Point-of-care assessment of platelet reactivity in the emergency department may facilitate rapid rule-out of acute coronary syndromes: a prospective cohort pilot feasibility study

The Harvard community has made this article openly available. Please share how this access benefits you. Your story matters

Citation

Published Version
doi:10.1136/bmjopen-2013-003883

Citable link
http://nrs.harvard.edu/urn-3:HUL.InstRepos:11879652

Terms of Use
This article was downloaded from Harvard University’s DASH repository, and is made available under the terms and conditions applicable to Other Posted Material, as set forth at http://nrs.harvard.edu/urn-3:HUL.InstRepos:dash.current.terms-of-use#LAA
BMJ Open

Point-of-care assessment of platelet reactivity in the emergency department may facilitate rapid rule-out of acute coronary syndromes: a prospective cohort pilot feasibility study

Chad E Darling,1 Javier A Sala Mercado,2,3,4 Walter Quiroga-Castro,4 Gabriel F Tecco,4 Felix R Zelaya,4 Eduardo C Conci,4 Jose P Sala,4 Craig S Smith,5 Alan D Michelson,6 Peter Whittaker,1,2,7 Robert D Welch,2,7 Karin Przyklenk1,2,3,7

ABSTRACT

Objective: Accurate, efficient and cost-effective disposition of patients presenting to emergency departments (EDs) with symptoms suggestive of acute coronary syndromes (ACS) is a growing priority. Platelet activation is an early feature in the pathogenesis of ACS, thus, we sought to obtain an insight into whether point-of-care testing of platelet function: (1) may assist in the rule-out of ACS; (2) may provide additional predictive value in identifying patients with non-cardiac symptoms versus ACS-positive patients and (3) is logistically feasible in the ED.

Design: Prospective cohort feasibility study.

Setting: Two urban tertiary care sites, one located in the USA and the second in Argentina.

Participants: 509 adult patients presenting with symptoms of ACS.

Main outcome measures: Platelet reactivity was quantified using the Platelet Function Analyzer-100, with closure time (seconds required for blood, aspirated under high shear, to occlude a 150 µm aperture) serving as the primary endpoint. Closure times were categorised as ‘normal’ or ‘prolonged’, defined objectively as the 90th centile of the distribution for all participants enrolled in the study. Diagnosis of ACS was made using the standard criteria. The use of antiplatelet agents was not an exclusion criterion.

Results: Closure times for the study population ranged from 47 to 300 s, with a 90th centile value of 138 s. The proportion of patients with closure times ≥138 s was significantly higher in patients with non-cardiac symptoms (41/330; 12.4%) versus the ACS-positive cohort (2/105 (1.9%); p=0.0006). The specificity of ‘prolonged’ closure times (≥138 s) for a diagnosis of non-cardiac symptoms was 98.1%, with a positive predictive value of 95.4%. Multivariate analysis revealed that the closure time provided incremental, independent predictive value in the rule-out of ACS.

Conclusions: Point-of-care assessment of platelet reactivity is feasible in the ED and may facilitate the rapid rule-out of ACS in patients with prolonged closure times.

Strengths and limitations of this study

- The study suggests that a technically straightforward and cost-effective test, with minimal patient risk, may serve as a useful adjunct to the current, standard emergency department practices for the rule-out of acute coronary syndromes. Importantly, the results are generalisable: the observation of a higher incidence of prolonged closure times in patients with non-cardiac symptoms was seen in participants from two distinct healthcare systems and populations.

- Platelet Function Analyzer-100 testing will only contribute to the identification of a subset of patients with non-cardiac symptoms, with the size of the subset and the potential value of the test dependent on the threshold used to define the ‘prolonged’ closure times.

- Limitations of this pilot study include the enrolment of patients via convenience sampling, the fact that platelet outcomes were not monitored beyond hospital discharge and that a risk/benefit analysis was not included in the study design.

Accurate, efficient and cost-effective diagnosis of patients presenting to emergency departments (EDs) with symptoms suggestive of acute coronary syndromes (ACS)—and, in particular, the exclusion and early discharge of patients with non-cardiac chest pain—is a growing priority. In an effort to meet this challenge, interest has focused on the identification of new approaches to augment the standard ED procedures and facilitate the timely triage of patients with suspected ACS. For example, there is recent evidence that coronary CT angiography (CCTA) combined with routine clinical assessment may provide added prognostic value in the management of chest pain patients in the ED. Use of
CCTA in low-to-intermediate risk patients is reportedly safe and reduces ED costs and hospital length of stay. However, these benefits are accompanied by exposure to radiation and associated with increases in diagnostic testing and subsequent invasive procedures.

