Primary Outcomes From a Phase 3, Randomized, Double-Blind, Active-Controlled Trial of Surotomycin in Subjects With Clostridium difficile Infection

Vicente Boix,1 Richard N. Fedorak,2 Kathleen M. Mullen,3 Yves Pesant,4 Uschi Stoutenburgh,5 Mandy Jin,5 Adedayo Adedoyin,5
Laurent Chesnel,3 Dalya Guris,6 Kajal B. Larson,6 and Yoshihiko Murata6

1Unit of Infectious Diseases, Hospital General Universitario de Alicante, Instituto de Investigación Sanitaria de Alicante (ISABIAL) - Fomento de la Investigación Sanitaria y Biomédica de la Comunitat Valenciana (FISABIO), Spain; 2Division of Gastroenterology, University of Alberta, Edmonton, Canada; 3Department of Medicine, Section of Infectious Diseases and Global Health, University of Chicago, Illinois; 4St-Jerome Medical Research Inc., Saint-Jérôme, Quebec, Canada; and 5Merck & Co., Inc., Kenilworth, New Jersey

**Background.** Although the incidence of *Clostridium difficile* infection (CDI) is increasing, available CDI treatment options are limited in terms of sustained response after treatment. This phase 3 trial assessed the efficacy and safety of surotomycin, a novel bactericidal cyclic lipopeptide, versus oral vancomycin in subjects with CDI.

**Methods.** In this randomized, double-blind, active-controlled, multicenter, international trial, subjects with CDI confirmed by a positive toxin result were randomized to receive surotomycin (250 mg twice daily) or vancomycin (125 mg 4 times daily) orally for 10 days. The primary endpoints were clinical response at end of treatment and evaluation of surotomycin safety. The key secondary endpoints were clinical response over time and sustained clinical response through a 30- to 40-day follow-up period. *Clostridium difficile* infection recurrence during follow-up and time to diarrhea resolution were also analyzed.

**Results.** In total, 570 subjects were randomized and had confirmed CDI; 290 subjects received surotomycin and 280 subjects received vancomycin. Surotomycin clinical cure rates at end of treatment (surotomycin/vancomycin: 79.0%/83.6%; difference of −4.6%; 95% confidence interval, −11.0 to 1.9), clinical response over time (stratified log-rank test, P = .832), and sustained clinical response at end of trial (Day 40–50) (60.6%/61.4%; difference of −0.8%; 95% CI, −8.8 to 7.1) in the microbiological modified intent to treat population did not meet noninferiority or superiority criteria versus vancomycin. Both treatments were generally well tolerated.

**Conclusions.** Surotomycin failed to meet the criteria for noninferiority versus vancomycin for the primary and key secondary endpoints in this trial.

**Keywords.** antibiotic; *Clostridium difficile* infection; surotomycin; vancomycin.

*Clostridium difficile* infection (CDI) remains a leading cause of hospital-acquired diarrhea [1, 2]. The incidence of CDI has increased over the last decade in the United States from 4.5 to 8.2 incidents per 1000 hospital discharges [1], and epidemic strains, such as the North American pulsed-field type 1/restriction endonuclease analysis type B1/ribotype 027 (B1/NAP1/027) strain, have emerged [3, 4]. Furthermore, in a large US study, CDI recurrence occurred in ~14%–21% of patients after treatment of the initial CDI episode [5]. As a consequence of the incidence and impact of CDI, there remains a significant unmet need for additional CDI therapies, particularly those able to decrease posttherapy recurrence rates and increase sustained response.

Surotomycin ([SUR] CB-183,315; MK-4261) is a novel cyclic lipopeptide in development for the treatment of CDI. In phase 1 clinical trials, SUR ≤1000 mg twice daily (BID) had only modest disruptive effects on healthy gut microbiota, sparing the natural barrier to *C difficile* colonization [6]. In phase 2 clinical trials, CDI cure rates were similar for SUR 125 mg BID, 250 mg BID, and oral vancomycin (VAN) 125 mg 4 times daily (QID). *Clostridium difficile* infection recurrence rates were significantly lower after SUR 250 mg BID compared with VAN 125 mg QID [7]. The main objectives of this phase 3 clinical trial were to demonstrate the safety and efficacy of SUR 250 mg BID versus VAN 125 mg QID in adults with CDI.

