Meropenem in the treatment of complicated skin and soft tissue infections

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Abstract: Meropenem is a broad-spectrum carbapenem antibiotic with excellent activity against many pathogens associated with complicated skin and soft tissue infections (cSSTIs). At least three studies have shown meropenem to have good clinical efficacy and to be well tolerated in the treatment of cSSTIs. Two open-label studies compared meropenem 500 mg every 8 hours (total evaluable n=146) with imipenem/cilastatin 500 mg every 6 hours (n=147). Clinical efficacy rates in evaluable patients 7–14 days after end of treatment were similar, 92% and 100% in meropenem-treated groups versus 89% and 100% in groups receiving imipenem/cilastatin. An additional prospective, randomized, double-blind study evaluated meropenem 500 mg every 8 hours (261 evaluable patients) versus imipenem/cilastatin 500 mg every 8 hours (287 patients). Clinical efficacy rates of meropenem and imipenem/cilastatin 7–28 days after end of treatment were 86.2% and 82.9%, respectively. Meropenem was well tolerated in all studies. Carbapenems are currently recommended as appropriate for initial treatment of certain cSSTIs such as those likely to involve mixed and/or multidrug-resistant pathogens. Meropenem is an effective and safe alternative for monotherapy when used for appropriate types of cSSTIs. Higher doses (ie, 1 g every 8 hours) should be considered for treatment of cSSTIs in higher-risk patients where Pseudomonas aeruginosa is a suspected or documented pathogen.

Keywords: meropenem, carbapenems, wound infection, diabetic foot, antibiotics, nosocomial infection

Introduction

Skin and soft tissue infections (SSTIs) are very frequently encountered in clinical practice and are one of the most common sites of bacterial infections (Nichols 1999; Eron et al 2003; DiNubile and Lipsky 2004). These infections are also among the most common indications for antibiotic therapy and hospital admissions in the US (CDCP 2001). Complicated skin and soft tissue infections (cSSTIs) are those which involve abnormal skin or wounds, occur in a compromised host, or require substantial surgical intervention (DiNubile and Lipsky 2004). Because cSSTIs are by definition more severe in nature and are often associated with complications such as bacteremia and sepsis, patients with these infections are often hospitalized for management with intravenous antibiotics (Nichols 1999; Swartz 2000; Eron et al 2003; Stevens et al 2005). Many cSSTIs are also nosocomial in origin; it has been estimated that these infections are the most common nosocomial infection among surgical patients and occur in approximately 2.6% of all operations (Wilson 2003).

Meropenem (Merrem®, AstraZeneca Pharmaceuticals, Wilmington, Delaware, USA) is a broad-spectrum antibacterial agent of the carbapenem class. Meropenem has excellent activity against a broad range of bacteria including many Gram-positive and Gram-negative pathogens, including many potentially resistant strains such as Pseudomonas aeruginosa, as well as anaerobic organisms (Wiseman et al 1995; Fish and Singletary 1997). Because cSSTIs often involve a diverse range of both aerobic and anaerobic pathogens, meropenem was studied for the treatment of cSSTIs.
and was approved by the US Food and Drug Administration (FDA) for this indication in May 2005. The purpose of this article is to briefly review the clinical characteristics and bacterial etiology of cSSTIs, discuss pharmacological and clinical data pertaining to the use of meropenem for these infections, and provide recommendations regarding the role of meropenem in the treatment of cSSTIs.

Complicated skin and soft tissue infections

Clinical characteristics

The majority of SSTIs are classified as uncomplicated infections, ie, infections defined as involving only superficial layers of the skin (epidermis, dermis, and subcutaneous tissues) (Eron et al 2003; Stevens et al 2005). Uncomplicated infections such as impetigo, erysipelas, simple abscesses, and simple cellulitis are typically mild to moderate in severity and easily treated with local care with or without oral antibiotic therapy (Eron et al 2003; Raghavan and Linden 2004; Stevens et al 2005). By contrast, cSSTIs involve deeper skin structures such as fascia or muscle layers and require surgical intervention for effective management (Eron et al 2003; Fung et al 2003; DiNubile and Lipsky 2004). These complicated infections are often associated with significant comorbidities such as diabetes mellitus, peripheral vascular disease, critical illness, human immunodeficiency virus (HIV) infection, and other immunocompromised states such as malignancy or post-transplantation (Nichols 1999; Swartz 2000; Fung et al 2003; DiNubile and Lipsky 2004; Stevens et al 2005). Examples of complicated infections include more complex cellulitis and abscesses, perirectal abscesses, post-traumatic or surgical site infections, myositis, necrotizing fasciitis, and infected diabetic and vascular ischemic ulcers. These infections can be challenging to effectively manage because of the great diversity in pathophysiology, microbial etiology, clinical manifestations, severity, and potential for additional complications such as bacteremia, sepsis syndrome, and organ dysfunction (Swartz 2000; Eron et al 2003; Lipsky et al 2004; Stevens et al 2005).

The pathophysiology of cSSTIs is also significant in terms of potential effects on antimicrobial efficacy. Since the efficacy of antimicrobials depends on the ability to achieve adequate concentrations at a particular site of infection, good distribution to infected tissues is a key feature of any agent. However, cSSTIs are commonly associated with vascular insufficiency due to underlying comorbidities, disruption of normal blood and lymphatic flows, and/or areas of devitalized tissue, all of which may severely limit the penetration of drugs to the site of bacterial infection. The possibility of significantly decreased penetration of antimicrobials to tissues involved in cSSTIs thus may potentially lead to inadequate drug concentrations, clinical and microbiological treatment failure, and the development of resistance among bacterial pathogens. Successful antimicrobial management of cSSTIs therefore depends on the use of agents possessing the combination of good intrinsic activity against potential pathogens and intrinsic ability to distribute to tissues of the skin and skin structures, plus use of relatively high doses which increase the likelihood of sufficient tissue penetration.

Bacterial etiology

Complicated SSTIs are associated with a broad range of potential pathogens (Table 1) (Doern et al 1999; Jones et al 1999, 2003; Bowler et al 2001; Fung et al 2003; Rennie et al 2003; Jodrá et al 2006). Aerobic Gram-positive cocci are the predominant pathogens in most types of cSSTIs, particularly those that are community-acquired in origin. *Staphylococcus aureus*, other staphylococci, and various

<table>
<thead>
<tr>
<th>Organism</th>
<th>Frequency of isolation</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Gram-positive aerobes</strong></td>
<td></td>
</tr>
<tr>
<td><em>Staphylococcus aureus</em> (all)</td>
<td>29–46</td>
</tr>
<tr>
<td><em>Staphylococcus aureus</em> (MR)</td>
<td>3–15</td>
</tr>
<tr>
<td>Coagulase-negative staphylococci</td>
<td>3–5</td>
</tr>
<tr>
<td>Streptococci</td>
<td>7–20</td>
</tr>
<tr>
<td>Enterococcus spp.</td>
<td>3–8</td>
</tr>
<tr>
<td><strong>Gram-negative aerobes</strong></td>
<td></td>
</tr>
<tr>
<td><em>Pseudomonas aeruginosa</em></td>
<td>22–26</td>
</tr>
<tr>
<td><em>Escherichia coli</em></td>
<td>3–11</td>
</tr>
<tr>
<td>Enterobacter spp.</td>
<td>5–6</td>
</tr>
<tr>
<td>Klebsiella pneumoniae</td>
<td>3–4</td>
</tr>
<tr>
<td>Proteus spp.</td>
<td>4–6</td>
</tr>
<tr>
<td>Acinetobacter spp.</td>
<td>ND</td>
</tr>
<tr>
<td>Serratia spp.</td>
<td>2–3</td>
</tr>
<tr>
<td><strong>Anaerobes</strong></td>
<td></td>
</tr>
<tr>
<td>Bacteroides spp.</td>
<td>17–33</td>
</tr>
<tr>
<td>Peptostreptococcus spp.</td>
<td>5–13</td>
</tr>
<tr>
<td>Prevotella spp.</td>
<td>4–8</td>
</tr>
<tr>
<td>Clostridium spp.</td>
<td>3</td>
</tr>
</tbody>
</table>

