The Management of Hematologic Patients with Bloodstream Infections Due to Multi-Drug Resistant Bacteria: Where Do We Stand? From Antibacterial Prophylaxis to the Treatment of Septic Shock

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Abstract: Bloodstream infection (BSI) and septic shock represent one of the major limiting factors for the successful treatment of patients affected by hematologic malignancies. During the most recent years we have documented a shift in the epidemiology of bacterial infections toward a consistent rise of Gram-negative rods. In addition, the emergence of multi-drug-resistant bacteria is considered a life-threatening condition requiring a multi-disciplinary approach. Aim of present review is to summarize the most recent approaches in terms of anti-microbial prophylaxis and treatment of BSI in hematologic patients with neutropenic fever.

Keywords: bloodstream infections; multi-drug-resistant bacteria; septic shock

1. Introduction

Despite remarkable improvements have been achieved in the management of patients with malignant hematologic disorders, infectious complications still represent a leading cause of morbidity and mortality, particularly during the neutropenic phase after the chemotherapy treatment.

Bloodstream infection (BSI) affects 11% to 40% of neutropenic patients, with an associated mortality ranging from 5% to 60% in cases of multi-drug-resistant (MDR) organisms BSI [1–5]. The epidemiology of BSI has undergone substantial changes in the most recent years: during the 1990s, Gram-positive BSI were the leading cause of BSI due to the increased use of intravascular devices and the extensive use of prophylaxis with fluoroquinolones (FQ) [6]. This trend was followed by the gradual rise of Gram-negative rods, with extended-spectrum beta–lactamase (ESBL), multi-drug resistance (MDR) and carbapenem-resistant Enterobacteriales (CRE) as well as MDR P. aeruginosa and Acinetobacter spp. as critical issues in daily clinical practice. A recent review by the European Conference on Infections in Leukemia (ECIL) group showed a change of the Gram-positive to Gram-negative ratio to 55–45% from 60–40%, reported by studies published between 2005 and 2011 [7].

According to this observation, the potential benefit of fluoroquinolone (FQ) prophylaxis in terms of the reduced rate of BSI needs to be weighted against its impact in terms of increased risk of colonization with MDR strains. Aim of the present review is to summarize the current epidemiology of BSI in patients with hematologic malignancies and hematopoietic stem cell transplant (HSCT) recipients, to provide a critical appraisal of the usefulness of FQ prophylaxis, to review the contemporary...
epidemiological trends and to define the most appropriate antibiotic treatment in febrile neutropenia considering rising anti-microbial resistance.

1.1. Etiology

MDR bacteria are increasingly recognized in hematology patients, and the prevalence of different resistant species varies significantly between centers [7–9]. Bacteria have been defined as MDR if they are resistant “to three or more antibiotic classes” or according to other authors, resistant “to one key anti-microbial agent” [10]. The spectrum of MDR bacteria identified in hematologic patients is mostly limited to Gram-negative rods. Traditionally *P. aeruginosa* infections have been frequently encountered in hematological population and recently its management got complicated by the raising of MDR strains [7–9]. Several European countries did already show an increase in extended-spectrum β-lactamase (ESBL), with the recent addition of carbapenem-resistant *K. pneumoniae*, endemic in Italy and Greece, and MDR Acinetobacter baumannii [11]. Within Gram-positive infections, the prevalence is due to methicillin-resistant *S. aureus* (MRSA) and Enterococci. However, vancomycin-resistant enterococci, especially in HSCT recipients, remain a troublesome and actual nosocomial infection, with significant geographical variability [7–9,12,13]. A prospective study from the European Society for Blood and Marrow Transplantation (EBMT) analyzed the emergence of MDR Gram-negative rods during the first six months following HSCT among 65 centers in Europe, Australia, and Asia. Overall, 655 Gram-negative episodes in 591 patients were included in the final analysis: 414 in allogeneic and 241 in autologous hematopoietic stem cell (HSC) recipients. Enterobacteriaceae accounted for 73% of the bacteremia; non-fermentative rods, for 24%; and others, Gram-negative for 3% of the episodes. Overall, 50% of the isolates were FQ and non-carbapenem-resistant, 18.5% were carbapenem-resistant, and 35% were MDR. The rate of FQ-resistant (FQ-R) organisms was significantly higher in centers providing prophylaxis (79% vs. 50% in allogeneic HSCT and 74% vs. 25% in autologous HSCT) and in patients receiving an allogeneic HSCT compared to autologous HSCT (56% vs. 41%, *p* < 0.001). Similarly, the rate of MDR isolates was significantly higher in patients receiving FQ prophylaxis as compared to those who did not (35% vs. 8%, *p* < 0.001) [14].

Tricarichi et al. found a 58% rate of carbapenem resistant (CR) among 278 *K. pneumoniae* BSI (KP-BSI) in patients with hematologic malignancies; the higher incidence of *K. pneumoniae* carbapenem resistant (KP-CR) was identified in acute myeloid leukemia (AML) patients (74%) but acute lymphoblastic leukemia (ALL) and non-Hodgkin lymphoma (NHL) did not have a negligible rate of KP-CR (7% and 11% respectively). The presence of KP-CR translated into a high mortality rate compared to carbapenem-susceptible KP (52% vs. 14%) [15].

