Infection of hamsters with the UK *Clostridium difficile* ribotype 027 outbreak strain R20291

Anthony M. Buckley, Janice Spencer, Denise Candlish, June J. Irvine and Gillian R. Douce

Institute of Immunology, Infection and Inflammation, Glasgow Biomedical Research Centre, University of Glasgow, Glasgow G12 8QQ, UK

*Clostridium difficile* is the main cause of antibiotic-associated disease, a disease of high socio-economical importance that has recently been compounded by the global spread of the 027 (BI/NAP1/027) ribotype. *C. difficile* cases attributed to ribotype 027 strains have high recurrence rates (up to 36%) and increased disease severity. The hamster model of infection is widely accepted as an appropriate model for studying aspects of *C. difficile* host–pathogen interactions. Using this model we characterized the infection kinetics of the UK 2006 outbreak strain, R20291. Hamsters were orally given a dose of clindamycin, followed 5 days later with 10 000 *C. difficile* spores. All 100% of the hamsters succumbed to infection with a mean time to the clinical end point of 46.7 h. Colonization of the caecum and colon were observed 12 h post-infection reaching a maximum of approximately $3 \times 10^4$ c.f.u. per organ, but spores were not detected until 24 h post-infection. At 36 h post-infection *C. difficile* numbers increased significantly to approximately $6 \times 10^7$ c.f.u. per organ where numbers remained high until the clinical end point. Increasing levels of *in vivo* toxin production coincided with increases in *C. difficile* numbers in organs reaching a maximum at 36 h post-infection in the caecum. Epithelial destruction and polymorphonuclear leukocyte (PMN) recruitment occurred early on during infection (24 h) accumulating as gross microvilli damage, luminal PMN influx, and blood associated with mucosal muscle and microvilli. These data describe the fatal infection kinetics of the clinical UK epidemic *C. difficile* strain R20291 in the hamster infection model.

INTRODUCTION

The Gram-positive, spore-forming bacillus *Clostridium difficile* is a pathogen of nosocomial importance causing a range of symptoms from asymptomatic carriage to severe diarrhoea, pseudomembranous colitis and death. *C. difficile* infection (CDI) typically occurs following antibiotic therapy, where disruption of the gut flora leaves the intestine susceptible to *C. difficile* colonization. Risk factors include the use of proton-pump inhibitors, antiperistaltic drugs, chemotherapy or immunosuppressive agents and old age (>65 years) (Cartman et al., 2010).

In the last decade global incidence of CDI has increased dramatically, in the most part due to the emergence and spread of epidemic strains that are predominantly ribotype 027. Ribotype 027 strains are associated with more severe disease, increased mortality, higher relapse rates and increased resistance to fluoroquinolone antibiotics (Stabler et al., 2009). In 2003 the first outbreak of ‘hypervirulent’ *C. difficile* 027 in the UK was reported, followed by another outbreak in 2005 (HC, 2006). During these two outbreaks, a total of 498 cases were reported with 172 deaths (HC, 2006). Since this first incursion several outbreaks due to ribotype 027 have been documented in almost all UK health board regions.

In 2007 there were 57 255 laboratory-confirmed cases in the UK (55.3% were attributable to ribotype 027), but these numbers have steadily decreased to 28 458 cases in 2009 (36.1% attributable to ribotype 027) [http://www.hpa.org.uk at Topics—Infectious Diseases—Infections A–Z→*Clostridium difficile*→Epidemiological Data *Clostridium difficile*→Voluntary Surveillance of *Clostridium difficile*→Results of the Voluntary Reporting (http://www.hpa.org.uk/web/HPAweb&Page&HPAwebAutoListName/Page/1179745282413)]. The total identifiable cost of CDI was estimated to be £4000 per case in England in 1996 (Wilcox et al., 1996). On this basis such infections cost the UK £113 million in 2009 and £229 million in 2007.

