Impaired Quality of Life, Work, and Activities Among Adults with \textit{Clostridium difficile} Infection: A Multinational Survey

Kirstin Heinrich$^{1}$ · James Harnett$^{2}$ · Jeffrey Vietri$^{1}$ · Richard Chambers$^{1}$ · Holly Yu$^{1}$ · Marya Zilberberg$^{3,4}$

Received: 28 March 2018 / Accepted: 20 July 2018 / Published online: 3 August 2018
© The Author(s) 2018

Abstract

Background Evidence of humanistic detriments of \textit{Clostridium difficile} infection (CDI) remains limited.

Aims To assess humanistic burden associated with CDI.

Methods Self-reported National Health and Wellness Survey data between 2013 and 2016 were analyzed for the USA, five European countries, China, and Brazil. Outcome measures included SF-36v2® for health-related quality of life (HRQoL) and Work Productivity and Activity Impairment questionnaire. Respondents (≥ 18 years old) were classified as (1) currently treated doctor-diagnosed CDI (C-CDI), (2) doctor-diagnosed prior CDI (P-CDI), or (3) never experienced CDI (NO-CDI). Regression modeling assessed the association between CDI status and outcomes, adjusting for potential confounders.

Results Of 352,780 respondents, 299, 2111, and 350,370 met the criteria for C-CDI, P-CDI, and NO-CDI, respectively, with 45% of the total from the USA. C-CDI and P-CDI respondents were older, were less often employed and had more comorbidities than those with NO-CDI. After adjustment for covariates, C-CDI and P-CDI had significantly lower HRQoL relative to NO-CDI for mental (MCS 39, 43 vs. 46) and physical (PCS 39, 41 vs. 46) component summary scores, and health utility (SF-6D 0.58, 0.64 vs. 0.71) (all $p < 0.05$), meeting common thresholds for minimally important differences. Those with C-CDI and P-CDI reported missing more work (21, 16 vs. 8%), greater impairment while working (43, 34 vs. 22%), and more activity impairment (61, 49 vs. 34%) than those with NO-CDI (all $p < 0.05$), respectively.

Conclusions CDI is associated with meaningfully worse HRQoL and greater impairment to work and activities compared with NO-CDI. The impairment directly attributable to CDI requires further evaluation.

Keywords \textit{Clostridium difficile} · Quality of life · Work productivity · Outcomes · Questionnaire · Survey

Introduction

\textit{Clostridium difficile} is the most common cause of infectious diarrhea in hospitalized patients and the leading cause of healthcare-associated infections in hospitals [1–3]. A population study estimates that the annual volume of \textit{Clostridium difficile} infection (CDI) in the USA tops 450,000 cases [2]. Due to this statistic, the United States of America (US) Centers for Disease Control and Prevention has classified CDI as an urgent public health threat [4]. In addition to healthcare-acquired CDI, community-acquired CDI is increasingly a concern as rates increase [5–7]. Though lower in incidence than in the USA, CDI remains a large and growing concern in both Europe and Asia; however, published estimates may undercount the actual burden given the lack of testing, underdiagnoses, and potentially missed community-acquired cases [8–12].
Numerous studies have documented clinical and economic consequences of CDI. Patients with CDI have longer lengths of hospital stay [13–17], are more frequently readmitted to the hospital, and are less likely to be discharged directly home from the hospital [13, 18], and incur higher hospital costs [19, 20] than those without CDI. Nursing homes with CDI have been found to incur greater direct medical costs including hospital costs, nonhospital-based outpatient medical costs, and prescription costs [21] compared to those without CDI. The total economic impact is considerable, with CDI estimated to contribute nearly $5 billion in excess healthcare costs in the USA [19] and nearly €3 billion in Europe [22].

In contrast, few studies have examined the impact of CDI on health-related quality of life (HRQoL). Some of the studies published to date that demonstrate potentially serious negative HRQoL consequences of CDI [23–25] present multiple methodological challenges that may limit their interpretation. To address this evidence gap, we assessed the HRQoL, work productivity, and activity impacts of CDI using a large international survey.

