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Cost-Effectiveness of Competing Treatment Strategies for *Clostridium difficile* Infection: A Systematic Review

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Abstract

BACKGROUND—*Clostridium difficile* infection (CDI) presents a substantial economic burden and is associated with significant morbidity. While multiple treatment strategies have been evaluated, a cost-effective management strategy remains unclear.

OBJECTIVE—We conducted a systematic review to assess cost-effectiveness analyses of CDI treatment and to summarize key issues for clinicians and policy makers to consider.

METHODS—We searched PubMed and 5 other databases from inception to August 2016. These searches were not limited by study design or language of publication. Two reviewers independently screened the literature, abstracted data, and assessed methodological quality using the Drummond and Jefferson checklist. We extracted data on study characteristics, type of CDI, treatment characteristics, and model structure and inputs.

RESULTS—We included 14 studies, and 13 of these were from high-income countries. More than 90% of these studies were deemed moderate-to-high or high quality. Overall, 6 studies used a decision-tree model and 7 studies used a Markov model. Cost of therapy, time horizon, treatment cure rates, and recurrence rates were common influential factors in the study results. For initial CDI, fidaxomicin was a more cost-effective therapy than metronidazole or vancomycin in 2 of 3 studies. For severe initial CDI, 2 of 3 studies found fidaxomicin to be the most cost-effective therapy. For recurrent CDI, fidaxomicin was cost-effective in 3 of 5 studies, while fecal microbiota transplantation (FMT) by colonoscopy was consistently cost-effective in 4 of 4 studies.

CONCLUSIONS—The cost-effectiveness of fidaxomicin compared with other pharmacologic therapies was not definitive for either initial or recurrent CDI. Despite its high cost, FMT by

SUPPLEMENTARY MATERIAL

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colonoscopy may be a cost-effective therapy for recurrent CDI. A consensus on model design and assumptions are necessary for future comparison of CDI treatment.

Clostridium difficile infection (CDI) is one of the most common healthcare-associated infections in North America and Europe.¹ In 2011, the estimated incidence of CDI in the United States was approximately 453,000.² The management of CDI remains complicated because of epidemic strains (BI/NAP1/027) introduced in 2005 and because disease severity varies.^{3,4} In addition, patients often have multiple and frequent recurrences,⁵ which exacerbate the disease burden and increase medical costs. The most common risk factors for CDI recurrence include age 65 years, severe underlying comorbidities, and concomitant use of antibiotics.^{6,7} *Clostridium difficile* infection continues to impose a significant economic burden on the US healthcare system, estimated to be more than \$5.4 billion (2014 US dollars).⁸

The current guidelines for CDI management recommend either metronidazole, vancomycin, fidaxomicin, or fecal microbiota transplantation (FMT), depending on disease severity and the presence and number of recurrences.^{3,9–11} Current treatment choices and available algorithms make it difficult for physicians to tailor individualized therapies for patients. While newer drugs and therapies may be more effective, they are also more expensive. In the past few years, several cost-effectiveness analyses of different CDI treatment strategies have been conducted to support evidence-based decision making,^{12–17} but the results were mixed. A previous review summarized the economics of CDI treatments, but it did not include study quality assessments and based recommendations on partial costing or comparative effectiveness studies.¹⁸ Therefore, the aim of this systematic review was to critically assess the available literature on economic evaluations of various treatment modalities for initial and recurrent CDI. Based on model comparison, we summarized the findings about treatment modalities and key issues for clinicians to consider when treating patients with CDI, to inform health policy makers, and to identify important areas for future cost-effectiveness research.

METHODS

We conducted a systematic review following the Preferred Reporting Items for Systematic reviews and Meta-Analysis (PRISMA) reporting guideline¹⁹ and a measurement tool for the Assessment of Multiple Systematic Reviews (AMSTAR) standard for quality of execution.²⁰

Search Strategy

Studies were included if they (1) were original analyses; (2) were full cost-effectiveness analysis (CEA), cost-utility analysis (CUA), cost-benefit analysis (CBA), or a combination of CEA-CUA or CEA-CBA; and (3) examined treatment modalities that were approved for patient use. Studies were excluded if they (1) did not estimate cost per unit of health outcomes; (2) only addressed CDI diagnostic tests, prevention strategies, and hypothetical or under-investigation treatments; or (3) were an editorial, comment, review, letter to the editor, or conference abstract. In case of multiple publications using the same cost-effectiveness model and data, the more recent and comprehensive study was included. All studies using similar models for different treatments, populations, or types of CDI were included.

Independently, 2 investigators (P.L. and V.T.N.) identified relevant articles by searching PubMed/MEDLINE, Cochrane Library, Web of Science, EMBASE, and Scopus databases from inception through August 2016. We also searched the British National Health Service (NHS) Economic Evaluation database and the reference lists of included studies. The search terms were "*Clostridium difficile*," "*C. difficile*," "economic," "economic evaluation," "cost," "cost-effectiveness," "cost-utility," and "cost-benefit." The full PubMed search strategy is available as supplementary material. After reviewing the study title and abstract, P.L. and V.T.N. selected articles and independently reviewed the full text to determine inclusion. All disagreements were resolved through discussion with the third investigator (A.D.).

