vaccine addresses an important unmet medical need, ACAM-CDIFF has been fast-tracked by the FDA.

In parallel, Intercell (Vienna, Austria) is now testing the safety, tolerability and immunogenicity of an anti-CDI vaccine candidate in an open-label, dose-escalating Phase I study in healthy volunteers (NCT01296386). In this study, one cohort includes individuals older than 65 yr. The subunit vaccine developed by Intercell/Valneva consists of a recombinant fusion protein containing a part of the receptor-binding domain of toxins A and B administered with aluminum salt and given three times intramuscularly [71,72]. Although immunization appears promising, additional trials are warranted [73].

Tolevamer, non-antibiotic polymer

Tolevamer is a toxin-binding polymer that binds to the *C. difficile* toxin in the intestinal tract at doses of 3 to 6 g daily. It has no antibiotic properties, does not alter normal gut
Table 4. Studies of Human Monoclonal Antibodies Against *Clostridium difficile* Toxins A and B

<table>
<thead>
<tr>
<th>Candidates and sponsor</th>
<th>Therapeutic agent</th>
<th>Formulation</th>
<th>Development stage</th>
<th>Patient study cohorts</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antitoxin A and antitoxin B monoclonal antibodies (Medarex and Bristol-Myers Squibb-Merck)</td>
<td>Antitoxin A and B fully human monoclonal antibodies targeting the receptor-binding domain of the toxins</td>
<td>CDA1 (antitoxin A) plus CDB1 (antitoxin B) once at 10 mg/kg in parallel to standard antibiotics</td>
<td>Phase II (completed)</td>
<td>200 patients with symptomatic CDI</td>
<td>Lowy et al. N Engl J Med 2010;362:197–205.</td>
</tr>
<tr>
<td>Biologic</td>
<td>MK-3415, MKI-3415A, MK-6072, Merck</td>
<td>MK-3415 (antitoxin A) or MK-6072 (antitoxin B) or MK-3415A (combination of the two antibodies) at a single administration of 10 mg/kg, with antibiotic therapy</td>
<td>Phase III</td>
<td>1,600 patients with symptomatic CDI and under antimicrobial therapy</td>
<td>A Phase III, Randomized, Double-Blind, Placebo-Controlled, Adaptive Design Study of the Efficacy, Safety, and Tolerability of a Single Infusion of MK-3415 (Human Monoclonal Antibody to Clostridium Difficile Toxin A), MK-6072 (Human Monoclonal Antibody to Clostridium Difficile Toxin B), and MK-3415A (Human Monoclonal Antibodies to <em>C. difficile</em> Toxin A and Toxin B) in Patients Receiving Antibiotic Therapy for <em>C. difficile</em> Infection (MODIFY I) NCT01241552 Study enrollment ongoing: October 2011 – November 2014</td>
</tr>
<tr>
<td>Biologic</td>
<td>MK-6072, MK-3415A, Placebo, Merck</td>
<td>MK-6072 (antitoxin B) or MK-3415A (combination of the two antibodies) at a single administration of 10 mg/kg, with antibiotic therapy</td>
<td>Phase III</td>
<td>1,200 patients with symptomatic CDI receiving SOC therapy</td>
<td>A Phase III, Randomized, Double-Blind, Placebo-Controlled Study of the Efficacy, Safety and Tolerability of a Single Infusion of MK-6072 (Human Monoclonal Antibody to <em>C. Difficile</em> Toxin B), and MK-3415A (Human Monoclonal Antibodies to <em>C. difficile</em> Toxin A and B) in Patients Receiving Antibiotic Therapy for <em>C. difficile</em> Infection (MODIFY II) NCT01513239 Study enrollment ongoing: February 2012 – December 2014</td>
</tr>
</tbody>
</table>

CDI = *Clostridium difficile* infection; SOC = standard of care.
Surgical Management

Timing of surgical consultation

Surgical consultation should occur early in the course of disease in patients with severe and complicated *C. difficile* colitis, as surgical decision making is challenging and the clinical course of the disease is difficult to predict in any individual patient. Early surgical consultation is recommended for the critically ill patient with CDI [77]. A recent single-center study documented that survival rates were higher in patients cared for by surgical versus nonsurgical departments, possibly because of earlier and more frequent surgical intervention [78]. Furthermore, surgical consultation does not always recommend surgical intervention. A single-center case series report of 47 patients with severe *C. difficile* colitis with surgical consultation recommended initiation of an adjuvant intracolonic vancomycin enema (1 g/500 mL normal saline every 6 h) protocol and 70% of patients responded with complete resolution without need for surgery. Multivariable logistic regression confirmed that acidosis and hypoalbuninemia were significantly associated with incomplete response to intracolonic vancomycin enema treatment [79].

Total abdominal colectomy

The indications for surgical management of patients with CDI are not defined clearly. Most clinicians advocate surgical intervention in patients with severe disease, worsening clinical condition and in those with peritonitis or shock states. Total abdominal colectomy was the standard surgical procedure performed in these cases, but mortality rates were

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**Table 5. Proposed Indications for Ileostomy with Antegrade Colonic Lavage**

<table>
<thead>
<tr>
<th>Suspected or established diagnosis of CDI plus one or more of the following:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Need for intensive care unit admission</td>
</tr>
<tr>
<td>Hypotension necessitating vasopressor support</td>
</tr>
<tr>
<td>Neurologic changes</td>
</tr>
<tr>
<td>Respiratory failure necessitating mechanical ventilation</td>
</tr>
<tr>
<td>Increasing white blood cell count (&gt; 20,000/\text{mm}^3)</td>
</tr>
<tr>
<td>Lactate concentration (&gt; 5 \text{mmol/L})</td>
</tr>
<tr>
<td>Other signs of end-organ damage or new organ failure</td>
</tr>
</tbody>
</table>