Assessment of platelet activation, an early feature in the pathogenesis of ACS, has also been investigated as a possible benign strategy to expedite the diagnosis of ACS. Application of flow cytometry, the ‘gold standard’ for the quantification of molecular indices of platelet activation, is, however, impractical for routine use under emergent conditions. More importantly, classic molecular indices of platelet activation have not provided added benefit in the risk stratification of undifferentiated chest pain patients.

We hypothesised that rapid assessment of platelet reactivity using a technically straightforward point-of-care device—specifically, the Platelet Function Analyzer (PFA-100 (Siemens)—may represent a more feasible strategy to assist in the timely rule-out of ACS in the ED. Rather than quantifying molecular markers of platelet activation-aggregation, the output of the PFA-100 is the ‘closure time’: that is, the time required for whole blood, aspirated under high shear, to occlude a small, 150 μm aperture in a membrane coated with standard platelet agonists (collagen-ADP or collagen-epinephrine). Although the PFA-100 is typically utilised to investigate the responsiveness of patients to aspirin and other antiplatelet therapies and aid in the detection of platelet dysfunction, there is evidence to suggest that shortened closure times may be a marker of the acuity of coronary disease. Accordingly, our primary aim in this pilot study was to determine whether point-of-care testing of platelet function: (1) may assist in the rule-out of ACS; (2) may yield additional predictive value in identifying patients with non-cardiac symptoms versus ACS-positive patients and (3) is logistically feasible in an emergent setting. In addition, as a secondary analysis, we investigated whether the closure time may be of value in discriminating between ST-elevation myocardial infarction (STEMI) and non-ST-elevation myocardial infarction (NSTEMI)/unstable angina (UA).

METHODS
We conducted a prospective cohort feasibility study of patients presenting with potential ACS at two urban tertiary care sites: the ED at the University of Massachusetts (UMASS)-Memorial Medical Center, University Campus, Worcester, Massachusetts, USA, and the Cardiology Department, Instituto Modelo de Cardiologia Privado SRL, Cordoba, Argentina. The enrolment period was from January 2007 to December 2010. At each site, patients were entered on a convenience basis: that is, enrolments were not consecutive but, rather, coincided with the work schedules of the study investigators.

Patient population
Patients ≥18 years of age with a potential diagnosis of ACS (at the discretion of the treating physician and reflected by orders for an ECG and cardiac biomarkers) were considered for enrolment in the study. The prospective exclusion criteria were pregnancy, renal insufficiency (defined as serum creatine levels ≥1.5 mg/dL), anaemia (haematocrit <30%), platelet count <100,000/μL, major bleeding, any gastrointestinal bleeding, trauma or the inability to provide informed consent for any reason. The use of antiplatelet agents was not an exclusion criterion.

Study enrolment and protocol
As soon as possible after evaluation by ED staff, potential participants were approached for enrolment into the study. If the patient was too ill or otherwise unable to provide written consent, then proxy consent was attempted. If consent was obtained by proxy and the patient later became cognisant, they were given the option to continue or withdraw their participation in the study. If consent was withdrawn, all collected data were discarded.

Blood used in the PFA-100 was drawn within <1 h after presentation, at the same time as cardiac biomarker testing, and the closure time was measured within 3 h of collection. All samples were obtained via peripheral venipuncture and drawn into tubes containing 3.2% sodium citrate. Immediately before analysis, each sample was gently inverted; a 900 μL aliquot of whole blood was then applied to a collagen-ADP test cartridge and the closure time was quantified. Maximum test duration for the PFA-100 is 5 min; if the aperture is not occluded within this period, a closure time of ‘>300 s’ is displayed.

On enrolment, patients were questioned regarding a history of bleeding or platelet function disorders and ingestion of known antiplatelet agents within the past 7 days (aspirin, clopidogrel, non-steroidal anti-inflammatory agents). In addition, the medical record was reviewed and the standard cardiac risk factors (diabetes, hypertension, hypercholesterolaemia, smoking, family history of heart disease) were recorded and used in the calculation of thrombolysis in myocardial infarction (TIMI) risk score.