A more recent study investigated the changing epidemiology of BSI in HSCT recipients over a 25 years period: 1164 BSI have been included in the analysis occurring at a median time of 55 days post-HSCT. The rate of Gram-positive BSI decreased from 68% in 1993–2012 to 52% in 2013–2017 while there was the concomitant increase of Gram-negative BSI from 32% to 42% as well as of MDR Gram-negative bacteria from 1% up to 10% [16]. Table 1 shows the major risk factors for developing Multi-Drug Resistant bacterial infections.
Table 1. Risk factors for Multi-Drug resistant bacterial infections

<table>
<thead>
<tr>
<th>MRSA</th>
<th>VRE</th>
<th>CRE</th>
<th>MDR PA</th>
<th>AB</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (HA-MRSA &gt; 65 ys, CA-MRSA younger)</td>
<td>Advanced age (&gt;65 ys)</td>
<td>Advanced age (&gt;65 ys)</td>
<td>Advanced age (&gt;65 ys)</td>
<td>Advanced age (&gt;65 ys)</td>
</tr>
<tr>
<td>Prolonged or prior hospitalization (&gt;HA-MRSA)</td>
<td>Transplant recipient</td>
<td>Transplant recipient</td>
<td>Neutropenia/immunosuppression</td>
<td>Critically ill patients admitted to ICU</td>
</tr>
<tr>
<td>ICU admission</td>
<td>HD</td>
<td>Prior and prolonged ATB use last 90 days (notably glycopeptides, FQ, CIII-IV, anti-anaerobes, carbapenems)</td>
<td>Use of broad spectrum antibiotics</td>
<td>Prior and prolonged abt use last 90 days (notably AG, FQ, CIII and carbapenems)</td>
</tr>
<tr>
<td>HD</td>
<td>Prolonged or prior hospitalization</td>
<td>Presence of invasive devices</td>
<td>Malnutrition</td>
<td>Burns and surgical wounds</td>
</tr>
<tr>
<td>Indwelling line, catheters</td>
<td>ICU admission</td>
<td>ICU admission (with or without MV)</td>
<td>Structural lung disease</td>
<td>High CCI (notably diabetes, chronic lung, liver and renal diseases)</td>
</tr>
<tr>
<td>Invasive procedures or recent surgery</td>
<td>Surgery</td>
<td>Decubitus ulcer</td>
<td>Malignancy</td>
<td>Prolonged or prior hospitalization</td>
</tr>
<tr>
<td>Institutionalization</td>
<td>High CCI</td>
<td>Prolonged or prior hospitalization</td>
<td>Prolonged or prior hospitalization</td>
<td>Malignancy</td>
</tr>
<tr>
<td>High CCI</td>
<td>Invasive devices</td>
<td>Poor functional state</td>
<td>Prior and prolonged ATB use last 90 days (notably AG, FQ, CIII and carbapenems)</td>
<td>Transplant recipient</td>
</tr>
<tr>
<td>PWID</td>
<td>Prior and prolonged abt use last 90 days (notably vancomycin, FQ, CIII, anti-anaerobes, carbapenems)</td>
<td>Indwelling devices</td>
<td>Presence of invasive devices</td>
<td>Presence of invasive devices</td>
</tr>
<tr>
<td>Athletes (&gt;CA-MRSA)</td>
<td>Malignancy</td>
<td>Surgical interventions</td>
<td>Transplant recipient</td>
<td>HD</td>
</tr>
<tr>
<td>Prior and prolonged ATB use last 90 days (&gt;HA-MRSA)</td>
<td>Neutropenia/immunosuppression</td>
<td>Transfer between units</td>
<td>High CCI (notably diabetes, chronic lung, liver and renal diseases)</td>
<td>Severe underlying illness</td>
</tr>
<tr>
<td>Crowded Living Environment (&gt;CA-MRSA)</td>
<td>Institutionalization</td>
<td>High CCI (notably diabetes, chronic lung, liver and renal diseases)</td>
<td>ICU admission</td>
<td>Neutropenia/immunosuppression</td>
</tr>
</tbody>
</table>

**Abbreviations:** MRSA—Methicillin-Resistant Staphylococcus aureus; HA—hospital-acquired; CA—community-acquired; ICU—intensive care unit; HD—hemodialysis; CCI—Charlson Comorbidity Index; ATB—antibiotic; PWID—people who inject drugs; VRE—vancomycin-resistant enterococci; FQ—fluoroquinolones; CIII—third generation cephalosporins; CRE—carbapenem-resistant Enterobacteriaceae; MDR—multi-drug resistant; PA—Pseudomonas aeruginosa; ys.: years; AG—aminoglycosides; AB—Acinetobacter baumannii.
1.2. MDR Colonization and BSI

