Prevention of Infective Endocarditis. Guidelines From the American Heart Association. A Guideline From the American Heart Association Rheumatic Fever, Endocarditis, and Kawasaki Disease Committee, Council on Cardiovascular Disease in the Young, and the Council on Clinical Cardiology, Council on Cardiovascular Surgery and Anesthesia, and the Quality of Care and Outcomes Research Interdisciplinary Working Group


Circulation published online Apr 19, 2007; DOI: 10.1161/CIRCULATIONAHA.106.183095

Circulation is published by the American Heart Association. 7272 Greenville Avenue, Dallas, TX 75231
Copyright © 2007 American Heart Association. All rights reserved. Print ISSN: 0009-7322. Online ISSN: 1524-4539

The online version of this article, along with updated information and services, is located on the World Wide Web at:
http://circ.ahajournals.org

Subscriptions: Information about subscribing to Circulation is online at http://circ.ahajournals.org/subscriptions/

Permissions: Permissions & Rights Desk, Lippincott Williams & Wilkins, a division of Wolters Kluwer Health, 351 West Camden Street, Baltimore, MD 21202-2436. Phone: 410-528-4050. Fax: 410-528-8550. E-mail: journalpermissions@lww.com

Reprints: Information about reprints can be found online at http://www.lww.com/reprints
Prevention of Infective Endocarditis
Guidelines From the American Heart Association

Walter Wilson, MD, Chair; Kathryn A. Taubert, PhD, FAHA; Michael Gewitz, MD, FAHA; Peter B. Lockhart, DDS; Larry M. Baddour, MD; Matthew Levison, MD; Ann Bolger, MD, FAHA; Christopher H. Cabell, MD, MHS; Masato Takahashi, MD, FAHA; Robert S. Baltimore, MD; Jane W. Newburger, MD, MPH, FAHA; Buyan L. Strom, MD; Lloyd Y. Tani, MD; Michael Gerber, MD; Robert O. Bonow, MD, FAHA; Thomas Pallasch, DDS, MS; Stanford T. Shulman, MD, FAHA; Anne H. Rowley, MD; Jane C. Burns, MD; Patricia Ferrieri, MD; Timothy Gardner, MD, FAHA; David Goff, MD, PhD, FAHA; David T. Durack, MD, PhD

The Council on Scientific Affairs of the American Dental Association has approved the guideline as it relates to dentistry. In addition, this guideline has been endorsed by the Infectious Diseases Society of America and by the Pediatric Infectious Diseases Society.

Background—The purpose of this statement is to update the recommendations by the American Heart Association (AHA) for the prevention of infective endocarditis that were last published in 1997.

Methods and Results—A writing group was appointed by the AHA for their expertise in prevention and treatment of infective endocarditis, with liaison members representing the American Dental Association, the Infectious Diseases Society of America, and the American Academy of Pediatrics. The writing group reviewed input from national and international experts on infective endocarditis. The recommendations in this document reflect analyses of relevant literature regarding procedure-related bacteremia and infective endocarditis, in vitro susceptibility data of the most common microorganisms that cause infective endocarditis, results of prophylactic studies in animal models of experimental endocarditis, and retrospective and prospective studies of prevention of infective endocarditis. MEDLINE database searches from 1950 to 2006 were done for English-language papers using the following search terms: endocarditis, infective endocarditis, prophylaxis, prevention, antibiotic, antimicrobial, pathogens, organisms, dental, gastrointestinal, genitourinary, streptococcus, enterococcus, staphylococcus, respiratory, dental surgery, pathogenesis, vaccine, immunization, and bacteremia. The reference lists of the identified papers were also searched. We also searched the AHA online library. The American College of Cardiology/AHA classification of recommendations and levels of evidence for practice guidelines were used. The paper was subsequently reviewed by outside experts not affiliated with the writing group and by the AHA Science Advisory and Coordinating Committee.

Conclusions—The major changes in the updated recommendations include the following: (1) The Committee concluded that only an extremely small number of cases of infective endocarditis might be prevented by antibiotic prophylaxis for...
dental procedures even if such prophylactic therapy were 100% effective. (2) Infective endocarditis prophylaxis for dental procedures should be recommended only for patients with underlying cardiac conditions associated with the highest risk of adverse outcome from infective endocarditis. (3) For patients with these underlying cardiac conditions, prophylaxis is recommended for all dental procedures that involve manipulation of gingival tissue or the periapical region of teeth or perforation of the oral mucosa. (4) Prophylaxis is not recommended based solely on an increased lifetime risk of acquisition of infective endocarditis. (5) Administration of antibiotics solely to prevent endocarditis is not recommended for patients who undergo a genitourinary or gastrointestinal tract procedure. These changes are intended to define more clearly when infective endocarditis prophylaxis is or is not recommended and to provide more uniform and consistent global recommendations. (Circulation. 2007;115;1645–1668)

Key Words: AHA Scientific Statements • cardiovascular diseases • endocarditis • prevention • antibiotic prophylaxis

Infective endocarditis (IE) is an uncommon but life-threatening infection. Despite advances in diagnosis, antimicrobial therapy, surgical techniques, and management of complications, patients with IE still have high morbidity and mortality rates related to this condition. Since the last American Heart Association (AHA) publication on prevention of IE in 1997, many authorities and societies, as well as the conclusions of published studies, have questioned the efficacy of antimicrobial prophylaxis to prevent IE in patients who undergo a dental, gastrointestinal (GI), or genitourinary (GU) tract procedure and have suggested that the AHA guidelines should be revised. Members of the Rheumatic Fever, Endocarditis, and Kawasaki Disease Committee of the AHA Council on Cardiovascular Disease in the Young (“the Committee”) and a national and international group of experts on IE extensively reviewed data published on the prevention of IE. The Committee is especially grateful to a group of international experts on IE who provided content review and input on this document (see Acknowledgments). The revised guidelines for IE prophylaxis are the subject of this report.

The writing group was charged with the task of performing an assessment of the evidence and giving a classification of recommendations and a level of evidence (LOE) to each recommendation. The American College of Cardiology (ACC)/AHA classification system was used as follows.

Classification of Recommendations:
Class I: Conditions for which there is evidence and/or general agreement that a given procedure or treatment is beneficial, useful, and effective.
Class II: Conditions for which there is conflicting evidence and/or a divergence of opinion about the usefulness/efficacy of a procedure or treatment.
Class IIa: Weight of evidence/opinion is in favor of usefulness/efficacy.
Class IIb: Usefulness/efficacy is less well established by evidence/opinion.
Class III: Conditions for which there is evidence and/or general agreement that a procedure/treatment is not useful/effective and in some cases may be harmful.

Level of Evidence:
Level of Evidence A: Data derived from multiple randomized clinical trials or meta-analyses.
Level of Evidence B: Data derived from a single randomized trial or nonrandomized studies.

Level of Evidence C: Only consensus opinion of experts, case studies, or standard of care.

History of AHA Statements on Prevention of IE

The AHA has made recommendations for the prevention of IE for more than 50 years. In 1955, the first AHA document on this subject was published in Circulation. Table 1 shows a summary of the documents published from 1955 to 1997. The 1960 document called attention to the possible emergence of penicillin-resistant oral microflora as a result of prolonged therapy for prevention of IE, and pediatric patients were included for the first time. Chloramphenicol was recommended for patients who were allergic to penicillin. In 1965, the Committee published for the first time a document devoted solely to the prophylaxis of IE and recognized the importance of enterococci after GI or GU tract procedures. The revised recommendations published in 1972 were endorsed for the first time by the American Dental Association (ADA) and emphasized the importance of maintenance of good oral hygiene. This version introduced a recommendation for ampicillin in patients undergoing a GI or GU tract procedure. The 1977 revisions categorized both patients and procedures into high- and low-risk groups. This resulted in complex tables with many footnotes. The 1984 recommendations attempted to simplify prophylactic regimens by providing clear lists of procedures for which prophylaxis was and was not recommended and reduced postprocedure prophylaxis for dental, GI, and GU tract procedures to only 1 oral or parenteral dose. In 1990, a more complete list of cardiac conditions and dental or surgical procedures for which prophylaxis was and was not recommended was provided. These previous recommendations recognized the potential medical-legal risks associated with IE prophylaxis and suggested that the recommendations were intended to serve as a guideline, not as established standard of care. The most recent AHA document on IE prophylaxis was published in 1997. The 1997 document stratified cardiac conditions into high-, moderate-, and low-risk (negligible risk) categories, with prophylaxis not recommended for the low-risk group. An even more detailed list of dental, respiratory, GI, and GU tract procedures for which prophylaxis was and was not recommended was provided. The 1997 document was notable for its acknowledgment that most cases of IE are not attributable to an invasive procedure but corresponds to the classification of recommendations and the level of evidence.
TABLE 1. Summary of 9 Iterations of AHA-Recommended Antibiotic Regimens From 1955 to 1997 for Dental/Respiratory Tract Procedures*

