Treatment Strategies for Recurrent Clostridium difficile Infection

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ABSTRACT
Background: Recurrent Clostridium difficile infection represents a major clinical challenge. Treatment is often based on empiric selection from relatively few options supported by limited clinical evidence.

Objective: To review and evaluate the literature on therapeutic alternatives for recurrent C. difficile infection.

Data Sources: The MEDLINE, PubMed, Embase, and Cochrane databases were searched from inception to 2013 for published evidence in English on the treatment of recurrent C. difficile infection. The search terms were “Clostridium difficile”, “recurrent” or “relapse”, and “treatment”.

Study Selection and Data Extraction: Studies of any design were eligible for inclusion. Two reviewers assessed abstracts, full articles, and reference lists from retrieved articles and clinical practice guidelines to identify relevant literature.

Data Synthesis: The evidence to guide treatment of recurrent C. difficile infection is limited, with 24 studies meeting the inclusion criteria for this review. A repeat course of oral metronidazole or vancomycin is recommended for treatment of mild to moderate first recurrences and has not been found to influence the likelihood of subsequent recurrence. Oral vancomycin may be preferred for more severe infections; however, the severity score warrants further study and validation. For the treatment of second and subsequent recurrences, tapered or pulsed vancomycin regimens have been recommended in practice guidelines, despite very limited clinical evidence. Similarly, the potential benefits of longer treatment courses of oral vancomycin for second and subsequent recurrences warrant investigation. The potential role, including costs and benefits, of new agents such as fidaxomicin in the treatment of recurrent C. difficile infection remains to be determined. Although there is insufficient evidence to recommend probiotics as an adjunct to conventional treatment for recurrent infection, there may be benefit in terms of prevention.

Conclusions: This literature review identified significant limitations in currently recommended interventions for the treatment of recurrent C. difficile infection. It has also provided insight into the available evidence for determining the appropriateness of therapy for patients with recurrent infection.

Keywords: recurrent Clostridium difficile infection, relapse, vancomycin, metronidazole, fidaxomicin, literature review


RÉSUMÉ
Contexte: Les récidives d’infection à Clostridium difficile constituent un défi clinique majeur. Le traitement est souvent basé sur un choix empirique parmi relativement peu d’options étayées par des données cliniques probantes limitées.

Objectif: Examin er et évaluer les traitements de rechange des récidives d’infection à C. difficile dans la littérature.


 Sélection des études et extraction des données: Toutes les études, peu importe la méthodologie, étaient admissibles. Deux examineurs ont évalué les résumés, les articles complets, et les listes de références des articles et des guides de pratique clinique recensés afin de répertorier la littérature pertinente.

Synthèse des résultats: Les données guidant le traitement des récidives d’infection à C. difficile se limitent à vingt-quatre études qui ont satisfait aux critères d’admissibilité à cet examen documentaire. Une reprise du traitement par le métronidazole ou la vancomycine par voie orale est recommandée pour les premières récidives légères ou modérées et s’est révélée sans incidence sur le risque d’autres récidives. La vancomycine orale pourrait être mieux indiquée dans les cas d’infection plus grave; en revanche, le score de gravité mérite de faire l’objet d’études plus poussées et d’être validé. Pour le traitement des deuxièmes récidives et des récidives subséquentes, des traitements dégressifs ou intermittents par la vancomycine ont été recommandés dans les guides de pratique, malgré des données cliniques probantes très limitées. De même, les atouts potentiels des traitements de plus longue durée par la vancomycine orale dans les cas de deuxièmes récidives et de récidives subséquentes doivent être davantage étudiés. Le rôle potentiel, y compris les coûts et les bénéfices, des nouveaux agents comme la fidaxomicine dans le traitement des récidives d’infection à C. difficile reste à déterminer. Malgré les données probantes insuffisantes pour recommander les probiotiques comme adjuvant au traitement classique des récidives d’infection, ces agents pourraient exercer des effets préventifs favorables.

Conclusion: Cet examen de la littérature a relevé des limites considérables dans les interventions actuellement recommandées pour le traitement des récidives d’infection à C. difficile. Il a aussi jeté un nouvel éclairage sur les données disponibles permettant de déterminer la pertinence du traitement dans les cas de récidive de l’infection.

