Tigecycline: an evidence-based review of its antibacterial activity and effectiveness in complicated skin and soft tissue and intraabdominal infections

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Abstract

Introduction: There is an urgent need for novel agents to manage serious bacterial infections, particularly those contracted in healthcare facilities. Tigecycline is a novel broad-spectrum glycylcycline with good activity against Gram-positive, many Gram-negative, anaerobic, and some atypical pathogens that has been developed to address this need.

Aims: To review the evidence for the use of tigecycline in serious and complicated skin and soft tissue and intraabdominal infections.

Evidence review: There is substantial evidence that tigecycline is as effective as vancomycin plus aztreonam in complicated skin and skin structure infections (SSSIs) and as effective as imipenem plus cilastatin in intraabdominal infections. Limited evidence shows effectiveness in patients with resistant Acinetobacter infection in an intensive care unit, and the possibility that the use of tigecycline may reduce length of hospital stay. The drug is well tolerated, with nausea and vomiting as the major adverse effects.

Outcomes summary: The introduction of tigecycline should be beneficial at a time of increasing problems with bacterial resistance, and evidence to date has been sufficient for regulatory approval for complicated SSSIs and intraabdominal infections. Research into tigecycline's efficacy in other infectious diseases (notably pneumonia and bacteremia) is ongoing. Further good quality studies and ongoing surveillance for any emerging bacterial resistance will be needed to determine outcomes with tigecycline relative to other novel antibacterial agents, and to explore the economic implications of its adoption.

Key words: antibiotic resistance, bacterial infections, glycylcycline, nosocomial infections, review, tigecycline

Core evidence outcomes summary for tigecycline in complicated skin and soft tissue and intraabdominal infections

| Outcome measure | Evidence | Implications |
|-----------------|----------|--------------|
| Patient-oriented evidence |          |              |
| Good patient acceptability | Substantial | Good chance of success in severe infections without poor tolerability characteristic of some older antibacterials |
| Avoidance of morbidity or mortality due to empiric treatment failure in the most seriously ill patients | Limited | Patients needing intensive care and/or infected with multiresistant pathogenic strains resistant to other available antibacterials may respond to tigecycline. Further studies required |
| Disease-oriented evidence |          |              |
| Clinical cure in complicated skin and soft tissue and intraabdominal infections | Substantial | Effective in these infections, but superiority over other antibacterials in terms of response rates has not yet been demonstrated |
| Microbiologic eradication in patients infected with a range of Gram-positive and Gram-negative organisms | Substantial | Is likely to be widely effective in a range of patients with serious and difficult infections resistant to other agents |
| Economic evidence |          |              |
| More rapid discharge from hospital | Limited | Has considerable implications for both patient wellbeing and health economics; level 2 evidence required |
| Acquisition cost offset or incremental cost effectiveness/benefit | No evidence | Dependent on maintenance of clinical effectiveness and avoidance of empiric treatment failure. Studies are required |
Scope, aims, and objectives

This evidence-based review of the activity and clinical effectiveness of tigecycline, the first of the novel glycylcycline class of antibiotics, focuses on the use of the drug in complicated and serious skin and soft tissue and intraabdominal infections known to pose resistance problems with current therapy and associated with or requiring treatment in healthcare facilities.

The emergence of microorganisms resistant to current antibacterial agents is a matter of concern, particularly in patients with serious, complicated, and nosocomial infections. Virtually all common infectious bacteria have developed resistance to at least one class of antibiotics, and tigecycline has been developed as part of the effort to identify novel antibacterial agents in the face of the growing problem, particularly in hospitals, of potentially life-threatening multiresistant organisms and polymicrobial infections. A number of agents are available for the treatment of conditions of chief concern, including skin and intraabdominal infections (particularly those contracted in hospitals) and pneumonia, but available antibacterials have been widely affected by resistance, and some older agents used for serious infections (e.g. vancomycin) may be poorly tolerated by some patients. New drugs that have recently been introduced into clinical practice have been aimed mainly at multiresistant Gram-positive infections, but early resistance to, for example, the novel oxazolidinone linezolid (Mutnick et al. 2003; Halle et al. 2004; Meka et al. 2004) and the cyclic lipopeptide daptomycin (Mangili et al. 2005; Sabol et al. 2005) has already been reported. New agents that target broad ranges of both Gram-positive and Gram-negative organisms, particularly the emerging extended spectrum beta-lactamase (ESBL) producers, anaerobes, and methicillin- and/or oxacillin-resistant Gram-positive isolates, are urgently needed.

Tigecycline is one of a number of drugs under development with these needs in mind, and is designed to circumvent two of the most common resistance mechanisms, efflux and ribosomal protection. It is a semisynthetic derivative of minocycline that is mechanistically similar to the aminoglycosides, macrolides, streptogramins, and oxazolidinones in that it binds to the 30S ribosomal subunit (Chopra 2001; Guay 2004; Zhanel et al. 2004). This blocks access of aminoacyl tRNA to its acceptor site and leads to the prevention of bacterial protein synthesis and growth. Examination and review of tigecycline is timely because, at the time of writing, two phase III trials have been completed with several others underway, and Food and Drug Administration (FDA) approval has been granted in the USA (Anon. 2005).

Methods

The English language medical literature was searched for appropriate articles relating to tigecycline. The following databases were searched on June 16, 2005, and again on November 11, 2005, with the search terms “tigecycline OR GAR-936,” and no date limits:

- PubMed, http://www.ncbi.nlm.nih.gov/entrez
- EMBASE and BIOSIS, http://www.datastarweb.com
- Database of Abstracts of Reviews of Effects (DARE), National Health Service Economic Evaluation Database (NHSEED), Health Technology Assessment (HTA), http://www.york.ac.uk/inst/crd/crddatabases.htm
- Cochrane Database of Systematic Reviews (CDSR), http://www.cochrane.org
- Clinical Evidence (BMJ), http://www.clinicalevidence.com
- National Institute for Health and Clinical Excellence (NICE), http://www.nice.org.uk
- National Guidelines Clearing House, http://www.guideline.gov

The following websites were also searched for information including details of recent meetings (since 2003) hosted by societies specializing in medical microbiology:

- The annual congress of the European Society of Clinical Microbiology and Infectious Diseases (ESCMID), http://www.escmid.org
- The annual Interscience Conference on Antimicrobial Agents and Chemotherapy (ICAAC), http://www.icaac.org
- The Federation of European Microbiological Societies (FEMS), http://www.fems-microbiology.org
- The Infectious Diseases Society of America (IDSA), http://www.idsociety.org

Table 1 summarizes the levels of evidence for articles identified from the search strategy. The original search (June 2005) yielded 89 full papers, of which one was included as level 3 evidence. Two of four level 2 studies were available in abstract form only at that time. Updated searching (November 2005) yielded subsequent full publications of these articles, together with one further paper and an additional full report not included in the databases searched but provided by the company developing tigecycline (Table 1).

There are currently no systematic reviews on tigecycline, and there is a good deal of information on the drug that relates to its antimicrobial activity in vitro and in animal models. These references, together with nonsystematic general review articles in which tigecycline featured, were excluded from the 132 full papers identified. This left one fully published phase II study (level 3 evidence) and four phase III trials (level 2 evidence). Five studies available as meetings abstracts were also identified (from websites of major professional bodies and meetings as above) at the time of writing. One of these provided level 3 (phase II) evidence, one level 5 (a small case series from a single center), and three were economic studies classified as level 3 evidence.