<table>
<thead>
<tr>
<th>Year (Reference)</th>
<th>Primary Regimens for Dental Procedures</th>
</tr>
</thead>
<tbody>
<tr>
<td>1955 (6)</td>
<td>Aqueous penicillin 600 000 U and procaine penicillin 600 000 U in oil containing 2% aluminum monostearate administered IM 30 minutes before the operative procedure</td>
</tr>
<tr>
<td>1957 (7)</td>
<td>For 2 days before surgery, penicillin 200 000 to 250 000 U by mouth 4 times per day. On day of surgery, penicillin 200 000 to 250 000 U by mouth 4 times per day and aqueous penicillin 600 000 U with procaine penicillin 600 000 U IM 30 to 60 minutes before surgery. For 2 days after, 200 000 to 250 000 U by mouth 4 times per day.</td>
</tr>
<tr>
<td>1960 (8)</td>
<td>Step I: prophylaxis 2 days before surgery with procaine penicillin 600 000 U IM on each day</td>
</tr>
<tr>
<td></td>
<td>Step II: day of surgery: procaine penicillin 600 000 U IM supplemented by crystalline penicillin 600 000 U IM 1 hour before surgical procedure</td>
</tr>
<tr>
<td></td>
<td>Step III: for 2 days after surgery: procaine penicillin 600 000 U IM each day</td>
</tr>
<tr>
<td>1965 (9)</td>
<td>Day of procedure: procaine penicillin 600 000 U, supplemented by crystalline penicillin 600 000 U IM 1 to 2 hours before the procedure</td>
</tr>
<tr>
<td>1972 (10)</td>
<td>Procaine penicillin G 600 000 U mixed with crystalline penicillin G 200 000 U IM 1 hour before procedure and once daily for the 2 days after the procedure</td>
</tr>
<tr>
<td>1977 (11)</td>
<td>Aqueous crystalline penicillin G (1 000 000 U IM) mixed with procaine penicillin G (600 000 U IM) 30 minutes to 1 hour before procedure and then penicillin V 500 mg orally every 6 hours for 8 doses.</td>
</tr>
<tr>
<td>1984 (12)</td>
<td>Penicillin V 2 g orally 1 hour before, then 1 g 6 hours after initial dose</td>
</tr>
<tr>
<td>1990 (13)</td>
<td>Amoxicillin 3 g orally 1 hour before procedure, then 1.5 g 6 hours after initial dose</td>
</tr>
<tr>
<td>1997 (1)</td>
<td>Amoxicillin 2 g orally 1 hour before procedure</td>
</tr>
</tbody>
</table>

IM indicates intramuscularly.
*These regimens were for adults and represented the initial regimen listed in each version of the recommendations. In some versions, >1 regimen was included.

rather are the result of randomly occurring bacteremias from routine daily activities and for its acknowledgment of possible IE prophylaxis failures.

Rationale for Revising the 1997 Document
It is clear from the above chronology that the AHA guidelines for IE prophylaxis have been in a process of evolution more than 50 years. The rationale for prophylaxis was based largely on expert opinion and what seemed to be a rational and prudent attempt to prevent a life-threatening infection. On the basis of the ACC and AHA Task Force on Practice Guidelines’ evidence-based grading system for ranking recommendations, the recommendations in the AHA documents published during the past 50 years would be Class IIb, LOE C. Accordingly, the basis for recommendations for IE prophylaxis was not well established, and the quality of evidence was limited to a few case-control studies or was based on expert opinion, clinical experience, and descriptive studies that utilized surrogate measures of risk.

Over the years, other international societies have published recommendations and guidelines for the prevention of IE.14,15 Recently, the British Society for Antimicrobial Chemotherapy issued new IE prophylaxis recommendations.15 This group now recommends prophylaxis before dental procedures only for patients who have a history of previous IE or who have had cardiac valve replacement or surgically constructed pulmonary shunts or conduits.

The fundamental underlying principles that drove the formulation of the AHA guidelines and the 9 previous AHA documents were that (1) IE is an uncommon but life-threatening disease, and prevention is preferable to treatment of established infection; (2) certain underlying cardiac conditions predispose to IE; (3) bacteremia with organisms known to cause IE occurs commonly in association with invasive dental, GI, or GU tract procedures; (4) antimicrobial prophylaxis was proven to be effective for prevention of experimental IE in animals; and (5) antimicrobial prophylaxis was thought to be effective in humans for prevention of IE associated with dental, GI, or GU tract procedures. The Committee believes that of these 5 underlying principles, the first 4 are valid and have not changed during the past 30 years. Numerous publications have questioned the validity of the fifth principle and suggested revision of the guidelines, primarily for reasons as shown in Table 2.

Another reason that led the Committee to revise the 1997 document was that over the past 50 years, the AHA guidelines on prevention of IE became overly complicated, making it difficult for patients and healthcare providers to interpret or remember specific details, and they contained ambiguities and some inconsistencies in the recommendations. The decision to substantially revise the 1997 document was not taken lightly. The present revised document was not based on the results of a single study but rather on the collective body of

TABLE 2. Primary Reasons for Revision of the IE Prophylaxis Guidelines

IE is much more likely to result from frequent exposure to random bacteremias associated with daily activities than from bacteremia caused by a dental, GI tract, or GU tract procedure.

Prophylaxis may prevent an exceedingly small number of cases of IE, if any, in individuals who undergo a dental, GI tract, or GU tract procedure.

The risk of antibiotic-associated adverse events exceeds the benefit, if any, from prophylactic antibiotic therapy.

Maintenance of optimal oral health and hygiene may reduce the incidence of bacteremia from daily activities and is more important than prophylactic antibiotics for a dental procedure to reduce the risk of IE.
evidence published in numerous studies over the past 2 decades. The Committee sought to construct the present recommendations such that they would be in the best interest of patients and providers, would be reasonable and prudent, and would represent the conclusions of published studies and the collective wisdom of many experts on IE and relevant national and international societies.

**Potential Consequences of Substantive Changes in Recommendations**

Substantive changes in recommendations could (1) violate long-standing expectations and practice patterns; (2) make fewer patients eligible for IE prophylaxis; (3) reduce malpractice claims related to IE prophylaxis; and (4) stimulate prospective studies on IE prophylaxis. The Committee and others recognize that substantive changes in IE prophylaxis guidelines may violate long-standing expectations and practice patterns by patients and healthcare providers. The Committee recognizes that these new recommendations may cause concern among patients who have previously received antibiotic prophylaxis to prevent IE before dental or other procedures and are now advised that such prophylaxis is unnecessary. Table 2 includes the main talking points that may be helpful for clinicians in reeducating their patients about these changes. To recommend such changes demands due diligence and critical analysis. For 50 years, since the publication of the first AHA guidelines on the prevention of IE, patients and healthcare providers assumed that antibiotics administered in association with a bacteremia-producing procedure effectively prevented IE in patients with underlying cardiac risk factors. Patients were educated about bacteremia-producing procedures and risk factors for IE, and they expected to receive antibiotic prophylaxis; healthcare providers, especially dentists, were expected to administer them. Patients with underlying cardiac conditions that carry a lifetime risk of acquisition of IE, such as mitral valve prolapse (MVP), had a sense of reassurance and comfort that antibiotics administered in association with a dental procedure were effective and usually safe to prevent IE. Healthcare providers, especially dentists, felt a sense of obligation and professional and legal responsibility to protect their patients from IE that might result from a procedure. On the basis of recommendations in this revised document, substantially fewer patients will be recommended for IE prophylaxis.

Cases of IE either temporally or remotely associated with an invasive procedure, especially a dental procedure, have frequently been the basis for malpractice claims against healthcare providers. Unlike many other infections for which there is conclusive evidence for the efficacy of preventive therapy, the prevention of IE is not a precise science. Because previously published AHA guidelines for the prevention of IE contained ambiguities and inconsistencies and were often based on minimal published data or expert opinion, they were subject to conflicting interpretations among patients, healthcare providers, and the legal system about patient eligibility for prophylaxis and whether there was strict adherence by healthcare providers to AHA recommendations for prophylaxis. This document is intended to identify which, if any, patients may possibly benefit from IE prophylaxis and to define, to the extent possible, which dental procedures should have prophylaxis in this select group of patients. Accordingly, the Committee hopes that this document will result in greater clarity for patients, healthcare providers, and consulting professionals.

The Committee believes that recommendations for IE prophylaxis must be evidence based. A placebo-controlled, multicenter, randomized, double-blinded study to evaluate the efficacy of IE prophylaxis in patients who undergo a dental, GI, or GU tract procedure has not been done. Such a study would require a large number of patients per treatment group and standardization of the specific invasive procedures and the patient populations. This type of study would be necessary to definitively answer long-standing unresolved questions regarding the efficacy of IE prophylaxis. The Committee hopes that this revised document will stimulate additional studies on the prevention of IE. Future published data will be reviewed carefully by the AHA, the Committee on Rheumatic Fever, Endocarditis, and Kawasaki Disease, and other societies, and further revisions to the present document will be based on relevant studies.

**Pathogenesis of IE**

The development of IE is the net result of the complex interaction between the bloodstream pathogen with matrix molecules and platelets at sites of endocardial cell damage. In addition, many of the clinical manifestations of IE emanate from the host’s immune response to the infecting microorganism. The following sequence of events is thought to result in IE: formation of nonbacterial thrombotic endocarditis (NBTE) on the surface of a cardiac valve or elsewhere that endothelial damage occurs, bacteremia, adherence of the bacteria in the bloodstream to NBTE, and proliferation of bacteria within a vegetation.

**Formation of NBTE**

Turbulent blood flow produced by certain types of congenital or acquired heart disease, such as flow from a high- to a low-pressure chamber or across a narrowed orifice, traumatizes the endothelium. This creates a predisposition for deposition of platelets and fibrin on the surface of the endothelium, which results in NBTE. Invasion of the bloodstream with a microbial species that has the pathogenic potential to colonize this site can then result in IE.

**Transmucosal Bacteremia**

Mucosal surfaces are populated by a dense endogenous microflora. Trauma to a mucosal surface, particularly the gingival crevice around teeth, oropharynx, GI tract, urethra, and vagina, releases many different microbial species transiently into the bloodstream. Transient bacteremia caused by viridans group streptococci and other oral microflora occurs commonly in association with dental extractions or other dental procedures or with routine daily activities. Although controversial, the frequency and intensity of the resulting bacteremias are believed to be related to the nature and magnitude of the tissue trauma, the density of the microbial flora, and the degree of inflammation or infection at the site of trauma. The microbial species entering the circulation
depends on the unique endogenous microflora that colonizes the particular traumatized site.

