Abstract

Bloodstream infections (BSIs) are frequent in ICU and is a prognostic factor of severe sepsis. Community acquired BSIs usually due to susceptible bacteria should be clearly differentiated from healthcare associated BSIs frequently due to resistant hospital strains. Early adequate treatment is key and should use guidelines and direct examination of samples performed from the infectious source. Previous antibiotic therapy knowledge, history of multi-drug resistant organism (MDRO) carriage are other major determinants of first choice antimicrobials in healthcare-associated and nosocomial BSIs. Initial antimicrobial dose should be adapted to pharmacokinetic knowledge. In general, a high dose is recommended at the beginning of treatment.

If MDRO is suspected combination antibiotic therapy is mandatory because it increase the spectrum of treatment. Most of time, combination should be pursued no more than 2 to 5 days.

Given the negative impact of useless antimicrobials, maximal effort should be done to decrease the antibiotic selection pressure. De-escalation from a broad spectrum to a narrow spectrum antimicrobial decreases the antibiotic selection pressure without negative impact on mortality. Duration of therapy should be shortened as often as possible especially when organism is susceptible, when the infection source has been totally controlled.

Keywords: Bloodstream infection, Healthcare associated infections, Bacterial resistance, Intensive care unit

Introduction

Bloodstream infections (BSI) are a major cause of morbidity and mortality worldwide [1]. These infections may arise secondarily to localized infection at a specific body site, or may be classified as primary when no focus is identified. Severe sepsis, septic shock, and multisystem organ dysfunction related to BSI frequently require admission in ICU for appropriate management [2]. BSI, as a complication of critical illness, occurs in approximately 5% of all patients admitted to ICUs [3,4]. The case-fatality rate associated with BSI reaches 35-50% when associated with admission to ICU [1].

The mainstay of therapy for patients with bacteremia remains antimicrobial therapy, together with the optimal management of its consequences (e.g., shock or metastatic suppurative complications), and surgical treatment of an infection site (e.g., debridement, abscess drainage, or removal of intravascular devices), when appropriate.

Considering the paucity of new antimicrobials in the industry pipeline, and the emergence of strains resistant to recent antimicrobials, enhanced efforts must be made to decrease selection pressure to antimicrobials [5]. To contain the worldwide problem of antimicrobial resistance, intensivists have two major tools with regards to antimicrobials treatments: [1] antimicrobial de-escalation, and [2] shortened duration of antimicrobial therapy. Available data suggest that the treatment should be shortened systematically [6] or based on the evolution of procalcitonin levels [7,8].

We will review the main elements that should be taken into account for treating BSIs in ICU (figure).

At the initiation of antimicrobial therapy

Early appropriate antibiotic therapy is a fundamentally important aspect of therapy of patients with BSI. Adequate treatment requires [1] that all organisms isolated from blood are susceptible in vitro to the antimicrobials chosen; [2] a proper route and dose, and [3] an early administration after blood cultures collection. In a recent large hospital cohort study, Retamar et al showed that, even fully adjusted on confounding factors, inadequate
antibiotic therapy within 24 hours was associated with a significant OR of 3 for the 14-day mortality and 1.70 for the 30-day mortality, respectively [9]. In a large cohort of septic patients, Kumar et al found that inadequate therapy within 6 hours after onset of hypotension was associated with an increase of the risk of death by more than 5-fold in case of septic shock, and by more than 9-fold if a BSI is associated to the septic shock [10]. In a very recent analysis of the surviving sepsis campaign dataset involving 17,990 patients with severe sepsis who received appropriate antimicrobials, after a careful adjustment on severity, geographic region, admission source, the mortality increased steadily by about 1 percent per hour from 24.6% when treatment was adequate within one hour to 33.1% when treatment was adequate after the 6th hour [11]. In the Eurobact study [12], an increased mortality was observed among patients who had hospital-acquired bacteremia and were hospitalized in ICU but never appropriately treated (adjusted odds ratio 1.56; 95% CI 1.04-2.35, p = 0.03). Conversely, a very early treatment (<one day after having drawn the first positive blood culture) was not associated with a decreased risk of death as compared to rather delayed appropriate treatments (<2 days and <5 days). Some kind of underdosage, associated with the increase of the volume of distribution and the paradoxical increase in glomerular filtration frequently observed in case of septic shock, may explain this result.

