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Antistaphylococcal (MSSA, MRSA, MSSE, MRSE) antibiotics

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optimal clinical outcomes while minimizing resistance potential problems and applying pharmacoeconomic considerations to antibiotic selection.

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> BURKE A. CUNHA, MID Guest Editor

ANTIBIOTIC THERAPY, PART II

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ANTISTAPHYLOCOCCAL (MSSA, MRSA, MSSE, MRSE) ANTIBIOTICS

Franco Paradisi, MD, Giampaolo Corti, MD, and Daniela Messeri, MD

Coagulase-positive (Staphylococcus aureus) and coagulase-negative staphylococci are gram-positive cocci appearing in clusters that are important causes of infection, primarily of the bloodstream, native and prosthetic cardiac valves, other implanted devices, and skin. Their progressively reduced susceptibility to penicillin, methicillin, and glycopeptides makes treatment of staphylococcal infections difficult. This article reviews the current status and the future perspectives for the therapy of infections caused by S. aureus and coagulase-negative staphylococci, such as S. epidermidis.

STAPHYLOCOCCUS AUREUS

S. aureus has been known as a bacterial agent of infection since 1882, when Ogston⁴⁰ clarified its role in sepsis and abscess formation. A list of community-acquired and hospital-acquired infections caused by S. aureus follows:

Bacteremia and sepsis
Endocarditis of native and prosthetic cardiac valve
Surgical site infection

Toxic shock and scalded skin syndromes Localized skin infections (folliculitis, furuncle, abscess, cellulitis, impetigo, pyoderma, hydradenitis, mastitis)

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Bone and joint infections (osteomyelitis, septic arthritis)
Food poisoning

Bacterial meningitis

Localized brain infections (epidural abscess, brain abscess, subdural empyema)

Upper respiratory tract infections (otitis media, sinusitis)

Lower respiratory tract infections (acute exacerbation of chronic bronchitis, pneumonia, pleural empyema)

Most of these infections occur in subjects with predisposing factors, primarily congenital or acquired (rheumatoid arthritis) chemotaxis defects, chronic granulomatous diseases, diabetes mellitus, or presence of foreign bodies. At present, S. aureus is significant as a frequent and difficult agent of nosocomial infections. A report from the National Nosocomial Infections Surveillance System cited S. aureus as the most common cause of infection in U.S. hospitals, with an average 13% incidence. In particular, it was the first agent of surgical site infections and lower respiratory tract infections and the second cause of blood-stream infections and all other site infections excluding urinary tract infections (Fig. 1).37

Epidemiology of β-Lactam Resistance

Originally, *S. aureus* was uniformly susceptible to penicillin, but soon after its introduction into clinical practice, penicillin resistance emerged as the result of the production of inactivating enzymes (i.e., β-lactamases). ³¹ Currently, most *S. aureus* strains isolated from community-acquired lower respiratory tract infections are β-lactamase producers

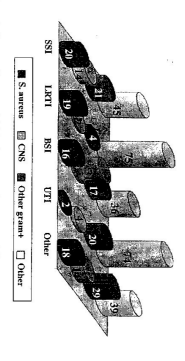


Figure 1. Causes of nosocomial infections: prevalence rates (%) of staphylococci and other micro-organisms. SSI = surgical site infection, LRTI = lower respiratory tract infection, BSI = blood stream infection, UTI = urinary tract infection, CNS = coagulase-negative staphylococci. (Data from National Nosocomial Infections Surveillance (NUIS) Report, data summary, October 1986–April 1996, May 1996. Am J Infect Control 24:380, 1996.)

(approximately 80% in Europe and 90% in the United States) and resistant to penicillin and ampicillin. ⁶⁰ These isolates usually remain susceptible to other β-lactams, such as penicillinase-resistant penicillins (nafcillin, oxacillin) penicillin-β-lactamase inhibitor combinations, and most cephalosporins.