**Bacterial Adherence**

The ability of various microbial species to adhere to specific sites determines the anatomic localization of infection caused by these microorganisms. Mediators of bacterial adherence serve as virulence factors in the pathogenesis of IE. Numerous bacterial surface components present in streptococci, staphylococci, and enterococci have been shown in animal models of experimental endocarditis to function as critical adhesins. Some viridans group streptococci contain a FimA protein that is a lipoprotein receptor antigen I (LraI) that serves as a major adhesin to the fibrin platelet matrix of NBTE. Staphylococcal adhesins function in at least 2 ways. In one, microbial surface components recognizing adhesive matrix molecules facilitate the attachment of staphylococci to human extracellular matrix proteins and to medical devices that become coated with matrix proteins after implantation. In the other, bacterial extracellular structures contribute to the formation of biofilm that forms on the surface of implanted medical devices. In both cases, staphylococcal adhesins are important virulence factors.

Both FimA and staphylococcal adhesins are immunogenic in experimental infections. Vaccines prepared against FimA and staphylococcal adhesins provide some protective effect in experimental endocarditis caused by viridans group streptococci and staphylococci. The results of these experimental studies are highly intriguing, because the development of an effective vaccine for use in humans to prevent viridans group streptococcal or staphylococcal IE would be of major importance.

**Proliferation of Bacteria Within a Vegetation**

Microorganisms adherent to the vegetation stimulate further deposition of fibrin and platelets on their surface. Within this secluded focus, the buried microorganisms multiply as rapidly as bacteria in broth cultures to reach maximal microbial densities of $10^7$ to $10^{10}$ colony-forming units per gram of vegetation within a short time on the left side of the heart, apparently uninhibited by host defenses in left-sided lesions. Right-sided vegetations have lower bacterial densities, which may be the consequence of host defense mechanisms active at this site, such as polymorphonuclear activity or platelet-derived antibacterial proteins. More than 90% of the microorganisms in mature left- or right-sided valvular vegetations are metabolically inactive rather than in an active growth phase and are therefore less responsive to the bactericidal effects of antibiotics.

**Rationale for or Against Prophylaxis of IE**

**Historical Background**

Viridans group streptococci are part of the normal skin, oral, respiratory, and GI tract flora, and they cause at least 50% of cases of community-acquired native valve IE not associated with intravenous drug use. More than a century ago, the oral cavity was recognized as a potential source of the bacteremia that caused viridans group streptococcal IE. In 1885, Osler noted an association between bacteremia from surgery and IE. Okell and Elliott in 1935 reported that 11% of patients with poor oral hygiene had positive blood cultures with viridans group streptococci and that 61% of patients had viridans group streptococcal bacteremia with dental extraction.

As a result of these early studies and subsequent studies, during the past 50 years, the AHA guidelines recommended antimicrobial prophylaxis to prevent IE in patients with underlying cardiac conditions who underwent bacteremia-producing procedures on the basis of the following factors: (1) bacteremia causes endocarditis; (2) viridans group streptococci are part of the normal oral flora, and enterococci are part of the normal GI and GU tract flora; (3) these microorganisms are usually susceptible to antibiotics recommended for prophylaxis; (4) antibiotic prophylaxis prevents viridans group streptococcal or enterococcal experimental endocarditis in animals; (5) a large number of poorly documented case reports implicated a dental procedure as a cause of IE; (6) in some cases, there was a temporal relationship between a dental procedure and the onset of symptoms of IE; (7) an awareness of bacteremia caused by viridans group streptococci associated with a dental procedure exists; (8) the risk of significant adverse reactions to an antibiotic is low in an individual patient; and (9) morbidity and mortality from IE are high. Most of these factors remain valid, but collectively, they do not compensate for the lack of published data that demonstrate a benefit from prophylaxis.

**Bacteremia-Producing Dental Procedures**

The large majority of published studies have focused on dental procedures as a cause of IE and the use of prophylactic antibiotics to prevent IE in patients at risk. Few data exist on the risk of or prevention of IE associated with a GI or GU tract procedure. Accordingly, the Committee undertook a critical analysis of published data in the context of the historical rationale for recommending antibiotic prophylaxis for IE before a dental procedure. The following factors were considered: (1) frequency, nature, magnitude, and duration of bacteremia associated with dental procedures; (2) impact of dental disease, oral hygiene, and type of dental procedure on bacteremia; (3) impact of antibiotic prophylaxis on bacteremia from a dental procedure; and (4) the exposure over time of frequently occurring bacteremia from routine daily activities compared with bacteremia from various dental procedures.

**Frequency, Nature, Magnitude, and Duration of Bacteremia-Producing Dental Procedures**

Transient bacteremia is common with manipulation of the teeth and periodontal tissues, and there is a wide variation in reported frequencies of bacteremia in patients resulting from dental procedures: tooth extraction (10% to 100%), periodontal surgery (36% to 88%), scaling and root planing (8% to 80%), teeth cleaning (up to 40%), rubber dam matrix/wedge placement (9% to 32%), and endodontic procedures (up to 20%). Transient bacteremia also occurs frequently during routine daily activities unrelated to a dental procedure, such as tooth brushing and flossing (20% to 68%), use of wooden toothpicks (20% to 40%), use of water irrigation devices (7%...
to 50%), and chewing food (7% to 51%).36–39 Considering that the average person living in the United States has fewer than 2 dental visits per year, the frequency of bacteremia from routine daily activities is far greater.

There has been a disproportionate focus on the frequency of bacteremia associated with dental procedures rather than on the species of bacteria recovered from blood cultures. Studies suggest that more than 700 species of bacteria, including aerobic and anaerobic Gram-positive and Gram-negative microorganisms, may be identified in the human mouth, particularly on the teeth and in the gingival crevices.24,37–40 Approximately 30% of the flora of the gingival crevice is streptococci, predominantly of the viridans group. Of the more than 100 oral bacterial species recovered from blood cultures after dental procedures, the most prevalent are viridans group streptococci, the most common microbiological cause of community-acquired native valve IE in non–intravenous drug users.21 In healthy mouths, a thin surface of mucosal epithelium prevents potentially pathogenic bacteria from entering the bloodstream and lymphatic system. Anaerobic microorganisms are commonly responsible for periodontal disease and frequently enter the bloodstream but rarely cause IE, with fewer than 120 cases reported.31 Viridans group streptococci are antagonistic to periodontal pathogens and predominate in a clean, healthy mouth.42

Few published studies exist on the magnitude of bacteremia after a dental procedure or from routine daily activities, and most of the published data used older, often unreliable microbiological methodology. There are no published data that demonstrate that a greater magnitude of bacteremia, compared with a lower magnitude, is more likely to cause IE in humans. The magnitude of bacteremia resulting from a dental procedure is relatively low (<10^5 colony-forming units of bacteria per milliliter), similar to that resulting from routine daily activities, and is less than that used to cause experimental IE in animals (10^6 to 10^7 colony-forming units of bacteria per milliliter).20,43–44 Although the infective dose required to cause IE in humans is unknown, the number of microorganisms present in blood after a dental procedure or associated with daily activities is low. Cases of IE caused by oral bacteria probably result from the exposures to low inocula of bacteria in the bloodstream that result from routine daily activities and not from a dental procedure. Additionally, the vast majority of patients with IE have not had a dental procedure within 2 weeks before the onset of symptoms of IE.2–4

The role of duration of bacteremia on the risk of acquisition of IE is uncertain.45,46 Early studies reported that sequential blood cultures were positive for up to 10 minutes after tooth extraction and that the number of positive blood cultures dropped sharply after 10 to 30 minutes.24,45–51 More recent studies support these data but report a small percentage of positive blood cultures from 30 to 60 minutes after tooth extraction.43,52,53 Intuitively, it seems logical to assume that the longer the duration of bacteremia, the greater the risk of IE, but no published studies support this assumption. Given the preponderance of published data, there may not be a clinically significant difference in the frequency, nature, magnitude, and duration of bacteremia associated with a dental procedure compared with that resulting from routine daily activities. Accordingly, it is inconsistent to recommend prophylaxis of IE for dental procedures but not for these same patients during routine daily activities. Such a recommendation for prophylaxis for routine daily activities would be impractical and unwarranted.

**Impact of Dental Disease, Oral Hygiene, and Type of Dental Procedure on Bacteremia**

It is assumed that a relationship exists between poor oral hygiene, the extent of dental and periodontal disease, the type of dental procedure, and the frequency, nature, magnitude, and duration of bacteremia, but the presumed relationship is controversial.23,29,38,45,54–61 Nevertheless, available evidence supports an emphasis on maintaining good oral hygiene and eradicating dental disease to decrease the frequency of bacteremia from routine daily activities.45,56–58,62,63 In patients with poor oral hygiene, the frequency of positive blood cultures just before dental extraction may be similar to that after extraction.62,63

More than 80 years ago, it was suggested that poor oral hygiene and dental disease were more important as a cause of IE than were dental procedures.64 Most studies since that time have focused instead on the risks of bacteremia associated with dental procedures. For example, tooth extraction is thought to be the dental procedure most likely to cause bacteremia, with an incidence ranging from 10% to 100%.6 However, numerous other dental procedures have been reported to be associated with risks of bacteremia that are similar to that resulting from tooth extraction.† A precise determination of the relative risk of bacteremia that results from a specific dental procedure in patients with or without dental disease is probably not possible.27,72,73

Bleeding often occurs during a dental procedure in patients with or without periodontal disease. Previous AHA guidelines recommended antibiotic prophylaxis for dental procedures in which bleeding was anticipated but not for procedures in which bleeding was not anticipated.1 However, no data show that visible bleeding during a dental procedure is a reliable predictor of bacteremia.62 These ambiguities in the previous AHA guidelines led to further uncertainties among healthcare providers about which dental procedures should be covered by prophylaxis.