Patient diagnosis
The final diagnosis (non-cardiac symptoms vs ACS-positive) was established via a standardised chart review. All reviews were conducted by physicians in a blinded manner, without knowledge of the closure time data. For participants enrolled at the Instituto Modelo de Cardiologia Privado, patients were seen by a cardiologist at the time of hospital presentation, and all reviews were performed by a team of cardiologists (WQC, GFT, FRZ, ECC and JPS). At UMASS, each patient was first seen by an Emergency Medicine physician at the time of hospital presentation, and each chart was reviewed by an ED physician (CED). For patients with a definitive diagnosis of non-cardiac symptoms (ie, patients with no ischaemic ECG changes, no positive markers or provocative cardiovascular test results, and no history of a diagnosis of ACS of any type in the chart), no additional
review was performed. However, for patients who: (1) had positive cardiac biomarkers during admission; (2) underwent cardiac catheterisation; (3) had a positive provocative cardiovascular test (eg, exercise stress test) or (4) had ischaemic ECG changes, the chart was reviewed by an ED physician and a cardiologist. In any case in which the diagnosis was uncertain, the decision was adjudicated by a cardiologist (CSS).

Patients considered to be ACS positive were categorised into one of the two groups, STEMI or NSTEMI/UA, in accordance with the standard guidelines. Patients who did not meet these criteria were classified as having a diagnosis of non-cardiac chest pain/symptoms.

**Statistical analysis**

Our target sample size in this exploratory study was empiric. We reasoned that, as an approximate order of magnitude, ~100 patients would be required in the ACS-positive group in order to have the likelihood of discerning a potential difference in the closure times between patients with non-cardiac symptoms and the ACS-positive cohort. Assuming that 75% of patients enrolled in the study would have a diagnosis of non-cardiac chest pain (and thus 25% would be ACS positive), a total of approximately 400 patients would be required. To account for exclusions, target enrolment was set at ~500 participants.

**Univariate comparisons**

Our primary analyses focused on the comparison of patients in the two main outcome groups: non-cardiac symptoms versus ACS positive. All categorical data were reported as percentages. For continuous data, the D’Agostino and Pearson test was applied to determine normality and the appropriate parametric (t test) or non-parametric alternative (Mann-Whitney) tests were utilised. Secondary analysis (comparison of closure times among non-cardiac chest pain, STEMI and NSTEMI/UA groups) was conducted using the Kruskal-Wallis test. Results are reported as means±SD or medians with associated 10th and 90th centile ranges.

**Sensitivity, specificity and multivariate analysis**

We made the prospective and arbitrary decision to categorise the closure time into ‘normal’ and “prolonged” values based on the 90th centile of the distribution for all patients enrolled in the study. Utilising the normal and prolonged values of closure time, the sensitivity, specificity, negative and positive predictive values and likelihood ratio for identifying patients with non-cardiac symptoms were determined. Two reasons contributed to our choice of non-cardiac chest pain (rather than ACS positive) as the main outcome of interest: (1) this approach is consistent with current guidelines for the improvement of diagnostic accuracy and (2) we reasoned that prolonged closure times may be associated with the absence of ACS, whereas shorter closure times may be manifest in either group of patients. Sensitivity, specificity, predictive values and the likelihood ratio are reported with associated 95% CIs.

To determine whether the PFA closure time provides an additional predictive value to the standard clinical diagnostic information, a logistic regression was developed to assess its independent association with the main outcome of non-cardiac symptoms. Independent predictor variables considered to be mandatory in the model were the closure time (utilising continuous rather than the categorised values; the main predictor of interest) and the TIMI risk score (representing standard clinical information to predict ACS outcomes). To account for potential differences between UMASS and Cordoba, the study site was also added to the regression model. Other non-mandatory variables considered for inclusion were sex, a medical history of diabetes, hypertension, hypercholesterolaemia, smoking and clopidogrel use. As age and aspirin use are individually incorporated in the TIMI risk score, these variables were not considered for separate inclusion in the model. Non-mandatory variables were left in the model only if they yielded an increase in predictive value as determined by receiver operating characteristic (ROC) analysis or were of additional importance based on the 2 log-likelihood ratio test.