Data Extraction

Independently, 2 investigators (P.L. and V.T.N.) extracted relevant data using a uniform data extraction tool (available as Supplementary Table 1). We extracted information on study characteristics (authors, publication year, country, funding sources), type of CDI (initial, recurrent), treatment characteristics (types, medication dose and administration route, and mode of delivery of FMT), model structure (design, population, perspective, time horizon, discount rate), epidemiological data related to CDI and treatment effectiveness, types of costs and values, cost year and currency, outcome measures, the incremental cost-effectiveness ratio, decision threshold, and sensitivity analyses. We summarized data by type of CDI. Cost-effectiveness findings were additionally stratified by funding source.

Quality Assessment

We assessed study quality using the *British Medical Journal's* Drummond and Jefferson checklist.²¹ We adapted the checklist to include 3 additional items: generalizability, source of funding, and conflict of interest based on the Consolidated Health Economic Evaluation Reporting Standards checklist.²² Each item in the checklist has a 'Yes', 'No' or 'Not applicable' (NA) option and was scored 1, 0, or no score, respectively (available as Supplementary Table 2). The overall quality score was then calculated as the percentage of 'Yes' responses out of the total criteria applicable to each individual study. For example, if a paper had 27 Yes, 7 No, and 4 NA, the quality score was calculated as 71% (27 of 34). Based on its quality score, each study was ranked as either low quality (<50%), moderate quality (50%–64%), moderate-to-high quality (65%–80%), or high quality (>80%).

Conversion of Outcomes to a Standard Metric

For US-based studies, we converted reported costs and incremental cost-effectiveness ratios into 2016 US dollars, using the medical care component of the Consumer Price Index. For other countries, we inflated data to 2016 using the country-specific Consumer Price Index²³ and converted the result to US dollars using relevant exchange rates.

RESULTS

Search Results

We retrieved 556 unique citations and screened all titles and abstracts, as well as full texts of 21 potentially relevant study reports. We excluded 7 studies after a full text review because

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they did not consider both cost and health outcomes, conducted burden-of-illness analyses, or did not report data with their analytical frameworks. A total of 14 eligible studies remained (Figure 1). No additional studies were found after we reviewed references of included studies and searched the NHS database.

Study Characteristics

Of the 14 studies reviewed, 13 were conducted in high-income countries within the past 5 years (Table 1). Federal or local governments sponsored 6 studies, and the pharmaceutical industry funded 5 studies. Furthermore, 7 studies evaluated treatments for initial CDI, 2 of which focused solely on severe infection. In addition, 4 studies considered treatments for recurrent CDI, while 2 others investigated both initial and recurrent CDI. The final study evaluated *C. difficile*–induced colitis unresponsive to metronidazole. All available treatment modalities approved for patient use were evaluated for initial and recurrent CDI, irrespective of guideline recommendation. Notably, 1 study examined FMT use for initial CDI,¹⁵ and 2 others investigated metronidazole use for recurrent CDI.^{13,14} Fidaxomicin was evaluated in 10 studies, while vancomycin was examined in all studies.

Model Design

Overall, 13 studies employed either a Markov or decision-tree model. The common Markov cycle length was 10 days (Table 1). Also, 2 studies used the same model to evaluate CDI treatments in different patient populations.^{24,25} The analytical perspective was that of the healthcare provider/health system or third-party payer for most studies (k = 12). Discounting was not applied for most studies because of the short time horizon. Furthermore, 2 studies that followed patients throughout their lives used appropriate discounting rates,^{14,26} but 1 of these studies had discordant time frames for cost and quality-adjusted life years (QALY; 18 weeks for costs vs lifetime for QALY).¹⁴ Comorbidities (eg, cancer, concomitant antibiotics, renal impairment) were accounted for in 3 studies.^{17,24,27}

Study Quality

Most of the studies were deemed moderate-to-high or high quality (k = 13). The mean and median quality scores were ~ 80% (data not shown but available upon request). Most studies provided detailed information on study design and population. In 1 study, the analytical perspective was societal, but indirect costs were not included.¹³ Another study did not specify the perspective,²⁶ and 2 studies lacked information on cost year.^{28,29}

Health Outcomes

Quality-adjusted life years was the most common health outcome reported (Table 1). Other outcome measures were CDI cases/recurrences avoided, clinical cure, life years, or bed days saved. Of the 10 studies that estimated QALY, 8 specified a cost-effectiveness decision threshold, but none conducted primary data collection for utility measurement. Because of the lack of CDI-specific utility weights,^{14–17,27} alternative weights for noninfectious diarrhea or for grade 3–4 diarrhea associated with chemotherapy were used. Utility weights generally varied substantially across studies; for example, utility for CDI was between 0.319 and 0.880.^{14,17,24–27}

Treatment Effectiveness

Table 2 shows how reviewed studies differed on treatment effectiveness across CDI episode and severity. Studies used a range of probabilities (0.65–0.84) as the metronidazole cure rate. Perras et al³⁰ used the lowest value (0.65) based on the success rate of metronidazole for initial severe CDI reported in a conference proceeding.³⁰ In contrast, Varier et al¹⁵ used a higher cure rate of 0.80 based on 1997 American College of Gastroenterology guidelines. Bartsch et al¹² derived the highest rate from a randomized clinical trial (RCT) and assumed it to be the same for both initial and recurrent CDI.