The initial antimicrobial therapy is necessarily empiric, targeting the most likely etiologic pathogens. When risk factors of antibiotic resistance are identified in patients with a serious infection, broad spectrum antimicrobials should be prescribed. The determinants of first-choice molecules when empirical treatment is described in Figure 1.

In a post hoc analysis among emergency department patients undergoing a quantitative resuscitation protocol for septic shock, Puskarich et al found that [13] the risk of death increased when antimicrobials were delayed until after the onset of shock (odds ratio = 2.59 95% CI 1.17-5.78). This suggests that, if an aggressive resuscitation is performed early, the timing of initial antibiotic therapy may have lesser importance.

On the opposite, when antimicrobial are started in the first hours of severe sepsis, the recent Protocolized Care for Early Septic Shock (ProCESS) trial have found no difference in mortality between an aggressive therapy of organ hypoperfusion that require the placement of a central venous catheter, administration of inotropes, or blood transfusions, a protocol-based standard therapy that did not, or an up-to-date usual care [14].

While controversy exists regarding which elements of sepsis resuscitation bundles are most beneficial, it is intuitive that good principles of care dictate that appropriate antimicrobials be prescribed promptly in the setting of severe infection.

Epidemiology of resistant bacteria
Epidemiological data about spread of MDRO has been updated by the WHO organization very recently. These data clearly found a global increase in the risk of bacterial resistance mainly gram negative bacteria with great variability between countries. The global report updated in april 2014, is available in the WHO website (http://www.who.int/drugresistance/documents/surveillancereport/en/; accessed august 22nd 2014). These general data collected mainly by hospital lab imperfectly reflect the pathogen responsible for bacteremia in ICU patients.

For community-acquired infections, recovered microorganisms are mainly Streptococcus pneumoniae and Staphylococcus aureus for Gram positive, and Escherichia coli for Gram negative. The betalactams active against these bacterias did not change in the past decade. However, extended spectrum beta-lactamase (ESBL)-producing E. coli and community acquired MRSA are more and more reported.

The fecal carriage of ESBL E. coli (mainly CTX M-15), which was unknown before the turn of the millennium, has since increased significantly everywhere, with developing countries being the most affected (Southeast Asia, Eastern Mediterranean region, Western Pacific region). Intercontinental travels may have emphasized and globalized the issue [15]. Among European subjects with traveler’s diarrhea, returning from diverse overseas areas, ESBL-E. coli carriage rates were observed in 1 out of five. These rates increased significantly after travel to Egypt, India, Southeast Asia, Thailand, and the Middle East [16]. In a prospective case-control study performed in 2008, European travelers had a 23% ESBL-E. coli carriage rate, which was significantly more than the 4% found in non travelers. Upon return from India, Africa, or Asia, the ESBL-E carriage rate reached 46% [17]. The duration of carriage after travel seems to be relatively short, lasting only a few weeks. The practical consequences of these epidemiological changes is that empirical therapy of septic shock probably due to Gram negative organisms (i.e., urinary tract infections or intra-abdominal infections) after returning from endemic areas must cover ESBL Enterobacteriaceae.

Community-acquired methicillin-resistant S. aureus (CA-MRSA) is another matter of concern. CA-MRSA responsible of community acquired pneumonia (CAP) is very uncommon in Europe [18] (less than 1% of CAP) but much more frequent (8-10%) in USA. MRSA treatment in case of CAP is not recommended [19], but
should be borne in mind in cases of severe necrotizing CAP with hemoptysis and leucopenia, often preceded by influenza infection [20].

For nosocomial and health-care-associated (HCA) bloodstream infections, resistance profile is highly dependent on national and local epidemiology. Hospital-acquired and HCA BSIs are associated with increased incidence of multi-resistant drug microorganisms (MDRO), such as MRSA, ESBL producing Enterobacteriaceae, Acinetobacter baumannii, and Pseudomonas aeruginosa. In these cases, it is more difficult for empiric treatment to be appropriate, especially in patients admitted in the ICU with a major incidence of multidrug-resistant microorganisms [21,22]. In these cases, in addition to applying guidelines recommendations, it is indispensable to know the local flora predominant in each area before initiating empiric antibiotic treatment.