Disease overview

The development of tigecycline and other novel antibacterials currently under investigation is aimed chiefly at the problems...
caused by resistant or complicated infections, with particular emphasis on those contracted in healthcare facilities. It is worth noting that, in the USA alone, nosocomial infections affect around 2 million persons admitted to hospitals for acute care each year (Anon. 2000). Many of these infections are caused by multiresistant Gram-positive and Gram-negative pathogens, and their incidence continues to increase because of failure of hospital hygiene procedures, selective pressures resulting from inappropriate use and overuse of antibiotics, and other factors that can lead to the encoding of bacterial resistance mechanisms. The costs associated with managing infections are clearly considerable. While it is difficult to estimate the global burden on healthcare systems of infective disease, some helpful indicators were provided in the 1990s by the UK Public Health Laboratory Service (Plowman 2000), which examined the socioeconomic consequences of hospital-acquired infection (HAI). Patients who presented with one or more HAIs during an inpatient stay incurred costs 2.9 times greater than those for uninfected patients. This represented an absolute increase of £3154 per case in the 1449 patients selected for follow-up in this study. On average, the length of hospital stay was 2.9 times (14 days) longer for patients with an HAI, and additional burdens were imposed on community healthcare services after discharge. The overall cost of HAIs to the UK National Health Service was estimated as £986.36 million.

Diseases of particular concern include the frequently encountered skin and skin structure infections (SSSIs) and intraabdominal infections. SSSIs are caused chiefly by Gram-positive organisms, and are increasing in incidence mainly because of aging of the general population, increases in the numbers of critically ill and immunocompromised patients, and the emergence of multiresistant pathogens. SSSIs include simple and uncomplicated or superficial infections such as erysipelas, cellulitis, simple abscesses, furuncles, and wound infections, but also encompass complicated and more serious disorders such as necrotizing fasciitis, myositis, and gas gangrene. In general, an SSSI is considered complicated if it involves structures beneath the skin, such as fascia or muscle, requires significant surgical intervention, or accompanies disorders such as diabetes mellitus or human immunodeficiency virus (HIV) infection. Common pathogens are *Staphylococcus aureus*, *Streptococcus pyogenes*, *Str. agalactiae*, and group C and G streptococci [see the full review on this subject by Raghavan & Linden (2004)].

Data from England and Wales indicate further the increasing scale of the problem of multiresistant infections. Crowcroft and Catchpole (2002) examined death registrations from 1993 to 1998 and found an increase in the proportion of certificates mentioning methicillin-resistant *Staph. aureus* (MRSA) from 7.5% in 1993 to 25% in 1998. The authors stated that improved rates of reporting were unlikely to explain this increase, and found the greatest rise in MRSA rates to be associated with death in which invasive staphylococcal infection was given as the final underlying cause.

Group D streptococci include the enterococci *Enterococcus faecalis*, *Ent. durans*, and *Ent. faecium*; these microorganisms cause endocarditis, urinary tract infections, intraabdominal infections, cellulitis, wound infections, and bacteremia. *Enterococci* are intrinsically resistant to many antimicrobial agents including cephalosporins, penicillins, cotrimoxazole, and clindamycin (Kauffman 2003). Since 1989, however, rapid increases in the incidences of infection and colonization with vancomycin-resistant enterococci (VRE) have been reported (Anon. 1995). Increased risk of VRE infection is associated with previous vancomycin or multiple antimicrobial therapy, severe underlying disease, or abdominal surgery (Anon. 1995). *Enterococci* can also be spread by direct or indirect contact within institutions, and between hospitals by contaminated personnel (Kauffman 2003).

### Current therapy options

Although the treatment of complicated, resistant, and/or nosocomial infections varies in accordance with national recommendations and local infection control guidelines, generally applicable prescribing information can be found in recent reviews and official guidelines from the UK and USA. It is obviously not possible to present a comprehensive review of current antif infective prescribing practice here, but a representative picture can be drawn by focusing on the agents currently in use for treating SSSIs and intraabdominal infections.

Current UK recommendations for skin and gastrointestinal infections can be found in the British National Formulary, and are summarized in Table 2 (BNF 2005). It should be noted that the general principals of antibiotic prescribing apply in all cases: benzyl- or phenoxymethylpenicillin is recommended for streptococcal infection, with the addition of fluclaxacillin where staphylococcal infection is suspected, or a macrolide for atypical infection (e.g. *Mycoplasma* or *Chlamydia* spp.). Topical mupirocin may be effective against MRSA. Peritonitis and related infections are covered generally by a cephalosporin or gentamicin with metronidazole or clindamycin.
### Table 2 | UK prescribing recommendations for skin and gastrointestinal infections (BNF 2005)

| Condition                      | Recommended treatment                                      |
|--------------------------------|------------------------------------------------------------|
| **Skin infections**            |                                                            |
| Cellulitis                     | Benzylenicillin + flucloxacillin (or erythromycin alone if penicillin-allergic) |
| **Gastrointestinal system**    |                                                            |
| Campylobacter enteritis        | Ciprofloxacin or erythromycin                             |
| Invasive salmonellosis         | Ciprofloxacin or trimethoprim                             |
| Shigellosis                    | Ciprofloxacin or trimethoprim                             |
| Pseudomembranous colitis       | Metronidazole or vancomycin                               |
| Biliary tract infection        | Cephalosporin or gentamicin                               |
| Peritonitis                    | Cephalosporin (or gentamicin) + metronidazole (or clindamycin) |

Comprehensive diagnostic and prescribing guidelines have been published in the USA for SSSIs (Stevens et al. 2005) and intraabdominal infections (Solomkin et al. 2003) by IDSA. Broadly, a penicillinase-resistant semisynthetic penicillin or first generation cephalosporin is recommended for cellulitis, with clindamycin or vancomycin being suggested for patients allergic to penicillin. For severe group A streptococcal and clostridial necrotizing infections, parenteral clindamycin and penicillin therapy is recommended, with a variety of single agents indicated for aerobic Gram-positive and Gram-negative infections as well as for anaerobes (see Table 3). SSSIs caused by community-acquired MRSA may be susceptible to non-beta-lactam antibiotics such as doxycycline, clindamycin, trimethoprim/sulfamethoxazole, quinolones, or rifampicin. Severe infections that require hospitalization and that have not responded to other interventions can be treated with linezolid, daptomycin, or vancomycin. Trimethoprim/sulfamethoxazole has also been used to treat serious staphylococcal infections (Stevens et al. 2005).