The studies that compared vancomycin and metronidazole generally used higher cure rate estimates for vancomycin, from 0.817 to 0.916.^{13,15,30,31} These rates were, however, lower than that of fidaxomicin, except for severe CDI (NAP1/BI/027 strain) or patients with renal impairment.^{17,27} For recurrent CDI, Varier et al used a vancomycin cure rate of 0.69,¹⁶ which was lower than the 0.889–0.926 range used in other studies.^{13,17,29} Furthermore, 2 studies assumed that vancomycin and fidaxomicin were similarly effective,^{25,29} and in 1 study, both drugs had much lower cure rates for *C. difficile*–induced colitis.²⁸ The fidaxomicin cure rates for the NAP1/BI/027 strain were considerably different in 2 studies, ^{12,27} whereas the cure rate of FMT was high (0.910–0.945) when delivered via colonoscopy but not other modes.¹³

Similarly, the probability of CDI recurrence after treatment varied significantly across studies. Recurrence rates after treatment with metronidazole ranged from 0.150 to 0.421 and were higher for recurrent CDI than for initial CDI. The CDI recurrence rate after vancomycin was lower than after metronidazole but higher than after fidaxomicin. While 2 studies modeled vancomycin with a higher recurrence rate for the NAP1/BI/027 strain than fidaxomicin,^{12,27,29} another study did the opposite.²⁷ The probability of recurrence after FMT via colonoscopy was comparable among studies but differed noticeably for other modes of delivery. Specifically, the recurrence rate of FMT by duodenal infusion or enema was 2–4 times higher in a study than in another, although the same reference source was cited in both.^{13,14} In some studies, recurrence rates were not stated explicitly.²⁶

Economic Parameters

Costs of CDI therapies and hospitalizations were included in all studies. Costs of laboratory tests were included in most studies, and costs of outpatient visits were included much less often (Table 2). Although excluding costs of treatment-related adverse events would bias results, only 3 studies included such costs.^{15,16,31} Most studies used official sources for cost estimates, and US studies had higher per-unit costs than studies in other countries. The cost of FMT therapy varied depending on the route of administration and often included associated pretreatment cost of oral vancomycin.

Cost-Effectiveness of CDI treatments

Table 2 summarizes the incremental cost-effectiveness ratios in 2016 US dollars per QALY gained stratified by type of CDI, wherever available. For initial CDI with no specific disease severity, fidaxomicin was cost-effective compared to vancomycin in 2 studies^{17,27} but not in the study accounting for severity.¹² For initial CDI in patients with concomitant antibiotics

use, cancer, or renal impairment, 2 studies found fidaxomicin to be cost-effective.^{17,24} Although FMT has not been recommended for initial CDI, the study that examined the use of colonoscopy-delivered FMT found it not cost-effective.¹⁵ Also, 2 studies found fidaxomicin cost-effective for severe initial CDI,^{17,25} but another concluded differently.²⁷ While many factors might have influenced results, a much higher cure rate of vancomycin (0.886) and the double cost for fidaxomicin,²⁷ compared with the other 2 studies, were notable. For recurrent CDI, studies consistently reported that FMT via colonoscopy was a cost-effective treatment, whereas findings on other FMT delivery routes were inconsistent. ^{13,14,16,26} When FMT was not available, fidaxomicin was a cost-effective option compared to other drugs in 3 studies^{14,17,25} but not in 2 other studies.^{12,13}

Stratified by funding source, all 5 industry-funded studies examined fidaxomicin, 3 of which concluded that fidaxomicin was either cost-effective or cost saving compared to metronidazole or vancomycin.^{17,24,25} The remaining 2 studies did not measure QALYs and made no conclusion about its cost-effectiveness.^{29,31} For studies with other types of or no funding, fidaxomicin was found cost-effective in one study²⁷ but not the other,¹² whereas FMT was favored in most of them.^{13,14,16,26}

Sensitivity Analysis

Most studies reported that treatment effectiveness was an important factor in 1-way sensitivity analysis (Table 2). For example, if the cure rate after vancomycin was >95.5%, it would be the preferred treatment for recurrent CDL.¹³ Cost of therapy was another influential parameter; FMT would no longer be dominant if its cost was > $3,205^{16}$ or if the fidaxomicin cost was < 1,359.¹³ Some other important variables were treatment duration, complication rates, and CDI mortality rate.