The golden Syrian hamster is regarded as an important model of *C. difficile* disease as many of the clinical symptoms in humans can be observed in this model. After clindamycin administration and challenge with toxigenic *C. difficile* strains, hamsters develop a haemorrhagic caecitis, which
manifests as 'wet tail' (a sign of diarrhoea in hamsters) and is followed by death (Bartlett et al., 1977; Sambol et al., 2001; Razaq et al., 2007). Using this model we characterized the phenotypic outcome of infection with the clinical C. difficile ribotype 027 strain R20291. This strain was originally isolated from a patient who presented with pseudomembranous colitis from the first UK outbreak in 2006 and is considered an archetypal 027 strain within the UK.

METHODS

**Bacterial strain and spore preparation.** C. difficile strain R20291 was obtained from Professor Brendan Wren (London School of Hygiene and Tropical Medicine, London, UK) and is classed as ribotype 027 and toxinoype III (Rupnik et al., 1998). Bacteria were grown on CCEY agar supplemented with cefoxitin–cycloserine, egg yolk emulsion (LabM) and erythromycin (100 μg ml⁻¹) at 37 °C under anaerobic conditions. Spores were made according to Lawley et al. (2009) except strains were grown in brain heart infusion broth. Serial 10-fold dilutions of the spore preparation were inoculated onto supplemented CCEY agar plates to determine the number of cfu ml⁻¹, from which the inocula dilution was calculated.

**Animal experiments.** Female golden Syrian hamsters that weighed approximately 100 g were purchased from Harlan Olac, UK. These animals were housed individually, and given water and food ad libitum. Telemetry chips (VitalView Emitter) were inserted by laparotomy into the body cavities of the animals at least 3 weeks before infection with C. difficile. Once the wounds healed, the animals were placed on receiver pads, and their body temperature and activity were monitored (VitalView software). Each animal received orogastriically 30 mg clindamycin phosphate kg⁻¹ 5 days before infection. Following administration of the antibiotic, all animals were placed in individual sterilized isolator cages with disposable filter top lids, and were provided with sterile food, water and bedding. Each hamster received approximately 10,000 spores of strain R20291 (determined by serial dilutions before and after challenge, as described above). Experiments using 1, 3, 5 or 10 days post-clindamycin treatment and infection of hamsters with 100–100,000 spores have been tested previously; however, only the combination described above resulted in 100 % infection of the animals. Following infection, each animal was placed in a second sterile cage with appropriate food, water and bedding. At 12 h (n=6), 24 h (n=6) and 36 h post-challenge (n=6), and when animals succumbed to infection (n=13), animals were culled for post-mortem analysis. At each time point, to establish the level of colonization, the caecum and colon were removed aseptically. To enumerate the total bacterial load (spores and vegetative cells), each section was opened longitudinally, and the contents were removed by gentle washing in 10 ml PBS (luminal-associated bacteria). The tissues were washed in 10 ml PBS and homogenized in 5 ml PBS for 1 min using a stomacher (tissue-associated bacteria), and viable counts were determined for the homogenates. Serial 10-fold dilutions were plated on supplemented CCEY agar plates. To determine the numbers of spores present in the samples, the samples were heated for 10 min at 56 °C, and the numbers of spores present were determined by the viable count method as described above. Results are given as the mean number of recovered bacteria in at least six animals per time point.

**Multilocus variable-number tandem-repeat analysis (MLVA).** To confirm the C. difficile isolated at post-mortem were the same strain originally used for infection, bacteria were isolated as single colonies, grown and the DNA isolated. PCR amplification of seven C. difficile repeat loci was performed using the primers and PCR amplification protocol described by Marsh et al. (2006). Following agarose gel electrophoresis MLVA patterns from spore inocula and recovered isolates were compared.

**Detection of in vivo toxin levels.** Production of toxins by R20291 was detected in vitro using filtered caecum and colon content from animals that was taken during the post-mortem. Confluent monolayers of Vero cells (kidney epithelial cells) were maintained in Eagle's minimal essential medium (EMEM; Sigma) supplemented with 1 % heat-inactivated fetal calf serum (Invitrogen), 2 mM l-glutamine (Invitrogen) and 100 μg penicillin/streptomycin ml⁻¹. Cells were washed with preheated sterile PBS before addition of serial diluted filtered gut content in supplemented EMEM and incubated for 18 h at 37 °C (5 % CO₂). Cells were washed with PBS, fixed in 1 % formalin for 10 min then washed again. Fixed adherent cells were stained with Giemsa for 30 min then washed before addition of 0.1 % SDS and left for 1 h. Optical density was measured using an EL 808 Ultra microplate reader (Bio-Tek Instruments) at 600 nm and compared to non-infected hamster caecal and colon gut contents as a negative control. If the toxin dilution is able to cause cell toxicity (cell rounding) this leads to the loss of cell adherence resulting in a reduced staining and hence optical density of the contents of the wells.