In the 1950s, resistance of *S. aureus* to chloramphenicol, erythromycin, and the tetracyclines emerged, ¹⁵ and in 1960, the first strains resistant to methicillin were isolated in London. ²⁴ Methicillin resistance of staphylococci is mediated by the *mecA* gene, which encodes for penicillin-binding protein (PBP) 2a; PBP2a has low affinity for all β-lactams, conferring resistance not only to methicillin and other penicillins, but also to all β-lactams, including cephalosporins and carbapenens. ⁸ Methicillin-resistant *S. aureus* (MRSA) frequently is cross-resistant to aminogly-cosides, lincosamides, macrolides, tetracyclines, trimethoprim, and sulfonamides. By the end of the 1970s, MRSA isolates spread worldwide in hospitals and communities. In U.S. hospitals, the percentage of MRSA associated with nosocomial infections rose from 2.4% in 1975 to 29% in 1991. ⁴¹ Similar results (29.3%) have been found in nosocomial bloodstream infections, with higher proportions of MRSA in the eastern United States (29.8% to 38.5%) than in the western United States (14.5% to 22.5%). ¹² Sharp differences in MRSA prevalence rates between the United States (26.2%) and Canada (2.7%) have been noted as well. ⁴⁵

Intensive care units (ICUs) generally report the highest percentages of MRSA: a laboratory-based surveillance program conducted in the United States found a significantly higher prevalence of MRSA in ICU patients (35.2%) than in non-ICU inpatients (31.9%; P<.01). A European multicenter study reported an average rate of prevalence of 12.8%, ranging from less than 1% in Scandinavia to greater than 30% in Mediterranean countries (Spain, France, Italy). Distribution of MRSA strains was much higher (32.5%) in the ICUs, ranging from 0 in Holland to 52.9% in Austria and Italy. The well-known European Prevalence of Infection in Intensive Care (EPIC) study reported an average rate of 59.6% and the highest prevalence (81%) found in Italy.

MRSA is significantly less prevalent among outpatients (17.7%) than among inpatients (31.9% to 35.2%).\(^{17}\) Current trends in shortening hospital stays, performing outpatient surgery, and providing home parenteral therapies as well as the use of broad-spectrum antibiotics for community-acquired infections, injection drug abuse, and contact with MRSA-colonized subjects all increase the risk of the spread of hospital-acquired MRSA to the community.\(^{53}\)

Epidemiology of Reduced Susceptibility to Glycopeptides

S. aureus was fully susceptible to vancomycin and teicoplanin (the latter drug being available for clinical use in Europe and in other parts of the world but not in the United States). Transfer of resistance genes

are directed against the enterococcal mechanism of resistance to vanco-

from vancomycin-resistant enterococci (VRE) to *S. aureus* has been achieved in the laboratory, however.³⁹ The concern that vancomycin resistance might emerge and spread in *S. aureus* gained ground among microbiologists and clinicians in the early 1990s, and well-known reports on the decreased susceptibility of *S. aureus* to vancomycin exist. The first case of vancomycin-intermediate *S. aureus* (VISA) was reported in 1996 from Japan.²² Two VISA strains were isolated in 1997 in Michigan and New Jersey⁶⁶ and another one in the New York area in 1998.⁶³ The first chronologic appearance of a VISA strain probably occurred in France in November 1995, however, when it was isolated from a two-year-old girl with leukemia and MRSA bacteremia.⁴⁶ These four strains shared the following factors: minimal inhibitory concentration (MIC) of vancomycin of 8 mg/L, presence of underlying disease, presence of indwelling medical devices, and previous therapy with vancomycin. MRSA strains with reduced susceptibility to teicoplanin (MIC ≥ 16 mg/L) have been isolated in Europe, the United States, and South America²⁵ so that the acronym *GISA* (glycopeptide-intermediate *S. aureus*) has been proposed as well.