Probabilistic sensitivity analysis was conducted in 79% of the studies, but final results were not reported in 2 of them.^{12,13} Some studies presented a cost-effectiveness acceptability curve, while others reported 95% CI around the mean cost and effectiveness. The probability of being cost-effective at a pre-specified willingness to pay, defined as the maximum amount of dollars spent for an additional QALY gained, was between 60% and 96% for fidaxomicin, depending on CDI severity and population.^{24,25,27} FMT was either dominant or had a probability of cost-effectiveness between 38% and 87%.^{14,15}

DISCUSSION

Our study is one of the first systematic reviews to critically assess the quality of studies and cost-effectiveness of CDI treatment modalities, and we found substantial differences among the included studies. Because fidaxomicin is a newer drug, it was examined extensively for use in treating initial CDI. Results for fidaxomicin were inconclusive, however, except being cost-effective in some special and/or selective populations. The 3 studies of fidaxomicin for severe, initial CDI treatment had divergent conclusions, as did the 5 studies of fidaxomicin for recurrent CDI. FMT by colonoscopy was cost-effective for recurrent infection, but not for initial CDI. These cost-effectiveness findings did not hold true when FMT was delivered by other routes.

We identified important differences in study design among the included studies. Although QALY has become the most common outcome measure, one-third of the studies reviewed did not estimate QALY. Furthermore, studies accounted for CDI complications differently, and while some included costs of treating adverse events, none accounted for complications such as renal failure, which might bias the results in either direction. Another source of divergence was differences in healthcare resource utilization and costs among different settings. In particular, assumptions about treatment effectiveness contributed significantly to the diverging results. Two randomized controlled trials examined fidaxomicin.^{32,33} Both trials were conducted by OPT-80-003 Clinical Study Group investigators, and although the times and settings differed, they reported comparable cure and recurrence rates. These studies excluded patients with > 1 CDI occurrence within 3 months before studies started, and only 16% of enrolled patients had 1 previous CDI. Therefore, it is possible that the results applied to patients with initial CDI and not to those with recurrence. To date, there has been no published RCT on fidaxomicin effectiveness in recurrent CDI. Similarly, 2 other RCTs investigated the efficacy and safety of FMT in patients with recurrent but not initial CDI.^{34,35} and there were no RCTs comparing delivery routes when conducting this systematic review. Therefore, any study that examined fidaxomicin for recurrent CDI or FMT for initial CDI or compared delivery routes for FMT would have assumed their effectiveness or used data sources other than the available RCTs.^{36–38} Previous studies showed that comorbidities (eg, cancer, inflammatory bowel disease, and surgical burden) were strongly associated with increased risks for development and recurrence of CDI.³⁹⁻⁴³ However, most included studies did not account for such comorbidities in their models, which potentially biased the results. Lastly, studies modeled various numbers of recurrences following the initial episode, which might be another reason the results differed.

Our study has several limitations. Although we searched a wide range of databases, we may have missed some unpublished studies. In addition, because these studies differed in terms of study design, target population, model structure and input, our conclusions on the cost-effectiveness of CDI treatments were speculative. Finally, because we included industry-sponsored studies, which tend to be published only when results are favorable,⁴⁴ our synthesis and interpretation of results might be biased toward positive findings.

Our review has highlighted certain areas that could be improved in future CDI costeffectiveness analyses. While some of the models followed patients in the short term, those examining the long-term impact would present a more comprehensive assessment of interventions. Because there has been no widely accepted decision threshold for costeffectiveness using effectiveness measures other than QALY, future studies should preferably estimate QALY change to facilitate comparison. The cost-effectiveness of fidaxomicin compared with other pharmacologic therapies was not definitive for either initial or recurrent CDI, and different studies have used different values for its effectiveness. Therefore, future research might include a comprehensive literature review and provide rationale for choosing specific effectiveness values. A wide range for effectiveness and threshold analyses could also help understand the impact of fidaxomicin in various treatment scenarios. More prospective studies are needed to establish the efficacy and safety of fidaxomicin for recurrent CDI. There is also an urgent need for specific CDI utility weights that consider different complications, other comorbidities, or infection/severity stages. Given

that a validated instrument for CDI-specific, health-related, quality-of-life assessment is now available,⁴⁵ future research on utility weights will facilitate a more precise estimate of QALY change across CDI treatments.

