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Evidence-Based Management of Anticoagulant Therapy : Antithrombotic Therapy and Prevention of Thrombosis, 9th ed: American College of Chest Physicians Evidence-Based Clinical Practice Guidelines

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Chest 2012;141;e152S-e184S DOI 10.1378/chest.11-2295

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Supplemental material related to this article is available at: http://chestjournal.chestpubs.org/content/suppl/2012/02/03/141.2_suppl. e152S.DC1.html

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ANTITHROMBOTIC THERAPY AND PREVENTION OF THROMBOSIS, 9TH ED: ACCP GUIDELINES

Evidence-Based Management of Anticoagulant Therapy

Antithrombotic Therapy and Prevention of Thrombosis, 9th ed: American College of Chest Physicians Evidence-Based Clinical Practice Guidelines

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Background: High-quality anticoagulation management is required to keep these narrow therapeutic index medications as effective and safe as possible. This article focuses on the common important management questions for which, at a minimum, low-quality published evidence is available to guide best practices.

Methods: The methods of this guideline follow those described in Methodology for the Development of Antithrombotic Therapy and Prevention of Thrombosis Guidelines: Antithrombotic Therapy and Prevention of Thrombosis, 9th ed: American College of Chest Physicians Evidence-Based Clinical Practice Guidelines in this supplement.

Results: Most practical clinical questions regarding the management of anticoagulation, both oral and parenteral, have not been adequately addressed by randomized trials. We found sufficient evidence for summaries of recommendations for 23 questions, of which only two are strong rather than weak recommendations. Strong recommendations include targeting an international normalized ratio of 2.0 to 3.0 for patients on vitamin K antagonist therapy (Grade 1B) and not routinely using pharmacogenetic testing for guiding doses of vitamin K antagonist (Grade 1B). Weak recommendations deal with such issues as loading doses, initiation overlap, monitoring frequency, vitamin K supplementation, patient self-management, weight and renal function adjustment of doses, dosing decision support, drug interactions to avoid, and prevention and management of bleeding complications. We also address anticoagulation management services and intensive patient education.

Conclusions: We offer guidance for many common anticoagulation-related management problems. Most anticoagulation management questions have not been adequately studied.

CHEST 2012; 141(2)(Suppl):e152S-e184S

Abbreviations: AMS = anticoagulation management service; aPTT = activated partial thromboplastin time; COX = cyclo-oxygenase; FFP = fresh frozen plasma; HR = hazard ratio; INR = international normalized ratio; LMWH = low-molecular-weight heparin; NSAID = nonsteroidal antiinflammatory drug; PCC = prothrombin complex concentrate; PE = pulmonary embolism; POC = point-of-care; PSM = patient self-management; PST = patient self-testing; RCT = randomized controlled trial; RR = risk ratio; SC = subcutaneous; TTR = time in therapeutic range; UFH = unfractionated heparin; VKA = vitamin K antagonist

SUMMARY OF RECOMMENDATIONS

Note on Shaded Text: Throughout this guideline, shading is used within the summary of recommendations sections to indicate recommendations that are newly added or have been changed since the publication of Antithrombotic and Thrombolytic Therapy: American College of Chest Physicians Evidence-Based Clinical Practice Guidelines (8th Edition). Recommendations that remain unchanged are not shaded.

2.1. For patients sufficiently healthy to be treated as outpatients, we suggest initiating vitamin K antagonist (VKA) therapy with warfarin 10 mg daily for the first 2 days followed by dosing based on international normalized ratio (INR) measurements rather than starting with the estimated maintenance dose (Grade 2C).

2.2. For patients initiating VKA therapy, we recommend against the routine use of pharmacogenetic testing for guiding doses of VKA (Grade 1B).

2.3. For patients with acute VTE, we suggest that VKA therapy be started on day 1 or 2 of LMWH or UFH therapy rather than waiting for several days to start (Grade 2C).

3.1. For patients taking VKA therapy with consistently stable INRs, we suggest an INR testing frequency of up to 12 weeks rather than every 4 weeks (Grade 2B).

3.2. For patients taking VKAs with previously stable therapeutic INRs who present with a single out-of-range INR of ≤ 0.5 below or above therapeutic, we suggest continuing the current dose and testing the INR within 1 to 2 weeks (Grade 2C).

3.3. For patients with stable therapeutic INRs presenting with a single subtherapeutic INR

Funding/Support: The Antithrombotic Therapy and Prevention of Thrombosis, 9th ed: American College of Chest Physicians Evidence-Based Clinical Practice Guidelines received support from the National Heart, Lung, and Blood Institute [R13 HL104758] and Bayer Schering Pharma AG. Support in the form of educational grants were also provided by Bristol-Myers Squibb; Pfizer, Inc; Canyon Pharmaceuticals; and sanofi-aventis US.

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DOI: 10.1378/chest.11-2295

value, we suggest against routinely administering bridging with heparin (Grade 2C).

3.4. For patients taking VKAs, we suggest against routine use of vitamin K supplementation (Grade 2C).

3.5. (Best Practices Statement) We suggest that health-care providers who manage oral anticoagulation therapy should do so in a systematic and coordinated fashion, incorporating patient education, systematic INR testing, tracking, follow-up, and good patient communication of results and dosing decisions.

3.6. For patients treated with VKAs who are motivated and can demonstrate competency in self-management strategies, including the self-testing equipment, we suggest patient selfmanagement (PSM) rather than usual outpatient INR monitoring (Grade 2B). For all other patients, we suggest monitoring that includes the safeguards in our best practice statement 3.5.

3.7. For dosing decisions during maintenance VKA therapy, we suggest using validated decision support tools (paper nomograms or computerized dosing programs) rather than no decision support (Grade 2C).

Remarks: Inexperienced prescribers may be more likely to improve prescribing with use of decision support tools than experienced prescribers.

3.8. For patients taking VKAs, we suggest avoiding concomitant treatment with nonsteroidal antiinflammatory drugs (NSAIDs), including cyclooxygenase (COX)-2-selective NSAIDs, and certain antibiotics (see Table 8) (Grade 2C).

For patients taking VKAs, we suggest avoiding concomitant treatment with antiplatelet agents except in situations where benefit is known or is highly likely to be greater than harm from bleeding, such as patients with mechanical valves, patients with acute coronary syndrome, or patients with recent coronary stents or bypass surgery (Grade 2C).

4.1. For patients treated with VKAs, we recommend a therapeutic INR range of 2.0 to 3.0 (target INR of 2.5) rather than a lower (INR < 2) or higher (INR 3.0-5.0) range (Grade 1B).

4.2. For patients with antiphospholipid syndrome with previous arterial or venous thromboembolism, we suggest VKA therapy titrated to a moderate-intensity INR range (INR 2.0-3.0) rather than higher intensity (INR 3.0-4.5) (Grade 2B).

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Revision accepted August 31, 2011.

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5.0. For patients eligible to discontinue treatment with VKA, we suggest abrupt discontinuation rather than gradual tapering of the dose to discontinuation (Grade 2C).

6.1. For patients starting IV unfractionated heparin (UFH), we suggest that the initial bolus and the initial rate of the continuous infusion be weight adjusted (bolus 80 units/kg followed by 18 units/kg per h for VTE; bolus 70 units/kg followed by 15 units/kg per h for cardiac or stroke patients) or use of a fixed dose (bolus 5,000 units followed by 1,000 units/h) rather than alternative regimens (Grade 2C).

6.2. For outpatients with VTE treated with subcutaneous (SC) UFH, we suggest weight-adjusted dosing (first dose 333 units/kg, then 250 units/kg) without monitoring rather than fixed or weight-adjusted dosing with monitoring (Grade 2C).

7.1. For patients receiving therapeutic LMWH who have severe renal insufficiency (calculated creatinine clearance < 30 mL/min), we suggest a reduction of the dose rather than using standard doses (Grade 2C).

8.1. For patients with VTE and body weight over 100 kg, we suggest that the treatment dose of fondaparinux be increased from the usual 7.5 mg to 10 mg daily SC (Grade 2C).

9.1.

(a) For patients taking VKAs with INRs between 4.5 and 10 and with no evidence of bleeding, we suggest against the routine use of vitamin K (Grade 2B).

(b) For patients taking VKAs with INRs > 10.0 and with no evidence of bleeding, we suggest that oral vitamin K be administered (Grade 2C).

9.2. For patients initiating VKA therapy, we suggest against the routine use of clinical prediction rules for bleeding as the sole criterion to withhold VKA therapy (Grade 2C).

9.3. For patients with VKA-associated major bleeding, we suggest rapid reversal of anticoagulation with four-factor prothrombin complex concentrate (PCC) rather than with plasma. (Grade 2C).

We suggest the additional use of vitamin K 5 to 10 mg administered by slow IV injection rather

than reversal with coagulation factors alone (Grade 2C).

This article deals with the evidence regarding managing anticoagulant therapy, that is, oral vitamin K antagonists (VKAs), heparins, and fondaparinux. Separate articles address the pharmacology of these drugs.¹ The questions that we address reflect those commonly posed in clinical practice.

1.0 Methods

The methods for the development of this article's recommendations follow those developed for the Antithrombotic Therapy and Prevention of Thrombosis, 9th ed: American College of Chest Physicians Evidence-Based Clinical Practice Guidelines.² Although we aimed to summarize and use randomized controlled trial (RCT) evidence to inform recommendations for clinicians, we found only lower-quality evidence to address most of our questions. At the onset of our review process, our panel decided to limit the recommendations to questions in which evidence met a minimum threshold for quality: at least one comparative study with \geq 50 patients per group with contemporaneous or historical controls reporting on patient-important outcomes or closely related surrogates. Despite this low threshold, evidence was unavailable for several important clinical management questions. When randomized trials were available, confidence in estimates often decreased because of indirectness (surrogate outcomes) and imprecision (wide CIs).

This article does not address anticoagulation management issues specific to pregnancy or to children. Issues believed to be specific to a particular diagnosis, such as VTE or atrial fibrillation, are dealt with in those specific articles of this supplement. Table 1 presents the questions for which we found evidence that met our quality threshold, including the relevant populations, interventions, comparators, and outcomes.

2.0 VKA—INITIATION OF THERAPY

2.1 Initial Dose Selection—Loading Dose

Loading doses of VKA may be worth considering where rapid attainment of therapeutic international normalized ratio (INR) is required and considered safe, primarily for patients with VTE. Predictable and timely achievement of therapeutic INRs without increased risk of bleeding or recurrent thromboembolic events avoids the inconvenience and pain of prolonged administration of subcutaneous (SC) lowmolecular-weight heparin (LMWH) and facilitates early patient discharge and eligibility for outpatient dosing nomograms. Two large case series^{5,6} involving a total of 1,054 outpatients suggest that a nomogram specifying a 10-mg loading dose is safe, with a recurrent VTE rate of 1.9% and a major bleeding rate of 1.0% at 3 months follow-up.⁵ However, pooling across both studies suggests that only 49.3% of participants followed the nomogram completely.

Table 2 and Table S1 (tables that contain an "S" before the number denote supplementary tables

			-	FILO		
Section	Informal Question	Population	Intervention	Comparator	Outcome	Comment
		2.(2.0 VKAs—initiation of therapy	À		
2.1. Loading doses of VKA	Is a loading dose of VKA superior to no loading dose?	Patients taking VKA	Loading dose	No loading dose	Hemorrhage, thromboembolic events, time to therapeutic range, rates of supratherapeutic or subtheramentic INR	
2.2. Dose by Pharmacogenetics	Should the initial dose of VKA be based on pharmacogenetic testing?	Patients taking VKA	Analysis of CYP2C9, VKORC1, and other polymorphisms	No pharmacogenetic testing	Hemorrhage, thromboembolic events, time to therapeutic range, rates of supratherapeutic or subtherapeutic INR	
2.3. Initiation overlap	Should VKA be started simultaneously with heparin rather than delayed a few days?	Patients treated for acute thromboembolism (or other high-risk situation requiring long-term VKA)	Simultaneous start	Initial heparin followed by overlap with VKA	Hemorrhage, thromboembolic events, time to therapeutic range, rates of supratherapeutic or subtherapeutic INR, resource utilization (hospital stay)	
		3.01	3.0 VKAs—maintenance treatment	ent		
3.1. INR monitoring frequency	How frequently should treatment be monitored initially and once dose and INR have been stable for months?	Patients taking VKA	Higher frequency	Lower frequency	Hemorrhage, thromboembolic events, time to therapeutic range	
3.2. Single out-of-range INR—dose adjustment	Should the VKA dose change for a single deviating INR in otherwise stable patients?	Patients taking VKA	Dose adjustment	Continue as usual	Hemorrhage, thromboembolic events, time in therapeutic range	
3.3. Bridging for subtherapeutic INR	Does bridging anticoagulant therapy improve outcomes for low INR?	Patients taking VKA with subtherapeutic INR	Dose management and overlapping with heparin	Only dose management	Hemorrhage, thromboembolic events, time in therapeutic range	
3.4. Vitamin K supplementation	Can outcomes be improved with low-dose vitamin K supplementation or dietary manipulation?	Patients taking VKA with variability of INR	Cotherapy with small- dose vitamin K or with dietary modification	No vitamin K	Hemorrhage, thromboembolic events, time in therapeutic range	
3.5. Dose management services	Dose management services: does a specialized AMS improve outcomes?	Patients taking VKA	AMS care	Usual care (primary care or regular hospital physicians)	Hemorrhage, thromboembolic events, time in therapeutic range, resource utilization	
3.6. Patient self-testing and self-monitoring	Does self-monitoring of anticoagulation improve outcomes	Patients taking VKA	Use of point-of-care monitor at home to measure INR and to adiust VKA dose	Usual care or AMS care	Hemorrhage, thromboembolic events, time in therapeutic range, resource utilization	

Table 1—Structured Clinical Questions

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Section	Informal Question	Population	Intervention	Comparator	Outcome	Comment
3.7. Dosing decision support	Does dosing decision support improve	Patients taking VKA	Computer software, manual algorithms	Usual care	Hemorrhage, thromboembolic events, time in therapeutic	
	outcomes				range, resource utilization	
3.8. Drug interactions	What anticoagulant drug	Patients taking	Patients starting or	Patients not taking	Hemorrhage, thromboembolic	Limited to randomized
to avoid	or food interactions are	anticoagulants	stopping potentially	potentially	events, time in therapeutic	trials of clinical
	important enough to		interacting drugs	interacting drugs	range	outcomes or large
	avoid the interacting					observational
	drug while patients take					studies
	anucoaguiants					
			4.0 VKAs—monitoring			
4.1. Optimal INR	What is the optimal INR	Patients taking VKA	Optimal INR range	Wider INR range	Hemorrhage, thromboembolic	
Iallge	outcomes?				CVCIILS	
4.2. Optimal INR range	Should high-risk groups	Patients with APS (or other	More intensive INR	Standard INR	Hemorrhage,	
for high-risk groups	(such as APS, cancer)	high-risk feature) and	therapeutic range	therapeutic	thromboembolic	
	be treated more	taking VKA	or alternative assay	range	events, time in	
	intensively?				therapeutic range	
		5.1	5.0 VKAs—discontinuing therapy	ıpy		
5.1 Tapering vs abrupt	How should VKA be	Patients discontinuing	Tapered discontinuation	Abrupt discontinuation	Hemorrhage, thromboembolic	
discontinuation	discontinued?	VKA			events, time to normal anticoaorulation status	
		6.0	6.0 Parenteral anticoagulants—UFH	JFH	с	
6.1. UFH—dose	Should the initial bolus	Patients treated with	Weight-adjusted dose	Fixed dose	Hemorrhage, thromboembolic	
adjustment by	dose or maintenance	IV UFH	`		events, time in therapeutic	
weight	dose be weight				range	
	adjusted?					
6.2. SC UFH dose	Should doses of SC UFH	Patients treated with SC	Weight-adjusted dose	Fixed dose with or	Hemorrhage, thromboembolic	
adjustment and	be adjusted for weight	UFH	with and without	without aPTT	events, time in therapeutic	
monitoring	and monitored by a pTTP		aPTT monitoring	monitoring	range	
		7.0 P	7.0 Parenteral anticoagulants—LMWH	HMM		
7.1. LMWH—dose	Should doses be	Patients with mild to	Dose adjustment	Dose adjustment only	Hemorrhage, thromboembolic	
modification by renal	modified for renal	moderate renal failure	according to renal	by body weight or	events	
function	function?	treated with LMWH	function	no dose adjustment		
7.2. LMWH—dose	Can doses be	Patients treated with	Doses administered daily	Doses administered	Hemorrhage, thromboembolic	Moved to Kearon
Frequency	administered daily	LMWH		bid	events	et al ³
	instead of twice daily?					in this supplement
7.3. LMWH dose	Should the dose be	Obese or significantly	Dose adjustment	Standard dose	Hemorrhage, thromboembolic	Gould et al ⁴ in this
modification by	weight adjusted?	underweight patients	according to body		events	supplement
weight tor		receiving prophylaxis	weight			
propriyraxis		MILLI LIM WELL				(Continued)
						1 manual 1

Table 1—Continued

Anticoagulant Therapy

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Section	Informal Question	Population	Intervention	Comparator	Outcome	Comment
		8.0 Pare	8.0 Parenteral anticoagulants—fondaparinux	parinux		
8.1. Fondaparinux dose management	Should the dose be weight adjusted?	Obese or significantly underweight patients receiving fondaparinux	Dose adjustment according to body weight	Standard dose	Hemorrhage, thromboembolic events	
		9.0 Prevention and	9.0 Prevention and management of anticoagulant complications	ant complications		
9.1. Vitamin K for high INR without bleeding	Does vitamin K improve outcomes for high INRs without bleeding?	Patients taking VKA with high INR (>4.5) without bleeding	KA dose management plus use of vitamin K	Only dose management (= holding the VKA until therapeutic)	Hemorrhage, thromboembolic events, time to therapeutic range, rates of overcorrection of INR	
9.2. Predicting anticoagulant- associated bleeding	Does a bleeding clinical prediction rule improve outcomes? Which prediction rule should be used?	Patients taking anticoagulant therapy or considering therapy	Use of a bleeding clinical prediction rule to guide therapy (dose and whether to give)	No clinical prediction rule or alternate prediction rule	Hemorrhage, thromboembolic events, choice of therapy	
9.3. Treatment of	What is the most	Patients actively bleeding	Vitamin K, FFP, PCC,	One of the other	Time to resolution of bleeding,	
anticoagulant-	effective and safe	from excessive	recombinant factor	treatments or	bleeding complications,	
related bleeding	urgent treatment of anticoagulant-related bleeding?	anticoagulation who need to have the bleeding stopped urgently	VIIa	vitamin K alone	thromboembolism rates, resource utilization	
9.4. Investigating anticoagulant- associated	When is it appropriate to investigate anticoagulant-	Patients taking VKAs with therapeutic INRs and major bleeding episodes	Patients who bleed	Patients who do not bleed	Incidence of malignancy, ulcer disease, other serious or treatable outcome	
Bitcouilg	associated Diceding:		10.0 Other			
10.1. Intensive patient education	Does additional structured patient education improve outcomes related to anticoaeulation?	Patients who are to take or are taking VKAs or parenteral anticoagulants	Patient education on benefits, harms, and use of anticoagulants	Usual care	Hemorrhage, thromboembolic events, time in therapeutic range, compliance	

Table 1—Continued

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VKORC1 = vitamin K epoxide reductase complex 1.

Table 2—[Section 2.1] Warfarin 10 mg Loading Dose Nomogram Compared With Warfarin 5 mg Loading Dose Nomogram for Warfarin Initiation^{7,8,10,11}

				Anticip	ated Absolute Effects
Outcomes	No. of Participants (Studies), Follow-up	Quality of the Evidence (GRADE)	Relative Effect (95% CI)	Risk With Warfarin 5 mg Loading Dose Nomogram	Risk Difference With Warfarin 10 mg Loading Dose Nomogram (95% CI)
Bleeding events	420 (3 studies ^{a-c}), 5-90 d ^d	Very low ^e g due to indirectness, imprecision	OR 1.90 (0.17-21.1)	5 per 1,000	0 more per 1,000 (from 10 fewer to 20 more) ^h
Recurrent VTE	420 (3 studies ^{a-c})	Very low ^{e-g} due to indirectness, imprecision	Not estimable	0 per 1,000	$\begin{array}{c} 10 \text{ more per } 1,000 \text{ (from 30 more} \\ \text{to 0 more})^i \end{array}$

GRADE = Grades of Recommendations, Assessment, Development, and Evaluation. See Table 1 legend for expansion of other abbreviation. ^aAll pooled studies included only patients with acute VTE. Studies from which data could be pooled are Kovacs et al,⁹ Quiroz et al,¹⁰ and Schulman et al.¹¹

^bMinimal loss to follow-up; adherence to intention-to-treat principle in two of three studies; follow-up period short but adequate for this outcome; any lack of blinding should not impact objective outcome (laboratory value, INR); adequate allocation concealment; sample size calculations reported for two of three studies.

eResults based on only three studies; one study shows no difference; one shows statistically significant reduction in time to therapeutic INR; and one had two parts to it, where one showed statistically significant reduction and the other did not.

^dMean follow-up period of 5 d for patients in the loading dose warfarin group from Schulman et al¹¹ (this was the shortest period, only mean is available). ^eData collectors unblinded.

Indirect given application aimed at outpatients with VTE; follow-up period is very short in two of three studies (5 d-2 wk).

^gNo studies were powered to detect differences in bleeding events between groups. Number of events is too sparse to draw any conclusions. ^hVery small number of events; risk difference calculated.

OR not estimable; absolute risk difference calculated.

not contained in the body of the article and available instead in an online data supplement; see the "Acknowledgments" for more information) summarizes our confidence in effect estimates and main findings from a meta-analysis of five RCTs of loading dose vs no loading dose of warfarin.⁷⁻¹¹ The table shows that clinical outcomes, where documented, were similar between the groups. The studies typically measured time to therapeutic range of anticoagulation as the primary outcome and the patients were mainly those starting treatment (not prophylaxis) for VTE. Many of those treated as inpatients at the time of the study would, in current practice, be treated as outpatients.

Two studies by a single group^{7,8} compared a 10-mg loading dose to 5 mg daily for the first 2 days. Both included primarily inpatients, and one did not report recurrent VTE.8 The concentrations of protein C and factor VII, but not those of factor II or X, decreased faster in the 10-mg group than in the 5-mg group⁸; an increased risk of recurrent thromboembolism, however, has not been demonstrated in any of the studies presumably because initiation overlaps with heparin or LMWH. Quiroz et al¹⁰ compared 5 vs 10 mg initial warfarin dosing in 50 inpatients and reported no difference in median time to two consecutive therapeutic INRs. This study had only a 2-week follow-up and excluded 322 of the 372 patients screened. Another study compared loading dose vs standard warfarin initiation for patients with VTE and showed a shorter time to a therapeutic INR (3.3 vs 4.3 days).¹¹ Finally, Kovacs et al⁹ found that the use of a 10- vs 5-mg initiation nomogram with 210 outpatients resulted in shorter mean time to therapeutic INR of 4.2 vs 5.6 days. The proportion therapeutic by day 5 was also significantly better at 86% vs 45% in the 10- vs 5-mg group, respectively. All studies followed the initiation period with INR-based dose adjustment.

Recommendation

2.1. For patients sufficiently healthy to be treated as outpatients, we suggest initiating VKA therapy with warfarin 10 mg daily for the first 2 days followed by dosing based on INR measurements rather than starting with the estimated maintenance dose (Grade 2C).