way, NJ) proved to be the most effective in detecting VISA strains.70 The same study suggested that MRSA isolates for which the vancomycin isolates and from 0 in 1995 to 0.4% for blood isolates.56 the United States, the percentage of MRSA strains with a vancomycin MIC of 4 mg/L increased from 0.2% in 1995 to 0.4% in 1997 for all reduced vancomycin susceptibility and should be carefully studied. In MIC was greater than or equal to 4 mg/L should be considered to have among commercial kits, MicroScan conventional panels (Dade Behring, the disk-diffusion test should not be used because it is less reliable; reported 17- or 18-mm zone sizes (full susceptibility),66 suggesting that the latter technique should not be used for testing staphylococci with vancomycin. A well-designed study verified that VISA isolates are best of vancomycin (intermediate susceptibility), whereas disk-diffusion test niques performed: Broth microdilution methods found an 8 mg/L MIC Michigan and New Jersey VISA strains varied according to the tech-Intermediate susceptibility to glycopeptides is difficult to detect with standard laboratory tests. Vancomycin susceptibility patterns of West Sacramento, CA) and E-test (AB Biodisk North America, Piscata-(NCCLS) reference method, after a 24-hour incubation period, whereas lution with the National Committee for Clinical Laboratory Standards detected by nonautomated quantitative tests, such as the broth microdi-

The mechanism of reduced susceptibility to glycopeptides is still unclear, but it seems not to be due to the *van* genes present in enterococci. To Some researchers have found a hyperproduction of PBPs that compete with vancomycin for binding to peptidoglycan precursors; other scientists have noted alteration in the structure of cell walls (thickening, hyperproduction of nonamidated muropeptides, reduced cross-linking of the peptidoglycan) resulting in reduced amounts of drug available at bacterial targets. Euclidation of resistance mechanisms is of great importance because current efforts in drug development

Prevention and Control of Antibiotic-Resistant

Prevention and Control of Antibiotic-Resistant Staphylococcus aureus Infection

Measures to prevent and control spread of MRSA and GISA are crucial and include early isolation of patients, use of contact precautions (hand washing and use of gloves, gowns, and masks), ward closure, and screening of patients and health personnel for carriage.^{7,76} Prudent use of antibiotics is mandatory above all, however, because it has been known for years that there is a causal association between antimicrobial usage and antimicrobial resistance.²⁵ Some authors have assumed that MRSA strains emerged and spread in hospitals after the use of prophylactic narrow-spectrum cephalosporins and use of therapeutic doses of third-generation cephalosporins and use of therapeutic doses of third-generation cephalosporins.⁵⁸ As a result of new control strategies decreasing administration of cephalosporins in favor of piperacillintazobactam, the prevalence of MRSA isolates declined in a U.S. hospital from 35% to 23%.⁵⁶ After the emergence of VRE and GISA strains, the Centers for Disease Control and Prevention published new exhaustive guidelines, and a prudent use of vancomycin was cited as the major factor contributing to limited spread of these troublesome multiresistant bacteria.^{23,52}

Antimicrobial Agents for Treatment of Staphylococcus aureus Infection

 β -Lactams are the cornerstone for treatment of susceptible *S. aureus* infections. Vancomycin must be reserved only for patients with a history of β -lactam allergy because it appears to be a less effective antistaphylococcal drug than β -lactams.²⁰ Treatment failures with vancomycin have been observed in 10% to 20% of patients with endocarditis compared with 5% to 10% of β -lactams.⁸ In Europe and in other parts of the world, teicoplanin is available as well.

Penicillin remains the drug of choice for therapy of infections caused by penicillin-susceptible *S. aureus* strains. Severe infections must be treated intravenously with penicillin G (Table 1); addition of gentamicin is advisable for endocarditis to profit by the synergistic effect of the combination β-lactam-aminoglycoside. Further addition of rifampin to the combination of penicillin (or a glycopeptide in allergic subjects) and gentamicin has been advocated for treatment of prosthetic-valve endocarditis, although the efficacy of rifampin as an adjunctive drug remains controversial. Patients with mild infections alternatively can be treated with less potent oral agents, such as penicillin V and others that are generally effective against penicillin-susceptible isolates (co-trimoxazole, lincosamides, macrolides, tetracyclines).