In conclusion, CDI is a complex condition with a high recurrence rate, resulting in a significant burden of morbidity and mortality, as well as economic costs. Metronidazole and vancomycin have long been standard CDI treatment, but they are often associated with high rates of recurrence. New medications, such as fidaxomicin, and novel treatment modalities, such as FMT, have opened a new arena in CDI management. Because new treatments often come with a high cost, cost-effectiveness analyses are important to aid clinicians in rational decision making and health policy makers. Our review has identified an important divergence in research findings, especially in cost-effectiveness of fidaxomicin for either initial or recurrent CDI, which arose from discrepancies in model design and methods. Finally, our review informs future research of areas that need improvement and may help policymakers and physicians to critically assess the cost-effectiveness of different CDI treatments.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Characteristics of Ir	icluded Econor	mic Evaluations					
				Model Design			
Author (Year) Country Funding	Comparisons	Population	Model Type No. of CDI recurrences	Perspective	Time Horizon Discount Rate	Decision Threshold, US\$/QALY	Health Outcomes
Initial CDI (no specific	disease severity)						
Gidengil ³¹ (2014) Unired States Industry	 Metronidazole, then metronidazole for 1st recurrence Metronidazole, then vancomycin for 1st recurrence Vancomycin for list recurrence Fidaxomicin, then 	Adult inpatients	Markov (no cycle length) 0	Healthcare provider/health system	• None None	NR.	 No. of CDI recurrences No. of persistent equiring tx change No. of readmissions No. of CDI-readmissions No. of CDI-readmissions No. of CDI-readmissions No. of VRE colonization cases No. of VRE
Rubio-Terres ²⁴ (2015) Spain Industry	FidaxomicinVancomycin	Pts with cancer, concomitant antibiotic tx, renal impairment	• Markov (10-d cycle) • 1	Healthcare provider/health system	• 1 y • None	\$31,800 (€0,000)	QALY
Stranges ²⁷ (2013) United States None	FidaxomicinVancomycin	Mean age, 59.9y	Decision tree2	Third-party payer	• 23 y • 3%	\$100,000	QALY
Varier ¹⁵ (2014) United States NGO	• Metronidazole • Vancomycin • FMT colonoscopy	Adult outpatients	• Decision tree • 2	Third-party payer	• 90 d • None	\$100 000	QALY
Watt ¹⁷ (2016) Germany Industry	Fidaxomicin Vancomycin	Pts with severe initial CDI, recurrent CDI, concomitant antibiotic tx, age 65 y, cancer, renal impairment	• Markov (10-d cycle) • 2	Healthcare provider/health system	• I year • None	\$63 000 (£0,000)	QALY No. of bed days saved No. of CDI recurrences avoided
Initial CDI (severe)							

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TABLE 1

				Model Design			
Author (Year) Country Funding	Tomnomicone	Donulotion	Model Type No. of CDI	Docensolities	Time Horizon Discount Data	Doricion Thusdaild 1164/0AIV	Hoolth Outcomos
Perras ³⁰ (2011)	- Comparisons	Pts with severe	Decision tree	Healthcare provider/health system	• 50 d	DECISION LINESHORD, USANCALI NR	Clinical cure
Canada Public	Metronidazole • Vancomycin • Fidaxomicin	CDI	·		• None		
Wagner ²⁹ (2014) Canada Industry	Vancomycin	Pts with severe CDI	• Decision tree • 1	Healthcare provider/health system	• 2 months • None	NR	No. of CDI recurrences avoided
Recurrent CDI							
Konijeti ¹³ (2014) United States Public	 Metronidazole Vancomycin Fidaxomicin FMT FMT Muodenal infusion FMT enema 	Pts age 65 y	• Decision tree • 2	Societal	• 1 y • None	\$50,000	QALY
Lapointe-Shaw ¹⁴ (2011) Canada Public	 Metronidazole and vancomycin for vancomycin for recurrences Vancomycin for vancomycin for vancomyci for vancomycin for vancomycin for vancomycin for vancomycin	Community- dwelling Porsons, age 70y	• Markov (6-week cycle) • 2	Healthcare provider/health system	 18 weeks (CDI-related costs and costs and costs); lifetime (QALYs) 5% (health benefits); none (costs) 	\$38,000 (CAD 50,000)	QALY

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				Model Design			
Author (Year) Country Funding	Comparisons	Population	Model Type No. of CDI recurrences	Perspective	Time Horizon Discount Rate	Decision Threshold, US\$/QALY	Health Outcomes
	• FMT by colonoscopy and repeated therapy using a different donor for recurrence						
Merlo ²⁶ (2016) Australia None	 Vancomycin Nasoduodenal FMT Colorectal FMT 	Age 65y	• Markov (10-d cycle) • 2	NR	• Lifetime • 5%	NR	• QALY • No. of life years saved
Varier ¹⁶ (2015) United States Public	Vancomycin FMT colonoscopy	Outpatient adults	• Decision tree • 1	Third-party payer	• 90d • None	NR	QALY
Initial and Kecurrert (-DI						
Bartsch ¹² (2013) United States Public	 Metronidazole (nonsevere) and vancomycin (severe) Fidaxomicin with strain typing 	Adults, age 18 y	• Microsimulation • 1	Third-party payer	• None • None (3% for cost conversion)	\$50,000	QALY
Nathwani ²⁵ (2014) Scotland Industry	FidaxomicinVancomycin	Adults, age 18 y	• Markov (10-d cycle) • 2	Governmental	• 1 year • None	\$25,400 (£20,000) \$38,100 (£30,000)	QALY
Other							
Markovic ²⁸ (2014) Serbia Public	FidaxomicinVancomycin	NR	• Markov (15-d cycle) • 2	Third-party payer	• 90 d • None	\$458,440/life saved (RSD 53,307,040/life saved)	 No. of lives saved No. of subtotal colectomies avoided
NOTE. CDI, <i>Clostridium</i> years; tx, treatment; VRE,	<i>difficile</i> infection; d, vancomycin-resista	, day; FMT, fecal mic nt enterococci, venou	robiota transplantation s thromboembolism; y	; NGO, nongovernmental organizatio ; years.	m; NR, not reported	or not available; pt, patients; QALY,	quality-adjusted life

