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1 **Impact of recurrent Clostridium difficile infection: Hospitalisation and patient quality of life**

2

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22 **Running title:** Impact of recurrent Clostridium difficile infection

23

24

25 **Abstract**

26

27 **Objectives**

28 Data quantifying outcomes of recurrent *Clostridium difficile* infection (rCDI) are lacking. We sought
29 to determine the UK hospital resource use and health-related quality of life (HrQoL) associated with
30 rCDI hospitalisations.

31 **Patients and methods**

32 A non-interventional study in 6 UK acute hospitals collected retrospective clinical and resource use
33 data from medical records of 64 adults hospitalised for rCDI and 64 matched inpatient controls with
34 a first episode only (f)CDI. Patients were observed from the index event (date rCDI/fCDI confirmed)
35 for 28-days (or death, if sooner); UK-specific reference costs were applied. HrQoL was assessed
36 prospectively in a separate cohort of 30 patients hospitalised with CDI, who completed the EQ-5D-3L
37 questionnaire during their illness.

38 **Results**

39 The median total management cost (post-index) was £7,539 and £6,294 for rCDI and fCDI,
40 respectively (cost difference, $p=0.075$); median length of stay (LOS) was 21 days and 15.5 days,
41 respectively ($p=0.269$). The median cost difference between matched rCDI and fCDI cases was £689
42 (IQR=£-1,873-£3,954). Subgroup analysis demonstrated the highest median costs (£8,542/patient) in
43 severe rCDI cases. CDI management costs were driven primarily by hospital LOS, which accounted
44 for >85% of costs in both groups. Mean EQ-5D index values were 46% lower in CDI patients
45 compared with UK population values (0.42 and 0.78, respectively); EQ-VAS scores were 38% lower
46 (47.82 and 77.3, respectively).

47 **Conclusions**

48 CDI has considerable impact on patients and healthcare resources. This multicentre study provides a
49 contemporaneous estimate of the real-world UK costs associated with rCDI management, which are
50 substantial and comparable to fCDI costs.

51

52 Introduction

53 *Clostridium difficile* infection (CDI) is a major public health challenge worldwide, and is associated
54 with significant morbidity, mortality and healthcare resource utilisation.¹⁻⁵ In the UK, although CDI
55 reports decreased by 61% between 2007/08 and 2010/11 following the introduction of national
56 surveillance, there was a 6% increase in CDI cases in England (from 24.8-26.3/100,000 population)
57 between 2013/14 and 2014/15.⁶⁻⁸

58 A recent study undertaken in 2012-13 across 482 hospitals in 20 European countries reported a CDI
59 incidence of 7 cases/10,000 patient-bed days, a 70% increase on rates recorded in 2008.
60 Furthermore, when diarrhoeal samples were re-tested by an optimised method for diagnosing CDI,
61 about a quarter of cases had been missed locally; consequently, the true rate of CDI in Europe is
62 probably much higher.⁹ In the US, *C. difficile* was recently reported to be the most common cause of
63 healthcare associated infection, with approximately half-a-million CDI cases and 29,000 deaths in
64 2011.¹⁰

65 It is estimated that recurrent CDI (rCDI) following initial resolution occurs in 20-30% of patients.¹¹⁻¹³
66 However, data on the burden and outcomes associated with rCDI are scarce. Surveillance systems
67 may fail to capture many rCDI cases given that re-testing of patients with symptoms suggestive of
68 rCDI may not occur. Notably, mandatory surveillance data in England largely exclude rCDI cases, as
69 the collected figures exclude repeat laboratory-positive results within 28-days from the same
70 patient.⁸ There is a lack of contemporaneous information quantifying the economic burden of CDI in
71 the UK, and particularly the resource use associated with recurrent episodes. Such costs have
72 growing relevance as new therapeutic options become available that reduce rCDI rates compared
73 with conventional treatments.^{14,15}

74 In addition to the economic burden, it is important to consider the impact of new CDI therapeutics
75 on health-related quality of life (HRQoL); this may be impaired in CDI patients due to decreased

76 functional capacity and anxiety about physical symptoms or complications.^{16,17} Despite the high
77 incidence of CDI, its impact on HRQoL has not been widely studied and therefore conventional
78 economic analyses may underestimate the true burden.

79 This industry-initiated study aimed to quantify the cost of hospital resource use (HRU) for patients
80 with rCDI and describe the impact of CDI on HRQoL. The study was initiated by Merck Sharp and
81 Dohme Limited (MSD) prior to the Phase 3 study for bezlotoxumab, which has subsequently been
82 approved for prevention of CDI recurrence and was designed to provide “real-world” data that can
83 be used to help determine the cost-effectiveness of new CDI management options.

84

85 **Methods**

86 A mixed-methodology non-interventional study was conducted between September-2013 and
87 September-2014 in six geographically-dispersed UK National Health Service hospitals. Potential
88 hospital sites likely to provide sufficient study participants and representation across NHS England
89 regions and Scotland were identified by the lead investigator, study sponsor, and by review of Health
90 Protection Agency (now PHE) mandatory surveillance data. Potential hospitals were approached and
91 subsequently recruited to the study on the basis of their ability/capacity to deliver the study data
92 collection requirements (including database systems that would allow identification of eligible
93 patients; availability of local clinical staff to seek informed consent and collect the required data; and
94 confirmed participant availability). Financial support for individual centres was provided in line with
95 the National Institute for Health Research (NIHR) costing template, as is standard for studies
96 implemented in the UK. Ethics committee (London-Brent, reference 13/LO/1046) and relevant local
97 approvals were obtained. The study was undertaken in two parts:

98 **Part 1: Matched retrospective cohort study**

99 ***Design***

100 A matched retrospective cohort study covering an observation period of 01-March-2012 to 02-June-
101 2014 gathered clinical and HRU data from the medical records of patients hospitalised for rCDI and
102 matched patients from the same centres, with a first episode of CDI but no recurrence (fCDI). The
103 design and flow is summarised in **Figure 1**.