These factors complicated recommendations in previous AHA guidelines on prevention of IE that suggested antibiotic prophylaxis for some dental procedures but not for others. The collective published data suggest that the vast majority of dental office visits result in some degree of bacteremia; however, there is no evidence-based method to decide which procedures should require prophylaxis, because no data show that the incidence, magnitude, or duration of bacteremia from any dental procedure increase the risk of IE. Accordingly, it is not clear which dental procedures are more or less likely to cause a transient bacteremia or result in a greater magnitude of bacteremia than that which results from routine daily activities such as chewing food, tooth brushing, or flossing.

---

†References 23, 27, 29, 45, 48, 52, 54, 57, and 65–67.
†References 27, 28, 47, 51, 54, 56, 58, and 68–71.
In patients with underlying cardiac conditions, lifelong antibiotic therapy is not recommended to prevent IE that might result from bacteremias associated with routine daily activities. In patients with dental disease, the focus on the frequency of bacteremia associated with a specific dental procedure and the AHA guidelines for prevention of IE have resulted in an overemphasis on antibiotic prophylaxis and an underemphasis on maintenance of good oral hygiene and access to routine dental care, which are likely more important in reducing the lifetime risk of IE than the administration of antibiotic prophylaxis for a dental procedure. However, no observational or controlled studies support this contention.

**Impact of Antibiotic Therapy on Bacteremia From a Dental Procedure**

The ability of antibiotic therapy to prevent or reduce the frequency, magnitude, or duration of bacteremia associated with a dental procedure is controversial. Some studies reported that antibiotics administered before a dental procedure reduced the frequency, nature, and/or duration of bacteremia, whereas others did not. Recent studies suggest that amoxicillin therapy has a statistically significant impact on reducing the incidence, nature, and duration of bacteremia from dental procedures, but it does not eliminate bacteremia. However, no data show that such a reduction as a result of amoxicillin therapy reduces the risk of or prevents IE. Hall et al reported that neither penicillin V nor amoxicillin therapy was effective in reducing the frequency of bacteremia compared with untreated control subjects. In patients who underwent a dental extraction, penicillin or ampicillin therapy compared with placebo diminished the percentage of viridans group streptococci and anaerobes in culture, but there was no significant difference in the percentage of patients with positive cultures 10 minutes after tooth extraction. In a separate study, Hall et al reported that cefaclor-treated patients did not have a reduction of postprocedure bacteremia compared with untreated control subjects. Contradictory published results from 2 studies showed reduction of postprocedure bacteremia by erythromycin in one but lack of efficacy for erythromycin or clindamycin in another. Finally, results are contradictory with regard to the efficacy of the use of topical antiseptics in reducing the frequency of bacteremia associated with dental procedures, but the preponderance of evidence suggests that there is no clear benefit. One study reported that chlorhexidine and povidone iodine mouth rinse were effective, whereas others showed no statistically significant benefit. Topical antiseptic rinses do not penetrate beyond 3 mm into the periodontal pocket and therefore do not reach areas of ulcerated tissue where bacteria most often gain entrance to the circulation. On the basis of these data, it is unlikely that topical antiseptics are effective to significantly reduce the frequency, magnitude, and duration of bacteremia associated with a dental procedure.

**Cumulative Risk Over Time of Bacteremias From Routine Daily Activities Compared With the Bacteremia From a Dental Procedure**

Guntheroth estimated a cumulative exposure of 5370 minutes of bacteremia over a 1-month period in dentulous patients resulting from random bacteremia from chewing food and from oral hygiene measures, such as tooth brushing and flossing, and compared that with a duration of bacteremia lasting 6 to 30 minutes associated with a single tooth extraction. Roberts estimated that tooth brushing 2 times daily for 1 year had a 154,000 times greater risk of exposure to bacteremia than that resulting from a single tooth extraction. The cumulative exposure during 1 year to bacteremia from routine daily activities may be as high as 5.6 million times greater than that resulting from a single tooth extraction, the dental procedure reported to be most likely to cause a bacteremia.

Data exist for the duration of bacteremia from a single tooth extraction, and it is possible to estimate the annual cumulative exposure from dental procedures for the average individual. However, calculations for the incidence, nature, and duration of bacteremia from routine daily activities are at best rough estimates, and it is therefore not possible to compare precisely the cumulative monthly or annual duration of exposure for bacteremia from dental procedures compared with routine daily activities. Nevertheless, even if the estimates of bacteremia from routine daily activities are off by a factor of 1000, it is likely that the frequency and cumulative duration of exposure to bacteremia from routine daily events over 1 year are much higher than those that result from dental procedures.

**Results of Clinical Studies of IE Prophylaxis for Dental Procedures**

No prospective, randomized, placebo-controlled studies exist on the efficacy of antibiotic prophylaxis to prevent IE in patients who undergo a dental procedure. Data from published retrospective or prospective case-control studies are limited by the following factors: (1) the low incidence of IE, which requires a large number of patients per cohort for statistical significance; (2) the wide variation in the types and severity of underlying cardiac conditions, which would require a large number of patients with specific matched control subjects for each cardiac condition; and (3) the large variety of invasive dental procedures and dental disease states, which would be difficult to standardize for control groups. These and other limitations complicate the interpretation of the results of published studies of the efficacy of IE prophylaxis in patients who undergo dental procedures.

Although some retrospective studies suggested that there was a benefit from prophylaxis, these studies were small in size and reported insufficient clinical data. Furthermore, in a number of cases, the incubation period between the dental procedure and the onset of symptoms of IE was prolonged. van der Meer and colleagues published a study of dental procedures in the Netherlands and the efficacy of antibiotic prophylaxis to prevent IE in patients with native or prosthetic cardiac valves. They concluded that dental or other procedures probably caused only a small fraction of cases of IE and that prophylaxis would prevent only a small number of cases even if it were 100% effective. These same authors performed a 2-year case-control study. Among patients for whom prophylaxis was recommended, 5 of 20 cases of IE
occurred despite receiving antibiotic prophylaxis. The authors concluded that prophylaxis was not effective. In a separate study, these authors reported poor awareness of recommendations for prophylaxis among both patients and healthcare providers. Strom and colleagues evaluated dental prophylaxis and cardiac risk factors in a multicenter case-control study. These authors reported that MVP, congenital heart disease (CHD), rheumatic heart disease (RHD), and previous cardiac valve surgery were risk factors for the development of IE. In that study, control subjects without IE were more likely to have undergone a dental procedure than those with cases of IE ($P=0.03$). The authors concluded that dental treatment was not a risk factor for IE even in patients with valvular heart disease and that few cases of IE could be prevented with prophylaxis even if it were 100% effective.

These studies are in agreement with a recently published French study of the estimated risk of IE in adults with predisposing cardiac conditions who underwent dental procedures with or without antibiotic prophylaxis. These authors reported that MVP, congenital heart disease (CHD), rheumatic heart disease (RHD), and previous cardiac valve surgery were risk factors for the development of IE. In that study, control subjects without IE were more likely to have undergone a dental procedure than those with cases of IE ($P=0.03$). The authors concluded that dental treatment was not a risk factor for IE even in patients with valvular heart disease and that few cases of IE could be prevented with prophylaxis even if it were 100% effective.

These studies are in agreement with a recently published French study of the estimated risk of IE in adults with predisposing cardiac conditions who underwent dental procedures with or without antibiotic prophylaxis. These authors reported that MVP, congenital heart disease (CHD), rheumatic heart disease (RHD), and previous cardiac valve surgery were risk factors for the development of IE. In that study, control subjects without IE were more likely to have undergone a dental procedure than those with cases of IE ($P=0.03$). The authors concluded that dental treatment was not a risk factor for IE even in patients with valvular heart disease and that few cases of IE could be prevented with prophylaxis even if it were 100% effective.

The estimated absolute risk of IE resulting from a dental procedure is exceedingly small. One would not expect antibiotic prophylaxis to be near 100% effective, is similarly small. One would not expect antibiotic prophylaxis to be near 100% effective, however, because of the nature of the organisms and choice of antibiotics.

**Risk of Adverse Reactions and Cost-Effectiveness of Prophylactic Therapy**

Nonfatal adverse reactions, such as rash, diarrhea, and GI upset, occur commonly with the use of antimicrobials; however, only single-dose therapy is recommended for dental prophylaxis, and these common adverse reactions are usually not severe and are self-limited. Fatal anaphylactic reactions were estimated to occur in 15 to 25 individuals per 1 million patients who receive a dose of penicillin. Among patients with a prior penicillin use, 36% of fatalities from anaphylaxis occurred in those with a known allergy to penicillin compared with 64% of fatalities among those with no history of penicillin allergy. These calculations are at best rough estimates and may overestimate the true risk of death caused by fatal anaphylaxis from administration of a penicillin. They are based on retrospective reviews or surveys of patients or on healthcare providers’ recall of events. A prospective study is necessary to accurately determine the risk of fatal anaphylaxis resulting from administration of a penicillin.