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Author (Year)	Eff	ectiveness	Cost E	Istimate			
Country Funding	Probability of Cure	Probability of Recurrence	Types Source Original Cost Year	Treatment, 2016 US\$	Incremental Cost- Effectiveness Ratio, 2016 US\$\QALY (Original Values)	Scenario Analysis, 2016 US\$/ QALY (Original Values)	Influential Variables
Initial CDI (no specific s	severity)						
Gidengil ³¹ (2014) United States Industry ^a	MET: 0.735 VAN: 0.817 FID: 0.841	MET Initial episode: 0.267 1 st recurrence: 0.330 2 recurrences: 0.308 VAN Initial episode: 0.253 1 recurrence: 0.355 2 recurrences: 0.308 FID Initial episode: 0.154 1 st recurrence: 0.197 2 recurrences: 0.308	 Hospitalization, tx, lab test, outpatient visit Manufacturer cost, expert interviews, litterviews, 2010 	MET: \$27 VAN: \$1,255 FID: \$3,316	NR	NR	 Recurrence probability for initial episode with MET and VAN tx All VRE clinically related probabilities Recurrence probability (beyond the 1st 2) with VAN tx Cost of an invasive VRE infection Cost of FID PSA: yes
Rubio-Terres ²⁴ (2015) Spain Industry			 Hospitalization, tx, outpt visit Spanish public healthcare prices, literature 2013 	VAN: \$40 FID: \$1,577	FID dominant	NR	 Duration of excess stay attributable to CDI, initial CDI or recurrent CDI PSA: yes
Stranges ²⁷ (2013) USA None	VAN Inpatient: 0.781 Outpatient: 0.975 Mild-to-moderate CDI: 0.839 Severe CDI: 0.886 Concomitant antimicrobials: 0.794 NAPI/BI/027: 0.807 FID Inpatient: 0.814 Outpatient: 0.814 Outpatient: 0.814 Outpatient: 0.814 Outpatient: 0.975 Mild-to-moderate CDI: 0.920 Severe CDI: 0.920 Severe CDI: 0.920 Severe CDI: 0.921 Ocoromitant antimicrobials: 0.900	VAN Inpatient: 0.274 Outpatient: 0.227 With previous CDI Wild-to-moderate CDI: 0.244 Severe CDI: 0.266 Concomitant NAPI/BI/027: 0.209 FID Inpatient: 0.179 Outpatient: 0.179 Outpatient: 0.179 Outpatient: 0.179 Outpatient: 0.128 With previous CDI episode: 0.214 Mild-to-moderate CDI: 0.168	Hospitalization, tx, lab test HCUP • 2011	VAN: \$1,335 FID: \$3,218	FID: \$77,661 (\$67,576)	ICER of FID • Severe CDI: \$405,676 (\$532,994) • Mild-to- moderate CDI: \$36,799 (\$32,020) (\$336,771) • Initial tx as • Initial tx as • Initial tx as • Sev.321 • Sav.321 • Sav.322 • S	 Recurrence rate of FID Recurrence rate of VAN VaN First episode inpatient FID cure rate of Rate of Probability of CDI mortality PSA: yes

Effectiveness, Costs, Incremental Cost-Effectiveness Ratio, and Sensitivity Analyses of Included Economic Evaluations