**Histology.** Samples from different regions of the gut were prepared for simple histology as described by Goulding et al. (2009).

**Scanning electron microscopy (SEM).** SEM samples were prepared as described by Goulding et al. (2009).

**Statistical analysis.** All statistical analyses were performed using the GraphPad Prism 5 (GraphPad Prism Software). A Student’s one-tailed t-test was used to determine significant differences in bacterial recoveries between all time points examined. P values ≤ 0.05 were considered significant.

**RESULTS**

**Telemetry monitoring of infected hamsters**

If infection was allowed to progress to the clinical end point then the infection of hamsters with R20291 resulted in 100 % mortality. Hamsters typically followed the temperature profile shown in Fig. 1(a), where a rapid decrease in body temperature was observed following onset of clinical signs that included a wet tail (diarrhoea; represented by the bar in Fig. 1a). The drop in body temperature from 37 °C to the clinical end point (35 °C) took a mean time of 76 ± 6.4 min SEM. In comparison hamsters infected with C. difficile strain 630 displayed a significantly longer body temperature drop time (145 ± 28.9 min; P=0.02). The mean time taken for hamsters to succumb to infection was 46.7 ± 2.8 h SEM; however, a range between 35.2 and 62.3 h was observed (n=13) (Fig. 1b). No significant difference was seen compared to hamsters infected with C. difficile strain 630 (43.7 ± 3.7 h SEM; n=11).

**Colonization kinetics of R20291**

To assess the colonization kinetics of C. difficile R20291 after inoculation, hamsters were culled at 12, 24 and 36 h...
post-challenge and at the clinical end point to measure the levels of R20291 in the caecum and colon. At 12 h after infection with $10^4$ spores per animal, *C. difficile* caecum levels reached approximately $3 \times 10^4$ c.f.u. per organ with higher bacterial levels associated with the lumen compared to those associated with the tissue (Fig. 2a). Levels in the colon were lower than in the caecum, reaching approximately $1 \times 10^4$ c.f.u. per organ. No *C. difficile* spores were detected in either the lumen or the tissue at this time. At 24 h post-challenge *C. difficile* levels increased in the caecum and colon (approximately $3 \times 10^5$ and $6 \times 10^4$ c.f.u. per organ, respectively) compared to 12 h, where more bacteria were associated with the lumen than tissue ($P=0.03$ caecum only; Fig. 2b). *C. difficile* spores were present in low levels in the caecum (approximately $5 \times 10^3$ c.f.u. per organ) but could not be detected in the colon. The rate of growth increased exponentially over the next 12 h, whereby at 36 h post-challenge mean caecal and colon levels were at approximately $5 \times 10^7$, $3 \times 10^6$, $1 \times 10^7$ and $3 \times 10^5$ c.f.u. per organ for caecum luminal associated (CAE-LA), caecum tissue associated (CAE-TA), colon luminal associated (COL-LA) and colon tissue associated (COL-TA) samples, respectively (Fig. 2c). Over the same 12 h period a significant increase in spores was observed at all sites sampled, with the highest levels observed at 36 h post-challenge in the lumen [60 (CAE-LA), 10 (CAE-TA), 80 (COL-LA) and 20% (COL-TA) of total bacteria isolated, $P=0.01$, 0.03, 0.04 and 0.06, respectively]. At the experimental end point (~46.7 h) the number of bacteria isolated from each site remained high and comparable to those observed at 36 h post-challenge (approximately $6 \times 10^7$, $3 \times 10^6$, $2 \times 10^7$ and $6 \times 10^5$ c.f.u. per organ for CAE-LA, CAE-TA, COL-LA and COL-TA samples, respectively; Fig. 2d). Compared to 36 h no significant differences in spore levels were seen at the clinical end point.

