

Hospitalist-Operated Compression Ultrasonography: a Point-of-Care Ultrasound Study (HOCUS-POCUS)

Ernest A. Fischer, MD¹, Benjamin Kinnear, MD^{2,3}, Dana Sall, MD²,
Matthew Kelleher, MD^{2,3}, Otto Sanchez, MD PhD⁴, Benji Mathews, MD^{4,5},
Daniel Schnobrich, MD^{4,6}, and Andrew P. J. Olson, MD^{4,6}



¹Department of Medicine, MedStar Georgetown University Hospital, Washington, DC, USA; ²Department of Medicine, University of Cincinnati, Cincinnati, OH, USA; ³Department of Pediatrics, University of Cincinnati, Cincinnati, OH, USA; ⁴Department of Internal Medicine, University of Minnesota Medical School, Minneapolis, MN, USA; ⁵HealthPartners, Minneapolis/St. Paul, MN, USA; ⁶Department of Pediatrics, University of Minnesota Medical School, Minneapolis, MN, USA.

BACKGROUND: Venous thromboembolism includes deep vein thrombosis (DVT) and pulmonary embolism. Compression ultrasonography is the most common way to evaluate DVT and is typically performed by sonographers and interpreted by radiologists. Yet there is evidence that ultrasound examinations can be safely and accurately performed by clinicians at the bedside.

OBJECTIVE: To measure the operating characteristics of hospital medicine providers performing point-of-care ultrasound (POCUS) for evaluation of DVT.

DESIGN: This is a prospective cohort study enrolling a convenience sample of patients. Hospital medicine providers performed POCUS for DVT and the results were compared with the corresponding formal vascular study (FVS) interpreted by radiologists.

PARTICIPANTS: Hospitalized non-ICU patients at four tertiary care hospitals for whom a DVT ultrasound was ordered.

MAIN MEASURES: The primary outcomes were the sensitivity, specificity, and predictive values of the POCUS compression ultrasound compared with a FVS. The secondary outcome was the elapsed time between order and the POCUS study compared with the time the FVS was ordered to when the formal radiology report was finalized.

KEY RESULTS: One hundred twenty-five limbs from 73 patients were scanned. The prevalence of DVT was 6.4% (8/125). The sensitivity of POCUS for DVT was 100% (95% CI 74–100%) and specificity was 95.8% (95% CI 91–98%) with a positive predictive value of 61.5% (95% CI 35–84%) and a negative predictive value of 100% (95% CI 98–100%). The median time from order to POCUS completion was 5.8 h versus 11.5 h median time from order until the radiology report was finalized ($p=0.001$).

CONCLUSION: Hospital medicine providers can perform compression-only POCUS for DVT on inpatients with accuracy similar to other specialties and settings, with results available sooner than radiology. The observed prevalence of DVT was lower than expected. POCUS may be reliable in excluding DVT but further study is required to determine how to incorporate a positive POCUS DVT result into clinical practice.

KEY WORDS: ultrasonography; thromboembolism; hospitalists.

J Gen Intern Med 34(10):2062–7
DOI: 10.1007/s11606-019-05120-5
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INTRODUCTION

Venous thromboembolism (VTE), including deep venous thrombosis (DVT) and pulmonary embolism (PE), is a major public health problem. Inpatient rates of incident and recurrent VTE range from 142 to 300+ per 100,000 person years versus 8 per 100,000 in community-dwelling patients.^{1, 2} VTE increases morbidity, mortality, and cost in hospitalized patients,³ and incidence of VTE has not improved with prophylaxis.⁴

Prompt recognition of DVT is paramount to reduce PE risk,⁵ yet diagnostic strategies remain suboptimal. While risk factors are known and guidelines delineate indications for ordering diagnostics,⁶ no prospectively validated risk stratification for inpatient DVT exists. For example, Wells score performs poorly on inpatients.⁷

DVT is typically diagnosed using ultrasonography: either compression (CUS) alone, duplex (CUS plus color Doppler), or triplex (duplex with continuous wave Doppler). One meta-analysis of radiology performed CUS reported a sensitivity of 94% for proximal DVTs and 57% for distal DVTs with an overall specificity of 98%.⁸ CUS has reduced accuracy with recurrent DVT but is considered appropriate for a suspected first episode.^{8, 9}