TABLE 2

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Author (Year)	EU	Tectiveness	Cost H	Estimate			
Country Funding	Probability of Cure	Prohability of Recurrence	Types Source Original Cost Year	Treatment, 2016 US\$	Incremental Cost- Effectiveness Ratio, 2016 US\$/QALY (Original Values)	Scenario Analysis, 2016 US\$/ QALY Values)	Influential Variables
	NAP1/BI/027: 0.787	Concomitant antimicrobials: 0.169 NAPI/BI/027: 0.271	0		(mar.	• Concomitant antimicrobials: \$1,709 (\$1,487)	
Varier ¹⁵ (2014) United States NGO ^C	MET: 0.800 VAN: 0.900 FMT colonoscopy: 0.910	MET: 0.168 VAN: 0.084 FMT colonoscopy: 0.076	 Hospitalization, cost of tx Medicare, literature 2011 	FMT: \$1,248 MET: \$66 VAN: \$1,548	FMT colonoscopy: \$143,614 (\$124,964)	NR	 Cure rate of MET Cost of FMT Cost of MET PSA: yes
Watt ¹⁷ (2016) Germany Industry	VAN I recurrences: 0.889 Severe CDI: 0.826 Concornitant antibiotics: 0.755 Age 65 y: 0.845 Cancer: 0.740 Renal impairment: 0.760 I recurrence: 0.898 Severe CDI: 0.800 Conconitant antibiotics: 0.843 Age 65 y: 0.848 Cancer: 0.851 Renal impairment: 0.731 Renal impairment:	VAN 9 1 recurrence: 0.325 Severe CDI: 0.283 Concomitant antibiotics: 0.255 Age 65 y: 0.293 Cancer: 0.296 Renal impairment: 0.316 FID 1 recurrence: 0.203 Sever CDI: 0.114 Concomitant antibiotics: 0.174 Age 65 y: 0.161 Cancer: 0.135 Renal impairment: 0.147	 Hospitalization, tx German drug tariff 2014 	VAN: \$69 FID: \$1,564	ICER of FID • Pts with 1 recurrence: \$49,482/QALY (\notin 3,900) • Pts with severe CDI: \$39,255/QALY (\notin 4,800) • Pts with concomitant antibiotics: \$34,603/QALY (\notin 0,700) • Age 65 y \$50,159/QALY (\notin 4,500) • Age 65 y \$50,159/QALY (\notin 4,500) • Pts with cancer: dominant • Pts with cancer: dominant	Z	 Recurrence rate Clinical cure rate CDI-attributable mortality rate PSA: no
Initial CDI (severe)							
Perras ³⁰ (2011) Canada Public ^d	MET: 0.649 VAN: 0.849	MET: 0.150 VAN: 0.150	 Hospitalization, tx, lab test, outpatient visit Provincial drug formularies 2010 	MET: \$3.10 VAN: \$279	Х	N	 Infection populations (with or without NAP1 strain), efficacy rates in initial therapy with MET Cost of VAN Complication rate among tx failures PSA: yes
Wagner ²⁹ (2014) Canada Industry ^e	VAN Severe CDI: 0.813 With recurrence: 0.926	VAN Severe CDI: 0.283 With recurrence: 0.323 NAP1/BI/027: 0.282 Non-NAP1/BI/027: 0.278	 Hospitalization, tx, outpt visit Canadian Agency for Drugs and Technologies in 	NR	NR	NR	 Recurrence rate of FID Duration of tx, 10-14 d PSA: no

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Author (Year)	Eff	fectiveness	Cost I	Estimate			
Country Funding	Probability of Cure	Probability of Recurrence	Types Source Original Cost Year	Treatment, 2016 US\$	Incremental Cost- Effectiveness Ratio, 2016 US\$/QALY (Original Values)	Scenario Analysis, 2016 US\$/ QALY (Original Values)	Influential Variables
	NAP1/BL/027: 0.813 0.813 Non-NAPI/BL/027: 0.912 FUD Severe CDI: 0.813 With recurrence: 0.926 NAP1/BL/027: 0.813 0.813 Non-NAP1/BL/027: 0.912	FID Severe CDI: 0.114 With recurrence: 0.203 NAP1/JBI/027: 0.248 Non-NAP1/JBI/027: 0.097	Health, Ontario Case Costing Initiative, Ontario Schedule for Physician Services • NR				
Recurrent CDI							
Konijeu ¹³ (2014) United States Public	MET: 0.710 VAN: 0.916 FID: 0.937 FMT colonoscopy: 0.945 FMT duodenal infusion: 0.813 FMT enema: 0.815	MET: 0.421 VAN: 0.355 FID: 0.197 FMT duodenal infusion: 0.063 FMT enema: 0.091	 Hospitalization, tx, outpt visit Clinical diagnostic laboratory fee schedule from Centers for Medicare and Medicare and Medicare and iterature 2012 	MET: \$25 VAN: \$754 FID: \$3,104 FMT colonoscopy: \$2,495 FMT duodenal infusion: \$2,386 FMT enema: \$2,048	FMT colonoscopy: \$18,865/ QALY (\$17,016)	FMT duodenal available • TMT • TMT • 5MT • 5MT • 5107,9277 • 352); • 7D2; • 7D2; • 7D2; • 7D2; • 7D2; • 709/ • 700/ • 700	 Cure and recurrence rate of oupt oral VAN, FMT VAN, FMT colonoscopy colonoscopy, FID, outpt oral VAN Probability of severe CDI if tx failure PSA: no

