Optimal infusion rate in antimicrobial therapy: explosion of evidence in the last five years

Ling-ling Zhu,1 Quan Zhou2
1VIP care ward, Division of Nursing, The Second Affiliated Hospital, School of Medicine, Zhejiang University, Hangzhou, People’s Republic of China; 2Department of Pharmacy, the Second Affiliated Hospital, School of Medicine, Zhejiang University, Hangzhou, People’s Republic of China

Background: Sporadic studies in antimicrobial therapy have evaluated the effects of infusion rates on therapeutic and economic outcomes, and new findings may challenge the regular infusion regimen.

Methods: Focusing on studies comparing the outcomes of different infusion regimens, the relevant literature was identified by searching PubMed, Web of Science, and Scopus from January 1, 2013 to March 1, 2018. Papers were finally chosen using a PRISMA flowchart.

Results: Antimicrobials with the superiority of prolonged infusion to standard infusion in terms of efficacy and safety include meropenem, doripenem, imipenem, cefepime, ceftazidime, piperacillin/tazobactam, linezolid, and vancomycin. The strategy of concomitantly reducing total daily dose and prolonging infusion time may cause treatment failure (eg, imipenem). Extended infusion of piperacillin/tazobactam has pharmacoeconomic advantage over standard infusion. Prolonged infusion of voriconazole is inferior to standard infusion because of lower efficacy caused by pharmacokinetic changes. Comparable outcomes following standard infusion and continuous infusion were observed with norvancomycin and nafcillin. Factors determining whether prolonged infusion has a benefit over standard infusion include MIC of bacterial pathogens, bacterial density, diagnosis, disease severity, total daily dose, and renal function.

Conclusion: To maximally preserve the effectiveness of current antimicrobials, effective interventions should be implemented to enhance the application of optimal infusion strategies. For reducing nephrotoxicity, prolonged infusion of meropenem is better than conventional infusion in neonates with Gram-negative late-onset sepsis, and continuous infusion of vancomycin is superior to intermittent infusion. For increasing efficacy, prolonged or continuous infusion of time-dependent antimicrobials (eg, meropenem, doripenem, imipenem, cefepime, ceftazidime, piperacillin/tazobactam, linezolid, and vancomycin) is an optimal choice. Nevertheless, such advantages may only be demonstrated in special clinical circumstances and special populations (eg, patients with a sequential organ failure assessment (SOFA) score ≥9, respiratory tract infections, urinary or intra-abdominal infections, or infections caused by less susceptible pathogens would benefit from prolonged infusion of piperacillin/tazobactam).

Keywords: antimicrobial resistance, antimicrobial stewardship, efficacy, infection, infusion regimen, nursing time, safety, pharmacoeconomics

Introduction

Infusion rate is one of essential elements that should be included in all intravenous orders or prescriptions according to the Joint Commission International (JCI). Accreditation Standards for Hospitals, sixth edition.1 Numerous sporadic clinical studies have been conducted to evaluate the effects of infusion rates on therapeutic and economic
outcomes of drug therapy, and the new findings have challenged the regular infusion regimen. Health care-associated infections, alongside increasing antimicrobial resistance, pose significant burdens to the hospital system. More aggressive dosing strategies, such as prolonged and continuous infusion of β-lactam antibiotics, are applied to address the tremendous challenges associated with the difficult-to-treat pathogens and increase the probability of attaining pharmacokinetic/pharmacodynamic (PK/PD) targets predictive of successful clinical outcomes. Clinicians who have a thorough understanding of appropriate antibiotic use and dosing regimens are essential in the battle to preserve the usefulness of antibiotics and prevent further antibiotic resistance. In this article, based on the literature in the latest 5 years, we share our perspectives in rational intravenous rates of antimicrobial agents and hope to bring some implications for clinical practice and research.

Methods

Search strategy
The relevant literature was identified by searching PubMed, Web of Science, and Scopus from January 1, 2013 to March 1, 2018. For PubMed, the search method was identifying papers with titles containing “infusion rate,” “infusion rates,” “short infusion,” “fast infusion,” “rapid infusion,” “slow infusion,” “extended infusion,” “prolonged infusion,” “continuous infusion,” “continuous administration,” “infusion time,” “infusion mode,” “dosing regimen,” “dosing regimens,” “continuous vs intermittent,” “extended vs intermittent,” “standard vs prolonged,” “extended vs standard,” or “intermittent vs prolonged,” with a filter of “text availability: abstract” and “languages: English.” For Scopus and Web of Science, the query was article title containing the same words as in the PubMed search, with a filter of “language: English; publication type: article,” followed by “search within results: antibiotics or antimicrobials.”

Selection criteria
Inclusion criteria included studies comparing the outcomes of different infusion regimens focusing on infusion rates. Exclusion criteria included documents with actually irrelevant topics on antimicrobial therapy, despite being retrieved using the search terms, studies without comparison of outcomes at different intravenous infusion rates of antimicrobial agents, studies lacking a control group with different infusion rates, and documents such as comments, letters, reviews, case reports, editorials, and descriptive studies without statistical analysis. Fifty-three papers were finally chosen according to the inclusion/exclusion criteria. The PRISMA flowchart is illustrated in Figure 1. The full text of each include article was critically reviewed, and valuable information was summarized by data interpretation.

Results and discussion

β-lactam antibiotics
Two meta-analyses confirmed the advantages of continuous or prolonged infusion of β-lactam antibiotics. Roberts et al performed a meta-analysis of 632 critically ill patients with severe sepsis from three randomized trials, and concluded that administration of β-lactam antibiotics by continuous infusion was associated with decreased hospital mortality (19.6% vs 26.3%; relative risk=0.74; P=0.045) and improved clinical cure (55.4% vs 46.3%; relative risk=1.20; P=0.021) compared with intermittent dosing. Teo et al conducted a meta-analysis of 29 studies with 2,206 patients and observed that use of prolonged infusion of β-lactam antibiotics appeared to be associated with a significant reduction in mortality (relative risk=0.66) and improvement in clinical success (relative risk=1.12) in hospitalized adult patients compared with intermittent boluses.