CUS can be performed as proximal only (femoral and popliteal veins) or whole-leg (proximal and distal veins). Serial two-point CUS (1-week apart) of the proximal venous system is equivalent to whole-leg examination for suspected first episode of DVT,^{10, 11} and proximal vein CUS is recommended as an initial test for suspected first episode of lower extremity DVT.¹² Despite this, formal ultrasounds are not universally available or technologists may only be available during weekday daytime hours, delaying formal read, diagnosis,^{13, 14} and potentially lifesaving anticoagulation.

Received April 30, 2018
Revised October 26, 2018
Accepted April 20, 2019
Published online August 6, 2019

Point-of-care ultrasound (POCUS) has emerged for rapid clinician diagnosis of DVT. Emergency Medicine (EM) physicians evaluated POCUS to diagnose DVT and reported a sensitivity and specificity of 96.1% and 96.8%, respectively.¹⁵ General practitioners had a sensitivity of 90% and specificity of 97.1% when assessing outpatients.¹⁶ In the ICU, critical care physicians had a sensitivity ranging 63–85% and specificity 96–97% for DVT diagnosis.^{13, 14} The variable DVT prevalence in these populations impacted positive and negative predictive values. In the Caronia et al. ICU study, the superficial femoral vein was not scanned, nor was the popliteal vein from its most proximal to its most distal extent. The authors attributed their lower sensitivity (63%) to these exclusions.¹³

The Hospitalist-Operated Compression Ultrasound Study – A Point-of-Care Ultrasound Study (HOCUS-POCUS) is a prospective, multi-centered trial to determine the accuracy and feasibility of hospitalist-performed POCUS in DVT evaluation of hospitalized, non-ICU patients. To our knowledge, POCUS for DVT diagnosis has not been evaluated on this population. We utilized a three-region compression protocol to attempt to identify all proximal DVTs. The primary outcomes of the study are the operating characteristics (sensitivity and specificity) of hospitalists performing DVT POCUS compared with radiology formal vascular studies (FVS) as the reference standard.

METHODS

Study Design and Population

This multicenter study was conducted at four institutions: The University of Minnesota, MedStar Georgetown University, HealthPartners/Regions, and The University of Cincinnati Medical Center. Each site's Institution Review Board approved the study. Patients were enrolled between May 2016 and June 2017.

Adults (≥ 18 years) were eligible for inclusion if they were admitted to a non-ICU general medicine ward and had a high enough pre-test probability that the primary team ordered a lower extremity DVT ultrasound of either or both legs. Patients were excluded if they (1) had critical illness precluding examination, (2) had DVT diagnosed within 90 days, (3) had pain limiting examination, (4) were unable to consent, or (5) were non-English speaking.

The study patients represent a convenience sample based on hospitalist availability. Hospitalist scanners were not necessarily contacted at the time of a FVS order and were never aware of FVS results prior to performing POCUS. Sonographers independently chose scanning days and each scanning day the ordering primary team would contact the hospitalist scanner, or the sonographer would use the medical record to find new

studies. Once identified, the protocol and FVS were completed regardless of the result (i.e., a negative POCUS could not be used to cancel the FVS). Neither potentially eligible nor excluded patients were tracked. Study flow is summarized in Figure 1.

Outcomes and Statistical Analysis

The pre-specified primary outcomes were the sensitivity and specificity of the POCUS CUS compared with a “gold standard” FVS, either the radiology performed duplex or triplex. The FVS is a practical standard as direct or CT/MRI venography is not routinely performed and duplex/triplex has excellent accuracy for proximal DVT.⁸ We recorded the time the POCUS was performed for comparison with the time the FVS was ordered and time the formal radiology report became available.

POCUS and FVS results were treated as dichotomous outcomes. POCUS study results were recorded as compressible or not, while for FVS, results were recorded as DVT present or absent. For safety, any non-compressibility was recorded as “positive” regardless of the hospitalist's suspicion for DVT. If an area was not visualized, this was noted but the study was still included in the analysis. For statistical analysis, legs were counted independently (a bilateral study was treated as two separate scans).