104 ***Patients***

105 rCDI cases were adult (≥ 18 years) inpatients with a positive CD toxin test after 01-May-2012 (the
106 index result) and any previous positive CD toxin test in the ≤ 12 -weeks before this result. Potential
107 patients were identified from microbiology records and eligibility confirmed by cross-referencing
108 with hospital administration systems and full CDI testing records. Patients were included in the final
109 dataset only where a matched (fCDI) control was identified (see below). May-2012 was chosen as
110 the start of the observation period because the UK Department of Health issued revised guidance on
111 *C. difficile* diagnosis and reporting in March-2012; hence, it was considered that there would be
112 greater uniformity between centres in CDI testing after this date.¹⁸ It was expected (although not
113 confirmed) that included CDI cases would have been diagnosed according to this guidance; although
114 all patients identified as eligible by the participating centres were included.

115 Matched fCDI controls were patients with a first CDI episode (community or healthcare-facility
116 acquired) but no subsequent positive CD toxin test within the 12-weeks following last CDI treatment,
117 who matched a rCDI case according to date of first positive CD toxin test (± 12 -weeks), age group
118 ($< 75 / \geq 75$ years) and gender.

119 Patients were excluded if they transferred hospital trusts or died before the end of CDI treatment.
120 All eligible paired patients were included. Patient consent was not required since this part of the
121 study involved only routinely-collected clinical data gathered in pseudo-anonymised form by
122 members of the direct care team.

123 ***Data collection***

124 Pseudo-anonymised data were collected retrospectively by local clinical staff from eligible patients'
125 hospital medical records using a standard data collection form. The dataset comprised baseline
126 demographics; co-morbidities; CDI strain and illness severity at first episode (both groups) and
127 recurrence (rCDI only); and HRU (hospital admission and discharge dates, length of stay [LOS] per
128 ward/side room, outpatient appointments, Emergency Department [ED] attendances, prescribing,
129 diagnostic tests, supplementary nutrition). HRU data were collected for the 'post-index' period,
130 defined as the period between the index event (date rCDI or fCDI first confirmed by positive CD toxin
131 test) and 28-days post-index or death, whichever was shorter. A 28-day observation period was
132 chosen to reduce the risk of the results being skewed by non-CDI-related resource. Due to the acute
133 nature of the disease, HRU occurring after 28-days was considered much less likely to be attributable
134 to CDI. For rCDI, data were also collected for the 'between-episode' period (from 72-hours after end
135 of treatment for the first CDI episode until the index event).

136 Resource costs were calculated using UK-specific reference costs (Supplementary appendix 1) and a
137 Market Forces Factor Index applied to the costs for each Trust.¹⁹⁻³⁰

138 ***Outcomes***

139 The primary outcome was the difference in total hospital management costs between patients with
140 rCDI and fCDI. Secondary outcomes included the difference in the number of days hospitalised
141 during the post-index period and total management costs for the between-episode period (rCDI
142 only).

143 ***Statistical analysis***

144 We aimed to include 75 rCDI and 75 fCDI patients in the study. As there are no UK estimates of rCDI
145 costs, the sample size was based on US data, which showed an average LOS for rCDI of 9-days and a
146 cost range of \$3,500-\$5,000/recurrence (1999 Dollars).³¹ Assuming similar UK costs and using the
147 mid-point of this estimate converted to UK pounds (£2,800), a sample of 75 patients provided a 95%
148 CI of £2,623-£2,977 (\pm £177 [6%]); this was considered to be interpretable to clinicians and payers

149 given the magnitude of the cost difference between established and more expensive new
150 therapeutics. Six study centres were used, with the expectation of achieving the recommended
151 sample size based on expected numbers of eligible paired patients.

152 Analysis was conducted using Microsoft Excel on the available data, with no imputation of missing
153 values apart from a set of pre-specified assumptions (Supplementary Appendix 1). The number of
154 patients available for each analysis is stated where data were missing. Descriptive endpoints are
155 presented using the mean (standard deviation, SD), median (IQR) or percentages, as appropriate.
156 The difference between rCDI and fCDI patients in the median total cost of treating CDI and the
157 median LOS during the post-index period was compared using the Wilcoxon rank-sum test.

158 HRU endpoints are presented overall and stratified by CDI severity (a planned subgroup analysis).
159 Severe CDI was defined by the presence of any of the following criteria: white cell count $>15 \times 10^9/L$,
160 acutely rising blood creatinine (e.g. $>50\%$ increase above baseline), temperature $>38.5^\circ C$ or evidence
161 of severe colitis (abdominal signs, radiology).³² When none of these was present, CDI was classified
162 as mild/moderate.

163 **Part 2: Prospective patient self-assessment of QoL**

164 ***Design***

165 As HRQoL is not routinely measured and documented in medical records, it was assessed
166 prospectively in a separate cohort of adult patients from the same centres, who were hospitalised
167 with CDI. Eligible patients completed the EQ-5D-3L questionnaire^{33,34} during their illness, within five-
168 days of symptom onset. Questionnaires were completed between 10-September-2013 and 07-
169 August-2014.