2.2 Initial Dose Selection and Pharmacogenetic Testing

Selection of the initial and maintenance doses of VKA therapy usually has been based on subjective estimates of patient age, size, nutritional status, and organ function. In section 2.1, we suggest a standard short loading dose for outpatients. Theoretically, individual patient pharmacogenetic testing of CYP2C9 (cytochrome P450 2C9), which is involved with VKA metabolism and VKORC1 (vitamin K epoxide reductase complex 1, the VKA target), might improve VKA therapy through more-accurate dose selection. There are four RCTs of pharmacogenetic testing-based dosing vs standard dosing; all addressed warfarin initiation.¹²⁻¹⁵ The studies included patients with

artificial heart valves, atrial fibrillation, or acute VTE. All studies were small (total n = 544). None showed any difference in thrombotic events, major bleeding, or survival (Table S2).

Hillman et al¹² conducted a pilot study of 38 patients. Caraco et al¹³ randomized 283 patients but excluded 92 for reasons such as failure to follow warfarin dosing instructions. Huang et al¹⁵ included 121 valve inpatients and showed improvement in time to therapeutic range; the control group, however, used a substandard 2.5-mg daily regimen. Anderson et al,¹⁴ who had the highest methodologic quality, studied inpatients in which the control group experienced close INR monitoring following a loading-dose strategy. The investigators found no difference in time in therapeutic range or time to therapeutic range. A systematic review also concluded that there is a lack of evidence to support using pharmacogenetic testing to guide VKA dosing.¹⁶

Several recent economic evaluations have assessed the cost-effectiveness of pharmacogenetic testing to guide VKA (warfarin) initiation.¹⁷⁻¹⁹ The results of these studies estimated the incremental cost at \sim \$50,000 to \$170,000 per quality-adjusted life year gained, but in sensitivity analyses, the incremental cost-effectiveness ratios were as high as \$200,000 to \$300,000 per quality-adjusted life year and included scenarios in which pharmacogenetic testing led to poorer patient outcomes. These results would be judged as not cost-effective by most drug policy experts.

Recommendation

2.2. For patients initiating VKA therapy, we recommend against the routine use of pharmacogenetic testing for guiding doses of VKA (Grade 1B).

2.3 Initiation Overlap for Heparin and VKA

Historically, clinicians administered IV unfractionated heparin (UFH) to inpatients for 5 to 7 days with subsequent initiation of a VKA, leading to a total duration of IV UFH of 10 to 14 days. More recently, VKA therapy has been initiated on the first or second day of heparin therapy, leading to shorter durations of heparin and earlier discharge from the hospital.

Table 3 (and Table S3) summarizes the evidence from a meta-analysis of 807 patients in four RCTs

				Anticipat	ed Absolute Effects
Outcomes	No. of Participants (Studies), Follow-up	Quality of the Evidence (GRADE)	Relative Effect (95% CI)	Risk With Late	Risk Difference With VKA Started Early (95% CI)
Death	807 (4 studies), 3-6 mo	Low ^{a-c} due to inconsistency and imprecision	RR 1.28 (0.43-3.85)	58 per 1,000	16 more per 1,000 (from 33 fewer to 166 more)
Recurrent thromboembolism DVT: venography, Doppler ultrasonography or impedance plethysmography. PE: lung scanning Left ventricle thrombus: 2-dimensional transthoracic echocardiography	807 (4 studies), 3-6 mo	Low ^{e,d} due to risk of bias and imprecision	RR 0.92 (0.46-1.82)	41 per 1,000	3 fewer per 1,000 (from 22 fewer to 33 more)
Major bleeding-required blood transfusion, bleeding in body cavity, bleeding that required anticoagulation withdrawal or intracranial or retroperitoneal, or bleeding that led to a hemoglobin level decrease of ≥ 2 g/dL or to death	807 (4 studies), 0.5-6 mo	Low ^{e,d} due to risk of bias and imprecision	RR 1.22 (0.58-2.56)	33 per 1,000	7 more per 1,000 (from 14 fewer to 51 more)
Hospital utilization	536 (3 studies)	High		The mean hospital utilization in the control groups was 14 d	The mean hospital utilization in the intervention groups was 4.07 lower (4.76 to 3.37 lower)

PE = pulmonary embolism; RR = risk ratio. See Table 2 legend for expansion of other abbreviation.

^aFor three out of four studies, concealment of allocation was unclear. Lack of blinding of health-care professionals in some studies.

^bThe value for I2 test for death was 55%, and therefore, it was rated down for inconsistency.

^eThe 95% confidence intervals around the absolute risk values were very wide for this outcome.

 d Potential limitations in design for this outcome: allocation sequence concealment was not reported in three out of four studies; health-care professionals blinded in only one study (Hull et al²⁰) (outcome assessors were blinded in three of four studies).

addressing this issue (F. Qayyum, unpublished data, 2011). These trials compared early start (day 1 or 2 of heparin) vs late start (days 3-10 of heparin) for the VKA therapy together with UFH or LMWH therapy. Two studies^{20,21} enrolled patients with DVT only, one enrolled patients with DVT or pulmonary embolism (PE),²² and the fourth included patients with left ventricular mural thrombosis.²² There were no differences between early vs late initiation of VKA for the outcomes of recurrent VTE, major bleeding, or death. Patients assigned to early initiation of VKA spent a mean of 4 fewer days in the hospital than patients assigned to late initiation of VKA. No studies have assessed early vs late initiation of VKA therapy in the outpatient setting, but we consider the results of the meta-analysis to be applicable to outpatients.

Recommendation

2.3. For patients with acute VTE, we suggest that VKA therapy be started on day 1 or 2 of LMWH or UFH therapy rather than waiting for several days to start (Grade 2C).

3.0 MAINTENANCE TREATMENT WITH VKAS

3.1 Monitoring Frequency for VKAs

The frequency of long-term INR monitoring is influenced by patient compliance, changes in health status, the addition or discontinuation of interacting medications, changes in diet, the quality of doseadjustment decisions, and whether the patient has demonstrated stable INRs.²³⁻²⁵ We define stable INRs as at least 3 months of consistent results with no need to adjust VKA dosing.26 Recall intervals for various clinical situations have not been extensively studied; rather, they evolved from routine clinical practice and expert opinion and differ substantially from one country to another.²⁷ For example, in North America, stable patients usually are tested every 4 weeks,²⁴ whereas in the United Kingdom, INR recall intervals of up to 90 days are routine.²⁸ This discussion does not apply to patients engaging in INR self-testing using portable finger-stick monitors in whom only weekly INR recall intervals have been adequately evaluated.

For patients receiving traditional laboratory-based INR monitoring, retrospective studies have found increasing INR recall intervals associated with both increased²⁹ and decreased^{26,30} time in therapeutic range (TTR). Other observational studies have suggested that for patients who demonstrate a consistent pattern of stable therapeutic INRs, allowing INR recall intervals of up to 8 weeks would not result in increased risk for bleeding or thromboembolism.³¹⁻³³

Three RCTs have evaluated the effectiveness of INR recall intervals exceeding the traditional North American standard of 4 weeks.^{23,34,35} One study compared 6- to 4-week recall intervals,³⁴ whereas another evaluated a flexible approach that allowed recall intervals of up to 12 weeks based on several factors, including the number of prior INRs, longitudinal INR variability, and the risk of adverse events expressed as a function of the INR.²³ The third study compared 4- to 12-week recall intervals using a blinded design.³⁵ None of the studies found a difference in rates of thromboembolism, bleeding, or INR control (Table 4, Table S4). The appropriate length of the recall interval depends on the duration of prior stability and foreseeable future changes in medications or disorders that affect the INR. Whatever maintenance dose interval is chosen, when adjustments to the VKA dose are required, a cycle of more-frequent INR monitoring should be completed until a consistent pattern of stable therapeutic INRs can be reestablished.³⁶

Recommendation

3.1. For patients taking VKA therapy with consistently stable INRs, we suggest an INR testing frequency of up to 12 weeks rather than every 4 weeks (Grade 2B).

3.2 Management of the Single Out-of-Range INR

A common dilemma encountered in clinical management of patients taking VKAs is what to do with an INR slightly outside the therapeutic range when

 Table 4—[Section 3.1] Prolonged INR Recall Intervals Compared With 4-Week Recall Intervals for Patients

 With a Stable INR^{23,34,35}

				Antici	pated Absolute Effects ^a
Outcomes	No. of Participants	Quality of the Evidence	Relative	Risk With 4-wk	Risk Difference With Prolonged
	(Studies), Follow-up	(GRADE)	Effect (95% CI)	Recall Intervals	INR Recall Intervals (95% CI)
Thromboembolism variously defined	994 (3 studies), 313 patient-y	Moderate ^b due to imprecision	OR 1.05 (0.28-3.97)	12 per 1,000	1 more per 1,000 (from 8 fewer to 33 more)
Major bleeding	994 (3 studies),	Moderate ^b due to	RR 1.12	33 per 1,000	4 more per 1,000 (from 14
variously defined	313 patient-y	imprecision	(0.57-2.23)		fewer to 41 more)

See Table1-3 legends for expansion of abbreviations.

^aTime frame in months.

^bWide CIs around the estimate of effect.

INRs were previously in the therapeutic range.²⁵ The question is whether the dose should be adjusted or left unchanged until the next INR is obtained.

This issue has been evaluated in two studies. An open-label RCT compared a one-time dose increase or hold vs continuing as is when the INR was slightly below or above the therapeutic range.³⁷ Randomized patients had been taking a stable warfarin dose for at least 3 months, the out-of-range INR was between 1.5 and 4.4, and the target ranges were 2.0 to 3.0 or 2.5 to 3.5. Reduced or boosted doses were usually 50% lower or higher, respectively, than the regularly scheduled dose. Results were similar at follow-up \sim 2 weeks later, with 44% outside the therapeutic range among patients randomized to a one-time dose change compared with 40% of those randomized to no dose change (OR, 1.17; 95% CI, 0.59-2.30; P = .75).

The other study evaluated the safety of not changing the usual warfarin maintenance dose in response to isolated, asymptomatic INRs of 3.2 to 3.4 in patients who had been taking warfarin for at least 30 days and had a targeted INR range of 2.0 to 3.0.³⁸ This was an observational study nested within an RCT evaluating anticoagulation management services (AMS) vs primary care management. The response to an isolated INR between 3.2 and 3.4 was to continue the same dose 78% of the time in AMS vs 47% in primary care. The proportion of patients with a therapeutic follow-up INR was not significantly different between the two groups (AMS, 63%; control, 54%). No major bleeding or thromboembolic events were observed during the 14 to 30 days follow-up in either of these studies.

The evidence from both studies suffers from relatively small sample sizes; lack of blinding; and in the second study, lack of randomization and a lack of uniformity in INR management between groups. Both studies were consistent with a dosing model developed from an observational study of 3,961 patients that suggested that warfarin doses did not need to be changed for INRs between 1.7 and 3.3.²⁵ It is reasonable to follow up with an INR after 1 to 2 weeks to exclude a progressive deviation from the therapeutic range.^{36,37}

Recommendation

3.2. For patients taking VKAs with previously stable therapeutic INRs who present with a single out-of-range INR of ≤ 0.5 below or above therapeutic, we suggest continuing the current dose and testing the INR within 1 to 2 weeks (Grade 2C).

3.3 Bridging for Low INRs

When the INR becomes subtherapeutic, there may be an increased risk of thrombosis. A 2008 retrospective study of 2,597 adult patients receiving warfarin mainly for atrial fibrillation or VTE matched 1,080 patients in the low-INR cohort with 1,517 patients in the therapeutic-INR cohort based on index INR date, indication for warfarin, and age.³⁹ All patients in the low-INR cohort had a subtherapeutic INR following two therapeutic INR measurements. There was no significant difference in thromboembolic events between the two groups, including the small number (99) of patients with artificial heart valves.

A second retrospective study addressed the same scenario in 294 patients with mechanical heart valves.⁴⁰ Bridging with LMWH was prescribed in 14 cases. The incidence of thromboembolic events was found to be 0.3% (95% CI, 0%-1.9%) for all patients included in the study and 0.4% (95% CI, 0%-2.0%) for all patients who did not receive bridging therapy. Both studies are limited by the observational study design and its potential for confounding. Unfortunately, this evidence only addresses the single low INR, not several consecutive low INRs.

Recommendation

3.3. For patients with stable therapeutic INRs presenting with a single subtherapeutic INR value, we suggest against routinely administering bridging with heparin (Grade 2C).

3.4 Vitamin K Supplementation

A low TTR as well as highly variable INR results are independent predictors of bleeding and thromboembolic complications during VKA therapy. One observational study using food diaries to quantify daily vitamin K intake showed that patients in the highest tertile of vitamin K intake had the most stable INR control over time, suggesting the possibility that daily vitamin K supplementation might improve anticoagulation control.⁴¹

Three randomized, placebo-controlled trials using pharmaceutically prepared vitamin K have addressed this issue.⁴²⁻⁴⁴ There are important differences among these RCTs, including the daily dose of vitamin K studied (100 μ g,^{42,44} 150 μ g,^{43,44} or 200 μ g⁴⁴), the study participants (general anticoagulation clinic patients^{42,44} or patients with unstable INR control⁴³), the width of targeted INR range (1.5^{42,44} or 1.0), and type of VKA (phenprocoumon or warfarin). Table 5 (and Table S5) shows the quality of evidence and main findings of our meta-analysis of the three RCTs. The absolute difference in TTR was a modest 3.54% (95% CI, 1.13%-5.96%). No difference in major bleeding or thromboembolic complications was seen.

The TTR observed in the control arms of these vitamin K RCTs indicates that studied patients had relatively stable INRs (TTR range, 78.0%-85.5%). It

Table 5—[Section 3.4] Low-Dose Vitamin K Supplementation Compared With Placebo for Patients
Taking VKAs To Stabilize INR ⁴²⁻⁴⁴

				A	anticipated Absolute Effects ^a
Outcomes	No. of Participants (Studies), Follow-up	Quality of the Evidence (GRADE)	Relative Effect (95% CI)	Risk With Placebo	Risk Difference With Low-Dose Vitamin K Supplementation (95% CI)
Thromboembolism	626 (3 studies), 6-12 mo	Very low ^{b-e} due to inconsistency, imprecision	RR 1.65 (0.08-34.03)	0 per 1,000	Not estimable ^f
Major bleeding	626 (3 studies), 6-12 mo	Very low ^{b-e} due to inconsistency, imprecision	RR 2.61 (0.34-20.28)	0 per 1,000	Not estimable ^f

^aTime frame in months.

^bAllocation concealment not reported; uncertain whether outcome adjudicators were blinded.

^cDefinition of thromboembolism and major bleeding different in each study.

^dStudies not powered to detect bleeding or thromboembolic events; total number of events is extremely low.

"Unable to rule out publication bias because not enough studies exist to populate a funnel plot.

^rTotal of two thromboembolic and three major bleeding events in low-dose vitamin K groups.

would be of greater interest to evaluate the effect of daily vitamin K supplementation in a population with unstable INRs that are not due to other correctable factors. In summary, current evidence does not support supplementation with vitamin K to increase TTR or to improve clinical outcomes.

Recommendation

3.4. For patients taking VKAs, we suggest against routine use of vitamin K supplementation (Grade 2C).

3.5 Anticoagulation Management Services for VKAs

In response to the recognized difficulty in coordinating oral anticoagulation therapy, AMS have evolved in both inpatient and outpatient settings. For the purposes of this review, an AMS was defined as having a designated, trained staff member responsible for patient INR monitoring and follow-up, the use of a standardized local procedure for VKA management (eg, dosing nomogram), and the management of regular INR testing. Further, usual care was defined as regular medical care that generally was provided by the patient's personal physician in the absence of an AMS.

Four prospective RCTs comparing usual care with the care of an AMS failed to show a significant difference in major bleeding, thromboembolism, or anticoagulation therapy-related mortality.⁴⁵⁻⁴⁸ None of these RCTs were blinded, only two studies clearly specified an intention-to-treat analysis,⁴⁶⁻⁴⁸ one study allowed patients to switch between treatment arms,⁴⁵ and all patients in two studies were stabilized in an AMS prior to randomization.^{45,48}

In contrast, the results of many low-quality observational studies have reported higher TTR and better outcomes in patients when anticoagulant therapy is managed by an AMS compared with usual care.⁴⁹⁻⁶⁵ The absolute difference in TTR between AMS and

community practices in a systematic review was $8.3\%~(95\%~{\rm CI},\,4.4\%\text{-}12.1\%),$ favoring AMS. 66

Given the conflicting results between randomized and nonrandomized studies and the lack of economic analysis or compelling patient preference data, the Antithrombotic Therapy and Prevention of Thrombosis, 9th ed: American College of Chest Physicians Evidence-Based Clinical Practice Guidelines committee decided to make the following best practice statement on this question:

3.5. We suggest that health-care providers who manage oral anticoagulation therapy should do so in a systematic and coordinated fashion, incorporating patient education, systematic INR testing, tracking, follow-up, and good patient communication of results and dosing decisions.

3.6 Patient Self-Testing and Self-Management

Patients using long-term oral anticoagulation therapy usually are monitored by going to a hospital or laboratory to provide blood by venipuncture for INR testing. Point-of-care (POC) devices allow INR testing to be performed by patients in their homes with a drop of blood from the finger. This is defined as patient self-testing (PST). If the patients who perform their own INR testing also adjust their anticoagulant dose, this is called patient self-management (PSM).⁶⁷ Several systematic reviews have evaluated RCTs of PST/PSM to determine whether these approaches to oral anticoagulation therapy result in better clinical outcomes than traditional laboratory-based INR monitoring.67-71 A recent individual patient metaanalysis clarified several aspects; our recommendations are based primarily on this more-detailed analysis.72

Pooled analyses show a significant reduction in the rate of thromboembolic complications with PST/PSM but not in the rate of major bleeding or overall mortality compared with usual laboratory-based INR

 Table 6—[Section 3.6] Patient Self-Testing/Self-Monitoring Compared With Usual Laboratory-Based Monitoring for VKA Therapy Management⁷²

				Anticipat	ted Absolute Effects ^a
Outcomes	No. of Participants (Studies), Follow-up	Quality of the Evidence (GRADE)	Relative Effect (95% CI)	Risk With Usual Laboratory-Based Monitoring	Risk Difference With Patient Self-Testing/Patient Self-Monitoring (95% CI)
Thromboembolism various methods ^b	6,417 (11 studies), 3-36 mo	Low ^{e,d} due to risk of bias, inconsistency	OR 0.51 (0.31-0.85)	48 per 1,000	23 fewer per 1,000 (from 7 fewer to 33 fewer)
Major bleeding various definitions ^b	6,417 (11 studies), 3-36 mo	Moderate ^c due to risk of bias	OR 0.88 (0.74-1.06)	77 per 1,000	9 fewer per 1,000 (from 20 fewer to 4 more)
Mortality all-cause mortality	6,417 (11 studies), 3-36 mo	Moderate ^c due to risk of bias	OR 0.82 (0.62-1.09)	87 per 1,000	15 fewer per 1,000 (from 32 fewer to 7 more)

^aTime frame in months.

^bDefined by individual studies.

°Flaws in study design, most commonly lack of blinding.

^dSignificant heterogeneity in pooled analysis ($I^2 = 52.6\%$).

monitoring (Table 6, Table S6). These benefits are seen most prominently in PSM rather than PST groups and possibly in patients with mechanical heart valves rather than other indications.⁷² The largest RCT of PST (n = 2,915), the Home INR Study (THINRS), demonstrated no advantage in clinical outcomes vs laboratory-based monitoring but did show modest, significant improvements in patient satisfaction with anticoagulant therapy and quality of life.⁷³ Data from a pooled analysis also show better patient satisfaction, quality of life, or both with PST/PSM, but these results are difficult to interpret because of the wide range and variable quality of the outcome measures used.⁷¹

Pooled results from RCTs show only modest (weighted mean difference, 1.50%; 95% CI, -0.63%-3.63%), nonsignificant improvement in TTR with ST/PSM compared with usual laboratory-based monitoring.⁷¹ The frequency of INR testing was considerably higher with PST/PSM compared with usual laboratory-based monitoring, with a mean of 22 to 24 more INR tests annually compared with control groups.⁷²

Resource utilization is relevant when considering whether to recommend widespread use of PST/PSM. Some analyses have deemed PST/PSM to be costeffective,⁷⁴⁻⁷⁶ whereas others have not.^{68,69,77} Higher costs with PST/PSM are driven largely by the cost of test strips and increased testing frequency.⁷⁷ However, the increased convenience that PST/PSM offers, particularly to those who travel frequently or who live remotely from testing facilities, can result in lower personal costs for individual patients.⁷⁷

Successful PST/PSM requires well-trained, highly motivated patients. In most RCTs, more than onehalf of patients were excluded because of physical limitations, inability to demonstrate competence with POC devices, apprehension about self-care, or patient refusal.⁷¹ Furthermore, up to 25% of patients randomized to PST/PSM withdrew prior to study completion.⁶⁷ THINRS was more promising in that \sim 80% were able to pass a PST competency assessment, but 16% switched from PST to the clinic testing group during the study.⁷³

Recommendation

3.6. For patients treated with VKAs who are motivated and can demonstrate competency in self-management strategies, including the selftesting equipment, we suggest PSM rather than usual outpatient INR monitoring (Grade 2B).

For all other patients, we suggest monitoring that includes the safeguards in our best practice statement 3.5.

3.7 Dosing Decision Support

There have been many reports of experience with paper nomograms and computer programs used to assist with VKA dosing.^{46,78-97} These dosing adjuncts have been studied at the initiation of therapy (no prior VKA doses) and during the maintenance phase of therapy and were compared with dose decisions made without the use of decision support (manual dosing). Both nomogram/computer-assisted and manual dosing were performed by experienced anticoagulation providers in some studies^{78,86,87,90,91} and by providers without specialized training (eg, trainee physicians, house staff, regular physician, nurses) in others.^{46,79-85,88,99,92-95}

Decision support-guided dosing (paper nomograms or computer programs) performed no better than manual dosing during initiation of VKA therapy in pooled analyses of available RCTs (Table 7, Table S7). Pooled analyses of RCTs evaluating decision supportguided dosing during maintenance therapy (all were computer-assisted dosing programs) revealed a mean TTR improvement of 4.5% (95% CI, 2.4%-6.7%) compared with no decision support. Although statistically

				Anticipa	ated Absolute Effects ^a
Outcomes	No. of Participants (studies), Follow-up	Quality of the Evidence (GRADE)	Relative Effect (95% CI)	Risk With Manual Dosing	Risk Difference With Dosing Decision Support (95% CI)
Thromboembolism, initiation variously defined	503 (4 studies), 3 mo	Low ^{b,c} due to risk of bias, imprecision	RR 0.61 (0.27-1.37)	63 per 1,000	63 fewer per 1,000 (from 46 fewer to 23 more)
Major bleeding, initiation variously defined	926 (7 studies ^d), 1-3 mo	Low ^{b,c} due to risk of bias, imprecision	RR 0.43 (0.17-1.09)	30 per 1,000	17 fewer per 1,000 (from 25 fewer to 3 more)
Mortality, initiation all-cause mortality	748 (5 studies ^e), 1-3 mo	Low ^{b,c} due to risk of bias, imprecision	RR 0.73 (0.36-1.46)	44 per 1,000	12 fewer per 1,000 (from 28 fewer to 20 more)
Thromboembolism, maintenance variously defined	14,213 (7 studies ^f), 1-12 mo	Moderate ^b due to risk of bias	RR 0.9 (0.7-1.17)	17 per 1,000	2 fewer per 1,000 (from 5 fewer to 3 more)
Major bleeding, maintenance variously defined	14,035 (5 studies ^g), 4.8-12 mo	Moderate ^b due to risk of bias	RR 0.92 (0.71-1.21)	15 per 1,000	1 fewer per 1,000 (from 4 fewer to 3 more)
Mortality, maintenance all cause mortality	14,044 (5 studies ^h), 4.8-12 mo	Moderate ^b due to risk of bias	RR 1.07 (0.78-1.48)	10 per 1,000	1 more per 1,000 (from 2 fewer to 5 more)

^aTime frame in days to months.