Table 1 displays the distribution of organisms from recent studies investigating the epidemiology of BSIs in ICUs. The most important change in the epidemiology of ICU-related BSI in recent years is the emergence of highly resistant organisms, particularly extensively

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**Figure 1** Decision tree including main determinants guiding the choice of the most appropriate antimicrobial therapy. (BSI: bloodstream infection; MDRO: multiresistant drug organism; TDM: therapeutic drug monitoring; MIC: minimum inhibitory concentration.)

| Type of BSI | Community acquired BSI | Health care associated BSI | Hospital-acquired BSI | ICU-acquired BSI |
|-------------|------------------------|---------------------------|----------------------|-----------------|

| Use guidelines | Catheter removal, surgical drainage or debridment, puncture or drainage of empyema? |
|----------------|-----------------------------------------------------------------|

| Monotherapy | Combination therapy |
|-------------|---------------------|
| Susceptible bacteria | To broaden the spectrum of probablist Tx |
| Source control achieved appropriately | Risk of MDRO |
| Poor tissue diffusion | Source control difficult or not complete |

- Use initial high doses
- TDM when high MIC or MDRO are suspected
- Consider short duration of treatment (7-10 days)
- Adequate source control
- Susceptible bacteria with low MIC

- Adapt initial dose to: antimicrobial clearance and TDM
| Type of BSI (number of cases) | HAB (1196) | ICU-AB (330) | CAB (343) | HCAB (131) | HAB (252) | CAB (431) | HAB (351) | ICU-AB (915) | HAB (279) | ICU-AB (877) | % MDROb |
|----------------------------|------------|--------------|-----------|-----------|-----------|-----------|-----------|-------------|-----------|-------------|--------|
| S. pneumoniae              |            |              | 21.4%     | 10.9%     | 1.4%      | 0.4%      |           |             |           |             |        |
| S. aureus                  | 12.8%      | 26.7%        | 10.3%a    | 13.8%a    | 9.6%a     | 22.3%     | 26.2%     | 23.6%       | 16.1%     | 8.4%        | 60%    |
| CoNS                       | 16.6%      | 24.3%        | 3.5%      | 6.2%      | 10.8%     | 16.5%     | 19.7%     | 29.6%       | 10%       | 12.7%       | –      |
| Enterococcus spp.          | 5.5%       | 17%          | 1.5%      | 4.6%      | 8.4%      | 7.2%      | 9.1%      | 11.4%       | 10%       | 12.9%       | 43.5%  |
| E. coli                    |            |              | Gram negative 28.2% | 20.5% | 26.9 | 18.5% | 19% | 9.7% | 5.6% | 19% | 5.1% | 66.7% |
| Enterobacter spp.          | 5.8%       |              | 2.3%      | 6.9%      | 3.6%      | 2.6%      | 4.8%      | 6.3%        | 7.2%      | 7.6%        | 50%    |
| Klebsiella spp.            | 11.8%      |              | 4.1%      | 4.6%      | 6.0%      | 4.6%      | 8.6%      | 8.5%        | 8.2%      | 15.2%       | 90.9%  |
| Serratia spp.              | 3.2%       |              |           |           |           |           |           |             |           |             |        |
| Proteus spp.               | 1.8%       |              |           |           |           |           |           |             |           |             |        |
| P. aeruginosa              | 10%        |              | 1.8%      | 4.6%      | 6.4%      | 6%        | 10.3%     | 9.7%        | 10.4%     | 12.5%       | 60%    |
| Acinetobacter spp.         | 11.8%      |              |           |           |           |           |           |             |           |             |        |
| Anaerobes                  | 3.2%       |              | 1.5%      | 1.6%      |           |           |           |             |           |             |        |
| Candida spp.               | 5.8%       | 15.5%        | 1.5%      | 3.8%      | 8.4%      | 4.9%      | 10.5%     | 6.5%        | 7.5%      | 8.2%        | –      |

Legend: HAB: hospital-acquired BSI, ICU-AB: ICU-acquired BSI, HCAB: healthcare-associated BSI, MDRO: multi drug resistant organisms.

a: MRSA represents 3% of CAB, 27.5% of HCAB and 54.1% of HAB.
b: proportion of multidrug resistant organisms for both HAP and ICU-AB.
resistant Gram negative bacteria including \(P. \text{aeruginosa}, A. \text{baumannii}\) and \(K. \text{pneumoniae}\) [26-28].