Reviewers of recent developments in antiinfective therapy have recommended antistaphylococcal penicillins (e.g. nafcillin, oxacillin) or cefazolin for simple community-acquired SSSIs (Raghavan & Linden 2004). Nosocomial infections, on the other hand, may be treated with a semisynthetic penicillin such as piperacillin/tazobactam with or without vancomycin or teicoplanin. Vancomycin and teicoplanin are preferred for suspected MRSA. Oral switching to linezolid can take place after

### Table 3 | Examples of US prescribing recommendations for skin and skin structure and intraabdominal infections (Solomkin et al. 2003; Stevens et al. 2005)

| Condition                              | Recommended treatment                                      |
|----------------------------------------|------------------------------------------------------------|
| **Skin and skin structure infections** |                                                            |
| Cellulitis or erysipelas               | Dicloxacillin, cephalaxin, clindamycin, or erythromycin. Parenteral therapy in severely ill patients/unable to take oral medication: nafcillin, cefazolin, clindamycin, or vancomycin MRSA (hospital): linezolid, daptomycin or vancomycin. Also trimethoprim/sulfamethoxazole, or quinolone with good Gram-positive activity. |
| Necrotizing infections of skin, fascia, and muscle | Mixed infection: ampicillin/subactam or piperacillin/tazobactam plus clindamycin plus ciprofloxacin. Also imipenem/cilastatin, meropenem, ertapenem, cefotaxime plus metronidazole or clindamycin Streptococcal infection: penicillin plus clindamycin Staphylococcus aureus: nafcillin, oxacillin, cefazolin, vancomycin, or clindamycin Clostridial infection: clindamycin or penicillin |
| **Surgical site infections**           |                                                            |
| Intestinal/genital tract, single agents: cefoxitin, cefotizoxime, ampicillin/subactam, ticarcillin/clavulanate, piperacillin/tazobactam, imipenem/cilastatin, meropenem, or ertapenem | Intestinal/genital tract, combination: quinolone, third generation cephalosporin, aztreonam, aminoglycoside (facultative and aerobic activity). Clindamycin, metronidazole, chloramphenicol, penicillin + beta-lactamase inhibitor (anaerobic activity) Nonintestinal: oxacillin, first generation cephalosporin (trunk and extremities). Cefoxitin, ampicillin/subactam, other single agents as above for intestinal/genital operations (axillary or perineal) |
| **Intraabdominal infections**          |                                                            |
| Mild to moderate community-acquired infections | Ampicillin/subactam, cefazolin or cefuroxime/metronidazole, ticarcillin/clavulanate, or ertapenem |
| Healthcare-acquired infections         | Aminoglycoside (first choice for empiric use; completion with oral quinolone plus metronidazole or oral amoxicillin/clavulanate) |
isolates are resistant to methicillin (Wenzel 2004). Staph. aureus, Ent. faecalis MRSA, spp.
early discharge of hospitalized patients with MRSA. Nosocomial Gram-negative infections can be treated with ... penicillin-resistant Str. pneumoniae; VRE, vancomycin-resistant enterococci; VRSA, vancomycin-resistant Staph. aureus.

For patients with community-acquired intraabdominal infections, IDSA recommends the use of agents with narrower spectra of activity that are not commonly used for nosocomial infections. These include ampicillin/sulbactam, cefazolin or cefuroxime plus metronidazole, ticarcillin/clavulanate, ertapenem, and a quinolone plus metronidazole. For patients with more serious infections, alternatives include meropenem, imipenem/cilastatin, third or fourth generation cephalosporins plus metronidazole, ciprofloxacin plus metronidazole, or piperacillin/tazobactam (see also Table 3). The more resistant flora associated with nosocomial intraabdominal infections require treatment with more complex multidrug regimens (Solomkin et al. 2003). It should be noted here that these and other guidelines for antinfecrive therapy are subject to ongoing revision as novel agents are developed and infection and resistance patterns change.

A number of novel antibacterial agents have been introduced in recent years or are under current development. The major drugs of interest are discussed in more detail in the following section and are summarized in Table 4.

**Unmet needs**

Resistance to conventional antiinfective agents and the continuing emergence of multiresistant organisms in healthcare institutions are currently major concerns for researchers and clinicians. Indeed, the IDSA is sufficiently worried about these problems to have endorsed the introduction of legislation in the US Senate to encourage the development of new antiinfecrices, and is currently urging its members to lobby Congress (IDSA 2005a). Key issues identified by IDSA include the “drying up of the pipeline of new antibiotics,” and a loss of momentum in antibiotic research because of cost and regulatory factors, time constraints, and reduced profitability relative to other disease areas (IDSA 2005b).

Epidemiologic data underline the reasons for these concerns. High or intermediate resistance to penicillin is exhibited by 50% of pneumococcal strains in the USA, and similar numbers of Staph. aureus isolates are resistant to methicillin (Wenzel 2004). Of the enterococci, 30% are resistant to vancomycin, and 20% of strains of Pseudomonas aeruginosa are resistant to quinolones and 15% to imipenem. A recent large survey of 670 US hospitals has shown resistance rates for oxacillin-resistant Staph. aureus (ORSA), VRE, quinolone-resistant Escherichia coli, and ESBL-producing Klebsiella spp. to be 36, 10, 6, and 5%, respectively (Diekema et al. 2004). These rates were reported to be on the increase in the majority of hospitals.

Several research initiatives have been underway in recent years, however, to identify new antinfector agents. These efforts have resulted in the introduction of a number of new drugs, most of which are aimed at Gram-positive infections. These agents are represented most prominently by the oxazolidinone linezolid and the cyclic lipopeptide daptomycin.

Linezolid has a comprehensive spectrum of activity against the major nosocomial Gram-positive pathogens, including those with multiresistant phenotypes, and can be given either parenterally or orally (Kaufman 2003; Raghavan & Linden 2004). However, as mentioned above, early resistance to linezolid has already been reported (Mutnick et al. 2003; Halle et al. 2004; Meka et al. 2004), notably in staphylococcal and enterococcal isolates. Other oxazolidinones are in early clinical development (Gravestock 2005).

Daptomycin has a mode of action that is not completely understood but that involves cell membrane binding and potassium efflux. The drug is rapidly bactericidal against most Gram-positive pathogens, which may have importance in the treatment of endocarditis and meningitis. Daptomycin has also

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**Table 4 | Major new antimicrobial agents under development or recently introduced for nosocomial or resistant infections (adapted from Raghavan & Linden 2004)**

| Drug         | Class          | Susceptible organisms                                                                 | Resistance                  |
|--------------|----------------|---------------------------------------------------------------------------------------|-----------------------------|
| Linezolid    | Oxazolidinone  | MSSA, group A, B, C, F, G, and viridans streptococci, CoNS, MRSA, GRSA, GISA, VRE, VRSA, PRSP, corynebacteria, Listeria | Enterococcus faecium, Staphylococcus aureus |
| Daptomycin   | Cyclic peptolide | Ent. faecalis (vancomycin-susceptible), MSSA, MRSA, Streptococcus agalactiae, Str. pyogenes | Staph. aureus, Ent. faecalis |
| Quinupristin/dalfopristin | Streptogramin | MSSA, group A, B, C, F, G, and viridans streptococci, CoNS, MRSA, GRSA, GISA, VRE, VRSA, PRSP, Propionibacterium acnes, clostridia, lactobacilli | Ent. faecalis, Staph. aureus |
| Ertapenem   | Carbapenem     | MSSA, Str. agalactiae, Str. pyogenes, Enterobacter spp.                                | MRSA, Str. agalactiae, Str. pyogenes |
| Moxifloxacin and gatifloxacin | Quinolone | MSSA, CoNS, PRSP, group A, B, C, F, and G streptococci | Staph. aureus |
| Dalbavancin  | Glycopeptide    | MSSA, MRSA, VRE, CoNS, most streptococci, anaerobes                                   | None to date                |

CoNS, coagulase-negative staphylococci; GISA, glycopeptide intermediate-resistant Staph. aureus; GRSA, glycopeptide-resistant Staph. aureus; MRSA, methicillin-resistant Staph. aureus; MRSA, methicillin-sensitive Staph. aureus; PRSP, penicillin-resistant Str. pneumoniae; VRE, vancomycin-resistant enterococci; VRSA, vancomycin-resistant Staph. aureus.
Tigecycline has proven useful against SSSIs since its approval (Shah 2005), but has no useful activity against Gram-negative organisms (Strahilevitz & Rubinstein 2002).

