





# A Desirability of Outcome Ranking Analysis of a Randomized Clinical Trial Comparing Seven Versus Fourteen Days of Antibiotics for Uncomplicated Gram-Negative Bloodstream Infection

Jessica Howard-Anderson,<sup>1,0</sup> Weixiao Dai,<sup>2</sup> Dafna Yahav,<sup>3,4</sup> Toshimitsu Hamasaki,<sup>2</sup> Adi Turjeman,<sup>4,5</sup> Fidi Koppel,<sup>6</sup> Erica Franceschini,<sup>7</sup> Carol Hill,<sup>8</sup> Zoë Sund,<sup>8</sup> Henry F. Chambers,<sup>9</sup> Vance G. Fowler Jr.,<sup>8,10</sup> Helen W. Boucher,<sup>11</sup> Scott R. Evans,<sup>2</sup> Mical Paul,<sup>6,12</sup> Thomas L. Holland,<sup>8,10</sup> and Sarah B. Doernberg<sup>9</sup>; on behalf of the Antibacterial Resistance Leadership Group

<sup>1</sup>Department of Medicine, Division of Infectious Diseases, Emory University School of Medicine, Atlanta, Georgia, USA, <sup>2</sup>The Biostatistics Center and Department of Biostatics and Bioinformatics, Milken Institute School of Public Health, George Washington University, Washington, District of Columbia, USA, <sup>3</sup>Unit of Infectious Diseases, Rabin Medical Center, Beilinson Hospital, Petah Tikva, Israel, <sup>4</sup>Sackler Faculty of Medicine, Tel Aviv University, Ramat-Aviv, Israel, <sup>5</sup>Department of Medicine E, Rabin Medical Center, Beilinson Hospital, Petah-Tikva, Israel, <sup>7</sup>Clinic of Infectious Diseases, Rambam Health Care Campus, Haifa, Israel, <sup>7</sup>Clinic of Infectious Diseases, University of Modena and Reggio Emilia, Modena, Italy, <sup>8</sup>Duke Clinical Research Institute, Durham, North Carolina, USA, <sup>9</sup>Department of Internal Medicine, Division of Infectious Diseases, University of California, San Francisco, San Francisco, California, USA, <sup>10</sup>Department of Medicine, Duke University Medical Center, Durham, North Carolina, USA, <sup>11</sup>Division of Geographic Medicine and Infectious Diseases, Tufts Medical Center, Boston, Massachusetts, USA, <sup>12</sup>Faculty of Medicine, Technion, Israel Institute of Technology, Haifa, Israel

**Background.** Although a short course (7 days) of antibiotics has been demonstrated to be noninferior to a conventional course (14 days) in terms of mortality and infectious complications for patients with a Gram-negative bacterial bloodstream infection (GNB), it is unknown whether a shorter treatment duration can provide a better overall clinical outcome.

*Methods.* We applied a bloodstream infection-specific desirability of outcome ranking (DOOR) analysis to the results of a previously completed, randomized controlled trial comparing short versus conventional course antibiotic therapy for hospitalized patients with uncomplicated GNB. We determined the probability that a randomly selected participant in the short course group would have a more desirable overall outcome than a participant in the conventional duration group. We performed (1) partial credit analyses allowing for calculated and variable weighting of DOOR ranks and (2) subgroup analyses to elucidate which patients may benefit the most from short durations of therapy.

**Results.** For the 604 patients included in the original study (306 short course, 298 conventional course), the probability of having a more desirable outcome with a short course of antibiotics compared with a conventional course was 51.1% (95% confidence interval, 46.7% to 55.4%), indicating no significant difference. Partial credit analyses indicated that the DOOR results were similar across different patient preferences. Prespecified subgroup analyses using DOOR did not reveal significant differences between short and conventional courses of therapy.

*Conclusions.* Both short and conventional durations of antibiotic therapy provide comparable clinical outcomes when using DOOR to consider benefits and risks of treatment options for GNB.

Keywords. antibiotics; desirability of outcome ranking; Gram-negative bacteremia; treatment duration.