Note: *Infections in patients requiring hospitalization for treatment. Abbreviations: MR, methicillin-resistant; ND, no data reported.*

Table 1 Bacterial pathogens associated with complicated skin and soft tissue infections (Doern et al 1999; Jones et al 1999, 2003; Bowler et al 2001; Fung et al 2003; Rennie et al 2003; Jodrá et al 2006)
streptococci (particularly of groups A and B) are the most
common organisms (Bowler et al 2001; Fung et al 2003;
Raghavan and Linden 2004). The incidence of infections
caused by methicillin-resistant \textit{S. aureus} (MRSA) has been
steadily increasing; MRSA now accounts for approximately
60% of all \textit{S. aureus} isolates from nosocomial infections in
the US (DPH 2004). Even more alarming is the growing
incidence of infections due to community-acquired MRSA.
Although MRSA has been traditionally regarded as a
hospital-acquired pathogen, this organism has also emerged
as a common cause of community-acquired infections with
approximately 30% of all MRSA isolates now community-
acquired in origin (Burkharie et al 2001; Eady and Cove
2003). A recent study of MRSA infections in three
geographic areas within the US reported that 17% of all
isolates were community-acquired in origin (Fridkin et al
2005). This study also found that 77% of these infections
involved skin and soft tissue, and the overall incidence of
community-acquired MRSA infections was significantly
higher among persons less than two years of age. Specifically
in cSSTIs, MRSA has been reported in up to
10%–15% of patients overall, and 5%–15% of patients with
nosocomial infections (Jones et al 1999, 2003; Fung et al
2003; Rennie et al 2003; DPH 2004; Raghavan and Linden
2004). Enterococci are also common pathogens in cSSTIs
and are isolated with somewhat greater frequency in
nosocomial infections (Jones et al 1999, 2003; Bowler et al
2001; Raghavan and Linden 2004). Up to 20% of
enterococci isolated from these infections are now
vancomycin-resistant (VRE) (Rennie et al 2003; Raghavan
and Linden 2004). \textit{S. aureus}, enterococci, and coagulase-
negative staphylococci are particularly important pathogens
in surgical wound infections and together account for
approximately 46% of all surgical site infections; both
MRSA and VRE occur more frequently in these infections
as well (DiNubile and Lipsky 2004; DPH 2004).

Gram-negative organisms which are frequently seen in
cSSTIs include the Enterobacteriaceae (particularly
\textit{Escherichia coli}, \textit{Enterobacter} spp., \textit{Klebsiella} spp., and
\textit{Proteus} spp.) and \textit{P. aeruginosa} (Jones et al 1999, 2003;
Swartz 2000; Fung et al 2003; Rennie et al 2003; Stevens et
al 2005). Complicated skin and soft tissue infections
commonly associated with \textit{P. aeruginosa} include lower
extremity infections (particularly in patients with
comorbidities such as vascular insufficiency or diabetes),
surgical wound infections, chronic renal disease, and
diabetes or other immunocompromising diseases (Bowler
et al 2001; Pellizer et al 2001; Eron et al 2003; Rennie et al
2003; DPH 2004; Lipsky et al 2004). Of particular concern
is the rapid spread of resistance mediated by extended-
spectrum \textit{\beta}-lactamases (ESBLs) among organisms such as
\textit{Klebsiella pneumoniae} and \textit{Escherichia coli}. Organisms that
produce ESBLs are usually resistant to multiple
antimicrobials including third-generation (eg, ceftriaxone,
cefotaxime, ceftazidime) and fourth-generation (ie,
cefepime) cephalosporins and aztreonam (Bush 2001;
Paterson et al 2003), and are also associated with high rates
of resistance to aminoglycosides and fluoroquinolones
(Paterson et al 2000, 2003). Although recent data for cSSTIs
are not readily available, a recent international study of
Gram-negative bacilli isolated from intra-abdominal
infections found 7% of \textit{E. coli}, 13% of \textit{Klebsiella} spp., and
18% of \textit{Enterobacter} spp. were ESBL producers (Chow et
al 2006). These data are consistent with data reported
specifically from the US, although rates may be higher in
certain geographical areas (DPH 2004; Pfaller and Segreti
2006). Resistance of \textit{P. aeruginosa} to fluoroquinolones,
ceftazidime, and imipenem/cilastatin has also increased
rapidly; nearly 10% of \textit{P. aeruginosa} isolates are now
resistant to multiple drug classes including cephalosporins,
carbapenems, aminoglycosides, and/or fluoroquinolones
(Karlowsky et al 2003). Reports of fluoroquinolone
resistance are becoming more common among organisms
such as \textit{E. coli} and \textit{Proteus} spp. that are usually considered
to be very susceptible to this class of drugs (Neuhauser et al

Anaerobic bacteria such as \textit{Bacteroides fragilis}, other
bacteria of the \textit{Bacteroides} group, \textit{Fusobacterium} spp.,
\textit{Prevotella} spp., \textit{Clostridium} spp., and \textit{Peptostreptococcus}
spp. are also relatively common causes of certain cSSTIs.
Types of infections in which anaerobes are commonly found
include traumatic wounds, infections in diabetic and other
immunocompromised patients, and infections associated
with ischemic tissue injury (Jones et al 2003). Increasing
resistance to common antibiotics such as clindamycin and
second-generation cephalosporins (eg, cefoxitin, cefotetan)
has been documented among anaerobic bacteria such as \textit{B.
fragilis} and organisms of the \textit{Bacteroides} group (Hecht
2004). However, agents such as metronidazole, piperacillin/
tazobactam, and the carbapenems still maintain excellent
activity against most clinically relevant anaerobic bacteria.

This broad range of potential pathogens is further
complicated by the fact that many cSSTIs are polymicrobial
and represent a variety of mixed infections, ie, presence of
aerobic Gram-positive and Gram-negative pathogens as well
as anaerobes (Nichols 1999; Fung et al 2003; Stevens et al
2005). Classic mixed aerobic and anaerobic infections are particularly common in surgical wound infections, necrotizing infections including fasciitis and myositis, complex and perirectal abscesses, diabetic and ischemic ulcers, and complicated cellulitis in diabetics and other immunocompromised patients (Nichols 1999; Pellizer et al 2001; Fung et al 2003; Lipsky et al 2004; Stevens et al 2005; Vinh and Embil 2005).