Mots clés: récidive d’infection à Clostridium difficile, rechute, vancomycine, métronidazole, fidaxomicine, examen de la littérature

[Traduction par l’éditeur]
INTRODUCTION

Recurrent Clostridium difficile infection, defined as symptoms and a positive toxin result within 8 weeks after the end of therapy for an initial episode, represents a major clinical challenge. A first recurrence occurs in 20% to 30% of patients treated for an initial episode, and subsequent (second or later) recurrences are observed in 40% to 60% of those cases. C. difficile infection, especially recurrent disease, is associated with significant patient morbidity and can lead to serious and even life-threatening complications such as pseudomembranous colitis, bowel perforation, and sepsis. Reports indicate that the incidence of C. difficile infection, including recurrent infection, is on the rise.

Recurrent episodes may be due to re-infection from persistent spores or new infection with a different strain of C. difficile. The risk of re-infection increases with continued use of antimicrobials, advanced age, history of recurrence, and deficiency in the immune response to C. difficile toxins. Recurrence may also be influenced by bacterial strain; for example, the toxin hyper-producing BI/NAP1/027 (NAP1) strain emerged during outbreaks a decade ago and now accounts for 31% to 53% of infections. The NAP1 strain is associated with significant patient morbidity and mortality and high rates of severe, refractory recurrent C. difficile infection. The relative risks of patient-related versus strain-related factors for recurrence are unknown, as is the comparative efficacy of first-line agents in treating infections associated with the NAP1 strain of C. difficile.

The treatment of recurrent C. difficile infection is often based on empiric selection from relatively few options supported by limited clinical evidence. Accordingly, this paper reviews the recommendations of the Society for Hospital Epidemiology of America and the Infectious Diseases Society of America (SHEA–IDSA; guidelines published in May 2010) and the American College of Gastroenterology (ACOG; guidelines published in February 2013), as well as other references describing treatment strategies for first and second or later recurrence of C. difficile infection.

METHODS

A literature search was conducted in the MEDLINE, PubMed, Embase, and Cochrane databases from inception to 2013 for studies in English that evaluated the treatment of recurrent C. difficile infection in adults. All relevant randomized controlled trials (RCTs), observational studies, case series, and case reports were identified using the MeSH (Medical Subject Heading) search terms “Clostridium difficile”, “recurrent” or “relapse”, and “treatment”. Reference lists from retrieved articles and clinical guidelines were also searched for relevant literature. Two reviewers (C.L. and S.Z.) assessed abstracts, full articles, and reference lists from retrieved articles. The reviewers worked collaboratively to determine the relevance of each article to the topic of interest (treatment of recurrent C. difficile infection).

RESULTS

The literature search yielded 24 studies with information about treating recurrent C. difficile infection.

Treatment of First Recurrence

A repeat course of oral metronidazole or oral vancomycin, or a switch from oral metronidazole to oral vancomycin, has been the mainstay treatment for a first recurrence of C. difficile infection. Although there is evidence to support the use of metronidazole or vancomycin for the initial episode of C. difficile infection, there is limited evidence to support the repeat use of either agent in the setting of recurrent infection. The SHEA–IDSA and ACOG guidelines both recommend repeating a 10- to 14-day course of oral metronidazole (500 mg every 8 h) or vancomycin (125 mg every 6 h). Vancomycin is the preferred antibiotic for severe recurrent C. difficile infection (Table 1). Failure to respond to metronidazole therapy within 5 to 7 days should prompt consideration of a change in therapy to vancomycin. Both guidelines acknowledge the lack of high-quality evidence to support these recommendations.