When selecting an antibiotic treatment duration, clinicians must balance the potential risks of stopping antibiotics with the potential benefits of this decision [1]. For more than a decade, many clinicians have advocated for shorter courses of antibiotic therapy [2–5]. There have been 3 published randomized controlled trials

Received 24 January 2022; editorial decision 14 March 2022; accepted 17 March 2022; published online 9 April 2022.

Correspondence: Jessica Howard-Anderson, MD, MSc, Emory University School of Medicine, Orr Building #1016, 550 Peachtree St. NE, Atlanta, GA 30303, USA (jrhowa4@emory.edu).

# Open Forum Infectious Diseases®2022

© The Author(s) 2022. Published by Oxford University Press on behalf of Infectious Diseases Society of America. This is an Open Access article distributed under the terms of the Creative Commons Attribution-NonCommercial-NoDerivs licence (https://creativecommons.org/licenses/by-nc-nd/4.0/), which permits non-commercial reproduction and distribution of the work, in any medium, provided the original work is not altered or transformed in any way, and that the work is properly cited. For commercial re-use, please contact journals.permissions@oup.com https://doi.org/10.1093/ofid/ofac140

(RCTs) assessing antibiotic duration in Gram-negative bacterial bloodstream infection (GNB) [6–8]. All studies demonstrated noninferiority with a short course (7 days) compared with a conventional course (14 days) of antibiotics with respect to a binary efficacy variable, and one study performed a post hoc desirability of outcome ranking (DOOR) analysis suggesting that a 7-day strategy might be preferred to 14 days [8]. More comprehensive benefit-to-risk analyses can help continue to elucidate effects of treatment duration on patient outcomes and inform the adoption of a treatment strategy for GNB.

Desirability of outcome ranking is an innovative analytic method that provides a global assessment of a patient's outcome, accounting for both benefits and harms, which is lacking with traditional noninferiority approaches. The DOOR analysis combines the benefits and risks of an intervention or new

therapy into a single outcome measure and allows for a superiority assessment [9]. Using a "partial credit" strategy, patients or clinicians can also choose the relative weighting of DOOR ranks, allowing for a more individualized analysis and approach to treatment [10].

The Antibacterial Resistance Leadership Group (ARLG) has used DOOR in studies of patients with multidrug-resistant infections [11–13] and developed a DOOR analysis plan for use in *Staphylococcus aureus* bloodstream infection trials [14]. In the current study, we performed a post hoc DOOR analysis of Yahav et al's [7] RCT comparing 7 versus 14 days of antibiotics for uncomplicated GNB. Our primary objective was to determine the probability that a patient assigned to a short course of antibiotics (7 days) would have a more desirable outcome than a patient assigned to a conventional course (14 days) of antibiotics.

## **METHODS**

# Study Design and Overview of the Original Study

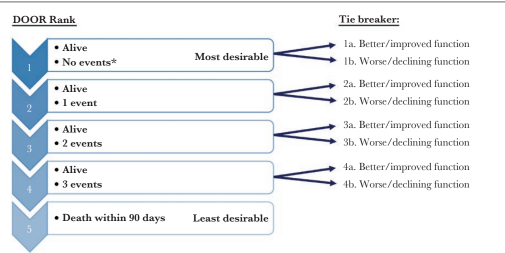
We performed a retrospective analysis of a multicenter, randomized, open-label, noninferiority trial of hospitalized patients

comparing 7 versus 14 days of antibiotics for GNB [7]. Patients enrolled in the original study were afebrile, hemodynamically stable for ≥48 hours before randomization, and had adequate source control. The primary outcome in the original study was a composite of 90-day all-cause mortality, clinical failure (included relapse, suppurative or distant complications), and readmission or extended hospitalization [7].

#### **Development of Desirability of Outcome Ranking Analysis Strategy**

We adapted the DOOR analysis strategy that was described and validated for *S aureus* bloodstream infection [14]. The DOOR analysis strategy was iteratively reviewed and edited by all study authors. With input from the investigators of the original RCT, we then determined which study variables would map to the ARLG DOOR analysis (Figure 1).