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**Table 6. Recommendations for Infection Control and Prevention of *Clostridium difficile* Infection (CDI)**

- A hospital-based infection control program can help to decrease the incidence of CDI. (Conditional recommendation, moderate-quality evidence)
- Routine screening for *C. difficile* in hospitalized patients without diarrhea is not recommended and asymptomatic carriers should not be treated. (Strong recommendation, low-quality evidence)
- Antibiotic stewardship is recommended to reduce the risk of CDI. (Strong recommendation, high-quality evidence)
- Contact precautions for a patient with CDI should be maintained at a minimum until the resolution of diarrhea. (Strong recommendation, high-quality evidence)
- Patients with known or suspected CDI should be placed in a private room or in a room with another patient with documented CDI. (Strong recommendation, high-quality evidence)
- Hand hygiene and barrier precautions, including gloves and gowns, should be used by all health-care workers and visitors entering the room of any patient with known or suspected CDI. (Strong recommendation, moderate-quality evidence)
- Single-use disposable equipment should be used for prevention of CDI transmission. Non-disposable medical equipment should be dedicated to the patient’s room and other equipment should be thoroughly cleaned after use in a patient with CDI. (Strong recommendation, moderate-quality evidence)
- Disinfection of environmental surfaces is recommended using an Environmental Protection Agency (EPA)-registered disinfectant with *C. difficile*-sporicidal label claim or 5000 p.p.m. chlorine-containing cleaning agents in areas of potential contamination by *C. difficile*. (Strong recommendation, high-quality evidence)
- Although there is moderate evidence that two probiotics (*Lactobacillus rhamnosus* GG and *Saccharomyces boulardii*) decrease the incidence of antibiotic associated diarrhea, there is insufficient evidence that probiotics prevent *C. difficile* infection. (Strong recommendation, low-quality evidence)

From Surawicz et al. [25]. (Reprinted by permission from Macmillan Publishers Ltd.)
high [80,81] and most survivors required permanent ileostomy with long-term complications related to fluid losses from the ileostomy. Common indications for colectomy include colonic perforation, toxic megacolon with impending perforation, peritonitis, severe sepsis or septic shock, need for vasopressor support, new end-organ dysfunction or failure, and colitis refractory to medical management. All of these factors also are significant predictors of mortality.

A systematic review and meta-analysis of outcomes following emergency surgery for *C. difficile* colitis included 31 studies (n = 1,433), and the most commonly performed surgery was total colectomy with end-ileostomy (89%; 1,247 of 1,401 surgical procedures). When total colectomy was not performed, additional bowel resection was required in 15.9% of patients. The 30-d mortality rate was 41.3%. The strongest predictors of post-operative death were those relating to pre-operative physiologic status: pre-operative intubation, acute renal failure, multiple organ failure, and shock requiring vasopressors [82].

Diverting loop ileostomy and colonic irrigation

More recently, an alternative surgical procedure to total abdominal colectomy has emerged with creation of a loop ileostomy, intraoperative colonic lavage via the ileostomy, and post-operative antegrade vancomycin flushes via the ileostomy (Fig. 6). The initial report of this procedure in 42 patients documented that the procedure was accomplished laparoscopically in 35 (83%) patients. The new treatment strategy resulted in reduced mortality, compared with the authors’ historical population who had undergone total abdominal colectomy (19% vs. 50%, OR 0.24; p = 0.006). Preservation of the colon was achieved in 39 (93%) patients [83].

This new procedure is a potential surgical advance in the treatment of severe life-threatening CDI. Proposed indications for this surgical procedure are listed in Table 5. A single-institution prospective randomized trial is ongoing at the Massachusetts General Hospital (NCT01441271) comparing ileal diversion and colonic lavage with total abdominal colectomy for the treatment of fulminant *C. difficile* colitis.

Infection Control

Transmission of *C. difficile* is usually person-to-person, and via a fecal-oral route. The hands of healthcare workers and environmental fomites (such as bedpans, commodes, and rectal probes) are key players for spread of the bacterium and spores in acute care facilities. Thus, the best mode of preventing transmission is with the use of full contact isolation precautions (e.g., gloves, barrier gowns, private rooms, and disposable or dedicated supplies and equipment), as well as good hand hygiene and environmental cleaning. Contact precautions should be maintained at least until diarrhea is resolved. It is important to note that alcohol-based solutions do not kill *C. difficile* spores, and thus hand washing with soap and water is required to prevent spread via healthcare workers’ hands [84]. Likewise, a bleach-based solution (5,000 ppm chlorine) or Environmental Protection Agency–registered disinfectant with a *C. difficile* sporicidal claim is required to kill spores in the environment. General guidelines for infection control and prevention are summarized in Table 6.

Summary

*C. difficile* infection incidence and severity has grown dramatically in the past two decades, gaining increased significance as a clinically important disease entity. Treatment modalities vary based on CDI classification with regards to severity and timing of onset. The mainstay of medical treatment remains metronidazole (preferred due to substantial cost differential) and oral/rectal vancomycin. Other treatment modalities may be used in special circumstances, and more trials are underway to evaluate adjunctive treatment options such as monoclonal antibodies, vaccination, and fecal transplant. Early surgical consultation in severe cases of CDI and in patients with deteriorating clinical condition may be life-saving. Diverting loop ileostomy for antegrade colonic irrigation may allow for colonic preservation in the appropriate patient population. Infection control measures both at the individual as well as institution levels are vital to mitigating the spread of CDI.

Author Disclosure Statement

No competing financial interests exist.

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