

## Complete Summary

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### **GUIDELINE TITLE**

Antithrombotic therapy in patients with saphenous vein and internal mammary artery bypass grafts: the Seventh ACCP Conference on Antithrombotic and Thrombolytic Therapy.

### **BIBLIOGRAPHIC SOURCE(S)**

Stein PD, Schunemann HJ, Dalen JE, Gutterman D. Antithrombotic therapy in patients with saphenous vein and internal mammary artery bypass grafts: the Seventh ACCP Conference on Antithrombotic and Thrombolytic Therapy. Chest 2004 Sep;126(3 Suppl):600S-8S. [49 references] [PubMed](#)

### **GUIDELINE STATUS**

This is the current release of the guideline.

This guideline updates a previous version: Stein PD, Dalen JE, Goldman S, Theroux P. Antithrombotic therapy in patients with saphenous vein and internal mammary artery bypass grafts. Chest 2001 Jan;119(1 Suppl):278S-282S.

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## SCOPE

### **DISEASE/CONDITION(S)**

Coronary artery disease: saphenous vein and internal mammary artery (IMA) bypass grafts

### **GUIDELINE CATEGORY**

Treatment

## **CLINICAL SPECIALTY**

Cardiology  
Family Practice  
Internal Medicine  
Surgery

## **INTENDED USERS**

Physicians

## **GUIDELINE OBJECTIVE(S)**

- To review the literature on the effects of treatment with antithrombotic agents on the prevention of saphenous vein and internal mammary artery (IMA) bypass graft occlusion following coronary artery bypass grafting (CABG)
- To provide evidence-based recommendations on the treatment of saphenous vein bypass grafts and internal mammary artery bypass grafts with antithrombotic agents for the purpose of maintaining graft patency

## **TARGET POPULATION**

Patients who have undergone saphenous vein and internal mammary artery (IMA) bypass graft procedures for coronary artery disease

## **INTERVENTIONS AND PRACTICES CONSIDERED**

### **Treatment**

1. Aspirin therapy
2. Clopidogrel, as an alternative to or in addition to aspirin therapy
3. Oral anticoagulants, including vitamin K antagonists (VKAs)
4. Aspirin therapy in combination with oral anticoagulants

**Note:** The following medications, or combinations of medications were considered but not recommended: dipyridamole alone or in combination with aspirin, indobufen, ticlopidine, sulfinpyrazone.

## **MAJOR OUTCOMES CONSIDERED**

- Effectiveness of treatment on graft patency as determined by:
  - Cardiovascular events
  - Death
  - Graft thrombosis
  - Graft patency
- Rates of bleeding complications

## **METHODOLOGY**

### **METHODS USED TO COLLECT/SELECT EVIDENCE**

Hand-searches of Published Literature (Primary Sources)  
Searches of Electronic Databases

## **DESCRIPTION OF METHODS USED TO COLLECT/SELECT THE EVIDENCE**

### **Process of Searching for Evidence**

Defining the clinical question provided the framework for formulating eligibility criteria that guided the search for relevant evidence. Prior to searching for the evidence, methodological experts and librarians reviewed each question to ensure that the librarians could derive a comprehensive search strategy.

In specifying eligibility criteria, authors not only identified patients, interventions, and outcomes, but also methodological criteria. For most therapeutic studies, authors restricted eligibility to randomized controlled trials (RCTs).

For many questions, RCTs did not provide sufficient data, and article authors also included observational studies. This was also true when randomized trials were not the most appropriate design to use for addressing the research question. In particular, randomized trials are not necessarily the best design to understand risk groups (e.g., the baseline or expected risk of a given event for certain subpopulations). Because there are no interventions examined in questions about prognosis, one replaces interventions by the exposure, which is time.

### **Identifying the Evidence**

To identify the relevant evidence, a team of librarians at the University at Buffalo conducted comprehensive literature searches. For each question the authors provided, the librarians developed sensitive (but not specific) search strategies, including all languages, and conducted separate searches for systematic reviews, RCTs, and, if applicable, observational studies. The librarians searched the Cochrane Database of Systematic Reviews, the Database of Abstracts of Reviews of Effectiveness and Cochrane Register of Controlled Trial, the ACP Journal Club, MEDLINE, and Embase for studies published between 1966 and June 2002 in any language. To filter MEDLINE and Embase search results for RCT evidence, the librarians used the search strategy developed by the Cochrane Collaboration (full strategy available in Appendix online at:

[http://www.chestjournal.org/content/vol126/3\\_suppl\\_1](http://www.chestjournal.org/content/vol126/3_suppl_1)).