Several studies evaluated the relationship between MDR colonization and the risk of developing BSI. Girmenia et al. found that a KP-CR colonization before or after the transplant was followed by a BSI in 25% of the cases of autologous and 39% of the cases of allogeneic HSCT [17]; infection-related mortality was 16% among autologous and 70% among allogeneic HSCT recipients. A subsequent Italian prospective study analyzed the epidemiology of pre-engraftment Gram-negative BSIs in patients receiving allogeneic (n = 1118) and autologous (n = 1625) HSCT. Overall, the cumulative incidence of pre-engraftment Gram-negative BSI was 17% in allogeneic and 9% in autologous HSCT recipients. In allogeneic HSCT patients, *E. coli* accounted for 52% of Gram-negative BSI, KP for 19% and *P. aeruginosa* (PA) for 14% of BSI. The presence of pre-HSCT colonization from KP−CR was associated with a 28% probability of developing a post-HSCT BSI from the same isolate, while the risk of KP-CR BSI for patients who were not colonized before the graft was only 0.4%. Similar results were shown for MDR PA (28% vs. 0.6%), ESBL *E. coli* (9% vs. 1.3%) and ESBL *K. pneumoniae* (7% vs. 0.4%) [18].

Staphylococci are the most frequent Gram-positive pathogen isolated in BSI and coagulase-negative Staphylococci (CoNS) are responsible for approximately a quarter of BSI, while *S. aureus* is found in around 5% of cases [2]. *S. aureus* is more virulent than CoNS; in particular, if methicillin resistance occurs, mortality is higher. Risk factors for MRSA are, among the others, previous nasal colonization and the presence of a central venous line.

Enterococci are commensals of the gastrointestinal (GI) tract and can affect 10–12% of transplant recipients [3]. *E. faecalis* used to be the main strain encountered even if in some centers *E. faecium* is replacing it. For both strains, *E. faecalis* and *E. faecium* vancomycin resistance has increased, causing a higher mortality rate [6]. Viridans streptococci are found in approximately 5% of BSI with a mortality and morbidity not negligible [4] with consequent high mortality. The frequent use of vancomycin in the empiric therapy is possibly associated with a higher rate of pharmacological pressure and secondary risk of resistant pathogen selection.

2. Antibiotic Prophylaxis

Several meta-analyses addressing the issues of anti-microbial prophylaxis in neutropenic patients have been published. The Cochrane evaluated 109 trials between 1973 and 2010, including 70 studies in patients with hematologic malignancies [19]. Different antibiotic prophylactic regimens have been administered, namely FQ, trimethoprim-sulfamethoxazole (TMP−SMZ), non-absorbable antibiotics, rifampicin, and even IV prophylaxis. All-cause mortality resulted in being significantly reduced with FQ prophylaxis compared to placebo or no treatment as well as infection-related mortality, the number of febrile episodes and Gram-negative/Gram-positive BSI. On the other hand, compared to placebo, patients receiving prophylaxis were more likely to harbor resistant bacteria to the specific drug.

Kern et al. prospectively analyzed the impact of FQ prophylaxis on BSI in a large number of patients who received different treatments, for instance high-dose chemotherapy for acute leukemia, autologous or allogeneic HSCT. Overall, the use of prophylaxis was associated with less Gram-negative BSI but a greater incidence of Gram-positive BSI. In addition, patients receiving FQ prophylaxis had significantly more BSI due to ESBL-positive Enterobacteriaceae. All-cause mortality was positively affected by prophylaxis only in patients receiving high-dose chemotherapy for acute leukemia [20].

A recent meta-analysis investigated the efficacy of levofloxacin in patients with acute leukemia. Prophylaxis reduced the incidence of febrile neutropenia and microbiologically documented infections but did not have effect on mortality [21].

Kimura et al. reported the results of a meta-analysis evaluating 17 trials including 1453 patients who received autologous (n = 842) or allogeneic HSCT (n = 407) [22]. Similarly, to the previous studies, systemic antibiotic prophylaxis reduced the number of febrile episodes, clinically or microbiologically documented infections and BSI, while all-cause mortality and infection-related mortality did not result affected by the antibiotic prophylaxis.
An Italian study analyzed 342 cases of *E. coli* BSIs occurring in onco-hematologic patients between 2016 and 2017. In multivariate analysis, FQ prophylaxis emerged as a risk factor for the emergence of ESBL *E. coli* [7].

**Guidelines**

The ECIL guideline published in 2007 recommended FQ prophylaxis based on two randomized trials and one meta-analysis. Subsequently, indications for antibiotic prophylaxis were rather discordant with some groups still maintaining the indication and others discouraging the use of prophylaxis. Nevertheless, over the most recent years the emergence of MDR bacteria strongly correlated with the wide use of prophylaxis, raised concerns for usefulness of this strategy. According to these observations, the American Society of Clinical Oncology (ASCO) and the Infectious Diseases Society of America (IDSA) guidelines published in 2018 recommend FQ prophylaxis in high-risk patients, namely those with acute leukemia and HSCT recipients [23].