The recent emergence and spread of antibiotic resistance among both Gram-positive and Gram-negative pathogens have created a significant clinical challenge in the appropriate management of cSSTIs (Colsky et al 1998; Sader et al 2002; Rennie et al 2003). Although a number of drugs for treatment of multidrug-resistant organisms such as MRSA and VRE have become available or are in development (eg, linezolid, daptomycin, tigecycline, dalbavancin), there are few drugs under development for treatment of Gram-negative or anaerobic pathogens which are resistant to the currently available agents. Thus there is a clear need for new antibiotic options for treatment of cSSTI as a result of increasing bacterial resistance. What is equally clear is that the use of currently available agents also needs to be optimized to increase or improve the effective therapeutic options.

Considerations in antibiotic selection

Appropriate therapeutic strategies for cSSTIs depend on a number of factors including whether the infection is community- versus hospital-acquired in origin, virulence of the infecting pathogens, severity, and presence and type of comorbidities. Surgical intervention will often be required and antibiotic therapy is often considered an adjunctive rather than a primary therapy for certain infections, eg, diabetic foot infections or necrotizing fasciitis (Eron et al 2003; Lipsky et al 2004; Stevens et al 2005). However, adequate antibiotic therapy remains a crucial component of appropriate management of cSSTIs. Empirical antibiotic therapy should usually be promptly initiated after obtaining samples for culture in order to begin appropriate treatment and slow the progression of the infection (Nichols 1999; Eron et al 2003; Stevens et al 2005).

Selection of a specific antibiotic regimen for treatment of cSSTIs depends on many factors such as etiology of the infection (ie, community- or hospital-acquired), knowledge of local or institutional antibiotic susceptibility patterns, availability of suitable antibiotics which provide appropriate spectrum of activity for known or presumed pathogens, pharmacokinetic and pharmacodynamic considerations such as penetration of infected tissues and ability to achieve suitable drug concentrations, adverse effect and drug interaction profiles, and drug cost (Nichols 1999; Eron et al 2003; Fung et al 2003; DiNubile and Lipsky 2004). Because of the broad range of pathogens associated with cSSTIs and the frequency of mixed infections involving both aerobic and anaerobic bacteria, empirical antibiotic regimens are usually broad in spectrum and provide coverage against both Gram-positive and Gram-negative pathogens, with the addition of anaerobic activity also often desired (Nichols 1999; Eron et al 2003; Fung et al 2003; Lipsky et al 2004; Stevens et al 2005). Although streamlining of antibiotic therapy to narrower-spectrum regimens is appropriate once results of microbiological cultures are known, initial antibiotics must be sufficiently broad to cover potential pathogens until other management modalities (ie, surgical intervention) can be instituted.

It has been well documented that increasing resistance among bacterial pathogens results in inappropriate selection of antibiotic therapy for hospital-acquired pneumonia and bloodstream infections (Rello et al 1993; Alvarez-Lerma et al 1996; Luna et al 1997; Leibovici et al 1998; Carmeli et al 1999; Kollef et al 1999; Garnacho-Montero et al 2003). Furthermore, the relationship between inappropriate antibiotic selection and resultant mortality in pneumonia and sepsis has been clearly shown (Rello et al 1993; Alvarez-Lerma et al 1996; Luna et al 1997; Leibovici et al 1998; Carmeli et al 1999; Kollef et al 1999). Similar data specifically addressing the association between inappropriate antibiotic selection and subsequent adverse outcomes are not as plentiful for cSSTIs. However, one retrospective study of 137 patients with intra-abdominal infections and SSTIs treated with cefotetan, cefoxitin, or ampicillin/sulbactam monotherapy found that isolation of a pathogen resistant to the antibiotic treatment regimen (particularly *P. aeruginosa* was an independent risk factor for clinical treatment failure with multivariate analysis (odds ratio [OR] = 14.9; p=0.001) (Falagas et al 1996). Improper antibiotic selection thus has adverse consequences in cSSTIs as well. Because the bacterial etiology of cSSTIs often involves polymicrobial infections and because of the changing patterns of antibiotic susceptibilities among both Gram-positive and Gram-negative pathogens, the empirical use of antibiotics with a broad spectrum of activity which includes multidrug-resistant organisms is becoming increasingly common and ever more appropriate (Nichols 1999; Eron et al 2003; Fung et al 2003; Lipsky et al 2004; Stevens et al 2005).
Meropenem in the treatment of cSSTIs

Antibacterial spectrum of activity

Meropenem is a broad-spectrum carbapenem antibiotic that possesses excellent activity against both aerobic Gram-positive and aerobic Gram-negative bacteria, and also covers common anaerobes. Based on published MIC$_{90}$s (the minimum inhibitory concentration at which 90% of tested strains are inhibited), meropenem is generally similar to or slightly less active than imipenem/cilastatin or ertapenem against Gram-positive aerobic bacteria. However, these differences are not likely to be clinically significant for most pathogens because MIC$_{90}$s are usually well below recommended susceptibility breakpoints (Wiseman et al 1995; Fish and Singletary 1997; Merrem® product package insert 2005). Meropenem is active against methicillin-susceptible S. aureus and most strains of methicillin-susceptible coagulase-negative staphylococci. However, as with other carbapenems, meropenem has poor activity against MRSA and methicillin-resistant coagulase-negative staphylococci. Meropenem also has excellent activity against most streptococci, including Streptococcus pyogenes and S. agalactiae. The activity of meropenem against enterococci varies considerably among different species. Most strains of Enterococcus faecalis, E. avium and E. liquefaciens are susceptible or moderately susceptible to meropenem, but most strains of E. faecium and VRE of any species are resistant.

Meropenem has impressive in vitro activity against Gram-negative aerobic bacteria and compares very favorably against both imipenem/cilastatin and ertapenem. Meropenem is generally slightly more active than imipenem/cilastatin (based on MIC$_{90}$s) and similar in activity to ertapenem against clinical isolates of Enterobacteriaceae. Meropenem has shown excellent efficacy against clinical isolates resistant to other β-lactam agents, including many strains which were highly resistant to third-generation cephalosporins, piperacillin, and aminoglycosides. Meropenem and other carbapenems are considered to be drugs of choice for the treatment of the ESBL-producing Gram-negative bacilli (Paterson et al 2000, 2003). Although meropenem generally has 1- to 2-fold lower MIC$_{90}$s than imipenem/cilastatin against P. aeruginosa, these two drugs appear to be relatively similar in their in vitro activity against clinical isolates of this organism. However, meropenem and imipenem/cilastatin are affected by somewhat different mechanisms of resistance among P. aeruginosa strains and cross-resistance among the two drugs is not complete (Fish and Singletary 1997; Rhomberg et al 2005; Quale et al 2006). The Meropenem Yearly Susceptibility Test Information Collection (MYSTIC) surveillance database of antibiotic susceptibilities reports that approximately 7% of P. aeruginosa strains are resistant to imipenem/cilastatin but susceptible to meropenem, while only approximately 1% of strains were susceptible to imipenem/cilastatin and resistant to meropenem (AstraZeneca 2006). Such discordant susceptibilities are consistent with data published elsewhere (Wiseman et al 1995; Fish and Singletary 1997; Mutnick et al 2004), although the drugs are considered to be equivalent in P. aeruginosa activity under most clinical circumstances. Ertapenem is not reliably active against P. aeruginosa and is not generally considered to be clinically useful for infections caused by this organism (Fuchs et al 2001; Livermore et al 2001).