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	Scenario Analysis, 2016 US\$/ (Original Values) Influential Variables	ant • Age 10 y: • Probability of FMT recurrence following colonoscopy fecal transplantation dominant by enema • FID off- patent: FMT colonoscopy dominant	o FMI available • FID \$20 757/ QALY (CAD 25,968) • No FMT colonoscopy: FMT by enema \$1,365/ QALY (CAD 1,708) 2 recurrences considered • FMT colonoscopy: \$411/QALY (CAD514)	No FMI available • FID \$20757/ QALY (CAD 25,968) • No FMT colonoscopy: FMT by colonoscopy: FMT by considered • FMT • FMT • FMT • CAD514) NR • PSA: yes
I	Incremental Cost- Effectiveness Ratio, 2016 US\$/QALY (Original \$ Values)	FMT colonoscopy dominan		Nasoduodenal FMT and colorectal FMT dominant
st Estimate	ar Treatment, 2016 US\$	MET: \$31.20 VAN: \$278 FID: \$1,923 FMT enema: \$6,504 h FMT nasogastric: \$1,040 FMT colonoscopy: \$4,083		VAN: \$494 Colorectal FMT: \$1,688 Nasoduodenal FMT: \$1,637
Cos	Types Source Coriginal Cost Yea	 Hospitalization, tx, lab test, outpt visit, capital cost (equipment) University Health Network outpt database, Ontario Schedule of Benefits 2014 		 Hospitalization, tx, lab test tx, lab test National databases, market pharmaceutical benefits schedule. Queensland health wage rate 2015
fectiveness	Probability of Recurren	MET: 0.400 VAN oral: 0.517 VAN pulse/taper: 0.178 FDD: 0.321 FMT colonoscopy: 0.078 FMT duodenal infusion: 0.233 FMT enema: 0.185		
ER	Probability of Cure	MET: 0.776		VAN: 0.308 Colorectal FMT: 0.939 Nasoduodenal FMT: 0.939
Author (Year) Country Funding		Lapointe-Shaw ¹⁴ (2011) Canada Public ^f		Merlo ²⁶ (2016) Australia None <i>ĥ.g</i>

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Initial and recurrent CDI

Country Funding	Probability of Cure	Probability of Recurrence	Types Source Original Cost Year	Treatment, 2016 US\$	Incremental Cost- Effectiveness Ratio, 2016 US\$/QALY (Original Values)	Scenario Analysis, 2016 US\$/ QALY (Original Values)	Influential Variables
Bartsch ¹² (2013) United States Public	MET: 0.835 VAN NAP1/BI/027: 0.820 Non-NAPI/BI/027: 0.897 FID NAP1/BI/027: 0.859 Non-NAP1/BI/027: 0.926	MET: 0.136 VAN NAPI/BI/027: 0.295 Non-NAPI/BI/027: 0.278 FID NAPI/BI/027: 0.247 Non-NAPI/BI/027: 0.098	• Hospitalization, tx, lab test • HCUP, Redbook • 2012	MET: \$65 VAN: \$1,144 FID: \$3,725	NR, "No FID" was best	NR	 Proportion of C. diffinfections caused by NAP1 strain versus other strains Cost of FID To guidelines recommends VAN as the first-line tx PSA: yes
Nathwani ²⁵ (2014) Scotland Industry ^{C.e}	VAN Severe CDI: 0.853 Recurrence: 0.889 FID Severe CDI: 0.853 Recurrence: 0.889	VAN Severe CDI: 0.267 Recurrence: 0.325 FUD Severe CDI: 0.172 Recurrence: 0.172	 Hospitalization, tx, outpatient visit British National Formulary 2011 	VAN: \$312 FID: \$1,568	For severe CDI: FID \$27 225/QALY (£16 529) For recurrence CDI: FID dominant	X	 OR of experiencing a recurrence with FID in pis who had already experienced > 1 recurrence OR of experiencing a recurrence with FID in pis with severe CDI OR of having recurrent CDI if reverte CDI Probability of a recurrence being treated in hospital PSA: yes
Other							
Markovic ²⁸ (2014) Serbia Public	VAN: 0.652 FID: 0.741	VAN: 0.221 FID: 0.130	 Hospitalization, tx, lab test, surgery Literature NR 	NR	NR	NR	• Cost of FID • PSA: no
NOTE. C. diff, Clostridiun	n difficile; CDI, Clostridi	<i>ium difficile</i> infection; d, day; F	ID, fidaxomicin; FMT, 1	ècal microbiota transplan	tation; HCUP, Healthcare Cost	and Utilization Pr	oject; ICER,

incremental cost-effectiveness ratio; MET, metronidazole; NGO, nongovernmental organization; NR, not reported or not available; outpt, outpatients; OR, odds ratio; PSA, probabilistic sensitivity analysis; pt, patients; QALY, quality-adjusted life-years; tx, treatment; VAN, vancomycin; VRE, vancomycin-resistant enterococci, venous thromboembolism; y, years.

 a Probabilities of cure estimated from published reports.

 b_{III} is the second se

 $\boldsymbol{c}^{}$ Probabilities of recurrence estimated from published reports.

 $\boldsymbol{d}_{\text{Probabilities}}$ of recurrence assumed to be similar for both treatments.

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Cost Estimate

Effectiveness

Author (Year)

 $f_{\mbox{Probability}}$ of cure not listed for other treatments in the report.

 $\ensuremath{\mathcal{E}}$ Probability of recurrence not available from the published report.

hThe vancomycin probability of cure for 10-d cycles. The FMT probability of cure assumed to be the same regardless of delivery modes.