For 50 years, the AHA has recommended a penicillin as the preferred choice for dental prophylaxis for IE. During these 50 years, the Committee is unaware of any cases reported to the AHA of fatal anaphylaxis resulting from the administration of a penicillin recommended in the AHA guidelines for IE prophylaxis. The Committee believes that a single dose of amoxicillin or ampicillin is safe and is the preferred prophylactic agent for individuals who do not have a history of type 1 hypersensitivity reaction to a penicillin, such as anaphylaxis, urticaria, or angioedema. Fatal anaphylaxis from a cephalosporin is estimated to be less common than from penicillin, at approximately 1 case per 1 million patients. Fatal reactions to a single dose of a macrolide or clindamycin are extremely rare. There has been only 1 case report of...
is similar to that reported in other studies.100–103 Previously, incidence has remained stable during the past 4 decades and condition. Steckelberg and Wilson90 reported the lifetime risk of IE associated with a specific underlying cardiac condition. They estimated that the lifetime risk of IE associated with mitral valve prolapse (MVP) was 5 to 7 cases per 100 000 person-years.99 This estimation was based on data from Steckelberg and Wilson’s study. In Olmsted County, Minnesota, the incidence of IE in adults 50 years of age and older was calculated to be 271 per 100 000 patient-years in patients who underwent replacement of an infected prosthetic cardiac valve. In that study,90 the risk of IE per 100 000 patient-years was 4.6 in patients with MVP without a audible cardiac murmur and 52 in patients with MVP with an audible murmur of mitral regurgitation. Per 100 000 patient-years, the lifetime risk (380 to 440) for RHD was similar to that (308 to 383) for patients with a mechanical or bioprosthetic cardiac valve. The highest lifetime risks per 100 000 patient-years were as follows: cardiac valve replacement surgery for native valve IE, 630; previous IE, 740; and prosthetic valve replacement done in patients with prosthetic valve endocarditis, 2160. In a separate study, the risk of IE per 100 000 patient-years was 271 in patients with congenital aortic stenosis and 145 in patients with ventricular septal defect.105 These data indicate that the risk of IE before closure of a ventricular septal defect was more than twice that after closure. Although these data provide useful ranges of risk in large populations, it is difficult to utilize them to define accurately the lifetime risk of acquisition of IE in an individual patient with a specific underlying cardiac risk factor. This difficulty is based in part on the fact that each individual cardiac condition, such as RHD or MVP, represents a broad spectrum of pathology from minimal to severe, and the risk of IE would likely be influenced by the severity of valvular disease.

Cardiac Conditions and Endocarditis

Previous AHA guidelines categorized underlying cardiac conditions associated with the risk of IE as those with high risk, moderate risk, and negligible risk and recommended prophylaxis for patients in the high- and moderate-risk categories.1 For the present guidelines on prevention of IE, the Committee considered 3 distinct issues: (1) What underlying cardiac conditions over a lifetime have the highest predisposition to the acquisition of endocarditis? (2) What underlying cardiac conditions are associated with the highest risk of adverse outcome from endocarditis? (3) Should recommendations for IE prophylaxis be based on either or both of these 2 conditions?

Underlying Conditions Over a Lifetime That Have the Highest Predisposition to the Acquisition of Endocarditis

In Olmsted County, Minnesota, the incidence of IE in adults ranged from 5 to 7 cases per 100 000 person-years.99 This incidence has remained stable during the past 4 decades and is similar to that reported in other studies.100–103 Previously, RHD was the most common underlying condition predisposing to endocarditis, and RHD is still common in developing countries.99 In developed countries, the frequency of RHD has declined, and MVP is now the most common underlying condition in patients with endocarditis.104 Few published data quantify the lifetime risk of acquisition of IE associated with a specific underlying cardiac condition. Steckelberg and Wilson90 reported the lifetime risk of acquisition of IE, which ranged from 5 per 100 000 patient-years in the general population with no known cardiac conditions to 2160 per 100 000 patient-years in patients with MVP. This estimated risk is based on data from Steckelberg and Wilson’s study. In Olmsted County, Minnesota, the incidence of IE in adults 50 years of age and older was calculated to be 271 per 100 000 patient-years in patients who underwent replacement of an infected prosthetic cardiac valve. In that study,90 the risk of IE per 100 000 patient-years was 4.6 in patients with MVP without an audible cardiac murmur and 52 in patients with MVP with an audible murmur of mitral regurgitation. Per 100 000 patient-years, the lifetime risk (380 to 440) for RHD was similar to that (308 to 383) for patients with a mechanical or bioprosthetic cardiac valve. The highest lifetime risks per 100 000 patient-years were as follows: cardiac valve replacement surgery for native valve IE, 630; previous IE, 740; and prosthetic valve replacement done in patients with prosthetic valve endocarditis, 2160. In a separate study, the risk of IE per 100 000 patient-years was 271 in patients with congenital aortic stenosis and 145 in patients with ventricular septal defect.105 These data indicate that the risk of IE before closure of a ventricular septal defect was more than twice that after closure. Although these data provide useful ranges of risk in large populations, it is difficult to utilize them to define accurately the lifetime risk of acquisition of IE in an individual patient with a specific underlying cardiac risk factor. This difficulty is based in part on the fact that each individual cardiac condition, such as RHD or MVP, represents a broad spectrum of pathology from minimal to severe, and the risk of IE would likely be influenced by the severity of valvular disease.

Cardiac Conditions and Endocarditis

Previous AHA guidelines categorized underlying cardiac conditions associated with the risk of IE as those with high risk, moderate risk, and negligible risk and recommended prophylaxis for patients in the high- and moderate-risk categories.1 For the present guidelines on prevention of IE, the Committee considered 3 distinct issues: (1) What underlying cardiac conditions over a lifetime have the highest predisposition to the acquisition of endocarditis? (2) What underlying cardiac conditions are associated with the highest risk of adverse outcome from endocarditis? (3) Should recommendations for IE prophylaxis be based on either or both of these 2 conditions?

Underlying Conditions Over a Lifetime That Have the Highest Predisposition to the Acquisition of Endocarditis

In Olmsted County, Minnesota, the incidence of IE in adults ranged from 5 to 7 cases per 100 000 person-years.99 This incidence has remained stable during the past 4 decades and is similar to that reported in other studies.100–103 Previously, RHD was the most common underlying condition predisposing to endocarditis, and RHD is still common in developing countries.99 In developed countries, the frequency of RHD has declined, and MVP is now the most common underlying condition in patients with endocarditis.104 Few published data quantify the lifetime risk of acquisition of IE associated with a specific underlying cardiac condition. Steckelberg and Wilson90 reported the lifetime risk of acquisition of IE, which ranged from 5 per 100 000 patient-years in the general population with no known cardiac conditions to 2160 per 100 000 patient-years in patients with MVP. This estimated risk is based on data from Steckelberg and Wilson’s study. In Olmsted County, Minnesota, the incidence of IE in adults 50 years of age and older was calculated to be 271 per 100 000 patient-years in patients who underwent replacement of an infected prosthetic cardiac valve. In that study,90 the risk of IE per 100 000 patient-years was 4.6 in patients with MVP without an audible cardiac murmur and 52 in patients with MVP with an audible murmur of mitral regurgitation. Per 100 000 patient-years, the lifetime risk (380 to 440) for RHD was similar to that (308 to 383) for patients with a mechanical or bioprosthetic cardiac valve. The highest lifetime risks per 100 000 patient-years were as follows: cardiac valve replacement surgery for native valve IE, 630; previous IE, 740; and prosthetic valve replacement done in patients with prosthetic valve endocarditis, 2160. In a separate study, the risk of IE per 100 000 patient-years was 271 in patients with congenital aortic stenosis and 145 in patients with ventricular septal defect.105 These data indicate that the risk of IE before closure of a ventricular septal defect was more than twice that after closure. Although these data provide useful ranges of risk in large populations, it is difficult to utilize them to define accurately the lifetime risk of acquisition of IE in an individual patient with a specific underlying cardiac risk factor. This difficulty is based in part on the fact that each individual cardiac condition, such as RHD or MVP, represents a broad spectrum of pathology from minimal to severe, and the risk of IE would likely be influenced by the severity of valvular disease.

Cardiac Conditions and Endocarditis

Previous AHA guidelines categorized underlying cardiac conditions associated with the risk of IE as those with high risk, moderate risk, and negligible risk and recommended prophylaxis for patients in the high- and moderate-risk categories.1 For the present guidelines on prevention of IE, the Committee considered 3 distinct issues: (1) What underlying cardiac conditions over a lifetime have the highest predisposition to the acquisition of endocarditis? (2) What underlying cardiac conditions are associated with the highest risk of adverse outcome from endocarditis? (3) Should recommendations for IE prophylaxis be based on either or both of these 2 conditions?

Underlying Conditions Over a Lifetime That Have the Highest Predisposition to the Acquisition of Endocarditis

In Olmsted County, Minnesota, the incidence of IE in adults ranged from 5 to 7 cases per 100 000 person-years.99 This incidence has remained stable during the past 4 decades and is similar to that reported in other studies.100–103 Previously, RHD was the most common underlying condition predisposing to endocarditis, and RHD is still common in developing countries.99 In developed countries, the frequency of RHD has declined, and MVP is now the most common underlying condition in patients with endocarditis.104 Few published data quantify the lifetime risk of acquisition of IE associated with a specific underlying cardiac condition. Steckelberg and Wilson90 reported the lifetime risk of acquisition of IE, which ranged from 5 per 100 000 patient-years in the general population with no known cardiac conditions to 2160 per 100 000 patient-years in patients who
replacement surgery, and they have a higher mortality rate of IHF and increased need for cardiac valve replacement, perivalvular extension of infection, and other complications.

Valve replacement surgery, perivalvular extension of infection, and other complications.108,110–116

Similarly, the mortality of enterococcal prosthetic valve endocarditis is higher than that of native valve enterococcal endocarditis.107,108,114,117 Moreover, patients with prosthetic valve endocarditis are more likely than those with native valve endocarditis to develop heart failure, the need for cardiac valve replacement surgery, perivalvular extension of infection, and other complications.