Based on previous literature, we expected the prevalence of DVT in the eligible population to be 22%.^{13–16} We anticipated POCUS to have a minimum sensitivity of 95%, requiring 317 extremities to provide a confidence interval of 5%.¹⁷ To reach the goal, each of the four centers would scan approximately 80 extremities. We did not meet this goal due to slow accrual and the study was stopped after 125 extremities.

All data were entered into the RedCap database. All POCUS data were recorded at the bedside and formal vascular study results and clinical information were documented. The elapsed times from FVS order to POCUS completion and to FVS report completion were tabulated. These were compared via the Wilcoxon signed-rank test. Performance of diagnostic testing was assessed by sensitivity, specificity, negative and positive predictive values (NPV and PPV, respectively), and negative and positive likelihood ratios, (LR+ and LR–, respectively). The 95% confidence intervals were calculated using the Bayesian Jeffreys method.^{18, 19} Statistical analyses were performed using SAS 9.4 (Copyright 2002–2012, SAS Institute Inc., Cary, NC, USA) and a Clinical Decision Making Spreadsheet Calculator designed by Robert M. Hamm (oumedicine.com/familymedicine/academic-information/research/research-faculty/rob-hamm/download-computer-programs-for-cdm-calculations/cdm-calculators, accessed September, 2018).