The use of either metronidazole or vancomycin for treatment of an initial episode has not been found to influence the likelihood of subsequent recurrence. One early prospective trial found that metronidazole was noninferior to vancomycin, with cure rates exceeding 90% after treatment of the initial C. difficile infection. However, more recent data, including results for patients with NAP1 infection, have indicated treatment failure rates up to 50% with metronidazole. In one prospective observational study of 52 patients with C. difficile infection, with 60% of cases associated with the NAP1 strain, those treated with vancomycin were more likely to have undetectable levels of C. difficile toxin (adjusted hazard ratio [HR] 3.99, 95% confidence interval [CI] 1.41–11.3; p = 0.09) and resolution of diarrhea (adjusted HR 4.17, 95% CI 1.53–11.4; p = 0.05) during the first 5 days of therapy than those who received metronidazole. Furthermore, therapy was changed because of persistent symptoms in 29% (10/34) of patients treated with metronidazole but none of the 18 treated with vancomycin. The effect of the NAP1 strain on recurrence was investigated in a retrospective cohort study of 1616 patients with C. difficile infection. The 60-day probability of recurrence in the pre-epidemic period, from 1991 to 2002, was 19.6% (108/551) and 20.3% (13/64) in those treated with metronidazole and vancomycin, respectively. In the post-epidemic period, from 2003 to 2004, the recurrence rates were
44.6% (157/352) with metronidazole and 38.7% (29/75) with vancomycin. However, the statistical significance of these findings was not determined, and the authors noted that the higher risk of recurrence during the 2003–2004 period may have resulted from re-infections rather than relapses. Overall, these findings suggest a possible association between the use of vancomycin and lower rates of subsequent recurrence in the setting of NAP1 infections.

No studies evaluating treatment alternatives in the setting of severe recurrent \textit{C. difficile} infection were found. However, patients with a severe first recurrence are to be treated similarly to those with an initial episode of severe \textit{C. difficile} infection.\textsuperscript{11} As such, there are some limitations worth noting in the evidence related to treatment of an initial episode of severe infection. For instance, the use of vancomycin for initial episodes of “severe” \textit{C. difficile} infection was supported by an RCT with 150 patients stratified as having mild or severe disease.\textsuperscript{23} Those with severe disease experienced a higher rate of cure (defined as resolution of diarrhea and negative results on testing for \textit{C. difficile} toxin) with vancomycin (125 mg liquid 4 times daily; cure rate 97% [30/31]) than with metronidazole (250 mg tablet 4 times daily; cure rate 76% [29/38]) (\(p = 0.02\)), after 10 days of treatment. Among those with mild disease, no significant difference was observed between vancomycin and metronidazole (98% [39/40] versus 90% [37/41]; \(p = 0.36\), and the authors concluded that vancomycin may be preferred for more severe infection. However, the severity score and criteria for “severe” disease were relatively comprehensive (Table 1) including, for example, all patients over 60 years of age with fever or leukocytosis. The severity score and its ability to predict severe clinical infection require prospective validation. In addition, the total daily dose of metronidazole used in this study (1000 mg per day) was lower than that recommended by clinical guidelines (1500 mg per day).\textsuperscript{1,2}

Although there were no studies specifically examining patients with severe recurrent \textit{C. difficile} infection or patients with intolerance to oral medication (e.g., those with impaired gastrointestinal absorption or ileus requiring alternate routes of administration), it is useful to examine the interventions that have been used in this setting. Application of retention enemas containing 500 mg vancomycin in 500 mL normal saline (0.9% sodium chloride) every 6 h is recommended\textsuperscript{1} on the basis of a case series of 9 patients with \textit{C. difficile} infection (4 of whom had a history of infection 6 weeks before admission).\textsuperscript{26} A volume of at least 500 mL (more than that previously recommended by the SHEA-IDSA guidelines\textsuperscript{*}) is thought to improve delivery to the ascending and transverse colon.\textsuperscript{1} However, since rectal administration may not deliver sufficient concentrations of vancomycin to the ascending and transverse colon, concurrent treatment with parenteral metronidazole is recommended.\textsuperscript{1} Because of limited transport across the intestinal membranes, parenteral vancomycin is not used. Notably, the use of parenteral metronidazole as standard treatment for severe infection in those who cannot tolerate oral therapy is also based on limited studies. The relationship between inflammation and drug transport across intestinal

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### Table 1. Criteria for Severe \textit{Clostridium difficile} Infection