**MATERIAL AND METHODS**

**Trial Design**

This randomized, double-blind, active-controlled trial, consisting of 2 treatment arms (protocol MK-4261-005; ClinicalTrials.gov identifier NCT01597505), was conducted between July 28, 2012 and March 20, 2015 in accordance with the Declaration of Helsinki, Good Clinical Practice Guidelines, and the approval of local institutional review boards. All subjects provided written informed consent. The trial consisted of a 10-day treatment...
period (Day 1–10) and follow-up visits 2 days after end of treatment ([EOT] Day 10–13), Day 24 ± 3 days, and Day 40–50. During the treatment period, subjects received oral SUR 250 mg BID with alternate dummy tablets or the active comparator, oral VAN 125 mg QID. The primary endpoint of the trial was clinical response at EOT. Key secondary endpoints were clinical response over time and sustained clinical response 30–40 days post-EOT.

**Trial Population**

Subjects were enrolled at 115 sites in North America (51 sites), Europe (62 sites), and the Middle East (2 sites). Eligible subjects were ≥18 and <90 years of age, had diarrhea with a minimum of 3 unformed bowel movements (UBMs), or >200 mL volume of stool (ie, for those with a collection device) over a 24-hour period before randomization/treatment administration. Furthermore, subject stool samples must have been *C. difficile* toxin-positive within 48 hours of treatment administration, as identified by enzyme immunoassay (EIA), polymerase chain reaction (PCR), or cell culture cytotoxin neutralization assay. Subjects with toxic megacolon and/or small bowel ileus or those who had received a fecal transplant or treatment for the current episode of CDI were excluded. Other exclusion criteria included subjects with >2 episodes of CDI within 90 days of trial therapy, a history of inflammatory bowel disease (eg, ulcerative colitis, Crohn’s disease, or microscopic colitis), a positive stool culture for other enteropathogens, life-threatening illness at the time of enrollment as measured by a score of 4 using a modified Horn’s index [8], or a life expectancy of <8 weeks.

**Randomization and Treatment**

Subjects were randomly assigned to treatment arms (1:1; Figure 1) based on a centralized computer-generated randomization schedule and stratified by age (<75 or ≥75 years of age) and number of prior CDI episodes (0 or ≥1) in the last 90 days. Subjects, trial staff, and the sponsor remained blinded until the database was locked. An interactive voice/web response system was used for allocation of identification numbers and treatment kits.

Either SUR 250 mg BID or VAN 125 mg QID was administered according to the randomized allocation. In the SUR treatment arm, additional placebo dummy treatments were included to match the QID dosing regimen of VAN. Commercially available oral VAN tablets were backfilled with microcrystalline cellulose as a filler to prevent rattling and maintain the blind. During the 10-day treatment period, a single capsule was administered at breakfast, lunch, dinner, and bedtime.

**Clinical Outcomes**

All subjects were evaluated for clinical response 2 days after the last dose of study drug, and all subjects continued to be assessed throughout the course of the study (ie, through the late follow-up visit, on Day 40–50). A favorable clinical response (cure) was defined as resolution of diarrhea (ie, ≤2 loose stools per 24 hours for 2 consecutive days) and no need for additional CDI treatment after the trial treatment period. If the treatment did not achieve a favorable clinical response, the outcome was deemed a failure. After initial cure, sustained clinical response was identified if there was no recurrence of CDI before the final follow-up visit (Day 40–50). Recurrence was identified by a minimum of 3 UBMs over a 24-hour period (or >200 mL stool volume over 24 hours—for those with a collection device) and positive detection of toxigenic *C. difficile* from a stool sample.

**Microbiology**

Baseline and recurrent diarrhea stool samples were analyzed onsite for the detection of *C. difficile* toxin. Stool samples were also analyzed for the presence of B1/NAP1/027-positive epidemic strains of *C. difficile*.

**Pharmacokinetic Evaluations**

A selection of sites participated in an intensive pharmacokinetic (PK) substudy. At these sites, intensive blood and stool sampling occurred on Day 5–7 over a dosing interval. Plasma and fecal SUR and VAN concentrations were determined by a validated liquid chromatography with tandem mass spectrometry method (Tandem Laboratories, West Trenton, NJ and Salt Lake City, UT).