^bMost studies were unblinded, including patients, health-care providers, and outcome adjudicators.

°CI of relative effect encompasses wide range of benefit and harm.

^dAsnis et al,⁷⁹ Doecke et al,⁸² Kovacs et al,⁸⁵ Landefeld and Anderson,⁴⁶ Vadher et al,⁹² van den Bemt et al,⁹⁴ and White et al.⁹⁵

eAsnis et al,⁷⁹ Doecke et al,⁸² Kovacs et al,⁸⁵ Landefeld and Anderson,⁴⁶ and Vadher et al.⁹²

⁶Claes et al,⁸¹ Fitzmaurice et al,⁸³ Fitzmaurice et al,⁸⁴ Mitra et al,⁸⁸ Poller et al,⁹¹ Vadher et al,⁹² and Vadher et al.⁹³

^gClaes et al,⁸¹ Fitzmaurice et al,⁸³ Fitzmaurice et al,⁸⁴ Poller et al,⁹¹ and Vadher et al.⁹³

^hClaes et al,⁸¹ Fitzmaurice et al,⁸³ Fitzmaurice et al,⁸⁴ Poller et al,⁸⁹ and Poller et al.⁹¹

significant, this did not result in improvements in thromboembolism, major bleeding, or mortality outcomes (Table 7). The magnitude of TTR improvement with decision support-guided dosing was smaller when manual dosing in control groups was performed by experienced anticoagulation providers vs providers without specialized training (2.04% vs 8.22%, respectively; no *P* value provided). Higher TTR also has been associated with a paper nomogram in an observational study.⁹⁶

The use of computerized VKA dosing decision support reduces the time taken to dose each patient (mean time for computer-assisted dosing, 94 s; [95% CI, 66-123 s]; manual dosing, 149 s [95% CI, 102-196 s]).⁹⁷ This difference is unlikely to be clinically meaningful except in high-volume AMS locations.^{84,92,93} Inexperienced anticoagulation providers have safely used decision support-guided dosing.^{84,92,93} Although the computer-assisted dosing software is expensive, an economic analysis of the largest computer-assisted dosing RCT concluded that investment in computer-assisted dosing could represent good value if perpatient costs of dosing were reduced.⁹⁷

Recommendation

3.7. For dosing decisions during maintenance VKA therapy, we suggest using validated decision support tools (paper nomograms or computerized dosing programs) rather than no decision support (Grade 2C). *Remarks:* Inexperienced prescribers may be more likely to improve prescribing with use of decision support tools than experienced prescribers.

3.8 VKA Drug Interactions to Avoid

Previous systematic reviews addressing drug interactions with VKAs have examined INR results as outcomes and included case reports as evidence.98 Through a literature review, we sought evidence generated from 1996 to early 2011, looking for randomized trials with >50 patients per group or for large observational studies reporting on clinical outcomes (hemorrhage or VTE) related to drug interactions with VKAs. Our research identified 21 relevant studies. One meta-analysis of RCTs, one prospective cohort study, and many large health database studies were included. A meta-analysis of 10 RCTs (n = 4,180) compared VKA plus aspirin vs VKA alone and showed a reduced rate of arterial thromboembolism (OR, 0.66; 95% CI, 0.52-0.84). However, these benefits were limited to patients with a mechanical heart valve (OR, 0.27; 95% CI, 0.15-0.49), whereas the five studies that dealt with atrial fibrillation and cardiac disease showed no benefit with the combination.99 Major bleeding was increased in the meta-analysis regardless of the indication for the combination of VKA plus aspirin vs VKA alone (OR, 1.43; 95% CI, 1.00-2.02).

The remaining nonexperimental studies, which varied in size from 53 bleeding events to > 13,000 events,

measured hemorrhage as the clinical outcome. In general, the quality of evidence from these studies was low. The VKA studied in \sim 70% of the reports was warfarin. There was sufficient consistency in statistically significant increased rates of bleeding to be concerned about three main therapeutic drug categories. As noted in Table 8, nonsteroidal antiinflammatory drugs (NSAIDs), both nonselective and cyclooxygenase (COX)-2 selective; antiplatelet agents; and some antibiotics are associated with an increased risk of bleeding in patients taking VKAs.

For nonselective NSAIDs, studies reported ORs or risk ratios (RRs) from 1.9 (95% CI, 1.4-3.7) to 4.6 (95% CI, 3.3-6.5).^{100-103,105,118} In addition, two studies reported a higher risk of bleeding with nonselective NSAIDs compared with COX-2-selective NSAIDs.^{101,104} There was less consistency in the relationship between COX-2-selective NSAIDs plus VKAs vs VKA alone and bleeding outcomes, varying from a nonsignificant RR of 1.4 (95% CI, 0.44-4.30) to a significant OR of 3.1 (95% CI, 1.4-6.7).¹⁰⁰⁻¹⁰³ Antiplatelet agents, either undifferentiated, aspirin alone, or clopidogrel alone, were associated with increased rates of bleeding, with estimates of risk from an OR of 1.5 (95% CI, 1.05-2.22) to a hazard ratio (HR) of 3.1 (95% CI, 2.3-3.9).^{102,105,105-111} Aspirin plus clopidogrel plus VKA compared with VKA alone was associated with an HR of 3.70 (95% CI, 2.89-4.76).108

Data addressing interactions of antibiotics from multiple large database studies present a somewhat confusing picture. However, there are sufficient studies to suggest a risk of increased bleeding with cotrimoxazole (OR, 2.54 [95% CI, 2.08-3.10]; RR, 5.1 [95% CI, 2.1-12.3])^{112,113,115} and quinolones (OR, 1.55 [95% CI, 1.30-1.86]; RR, 5.9 [95% CI, 1.9-18.6]).^{105,112,113,115} There is a suggestion that cephalosporins (ignoring the anomalously high RR provided for cefradine), metronidazole, amoxicillin, amoxicillin/clavulanic acid, doxycycline, and fluconazole may have some impact on bleeding risk, but these drugs in general are insufficiently studied.¹¹²⁻¹¹⁴ Similarly, some studies suggest that selective serotonin reuptake inhibitors, tramadol, acetaminophen, coenzyme Q, and ginger may increase the risk of bleeding, but these also require confirmation.^{103,105,106,116,117}

Recommendations

3.8. For patients taking VKAs, we suggest avoiding concomitant treatment with NSAIDs, including COX-2-selective NSAIDs, and certain antibiotics (Grade 2C).

For patients taking VKAs, we suggest avoiding concomitant treatment with antiplatelet agents except in situations where benefit is known or is highly likely to be greater than harm from bleeding, such as patients with mechanical valves, patients with acute coronary syndrome, or patients with recent coronary stents or bypass surgery (Grade 2C).

4.0 VKA-Monitoring

4.1 Optimal Therapeutic INR Range

The desired effect of VKA on the prothrombin time, expressed as INR, can be provided as a therapeutic range (eg, INR 2.0-3.0) or a therapeutic target (eg, INR 2.5). The former provides information on INR values considered acceptable for the patient, whereas the latter is intended to induce those managing anticoagulant therapy to strive for an ideal level.

In a systematic review of 19 studies (one RCT, five with analysis of INR-specific outcomes from RCTs, and 13 observational studies) reporting clinical outcomes in at least three discrete INR ranges and including > 80,000 patients, the lowest rate of a composite outcome of major hemorrhage and symptomatic thromboembolism was seen with INR 2.0 to 3.0.119 Compared with INR 2.0 to 3.0, the RR for the composite outcome was 2.4 (95% CI, 1.9-3.1) for INR < 2and 1.8 (95% CI, 1.2-2.6) for INR 3.0 to 5.0. For INR > 5, the RR was 11.9 (95% CI, 6.0-23.4) based on 13 studies for bleeding and only one study for thromboembolism. The evidence profiles are shown separately for comparisons of INR 2.0 to 3.0 vs INR 3.0 to 5.0 (Table 9, Table S8) vs INR < 2.0 (Table 10, Table S9). The definition of major bleeding differed among studies, and the type of thromboembolic events varied according to the studied indication for VKA. However, the pattern of relative risks was consistent among atrial fibrillation, valvular heart disease, and other indications taken together.

Patients with an increased risk of thromboembolic complications are those with (1) a mechanical mitral valve; (2) a mechanical aortic valve in combination with atrial fibrillation, anterior-apical ST-segment elevation myocardial infarction, left atrial enlargement, low ejection fraction, or hypercoagulable state; and (3) caged-ball or caged-disk valve or thromboembolic complications while in INR 2.0 to 3.0. These subsets of patients are traditionally, although with lack of evidence, treated at a higher-intensity INR 2.5 to 3.5 (see Whitlock et al¹²⁰ in this supplement).

4.1.1 Low-Intensity VKA for Patients With VTE: Low-intensity treatment with VKA corresponds to INR 1.5 to 1.9/2.0 and is of interest because of the possibility that it might cause less bleeding than conventional intensity (INR 2.0-3.0). In addition, given a wider margin of safety from excessive anticoagulation, laboratory

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Interacting Drug	Summary Effect on Bleeding (95% $\rm CI)^b$	Study
NSAIDS		
NSNSAIDs	OR 1.9 (1.4-3.7)	Battistella et al ¹⁰⁰
	HR 3.6 (2.3-5.6)	Cheetham et al ¹⁰¹
	RR 1.33 (0.78-2.25)	Delaney et al ¹⁰²
	OR 2.6 (1.6-4.2)	Hauta-Ato et al ¹⁰³
	OR 3.01 (1.42-6.37)	Knijff-Dutmer et al ^{104,a}
	RR 2.6-6.5°	Penning-van Beest et al ^{105,}
	OR 4.6 (3.3-6.5) ^d	Schalekamp et al ¹⁰⁶
	NSNSAID vs COX-2 OR 3.07 (1.18-8.03)	Knijff-Dutmer et al ^{104,a}
	NSNSAIDs vs COX-2 HR 3.7 (1.4-9.6)	Cheetham et al ¹⁰¹
COX-2-selective NSAIDs	OR 1.7-2.4 ^e	Battistella et al ¹⁰⁰
	HR 1.7 (0.6-4.8)	Cheetham et al ¹⁰¹
	RR 1.37 (0.44-4.30)	Delaney et al ¹⁰²
	OR 3.1 (1.4-6.7)	Hauta-Aho et al ¹⁰³
Antiplatelet agents		
Aspirin	OR 1.43 (1.00-2.02) ^f	Dentali et al ⁹⁹
	RR 2.23 (1.46-3.41)	Delaney et al ¹⁰²
	IR 0.08/patient-y vs 0.06 for warfarin alone	Buresly et al ¹⁰⁷
	HR 1.83 (1.72-1.96)	Hansen et al ¹⁰⁸
	RR 3.0 (1.0-9.4)	Penning-van Beest et al ^{105,}
Clopidogrel	HR 3.08 (2.3-3.9)	Hansen et al ¹⁰⁸
Aspirin plus clopidogrel	HR 3.70 (2.89-4.76)	Hansen et al ¹⁰⁸
Antiplatelet agents (any antiplatelet)	OR 2.06 (1.01-4.36)	Johnson et al ¹⁰⁹
	OR 1.53 (1.05-2.22)	Shireman et al ¹¹⁰
	RR 1.76 (1.05-2.95)	Toyoda et al ¹¹¹
Antibiotics		
Cephalexin	OR 1.38 (1.10-1.73)	Schelleman et al ¹¹²
Cefradine	RR 43.0 (10.7-172.4)	Penning-van Beest et al ^{113,}
Cephalosporins	OR 1.16 (1.04-1.29)	Zhang et al ¹¹⁴
Metronidazole	OR 1.58 (1.32-1.89)	Zhang et al ¹¹⁴
Cotrimoxazole	OR 3.84 (2.33-6.33)	Fischer et al ¹¹⁵
	OR 2.54 (2.08-3.10)	Schelleman et al ¹¹²
	RR 5.1 (2.1-12.3)	Penning-van Beest et al ^{113,a,}
	Cotrimox vs cephalexin OR 1.68 (1.21-2.33)	Schelleman et al ¹¹²
Ciprofloxacin	OR 1.94 (1.28-2.95)	Fischer et al ¹¹⁵
	OR 1.62 (1.31-1.99)	Schelleman et al ¹¹²
	RR 3.2 (1.3-7.7)	Penning-van Beest et al ^{113,}
Levofloxacin	OR 1.55 (1.30-1.86)	Schelleman et al ¹¹²
Norfloxacin	RR 5.9 (1.9-18.6)	Penning-van Beest et al ^{105,}
Amoxycillin	OR 1.28 (1.03-1.58)	Schelleman et al ¹¹²
	RR 3.1 (1.6-6.3)	Penning-van Beest et al ^{113,}
Amoxycillin/clavulanic acid	RR 4.4 (2.5-7.8)	Penning-van Beest et al ^{113,a}
Doxycycline	RR 2.6 (1.2-4.8)	Penning-van Beest et al ^{113,}
Fluconazole	OR 1.89 (1.35-2.64)	Schelleman et al ¹¹²
	Fluconazole vs cephalexin OR 2.09 (1.34-3.26)	Schelleman et al ¹¹²
Other		
SSRIs	OR 2.6 (1.5-4.3)	Hauta-Aho et al ¹⁰³
	OR 1.7 (1.1-2.5) ^h	Schalekamp et al ^{106,a}
	OR 1.1 (0.9-1.4) to 1.2 (0.8-1.7); NS	Kurdyak et al ¹¹⁶
Tramadol	RR 3.3 (1.1-10.4)	Penning-van Beest et al ^{105,4}
Complementary medicines	Coenzyme Q10 (OR 3.69, 95% CI 1.88-7.24) and	Shalansky et al ¹¹⁷
	ginger (OR 3.20, 95% CI 2.42-4.24).	

COX = cyclooxygenase; IR = incidence rate; NSNSAID = nonselective nonsteroidal antiinflammatory drug; SSRI = selective serotonin reuptake inhibitor. See Table 1 and 3 legends for expansion of other abbreviations.

aStudy VKAs were warfarin, phenprocoumon, and acenocoumarol.

^bUnless stated, refers to drug plus VKA vs VKA alone.

^cDiclofenac (RR, 2.6) and naproxen (RR, 6.5) studied separately.

^dOR is for GI bleeding, whereas OR for non-GI bleeding is 1.7 (95% CI, 1.3-2.2).

^eSeparate OR given for celecoxib (1.7) and rofecoxib (2.4); both statistically significant.

^fDentali et al⁹⁹ meta-analysis of randomized clinical trials.

gData duplication between two study publications; therefore, more conservative estimate used.

^hOnly statistically significant for non-GI bleeding; not significant for GI bleeding or intracranial bleeding.

				Anticipa	ated Absolute Effects ^a
Outcomes	No. of Participants, (Studies) Follow-up	Quality of the Evidence (GRADE)	Relative Effect (95% CI)	Risk With INR 2-3	Risk Difference With INR 3-5 (95% CI)
Major hemorrhage per 100 patient-y, various definitions	76,646 (17 studies ^b), 1.8 y	Low ^{c,d} due to risk of bias, dose-response gradient	RR 2.7 (1.8-3.9)	6 per 1,000	10 more per 1,000 (from 5 more to 17 more)
Thromboembolism per 100 patient-y, various definitions	835 (10 studies ^e)	Very low ^{fg} due to risk of bias, inconsistency	RR 0.9 (0.6-1.3)	50 per 1,000	tudy population 5 fewer per 1,000 (from 18 fewer to 14 more) Moderate 5 fewer per 1,000 (from 20 fewer to 15 more)

Table 9—[Section 4.1.1] Optimal Therapeutic INR Range: Higher Target vs 2 to 3119

^aTime frame in days to months.

^bSix studies had a randomized controlled trial design.

^cThe majority of studies (eight) were retrospective cohorts.

^dIt is biologically plausible that with increased intensity there will be more bleeding.

^eOne study had a randomized control design.

'Three of four studies had a retrospective cohort design.

^gThromboembolic events were more frequent with an INR of 2 to 3 in two studies, less frequent in one study, and similar in one study.

monitoring intervals could perhaps be increased to decrease the burden of therapy on the patient. Two RCTs, both blinded, investigated the efficacy and safety of low-intensity VKA in patients with unprovoked VTE.^{121,122} Patients were recruited after having received initial conventional-intensity anticoagulation for months to years. Kearon et al¹²¹ compared low intensity with conventional intensity in 738 patients and found a higher risk of recurrent VTE without any reduction of bleeding events in patients treated with low-intensity VKA. Ridker et al¹²² compared low-intensity warfarin with placebo in 508 patients and observed a reduction of recurrent VTE with active treatment without any significant increase in bleeding.

In conclusion, the benefit of low-intensity VKA in terms of reduced risk of bleeding is uncertain because of these inconsistent results. The second benefit of reduced frequency of monitoring is attainable also with conventional-intensity VKA for patients with a stable INR, as reviewed in section 2.1. Thus, the proposed advantage of lower-intensity VKA therapy in the extended-treatment phase is questionable.

4.1.2 Low-Intensity VKA for Patients With Atrial Fibrillation: For stroke prophylaxis in atrial fibrillation, two less-intensive alternatives to conventional-intensity VKA have been studied. Minidose or low-intensity fixed-dose VKA, usually corresponding to 1.25 mg (0.5-3 mg) warfarin daily, was given with the intention to minimize the need for laboratory monitoring. A meta-analysis of four randomized trials with 2,753 patients showed that minidose warfarin was inferior to conventional-intensity VKA with regard to thrombotic events (RR, 0.50; 95% CI, 0.25-0.97). Results were uncertain for major hemorrhage (RR, 1.23; 95% CI, 0.67-2.27) or fatal bleeding (RR, 0.97; 95% CI, 0.27-3.54).¹²³

Low-intensity VKA with a therapeutic range of INR 1.5 to 2.0 (or 2.1 in one study) has been compared head to head with conventional intensity, without the addition of aspirin, in two randomized trials.^{124,125} One study from Japan was stopped prematurely after an excess of major hemorrhages in the conventional-intensity group.¹²⁴ A similar trend was seen in a separate study from Italy.¹²³ Neither study showed a difference in stroke or deaths. The mean age of the patients differed; 65 years in the Japanese study¹²⁴ and 80 years in the Italian trial.¹²⁵ The pooled results show that there is a significant reduction of nonfatal extracranial hemorrhages with low-intensity VKA (OR, 0.21; 95% CI, 0.06-0.6) without any appreciable increase in the rate of stroke or mortality.

A case-control study in patients with atrial fibrillation suggested that the risk of stroke increases at INR < 2.0.¹²⁶ Compared with an INR of 2.0, the OR for stroke was 2.0 (95% CI, 1.6-2.4) at an INR of 1.7 and 3.3 (95% CI, 2.4-4.6) at an INR of 1.5. There is a trade-off that pits a substantial relative risk reduction of stroke (~80%) with INR 1.5 to 2.0 compared with INR < 1.2,^{127,128} with a greater risk of thromboembolic events with INR 1.4 to 1.7 compared with INR 2.0 to 2.5 (OR, 3.72; 95% CI, 2.67-5.19) (Anticoagulation and Risk Factors in Atrial Fibrillation [ATRIA] cohort).¹²⁹ In this study, there was no evidence for a reduced risk for intracranial hemorrhage at INR < 2.0 compared with 2.0 to 3.5. The event

				Anticipa	ted Absolute Effects
Outcomes	No. of Participants (Studies)	Quality of the Evidence (GRADE)	Relative Effect (95% CI)	Risk With INR 2-3	Risk Difference With $INR < 2 (95\% CI)$
Major hemorrhage	78,493 (17	Very low ^{a,b} due to risk	RR 1.1 (0.7-1.7)	St	udy population
per 100 patient-y,	studies ^a)	of bias, inconsistency		6 per 1,000	1 more per 1,000 (from
various definitions				1	2 fewer to 4 more)
					Moderate
				23 per 1,000	2 more per 1,000 (from
				-	7 fewer to 16 more)
Thromboembolism	827 (4 studies ^c)	Moderate ^{d-f} due to risk	RR 3.5 (2.8-4.4)	St	udy population
per 100 patient-y		of bias, large effect,		46 per 1,000	115 more per 1,000
		dose-response gradient			(from 83 more to
					157 more)
					Moderate
				40 per 1,000	100 more per 1,000
					(from 72 more to 136
					more)

^aEight of the studies were retrospective cohorts.

 $^{\mathrm{b}}$ Four studies showed higher risk of bleeding, with INR < 2.

°Only one study had a randomized control design.

^dNo explanation was provided.

eAt least 2.8 times more frequent thromboembolism.

fIt is biologically plausible with more thromboembolism at a lower INR.

rate of intracranial hemorrhage is low with long-term VKA therapy (0.3% per year),¹³⁰ thus very large numbers are required to detect a difference. There was a reduction of major, nonfatal extracranial hemorrhage with low- vs standard-intensity VKA in the two RCTs (OR, 0.21; 95% CI, 0.06-0.6), and this could be important for patients with a documented bleeding diathesis.

Recommendation

4.1. For patients treated with VKAs, we recommend a therapeutic INR range of 2.0 to 3.0 (target INR of 2.5) rather than a lower (INR < 2) or higher (INR 3.0-5.0) range (Grade 1B).

4.2 Therapeutic Range for High-Risk Groups

The most common therapeutic range for treatment with VKAs is INR 2.0 to 3.0, as discussed previously. Higher intensity for patients with a mechanical mitral valve or with a mechanical aortic valve in combination with other risk factors is discussed in Whitlock et al^{120} in this supplement.

Patients with severe thrombophilia (antiphospholipid syndrome, deficiency of protein C, protein S, or antithrombin homozygous factor V Leiden) who have thromboembolic events have an increased risk of recurrent VTE compared with those without thrombophilia or with mild defects (eg, heterozygous factor V Leiden) in the absence of anticoagulant treatment. It is not clear to what extent this is true while taking VKAs. Case series of patients with deficiency of any of the natural inhibitors (protein C, protein S, antithrombin) or with the common factor V Leiden or prothrombin gene polymorphisms have not provided any indication that moderate intensity (INR 2.0-3.0) is inadequate for these conditions.

In retrospective studies, moderate-intensity anticoagulation often was insufficient to prevent arterial or venous thrombosis in patients with antiphospholipid antibodies. Many of the patients in these studies were recruited from specialized centers for patients with rheumatic disease,¹³¹⁻¹³³ which may be a different population than those with primary antiphospholipid syndrome (ie, thromboembolism without identified underlying disease).