Meropenem is also active against most strains of clinically significant anaerobes. Meropenem displayed MIC$_{90}$s which were quite similar to those for imipenem/cilastatin and ertapenem in Gram-negative isolates including Bacteroides fragilis, Bacteroides group, and Fusobacterium spp. Meropenem MICs for these organisms are usually substantially lower than those of metronidazole, clindamycin, cefoxitin, or piperacillin/tazobactam. Against Gram-positive anaerobic bacteria such as Clostridium perfringens, C. difficile and Veillonella spp., meropenem is equal to or slightly more active than imipenem/cilastatin, ertapenem, metronidazole, clindamycin, cefoxitin or piperacillin/tazobactam. Meropenem also has excellent activity against clinical isolates of Peptostreptococcus spp., Propionibacterium acnes, Actinomyces spp., and Actinobacillus spp. (Wiseman et al 1995; Fish and Singletary 1997).

Considerations regarding pharmacokinetics/pharmacodynamics and tissue penetration

Meropenem achieves mean peak plasma concentrations (C$_{max}$) and minimum plasma concentrations (C$_{min}$) of approximately 23–25 mg/L and 0.1–0.2 mg/L, respectively, following the administration of single or multiple 30-minute intravenous infusions of 500 mg to healthy adult volunteers (Maglio et al 2003; Merrem® product package insert 2005). Although C$_{min}$ values for meropenem are relatively low, the drug is extremely active against many pathogens commonly found in cSSTIs with MIC$_{90}$s of ≤0.12 mg/L for most
Enterobacteriaceae, methicillin-susceptible *S. aureus* and coagulase-negative staphylococci, and streptococci (Rhomberg et al 2005). Studies of carbapenems indicate that the percentage of free (ie, not protein bound) drug remain above the MIC of the pathogen (referred to as the percent time above MIC, or % T>MIC) is the key pharmacodynamic parameter which is related to clinical and microbiological efficacy of the drugs. A ≥30% T>MIC is associated with bacteriostatic activity of the carbapenems, while a ≥40% T>MIC is more predictably bactericidal and associated with prevention of the development of resistance (Craig 1998; Turnidge 1998). Meropenem 500 mg every 8 hours achieves T>MIC of ≥40% for most susceptible pathogens and would be expected to be effective in the treatment of cSSTIs caused by these organisms at this dose.

Meropenem appears to be widely distributed in various body tissues and fluids (Hutchison et al 1995; Mouton et al 1995; Fish and Singletary 1997). Specifically regarding the use of meropenem in the treatment of cSSTIs, several early studies demonstrated that meropenem achieves concentrations adequate for the treatment of infections caused by susceptible bacteria in skin, burned skin, fascia, skeletal muscle, and blister fluid (Table 2). Meropenem concentrations in these various tissues were quite variable and ranged from 9% to 85% of simultaneous serum concentrations; however, the absolute concentrations were nevertheless in excess of the susceptibility breakpoints (MIC ≤4 mg/L) of bacteria commonly found in cSSTIs (Hutchison et al 1995; Mouton et al 1995; Fish and Singletary 1997). A more recent study also evaluated the pharmacokinetic disposition of meropenem in plasma and cantharidin-induced blister fluid (Maglio et al 2003). The mean penetration of meropenem into blister fluid was determined to be 67% when calculated by comparing the pharmacokinetic area under the concentration-time curve (AUC) of the drug in blister fluid with the AUC of meropenem in plasma. Additionally, this study found that meropenem doses of 500 mg every 8 hours maintains a ≥50% T>MIC in blister fluid for most susceptible pathogens. This study thus indicated that meropenem 500 mg every 8 hours achieves concentrations in blister fluid sufficient for the treatment of SSTIs caused by susceptible organisms (Maglio et al 2003).

Another recent study used data concerning pharmacokinetics and penetration into blister fluid in order to model the pharmacodynamics of meropenem, imipenem/cilastatin and piperacillin/tazobactam in the treatment of cSSTIs (Ong et al 2005). This study utilized Monte Carlo simulation, a statistical modeling tool which combines both pharmacokinetic and MIC information to predict the probability of achieving desired pharmacodynamic targets with various drug dosing regimens and against specific pathogens. Optimal pharmacodynamic targets with common dosing regimens of each specific agent were defined as ≥90% probability of ≥40% T>MIC (meropenem and imipenem/cilastatin) or ≥50% T>MIC (piperacillin/tazobactam) in serum and blister fluid. Meropenem 500 mg every 8 hours, imipenem/cilastatin 500 mg every 8 hours, and piperacillin/tazobactam 3.375 g every 6 hours were all determined to be optimal choices for empiric treatment of cSSTIs. This study offers further support for the suitability of meropenem as an appropriate agent for the treatment of these infections (Ong et al 2005).

The further application of Monte Carlo simulation to meropenem pharmacodynamics raises important questions related to the optimal dosing of this agent for cSSTIs. Although the previously cited studies provide pharmacokinetic and pharmacodynamic data indicating that meropenem 500 mg every 8 hours is an adequate regimen for infections involving *most* susceptible pathogens, the adequacy of this regimen specifically for infections

<table>
<thead>
<tr>
<th>Body tissue or fluid</th>
<th>N</th>
<th>Dose (g)</th>
<th>Sample time (hrs)</th>
<th>Mean drug concentration (mg/L) Tissue/Fluid</th>
<th>Serum</th>
<th>Tissue/Fluid:Serum Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Skeletal muscle</td>
<td>14</td>
<td>1</td>
<td>0.5–4.5</td>
<td>0.93–6.10</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Fascia</td>
<td>18</td>
<td>1</td>
<td>0.5–5.5</td>
<td>0.99–8.76</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Skin</td>
<td>3</td>
<td>0.5</td>
<td>0</td>
<td>3.97</td>
<td>42.6</td>
<td>9.6%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.5</td>
<td>1</td>
<td>3.4</td>
<td>11.9</td>
<td>33.5%</td>
</tr>
<tr>
<td>Burned skin</td>
<td>2</td>
<td>1</td>
<td>0</td>
<td>5.5</td>
<td>64.3</td>
<td>8.8%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1</td>
<td>1</td>
<td>9.2</td>
<td>14.8</td>
<td>61.6%</td>
</tr>
<tr>
<td>Blister fluid</td>
<td>8</td>
<td>10 mg/kg</td>
<td>3.5–4.5</td>
<td>1.36</td>
<td>1.44</td>
<td>85%</td>
</tr>
<tr>
<td>Blister fluid</td>
<td>6</td>
<td>1</td>
<td>3.5–4.5</td>
<td>4.94</td>
<td>2.07</td>
<td>&gt;38%</td>
</tr>
</tbody>
</table>
involving *P. aeruginosa* is less clear. Other published Monte Carlo evaluations of meropenem pharmacodynamics indicate that even regimens of 1 g every 8 hours often fail to produce desired (≥90% probability), and sometimes not even acceptable (≥80% probability) likelihood of attaining the desired pharmacodynamic target of ≥40% T>MIC for *P. aeruginosa* (Kuti et al 2004; Masterton et al 2005). While the probabilities of achieving specified pharmacodynamic targets for *P. aeruginosa* isolates from different geographic regions ranged from 63% to 91% with meropenem 1 g every 8 hours, probabilities of target attainment with the regimen of 500 mg every 8 hours ranged from only 59% to 81% (Kuti et al 2004; Masterton et al 2005). These analyses were performed utilizing MIC data from the MYSTIC database, but *P. aeruginosa* susceptibilities of isolates obtained from different geographic regions, specific institutions, and even individual patients may be very different from those in MYSTIC and would therefore yield quite different probabilities of pharmacodynamic target attainment (either higher or lower). Although a previous analysis of meropenem for cSSTIs indicated a high probability of achieving desired pharmacodynamic goals with the regimen of 500 mg every 8 hours, this analysis was based on pooled susceptibility data in which MIC data were weighted according to the prevalence of causative pathogens encountered in cSSTIs and did not report probability of target attainment specifically for *P. aeruginosa* (Ong et al 2005). Based on pharmacodynamic considerations, it appears that meropenem 500 mg every 8 hours is not necessarily optimal for the treatment of cSSTIs or other infections when *P. aeruginosa* is suspected or documented. Meropenem 1 g every 8 hours may be a more preferred regimen in these cases.