Patients with relapsing or recurrent IE are at greater risk of congestive heart failure and increased need for cardiac valve replacement surgery, and they have a higher mortality rate than patients with a first episode of native valve IE.118–124 Additionally, patients with multiple episodes of native or prosthetic valve IE are at greater risk of additional episodes of endocarditis, each of which is associated with the risk of more serious complications.90

Published series regarding endocarditis in patients with CHD are underpowered to determine the extent to which a specific form of CHD is an independent risk factor for morbidity and mortality. Nevertheless, most retrospective case series suggest that patients with complex cyanotic heart disease and those who have postoperative palliative shunts, conduits, or other prostheses have a high lifetime risk of acquiring IE, and these same groups appear at highest risk for morbidity and mortality among all patients with CHD.125–129

In addition, multiple series and reviews reported that the presence of prosthetic material110,113 and complex cyanotic heart disease in patients of very young age (newborns and infants <2 years of age)122,132 are 2 factors associated with the worst prognoses from IE. Some types of CHD may be repaired completely without residual cardiac defects. As shown in Table 3, the Committee recommends prophylaxis for dental procedures for these patients during the first 6 months after the procedure. In these patients, endocardialization of prosthetic material occurs within 6 months after the procedure.134 The Committee does not recommend prophylaxis for dental procedures more than 6 months after the procedure provided that there is no residual defect from the repair. In most instances, treatment of patients who have infected prosthetic materials requires surgical removal in addition to medical therapy with associated high morbidity and mortality rates.

Should IE Prophylaxis Be Recommended for Patients With the Highest Risk of Acquisition of IE or for Patients With the Highest Risk of Adverse Outcome From IE?

In a major departure from previous AHA guidelines, the Committee no longer recommends IE prophylaxis based solely on an increased lifetime risk of acquisition of IE. It is noteworthy that patients with the conditions listed in Table 3 with a prosthetic cardiac valve, those with a previous episode of IE, and some patients with CHD are also among those patients with the highest lifetime risk of acquisition of endocarditis. No published data demonstrate convincingly that the administration of prophylactic antibiotics prevents IE associated with bacteremia from an invasive procedure. We cannot exclude the possibility that there may be an exceedingly small number of cases of IE that could be prevented by prophylactic antibiotics in patients who undergo an invasive procedure. However, if prophylaxis is effective, such therapy should be restricted to those patients with the highest risk of adverse outcome from IE who would derive the greatest benefit from prevention of IE. In patients with underlying cardiac conditions associated with the highest risk of adverse outcome from IE (Table 3), IE prophylaxis for dental procedures may be reasonable, even though we acknowledge that its effectiveness is unknown (Class IIb, LOE B).
prophylaxis. MVP is the most common underlying condition that predisposes to acquisition of IE in the Western world; however, the absolute incidence of endocarditis is extremely low for the entire population with MVP, and it is not usually associated with the grave outcome associated with the conditions identified in Table 3. Thus, IE prophylaxis is no longer recommended for this group of individuals.

Finally, the administration of prophylactic antibiotics is not risk free, as discussed above. Additionally, the widespread use of antibiotic therapy promotes the emergence of resistant microorganisms most likely to cause endocarditis, such as viridans group streptococci and enterococci. The frequency of multidrug-resistant viridans group streptococci and enterococci has increased dramatically during the past 2 decades. This increased resistance has reduced the efficacy and number of antibiotics available for the treatment of IE.

Regimens Recommended

General Principles

An antibiotic for prophylaxis should be administered in a single dose before the procedure. If the dosage of antibiotic is inadvertently not administered before the procedure, the dosage may be administered up to 2 hours after the procedure. However, administration of the dosage after the procedure should be considered only when the patient did not receive the pre-procedure dose. Some patients who are scheduled for an invasive procedure may have a coincidental endocarditis. The presence of fever or other manifestations of systemic infection should alert the provider to the possibility of IE. In these circumstances, it is important to obtain blood cultures and other relevant tests before administration of antibiotics intended to prevent IE. Failure to do so may result in delay in diagnosis or treatment of a concomitant case of IE.

Regimens for Dental Procedures

Previous AHA guidelines on prophylaxis listed a substantial number of dental procedures and events for which antibiotic prophylaxis was recommended and those procedures for which prophylaxis was not recommended. On the basis of a critical review of the published data, it is clear that transient viridans group streptococcal bacteremia may result from any dental procedure that involves manipulation of the gingival or periapical region of teeth or perforation of the oral mucosa. It cannot be assumed that manipulation of a healthy-appearing mouth or a minimally invasive dental procedure reduces the likelihood of a bacteremia. Therefore, antibiotic prophylaxis is recommended for patients with the conditions listed in Table 3 who undergo any dental procedure that involves manipulation of the gingival or periapical region of a tooth or for those procedures that perforate the oral mucosa (Table 4). Although IE prophylaxis may be reasonable for these patients, its effectiveness is unknown (Class IIb, LOE C). This includes procedures such as biopsies, suture removal, and placement of orthodontic bands, but it does not include routine anesthetic injections through noninfected tissue, the taking of dental radiographs, placement of removable prosthetic or orthodontic appliances, placement of orthodontic brackets, or adjustment of orthodontic appliances. Finally, there are other events that are not dental procedures and for which prophylaxis is not recommended, such as shedding of deciduous teeth and trauma to the lips and oral mucosa.

TABLE 4. Dental Procedures for Which Endocarditis Prophylaxis Is Recommended for Patients in Table 3

<table>
<thead>
<tr>
<th>All dental procedures that involve manipulation of gingival tissue or the periapical region of teeth or perforation of the oral mucosa*</th>
</tr>
</thead>
</table>

*The following procedures and events do not need prophylaxis: routine anesthetic injections through noninfected tissue, taking dental radiographs, placement of removable prosthetic or orthodontic appliances, adjustment of orthodontic appliances, placement of orthodontic brackets, shedding of deciduous teeth, and bleeding from trauma to the lips or oral mucosa.

In this limited patient population, prophylactic antimicrobial therapy should be directed against viridans group streptococci. During the past 2 decades, there has been a significant increase in the percentage of strains of viridans group streptococci resistant to antibiotics recommended in previous AHA guidelines for the prevention of IE. Prabhu et al135 studied susceptibility patterns of viridans group streptococci recovered from patients with IE diagnosed during a period from 1971 to 1986 and compared these susceptibilities with those of viridans group streptococci from patients with IE diagnosed from 1994 to 2002. In that study, none of the strains of viridans group streptococci were penicillin resistant in the early time period compared with 13% of strains that were intermittently or fully penicillin resistant during the later time period. In that study, macrolide resistance increased from 11% to 26% and clindamycin resistance from 0% to 4%.

Among 352 blood culture isolates of viridans group streptococci, resistance rates were 13% for penicillin, 15% for amoxicillin, 17% for ceftriaxone, 38% for erythromycin, and 96% for cephalexin.136 The rank order of decreasing level of activity of cephalexin in that study was cefotaxime equal to ceftriaxone, greater than cefprozil, and equal to cefuroxime, and cephalexin was the least active. In other studies, resistance of viridans group streptococci to penicillin ranged from 17% to 50%,137-142 and resistance to ceftriaxone ranged from 22% to 42%.131,140 Ceftriaxone was 2 to 4 times more active in vitro than cefazolin.131,140 Similarly high rates of resistance were reported for macrolides, ranging from 22% to 58%.137,141,143,144 Resistance to clindamycin ranged from 13% to 27%.128,129,131,137,138,140

Most of the strains of viridans group streptococci in the above-cited studies were recovered from patients with serious underlying illnesses, including malignancies and febrile neutropenia. These patients are at increased risk of infection and colonization by multiple-drug–resistant microorganisms, including viridans group streptococci. Accordingly, these strains may not be representative of susceptibility patterns of viridans group streptococci recovered from presumably normal individuals who undergo a dental procedure. Diekema et al137 reported that 32% of strains of viridans group streptococci were resistant to penicillin in patients without cancer. King et al144 reported erythromycin resistance in 41% of streptococci recovered from throat cultures in otherwise healthy individuals who presented with mild respiratory tract infections. In that study, after treatment with either azithromycin or clindamycin, the percentage of resistant streptococci increased to 82% and 71%, respectively. Accordingly, the
resistance rates of viridans group streptococci are similarly high in otherwise healthy individuals and in patients with serious underlying diseases.

The impact of viridans group streptococcal resistance on antibiotic prevention of IE is unknown. If resistance in vitro is predictive of lack of clinical efficacy, the high resistance rates of viridans group streptococci provide additional support for the assertion that prophylactic therapy for a dental procedure is of little, if any, value. It is impractical to recommend prophylaxis with only those antibiotics, such as vancomycin or a fluoroquinolone, that are highly active in vitro against viridans group streptococci. There is no evidence that such therapy is effective for prophylaxis of IE, and their use might result in the development of resistance of viridans group streptococci and other microorganisms to these and other antibiotics.

In Table 5, amoxicillin is the preferred choice for oral therapy because it is well absorbed in the GI tract and provides high and sustained serum concentrations. For individuals who are allergic to penicillins or amoxicillin, the use of cephalaxin or another first-generation oral cephalosporin, clindamycin, azithromycin, or clarithromycin is recommended. Even though cephalaxin was less active against viridans group streptococci than other first-generation oral cephalosporins in 1 study,136 cephalaxin is included in Table 5. No data show superiority of 1 oral cephalosporin over another for prevention of IE, and generic cephalosporin is widely available and relatively inexpensive. Because of possible cross-reactions, a cephalosporin should not be administered to patients with a history of anaphylaxis, angioedema, or urticaria after treatment with any form of penicillin, including ampicillin or amoxicillin. Patients who are unable to tolerate an oral antibiotic may be treated with ampicillin, ceftriaxone, or cefazolin administered intramuscularly or intravenously. For ampicillin-allergic patients who are unable to tolerate an oral agent, therapy is recommended with parenteral cefazolin, ceftriaxone, or clindamycin.