Newer quinolones, including levofloxacin, moxifloxacin, and gatifloxacin, have superior activity relative to ciprofloxacin against respiratory Gram-positive pathogens (Abbanat et al. 2003), but they do not cover MRSA, and may be compromised by cross-resistance to older agents (Shah 2005). A number of novel cephalosporins are in development; these have shown promising activity against staphylococci but not against enterococci. However, some of these agents have shown good activity against Gram-negative pathogens such as Haemophilus influenzae and Moraxella catarrhalis. According to recent comment (Nathwani 2005), the novel carbapenem ertapenem was the last agent introduced that offered broad antimicrobial cover, but this drug is not active against methicillin- or oxacillin-resistant Gram-positive isolates and has limited activity against Pseudomonas species and ESBL-producing Gram-negative bacteria.

The novel streptogramin combination of quinupristin/dalfopristin has proven useful in patients with serious VRE infections and in SSSIs, including those with MRSA (Kauffman 2003; Raghavan & Linden 2004). However, the combination is not active against infections caused by Ent. faecalis, and resistance has been reported (Chow et al. 1997; Dowzicky et al. 2000). Quinupristin/dalfopristin also has to be given via a central intravenous catheter to avoid phlebitis, and can cause painful arthralgia and myalgia (Kauffman 2003).

The glycopeptides vancomycin and teicoplanin (the latter is not available in the USA) have featured prominently in the management of multiresistant Gram-positive infection in the past, but around 20% of enterococci in the USA are now resistant to vancomycin (Abbanat et al. 2003). Moreover, vancomycin is ototoxic and nephrotoxic (BNF 2005). These considerations, and the threat of global acquisition of resistance to glycopeptides by MRSA, have led to the development of newer derivatives. These include ramoplanin, dalbavancin, and telavancin, all of which show promise and are in clinical development (Shah 2005; Bosso 2004).

There is therefore a need for research to discover and develop agents with activity against a range of both Gram-positive and Gram-negative pathogens, and against atypical pathogens, particularly those most frequently associated with institutional, resistant, or complicated infections. Moreover, research should be ongoing and aimed at the development of “pipelines” of new drugs, as resistance patterns are continually changing over time.

Clinical evidence with tigecycline

Antimicrobial activity

Tigecycline is active against most Gram-positive and Gram-negative aerobes in addition to anaerobes and atypical organisms. In preclinical studies, the drug was shown to have activity against a broad range of Gram-positive pathogens including MRSA, vancomycin-resistant Ent. faecalis and Ent. faecium, and penicillin-resistant Str. pneumoniae; against Gram-negative organisms such as ESBL-producing E. coli and Klebsiella pneumoniae; and anaerobes such as the Bacteroides fragilis group, and atypical organisms including Mycoplasma spp., Chlamydia spp., and rapidly growing mycobacteria (Edlund & Nord 2000; Goldstein et al. 2000; Robin & Hammerschlag 2000; Kenny & Cartwright 2001; Wallace et al. 2002; Jacobus et al. 2004). Tigecycline has also been shown in animal models to be active against a range of glycopeptide- and tetracycline-susceptible and -resistant enterococci (Lefort et al. 2003; Nannini et al. 2003), and the drug’s antistaphyloccocal activity has been confirmed against MRSA, being comparable to vancomycin and quinupristin/ dalfopristin, and superior to linezolid, imipenem, and beta-lactam antimicrobials (Johnson et al. 2003). Other recent data confirm antistaphyloccocal activity against strains resistant to tetracycline, minocycline, erythromycin, clindamycin, gentamicin, levofloxacin, or rifampicin (Picazo et al. 2003), and against tetracycline- and minocycline-resistant bacteremia isolates (Reynolds & Potz 2003).

The activity of tigecycline in clinical isolates was summarized on the basis of minimum inhibitory concentration (MIC) breakpoints of ≤2 mg/L and ≥8 mg/L for susceptibility and resistance, respectively (Garrison et al. 2005), but inconsistencies have been noted between MIC limits obtained in different studies during the establishment of quality control ranges for tigecycline. Because of this, further investigations have been carried out, and have identified discrepancies between results obtained in fresh Mueller–Hinton broth (MHB) and those in aged medium that have been attributed to acceleration of oxidative inactivation of tigecycline in the latter (Bradford et al. 2005a; Petersen & Bradford 2005). As a result, the Clinical Laboratory Standards Institute has approved quality control ranges for tigecycline tested with fresh MHB, and this is now considered the reference method for MIC testing of the drug (Clinical and Laboratory Standards Institute 2005).

These standards have been applied to in-vitro susceptbility testing of tigecycline with isolates from patients participating in phase III studies assessing the clinical efficacy of the drug in complicated SSSIs and intraabdominal infections (Bradford et al. 2005b). The results, which are summarized in Table 5, show a similar pattern to that seen in preclinical studies. Tigecycline was active (MIC ≤2 mg/L in most cases) against the most prevalent pathogens in these patients, which included both Gram-positive and Gram-negative strains of aerobic and anaerobic bacteria. Staphylococci were inhibited regardless of their susceptibility or otherwise to methicillin or oxacillin, and all enterococci were inhibited by tigecycline at a concentration of 0.5 mg/L or less. Good activity was also observed against beta-hemolytic streptococci, whereas MICs for 90% of isolates (MIC90) for tetracycline and minocycline were above resistance breakpoints.

Activity against Gram-negative pathogens, particularly the Enterobacteriaceae, was also good (Table 5). All strains of Citrobacter freundii were inhibited (MIC ≤2 mg/L), in contrast to minocycline, tetracycline, aztreonam, and ceftazidime, MIC90 values for all of which were above resistance breakpoints. All isolates of E. coli were inhibited by tigecycline at concentrations of
**Table 5**  *In-vitro* susceptibility data for tigecycline. Activity against isolates from phase III clinical studies in patients with skin and skin structure infections or intraabdominal infections (Bradford et al. 2005b)