Two early studies did not show advantages of continuous or prolonged infusion of β-lactam antibiotics. A before–after study showed that routine use of prolonged infusion (over 3 hours) for the empiric treatment of Gram-negative bacterial infections in the intensive care unit (ICU) with β-lactams could not offer advantage over intermittent infusion (over 30 minutes) in terms of treatment success, mortality, or hospital length of stay (LOS). A multicenter randomized trial showed no difference in outcomes between continuous infusion and 30-minutes intermittent infusion in critically ill patients with severe sepsis receiving β-lactam antibiotics (eg, ICU-free day, 90-day survival, clinical cure, organ failure-free days, and duration of bacteremia).

A prospective, two-center randomized controlled trial compared clinical and PK/PD outcomes of β-lactams following continuous infusion or intermittent bolus dosing in adult critically ill patients with severe sepsis not receiving renal replacement therapy. Although two infusion methods did not exhibit a difference in 14 or 30-day survival, continuous infusion resulted in better outcomes in terms of clinical cure rates (56% vs 34%, P=0.011), median ventilator-free days (22 vs 14 days, P<0.043), PK/PD target attainment rates at 100% free drug levels above the minimal inhibitory concentration (fT>MIC) on day 1 (97% vs 70%, P<0.001) and day 3 (97% vs 68%, P<0.001) post-randomization compared with intermittent infusion.
There were nine studies of extended or continuous infusion of meropenem vs intermittent administration (Table 1). Among these, two studies were performed in neonates and children, respectively. Six studies were conducted in adults. All studies showed the benefits of prolonged or continuous infusion mode, although they were related with diagnosis, disease severity, and MIC of bacterial pathogens. Only one study revealed the advantage of prolonged infusion mode regarding drug toxicities.

**Doripenem**

A retrospective study confirmed that adult critically ill patients receiving standard 1-hour infusion of doripenem for treatment of Gram-negative bacterial infections were significantly more likely to experience clinical failure (odds ratio=5.5) and recurrence of infection or death within 90 days compared with the 4-hour infusion regimen, despite no significant differences in clinical success, LOS, and duration of treatment.

**Imipenem**

Ibrahim et al compared the efficacy of imipenem in adult ICU patients with ventilator-associated pneumonia (VAP) following regular infusion over 30–60 minutes (first year of the study) or 3-hour extended infusion (second year of the study). Extended infusion resulted in a significant decrease in mortality, number of recurrent infections, ICU LOS, and the number of mechanical ventilator days.

Suchánková et al evaluated the population pharmacokinetics of imipenem in adult critically ill patients with hospital-acquired pneumonia (HAP). The 3-hour infusions
Table 1 Extended or continuous infusion of meropenem vs intermittent administration

<table>
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<th>Study design</th>
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<td>Prospective randomized clinical trial</td>
<td>Prolonged infusion of meropenem over 4 h vs conventional infusion (30 min); 20 mg/kg/dose every 8 h and 40 mg/kg/dose every 8 h</td>
<td>Meningitis and Pseudomonas infection in neonates with Gram-negative late-onset sepsis</td>
<td>Higher clinical improvement (61% vs 33%, P=0.009) and microbiologic eradication at the 7th day of meropenem therapy (62% vs 56.8%, P=0.009), lower neonatal mortality (14% vs 31%, P=0.03), shorter duration of respiratory support (12.5 days vs 4 days, P=0.03), and less acute kidney injury (6% vs 23.5%, P=0.02).10</td>
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<td>Retrospective observational study</td>
<td>4 h extended infusion of meropenem (1 g/8 h) vs conventional 30 min short infusion at the same dose</td>
<td>Adult patients with febrile neutropenia</td>
<td>Better clinical outcome (ie, fewer additional antibiotics during the first 5 days of treatment, a more prompt defervescence and a faster decrease in C-reactive protein level) (P&lt;0.05) despite comparable LOS in the hospital and mortality rate between two groups.11</td>
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<td>Randomized clinical trial</td>
<td>Continuous group: a loading dose of 0.5 g of meropenem infused over 30 min followed by continuous infusion of 3 g/day, which was divided into six consecutive 4 h continuous infusions of meropenem of 0.5 g; intermittent group: an initial dose of 1.5 g followed by 1 g infused over 30 min every 8 h.</td>
<td>Adult patients with severe sepsis and septic shock</td>
<td>Continuous infusion of meropenem provided significantly shorter treatment duration (7.6 vs 9.4 days; P=0.035) while clinical success was similar between both groups. For medium-susceptibility pathogens, concentrations above the MIC in the continuous infusion group were 100%, which was better than that in the intermittent group.12</td>
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<td>Observational study using a population PK/PD model</td>
<td>3 h and 30 min infusion regimens at a dose of meropenem 1 g (twice or thrice daily)</td>
<td>Adult critically ill patients</td>
<td>The PTA, T&gt;MIC of 100%, was higher for 3 h infusion regimen compared with 30 min infusion regimen for all ranges of creatinine clearance.13</td>
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<td>Comparative study using an in vitro PK/PD model</td>
<td>Short (0.5 h) and prolonged (3 h) infusion regimens of 1 g meropenem every 8 h</td>
<td>Different adult patient groups with CPKP isolates</td>
<td>ICU patients exhibited the lowest target attainment rates, whereas internal medicine patients achieved the highest target attainment rates.14 The PTA (ie, 40% T&gt;MIC) with short infusion was higher than those with prolonged infusion for isolates with MIC 4–8 mg/L rather than MIC ≥16 mg/L or ≤2 mg/L (MIC 4 mg/L: 98%–99% vs 61%–83%; MIC 8 mg/L: 55%–79% vs 23%–33%).14</td>
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<td>Observational pharmacokinetic Monte Carlo simulations</td>
<td>Meropenem as sole agent or in combination with other antimicrobials</td>
<td>Acinetobacter baumannii and Pseudomonas aeruginosa infections in adult patients with septic shock</td>
<td>Intermittent dosing of 1 g/8 h (30 min) seemed sufficient in patients with normal renal function, whereas increasing doses (2 g/6 h for A. baumannii, 2 g/8 h or 1 g/6 h for P. aeruginosa) by intermittent (30 min) or prolonged (3 h) infusion or continuous infusion (6 g over 24 h) could increase the possibility of achieving therapeutic drug concentrations in patients with septic shock and possible augmented renal clearance.15</td>
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<td>Prospective, multicenter pharmacokinetic point-prevalence study with a post hoc analysis</td>
<td>Intermittent-bolus administration vs prolonged infusion</td>
<td>Critically ill patients</td>
<td>Compared with intermittent-bolus administration, prolonged infusion demonstrated significantly better 30-day survival in the subgroup of patients with respiratory infection (86.2% vs 56.7%; P=0.012), and higher clinical cure rate (73.3% vs 35.0%; P=0.035) and survival rates (73.3% vs 25.0%; P=0.025) in patients with a SOFA score of ≥9. However, two infusion methods did not exhibit a significant difference in 30-day survival rate in patients receiving antimicrobial treatment and in the subgroup of patients with abdominal infection.16</td>
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and continuous infusion achieved over 97% of cumulative fraction of response (CFR) against all pathogens, with the exception of *Pseudomonas aeruginosa* and *Acinetobacter* spp., where the maximum CFRs were 85.5% and 88.4%, respectively. The 3-hour extended infusions and continuous infusion achieved higher probability of target attainment (PTA) for the targets of achieving \(fT_{\geq MIC}\) for 40% and 100% of the dosing interval as PK/PD targets compared with 0.5-hour infusions; however, continuous infusion carried a risk of not reaching the MIC of less susceptible pathogens in a higher proportion of patients. Therefore, it may be necessary to use maximum doses administered as extended infusions in critically ill patients with HAP and risk factors for Gram-negative non-fermenting bacteria.