A systematic review compared the efficacy and safety of different approaches of secondary prophylaxis against thromboembolism in patients with antiphospholipid antibodies based on 16 studies (two RCTs, two subgroup analyses from RCTs, three prospective cohorts or subgroup analysis, and nine retrospective cohorts or subgroup analyses).¹³⁴ There were more fatal thromboembolic events than fatal hemorrhages (18 vs one), and the risk of thrombotic events was inversely related to the INR value in the observational studies but not in the RCTs. In many of the studies, only a single laboratory test had been used to confirm the syndrome, whereas according to current criteria (revised Sapporo criteria), at least two positive tests should be recorded with an interval of at least 12 weeks.135

The results of the two RCTs^{136,137} are shown in Table 10 (Table S10). Both studies were small, with 110 patients randomized to higher-intensity (INR 3.0-4.0 or INR 3.0-4.5) and 110 randomized to moderate-intensity (INR 2.0-3.0) warfarin therapy. Three patients with nonembolic arterial disease were assigned to aspirin alone (not included in Table 11¹³⁸). Because the CIs for the relative risk are wide and risk of bias is substantial, the quality of evidence is low.

Patients with cancer and VTE have a higher risk of recurrent events during anticoagulant therapy than patients without cancer.^{139,140} When such a breakthrough event occurs, an intensification of treatment sometimes is suggested.¹⁴¹ There are no published aggregate data on the effectiveness and safety of intensified treatment with VKA, only single-patient case reports. Dose escalation of LMWH appeared effective to prevent further recurrence in a retrospective review of 70 patients.¹⁴²

Recommendation

4.2. For patients with antiphospholipid syndrome with previous arterial or venous thromboembolism, we suggest VKA therapy titrated to a

moderate-intensity INR range (INR 2.0-3.0) rather than higher intensity (INR 3.0-4.5) (Grade 2B).

5.0 VKA—DISCONTINUATION OF THERAPY

There is a theoretical concern that abrupt VKA discontinuation may result in a temporary hypercoagulable state due to an imbalance in the rates of normalization of activity of the coagulation factors II, VII, IX, and X on the one hand and the natural inhibitors protein C and protein S on the other.¹⁴³ Five small controlled trials (total n = 217) have addressed this issue.¹⁴⁴⁻¹⁴⁷ The primary outcomes of four of the studies were laboratory results suggestive of a hypercoaguable state^{144,145,147,148} and produced inconsistent results. Elevations tended to persist for 8 to 9 weeks, regardless of discontinuation strategy, suggesting an unmasked prothrombotic state in the absence of anticoagulant protection rather than a rebound phenomenon associated with abrupt discontinuation.

The thromboembolism event rate appeared similar between groups across the five studies (Table 12, Table S11).¹⁴⁴⁻¹⁴⁸ The only major hemorrhage occurred

Table 11—[Section 4.2] High-Intensity VKA Compared With Moderate-Intensity VKA for Patients
With Antiphospholipid Syndrome ^{136,137}

				Anticipated	Absolute Effects
Outcomes	No. of Participants (Studies), Follow-up	Quality of the Evidence (GRADE)	Relative Effect (95% CI)	Risk With Moderate-Intensity VKA	Risk Difference With High-Intensity VKA (95% CI)
Thromboembolism	220 (2 studies ^a), 3 y	Very low ^{b,c} due to risk of	OR 2.33 (0.82-6.66)	Study	population ^d
objective confirmation		bias, indirectness, and imprecision		45 per 1,000 ²	54 more per 1,000 (from 8 fewer to 195 more)
		-			Low ^d
				50 per 1,000ª	59 more per 1,000 (from 9 fewer to 210 more)
]	Highd
				700 per 1,000ª	145 more per 1,000 (from
				*	43 fewer to 240 more)
Major bleeding	220 (2 studiese), 3 y	Moderate ^f due to	OR 0.70 (0.23-2.16)	Study	population
		imprecision		64 per 1,000ª	18 fewer per 1,000 (from 48 fewer to 64 more)
					Low
				25 per 1,000ª	7 fewer per 1,000 (from
				*	19 fewer to 27 more)
					High
				100 per 1,000ª	28 fewer per 1,000 (from
				_	75 fewer to 94 more)
Mortality all-cause	220 (2 studies), 3 y	Moderate ^f due to	OR 1.51 (0.3-7.72)	18 per 1,000	9 more per 1,000 (from
mortality		imprecision			13 fewer to 107 more)

See Table 1 and 2 legends for expansion of abbreviations.

^aIn the study by Finazzi et al,¹³⁷ three patients with nonembolic arterial thrombosis received, as planned, only aspirin. They had no events and have not been included here.

^bThe study by Finazzi et al¹³⁷ was open label.

^cBoth studies were designed to show superiority of the more intensive regimen, not equivalence. The 95% CI includes both benefit and significant harm. ^dLow of 5% from Schulman et al¹³⁸; high of 70% from Khamashta et al.¹³¹

^eThe types of major hemorrhage were not disclosed.

'The 95% CI includes both benefit and significant harm.

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				Anticipated	l Absolute Effectsª
Outcomes	No. of Participants (Studies), Follow-up	Quality of the Evidence (GRADE)	Relative Effect (95% CI)	Risk With Abrupt Withdrawal	Risk Difference With Gradual Withdrawal (95% CI)
Thromboembolism imaging diagnostics	217 (5 studies), 3 mo	Low ^{b,c} due to risk of bias, imprecision	OR 0.96 (0.42-2.18)	$126 \text{ per } 1,000^{d}$	4 fewer per 1,000 (from 69 fewer to 113 more)
Mortality all-cause mortality	217 (5 studies), 1 mo	Very low ^{b,c} due to risk of bias, imprecision	OR 0 (0.01-5.6)	9 per 1,000	9 fewer per 1,000 (from 9 fewer to 39 more) ^d
Major hemorrhage	217 (5 studies), 1 mo	Very low ^{b,c} due to risk of bias, imprecision	OR 1 (0.1-5.6)	9 per 1,000	0 fewer per 1,000 (from 8 fewer to 39 more) ^d

^aTime frame is weeks.

^bUnclear whether allocation was adequate in Tardy et al,¹⁴⁸ de Groot et al,¹⁴⁵ and Ascani et al.¹⁴⁴ In Michaels and Beamish,¹⁴⁶ it was according to year of birth. Unclear whether allocation was concealed in Tardy, de Groot, and Ascani; it was not concealed in Michaels. Clinicians and patients were not blinded in de Groot, Michaels, Palareti et al,¹⁴⁷ or Ascani.

eVery small patient groups and few events.

^dThere is no better source than these trials, so low or high estimates are not provided.

in the gradual withdrawal group. Gradual discontinuation of VKA is likely to be more confusing and inconvenient for the patient.

Recommendation

5.0. For patients eligible to discontinue treatment with VKA, we suggest abrupt discontinuation rather than gradual tapering of the dose to discontinuation (Grade 2C).

6.0 PARENTERAL ANTICOAGULANTS

6.1 UFH—Dose Adjustment by Weight

Five RCTs compared initial IV UFH dosing according to a weight-based nomogram with a fixed-dose approach.¹⁴⁹⁻¹⁵³ The study by Jaff et al¹⁵¹ was excluded because no weight-adjusted group for the initial bolus was included. The study by Toth and Voll¹⁵³ was excluded because the fixed dose varied by treating physician, and thromboembolic or bleeding complications were not specified. In the remaining three RCTs a total of 292 patients were randomized to either weight based or fixed dose initially. The fixed dose was a bolus of 70 to 80 units/kg followed by an infusion rate of 15 to 18 units/kg per h. Activated partial thromboplastin time (aPTT) values were monitored, and UFH dose titrated to the therapeutic range.^{149,150,152} In one of the studies, a POC device for measuring aPTT was used.¹⁴⁹ Patients with acute coronary syndromes¹⁵⁰ or mixed diagnosed conditions, including VTE,^{149,152} were recruited. Study follow-up periods ranged from 48 h^{149,150} to 3 months.¹⁵² The weight-based and fixed-dose approaches achieved similar therapeutic aPTTs during the first 24 to 48 h. Patient-important adverse events, which were not well defined, were few; thromboembolism in eight vs two (OR, 0.22; 95% CI, 0.02-1.13) in the fixeddose vs weight-adjusted group and only one major bleed (fixed-dose group) (Table 13, Table S12). These results suggest that weight-adjusted dosing and fixed dosing of IV UFH are similar in outcomes. Small numbers of clinical events and failure to specify the timing of thromboembolic complications are major limitations of available studies.

Either regimen can be monitored with plasma heparin levels, but there is no evidence to suggest that monitoring improves clinical outcomes. The evidence linking plasma heparin levels of 0.3 to 0.7 International Units/mL anti-Xa activity by the amidolytic assay to the occurrence of either bleeding or thrombosis is also of low quality.¹⁵²

Recommendation

6.1. For patients starting IV UFH, we suggest that the initial bolus and the initial rate of the continuous infusion be weight adjusted (bolus 80 units/kg followed by 18 units/kg per h for VTE; bolus 70 units/kg followed by 15 units/kg per h for cardiac or stroke patients) or a fixed-dose (bolus 5,000 units followed by 1,000 units/h) rather than alternative regimens (Grade 2C).

6.2 UFH—Dose Management of SC UFH

Treatment with UFH has traditionally been monitored with aPTT plasma tests, whether administered by IV or SC. The SC treatment regimens for UFH generally were based on a fixed initial dose.¹⁵⁴ In contrast, short-term treatment with LMWH is given without any laboratory monitoring because the pharmacokinetic characteristics are believed to be more predictable than for UFH. Studies of SC UFH have not compared weight-based dosing

Table 13—[Section 6.1] UFH: Weight-Based Nomogram Compared With Fixed Initial Dose for Patients With Thromboembolic Disease^{149,150,152}

				Aı	nticipated Absolute Effects ^a
Outcomes	No. of Participants (Studies), Follow-up	Quality of the Evidence (GRADE)	Relative Effect (95% CI)	Risk With Fixed Initial Dose	Risk Difference With UFH-Weight-Based Nomogram (95% CI)
Thromboembolism	292 (3 studies), 2-90 d ^b	Low ^{c,d} due to risk of bias and imprecision	$\begin{array}{c} \text{OR 0.22} \\ (0.02\text{-}1.13)^{\text{e}} \end{array}$	$57 \ \mathrm{per} \ 1{,}000^{\mathrm{f}}$	44 fewer per 1,000 (from 56 fewer to 7 more)
Major hemorrhage	179 (2 studies ^g), 1 wk	Very low ^{c,d} due to risk of bias and imprecision	Not estimable ^h	11 per 1,000	10 fewer per 1,000 (from 30 fewer to 10 more)

See Table 1 and 2 legends for expansion of abbreviations.

^aTime frame is days to weeks.

^bOnly Raschke et al¹⁵² collected data over a 3-mo period.

^cThe studies did not use blinding.

^dNone of the studies was powered for clinical outcomes, which were few and poorly reported with regard to type and timing. ^eFisher exact test.

'Two of the eight events occurred after discontinuation of warfarin.

^gBecker et al¹⁴⁹ reported 2% bleeding without specifying allocation group or type of bleeding.

^hZero events in control group; 95% CI on OR not estimable.

vs fixed dosing with or without the use of aPTT monitoring. Weight-adjusted SC UFH monitored with aPTT has been compared with SC LMWH in three RCTs (n = 937) with similar clinical outcome results as follows: recurrent VTE (OR, 1.13; 95% CI, 0.52-2.46), major bleeding (OR, 1.28; 95% CI, 0.42-4.09), and death (OR, 1.34; (95% CI, 0.62-2.93).¹⁵⁵

One RCT in patients with VTE has compared the use of weight-adjusted dosing of SC UFH to weightbased dosing of LMWH without monitoring.¹⁵⁶ The SC UFH was administered at an initial dose of 333 units/kg followed by a dose of 250 units/kg bid; subsequent UFH dosing was kept constant. Clinical outcomes were similar between the SC UFH and LMWH groups (Table 14, Table S13). Because all of the evidence for initial dosing and monitoring of SC UFH is indirect, the quality of evidence for any recommendation is very low. Outpatient use of SC UFH while transitioning to VKA treatment derives some benefit from the elimination of daily blood work. Treatment with UFH often is preferred for patients with severe renal insufficiency, where there is a risk for accumulation of LMWH or fondaparinux.

Recommendation

6.2. For outpatients with VTE treated with SC UFH, we suggest weight-adjusted dosing (first dose 333 units/kg, then 250 units/kg) without monitoring rather than fixed or weight-adjusted dosing with monitoring (Grade 2C).

				Anticipated	Absolute Effects ^a
Outcomes	No. of Participants (Studies), Follow-up	Quality of the Evidence (GRADE)	Relative Effect (95% CI)	Risk With Weight- Adjusted Nonmonitored LMWH SC	Risk Difference With Weight- Adjusted Nonmonitored UFH SC (95% CI)
Recurrent VTE objectively measured with same method as for index event	697 (1 study), 3 mo	Low ^{h.c} due to indirectness and imprecision	OR 1.11 (0.49-2.52)	34 per 1,000	4 more per 1,000 (from 17 fewer to 48 more)
Major bleeding by ISTH criteria	697 (1 study), 3 mo	Low ^{b,c} due to indirectness and imprecision	OR 0.5 (0.17-1.34)	34 per 1,000	17 fewer per 1,000 (from 28 fewer to 11 more)
Mortality	697 (1 study), 3 mo	Low ^{b,c} due to indirectness and imprecision	OR 0.83 (0.43-1.57)	62 per 1,000	10 fewer per 1,000 (from 35 fewer to 32 more)

 Table 14—[Section 6.2] UFH: Weight-Adjusted Nonmonitored UFH SC Compared With Weight-Adjusted Nonmonitored LMWH SC for Outpatients With Acute VTE¹⁵⁶

ISTH = International Society on Thrombosis and Haemostasis; SC = subcutaneous. See Table 1 and 2 legends for expansion of other abbreviations.

^aTime frame is days to weeks.

^bThe comparison should actually be vs fixed-dose UFH SC with monitoring, but weight-adjusted UFH SC has only been compared directly with weight-adjusted LMWH. Thus, the comparison is indirect.

^cBecause of premature discontinuation, the study was not powered to demonstrate equivalence.

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7.0 LMWH-Dosing

7.1 Should the Therapeutic Dose of LMWH Be Modified for Decreased Renal Function?

LMWH, as opposed to UFH, is primarily eliminated through renal excretion. We found no RCTs comparing a standard, body-weight-adjusted dose to a reduced dose of LMWH in severe renal insufficiency, defined as creatinine clearance < 30 mL/min.

A meta-analysis of 18 observational studies or subgroup analyses of studies using therapeutic doses of LMWH provides some indirect evidence on this patient population.¹⁵⁷ On the basis of four of the studies, this review suggested that standard doses of LMWH led to higher peak levels of anti-factor Xa in patients with a creatinine clearance < 30 mL/min compared with those with a creatinine clearance > 30 mL/min. On the basis of three studies, when the dose of LMWH was reduced for severe renal failure, no such difference in peak level was observed. All of these seven studies used enoxaparin, so there are insufficient data to comment on other LMWHs. In addition, the relevance of anti-factor Xa levels is unclear: several studies have failed to show a relationship between the anti-Xa levels and bleeding.158-160

For patients treated with LMWH, the risk of bleeding was generally higher in patients with a creatinine clearance < 30 mL/min compared with patients with a creatinine clearance >30 mL/min (5.0% vs 2.4%; OR, 2.25; 95% CI, 1.19-4.27; P = .013).¹⁵⁷ However, because the risk of bleeding is also increased when patients with severe renal failure are treated with UFH,¹⁶¹ the problem may be the renal function rather than the dosing regimen. Four observational studies in the review using enoxaparin suggested that lowering doses for severe renal impairment may reduce the incidence of bleeding (Table 15).¹⁵⁷ The dose adjustment was either empirical or to 0.5 vs the standard 1 mg/kg bid of enoxaparin. There are insufficient data on VTE outcomes. Overall, the evidence is indirect and from studies of low quality and provides no advice on how LMWH should be reduced if the decision is to reduce.

Recommendation

7.1. For patients receiving therapeutic LMWH who have severe renal insufficiency (calculated

creatinine clearance < 30 mL/min), we suggest a reduction of the dose rather than using standard doses (Grade 2C).

8.0 Fondaparinux—Dosing

8.1 Fondaparinux Dose Management by Weight

Doses of heparins for the treatment of thrombosis often are administered according to patient body weight for both LMWH and UFH. Both total body weight and lean body weight have been used. In clinical trials, patients with morbid obesity (>120-130 kg) often have been excluded. We did not identify any studies comparing weight-adjusted dosing of fondaparinux to standard doses not adjusted for weight. Two randomized trials for symptomatic venous thrombosis^{162,163} used doses adjusted for the total body weight of the patient (5.0, 7.5, or 10 mg in patients)weighing <50, 50-100, or >100 kg, respectively). These trials—one in DVT,¹⁶² one in PE¹⁶³—compared fondaparinux to enoxaparin and UFH, respectively. A separate study was a subgroup analysis comparing the 3-month incidence of recurrent VTE or major bleeding events in a subset of patients weighing <100 kg and >100 kg.¹⁶⁴ The incidences of recurrence and major bleeds appeared to be similar for each patient subset of weight and BMI for patients treated with fondaparinux; VTE occurred in 75 of 1,946 (3.9%) nonobese patients vs 10 of 251 (4%) obese patients, and major bleeds occurred in 25 of 1,993 (1.3%) nonobese patients vs in one of 248 (0.4%) in obese patients. This subgroup analysis has several limitations (no tests for interaction, small number of obese patients, unclear definitions of major bleeds) and provides only low-quality evidence. There are insufficient data on patient s with low body weight to make any recommendation or suggestion regarding dose adjustment for these patients.

Recommendation

8.1. For patients with VTE and body weight over 100 kg, we suggest that the treatment dose of fondaparinux be increased from the usual 7.5 mg to 10 mg daily subcutaneously (Grade 2C).

Table 15—[Section 7.1] Risk of Bleeding With Enoxaparin According to the Calculated Creatinine Clearance

	Bleeding Rate CalCrCl \leq 30, n/N (%)	Bleeding Rate CalCrCl $\!>\!30,$ n/N (%)	${\rm OR}~(95\%~{\rm CI})$
Studies where dose was unadjusted for CalCrCl	17/206 (8.3)	96/4,081 (2.4)	3.88 (1.78-8.45)
Studies where dose was adjusted for CalCrCl	1/106 (0.9)	5/265 (1.9)	0.58 (0.09-3.78)

CalCrCl = calculated creatinine clearance.

Anticoagulant Therapy

9.0 Prevention and Management of Anticoagulant Complications

9.1 Vitamin K for Patients Taking VKAs With High INRs Without Bleeding

The risk of bleeding increases significantly when the INR exceeds 4.5.¹⁶⁵ In a retrospective review, patients with mechanical heart valves had a risk of adverse events that increased logarithmically from two per 100 patient-years at INR 2.5 to 4.9, to 4.8 per 100 patient-years for INR 5 to 5.5, then to 75 per 100 patient-years for INR \geq 6.5.¹⁶⁶ Similarly, a casecontrol analysis of adults sustaining intracerebral bleeding while on warfarin noted a doubling of intracerebral bleeding for every 0.5-s increment in prothrombin time (approximately every 1-point increase in INR).¹⁶⁷

When the INR is supratherapeutic without evidence of bleeding, strategies used to lower the INR have included withholding VKA, adjusting the dose of VKA, and providing some dose of vitamin K. Vitamin K shortens the time to return to normal INR.¹⁶⁸⁻¹⁷⁰ A 2006 meta-analysis found that administration of vitamin K orally or by IV was more likely to reverse overanticoagulation (INR > 4) at 24 h compared with simply withholding VKA.¹⁷¹

INR 4.5 to 10 Without Bleeding: Four RCTs compared vitamin K with placebo for patients with INR 4.5 to 10, and all reported on major bleeding as an outcome (Table 16, Table S14).^{168,169,172,173} Pooled analysis suggests that rates of major bleeding were similar over 1 to 3 months of follow-up (2% [10 of 452] of patients receiving vitamin K vs 0.8% [four of 4/471] in the placebo group). Thromboembolism as reported in three of the studies^{168,169,172} and occurred in five of 423 patients in the vitamin K group vs four of 441 patients in the placebo group. In summary, although vitamin K use may reverse supratherapeutic INRs more rapidly, there is no evidence of benefit for patient-important outcomes.

INR > 10 *Without Bleeding*: We found no randomized trials that tested treatment strategies in this patient group. A prospective case series of 107 patients with INR > 10 and without evidence of bleeding showed that 2.5 mg of oral vitamin K resulted in a low rate of observed major bleeding by 90 days (3.9%; 95% CI, 1.1-9.7).¹⁷⁴ Another retrospective study of 89 patients found that such patients given oral vitamin K 2 mg were less likely to still have an INR > 5 by day 3 compared with those who only had warfarin withheld (11.1% vs 46.7%).¹⁷⁵ Patient preferences and clinical assessment of risks of thrombosis and bleeding are likely important factors in determining whether to give vitamin K. In summary, the benefit and harm of vitamin K administration for patients with an INR>10 and no bleeding are unclear, although the risk of bleeding may be substantial.

Recommendations

9.1.

(a) For patients taking VKAs with INRs between 4.5 and 10 and with no evidence of bleeding, we suggest against the routine use of vitamin K (Grade 2B).

 Table 16—[Section 9.1] Vitamin K vs Only Withholding VKA for Patients Taking Warfarin With an Elevated INR (4.5-10) Without Evidence of Bleeding^{a,168,169,172,173}

				Anticipated	Absolute Effects ^a
Outcomes	No. of Participants (studies), Follow-up	Quality of the Evidence (GRADE)	Relative Effect (95% CI)	Risk With Only Holding VKA	Risk Difference With Vitamin K (95% CI)
Major bleeding	923 (4 studies ^b), 1-3 mo ^c	Moderate ^{d.e} due to imprecision	OR 2.6 (0.8-9.8)	8 per 1,000	13 more per 1,000 (from 2 fewer to 69 more)
Thromboembolism	864 (3 studies ^f), 1-3 mo ^c	Moderate ^{d,e} due to imprecision	OR 1.3 (0.3-6.6)	9 per 1,000	3 more per 1,000 (from 6 fewer to 48 more)
Mortality all-cause mortality	863 (3 studies ^f), 1-3 mo ^c	Moderate ^{d,e} due to imprecision	OR 1.3 (0.6-2.9)	29 per 1,000	9 more per 1,000 (from 12 fewer to 51 more)

See Table 1 and 2 legends for expansion of other abbreviations. See Table 1 through 3 legends for expansion of other abbreviations. «Time frame is days.

INR 6.0-12.0 in Ágeno et al.¹⁷³

^bNone of the studies specified whether any bleeding events were fatal or intracranial.

°Follow-up was 3 mo in both studies by Crowther et al.^{168,169}

^dTwo small studies, Ageno et al¹⁷² and Ageno et al,¹⁷³ were open label.

eWide CIs encompass both benefit and significant harm.

^fAgeno et al¹⁷³ did not report thromboembolism, and Ageno et al¹⁷² did not report deaths.

(b) For patients taking VKAs with INRs > 10.0 and with no evidence of bleeding, we suggest that oral vitamin K be administered (Grade 2C).

9.2 Clinical Prediction Rules for Bleeding While Taking VKA

The annual incidence of warfarin-associated major bleeding is estimated at 1% to 3%.¹⁷⁶ Clinicians continually struggle with estimating and weighing patient risk of thromboembolic events with risk of major bleeding. A clinical prediction rule for an individual's risk of bleeding while taking warfarin or other VKAs would be very useful if prediction of low risk reassured patients sufficiently to start VKA therapy or, more importantly, if prediction of high risk of bleeding was sufficiently accurate to withhold VKA therapy.