**Safety Assessments**

Adverse events (AEs), defined by the protocol as nonserious, were collected from first dose of trial treatment until 7 days...
after the last dose. Serious AEs (SAEs) were collected from the first dose until 30 days after the last dose. Vital signs (eg, heart rate, blood pressure, and temperature), physical examination, electrocardiogram, and laboratory safety tests were measured throughout the trial. The safety population consisted of all randomized subjects who received any amount of study drug.

Data Analysis and Statistics
The 95%–95% fixed margin approach was used to justify the noninferiority based on a meta-analysis of the medical literature. The selected clinically relevant noninferiority margin of 10% was demonstrated to preserve 51.7% of the active control effect. Assuming an 83% clinical response rate for both treatment arms, 258 subjects per arm would have at least 85% power to demonstrate SUR noninferiority versus VAN based on the 10% noninferiority margin and at a 1-sided significance interval of 0.025. Given that an expected 85% of randomized subjects would meet criteria to be included in the microbiological modified intent-to-treat population (mMITT) all randomized subjects with a confirmed diagnosis of CDI, a total of 304 subjects per arm were enrolled.

Noninferiority and superiority tests were conducted for clinical outcomes in the mMITT population. Two-sided 95% confidence intervals (CIs) were calculated for favorable clinical outcomes associated with the treatments. Noninferiority was identified when the difference between SUR and VAN arms 95% CI was ≥−10% and superior when the difference was >0%. Analyses were stratified by age (<75 or ≥75 years of age) and previous episodes of CDI in the last 90 days (0 or ≥1). The Kaplan-Meier method with a stratified log-rank P value was used to determine clinical response over time. All statistical tests were 2-sided and conducted at the 0.05 significance level. Descriptive statistics were provided for continuous data. The PK analysis population consisted of subjects who provided at least 3 serial plasma samples postdosing on Day 5–7. Individual plasma PK parameters were determined by noncompartmental analysis using WinNonlin (version 6.3 or higher; Pharsight Corporation, Mountain View, CA).

RESULTS

Trial Population
A total of 606 subjects were enrolled and randomized, 570 of whom had confirmed CDI (ie, 290 in the SUR arm and 280 in the VAN arm) and were included in the mMITT population (Figure 2). The baseline and demographic characteristics of the mMITT population are shown in Table 1. The characteristics of the treatment groups were similar. Study drug completion rates in the mMITT population were 82.8% and 87.1% for SUR and VAN, respectively.

Efficacy
Clinical cure at EOT (Day 1–13) was observed in 79.0% (229 of 290) of subjects after SUR treatment and 83.6% (234 of 280) of subjects after VAN. The primary noninferiority endpoint, clinical cure at EOT, was not met. The difference between clinical response rates (SUR–VAN) was −4.6% (95% CI, −11.0 to 1.9) (Figure 3A). Neither of the 2 key secondary endpoints were met; SUR did not demonstrate superiority over VAN for the sustained clinical response at the end of the trial (SUR 60.6% vs VAN 61.4%; difference: −0.8%; 95% CI, −8.8 to 7.1) (Figure 3B) or for clinical response over time (defined as clinical response through EOT and the sustained clinical response from EOT to Day 40; stratified log-rank test P = .832) (Figure 3C).

Recurrence (ie, subjects who were cured and had a CDI recurrence or were lost to follow-up, died, or had their last follow-up contact before Day 40) occurred in 53 subjects (17.7%) in the SUR arm and in 63 subjects (21.2%) in the VAN arm (difference −3.5%; 95% CI, −10.0 to 3.0). A Kaplan-Meier analysis

Figure 2. Subject flow diagram. CDI, Clostridium difficile infection; mMITT, microbiological modified intent to treat.
demonstrated that time to resolution of diarrhea (ie, ≤2 UBM per 24-hour period) was similar between groups (SUR vs VAN; \( P = .431 \)). Although the Kaplan-Meier analysis of time to diarrhea reappearance (ie, ≥3 UBM per 24-hour period) in subjects who had a clinical response with SUR versus those with VAN initially supported SUR superiority (\( P = .011 \)), when adjusted for multiplicity, this efficacy endpoint was not met. In subjects infected with the \( C. difficile \) BI/NAP1/027 strain at baseline, the cure rate and sustained clinical response rate were numerically higher and recurrence rates were lower with SUR versus VAN (Figure 4A). However, none of these comparisons were statistically significant. In subjects infected at baseline with a non-BI/NAP1/027 strain, cure at the EOT and sustained clinical response were lower and recurrence was higher with SUR compared with VAN, and the upper limit of the 95% CI was <0 (Figure 4B). Response rates at EOT were numerically lower for both treatments in the population identified as toxin positive by EIA (cure rates: SUR, 77.2% [95% CI, 69.1%–83.8%]; VAN, 79.8% [95% CI, 71.3%–86.3%]) versus PCR (cure rates: SUR, 80.2% [95% CI, 73.4%–85.6%]; VAN, 86.0% [95% CI, 80.0%–90.4%]).