170 Demographic and disease history data including gender, age and CDI severity were recorded from
171 the patients' medical records.

172 ***Patients***

173 Patients were included if they had a positive CDI test, were hospital inpatients and aged ≥ 18 years at
174 the date of the positive CDI test and consented to complete the questionnaire. Owing to the lack of
175 data on HRQoL in CDI patients in general, part 2 was not restricted to rCDI and all patients with CDI
176 (both first and recurrent episodes) were eligible for inclusion. Potentially eligible patients were
177 identified from microbiology records and, if considered by clinical staff to be competent to consent,
178 they were approached by a member of their care team with study information and asked if they
179 wished to participate. Only consenting patients were included. Consecutive eligible patients were
180 invited until the recruitment target was met (30 patients, maximum 10 patients/centre). A sample
181 size of 30 patients was recommended, based on the Central Limit Theorem, assuming the results
182 would be normally distributed.

183 ***Outcomes***

184 The main outcomes were the mean (SD) EQ-5D index and visual analogue scale (VAS) scores.

185 ***Statistical analysis***

186 The EQ-5D descriptive system was scored according to the published instructions.³⁵ EQ-5D index and
187 VAS scores in patients with CDI were compared with published norms for the UK general population
188 using Welch's t-test.³⁶ The EQ-5D population norm for patients aged 65-74 years was used for
189 comparison, since the median age of patients in our study (70-years) was within this range.

190 **Results**

191 ***Part 1: Matched retrospective cohort study***

192 ***Demographics and CDI characterisation***

193 Sixty-four rCDI patients and 64 matched fCDI controls were included (range 8-14 pairs/centre). The
194 pre-planned sample size of 75 matched pairs was not achieved owing to challenges with matching

195 patients (as described in Figure 1). Patients' demographic and clinical characteristics are summarised
196 in **Table 1**.

197 Thirty-three percent (21/64) of rCDI patients had severe CDI at the recurrent episode; 52% (33/64)
198 had severe infection at their first episode, compared with 41% (26/64) of the matched fCDI controls.
199 There was considerable heterogeneity in *C. difficile* strains identified, with 27 different ribotypes
200 identified overall. Nine percent (6/64) of rCDIs, 11% (7/64) of the first episodes (in rCDI cases) and
201 8% (5/64) of fCDIs were attributable to the hypervirulent ribotypes 078 and 027, with other strains
202 (most commonly 002, 014, 015) accounting for the majority of CDI cases. Thirteen rCDI patients
203 (20%) had a different CDI strain compared with the isolate recovered from their first CDI episode (i.e.
204 re-infection).

205 Six percent (4/64) of rCDI cases (all with mild/moderate CDI) and 14% (9/64) of matched fCDI
206 controls (5 severe, 4 mild/moderate CDI) died within the 28-day post-index period. The median
207 duration of the post-index period in deceased patients was 13-days (IQR=7.3-18.8) for rCDI and 12-
208 days (IQR=9.0-16.0) for fCDI.

209 **Resource utilisation and costs**

210 The total costs of treating rCDI and fCDI patients during the 28-day post-index period are shown in
211 **Table 2**. The median cost per patient was £7,539 (IQR=£5,617-£9,730) for rCDI and £6,294
212 (IQR=£2,700-£9,216) for fCDI (cost difference, p=0.075). There were some outliers in the fCDI group,
213 with three patients having total costs >£20,000.

214 Because more fCDI than rCDI patients (9 versus 4, respectively) died during the post-index period, a
215 post-hoc sensitivity analysis was conducted on data from the subgroup of 52 matched pairs where
216 both patients survived to the end of the observation period (i.e. excluding both patients from pairs
217 in which one died). In this group, median costs were similar to those for the overall sample: £7,888
218 (IQR=£6,047-£9,866) and £6,719 (IQR=£3,329-£9,216) for rCDI and fCDI, respectively (**Table 2**).

219 The differences in costs between matched rCDI and fCDI patients (cost for rCDI case minus cost for
220 fCDI control) ranged from -£38,163 (fCDI>rCDI) to £11,841 (rCDI>fCDI), with a median difference of
221 £689 (IQR=-£1,873-£3,954) (rCDI>fCDI) (**Figure 2**).

222 **Table 3** shows the breakdown of total costs. The cost of hospital admissions and ED visits accounted
223 for the majority (>85%) of costs for both groups. The median cost for CDI-specific medicines was
224 higher in rCDI patients (£376 per patient [IQR=£31-£1,521]) compared with fCDI (£46 [IQR=£2-£286])
225 (**Table 3**).

226 When stratified by severity, the median cost of CDI treatment per patient with severe infection was
227 £5,631 (IQR=£2,910-£9,453) for fCDIs and £8,542 (IQR=£7,463-£10,532) for rCDIs (cost difference,
228 p=0.039). When deceased patients were excluded, median costs were £6,961 (IQR=£4,464-£10,138)
229 and £9,030 (IQR=£7,463-£10,288) for severe fCDIs and rCDIs, respectively (**Table 2**).

230 The cumulative total number of bed days (median) in rCDI patients during the post-index period was
231 1,171 (21) days compared with 1,027 (15.5) days for fCDI (difference, p=0.269). The highest median
232 number of bed days (25.5) was observed in patients with a severe rCDI.