Severe infections:

infusion)

Severe infections:

Cefazolin 2 g tid IV

First Choice

Penicillin G 4 million units every 4 h
IV (24 million U/d in IV continuous

Nafcillin or oxacillin 2 g every 4 h IV

Mild infections:
Amoxicillin-clavulanate 1 g bid or tid

Amoxiculin-clavulanate 1 g bid or tid orally
Vancomycin 1 g bid IV
Teicoplanin 3 mg/kg (mild infections),
6 mg/kg (severe infections), 12 mg/kg
(endocarditis) bid for the first 2-5 d

(loading dose), followed by 3-12 mg/kg qd IV

Quinupristin-dalfopristin 7.5 mg/kg bid

modifying enzymes produced by many MRSA strains generally make be active in vitro and bactericidal for some isolates. Aminoglycoside-

dosage tetracyclines, such as doxycycline and

in combination with another effective antistaphylococcal agent. Lowoccur early in vivo if it is used alone so that rifampin must be used only and bactericidal antistaphylococcal agent, but high-level resistant strains to 97% in Europe and 92% in the United States).73,74 Rifampin is a potent to 97% in Europe and 98% in the United States) and erythromycin (38% respectively; similar results have been obtained with clindamycin (30% but studies on its efficacy against clinical MRSA isolates in Europe and the United States have reported resistance rates of 47% to 76% and 100%, serious infections have been treated successfully with co-trimoxazole, experimental endocarditis, but there is no human experience.16 Some

minocycline, seem to

Mild infections: Penicillin V 250 mg tid orally

Alternatives

Vancomycin 1 g bid IV
Teicoplanin 3 mg/kg (mild infections),
6 mg/kg (severe infections),
12 mg/kg (endocarditis) bid for the
first 2–5 d (loading dose), followed
by 3–12 mg/kg qd IV
Mild infections:
Co-trimoxazole 960 mg (1 doublestrength tablet) bid orally
Doxycycline 100 mg bid orally
Erythromycin 500 mg qid orally

Erythromycin 500 mg qid orally Clindamycin 300 mg qid orally

Co-trimoxazole 60 mg/kg/d in 2-4 divided doses IV (960 mg bid orally) Minocycline (or doxycycline) 100 mg bid IV or orally Ciprofloxacin 400 mg bid IV (500 mg bid

orally)
Levofloxacin 500 mg qd IV or orally
Quinupristin-dalfopristin 7.5 mg/kg bid

Linezolid 600 mg bid IV or orally Based on antimicrobial susceptibility

lactamase inhibitor combinations have been effective in vitro and in

Severe infections:

See above

or tid IV

Vancomycin 1 g bid IV

Susceptibility of Isolate

Penicillin-susceptible

Penicillin-resistant, methicillin

Methicillin (oxacillin)-resistant

Oxacillin-resistant, with reduced

susceptibility to vancomycin

(oxacillin)-susceptible

above			
above			
	* 1.20 m		

Comments

Addition of an aminoglycoside (gentamicin 80-120 mg tid IV) advisable for endocarditis and of rifampin (600 mg qd orally) for prosthetic-valve endocarditis

asceptionity to vancomycin	or tid IV Linezolid 600 mg bid IV or orally	pattern Investigational drugs (?): LY 333328, SCH 27899, glycylcyclines
*Dosage must be adjusted in patients	with altered renal function treated with aminoglycos	ides or glyconeptides (or both).

clinafloxacin appears to possess the best activity against ciprofloxacin-resistant strains (Table 2).26 There is concern, however, that bacterial aminoglycosides not useful in this setting.^{8, 11}
Fluoroquinolones initially were active against MRSA, but soon after antimicrobials primarily active against difficult gram-positive cocci, such against gram-positive cocci are now (or will soon be) available, and therapy is desirable. Newer fluoroquinolones with enhanced combination with rifampin to prevent emergence of resistance during group. 4 Provided that a MRSA strain is susceptible to a fluoroquinolone, their introduction into clinical practice resistance emerged and spread resistance can emerge early and spread widely with use of these molethe United States, 10, 33 become resistant to ciprofloxacin in 70% to 80% of cases in Europe and primarily among Pseudomonas aeruginosa and MRSA.38 New enthusiasm has been generated by the development of potent and it is cross-resistant to the other agents of the pencilin-resistant pneumococci: streptogramins The latter has

activity

(quinupristin-dalfopristin), oxazolidinones (linezolid), glycopeptides (LY

isolates, glycopeptides and miscellaneous oral agents are alternatives in indicated for infections caused by penicillin-resistant, methicillin-suscepalternatively, narrow-spectrum cephalosporins, patients with allergy to β -lactams and mild infection. Penicillinase-resistant penicillins, such as nafcillin and oxacillin As in the case of penicillin-susceptible such as cetazolin,