| Organism and no. of isolates (skin and skin structure infections; intraabdominal infections) | Skin and skin structure infections | Intraabdominal infections |
|---|---|---|
|  | MIC<sub>50</sub> (mg/L) | MIC<sub>50</sub> (mg/L) | MIC<sub>50</sub> (mg/L) | MIC<sub>50</sub> (mg/L) |
| **Gram-positive** |  |  |  |  |
| Staphylococcus aureus: methicillin-resistant (127; 16) | 0.12 | 0.25 | 0.25 | 0.25 |
| Staphylococcus aureus: methicillin-susceptible (373; 61) | 0.12 | 0.25 | 0.12 | 0.25 |
| Staphylococcus capitis (11; 7) | 0.12 | 0.25 | NA | NA |
| Staphylococcus epidermidis: methicillin-resistant (44; 30) | 0.12 | 0.25 | 0.25 | 0.5 |
| Staphylococcus epidermidis: methicillin-susceptible (36; 40) | 0.12 | 0.25 | 0.12 | 0.25 |
| Staphylococcus haemolyticus (26; 18) | 0.25 | 1 | 0.25 | 0.5 |
| Staphylococcus hominis (16; 19) | 0.12 | 0.12 | 0.12 | 0.25 |
| Staphylococcus warneri (10; 7) | 0.12 | 0.25 | NA | NA |
| Enterococcus avium (0; 57) | NA | NA | 0.12 | 0.12 |
| Enterococcus faecalis: vancomycin-susceptible (59; 109) | 0.12 | 0.25 | 0.12 | 0.25 |
| Enterococcus faecium: vancomycin-susceptible (11; 50) | 0.06 | 0.12 | 0.06 | 0.12 |
| Enterococcus hirae (3; 12) | NA | NA | 0.06 | 0.06 |
| Streptococcus pyogenes (84; 2) | 0.06 | 0.12 | NA | NA |
| Streptococcus agalactiae (32; 5) | 0.06 | 0.25 | NA | NA |
| Streptococcus dysgalactiae (14; 3) | 0.12 | 0.25 | NA | NA |
| Streptococcus anginosus (29; 154) | 0.06 | 0.12 | 0.06 | 0.12 |
| Streptococcus constellatus (2; 61) | NA | NA | 0.06 | 0.06 |
| Streptococcus equisimilis (11; 0) | 0.12 | 0.5 | NA | NA |
| Streptococcus intermedius (2; 10) | NA | NA | 0.06 | 0.12 |
| Streptococcus mitis (6; 16) | NA | NA | 0.06 | 0.12 |
| Streptococcus oralis (11; 37) | 0.06 | 0.12 | 0.06 | 0.12 |
| Streptococcus salivarius (2; 20) | NA | NA | 0.06 | 0.12 |
| **Gram-negative** |  |  |  |  |
| Citrobacter braakii (0; 15) | NA | NA | 0.5 | 0.5 |
| Citrobacter freundii (2; 29) | NA | NA | 0.5 | 1 |
| Enterobacter cloacae (41; 41) | 0.5 | 1 | 1 | 1 |
| Escherichia coli (90; 964) | 0.25 | 0.5 | 0.25 | 0.5 |
| Klebsiella oxytoca (18; 52) | 0.25 | 0.5 | 0.5 | 0.5 |
| Klebsiella pneumoniae (32; 152) | 0.5 | 1 | 0.5 | 1 |
| Morganella morganii (3; 19) | NA | NA | 2 | 2 |
| Proteus mirabilis (39; 46) | 4 | 4 | 2 | 4 |
| Proteus vulgaris (2; 10) | NA | NA | 1 | 2 |
| Serratia marcescens (10; 12) | 1 | 2 | 2 | 4 |
| Acinetobacter calcoaceticus/baumanii complex (16; 16) | 0.25 | 1 | 0.5 | 2 |
| Pseudomonas aeruginosa (32; 126) | 16 | 32 | 16 | 32 |
| Comamonas testosteron (0; 10) | NA | NA | 0.06 | 0.12 |
| **Anaerobes** |  |  |  |  |
| Bacteroides fragilis (24; 244) | 0.12 | 0.25 | 0.5 | 4 |
| Bacteroides caccae (2; 19) | NA | NA | 0.25 | 8 |
| Bacteroides distasonis (5; 23) | NA | NA | 2 | 2 |
| Bacteroides ovatus (6; 21) | NA | NA | 0.25 | 4 |
| Bacteroides thetaiomicron (3; 108) | NA | NA | 0.5 | 2 |
| Bacteroides uniformis (4; 53) | NA | NA | 0.25 | 1 |
| Bacteroides vulgatus (0; 33) | NA | NA | 0.25 | 2 |
| Clostridium innocuum (1; 12) | NA | NA | ≤0.06 | ≤0.06 |
| Clostridium perfringens (5; 51) | NA | NA | 0.12 | 1 |
| Eubacterium lentum (0; 14) | NA | NA | 0.12 | 0.5 |
| Propionibacterium acnes (8; 17) | NA | NA | ≤0.06 | 0.12 |
| Peptostreptococcus micros (1; 39) | NA | NA | ≤0.06 | ≤0.06 |
| Fusobacterium nucleatum (4; 11) | NA | NA | 0.12 | 0.12 |
| Veillonella spp. (0; 12) | NA | NA | 0.25 | 0.5 |

MIC<sub>50</sub>, minimum inhibitory concentration for 50% of isolates; MIC<sub>90</sub>, minimum inhibitory concentration for 90% of isolates; NA, not applicable.
Tigecycline | outcomes review

2 mg/L or less, and good activity was noted against *K. pneumoniae* isolates (including those that showed resistant to ceftazidime and aztreonam) and *K. oxytoca*. MICs ranged from 0.5 to 8 mg/L for *Serratia marcescens*, but lower values for most strains were reflected by the MIC50 values reported of 1 or 2 mg/L. There was a trend towards higher MICs for the Proteaeae than for other Enterobacteriaceae. For strains from patients with intraabdominal infections, MIC50 was 2 mg/L for *Morganella morgani* and *Proteus vulgaris*. Tigecycline was also active against *Acinetobacter baumanii*, but was less active against *P. aeruginosa* (Table 5).

Tigecycline was also active against anaerobic pathogens, and all Gram-positive anaerobes were inhibited by tigecycline 2 mg/L or less. There was a wide range of MICs against members of the *B. fragilis* group (0.06–16 mg/L), but MIC50 values for all species of *Bacteroides were 2 mg/L or less.

**Clinical cure and microbiologic eradication in patients**

Clinical studies of tigecycline have focused to date on clinical cure and microbiologic eradication in patients with complicated SSSIs or intraabdominal infections. The primary endpoint of choice in such trials should be cure and eradication at a predefined test-of-cure visit that postdates the end of treatment. Cures that are apparent when treatment ceases may later relapse, with subsequent requirement for further therapy. Clinical evidence for the efficacy of tigecycline against the serious and complicated infection types examined to date appears good, and overall clinical activity is as might be expected on the basis of the drug's *in-vitro* bacteriologic profile.

In all studies reviewed, except for one phase II trial in which two dosages were compared, tigecycline was given intravenously as a 100 mg loading dose followed by 50 mg every 12 hours for at least 5 days (Table 6). Two large, randomized, multicenter, double-blind phase III studies (level 2 evidence) have recently compared tigecycline with intravenous combination therapy with vancomycin 1 g and aztreonam 2 g every 12 hours for up to 14 days in 1057 patients (clinically modified intent-to-treat (ITT) populations) with complicated SSSIs (Breedt et al. 2005; Sacchidanand et al. 2005). Both studies showed no significant differences between either regimen in terms of clinical cure or microbiologic eradication rates (see also Table 6). Clinical cure rates (where treatment was observed to resolve the infection process) were measured in the clinically modified ITT populations (patients who received at least one dose of study drug and who showed clinical signs of a complicated SSSI) and microbiologic eradication in the microbiologically evaluable populations. Patients were deemed to be microbiologically evaluable if they satisfied all protocol requirements and had a baseline culture containing at least one causative isolate that was susceptible to both study treatments. Across the difference treatments in both studies, clinical cure rates ranged from 75.5 to 86.9% (Table 6).