The elimination half-life of meropenem is approximately one hour in patients with normal renal function. The major route of meropenem elimination is urinary excretion of unchanged drug, with renal clearance (CL\textsubscript{R}) accounting for 58%–83% of total systemic clearance of the drug and up to 98% of each dose eventually excreted in the urine as unchanged meropenem and as a pharmacologically inactive, open β-lactam metabolite (Wiseman et al 1995; Fish and Singletary 1997). It appears that meropenem pharmacokinetics in children ≤6 months of age are very similar to those seen in adults.

Because many cSSTIs occur in the setting of diabetes mellitus and other acute or chronic illnesses associated with renal dysfunction, the disposition of meropenem in patients with renal impairment is of relevance. Following intravenous administration, meropenem AUC, CL\textsubscript{R}, and total systemic clearance are altered in proportion to the degree of renal impairment present. Dosage adjustments are therefore required in patients with moderate to severe renal insufficiency and calculated creatinine clearances of <50 mL/minute in order to prevent excessive drug accumulation. Dosage adjustments of meropenem in elderly patients are also based on renal function with no other apparent age-related pharmacokinetic alterations. The pharmacokinetics of meropenem are not significantly altered in patients with hepatic impairment and no dosage adjustments are required (Wiseman et al 1995; Fish and Singletary 1997).

**Clinical data**

At least three studies have evaluated meropenem in the treatment of cSSTIs (Table 3) (Lami et al 1991; Nichols et al 1995; Fabian et al 2005). The first of these was a small prospective, open-label study comparing meropenem with imipenem/cilastatin in patients with soft tissue infections requiring hospitalization, surgical drainage, and/or debridement in addition to antibiotic therapy (Lami et al 1991). Pertinent among the exclusion criteria were patients with neutropenia or diabetic foot infections requiring amputation; clinically evaluable patients were required to have documented bacterial pathogens. Patients were randomized to receive either meropenem 500 mg every 8 hours (23 evaluable patients) or imipenem/cilastatin 500 mg every 6 hours (21 patients); patients in both groups were treated for a mean duration of approximately 7 days. Patients in the two treatment groups were similar with regards to demographics, severity and type of infections. *S. aureus*, streptococci, and *E. coli* were the most common pathogens with a mean of three bacterial isolates per patient. Satisfactory clinical responses were demonstrated in 100% of patients in both treatment groups; microbiological response rates and the number of infections caused specifically by *P. aeruginosa* were not reported. Both antibiotic regimens were well tolerated with an apparently low reported incidence of adverse events.

A second study also evaluated the efficacy of meropenem compared with imipenem/cilastatin in the treatment of severe infections (Nichols et al 1995). This was a prospective, randomized, multicenter, open-label study. The primary inclusion criteria were requirement for hospitalization and treatment with parenteral antibiotics; the specific types of infections eligible for inclusion in the study were not well defined. Relevant exclusion criteria included neutropenia,
Fish rapidly progressive disease, and severe peripheral vascular
disease likely to require amputation. Patients received either
meropenem 500 mg every 8 hours (123 evaluable patients)
or imipenem/cilastatin 500 mg every 6 hours (126 patients);
patients in both groups were treated for a mean duration of
approximately 6 days. Primary study endpoints were clinical
and bacteriologic responses at the end of therapy and at a
follow-up visit 2 to 4 weeks after the end of therapy.
Treatment groups were similar with respect to patient
demographics, diagnoses, or required surgical procedures
during treatment. Greater than 97% of infections were
community-acquired and included patients with diabetes,
peripheral vascular disease, and injection drug use-related
infections. Approximately half of all infections were
monomicrobial and involved a Gram-positive organism in
86% of these patients. Polymicrobial infections were
documented in 48% of meropenem-treated patients and 45%
of patients receiving imipenem/cilastatin. The overall
distribution of pathogens in these patients was 58% aerobic
Gram-positive bacteria, 22% aerobic Gram-negatives, and
20% anaerobes. Satisfactory clinical responses at end of
therapy were similar between the groups treated with
meropenem and imipenem/cilastatin (98% versus 95%,
respectively; 95% confidence interval [CI], −2.29 to 6.93).
Microbiological response rates at end of treatment in the
meropenem and imipenem/cilastatin groups were also
similar (94% and 91%, respectively; 95% CI, −2.73 to
10.39); however, meropenem was statistically better than
imipenem/cilastatin in overall eradication of pathogens (96% vs
90%, respectively; p=0.007) and eradication of gram-
positive aerobes (96% vs 88%, respectively; p=0.01).
Although P. aeruginosa was reported to be eradicated in 9
of 11 patients (82%), this rate was somewhat lower than
that reported for other Gram-negative pathogens (48 of 50,
96%). Clinical efficacy rates in meropenem and imipenem/
cilastatin groups at the 2- to 4-week follow-up visit were
92% and 89%, respectively (p=0.60), while microbiological
efficacy at follow-up was 92% versus 82%, respectively
(p=0.11). Superinfection due to S. aureus was
documented in one meropenem-treated patient; the
incidence of other drug-related adverse events was similar
between groups.

Although these first two studies provide support for
meropenem’s efficacy in the treatment of cSSTIs, they had
potential limitations in assessing the role of meropenem
specifically for complicated infections. First, it is not clear
whether these studies actually evaluated the treatment of
complicated infections as these are currently defined. Both
studies included patients with infections requiring
hospitalization and surgical intervention, so these infections
do appear to have been more severe or complicated in nature,
or both. However, the sites and types of infections eligible
for inclusion in the studies were not always well defined.
The 100% efficacy rates reported in the first study are also
somewhat suspect for complicated infections (Lami et al
1991); this may reflect the unblinded nature of the study
rather than the possibility that the infections were not truly
complicated in nature. In the second study (Nichols et al
1995), the infections were overwhelmingly community-
acquired in origin. Although these were appropriate for
evaluation, the study did not represent the full spectrum of
infections (eg, surgical wound infections, other nosocomial
infections) of importance in cSSTIs. Thus although the
overall results of these two studies were favorable for
meropenem, they both had relative limitations regarding
study design and/or enrolled patient populations which make

<table>
<thead>
<tr>
<th>Reference</th>
<th>Study type</th>
<th>Evaluable patients (N)</th>
<th>Regimen</th>
<th>Mean duration of therapy (days)</th>
<th>Response</th>
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</thead>
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<tr>
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<td>M 500 mg q8h</td>
<td>6.4</td>
<td>100</td>
</tr>
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<td></td>
<td></td>
<td>21</td>
<td>I/C 500 mg q6h</td>
<td>6.5</td>
<td>NR</td>
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<tr>
<td>Nichols et al 1995</td>
<td>P, MC, NB</td>
<td>123</td>
<td>M 500 mg q8h</td>
<td>7.1</td>
<td>98</td>
</tr>
<tr>
<td></td>
<td></td>
<td>126</td>
<td>I/C 500 mg q6h</td>
<td>7.3</td>
<td>95</td>
</tr>
<tr>
<td>Fabian et al 2005</td>
<td>P, R, MC, DB</td>
<td>261</td>
<td>M 500 mg q8h</td>
<td>5.8 *</td>
<td>86.2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>287</td>
<td>I/C 500 mg q8h</td>
<td>6.0 *</td>
<td>82.9</td>
</tr>
</tbody>
</table>

**Note:** Numbers in table represent duration of parenteral therapy only. Approximately 50% of patients in each treatment group were switched to oral therapy and treated for an additional mean of 9.3 and 9.0 days in meropenem and I/C groups, respectively.