**Methods**

**Data Source**

We used data from the National Health and Wellness Survey (NHWS; Kantar Health, New York, NY, USA), a cross-sectional, primarily Internet-based commercial survey of adults (aged 18 years and older) fielded annually in the USA and periodically in nine other countries. The majority of potential respondents to NHWS are identified through commercial survey panels (e.g., MySurvey.com and Lightspeed GMI), which recruit their members through convenience methods such as targeted advertisements, e-newsletters, and online banner placements. The NHWS sample is derived from these panels through random sampling stratified by key characteristics to attempt a sample derivation that is representative of the adult population of each participating country. These include age and sex for all countries with the addition of white/nonwhite race in the USA. Online recruitment is supplemented by offline recruitment for certain age-groups in countries with lower Internet penetration among the older population (Brazil, China; 2013 France, Germany, Italy, Spain; 2016 Germany, Italy). The 2013 surveys received Institutional Review Board (IRB) approval from the Essex IRB (Lebanon, New Jersey, USA). The 2015 and 2016 surveys were approved by the Pearl IRB (Indianapolis, Indiana, USA). All respondents provided informed consent prior to participating.

**Study Cohort**

The surveys used to create the cohort were from the USA (2013, 2015), Europe (EU5: UK, France, Germany, Italy, and Spain; 2013, 2016), Brazil (2015), and China (2013, 2015). Depending on the year and country of the survey, 4–16% of those who were invited to the survey completed the entire NHWS. These response rates were largely due to individuals not opening the invitation or not clicking on the survey link; most of those who clicked the link and completed the screening questions (61–79%, again depending on country and year) completed the survey. Because the NHWS is a cross-sectional survey and multiple years of data are included, a few respondents might have participated in multiple years. To avoid having these respondents unduly impact the results, only 1 year of data for such respondents were included in the analysis. For respondents who completed the survey twice, data of the first year were used if they reported a history of CDI in the first year. The data of the second (i.e., most recent) year of survey were used if the respondent reported a history of CDI in only the second year or if the respondent indicated no history of CDI in either year.

The cohort was divided into three groups according to CDI status based on self-reports: current CDI (C-CDI), prior CDI (P-CDI), and never experienced CDI (NO-CDI) (Fig. 1). To be categorized as C-CDI, the respondent had to say “yes” to all of the following: (1) having ever experienced “C. diff (antibiotic associated colitis which is diarrhea from antibiotic use),” (2) having a physician diagnosis of CDI, and (3) currently using prescription medication to treat CDI. P-CDI respondents reported all of the following: (1) ever experienced CDI, (2) a physician diagnosis for CDI, and (3) not currently using prescription medication to treat CDI. Treatment status was used in an attempt to differentiate respondents whose CDI had resolved versus those who were currently experiencing infection. NO-CDI respondents were limited to those who did not report ever experiencing CDI.

**Demographic and Health Characteristic Measures**

Demographic variables from the NHWS included age, education, gender, insurance status, employment status, and ethnicity (USA, Brazil, and China only). As response options for ethnicity, health insurance, and education vary among countries and across different years of the NHWS, these variables were harmonized to allow for inclusion in the analysis. Ethnicity was categorized into either majority or minority categories at the country level, with majority defined as the most frequently reported ethnicity within
that specific country, with all others categorized as minority. Insurance was dichotomized as either reporting having any kind of insurance (public and/or private), or reporting having no insurance of any kind. The education variable was categorized as the highest level of education completed based on the 2011 International Standard Classification of Education developed by the United Nations Educational, Scientific and Cultural Organization (UNESCO) as below upper secondary education, upper secondary or post-secondary nontertiary, tertiary, or decline to answer [26].

Patient health characteristics included the Charlson comorbidity index (CCI) calculated from self-reported comorbidities [27], antibiotic use, and healthcare resource utilization (HCRU). As the NHWS does not collect information on paraplegia or liver disease severity, the CCI used herein did not include these conditions. All HCRU variables used 6-month recall except in the case of surgery, where 12-month recall was required. The descriptive statistics for the number of HCRU visits were calculated to include respondents with no reported visits.

Outcomes Measures

The Medical Outcomes Study 36-Item Short-Form Health Survey version 2 (SF-36v2®) standard (4-week recall) form was used in the NHWS to measure HRQoL [28]. The SF-36v2® provides scores for 8 subscales (domains) which are comprised of mutually exclusive questions: general health (GH), social functioning (SF), vitality (VT), bodily pain (BP), mental health (MH), physical functioning (PF), role-emotional (RE) and role-physical (RP), which are further combined into two scores summarizing mental (mental component summary score [MCS]) and physical (physical component summary score [PCS]) health. Consistent with usual practice, the scores presented here are the norm-based scores, which are standardized so that each scale has a mean score of 50 and standard deviation of 10 in the US general population. Lower values indicate worse outcomes. The SF-36v2® also provides health utility scores with the SF-6D, a single preference-based measure of health using general population values.