For observational studies, they restricted their searches to human studies. Searches were not further restricted in terms of methodology. While increasing the probability of identifying all published studies, this sensitive approach resulted in large number of citations for many of the defined clinical questions. Therefore, trained research assistants screened the citation list developed from the search and removed any apparently irrelevant citations. These irrelevant citations included press news, editorials, narrative reviews, single case reports, animal studies (any nonhuman studies), and letters to the editor. Authors included data from abstracts of recent meetings if reporting was transparent and all necessary data for the formulation of a recommendation were available. The guideline developers did not explicitly use Internet sources to search for research data.

## **NUMBER OF SOURCE DOCUMENTS**

Not stated

## **METHODS USED TO ASSESS THE QUALITY AND STRENGTH OF THE EVIDENCE**

Weighting According to a Rating Scheme (Scheme Given)

## **RATING SCHEME FOR THE STRENGTH OF THE EVIDENCE**

The rating scheme framework captures the trade-off between benefits and risks (1 or 2) (and the methodological quality of the underlying evidence (A, B, C+, or C). See "Rating Scheme for the Strength of the Recommendations."

## **METHODS USED TO ANALYZE THE EVIDENCE**

Meta-Analysis of Randomized Controlled Trials  
Systematic Review with Evidence Tables

## **DESCRIPTION OF THE METHODS USED TO ANALYZE THE EVIDENCE**

### **Summarizing Evidence**

The electronic searches also included searching for systematic reviews. If authors were satisfied with a recent high-quality systematic review, evidence from that review provided a foundation for the relevant recommendation.

Pooled analyses from high-quality systematic reviews formed, wherever possible, the evidence base of the recommendations. Pooling offers the advantage of obtaining more precise estimates of treatment effects and allows for a greater generalizability of results. However, pooling also bears the risk of spurious generalization. In general, the summary estimates of interest were the different types of outcomes conveying benefit and downsides (i.e., risk, burden, and cost).

## **METHODS USED TO FORMULATE THE RECOMMENDATIONS**

Expert Consensus (Consensus Development Conference)

## **DESCRIPTION OF METHODS USED TO FORMULATE THE RECOMMENDATIONS**

The strength of any recommendation depends on the following two factors: the trade-off between the benefits and the risks, burdens, and costs; and the strength of the methodology that leads to the treatment effect. The guideline developers grade the trade-off between benefits and risks in the two categories: 1, in which the trade-off is clear enough that most patients, despite differences in values, would make the same choice; and 2, in which the trade-off is less clear, and individual patients' values will likely lead to different choices.

When randomized trials provide precise estimates suggesting large treatment effects, and the risks and costs of therapy are small, treatment for average patients with compatible values and preferences can be confidently recommended.

If the balance between benefits and risks is in doubt, methodologically rigorous studies providing **Grade A** evidence and recommendations may still be weak (**Grade 2**). Uncertainty may come from less precise estimates of benefit, harm, or costs, or from small effect sizes.

There is an independent impact of validity and consistency, and the balance of positive and negative impacts of treatment on the strength of recommendations. In situations in which there is doubt about the value of the trade-off, any recommendation will be weaker, moving from **Grade 1** to **Grade 2**.

**Grade 1** recommendations can only be made when there is a relatively clear picture of both the benefits and the risks, burdens, and costs, and when the balance between the two clearly favors recommending or not recommending the intervention for the typical patient with compatible values and preferences. A number of factors can reduce the strength of a recommendation, moving it from **Grade 1** to **Grade 2**. Uncertainty about a recommendation to treat may be introduced if the following conditions apply: (1) the target event that is trying to be prevented is less important (confident recommendations are more likely to be made to prevent death or stroke than asymptomatic deep vein thrombosis); (2) the magnitude of risk reduction in the overall group is small; (3) the probability of the target event is low in a particular subgroup of patients; (4) the estimate of the treatment effect is imprecise, as reflected in a wide confidence interval (CI) around the effect; (5) there is substantial potential harm associated with therapy; or (6) there is an expectation for a wide divergence in values even among average or typical patients. Higher costs would also lead to weaker recommendations to treat.

The more balanced the trade-off between benefits and risks, the greater the influence of individual patient values in decision making. Virtually all patients, if they understand the benefits and risks, will take aspirin after experiencing a myocardial infarction (MI) or will comply with prophylaxis to reduce the risk of thromboembolism after undergoing hip replacement. Thus, one way of thinking about a **Grade 1** recommendation is that variability in patient values is unlikely to influence treatment choice in average or typical patients.