**Timing of BSI onset versus hospital admission**

BSIs are generally classified as either being community-acquired (CA) or nosocomial infections [29]. However, more and more patients with complex medical problems are now managed in the community setting. Those patients managed in the outpatient setting have infections that share many clinical, microbiologic, and outcome characteristics with nosocomial infections. As a result, a new category of healthcare-associated (HCA) community-onset infections has become widely recognized [30].

Traditional risk factors of HCA infections are [1] residency in a nursing home or long-term care facility in the 30 days before the BSI. [2] hospitalization in an acute care hospital for 48 h or longer in the 90 days before the BSI. [3]. Attended a hospital or dialysis clinic or received intravenous chemotherapy in the 30 days before the BSI. [4] Received intravenous therapy, wound care, enteric nutrition, or specialized nursing care at home, in the 30 days before the BSI but the respective impact of each risk factor has never been fully investigated and probably vary between countries [31].

In hospital settings, many recent publications showed that \(S. \text{aureus}, P. \text{aeruginosa}\) is much more common in HCA-BSI than in CA BSI. \(E. \text{coli}\) is more frequent in CA BSI but the rate of ESBL E coli is similar between CA and HCA BSIs [31-37]. To the best of our knowledge there is no study that has compared microorganisms responsible from HCA-BSI and CA or HA BSI admitted in ICUs.

**Patient’s prior antimicrobial history**

If a prior antibiotic use is known, it should be considered in conjunction with previous bacterial culture and antimicrobial susceptibility testings, for setting up an appropriate treatment [38]. Most of the time, the prior use of one antibiotic is associated with an increase in the risk of infection with a microorganism resistant to it [39,40]. In area where the rate of carbapenem-resistant micro-organism is important, previous antimicrobial therapy is of key importance. The duration of exposure to carbapenems (OR \(1.079\) per day of exposure, 95% CI \(1.022-1.139\), \(p = 0.006\)) and to colistin (OR \(1.113\) per day of exposure, 95% CI \(1.046-1.184\), \(p = 0.001\)) were independent risk factors for acquisition of CR-GNB bacteremia in a Greek ICU [41]. Indeed, previous antimicrobial therapy strongly modified the gut microbiota, the main reservoir of potential pathogens in ICU patients. As an example, in our ICU, Armand-Lefevre and colleagues [42] found that imipenem resistant Gram negative bacteria (IRGNB) in the digestive flora increased regularly from 5.6% after 1 week to 58.6% after 6 weeks in the ICU.

Imipenem exposure was the main risk factor of IRGNB. As compared to patients that did not received imipenem, the odds ratio for colonization was already as high as 5.9 (95% confidence interval [43], 1.5 to 25.7) after only 1 to 3 days of exposure and increased up to 7.8 (95% CI, 2.4 to 29.8) thereafter.

**Source of infection**

The source of infection is another key element to choose empirical therapy with two objectives. We need to adapt the choice of antimicrobials to the probable agents responsible for infection and to the diffusion of antimicrobials to infected tissues.

For CA infections, pneumonia is associated with bacteremia in 15% cases. \(S. \text{pneumoniae}\) represents the main cause of bacteremia of respiratory origin, followed by \(S. \text{aureus}\), as showed by a recent large multicenter ICU cohort study [44]. Enterobacteriaceae and \(P. \text{aeruginosa}\) are encountered in less than 5% of the CA pneumonia, mainly HCA-pneumonia, but is associated with bacteremia in about half of the cases [45]. For digestive and urinary tract infection, \(E. \text{coli}\) is by far the main cause of CA BSI.

In the French national network REA RAISIN in 2013 (213 ICUs, 34 278 patients), the main causes of ICU-acquired bacteremia were intravascular catheters (29.2%), lower respiratory tract infections (18.0%) and digestive tract (13.6%). The source of ICU-acquired bacteremia remained unknown in 24.9% cases. In the same network, the main causes of catherer-related BSIs were coagulase negative staphylococci (28%), \(S. \text{aureus}\) (23%), \(P. \text{aeruginosa}\) (12%), \(E. \text{coli}\) (8%) and multiple causes (11.5%) [46].