The benefit of prolonged infusion vs standard infusion of imipenem may be related with drug dose and MIC of bacterial pathogens. Lipš et al.\(^2\) compared the PK/PD indices of imipenem administered by standard infusion (1 g imipenem/1 g cilastatin over 30 minutes every 8 hours) and extended infusion with a reduced total dose (0.5 g imipenem/0.5 g cilastatin over 3 hours every 6 hours) in critically ill patients with nosocomial pneumonia. Concentrations of imipenem were above the target concentration (4×MIC of 2 mg/L) for ≥40% of the dosing interval in every patient in the bolus group, whereas in the extended group this PK/PD index (40% \(fT_{\geq4\times MIC}\)) was achieved in only 20% of patients. Moreover, 70% of the patients in the extended group did not reach the desired drug concentrations at all. For a MIC of 4 mg/L, only one patient in the bolus group achieved 40% \(fT_{\geq4\times MIC}\), and the rest of the patients still at least reached concentrations above 4×MIC, whereas none of the patients in the extended group reached these concentrations. Therefore, administration of 0.5 g of imipenem by a 3-hour infusion every 6 hours could not provide adequate drug concentrations to control infections caused by pathogens with MIC≥2 mg/L. In empirical treatment where the presence of a less sensitive micro-organism cannot be excluded, the strategy of concomitantly reducing total daily dose and prolonging infusion time may cause treatment failure.

### Biapenem

The usual biapenem infusion regimen (0.3 g every 12 hours, 0.5-hour infusion) may result in unsatisfactory clinical outcomes in Chinese adult patients with lower respiratory tract infections, because PTA could not achieve ≥90% with an MIC of 4 µg/mL. With prolonging infusion time of biapenem from 0.5 hour to 3 hours with a dose of 0.3 g every 6 hours, the PTAs for achieving 40% \(T_{>\text{MIC}}\) with an MIC of 4 µg/mL, 60% \(T_{>\text{MIC}}\) with an MIC of 2 µg/mL, and 80% \(T_{>\text{MIC}}\) with an MIC of 1 µg/mL were increased from 53.8% to 94.3%, from 60.3% to 98.4%, and from 62.8% to 96.2%, respectively. Higher doses and longer infusion time of biapenem would be appropriate for empirical therapy, and combination therapy with other antibacterials may be more appropriate if *P. aeruginosa* or *Acinetobacter baumannii* infection is suspected.\(^2\)
Ertapenem
Prescribing information for ertapenem specifies that the drug should be infused over a period of 30 minutes when administered intravenously. Wiskirchen et al\(^2^{24}\) showed that ertapenem 1 g administered as a rapid 5-minute infusion could provide a well-tolerated, bioequivalent, and pharmacodynamically equivalent regimen to the 30-minute infusion at clinically relevant MICs in healthy adult volunteers. The estimated PTA was greater than 90% when using a bactericidal target of 40% FT>MIC up to an MIC of 0.25 mg/L for both infusion times. Additional studies in infected patients are needed to determine the safety and value of rapid 5-minute infusion of ertapenem.

Cefepime
A retrospective quasi-experimental study showed that the extended infusion (a 4-hour infusion of 2 g cefepime every 8 hours) rather than standard infusion (a 30-minute infusion of 2 g cefepime every 8 hours) resulted in significantly lower overall mortality, reduced hospital costs, and significantly shorter LOS in the treatment of \(P\) aeruginosa\) infections in adult ICU patients.\(^2^{25}\) However, a newer study did not confirm the superiority of extended infusion of cefepime.\(^2^{26}\) Wrenn et al\(^2^{26}\) compared the outcomes between extended infusion and standard infusion of cefepime as empiric treatment of febrile neutropenia in adult patients admitted to a hematology/oncology service and observed no statistically significant differences in defervescence by 72 hours and secondary outcome measures (time to defervescence, clinical success, in-hospital mortality, hospital LOS, and need for additional antimicrobials).