The German guidelines reinforce the concept that FQ prophylaxis demonstrated its efficacy in reducing the incidence of Gram-negative sepsis during neutropenia without significant changes in mortality [24]; in particular at institutions with a low rate of MDR Gram-negative bacteria, antibacterial prophylaxis remains a reasonable choice to reduce the incidence of Gram-negative sepsis during neutropenia (AII). In patients with chronic graft versus host disease (cGVHD) on immunosuppressive therapy, antibiotic prophylaxis against encapsulated bacteria might be useful until immunizations can be applied (BIII). A summary of recommendations is given in Figure 1 according to the main international guidelines in hematological patients.

**Figure 1.** Guidelines recommendations of antibiotic prophylaxis in hematologic patients. Abbreviations: GN—Gram-negative; FQ—fluoroquinolone.

### 3. Sepsis/Septic Shock

Sepsis and septic shock represent a major challenge for physicians treating neutropenic hematological patients. The high rate of morbidity and mortality and the need for prompt intensive care, related to the peculiar clinical characteristics and the immunocompromised status of the patients, are of particular concern and require a multi-disciplinary approach. The prevalence of sepsis in neutropenic hematological patients is not precisely defined, and most studies do not analyze specific subgroups as immunocompromised patients.
3.1. Definition

Since 1991, sepsis has been described as a host’s systemic inflammatory response syndrome (SIRS) to infection, defined by two or more of the following criteria: temperature >38° or <36°, heart rate >90/min, respiratory rate >20/min, white blood cell count >12,000/mm³ or <4000/mm³ or >10% immature bands. The 2016 Surviving Sepsis Campaign (SSC), overcoming the previous SIRS criteria that can reflect an adaptive and appropriate response to infection, defined sepsis as a life-threatening organ dysfunction caused by a dysregulated host response to infection and septic shock as a subset of sepsis with circulatory and cellular/metabolic dysfunction associated with a higher risk of mortality. Both the diagnosis and treatment of sepsis and septic shock in hematological patients should not differ from those used in the general population [25–27].

The SSC and especially the 2018 Update Bundle remark that sepsis is a medical emergency and resuscitation, and management must start immediately (hour-1 Bundle) [28].

A bundle is a series of interventions that together or at a multi-disciplinary level, result in a reduction of an event or an improvement in the quality of care or in the prevention efforts.

3.2. Diagnosis

In the setting of a febrile neutropenic hematological patient, the diagnosis of sepsis may be challenging, and fever can be the only manifestation of an underlying critical condition.

Sepsis is a clinical diagnosis, but different scores have been developed, mainly in the settings of emergency medicine or the intensive care unit (ICU), to identify patients likely to have sepsis. The most used scores are the sequential organ failure assessment (SOFA) score that comprehends a set of laboratory data (as platelet count, creatinine and hepatic function) and the quick SOFA, a clinical score that provides simple bedside criteria (respiratory rate ≥22/min, GCS < 15 and a systolic blood pressure ≤100 mmHg). Compared to SIRS criteria, these scores can be altered by a massive dysregulated response of the organism to infection and represent the mainstay of a rapid diagnosis. Sepsis is defined as an infection with a qSOFA and a SOFA score ≥2. Septic shock is considered a subset of sepsis with persistent hypotension requiring a vasopressor to maintain a mean arterial pressure (MAP) ≥65 mmHg despite adequate fluid resuscitation and an increase in serum lactate ≥2 mmol/L [29].

The value of these scores in the neutropenic hematological population is less established. A recent study considering both hematological and solid cancer patients shows that the SSC’s new definition of sepsis and septic shock is applicable in an ICU setting with the same reliability as in the general population. In the ICU, the SOFA score seems more accurate than qSOFA and SIRS criteria to predict hospital mortality; however, outside of the ICU, qSOFA is useful and easily calculated [30].

4. Treatment

The following issues should be taken into account when we consider the initial antibiotic treatment of a neutropenic hematological patient with fever:

- High-risk patients (those with an expected severe neutropenia duration ≥7 days, for example HSCT recipients or this candidate for intensive chemotherapy);
- Hemodynamic stability and illness severity;
- Previous colonization by MDR bacteria, detected on routine swabs (ESBL or carbapenemases-producing Enterobacteriales (CPE), MRSA, VRE);
- Previous infections by MDR bacteria;
- Local epidemiological resistance patterns [31].

The initial treatment of neutropenic fever should consider all these variables. Important risk factors for developing MDR infections are exposure to broad-spectrum antibiotic therapy, especially within the last month before the infectious episode, in-hospital/ICU staying, prior colonization, indwelling devices [32,33], all frequently found in HSCT recipients and hematological patients. BSI from MDR
Gram-negative bacteria is associated with significantly lower overall survival (OS) compared to hematological patients with non-MDR Gram-negative BSI [15,34,35].