233 The median cost for the between-episode period (rCDI only) was £2,973 (IQR=£778-£4,610) (**Table**
234 **2**).

235 ***Part 2: Prospective patient self-assessment of QOL***

236 **Demographics**

237 Thirty patients completed the EQ-5D-3L questionnaire during a CDI hospitalisation, of whom 63%
238 (19/30) were male. The median age was 70.2 years (IQR=52-77). CDI was severe in 27% of patients
239 (8/30) and mild/moderate in 73% (22/30).

240 **EQ-5D scores**

241 EQ-5D index and VAS scores for patients hospitalised with CDI compared with age-matched
242 population norms are shown in **Figure 3**. The mean EQ-5D index score in CDI patients (0.42
243 [SD±0.29]) was 46% lower than the value for patients of similar age (65-74 years) in the UK general
244 population (0.78) (difference, $p<0.001$); similar reduced scores were observed for the VAS (mean
245 47.82 [SD±21.93] for CDI, 38% lower than the general population score of 77.3, $p<0.001$). EQ-5D
246 dimension scores are shown in **Table 4**.

247 **Discussion**

248 This non-interventional study used a matched retrospective cohort design to estimate the current
249 costs associated with treatment of rCDI in hospitalised patients in the UK. It provides
250 contemporaneous cost-burden data to aid decision-making by payers and clinicians on the targeting
251 of resources for CDI treatment. The study also demonstrates the adverse impact of CDI on HrQoL,
252 which has to-date been a largely neglected area of research. Taken together, the findings highlight
253 the considerable burden that CDI places on patients and healthcare resources and the substantial
254 financial costs associated with both fCDI and rCDI.

255 In this study, there was considerable heterogeneity of strains causing CDI with 27 different strains
256 identified overall. This pattern is consistent with the epidemiology of CDI in the UK, where no
257 particular ribotypes are dominant and suggests an endemic (non-outbreak) population.¹² It is
258 therefore more representative to the wider UK patient population than data derived during a CDI
259 epidemic.

260 We found the total cost of treating CDI and hospital LOS to be higher for patients with rCDI
261 compared with fCDI, although these differences were not statistically significant. This may be due to
262 lack of power as a consequence of not meeting the planned sample size, but may also reflect the
263 wide variation in costs between individual patients; this is typical in analyses of healthcare costs and
264 has been observed in previous studies.^{1,37} It is also acknowledged that the differences between rCDI
265 and fCDI costs may in part be due to the higher number of deaths in the fCDI group. When deceased
266 patients were excluded, the difference between the groups was smaller than for the whole study
267 sample but the cost remained higher for rCDI than fCDI. Recent systematic reviews have
268 demonstrated incremental costs of \$2,871-\$4,846/case for primary CDI in US-based² studies and
269 £4,577-£8,843 in European studies.¹ Although the median total cost for fCDI in the present study
270 (£6,294) is consistent with these previous estimates, direct comparison is problematic due to
271 methodological differences and variability in costs between different studies, partly due to

272 differences in healthcare systems. Few studies have estimated the costs associated with rCDI
273 specifically. One US study found that the cost of treating rCDI was \$4,948 per-episode,^{31,38} which is
274 broadly similar to the £7,539 observed in our study. However, our costs are lower than a recent
275 single centre US study reported by Dubberke *et al.* (attributable costs \$11,631 over 180-days)³⁷ and
276 those reported in a recent abstract (£20,249) presenting the results of a costing analysis in a single
277 UK centre.³⁹ This may reflect the use of different reference costs or the fact that we used a fixed 28-
278 day observation period. The results of the present study provide an updated estimate of the UK
279 costs associated with treating rCDI, which is important given the lack of contemporaneous data.
280 Recently, new CDI medicines (fidaxomicin and bezlotoxumab) have been developed that have been
281 shown to reduce CDI recurrence rates compared with conventional treatments.^{14,15} Our results will
282 help clinical decision makers to evaluate the cost-effectiveness of investing in these new medicines
283 and the potential impact on local service provision. The resource utilisation between CDI episodes in
284 rCDI patients (median £2,973) may represent previously un-recognised costs associated with
285 treatment of patients with CDI once specific CDI treatment has ceased.

286 Consistent with previous studies, the costs of treating CDI in this study were driven largely by the
287 costs associated with the duration of hospital admissions, which accounted for >85% of total costs in
288 both groups.^{40,1} However, the costs of CDI medicines were considerably higher in rCDI patients,
289 reflecting the use of fidaxomicin in recurrent but not fCDI. The median LOS in rCDI cases was 21
290 days, which is similar to the 20.5 days reported previously in the UK.⁴⁰ This similarity is perhaps
291 surprising given that the previous UK study was reported in 1996, and significant efforts have been
292 made to shorten hospital admissions in the intervening period.⁴¹

293 The results presented here on the cost of treating CDI according to disease severity are important as
294 PHE recommends different treatment strategies for patients with severe and mild/moderate
295 disease.⁴² The highest median total cost was observed in severe rCDI cases (£8,542) and the lowest
296 in patients with severe fCDI (£5,631), although as before, this may be explained partly by the higher
297 number of deaths among fCDI patients with severe disease.