Regarding continuous infusion of cefepime, this infusion mode could significantly enhance the antibacterial effect toward intracranial infections. Compared to intermittent infusion of 2 g cefepime every 12 hours, continuous infusion of 4 g/day cefepime over 24 hours could significantly reduce the treatment duration in adult neurosurgical patients with postoperative intracranial infections (6.6±1.9 days vs 7.8±2.6 days; \(P=0.036\)), and achieve higher %T>MIC for cefepime concentrations in cerebrospinal fluid (for MICs of 4 µg/mL, 83.3% vs 25%; for MICs of 8 µg/mL, 75% vs 0%).\(^2^{27}\)

Ceftazidime
Cousson et al\(^2^{28}\) confirmed the advantages of continuous infusion (loading dose of 20 mg/kg followed by 60 mg/kg/day) over intermittent infusion (20 mg/kg over 30 minutes every 8 hours) in terms of pharmacodynamics and predictable efficacy in adult patients with VAP due to Gram-negative bacilli. The %T>MIC of ceftazidime (MIC=20 mg/L) was 100% and 46%, respectively, following continuous infusion and intermittent infusion (\(P<0.003\)).

Piperacillin/tazobactam
Efficacy issue
Compared to the conventional intermittent dosage regimen, prolonged piperacillin/tazobactam infusion resulted in shorter ICU and hospital LOS among adult critically ill patients (ie, saving 0.6 days and 5.6 days, respectively),\(^2^{29}\) a significantly lower rate of readmission within 30 days of hospital discharge (1.2% vs 13.7%, \(P=0.002\)),\(^3^{30}\) and greater PTA in critically ill patients receiving continuous renal replacement therapy or critically ill patients with severe sepsis or septic shock undergoing continuous venovenous hemofiltration.\(^3^{31,32}\)

Safety issues
There were two studies specially comparing safety issues of extended infusion vs intermittent infusion of piperacillin/tazobactam. Both intermittent and extended-infusion groups experienced similar incidence of acute renal injury in hospitalized patients,\(^3^{33}\) and comparable nephrotoxicity in patients with a creatinine clearance (CrCl) ≥40 mL/min receiving at least 96-hour combination therapy of vancomycin and piperacillin/tazobactam.\(^3^{34}\)

Pharmacoeconomic issues
There were four studies involving the pharmacoeconomic issues of extended infusion vs intermittent infusion of piperacillin/tazobactam. Brunetti et al\(^3^{35}\) showed that automatic substitution of extended infusion for standard infusion of piperacillin/tazobactam in adult patients could reduce total cost per treatment course by 13%, with no difference in LOS and hospital mortality between two groups. Compared with regular infusion in the pre-intervention phase, post-implementation of an extended infusion dosing protocol resulted in a significant shorter ICU and hospital LOS and a significantly reduced cost of therapy ($120.21 vs $155.17; \(P=0.035\)), although there was no significant difference in the primary end-point of mortality and adverse drug effects between the two phases.\(^3^{36}\) Bao et al\(^3^{37}\) showed better PK/PD and pharmacoeconomic parameters following 3-hour extended infusion of piperacillin/tazobactam vs 30-minute intermittent infusion every 6 hours in the treatment of HAP in adult critically ill patients with low illness severity. Extended infusion achieved a lower therapy cost ($1,351.72 vs $1,782.04, \(P=0.001\)), and higher %T>MIC (−100, 98.73, and 93.04% in the extended infusion group vs 81.48, 53.29, and 42.15% in the intermittent infusion group, respectively,
for the pathogen with an MIC of 4, 8, and 16 mg/L), although the clinical success rate, clinical failure rate, and drug-related adverse events did not significantly differ between groups. Cotrina-Luque et al\(^\text{38}\) confirmed similar efficacy and safety of continuous infusion of piperacillin/tazobactam to intermittent infusion in treating \textit{P. aeruginosa} infection; however, continuous infusion (an initial loading dose of piperacillin/tazobactam 2/0.25 g in 100 mL of saline solution for 30 minutes, immediately followed by continuous infusion of piperacillin/tazobactam 8/1 g in 500 mL of saline over 24 hours) could lead to system cost savings due to utilizing a 30% lower dose of piperacillin/tazobactam than intermittent infusion (an initial dose of piperacillin/tazobactam 4/0.5 g in 100 mL of saline solution for 30 minutes, followed by piperacillin/tazobactam 4/0.5 g in 100 mL of saline for 30 minutes every 8 hours).

**Whether prolonged infusion has a benefit over standard infusion depends on the MICs of bacterial pathogens**

Zelenitsky et al\(^\text{39}\) characterized the pharmacodynamics of prolonged-infusion piperacillin/tazobactam in an in vitro pharmacodynamic model of \textit{P. aeruginosa}. The benefits of prolonged infusion were selective and most likely observed in patients with less susceptible pathogens, ie, prolonged infusion had no advantages over standard infusion against isolates with susceptible MICs of 8 or 16 mg/L, whereas it produced more than twice the final bacterial kill against less susceptible isolates with an intermediate MIC of 32 mg/L.

Nichols et al\(^\text{40}\) evaluated the population PK/PD profiles for dosing regimens of 80–100 mg/kg of the piperacillin component given every 6–8 hours and infused over 0.5, 3, or 4 hours in critically ill children. At the pharmacodynamic target of a T>MIC of \(\geq 50\%\), all simulated dosing regimens achieved a PTA of \(>90\%\) at MICs of \(\leq 8\) mg/L. Only 0.5-hour infusion regimens of 80–100 mg/kg every 8 hours did not achieve a PTA of \(>90\%\) at a MIC of 16 mg/L. All extended-infusion regimens achieved PTAs of \(>90\%\) at MICs of \(\leq 16\) mg/L; however, only the 3-hour infusion regimens given every 6 hours achieved PTAs of \(>90\%\) at a MIC of 32 mg/L. At the pharmacodynamic target of a T>MIC of 100%, none of the regimens achieved a PTA of \(>90\%\) at a MIC of \(\geq 16\) mg/L, and only 100 mg/kg given every 6 hours and infused over 3 hours achieved a PTA of \(>90\%\) at a MIC of 8 mg/L. For pathogens with MICs of \(\leq 8\) mg/L, extended infusion did not substantially improve PTA, and standard infusion was likely sufficient. However, piperacillin/tazobactam 100/12.5 mg/kg given as an extended infusion every 6–8 hour may be optimal for empirical or directed therapy in critically ill pediatric patients with infections caused by less susceptible pathogens or if the desirable T>MIC is greater than 50%.