There are two possible approaches when starting an empiric antibiotic treatment in patients with neutropenic fever: escalation or de-escalation.

- The 2010 IDSA guidelines recommend that if no risk factors for complicated fever or MDR infection are present or suspected and the patient is stable, an escalation strategy should be performed, starting from a monotherapy with an anti-pseudomonal B-lactam agent, such as cephalosporin (e.g., cefepime), carbapenems (meropenem or imipenem–cilastatin) or piperacillin–tazobactam, covering the most common bacteria (AI) [36]. The guidelines report rising resistance rates among Gram-negative bacteria with the use of ceftazidime monotherapy, with the limited activity of this agent against Gram-positive bacteria. The latest ECIL guidelines are in line with this escalation approach in non-colonized, uncomplicated patients. Monotherapy with piperacillin/tazobactam or cefepime and ceftazidime (the latter not to be used as empirical monotherapy in a setting of high ESBL prevalence) is a valid option (AI). Altogether, in patients without critical presentation and no risk factors for resistance to other agents, ECIL recommends a carbapenem-sparing approach [32].

- Antibacterial spectrum should be broadened in cases of clinical worsening, documented infection, or isolation of resistant microorganisms. An anti-Gram-positive agent should be added if such an infection is suspected: evidence of central venous catheter (CVC) dysfunction or infection, pneumonia, hemodynamic instability, soft tissue, or skin infection.

- In case of known colonization with MDR bacteria, mostly consistent with the use of whatever strategy of colonization study, a previous infection with resistant bacteria or a critical clinical presentation, a de-escalation approach is preferred, aiming at covering a wide spectrum of bacteria while awaiting the final etiological diagnosis and the possibility of de-escalate, then reducing the number of antibiotics or the spectrum of antibacterial treatment. Anti-microbial de-escalation is thought to decrease exposure to broad-spectrum antimicrobials and to prevent the emergence of anti-microbial resistance. Valid options of treatment are carbapenem monotherapy, for example in case of suspected ESBL Gram-negative infection, or combinations, as β Lactam/β-lactamase inhibitor (BLBLI) plus aminoglycoside. A randomized trial on 379 patients with ESBL Gram-negative BSI (13% neutropenic) showed a survival advantage of meropenem with piperacillin/tazobactam monotherapy, although patients were exposed to very heterogeneous empirical antibiotic regimens before randomization [37]. In confirmed MDR BSI a combination treatment is required and old antibiotics (like fosfomycin and colistin) should not be given as monotherapy but with other drugs regarding the potential risk of generating resistant strains. Figure 2 summarizes bacterial susceptibility to the most relevant antibiotics.

Early reassessment at 48–72 h and narrowing of the antibiotic spectrum until discontinuation are advised when the patient is hemodynamically stabilized or a targeted pathogen is isolated. Early de-escalation and discontinuation also applies for HSCT recipients. A study from Snyder et al. on 120 HSCT recipients showed that de-escalation after five days of broad-spectrum therapy and defervescence did not increase the rate of recurrent fever and the clinical outcome, compared to those who continued antibiotic treatment until neutropenia recovery [38]. These considerations are confirmed by a study of Gustinetti et al. on 110 HSCT patients in the pre-engraftment phase, where de-escalation therapy was feasible in 60% of patients. Failures of early or late de-escalation occurred in 10 patients (17.5%); however, no cases of septic shock or death have been reported [39]. This approach is safe even in critically ill neutropenic patients in the ICU setting [40].
**Figure 2.** First-line antibiotic treatment in hematologic patients with neutropenic fever. Abbreviations: ESBL: extended-spectrum beta-lactamase; CRE: carbapenem-resistant Enterobacteriaceae; MDR PA: multi-drug-resistant Pseudomonas aeruginosa. C/A: ceftazidime/avibactam; C/T: ceftolozane/tazobactam; AG: aminoglycosides; PTZ: piperacillin-tazobactam; M/V: meropenem/vaborbactam.

<table>
<thead>
<tr>
<th>Patient Not Colonized</th>
<th>Patient Colonized</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>ESBL</td>
</tr>
<tr>
<td><strong>Carba-sparing strategies</strong></td>
<td></td>
</tr>
<tr>
<td>C/T ± AG</td>
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<tr>
<td>C/T ± Fosfomycin</td>
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<td>C/A ± AG</td>
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<td>C/A ± Fosfomycin</td>
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<td>PTZ + Fosfomycin</td>
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<td>PTZ ± AG</td>
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<tr>
<td><strong>Carba-based strategies</strong></td>
<td></td>
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<tr>
<td>Meropenem (or Imipenem or Ertapenem) ± AG</td>
<td></td>
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</tbody>
</table>

**To be avoided: «old» cephalosporins (i.e., ceftriaxone, ceftazidime)**

**Carba-sparing strategies**
- C/T + Colistin (± Fosfomycin)
- C/A + Colistin (± Fosfomycin)
- C/A or C/T + Aztreonam
- Cefiderocol