**Abbreviations:** AE, adverse events; DB, double-blind; I/C, imipenem/cilastatin; M, meropenem; MC, multicentered; NB, non-blinded; NR, not reported; P, prospective; R, randomized.
the significance and applicability of the results difficult to fully evaluate.

The best and most recent study of meropenem in the treatment of cSSTIs overcame the deficiencies of the two earlier, previously-described studies. This large prospective, randomized, multicenter, international, double-blind study compared the efficacy and safety of meropenem 500 mg every 8 hours versus imipenem/cilastatin 500 mg every 8 hours in hospitalized patients with cSSTIs (Fabian et al 2005). Infections classified as complicated for purposes of this study included the following: complicated cellulitis (ie, in the setting of diabetes, involving the perineum, or deep tissue involvement evidenced by necrosis, tissue fluctuance, bacteremia, or need for surgical incision); surgical site infections; traumatic infections; infected diabetic and ischemic ulcers; complex or perirectal abscesses, or other bacterial infections requiring hospitalization, surgical intervention, and parenteral antibiotics. In addition, all patients were required to have had specimens for culture and susceptibility testing obtained before study enrollment; any pathogens known to be present at study enrollment were required to be susceptible to study drugs. Pertinent exclusion criteria included neutropenia, underlying osteomyelitis, severe peripheral vascular disease likely to require amputation, necrotizing fasciitis, pressure ulcerations, or infected prosthetic materials. Concomitant therapy with vancomycin or other systemic antibiotics was not permitted; patients could be switched to oral antibiotics of the investigators’ choice after a minimum of three days of parenteral therapy. The primary study endpoints were clinical efficacy (cure versus failure or relapse) in clinically evaluable (CE) and modified intent-to-treat (MITT, patients meeting all eligibility criteria and receiving ≥1 dose of drug) populations at a follow-up test-of-cure (TOC) visit 7 to 28 days after completion of all antibiotics. Secondary endpoints included clinical efficacy in the intent-to-treat (ITT, all patients receiving ≥1 dose of drug whether or not they met eligibility criteria) population at the TOC follow-up visit, as well as clinical efficacy in the ITT, MITT, and CE populations at the end-of-therapy visit occurring within 24 hours of completion of antibiotic therapy.

A total of 1076 patients were enrolled into the study. Of these, final study populations treated with meropenem and imipenem/cilastatin included 510 and 527 patients, respectively, in the ITT groups; 334 and 358 patients, respectively, in the MITT groups; and 261 and 287 patients, respectively, in the CE groups. Treatment groups were similar in respect to patient demographics, types of infection, and underlying medical conditions. Approximately 93% of patients in each group had infections classified as moderate or severe in nature, and 80% in each treatment arm had deep tissue involvement. Additionally, 68% of patients treated with meropenem and 70% of those receiving imipenem/cilastatin required surgical intervention. In contrast to the two earlier studies, approximately 18% of patients in both groups had surgical site or traumatic wound infections. Isolated pathogens were also evenly represented among the two treatment groups. Aerobic Gram-positive organisms accounted for 57% of pre-treatment pathogens, while aerobic Gram-negative pathogens were found in 26% of patients and anaerobes in 17% (Table 4). Polymicrobial infections were microbiologically documented in 38% of patients at pre-treatment baseline.

The mean duration of therapy with intravenous therapy was approximately 6 days in both groups, similar numbers of patients in each group were switched to oral antibiotics (approximately 50%), and the mean duration of oral antibiotics was also similar between groups (approximately 9 days). Evaluation of the primary study endpoints demonstrated equivalent efficacy between meropenem and imipenem/cilastatin within both the CE (86.2% vs 82.9%, respectively; 95% CI, −2.8 to 9.3) and MITT populations (73.1% vs 74.9%, respectively; 95% CI, −8.4 to 4.7). There were also no statistically significant differences between meropenem and imipenem/cilastatin among the secondary endpoints. Clinical efficacy rates in the meropenem and imipenem/cilastatin groups within the ITT population at the

<table>
<thead>
<tr>
<th>Table 4 Pre-treatment pathogens isolated from patients with complicated skin and soft tissue infections (Fabian et al 2005)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Total pathogens</strong></td>
</tr>
<tr>
<td><strong>Gram-positive aerobes</strong></td>
</tr>
<tr>
<td>Staphylococcus aureus (MS)</td>
</tr>
<tr>
<td>Staphylococcus aureus (MR)</td>
</tr>
<tr>
<td>Streptococcus pyogenes</td>
</tr>
<tr>
<td>Streptococcus agalactiae</td>
</tr>
<tr>
<td>Enterococcus faecalis</td>
</tr>
<tr>
<td><strong>Gram-negative aerobes</strong></td>
</tr>
<tr>
<td>Escherichia coli</td>
</tr>
<tr>
<td>Pseudomonas aeruginosa</td>
</tr>
<tr>
<td>Proteus spp.</td>
</tr>
<tr>
<td>Klebsiella spp.</td>
</tr>
<tr>
<td><strong>Anaerobes</strong></td>
</tr>
<tr>
<td>Bacteroides fragilis</td>
</tr>
<tr>
<td>Peptostreptococcus spp.</td>
</tr>
<tr>
<td>Prevotella spp.</td>
</tr>
<tr>
<td><strong>Monomicrobial infections</strong></td>
</tr>
<tr>
<td>Polymicrobial infections</td>
</tr>
</tbody>
</table>

Abbreviations: MS, methicillin-susceptible; MR, methicillin-resistant.
TOC visit (approximately 57% vs 61%, respectively; 95% CI, −9.8 to 2.1), and within the CE (93.5% vs 92.3%, respectively; approximate 95% CI, −3.1 to 5.4), MITT (91.0% vs 91.1%, respectively; approximate 95% CI, −4.3 to 4.2), and ITT (81.0% vs 83.5%, respectively; approximate 95% CI, −7.1 to 2.0) populations at the end-of-treatment evaluations were all similar among the two treatment groups. Again, no significant differences in efficacy among the meropenem and imipenem/cilastatin groups were observed when patients were divided into subgroups according to demographic parameters, types of infection, or infecting pathogens. Clinical cure rates specifically in patients infected with P. aeruginosa and treated with meropenem and imipenem/cilastatin were 73.3% (11/15) and 86.7% (13/15), respectively (p=0.652). The incidence of drug-related adverse events was similar between meropenem and imipenem/cilastatin groups (9.0% vs 10.8%, respectively).