**Whether prolonged infusion has a benefit over standard infusion depends on the diagnosis and disease severity**

Lyu et al\(^\text{41}\) investigated the infusion modes of piperacillin/tazobactam guided by SOFA score in cancer patients with postoperative HAP. Compared to conventional infusion (30 minutes), prolonged infusion improved patient survival, increased clinical efficacy and bacteriologic efficacy, and reduced 28-day mortality \((P<0.05)\). However, patients with severe disease (SOFA score \(\geq 9\)), rather than those with mild disease (SOFA score <9), benefited from prolonged infusion (eg, shorter days of antibiotics use and ventilator time, longer survival, better clinical efficacy and lower 28-day mortality rate).

Extended infusion of piperacillin/tazobactam over 4 hours for presumed sepsis syndromes exhibited a shorter duration of piperacillin/tazobactam therapy than regular infusion over 30 minutes \((P<0.001)\), with no significant difference in inpatient mortality rates, overall LOS and clinical failure rates between two infusion methods. Subgroup analysis revealed that ICU patients exhibited no significant differences in outcomes following two dosing methods, whereas patients with urinary or intra-abdominal infections had lower mortality and clinical failure rates following extended-infusion treatment.\(^\text{42}\)

Critically ill patients who had a diagnosis of either bacterial infection or neutropenic fever receiving a 4-hour extended-infusion piperacillin/tazobactam demonstrated similar 14-day mortality compared with patients receiving 30-minute intermittent infusions. However, post hoc subgroup analysis showed a lower mortality rate in the extended-infusion group who had infectious organisms identified \((9.3\% \text{ vs } 22.4\%, P=0.01)\) or were diagnosed with respiratory tract infections \((8.9\% \text{ vs } 18.7\%, P=0.02)\).\(^\text{43}\)

**Whether prolonged infusion has a benefit over standard infusion depends on bacterial density**

Bacterial burden may exert a significant influence on the ultimate outcome of antimicrobial therapy. Felton et al\(^\text{44}\) used a hollow-fiber infection model with \textit{P. aeruginosa} to examine the impact of administering piperacillin/tazobactam by 0.5 hour bolus dosing or 4 hours extended infusion every 8 hours on the emergence of piperacillin resistance. Both infusion methods resulted in comparable antibacterial activities and rates of emergence of antimicrobial resistance. For the lower initial bacterial density, trough total plasma piperacillin...
concentration/MIC ratios of 3.4 and 10.4 following bolus and extended-infusion regimens, respectively, were able to suppress the emergence of piperacillin resistance. For the higher initial bacterial density, both infusion regimens were associated with progressive growth of a resistant subpopulation, and thus combination therapy may be required to maximize bacterial killing and prevent antimicrobial resistance.

Nafcillin
Prescribing information for nafcillin specifies that the drug should be administered slowly as an intravenous infusion over at least 30–60 minutes at the adult dose of 500 mg to 1 g every 4 hours. Bauer et al4 compared the clinical efficacy and incidence of acute interstitial nephritis or acute renal failure between nafcillin intermittent (2 g every 4 hours for 30 minutes) vs continuous (6 g every 12 hours twice daily) infusion in patients with methicillin-sensitive *Staphylococcus aureus* bacteremia. Results showed no difference in the safety and efficacy between two infusion methods.

Linezolid
Linezolid injection (Zyvox®; Pfizer Inc., New York, NY, USA) should be administered by intravenous infusion over a period of 30–120 minutes. A clinical trial showed that 2-hour infusion of linezolid (600 mg) was the most effective regimen among three different infusion regimens. The PTA of the 4-hour infusion regimen was much lower compared with the 0.5 and 2-hour infusion regimens in terms of the ratio of the area under the curve to the MIC (AUC/MIC), whereas the PTA of 2-hour and 4-hour regimens rather than the 0.5-hour regimen remained 100% in terms of %T > MIC.46

De Pascale et al47 compared linezolid concentrations in plasma and epithelial lining fluid following administration in obese critically ill patients with VAP by 60-minute intermittent (600 mg every 12 hours) and continuous infusion (600 mg intravenous loading dose infused over 60 minutes, followed by 1,200 mg continuous infusion over 24 hours). Continuous infusion resulted in a significantly longer median time of linezolid plasma concentration persisted above MIC ranging from 1–4 mg/L and a higher epithelial lining fluid/plasma penetration ratio (98.8% vs 87.1%; *P*<0.001) compared with intermittent infusion.

Glycopeptide antibiotics
Vancomycin
Tafelski et al48 compared the effects of different dosing regimens on vancomycin target levels in critically ill patients. Intermittent administration in adults was as follows: 500 mg vancomycin every 6 hours given over 1 hour or 1 g every 12 hours given over 1 hour. The continuous infusion regimen was as follows: initiation of therapy with 1 g vancomycin as an infusion given over 1 hour, followed by 24-hour-continuous infusion of 1 g vancomycin in 50 mL of solution at different infusion rates (normal renal function: 4.2 mL/h; impaired renal function and CrCl of <50 mL/min: 2.1 mL/h; impaired renal function and CrCl of <20 mL/min: 1 mL/h). Patients with continuous administration achieved target serum levels significantly earlier and showed fewer sub-therapeutic serum levels (41% vs 11%, *P*<0.001).