Level 2 evidence is also available from two randomized, double-blind, and multicenter studies comparing tigecycline with imipenem/cilastatin (500 mg/500 mg every 6 hours intravenously, weight-adjusted where necessary) for 5 to 14 days for complicated intraabdominal infections (Fomin et al. 2005; Oliva et al. 2005). Of 825 (Oliva et al. 2005) and 817 (Fomin et al. 2005) patients in the modified ITT populations who received more than one dose of study medication, 621 and 641, respectively, were included in the microbiologically modified ITT populations of patients who had clinical evidence of an intraabdominal infection, met minimal disease criteria, and who had a confirmed baseline isolate. Of these, 502 (Oliva et al. 2005) and 523 (Fomin et al.

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### Table 6 | Summary of outcomes evidence for tigecycline: clinical and microbiologic cure in patients with complicated skin and skin structure or intraabdominal infections

| Level of evidence | Outcomes | Comparators | Study population | Reference |
|-------------------|----------|-------------|------------------|-----------|
| 2                 | CC in 84.3% (TIG) and 86.9% (VAN/AZT) of patients (clinically modified ITT population). ME in 84.8% and 93.2% for microbiologically evaluable population of 312 (criteria for noninferiority of TIG met) | TIG 50 mg q 12 h vs VAN 1 g + AZT 2 g q 12 h | Clinically modified ITT population of 520 with complicated skin and skin structure infections | Breedt et al. 2005 |
| 2                 | CC in 75.5% (TIG) and 76.9% (VAN/AZT) of patients (clinically modified ITT population). ME in 80.9% and 77.8% for microbiologically evaluable population of 228 (criteria for noninferiority of TIG met) | TIG 50 mg q 12 h vs VAN 1 g + AZT 2 g q 12 h | Clinically modified ITT population of 537 with complicated skin and skin structure infections | Sacchidanand et al. 2005 |
| 2                 | CC in 73.5% (TIG) and 78.2% (IMI/CIL). ME in 80.6% and 82.4% for microbiologically evaluable population of 502 (criteria for noninferiority of TIG met) | TIG 50 mg q 12 h vs IMI/CIL 500 mg/500 mg q 6 h | Microbiologically modified ITT population of 621 patients with complicated intraabdominal infections | Oliva et al. 2005 |
| 2                 | CC in 86.6% (TIG) and 84.6% (IMI/CIL). ME in 91.3% and 89.9% for microbiologically evaluable population of 523 (criteria for noninferiority of TIG met) | TIG 50 mg q 12 h vs IMI/CIL 500 mg/500 mg q 6 h | Microbiologically modified ITT population of 641 patients with complicated intraabdominal infections | Fomin et al. 2005 |
| 3                 | CC in 67% (TIG 25 mg) and 74% (TIG 50 mg) of patients. ME in 56% and 69% | TIG 25 mg q 12 h vs TIG 50 mg q 12 h | 160 with complicated skin and skin structure infections | Poster et al. 2004 |
| 3                 | CC in 55% of patients | TIG 50 mg q 12 h | 111 with complicated intraabdominal infections | Murray et al. 2003 |
| 5                 | Full recovery with TIG (n=2); no response with colistin (n=5) | TIG (dosage not stated) | 7 ventilator-assisted patients with resistant *Acinetobacter baumanii* pneumonia | Wilson 2004 |

AZT, aztreonam; CC, clinical cure; IMI/CIL, imipenem/cilastatin; ITT, intent-to-treat; ME, microbiologic eradication; q 6 h, every 6 hours; q 12 h, every 12 hours; TIG, tigecycline; VAN, vancomycin.
2005) were microbiologically evaluable. The mean duration of therapy across both studies was approximately 8 days. At the test-of-cure assessment (12–44 days after therapy), clinical cure rates were 73.5 (Oliva et al. 2005) and 86.6% (Fomin et al. 2005) for tigecycline, and 78.2 (Oliva et al. 2005) and 84.6% (Fomin et al. 2005) for imipenem/cilastatin (Table 6). At approximately 80 (Oliva et al. 2005) and 90% (Fomin et al. 2005), rates of microbiologic eradication were also similar between treatments in both studies for microbiologically evaluable populations. Tigecycline met statistical criteria for noninferiority to imipenem/cilastatin for all endpoints in both studies.

The most common diagnosis in both studies was complicated appendicitis (around half of all patients overall), followed by perforated intestine and gastric or duodenal ulcer (Oliva et al. 2005), or by cholecystitis or intraabdominal abscess (Fomin et al. 2005). *E. coli* was the most commonly isolated aerobe, followed by *Klebsiella* spp., in both trials. Oliva et al. (2005) reported identification of six ESBL-producing *E. coli* and seven ESBL-producing *K. pneumoniae* isolates before therapy in their study.

Postier et al. (2004) have reported fully their phase II study (level 3 evidence) in 160 ITT patients with complicated SSIs who received tigecycline 25 mg (n=79) or 50 mg (n=81) every 12 hours for 7–14 days. The primary endpoint in this well-designed and executed study was clinically observed cure (defined as resolution or improvement of all signs and symptoms to the extent that no further antibiotic therapy was necessary) at the test-of-cure visit approximately 3 weeks after the start of therapy. Bacteriologic responses were assessed according to the National Committee for Clinical Laboratory Standards (NCCLS; now known as the Clinical Laboratory Standards Institute) criteria and clearly defined MIC breakpoints; pharmacokinetic and safety analyses were also included. The main diagnoses were ulcer with acute infection (35% of patients) and major abscesses (31%). Of all patients, 82% were infected with at least one of the pathogens selected for bacteriologic analysis: *m* -resistant or -sensitive *Staph. aureus*, *Str. pyogenes*, *E. coli*, *Ent. faecalis*, or *Ent. faecium*. As shown in Table 6, clinical cure as specified was noted in 67 and 74% of patients in the 25 mg and 50 mg groups, respectively, at the test-of-cure visit (six patients in each group required further treatment between the end-of-treatment and test-of-cure visits). In the 25 mg group, 56% of patients had microbiologic eradication, which was compared with 69% in the 50 mg group. The MICs required for 90% of all selected isolates ranged from 0.06 to 0.5 mg/L (Postier et al. 2004).

Tigecycline was also effective in patients with complicated intraabdominal infections (mostly perforated or gangrenous appendicitis, complicated cholecystitis, perforated diverticulitis, or peritonitis) in a noncomparative phase II study (Murray et al. 2003). In this trial, presented as an abstract and summarized in Table 6, 111 ITT patients received tigecycline for up to 14 days, after which 55% were assessed as clinically cured at the test-of-cure visit. These patients represented a challenging population, meeting inclusion criteria that included a need for surgical extirpation of the source of infection plus antibiotic therapy, and subsequent isolation of both aerobic and anaerobic cultures. Although not considered strong evidence, a small descriptive study (level 5 evidence) in a series of seven patients requiring intensive care in a UK district general hospital has indicated that tigecycline is likely to be useful in persons with ventilator-assisted pneumonia associated with resistant *Acinetobacter* infection (Table 6) (Wilson 2004). Tigecycline was tried successfully in two patients in this unit after increasing problems with resistance to carbapenems, tobramycin, amikacin, colistin, and minocycline, to the point where one patient had already died with *A. baumannii* infection resistant to all available antifective agents. The other five patients in this series were treated with colistin and failed to respond, but it was not clear from the abstract presented as to how these patients were managed thereafter.