Impairment to work and other activities was measured using the Work Productivity and Activity Impairment questionnaire, general health version (WPAI-GH), with a 7-day recall [29]. Outcomes calculated from responses to the WPAI-GH included degree of activity impairment due to health (activity impairment), degree of impairment while working due to health (presenteeism), percent of work missed due to health (absenteeism), and degree of overall work impairment due to health (overall work impairment, a combination of both presenteeism and absenteeism). Presenteeism, absenteeism, and overall work impairment assessments were restricted to respondents who reported part-time, full-time, or self-employment. Activity impairment was assessed among all respondents. Higher values indicate a greater degree of impairment.

Statistical Analysis

Unadjusted descriptive summary statistics were calculated for the aggregated cohort by CDI status for all demographic and outcome variables. Demographic and outcome variables were compared using the t test for continuous variables and the Chi-square test for categorical variables. Mean norm-based health domain scales were described and compared between CDI groups for the aggregated cohort using the equal or unequal variance t-test depending on the variance equality test result.

Regression modeling assessed the association between CDI status and outcomes after adjusting for CCI category (0, 1 or 2, 3 or more), education level, age category (18–49, 50–64, and ≥ 65 years), sex, and country. Estimates and standard errors for activity impairment, presenteeism, absenteeism, and overall work impairment were exponentiated and derived from a generalized linear model with negative binomial distribution with log link function. All HRQoL outcomes were assessed between groups using a general linear model with normal distribution.
For all comparisons, *p* values less than 0.05 were considered statistically significant. Minimally important differences (MIDs) were defined based on criteria from the developers of each measure. For the SF-36v2, MIDs are defined as differences in norm-based scores of at least 2 points for PCS, GH, and VT; 3 points for each of MCS, PF, BP, RP, SF, and MH; and 4 points for RE [28]. For SF-6D, MID was defined as the difference in average health utility of at least 0.041 [30].

All analyses were performed using Statistical Analysis System (SAS), version 9.4 (SAS Institute Inc., Cary, North Carolina, USA).

**Results**

Among the 352,780 respondents included in the analysis, 299 (0.08%) met the criteria for C-CDI and 2111 (0.60%) for P-CDI. A plurality of respondents (45%) were from the USA, 39% from the EU5, 13% from China, and 3% from Brazil (Table 1). Of those reporting C-CDI, nearly one-half were from the EU5 (48%), while the majority of the P-CDI respondents were in the USA (59%). Compared to those with NO-CDI and P-CDI, respondents with C-CDI were older, were less often employed, and had a higher mean CCI score (Table 1). Fewer people in the C-CDI and NO-CDI groups were female than in P-CDI (49, 53 vs. 67%; *p* values < 0.0001). Education levels differed significantly among groups, with P-CDI responders less likely than the other two groups to have tertiary education (Table 1). There were a significantly greater number of healthcare provider visits, emergency room visits, and hospitalizations among persons with C-CDI than among those with NO-CDI or P-CDI (Table 2). A greater proportion of the C-CDI group reported surgeries and at least one physician visit, with the mean number of healthcare encounters among those with at least one visit also significantly higher in C-CDI than in NO-CDI or P-CDI groups (Table 2).

Respondents with C-CDI and P-CDI reported significantly lower unadjusted mean scores across the eight SF-36 domains compared with those with NO-CDI (Fig. 2). The C-CDI group had significantly lower scores than the P-CDI group on all domains except vitality. Results of country-level unadjusted analyses mirrored the aggregated analysis (data not shown). Unadjusted differences in individual health domains between groups are shown in Fig. 2.

After adjustment for CCI scores, age, sex, education level, and country, C-CDI and P-CDI were associated with significantly lower (all *p* values <0.05) mean scores for MCS (39.5, 42.7 vs. 45.7), PCS (38.9, 40.9 vs. 45.5), and health utility (0.58, 0.64 vs. 0.71) vs. NO-CDI, respectively (Figs. 3 and 4). Persons with C-CDI had significantly lower mean scores on the MCS, PCS, and health utility versus those with P-CDI. C-CDI and P-CDI groups reported significantly greater percent of work time missed (20.8, 16.0 vs. 8.4%), degree of impairment while working (43.1, 33.7 vs. 22.2%), degree of overall work impairment (52.1, 39.3 vs. 26.0%), and degree of activity impairment (60.7, 49.0 vs. 34.4%) due to health than NO-CDI group, respectively (C-CDI or P-CDI vs. NO-CDI, all *p* values < 0.05; Fig. 5). There were no significant differences between C-CDI and P-CDI groups with respect to work productivity and activity impairment measures (Fig. 5).