Compared to intermittent dosing, continuous infusion of vancomycin could result in the following outcomes: (1) higher target attainment rate for therapeutic vancomycin exposure (48% vs 19%, *P*<0.001) and less between-patient variation in vancomycin serum level in critically ill patients;49 (2) greater (40% vs 21.5%, *P*=0.02) and more rapid achievement of goal plasma concentrations (2.04 vs 3.76 days, *P*<0.0001), higher pharmacodynamic target attainment (92.3% vs 30.8%, *P*<0.0001), and less dose adjustments, days of therapy, and mean total daily dose requirements per patient without significant difference in nephrotoxicity in neurosurgical ICU patients;50 (3) more quickly achieving the target level, greater percentage of keeping therapeutic 24-hour levels (100% vs 4.4%, *P*<0.001), and higher mean 24-hour vancomycin levels (20.35±2.78 µg/mL vs 9.7±3.52 µg/mL, *P*<0.001) in adult ICU patients during continuous venovenous hemofiltration;51 (4) higher serum vancomycin level (19.8±6.13 µg/mL vs 8.9±3.9 µg/mL, *P*<0.0001) and percentage of keeping drug level in the therapeutic range (57.1% vs 7.4%, *P*<0.0001) in trauma patients with suspected VAP. To avoid inadequate vancomycin level, continuous vancomycin infusion should be the optimal dosing strategy.52

Hanrahan et al53 conducted a meta-analysis of seven studies of vancomycin administration by continuous vs intermittent infusion and identified a non-significant trend of reduced nephrotoxicity in those who received vancomycin by continuous infusion (risk ratio=0.799, *P*=0.299). However, a newer meta-analysis of eleven studies by Hao et al54 revealed that patients treated with continuous infusion of vancomycin had a significantly lower incidence of nephrotoxicity compared with patients receiving intermittent infusion (relative risk=0.61, *P*<0.001), whereas the clinical efficacy was not significantly different.

Norvancomycin
Wu et al55 failed to observe significant advantages of a continuous infusion method of norvancomycin (ie, 0.8 g norvancomycin for 1 hour, and then another 0.4 g for 11 hours with...
extended infusion, followed by continuous infusion of 0.4 g for 12 hours) over the conventional infusion method (ie, 0.8 g norvancomycin for 1 hour every 12 hours) in an early stage after neurosurgery. The cerebral spinal fluid concentration in both groups reached or exceeded the MIC90 (2 mg/L) of target bacteria such as methicillin-resistant Staphylococcus aureus (MRSA). This pilot study of norvancomycin in a neurosurgical individual needs to be confirmed by further large-scale studies.

**Fosfomycin**

Alibero et al\(^5^6\) evaluated treatment regimens of the meropenem-fosfomycin combination against carbapenemase-producing Klebsiella pneumoniae (CPKP). For the 0.5-hour infusion, fosfomycin monotherapy regimens did not achieve a 90% PTA of >70% fT>MIC against the MIC50 or MIC90 in patients with normal renal function. Prolonged infusion of fosfomycin over 3 hours improved the PTA over that of a 0.5-hour infusion, however, fosfomycin monotherapy regimens did not reach 90% PTA against either the MIC50 or the MIC90 in patients with normal renal function. In combination therapy of meropenem, fosfomycin 6 g every 6 hours or 8 g every 8 hours as a 3-hour infusion had better chances of achieving PTA.

**Voriconazole**

Prescribing information for voriconazole (Vfend\(^5^8\); Pfizer Inc.) requires a maximum infusion rate of 3 mg/kg per hour over 1–2 hours. Hohmann et al\(^3^7\) observed that infusion rate could modulate voriconazole pharmacokinetics. Maximum concentration after a 4-hour infusion, and after a 6-hour infusion was 29% and 51% lower compared to a 400 mg/2-hour infusion, and fell outside of the bioequivalence margins (0.80–1.25). The AUC of 400 mg intravenous voriconazole was 16% lower when administered over 6 hours compared to 2 hours infusion. Metabolic clearance of CYP3A probe midazolam was 516 mL/min following 100 mg/4 hours voriconazole, 152 mL/min for 400 mg/6 hours, 192 mL/min for 400 mg/4 hours, and 202 mL/min for 400 mg/2 hours, which reflected an auto-inhibitory effect on CYP3A4-mediated voriconazole metabolism. Voriconazole efficacy correlates well in vivo with the AUC/MIC or mean unbound voriconazole concentration/MIC ratio.\(^5^8\) Thus, to avoid reduced exposure and potential efficacy, infusion rate of voriconazole should not deliberately be prolonged.

**Practical issues in the choice of infusion rate**

When implementing the prolonged or continuous infusion of antimicrobial therapy, several pragmatic issues should be considered, including availability of intravenous catheters, compatibility of intravenous drugs, drug stability, safety concerns, monitoring for efficacy and toxicity, and timing of administration.

**Compatibility**

Critically ill patients often receive multiple drugs via continuous intravenous infusion. Intravenous drugs may be infused concurrently via Y-site when compatibility data are available. For example, vancomycin was stable during the whole process and also during 72 hours exposure of concentrated solutions at temperatures up to 37°C, but careful attention must be paid to incompatibilities with β-lactams (temocillin, piperacillin/tazobactam, ceftazidime, imipenem, cefepime, and fluoroceillin), moxifloxacin, propofol, valproic acid, phenytoin, theophylline, methylprednisolone, and furosemide.\(^5^9\) For combination therapy of cefepime and vancomycin, worries about incompatibility between two antibiotics would result in conventional practice (ie, flush the tube with diluent and initiate vancomycin infusion when cefepime infusion over 30–60 minutes is completed). Tran et al\(^6^0\) evaluated whether an intravenous push of cefepime (a mixture of 10 mL normal saline injected over 2–5 minutes) was associated with a shorter time to vancomycin administration in the emergency department compared with intravenous piggyback infusion (a mixture of 100 mL normal saline infused over 30–60 minutes). The transition from intravenous piggyback infusion to intravenous push of cefepime could statistically significantly decrease the time to initiation of vancomycin administration by more than 1 hour to patients with suspected infections who were ordered both antibiotics. To prevent delay to MRSA coverage with vancomycin in the emergency department, it is reasonable to administer the first dose of cefepime as an intravenous push.

