Strategies for appropriate antibiotic use in intensive care unit

Estratégias para uso adequado de antibioticoterapia em unidade de terapia intensiva

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ABSTRACT

The consumption of antibiotics is high, mainly in intensive care units. Unfortunately, most are inappropriately used leading to increased multi-resistant bacteria. It is well known that initial empirical therapy with broad-spectrum antibiotics reduce mortality rates. However the prolonged and irrational use of antimicrobials may also increase the risk of toxicity, drug interactions and diarrhea due to Clostridium difficile. Some strategies to rational use of antimicrobial agents include avoiding colonization treatment, de-escalation, monitoring serum levels of the agents, appropriate duration of therapy and use of biological markers. This review discusses the effectiveness of these strategies, the importance of microbiology knowledge, considering there are agents resistant to Staphylococcus aureus and Klebsiella pneumoniae, and reducing antibiotic use and bacterial resistance, with no impact on mortality.

Keywords: Antibacterial agents; Drug utilization; Intensive care units

INTRODUCTION

Antimicrobials are one of the major drugs used in intensive care units (ICU), although their indiscriminating and prolonged use is one of the main factors involved in the emergence of multidrug-resistant bacteria, whose incidence has grown in all continents.¹

One example from data collected between 2004 and 2009 on bloodstream infections in the ICU of several countries showed that Staphylococcus aureus isolated were methicillin-resistant in 84.4% of cases; 100% of cases of Pseudomonas aeruginosa were resistant to cefepime and 47.2% to carbapenems; 76.3% of cases of Klebsiella pneumoniae and 66.7% of Escherichia coli were resistant to ceftriaxone; and 55.3% of Acinetobacter baumannii cases were resistant to carbapenems.²

Meanwhile, there has been a reduction in the availability of new antimicrobials in the market. This is also due to the fast emergence of strains resistant to new drugs, which possibly dampens more investments.²

Thus, the best way to reduce the emergence of resistant strains, especially in ICUs, is the rational use of antimicrobial strategies, such as practicing de-escalation; avoiding colonization treatment; evaluating serum levels of antimicrobials and adequate antibiotic treatment duration; and using biological markers such as procalcitonin, for example, that enable differentiating cases of infectious from non-infectious etiology.³,⁴

De-escalation is the adjustment of the antimicrobial regimen according to culture results, that is, changing a regimen with more drugs and/or drugs with a broader spectrum by using another regimen with fewer drugs and/or drugs with a more narrow spectrum, but more sensitive on the antibiogram.⁵,⁶
COLONIZATION VERSUS INFECTION

A common problem observed in the abuse of antimicrobials is treating colonized patients. This happens when antibiotic treatment is introduced due to a positive culture, despite no signs or symptoms of infection. This is the case, for example, of tracheal secretion, urine or tip of central venous catheter cultures.

Knowing the microbiota that colonizes the patient may help in cases in which empirical treatment is necessary, as shown by Blot et al.\(^7\). The acknowledgement of colonization previous to infection was associated with higher rates of adequate treatment in patients that developed bacteremia.\(^7\)

KLEBSIELLA PNEUMONIAE CARBAPENEMASE-PRODUCING BACTERIA

In the multiresistance scenario, carbapenemase-producing Gram-negative (carbapenem inactivating enzymes) stand out, especially *Klebsiella pneumoniae* (KPC). Infections caused by these agents are important in terms of morbidity and mortality and their treatment is challenging.\(^8\)

Automated systems frequently report antibiogram results with agents susceptible to imipenem and meropenem, *in vitro*. Therefore, confirmatory tests are recommended in the presence of strains with decreased sensitivity to carbapenems or resistance to most other beta-lactam drugs.\(^8\)

ANTIMICROBIAL DE-ESCALATION

Empirical broad-spectrum treatment aims to use drugs with wider antimicrobial coverage, boosting therapeutics, and reducing mortality and bacterial resistance.\(^3\) On the other hand, the high consumption of antimicrobials leads to higher risk of toxicity, drug interaction, and in the long term, higher incidence of diarrhea due to *Clostridium difficile*.\(^3\)

Unfortunately, there are no protocols guiding which is the best association of antimicrobials according to each infection site. Therefore, the empirical association of antibiotics occurs according to local epidemiology, patient clinical presentation and risk factors.\(^3\)

For empirical treatment protocols in septic patients, broad-spectrum treatment is recommended to reduce mortality,\(^5\) with the recommendation of collecting cultures before beginning treatment and de-escalation, as soon as possible, after microbiology results.\(^5\)

There are countless studies showing safety and efficacy of de-escalation, although there are no randomized studies.\(^11\)

Shime et al.\(^12\) assessed the efficacy and safety of de-escalation in a study on Gram-negative bacteremia. Although de-escalation was indicated to all patients, it was conducted only in 57% of cases. There were no cases of treatment failure or reduction in costs.

Joung et al.\(^13\) assessed the impact of de-escalation in mortality of patients with pneumonia, in ICU. The practice aimed to evaluate the decrease in number and spectrum of antimicrobials. There was a trend toward lower pneumonia-related mortality in 14 days, but without statistical significance. However, 30-day mortality was significantly lower in patients submitted to de-escalation.