The reported estimate and adjusted OR for closure time were calculated for an increase in platelet closure time of 10 s (rather than 1 s), a choice based on the premise that a 1 s increase would be of limited clinical usefulness. The final model was evaluated by c-statistic (area under the ROC curve) and the Hosmer and Lemeshow fit test. As a sensitivity test, the results were calculated using the general estimating equation (PROC GENMOD) with study site evaluated as a subject factor (cluster) rather than a term in the model. The Net Reclassification Index (NRI) and Integrated Discrimination Improvement (IDI) statistic were not included in the analyses, given the scepticism and criticisms that have been raised concerning their value in predicting the potential, incremental prognostic impact of novel biomarkers.

Analyses were performed using GraphPad Prism V.5.04 (San Diego, California, USA) and SAS V.9.3 (Cary, North Carolina, USA).

**RESULTS**

**Enrolment and exclusions**

A combined total of 509 patients were enrolled at the two study sites (figure 1). Sixty-one participants were excluded because the closure time data were not available. The reasons included technical errors (n=14), closure times >300 s (possibly due to mild thrombocytopenia, anaemia or inadequate mixing of the blood sample prior to testing; n=36) and failure to measure the closure time despite obtaining consent (n=11). An additional five participants were removed from analysis because exclusion criteria were...
identified after consent was obtained (n=4), or consent was revoked after the blood sample was collected (n=1). For the remaining 443 patients in whom the closure time was quantified, 8 were diagnosed with thrombotic events that were non-cardiac in origin and thus were excluded from further analysis. Accordingly, results are reported for 435 participants (324 enrolled at UMASS and 111 enrolled in Cordoba): 105 were diagnosed with ACS and 330 had non-cardiac symptoms.

Demographics
As expected,29 the ACS-positive group had higher TIMI scores, was older, and had a higher proportion of male participants versus patients with non-cardiac symptoms (table 1). In addition, the incidence of hypercholesterolaemia and the use of clopidogrel were significantly higher in the ACS group. There were, however, no differences in the use of aspirin, incidence of hypertension or diabetes, proportion of smokers or a reported family history of heart disease between the two groups.

---

**Table 1** Demographics: all patients

<table>
<thead>
<tr>
<th></th>
<th>Non-cardiac symptoms (total n=330)</th>
<th>ACS positive (total n=105)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years): mean±SD</td>
<td>57±14 (n=328)</td>
<td>61±13 (n=104)</td>
<td>0.034</td>
</tr>
<tr>
<td>Male</td>
<td>65% (n=330)</td>
<td>80% (n=105)</td>
<td>0.004</td>
</tr>
<tr>
<td>TIMI score: mean±SD</td>
<td>1.9±1.4 (n=320)</td>
<td>3.1±1.4 (n=104)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Aspirin</td>
<td>71% (n=321)</td>
<td>64% (n=104)</td>
<td>0.222 (ns)</td>
</tr>
<tr>
<td>Clopidogrel</td>
<td>11% (n=325)</td>
<td>20% (n=104)</td>
<td>0.031</td>
</tr>
<tr>
<td>Smoker</td>
<td>29% (n=322)</td>
<td>28% (n=105)</td>
<td>0.901 (ns)</td>
</tr>
<tr>
<td>Hypertension</td>
<td>57% (n=322)</td>
<td>62% (n=105)</td>
<td>0.425 (ns)</td>
</tr>
<tr>
<td>Hypercholesterolaemia</td>
<td>57% (n=322)</td>
<td>69% (n=105)</td>
<td>0.030</td>
</tr>
<tr>
<td>Diabetes</td>
<td>24% (n=322)</td>
<td>27% (n=105)</td>
<td>0.601 (ns)</td>
</tr>
<tr>
<td>Family history</td>
<td>40% (n=319)</td>
<td>41% (n=105)</td>
<td>0.909 (ns)</td>
</tr>
</tbody>
</table>

ACS, acute coronary syndromes; TIMI, thrombolysis in myocardial infarction.
There was a modest but significant difference in the median closure times in patients with non-cardiac symptoms versus the ACS-positive group: 91 vs 87 s, respectively (p=0.0061; figure 2A). When the ACS-positive group was divided into STEMI and combined NSTEMI/UA cohorts, the modest differences in the closure time in the secondary three-group analysis remained significant (p=0.0001; figure 2C).