Additional data from this study regarding the bacteriology and clinical efficacy of meropenem and imipenem/cilastatin specifically in diabetic patients were recently presented at an international meeting (Embil et al 2005). A total of 398 patients (38%) enrolled into the study had diabetes mellitus. Of these patients, 254 (82%) had infections involving the leg or foot, or both, most commonly complicated cellulitis, complex abscess, or infected ulcers. The distribution of pathogens in these patients was 58% aerobic Gram-positive bacteria, 27% aerobic Gram-negatives, and 15% anaerobes. This distribution of pathogens was similar to that seen in non-diabetic patients, although the frequency of polymicrobial infection was greater in diabetic versus non-diabetic patients (44% versus 34%, respectively; p=0.002). Clinical efficacy rates among all enrolled patients at follow-up were significantly higher with meropenem vs. imipenem/cilastatin in diabetic patients (86% vs 72%, respectively; p<0.0001) but not in non-diabetic patients (87% vs 89%, respectively; p=0.322). Clinical efficacy rates among microbiologically evaluable diabetic patients were also significantly higher for infections caused by various bacteria among meropenem-treated patients compared with those receiving imipenem/cilastatin (Gram-positive aerobes, 87% vs 70%; Gram-negative aerobes, 74% vs 60%; anaerobes, 94% vs 71%; and polymicrobial infections, 88% vs 65%, respectively; p<0.0001 for each of the four comparisons). Possible reasons for these reported differences in efficacy are not clear. Although previous studies used imipenem/cilastatin doses of 500 mg every 6 hours (Lami et al 1991; Nichols et al 1995), the lower dose of 500 mg every 8 hours used in this third study has been shown to be apparently adequate for the treatment of SSTIs based on pharmacodynamic modeling (Ong et al 2005). In addition, since this abstract reported data from an unplanned subgroup analysis of the original study, the post-hoc statistical comparisons do not necessarily indicate that meropenem is superior to imipenem/cilastatin in the treatment of SSTIs in diabetic patients. However, the data do indicate that meropenem is at least comparable with imipenem/cilastatin and is quite an effective agent in the treatment of such infections.

Based on available data as reviewed in this article, meropenem was approved by the FDA on May 25, 2005 for the treatment of SSTIs at a dose of 500 mg intravenously every 8 hours. Meropenem is specifically indicated for SSTIs caused by the following pathogens: S. aureus (methicillin susceptible only), S. pyogenes, S. agalactiae, viridans group streptococci, E. faecalis (vancomycin susceptible only), E. coli, P. mirabilis, P. aeruginosa, B. fragilis, or Peptostreptococcus spp.

**Role of meropenem in SSTIs**

Complicated skin and soft tissue infections continue to represent a considerable clinical challenge. The underlying conditions such as diabetes mellitus, peripheral vascular disease, surgery, and immunosuppression that predispose to SSTIs also make them potentially less responsive to traditional antibiotic therapy, thus necessitating hospital admission for parenteral antibiotics and surgical interventions. The broad range of bacterial pathogens likely to be encountered in SSTIs, together with the changing epidemiology of susceptibilities to currently available antibiotics, adds to the complexity of managing these infections. As previously discussed, meropenem has excellent activity against most bacteria commonly associated with SSTIs, often including P. aeruginosa and ESBL-producing organisms which are resistant to other β-lactam antibiotics including third-generation cephalosporins. The status of carbapenems as the drugs of choice in the treatment of infection due to ESBL-producing organisms makes meropenem and other agents of this class particularly attractive in geographic areas with high prevalence of ESBL producers. Meropenem’s broad antimicrobial spectrum of action is thus well suited to either empiric treatment of SSTIs when specific pathogens are unknown, as well as definitive treatment of mixed bacterial infections and/or those caused by multidrug-resistant organisms once specific pathogens and susceptibilities have been documented. However, because of the recent emergence of community-
acquired MRSA as an important pathogen, the addition of vancomycin, linezolid, or some other agent with good activity against MRSA should be considered for community-acquired infections in which *S. aureus* is a suspected pathogen.

The use of broad-spectrum antibiotics for the initial treatment of cSSTIs has been addressed in recent practice guidelines from the Infectious Diseases Society of America (Lipsky et al 2004; Stevens et al 2005). Broad-spectrum agents which provide good activity against Gram-positive aerobes, Gram-negative aerobes, and anaerobes have been recommended as appropriate initial choices for a variety of cSSTIs including complicated cellulitis, severe diabetic foot infections, surgical wound infections, infections in immunocompromised hosts, severe animal or human bite wounds, and necrotizing infections (Lipsky et al 2004; Stevens et al 2005). There are currently a number of agents that could potentially be appropriate for use in these types of infections. β-Lactam/β-lactamase inhibitor combinations such as piperacillin/tazobactam have been extensively used for these indications due to their favorable spectrum of activity and proven efficacy in a wide variety of infection types (Tan et al 1993; Tassler et al 1993; Lipsky et al 2004, 2005). The new glycyrlcycline-class drug, tigecycline, has also been shown to have good efficacy in the treatment of cSSTIs and was approved in the US for this indication in June 2005 (Breedt et al 2005; Wilcox 2005). Carbapenems such as imipenem/cilastatin and, more recently, ertapenem have also been shown to be very effective in the initial treatment of severe cSSTIs (Grayson et al 1994; Graham et al 2002; Lipsky et al 2005). Although the carbapenems are too broad in spectrum for routine first-line use in all types of infections, they are particularly well suited and recommended as appropriate agents for infections likely to involve mixed and/or multidrug-resistant Gram-negative pathogens such as surgical wound and other hospital-acquired infections, severe diabetic foot infections, and infections in immunocompromised patients (Fung et al 2003; Dinubile and Lipsky 2004; Lipsky et al 2004; Stevens et al 2005).

Meropenem has shown excellent efficacy in controlled clinical studies in a variety of types of moderate to severe infections in seriously ill patients (Wiseman et al 1995; Fish and Singletary 1997). Clinical studies directly comparing meropenem versus imipenem/cilastatin in the treatment of intra-abdominal, lower respiratory tract, SSTIs, urinary tract infections, and exacerbations of chronic obstructive pulmonary disease (COPD) generally found these two agents to be comparable in clinical and microbiological efficacy (Colardyn and Faulkner 1996; Garau et al 1997). As discussed in this article, at least three clinical studies have also shown meropenem to be comparable with imipenem/cilastatin in the treatment of cSSTIs, one of these being among the largest and best-designed studies of cSSTIs ever performed (Fabian et al 2005). While meropenem and the other carbapenems are not generally recommended for initial treatment of all cSSTIs because their extremely broad spectrum of activity is not necessarily required and excessive use of these agents is undesirable, their use is appropriate and recommended for initial treatment in certain selected patients as previously discussed. Meropenem is an appropriate antibiotic choice in those patients in whom use of a carbapenem is recommended for infections involving pathogens known to be resistant to other narrower-spectrum, more preferred agents.