<table>
<thead>
<tr>
<th>SHEA-IDSA\textsuperscript{*}</th>
<th>American College of Gastroenterology\textsuperscript{5}</th>
<th>Zar et al.\textsuperscript{22} *</th>
</tr>
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<tbody>
<tr>
<td>WBC\textsuperscript{†} (\geq 15000) cells/µL OR Serum creatinine (\geq 1.5 \times) premorbid value</td>
<td>Severe disease: Hypoalbuminemia (&lt; 30 g/L), plus ONE of the following: • Abdominal distension • Elevated WBC\textsuperscript{†} ((\geq 15000) cells/mm(^3))</td>
<td>Age (&gt; 60) years (1 point)</td>
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<td>Severe and complicated disease (any ONE of the following): • Admission to ICU • Hypotension (\pm) vasopressor use • Fever (temperature &gt; 38.5°C) • Ileus or significant abdominal distension • Mental status changes • WBC\textsuperscript{†} (\geq 35000) cells/mm(^3) or &lt; 2000 cells/mm(^3) • Serum lactate &gt; 2.2 mmol/L. • End organ failure</td>
<td>Temperature &gt; 38.3°C (1 point)</td>
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\(\text{ICU}\) = intensive care unit, SHEA-IDSA = Society for Hospital Epidemiology of America and the Infectious Diseases Society of America, WBC = white blood cells.

\textsuperscript{*}Severe disease defined as a score of 2 or greater.

\textsuperscript{†}WBC counts in conventional units, as presented in the original sources; for conversion to SI unit (\(\times 10^9/L\)), multiply count per cubic millilitre or per microlitre by 0.001 (e.g., 15 000 cells/mm\(^3\) = 15 \(\times 10^9/L\)).
membranes was examined in a case series of 9 patients with documented *C. difficile* diarrhea; this study showed that fecal metronidazole concentrations decreased as the diarrheal symptoms improved.37 No significant difference was found between oral and parenteral routes of administration. In a more recent observational study, Wenisch and others38 found that parenteral metronidazole was inferior to oral metronidazole or vancomycin, with 30-day all-cause mortality rates of 38% (16/42), 7% (9/121), and 10% (4/42), respectively. The authors suggested that concentrations of metronidazole in the colon might be lower with parenteral administration. A few possible explanations for this finding include inadequate transit of metronidazole in the absence of mucosal inflammation during mild disease or occurrence of higher colon concentrations with oral metronidazole through metabolic auto-inhibition or enterohepatic recirculation in the intestine.29,30 Limitations of the study included enrolment of patients with mild disease, defined as those with fewer than 4 stools per day and no signs of severe colitis. Furthermore, the mortality rate among those receiving parenteral metronidazole was considerably higher than that reported for mild disease.39 Although the authors ensured similar baseline age and comorbidities between the groups, selection bias and the influence of unknown confounders on the primary outcome of all-cause mortality cannot be ruled out. The observations, however, draw attention to the relative lack of study of the pharmacokinetics and efficacy of parenteral metronidazole for the treatment of severe *C. difficile* infection.

**Treatment of Second or Later Recurrence**

The treatment options for patients who experience second or later recurrences of *C. difficile* infection are limited. Repeat courses of metronidazole are discouraged because of the risk of neurotoxicity associated with prolonged exposure to this drug.35 For example, peripheral neuropathy has been associated with metronidazole use for periods of weeks to months.35 Cerebellar dysfunction, altered mental status, and seizures were also described in 64 patients exposed to metronidazole at a mean cumulative dose of 93.4 g (range 250 mg to 1095 g) for a median duration of 54 days.31 Therefore, oral vancomycin may be preferred for patients with a second or later recurrence of *C. difficile* infection or predisposing neurologic conditions. A standard 10- to 14-day course of vancomycin (125 mg every 6 h) may be used for those with failure of standard metronidazole treatment during the initial episode and first recurrence of *C. difficile* infection.1,2

Although the SHEA–IDSA and ACOG guidelines proposed pulsed or tapered vancomycin regimens for those with multiple recurrences of *C. difficile* infection,1,2 this approach is largely theoretical. One example of such therapy is the standard 10- to 14-day course of vancomycin 125 mg orally 4 times daily followed by 125 mg twice daily for 1 week, 125 mg once daily for 1 week, and then 125 mg daily pulsed every 2 or 3 days for 2 to 8 weeks.3 Pulsed or tapered regimens are thought to be beneficial in suppressing formation of *C. difficile* spores and restoring normal gut flora. It is important to note that there are limited data to support the guideline’s claim that “A substantial proportion of patients with a second recurrence will be cured with a tapering and/or pulsed regimen of oral vancomycin”1.