**Discussion**

In this large and geographically diverse analysis of the impact of self-reported CDI on HRQoL, we have found that individuals either currently treated for CDI or with a past history but no current treatment for CDI have significantly worse HRQoL, greater impairment in their daily activities, and reduced work productivity than those with no history of CDI. These differences persist even after adjusting for age, sex, CCI scores, education, and country. In addition to worsened HRQoL, C-CDI respondents suffer diminished work productivity, with the rate of absenteeism 2.5 times higher among those with no history of CDI. The productivity loss among those attending work (i.e., presenteeism) associated with C-CDI is nearly double that of respondents with no CDI history. Additionally, after adjustment for covariates, differences between groups in PCS, MCS, and health utility met MID criteria (Figs. 3 and 4). Compared to NO-CDI respondents, C-CDI and P-CDI persons reported significantly lower unadjusted mean scores across the eight SF-36v2 domains meeting MID criteria (Fig. 2). When comparing P-CDI and C-CDI persons, MIDs persisted for all health domains except for VT and GH where the difference did not meet the thresholds for MID (Fig. 2).

One prior study has noted a substantial degree of impairment to HRQoL, work, and activities in CDI using the same instruments. In an analysis of the 2010 US NHWS, Peery and colleagues noted that patients who had ever experienced CDI had a mean MCS of 45, mean PCS of 39, mean degree of activity impairment of 49%, and mean degree of overall work impairment of 50% [24]. Although difficult to interpret without a comparator, this study’s results hinted at the magnitude of CDI-associated impairment. Our results build on Peery’s work by confirming the overall CDI-related decrements and further clarify the differential impacts of prior and current CDI on people in comparison with those who have never experienced this affliction. In addition, whereas Peery’s analysis did not attempt to deal with confounding, our adjusted analyses demonstrate persistent magnitude of CDI-associated HRQoL impairment.

© Springer
Table 1 Descriptive characteristics

| Country (n, %) | Never experienced CDI (NO-CDI) | Experienced doctor-diagnosed CDI | p value C-CDI versus P-CDI |
|---------------|--------------------------------|----------------------------------|---------------------------|
|               | n = 350,370                    | Not currently treated (P-CDI)    | Currently treated (C-CDI) |
|               | n = 2111                       | n = 299                          |                           |
| Brazil        | 11,933 3.4%                    | 12 0.6%                          | < .0001                   | 14 4.7%                       | 0.1243 < .0001 |
| China         | 45,833 13.1%                   | 16 0.8%                          | < .0001                   | 21 7.0%                       | 0.0019 < .0001 |
| USA           | 157,285 44.9%                  | 1245 58.9%                       | < .0001                   | 121 40.5%                     | 0.2239 < .0001 |
| EU            | 135,319 38.6%                  | 838 39.7%                        | 0.3118                    | 143 47.8%                     | 0.0011 0.0074 |
| Age [mean; median (SD)] | 45.9 45.0 (16.47) | 52.5 54.0 (15.97) | < .0001 | 48.4 48.0 (16.44) | 0.0122 < .0001 |
| Age group (n, %) | < .0001                     | < .0001                          |                           | < .0001 | 0.15 0.0002 |
| 18–49         | 203,673 58.1%                  | 887 42.0%                        | 164 54.9%                 | 68 22.7%                      | 0.0012 < .0001 |
| 50–64         | 83,364 23.8%                   | 622 29.5%                        | 67 22.4%                  |                           |                |
| ≥ 65          | 63,333 18.1%                   | 602 28.5%                        |                           |                |                |
| Employed (n, %) | 206,821 59.0%                | 909 43.1%                        | < .0001                   | 149 49.8%                     | 0.0012 < .0001 |
| Education level (n, %) | < .0001                   | 0.2663 0.0056                    |                           |                |                |
| Below upper secondary | 39,899 11.4%                 | 202 9.6%                         | 43 14.4%                  |                           |                |
| Upper secondary or post-secondary | 124,778 35.6%              | 893 42.3%                        | 103 34.5%                 |                           |                |
| Tertiary      | 184,407 52.6%                  | 1011 47.9%                       | 152 50.8%                 |                           |                |
| Female (n, %) | 183,939 52.5%                  | 1420 67.3%                       | < .0001                   | 146 48.8%                     | 0.2041 < .0001 |
| Majority ethnicitya (n, %) | 167,478 78.1%              | 1143 89.9%                       | < .0001                   | 134 85.9%                     | 0.0181 0.1289 |
| Health insurance status—insured (n, column %) | 307,005 87.6% | 1985 94.0% | < .0001 | 273 91.3% | 0.1316 0.1294 |
| Adapted CCI score [mean; median (SD)] | 0.2 0.0 (0.7) | 0.9 0.0 (1.6) | < .0001 | 2.6 1.0 (4.6) | < .0001 < .0001 |
| Adapted CCI score category (n, %) | < .0001                   | < .0001                          | < .0001                   | < .0001 | < .0001 |
| 0             | 309,829 88.4%                  | 1255 59.5%                       | 128 42.8%                 |                           |                |
| 1 or 2        | 34,980 9.9%                    | 607 28.8%                        | 92 30.8%                  |                           |                |
| ≥ 3           | 5561 1.6%                      | 249 11.8%                        | 79 26.4%                  |                           |                |