Pharmacokinetics of Surotomycin
A total of 18 subjects receiving SUR from selected clinical sites were included in the PK substudy population. The median (range) fecal concentration of SUR was 1216 µg/g (range, 0–3780 µg/g). Most subjects had SUR fecal concentrations greater than the minimum inhibitory concentration of 1 µg/mL at which the growth of 90% of the \( C. difficile \) is inhibited, suggesting that adequate therapeutic fecal SUR concentrations were achieved \([9]\). Plasma PK parameters of SUR (\( n = 18 \) subjects) and VAN (\( n = 6 \)) were also calculated and showed rapid

| Table 1. Trial Population Demographic and Baseline Characteristics<sup>a</sup> |
|-----------------|-----------------|-----------------|
| Characteristic | Surotomycin (N = 290) | Vancomycin (N = 280) | Total (N = 570) |
| Female, n (%)   | 117 (40.3)       | 114 (40.7)       | 231 (40.5)       |
| Race, n (%)     |                 |                 |                 |
| Black or African American | 15 (5.2)       | 22 (7.9)       | 37 (6.5)       |
| White           | 260 (89.7)      | 248 (88.6)      | 508 (89.1)      |
| Ethnicity, n (%)|                 |                 |                 |
| Hispanic or Latino | 18 (6.2)       | 9 (3.2)        | 27 (4.7)        |
| Not Hispanic or Latino | 266 (91.7)   | 264 (94.3)     | 530 (93.0)      |
| Age at first dose (years) |                   |                 |                 |
| Mean (SD)       | 61.1 (17.6)     | 61.5 (18.4)     | 61.3 (18.0)     |
| Median (range)  | 64.0 (18–89)    | 64.5 (18–89)    | 64.0 (18–89)    |
| <75 years, n (%)| 211 (72.8)      | 200 (71.4)      | 411 (72.1)      |
| Body mass index (kg/m²) |               |                 |                 |
| Mean (SD)       | 26.4 (7.2)      | 27.5 (6.7)      | 26.9 (7.0)      |
| Subject’s hospitalization status at baseline, n (%) |                   |                 |                 |
| Inpatient       | 178 (61.4)      | 181 (64.6)      | 359 (63.0)      |
| Outpatient      | 108 (37.2)      | 90 (32.1)       | 198 (34.7)      |
| ICU status at baseline, n (%) |               |                 |                 |
| Yes             | 11 (3.8)        | 8 (2.9)         | 19 (3.3)        |
| CDI Epidemiologic Classification, n (%) |                   |                 |                 |
| 1 – HC-Fenst, HCF-associated CDI | 92 (31.7)      | 91 (32.5)      | 183 (32.1)      |
| 2 – Community-onset, HCF-associated CDI | 53 (18.3)      | 53 (18.9)      | 106 (18.6)      |
| 3 – Community-associated CDI | 121 (41.7)     | 103 (36.8)     | 224 (39.3)      |
| 4 – Indeterminate disease | 24 (8.3)       | 33 (11.8)      | 57 (10.0)       |
| Severe disease, n (%) |               |                 |                 |
| ESCMID Comprehensive Criteria | 215 (74.1)    | 209 (74.6)     | 424 (74.4)      |
| ESCMID Abbreviated Criteria | 1 (0.3)        | 0              | 1 (0.2)         |
| IDSA Criteria   | 9 (3.1)         | 15 (5.4)       | 24 (4.2)        |
| UBM and WBC Criteria | 95 (32.9)      | 96 (34.3)      | 191 (33.6)      |
| Horn’s Index    | 47 (16.4)       | 45 (16.3)      | 92 (16.4)       |
| ≥1 previous episode of CDI | 49 (17.1)      | 51 (18.5)      | 100 (17.8)      |
| BI/NAP1/027 strain-positive | 59 (23.1)     | 67 (27.2)      | 126 (25.1)      |
| Mean number of UBM(s) at baseline (SD) | 7.1 (4.6)     | 6.6 (4.3)      | 6.9 (4.5)       |

Abbreviations: CDI, Clostridium difficile infection; ESCMID, European Society of Clinical Microbiology and Infectious Diseases; HCF, healthcare facility; ICU, intensive care unit; IDSA, Infectious Diseases Society of America; SD, standard deviation; UBM, unformed bowel movements; WBC, white blood cells.