McFarland and others32 conducted a secondary analysis of 2 RCTs comparing *Saccharomyces boulardii* with placebo as adjunct therapy to various regimens of oral vancomycin or metronidazole. For their analysis, they selected 163 patients from the placebo arms of the 2 trials; these patients had recurrent *C. difficile* infection and an average of 3.2 episodes (range 1 to 14). A fixed vancomycin dose (1000 mg per day) was used as the comparator for 4 other vancomycin regimens (< 1000 mg per day, ≥ 2000 mg per day, tapered, or pulsed), 3 metronidazole regimens (≤ 1000 mg per day, 1500 mg per day, or 2000 mg per day), and a miscellaneous group of regimens involving vancomycin plus rifampin or vancomycin plus metronidazole. Treatment durations ranged from 1 to 2 weeks for the fixed dose, from 19 to 25 days for the tapered dose, and from 9 to 20 days for the pulsed regimens. The main observation was a lower recurrence rate, relative to the fixed vancomycin dose (10/14 [71%]) with the tapered regimen (9/29 [31%]; p = 0.01) and the pulsed regimen (1/7 [14%; p = 0.02). Of note was the exceptionally high recurrence rate for the primary comparator (i.e., fixed vancomycin dose). Other significant limitations included the nonrandomized study design, variable treatment regimens, small sample sizes, and apparent use of multiple comparisons among regimens without appropriate statistical corrections.33 The influence of important factors such as duration of therapy were not adequately considered, even though the tapered vancomycin regimens were significantly longer than the fixed-dose regimens (mean of 21.5 days versus 7 to 14 days). At best, the study by McFarland and others32 described tapered and pulsed vancomycin regimens in a small number of patients but provided limited evidence as to the effectiveness of these regimens in the treatment of recurrent *C. difficile* infection.

**Fidaxomicin**

Fidaxomicin (Dificid) received notice of compliance for the treatment of *C. difficile* infection from Health Canada in June 2012.34 It is a narrow-spectrum macrocyclic antibiotic that exhibits minimal systemic absorption. It is bactericidal and demonstrates prolonged post-antibiotic effects against *C. difficile*.35 Fidaxomicin has been shown to be safe and effective for the first episode of *C. difficile* infection36,37; however, there is limited evidence for recurrent infection.
Fidaxomicin was compared with vancomycin in 2 RCTs involving a total of more than 500 patients with mild to moderate *C. difficile* infection. Approximately 16% of the patients had had a previous episode of infection. Both trials found that 10 days of oral fidaxomicin (200 mg twice daily) was noninferior to 10 days of oral vancomycin (125 mg 4 times daily) in terms of clinical cure. In the first trial, by Louie and others, 88.2% (253/287) of patients in the fidaxomicin group and 85.8% (265/309) of those in the vancomycin group experienced clinical cure (one-sided 97.5% confidence limit −3.1%). In the second trial, Cornely and others reported a clinical cure rate of 87.7% (221/252) for the fidaxomicin group and 86.8% (223/257) for the vancomycin group (one-sided 97.5% confidence limit −4.9%). Fidaxomicin was also associated with a lower recurrence rate within 28 days of clinical cure relative to vancomycin (15.0% versus 25.0%, respectively; \( p = 0.005; \) relative risk [RR] 0.39). Intestinal effects, such as nausea and abdominal discomfort, were the most commonly reported adverse effects in patients receiving fidaxomicin.

Cornely and others conducted a retrospective subgroup analysis in the 128 patients with recurrent *C. difficile* infection in the aforementioned studies. In these patients, initial response, defined as clinical cure after 8 or more days of therapy, was similar for fidaxomicin and vancomycin. Recurrence within 28 days, however, was significantly less likely among patients who received fidaxomicin (20% [13/66] versus 35% [22/62], \( p = 0.045 \))

Although this was a retrospective subgroup analysis with small sample size, these findings suggest that fidaxomicin may have a role in treating recurrent *C. difficile* infection. Of note, fidaxomicin has not been compared with metronidazole, and additional prospective studies are also needed in patients with severe *C. difficile* infection, as well as those with multiple episodes of recurrent infection.