A 2007 systematic review by Dahri and Loewen¹⁷⁷ examined studies developing clinical prediction rules for bleeding while taking warfarin for any indication. Seven studies were included, with the primary outcome being the ability of the clinical prediction rule to distinguish between patients at high vs low risk of experiencing major bleeding.^{6,178-183} The performance of a rule was considered moderate if the likelihood ratio for a high score to predict major bleeding was > 5.0 and strong if it was > 10.0.^{184,185} Two variants of the same clinical prediction rule had a likelihood ratio of $\sim 9.178,179$ The independently validated mOBRI (modified Outpatient Bleeding Risk Index)179 includes the following predictors: age ≥ 65 years, history of stroke, GI bleed in the past 2 weeks, and at least one of the following comorbidities: recent myocardial infarction, hematocrit level < 30%, creatinine level >1.5 mg/dL, or diabetes mellitus. One point is given for each of the four risk factor categories, with high risk defined as ≥ 3 points.

Since the 2007 systematic review, two additional clinical prediction rules have been published.¹⁸⁶⁻¹⁸⁸ Table 17^{179-183,186-190} summarizes the clinical prediction rules according to (1) the proportion of patients classified as high risk, (2) the risk of major bleeding measured in that subset, and (3) the annual risk of stroke required to prefer an alternative therapy with a lower risk of bleeding for patients with atrial fibrillation. The column on stroke risk required is based on the assumption of a stronger preference for avoiding stroke compared with avoiding a major bleeding event by a factor of 3:1.² Using this metric, most of the rules would suggest a prohibitively high risk of major bleeding only for patients with a CHADS₂ (congestive heart failure, hypertension, age ≥ 75 years, diabetes mellitus, prior stroke or transient ischemic attack) score of 0, a group for whom VKA therapy might not be preferred anyway. However, for patients with a greater preference of avoiding bleeding events compared with stroke, use of CHADS₂ score along with a clinical prediction rule, such as mOBRI, may provide some prognostic guidance. Similarly, the studies involving a population treated for VTE do not identify a group with a risk of bleeding sufficiently high to preclude secondary prophylaxis with VKA. A clinical prediction rule that could predict an individual's risk of both benefit and harm at the time of initiation of VKA therapy would be desirable, but none has been validated.¹⁹¹

Recommendation

9.2. For patients initiating VKA therapy, we suggest against the routine use of clinical prediction rules for bleeding as the sole criterion to withhold VKA therapy (Grade 2C).

9.3 Treatment of Anticoagulant-Related Bleeding

When patients present with major bleeding due to VKA use, rapid reversal of anticoagulation is desirable, particularly if the bleeding is life threatening. Several products are available to assist, with treatment, often combining vitamin K with one of prothrombin complex concentrate (PCC), fresh frozen plasma (FFP), or recombinant factor VIIa. FFP has the disadvantage of potential allergic reaction or transmission of infection, preparation time, and higher volume. PCC and recombinant factor VIIa are more rapidly concentrated with less infection transmission risk but have not been compared with FFP in adequately powered RCTs.

Vitamin K is given to sustain the effects of the other products because of the relatively short half-lives of the latter. In emergency situations, vitamin K 10 mg IV instead of given orally is recommended because of its more rapid onset.^{24,171,192} IV injection of vitamin K is reported to cause anaphylaxis in three of 100,000 patients, resulting in advice to infuse slowly.¹⁹³ In one RCT of patients with INR 6 to 10 without bleeding, IV injection (0.5 mg) compared with po (2.5 mg) phytonadione more rapidly brought the INR back to therapeutic range (11 of 24 patients vs 0 of 23 patients at 6 h).¹⁹² However, by 24 h, the mean INR in both groups was similar. In a second RCT of patients with INR 6 to 10, vitamin K 0.5 mg IV led to faster resolution than vitamin K 3 mg SC, with an INR < 5 in 95% vs 45% of patients and a mean INR of 3.7 vs 5.4 at 24 h.194 Accordingly, SC injection is not recommended.

Several studies have compared products in addition to vitamin K, three of which reported rates of intracranial hemorrhage. A small case series of 17 patients compared the use of FFP and three-factor PCC; all patients received vitamin K.¹⁹⁵ The mean INR decreased from 2.83 to 1.22 within 4.8 h in patients receiving PCC vs from 2.97 to 1.74 within 7.3 h for those receiving FFP (P < .001). The reaction level

Study Acronym or Authors	Sample, No.	Population	Follow-up Duration, Mean	Proportion With High Risk	Major Bleeding Events in High-Risk Group	Stroke Risk Required to Avoid VKA ^a
mOBRI179	Derivation: 565 Validation: 264	VTE, valves, other	Derivation: 2 y Validation: 6-7 y	Derivation: 6.1% Validation: 6.9% ($\ge 3 p$)	Derivation: 3 m: 23% 12 m: 48% Validation: 3 m: 6% 12 m: 30%	N/A ^b
mOBRI—validation ¹⁵⁰	Validation 1,269	50% AF, 50% other diagnoses	1 y	15.4%	Patients with AF: 12.3%/y	<4%/y
mOBRI-validation ⁶	Validation: 222	VTE	1.5 y	1%	0%0	N/A ^b
Kuijer et al ¹⁸²	Derivation: 241 Validation: 780	VTE	3 mo	21% 19% (> 3 p)	Derivation: 14%/3 mo Validation: 7%/3 mo	N/A^b
HEMORR ₂ HAGES ¹⁸¹	1,604 discharged on warfarin	AF	$0.83 \mathrm{y}$	16.3% (3.4 p)	8.8%/y	<3%/y
Shireman et al ¹⁵³	Derivation: 19,875 Validation: 6,470	AF, warfarin naïve	3 mo	Validation: 3.4% (score ≥ 2.19)	Validation: $5.4\%/3$ mo ^c	<1.8% first 3 mo
RIETE registry ¹⁸⁷	Derivation: 13,057 Validation: 6,572	VTE	3 mo	Derivation: 5.8% Validation: 5.2%	Derivation: 7.3%/3 mo Validation: 6.2%/3 mo	N/A^b
HAS-BLED ¹⁸⁶	Derivation: 3,381 Validation: 3,071	AF	1 y	Derivation: 1.7% Validation: $7.9\% (\ge 3 p)$	Derivation: 20%/y Validation: 4.9%/y	Validation: $< 1.7\%/y$
HAS-BLED—separate validation ¹⁸⁸	Validation: 3,665	AF	499 d?	18.7%	6.7%/y	<2%/y
HAS-BLED—separate validation ¹⁸⁵ Validation: 3,665 AF 499 d? 18.7% 6.7% , 5.7% 5.7% 5.7% y -2% /y -2% /y AE = atrial fibrillation; HAS-BLED = hypertension, abnormal renal/liver function, stroke, bleeding history or predisposition, labile INR, elderly (>65), drugs/alcohol concomitantly; HEMORR ₂ HAGES = hepatic or renal disease, ethanol use, malignancy, reduced platelet count. re-bleeding, hypertension, anemia, genetic factors, elevated risk of fall including neuropsychiatric disease, stroke: mOBRI = modified	Validation: 3,665 ypertension, abnormal ren nev. reduced platelet co	AF aal/liver function, stroke, bleu unt re-bleeding hynertens	499 d? eding history or predisp sion memia genetic f	18.7% osition, labile INR, elderly (>65) orders elevated rick of fall inclu	6.7%/y , drugs/alcohol concomitantly; F	

Table 17–[Section 9.2] Clinical Prediction Rules for VKA-Associated Major Bleeding

Outpatent Bieeding Kisk Index; p = total points within the CFR; KLETE = Computerized Registry of Patients with Venous Thromboembolism. See Table 1 legend for expansion of abbreviation. Based on assumption that the dysutility of a stroke is three times that of a major bleeding event, where most major bleeding is GI.

^bFor patients with VTE, the alternative would be no therapy, which can be estimated to result in a risk of recurrence of 22% to 29%^{189,190} during the first 3 mo. With the assumption that the dysutility of a major bleeding event, where the majority consists of GI bleeding, the risk of major bleeding would have to be at least the same during the first 3 mo to avoid VKA.

"This risk normally will decrease after the first 3 mo of treatment.

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grade, used to assess symptoms and signs of intracerebral hemorrhage, suggested less progression in those receiving PCC (0.2 vs 1.9 grades on a scale of 1-8) (P<.05). Another small before-after study of 12 patients reported that the six patients receiving three-factor PCC compared with six age- and sex-matched historical controls given FFP had a mean INR correction time of 41 min for PCC vs 115 min for FFP.¹⁹⁶

Finally, a small RCT compared factor IX complex concentrate (four-factor PCC) plus FFP vs FFP alone in 13 patients (five in factor IX concentrate and eight in FFP).¹⁹⁷ Factor IX concentrate plus FFP corrected the INR more quickly than FFP alone (2.95 vs 8.9 h, P < .01). In addition, five of eight patients in the FFP-alone group experienced significant fluid overload complications, despite monitoring of central venous pressure and the use of furosemide, compared with no reported complications in the combination group.

FFP has also been compared with four-factor PCC in patients undergoing cardiopulmonary bypass surgery.¹⁹⁸ Forty patients admitted to the hospital for urgent or semiurgent cardiac surgery who were taking oral anticoagulants (INR 2.1-7.8) were randomized, 20 to each treatment. Seven PCC patients vs no FFP patients had an INR < 1.5 by 15 min (P = .007); an additional six PCC vs four FFP patients had this level an hour later (P = .70).

Three very small case series addressed the use of recombinant factor VIIa. In a series of 13 patients presenting with bleeding (four patients), requiring rapid reversal for interventions (five patients), or with an INR > 10 and not good candidates for FFP (four patients), all had a reduction in INR after administration but to variable degrees.¹⁹⁹ Use in four patients presenting with major bleeding (two with spinal cord hemorrhages and two with intracerebral hemorrhages) resulted in a normal INR within 2 h, with no complications reported.²⁰⁰ Finally, in a series of seven patients with acute intracranial hemorrhage while taking warfarin, the mean INR was reduced from 2.7 prerecombinant factor VIIa to 1.1 afterward. Several of the patients also received vitamin K and FFP. Five of the patients survived with severe disability, and two died.²⁰¹ Factor concentrates including PCC are expensive and, therefore, not available in some jurisdictions.

Recommendations

9.3. For patients with VKA-associated major bleeding, we suggest rapid reversal of antico-agulation with four-factor PCC rather than with plasma (Grade 2C).

We suggest the additional use of vitamin K 5 to 10 mg administered by slow IV injection rather than reversal with coagulation factors alone (Grade 2C).

9.4 Investigating Anticoagulant-Associated Bleeding

No randomized trials have addressed different strategies of investigating bleeding in patients taking anticoagulants. The topic is of great practical importance in patient management, but the evidence found was not of sufficient quality to make a recommendation. One small case-control study found the monthly incidence and prevalence of hematuria to be 0.05% and 3.2% in those taking anticoagulants vs 0.08% and 4.8% for those in the control group.²⁰² Subsequent diagnosis of cancer was also similar at two of 32 patients in the anticoagulation group compared with one of 11 patients in the control group. Two small case series of patients investigated for anticoagulant-associated hematuria found two of 30 and four of 24, respectively, had neoplastic disease.^{203,204} A retrospective analysis of all patients presenting with gross hematuria over a 9-year period while taking anticoagulant or aspirin therapy found that 25% (six of 25) of those patients presenting with hematuria were found to have a tumor.²⁰⁵

Several studies addressed the question of GI bleeding. A retrospective series of 166 patients presenting with lower GI bleeding, with 100 of the patients taking an antiplatelet or anticoagulant and 66 not, found that nine of 88 (10.2%) patients taking anticoagulants had colon cancer compared with two of 62 (3.2%) not taking anticoagulants.²⁰⁶ Another analysis of 98 patients taking warfarin who presented to a Veterans Affairs hospital with acute GI bleeding found on endoscopy that 52 of the 71 had upper-GI lesions, whereas on colonoscopy, 26 of 41 had lesions, including five cancers.²⁰⁷ In summary, although the data are of low quality, they suggest that there might be sufficient incidence of pathologic causes for VKA-associated hematuria or GI bleeding to warrant investigation.

10.0 OTHER

10.1 Intensive Patient Education and Anticoagulation Outcomes

Intensive patient education (defined as dedicated patient education sessions beyond the usual VKA information distributed by pamphlet or the patient's usual provider) has been proposed to reduce adverse events related to anticoagulation and to improve TTR. Although better patient knowledge of anticoagulation has been associated with improved INR control, these were no randomized trials, and INRs were surrogate outcomes.^{205,209}

Seven RCTs (n = 1,195) compared supplemental patient education with usual care and provided some data on clinical outcomes.²¹⁰⁻²¹⁶ Patient age varied widely (18-91 years), and the indications for VKA therapy included atrial fibrillation and VTE. Six of

the studies were based in anticoagulation clinics. Educational interventions varied among studies. Several allowed for only one teaching session delivered in person by a health-care professional, by means of a video presentation of a physician-patient interaction, or by a patient-administered self-guided instruction booklet.^{215,216} Others had repeated interaction with patients at daily intervals on a ward until discharge or at weekly or bimonthly intervals in outpatient clinics.^{210,212,213} The curricula covered similar content, including indications for VKAs, benefits and risks, the importance of INR surveillance, drug interactions, the effect of diet, and information on dose management. The amount and type of education in the control arms were unclear. The length of follow-up ranged from 3 to 6 months.

The quality of evidence based on these studies is low primarily because of limitations in design and imprecision for the clinical outcomes. In pooling data from three of the studies that reported clinical outcomes in a similar manner, there was no significant difference between supplemental patient education and usual care (VTE RR, 0.61 [95% CI, 0.06-6.56]; hemorrhage RR, 0.92 [95% CI, 0.04-20.56]).^{210,212,213} TTR was reported in four trials and was similar between groups (mean difference, 2.03%; 95% CI, -2.79-6.86).^{210-212,214} In the single study where the difference in intensity of education was marked (described as minimal vs daily intensive education for mean of 8 days), there was no difference in outcomes, including TTR.²¹² Although we found no compelling evidence favoring intensive patient education over standard patient education practices, the panel believed that a specific recommendation could not be made at this time.

Acknowledgments

Author contributions: As Topic Editor, Dr Holbrook oversaw the development of this article, including the data analysis and subsequent development of the recommendations contained herein

- Dr Holbrook: served as Topic Editor.
- Dr Schulman: served as Deputy Editor.
- Dr Witt: served as a panelist
- Dr Vandvik: served as a panelist.
- Dr Fish: served as a frontline clinician.
- Dr Kovacs: served as a panelist.
- Dr Svensson: served as a panelist.
- Dr Veenstra: served as a resource consultant.
- Dr Crowther: served as a panelist.

Dr Guyatt: served as guideline editor and contributed to the editing of this manuscript.

Financial/nonfinancial disclosures: The authors of this guideline provided detailed conflict of interest information related to each individual recommendation made in this article. A grid of these disclosures is available online at http://chestjournal.chestpubs. org/content/141/2_suppl/e152S/suppl/DC1. In summary, the authors have reported to *CHEST* the following conflicts of interest: Dr Crowther has served on various advisory boards, has assisted in the preparation of educational materials, and has sat on data safety and monitoring boards. His institution has received research funds from the following companies: Leo Pharma A/S, Pfizer Inc, Boerhinger Ingelheim GmbH, Bayer Healthcare Pharmaceuticals, Octapharm AG, CSL Behring, and Artisan Pharma. Personal total compensation for these activities over the past 3 years totals less than US \$10,000. Dr Guyatt is co-chair of the GRADE Working Group and Dr Vandvik is a prominent contributor to the GRADE Working Group. Drs Holbrook, Schulman, Witt, Fish, Kovacs, Svensson, and Veenstra have reported that no potential conflicts of interest exist with any companies/organizations whose products or services may be discussed in this article.

Role of sponsors: The sponsors played no role in the development of these guidelines. Sponsoring organizations cannot recommend panelists or topics, nor are they allowed prepublication access to the manuscripts and recommendations. Guideline panel members, including the chair, and members of the Health & Science Policy Committee are blinded to the funding sources. Further details on the Conflict of Interest Policy are available online at http://chestnet.org.

Endorsements: This guideline is endorsed by the American Association for Clinical Chemistry, the American College of Clinical Pharmacy, the American Society of Health-System Pharmacists, the American Society of Hematology, and the International Society of Thrombosis and Hematosis.

Additional information: The supplement Tables can be found in the Online Data Supplement at http://chestjournal.chestpubs. org/content/141/2_suppl/e152S/suppl/DC1.

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Evidence-Based Management of Anticoagulant Therapy

Antithrombotic Therapy and Prevention of Thrombosis, 9th ed: American College of Chest Physicians Evidence-Based Clinical Practice Guidelines

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		Q	Quality Assessment	nent					Summary of Findings	lings	
						_	Study Event Rates (%)	: Rates $(\%)$		Anticipated A	Anticipated Absolute Effects
Participants (Studies), Follow-up	Risk of Bias	Inconsistency	Indirectnes	Indirectness Imprecision	Publication Bias	Overall Quality of Evidence	With Warfarin 5-mg Loading Dose Nomogram	With Warfarin 10-mg Loading Dose Nomogram	Relative Effect (95% CI)	Risk With Warfarin 5-mg Loading Dose Nomogram	Risk Difference With Warfarin 10-mg Loading Dose Nomogram (95% CI)
					Bleeding even	Bleeding events (critical outcome)	ie)				
$\frac{420}{5-90} \frac{(3 \text{ studies}^{\text{s-c}})}{d^d},$	Serious	No serious inconsistency	Serious ^f	Very serious ^g	Undetected	Very low ^{est} due to risk of bias, indirectness, and imprecision	1/204 (0.49)	2/216 (0.93) ^h	OR 1.90 (0.17-21.1)	Mod 10 per 1,000	Moderate 00 9 more per 1,000 (from 8 fewer to 166 more)
					Recurrent VT	Recurrent VTE (critical outcome)	(e)				(
420 (3 studies ^{a.c.}), 5-90 d	Serious	No serious inconsistency	Serious ^f	Very serious ^g	Undetected	Very low ^{eg} due to risk of bias, indirectness, and	0/204 (0)	3/216 (1.4) ⁱ	OR 6.72 (0.34-131.88)	Mod 0 per 1,000	Moderate 0
Biblioambur Cue	Mar MA	Cinchard I Vacua	n C ot ol A	mandomized tri	ol comparing 5 m	imprecision	forting looding doe	ac Arch Intan	Med 1000.150	(1).46.48 Hourison	n I Iohneton M
^a biolography: Cri Massicotte MP, C Comparison of 1 2003;138:714-415 2006;98:535-537. (^a All pooled studie:	owther MA, ' frowther M, N (0-mg and 5- (0-mg and 5- (0-mg and 5- (0-mg and 5- (0-mg and 5) (0-mg and 2) (0-mg and 3) (0-mg and 3) (0-m	biolography: Crowther MA, Consperg J, Kearon C, et al. A rando Massicotte MP, Crowther M, Moffat K, Hirsh J. Comparison of 5-mg a Comparison of 10-mg and 5-mg warfarin initiation nomograms - 2003;138:714-419. Quiroz R, Gerhard-Herman M, et al. Compariso 2006;98:535-537. Schulman S, Lockner D, Bergstrom K, Blomback M ^ All pooled studies included only patients with acute VTE. Studies fro	n C, et al. A comparison of ation nomog M, et al. Conr rom K, Blomh te VTE. Stud	Frandomized tr. 5-mg and 10-mg rams together parison of a sin ack M. Intensiv ies from which (Biolography: Crowther MA, Ginsberg J, Keaton C, et al. A randomized trial comparing 5-mg and 10-mg warfarin herapy. Am Intern Med. 1997;136:133-136. Kovacs MJ, Rodger M, Anderson DR, et al. Massicotte MP, Crowther M, Moffat K, Hirsh J. Comparison of 5-mg and 10-mg loading doses in initiation of warfarin therapy. Am Intern Med. 1997;126:133-136. Kovacs MJ, Rodger M, Anderson DR, et al. Comparison of 5-mg and 10-mg loading doses in initiation of warfarin therapy. Am Intern Med. 1997;126:133-136. Kovacs MJ, Rodger M, Anderson DR, et al. Comparison of 5-mg and 10-mg loading doses in initiation of warfarin therapy. Am Intern Med. 1997;126:133-136. Kovacs MJ, Rodger M, Anderson DR, et al. 2003;138:714-419. Quiroz R, Gerhard-Herman M, et al. Comparison of a single end point to determine optimal initial warfarin dosing (5 mg vs 10 mg) for venous thromboembolism. Am Intern Med. 2006;98:535-537. Schulman S, Lockner D, Bergstrom K, Blomback M. Intensive initial oral anticoagulation and shorter heparin treatment in deep vein thrombosis. Thromb Haemost. 1984:52:276-280.	g and 10-mg war uitiation of warfarin ar-weight heparin etermine optimal agulation and shor od are Kovacs et al	tarm loading do: therapy. Ann Int. 1 for outpatient initial warfarin c ter heparin treati , Quiroz et al, and	ies. Arch Interv im Med. 1997;1: treatment of losing (5 mg vs ment in deep ve 1 Schulman et a	n Med. 1999;103 26:133-136. Kovac acute venous thi 10 mg for venou sin thrombosis. Th l.	(1):40-45. Harriso s MJ, Rodger M, A romboembolism. <i>z</i> is thromboembolis <i>tromb Haemost</i> . 19	n L., Johnston M., nderson DR, et al. <i>Ann Intern Med.</i> :m. <i>Am J Cardiol.</i> 84:52:276-280.
^b Minimal loss to ft (laboratory value, ^e Results based on it, with one part sh dFive days is the m	ollow-up; adh international only three stu nowing statisti	^b Minimal loss to follow-up; adherence to intention-to-treat principle in two of th (laboratory value, international normalized ratio); adequate allocation concealme ^c Results based on only three studies; one study shows no difference; one study s it, with one part showing statistically significant reduction and the other did not. ^d Five days is the mean follow-un period for patients in the loading dose warfant.	-to-treat prin- adequate allo ows no differ duction and the fs in the loadi	ciple in two of th cation concealm ence; one study is other did not no dose warfarii	^b Minimal loss to follow-up; adherence to intention-to-treat principle in two of three studies; follow-up period is short but adequate for this outcome; any lack of blinding should not affect objective outcome (laboratory value, international normalized ratio); adequate allocation concealment; and sample size calculations reported for two of three studies. ^c Results based on only three studies; one study shows no difference; one study shows statistically significant reduction in time to therapeutic international normalized ratio; and one study had two parts to it, with one part showing statistically significant reduction and the other did not.	-up period is short se calculations rep ignificant reductio lman et al (this wa	but adequate for orted for two of t on in time to ther s the shortest ner	this outcome; a hree studies. apeutic internat	ny lack of blindin _ų tional normalized is available)	g should not affect of ratio; and one stud	objective outcome y had two parts to
Adequate allocation concealmer	ion concealm	ent; adjudicators bl	s blinded in two	vo of three studies	Adequate allocation concealing the restored to be a set of the studies of a construction of three studies); minimal loss to follow-up; intention to treat followed in 	l data collectors bl	linded in zero of 1	three studies); n	ninimal loss to fol	low-up; intention to	o treat followed in

ώ two of three studies, but follow-up period is very short in two of three studies (5 d-2 wk).