Of potentially greater importance, patients with non-cardiac symptoms versus ACS-positive patients were distinguished by differences in the proportion of the prolonged closure times. We found that 41/330 (12.4%) of patients with non-cardiac symptoms had closure times ≥138 s (ie, the 90th centile of the distribution for all patients enrolled in the study) while, in contrast, 2/105 (1.9%) of patients in the ACS-positive group had closure times ≥ this value (p=0.0006; figure 2B and table 2). The specificity and sensitivity of ‘prolonged’ closure times (≥138 s) for a diagnosis of non-cardiac symptoms were 98.1% and 12.4%, respectively, with a positive predictive value of 95.4%, a negative predictive value of 26.3% and a likelihood ratio of 6.52 (table 3).

Site differences: UMASS versus Cordoba

Patients at both sites displayed the expected differences in TIMI score, age and sex between non-cardiac chest pain and ACS-positive groups (data not shown). However, the participants enrolled in Cordoba were significantly younger, with a higher proportion of males.
and smokers but lower proportion of patients with diabetes, when compared with UMASS (table 4). In addition, and as expected, aspirin use was significantly lower among the cohort in Cordoba versus the UMASS population (40% vs 80%, p<0.0001, table 4). Despite these demographic differences, a higher incidence of prolonged closure times in patients with non-cardiac symptoms was observed at both sites: that is, the proportion of participants with closure times ≥138 s was 13% vs 10% in patients with non-cardiac chest pain and 1.9% vs 2% in ACS-positive patients at UMASS versus Cordoba, respectively.

Closure time as an independent predictor of diagnosis

Logistic regression analysis was first performed with mandatory variables (closure time, TIMI risk score) and non-mandatory variables (study site, sex, diabetes, hypertension, hypercholesterolaemia, smoking, clopidogrel use) included in the regression model (data not shown). ROC analysis revealed that, among the non-mandatory variables, only study site (UMASS vs Cordoba) contributed to an increase to the predictive value of the model for a diagnosis of non-cardiac symptoms. Comparison of ROC curves obtained by including the closure time, TIMI score and site in the model versus TIMI score and site alone showed a significant, incremental increase in area under the curve with the addition of closure time (0.818 vs 0.795; p=0.009; figure 3). Accordingly, the final multivariate logistic regression model incorporated these three variables (table 5).

For every 10 s increase in the closure time, the adjusted odds of a diagnosis of non-cardiac symptoms was 1.17 (95% CI 1.06 to 1.29; table 5). That is, when controlling for TIMI risk score and study site: for every 10 s increase, the prolonged closure time was associated with a 17% increase in the patient having non-cardiac chest pain. These results demonstrate that, irrespective of TIMI score and site, the closure time was associated with the diagnosis of non-cardiac symptoms, with closure time providing additional, incremental predictive value beyond that obtained by TIMI score. The model demonstrated good predictive characteristics (c=0.82) and model fit (χ²=5.24, df=8; p=0.73). Finally, the sensitivity analysis that utilised a general estimating equation accounting for potential correlations among sites did not result in any important changes in the direction of the effects (adjusted odds for every 10 s increase in closure time: 1.15 with 95% CI 1.10 to 1.23) or conclusions.

**DISCUSSION**

In this study, we demonstrate that point-of-care testing of platelet reactivity using the PFA-100 is feasible in the environment of the ED. The results of our primary analysis revealed differences in closure times between patients with non-cardiac symptoms versus ACS-positive cohorts, with the most notable, discriminating feature being the higher incidence of prolonged closure times in the group with non-cardiac chest pain. Finally, the outcome of our multivariate analysis is consistent with the concept that the assessment of closure time provides incremental, independent prognostic value beyond that obtained using the standard clinical predictor of TIMI score.

**Assessment of platelet reactivity as a diagnostic tool for ACS in emergent settings**

It is well-established that heightened platelet activity occurs in the setting of ACS. Indeed, measurement of platelet reactivity has been utilised in an effort to discern the stability of coronary disease, predict the future incidence and outcomes of major adverse cardiovascular events, and, although results have been disappointing, guide the dosing of antiplatelet agents with the goal of improving outcomes. Our current study differs from these previous reports, in that it focused on the assessment of platelet reactivity as an adjunct strategy to risk stratify patients with potential ACS in emergent conditions. Accordingly, the novelty of our study lies in our expanded analysis of the closure time, and the identification of a more practical and feasible application of these data in the prognosis of ACS.