Unless otherwise noted, the percentage is calculated as the respondents for that category divided by the total nonmissing or declined-to-answer responses for that question.

a Respondents from Europe are not included in ethnicity as it is not collected for European countries in the NHWS.
Taken in concert, our results may indicate that the impact of CDI on HRQoL is different from other diarrheal diseases. A study conducted in the US sample of NHWS found, after adjusting for confounders, that patients with irritable bowel syndrome (IBS), diarrhea subtype (IBS-D), reported significantly lower HRQoL (mean MCS 45.2 vs. 49.5 \( p < 0.001 \); mean PCS 47.3 vs. 50.7; \( p < 0.001 \); mean SF-6D 0.68 vs. 0.74; \( p < 0.001 \)) and greater absenteeism (5.1 vs. 2.9%; \( p = 0.004 \)), presenteeism (17.9 vs. 11.3%; \( p < 0.001 \)), overall work productivity loss (20.7 vs. 13.2% \( p < 0.001 \)), and activity impairment (29.6 vs. 18.9%; \( p < 0.001 \)) than those without IBS-D [31]. Another study using the US NHWS found that persons with any form of IBS reported mean unadjusted MCS, PCS, activity impairment, and overall work impairment scores of 43, 42, 43, and 30%, respectively [24]. A key threat to the validity of comparing these studies to ours is the chronicity of IBS, which contrasts with the generally acute nature of CDI. In a small hospital-based matched case–control study, domains of the SF-36 were found to be lower among patients with antibiotic-associated diarrhea (\( n = 18 \)) than among those without (\( n = 36 \)); however, while the differences met MID criteria, no statistically significant differences were found, most likely due to the small sample size [32]. The same study also assessed HRQoL using a validated gastrointestinal disease-specific instrument, finding significantly worse HRQoL among those in the hospital with antibiotic-associated diarrhea, when compared to the published norms for active IBS patients in the USA [32]. Since diarrhea is the most frequent symptom of CDI [33] and diarrheal diseases are known to impair HRQoL, CDI’s impact on HRQoL is not surprising. Multiple patient populations, such as those with renal transplant or patients with HIV, have reported worsened HRQoL in association with acute diarrhea, some showing a dose–response effect, where the HRQoL is diminished in proportion to greater diarrheal symptoms [34, 35].

Health utilities for noninfectious diarrhea from symptomatic prostate cancer, colorectal cancer chemotherapy-induced diarrhea, or ulcerative colitis have been used as proxies for CDI in cost-effectiveness modeling [36–39]. Given that our results suggest greater impairments to HRQoL than have been noted in other gastrointestinal afflictions, a CDI-specific instrument may be useful in improving the accuracy and precision of these estimates. To date, no CDI-specific HRQoL instrument published has been fully validated, although CDI-DaySymSTM and Cdiff32 are known to be in development [23, 40].