^fIndirect given application aimed at VTE outpatients.

*No studies were powered to detect differences in bleeding events between groups. Number of events is too sparse to draw any conclusions.

¹One major bleeding event in the 10-mg group vs none in the 5-mg in Quiroz et al; no bleeding events in either group in Schulman et al; one major bleeding event per group in Kovacs et al. No recurrent VTE in either group for Quiroz et al or Schulman et al; three in the 10-mg group vs none in the 5-mg group in Kovacs et al.

		Q	Quality Assessment	t					Summary of Findings	ndings	
						_	Study E	Study Event Rates $(\%)$		Anticipate	Anticipated Absolute Effects
Participants (Studies), Follow-up	Risk of Bias	Inconsistency	Indirectness	Imprecision	Publication Bias	Overall Quality of Evidence	With Usual Dosing Strategies	With Pharmacogenetic- Based Testing	Relative Effect (95% CI)	Risk With Usual Dosing strategies	Risk Difference With Pharmacogenetic- Based Testing (95% CI)
			Majo	or bleeding (noi	nfatal) (critical	Major bleeding (nonfatal) (critical outcome; assessed with clinical evaluation ^a)	l with clinical	evaluation ^a)			
468 (3 studies), 1-3 mo	468 (3 studies),No serious riskNo serious1-3 moof biasinconsist	No serious inconsistency	No serious indirectness	Serious ^b	Undetected [°]	Undetected [®] Moderate ^{lo,c} due to imprecision	6/234 (2.6) ^d	4/234 (1.7) ^e	OR 0.66 (0.16-2.45)	26 per 1,000 ^e	9 fewer per 1,000 (from 21 fewer to 35 more)
				Thromboem	bolism (critica	Thromboembolism (critical outcome; assessment not reported)	nent not repor	ted)			
163 (3 studies), 1-3 mo	463 (3 studies), No serious risk No serious 1-3 mo of bias inconsist	No serious inconsistency	No serious indirectness	Serious ^b			2/231 (0.87) ^{d,f}	0/232(0)	OR 0 (0-3.4) ^f	$OR \ 0 \ (0-3.4)^f 9 \ per \ 1,000^{d.f}$	9 fewer per 1,000 (from 9 fewer to 20 more)
Bibliography: A Circulation. 20 Clin Med Res 20 Clin Med Res rolled study C tudy in Chines Bleeding rate : Total sample s All studies wer Control rate is Untervention r Hillman et al r to events in bot	nderson JL, Hor 07;116(22):2563 2005;3(3):137-14; <i>lin Pharmacol Th</i> , <i>e</i> patients. <i>Pharm</i> <i>n</i> Hillman et al w rize below optimal <i>e</i> funded by publ median percenta the is mean percent eported one DV7 h groups. Huang	Bibliography: Anderson JL, Horne BD, Stevens SM, et al; for Couma-Gen Investigators. Randomized trial <i>Circulation</i> . 2007;116(22):2563-2570. Hillman MA, Wilke RA, Yale SH, et al. A prospective, randomized J <i>Clin Med Res</i> . 2005;3(3):137-145. Caraco Y, Blotnick S, Muszkat M. CYP2C9 genotype-guided warfarin pru trolled study. <i>Clin Pharmacol Ther</i> : 2008;53(3):460-470. Huang SW, Chen HS, Wang XQ, et al. Validation of V study in Chinese patients. <i>Pharmacol Ther</i> : 2008;19(3):226-234. 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Huang SW cs. 2009;19(3):222 severity was not ganizations. e usual dosing gr ross all intervent coembolism (unc VTE in either gr VTE in either gr	ima-Gen Invest lue SH, et al. A M. CYP2C9 gen M. CYP2C9 gen (Chen HS, Wa 6-234. VKA = v reported; inclue onp across all st ion groups (unw lear whether th oup. Anderson	igators. Rando prospective, ri potype-guided ng XQ, et al. V ritamin K antag ded one case h ded one case h udies reportin, veighted). te same or diff et al did not se	-Gen Investigators. Randomized trial of gen SH, et al. A prospective, randomized pilot t ZYP2C9 genotype-guided warfarin prescrib nen HS, Wang XQ, et al. Validation of VKOR 34. VKA = vitamin K antagonist. orted: included one case hematuria, two case orted: included one case hematuria, two case across all studies reporting that outcome. groups (unweighted). . whether the same or different patients) ar o. 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[•] Hillman et al reported one DVT and one thromboenbolism (unclear whether the same or different patients) among 20 control patients and no events in the intervention events in both groups. Huang et al reported no VTE in either group. Anderson et al did not separate VTE.	farin dosing in J i mitiation using ety of anticoagu terindividual wan ae control group ae control group	patients initiatin g CYP2C9 geno lation: a prospec rfarin maintenar	Bibliography: Anderson JL. Horne BD, Stevens SM, et al. for Couma-Gen Investigators. Randomized trial of genotype-guided versus standard warfarin dosing in patients initiating oral anticoagulation. <i>Circulation</i> . 2007;116(22):2563-2570. Hillman MA, Wilke RA, Yale SH, et al. A prospective, randomized pilot trial of model-based warfarin dose initiation using CYP2C9 genotype and clinical data. <i>Clin Med Res</i> . 2005;3(3):137-145. Caraco Y, Blotnick S, Muszkat M. 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Validation of VKORC1 and CYP2C9 genotypes on interindividual warfarin maintenance dose: a prospective study in Chinese patients. <i>Pharmacol Ther</i> . 2008;83(3):460-470. Huang SW, Chen HS, Wang XQ, et al. Validation of WCORC1 and GYP2C9 genotypes on interindividual warfarin maintenance dose: a prospective study in Chinese patients. <i>Pharmacol Ther</i> . 2008;83(3):460-470. Huang SW, Chen HS, Wang XQ, et al. Validation of WCORC1 and CYP2C9 genotypes on interindividual warfarin maintenance dose: a prospective study in Chinese were funded by public or norportis organized metal vas four of 20, but severity was not reported, included one case hematuria, two cases epistaxis, and one GI bleed in the control group. ^o Total sumple size head provemotine isize. <i>Clo</i>

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Table S3—[Section 2.3] Evidence Profile: VKA Started Early vs Late in Heparin Patients With Acute Thromboembolism

									ļ		
		ت ت	Quality Assessment						Summary of Findings	sguipt	
_						-	Study Event Rates (%)	t Rates $(\%)$		Anticipated A	Anticipated Absolute Effects
Participants (Studies), Follow-up	Risk of Bias	Inconsistency	Indirectness	Imprecision	Publication Bias	Overall Quality of Evidence	With Late	With VKA Started Early	Relative Effect (95% CI)	Risk With Late	Risk Difference With VKA Started Early (95% CI)
					Death (crit	Death (critical outcome)					
807 (4 studies), 3-6 mo	No serious risk of biasª	Serious ^b	No serious indirectness	Serious	Undetected	Low ^{a.e.} due to inconsistency and imprecision	23/394 (5.8)	27/413 (6.5)	RR 1.28 (0.43-3.85)	58 per 1,000	16 more per 1,000 (from 33 fewer to 166 more)
Recurrent tl	hromboembolisn	Recurrent thromboembolism (critical outcome; DVT assessed with	DVT assessed w.	ith venography, assessed with	Doppler ultraso two-dimensions	h venography. Doppler ultrasonography, or impedance plethysmography; PE assessed with lung scanning; left ventricle thrombus assessed with two-dimensional transthoracic echocardiography)	lance plethysm ocardiography)	ography; PE as	sessed with lung :	scanning; left vent	ricle thrombus
807 (4 studies), 3-6 mo	Serious ^d	No serious inconsistency	No serious indirectness	Serious°	Undetected	Low ^{e,d} due to risk of bias and imprecision	16/394 (4.1)	15/413 (3.6)	RR 0.92 (0.46-1.82)	41 per 1,000	3 fewer per 1,000 (from 22 fewer to 33 more)
Major bleedin£	g (critical outcon	ne; assessed with re	quired blood trar	nsfusion or bleec led to hem	ling in body cavior of the second sec	Major bleeding (critical outcome; assessed with required blood transfusion or bleeding in body cavity or bleeding that required anticoagulation withdrawal for intracranial or retroperitoneal bleeding or level decrease of ≥ 2 g/dL or to death)	required anticon or to death)	oagulation with	drawal for intracr	anial or retroperit	oneal bleeding or
807 (4 studies), 0.5-6 mo	Serious ^d	No serious inconsistency	No serious indirectness	Serious°	Undetected	Low ^{e,d} due to risk of bias and imprecision	13/394 (3.3)	16/413 (3.9)	RR 1.22 (0.58-2.56)	33 per 1,000	7 more per 1,000 (from 14 fewer to 51 more)
			Hospital	al utilization (day	vs) (important o	utilization (days) (important outcome; better indicated by lower values)	icated by lower	values)			
536 (3 studies), follow-up for hospital, 2 d to 6 mo	No serious risk of bias	No serious inconsistency	No serious indirectness	No serious imprecision	Undetected	High	263	273	:	The mean hospital utilization in the control groups was 14 d	The mean hospital utilization in the intervention groups was 4.07 d lower (4.76-3.37 d lower)
Bibliography: Q ^a For three of for sionals in some s ^b The value for th ^c The 95% CIs an ^d Potential limita (outcome assesss	ayyum F, Holbrc ur studies, conce studies is not like he I^2 test for dea round the absolu tions in design f ors were blinded	Bibliography: Qayyum F, Holbrook A, Lam J, Kovacs MJ, Schulman S, unpublished data, ²⁰ ^a For three of four studies, concealment of allocation was unclear. However, this alone was 1 sionals in some studies is not likely to affect incidence of this outcome. ^b The value for the I ² test for death was 55%; therefore, it was rated down for inconsistency: ^c The 95% CIs around the absolute risk values were very wide for this outcome. ^d Potential limitations in design for this outcome, including allocation sequence concealmen (outcome assessors were blinded in three of four studies).	cs MJ, Schulman n was unclear. Hc cce of this outcom ore, it was rated d very wide for this cluding allocatior rdies).	S, unpublished wever, this alon e. lown for inconsi. s outcome. t sequence conc	data, 2011. PE : e was not seen : stency. ealment not rep	Bibliography: Qayyum F, Holbrook A, Lam J, Kovacs MJ, Schulman S, unpublished data, 2011. PE = pulmonary embolism; RR = risk ratio. See Table S2 legend for expansion of other abbreviation. "For three of four studies, concealment of allocation was unclear. However, this alone was not seen as a compelling reason to downgrade evidence for this outcome. Lack of blinding of health-care profes- sionals in some studies is not likely to affect incidence of this outcome. "The value for the <i>I</i> ² test for death was 55%; therefore, it was rated down for inconsistency. "The 95% CIs around the absolute risk values were very wide for this outcome." ("The 95% CIs around the absolute risk values were very wide for this outcome." (Hull et al 1990) (outcome assessors were blinded in three of four studies).	lism; RR = ris son to downgra our studies and	de evidence foi de evidence foi health-care pr	le S2 legend for ε r this outcome. Lá ofessionals blinde	xpansion of other tek of blinding of d in only one stud	abbreviation. nealth-care profes- y (Hull et al 1990)

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			Quality Assessment	at				Sı	Summary of Findings	sßt	
							Study Event Rates (%)	t Rates (%)		Anticipated Al	Anticipated Absolute Effects ^a
Participants (Studies), Follow-up	Risk of Bias	Risk of Bias Inconsistency	Indirectness	Imprecision	Imprecision Publication Bias	Overall Quality of Evidence	With 4-wk Recall Intervals	- With Prolonged INR Recall Intervals	Relative Effect (95% CI)	Relative Effect Risk With 4-wk (95% CI) Recall Intervals	Risk Difference With Prolonged INR Recall Intervals (95% CI)
				Thromboen	nbolism (critical or	utcome; assessmen	Thromboembolism (critical outcome; assessments variously defined)	(p)			
744 (2 studies), 163 patient-y	Serious ^b	No serious inconsistency	No serious indirectness	Serious	Undetected	Low ^{b,c} due to risk of bias and imprecision	5/384 (1.3)	6/360 (1.7)	RR 1.02 (0.13-8.32)	13 per 1,000	0 more per 1,000 (from 11 fewer to 95 more)
				Major ble	seding (critical outc	come; assessment:	Major bleeding (critical outcome; assessments variously defined)	-			
744 (2 studies), 163 patient-y	Serious ^b	No serious inconsistency	No serious indirectness	Serious	Undetected	Low ^{b,c} due to risk of bias and imprecision	16/384 (4.2)	16/360 (4.4)	RR 1.06 (0.54-2.1)	42 per 1,000	2 more per 1,000 (from 19 fewer to 46 more)
Bibliography: Fihn SD, M Intern Med. 1994;9:131-1: Clin Pathol. 2003;120:944. * Time frame is in months. ^b Lack of blinding; intentio	In SD, McD 4(9:131-139, 3;120:944-947 a months. g; intention tc	Bibliography: Fihn SD, McDonell MB, Vermes D, et al. A compute Intern Med. 1994;9:131-139; Pengo V, Barbero F, Biasiolo A, Pegora Clin Pathol. 2003;120:944-947. INR = international normalized ratio ^a Time frame is in months. ^b Lack of blinding; intention to treat not specified in Pengo et al; and i	D, et al. A com F, Biasiolo A, Pe _i onal normalized r l in Pengo et al; a	puterized inte goraro C, Cuc ratio. See Tabl md adherence	ized intervention to improve timing of outpatient follow o C, Cucchini U, Iliceto S. A comparison between six- ai See Table S3 legend for expansion of other abbreviation. dherence to recommended INR recall intervals was not τ	ve timing of outpi A comparison be pansion of other a INR recall interv	Bibliography: Film SD, McDonell MB, Vermes D, et al. A computerized intervention to improve timing of outpatient follow-up: a multicenter randomized trial in patients treated with warfarin. <i>J Gen Intern Med.</i> 1994;9:131-139; Pengo V, Barbero F, Biasiolo A, Pegoraro C, Cucchini U, Iliceto S. A comparison between six- and four-week intervals in surveillance of oral anticoagulant treatment. <i>Am J Clin Pathol.</i> 2003;120:944-947. INR = international normalized ratio. See Table S3 legend for expansion of other abbreviation. "Time frame is in months."	multicenter rand r-week intervals i ed in Fihn et al.	omized trial in pr in surveillance of	atients treated wit oral anticoagulan	h warfarin. <i>J Gen</i> t treatment. <i>Am J</i>

Table S4—[Section 3.1] Evidence Profile: Prolonged INR Recall Intervals Compared With Four-Week Recall Intervals

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• Wide CIs around the estimate of effect.

			Quality A	Quality Assessment				Su	Summary of Findings		
						_	Study Ev	Study Event Rates (%)		Anticipated	Anticipated Absolute Effects ^a
Participants (Studies), Follow-up	Risk of Bias	Inconsistency	Indirectness	Imprecision	Publication Bias	Overall Quality of Evidence	With Placebo	With Low- Dose Vitamin K Supplementation	Relative Effect (95% CI)	Risk With Placebo	Risk Difference With Low-Dose Vitamin K Supplementation (95% CI)
					Major bleeding (Major bleeding (important outcome)	le)				
626 (3 studies), No serious 168-180 d risk of bi	No serious risk of bias ^d	Serious ^b	No serious indirectness	Serious	Reporting bias strongly suspected ^d	Very lowb-e due to inconsistency, imprecision, and publication bias	0/219 (0)	3/407 (0.74)	2.61 (0.34-20.28) 0 per 1,000	0 per 1,000	Not estimable
				L	Thromboembolism (important outcome)	n (important outce	ome)				
626 (3 studies), 168-180 d	626 (3 studies), No serious risk No serious 168-180 d of bias ^e inconsist	No serious inconsistency	No serious indirectness	Very serious ^c	Reporting bias strongly suspected ^d	See comment	0/219 (0)	0/407 (0)	1.65 (0.08-34.03) 0 per 1,000	0 per 1,000	Not estimable
Bibliography: Rombouts F Kamali F. Vitamin K supp van Meegen E, van der M See Table S2 and S4 legen "Time frame is in morths.	ombouts EK, Rc iin K supplemen van der Meer F d S4 legends for in months.	Bibliography: Rombouts EK, Rosendaal FR, Van Der Meer Kamali F. Vitamin K supplementation can improve stability van Meegen E, van der Meer FJM. Vitamin K1 supplemen See Table S2 and S4 legends for expansion of abbreviations. "Time frame is in months.	Der Meer FJM. e stability of ant upplementation eviations.	Daily vitamin K ticoagulation fo to improve the	c supplementation r patients with un stability of anticos	i improves anticos explained variabil igulation therapy	ıgulant stability. ity in response t with vitamin K a	Bbliography: Rombouts EK, Rosendaal FR, Van Der Meer FJM. Daily vitamin K supplementation improves anticoagulant stability. J Thromb Haemost. 2007;5:2043-2048. Sconce E, Avery P, Wynne H, Kamali F. Vitamin K supplementation can improve stability of anticoagulation for patients with unexplained variability in response to warfarin. Blood. 2007;109:2419-2423. Gebuis EPA, Rosendaal FR, van Meegen E, van der Meer FJM. Vitamin K1 supplementation to improve the stability of anticoagulation therapy with vitamin K antagonists: a dose-finding study. Haematologica. 2011;96(4):583-589. See Table S2 and S4 legends for expansion of abbreviations.	. 2007;5:2043-2048. 2007;109:2419-242: înding study. <i>Haem</i>	. Sconce E, Av 3. Gebuis EPA natologica. 201	(ery P, Wynne H, 1, Rosendaal FR, 11;96(4):583-589.
^b Full definition ^c Studies not pov ^d Unable to rule	of major bleedin wered to detect l out because not	^b Full definition of major bleeding not provided in the Sconce et al study. Definition of major bleeding different in each study. ^c Studies not powered to detect bleeding or thromboembolic events. The sample sizes in trials by both Sconce et al and Romb ^d Unable to rule out because not enough studies exist to populate funnel plot.	the Sconce et al poembolic event ist to populate fi	study. Definitio s. The sample s unnel plot.	m of major bleedin izes in trials by bo	ng different in eac th Sconce et al an	sh study. d Rombouts et a	udy. Definition of major bleeding different in each study. The sample sizes in trials by both Sconce et al and Rombouts et al were small. The total number of events was extremely low. mel plot.	otal number of even	ıts was extrem	ely low.

Table S5-[Section 3.4] Evidence Profile: Low-Dose Vitamin K Supplementation Compared With Placebo for Patients Taking VKAs To Stabilize INR

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^e Allocation concealment not reported; uncertain whether outcome adjudicators were blinded.

			Ouality Assessment	lent					Summary of Findings	Soc.	
			Audity measure	ICIII				B C		cQ1	
							Study Event Rates (%)	Rates (%)		Anticipated	Anticipated Absolute Effects ^a
Participants (Studies), Follow-up	Risk of Bias	s Inconsistency	Indirectness	Imprecision	Publication Bias	Overall Quality of Evidence	With vs Usual Laboratory-Based Monitoring	With Patient Self-Testing/ Patient Self- Monitoring	Relative Effect (95% CI)	Risk with vs Usual Laboratory- Based Monitoring	Risk Difference With Patient Self- Testing/Patient Self-Monitoring (95% CI)
				Thromboem	nbolism (critical ou	itcome; assessed w	Thromboembolism (critical outcome; assessed with various methods $^{\mathrm{b}})$	[S ^b]			
7,759 (14 studies), 4.6-57 mo	Serious ^c	No serious inconsistency	No serious / indirectness	Ž	Undetected	Moderate ^e due to risk of bias	149/3,755 (4)	99/4,004 (2.5)	OR 0.58 (0.45-0.75)	40 per 1,000	16 fewer per 1,000 (from 10 fewer to 21 fewer)
				Major blee	ding (critical outco	ome; assessed with	Major bleeding (critical outcome; assessed with various definitions ^d)	(р			
7,867 (16 studies), 4.6-57 mo	Serious ^c	No serious inconsistency	No serious / indirectness	No serious imprecision	Undetected	Moderate [°] due to risk of bias	300/3,806 (7.9)	283/4,061 (7)	OR 0.87 (0.75-1.05)	79 per 1,000	10 fewer per 1,000 (from 19 fewer to 4 more)
				Mortalit	Mortality (critical outcome; assessed with all-cause mortality)	e; assessed with al	l-cause mortality)				
6,370 (13 studies),	Serious	Serious ^e	No serious indirectness	No serious imprecision	Undetected	Low ^{c,e} due to risk of bias,	369/3,123 (11.8)	298/3,247 (9.2)	OR 0.74 (0.63-0.87)	118 per 1,000	118 per 1,000 28 fewer per 1,000 (from 14 fewer
Bibliography:]	Bloomfield H	E, Krause A, Gré	eer N, et al. Met	ta-analysis: effec	ot of patient self-to	esting and self-m	anagement of long	-term anticoagula	ttion on major c	linical outcome	Bibliography: Bloomfield HE, Krause A, Greer N, et al. Meta-analysis: effect of patient self-testing and self-management of long-term anticoagulation on major clinical outcomes. Ann Intern Med.
2011;124:4/24.26 aTime frame is in years.	482. See 1able in years.	2011;1:04:4:4:2-452. See Table 52 through 54 legends for expansion of abbreviations. "Time frame is in years.	egends tor expans.	ion of abbreviat	lons.						
^b Designated a: ^e Flaws in study ^d Categorized a ^e Evidence of h tigators. Effect	s major by the y design, most is major by thu eterogeneity of home test	 study or categori: t commonly an abs e study or that me among studies thay among studies that 	zed as strokes, ne sence of informat st the ISCOAT (It it was probably at al normalized ratio	w or recurrent : tion about the al falian Study of C tributable to the o on clinical eve	^b Designated as major by the study or categorized as strokes, new or recurrent symptomatic DVT, FE, or arterial embolism. ^e Flaws in study design, most commonly an absence of information about the allocation concealment procedure or blinding. ^d Categorized as major by the study or that met the ISCOAT (Italian Study of Complications of Anticoagulant Therapy) critt ^e Evidence of heterogeneity among studies that was probably attributable to the THINRS (The Home INR Study) Machar figators. Effect of home testing of international normalized ratio on clinical events. <i>N Engl J Med.</i> 2010;363(17):1608-1620.	, PE, or arterial er ent procedure or l nticoagulant Ther fome INR Study) . 2010;363(17):160	^b Designated as major by the study or categorized as strokes, new or recurrent symptomatic DVT, PE, or arterial embolism. ^e Flaws in study design, most commonly an absence of information about the allocation concealment procedure or blinding. ^d Categorized as major by the study or that met the ISCOAT (Italian Study of Complications of Anticoagulant Therapy) criteria for major bleeding. ^e Evidence of heterogeneity among studies that was probably attributable to the THINRS (The Home INR Study) Matchar DB, Jacobson A, Dolor tigators. Effect of home testing of international normalized ratio on clinical events. N Engl J Med. 2010;363(17):1608-1620.	jor bleeding. son A, Dolor R, e	t al; THINRS E	xecutive Commi	Designated as major by the study or categorized as strokes, new or recurrent symptomatic DVT, PE, or arterial embolism. Flaws in study design, most commonly an absence of information about the allocation concealment procedure or blinding. Categorized as major by the study or that met the ISCOAT (Italian Study of Complications of Anticoagulant Therapy) criteria for major bleeding. Evidence of heterogeneity among studies that was probably attributable to the THINRS (The Home INR Study) Matchar DB, Jacobson A, Dolor R, et al; THINRS Executive Committee and Site Inves- gators. Effect of home testing of international normalized ratio on clinical events. <i>N Engl J Med.</i> 2010;363(17):1608-1620.
The reduction	n in mortality .	I he reduction in mortainty from all causes was largely influenced by one study.	as largely minuenc	ea by one study.							