**MLVA**

MLVA patterns for *C. difficile* isolated at the post-mortem analysis matched those MLVA patterns for the inoculating bacterial spores (data not shown).

**In vivo toxin levels**

*In vivo* toxin production was measured semiquantitatively *in vitro* using a Vero cell toxicity assay by serially diluting filtered luminal contents and calculating the maximum fold-dilution at which cell toxicity was still detected (cell rounding). Caecal and colon contents from animals culled at 12 and 24 h showed similar levels of toxin, approximate maximum 10- and 5-fold dilution required for cell rounding (caecum and colon samples, respectively; Table 1). In contrast, luminal contents taken at 36 h and at the clinical end point showed increased toxin abundance, approximate maximum 100000- and 625-fold dilution required for cell rounding (caecum and colon samples, respectively; Table 1).

**Histological changes**

Gut pathology at 12, 24 and 36 h post-infection, and at the clinical end point was assessed by staining sections of gut tissue with haematoxylin and eosin, and comparing to uninfected controls. At 12 h post-infection, caecal and colon samples showed no change from uninfected controls (data not shown). By 24 h after infection an increase in polymorphonuclear leukocytes (PMNs) in the submucosa and microvilli of the caecum was seen along with the destruction of the greater part of the epithelial brush border exposing the circulating PMNs (Fig. 3b, indicated by arrows). Caecal and colon (data not shown) histology samples at 36 h and at the clinical end point showed severe destruction of the epithelial layer and extensive PMN influx in the lumen through ‘volcanic-like’ eruptions on the microvilli (Fig. 3c, d). A high association of red blood cells with the mucosal muscle and amongst the microvilli was observed, which is an indication of rapid and extensive damage presumably caused by the *C. difficile* toxins.
The detailed characterization of host–bacterial interactions was confirmed by SEM on caecum and colon samples taken at 12, 24 and 36 h post-infection and at the time of culling. At 12 h post-infection individual *Clostridium*-like bacterial cells could be seen sporadically on caecal surfaces. By 24 h post-infection a greater number of *Clostridium*-like bacteria were seen associated with mucus over both the tissue surface and crypt crevasses (Fig. 4a). During the height of bacterial recovery from the caecum and colon (36 h post-infection), aggregates of *Clostridium*-like bacteria could be seen on the tissue surface forming microcolony plaques typically associated over the crypt crevasses (Fig. 4b). Intimate tissue–bacterial and bacterial–bacterial interactions could be seen via flagella-like appendages; however, these structures could potentially be host derived (Fig. 4, indicated by filled arrows). In agreement with histology observations red blood cells were also observed in the same vicinity as *Clostridium*-like bacteria (Fig. 4c, indicated by open arrow).

**DISCUSSION**

Here we present a detailed study of the colonization kinetics of the UK *C. difficile* outbreak strain R20291 in
hamsters. After infection with R20291 100% of the animals succumbed to the infection, although the times to the clinical end point were spread over 20 h. The morbidity and mortality we observed are consistent with those seen with infection with high virulence strains in humans (Warny et al., 2005). The diversity in times to the clinical end point could reflect the sensitivity of R20291 to the clindamycin used in the hamster model (MIC = 16 μg ml⁻¹). These results are similar to those observed by Razaq et al. (2007) where infection with a clindamycin-resistant 027 ribotype isolate killed hamsters more quickly (40 h) compared to clindamycin-susceptible isolates (48 and 69 h). However, the mean time to the clinical end point seen with R20291 is very similar to ribotype 027 strains isolated in North America (Razaq et al., 2007). Observed clinical-end points for C. difficile strain 630 (ribotype 012) were very similar to R20291; however, R20291 displayed a quicker temperature drop time compared to strain 630. This rapid temperature drop could be an indicator of enhanced virulence of these strains, an observation shared by Razaq et al. (2007), where strain 630 displayed a significantly longer time from C. difficile colonization to death compared to clindamycin-resistant 027 isolates.