Morel et al.\(^14\) also reported that there was no increase in mortality with de-escalation in general infections at the ICU.

ANTIMICROBIAL SERUM LEVELS

Using antimicrobials without the availability of serum dosages may be complicated in several scenarios, because of obesity, renal failure, hemodynamic instability and severe infections. If one cannot assess if a drug is used at a therapeutic level, there may be treatment failure, toxicity and adverse events.\(^15\)

Several studies on sepsis and septic shock have already made it clear that the early administration of antimicrobials reduces mortality rates, although there is little information available on adequate dosage regimens and clinical outcomes.\(^5,6\)

Van Lent-Evers et al. published a randomized controlled study that monitored serum aminoglycosides, with an impact on length of stay.\(^16\)

VANCOMYCIN SERUM LEVELS

Throughout the years, vancomycin has been one of the most studied antimicrobials with several pharmacokinetic analyses, and in a variety of populations. Serum concentration aims to minimize adverse events and to reach adequate concentrations, decreasing treatment failure rates.\(^17\)

Vancomycin is a glycopeptide antimicrobial available for clinical use for over 50 years, when there were not many options to treat penicillin-resistant *Staphylococcus* infections.\(^20\)

Initially, vancomycin was associated with countless adverse events, including toxicities related to infusion, nephrotoxicity and ototoxicity.\(^20\) The events were possibly related to initial formulations, considered more impure.\(^20\) The use of vancomycin was significantly reduced with the emergence of semisynthetic penicillins
(methicillin and oxacillin, for example), considered less toxic.\(^{(20)}\)

However, as of the 1980’s, there was an expressive growth in the use of vancomycin.\(^{(21)}\) This happened due to the progressive increase in infections due to resistant agents, mainly methicillin-resistant \textit{Staphylococcus aureus} (MRSA) infections.\(^{(21)}\)

In a systematic review of 15 studies that evaluated nephrotoxicity while using vancomycin, the incidence ranged between 5 and 43\%. Higher serum levels of the drug (\(\geq 15\text{mg/dL}\)) were associated with increased renal toxicity, occurring more frequently in patients with other risk factors, such as admission to an intensive care unit.\(^{(22)}\)

Forstner et al.\(^{(23)}\) performed a retrospective analysis of patients with methicillin-resistant \textit{Staphylococcus} bacteremia, and found the level of vancomycin as a predictor factor of mortality; in that, only 22.6\% of patients reached recommended serum levels of the drug.

In 2013, a Chinese group published a revision and meta-analysis of studies that assessed the benefit of monitoring serum vancomycin, with significantly higher rates of clinical efficacy and less nephrotoxicity in patients submitted to the practice.\(^{(24)}\)

In 2006, the Clinical and Laboratory Standards Institute (CLSI) decreased the minimum inhibitory concentration (MIC) \textit{breakpoint} of vancomycin susceptibility from 4\text{mg/L} to \(<2\text{mg/L}\) for MRSA. Despite this fact, there is frequent concern, given there is a historical decrease in sensitivity of MRSA to glycopeptides, a phenomenon called “\textit{MIC creep}”\(^{(25,26)}\). This means that there is a change in the MIC pattern of vancomycin among populations of \textit{Staphylococcus aureus} considered susceptible, generating a subpopulation with reduced sensitivity to the drug. It is essential that individuals with suspected infection by these strains be evaluated judiciously, with advanced microbiology knowledge, for adequate antimicrobial treatment.

Several studies reported worse outcomes in patients who developed MRSA infections with higher MIC.\(^{(27-29)}\) A systematic review and meta-analysis published in 2012 observed that the higher the level of vancomycin, the higher the mortality rate, regardless of the infection source.\(^{(27,28)}\)

Sakoulas et al.\(^{(29)}\) evaluated the relation between MIC and the bactericidal efficacy of vancomycin to treat MRSA bacteremia, and found a positive correlation between \textit{in vitro} bactericidal activity and clinical success.

Strains of glycopeptide intermediate-resistant \textit{S. aureus} (GISA) have been described in the past years. Infections by these agents are still not very frequent, but the concern is that the prevalence can grow due to high use and to exposure to vancomycin.\(^{(21)}\)

The most recent recommendations for vancomycin use in adults by the Infectious Diseases Society of America (IDSA) were published in 2009 and are displayed in chart 1. The serum concentration is described as the most accurate and practical way to monitor drug efficacy. A higher level is recommended, with aggressive monitoring, depending on the kind of infection, such as 15 to 20\text{mg/dL} for bacteremia, endocarditis, osteomyelitis, meningitis and nosocomial \textit{Staphylococcus aureus} pneumonia.\(^{(18,19)}\)

### Chart 1. Recommendations by the Infectious Diseases Society of America (2009) to adjust therapeutic dose of vancomycin

| Dosage      | Initial dose should be calculated according to weight, as 25-30\text{mg/kg} |
|-------------|--------------------------------------------------------------------------------|
| Serum level | More effective and practical method                                             |
|             | Should be obtained immediately before the fourth dose, for patients with      |
|             | normal kidney function                                                        |
|             | Recommendation: 15-20\text{mg/dL} every 8 to 12 hours                       |

### LENGTH OF TREATMENT PERIOD AND USE OF PROCALCITONIN

There has been ample discussion in the literature on length of treatment of both community and healthcare-related infections.