The problem arises in patients who cannot tolerate glycopeptides, although many alternative agents are, in theory, available 51 β -Lactam- β although some European researchers have found a MIC of vancomycin similar in vitro and in vivo activity of vancomycin and pin is advisable because this combination has proved more effective.8 patients who do not respond initially to vancomycin, addition of rifam (1 mg/L) significantly higher than that of teicoplanin (0.5 mg/L)tions is more restricted. Glycopeptides are the drugs of choice, with aureus isolates regardless of their susceptibility to methicillin.59 The range of antimicrobials available for treatment of MRSA infec teicoplanin,78

Table 2. COMPARATIVE IN VITRO ACTIVITIES (MICs, IN mg(L)) OF NEWER FLUOROQUINOLONES AGAINST METHICILLIN-RESISTANT S. AUREUS BASED ON CIPROFLOXACIN SUSCEPTIBILITY

Agent	Ciprofloxacin- Susceptible Strains	Ciprofloxacin- Resistant Strains
Clinafloxacin	≤0.06	,
Moxifloxacin	≤0.06	2
Trovafloxacin	≤0.06	4
Grepafloxacin	≤0.06	32
Sparfloxacin	0.12	16
Levofloxacin	0.25	16
Ofloxacin	0.5	32
Ciprofloxacin	1	64

MIC₀₀ = Minimal inhibitory concentration that prevents growth of 90% of the strains.

Data from Jones ME, Vissen MK, Klootwijk M, et al: Comparative activities of clinafloxacin, grepafloxacin, tevofloxacin, moxifloxacin, ofloxacicn, sparfloxacin, and trovafloxacin and nonquinolones linezolid, quinupristin-dalfopristin, gentamicin, and vancomycin against clinical isolates of ciprofloxacin-resistant and—susceptible Staphylococcus aureus strains. Antimicrob Agents Chemother 43:421, 1999;

derivatives of pristinomycin IA and IIA, respectively, in a natural 30:70 ratio; it is active in vitro against MRSA, with MIC₉₀ of 0.25 to 2 mg/L^6 , bacteremia, nosocomial pneumonia, and skin and soft tissue infection. 43 and it has proved effective in the therapy of MRSA infections such as bination of quinupristin and dalfopristin (two semisynthetic bacteriostatic injectable streptogramin consisting of a synergistic and bactericidal comthetic tetracyclines (glycylcylines). Quinupristin-dalfopristin is a new 333328), everninomicin derivatives (SCH 27899), and newer semisyn-

active (MIC, <0.25 to 2 mg/L), but also bactericidal (minimal bactericidal concentration, 1 to 2 mg/L) against *S. aureus*, and it appears to the favorable possibility of switch therapy from intravenous to oral administration. IX 333328 is a glycopeptide antibiotic that is not only ity of a more recent derivative of minocycline, GAR-936, is 2 dilutions lower than that of CL 329,998 and CL 331,002 against MRSA, with considerably more active than minocycline and slightly more effective of minocycline and demethyldeoxytetracycline, which, respectively, are of 0.25 mg/L.28 CL 329,998 and CL 331,002 are semisynthetic derivatives everninomicin antibiotic that is highly active against MRSA, with MIC₉₀ of bacteremias and skin and soft tissue infections caused by MRSA, with evaluation. Oxazolidinones such as linezolid are synthetic bacteriostatic guidelines have been published until now. The first two U.S. isolates MIC₉₀ of 0.5 mg/L.* Regarding treatment of GISA infections, no definite than glycopeptides against MRSA, with MIC₉₀ of 0.25 mg/L, 62 The activbe bactericidal against GISA as well. 20 SCH 27899 is an oligosaccharide from clinical trials indicate the good efficacy of linezolid in the treatment higher MIC₉₀ (4 mg/L) than that of vancomycin.⁵⁵ Preliminary results antimicrobials that are active in vitro against MRSA, although with were susceptible to quinupristin-dalfopristin, tetracycline, chlorampheni-Other investigational drugs are in preclinical or clinical phase II-III