Our study has a number of limitations, most notably the likelihood of selection bias. While the NHWS is constructed to be representative of the general population of each country in terms of age, sex and, in the USA, racial and ethnic mix, the self-selected nature of the respondents due to the opt-in nature and the low response rates almost
guarantee some degree of selection bias, likely leaving out in particular people who are either too sick to participate, or too healthy to bother. Therefore, it is likely that our results cannot be generalized to persons suffering a current bout of more severe CDI. However, it is also likely that those people, if included in our study, would have substantially
increased the magnitude of HRQoL impairments associated here with current CDI. The fact that we identified an important HRQoL impairment in a group of patients with what is most likely mild-to-moderate disease demonstrates CDI’s detrimental impact on these outcomes. Additionally, the majority of respondents in our analysis are from the USA or EU5, which may limit the generalizability of results outside of these geographic areas. Classification of respondents by CDI status relied on individuals being aware of and reporting their medical diagnosis. It is possible that respondents may not have identified their antibiotic-associated diarrhea as CDI. To partially address this, respondents who indicated “C. difficile (antibiotic associated colitis which is diarrhea from antibiotic use),” but did not indicate their condition was doctor-diagnosed were excluded; however, the possibility remains that true CDI respondents were misclassified as NO-CDI leading to more conservative results. Our results may also be subject to residual confounding as, though we adjusted for age, comorbidities, sex, education, and country, we could not adjust for differences in HCRU between groups due to unknown temporality. Additionally, there were likely unmeasured or unknown confounders that could have affected the results. As CDI diagnosis and treatment status were self-reported and not independently verified, the possibility of misclassification due to a recall bias exists. Variable mapping allowed us to combine data across countries for a larger sample size, but may have introduced measurement error in the covariates as the variable categories in the aggregated cohort were collapsed into fewer categories for health insurance, race/ethnicity, and education level. It should be noted that this increase in error variance would tend to reduce power (type 2 error) in the adjusted analysis rather than result in spurious findings (type 1 error). As clinically important differences have not been defined for diarrheal diseases, this study used common benchmarks for MIDs as defined by the developers of the measures; however, MIDs and clinically important differences are known to vary between populations and at different levels of HRQoL. Finally, as with any cross-sectional study, causality should not be inferred. That is, it is possible that at least in a subsample of the responders it was the worsened HRQoL that led to a positive response on the CDI questions, at least in the absence of current treatment.

Its limitations notwithstanding, our study highlights the global humanistic burden of CDI and raises important concerns regarding the degree to which CDI is associated with HRQoL, work, and activity impairments. Understanding the precise extent of CDI’s impact on its sufferers’ HRQoL and

**Fig. 4** Adjusted mean health utility by CDI status. *Statistically significant difference meeting MID criteria versus NO-CDI, † statistically significant difference meeting MID criteria versus P-CDI

**Fig. 5** Adjusted mean work and activity impairment by CDI status. *Statistically significant difference versus NO-CDI, differences between P-CDI and C-CDI were not statistically significant
impairment to work is essential in order to quantify the indirect costs and benefits of treatment and prevention strategies for this potentially debilitating infection.

**Funding** This study was sponsored by Pfizer, Inc.

**Compliance with ethical standards**

**Conflict of interest** HK, YH, VJ, CR, and HJ are employees and stockholders of Pfizer, Inc., the study sponsor. ZM has received compensation as a consultant to Pfizer, Inc., including for the development of this manuscript. She has also received research and/or consulting funding from Merck, The Medicines Company, Astellas, Melinta, Tetraphase, and Paratek. Editorial support was provided by Scott Vuocolo of Pfizer, Inc.

**Open Access** This article is distributed under the terms of the Creative Commons Attribution-NonCommercial 4.0 International License (http://creativecommons.org/licenses/by-nc/4.0/), which permits any noncommercial use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made.