**Carba-based strategies**
- Meropenem (or Imipenem) ± AG or Colistin or Fosfomycin or Aztreonam
- M/V or I/R ± Fosfomycin or Tigecyclin or AG or Aztreonam or Colistin

**To be avoided: Tigecycline, Ertapenem**

*According to local epidemiology, previous ATB regimens and phenotypic and genotypic pathogen susceptibility.*

Based on previous infections/severity of current infection/comorbidities/PK-PD.
A study from Gutierrez−Gutierrez on 480 patients with CPE−BSI showed that initial appropriate anti-microbial therapy was associated with 30-day lower mortality compared to patients who received inappropriate therapy (38.5% vs. 60.6%). Among those receiving appropriate therapy, 39% received combination therapy, and 61% received monotherapy. Overall mortality was not different between those receiving combination therapy or monotherapy (35% vs. 41%). However, combination therapy was associated with lower mortality than was monotherapy in the high-mortality-score stratum (patients with sepsis/septic shock, Pitt score > 6, Charlson score > 2), but not in the low-mortality score stratum [41].

Even in neutropenic septic patients, the first option is represented by an anti-pseudomonal broad-spectrum agent (e.g., piperacillin/tazobactam or meropenem). The use of a combination therapy with aminoglycosides in neutropenic patients with sepsis is a matter of debate, with some studies showing any significant advantage and increasing renal toxicity [42]. By contrast, Krumar et al. showed that early combination antibiotic therapy (β-lactam plus aminoglycoside, FQ or macrolides/clindamycin) improved survival in critically ill neutropenic patients compared to monotherapy [43]. One recent study showed that in neutropenic hematological patients with sepsis or septic shock, aminoglycoside combination therapy was not associated with increased renal toxicity and led to a reduction in the mortality rate [44]. According to these observations, the main international guidelines recommend the use of combination therapy including aminoglycosides in critically ill patients [25,32,36].

A de-escalation policy may be considered wise, as soon as a specific pathogen is identified or in case the absence of any isolate and clinical improvement is documented [25,45–47].

In patients with septic shock, recommendations include initial fluid resuscitation with 30 mL/kg of crystalloid within the first 3 h started within the first hour from the beginning of sepsis symptoms, to obtain a MAP ≥ 65 mmHg, which is the threshold below at which tissue perfusion becomes linearly dependent on arterial pressure. A continuous re-evaluation of the patient is warranted to assess the responsiveness of fluid therapy. If fluid resuscitation is not enough to maintain a permissive MAP in a shocked patient, a vasoactive medication is needed, with norepinephrine as the agent of first choice.

Steroid therapy should not be systematically used in patients responsive to effective fluid or vasopressor therapy. However, a conservative strategy regarding red blood cells transfusion should be used, with a transfusion threshold <7 g/dL if no risk factors are present.

Neutropenic septic patients should not be treated differently from the non-neutropenic population concerning all other intensive care procedures (e.g., sedation, analgesia, glucose control, bicarbonate therapy and nutrition) [25,26].

4.1. Carbapenemases, Carbapenem−Resistant Enterobacteriacae, and Novel Antibiotics

Several mechanisms of antibiotic resistance have been described for Gram-negative bacteria, which are not uncommon in the hematological setting. β-Lactamases can be subdivided according to Ambler’s classification into four main classes, each with its own peculiarities:

1. Class A β-lactamases include penicillinases and cephalosporinases, such as TEM, SHV, CTX-M, PER, VEB, GES, and IBC, with development of resistance to carbapenem, ureidopenicillins, and aztreonam [48,49]. This class also comprises Klebsiella pneumoniae carbapenemases (KPC) enzymes and GES-2, which can hydrolyze carbapenems [49,50].

2. Class B are defined as metallo-β-lactamases (MBL), including IMP, verone integron-encoded metallo-β-lactamase (VIM), New Delhi metallo-β-lactamases (NDM), SPM, and GIM, which initially emerged in Asian countries and later disseminated among Enterobacteriaceae worldwide [50–52].

3. Class C comprises intrinsic and inducible resistance mechanisms, such as AmpC-type cephalosporinase, not inhibited by clavulanic acid, tazobactam, and sulbactam. The structural modifications of AmpC may confer reduced susceptibility to new β-lactam–β-lactamase inhibitors (BLBLIs), ceftolozane/tazobactam, and ceftazidime/avibactam [50–53].
4. Class D oxacillinases provide resistance to all penicillins, 3- and 4-generation cephalosporines (3GCs) and aztreonam. OXA enzymes such as OXA-198 are known mechanisms of carbapenem resistance [50,54]; other OXA b-lactamases (mainly OXA-48) express low activity against carbapenems, hydrolyzing imipenem better than meropenem [50,54].