<sup>a</sup>Baseline values were taken as the last nonmissing result before first administration of the study drug. Therefore, some numbers do not represent the total number of initially enrolled subjects.
SUR absorption with median time to maximum concentration of 1 hour postdose with detectable plasma concentrations until ~12 hours after dosing (Supplementary Table 1).

Safety

Treatment-emergent AEs (TEAEs) were experienced by 148 (48.5%) of SUR-treated subjects and 158 (55.2%) of VAN-treated subjects. The proportion of subjects with at least 1 AE, study drug-related AEs, or at least 1 SAE was generally similar between treatments (Table 2). All except 2 SAEs (perforated jejunal ulcer and cardiac arrest with ventricular fibrillation; both in the SUR group) were considered unrelated to the study drug. The percentage of subjects with AEs leading to discontinuation or death was numerically higher in the SUR arm compared with VAN; however, none of the deaths were considered treatment-related. The majority of TEAEs in both treatment arms (SUR, 75%; VAN, 86%) were mild or moderate in intensity.

The highest frequency of AE (System Organ Class) was in gastrointestinal disorders followed by infections and infestations, the majority of which were urinary tract infections (Table 3).

The most common TEAEs (Preferred Term) that were deemed drug-related by the investigator were nausea (SUR, 2.0%; VAN, 1.7%), vomiting (SUR, 1.3%; VAN, 0%), and increased alanine aminotransferase (ALT) (SUR, 1.0%; VAN, 2.1%).

Nine subjects experienced protocol-specified Closely Monitored Events: 1 subject in the SUR arm with creatine phosphokinase (CPK) levels >1000 U/L post-baseline; 1 subject in the VAN arm with ALT or aspartate aminotransferase (AST) >8 x upper limit of normal (ULN); 2 subjects in each treatment arm with ALT or AST >3 × ULN and total bilirubin >2 × ULN; and 5 subjects (SUR arm: 2, VAN arm: 3, of which 1 subject in each group had concurrent total bilirubin >2 × ULN, as described above) demonstrated ALT or AST >3 × ULN with concurrent fatigue, nausea, vomiting, right upper-quadrant pain or tenderness, fever, rash, or eosinophilia. No individual or population trends in ALT, AST, total bilirubin, or CPK levels were noted.

None of the 4 subjects with ALT or AST >3 × ULN and total bilirubin >2 × ULN were deemed related to the study drug or were consistent with fulfilling Hy's Law [10].
There were no clinically relevant trends in mean change from baseline for vital signs or hematology or chemistry parameters in either treatment group. The proportion of subjects with QTcF or QTcB (Fridericia’s or Bazett’s corrected QT interval, respectively) values ≥500 ms during the trial was comparable across groups (QTcF: SUR 3.0%, VAN 3.4%; QTcB: SUR 5.9%, VAN 5.0%).

**DISCUSSION**

Given the burden of CDI and its recurrence, clinical development of new CDI therapeutics remains an urgent unmet medical need. The data from this phase 3 trial, which was conducted to assess the safety and efficacy of SUR relative to VAN, demonstrated that SUR did not meet the primary efficacy endpoint (clinical cure at EOT) of noninferiority compared with VAN, nor did it meet the key secondary efficacy endpoints of clinical response over time and sustained clinical response superiority over VAN. Surotomycin demonstrated reduced CDI recurrence rates and improved clinical response for subjects with baseline BI/NAP1/027-positive samples compared with VAN, but these differences were not statistically significant.

