Principles of antimicrobial stewardship for bacterial and fungal infections in ICU

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Appropriate use of antibiotics

Selecting antimicrobial therapy for suspected infection in critically ill patients is an important decision-making process for intensivists. In this current age of multi-drug-resistant organisms (MDROs), intensivists must balance the increased mortality associated with delaying therapy of microbiologically documented infections with the potential ecological damage caused by antimicrobial medications, including the selection and development of MDROs [1]. In patients with septic shock, even delays of few hours in the administration of appropriate antimicrobial therapy, defined as the antibiotic regimen to which the causative pathogen is sensitive, has been associated with increased mortality [2]. The same concept applies to other types of documented infections, such as pneumonia or bloodstream infections, where administration of an inappropriate initial antibiotic regimen is associated with an increased risk of death that is not mitigated by treatment escalation [1, 2]. Unfortunately, the “appropriateness” of initial antibiotic regimens is becoming increasingly difficult due to the rising levels of bacterial resistance.

Escalating rates of antibiotic resistance, including resistance to carbapenems and colistin, are becoming increasingly prevalent and significantly impact morbidity, mortality, and costs related to infection in hospitalized patients, especially in the intensive care unit (ICU). In Europe, the prevalence of carbapenemase-producing Enterobacteriaceae is increasing, in particular with the rapid spread of carbapenem-hydrolysing oxacillinase-48 and New Delhi metallo-beta-lactamase-producing Enterobacteriaceae [3]. Antimicrobial use is a key determinant in the selection of resistant bacteria, whereby increasing daily cumulative exposure, especially for carbapenems, increases the likelihood of intestinal carriage of MDROs [3]. The rapid evolution of antibiotic resistance impedes efforts to insure that initial appropriate antibiotic therapy is delivered to critically ill infected patients. The rising rates of antibiotic resistance have likely contributed to the increase in observed mortality attributed to antibiotic-resistant bacteria, despite the overall reduction in deaths ascribed to infectious diseases in the last century [4]. Acinetobacter baumannii and Pseudomonas aeruginosa represent two of the most important MDROs due to the extent of their antimicrobial resistance, intrinsic virulence, especially for Pseudomonas, and ability to establish themselves within hospital environments such as ICUs and to infect immune suppressed hosts [1, 5]. Indeed, a recent European task force now recommends the empiric use of polymixin or colistin as part of the empiric antimicrobial regimen when infection with A. baumannii is suspected due to the high rates of carbapenem resistance in this pathogen [5]. An important contributor to the escalating rates of infection with these MDROs, especially in Europe and North America, has been population migration from the Middle East and other parts of Asia and Africa. Similarly, the movement of individuals across borders is also recognized as an important source of spread of other emerging infections such as Dengue fever, Zika virus, and Middle East respiratory syndrome coronavirus [6]. Reducing the unnecessary use of antibiotics is seen as a pivotal strategy for curtailing the further emergence and spread of MDROs [7]. Figure 1 provides a summary of the strategy of antimicrobial de-escalation

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that can assist intensivists in balancing the desire to treat with an appropriate antibiotic regimen with the need to avoid unnecessary antibiotic exposure.

**Catheter-related infections**

There is accumulating recent evidence that rates of catheter-related infections (CRI) can be substantially reduced towards zero with the correct application of care bundles. As recently summarized in *Intensive Care Medicine*, the cornerstone of prevention relies on the correct skin preparation [8, 9]. Alcohol seems mandatory in antisepsis solutions due to its rapid activity; chlorhexidine gluconate ensures a prolonged bactericidal effect. There are actually some important certainties/recommendations on prevention of CRI like: (1) application of sterile 2% CHG–alcohol to disinfect the skin prior to insertion of a vascular catheter, (2) no need for cleaning the skin with soap or detergent before antiseptic application when it is not obviously dirty or contaminated, (3) rinsing with sterile water is not recommended after antiseptic application, (4) single-use vials of disinfectant solutions are preferable to multiple-use bottles, (5) daily evaluation of the necessity of retaining central line(s) and removal if they are clinically no longer necessary, (6) non-adherent, soiled, or moistened dressings should be promptly replaced, (7) a subclavian site of insertion is preferable unless otherwise contraindicated, (8) full barrier precautions should be used at insertion, (9) hand hygiene with alcoholic solution is mandatory before every manipulation, and (10) every unit should adopt a written protocol for catheter insertion and maintenance. Certainly, there is room for improvement, with behavioral approaches requiring well-designed multinational trials. However, mortality and CRI rates were not reduced in a prospective randomized controlled study from Brazil employing a behavioral approach for retention or withdrawal of central lines; the rather short evaluation period might have contributed to this. Nevertheless, the feedback from administrators and caregivers was positive, prompting universal adoption of the program after the experimental period of 6 months [10]. Transparent antimicrobial dressings, being significantly more efficacious in preventing CRBSIs when compared to the reference dressings and the use of sterile 2% CHG/70% isopropyl alcohol for skin antisepsis represent new standards of care [8, 9, 11].