In addition to the introduction of novel but now well-known cephalosporins such as ceftolozane/tazobactam and ceftazidime/avibactam, therapeutic options for CRE infections have been added to the equipment. The combination of meropenem and a boronic acid b-lactamase, vaborbactam (M/V), has shown increased in vitro activity against Class A and Class C serine b-lactamases; however, Class B (e.g., NDM, VIM) and Class D (e.g., OXA-48) carbapenemases are not inhibited by vaborbactam [55]. The efficacy, tolerability, and safety of M/V was first assessed in TANGO I [55], a multi-center, double-blind, randomized, Phase III, noninferiority trial versus P/T for the treatment of cUTIs and TANGO II [56], in which M/V was compared to the best available treatment (BAT) in a series of infections because of suspected or known CRE (notably cUTI, hospital-acquired or ventilator-associated bacterial pneumonia [HABP/VABP], complicated intrabdominal infections [cIAI], and bacteremia). Because of the superiority of M/V compared to the BAT, this study was terminated prematurely. A Phase III trial called TANGO III, in which M/V was compared to PTZ for the treatment of HABP/VABP, was withdrawn by sponsor decision [57]. Imipenem/cilastatin with relebactam (I/R) combines an approved carbapenem with a novel b-lactamase inhibitor [58]. The in vitro addition of relebactam showed increased activity of imipenem against ESBLs, serine b-lactamases, and \textit{P. aeruginosa}; however, relebactam cannot inhibit Class D OXA-48 [58]. A Phase II study assessed the efficacy, tolerability, and safety of I/R versus imipenem/cilastatin in cIAI and cUTI [58]. In addition, two Phase III studies, RESTORE-IMI 1 [59] and RESTORE-IMI 2 [60], are currently evaluating I/R in different settings. RESTORE-IMI 1 is a multi-center, randomized, double-blind, comparator-controlled trial that compares the efficacy and safety of I/R versus colistin and imipenem/cilastatin (COL þ IMI) in patients with imipenem-nonsusceptible bacterial infections, including HABP, VABP, cIAI, and cUTIs [59]. In the RESTORE-IMI 2 trial, I/R was compared to PTZ (both arms with the addition of linezolid) in HABP/VABP [60].

Cefiderocol is a novel siderophore cephalosporin designed to target CR pathogens, including CR strains of Enterobacteriaceae (CRE), \textit{Pseudomonas aeruginosa}, \textit{Acinetobacter baumannii}, and \textit{Stenotrophomonas maltophilia}, which is intrinsically CR [61,62]. CREDIBLE CR is a randomized, open-label, prospective, Phase III clinical study and was designed to evaluate cefiderocol in patients with CR Gram-negative infections regardless of species or infection site source, even including sepsis and BSIs [61,62]. Hsueh et al. evaluated the in vitro microbiological profiles of cefiderocol, ceftolozane/tazobactam, and ceftazidime/avibactam against \textit{P. aeruginosa}, S. and A. baumannii, and S. maltophilia isolates within the bloodstream. Cefiderocol exhibited significantly more potent in vitro activity when compared to comparators with MICs ≤ 4 mg/L for the five colistin- and imipenem-resistant \textit{P. aeruginosa} isolates and 70% of the ten colistin- and imipenem-resistant isolates [63]. Currently, high-dose and combination strategies that could have the new \beta-lactam/\beta-lactamase inhibitors should be considered in severe CRE infections to maximize treatment success [64]. Cefiderocol is added to these therapeutic options, widening on the A. baumannii and S. maltophilia sides, which have often been excluded from the spectrum of new molecules over the last decade [64,65].

Figure 2 summarizes the first-line antibiotic treatment that might be suggested in hematologic patients with neutropenic fever. In addition to the use of antibiotics, source control represents an aspect of remarkable importance for the successful treatment strategy of neutropenic patients with fever. Roughly all patient candidates in an intensive chemotherapy treatment have a central line catheter (CVC): when patients become febrile, clinicians should carefully evaluate CVC as a possible source of infection. According to these observations Table 2 shows main indications for CVC removal in hematologic patients.
Table 2. Summary of the indications for catheter removal in hematologic patients.

<table>
<thead>
<tr>
<th>Bloodstream infections due to</th>
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<tbody>
<tr>
<td>- <em>S. aureus</em>, VRE, <em>P. aeruginosa</em>, stenotrophomonas maltophilia, Acinetobacter, MDR bacteria</td>
</tr>
<tr>
<td>- Candida</td>
</tr>
<tr>
<td>- Mycobacteria</td>
</tr>
<tr>
<td>- Less virulent pathogens if ≥2 blood cultures revealing the same pathogen</td>
</tr>
<tr>
<td>Persistent infection despite ≥72 h of effective therapy</td>
</tr>
<tr>
<td>Clinical instability despite anti-microbial therapy</td>
</tr>
<tr>
<td>Sepsis and septic shock</td>
</tr>
<tr>
<td>Severe complications (Endocarditis, septic thrombosis, abscess)</td>
</tr>
<tr>
<td>Tunnel infection or port pocket site infection</td>
</tr>
</tbody>
</table>

4.2. Treatment Duration

Antibiotic duration in neutropenic patients with fever of unknown origin (FUO) is a matter of debate. While the Infectious Diseases Society of America (IDSA) and the European Society for Medical Oncology (ESMO) guidelines recommend continuing antibiotic treatment until neutropenia recovery (ANC ≥ 500 cells/mm³), ECIL guidelines recommend discontinued antibiotics if patients are stable for at least 72 h and remain afebrile for 48 h despite the neutrophil count <0.5 × 10⁹/L. This approach has been demonstrated to be safe in several studies [66,67]. AGIHO/DGHO guidelines recommend a semi-conservative approach, discontinuing empiric antibiotic treatment in severe neutropenic patients afebrile for ≥7 days without signs of infection [68].