Although meropenem is indicated by the US FDA for treatment of cSSTIs caused by *P. aeruginosa*, the approved regimen of 500 mg every 8 hours may not necessarily be the preferred regimen for empiric treatment of cSSTIs in higher-risk patients in which *P. aeruginosa* is more likely to occur, ie, hospital-acquired infections, severe diabetic foot infections, and infections in immunocompromised patients. As previously discussed, pharmacokinetic/pharmacodynamic analyses indicate that this regimen may not be as favorable as higher doses, ie, 1 g every 8 hours, in reliably achieving desired pharmacodynamic targets for treatment of *P. aeruginosa* infections (Kuti et al 2004; Masterton et al 2005). Additionally, although clinical studies of meropenem in the treatment of cSSTIs were overall very favorable, relatively few *P. aeruginosa* were isolated in these studies and the overall response rate in these patients (20/26, 77%) was somewhat lower than the overall clinical and microbiological efficacy rates reported in these three studies (86% to 100% and 88% to 94%, respectively) (Lami et al 1991; Nichols et al 1995; Fabian et al 2005). These findings, together with considerations regarding potential alterations in drug pharmacokinetics and tissue penetration, increasing *P. aeruginosa* resistance, and potential differences in local and institutional susceptibility patterns, strongly suggest that the use of higher daily doses of meropenem may be more prudent when *P. aeruginosa* is suspected or until specific susceptibilities can be determined in documented strains.

In addition to meropenem’s favorable spectrum of activity and proven clinical efficacy, meropenem is also well tolerated and has few serious toxicities associated with its
Fish et al. 1995; Fish and Singletary 1997; Norrby and Gildon 1999). Although carbapenems have been associated with seizures and other central nervous system (CNS) toxicities, the rate of meropenem-associated CNS toxicity appears to be less than that of imipenem/cilastatin and probably similar to ertapenem (Wiseman et al. 1995; Fish and Singletary 1997; Norrby and Gildon 1999; Teppler et al. 2004). The incidence of seizures during meropenem therapy is very low and appears to be 0.05% to 0.08% (Fish and Singletary 1997; Norrby and Gildon 1999). This potential difference in CNS toxicities between meropenem and imipenem/cilastatin may be most relevant in higher-risk patients such as those with head trauma, seizure disorders, or other underlying CNS pathology. The overall safety and tolerability of meropenem appears to be similar to other carbapenems (Fish and Singletary 1997; Norrby and Gildon 1999).

The average wholesale price (AWP) of meropenem and other very broad-spectrum agents which are currently used as monotherapy for the treatment of cSSTIs are shown in Table 5 (Red Book 2005). Prices shown are based on a 7-day course of therapy at FDA-approved doses of the respective drugs in patients with good renal function. Although meropenem is most easily compared against other carbapenems, piperacillin/tazobactam and tigecycline are also appropriate for treatment of certain cSSTIs and could potentially be used as alternatives to carbapenems in many patients (Fung et al. 2003; Lipsky et al. 2004; Breedt et al. 2005; Stevens et al. 2005; Wilcox 2005). Although ertapenem and tigecycline have poor activity against P. aeruginosa and are not strictly interchangeable with meropenem for all infections, ertapenem does offer potential cost advantages in the treatment of those infections for which it is appropriate (eg, when P. aeruginosa is not a likely pathogen). Compared with imipenem/cilastatin, a drug which is quite similar to meropenem in terms of activity and clinical efficacy, meropenem may in fact be more cost-effective. However, actual costs at individual institutions will vary considerably according to specific contract pricing agreements and costs within countries outside of the US. The selection of one agent over another may very well depend on prices available to the specific institution. Local antibiograms are also important to consider in the selection of a specific agent.

Meropenem possesses many favorable qualities and some possible advantages relative to other agents which are commonly used in the treatment of cSSTIs. These advantages may include good activity against pathogens (particularly Gram-negative aerobes) with increasing resistance to cephalosporins and extended-spectrum penicillins; the ability to effectively treat many cSSTIs with a single-drug regimen rather than a combination of agents in order to provide adequate coverage of suspected or documented pathogens; incomplete cross-resistance with imipenem/cilastatin and slightly greater activity against P. aeruginosa; potentially decreased CNS toxicity compared with imipenem/cilastatin; and potential cost advantages. Conversely, potential disadvantages of meropenem use include an excessively broad spectrum of activity for treatment of many cSSTIs in which P. aeruginosa, anaerobes, and/or multidrug-resistant pathogens are not likely to be present; a broad spectrum of activity which may increase selective pressure for development of resistance among organisms such as P. aeruginosa; the need for addition of another agent when MRSA is possibly present in community-acquired infections; the need for three-times-daily dosing; and potentially higher costs relative to some alternative agents. Based on available data, meropenem should be considered relatively interchangeable with imipenem/cilastatin for treatment of cSSTIs and would be preferred over ertapenem for certain patients who are at high risk of infection with P. aeruginosa. Meropenem is an

<table>
<thead>
<tr>
<th>Drug</th>
<th>US trade name and manufacturer</th>
<th>Dosing regimen</th>
<th>Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Meropenem</td>
<td>Merrem®, AstraZeneca, Wilmington, Delaware, USA</td>
<td>500 mg every 8 hours</td>
<td>$630</td>
</tr>
<tr>
<td>Imipenem/cilastatin</td>
<td>Primaxin®, Merck &amp; Co., Whitehouse Station, New Jersey, USA</td>
<td>500 mg every 8 hours</td>
<td>$722</td>
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<tr>
<td>Ertapenem</td>
<td>Invanz®, Merck &amp; Co., Whitehouse Station, New Jersey, USA</td>
<td>1 g every 24 hours</td>
<td>$360</td>
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<tr>
<td>Other agents</td>
<td></td>
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<tr>
<td>Piperacillin/tazobactam</td>
<td>Zosyn®, Wyeth Pharmaceuticals, Philadelphia, Pennsylvania, USA</td>
<td>3.375 g every 6 hours</td>
<td>$482</td>
</tr>
<tr>
<td>Tigecycline</td>
<td>Tygaci®, Wyeth Pharmaceuticals, Philadelphia, Pennsylvania, USA</td>
<td>50 mg every 12 hours</td>
<td>$735</td>
</tr>
</tbody>
</table>

Note: Figures are based on 2005 Average Wholesale Price (AWP) in the US and reflect drug acquisition costs only (Red Book 2005); a After an initial loading dose of 100 mg.
effective and safe agent for treatment of cSSTIs for which carbapenems are deemed appropriate.

**Summary**

Meropenem is a broad-spectrum carbapenem antibiotic with excellent activity against many aerobic Gram-positive, aerobic Gram-negative, and anaerobic organisms commonly associated with cSSTIs. At least three clinical studies, including one of the largest prospective, randomized, double-blind studies of cSSTIs yet conducted, have shown that meropenem is comparable in clinical efficacy with imipenem/cilastatin and is well tolerated in the treatment of these infections. Carbapenems are currently recommended as appropriate agents in the initial treatment of certain types of cSSTIs including those likely to involve mixed and/or multidrug-resistant pathogens, ie, surgical wound and other hospital-acquired infections, severe diabetic foot infections, and infections in immunocompromised patients. When used for appropriate types of cSSTIs in carefully selected patients, and with the addition of a second agent for coverage of MRSA when appropriate, meropenem is an effective and viable alternative for monotherapy of these challenging infections. The regimen of meropenem 500 mg every 8 hours is appropriate for the treatment of infections caused by most pathogens. However, this author recommends the use of higher doses (ie, 1 g every 8 hours) for empiric treatment of patients at higher risk of infection with *P. aeruginosa* or when this specific pathogen is actually documented to be present.

**References**


Merrem (meropenem) product package insert. 2005 (October). AstraZeneca Pharmaceuticals, Wilmington, Delaware, USA.