When the trade-off between benefits and risks is less clear, individual patient values may influence treatment decisions even among patients with average or typical preferences.

**Grade 2** recommendations are those in which variation in patient values or individual physician values will often mandate different treatment choices, even among average or typical patients. An alternative, but similar, interpretation is that a **Grade 2** recommendation suggests that clinicians conduct detailed conversations with patients to ensure that their ultimate recommendation is consistent with the patient's values.

## **RATING SCHEME FOR THE STRENGTH OF THE RECOMMENDATIONS**

<b>Grade of Recommendation</b>	<b>Clarity of Risk/Benefit</b>	<b>Methodological Strength of Supporting Evidence</b>	<b>Implications</b>
<b>1A</b>	Clear	Randomized controlled trials (RCTs) without important limitations	Strong recommendation; can apply to most patients in most circumstances without reservation
<b>1C+</b>	Clear	No RCTs, but strong RCT results can be unequivocally extrapolated, or overwhelming evidence from observational studies	Strong recommendation; can apply to most patients in most circumstances
<b>1B</b>	Clear	RCTs with important limitations (inconsistent results, methodological flaws*)	Strong recommendation; likely to apply to most patients
<b>1C</b>	Clear	Observational studies	Intermediate-strength recommendation; may change when stronger evidence is available
<b>2A</b>	Unclear	RCTs without important limitations	Intermediate-strength recommendation; best action may differ depending on circumstances or patients' or societal values
<b>2C+</b>	Unclear	No RCTs, but strong RCT results can be unequivocally	Weak recommendation; best action may differ depending on

Grade of Recommendation	Clarity of Risk/Benefit	Methodological Strength of Supporting Evidence	Implications
		extrapolated, or overwhelming evidence from observational studies	circumstances or patients' or societal values
<b>2B</b>	Unclear	RCTs with important limitations (inconsistent results, methodological flaws*)	Weak recommendation; alternative approaches likely to be better for some patients under some circumstances
<b>2C</b>	Unclear	Observational studies	Very weak recommendation; other alternatives may be equally reasonable

*\*These situations include RCTs with both lack of blinding and subjective outcomes, where the risk of bias in measurement of outcomes is high, or RCTs with large loss to follow-up.*

## COST ANALYSIS

While conference participants agreed that recommendations should reflect economic considerations, incorporating costs is fraught with difficult challenges. For most recommendations, formal economic analyses are unavailable. Even when analyses are available, they may be methodologically weak or biased. Furthermore, costs differ radically across jurisdictions, and even sometimes across hospitals within jurisdictions.

Because of these challenges, the guideline developers consider economic factors only when the costs of one therapeutic option over another are substantially different within major jurisdictions in which clinicians make use of these recommendations. As a result, in jurisdictions in which resource constraints are severe, alternative allocations may serve the health of the public far better than some of the interventions that are designated as **Grade 1A**. This will likely be true for all less industrialized countries and, with the increasing promotion of expensive drugs with marginal benefits, may be increasingly true for wealthier nations. Furthermore, recommendations change (either in direction or with respect to grade) only when the guideline developers believe that costs are high in relation to benefits. Instances in which costs have influenced recommendations

are labeled in the "values and preferences" statements associated with the recommendation.

## METHOD OF GUIDELINE VALIDATION

External Peer Review  
Internal Peer Review

## DESCRIPTION OF METHOD OF GUIDELINE VALIDATION

The guideline authors formulated draft recommendations prior to the conference that served as the foundation for authors to work together and critique the recommendations. Drafts of all articles including draft recommendations were available for review during the conference. A representative of each article presented potentially controversial issues in their recommendations at plenary meetings. Article authors met to integrate feedback, to consider related recommendations in other articles, and to revise their own guidelines accordingly. Authors continued this process after the conference until they reached agreement within their groups and with other author groups who had provided critical feedback. Finally, the editors of this supplement harmonized the articles and resolved remaining disagreements through facilitated discussion.

## RECOMMENDATIONS

### MAJOR RECOMMENDATIONS

The rating scheme is defined at the end of the "Major Recommendations" field.