298 HRQoL in hospitalised CDI patients is dramatically reduced compared with people of similar age in
299 the general population, although from our descriptive study it is unclear whether this is directly
300 attributable to CDI or more generally to the effects of the hospitalisation and associated co-
301 morbidities. Published EQ-5D norms for the general hospital population are not available and
302 comparison with previous studies of hospitalised patients is problematic since most were conducted
303 in patients with specific health conditions or those undergoing surgery, often in single centres or
304 countries outside the UK. In one recent single centre study of patients (of similar age and gender
305 distribution) on adult medical wards in a single UK hospital, the mean EQ VAS at the time of
306 admission was 55.9, 14% higher than the CDI patients in our cohort.⁴³ This would seem to suggest
307 that the reduced HRQoL is at least partly attributable to CDI, but warrants further research. In our
308 study the usual activities, mobility and self-care EQ-5D dimensions were most affected, which is
309 unsurprising for a group of hospitalised patients. However, the anxiety and depression and pain
310 dimensions were also impaired (63% reported moderate-extreme anxiety/depression; 67% reported
311 moderate-extreme pain). Recent research has demonstrated that anxiety is common in patients
312 hospitalised with CDI, with a number of CDI-specific concerns identified, including worry about
313 future complications, physical concerns about ongoing symptoms and social concerns including
314 interference with daily activities and finances.¹⁶ A limitation of the HRQoL data is that we did not
315 collect information about co-morbidities or whether the patients had a first or recurrent CDI
316 episode; further research is needed to fully understand the impact of each on HRQoL, as well as
317 changes in HRQoL over time.

318 **Strengths and limitations**

319 The primary strength of this study is the matched design for estimation of costs, and the inclusion of
320 descriptive QoL data; the latter is important to enable healthcare providers to determine the overall
321 burden of CDI and has not been widely studied. The study was designed to minimise the impact of
322 bias and confounding factors, however, there are limitations. The quality of the retrospectively-

323 sourced data relies upon the accuracy and completeness of patients' medical records and there
324 were instances (including medication details) where data were missing or incomplete. This is an
325 inherent limitation of retrospective observational research, however, the impact of missing data in
326 this study should be low because the primary endpoint is driven primarily by LOS, which was well-
327 recorded. Furthermore, cases and controls would be affected equally. Despite age-matching, there
328 were more deaths among fCDI patients, particularly in those with severe CDI; this suggests that
329 either the rCDI patients in this study are a population of patients with less severe disease or that for
330 the healthcare-facility-acquired fCDI cases, the primary reason for hospitalization (not CDI) may be
331 the main determinant of mortality. Although there were some differences in patient characteristics
332 (particularly co-morbidities) between the two cohorts, we did not adjust for these factors in the
333 analysis as they were not considered to be of sufficient magnitude to have introduced major bias
334 into the results. Furthermore, it is not uncommon in CDI cohorts to observe modest imbalances in
335 co-morbidities. We also used clear eligibility criteria and matched patients on the key characteristics
336 related to the disease. The total costs associated with treating CDI may be underestimated because
337 the post-index period was fixed at 28-days; also, the observation period started when CDI was
338 confirmed and patients may have received CDI treatment before this. Only patients with a CDI that
339 was confirmed by testing were included and consequently, the patient population may not be
340 representative of all CDI cases. Testing practices and treatment protocols may have varied between
341 the participating hospitals. These differences were not explored in the analysis owing to the small
342 number of patients per centre and expected variability between individual patients. Despite all
343 available eligible patients being included, the planned sample size of 75 matched pairs was not met
344 due to challenges of matching patients. This may have affected the reliability of the cost estimates
345 and limited the study's ability to identify true differences between the groups. Furthermore, the
346 formal sample size calculation applied only to the overall sample, not to subgroups.

347 **Conclusions**

348 This multicentre study demonstrates that CDI has a considerable impact on both patients and
349 healthcare resources. The data provide an updated estimate of the “real-world” costs associated
350 with rCDI management in the UK. These costs are largely driven by the duration of hospital
351 admissions and are comparable to fCDI costs. The study also indicates increased costs associated
352 with the treatment of patients with severe rCDI; this is important in light of PHE guidance, which
353 recommends different treatment strategies for patients with severe and mild/moderate disease.
354 Overall, the study provides contemporaneous data on the burden of CDI to patients and the
355 healthcare system, which can be used to help clinical decision makers evaluate the cost-
356 effectiveness of new CDI therapeutics, particularly those associated with reduced risk of recurrence.

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366 assistance; and Cheryl Donnelly from Merck & Co., Inc., Kenilworth, NJ USA.

367 **Contributorship statement**

368 MHW was involved in the design of the study and the acquisition, analysis and interpretation of the
369 study data. HA was involved in the analysis and interpretation of the study data. JEC and CS were
370 involved in the acquisition and interpretation of the study data. AD, SH and ML were involved in the
371 acquisition, analysis and interpretation of the study data. SMS analysed the data. SWM was involved

372 in the interpretation of the study data. All authors reviewed the draft manuscript and approved the
373 final version for submission.

374 **Transparency declarations**

375 MHW has received: consulting fees from Actelion, Astellas, bioMerieux, MedImmune, MSD, Pfizer,
376 Qiagen, Sanofi-Pasteur, Seres, Summit, Synthetic Biologics and Valneva; lecture fees from Alere,
377 Astellas, MSD & Pfizer; and grant support from Actelion, Astellas, bioMerieux, Da Volterra, MSD,
378 Sanofi-Pasteur, Seres and Summit.

379 HA and SWM are employees for the sponsoring company (Merck Sharp & Dohme Corp., a subsidiary
380 of Merck & Co., Inc., Kenilworth, NJ USA (known as MSD outside the United States and Canada)) that
381 produces a product within the disease area. The funder (and these employees) initiated the study
382 and worked collaboratively with the primary investigator in some of the study design and data
383 analysis. SWM owns stock in Merck & Co., Inc., Kenilworth, NJ USA as part of his compensation.