Table S6-[Section 3.6] Evidence Profile: Patient Self-Testine/Self-Monitoring Compared With Usual Laboratoru-Based Monitoring for VKA Therapu Management

© 2012 American College of Chest Physicians. Reproduction of this article is prohibited without written permission from the American College of Chest Bowinioadetorform whestpolynate rites provides provide the American College of Chest Bowinioadetorform whestpolynate rites provide the American College of Chest Bowinioadetorform whestpolynate rites provide the American College of Chest Bowinioadetorform whestpolynate rites provide the American College of Chest Bowinioadetorform whestpolynate rites provide the American College of Chest Bowinioadetorform whestpolynate rites provide the American College of Chest Bowinioadetorform whestpolynate rites and the American College of Chest Bowinioadetorform and the American College of Chest Physicians

		Qué	Quality Assessment					Sı	Summary of Findings	ngs	
_						_	Study Event Rates (%)	: Rates $(\%)$		Anticipated /	Anticipated Absolute Effects ^a
Participants (Studies), Follow-up	Risk of Bias	Inconsistency	Indirectness	Imprecision	Publication Bias	Overall Quality of Evidence	With Manual Dosing	With Dosing Decision Support	Relative Effect (95% CI)	Risk With Manual Dosing	Risk Difference With Dosing Decision Support (95% CI)
			Throm	boembolism—in	itiation (critical	Thromboembolism—initiation (critical outcome; assessed with: variously defined)	d with: variously	defined)			
503 (4 studies), 3 mo	Serious ^b	No serious inconsistency	No serious indirectness	Serious	Undetected	Undetected Low ^{h,c} due to risk of bias, imprecision	16/255 (6.3)	9/248 (3.6)	RR 0 (0.27-1.37)	63 per 1,000	63 fewer per 1,000 (from 46 fewer to 23 more)
			Maj	or bleeding—ini	tiation (critical	Major bleeding—initiation (critical outcome; assessment variously defined)	ent variously def	ined)			
926 (7 studies ^d), 1-3 mo	Serious ^b	No serious inconsistency	No serious indirectness	Serious	Undetected	Undetected Low ^{b,c} due to risk of bias, imprecision	14/473 (3)	5/453 (1.1)	RR 0.43 (0.17-1.09)	30 per 1,000	17 fewer per 1,000 (from 25 fewer to 3 more)
			Me	ortality—initiatic	m (critical outc	Mortality—initiation (critical outcome; assessed with all-cause mortality)	h all-cause morta	lity)			
748 (5 studies ^e), 1-3 mo	Serious ^b	No serious inconsistency	No serious indirectness	Serious	Undetected	Undetected Low ^{h,e} due to risk of bias, imprecision	17/383 (4.4)	12/365 (3.3)	RR 0.73 (0.36-1.46)	44 per 1,000	12 fewer per 1,000 (from 28 fewer to 20 more)
			Thromb	oembolism—ma	iintenance (criti	Thromboembolism—maintenance (critical outcome; assessments variously defined)	ssments variously	/ defined)			
14,213 (7 studies ⁽⁾), 1-12 mo	Serious ^b	No serious inconsistency	No serious indirectness	No serious imprecision	Undetected	Undetected Moderate ^b due 122/7,091 (1.7) 109/7,122 (1.5) to risk of bias	122/7,091 (1.7)	109/7,122 (1.5)	RR 0.9 (0.7-1.17)	17 per 1,000	2 fewer per 1,000 (from 5 fewer to 3 more) (Continued)

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		δ	Quality Assessment					S	Summary of Findings	ngs	
_						_	Study Event Rates (%)	t Rates (%)		Anticipated 2	Anticipated Absolute Effects ^a
Participants (Studies), Follow-up	Risk of Bias	Inconsistency	Indirectness	Imprecision	Publication Bias	Overall Quality of Evidence	With Manual Dosing	With Dosing Decision Support	Relative Effect (95% CI)	Risk With Manual Dosing	Risk Difference With Dosing Decision Support (95% CI)
			Major	bleeding—main	tenance (critica	Major bleeding—maintenance (critical outcome; assessments variously defined)	sments variously c	lefined)			
14,035 (5 studies ^g), 4.8-12 mo	Serious ^b	No serious inconsistency	No serious indirectness	No serious imprecision	Undetected	Moderate ^b due to risk of bias	108/6,999 (1.5)	108/6,999 (1.5) 101/7,036 (1.4)	RR 0.92 (0.71-1.21)	15 per 1,000	1 fewer per 1,000 (from 4 fewer to 3 more)
			Mort	ality—maintena	nce (critical ou	Mortality—maintenance (critical outcome; assessed with all-cause mortality)	vith all-cause mor	tality)			
14,044 (5 studies ^h), 4.8-12 mo	Serious ^b	No serious inconsistency	No serious indirectness	No serious imprecision	Undetected	Moderate ^b due to risk of bias	70/6,973 (1)	75/7,071 (1.1)	RR 1.07 (0.78-1.48)	10 per 1,000	1 more per 1,000 (from 2 fewer to 5 more)
 Biography: Stais PD. Cardner MJ, Ranwart A et al. The effectiveness of wardiran dosing function a mongared with house staff dosing <i>J</i> Atthrophasty. 2007;32:2312.53. Ageno W. Turpie G. A machine domized compareston of a computer-based dosing program with a manual system to monitor onla mitcogalular therapy. <i>Thromb Res.</i> 1998;91:237-240. Mater 67, Moia M, Palareti G, et al. Effect of computer-aided management on the quality of treatments in anticogaluate patients: a prospective, randomized, multicenter trial of APBOXI (Automated PRogram for Oral Anticogaluat Treatment). <i>Haranatologica</i>. 2001;86:1060-1070. Marco F, Sedano C, Bernudez A, et al. A prospective, randomized, multicenter trial of APBOXI (Automated PRogram for Oral Anticogaluat Treatment). <i>Haranatologica</i>. 2001;86:1060-1070. Marco F, Sedano C, Bernudez A, et al. A prospective, randomized tareholdistical of a computer-assisted acenocoumarol dosage program. <i>Pathophysiol Haemost Thromb.</i> 2003;33:59:63. Mitra R, Marciello MA, Bania C, Janagar B, Burke DT: Effacey of computer-assisted acenocoumarol dosage program. <i>Pathophysiol Haemost Thromb.</i> 2003;35:59:63. Hold C, Bania C, Shangar B, Burke DT: Effacey of computer-assisted acenocoumarol dosage program. <i>Pathophysiol Haemost Thromb.</i> 2003;35:59:63. Hold C, Bania CR, Malanett C, Roia M, Fularett C, Sedan CR, Matcollan CR, Mala K, Bania CR, Malanett C, Mala K, Stata CR, Matcollan PK, Benia CO, Seyano S, Segano S, Satadardized J, Manda J, Sata S, Satadardized J, Manda J, Sata S, Satadardized J, Manda J, Satak BD, Futtrason DL, Landor Tardet J, Sota S, Satadardized J, Manda J, 1991;24:139-247. Palaetti C, Sata S, Satadardized J, Marco S, Satadardized J, Marco S, Satadardized J, Manda J, 1992;24:139-247. Palaetti C, Satadardized J, Satadardized J, Satadardized J, Satadardized J, Palaetti C, Rada M, Satadardized J, Satak BD, Harnado J, Satak BD, Satadardized J, 1992;63:19:2032;420. Palaetti C, Satadardized J, 1992;63:	PD, Cardner MJ 1 of a computer-J on the quality of Marco F, Sedam n C, Ahangar B, al. Multicenter randc anticoagulant c dized initial war dized initial war 091;21:319-324. 28:62-69. van de 28:62-69. van de 28:62-69. van de 28:62-69. van de 19:21:319-325. 28:62-69. van de 19:21:319-325. 1 compasses w e 1991, Kovacs 1 e 1991, Kovacs 1 unice 1996, Fitz	MJ, Ranawat A, et er-based dosing pru- of treatment in an lano C, Bermudez B, Burke DT. Eff B, Burke DT. Eff andomized study of idomized study of t control by a nur varfarin treatmen 4. Kovacs MJ, Cr den Bemt PM, Bé 33-208. Carter BL. Sypansion of abbre cluding patients, P s wide range of be s 1998, Landefeld itzmaurice 2000, N	at A, et al. The effectivene sing program with a manu- ent in anticoagulated patien mudez A, et al. A prospee DT. Efficacy of computer- ed study of computer-assiste- atudy of computer-assiste- y a nurse-practitioner usin- attment: evaluation of ini MJ, Cruickshank M, Well PM, Beinema M, van Roo rter BL, Taylor JW, Becke of abbreviations. Itents, health-care provide tients, health-care provide of benefit and harm. ndefeld 1992, Vadher 199 ndefeld 1992, Vadher 199 ndefeld 1992, Vadher 199	ness of warfarin c ual system to mo ients: a prospectiv bective controlled ective controlled ar-aided dosing o anticoagulant do ted oral anticoag sing a computer itital treatment anticoagulant do anticoagulant do anticoagulant do anticoagulant do sof (BMJ), van do 907 (BMJ), van do 977 (BMJ).	s of warfarin dosing from a nomogram co l system to monitor oral anticoagulant ther its: a prospective, randomized, multicentes ive controlled study of a computer-assist ided dosing of warfarin among patients i ticoagulant dosage. European Concerted. oral anticoagulant dosage vs medical st a computer decision-support system w ial treatment response and maintenance PS, et al. Randomized assessment of a n EN, et al. Initiation of oral anticoagulan A. Evaluation of three dosage-prediction a, and outcome adjudicators. (<i>BMJ</i>), van den Bemt 2002, White 1997. (<i>BMJ</i>). 008, Vadher 1997, (<i>BMJ</i>).	nomogram comparation of the comp	red with house strand with house strand with house strands $Thromb Res.$ 1999. I of APROAT (Automore and the comparisation on Anticoagulat losage. J Thromb hat by clinicians. Thromb hat by clinicians for the strand in orthoped for initial in the strand for the	uff dosing. J Arthri 3,91:237-240. Mar tomated PRograr age program. Path age program. Path 2014. Am J Phys l ion. Lancet. 1998; ion. Lancet. 1998; Clin Laboratory andomized trial, or initial oral ant bit and surgical p ihospital stabiliz i-hospital stabiliz	plasty. 2007;22:2 otti C, Moia M, I ofr Oral Anticoa neophysiol Haemos ded Rehabil. 200 352:1505-1509. P 3:935-943. Vadhe Haematol. 1997; and risk factors f and risk factors f icoagulation afte icoagulation afte atients: an algorit tion of warfarin tion of warfarin	13-218. Ageno V alareti G, et al gulant Treatme # Thromb. 2003 5;84:423-427. Po oller L, Keown I oller L, Keown I s: BD, Patterso 19:203-207. Doc or an excessive r venous throml hm compared w therapy. <i>Clin Ph</i>	 so f warfarin dosing from a nomogram compared with house staff dosing <i>J</i> Arthroplasty. 2007;22:213-218. Ageno W, Turpie G. A ran-l system to monitor oral anticoagulant therapy. <i>Thronb Res.</i> 1998;91:337-240. Manotti C, Moia M, Palareti G, et al. Effect of computer- its: a prospective, randomized multicenter trial of APROAT (Automated PRogram for Oral Anticoagulant Treatment). <i>Haemotologica</i>. tise controlled study of a computer-assisted acenocoumarol dosage program. <i>Pathophysiol Haemost Thronb.</i> 2003;33:59-63. Mitra R, uice addited dosing of warfarin among patients in a rehabilitation hospital. <i>Am J Phys Med Rehabil.</i> 2005;84:423-427. Poller L, Shiach CR, ticoagulant dosage. <i>European Concerted Action on Anticoagulation. Lancet.</i> 1998;53:1505-1509. Poller L, Keown M, Ibrahim S, et al. oral anticoagulant dosage. <i>European Concerted Action on Anticoagulation. Lancet.</i> 1998;53:21505-1509. Poller L, Keown M, Ibrahim S, et al. corral anticoagulant dosage. <i>European Concerted Action on Anticoagulation. Lancet.</i> 1998;53:21505-1509. Poller L, Keown M, Ibrahim S, et al. a computer decision-support system with that by clinicians. <i>Clin Laboratory Haemost J.</i> 1997;19:203-207. Doecke CJ, Cosh DG, ial treatment response and maintenance dose prediction by randomized trial, and risk factors for an excessive warfarin response. i PS, et al. Randomized assessment of a warfarin nomogram for initial oral anticoagulation after venous thromboembolic disease. n EN, et al. Initiation of oral anticoagulant therapy in orthopedic and surgical patients: an algorithm compared with routine dosing. A. Evaluation of three dosage-prediction methods for initial in-hospital stabilization of warfarin therapy. <i>Clin Pharm.</i> 1987;6:37-45. s, and outcome adjudicators. (<i>BMJ</i>), van den Bemt 2002, White 1997. (<i>BMJ</i>). 008, Vadher 1997, Vadher 1997. (<i>BMJ</i>).

Table S7—Continued

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*Claes 2005, Fitzmaurice 1996, Fitzmaurice 2000, Poller 2008, Vadher 1997.
^hClaes 2005, Fitzmaurice 1996, Fitzmaurice 2000, Poller 1993, Poller 2008.

			•	,	, 	-		0	
			Quality Assessment	ment				Summary of Findings	ngs
							Study Event Rates (%)	Rates $(\%)$	Anticipated Absolute Effects ^a
Participants (Studies), Follow-up	Risk of Bias	Risk of Bias Inconsistency	Indirectness	Imprecision	Publication Bias	Overall Quality of Evidence	With INR 2-3	Relative Effect With INR 3-5 (95% CI)	Risk Difference Risk With With INR 3-5 INR 2-3 (95% CI)
			Ma	ijor hemorrhage) (critical outcome	Major hemorrhage (critical outcome; assessed per 100 patient-y; various definitions)	tient-y; various def	finitions)	
76,646 (17 studies ^b)	Serious ^c	No serious inconsistency	No serious indirectness	No serious imprecision	Undetected	Low ^{ed} due to risk of bias and dose- response gradient		299/17,278 (1.7) RR 2.7 (1.8-3.9)	357/59,368 (0.6) 299/17,278 (1.7) RR 2.7 (1.8-3.9) 6 per 1,000 10 more per 1,000 (from 5 more to 17 more to 17 more)
			Thu	romboembolisn	1 (critical outcome	Thromboembolism (critical outcome; assessed per 100 patient-y; various definitions)	ttient-y; various de	finitions)	
835	Serious ^f	Seriouss	No serious		Undetected	Very low ^{f,g} due to	24/519 (4.6)	15/316(4.7) RR 0.9 (0.6-1.3)) Study population
(10 studies ^e)			indirectness	imprecision		risk of bias and inconsistency			46 per 1,000 5 fewer per 1,000 (from 18 fewer
									to 14 more)
									Moderate
									50 per 1,000 5 fewer per 1,000 (from 20 fewer
									to 15 more)
Bibliography: Oake N, Jennings A, and meta-analysis. $CMAJ$. 2008;17 ^a Time frame is in months to years.	Jake N, Jennir sis. <i>CMAJ</i> . 20 in months to y	Bibliography: Oake N, Jennings A, Forster AJ, Fergusson D, Doucette and meta-analysis. <i>CMAJ</i> . 2008;179(3):235-244. See Table S3 and S4 le "Time frame is in months to years.	Fergusson D, D 4. See Table S3 a		S, van Walraven C. Anticoagulation in egends for expansion of abbreviations.	oagulation intensity a bbreviations.	nd outcomes amor	ig patients prescribed oral anticoa	S, van Walraven C. Anticoagulation intensity and outcomes among patients prescribed oral anticoagulant therapy: a systematic review egends for expansion of abbreviations.
^b Six studies had	a randomize	^b Six studies had a randomized controlled trial design.	design.						
d It is biologicall	y plausible th	 The majority of studies (eight) were retrospective conorts. It is biologically plausible that with increased intensity there will be more bleeding. 	cuve conorts. intensity there v	vill be more ble	eding.				

Table S8–[Section 4.1.1] Evidence Profile: Optical Therapeutic INR Range—Higher Target vs 2 to 3

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sThromboembolic events were more frequent with INR 2 to 3 in two studies, less frequent in one study, and similar in one study.

«One study had a randomized controlled design. "Three of four studies had a retrospective cohort design.

			Quality Assessment	sment				Sum	Summary of Findings		
							Study Event Rates (%)	t Rates (%)		Anticipated /	Anticipated Absolute Effects ^a
Participants (Studies), Follow-up	Risk of Bias	Inconsistency	Indirectness	Imprecision	Publication Bias	Overall Quality of Evidence	With INR 2-3	With INR <2	Relative Effect (95% CI)	Risk With INR 2-3	Risk Difference With INR < 2 (95% CI)
			Majo	r hemorrhage (cr	itical outcome	; assessed per 100	Major hemorrhage (critical outcome; assessed per 100 patient-y; various definitions)	lefinitions)			
78,493	Serious ^b	Serious	No serious		Undetected	Very low ^{b,c} due	357/59,369 (0.6)	$357/59,369\ (0.6)\ 123/19,124\ (0.6)$	RR 1.1	Study I	Study population
(17 studies ^b)			indirectness	imprecision		to risk of bias, inconsistency			(0.7-1.7)	6 per 1,000	1 more per 1,000 (from 2 fewer to 4 more)
										Mo	Moderate
										23 per 1,000	2 more per 1,000 (from 7 fewer to 16 more)
				Thromboem	bolism (critica	d outcome; assesse	Thromboembolism (critical outcome; assessed per 100 patient-y)	(2)			
$827 (4 studies^d)$	Seriouse	No serious	Z	Z	Undetected	Moderate ^{e-g} due	24/520 (4.6)	$42/307\ (13.7)$	RR 3.5	Study I	Study population
		inconsistency	indirectness	imprecision		to risk of bias, large effect.			(2.8-4.4)	46 per 1,000	115 more per
						dose-response					1,000 (from
						gradient					83 more to 157 more)
										Mo	Moderate
										40 per 1,000	40 per 1,000 100 more per 1,000 (from
											72 more to
											136 more)
Bibliography: Oakes N. Jennings A. and meta-analysis. <i>CMAJ</i> . 2008,176 ^a Time frame is in months to years. ^b Eight of the studies were retrospe ^c Four studies showed higher risk o	Jakes N, Jenni sis. CMAJ. 200 in months to udies were ret nowed higher	Bibliography: Oakes N, Jennings A, Forster AJ, Fergusson D, Doucette and meta-analysis. <i>CMAJ</i> . 2008;179(3):235-244. See Table S3 and S4 le ^a Time frame is in months to years. ^b Eight of the studies were retrospective cohorts. ^e Four studies showed higher risk of bleeding with INR < 2.	Fergusson D, Do . See Table S3 an. th INR < 2.		S, van Walraven C. Anticoagulation ir sgends for expansion of abbreviations.	oagulation intensit bbreviations.	y and outcomes am	S, van Walraven C. Anticoagulation intensity and outcomes among patients prescribed oral anticoagulant therapy: a systematic review gends for expansion of abbreviations.	bed oral anticoagu	ulant therapy: a	systematic review
^d Only one study had a random [•] No explanation was provided.	y had a randoi 1 was providec	^d Only one study had a randomized controlled design. ^e No explanation was provided.	esign.								

Table S9–[Section 4.1.2] Evidence Profile: Optimal Therapeutic INR Range—Lower Target vs 2 to 3

^fAt least 2.8 times more frequent thromboembolism. sIt is biologically plausible with more thromboembolism at lower INR.

			Quality Assessment	lent					Summary of Findings	ings	
						-	Study Event Rates (%)	Rates (%)		Anticipated	Anticipated Absolute Effects
Participants (Studies), Follow-up	Risk of Bias	Inconsistency	Indirectness Imprecision	Imprecision	Publication Bias	Overall Quality of Evidence	With Moderate- Intensity VKA	With High- Intensity VKA	Relative Effect (95% CI)	Risk With Moderate- Intensity VKA	Risk Difference With High-Intensity VKA (95% CI)
			Ē	hromboembc	lism (importan	Thromboembolism (important outcome; assessed with objective confirmation)	with objective cont	1rmation)			
$220 (2 \text{ studies}^{a}),$	Serious ^b	No serious	Serious ^b	Serious	Undetected	Very low ^{b,c} due	$5/110 (4.5)^{a}$	11/110(10)	OR 2.33	Study	Study population ^d
ö.y		inconsistency				to risk of bias, indirectness, and imprecision			(0.32-6.66)	45 per 1,000 ³	54 more per 1,000 (from 8 fewer to 195 more)
											Low ^d
										$50 \text{ per } 1,000^{a}$	59 more per 1,000 (from 9 fewer to 210 more)
											$\operatorname{High}^{\operatorname{d}}$
										700 per 1,000 ^a	145 more per 1,000 (from 43 fewer to 240 more)
					Major ble	Major bleeding (important outcome ^e)	itcome ^e)				
$220 (2 \text{ studies}^{e}),$	Z	No serious	No serious	Serious ^f	Undetected	Moderate ^f due to	$7/110 \ (6.4)^{a}$	5/110(4.5)	OR 0.70	Study	Study population
3.y	risk of bias	inconsistency	indirectness			imprecision			(0.23-2.16)	$64 \text{ per } 1,000^{a}$	18 fewer per 1,000 (from 48 fewer to 64 more)
											Low
										$25 \text{ per } 1,000^{a}$	7 fewer per 1,000 (from 19 fewer to 27 more)
											High
										100 per 1,000 ^a	28 fewer per 1,000 (from 75 fewer to 94 more)
											(Continued)
			Î								

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Quality Assessment Summary of Findings	Study Event Rates (%) Anticipated Absolute Effects	Publication Overall Quality With Moderate- With High- Relative Effect Moderate- With High-Intensity Indirectness Imprecision Bias of Evidence Intensity VKA Intensity VKA (95% CI) Intensity VKA VKA (95% CI)	Mortality (assessed with all-cause mortality)	No serious Serious ^c Undetected Moderate ^c due to 2/110 (1.8) 3/110 (2.7) OR 1.51 18 per 1,000 9 more per 1,000 indirectness indirectness (0.3-7.72) (from 13 fewer to 10.7 more) 107 more)	Bibliography: Crowther MA, Ginsberg JS, Julian J, et al. A comparison of two intensities of warfarin for the prevention of recurrent thrombosis in patients with the antiphospholipid antibody syndrome. <i>N Engl J Med.</i> 2003;349:1133-1138. Finazzi G, Marchioli R, Brancaccio V, et al. A randomized clinical trial of high-intensity warfarin vs conventional antithrombotic therapy for the prevention of recurrent thrombosis in patients with the antiphospholipid syndrome (WAPS). J Thromb Haemost. 2005;3:848-853. Schulman S, Svenungsson E, Granqvist S. Anticardiolipin antibodies predict early recurrence of thrombosis in patients with the antiphospholipid syndrome (WAPS). J Thromb Haemost. 2005;3:848-853. Schulman S, Svenungsson E, Granqvist S. Anticardiolipin antibodies predict early recurrence of thrombosis in patients with venous thromboembolism following anticoagulant therapy. Duration of Anticoagulation Study Group. <i>Am J Med.</i> 1995;332(15):993-997. See Table S2 Khamashta MA, Cuadrado MJ, Mujic F, Taub NA, Hunt BJ, Hughes GR. The management of thrombosis in the antiphospholipid-antibody syndrome. <i>N Engl J Med.</i> 1995;332(15):993-997. See Table S2 legend for expansion of abbreviation.
uality Assessment			Mortality	Serious ^f less	, et al. A comparison of two intensities archioli R, Brancaccio V, et al. A rando bid syndrome (WAPS). <i>J Thromb Haem</i> th venous thromboembolism following Hunt BJ, Hughes GR. The managem onembolic arterial thrombosis received
Q		Inconsistency I		ency	Ginsberg JS, Julian J -1138. Finazzi G, M. th the antiphospholi th among patients wi J. Mujic F, Taub NA, viation. three patients with n
		Risk of Bias		No serious risk of bias	owther MA, 1003;349:1133 003;349:1133 in patients wii dism and deat Cuadrado MJ sion of abbrev Finazzi et al, t
	_	Participants (Studies), Follow-up		220 (2 studies), No serious No serious 3 y risk of inconsist bias	Bibliography: Crowther MA, Ginsberg JS, N Engl J Med. 2003;349:1133-1138. Finazz rent thrombosis in patients with the antiphe of thromboenbolism and death among pati Khamashta MA, Cuadrado MJ, Mujic F, Ti legend for expansion of abbreviation.