Drug cost is an important consideration in defining the role of fidaxomicin in the treatment of *C. difficile* infection, including recurrent infection. The cost of this drug is approximately $2200 for 10 days of treatment, whereas the corresponding costs are $550 for vancomycin and $25 for metronidazole. The Canadian Drug Expert Committee of the Canadian Agency for Drugs and Technologies in Health has recommended that fidaxomicin not be listed on formulary at this price. In addition, one economic analysis concluded that the use of fidaxomicin is not a cost-effective option for first-line treatment of *C. difficile* infection. According to the authors of this study, the cost of a course of treatment with fidaxomicin would have to be no more than $150 to be considered cost-effective. Interestingly, selective treatment of patients with the non-NAP1 strain of *C. difficile* is more cost-effective; however, the incremental cost-effectiveness ratio was estimated to be greater than $43.7 million per quality-adjusted life-year in the setting of screened patients.

### Other Alternatives

Other alternatives for treating recurrent *C. difficile* infection that have been investigated in clinical trials include anion exchange resins, other antibiotics (such as rifaximin), probiotics, and fecal microbiota transplant. The clinical guidelines do not support the use of anion exchange resins or other antibiotics in the management of recurrent infection. Anion exchange resins, such as cholestyramine, are thought to bind toxins produced by *C. difficile* without altering the bowel flora. However, they have also been found to bind oral vancomycin, which results in reduced efficacy of the drug, and concurrent use of resins with vancomycin should therefore be avoided.

There have been no studies in which binding of anion exchange resins to metronidazole was observed; however, this theoretical concern should not be ruled out. Rifamixin is a nonabsorbable rifampicin antibiotic that is not available in Canada but has been used in the United States for recurrent *C. difficile* infection. In one pilot study of 68 patients with *C. difficile* infection, administration of rifaximin following standard antibiotic therapy (with 56 patients [82%] receiving metronidazole and 12 [18%] receiving oral vancomycin for 10 to 14 days; standard doses not specified) resulted in a reduction in recurrent *C. difficile* infection relative to placebo; however, this finding was not statistically significant (15% [5/33] versus 31% [11/35]; \( p = 0.11 \)). In this study, the mean age was 61 years, 34 (50%) of the patients were female, and 13 (19%) had had a previous episode of *C. difficile* infection.

There is insufficient evidence to recommend probiotic therapy as an adjunct to conventional treatment for recurrent *C. difficile* infection. One Cochrane review of 4 small randomized, prospective studies found no overall benefit of probiotics in conjunction with conventional treatment for initial or recurrent episodes of *C. difficile* infection. In one of the studies included in the Cochrane review, there was a significant reduction in recurrent *C. difficile* diarrhea with *S. boulardii* in addition to conventional therapy relative to placebo (RR 0.59, 95% CI 0.35–0.98). In another study included in the Cochrane review, there was a significant reduction in recurrent *C. difficile* infection with *S. boulardii* (relative to placebo) only when used in conjunction with high-dose vancomycin (2 g per day) (RR 0.33, 95% CI 0.10–1.06). However, there was no significant difference in recurrent *C. difficile* infection when *S. boulardii* was used in combination with low-dose vancomycin (500 mg per day) or metronidazole. No statistically significant benefit was found with *Lactobacillus plantarum* or *Lactobacillus rhamnosus* GG in the remaining 2 studies.

Despite the limitations of probiotics in treating *C. difficile* infection, there is some evidence to support their value in...
preventing such infection. In one meta-analysis of 22 trials (n = 3818), the incidence of C. difficile–associated diarrhea was approximately 66% lower among patients receiving probiotics than among those receiving placebo or no treatment (pooled RR 0.34, 95% CI 0.24–0.49; I² = 0%). These agents are not recommended for immunocompromised, critically ill, or elderly patients or for patients with inflamed or leaky gastrointestinal mucosa, as there have been cases of invasive infection in such patients. For most probiotics, there are also some concerns about quality control and inconsistency of the quantity of live organisms. The clinical guidelines acknowledge the lack of convincing evidence for the use of adjunctive probiotics to reduce recurrent C. difficile infection.