Short-term treatments have several advantages, but may not eradicate the microorganism, increasing the likelihood of relapses, mainly for non-fermenting Gram-negative bacteria.\(^{(30-33)}\) Longer treatments may be related to higher toxicity, adverse events, higher risk of candidemia diarrhea due to \textit{Clostridium difficile}.\(^{(30)}\)

Many factors influence the medical decision regarding length of antimicrobial treatment, such as characteristics of the microorganism, of the patient, infection and of the drugs available for treatment (chart 2).\(^{(30)}\)

### Chart 2. Factors that influence length of antimicrobial treatment

| Microorganism characteristics | Sensitivity profile |
|------------------------------|--------------------|
|                              | Biofilm formation capacity |
|                              | Metastatic focus potential |
|                              | Virulence |
| Patient characteristics      | Immunological status (age, comorbidity and immunosuppressing treatments) |
|                              | Presence of foreign body (metal prosthesis, valve implant, catheter, etc.) |
| Infection characteristics    | Duration |
|                              | Location |
|                              | Severity and response to treatment |
| Antimicrobial characteristics | Profile of microorganism sensitivity |
|                              | Bactericidal versus bacteriostatic |
|                              | Monotherapy versus combined treatment |
Chastre et al.,(31) compared 8 versus 15 days of treatment for mechanical ventilation-associated pneumonia (MVAP) in a prospective and randomized study. They demonstrated that patients submitted to shorter treatment courses did not differ in mortality or infection recurrence, except for pneumonias caused by non-fermenting Gram-negative bacilli, including Pseudomonas.

Corona et al.(32) also evaluated patients with community and hospital bacteremia; 72.5% received antimicrobial treatment for a short period, with a median of 5 days, many of which with septic shock, and without harming patient outcome. Pugh et al.(33) evaluated short-term versus long-term antimicrobial treatment efficacy in patients with hospital pneumonia, including MVAP in a systematic review. A 7-to-8 day antibiotic course reduced recurrence of MVAP by multiresistant agents, without harming treatment, except in cases by non-fermenting Gram-negative bacilli.

A new approach to try to determine the real need for antimicrobial treatment and even help in length of treatment, is to use biomarkers such as procalcitonin.(34-39)

Procalcitonin is a calcitonin peptide precursor produced as a response to bacterial toxins and, mainly, to pro-inflammatory mediators such as interleukin 1b, tumor necrosis factor and interleukin 6. The levels of procalcitonin rise within 6 to 12 hours from the beginning of bacterial infection, and are expected to drop after control of the infectious presentation.

The authors of the prospective study PRORATA found a 23% reduction in the consumption of antimicrobials for patients submitted to a procalcitonin collection protocol, without statistical difference in mortality.(34)

A systematic review by Schuetz et al.(35) described safety and efficacy of using procalcitonin for respiratory infections and sepsis. The authors observed an important reduction in exposure to antimicrobials in several scenarios, with no increase in mortality rate: in clinical presentations with little evidence of infection, such as exacerbation of chronic obstructive pulmonary disease and bronchitis; and in shorter treatments in patients with pneumonia and sepsis.

There are several other studies that found lower rates of antimicrobial use in patients submitted to procalcitonin collecting protocols, although there is also some criticism: most investigated respiratory infections; many are inconclusive; standardized criteria are lacking for collecting procalcitonin; and there is no defined cut-off value. In severe patients, the levels of procalcitonin were not used for initiating treatment, but for discontinuing the antimicrobial. However, the drop in biomarker levels correlates with clinical resolution of the infectious process, which allows the safe withdrawal of the antimicrobial agent.(36-39)

A recent study by Huang et al.(38) prospectively evaluated the use of procalcitonin for withdrawal of antimicrobial regimens in patients with intra-abdominal infection, with an 87% reduction in duration of treatment, with no increase in risk of adverse events.

A meta-analysis by Prkno et al.(39) evaluated studies involving patients with severe sepsis treated in the ICU. The authors described a significant reduction in duration of antimicrobial treatment in patients whose therapy was guided by procalcitonin, with no impact on mortality.

Despite several publications showing a possible benefit in using procalcitonin to decrease duration of antimicrobial treatment, there are few cases, many studies do not differentiate clinical from surgical patients, and there are no standardized criteria.(34-39)

**CONCLUSION**

Bacterial resistance is a growing and global concern, as are high mortality rates in septic patients treated inadequately.

Several strategies for adequate use of antibiotic treatment have been proposed and discussed exhaustively in the literature. There are several benefits, such as not treating colonization, antimicrobial de-escalation, serum dosage of drugs, use of biomarkers, and shorter treatment with no difference in mortality.

Some issues, such as procalcitonin cut-off values, still need to be explained but the meticulous use of this and other strategies can optimize success of treatment and minimize risks due to prolonged and inadequate use of antimicrobials.

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