> a prominent role in this field. tional drugs such as quinupristin-dalfopristin and linezolid could have otic therapy of these difficult-to-treat infections must be guided by plus arbekacin (an aminoglycoside available in Japan,22 and the French was treated successfully with co-trimoxazole and rifampin),66 whereas col, and co-trimoxazole (one strain also to rifampin, and the infection laboratory tests; however, it is possible that the new potent investigapatient was treated successfully with quinupristin-dalfopristin). Antibithe infection in the Japanese patient responded to ampicillin-sulbactam

COAGULASE-NEGATIVE STAPHYLOCOCCI STAPHYLOCOCCUS EPIDERMIDIS AND OTHER

species being S. hominis. Other species are less frequent (S. haemolyticus, S. warneri), transient residents of the skin (S. xylosus, S. simulans, S. 65% to 90% of all staphylococci recovered, the second most frequent cohnii) or present in specific sites (S. auricolaris in the ear canal, S. capitis man skin. S. epidermidis is the most prevalent species, accounting for Coagulase-negative staphylococci are natural inhabitants of the hu-

tance to at least six antimicrobials, (2) *S. epidermidis* species and biotype, (3) more than one positive blood culture, and (4) mean colony count of 33.2 per milliliter of blood as determined by quantitative blood cultures.21 ered contaminants. It often is difficult to determine whether these organon the head, S. saprophyticus on the genitourinary skin).

For many years, S. epidermidis and other coagulose-negative staphylococci isolated from the blood of hospitalized patients have been consid-Factors that have been shown to predict a true bacteremia are (1) resisisms cause true bacteremia or they are only contaminants, however,

vascular access devices together with urinary, peritoneal, and ventricular indwelling devices in the management of a variety of life-threatening dermidis and other coagulase-negative staphylococci follows: community-acquired and hospital-acquired infections caused by S. epiand otherwise immunocompromised patients in hospitals.47 A list of in hospitalized patients; and (3) the high proportion of neutropenic and chronic diseases; (2) the increasing use of broad-spectrum antibiotdue to many factors, including (1) the widespread use of indwelling emergence of S. epidermidis as a major cause of nosocomial bacteremia is leading cause of nosocomial bloodstream infections (see Fig. 1).37 The cocci in nosocomial bacteremia increased from 9% in 1980 to 27% in ics, especially β-lactams, in the prevention and treatment of infections 1989,57 and coagulase-negative staphylococci have been found to be the In the United States, the incidence of coagulase-negative staphylo-

Urinary tract infections Female outpatients (S. saprophyticus) Hospital-acquired (S. epidermidis)

Osteomyelitis
Sternal wound
Hematogenous
Endocarditis of the native valve
Bacterenia in immunosuppressed patients
Endophthalmitis after ocular surgery
Infections of indwelling foreign devices
Intravenous catheters
Hemodialysis shunt and grafts
Peritomeal dialysis catheters
Pacemaker wires and electrodes
Prosthetic joints
Vascular grafts
Prosthetic cardiac valves
Breast implants

Epidemiology of β-Lactam Resistance

found to be oxacillin-resistant, but higher rates (80% to 85%) have been reported with S. epidermidis. In Finland, the incidence of methicillinclinical isolates of S. epidermidis.68 The mechanism of methicillin resis-1994, simultaneously with an increasing use of penicillinase-stable β-lactams. The EPIC study revealed a 70.1% rate of MRSE among ICU coagulase-negative staphylococci increased from 20% in 1980 to 60% in 3 decades because of the selective pressure of the increasing use of ally acquired S. epidermidis isolates.54 The prevalence of methicillin-resisand Europe.27,72 Resistance to methicillin is common among nosocomiceptible penicillins) has reached rates of 70% to 90% in the United States staphylococci to penicillin G (and consequently to all β-lactamase-susresistant S. epidermidis (MRSE) increased from 28% in 1983 to 77% in 1989,57 65% of coagulase-negative staphylococci blood isolates have been β-lactams. In the United States, the incidence of methicillin-resistant tant coagulase-negative staphylococci has increased rapidly over the last VRE have not been found in either species.70 tance of S. epidermidis is the same as S. aureus,8 and van genes typical of Currently, \(\beta\)-lactamase-mediated resistance of coagulase-negative