Fecal microbiota transplant (FMT) may have a role for patients who have had 3 or more recurrent infections, and the ACOG guidelines recommend consideration of this approach after a third recurrence of C. difficile infection following a pulsed vancomycin regimen. FMT involves transplanting stool from a healthy donor into the patient with recurrent C. difficile infection. Patients with recurrent infection may have an imbalance in the amount of healthy colonic flora, and introducing normal fecal bacteria from a healthy donor may be restorative. Various methods have been used for FMT, including retention enemas, nasoduodenal tube, and colonoscopy. Approximately 325 cases of FMT have been reported in the literature. Kassam and others performed a systematic review of 11 studies (with no RCTs identified) and found that 245 of 273 patients with C. difficile infection experienced clinical resolution of diarrhea with FMT treatment (weighted pooled resolution rate 89.1%, 95% CI 84%–93%), with no statistically significant heterogeneity among the studies (I² = 33.7%). In a retrospective case series of 70 patients with recurrent C. difficile infection, symptom resolution was reported at 12 weeks for all patients without the NAP1 strain and for 89% (32/36) of those with the NAP1 strain. The US Food and Drug Administration recently categorized FMT as a “drug and biologic product”, and hence an Investigational New Drug application is required for its use.

One open-label RCT involving adults with recurrent C. difficile compared a standard 14-day course of vancomycin (500 mg 4 times daily) with (1) a 14-day course of the same drug therapy with bowel lavage on day 4 or 5 and (2) a 5-day course of drug therapy with bowel lavage on the last day followed by FMT via nasoduodenal infusion the next day. Thirteen (81%) of the 16 patients in the FMT group, 4 (31%) of the 13 patients in the vancomycin-only group, and 3 (23%) of the 13 patients in the vancomycin plus bowel lavage group experienced resolution of C. difficile–associated diarrhea without recurrence after 10 weeks (p < 0.001; rate ratio 3.05 [99.9% CI 1.08–290.05] between the FMT and vancomycin-only group and 4.05 [99.9% CI 1.21–290.12] between the FMT and vancomycin with bowel lavage group). Mild diarrhea and abdominal cramping were reported more often in the infusion group; however, no significant difference was observed relative to the other treatment groups. A major limitation of concern was early termination because of lower-than-expected cure rates in the control group (patients receiving vancomycin). Given the small sample size, it is difficult to determine the effect of baseline differences on the results of this study. Long-term data for this intervention are limited, and the potential for transmission of infectious organisms is of concern.

**CONCLUSION**

Recurrent C. difficile infection remains a challenging illness for which an optimal treatment approach has not been well established. Recommendations presented in formal guidelines are largely supported by expert opinion and small case series.

The literature review reported here indicates that a repeat course of metronidazole or vancomycin is considered appropriate for the treatment of mild to moderate first recurrence of C. difficile infection. Although oral vancomycin is recommended for more severe C. difficile infection, a validated severity score has yet to be established, and comparative data with recommended metronidazole doses are not available. Parenteral metronidazole is a suggested addition to oral vancomycin for critically ill patients with ileus. However, in one observational study, the mortality rate was higher with parenteral metronidazole than with oral alternatives in patients with mild to moderate disease. The clinical impact of parenteral metronidazole in severe disease has not been studied. For the treatment of second or later recurrence, a tapered or pulsed regimen is recommended, but the best available evidence in support of this practice is limited. A longer (3-week) treatment course of oral vancomycin may have some benefit for patients with a second or later recurrence; however, further studies are needed to support this practice. Fidaxomicin has shown promise in patients with mild to moderate C. difficile infection, and new data are emerging to support its efficacy in patients with recurrent C. difficile infection. FMT may have a role in patients who have experienced 3 or more recurrences; however, RCTs and long-term efficacy and safety data are still needed.

There has been recent concern regarding the toxin hyper-producing NAP1 strain of C. difficile; however, in the majority of studies examining patients with recurrent infection, this strain has not been implicated.

Only a few studies have exclusively examined patients with second or later recurrence. More research in this area is needed given the limited options available at this stage.

This literature review has identified substantial limitations in currently recommended interventions for the treatment of recurrent C. difficile infection. It also provides insight into
the available evidence for determining the appropriateness of therapy for patients presenting with recurrent infection of this type.

References


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