In a post hoc analysis of the Eurobact study among 230 patients with HA-BSI respiratory infections (HABSI requiring ICU admission, \(n = 40\); ICU-acquired, non-invasively ventilated respiratory BSI, \(n = 30\); and ICU-acquired, invasively ventilated BSI, \(n = 160\)) were compared to 749 patients with HA-BSI not related to respiratory infections. HA-BSI respiratory infections were more frequently due to Gram negative pathogens (76.3% vs. 56.7%, \(P < 0.0001\)), mainly \(A. \text{baumannii}\) (18.3% vs. 10.4%, \(P = 0.0007\)), \(K. \text{lebelsiella}\) spp. (18.7% vs. 11%, \(P = 0.0013\)), and were less frequently due to Gram-positive cocci (23.3% vs. 41.2%, \(P < 0.0001\)), with the exception of \(S. \text{aureus}\) (11.3% vs. 9.5%, \(P = 0.39\)). HA-BSI respiratory infections were more frequently associated with MDRO (53.9% vs. 42.7%, \(P = 0.0003\)) [47].

In a retrospective single-center analysis of 96 patients who developed BSI of abdominal origin admitted in ICU, de Waele et al showed that secondary peritonitis, intra-abdominal abscesses and cholangitis were the sources of the BSI in the majority of patients. Gram-negative, especially \(E. \text{coli}\), were frequently recovered. BSI was polymicrobial in 22% of the cases [48]. Based on
epidemiological data and guidelines, Table 2 proposed empirical antibiotic therapy according to situations.

**Mono- or bi-antimicrobial therapy?**

Although combination therapy has been controversial in regard to a synergistic effect, it is becoming increasingly important to achieve adequate empirical therapy. Aminoglycoside use is associated with an increased risk of impaired renal function; more generally, the use of combination of antimicrobial exposed to more iatrogenic and allergic complications than a single drug. The impact of combination therapy as compared to monotherapy in resistant bacteria selection from the gut flora is also debated. Theoretically, and in experimental studies, combination therapy exposes the gut microbiota to a highest antibiotic selection pressure than monotherapy. However, in a randomized trial among ICU patients with severe sepsis, a combination of meropenem and

| Site                        | Pathogens                                                                 | Suggested empirical antibiotic                                                                 |
|-----------------------------|---------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------|
| **Urinary tract infection** | Enterobacteriaceae including                                               | - Ceftriaxone for CAB or ceftazidime for HAB or ICU AB (if suspicion of *P. aeruginosa*)       |
|                             | - *Escherichia coli*                                                       | ± aminoglycosides                                                                             |
|                             | Other enterobacteriaceae                                                  | **NB:** Consider the risk of ESBL and administer a penem in case of high risk (CA and recent travel to high risk countries, other situations in case of previous colonization, invasive procedure use, previous antibiotic therapy especially in case of septic shock) |
|                             | *Pseudomonas aeruginosa*                                                  |                                                                                                 |
|                             | *Enterococcus* sp.                                                        |                                                                                                 |
|                             | Staphylococcus sp.                                                        |                                                                                                 |
| **Intra-abdominal sepsis**  | Gram negative bacilli including                                            | - Piperacillin-tazobactam                                                                     |
|                             | - *Escherichia coli* (all BSIs)                                           | - Cephalosporin active against *P. aeruginosa* + metronidazole                                |
|                             | - other enterobacteriaceae (all BSIs)                                     | - Carbapenem (high-risk patients)                                                             |
|                             | - *Pseudomonas aeruginosa* (HA and ICU BSIs)                              | ± fluconazole                                                                                  |
|                             | *Enterococcus* sp.                                                        | ± aminoglycoside                                                                                |
|                             | Anaerobes including *Bacteroides*                                         |                                                                                                 |
|                             | *Candida*                                                                 |                                                                                                 |
| **CA and HCA pneumonia**    | *S. pneumoniae* (CA BSI)                                                  | - Third generation cephalosporin (macrolides in case of suspicion of intracellular bacteria will be added initially). |
|                             | *S. aureus* (HCA BSI)                                                    | - NB: In case of HCA, consider the risk of MRSA or ESBL E colii.                              |
|                             | *E. coli* (HCA BSI)                                                      | - Beta-lactam active against *P. aeruginosa* ± aminoglycoside ± glycopeptide or linezolid if suspicion of methicillin-resistant *Staphylococcus aureus* |
| **Hospital acquired pneumonia** | Enterobacteriaceae                                                       | - Third generation cephalosporins                                                             |
|                             | *Pseudomonas aeruginosa*                                                  | ± aminoglycosides or fluoroquinolones                                                          |
|                             | *Staphylococcus aureus*                                                   |                                                                                                 |
|                             | *Streptococcus pneumoniae*                                                |                                                                                                 |
|                             | *Haemophilus influenzae*                                                  |                                                                                                 |
|                             | Anaerobes                                                                |                                                                                                 |
| **Ventilator-associated pneumonia (no risk factor for multi-drug resistant pathogen)** | *Staphylococcus aureus* | - Glycopeptide or linezolid + beta-lactam active against *Pseudomonas aeruginosa* |
|                             | *Streptococcus pneumoniae*                                                | **NB:** Consider the risk of candidaemia in case of previous *Candida* colonization,          |
|                             | *Haemophilus influenzae*                                                  |                                                                                                 |
|                             | Anaerobes                                                                |                                                                                                 |
| **Catheter-related infection** | *Staphylococcus* sp.                                                      |                                                                                                 |
|                             | Enterobacteriaceae                                                       |                                                                                                 |
|                             | *Pseudomonas aeruginosa*                                                  |                                                                                                 |
|                             | *Candida* sp.                                                            |                                                                                                 |
moxifloxacin was associated to less emergence of resistance at the infection site to meropenem and aminoglycosides than meropenem alone [56]. Underdosage of antimicrobials and poor pharmacodynamic are certainly the main risk factors for bacterial resistance emergence among Gram negative microbiota.