**References**

1. Keller JM, Surawicz CM. *Clostridium difficile* infection in the elderly. Clin Geriatr Med. 2014;30:79–93.
2. Lessa FC, Mu Y, Bamberg WM, et al. Burden of *Clostridium difficile* infection in the United States. N Engl J Med. 2015;372:825–834.
3. Miller BA, Chen LF, Sexton DJ, Anderson DJ. Comparison of the burdens of hospital-onset, healthcare facility-associated *Clostridium difficile* infection and of healthcare-associated infection due to methicillin-resistant *Staphylococcus aureus* in community hospitals. Infect Control Hosp Epidemiol. 2011;32:387–390.
4. Centers for Disease Control and Prevention [CDC]. *Antibiotic resistance threats in the United States*. Washington, DC: Centers for Disease Control and Prevention; 2013.
5. Chitinis A, Holzbauer S, Belflower R, et al. Epidemiology of community-associated *Clostridium difficile* infection, 2009–2011. JAMA Intern Med. 2013;173:1359–1367.
6. Centers for Disease Control and Prevention. 2015 Annual Report from the Emerging Infections Program for *Clostridium difficile* Infection. Department of Health and Human Services; 2015. https://www.cdc.gov/hai/eip/pdf/cdiff/2015-CDI-Report.pdf. Accessed 21 Mar 2018.
7. Khanna S, Pardi D, Aronson S, Kammer P, Orenstein R, Sauver J. The epidemiology of community-acquired *Clostridium difficile* infection: a population-based study. The American Journal of Gastroenterology. 2012;107:89–95.
8. Barbut F, Mastrantonio P, Delmee M, et al. Prospective study of *Clostridium difficile* infections in Europe with phenotypic and genotypic characterisation of the isolates. Clin Microbiol Infect. 2007;13:1048–1057.
9. Borren NZ, Ghadermarzi S, Huttlless S, Ananthakrishnan AN. The emergence of *Clostridium difficile* infection in Asia: a systematic review and meta-analysis of incidence and impact. PLoS ONE. 2017;12:e0176797.
10. Collins DA, Hawkey PM, Riley TV. Epidemiology of *Clostridium difficile* infection in Asia. Antimicrob Resist Infect Control. 2013;2:21.
11. Davies KA, Longshaw CM, Davis GL, et al. Underdiagnosis of *Clostridium difficile* infection across Europe: the European, multicentre, prospective, biannual, point-prevalence study of *Clostridium difficile* infection in hospitalised patients with diarrhoea (EUCLID). Lancet Infect Dis. 2014;14:1208–1219.
12. Lopardo G, Morfin-Otero R, Moran V II, et al. Epidemiology of *Clostridium difficile*: a hospital-based descriptive study in Argentina and Mexico. Braz J Infect Dis. 2015;19:8–14.
13. Elixauser A (AHRQ), Steiner C (AHRQ), Gould C (CDC). Readmissions following hospitalizations with *Clostridium difficile* infections. HCUP Statistical Brief #145. December 2012. Agency for Healthcare Research and Quality, Rockville, MD: 2009. Available at http://www.hcup-us.ahrq.gov/reports/statbriefs/sb145.pdf. Accessed 21 Mar 2018.
14. Lucado J, Gould C, Elixauser A. *Clostridium difficile* infections (CDI) in Hospital Stays. HCUP Statistical Brief #124. January 2012. Agency for Healthcare Research and Quality, Rockville, MD; 2009. http://www.hcup-us.ahrq.gov/reports/statbriefs/sb124.pdf. Accessed 21 Mar 2018.
15. Eckmann C, Wasserman M, Latif F, Roberts G, Beriot-Mathiot A. Increased hospital length of stay attributable to *Clostridium difficile* infection in patients with four co-morbidities: an analysis of hospital episode statistics in four European countries. Eur J Health Econ. 2013;14:835–846.
16. Kwon JH, Olsen MA, Dubberke ER. The morbidity, mortality, and costs associated with *Clostridium difficile* infection. Infect Dis Clin North Am. 2015;29:123–134.
17. Tabak YP, Zilberberg MD, Johannes RS, Sun X, McDonald LC. Attributable burden of hospital-onset *Clostridium difficile* infection: a propensity score matching study. Infect Control Hosp Epidemiol. 2013;34:588–596.
18. Dubberke ER, Butler AM, Reske KA, et al. Attributable outcomes of endemic *Clostridium difficile*-associated disease in nonsurgical patients. Emerg Infect Dis. 2008;14:1031–1038.
19. Dubberke ER, Olsen MA. Burden of *Clostridium difficile* on the healthcare system. Clin Infect Dis. 2012;55:S88–S92.