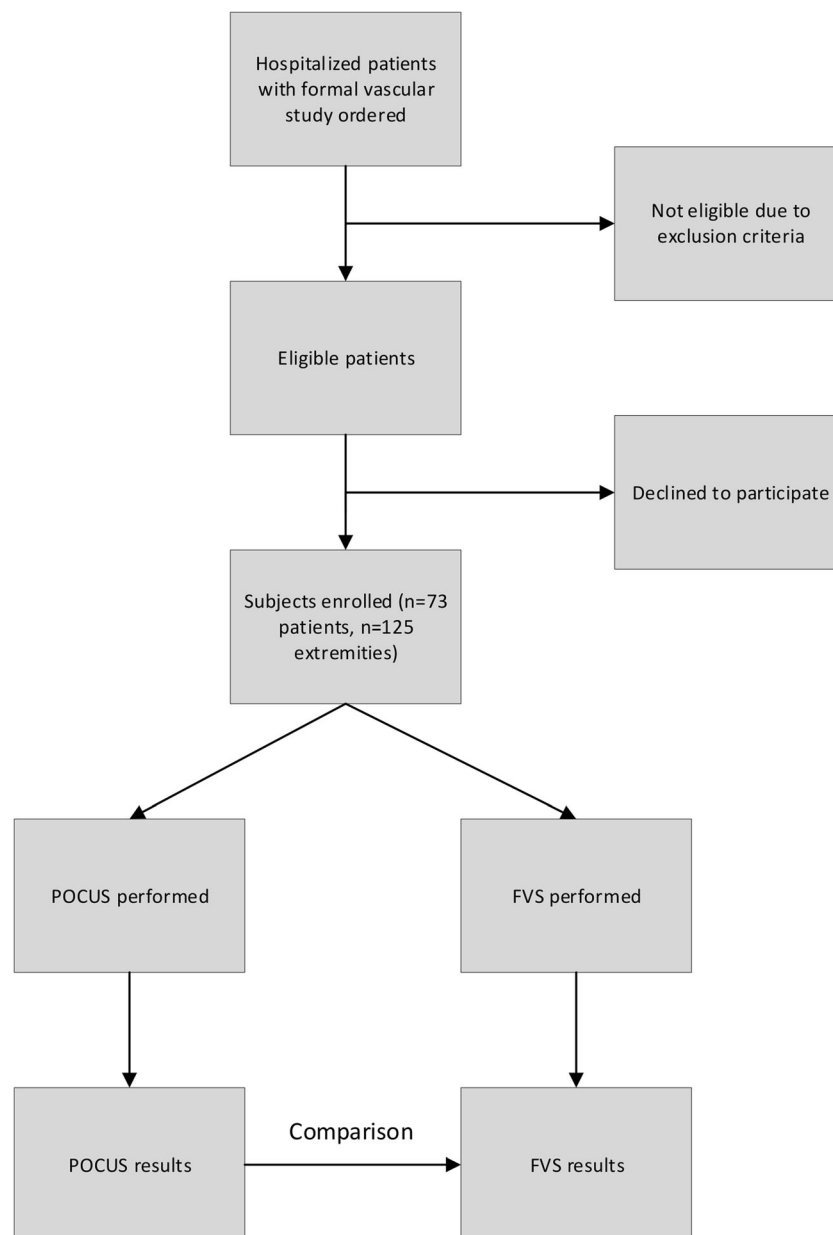


Fig. 1 HOCUS-POCUS study design and flow.

Ultrasound Training and Proficiency

All participating hospitalist scanners completed didactic and hands-on training²⁰ totaling 2 h. Each hospitalist had to successfully complete ten normal proctored DVT studies on standardized patients in order to begin enrolling patients. The didactic contained “abnormal” examples.

POCUS Protocol

POCUS was standardized as follows: The hospitalist scanner begins at the common femoral vein and compresses, moves distally both noting and compressing the greater saphenous, the common femoral arterial bifurcation, and venous bifurcation (zone 1) before reaching the proximal superficial femoral vein (SFV, zone 2). Our

protocol includes the entire SFV that is visible with a standard linear probe. The sonographer proceeds distally along the SFV compressing at 1–2-cm intervals until it is no longer visible, a variable distance dependent on patient anatomy. Finally, the popliteal vein is compressed (zone 3). Color or spectral Doppler were not used.

Ultrasound Machines

Each institution used the equipment available to them. MedStar Georgetown used a Sonosite M-turbo provided by the company for the study as well as a Sonosite Nanomaxx. The University of Cincinnati used a Sonosite S-Fast; HealthPartners used Sonosite Edge I, II, and Export; and the University of Minnesota used a MindRay T-7.

Table 1 Baseline Characteristics of Patients Enrolled

Characteristic	Value
Age, years (median, range)	61 (26–92)
BMI, kg/m ² (mean, standard deviation)	31.4 (8.4)
Male sex, %	49
Primary discharge diagnoses (n)	
Other*	15
Infection	14
Cancer	9
Respiratory	8
VTE	8
Cardiac/heart failure	7
AKI/ESRD	4
Trauma/wound	4
CVA/ICH	2
Hepatic/cirrhosis	2

*Other includes the following: abnormal vaginal bleeding, bilateral LE swelling, chest pain, complication of dialysis catheter, DKA, frequent falls, hypertensive emergency, hyponatremia, leukocytosis, periumbilical abdominal pain, pleuritic chest pain, right pneumothorax, SBO, UC flare, vocal cord dysfunction

RESULTS

Eight providers performed the POCUS DVT assessments across the four institutions.

A total of 73 patients representing 125 extremities were enrolled. Subject characteristics are shown in Table 1. The hospitalists had performed between 2 and 40 studies prior to the required training. The prevalence of DVT was 6.4% of limbs scanned (8/125) and 8.2% (6/73) of patients. Bilateral studies made up the majority of those ordered (71.2%, 52/73). In patients with bilateral studies ordered, the prevalence was 9.6% (5/52). In 21 patients with unilateral studies, only one DVT was found. The POCUS results compared with FVS are summarized in Table 2. When compared with the radiology FVS, the sensitivity of hospitalist-performed POCUS for DVT was 100% (95% CI 74–100%) and the specificity was 95.8% (95% CI 90–98%). Table 3 shows all operating characteristics of the hospitalist-performed POCUS. Details for the five false positive studies are shown in Table 4.

There was no statistical difference in the median time from FVS order to POCUS completion versus time to technician performing the FVS (5.8 vs 4.7 h, *p* = 0.36); however, the POCUS exam was completed before the FVS report 68% of the time. The median time until the FVS report became available was 11.5 h, significantly longer than POCUS completion (*p* = 0.001).