### Regimens for Respiratory Tract Procedures

A variety of respiratory tract procedures reportedly cause transient bacteremia with a wide array of microorganisms; however, no published data conclusively demonstrate a link between these procedures and IE. Antibiotic prophylaxis with a regimen listed in Table 5 may be considered (Class IIb, LOE C) for patients with the conditions listed in Table 3 who undergo an invasive procedure of the respiratory tract that involves incision or biopsy of the respiratory mucosa, such as tonsillectomy and adenoidectomy. We do not recommend antibiotic prophylaxis for bronchoscopy unless the procedure involves incision of the respiratory tract mucosa. For patients listed in Table 3 who undergo an invasive respiratory tract procedure to treat an established infection, such as drainage of an abscess or empyema, we recommend that the antibiotic regimen administered to these patients contain an agent active against viridans group streptococci (Table 5). If the infection is known or suspected to be caused by *Staphylococcus aureus*, the regimen should contain an agent active against *S. aureus*, such as an antistaphylococcal penicillin or cephalosporin, or vancomycin in patients unable to tolerate a β-lactam. Vancomycin should be administered if the infection is known or suspected to be caused by a methicillin-resistant strain of *S. aureus*.

### Recommendations for GI or GU Tract Procedures

Enterococci are part of the normal flora of the GI tract. These microorganisms may cause intra-abdominal infection or infection of the hepatobiliary system. Such infections are often polymicrobial, with a mix of aerobic and anaerobic Gram-negative and Gram-positive microorganisms, but among these varied bacteria, only enterococci are likely to cause IE. Enterococci may cause urinary tract infections, particularly in older males with prostatic hypertrophy and obstructive uropathy or prostatitis.

The administration of prophylactic antibiotics solely to prevent endocarditis is not recommended for patients who...
undergo GU or GI tract procedures, including diagnostic esophagogastroduodenoscopy or colonoscopy (Class III, LOE B). This is in contrast to previous AHA guidelines that listed GI or GU tract procedures for which IE prophylaxis was recommended and those for which prophylaxis was not recommended.1 A large number of diagnostic and therapeutic procedures that involve the GI, hepatobiliary, or GU tract may cause transient enterococcal bacteremia. The possible association between GI or GU tract procedures and IE has not been studied as extensively as the possible association with dental procedures.145 The cases of IE temporally associated with a GI or GU tract procedure are anecdotal, with either a single or very small number of cases reported.83 No published data demonstrate a conclusive link between procedures of the GI or GU tract and the development of IE.145 Moreover, no studies exist that demonstrate that the administration of antimicrobial prophylaxis prevents IE in association with procedures performed on the GI or GU tract.

There has been a dramatic increase in the frequency of antimicrobial-resistant strains of enterococci to penicillins, vancomycin, and aminoglycosides.146–151 These antibiotics were recommended for IE prophylaxis in previous AHA guidelines.1 The significance of the increased frequency of multiresistant strains of enterococci on prevention of IE in patients who undergo GI or GU tract procedures is unknown. The high prevalence of resistant strains of enterococci adds further doubt about the efficacy of prophylactic therapy for GI or GU tract procedures.

Patients with infections of the GI or GU tract may have intermittent or sustained enterococcal bacteremia. For patients with the conditions listed in Table 3 who have an established GI or GU tract infection or for those who receive antibiotic therapy to prevent wound infection or sepsis associated with a GI or GU tract procedure, it may be reasonable that the antibiotic regimen include an agent active against staphylococci and enterococci, such as an antistaphylococcal penicillin or a cephalosporin (Class IIb, LOE B). However, no published studies demonstrate that such therapy would prevent enterococcal IE.

For patients with the conditions listed in Table 3 scheduled for an elective cystoscopy or other urinary tract manipulation who have an enterococcal urinary tract infection or colonization, antibiotic therapy to eradicate enterococci from the urine before the procedure may be reasonable (Class IIb, LOE B). If the urinary tract procedure is not elective, it may be reasonable that the empiric or specific antimicrobial regimen administered to the patient contain an agent active against enterococci (Class IIb, LOE B).

Amoxicillin or ampicillin is the preferred agent for enterococcal coverage for these patients. Vancomycin may be administered to patients unable to tolerate ampicillin. If infection is caused by a known or suspected strain of resistant enterococcus, consultation with an infectious diseases expert is recommended.

### Recommendations for Procedures on Infected Skin, Skin Structure, or Musculoskeletal Tissue

These infections are often polymicrobial, but only staphylococci and β-hemolytic streptococci are likely to cause IE. For patients with the conditions listed in Table 3 who undergo a surgical procedure that involves infected skin, skin structure, or musculoskeletal tissue, it is reasonable that the therapeutic regimen administered for treatment of the infection contain an agent active against staphylococci and β-hemolytic streptococci, such as an antistaphylococcal penicillin or a cephalosporin (Table 5 for dosage: Class IIb, LOE C). Vancomycin or clindamycin may be administered to patients unable to tolerate a β-lactam or who are known or suspected to have an infection caused by a methicillin-resistant strain of staphylococcus.

A summary of the major changes in these updated recommendations for prevention of IE compared with previous AHA recommendations is shown in Table 6.

### Specific Situations and Circumstances

#### Patients Already Receiving Antibiotics

If a patient is already receiving long-term antibiotic therapy with an antibiotic that is also recommended for IE prophylaxis for a dental procedure, it is prudent to select an antibiotic from a different class rather than to increase the dosage of the current antibiotic. For example, antibiotic regimens used to prevent the recurrence of acute rheumatic fever are administered in dosages lower than those recommended for the prevention of IE. Individuals who take an oral penicillin for secondary prevention of rheumatic fever or for other purposes are likely to have viridans group streptococci in their oral cavity that are relatively resistant to penicillin or amoxicillin. In such cases, the provider should select either clindamycin, azithromycin, or clarithromycin for IE prophylaxis for a dental procedure, but only for patients shown in Table 3. Because of possible cross-resistance of viridans...

---

**TABLE 6. Summary of Major Changes in Updated Document**

<table>
<thead>
<tr>
<th>Recommendation</th>
<th>Changed from Previous AHA Guidelines</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antibiotic prophylaxis is no longer recommended for any other form of CHD, except for the conditions listed in Table 3.</td>
<td></td>
</tr>
<tr>
<td>Antibiotic prophylaxis is recommended for all dental procedures that involve manipulation of gingival tissues or periapical region of teeth or perforation of oral mucosa only for patients with underlying cardiac conditions associated with the highest risk of adverse outcome from IE (Table 3).</td>
<td></td>
</tr>
<tr>
<td>Antibiotic prophylaxis is recommended for procedures on respiratory tract or infected skin, skin structures, or musculoskeletal tissue only for patients with underlying cardiac conditions associated with the highest risk of adverse outcome from IE (Table 3).</td>
<td></td>
</tr>
<tr>
<td>Antibiotic prophylaxis solely to prevent IE is not recommended for GU or GI tract procedures.</td>
<td></td>
</tr>
<tr>
<td>The writing group reaffirms the procedures noted in the 1997 prophylaxis guidelines for which endocarditis prophylaxis is not recommended and extends this to other common procedures, including ear and body piercing, tattooing, and vaginal delivery and hysterectomy.</td>
<td></td>
</tr>
</tbody>
</table>

We concluded that there are relatively few cases of IE caused by daily activities compared with those associated with dental procedures. We concluded that bacteremia resulting from daily activities is much more likely to cause IE than bacteremia associated with a dental procedure. We concluded that only an extremely small number of cases of IE might be prevented by antibiotic prophylaxis even if prophylaxis is 100% effective.
group streptococci with cephalosporins, this class of antibiotics should be avoided. If possible, it would be preferable to delay a dental procedure until at least 10 days after completion of the antibiotic therapy. This may allow time for the usual oral flora to be reestablished.

Patients receiving parenteral antibiotic therapy for IE may require dental procedures during antimicrobial therapy, particularly if subsequent cardiac valve replacement surgery is anticipated. In these cases, the parenteral antibiotic therapy for IE should be continued and the timing of the dosage adjusted to be administered 30 to 60 minutes before the dental procedure. This parenteral antimicrobial therapy is administered in such high doses that the high concentration would overcome any possible low-level resistance developed among mouth flora (unlike the concentration that would occur after oral administration).

Patients Who Receive Anticoagulants
Intramuscular injections for IE prophylaxis should be avoided in patients who are receiving anticoagulant therapy (Class I, LOE A). In these circumstances, orally administered regimens should be given whenever possible. Intravenously administered antibiotics should be used for patients who are unable to tolerate or absorb oral medications.

Patients Who Undergo Cardiac Surgery
A careful preoperative dental evaluation is recommended so that required dental treatment may be completed whenever possible before cardiac valve surgery or replacement or repair of CHD. Such measures may decrease the incidence of late prosthetic valve endocarditis caused by viridans group streptococci.