In the DOOR analysis, patients were assigned a mutually exclusive rank 1 through 5. Rank 1 represented the most desirable outcome and included anyone who was alive and did not experience any of the prespecified undesirable events within the follow-up period, whereas rank 5 indicated the least desirable



## Functional status scale:

- Independent
- · Needs assistance in ADLs
- Dependent in ADLs
- Bedridden

DOOR Events*	ARLG DOOR Definition	Variables used from original Yahav et al. study
Treatment failure <sup>a</sup>	Lack of resolution or recurrence of index infection prior to test of cure	Recurrent bacteremia due to the same organism as index infection
Infectious complications <sup>a</sup>	Newly identified complications or progression of the index infection not present at enrollment	<ul> <li>Local suppurative complications (e.g., renal abscess, empyema)</li> <li>Distant complication (growth of the same bacteria as in the initial bacteremia)</li> <li>Resistance development in secondary infections (clinical isolates resistant to ≥1 of the antibiotics used to treat index gram-negative bacteremia</li> </ul>
Adverse events	Unfavorable sign, symptom, or outcome that is not necessarily related to index infection (no assessment of causality)	Hospital stay >14 days from randomization Clostridioides difficile infection (within 30 days) Acute kidney injury <sup>b</sup> (within 30 days) Antibiotic rash (within 30 days) Hospital readmission (within 90 days) New clinically or microbiologically documented infection other than gram-negative bacteremia (within 90 days) Severe diarrhea <sup>c</sup> (within 90 days)

**Figure 1.** Primary desirability of outcome ranking (DOOR) analysis strategy and definitions of the components used in the analysis. a. Within 90-days postrandomization. b. Increased creatinine level ≥1.5× from baseline or glomerular filtration rate decrease >25% or urine output of <0.5 mL/kg per hour for 6 hours. c. Defined as 3 episodes of diarrhea/day for ≥2 days. ADL, activities of daily living; ARLG, Antibacterial Resistance Leadership Group.

outcome and included all patients who died within 90 days of randomization. Ranks 2 through 4 included patients who were alive but had 1, 2, or 3 undesirable events occur during the follow-up period, respectively (Figure 1). The undesirable events included in the analysis were categorized as follows: (1) treatment failure (including recurrence of infection), (2) infectious complications thought to be direct complications or progression of the original infection (not present at enrollment), and (3) other significant adverse events, not necessarily related to the original infection. Figure 1 displays which variables from the original RCT were mapped to each DOOR event. As a surrogate for health-related quality of life, we used both the functional status at 90-days and the change in functional status (from baseline to 90 days) as a "tiebreaker" in the DOOR analysis for patients with the same rank (eg, if 2 patients both had 1 undesirable event occur, then the patient with the better functional status at 90 days or the most improved functional status at 90 days would be given the better rank). Functional status in the original RCT was assessed at baseline and 90 days postrandomization and was categorized as follows: independent, needs assistance in activities of daily living (ADLs), dependent in ADLs or bedridden.

In a post hoc sensitivity analysis, we removed diarrhea from the other adverse event category because the primary study definition of diarrhea (3 episodes of diarrhea/day for ≥2 days) may not be considered severe. The DOOR probability was not significantly different from the primary analysis and these results were not shown.

## Statistical Analysis

We analyzed all randomized participants using an intention-to-treat strategy. The distribution of DOOR ranks was compared between the short and conventional duration strategies, and we determined the probability that a randomly selected participant assigned to a short course of therapy would have a better rank, or more desirable outcome, than a randomly selected participant assigned to a conventional course of therapy. A DOOR probability of 50% indicates no difference between groups. We also explored the difference in duration of actual antibiotics received between the 2 treatment groups and plotted this as function of the DOOR probability.