Table 2 Two by Two Table of Results Showing Bedside Sonographer POCUS Findings Compared with the Formal Vascular Study

		Formal vascular study	
		DVT present	DVT absent
POCUS result	Not compressible	8	5
	Compressible	0	112

Table 3 HOCUS-POCUS Study Outcomes Including Operating Characteristics of Hospitalist-Performed CUS for Diagnosis of DVT

Operating characteristic	Value	95% CI
Prevalence (125 extremities)	6.4%	
Prevalence (73 patients)	8.2%	
Sensitivity	100%	74–100%
Specificity	95.8%	91–98%
LR+	23.4	9.9–55.1
LR–	0	0–0.32
PPV	61.5%	35–84%
NPV	100%	98–100%

NPV¥ A range for the post-test probability of DVT was assumed based on the 95% CI for the NPV. Using this, the upper limit for 193 the LR– confidence interval could be estimated according to the Bayes theorem (LR = post-test odds/pre-test odds)

DISCUSSION

In HOCUS-POCUS, we have shown via a prospective, multicenter trial on hospitalized non-ICU patients that hospitalist performed POCUS for DVT has excellent specificity (95.7%) and that results were available over 5 h before completed radiology reports. The sensitivity and specificity are similar to POCUS studies in the general outpatient clinic,¹⁶ emergency department,¹⁵ and ICU.^{13, 14} Though our observed sensitivity was 100%, the confidence interval does not exclude that the true sensitivity could have been as low as 74%. Our negative predictive value of 100% (95% CI 98–100%) is also similar to others and suggests that trained hospitalists accurately exclude DVT in patients for whom there is reason to order an FVS. The 6.4% observed extremity prevalence of DVT resulted in a lower positive predictive value than previous POCUS studies (61.5% vs others at 80–90%). Prior POCUS DVT study results are summarized in Table 5.

Our study did have a number of limitations. One was the sample size. We did not reach our enrollment goal due to slow enrollment and lower than expected FVS frequency. The study was stopped prematurely as it became clear that the false negative rate would be extremely low and that slow

Table 4 False Positive Results within HOCUS-POCUS

Age, gender laterality	Description of findings
74-year-old male, unilateral (left)	1. Unable to compress left mid SFV, no clot identified. Unable to visualize left distal SFV. Patient sitting up in chair at time of exam; back pain limited ability to optimally position patient for duration of exam.
56-year-old female, unilateral (right)	2. Unable to compress branch point tributary near R CFV bifurcation.
58-year-old female, unilateral (right)	3. Unable to completely compress CFV bifurcation, no clot visualized. Scanner had low suspicion of true DVT as the vein immediately proximal was almost fully compressible and appeared issue seemed related to artifact, but this was called a positive exam due to the predetermined criteria.
76-year-old male, bilateral	4. Left: dialysis fistula present, unable to compress 5. Right: examiner indicated visible clot

Table 5 Summary of HOCUS POCUS and other POCUS DVT Study Outcomes Including Operating Characteristics

Study	Location	n	Prevalence	Sensitivity	Specificity	PPV	NPV
HOCUS-POCUS	Wards	125	6.4	100	95.7	61.5	100
PRACTICUS ¹⁶	Outpatient	1107	18.1	90	97.1	87.4	97.8
PD Kory, et al. ¹⁴	ICU	128	20	86	96	84.3	96.5
J Caronia, et al. ^{13*}	ICU	75	16	63†	97	80	93
F. Pomeroy, et al. ^{15†}	ED	2379	23.1	96.1	96.8	90	98.8

*Used a 2-point compression protocol and missed SFV and possibly popliteal DVTs

†Meta-analysis of 16 cohorts

enrollment would preclude timely completion. We do not believe the operating characteristics would significantly differ had we reached our goal, though the confidence intervals would likely be narrower.

Additionally, patients were only enrolled during pre-specified scanning days, leading to a convenience rather than consecutive sample. Yet median POCUS results were available over 5 h before median FVS reports. To limit selection bias, scanners chose scanning days in advance, and all eligible patients were approached to participate. Still, most scans were performed during the weekday, daytime hours, and it is possible that this patient subset differed from that during off-hours. Even though we cannot say included patients were randomly selected, there was no pattern to our scanning days and no reason to believe this led to a population bias, though it may limit generalizability.