Epidemiology of Resistance to Other Antimicrobial Agents

Methicillin-resistant coagulase-negative staphylococci strains generally are cross-resistant to all β-lactams as well as to many other classes of antimicrobials, with wide differences linked to local variations in their use. Isolates from different geographic areas of the United States were shown to be 23% resistant to chloramphenicol; 35%, to tetracycline, 50%, to trimethoprim; 60%, to clindamycin; 61%, to gentamicin; and 75%, to

erythromycin.³ Figures of 31%, 39%, 60%, 30%, 71%, and 60% have been found in the Netherlands.¹¹

As in the case of *S. aureus*, fluoroquinolones initially were shown to be highly active against not only methicillin-susceptible *S. epidermidis* (MSSE), but also MRSE. Quinolone-resistant *S. epidermidis* strains emerged early in relation to the wide use of these antimicrobials in the community as well as in the hospital, however.³² For example, in Europe, more than 50% of methicillin-resistant coagulase-negative staphylococci are resistant to ciprofloxacin.³³ Among newer fluoroquinolones with enhanced activity against gram-positive cocci, clinafloxacin and moxifloxacin appear to possess the best activity, with MIC₉₀ of 0.06 to 1 mg/L.⁴

and 7% of *S. haemolyticus* and *S. epidermidis* isolates, respectively.²⁷ More recently, vancomycin was uniformly active against methicillin-resistant emia⁵⁰; these devices must be removed from patients with S. epidermidis embedded in biofilm has been attributed to poor penetration of glycoin suspension, and only 5% were tolerant, whereas 100% of the strains were tolerant in the biofilm environment.³⁰ Tolerance of S. epidermidis known as a high-risk factor for recurrence of catheter-related bacterserves as a barrier to antibiotic diffusion.14 Vascular catheter retention is vascular catheters were highly susceptible to vancomycin when tested ated with device-related infections produce a slimy material, structurally irregularity of the cell wall or to the increased capacity of the cell wall to bind and sequester glycopeptides. 5 epidermidis organisms associcoagulase-negative staphylococci, whereas full resistance to teicoplanin peptides through the biofilm matrix, suggesting that the slime itself has been reported that 134 S. epidermidis isolates from tips of indwelling isms become tolerant to it when tested in vitro as dispersed bacteria. It biofilm.9 In a biofilm environment, highly vancomycin-susceptible organbacteria to smooth surfaces of prosthetic devices and forms a microbial defined as an exopolysaccharide (slime), which facilitates adherence of known, although resistance has been attributed to the thickening and ity to teicoplanin (MIC, 16 mg/L) was found in the United States in 21% haemolyticus and, to a lesser extent, S. epidermidis.59 Reduced susceptibilvancomycin against coagulase-negative staphylococci, in particular, S. The mechanism of glycopeptide resistance in S. epidermidis still is un-(MIC, \geq 32 mg/L) was observed in 1.3% to 8.6% of strains worldwide.²⁵ In contrast to S. aureus, teicoplanin is significantly less active than

Antimicrobial Agents for Treatment of *Staphylococcus* epidermidis and Other Coagulase-Negative Staphylococcal Infections