Compared with the SUR phase 2 trial results [7], the CDI clinical cure rates in the current trial were lower across treatment groups (86.6% and 89.4% for SUR 250 mg and VAN, respectively, in the phase 2 trial; 79.0% and 83.6%, respectively, in the current trial). In addition, the CDI recurrence rates were lower and clinical failure rates were higher in the current trial for both treatments compared with the SUR phase 2 trial [7]. The observed differences in these efficacy parameters may be due to the different inclusion criteria of the trials (phase 2 trial: ≥4 UBM s were required for inclusion; current trial: ≥3 UBM s) as well as differences in the definition of cure (phase 2 trial: defined as <4 UBM s per 24-hour period for at least 2 consecutive days; current trial: ≥2 UBM s per 24-hour period for at least 2 consecutive days). Furthermore, in both the phase 2 and current trials, CDI diagnosis required only 1 toxin-positive result. More importantly, these diagnostic 1-step procedures reflected recommended methods at the time of protocol development. The most current CDI diagnosis recommendations include a 2-step procedure [11].

Overall, the cure rates at EOT, sustained clinical response, and CDI recurrence for VAN were similar to those of a previous phase 3 noninferiority trial for fidaxomicin [12]. In this current trial, the sustained clinical response for both VAN and SUR was ~61% by the end of the trial (Day 40–50); however, in the SUR phase 2 trial, sustained clinical response rates of 66.7%
and 70.1% were reported for SUR 125 mg BID and 250 mg BID, respectively [7, 12]. The lower sustained clinical response rate in the current study may be due to the longer follow-up period used in this trial (30–40 days) compared with previous investigations (28 days) [7, 12].

The proportion of subjects from whom the epidemic BI/NAP1/027 strain was isolated (~25%) was slightly lower than in previous trials, which report values closer to one third [7, 12]. In the current trial, among BI/NAP1/027 strain-infected subjects, there was a higher clinical cure rate at EOT, sustained clinical response at the end of the trial, and lower recurrence rate in the SUR group than the VAN group. Vancomycin administration was associated with more favorable outcomes for these 3 efficacy parameters than SUR for non-BI/NAP1/027-infected subjects (Figure 4A and B). In contrast, in the phase 2 trial, a numerically higher clinical cure rate was seen in the VAN group than in the SUR 250 mg group for subjects infected with BI/NAP1/027 strain (91.3% vs 71.4%); however, the clinical cure rates were comparable for non-BI/NAP1/027 strain-infected subjects in both groups (SUR, 91.5%; VAN, 89.5% [7]). The differences in the BI/NAP1/027 strain-specific efficacy results may have contributed to the overall differences between the phase 2 and phase 3 SUR trial outcomes.

### Table 2. Overall Summary of Treatment-Emergent AEs (All Randomized Subjects, Irrespective of CDI Confirmation Status)

| Category                                      | Surotomycin (N = 305) n (%) | Vancomycin (N = 286) n (%) |
|-----------------------------------------------|-----------------------------|-----------------------------|
| Subjects with at least 1 AE                   | 148 (48.5)                  | 158 (55.2)                  |
| AEs by maximum severity                       |                             |                             |
| Mild                                          | 68 (22.3)                   | 84 (29.4)                   |
| Moderate                                      | 43 (14.1)                   | 52 (18.2)                   |
| Severe                                        | 37 (12.1)                   | 22 (7.7)                    |
| AEs by strongest relationship to study drug   |                             |                             |
| Not related                                   | 124 (40.7)                  | 132 (46.2)                  |
| Related                                       | 24 (7.9)                    | 26 (9.1)                    |
| At least 1 SAE                                | 44 (14.4)                   | 37 (12.9)                   |
| At least 1 treatment-related SAE              | 2 (0.7)                     | 0                           |
| AE causing discontinuation of study drug      | 17 (5.6)                    | 8 (2.8)                     |
| Treatment-related AE causing discontinuation of study drug | 4 (1.3) | 0 |
| Death                                         | 15 (5.9)                    | 9 (3.1)                     |
| Treatment-related death                       | 0                           | 0                           |

Abbreviations: AE, adverse events; CDI, Clostridium difficile infection; SAE, serious AEs.