#### **Prevention of Saphenous Vein Graft Occlusion Following Coronary Artery Bypass Grafting (CABG)**

##### **Treatment with Antiplatelet Agents**

1. For all patients with coronary artery disease, the guideline developers recommend aspirin, 75 to 162 mg/day, indefinitely (**Grade 1A**).
2. For patients undergoing CABG, the guideline developers recommend aspirin, 75 to 162 mg/day, starting 6 hours after operation over preoperative aspirin (**Grade 1A**).
3. In patients in whom bleeding prevents the administration of aspirin at 6 hours after CABG, the guideline developers recommend starting aspirin as soon as possible thereafter (**Grade 1C**).

*Underlying values and preferences:* This recommendation places a relatively high value on avoiding cardiovascular complications and a relatively low value on avoiding bleeding complications.

4. For patients undergoing CABG, the guideline developers recommend **against** the addition of dipyridamole to aspirin therapy (**Grade 1A**).
5. For patients with coronary artery disease undergoing CABG who are allergic to aspirin, the guideline developers recommend clopidogrel, 300 mg, as a



loading dose 6 hours after operation followed by 75 mg/day orally (po) (**Grade 1C+**).

6. In patients who undergo CABG for non-ST-segment elevation acute coronary syndrome (ACS), the guideline developers recommend clopidogrel, 75 mg/day, for 9 to 12 months following the procedure in addition to treatment with aspirin (**Grade 1A**).

*Underlying values and preferences:* This recommendation places a relatively high value on avoiding myocardial infarction and a relatively low value on avoiding bleeding complications.

7. For patients who received clopidogrel for ACS and are scheduled for coronary bypass surgery, the guideline developers recommend discontinuing clopidogrel for 5 days prior to the scheduled surgery (**Grade 2A**).

### **Treatment with Oral Anticoagulants**

1. For patients undergoing CABG who have no other indication for vitamin K antagonists (VKAs), the guideline developers suggest clinicians **not** administer VKAs (**Grade 2B**).
2. For patients undergoing CABG in whom oral anticoagulants are indicated, such as those with heart valve replacement, the guideline developers suggest clinicians administer VKAs in addition to aspirin (**Grade 2C**).

### **Prevention of Internal Mammary Bypass Graft Occlusion Following CABG**

#### **Aspirin with and without Dipyridamole**

1. For all patients with coronary artery disease who undergo internal mammary artery (IMA) bypass grafting, the guideline developers recommend aspirin, 75 to 162 mg/day, indefinitely (**Grade 1A**).

*Remarks:* This recommendation reflects that aspirin is indicated in all patients with coronary artery disease, irrespective of its effects on graft patency (Refer to the National Guideline Clearinghouse (NGC) summary of the American College of Chest Physicians (ACCP) guideline [Antithrombotic Therapy for Coronary Artery Disease](#)).

#### **VKAs**

1. For all patients undergoing IMA bypass grafting who have no other indication for VKAs, the guideline developers suggest clinicians **not** use VKAs (**Grade 2C**).

### **Definitions**

<b>Grade of Recommendation</b>	<b>Clarity of Risk/Benefit</b>	<b>Methodological Strength of Supporting Evidence</b>	<b>Implications</b>

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*\*These situations include RCTs with both lack of blinding and subjective outcomes, where the risk of bias in measurement of outcomes is high, or RCTs with large loss to follow-up.*

### CLINICAL ALGORITHM(S)

None provided

## EVIDENCE SUPPORTING THE RECOMMENDATIONS

### TYPE OF EVIDENCE SUPPORTING THE RECOMMENDATIONS

The type of supporting evidence is identified and graded for each recommendation (see "Major Recommendations").

## BENEFITS/HARMS OF IMPLEMENTING THE GUIDELINE RECOMMENDATIONS

### POTENTIAL BENEFITS

Antithrombotic therapy has the potential for improving patency rates in saphenous vein and internal mammary bypass grafts, and thus preventing vein graft occlusion.

## POTENTIAL HARMS

Bleeding is the primary complication of antithrombotic therapy in patients with saphenous vein bypass grafts.

## QUALIFYING STATEMENTS

### QUALIFYING STATEMENTS

#### Interpreting the Recommendations

Clinicians, third-party payers, institutional review committees, or the courts should not construe these guidelines in any way as absolute dictates. In general, anything other than a **Grade 1A** recommendation indicates that the article authors acknowledge that other interpretations of the evidence, and other clinical policies, may be reasonable and appropriate. Even **Grade 1A** recommendations will not apply to all circumstances and all patients. For instance, the guideline developers have been conservative in their considerations of cost and have seldom downgraded recommendations from **Grade 1** to **Grade 2** on the basis of expense. As a result, in jurisdictions in which resource constraints are severe, alternative allocations may serve the health of the public far better than some of the interventions that are designated as **Grade 1A**. This will likely be true for all less industrialized countries and, with the increasing promotion of expensive drugs with marginal benefits, may be increasingly true for wealthier nations.