 $^{\rm d} {\rm Low}$ of 5% from Schulman et al. High of 70% from Khamashta et al. $^{\circ} {\rm The}$ types of major hemorrhage were not disclosed. 'The 95% CI includes both benefit and significant harm.

Table S10-Continued

			Quality Assessment	ment					Summary of Findings	dings	
						-	Study Even	Study Event Rates $(\%)$		Anticipated	Anticipated Absolute Effects ^a
Participants (Studies), Follow-up	Risk of Bias	Inconsistency	Indirectness	Imprecision	Publication Bias	Overall Quality of Evidence	With Abrupt Withdrawal	With Gradual Withdrawal	Relative Effect (95% CI)	Risk With Abrupt Withdrawal	Risk Difference With Gradual Withdrawal (95% CI)
				Thromboembol	lism (important ou	Thromboembolism (important outcome; assessed with imaging diagnostics)	h imaging diagne	ostics)			
217 (5 studies), 3 mo	Serious ^b	No serious inconsistency	No serious / indirectness	Serious	Undetected	Low ^{b.c} due to risk of bias, imprecision	14/111 (12.6) ^d	14/111 (12.6) ^d 13/106 (12.3)	OR 0.96 (0.42-2.18)	126 per 1,000 ^d	4 fewer per 1,000 (from 69 fewer to 113 more)
				Mortality (n	ot important outco	Mortality (not important outcome; assessed with all-cause mortality)	all-cause mortalit	ty)			
217 (5 studies), 1 mo	Serious ^b	No serious inconsistency	No serious / indirectness	Very serious ^e	Undetected	Very low ^{b,c} due to risk of bias, imprecision	1/111 (0.9)	0/106 (0)	OR 0 (0.01-5.6)	9 per 1,000	9 fewer per 1,000 (from 9 fewer to 39 more)d
					Major hemorrha	Major hemorrhage (important outcome)	ome)				
217 (5 studies), 1 mo	Serious ^b	No serious inconsistency	No serious / indirectness	Very serious ^c		Very low ^{b,c} due to risk of bias, immerision	1/111 (0.9)	0/106 (0)	OR 1 (0.1-5.6)	9 per 1,000	0 fewer per 1,000 (from 8 fewer to 39 more) ^d
Bibliography: Ascani A, de Groot MR, et al. Abru 2000;46(11-12):575-581. tion of treatment. $Am \int 1994;72(2):222-226.$ Tarc randomized study. $Br \int H$	scani A, et : : al. Abrupt w (75-581. Micl t. Am J Card 26. Tardy B, y. Br J Haem	Bibliography: Ascani A, et al. Withdrawal of warfarin after deep de Groot MR, et al. Abrupt vs gradual withdrawal of vitamin K antago 2000;46(11-12):575-581. Michaels L, Beamish RE. Relapses of thron tion of treatment. Am J Cardiol. 1967;20(5):670-673. Palareti G, et al. 1994;72(2):222-226. Tardy B, et al. Evolution of blood coagulation and randomized study. Br J Haematol. 1997;96(1)174-178. See Table S2 le "Time frame is in weeks.	warfarin after val of vitamin K RE. Relapses of 0-673. Palareti (blood coagulatic 74-178. See Table	deep vein thro antagonist treatn f thromboembol G, et al. Activati on and fibrinolys $^{\rm b}$ S2 legend for ϵ	Bibliography: Ascani A, et al. Withdrawal of warfarin after deep vein thrombosis: effects of a low de Groot MR, et al. Abrupt vs gradual withdrawal of vitamin K antagonist treatment in patients with veno 2000;46(11-12):575-581. Michaels L, Beamish RE. Relapses of thromboembolic disease after discontinution of treatment. Am J Cardiol. 1967;20(5):670-673. Palareti G, et al. Activation of blood coagulation is 1994;72(2):222-226. Tardy B, et al. Evolution of blood coagulation and fibrinolysis parameters after abrupt randomized study. Br J Haematol. 1997;96(1)174-178. See Table S2 legend for expansion of abbreviation.	Bibliography: Ascani A, et al. Withdrawal of warfarin after deep vein thrombosis: effects of a low fixed dose on rebound thrombin generation. <i>Blood Coagul Fibrinolysis</i> . 1999;10(5):291-295. de Groot MR, et al. Abrupt vs gradual withdrawal of vitamin K antagonist treatment in patients with venous thromboembolic disease: assessment of hypercoagulability and clinical outcome. <i>Clin Laboratory</i> . 2000;46(11-12):575-581. Michaels L, Beamish RE. Relapses of thromboembolic disease after discontinued anticoagulant therapy. A comparison of the incidence after abrupt and after gradual termination of treatment. <i>Am J Cardiol</i> . 1967;20(5):670-673. Palareti G, et al. Activation of blood coagulation after abrupt or stepwise withdrawal of oral anticoagulants-a prospective study. <i>Thromb Haemost</i> . 1994;72(2):222-226. Tardy B, et al. Evolution of blood coagulation after abrupt vs gradual withdrawal of acenocoumarol in patients with venous thromboembolism: a double-blind randomized study. <i>Br J Haematol</i> . 1997;96(1)174-178. See Table S2 legend for expansion of abbreviation.	on rebound th embolic disease: <i>i</i> ulant therapy. A or stepwise with <i>i</i> thdrawal of ace	rombin general assessment of hy comparison of ndrawal of oral nocoumarol in <u>F</u>	tion. Blood Coagy percoagulability ar the incidence afte anticoagulants-a F attients with venou	<i>ul Fibrinolysis.</i>] Id clinical outcom er abrupt and aft prospective study us thromboembol	
^b Unclear whethe de Groot, and As	er allocation scani; not con	^b Unclear whether allocation was adequate in Tardy et al, de Groot et de Groot, and Ascani; not concealed in Michaels and Beamish. Clinici	ardy et al, de G ls and Beamish.	root et al, and A Clinicians and p	scani et al. In Mic atients were not bl	^b Unclear whether allocation was adequate in Tardy et al, de Groot et al, and Ascani et al. In Michaels and Beamish, it was according to year of birth. Unclear whether allocation was concealed in Tardy, de Groot, and Ascani; not concealed in Michaels and Beamish. Clinicians and patients were not blinded in de Groot, Michaels, Palareti et al, or Ascani.	, it was according Michaels, Palare	g to year of birt ti et al, or Ascar	h. Unclear whethé ai.	er allocation was	concealed in Tardy,

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^dThere is no better source than these trials, so low or high estimates are not provided.

° Very small patient groups and few events.

Inconsistency Indirectness Imprecision No serious No serious Serious ^d inconsistency indirectness Serious inconsistency indirectness Very seriou inconsistency indirectness Very seriou f. Eisenberg P, et al; Antithrombotic Therapy C ospitalized patients with active thromboemboli arin nomogram in patients with active thromboemboli arin dosing nomogram compared with a "stand	Quality Assessment			Summary of Findings	ndings	
Participants (Studies), Follow-up Risk of Bias Inconsistency Indirectness Imprecision Publ Follow-up Risk of Bias Inconsistency Indirectness Imprecision Publ 2-90 d ^b Serious ^e No serious No serious Serious ^d Ur 2-90 d ^b Inconsistency indirectness <u>Major he</u> 2-90 d ^b Inconsistency indirectness <u>Major he</u> 179 (2 studies ^e), Serious ^e No serious No serious Very serious ^d Ur 1 wk Inconsistency indirectness Bibliography: Becker RC, Ball SP, Eisenberg P, et al: Antithrombotic Therapy Consortiu care cognidation monitoring in hospitalized patients with active thrombotic Therapy Consortiu care cognidation monitoring in hospitalized patients with active thrombotic Therapy Consortiu care cognidation monitoring in hospitalized patients with active thrombotic Therapy Consortiu care cognidation monitoring in hospitalized patients with active thrombotic Therapy Consortiu care coronary syndrome Srinivas S. The weight-based heparin dosing nonogram to patients with active thrombotic Therapy Consortiu		_	Study Event Rates (%)		Anticipated	Anticipated Absolute Effects ^a
Thron 292 (3 studies), Serious ^e No serious No serious Serious ^d Ur 2-90 d ^b inconsistency indirectness 2-90 d ^b Major he 779 (2 studies ^c), Serious ^e No serious No serious Very serious ^d Ur 1 wk inconsistency indirectness 8ibliography: Becker RC, Ball SP, Eisenberg P, et al: Antithrombotic Therapy Consortiu care cogulation monitoring in hospitalized patients with active thromboembolic disease tion with a weight-adjusted heparin homogram in patients with active thromboembolic disease tion with a weight-based heparin dosing nonogram compared with a "standard care"	Inconsistency Indirectness	Overall Quality Bias of Evidence	With Fixed Based Initial Dose Nonogram	ht- Relative Effect m (95% CI)	Risk With Fixed Initial Dose	Risk Difference With Weight- Based Nomogram (95% CI)
 292 (3 studies), Serious^e No serious No serious Serious^d Ur 2-90 d^b inconsistency indirectness 2-90 d^b inconsistency indirectness Major he 179 (2 studies^s), Serious^e No serious No serious Very serious^d Ur 1 uk Bibliography: Becker RC, Ball SP, Eisenberg P, et al; Antithrombotic Therapy Consortiu care coagulation monitoring in hospitalized patients with active thromboembolic disease tion with a weight-based heparin nonogram in patients with active thromboembolic disease tion weight-based heparin dosing nonogram compared with a "standard care" 	Thromboen	Thromboembolism (critical outcome)				
Major he 179 (2 studies ^g), Serious ^e No serious No serious Very serious ⁴ Ur 1 wk inconsistency indirectness Bibliography: Becker RC, Ball SP, Eisenberg P, et al; Antithrombotic Therapy Consortiucare coagnition monitoring in hospitalized patients with active thromboembolic disease tion with a weight-adjusted heparin nonogram in patients with acute coronary syndrome Srinivas S. The weight-based heparin dosing nonogram compared with a "standard care"	No serious No serious inconsistency indirectness	ed Low ^{ed} due to risk of bias and imprecision	8/140 (5.7)° 2/152 (1.3)	3) OR 0.22 (0.02-1.13) ^f	$57 \text{ per } 1,000^{\circ}$	57 per 1,000° 44 fewer per 1,000 (from 56 fewer to 7 more)
179 (2 studies ⁵), Serious ^e No serious No serious Very serious ^d Ur 1 wk inconsistency indirectness Very serious ^d Ur Bibliography: Becker RC, Ball SP, Eisenberg P, et al: Antithrombotic Therapy Consortiu care coagulation monitoring in hospitalized patients with active thromboembolic disease tion with a weight-adjusted heparin nomogram in patients with acute coronary syndrome Srinivas S. The weight-based heparin dosing nomogram compared with a "standard care"	Major hemorrh	Major hemorrhage (not important outcome)	ne)			
Bibliography: Becker RC, Ball SP, Eisenberg P, et al. Antithrombotic Therapy Consortiu care coagulation monitoring in hospitalized patients with active thromboembolic disease tion with a weight-adjusted heparin nonnogram in patients with acute coronary syndrome Srinivas S. The weight-based heparin dosing nonnogram compared with a "standard care" are able to according to the set of the standard care".	No serious No serious inconsistency indirectness	ed Very low ^{ed} due to 1/88 (1.1) risk of bias and imprecision	1/88 (1.1) 0/91 (0)	OR 0 (0-37.7) ^t 11 per 1,000		11 fewer per 1,000 (from 11 fewer to 291 more)
* Time traine is in tays to weeks. • Only Raschke et al collected data over a 3-mo period. • CThe studies did not use blinding. Primary outcome was a surrogate marker: time to reach therapeutic or stable therapeutic activated partial thromboplastin time. • Along of the outcome was a surrogate marker: time to reach therapeutic or stable therapeutic activated partial thromboplastin time.	RC, Ball SP, Eisenberg P, et al; Antithrombotic Therapy Consortium Investigators A randomized, m toring in hospitalized patients with active thromboembolic disease. Am Heart J. 1999:137(1):59-71. usted heparin nomogram in patients with acute coronary syndromes: a randomized trial. <i>J Thromb T</i> -based heparin dosing nomogram compared with a "standard care" nomogram. A randomized contro it to weeks. Dilected data over a 3-mo period.	stigators A randomized, i teart J. 1999:137(1):59-7. adomized trial. J Thromb gram. A randomized cont gram. A randomized cont apeutic or stable theraper	nulticenter trial of weight- Hassan WM, Flaker GC <i>Thrombolysis</i> . 1995:2(3):2 rolled trial. <i>Ann Intern Me</i> rtic activated partial throm	adjusted intravenou: Feutz C, Petroski C 45-249. Raschke RA d. 1993;119(9):874-6 boplastin time.	s heparin dose ti Fr, Smith, D. In Freilly BM, Gu S81. UFH = unl	tration and point-of- proved anticoagula - idry JR, Fontana JR, fractionated heparin.

Table S12—[Section 6.1] Evidence Profile: UFH Weight-Based Nomogram Comnared With Fixed Initial Dose for Patients With Thromboembolic Disease

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sBecker et al reported 2% bleeding without specifying allocation group or type of bleeding.

^eTwo of the eight events occurred after discontinuation of warfarin.

^fFisher exact test.

		-	Quality Assessment	nent					Summary of Findings	ngs	
						-	Study Event Rates (%)	t Rates (%)		Anticipated A	Anticipated Absolute Effects ^a
Participants (Studies), Follow-up	Risk of Bias	Inconsistency	Indirectness	Imprecision	Inconsistency Indirectness Imprecision Publication Bias	Overall Quality of Evidence	With Weight- Adjusted Nonmonitored LMWH SC	With Weight- Adjusted Nonmonitored UFH SC	Relative Effect (95% CI)	Risk With Weight- Adjusted Nonmonitored LMWH SC	Risk Difference With Weight- Adjusted Non- monitored UFH SC (95% CI)
			Recurrent VTI	VTE (critical	outcome; assessec	E (critical outcome; assessed objectively with same method as for index event)	same method as f	or index event)			
697 (1 study), 1 3 mo	No serious risk No serious of bias inconsist	No serious inconsistency	Serious ^b	Serious	Undetected	Low ^{b,e} due to indirect- ness and imprecision	12/352 (3.4)	13/345 (3.8)	OR 1.11 (0.49-2.52)	34 per 1,000	4 more per 1,000 (from 17 fewer to 48 more)
				Major ble	eding (critical out	Major bleeding (critical outcome; assessed with ISTH criteria)	ith ISTH criteria)				
697 (1 study), 1 3 mo	No serious risk No serious of bias inconsist	No serious inconsistency	Serious ^b	Serious	Undetected	Low ^{b,c} due to indirect- ness and imprecision	12/352 (3.4)	6/345 (1.7)	OR 0.5 (0.17-1.34)	34 per 1,000	17 fewer per 1,000 (from 28 fewer to 11 more)
					Mortality (not	Mortality (not important outcome)	me)				
697 (1 study), 1 3 mo	No serious risk No serious of bias inconsist	No serious inconsistency	Serious ^b	Serious	Undetected	Low ^{b,c} due to indirect- ness and imprecision	22/352 (6.3)	18/345 (5.2)	OR 0.83 (0.43-1.57)	62 per 1,000	10 fewer per 1,000 (from 35 fewer to 32 more)

^bThe comparison should actually be vs fixed-dose UFH SC with monitoring, but weight-adjusted UFH SC has only been compared directly with weight-adjusted LMWH. Thus, the comparison is indirect.

^e Due to premature discontinuation, the study was not powered to demonstrate equivalence.

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		ð	Quality Assessment	it				<u>s</u>	Summary of Findings	ings	
_						_	Study Eve.	Study Event Rates (%)		Anticipated A	Anticipated Absolute Effects ^b
Participants (Studies), Follow-up	Risk of Bias	Inconsistency	Indirectness	Imprecision	Imprecision Publication Bias	Overall Quality of Evidence	With Only Holding VKA	With Only Holding VKA With Vitamin K	Relative Effect (95% CI)	Relative Effect Risk With Only (95% CI) Holding VKA	Risk Difference With Vitamin K (95% CI)
					Major bleedin	Major bleeding (critical outcome)					
$\frac{923}{1-3} (4 \text{ studies}^{c})$	923 (4 studies ^e), No serious risk No serious 1-3 mo ^d of bias ^e inconsist	: No serious inconsistency	No serious / indirectness	Serious ^f	Undetected	Moderate ^{e,f} due to imprecision	4/471 (0.8)	10/452 (2.2)	OR 2.6 (0.8-9.8)	8 per 1,000	13 more per 1,000 (from 2 fewer to 69 more)
					Thromboembolis	Thromboembolism (important outcome)	ome)				
864 (3 studies ^g) 1-3 mo ^d	864 (3 studies ^s), No serious risk No serious 1-3 mo ^d of bias ^e inconsist	: No serious inconsistency	No serious indirectness	Serious ^f	Undetected	Moderate ^{e,f} due to imprecision	4/441 (0.91)	5/423~(1.2)	OR 1.3 (0.3-6.6)	9 per 1,000	3 more per 1,000 (from 6 fewer to 48 more)
				Mortality (not	important outcor	Mortality (not important outcome; assessed with all-cause mortality)	Il-cause mortali	ty)			
863 (3 studies ^g) 1-3 mo ^d	863 (3 studies ^g), No serious risk No serious 1-3 mo ^d of bias ^e inconsist	No serious inconsistency	No serious / indirectness	Serious ^f	Undetected	Moderate ^{e,f} due to imprecision	13/441 (2.9)	16/422 (3.8)	OR 1.3 (0.6-2.9)	29 per 1,000	9 more per 1,000 (from 12 fewer to 51 more)
Bibliography: Crowther Crowther MA, Julian J, Low dose oral vitamin A randomized trial con 730-742. See Table S2 a INR 6,0-12.0 in Agenc ^b Time frame is in days. ^c None of the studies sp ^d Follow-up was 3 m oi e Two small studies Ao oi	Bibliography: Crowther MA, Ageno Crowther MA, Julian J, McCarty D, Low dose oral vitamin K to reversa A randomized trial comparing 1 m 730-742. See Table S2 and S4 for er a INR 6.0-12.0 in Ageno et al 2005. ^b Time frame is in days. ^c None of the studies specified whet ^d Follow-up was 3 mo in both studie ^c Two small studies Ageno 2002 and	Bibliography: Crowther MA, Ageno W, Garcia D, et al. Oral vitamin K v Crowther MA, Julian J, McCarty D, et al. Treatment of warfarin-associa Low dose oral vitamin K to reverse acenocoumarol-induced coagulop A randomized trial comparing 1 mg of oral vitamin K with no treatme 730-742. See Table S2 and S4 for expansion of abbreviations. ^a INR 6.0-12.0 in Ageno et al 2005. ^b Time frame is in days. ^c Tonov-up was 3 mo in both studies by Crowther et al. ^c Tonov-up was 3 mo in both studies by Crowther et al.	et al. Oral vitamin tt of warfarin-assc ol-induced coagu in K with no trea previations. ng events were fa et al.	K vs placebo to co cciated coagulopath lopathy: a random tment in the mana trant in the mana tral or intracranial.	o correct excessiv pathy with oral vi lomized controlle- nanagement of w nial.	Bibliography: Crowther MA, Ageno W, Garcia D, et al. Oral vitamin K vs placebo to correct excessive anticoagulation in patients receiving warfarin: a randomized trial. <i>Ann Intern Med.</i> 2009;150(5):293-300. Crowther MA, Julian J, McCarty D, et al. Treatment of warfarin-associated coagulopathy with oral vitamin K: a randomized controlled trial. <i>Lancet.</i> 2000;356:1551-1553. Ageno W, Crowther M, Steidl L, et al. Low dose oral vitamin K to reverse acenocoumarol-induced coagulopathy: a randomized controlled trial. <i>Lancet.</i> 2000;356:1551-1553. Ageno W, Crowther M, Steidl L, et al. A randomized trial comparing 1 mg of oral vitamin K with no treatment in the management of warfarin-associated coagulopathy in patients with mechanical heart valves. <i>J Am Coll Cardiol.</i> 2005;46(4): a NR 6.0-12.0 in Ageno et al 2005. INR 6.0-12.0 in Ageno et al 2005. In the frame is in days. ^e Fonder S and she for expansion of abbreviations. ^b Time frame is a low so in both studies by Crowther et al.	ı patients receiv zed controlled t <i>Haemostasis.</i> 20(soagulopathy in	ing warfarin: a ram ial. <i>Lancet.</i> 2000;5 2;88:48-51. Ageno patients with meo patients with meo	domized trial. <i>An</i> 356:1551-1553. A 356:1551-1553. A 356:1551-1553. A shanical heart val shanical heart val	n Intern Med. 200 geno W, Crowthe ilingardi M, Gall ves. J Am Coll C ves. J Am Coll C	9,150(5):293-300. : M, Steidl L, et al. i M, Crowther M. <i>ardiol.</i> 2005;46(4):

Table S14—[Section 9.1] Evidence Profile: Vitamin K vs Only Withholding VKA for Patients on Warfarin With Elevated INR (4.5-10) Without Evidence of Bleeding^a

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¹Wide CIs encompassing both benefit and significant harm. ⁸Ageno et al 2005 did not report thromboembolism, and Ageno et al 2002 did not report deaths.

Evidence-Based Management of Anticoagulant Therapy : Antithrombotic Therapy and Prevention of Thrombosis, 9th ed: American College of Chest Physicians Evidence-Based Clinical Practice Guidelines

Anne Holbrook, Sam Schulman, Daniel M. Witt, Per Olav Vandvik, Jason Fish, Michael J. Kovacs, Peter J. Svensson, David L. Veenstra, Mark Crowther and Gordon H. Guyatt *Chest* 2012;141; e152S-e184S DOI 10.1378/chest.11-2295

This information is current as of March 4, 2012

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