Caecal and colon contents at 12 and 24 h post-infection showed lower recovery of C. difficile compared to clindamycin-resistant isolates, such as C. difficile M68 (data not shown), which suggests antibiotic killing of C. difficile by residual clindamycin levels. Larson & Borriello (1990) found that after a single 3 mg clindamycin dose C. difficile could still be detected in the caecum of hamsters after 11 days, which was enough to inhibit the growth of a clindamycin-susceptible strain. In this study treatment of hamsters with clindamycin was optimized to 5 days prior to challenge to give 100% infection (resulting in 100% mortality). Once gut colonization was established (24 h post-infection) rapid increases in bacterial numbers over a 12 h time period were observed (>2 log increase). This was quickly followed by morbidity, loss of body temperature and death (mean time to the clinical end point was 46.7 h). Razaq et al. (2007) reported mean times of colonization to death of between 7 and 16 h with other ribotype 027

![Fig. 3. Caecal histology of hamsters infected with R20291. Samples were taken from uninfected controls (a), at 24 h post-infection (b) and at the clinical end point [(c) and inset (d)]. Arrows denote epithelial brush broader damage.](image)
isolates; however, in this case, *C. difficile* faecal shedding was used as an indication of colonization. It is currently unknown whether colonization of the hamster gut is needed to cause pathology and disease (or mortality). Using the early time points in infection we would like to be able to investigate possible colonization factors of *C. difficile* and compare mutants defective in such factors.

Histology samples at 24 h post-infection show some pathology mainly to the epithelial brush border even though very little toxin activity could be measured using a Vero cell toxicity assay. Rolfe & Song (1993) showed that carbohydrates from brush border membranes of hamster ileum can bind toxin A. It is hypothesized that TcdA acts to disrupt the epithelial layer allowing access of TcdB to its receptor on the basolateral membrane (Stubbe et al., 2000). Potentially this could be reflected *in vivo* by variation in the temporal expression of these two genes (this is currently being investigated). Interestingly gut damage at 24 h (presumably due to toxin production) occurred at the same time as sporation. Whether *in vivo* toxin production and sporation are linked is unclear; however, Underwood et al. (2009) reported that disruption of the sporation initiation pathway (via spo0A inactivation) also resulted in decreased toxin production in *C. difficile* strain 630.

Most pathological damage was seen at 36 h post-infection and onwards, and this was associated with increased bacterial numbers and greater toxin production. *C. difficile* harbours a putative quorum-sensing pathway, akin to *Vibrio harveyi*, that may control toxin production *in vivo* through cell-to-cell signalling; although, Carter et al. (2005) showed that exogenous AI-2 did not affect toxin production. Studies have shown that intestinal damage occurs mostly through the actions of TcdA/B; however, *C. difficile* R20291 also produces the binary toxin CDT. CDT has been shown to cause fluid accumulation in a rabbit ileal loop model but not to cause disease in hamsters (Geric et al., 2006). Through the induction of microtubule-based protrusions, CDT can enhance the adherence of *C. difficile* bacteria to host cells (Schwan et al., 2009). The activity of CDT was not measured in this study and could play an adjunctive role to TcdA/B.

CDIs exert a substantial burden on human health and economy, and there is a pressing need for vaccines and/or treatment. As infection of the hamster with *C. difficile* mirrors many symptoms of CDI in humans this model is ideally placed to test such interventions. *C. difficile* host–pathogen kinetics are essential to determine the impact of such interventions, here we detail the infection process of a clinically relevant *C. difficile* strain, R20291. In conclusion, the epidemic *C. difficile* strain R20291 caused rapid death in the hamster model of infection, where high bacterial (including spores) and toxin levels caused significant tissue damage in the caecum and colon.

**Fig. 4.** SEM of hamsters infected with R20291. Samples were taken from the caecum at 24 h post-infection (a), and from the caecum (b) and colon (c) at 36 h post-infection. Bacteria appeared to be interacting with the tissue surface and other bacteria via flagella-like appendages (filled arrows). In agreement with histology observations red blood cells were also observed in the same vicinity as bacteria (open arrow).
ACKNOWLEDGEMENTS

We acknowledge the help of M. Mullin (Integrated Microscopy, University of Glasgow) and H. Thompson (Division of Pathological Sciences, University of Glasgow Veterinary School, Bearsden, Glasgow) for SEM sample preparation and examination of histological samples by light microscopy. The authors gratefully acknowledge the support of the Wellcome Trust (grant number 086418 awarded to G. R. D).

REFERENCES