### Table 3. Summary of TEAEs by MedDRA System Organ Class (Incidence ≥5%), Preferred Term (Incidence ≥2%), and Relationship to Study Drug (All Randomized Subjects, Irrespective of CDI Confirmation Status)

| System Organ Class          | Preferred Term                                      | Related n (%) | Not Related n (%) | Related n (%) | Not Related n (%) |
|-----------------------------|------------------------------------------------------|---------------|-------------------|---------------|-------------------|
| Subjects with at least 1 TEAE|                                                      |               |                   | 24 (7.9)      | 124 (40.7)        | 26 (9.1) | 132 (46.2) |
| Cardiac disorders           |                                                      |               |                   | 1 (0.3)       | 17 (5.6)          | 0        | 11 (3.8)  |
| Gastrointestinal disorders  |                                                      |               |                   | 13 (4.3)      | 59 (19.3)        | 12 (4.2) | 52 (18.2) |
| Abdominal pain              |                                                      |               |                   | 2 (0.7)       | 14 (4.6)         | 3 (1.0)  | 3 (1.0)   |
| Constipation                |                                                      |               |                   | 0             | 6 (2.0)          | 1 (0.3)  | 3 (1.0)   |
| Diarrhea                    |                                                      |               |                   | 0             | 10 (3.3)         | 0        | 14 (4.9)  |
| Nausea                      |                                                      |               |                   | 6 (2.0)       | 14 (4.6)         | 5 (1.7)  | 17 (5.9)  |
| Vomiting                    |                                                      |               |                   | 4 (1.3)       | 6 (2.0)          | 0        | 10 (3.5)  |
| General disorders and administration-site conditions | |               |                   | 1 (0.3)       | 24 (7.9)        | 1 (0.3)  | 24 (8.4)  |
| Edema peripheral            |                                                      |               |                   | 0             | 4 (1.3)          | 0        | 7 (2.4)   |
| Infections and infestations |                                                      |               |                   | 0             | 45 (14.8)        | 1 (0.3)  | 40 (14.0) |
| Urinary tract infection     |                                                      |               |                   | 0             | 15 (4.9)         | 0        | 7 (2.4)   |
| Investigations              |                                                      |               |                   | 5 (1.6)       | 18 (5.9)         | 10 (3.5) | 19 (6.6)  |
| Alanine aminotransferase increased |                                                |               |                   | 3 (1.0)       | 4 (1.3)          | 6 (2.1)  | 5 (1.7)   |
| Metabolism and nutrition disorders |                                                |               |                   | 1 (0.3)       | 21 (6.9)         | 2 (0.7)  | 20 (7.0)  |
| Dehydration                 |                                                      |               |                   | 0             | 4 (1.3)          | 0        | 6 (2.1)   |
| Hypokalemia                 |                                                      |               |                   | 0             | 5 (1.6)          | 0        | 7 (2.4)   |
| Musculoskeletal and connective tissue disorders | |               |                   | 0             | 8 (2.6)         | 2 (0.7)  | 17 (5.9)  |
| Arthralgia                  |                                                      |               |                   | 0             | 0               | 0        | 6 (2.1)   |
| Nervous system disorders    |                                                      |               |                   | 1 (0.3)       | 21 (6.9)         | 4 (1.4)  | 26 (9.1)  |
| Headache                    |                                                      |               |                   | 1 (0.3)       | 11 (3.6)         | 1 (0.3)  | 17 (5.9)  |

Abbreviations: CDI, Clostridium difficile infection; MedDRA, Medical Dictionary for Regulatory Activities; TEAEs, treatment-emergent adverse events.

AAs with the relationship to study drug missing were considered to be related in this table. Subjects who experienced more than 1 event were counted only once per System Organ Class and Preferred Term using the greatest relationship to study drug. System Organ Classes and Preferred Terms were based on MedDRA dictionary, version 15.0.
Throughout the current trial, SUR was generally well tolerated. The percentage of subjects with at least 1 AE and of drug-related AEs in this trial was similar between the 2 treatment groups. The number of deaths and discontinuations due to AEs was numerically higher in the SUR group than in the VAN group. None of the deaths were deemed treatment-related by the investigators.

Sparing of the Gram-negatives and bacteroidites during CDI therapy is hypothesized to favor recovery of a healthy microbiota and to lower CDI recurrence [13]. However, in this trial, the in vitro SUR spectrum of activity and modest perturbation of gut microbiota in vivo [6] was not associated with decreased CDI recurrence rates. After the unfavorable phase 3 data, the SUR development program was discontinued.