384 JEC has participated in an advisory board for MSD (May 2016).

385 SMS is an employee of pH Associates, an independent research consultancy which was
386 commissioned by the sponsor to provide support with the design and conduct of the study, data
387 analysis and medical writing.

388 Medical writing services were provided by Laura Baldock from pH Associates, funded by Merck Sharp
389 & Dohme Corp., a subsidiary of Merck & Co., Inc., Kenilworth, NJ USA (known as MSD outside the
390 United States and Canada).

391 AD, CS, SH and ML have no conflicts of interest.

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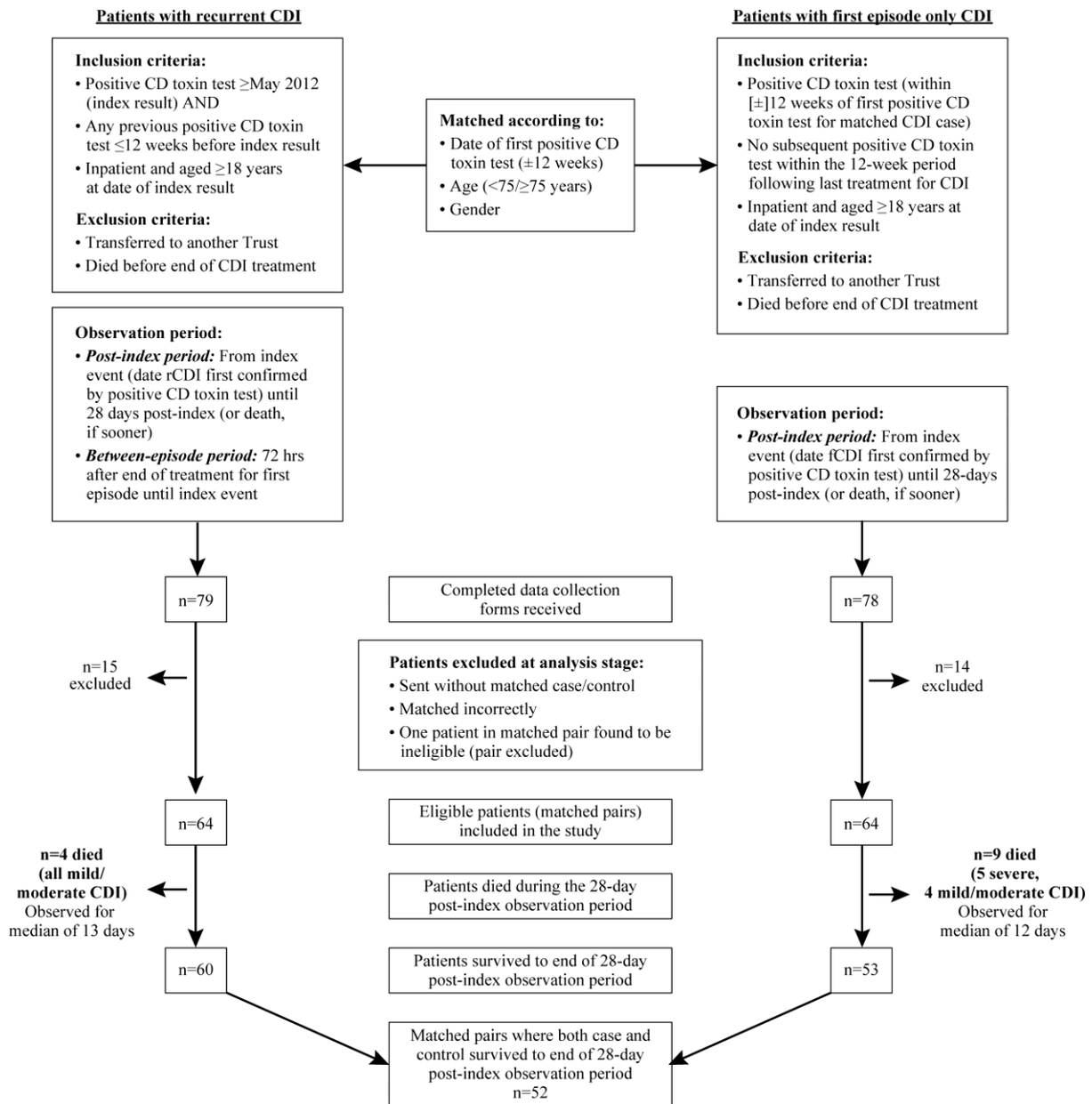
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508 **Figure 1: Summary of matched retrospective cohort design and flow of patients through the study**