Our observation of a modest but significant reduction in the median/mean closure times in patients with ACS is consistent with the outcomes of two previous, small ED studies. However, this ~4 s difference is of a limited practical significance given the broad and overlapping distributions of closure times for patients with non-cardiac symptoms and the ACS-positive cohort (figure 2). Rather, we propose that the clinical utility of

### Table 2  Incidence of prolonged closure time (≥138 s, defined as the 90th centile of the distribution of the study population)

<table>
<thead>
<tr>
<th></th>
<th>Non-cardiac symptoms</th>
<th>ACS positive</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prolonged closure time</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>41</td>
<td>2</td>
<td>43</td>
</tr>
<tr>
<td>No</td>
<td>289</td>
<td>103</td>
<td>392</td>
</tr>
<tr>
<td>Total</td>
<td>330</td>
<td>105</td>
<td>435</td>
</tr>
</tbody>
</table>

ACS, acute coronary syndromes.

### Table 3  Sensitivity, specificity, positive and negative predictive values and likelihood ratio of prolonged closure time for a diagnosis of non-cardiac symptoms

<table>
<thead>
<tr>
<th></th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sensitivity (%)</td>
<td>12.4</td>
</tr>
<tr>
<td>Specificity (%)</td>
<td>98.1</td>
</tr>
<tr>
<td>Positive predictive value (%)</td>
<td>95.4</td>
</tr>
<tr>
<td>Negative predictive value (%)</td>
<td>26.3</td>
</tr>
<tr>
<td>Likelihood ratio</td>
<td>6.52</td>
</tr>
</tbody>
</table>

95% CI = 9.1 to 16.5; 93.3 to 99.8; 84.2 to 99.4; 22.0 to 30.9; 1.61 to 26.51.
this test lies in the identification of patients with a prolonged closure time, a threshold that we objectively defined as the 90th centile of the distribution of the study population. In this regard, we found that the prolonged closure time (≥138 s) had a high specificity, positive predictive value and likelihood ratio for a diagnosis of non-cardiac chest pain.

**Strengths and weaknesses**

The results of our pilot study are consistent with the hypothesis that point-of-care testing of platelet reactivity may assist in the timely rule-out of ACS in the ED. Strengths of this approach include the fact that the assessment of closure time is technically straightforward and cost effective, with minimal patient risk (ie, does not involve exposure to radiation or additional invasive testing). In addition, our data suggest that the concept is generalisable. We observed significant differences in demographics (age and gender), the prevalence of specific risk factors (diabetes, smoking) and the use of anti-platelet therapy (aspirin: table 4) between UMASS and Cordoba—findings that are in agreement with the published reports focused on Argentine cohorts,30 31 42 and may underlie the robust contribution of study site as a predictor of outcome in our logistic regression model. Nonetheless, our results, obtained from two distinct healthcare systems and populations, consistently revealed a higher proportion of prolonged closure times in patients with non-cardiac symptoms.

There is, however, an important caveat to this strategy. PFA testing will only contribute to the identification of a subset of patients with non-cardiac symptoms, with the size of the subset and the potential value of the test dependent on the threshold used to define ‘prolonged’ closure times (figure 2 and table 6). For example, the prospective and arbitrary criterion used in our analysis (the 90th centile of closure times for all patients enrolled in the study) discerned an arguably modest 12.4% of patients with a diagnosis of non-cardiac symptoms. However, expediting the discharge of even a small proportion of patients with a non-cardiac diagnosis would limit the costs, potential risks and patient stress associated with unneeded advanced diagnostic testing and possibly invasive procedures.1 2 Reducing the threshold (ie, to the 80th centile of the distribution) would