We performed a partial credit analysis in which DOOR ranks are scored like an academic exam. This method allows individuals to provide a personalized scoring key for how to grade the different DOOR ranks based on individual preferences. Survival without any events (rank 1) is always assigned a score of 100, and death (rank 5) is always assigned a 0, but ranks 2–4 can be given any score between 0 and 100 as long as the original order is maintained [10]. We chose 3 different partial credit scenarios that represented a range of theoretical patient preferences. For each scenario, we subtracted the mean partial credit score for the conventional group from the mean partial credit

score for the short group. A difference of zero indicates no difference between groups. To visually display contours of the difference in mean partial credit scores in a 2-dimensional plot, we combined being alive with 2 or 3 events in Figure 4.

Fianlly, we compared the DOOR distributions by treatment strategy for prespecified subgroups defined by the primary source of bacteremia (urinary tract infection [UTI] versus nonurinary source) and the presence of a ceftazidime-resistant Enterobacterales pathogen, indicative of an extended-spectrum beta-lactamase (ESBL)-producing Enterobacterales in this trial. These subgroups were similar to the ones performed in the original study and were chosen because we thought clinicians may have more concern with using a short course of antibiotics in patients with a nonurinary source of infection or in those with a multidrug-resistant organism. Our hypothesis was that outcomes would be similar in patients randomized to the short course compared with those randomized to the conventional course, even in these potentially more challenging infectious scenarios.

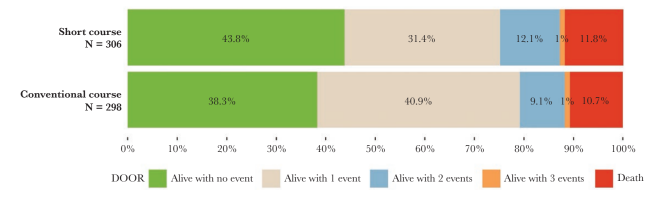
#### **Patient Consent Statement**

This study was reviewed by the Duke University Health System Institutional Review Board, which determined that this study was exempt from human subjects research review. Because this study was a secondary analysis of previously collected data, reconsent of participants was not required. As part of the initial trial, participants provided written informed consent, as described in the primary manuscript by Yahav et al [7].

# **RESULTS**

All 604 patients included in the original RCT were analyzed; 306 were assigned to the short course (7 days) and 298 were assigned to the conventional course (14 days). Descriptive characteristics of the patient population have been previously published [7]. Most patients (n = 411, 68%) had a UTI as the primary source of GNB. Most patients (n = 543, 90%) had Enterobacterales as the primary pathogen, 105 (19%) of which were ESBL-producing organisms. Most patients (n = 375, 62%) were independent in their ADLs but 40 (7%) were bedridden at baseline.

Overall, 248 (41%) patients were alive without any undesirable events, 218 (36%) with 1 event, 64 (11%) with 2 events, 6 (1%) with 3 events, and 68 (11%) died. The DOOR distribution was similar between the short and conventional treatment groups with probability of having a more desirable outcome with the short versus conventional course of 51.1% (95% confidence interval [CI], 46.7%–55.4%), indicating no significant difference (Figure 2). In addition, no significant differences were observed between the short and conventional groups for any of the component undesirable events contributing to the DOOR rank (Figure 3 and Supplemental Table 1). The median duration



**Figure 2.** Desirability of outcome ranking (DOOR) distribution by treatment groups. The DOOR probability of having a more desirable outcome when assigned to the short course of antibiotics was 51.1% (95% confidence interval, 46.7%–55.4%).

of appropriate covering antibiotics received for the index bacteremia was 7 days in the short course group and 14 days in the conventional course group [7]. However, when looking at the mean difference in antibiotic days received, the difference between the 2 treatment groups was closer to 6 days, and this is relatively consistent across the 95% confidence region of the DOOR probability (Supplemental Figure 1).

When the patient's functional status at 90 days was used as a tiebreaker in the DOOR analysis, the probability of having a more desirable outcome in the short versus conventional group remained similar (51.5%; 95% CI, 46.9%–56.0%). There was also no difference when using the change in functional status (from baseline to 90 days) as a tiebreaker (data not shown).