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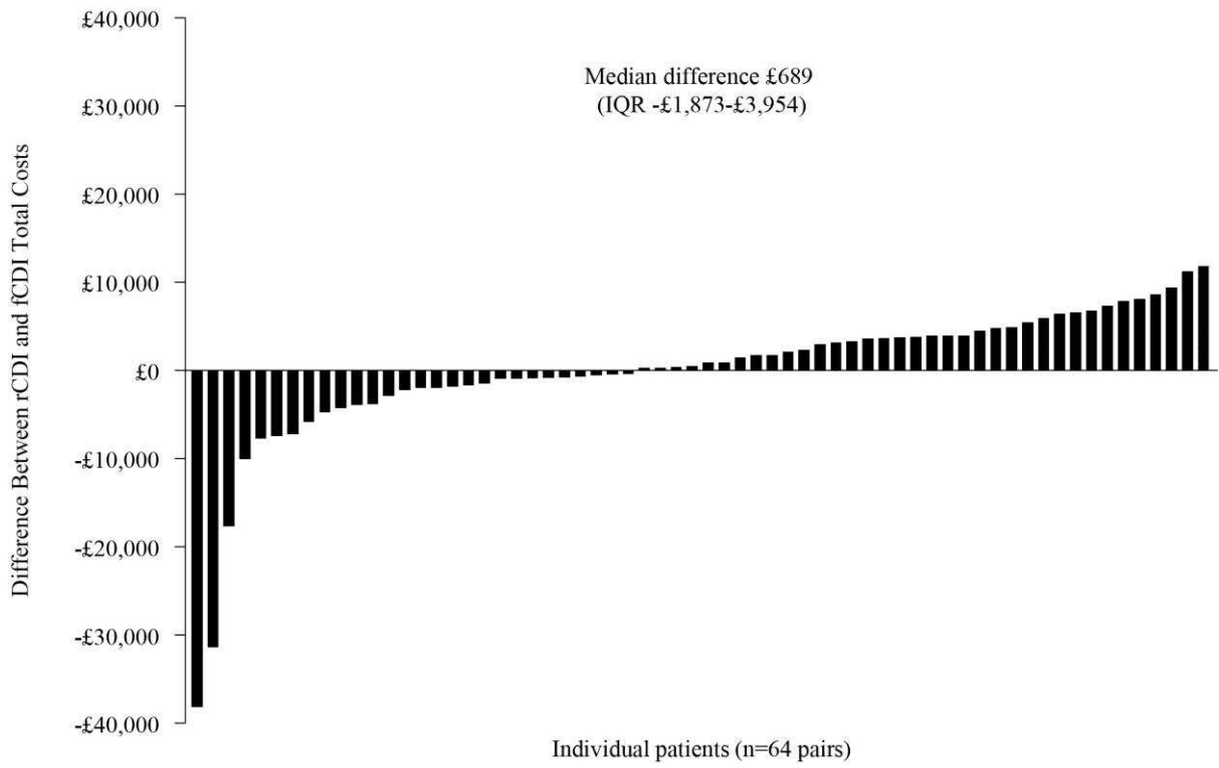
518 **Table 1: Demographic characteristics of patients with rCDI and fCDI**

Characteristic	Patients with recurrent CDI		Patients with first episode only CDI
	First episode	Recurrent episode	First episode
N	64		64
Gender (n, %)			
Male	28 (44%)		28 (44%)
Female	36 (56%)		36 (56%)
Age (years)			
Median	77.0		76.5
Interquartile range	68.5-84.1		66.9-84.1
Co-morbidities (n, %)			
Cardiac disease	18 (28%)		17 (27%)
COPD	19 (30%)		9 (14%)
Hypertension	24 (38%)		19 (30%)
Inflammatory bowel disease	3 (5%)		3 (5%)
Renal disease	5 (8%)		10 (16%)
Alzheimer's/Dementia	9 (14%)		3 (5%)
Atrial fibrillation	6 (9%)		6 (9%)
Osteoarthritis	7 (11%)		6 (9%)
Diabetes	14 (22%)		6 (9%)
Diverticular disease	8 (13%)		9 (14%)
Hypercholesterolemia	1 (2%)		4 (6%)
Hypothyroidism	3 (5%)		2 (3%)
Hyperparathyroidism	1 (2%)		1 (2%)
Cancer	7 (11%)		15 (23%)
Other	56 (88%)		52 (81%)
Setting where CDI acquired			
Community	12 (19%)	4 (6%)	12 (19%)
Healthcare facility	52 (81%)	60 (94%)	52 (81%)
Severity of CDI*			
Mild/moderate	31 (48%)	42 (67%)	38 (59%)
Severe	33 (52%)	21 (33%)	26 (41%)
Strain of CDI			
<i>Hypervirulent ribotypes</i>			
078	6 (9%)	5 (8%)	4 (6%)
027	1 (2%)	1 (2%)	1 (2%)
Other strains (not hypervirulent)	42 (66%)	43 (67%)	44 (69%)
Unassigned/unable to grow	14 (22%)	15 (23%)	10 (16%)
Not done or result unavailable	1 (2%)	0 (0%)	5 (8%)
<i>Abbreviations: CDI, Clostridium difficile infection; COPD, chronic obstructive pulmonary disease</i>			

519 * Unavailable for 1 patient at recurrence; first episode was severe and therefore classified as severe
 520 for subsequent analyses
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523 **Figure 2: Difference in total costs between individual rCDI cases and matched fCDI controls**

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Table 2: Resource utilisation and costs, overall and by CDI severity

	Patients with recurrent CDI			Patients with first episode only CDI		
	Mild/Moderate	Severe	Overall	Mild/Moderate	Severe	Overall
Total costs (post-index period)*						
All patients	n=42 £6,675 (£4,419-£8,960)	n=22 £8,542 (£7,463-£10,352)	n=64 £7,539 (£5,617-£9,730)	n=38 £6,518 (£2,652-£9,086)	n=26 £5,631 (£2,910-£9,453)	n=64 £6,294 (£2,700-£9,216)
Excluding patients who died (both patients in matched pair excluded if one died)	n=35 £6,907 (£5,088-£9,290)	n=17 £9,030 (£7,463-£10,288)	n=52 £7,888 (£6,047-£9,866)	n=33 £6,590 (£2,392-£8,906)	n=19 £6,961 (£4,464-£10,138)	n=52 £6,719 (£3,329-£9,216)
Total costs (between-episode period)*						
All patients	n=40 [‡] £2,683 (£737-£5,351)	n=22 £3,280 (£1,028-£4,159)	n=62 [‡] £2,973 (£778-£4,610)	Not applicable		
Hospital bed days (post-index observation period)*						
All patients	n=42 15.5 (10-27)	n=22 25.5 (21-27)	n=64 21.0 (12-27)	n=38 20.0 (7-27)	n=26 14.5 (8-27)	n=64 15.5 (7-27)
Excluding patients who died (both patients in matched pair excluded if one died)	n=35 18.0 (12-27)	n=17 26.0 (22-27)	n=52 21.0 (13-27)	n=33 21.0 (6-27)	n=19 18.0 (11-27)	n=52 19.5 (7-27)