In a retrospective propensity matched study, in case of septic shock, Kumar et al found a 23% reduction in the risk of death with the use of immediately adequate combination therapy as compared to immediately adequate monotherapy [57]. But in a cohort of 593 ICU patients with *P. aeruginosa*, BSI, there was no demonstrated benefit of a combination therapy if the monotherapy is adequate [58]. In a cohort of 760 patients with severe sepsis or shock, inappropriate initial therapy increased the risk of death (adjusted OR: 2.3); combination therapy decreased the rate of inappropriate therapy from 36% to 22.2% [59].

The appropriate combination is also debated. The benefit of combination with aminoglycosides has not been found in the study by Kumar et al. [57]. On another cohort of Gram negative bacteremia, combination with aminoglycosides improved the appropriateness of antimicrobial therapy in episodes due to ESBL or AmpC producing *Enterobacteriaceae* and to *P. aeruginosa*. Combination with aminoglycosides also improve prognosis of patients with neutropenia and septic shock [60].

**Right dosage**

Prescription of an antibiotic for which the microorganism is supposed to be susceptible *in vitro* is not synonymous of efficacy. It is especially the case for the most severe patients (with increase in the volume of distribution of antimicrobials) and for the less susceptible bugs, for which available antimicrobials have MICs at the border of susceptibility breakpoints.

Pharmacokinetic/pharmacodynamic (PK/PD) targets need to be achieved for decreasing the risk of treatment failure and selection of resistant pathogens [61].

The main pharmacodynamic properties that correlate with efficacy for largely used antimicrobials are the time above the MIC for beta-lactams and carbapenems, the ratio maximal concentration/MIC (Cmax/MIC) for fluoroquinolones, aminoglycosides, daptomycin, and colistin, and the area under the curve above the MIC (AUIC) for glycopeptides.

Estrely et al [62] studied the impact of MICs to carbapenems of multiresistant Gram negative bacteria (*P. aeruginosa*, *A. baumannii* and Beta-lactamases producer *Enterobacteriaceae*) on the outcome of 71 patients with BSIs. Overall, 52 patients survived, and 19 died. This study revealed that patients with organisms that had a MIC of >4 mg/L had worse outcomes than patients whose isolates had a MIC of <2 mg/L, even after adjustment for confounding variables (16.1% versus 76.9%; P < 0.01). Indeed, it has been demonstrated that beta-lactams used in critically ill infected patients achieved infrequently their pharmacodynamic target. Usual targets for beta lactams are free antibiotic concentration above the MIC (fT > MIC) of the pathogen at both 50% and 100% of the dosing interval. In the recent DALI study, involving 248 patients with infection, 16% of the beta-lactam administration did not achieve the 50% target and 32% did not achieve the 100% target. Importantly, a positive clinical outcome was associated with 50% and 100% fT > MIC (OR = 1.02 and 1.56, p < 0.03) [63]. In practice, we should know the MIC50 and MIC90 of the suspected pathogens, and ask for specific MICs after pathogen identification, especially for MDRO and extensively resistant pathogens.