We assumed observations were independent. Limbs are not biologically independent but it is unclear if they are statistically independent. Of the five positive bilateral studies, three (60%) were positive on only one limb. There are other potential factors that could have led to non-independent observations such as the scanner, institution, or sonographer experience. Our study was too small to explore these.

As we did not want to influence a patient's decision to proceed with a FVS, we excluded patients with significant pain, though these patients could have a higher prevalence of DVT. Furthermore, it is possible that patients with pain or higher degree of illness were more likely to decline participation. In contrast, we did not exclude those with prior DVT as part of the pragmatic study design, even though CUS has reduced accuracy in recurrence. Duplex or triplex is still the FVS ordered for recurrent DVT in the inpatient setting, and we were comparing hospitalist-performed POCUS to real-world practice.

The positive predictive value (61.5%) was lower than that in previous studies, owing to the lower DVT prevalence in our study. Our scanning protocol favored sensitivity over specificity, requiring categorical responses (compressible, non-compressible, or visualized clot) independent of the scanner's clinical suspicion. Though not representative of typical POCUS practice, this decision allowed evaluation of hospitalist POCUS technique and prioritized not missing critical DVT diagnoses. Not surprisingly, there were false positives in the study; however, many probably would not have been deemed positive in practice (see Table 4). We believe additional review

of clinical information would have decreased the pre-test probability of DVT or even negated some FVS orders (i.e., bilateral FVS for fever or bilateral swelling in heart failure).

The low overall DVT prevalence in our sample was unexpected. The predicted DVT prevalence was consistent with outpatient, ED, and ICU studies (see Table 5) and published FVS rates. However, FVS protocols often include distal deep veins and distal DVTs represent nearly half of all positive results.¹² Additionally, many FVSs are obtained in the high pretest scenario of known VTE (i.e., known PE). Known VTE and concern for distal DVT are not always scenarios to implement POCUS, which we advocate should be primary for rapid assessment that impacts clinical decision-making. We suspect the PPV would have been higher in a pragmatic study design in which the hospitalist could choose subjects based on pre-test probability. The logistics of such a study would invariably be more difficult.

CONCLUSIONS

HOCUS-POCUS adds to the literature on POCUS for DVT in several ways. It is the first study in non-ICU hospitalized patients and shows that hospitalists can be trained to competently perform POCUS with good sensitivity and excellent specificity. Second, the use of a three-region compression protocol as opposed to more common 2- or 3-point compression protocols resulted in no false negatives and a high NPV. Two-point compression protocols, though possibly sufficient in ambulatory patients, have missed clinically significant DVTs in the SFV in inpatients¹³.

In addition to establishing operating characteristics, this study showed the feasibility of training hospitalists in POCUS for DVT. Our hospitalists received only 2 h of training and completed ten practice scans on standardized patients. Many of the hospitalist scanners were new to POCUS or had no previous formal training. The training is both relatively fast and reproducible with minimal resources.

POCUS is an increasingly important skill for the modern hospitalist and we foresee the results from this study informing hospital medicine practice in myriad ways: (1) we present a learnable, effective protocol that improves decision-making for hospitalists working in settings where FVS cannot be obtained or interpreted in a reasonable timeframe, (2) this

protocol is safe and accurate, (3) our findings suggest that a negative POCUS scan could be sufficient to rule out DVT in a hospitalized patient, and (4) there is potential to develop a clinical workflow and encourage hospitalists to consider other pathologies after DVT is excluded via POCUS, before or possibly without a FVS. Further study is needed to ensure generalizability of our findings, understand how to integrate this into a hospitalist's workflow, and consider how this approach might be shared with trainees.

Acknowledgments: The authors wish to acknowledge all of their colleagues who participated in this study by performing POCUS scans.

Corresponding Author: Ernest A. Fischer, MD; Department of Medicine MedStar Georgetown University Hospital, Washington, DC, USA (e-mail: Ernest.A.Fischer@gunet.georgetown.edu).

Compliance with Ethical Standards:

Conflict of Interest: The authors declare that they have no conflict of interest.

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