Using a partial credit analysis, we evaluated the effect of ascribing different partial credit scores to the DOOR ranks based on hypothetical patient preferences, rather than having each rank receive equal weight (Figure 4). Scenario A represents a patient who places value only on avoiding mortality, regardless of the complications that arise on the path to survival (this is equivalent to using 90-day mortality as the primary outcome). Alternatively, a patient represented by scenario B places more value on quality of life and would consider anything other than an uncomplicated survival to be objectionable. In scenario C, the representative patient places significant value on surviving their illness but does place some value on avoiding complications, unlike the patient in scenario A. In all scenarios, we did not observe a significant difference in the mean partial credit scores between the short and conventional duration groups (Figure 4). The DOOR distribution was also similar comparing short versus conventional treatment duration, stratified by the primary source of bacteremia (UTI vs other source) or by the presence of an ESBL-producing pathogen (Supplemental Figures 2 and 3).

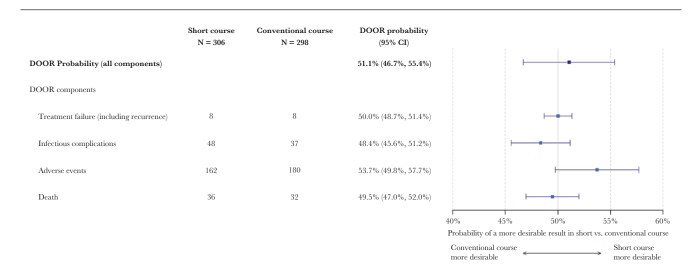
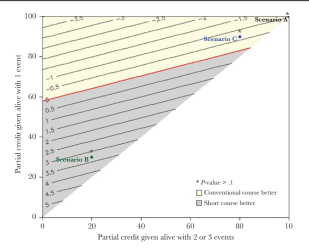


Figure 3. Forest plot demonstrating the desirability of outcome ranking (DOOR) probabilities for each individual DOOR component (treatment failure, infectious complications, adverse events, and death) as well as the overall DOOR probability of having a more desirable outcome with a short course of treatment compared with a conventional duration. The individual components are not mutually exclusive, and patients can have more than one of these events. CI, confidence interval.

	DOOR Distribution			Partial credit scoring key		
DOOR Rank	Description	Short course (n = 306)	Conventional course (n = 298)	Scenario A (mortality)	Scenario B (quality of life)	Scenario C (balanced)
1	Alive with no events	134 (43.8%)	114 (38.3%)	100	100	100
2	Alive with 1 event	96 (31.4%)	122 (40.9%)	100	30	90
3	Alive with 2 events	37 (12.1%)	27 (9.1%)	100	20	80
4	Alive with 3 events	3 (1.0%)	3 (1.0%)	100	10	70
5	Death	36 (11.8%)	32 (10.7%)	0	0	0

A			В		C	
Short 88.2	Conventional 89.3	Short 55.7	Conventional 52.4	Short 82.4	Conventional 83.1	
	Difference	e in mean p	artial credit scores	(95% CI)		



**Figure 4.** Desirability of outcome ranking (DOOR) partial credit scenarios. The top panel provides the partial credit scoring key. Scenario A represents a patient who only places value on surviving (equivalent to a 90-day mortality outcome). Scenario B represents a patient who places more value on quality of life and considers any adverse event as very undesirable. Scenario C represents a patient who places significant value on survival but also tries to balance this with avoiding complications. For each scenario, the difference in mean partial credit scores is calculated by subtracting the mean score for the conventional group from the mean score for the short group. The bottom panel displays contours of the difference in mean partial credit scores. The partial credit score assigned to being alive with 2 or 3 events is combined on the horizontal axis, and the partial credit score assigned to being alive with one event is on the vertical axis. The red line at zero indicates no difference between the mean partial credit scores. Positive differences (shaded in gray) suggest the short course could provide a more desirable outcome, and negative differences (shaded in yellow) suggest the conventional course could provide a more desirable outcome. However, none of the theoretical partial credit scenarios (A–C) demonstrated a significant difference between treatment groups (*P*> .1 for all scenarios).