For hydrophilic antibiotics such as beta-lactams, aminoglycosides, glycopeptides or colistin, the increase in the volume of distribution observed in severe patients leads to a decrease of the antibiotic concentrations. Furthermore, in young people without immediate renal dysfunction, a paradoxical increase in the glomerular filtration is observed. Both mechanisms argue for the administration of an important initial dose of antimicrobials (loading dose), in order to reach pharmacodynamic targets at the initiation of treatment (suggested initial doses are in Table 3). For lipophylic antibiotics such as fluoroquinolones, the volume of distribution is grossly unchanged and clearance depends on hepatic function.

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**Table 3 Suggested intravenous of initial selected antibiotic doses in critically ill patients during the first 24 hours of treatment of bacteremia with severe sepsis in ICU (to be adapted in case of kidney injury and renal replacement therapy) (adapted from [61,63-66])**

| Antibiotic class   | Amikacin 25 mg/kg | Gentamicin 7 mg/kg | Interval administration and doses adjusted according to TDM |
|--------------------|-------------------|-------------------|-------------------------------------------------------------|
| Aminoglycosides    |                   |                   |                                                             |
| Fluoroquinolones   | Ciprofloxacin 400 8 hourly |                   |                                                             |
| Colistin           | 9-12 MU (720-960 mg) loading dose followed by 480 mg 12 hourly if patient without kidney injury. |                   |                                                             |
| Beta-lactams       | Cefepime 2 g 8 hourly | Ceftazidime 2 g 6 hourly |                                                             |
|                    | Imipenem 1 g 6-8 hourly | Meropenem 1 g 6-8 hourly |                                                             |
|                    | Ertapenem 1 g 12 hourly | Piperacillin-Tazobactam 45 g 6 hourly |                                                             |
| Glycopeptides      | Vancomycin 35 mg/kg in a 1 hour infusion loading dose followed by 30 mg/kg continuous infusion |                   |                                                             |
| Daptomycin         | 8-12 mg/kg 24 hourly |                   |                                                             |
| Tigecyclin         | 200 mg followed by 100 mg 12 hourly when borderline susceptibility is suspected |                   |                                                             |
For time-dependent antimicrobials such as beta-lactams or glycopeptides, the use of continuous or extended infusions of antimicrobial is able to increase the $\text{fT} > \text{MIC}$. Continuous infusion required the antimicrobial to be stable after preparation at room air temperature. It has been tested successfully with ceftazidime, cefepime, piperacillin-tazobactam, doripenem, meropenem and vancomycin. The clinical proof of superiority of extended or continuous infusion of beta-lactams or vancomycin is suggested by few studies [64,67-71], and required further trials.

For concentration dependent antimicrobials, such as aminoglycosides [72,73] or colistin [65,66], the use of increased doses increased the probability to reach the pharmacodynamic targets.

Therapeutic drug monitoring (TDM) is common place, given uncertainties on the antimicrobial concentration actually achieved, especially for aminoglycosides and glycopeptides. Recent data have also shown that TDM of beta-lactams leads to dose adjustment in about half of cases [63,74].

In case of kidney injury, hydrophylic antibiotics clearance will be largely modified. Intensity of continuous renal replacement therapy and degree of residual renal function are crucial factors in accurately determining the antimicrobial requirement.

Right after the initiation of the antimicrobial therapy

Results of microbiological exams

Despite all previous knowledge, antibiotic therapy remained initially inadequate in 20-40% of the cases, even if broad spectrum combination therapy is used. Microbiological samples, collected just before antimicrobial start, may help to document the infection and decrease the risk of inadequate treatment. Good collaboration with the microbiological laboratory, Gram stain examination and new molecular techniques able to detect early various antimicrobial resistance mechanisms are important strategies to be developed in order to shorten the delay for appropriate treatment and subsequent de-escalation [75-78].

Source control

Source control is recognized as an important part of the treatment of BSIs and has been recently shown to be independently related with outcome [12,79]. The source control should be optimal, especially when microorganisms are less susceptible to antimicrobials or when antimicrobial diffusion is hazardous.

During the antimicrobial therapy

De-escalation

Antimicrobial de-escalation is a clinical approach to empirical antibiotic treatment of bacteremia that attempts to balance the need for appropriate initial therapy with the need to limit unnecessary antimicrobial exposure, in order to curtail the emergence of resistance.