In cases of documented infection or critically ill neutropenic patients, antibiotic treatment should continue until the infection is microbiologically eradicated or until all clinical signs are resolved for at least 7 to 10 days, depending on the patient’s condition and the infection site.

However, a recent survey from the EBMT work group analyzing current anti-microbial practices in febrile neutropenia across the world showed that ECIL guidelines are not widely implemented in clinical practice [69]. The major deviations from guidelines resulted in the overuse of combination therapy (1/3 of centers) or the use of carbapenems (10% of centers) as first-line therapy in non-complicated/colonized hematological patients, without further de-escalation and no early discontinuation in case of uncomplicated FUO (1/3 of centers) [69]. Perhaps clinicians feel uncomfortable when dealing with resistant pathogens, especially in centers with a high rate of resistance.

4.3. Management of Critically Ill Neutropenic Patients

Since the survival of hematological patients has increased over time, due to new effective therapies and better supportive care, the number of patients admitted to the ICU has increased proportionally. Compared to the past, mortality in critically ill cancer patients with sepsis or septic shock admitted to the ICU has significantly decreased [70]. Several studies confirm that the delayed alert of an ICU team correlates with lower survival, making it more difficult to manage critical situations [71–74].

ICU admission policies are lacking, partly because of a lack of universal scores able to identify deteriorating patients likely to benefit from ICU treatment [75]. Typically, a full code with no restrictions in intensive treatment should be applied to patients in complete remission of the malignancy, in patients receiving first-line therapy and in patients with curative therapeutic options, considering performance status, prognosis and the concept that critical care survival is largely determined by the acute critical illness rather than the underlying hematological malignancy. A full code status with a time-limited trial and frequent reassessment should be applied in patients with a potentially reversible cause of deterioration or uncertain prognosis [76,77]. Neutropenia should no longer be considered a negative prognostic factor in managing these patients, even in allo–HSCT recipients [78–80]. A meta-analysis of more than 2000 HSCT patients in the ICU showed the main prognostic factors of mortality were multi-organ failures requiring support and uncontrolled graft versus host disease (GVHD), not septic
shock, underlying malignancy or disease status, suggesting that they should not be used in the ICU transfer decision process [81]. Therefore, strict collaboration between intensivists and hematologists should be advised.

5. Conclusions and Key Points

The management of hematologic patients who exhibit neutropenic fever represents a challenging issue in the daily clinical practice. The best approach has not yet been defined, but certainly should include a multi-disciplinary strategy. In this respect, the co-operation with ICU, infectious disease specialists, and microbiology seems of paramount importance.

It is extremely important to know the epidemiology of own Center: this allows to set up antibiotic treatment strategies tailored to the pathogens most frequently identified.

Infections due to MDR bacteria represent a life-threatening complication for hematologic patients. To tackle this problem is important to preserve the activity of the old compounds as well as to limit the uncontrolled use of the new antibiotics.

Table 3 describes the most relevant strategies to optimize the antibiotic treatment in hematologic patients with BSI.

<table>
<thead>
<tr>
<th>Main Objective</th>
<th>Intervention</th>
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<tbody>
<tr>
<td>Identify colonized patients with MDR bacteria</td>
<td>Monitor patients with rectal swab for ESBL, KPC, and PA MDR bacteria</td>
</tr>
<tr>
<td>Initial anti-microbial treatment: escalation vs. de-escalation strategy</td>
<td>Knowledge of the local epidemiology</td>
</tr>
<tr>
<td>Source control</td>
<td>Consider CVC removal timely radiologic evaluation (CT scan, MRI etc.)</td>
</tr>
<tr>
<td>Antibiotic stewardship programs</td>
<td>Education, audit, guidelines implementation, antibiotic cycling/restriction</td>
</tr>
</tbody>
</table>

**Table 3.** Strategies to optimize the anti-microbial treatment in hematologic patients with BSI.

**Abbreviations:** ESBL: extended-spectrum beta-lactamase; KPC: Klebsiella pneumoniae carbapenemase producing; MDR PA: multi-drug-resistant Pseudomonas aeruginosa; CT: computed tomography; MRI: magnetic resonance imaging.

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47. Vossen, M.G.; Milacek, C.; Thalhammer, F. Empirical antimicrobial treatment in haematopo-oncological patients with neutropenic sepsis. ESMO Open 2018, 3, e000348. [CrossRef]


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