Meropenem/vaborbactam is administered as a 3-hour infusion given every 8 hours, thereby potentially restricting an intravenous line for 9 hours/day. Of the 88 medications tested, meropenem/vaborbactam was compatible with 73 (83%), including antibiotics such as aminoglycosides, colistin, fosfomycin, linezolid, tigecycline, and vancomycin. Physical incompatibility was observed with albumin, amiodarone, anidulafungin, calcium chloride, cefaroline, ciprofloxacin, daptofmycin, dobutamine, isavuconazole, midazolam, nocardipine, ondansetron, and phenytoin.\(^6^1\)

**Stability**

Continuous infusion of meropenem may be applied in clinical settings at normal and elevated temperatures. The 5 mg/mL...
aqueous reconstituted solution of meropenem was stable for up to 8 hours in the temperature range between 25°C and 35°C, and for up to 5 hours at 40°C. The 40 mg/mL solution of meropenem reconstituted in isotonic saline was stable at 25°C for a maximum of 8 hours. Therefore, continuous infusion of meropenem 3 g/day could be divided into six consecutive 4-hour continuous infusions of meropenem 0.5 g in consideration of meropenem stability. Nafcillin sodium was stable for at least 7 days at 25°C after reconstitution in 0.9% sodium chloride injection and storage in polypropylene syringes, which could support the application of continuous infusion (12 hours) of nafcillin.

**Infusion rate-related medication administration errors**

It is important to make sure that the fluid will be infused at the prescribed rate. Adverse events can occur if a large continuous infusion dose is administered via accidental rapid infusion. An observational study showed that errors in infusion rates accounted for 10.9% of drug-related problems associated with intravenous medication administration. A questionnaire survey showed that the wrong infusion rate (33.3%) was the most prevalent type of medication errors of nurses in the emergency department.

**Further opportunities**

There are many further opportunities in research and clinical practice with respect to optimal infusion rate in antimicrobial therapy. First, it is necessary to conduct head-to-head comparative studies of infusion regimens with different infusion rates from multiple perspectives (eg, clinical outcomes, resources utilization, nurse satisfaction, cost-effectiveness). Second, dosing and administration strategies should be individualized on the basis of the pathogen–drug–patient interplay to optimize outcomes. For time-dependent antimicrobials, the factors affecting the indice %T>MIC or %T<MIC (eg, dose, dosing frequency, infusion rate, MIC value, protein binding, clearance alternations) may have an influence on the difference between two infusion methods. In this review, the factors determining whether prolonged or continuous infusion has a benefit over standard infusion are summarized in Table 2. Therefore, subgroup analysis in specific circumstances and populations should be encouraged, and their findings may provide explanations for the inconsistency of conclusions derived from outcome comparison between extended infusion and standard infusion of antimicrobials. Also, there may be a very pronounced difference in PK/PD of antibiotics in neonates and adults; therefore, the conclusion drawn from adult studies could not be simply extrapolated to neonates or pediatric patients.

Third, implementing effective interventions is required to enhance the application of optimal infusion strategies. A survey of Australia and New Zealand ICU doctors and pharmacists showed that a greater proportion of pharmacists compared with doctors believed continuous infusion to be more effective than intermittent administration (85.4% vs 34.3%, respectively; \( P<0.001 \)). The pharmacy and therapeutics committee should draft an institutional protocol for the optimal infusion rate of some drugs and implement extensive quality assurance monitoring (eg, retrospective analysis of guideline compliance). Key elements contributing to the successful implementation of a piperacillin/tazobactam extended-infusion guideline included consistent pharmacy leadership, multidisciplinary involvement, thorough in servicing to health care professionals, hospital-wide implementation, and extensive quality assurance monitoring. Sanford Medical Center successfully achieved a great total cost savings of about 2 million US dollar to the hospital (ie, reduction in piperacillin/tazobactam expenditures and additional cost savings due to reduced LOS by 0.6 days) in the first 2 years through automatic conversion from a 30-minute infusion to a 4-hour extended infusion of piperacillin/tazobactam along with staff education and use of smart pump technology.

Fourth, prescribing information on the infusion rate of some medications should be revised according to the updated evidence. Clinicians may not dare to prescribe and administer antimicrobial agents with new infusion strategies due to potential medical disputes and hidden dangers caused by off-label use.

**Conclusion**

To maximally preserve the effectiveness of current antimicrobials and prevent further antimicrobial resistance, effective interventions should be implemented to enhance the application of optimal infusion strategies. For reducing nephrotoxicity, prolonged infusion of meropenem over 4 hours is better than the conventional infusion strategy (30 minutes) for meningitis and Pseudomonas infection in neonates with Gram-negative late-onset sepsis; and continuous infusion of vancomycin is superior to intermittent infusion. For increasing drug efficacy, prolonged or continuous infusion of time-dependent antibiotics (eg, meropenem, doripenem, imipenem, cefepime, ceftazidime, piperacillin/tazobactam, linezolid, and vancomycin) is an optimal choice. Extended infusion of piperacillin/tazobactam has pharmacoeconomic
advantage over standard infusion. Nevertheless, the advantage of prolonged or continuous infusion vs regular infusion of time-dependent antibiotics may only exhibit in special clinical circumstance and special populations (eg, patients with SOFA score ≥9, respiratory tract infections, urinary or intra-abdominal infections, or infections caused by less susceptible pathogens, benefited from prolonged infusion (eg, shorter days of antibiotics use and ventilator time, longer survival, better clinical efficacy, and lower 28-day mortality rate).41

Piperacillin/tazobactam: For pathogens with MICs of ≤8 mg/L, extended infusion did not substantially improve PTA, and standard infusion was likely sufficient. However, piperacillin/tazobactam 100/12.5 mg/kg given as an extended infusion every 6–8 hours may be optimal for empirical or directed therapy in critically ill pediatric patients with infections caused by less susceptible pathogens.40

Piperacillin/tazobactam: Patients with severe disease (SOFA score ≥9), rather than those with mild disease (SOFA score < 9), benefited from prolonged infusion (eg, shorter days of antibiotics use and ventilator time, longer survival, better clinical efficacy, and lower 28-day mortality rate).41

