Critical Review Form Clinical Practice Guidelines

Clinical Policy: Critical Issues in the Evaluation and management of Adult Patients Presenting with Suspected Pulmonary Embolism, *Ann Emerg Med* 2003; 41:257-270

<u>Objectives:</u> To revise the 1995 ACEP "Clinical Policy for the Initial Approach to Adults Presenting with a Chief Complaint of Chest Pain, with No History of Trauma" as it relates to the initial approach to patients with signs and symptoms of PE.

Methods: A MEDLINE search was conducted from Jan 1995 through April 2001 using the keyword "pulmonary embolism". In addition, practice guidelines were reviewed from the American Heart Association, British Thoracic Society, American College of Chest Physicians, American Thoracic Society, and European Heart Association (the last of which was updated in 2008). After the PE subcommittee reviewed the available evidence four topics were selected for this guideline:

- 1) Diagnostic utility of D-dimer for PE;
- 2) Diagnostic utility of ventilation-perfusion scanning for PE;
- 3) Diagnostic utility of spiral computed tomography for PE;
- 4) Therapeutic implications for fibrinolytic agents in PE.

Each manuscript was graded for quality by at least two sub-committee members based on study design. Design 1 represents the strongest evidence and design 3 the weakest evidence

Literature classification schema.*

Design/ Class	Therapy [†]	Diagnosis [‡]	Prognosis [§]
1	Randomized, controlled trial or meta-	Prospective cohort using a criterion	Population prospective cohort
	analyses of randomized trials	standard	r
2	Nonrandomized trial	Retrospective observational	Retrospective cohort Case control
3	Case series	Case series	Case series
	Case report	Case report	Case report
	Other (eg, consensus, review)	Other (eg, consensus, review)	Other (eg, consensus, review)

^{*}Some designs (eg, surveys) will not fit this schema and should be assessed individually.

[§]Objective is to predict outcome including mortality and morbidity.



Objective is to measure therapeutic efficacy comparing ≥ 2 interventions.

^{*}Objective is to determine the sensitivity and specificity of diagnostic tests.

Next, subcommittee members graded the evidence on 6 dimensions felt to be most important for guideline development: blinding of outcome assessment, blinded/randomized allocation, direct or indirect outcome measures, bias (selection, detection, transfer), external validity and sample size. The recommendations generated were then classified as Level A, B, or C.

Approach to downgrading strength of evidence.

	Design/Class			
Downgrading	1	2	3	
None	I	II	III	
None 1 level	II	III	X	
2 levels	III	X	X	
Fatally flawed	X	X	X	

Level A – high degree of clinical certainty based on Class 1 strength of evidence or preponderance of Class II studies.

Level B – moderate clinical certainty based on Class II studies or strong consensus of Class III studies.

Level C – preliminary or inconclusive evidence or panel consensus.

This guideline excluded studies of pregnant patients.

	Guide	Comments	
I.	Are the Recommendations Valid?	Answer questions IA-D below	
A.	Did the recommendations consider all	No. This guideline development	
	relevant patient groups, management	process excluded pregnant patients	
	options, and possible outcomes?	(and asymptomatic patients) (p.260)	
В.	If necessary, was an explicit, systematic, and	Uncertain. The process used to select	
	reliable process used to tap expert opinion?	ACEP Guideline Committee or PE	
		Subcommittee members was not	
	You should look for a clear description of how	explained. Also, no conflicts of	
	the panel was assembled along with the	interest are explained. The authors	
	members' specialties and any organizations	did seek guidance from pertinent	
	they are representing.	shareholders such as ATS, BTS,	
		AHA, etc.	

C. Is there an explicit, systematic specification of values or preferences?

Panelists' ratings presumably reflect the riskbenefit trade-offs of specific interventions, but whether other physicians or patients themselves would make the same decisions remains uncertain. Whether given options are value or preference related should be clearly stated in the guideline. No. The authors do not provide any description of clinician risk aversion variability, policy-maker, medicolegal or cost implications, or mechanisms to incorporate unique patient preferences into the diagnostic and therapeutic decision making for PE.

D. If the quality of the evidence used in originally framing the criteria was weak, have the criteria themselves been correlated with patient outcomes?

When the studies utilized to produce guidelines are less than randomized-controlled trials, conclusions can be strengthened by noting how outcomes can be correlated with adherence to the guidelines.

In some instances the guideline authors provide outcomes-based trial evidence to support recommendations. Examples:

- 1) "In a retrospective study,
 Rajendran and Jacobson
 investigated 536 patients with
 a low probability V/Q scan
 and found no evidence of PE
 on 6-month follow-up" (p.
 263)
- 2) "In the largest study to date, <u>Swensen et al</u> retrospectively studied 1512 consecutive patients undergoing CT angiography for suspected PE for 3-month outcome. The incidence of DVT or PE on follow-up was 0.5% and fatal PE 0.3% in the 1,010 patients with negative findings on a spiral CT." (p. 264)

II.	What are the recommendations?	Answer questions II A-B below
A.	Are practical, clinically important	Yes, within the limitations of the pre-
	recommendations made?	2001 search strategy being applied in
		2011.
		Question 1: Can a negative D-dimer
		exclude PE?
		Level A Recommendations: None
		Level B Recommendations: In
		patients with a low pretest probability
		of PE, use the following tests to
		exclude PE:
		1) A negative quantitative D-
		dimer assay (turbidimetric or
		ELISA).
		2) A negative whole blood cell
		qualitative D-dimer assay in
		conjunction with wells score of 2 or less.
		Level C Recommendations: In
		patients with a low pretest probability
		of PE, negative findings on a whole
		blood D-dimer assay (when not used
		with well's scoring system) or
		immunofiltration D-dimer assay can
		be used to exclude PE.
		Guideline authors offer the following
		evidence to support those
		recommendations.
		The qualitative whole blood
		assay requires 5-minutes to
		administer and when used with
		Well' criteria has a NPV of
		<u>99.5%</u> .
		• Two rapid (~ 2 hours)
		quantitative D-dimer tests are
		available (ELISA,
		turbidimetric) with <u>negative</u> LR 0.07 at cutoff 0.05 ng/mL.
		LIX 0.07 at cutoff 0.03 fig/file.
		Question 2: Can V/Q scan alone or
		in combination with venous Doppler
	W. 1 II	or D-dimer exclude PE?