* Reported as median (IQR) per patient

[‡] Two patients excluded (insufficient information to determine inter-episode period)

Table 3: Breakdown of costs associated with treatment of rCDI and fCDI (post-index observation period)

Breakdown of costs	Patients with recurrent CDI (n=64)*			Patients with first episode only CDI (n=64)*		
	% of total costs	Median	IQR	% of total costs	Median	IQR
Hospital bed days	86.7%	£6,033	£4,002 - £7,767	88.6%	£4,521	£2,240 - £7,767
CDI-specific medicine[‡]	5.4%	£376	£31 - £1,521	0.9%	£46	£2 - £286
Other medicine	2.4%	£170	£57 - £350	2.3%	£119	£58 - £299
Lab costs[†]	4.6%	£319	£190 - £462	5.9%	£304	£142 - £404
Procedures[†]	0.8%	£54	£0 - £204	2.2%	£111	£0 - £277
IV /nutritional support	0.1%	£6	£0 - £24	0.1%	£4	£0 - £28
Outpatient visits	0%	£0	£0-£0	0%	£0	£0 - £0

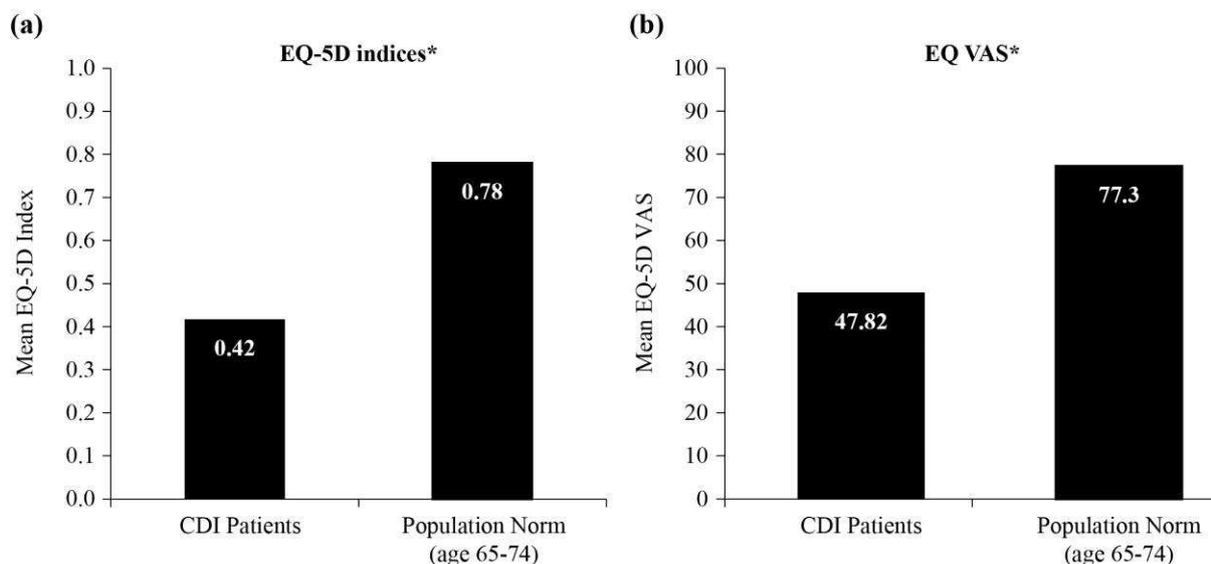
Abbreviations: CDI, Clostridium difficile infection; IV, intravenous; IQR, interquartile range

* Unless otherwise specified

[‡] 23 patients with rCDI were treated with fidaxomicin (median treatment duration 11 days); no patients with fCDI were treated with fidaxomicin

[†] n=54 (one hospital excluded from analysis due to missing data)

Figure 3: EQ-5D of patients hospitalised with CDI compared with UK general population norms for people aged 65-74



* EQ-5D index: maximum score 1 (indicating full health). Lower scores indicate poorer HRQoL; EQ VAS: score range 0-100 (0=Worst imaginable health state, 100=Best imaginable health state)

UK population norms (age 65-74) as published³⁶

Table 4: EQ-5D Dimension scores

EQ-5D Dimension		Patients with CDI n=30	
		n	%
Mobility	No problems (1)	11	37%
	Some problems (2)	11	37%
	Confined to bed (3)	8	27%
Self-care	No problems (1)	14	47%
	Some problems (2)	9	30%
	Unable (3)	7	23%
Usual activities	No problems (1)	6	20%
	Some problems (2)	7	23%
	Unable (3)	17	57%
Pain/discomfort	No (1)	10	33%
	Some (2)	16	53%
	Extreme (3)	4	13%
Anxiety/depression	No (1)	11	37%
	Some (2)	15	50%
	Extreme (3)	4	13%