Piperacillin/tazobactam: ICU patients exhibited no significant differences in outcomes following two dosing methods (extended infusion over 4 hours vs regular infusion over 30 minutes), whereas patients with urinary or intra-abdominal infections had lower mortality and clinical failure rates following extended infusion treatment.42

Piperacillin/tazobactam: There was a lower mortality rate in the extended-infusion group who had infectious organisms identified (9.3% vs 22.4%, P = 0.01) or were diagnosed with respiratory tract infections (8.9% vs 18.7%, P = 0.02).43

Meropenem: ICU patients exhibited the lowest target attainment rates, whereas internal medicine patients achieved the highest target attainment rates following short (0.5-hour) and prolonged (3-hour) infusion regimens of 1 g meropenem every 8 hours.14

Meropenem: In adult patients with septic shock and possible increased renal clearance, meropenem doses should be increased and/or administration by prolonged or continuous infusion should be considered to increase the likelihood of obtaining therapeutic drug concentrations. In patients with normal renal function, however, standard intermittent dosing (30 minutes) seems to be sufficient.15

Imipenem/cilastatin: In critically ill patients with nosocomial pneumonia, concentrations of imipenem were above the target concentration (4×MIC of 2 mg/L) for ≥40% of the dosing interval in every patient in the bolus group (1 g imipenem/1 g cilastatin over 30 minutes every 8 hours), whereas in the extended group (0.5 g of imipenem by a 3-hour infusion every 6 hours) this PK/PD index (40% TT > 4×MIC) was achieved in only 20% of patients. Moreover, 70% of the patients in the extended group did not reach the desired drug concentrations at all.35

Rational infusion rates

***Table 2*** The factors determining whether prolonged or continuous infusion has a benefit over standard infusion

<table>
<thead>
<tr>
<th>Factors</th>
<th>Descriptions on whether prolonged or continuous infusion has a benefit over standard infusion</th>
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| MIC of bacterial pathogens           | Meropenem: The PTA (ie, 40% T>MIC) with short (0.5-hour) infusion of 1 g meropenem every 8 hours was higher than those with prolonged (3-hour) infusion regimens for CPKP isolates with MIC 4–8 mg/L rather than MIC ≥16 mg/L or ≤2 mg/L (MIC 4 mg/L: 98%–99% vs 61%–83%; MIC 8 mg/L: 55%–79% vs 23%–33%).14
Piperacillin/tazobactam: The benefits of prolonged infusion were selective and most likely observed in patients with less susceptible pathogens, ie, prolonged infusion had no advantages over standard infusion against isolates with susceptible MICs of 8 or 16 mg/L, whereas it produced more than twice the final bacterial kill against less susceptible isolates with an intermediate MIC of 32 mg/L.39 |
| Diagnosis and disease severity       | Piperacillin/tazobactam: Patients with severe disease (SOFA score ≥9), rather than those with mild disease (SOFA score < 9), benefited from prolonged infusion (eg, shorter days of antibiotics use and ventilator time, longer survival, better clinical efficacy, and lower 28-day mortality rate).41
Piperacillin/tazobactam: ICU patients exhibited no significant differences in outcomes following two dosing methods (extended infusion over 4 hours vs regular infusion over 30 minutes), whereas patients with urinary or intra-abdominal infections had lower mortality and clinical failure rates following extended infusion treatment.42
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Meropenem: ICU patients exhibited the lowest target attainment rates, whereas internal medicine patients achieved the highest target attainment rates following short (0.5-hour) and prolonged (3-hour) infusion regimens of 1 g meropenem every 8 hours.14 |
| Bacterial density                    | Piperacillin/tazobactam: for the lower initial bacterial density, trough total plasma piperacillin concentration/MIC ratios of 3.4 and 10.4 following bolus and extended-infusion regimens, respectively, were able to suppress the emergence of piperacillin resistance. For the higher initial bacterial density, both infusion regimens were associated with progressive growth of a resistant subpopulation, and thus combination therapy may be required to maximize bacterial killing and prevent antimicrobial resistance.44
Imipenem/cilastatin: In critically ill patients with nosocomial pneumonia, concentrations of imipenem were above the target concentration (4×MIC of 2 mg/L) for ≥40% of the dosing interval in every patient in the bolus group (1 g imipenem/1 g cilastatin over 30 minutes every 8 hours), whereas in the extended group (0.5 g of imipenem by a 3-hour infusion every 6 hours) this PK/PD index (40% TT > 4×MIC) was achieved in only 20% of patients. Moreover, 70% of the patients in the extended group did not reach the desired drug concentrations at all.35 |
| Total daily dose                     | Meropenem: In adult patients with septic shock and possible increased renal clearance, meropenem doses should be increased and/or administration by prolonged or continuous infusion should be considered to increase the likelihood of obtaining therapeutic drug concentrations. In patients with normal renal function, however, standard intermittent dosing (30 minutes) seems to be sufficient.15 |
| Renal clearance                      | Imipenem/cilastatin: In critically ill patients with nosocomial pneumonia, concentrations of imipenem were above the target concentration (4×MIC of 2 mg/L) for ≥40% of the dosing interval in every patient in the bolus group (1 g imipenem/1 g cilastatin over 30 minutes every 8 hours), whereas in the extended group (0.5 g of imipenem by a 3-hour infusion every 6 hours) this PK/PD index (40% TT > 4×MIC) was achieved in only 20% of patients. Moreover, 70% of the patients in the extended group did not reach the desired drug concentrations at all.35 |

**Abbreviations**: CPKP, carbapenemase-producing Klebsiella pneumoniae; ICU, intensive care unit; MIC, minimum inhibition concentration; PK/PD, pharmacokinetic/pharmacodynamic; PTA, probability of target attainment; SOFA, sequential organ failure assessment; T>MIC, time for which drug concentrations exceed the MIC.

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**Disclosure**

The authors report no conflicts of interest in this work.

**References**


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