Similarly, following **Grade 1A** recommendations will at times not serve the best interests of patients with atypical values or preferences or of those whose risks differ markedly from those of the usual patient. For instance, consider patients who find anticoagulant therapy extremely aversive, either because it interferes with their lifestyle (e.g., prevents participation in contact sports) or because of the need for monitoring. Clinicians may reasonably conclude that following some **Grade 1A** recommendations for anticoagulation therapy for either group of patients will be a mistake. The same may be true for patients with particular comorbidities (e.g., a recent gastrointestinal bleed or a balance disorder with repeated falls) or other special circumstances (e.g., very advanced age) that put them at unusual risk.

The guideline developers trust that these observations convey their acknowledgment that no recommendations or clinical practice guidelines can take into account the often compelling and unique features of individual clinical circumstances. No clinician, and no body charged with evaluating a clinician's actions, should attempt to apply these recommendations in a rote or blanket fashion.

#### Limitations of Guideline Development Methods

The limitations of these guidelines include the possibility that some authors followed this methodology more closely than others, although the development process was centralized and supervised by the editors. Second, it is possible that the guideline developers missed relevant studies despite the comprehensive searching process. Third, the guideline developers did not centralize the

methodological evaluation of all studies to facilitate uniformity in the validity assessments of the research incorporated into these guidelines. Fourth, if high-quality meta-analyses were unavailable, the guideline developers did not statistically pool primary study results using meta-analysis. Finally, sparse data on patient preferences and values, resources, and other costs represent additional limitations that are inherent to most guideline development methods.

## IMPLEMENTATION OF THE GUIDELINE

### DESCRIPTION OF IMPLEMENTATION STRATEGY

#### Guideline Implementation Strategies

A full review of implementation strategies for practice guidelines is provided in the companion document titled "Antithrombotic and Antithrombolytic Therapy: From Evidence to Application." The review suggests that there are few implementation strategies that are of unequivocal, consistent benefit, and that are clearly and consistently worth resource investment. The following is a summary of the recommendations (see "Major Recommendations" for a definition of the recommendation grades).

To encourage uptake of guidelines, the guideline developers recommend that appreciable resources be devoted to distribution of educational material (**Grade 2B**).

They also suggest that:

- Few resources be devoted to educational meetings (**Grade 2B**)
- Few resources be devoted to educational outreach visits (**Grade 2A**)
- Appreciable resources be devoted to computer reminders (**Grade 2A**)
- Appreciable resources be devoted to patient-mediated interventions to encourage uptake of the guidelines (**Grade 2B**)
- Few resources be devoted to audit and feedback (**Grade 2B**)

### IMPLEMENTATION TOOLS

Patient Resources  
Personal Digital Assistant (PDA) Downloads  
Quick Reference Guides/Physician Guides  
Resources  
Slide Presentation  
Tool Kits

For information about [availability](#), see the "Availability of Companion Documents" and "Patient Resources" fields below.

## INSTITUTE OF MEDICINE (IOM) NATIONAL HEALTHCARE QUALITY REPORT CATEGORIES

### IOM CARE NEED

Getting Better

## **IOM DOMAIN**

Effectiveness

### **IDENTIFYING INFORMATION AND AVAILABILITY**

#### **BIBLIOGRAPHIC SOURCE(S)**

Stein PD, Schunemann HJ, Dalen JE, Gutterman D. Antithrombotic therapy in patients with saphenous vein and internal mammary artery bypass grafts: the Seventh ACCP Conference on Antithrombotic and Thrombolytic Therapy. Chest 2004 Sep;126(3 Suppl):600S-8S. [49 references] [PubMed](#)

#### **ADAPTATION**

Not applicable: The guideline was not adapted from another source.

#### **DATE RELEASED**

2001 Jan (revised 2004 Sep)

#### **GUIDELINE DEVELOPER(S)**

American College of Chest Physicians - Medical Specialty Society

#### **SOURCE(S) OF FUNDING**

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#### **GUIDELINE COMMITTEE**

American College of Chest Physicians Consensus Panel on Antithrombotic and Thrombolytic Therapy

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## **FINANCIAL DISCLOSURES/CONFLICTS OF INTEREST**

Dr. Stein received honoraria for speaking at educational events from Aventis Pharma and Dupont Pharma. He was a consultant for Shiley several years ago.