Paradoxically, although de-escalation is associated with no or a positive impact on mortality [80], its impact on overall antibiotic consumption has not been clearly demonstrated. It has even been associated with an increase in the number of days on antibiotics [81]. However, it clearly reduced the broad-spectrum antibiotic use. The value of such streamlining may seem intuitive to infectious diseases physicians, but for many other physicians, an attitude of “don’t change a winning team” is pervasive. The persistently reported figures of around 50% de-escalation delivery are unacceptable and must be improved. The requirement for multiple antimicrobials to achieve adequate initial therapy further emphasizes the need for obtaining both timely blood culture and early culture results through incorporation of molecular diagnostics for bacteremia [78,82] so that subsequent antimicrobial de-escalation can be performed in a rational manner [82].

Length of antimicrobial therapy

Up to half of the antibiotic use is unnecessary or inappropriate, and excessive duration is the greatest contributor to inappropriate use. A reduction in the length of antibiotic courses in ICU is an important strategy to minimize the consequences of antimicrobial overuse, including antibiotic resistance, Clostridium Difficile-associated diarrhea, adverse effects and costs. Usual length of therapy is about 10 days in routine, longer for S. aureus bacteremia and shorter for coagulase-negative staphylococci bacteremia [83].

Although large RCTs are missing, a meta-analysis of available data suggested that a short (5-7 days) therapy, vs. a longer one (7-21 days), is safe with respect to clinical or microbiologic cure and survival [6]. Of note, aminoglycosides treatment should be limited to 2-5 days to optimize efficacy and to limit potential toxicity.

Table 4 Arguments pro and against a short duration of antibiotic therapy

| Arguments for a short treatment | Arguments for a longer treatment |
|---------------------------------|---------------------------------|
| No comorbid conditions          | Immune depression               |
| Source control                  | No source control               |
| Low MICs, high bacterial titers | MDR, XDR bacteria               |
| Initial appropriate therapy     | Low bacterial titers            |
| Easy PK and tissue diffusion    | Poor PK and tissue diffusion    |
| Source control appropriate      | Foreign materials               |
| No foreign material             | Slow, partial clinical response |
| Rapid clinical improvement      |                                |
Of course, at the bedside, many factors should be taken into account to decide to shorten the duration of treatment (Table 4). A short duration of treatment should be more cautiously decided when the initial treatment was inadequate [84], when the therapeutic index was low, when the source control was not optimal. In immunocompromised patients, and when invasive devices or foreign material are left in place, a longer treatment is often recommended.

To reduce the duration of treatment, strategy should also be guided by the evolution of patients’ status under treatment and by biomarkers [85]. Procalcitonin (PCT) serum level has been used in randomized control trial to reduce the duration of therapy in severe infections. A PCT-driven approach was responsible for an increase of 3.5 of the number of days alive without antibiotics at day 28, with no adverse impact, as compared to the standard of care. Surprisingly, the PCT rules are not followed by clinicians in about half of the cases, although it suggests that regular monitoring of PCT may serve as a target to propose stopping antimicrobials.

The use of external rules of therapy, of systematic intervention of the antibiotic team of the hospital [86,87] or bundles [88], simple educational programs with audit, are also effective in reducing the antimicrobial consumption to a similar extent without adverse events.

Finally individual patient therapy needs to be coupled with adherence to optimal prevention practice to prevent spread of resistant bacteria from patient to patient.

Conclusions
Bacteremia is an independent risk factor of mortality in case of severe sepsis in ICU. Early adequate treatment of bloodstream infections is required and is based on previous knowledge and guidelines, rapid microbiological identification, adequate source control and administration of proper antimicrobials at the adequate dose regimen for the individual patient. These goals become difficult to achieve in case of BSI due to multi-drug resistant pathogens with high MICs to antimicrobials, in the most severe patients where low antibiotic concentrations are common, and when the source control is incomplete or impossible. In these situations, pharmacodynamic optimization and therapeutic drug monitoring should be recommended.

Competing interests
MW gave lecture for Pfizer, Novartis, GSK, Astra-Zeneca, BM gave lecture for Novartis, JFT gave lecture for Pfizer, Astellas, Merck, Novartis and received research grants from Merck, and Astellas. Other authors nothing to declare.

Authors’ contributions
JFT wrote the first draft. JFS and GV suggested major modifications. All authors made critical review of the manuscript. All authors read and approved the final manuscript.

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