# **DISCUSSION**

We applied a DOOR analysis to a previously published RCT assessing duration of antibiotics for GNB [7]. Using DOOR, we found that patients randomized to either treatment duration experienced similar rates of desirable outcomes. These results persisted after including a marker of functional status at 90 days. The DOOR results also remained similar in a partial credit analysis with various weights applied to each DOOR level to adjust for different patient values.

Our results confirm and extend those reported in the original RCT by incorporating both efficacy and safety concerns into the primary outcome [7]. We ultimately observed that the overall desirability of the outcome in patients receiving a short course of antibiotics was comparable to those receiving a conventional course, and the patients assigned to the short course

of antibiotics received, on average, 6 fewer days of antibiotics. At both the individual and the population level, this decrease in antibiotic duration can represent a significant advantage. Decreasing overall antibiotic usage can minimize the risk and global burden of antimicrobial resistance, and patients and clinicians should strongly consider this factor when deciding on an antibiotic duration [15, 16].

Our results are similar to the recently published study by Molina et al [8] who also performed a post hoc DOOR analysis of a trial comparing 7 versus 14 days of antibiotics for GNB. In the Molina et al [8] study, they used Response Adjusted for Duration of Antibiotic Risk (RADAR), a specific version of DOOR, which incorporates the total number of antibiotic days received in the DOOR endpoint (ie, if 2 participants had the same clinical outcome, the participant with the fewest antibiotic

days would be given the more desirable rank) [9]. We did not use RADAR in our study because we chose to focus more directly on the clinical outcomes most impactful to patients rather than duration of antibiotics received, which closely tracked with assigned study arm in this study. It should be noted, however, that the assigned duration of antibiotics does not always mirror the actual duration patients receive in clinical trials, and therefore we also analyzed the mean difference in antibiotics received between groups as a function of the DOOR probability and the 95% confidence region. In addition, in contrast to the Molina et al [8] study, the DOOR endpoint used in our analysis accounts for the cumulative nature of undesirable events that can occur throughout a patient's clinical course without assigning priority to the events. We favor this approach over the DOOR endpoint used by Molina et al [8] in which clinical efficacy was prioritized over safety (ie, a patient with a serious adverse event but who met the definition of cure was ranked higher than a patient with clinical failure but no adverse events). This prioritization might not be the same for all patients, clinicians, or antibiotics tested. In addition, our report incorporated a partial credit analysis, which is another unique aspect of this work. We are encouraged that DOOR is being used in more clinical trials. Further work should be done to standardize the DOOR analysis which will help clinicians compare results across clinical studies.

One of the advantages of the DOOR analysis is the ability to help elucidate when the potential risks of one therapy outweigh the benefits. In traditional registrational drug trials, the efficacy and safety analyses are usually performed in separate populations, and this can make it difficult to translate clinical trial results to individual patient care. Subgroup analyses using DOOR can incorporate both the potential benefits and harms of a new therapy or strategy and can help physicians individualize treatment decisions. In this study, we specifically addressed patients who had a UTI as the source of the bloodstream infection or those with an ESBL-producing pathogen, and we did not find that one antibiotic duration strategy was superior to the other. Performing similar, prespecified subgroup analyses in prospective clinical trials using DOOR could be a useful tool to analyze diverse patient populations, including patients who may be older, have poor functional status, or are immunocompromised. These subgroups are traditionally underrepresented in clinical trials, and inclusion should be encouraged to understand treatment effects in diverse populations [17, 18]. Differing partial credit scores using the DOOR analysis allow for individualized assessment of benefit/risk based on patient preferences.

This study is limited by the fact that we retrospectively fit the original study variables into our DOOR analysis strategy. Not all DOOR events were monitored for the same duration of follow-up. Recurrence and infectious complications were evaluated for 90 days, but many adverse events were only followed for 30 days. Other important clinical considerations such as the antibiotic susceptibly profile, requirements for intravenous antibiotics,

and cost were not included in the DOOR analysis. In addition, our current study did not include patient reported outcomes or include a comprehensive assessment of health-related quality of life. Understanding how to best incorporate both of these factors into future prospective studies using DOOR will be critical to ensure these studies remain relevant to patients [19].