Dr. Schünemann has received research funding from AstraZeneca, Boehringer Ingelheim, Pfizer, and Amgen Inc. He has received honoraria and consultant fees from AstraZeneca, Boehringer Ingelheim, and Amgen that were deposited into research accounts at the University of Buffalo and McMaster University.

Dr. Dalen has received honoraria as a consultant for DuPont Pharma (now Bristol-Myers Squibb), AstraZeneca and Sanofi-Organon.

Dr. Gutterman owns stock in Johnson & Johnson.

## **GUIDELINE STATUS**

This is the current release of the guideline.

This guideline updates a previous version: Stein PD, Dalen JE, Goldman S, Theroux P. Antithrombotic therapy in patients with saphenous vein and internal mammary artery bypass grafts. Chest 2001 Jan;119(1 Suppl):278S-282S.

## **GUIDELINE AVAILABILITY**

Electronic copies: Available from the [Chest - The Cardiopulmonary and Critical Care Journal](#).

Print copies: Available from the American College of Chest Physicians, Products and Registration Division, 3300 Dundee Road, Northbrook IL 60062-2348.

## **AVAILABILITY OF COMPANION DOCUMENTS**

The following are available:

- The Seventh ACCP Conference on Antithrombotic and Thrombolytic Therapy. Evidence-based guidelines. Northbrook, IL: ACCP, 2004 Sep.
- Methodology for guideline development for the Seventh American College of Chest Physicians Conference on Antithrombotic and Thrombolytic Therapy. Northbrook, IL: ACCP, 2004 Sep.
- Applying the grades of recommendation for antithrombotic and thrombolytic therapy: The Seventh ACCP Conference on Antithrombotic and Thrombolytic Therapy. Northbrook, IL: ACCP, 2004 Sep.
- Hemorrhagic complications of anticoagulant treatment: The Seventh ACCP Conference on Antithrombotic and Thrombolytic Therapy. Northbrook, IL: ACCP, 2004 Sep.
- Antithrombotic and thrombolytic therapy: from evidence to application: The Seventh ACCP Conference on Antithrombotic and Thrombolytic Therapy. Northbrook, IL: ACCP, 2004 Sep.
- Platelet-active drugs: the relationships among dose, effectiveness, and side effects: The Seventh ACCP Conference on Antithrombotic and Thrombolytic Therapy. Northbrook, IL: ACCP, 2004 Sep.

Electronic copies: Available from the [Chest - The Cardiopulmonary and Critical Care Journal Web site](#).

Print copies: Available from the American College of Chest Physicians (ACCP), Products and Registration Division, 3300 Dundee Road, Northbrook IL 60062-2348.

The following is also available:

- Seventh ACCP Conference on Antithrombotic and Thrombolytic Therapy: Evidence-based guidelines; quick reference guide. Northbrook, IL: ACCP, 2004 Sep. Personal Digital Assistant (PDA) download available at [ACCP Web site](#).

Additional implementation tools are also available:

- Clinical resource: antithrombotic and thrombolytic therapy. Northbrook, IL. ACCP, 2004. Ordering information: Available from the [ACCP Web site](#).

## **PATIENT RESOURCES**

The following is available:



- A patient's guide to antithrombotic and thrombolytic therapy. In: Clinical resource: antithrombotic and thrombolytic therapy. Northbrook (IL): American College of Chest Physicians (ACCP). 2004.

Ordering information is available from the [ACCP Web site](#).

Please note: This patient information is intended to provide health professionals with information to share with their patients to help them better understand their health and their diagnosed disorders. By providing access to this patient information, it is not the intention of NGC to provide specific medical advice for particular patients. Rather we urge patients and their representatives to review this material and then to consult with a licensed health professional for evaluation of treatment options suitable for them as well as for diagnosis and answers to their personal medical questions. This patient information has been derived and prepared from a guideline for health care professionals included on NGC by the authors or publishers of that original guideline. The patient information is not reviewed by NGC to establish whether or not it accurately reflects the original guideline's content.

## **NGC STATUS**

This summary was completed by ECRI on July 30, 2001. The information was verified by the guideline developer on September 27, 2001. This NGC summary was updated by ECRI on December 9, 2004. The updated information was verified by the guideline developer on January 12, 2005.

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