Level A Recommendations: In patients with a low-to-moderate pretest probability of PE, a normal perfusion scan reliably excludes clinically significant PE.

Level B Recommendations: In patients with a low-to-moderate pretest probability of PE and a non-diagnostic V/Q scan, the following tests can be used instead of a pulmonary arteriogram to exclude PE.

- 1) A negative quantitative D-dimer assay.
- 2) A negative whole blood qualitative D-dimer in conjunction with a Well's score <4.
- 3) For low-probability patients, a single negative bilateral venous ultrasonographic scan.
- 4) For moderate probability patients, serial (repeat at day 3-7) bilateral venous ultrasonographic scan.

Level C Recommendations: In patients with a low-to-moderate pretest probability of PE and a nondiagnostic V/Q scan, use a negative whole blood D-dimer assay (When not used with Well's score) or immunofiltration D-dimer assay to exclude PE.

Guideline authors offer the following evidence to support these recommendations.

- In the initial <u>PIOPED</u> analysis, PE rates for V/Q results of high, intermediate, low, and normal were 87%, 30%, 14%, and 4% with LR⁺ for "high" 18.3 and LR⁻ for normal 0.1.
- Unfortunately, 60% of PIOPED patients were non-diagnostic (intermediate or low probability V/Q scans).

- In patients with low probability V/Q scans, PE rates for low, intermediate, or high clinical probability patients were 4%, 16%, and 40% respectively.
- In one retrospective study, Rajendran and Jacobson investigated 536 patients with low probability V/Q scans and found no patient with evidence of PE at 6-month follow-up.
- <u>Sensitivity of LE Duplex</u> for patients with a nondiagnostic V/Q is 50%.

Question 3: Can spiral CT replace V/Q scanning in the diagnostic evaluation of PE?

Level A Recommendations: None Level B Recommendations: Thin collimation spiral CT scan of the thorax with 1- or 2-mm image reconstruction may be used as an alternative to V/Q in evaluating patients with suspected PE. Level C Recommendations: Spiral CT scan of the thorax with delayed CT venography may be used for increased detection of patients with significant thromboembolic disease.

Guideline authors offer the following evidence to support these recommendations:

- Meta-analysis of 9 prospective spiral CT trials indicates pooled sensitivity 77%, specificity 89% with increased sensitivity (95%), for segmental (or larger) PE's and lower for subsegmental PE (under 75%).
- <u>Goodman et al</u> evaluated 198 patients with negative CT and

		188 with a normal or low prob V/Q and 3% in those with low probability V/Q. • Swenson et al retrospectively evaluated 1512 consecutive patients with CT angiography for suspected PE with an incidence of PE or DVT 0.5% and fatal PE 0.3% in the 1010 patients with negative findings on spiral CT. Question 4: What are the indications
		for fibrinolytic therapy in patients with PE? See the <u>July 2010 Journal</u> Club.
B.	How strong are the recommendations?	See the Level A, B or C of
		recommendations above.
III.	How Can I Apply the Criteria to Patient Care?	
A.	Are the criteria relevant to your practice setting?	Yes. However, "recommendations offered in this policy are not intended to represent the only diagnostic and
	Medical practice is shaped by an amalgam of evidence, values and circumstances; clinicians should consider their local medical culture and practice circumstances before importing a particular set of audit criteria.	management options that the emergency physician should consider. ACEP clearly recognizes the importance of the individual clinician's judgment. Rather, they define for the clinician those strategies for which medical literature exists to provide strong support for their utility in answering the crucial questions addressed in this policy." (p. 258)
В.	Have the criteria been field-tested for feasibility of use in diverse settings, include settings similar to yours?	No, these recommendations (and guideline development methods) have not been field tested for feasibility in heterogeneous settings.

Limitations

- 1) Outdated guideline update pending which should incorporate concern below.
- 2) Failure to address frequent diagnostic and therapeutic issues such as the work-up threshold for symptomatic patients with known PE, indications for vena cava filter placement, and emergency physician Doppler ultrasound for DVT as part of PE work-up when V/Q is non-diagnostic.
- 3) Exclusion of pregnant patients.
- 4) Pre-dated <u>PERC criteria</u>.

Bottom Line

Well's criteria (below) or similar validated decision aids like the Wicki criteria or Kline criteria should be used prior to test-ordering to risk stratify ED patients with suspected PE. Once PE probability stratification is deduced, use of quantitative D-dimer (ELISA or turbidimetric) can exclude patients with low pre-test probability. In low to moderate pretest probability patients, a normal V/Q scan excludes clinically significant PE. The 3-month risk of DVT or PE in patients with a negative spiral CT is 0.5%.

Well's Criteria

Risk Factor	Points
Suspected DVT	3
Alternative Diagnosis Less Likely than PE	3
Heart Rate > 100	1.5
Immobilization or surgery previous 4 weeks	1.5
Previous DVT or PE	1.5
Hemoptysis	1
Malignancy (last 6 months or palliative)	1

Score	Mean Probability PE (%)	% with this Score	Risk Interpretation
0-2 points	3.6	40	Low
3-6 points	20.5	53	Moderate
>6 points	66.7	7	High