## **CONCLUSIONS**

In summary, we applied DOOR to a large RCT comparing 7 versus 14 days of antibiotics for patients with GNB and did not observe significant differences between groups using a global measure of clinical success. For most patients with uncomplicated GNB, a short course of antibiotics is equivalent to a conventional course. The DOOR can provide useful information regarding the totality of treatment effects on patient outcomes and provide clinicians with comprehensive analyses of the risks and benefits of an intervention to help guide therapeutic decision making. We believe DOOR should be prospectively included in future RCTs.

#### **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**

We are grateful for the study participants and all staff who participated in this trial

**Disclaimer.** The content is solely the responsibility of the authors and does not necessarily represent the official views of the National Institutes of Health.

Financial support. This work was funded by the National Institute of Allergy and Infectious Diseases of the National Institutes of Health (NIH) under Award Number UM1AI104681. J. H.-A. was supported by the Antibacterial Resistance Leadership Group fellowship (National Institute of Allergy and Infectious Disease UM1AI104681).

Potential conflicts of interest. H. W. B. declares consulting fees from Antimicrobial Agents and Chemotherapy, Sanford Guide, and ID Clinics of North America. H. F. C. participates in the Data Safety Monitoring Committee for a Merck SARS-CoV-2 antiviral agent and declares stock in Moderna. S. B. D. declares consulting fees from Genetech and Basilea Pharmaceutica. S. R. E. declares grants from Degruyter, royalties from Taylor & Francis, consulting fees from Genentech, AstraZeneca, Cardinal Health, Microbiotix, Stryker, Atricure, Roivant, Neovasc, Nobel Pharma, Horizon, International Drug Development Institute, and SVB Leernik, honoraria from Analgesic, Anesthetic, and Addiction Clinical Trial Translations, Innovations, Opportunities, and Networks (ACTTION), Osaka University, and National Cerebral and Cardiovascular Center of Japan. S. R. E. also participates on Data Safety Monitoring Boards for NIH, Breast International Group, University of Pennsylvania, Duke University, Roche, Pfizer, Takeda, Novartis, Amgen, Teva, Vir, Shire, Alexion, Gilead, Tracon, Rakuten, Abbvie, Nuvelution, Clover, FHI Clinical, Lung Biotech, SAB Biohparm, and Advantagene. S. R. E. is also a board member of the American Statistical Association, Society for Clinical Trials, and Frontier Science Foundation. V. G. F. reports personal fees from Novartis, Novadigm, Durata, Debiopharm, Genentech, Achaogen, Affinium, Medicines Co., Cerexa, Tetraphase, Trius, MedImmune, Bayer, Theravance, Basilea, Affinergy, Janssen, xBiotech, Contrafect, Regeneron, Basilea, Destiny, and Amphliphi Biosciences; Integrated Biotherapeutics; C3J, Armata, Valanbio; Akagera, Aridis; Grants from NIH, MedImmune, Allergan, Pfizer, Advanced Liquid Logics, Theravance, Novartis, Merck; Medical Biosurfaces; Locus; Affinergy; Contrafect; Karius; Genentech, Regeneron, Basilea, Janssen, and Akagera; Royalties from UpToDate; Stock options from Valanbio and ArcBio; a patent sepsis diagnostics pending; and a stipend from Infectious Diseases Society of America for service as Associate Editor on Clinical Infectious Diseases. T. H. declares consulting fees from Mitsubishis Tanabe Pharma and honoraria from Johnson & Johnson, KK, Duke University, and Chang Gung Medical Foundation (Taiwan). T. L. H. declares royalties from UpToDate and consulting fees from Basilea Pharmaceutica, MotifBio, Genentech, and Lysovant. All authors have submitted the ICMJE Form for Disclosure of Potential Conflicts of Interest. Conflicts that the editors consider relevant to the content of the manuscript have been disclosed.