Patients who undergo surgery for placement of prosthetic heart valves or prosthetic intravascular or intracardiac materials are at risk for the development of infection. Because the morbidity and mortality of infection in these patients are high, perioperative prophylactic antibiotics are recommended (Class I, LOE B). Early-onset prosthetic valve endocarditis is most often caused by *S aureus*, coagulase-negative staphylococci, or diphtheroids. No single antibiotic regimen is effective against all these microorganisms. Prophylaxis at the time of cardiac surgery should be directed primarily against staphylococci and should be of short duration. A first-generation cephalosporin is most often used, but the choice of an antibiotic should be influenced by the antibiotic susceptibility patterns at each hospital. For example, a high prevalence of infection by methicillin-resistant *S aureus* should prompt the consideration of the use of vancomycin for perioperative prophylaxis. The majority of nosocomial coagulase-negative staphylococci are methicillin-resistant. Nonetheless, surgical prophylaxis with a first-generation cephalosporin is recommended for these patients (Class I, LOE A). In hospitals with a high prevalence of methicillin-resistant strains of *S epidermidis*, surgical prophylaxis with vancomycin is reasonable but has not been shown to be superior to prophylaxis with a cephalosporin (Class IIB, LOE C). Prophylaxis should be initiated immediately before the operative procedure, repeated during prolonged procedures to maintain serum concentrations intraoperatively, and continued for no more than 48 hours postoperatively to minimize emergence of resistant microorganisms (Class IIa, LOE B). The effects of cardiopulmonary bypass and compromised renal function on antibiotic concentrations in serum should be considered and dosages adjusted as necessary before and during the procedure.

Other Considerations
There is no evidence that coronary artery bypass graft surgery is associated with a long-term risk for infection. Therefore, antibiotic prophylaxis for dental procedures is not needed for individuals who have undergone this surgery. Antibiotic prophylaxis for dental procedures is not recommended for patients with coronary artery stents (Class III, LOE C). The treatment and prevention of infection for these and other endovascular grafts and prosthetic devices are addressed in a separate AHA publication. There are insufficient data to support specific recommendations for patients who have undergone heart transplantation. Such patients are at risk of acquired valvular dysfunction, especially during episodes of rejection. Endocarditis that occurs in a heart transplant patient is associated with a high risk of adverse outcome (Table 3). Accordingly, the use of IE prophylaxis for dental procedures in cardiac transplant recipients who develop cardiac valvulopathy may be reasonable, but the usefulness is not well established (Class IIb, LOE C; Table 4). The use of prophylactic antibiotics to prevent infection of joint prostheses during potentially bacteremia-inducing procedures is not within the scope of this document.

Future Considerations
Prospective placebo-controlled, double-blinded studies of antibiotic prophylaxis of IE in patients who undergo a bacteremia-producing procedure would be necessary to evaluate accurately the efficacy of IE prophylaxis. Additional prospective case-control studies are needed. The AHA has made substantial revisions to previously published guidelines on IE prophylaxis. Given our current recommendations, we anticipate that significantly fewer patients will receive IE prophylaxis for a dental procedure. Studies are necessary to monitor the effects, if any, of these recommended changes in IE prophylaxis. The incidence of IE could change or stay the same. Because the incidence of IE is low, small changes in incidence may take years to detect. Accordingly, we urge that such studies be designed and instituted promptly so that any change in incidence may be detected sooner rather than later. Subsequent revisions of the AHA guidelines on the prevention of IE will be based on the results of these studies and other published data.

Acknowledgments
The writing group thanks the following international experts on infective endocarditis for their valuable comments: Drs Christa Gohlke-Bärwolf, Roger Hall, Jae-Hoon Song, Catherine Kilmartin, Catherine Leport, José M. Miró, Christoph Naber, Graham Roberts, and Jan T.M. van der Meer. The writing group also thanks Dr George Meyer for his helpful comments regarding gastroenterology. Finally, the writing group would like to thank Lori Hinrichs for her superb assistance with the preparation of this manuscript.
### Writing Group Disclosures

<table>
<thead>
<tr>
<th>Writing Group Member</th>
<th>Employment</th>
<th>Research Grant</th>
<th>Other Research Support</th>
<th>Ownership Interest</th>
<th>Consultant/Advisory Board</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>Walter Wilson</td>
<td>Mayo Clinic</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Larry M. Baddour</td>
<td>Mayo Clinic</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Robert S. Baltimore</td>
<td>Yale University School of Medicine</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Ann Bolger</td>
<td>University of California, San Francisco</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Robert O. Bonow</td>
<td>Northwestern University Feinberg School of Medicine</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Jane C. Burns</td>
<td>University of California, San Diego</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Christopher H. Cabell</td>
<td>Duke University National Institutes of Health†</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>Gloucester*; Shire*; Cubist*; Carbomedics*; GlaxoSmithKline*; Acusphere*; Endo*; Eli Lilly*; Watson*; Johnson &amp; Johnson*</td>
<td>None</td>
</tr>
<tr>
<td>David T. Durack</td>
<td>Becton Dickinson &amp; Co (manufactures medical devices and diagnostics)</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>Joint Commission Resources Board†</td>
<td>None</td>
</tr>
<tr>
<td>Patricia Ferrier</td>
<td>University of Minnesota Medical School</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Timothy Gardner</td>
<td>Christiana Care Health System</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Michael Gerber</td>
<td>Cincinnati Children’s Hospital Medical Center</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Michael Gewitz</td>
<td>Maria Fareri Children’s Hospital of Westchester, New York Medical College</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>David Goff</td>
<td>Wake Forest University School of Medicine</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Matthew Levison</td>
<td>Drexel University College of Medicine</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>Merck*</td>
<td>None</td>
</tr>
<tr>
<td>Peter B. Lockhart</td>
<td>Carolina Medical Center</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Jane W. Newburger</td>
<td>Boston Children’s Heart Foundation</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Thomas Pallasch</td>
<td>University of Southern California</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>Consultation and expert witness testimony on records of patients with endocarditis</td>
<td>None</td>
</tr>
<tr>
<td>Anne H. Rowley</td>
<td>Children’s Memorial Hospital, Chicago</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Stanford T. Shulman</td>
<td>Children’s Memorial Hospital, Chicago</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Brian L. Strom</td>
<td>University of Pennsylvania School of Medicine Pfizer*; Merck*; Novartis*; Wyeth*; Pfizer*</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>Abbott*; GlaxoSmithKline*; Eli Lilly*; Pfizer*; Sanofi Pasteur*; Johnson &amp; Johnson*; Schering AG*; Tap Pharma*; Wyeth*</td>
<td>None</td>
</tr>
<tr>
<td>Masato Takahashi</td>
<td>University of Southern California</td>
<td>Bristol-Myers Squibb Medical Imaging*</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Lloyd Y. Tani</td>
<td>University of Utah School of Medicine</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Kathryn A. Taubert</td>
<td>American Heart Association</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
</tbody>
</table>

This table represents the relationships of writing group members that may be perceived as actual or reasonably perceived conflicts of interest as reported on the Disclosure Questionnaire, which all members of the writing group are required to complete and submit. A relationship is considered to be “Significant” if (1) the person receives $10 000 or more during any 12-month period, or 5% or more of the person’s gross income; or (2) the person owns 5% or more of the voting stock or share of the entity or owns $10 000 or more of the fair market value of the entity. A relationship is considered to be “Modest” if it is less than “Significant” under the preceding definition.

*Modest.
†Significant.
### Reviewer Disclosures

<table>
<thead>
<tr>
<th>Reviewer</th>
<th>Employment</th>
<th>Research Grant</th>
<th>Other Research Support</th>
<th>Speakers’ Bureau/ Honoraria</th>
<th>Expert Witness</th>
<th>Ownership Interest</th>
<th>Consultant/ Advisory Board</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thomas Bashore</td>
<td>Duke University Medical Center</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Arnold Bayer</td>
<td>University of California, Los Angeles</td>
<td>Titan†</td>
<td>NIH†</td>
<td>Cubist†</td>
<td>June Baker Laird at McElroy, Deutsch, Mulvaney &amp; Carpenter, LLP (Denver, Colo)*</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Donald Falace</td>
<td>University of Kentucky</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Michael Freed</td>
<td>Boston Children’s Hospital</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Welton Gersony</td>
<td>Children’s Hospital of New York</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Loren Hiratzka</td>
<td>Bethesda North Hospital</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Patrick O’Gara</td>
<td>Brigham &amp; Women’s Hospital</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Lauren L. Patton</td>
<td>University of North Carolina</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Catherine L. Webb</td>
<td>Northwestern University</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>Amgen†</td>
<td>None</td>
<td>None</td>
</tr>
</tbody>
</table>

This table represents the relationships of reviewers that may be perceived as actual or reasonably perceived conflicts of interest as reported on the Disclosure Questionnaire, which all reviewers are required to complete and submit. A relationship is considered to be “Significant” if (1) the person receives $10,000 or more during any 12-month period, or 5% or more of the person’s gross income; or (2) the person owns 5% or more of the voting stock or share of the entity or owns $10,000 or more of the fair market value of the entity. A relationship is considered to be “Modest” if it is less than “Significant” under the preceding definition.

*Modest.
†Significant.

### References


Downloaded from circ.ahajournals.org by on October 3, 2007


42. Roberts GJ. Dentists are innocent! “Everyday” bacteremia is the real culprit: a review and assessment of the evidence that dental surgical procedures are a principal cause of bacteremia in children. Pediatr Cardiol. 1999;20:317–325.


52. Roberts GJ. Dentists are innocent! “Everyday” bacteremia is the real culprit: a review and assessment of the evidence that dental surgical procedures are a principal cause of bacteremia in children. Pediatr Cardiol. 1999;20:317–325.


119. Sexton DJ, Tenenbaum MJ, Wilson WR, Steckelberg JM, Tice AD, Gilbert D, Dismukes W, Drew RH, Durack DT; Endocarditis Treatment Consortium Group. Ceftriaxone once daily for four weeks compared with ceftriaxone plus gentamicin once daily for two weeks for treatment