Source control, recommended within 12 h from patient’s admission, includes primarily actions taken in the process of care to control the foci of infection. It

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**Suspected infection in critically ill patient**

- Prescribe an empiric antibiotic regimen employing a combination of agents targeting likely MDROs based on local epidemiology and patient risk factors
- Obtain the following: Microbiologic samples for culture and special stains, Biomarkers (e.g., procalcitonin), Identify risk factors for infection with DRPs, Rapid microbiologic tests
- Monitor clinical response to treatment (temperature, white blood cell count, hemodynamic parameters, organ function, biomarker levels)
- Microbiologic identification of pathogen(s) and antimicrobial susceptibility
- Significant clinical improvement after 48-96 hours of treatment

**Consider discontinuation of antibiotics**

- Review dosing of antibiotic to optimize drug levels
- Reculture for possible super infection
- Consider non-infectious causes of fever/inflammation
- Examine for abscess formation
- De-escalate empiric antibiotics based on microbiology results
- Discontinue antibiotics after 7-10 day course based on site of infection and pathogen

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**Fig. 1** Schematic outlining a practice of antimicrobial de-escalation. FiO₂ inspired oxygen fraction, PaO₂ partial pressure (or tension) of arterial oxygen, MDROs multi-drug resistant organisms
Invasive candidiasis (IC) comprises both bloodstream and other deep-seated invasive infections due to Candida species and represents a major cause of morbidity and mortality, which remains near 40% in most series. Over the past decades, incidence of IC has shown progressively increasing or stable rates in most regions [14–16], probably due to the growing complexity of surgical procedures, the presence of patient populations at higher risk of infection, and the changes in patient demographic characteristics [14]. Attributable mortality around 10% has been reported in several studies [16], while a recent well-designed study conducted in India has reported rates around 20% [14].

Infection-related morbidity and mortality associated with IC are in part associated with the critical illness itself, but they may also be attributed to suboptimal diagnostic tools. Standard mycological conventional blood culture is a gold standard, but may fail to diagnose IC in up to 25–50%, whereas time to positivity for yeast and species identification can take several days from the time the blood culture is drawn, depending on the implicated Candida species [16]. Unless antifungal therapy is promptly initiated by the confirmation of IC, effective antifungal therapy may be further delayed. Numerous studies have demonstrated that the delay of initiation of antifungal therapy was associated with significant increases in both in-hospital mortality and the cost of care for patients with IC [16].

Empiric and pre-emptive use of antifungals in Candida infections

In cases of IC, treatment should be given immediately, but thereafter it would be important to step down early to a triazole. De-escalation from an echinocandin to intravenous or oral fluconazole should be encouraged when the patient is clinically stable and the isolated strain is susceptible. The exact timing for shifting to fluconazole is not well established and may vary, depending on patient- and pathogen-related factors. The safety of de-escalation (within 5 days) in the case of proven or probable IC has been recently suggested for non-neutropenic patients treated in the ICU, since it was not associated with increased mortality and led to a subsequent and significant decrease in the antifungal consumption [18].

The most remarkable rules for developing antifungal stewardship program targeting antifungal treatments bundles are reported in the Electronic Supplement Material.

**Electronic supplementary material**

The online version of this article (doi:10.1007/s00134-017-4922-x) contains supplementary material, which is available to authorized users.

**Abbreviations**

CRI: Catheter-related infections; IAC: Intra-abdominal candidiasis; IC: Invasive candidiasis; ICU: Intensive care unit; MDROs: Multi-drug-resistant organisms.

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Compliance with ethical standards

Conflicts of interest
Dr. Bassetti has participated in advisory boards and/or received speaker honoraria from Achaogen, Angelini, Astellas, AstraZeneca, Bayer, Basilea, Gilead, Menarini, MSD, Pfizer, The Medicine Company, Tetraphase and Vifor. Dr. Kollef's effort was supported by the Barnes-Jewish Hospital Foundation. Dr Poulakou has participated in advisory boards and/or received speaker honoraria from MSD and Pfizer.

Received: 5 June 2017  Accepted: 14 June 2017  Published online: 12 September 2017

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