Choice of drug therapy for infections resulting from coagulase-negative staphylococci depends on several factors, including the severity of the infection, underlying disease condition, infecting pathogen, and presence of resistant strains. Recommendations for treatment of *S. aureus*

mial and device-related infections caused by MRSE and other coagulase-negative staphylococci has been described; in these cases, alternative infections summarized in Table 1 generally can be applied to the therapy of coagulase-negative staphylococcal infections. Because of the aboveagents, such as rifampin, minocycline, and others, must be considered. 47 Glycopeptide failure even with vancomycin in the treatment of nosocomycin is the drug of choice when a glycopeptide agent is required mentioned higher activity of vancomycin than that of teicoplanin, vanco-

reported, however, when rifampin is used not only in monotherapy but also in combination with nonlipophilic drugs active on the cell wall (β-lactams, glycopeptides). Epidermidis strains resistant to minocycline, the most active tetracycline against methicillin-resistant staphylococci, have been reported. The combination of minocycline and rifampin is glycopeptides, it improves their activity against *S. epidermidis* embedded in biofilm.⁴² The emergence of rifampin-resistant *S. epidermidis* has been of many drugs, including vancomycin. When used in combination with of increasing vancomycin resistance, however, there is now a clinical need for new types of antibiotics for the treatment of staphylococcal venting catheter-related bacteremia caused by S. epidermidis in highstaphylococci.49 In a multicenter randomized trial, minocycline and riin preventing the colonization of catheter surfaces with slime-producing more effective than vancomycin alone or in combination with rifampin infections. potential activity against methicillin-resistant staphylococci. 13, 51 Because risk patients.48 Chloramphenicol, imipenem, and ciprofloxacin all have fampin-coated central venous catheters were highly effective in pre-Antistaphylococcal in vitro activity of rifampin is superior to that

of linezolid in coagulase-negative staphylococcal infections of different sites show a positive clinical response in 10 of 11 patients. As in the case of S. aureus, the glycopeptide antibiotic LY 333328 is not only active (MIC, 0.5 to 2 mg/L), but also bactericidal (minimal bactericidal) Quinupristin-dalfopristin has good bactericidal activity against MSSE and MRSE, with MICs $_{90}$ of 0.25 to 1 mg/L 6 , and it has obtained susceptible and methicillin-resistant, coagulase-negative staphylococci concentration, 1 to 2 mg/L) against coagulase-negative staphylococci. ¹⁹ The everninomicin SCH 27899 is uniformly active against methicillinwith MIC_{∞} of 2 gm/L⁵⁵; it is possible that this new class of antimicrobials will be a valid alternative to glycopeptides in the treatment of systemic infections resulting from *S. epidermidis*. Preliminary results on the efficacy option in patients allergic to glycopeptides or receiving concomitant nephrotoxic drugs. The in vitro activity of oxazolidinones such as linezolid against MSSE and MRSE is comparable with that of vancomycin, negative staphylococci.43 Quinupristin-dalfopristin can be a therapeutic and soft tissue infections resulting from MRSE and other coagulasepreliminary studies on the treatment of bloodstream infections and skin satisfactory results in terms of clinical and microbiologic efficacy in with MIC₉₀ of 0.25 to 0.5 mg/L.28 Glycylcyclines CL 329, 998 and CL 331,002 show a similar activity to those of minocycline and vancomycin

against methicillin-resistant coagulase-negative staphylococci, with MIC $_{90}$ of 2 mg/L. 52 The activity of GAR-936 is 2 to 3 dilutions lower than that of CL 329,998 and CL 331,002 against MRSE, with MIC $_{90}$ of

SUMMARY

duced or experimental drugs, such as streptogramins (quinupristin-dalfopristin), oxazolidinones (linezolid), carbapenems (LY 333328), everninomicins (SCH 27899), and derivatives of tetracyclines (glycylcydifficult by the increasing emergence of resistance to β -lactams and other antimicrobials, including reduced susceptibility to glycopeptides. are important causes of infection of the bloodstream, cardiac valves, implanted devices, and skin, with repercussions on mortality and increased economic costs. Treatment of staphylococcal infections is made clines), could be useful for therapy of infections caused by multiresistan native agents, primarily in subjects allergic to β-lactams. Newly introlin-resistant staphylococci, and glycopeptides are the drugs of choice for infections caused by methicillin-resistant strains. Co-trimoxazole, lincosamides, macrolides, tetracyclines, and fluoroquinolones are alteroxacillin and nafcillin are to be considered the major option for penicil-Penicillin must be used for infrequent penicillin-susceptible isolates, S. aureus and coagulase-negative staphylococci such as S. epidermidis

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