CONCLUSIONS

Although SUR 250 mg BID was generally well tolerated, it did not meet the primary efficacy endpoint of noninferiority versus VAN for clinical cure at EOT or either of the key secondary endpoints, ie, superiority of SUR to VAN for clinical response over time and sustained clinical response at the end of the trial. Approximately 40% of subjects ultimately failed CDI therapy for both SUR and VAN at the end of the follow-up period.

Supplementary Data

Supplementary materials are available at Open Forum Infectious Diseases online. Consisting of data provided by the authors to benefit the reader, the posted materials are not copyedited and are the sole responsibility of the authors, so questions or comments should be addressed to the corresponding author.

Acknowledgments

The trial and statistical analyses were performed by INC Research and Cubist Legacy before acquisition by Merck & Co., Inc. Medical writing assistance was provided by Dr. Edward Rochford (Complete Medical Communications, Hackensack, NJ). This assistance was funded by Merck & Co., Inc., Kenilworth, NJ.

Financial support. This work was supported by Merck & Co., Inc. Kenilworth, NJ.

Potential conflicts of interest. V. B. has acted as a consultant for, and received payments for talks from, AbbVie, Bristol-Myers Squibb, Gilead Sciences, Janssen, Merck Sharp & Dohme, and ViiV Healthcare. K. M. M. has been a clinical trial investigator and participated in advisory boards for Merck & Co., Inc., for surotomycin and other products. U. S., M. J., A. A., L. C., D. G., K. B. L., and Y. M. are current or former employees of Merck Sharp & Dohme Corp., a subsidiary of Merck & Co., Inc., Kenilworth, NJ, and may own stock and/or stock options.

All authors have submitted the ICMJE Form for Potential Conflicts of Interest. Conflicts that the editors consider relevant to the content of the manuscript have been disclosed.

References

1. Reveles KR, Lee GC, Boyd NK, Frei CR. The rise in Clostridium difficile infection incidence among hospitalized adults in the United States: 2001–2010. Am J Infect Control 2014; 42:1028–32.
2. Lessa FC, Gould CV, McDonald LC. Current status of Clostridium difficile infection epidemiology. Clin Infect Dis 2012; 55(Suppl 2):S65–70.
3. Kelly CP, LaMont JT. Clostridium difficile–more difficult than ever. N Engl J Med 2008; 359:1932–40.
4.Connor JR, Johnson S, Gerding DN. Clostridium difficile infection caused by the epidemic BI/NAP1/027 strain. Gastroenterology 2009; 136:1913–24.
5. Lessa FC, Mu Y, Ramberg WM, et al. Burden of Clostridium difficile infection in the United States. N Engl J Med 2015; 372:825–34.
6. Citron DM, Tyrrell KL, Dale SE, et al. Impact of surotomycin on the gut microbiota of healthy volunteers in a phase 1 clinical trial. Antimicrob Agents Chemother 2016; 60:2069–74.
7. Lee CH, Patino H, Stevens C, et al. Surotomycin versus vancomycin for Clostridium difficile infection: phase 2, randomized, controlled, double-blind, non-inferiority, multicentre trial. J Antimicrob Chemother 2016; 71:2964–71.
8. Arora V, Kachroo S, Ghanoty SS, et al. High Horr’s index score predicts poor outcomes in patients with Clostridium difficile infection. J Hosp Infect 2011; 79:23–6.
9. Snyderman DR, Jacobus NV, McDermott LA. Activity of a novel cyclic lipopeptide, CB-183,315, against resistant Clostridium difficile and other Gram-positive aerobic and anaerobic intestinal pathogens. Antimicrob Agents Chemother 2012; 56:3448–52.
10. US Food and Drug Administration. Guidance for Industry: Drug-Induced Liver Injury: Premarketing Clinical Evaluation. Available at: http://www.fda.gov/downloads/Drugs/.../Guidances/UCM174090.pdf. Accessed 25 August 2016.
11. Crobach MJ, Planche T, Eckert C, et al. European Society of Clinical Microbiology and Infectious Diseases: update of the diagnostic guidance document for Clostridium difficile infection. Clin Microbiol Infect 2016; 22(Suppl 4):S63–81.
12. Louie TJ, Miller MA, Mullane KM, et al. Fidaxomicin versus vancomycin for Clostridium difficile infection. N Engl J Med 2011; 364:422–31.
13. Seekatz AM, Young VB. Clostridium difficile and the microbiota. J Clin Invest 2014; 124:4182–9.