#### References

- Spellberg B. The maturing antibiotic mantra: "shorter is still better". J Hosp Med 2018; 13:361.362.
- Spellberg B, Rice LB. Duration of antibiotic therapy: shorter is better. Ann Intern Med 2019; 171:210–1.
- Rice LB. The Maxwell Finland Lecture: for the duration-rational antibiotic administration in an era of antimicrobial resistance and Clostridium difficile. Clin Infect Dis 2008; 46:491–6.
- Royer S, DeMerle KM, Dickson RP, Prescott HC. Shorter versus longer courses of antibiotics for infection in hospitalized patients: A systematic review and metaanalysis. J Hosp Med 2018; 13:336–42.
- Wald-Dickler N, Spellberg B. Short-course antibiotic therapy—replacing Constantine units with "Shorter Is Better". Clin Infect Dis 2019; 69:1476–9.
- von Dach E, Albrich WC, Brunel A-S, et al. Effect of C-reactive protein-guided antibiotic treatment duration, 7-day treatment, or 14-day treatment on 30-day clinical failure rate in patients with uncomplicated Gram-negative bacteremia: a randomized clinical trial. JAMA 2020; 323:2160-9.
- Yahav D, Franceschini E, Koppel F, et al. Seven versus 14 days of antibiotic therapy for uncomplicated Gram-negative bacteremia: a noninferiority randomized controlled trial. Clin Infect Dis 2019; 69:1091–8.

- Molina J, Montero-Mateos E, Praena-Segovia J, et al. Seven-versus 14-day course of antibiotics for the treatment of bloodstream infections by Enterobacterales: a randomized, controlled trial. Clin Microbiol Infect 2021; 28:550-7.
- Evans SR, Rubin D, Follmann D, et al. Desirability of outcome ranking (DOOR) and response adjusted for duration of antibiotic risk (RADAR). Clin Infect Dis 2015; 61:800–6.
- Evans SR, Follmann D. Using outcomes to analyze patients rather than patients to analyze outcomes: a step toward pragmatism in benefit:risk evaluation. Stat Biopharm Res 2016: 8:386–93.
- van Duin D, Lok JJ, Earley M, et al. Colistin versus ceftazidime-avibactam in the treatment of infections due to carbapenem-resistant enterobacteriaceae. Clin Infect Dis 2018; 66:163–71.
- Lodise TP, Rosenkranz SL, Finnemeyer M, et al. The emperor's new clothes: PRospective observational evaluation of the association between initial vancomycin exposure and failure rates among ADult HospitalizEd Patients With Methicillin-resistant Staphylococcus aureus Bloodstream Infections (PROVIDE). Clin Infect Dis 2020; 70:1536–45.
- van Duin D, Arias CA, Komarow L, et al. Molecular and clinical epidemiology of carbapenem-resistant Enterobacterales in the USA (CRACKLE-2): a prospective cohort study. Lancet Infect Dis 2020; 20:731–41.
- 14. Doernberg SB, Tran TTT, Tong SYC, et al. Good studies evaluate the disease while great studies evaluate the patient: development and application of a desirability of outcome ranking endpoint for *Staphylococcus aureus* bloodstream infection. Clin Infect Dis 2019; 68:1691–8.
- Septimus EJ. Antimicrobial resistance: an antimicrobial/diagnostic stewardship and infection prevention approach. Med Clin North Am 2018; 102:819–29.
- Kadri SS, Boucher HW. U.S. efforts to curb antibiotic resistance Are we saving lives? N Engl J Med 2020; 383:806–8.
- Van Spall HGC, Toren A, Kiss A, Fowler RA. Eligibility criteria of randomized controlled trials published in high-impact general medical journals: a systematic sampling review. JAMA 2007; 297:1233–40.
- He J, Morales DR, Guthrie B. Exclusion rates in randomized controlled trials of treatments for physical conditions: a systematic review. Trials 2020; 21:228.
- 19. King HA, Doernberg SB, Miller J, et al. Patients' experiences with Staphylococcus aureus and Gram-negative bacterial bloodstream infections: a qualitative descriptive study and concept elicitation phase to inform measurement of patient-reported quality of life. Clin Infect Dis 2020; 73:237–47.