Echinocandins – first line in invasive candidiasis: how strong is this ‘strong’ evidence?

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In the previous issue of Critical Care, Kett and colleagues [1] published a post hoc analysis of a randomized controlled trial comparing the efficacy of anidulafungin versus fluconazole in non-neutropenic critically ill patients with invasive Candida infections (89% had candidemia). But the authors’ claim that their data support the superiority of anidulafungin may be misleading and raises several concerns. First, the primary endpoint of the study was clinical and microbiological success at the end of intravenous therapy. However, surrogate endpoints must be predictive of the clinically relevant endpoint that is mortality [2]. That was not the case, and no difference in 28-day mortality was noted (20.2% versus 24.3%; \( P = 0.57 \)). Second, in the present study [1], the duration of intravenous therapy was unclear, but in their original study [3], patients on anidulafungin received, on average, 3 more days of intravenous therapy than the fluconazol group. Besides, more patients in the anidulafungin arm had their central venous catheter removed. These facts markedly biased the results and could explain the observed differences [4]. Third, this was a non-inferiority study [3]. Therefore, from a statistical point of view, any conclusions regarding superiority must be interpreted with extreme caution [5]. Finally, at the time of the study design [3], the use of amphotericin B, and not fluconazole, was recommended in unstable patients with invasive Candida infections. Therefore, the choice of fluconazole as a comparator limits the study conclusions even further. We believe that, at present, there is no evidence to support the selection of a specific antifungal class in invasive Candida infections [4].

Authors’ response

Daniel H Kett, Annette C Reboli, Andrew F Shorr and Haran T Schlamm

In response to the letter by Gonçalves-Pereira and Póvoa, we would like to point out that, in our post hoc analysis of seriously ill patients with invasive candidiasis, anidulafungin was more effective than fluconazole in terms of global (combined clinical and microbiological) response at the end of treatment. Recognizing the limitations of our analysis, however, we were unable to conclude that anidulafungin was superior to fluconazole.

Many of the comments of Gonçalves-Pereira and Póvoa were directed at the original study. That study was a prospective, randomized, double-blind phase III trial [3] that incorporated a pre-specified, two-step test for non-inferiority and then superiority, an accepted statistical method [6]. The primary endpoint of that study was investigator-assessed global response, a commonly accepted endpoint in candidemia studies. The difference in global response between groups remained significant after adjustment for potential imbalances, including duration of treatment and central line status [3].

We disagree with the claim by Gonçalves-Pereira and Póvoa that previous guidelines did not support the use of fluconazole as a first-line treatment in unstable patients with candidemia. The 2000 guidelines of the Infectious Diseases Society of America (IDSA) recommended either fluconazole or amphotericin B, and not fluconazole, was recommended in unstable patients with invasive Candida infections. Therefore, the choice of fluconazole as a comparator limits the study conclusions even further. We believe that, at present, there is no evidence to support the selection of a specific antifungal class in invasive Candida infections [4].

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we called for additional studies in critically ill patients with invasive candidiasis.

Abbreviation
IDSA, Infectious Diseases Society of America.

Competing interests
JG-P and PP have each received honoraria from and served as advisor to Gilead (Foster City, CA, USA), Merck Sharp & Dohme (Whitehouse Station, NJ, USA), and Pfizer Inc (New York, NY, USA).

Authors’ contributions
JG-P and PP both searched the literature, analyzed data, and wrote the manuscript. Both authors read and approved the final manuscript.

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