**Tolerability and adverse events**

Clinical results to date bear out preclinical findings with tigecycline in healthy persons and those with renal disease that showed linear pharmacokinetics and good tolerability of the drug with no need for dosage adjustments because of age, gender, or renal impairment (Troy et al. 2003; Meagher et al. 2004; Muralidharan et al. 2005a,b). Nausea and vomiting appear to be the treatment-emergent adverse events of chief concern, and affected 35% of patients who received tigecycline 50 mg every 12 hours in the phase II study of Postier et al. (2004). Of the five treatment withdrawals, two were associated with nausea and vomiting and one with rash; none of these events were life threatening. Laboratory abnormalities (elevated hepatic enzymes, raised blood urea nitrogen levels, or anemia) considered likely to be associated with tigecycline therapy were also noted in nine patients, but none led to treatment discontinuation. Overall, nausea and vomiting were cited as the most common events with tigecycline in all other studies. Rates were higher than with vancomycin/aztreonam in patients with complicated SSIs in phase III trials (Breedt et al. 2005; Sacchadinand et al. 2005), although vancomycin/aztreonam was associated with higher rates of hepatic enzyme elevation, pruritus, and rash (Table 7). In one phase III study in patients with intraabdominal infections, the increases in rates of nausea and vomiting with tigecycline over imipenem/cilastatin approached or attained statistical significance (Table 7), but there were no statistically significant differences between groups in this respect in the other phase III intraabdominal infection trial (Fomin et al. 2005).

**Ongoing studies**

The studies reported so far indicate the likely utility of tigecycline in complicated and/or nosocomial infections of particular interest at the present time, but trials are also in progress to examine the activity of the drug in other serious and problematic infections. These include:

- an open-label and noncomparative trial in patients with serious resistant Gram-negative infections (e.g. *A. baumannii*, *Enterobacter* spp., *K. pneumoniae*, etc.) (evidence level 3)
- randomized and double-blind comparisons with levofloxacin in patients with community-acquired pneumonia (evidence level 2)
Table 7 | Summary of outcomes evidence for tigecycline: adverse events

| Level of evidence | Outcomes | Comparators | Study population | Reference |
|-------------------|----------|-------------|------------------|-----------|
| 2                 | Similar rates of TEAEs in both groups. More (P<0.05) nausea (25.2 vs 5.2%) and vomiting (12 vs 2.2%) with TIG; more increased AST (5.2 vs 1.5%) or ALT (6.7 vs 1.8%) with VAN/AZT | TIG 50 mg q 12 h vs VAN 1 g + AZT 2 g q 12 h | Safety population of 453 with complicated skin and skin structure infections | Breedt et al. 2005 |
| 2                 | Similar rates of TEAEs in both groups. More (P<0.05) nausea (43.2 vs 11%), vomiting (26.7 vs 5%), dyspepsia (5.5 vs 1.8%), and anorexia (4.1 vs 0.7%) with TIG; more abnormal LFTs (5.7 vs 1%), puritus (10.7 vs 4.5%), and rash (7.8 vs 2.7%) with VAN/AZT | TIG 50 mg q 12 h vs VAN 1 g + AZT 2 g q 12 h | Safety population of 573 patients with complicated skin and skin structure infections | Sacchidanand et al. 2005 |
| 2                 | Most common TEAEs: nausea (31% TIG and 24.8% IMI/CIL; P=0.052) and vomiting (25.7% TIG and 19.4% IMI/CIL; P=0.037) | TIG 50 mg q 12 h vs IMI/CIL 500 mg/500 mg q 6 h | Modified ITT cohort of 825 patients with complicated intraabdominal infections | Oliva et al. 2005 |
| 2                 | Most common TEAEs: nausea (17.6% TIG and 13.3% IMI/CIL; P=0.1) and vomiting (12.6% TIG and 9.2% IMI/CIL; P=0.144) | TIG 50 mg q 12 h vs IMI/CIL 500 mg/500 mg q 6 h | Modified ITT cohort of 817 patients with complicated intraabdominal infections | Fomin et al. 2005 |
| 3                 | Commonest TEAEs: nausea (22% with 25 mg; 35% with 50 mg). 5 discontinuations with 50 mg. No life-threatening reactions | TIG 25 mg q 12 h vs TIG 50 mg q 12 h | 160 with complicated skin and skin structure infections | Postier et al. 2004 |
| 3                 | Nausea and vomiting most common TEAEs | TIG 50 mg q 12 h | 111 with complicated intraabdominal infections | Murray et al. 2003 |
| 5                 | No adverse events with TIG therapy | TIG (dosage not stated) | 7 ventilator-assisted patients with resistant Acinetobacter baumanii pneumonia; 2 received TIG, 5 received colistin | Wilson 2004 |

ALT, alanine aminotransferase; AST, aspartate aminotransferase; AZT, aztreonam; IMI/CIL, imipenem/cilastatin; ITT, intent-to-treat; LFT, liver function test; q 6 h, every 6 hours; q 12 h, every 12 hours; TEAE, treatment-emergent adverse event; TIG, tigecycline; VAN, vancomycin.

Completion of these large-scale studies will provide a more complete picture of the overall activity of tigecycline in patients with infections associated with major management problems at the present time, and results are therefore awaited.

Economic evidence

Early level 3 evidence, presented as an abstract, suggests that reduction of length of hospital stay may accrue from the use of tigecycline in patients with VRE, and with vancomycin for serious infections in patients with MRSA. Conditions covered include bacteremia, pneumonia, and bacterial SSSIs (evidence level 2).

- a randomized and double-blind comparison with linezolid for serious infections in patients with VRE, and with vancomycin for serious infections in patients with MRSA. Conditions covered include bacteremia, pneumonia, and bacterial SSSIs (evidence level 2).

Yu et al. (2005) have reported a nonsignificant trend towards reduced duration of treatment relative to vancomycin/aztreonam when patients with complicated SSSIs were treated with tigecycline (Table 8). This analysis was based retrospectively on the same two clinical studies as above (Breedt et al. 2005; Sacchidanand et al. 2005), and included 1041 patients from the modified ITT population with complete hospitalization data. Independent risk factors for prolonged hospitalization that were identified and adjusted for were diabetes, infected ulcer, female gender, and concomitant use of antibiotics. It should be noted here that length of stay and duration of treatment were only two of several measured health outcomes, and that their reduction were not prospectively defined in the protocols of the two clinical trials on which these economic analyses were based.