20. Ghanotji SS, Sail K, Lairson DR, DuPont HL, Garey KW. Economic healthcare costs of *Clostridium difficile* infection: a systematic review. J Hosp Infect. 2010;74:309–318.
21. Yu H, Baser O, Wang L. Burden of *Clostridium difficile*-associated disease among patients residing in nursing homes: a population-based cohort study. BMC Geriatr. 2016;16:193.
22. European CDC [ECDC]. *Clostridium difficile* basic facts. http://www.ecdc.europa.eu/en/HEALTHTOPICS/CLOSTRIDIUM_DIFFICILE_INFECTION/BASIC_FACTS/Pages/basic_facts.aspx. Accessed 20 May 2013.
23. Garey KW, Aitken SL, Glischwind L, et al. Development and validation of a *Clostridium difficile* health-related quality-of-life questionnaire. J Clin Gastroenterol. 2016;50:631–637.
24. Peery AF, Dellon ES, Lund J, et al. Burden of gastrointestinal disease in the United States: 2012 update. Gastroenterology. 2012;143:1179–1187.
25. Wilcox MH, Ahir H, Coia JE, et al. Impact of recurrent *Clostridium difficile* infection: hospitalization and patient quality of life. J Antimicrob Chemother. 2017;72:2647–2656.
26. Organisation for Economic Co-operation and Development (OECD). Education at a glance 2017: OECD indicators. Paris: OECD Publishing; 2017. https://doi.org/10.1787/eag-2017-en.
27. Charlson ME, Pompei P, Ales KL, MacKenzie CR. A new method of classifying prognostic comorbidity in longitudinal studies: development and validation. J Chronic Dis. 1987;40:373–383.
28. Maruish ME, ed. User’s manual for the SF-36V2®/2 health survey. 3rd ed. Lincoln, RI: Quality Metric Inc; 2011.
29. Reilly MC, Zbrozek AS, Dukes EM. The validity and reproducibility of a work productivity and activity impairment instrument. *Pharmacoconomics*. 1993;4:353–365.
30. Walters SJ, Brazier JE. Comparison of the minimally important difference for two health state utility measures: EQ-5D and SF-6D. *Qual Life Res*. 2005;14:1523–1532.
31. Buono JL, Carson RT, Flores NM. Health-related quality of life, work productivity, and indirect costs among patients with irritable bowel syndrome with diarrhea. *Health Qual Life Outcomes*. 2017;15:35.
32. Gbito KE. The impact of antibiotic associated diarrhea on health-related quality of life in hospitalized patients (Unpublished masters thesis). The University of Texas School of Public Health, Houston, Texas; 2009. https://search.proquest.com/openview/2c025c7da2ca63ef25aaac0ea0939017/1.pdf?pq-origsite=gscholar&cbl=18750&diss=y. Accessed 21 Mar 2018.
33. Kelly CP, Lamont JT. Patient education: antibiotic-associated diarrhea caused by *Clostridium difficile*. In: EL Baron (ed) Uptodate. https://www.uptodate.com/contents/antibiotic-associated-diarrhea-caused-by-clostridium-difficile-beyond-the-basics. Accessed 1 Nov 2017.
34. Ekberg H, Kyllonen L, Madsen S, Grave G, Solbu D, Holdaas H. Increased prevalence of gastrointestinal symptoms associated with impaired quality of life in renal transplant recipients. *Transplantation*. 2007;83:282–289.
35. Siddiqui U, Bini EJ, Chandarana K, et al. Prevalence and impact of diarrhea on health-related quality of life in HIV-infected patients in the era of highly active antiretroviral therapy. *J Clin Gastroenterol*. 2007;41:484–490.
36. Bartsch SM, Curry SR, Harrison LH, Lee BY. The potential economic value of screening hospital admissions for *Clostridium difficile*. *Eur J Clin Microbiol Infect Dis*. 2012;31:3163–3171.
37. Konijeti GG, Sauk J, Shriime MG, Gupta M, Ananthakrishnan AN. Cost-effectiveness of competing strategies for management of recurrent *Clostridium difficile* infection: a decision analysis. *Clin Infect Dis*. 2014;58:1507–1514.
38. Lee BY, Popovich MJ, Tian Y, et al. The potential value of *Clostridium difficile* vaccine: an economic computer simulation model. *Vaccine*. 2010;28:5245–5253.
39. Stranges PM, Hutton DW, Collins CD. Cost-effectiveness analysis evaluating fidaxomicin versus oral vancomycin for the treatment of *Clostridium difficile* infection in the United States. *Value Health*. 2013;16:297–304.
40. Talbot GH, Kleinman L, Davies EW, Hunsche E, Roberts L, Nord CE. The *Clostridium difficile* infection—daily symptoms (CDI-DaySyms™) patient-reported outcome (PRO) questionnaire: final validation and responder thresholds. *Open Forum Infectious Diseases*. 2017;4:S394.