Resource utilization

Apart from the preliminary level 3 evidence described in the previous section, there are as yet no other published economic data relating to the use of tigecycline at this stage in the drug’s development, but results indicate the potential of novel antiinfectives such as tigecycline in complicated, nosocomial, and/or resistant infections. The pace at which resistance to
antimicrobial therapy develops continues to increase, but the magnitude of the effect of this resistance on outcomes and consumption of healthcare resources remains poorly understood. A review of this subject by Cosgrove and Carmeli (2003) has attempted to address these issues. Resistance often leads to a mismatch between empiric treatment and susceptibility, and thus to a significant delay in the delivery of effective antimicrobial therapy. Such delay resulted in significantly extended duration of hospitalization and increased hospital charges for patients with ESBL-producing strains of *K. pneumoniae* or *E. coli* relative to controls in one study (Lautenbach et al. 2001). Resistant organisms may also require treatment with more toxic or aggressive therapy than might usually be considered: colistin for resistant *Pseudomonas* or *Acinetobacter* strains may cause renal dysfunction (Levine et al. 1991), and total resistance may lead to the need for surgery, with high rates of mortality being seen in patients whose locus of infection cannot be surgically removed (Harris et al. 1999).

From the hospital perspective, most available studies have shown a link between antibiotic resistance and increased mortality, morbidity, and cost (Cosgrove & Carmeli 2003). These results, however, are limited because they do not include costs associated with care in the community and in rehabilitation facilities other than hospitals. The long-term effects of a resistant infection on future health, and on loss of work and family time caused by prolonged hospitalization or incapacity, are also difficult to estimate. The US Congress Office of Technology Assessment (1995) estimated the national cost of antibiotic resistance in that country to be $US4 billion for the year 1995, but this conservative assessment included only direct patient effects without wider societal costs.

The chief value of antiinfective drugs introduced at the present time for the types of infection under discussion lies in their potential in empiric therapy. Any agent that can be given with a good chance of success early in the course of disease will contribute to the avoidance of much morbidity and need for additional therapy following on from treatment failure as described above. Clinical results show that tigecycline is likely to fall into this category, but reliable outcomes and economic data pertaining to the use of the drug will have to be assessed in direct comparisons with other therapies in specified institutions, as treatment and resistance patterns vary between locations within and between countries. The cost to providers of drugs used in any study will also have to be known for reliable comparisons to be made, as acquisition costs for novel broad-spectrum antibiotics are typically much higher than those of established agents. Nevertheless, the activity of tigecycline in preclinical and clinical studies to date suggest that the drug does indeed have the potential to circumvent adverse outcomes and subsequent costs resulting from treatment failure in complicated and resistant infections.

### Patient group/population

Current product registration in the USA limits the use of tigecycline to adults with intraabdominal and complicated SSSIs, but the range of clinical trials still underway indicates that the drug is expected to contribute to the treatment of patients with a much wider range of conditions caused by multiresistant pathogens. Chief among these are likely to be patients with pneumonia (both hospital- and community-acquired), including some with atypical infections, patients with bacteremia/sepsis, and those with Gram-negative infections. Patients in the last group may benefit in particular, as tigecycline has good activity against a number of Gram-negative pathogens that are resistant to other novel antibacterials, most of which are directed against Gram-positive infections (Abbanat et al. 2003; Nathwani 2005). There is no oral formulation of tigecycline, and the drug must be given intravenously, but this consideration is of little consequence in light of the characteristics of the target patient group, the large majority of whom will have undergone surgery or will be sufficiently unwell to require care in hospital or some other institutional setting.

Preliminary level 5 evidence indicates that tigecycline is also likely to be useful in high-risk patients in intensive care who are particularly vulnerable to multiresistant pathogens (Wilson 2004). Ethical considerations make the conduct of randomized trials in patients of this type more difficult than in other groups, but accumulating clinical experience with tigecycline is likely to shed more light on the utility of the drug in this setting.

The use of tigecycline is likely to be empiric in the first instance in many patients, as early treatment is important in the management of serious infections. The broad spectrum of activity, which includes Gram-negative pathogens and some atypical organisms, suggests that this empiric therapy will be continued and should be successful in most patients after identification of the infective
agent. This situation could change, however, if resistance were to emerge, and it will therefore be important to combine appropriate use of tigecycline with effective infection control measures and close monitoring of susceptibility patterns.

Tigecycline has been approved in the USA, Mexico, Venezuela, and Brazil for the treatment of complicated intraabdominal infections and complicated skin and skin structure infections in adults (Anon. 2005). It is likely that these indications will be expanded as the results of ongoing phase III studies are released (filings are expected for hospital- and community-acquired pneumonia in September 2006). Registration is awaited in Australia, Canada, the European Union, and Switzerland. The use of tigecycline in children has not yet been evaluated.

Outcomes summary

Tigecycline has recently been introduced into clinical practice as part of the effort to combat the growing problem of bacterial resistance to antinfective therapy, particularly in patients with serious or complicated infections and those in hospitals and other institutions. Early reports of the first of the large-scale randomized clinical trials to be conducted have led to the approval of tigecycline for the treatment of complicated SSSIs and intraabdominal infections in the USA, Mexico, Venezuela, and Brazil, with approvals expected to follow in other countries, and other indications added, as ongoing studies report. Published or impending trials focus on comparisons of tigecycline with established agents that are already in common use for serious infections. Substantial evidence so far shows that tigecycline is as effective as these other agents, but superiority has not been shown. As resistance patterns vary widely between centers and countries, more research and clinical experience will be necessary before a more complete picture begins to emerge. Comparisons of tigecycline with other new agents developed specifically for use against the modern generation of multiresistant pathogens will also be needed. These will be of special interest because of tigecycline’s activity against Gram-negative and some atypical pathogens.

The possibility of development of resistance to any new antimicrobial is a significant source of concern. Tigecycline appears to have the potential to resist common mechanisms of resistance, as shown by unsuccessful attempts to create tigecycline-resistant isolates in the laboratory by exposing pathogens to suboptimal concentrations of the drug (Projan 2000). In addition, tigecycline is not affected by efflux, ribosomal protection, DNA gyrase mutation, binding site modification, or beta-lactamases (Bradford 2004).

Clinical endpoints being measured in trials are as expected for a new antinfective agent. The outcomes of clinical cure and bacteriologic eradication have immediate clinical relevance and show how well the drug works in the setting in which it is given. However, the economic implications attached to the introduction of tigecycline to clinical practice are less clear at present. For economic benefits to be realized, tigecycline will have to maintain high rates of clinical success against organisms resistant to other drugs, with sustained efficacy as empiric therapy in cases where other agents may fail. Longer term evaluation will be needed to show how these factors translate into other (e.g. quality-of-life and societal) benefits beyond the perspective of the institution where treatment is administered. Limited evidence suggests that the use of tigecycline may result in more rapid discharge of patients from hospital back into the community; ultimately, this will require confirmation with evidence from properly designed economic studies. It would also be of value to determine any effect of tigecycline on length of stay in intensive care. Tolerability in clinical studies appears good, and similar to the beta-lactam comparators used to date (imipenem/cilastatin and aztreonam are associated chiefly with nausea, vomiting, and diarrhea). Vancomycin may be ototoxic and hepatotoxic in some patients, but tigecycline does not appear to be associated with potentially serious effects such as organ toxicity.

The introduction of tigecycline is timely in an era of increasing bacterial resistance and relative shortage of new antibacterial agents. Current evidence shows that this novel glycycline is effective in the patient population in which it is being targeted, but it is not yet clear whether it is more effective or delivers better outcomes than other agents of interest. Continuing evolution of resistance patterns worldwide will no doubt affect the relative placing of tigecycline and other antinfectives (both old and new), and further evidence will need to be gathered to show whether overall benefits outweigh high acquisition costs (as with all novel antibacterials) relative to older agents.

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