<table>
<thead>
<tr>
<th>Demographics: UMASS vs Cordoba</th>
<th>UMASS all (total n=324)</th>
<th>Cordoba all (total n=111)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years): mean±SD</td>
<td>59±14 (n=324)</td>
<td>56±12 (n=108)</td>
<td>0.036</td>
</tr>
<tr>
<td>Male</td>
<td>65% (n=324)</td>
<td>78% (n=111)</td>
<td>0.009</td>
</tr>
<tr>
<td>TIMI score: mean±SD</td>
<td>2.3±1.5 (n=313)</td>
<td>2.0±1.4 (n=111)</td>
<td>0.142 (ns)</td>
</tr>
<tr>
<td>Aspirin</td>
<td>80% (n=314)</td>
<td>40% (n=104)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Clopidogrel</td>
<td>15% (n=319)</td>
<td>9% (n=111)</td>
<td>0.146 (ns)</td>
</tr>
<tr>
<td>Smoker</td>
<td>26% (n=316)</td>
<td>37% (n=111)</td>
<td>0.038</td>
</tr>
<tr>
<td>Hypertension</td>
<td>61% (n=316)</td>
<td>51% (n=111)</td>
<td>0.093 (ns)</td>
</tr>
<tr>
<td>Hypercholesterolaemia</td>
<td>62% (n=316)</td>
<td>53% (n=111)</td>
<td>0.115 (ns)</td>
</tr>
<tr>
<td>Diabetes</td>
<td>28% (n=313)</td>
<td>14% (n=111)</td>
<td>0.005</td>
</tr>
<tr>
<td>Family History</td>
<td>44% (n=313)</td>
<td>31% (n=51)</td>
<td>0.014</td>
</tr>
</tbody>
</table>

TIMI, thrombolysis in myocardial infarction.

<table>
<thead>
<tr>
<th>Multivariable logistic regression model (outcome modeled: non-cardiac symptoms)</th>
<th>Predictor</th>
<th>Adjusted OR</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Closure time</td>
<td>1.17</td>
<td>1.06 to 1.29</td>
</tr>
<tr>
<td></td>
<td>TIMI risk score</td>
<td>0.48</td>
<td>0.40 to 0.59</td>
</tr>
<tr>
<td></td>
<td>Study site</td>
<td>7.21</td>
<td>4.05 to 12.86</td>
</tr>
</tbody>
</table>

(UMASS vs Cordoba).

TIMI, thrombolysis in myocardial infarction.

Figure 3  Receiver operating characteristic (ROC) analysis. Comparison of ROC curves obtained by including the closure time, thrombolysis in myocardial infarction (TIMI) score and study site in the regression model versus the TIMI score and site alone showing a significant, incremental increase in area under the curve with the addition of closure time (0.818 vs 0.795; p=0.009).
increase the size of the subset identified as ACS negative, with an accompanying (and increasingly unacceptable) loss in specificity because of the growing proportion of false positives (ACS-positive patients with prolonged closure times: figure 2 and table 6). In contrast, increasing the threshold (ie, to the 95th centile) would identify a diminishing proportion of patients with non-cardiac symptoms with an increasing specificity (figure 2 and table 6). The appropriate definition of ’prolonged’ closure time will therefore require refinement based on risk/benefit analysis.

Finally, we emphasise that point-of-case assessment of platelet reactivity and identification of patients with ’prolonged’ closure times clearly cannot function as a stand-alone test for the rule-out of ACS. Rather, measurement of PFA closure times may serve as an adjunct to the current, standard ED practices. In support of this concept, our multivariate logistic regression model revealed that, irrespective of the TIMI score and study site (and, thus, irrespective of differences in demographics and aspirin use between sites), closure time was an independent predictor of the diagnosis of non-cardiac symptoms.

Summary, limitations and future directions
We report that measurement of the closure time using the PFA-100 provides additional and independent, incremental predictive value in the rule-out of ACS. Limitations of this pilot feasibility study include the enrolment of patients via convenience sampling, and differences in the logistics of the chart review process between the two sites. In addition, neither monitoring of patient outcomes beyond hospital discharge (raising the possibility of potential misclassification of some patients) nor risk/benefit analysis of PFA testing was incorporated into the study design. We emphasise that point-of-care assessment of platelet reactivity cannot serve as a stand-alone test to either discern patients with non-cardiac symptoms versus ACS-positive patients or distinguish between STEMI versus NSTEMI/UA in the emergent setting—limitations that in all likelihood reflect the complex and multifactorial pathophysiology of acute myocardial ischaemia and infarction. Rather, our results suggest that the assessment of closure times may provide benefit by augmenting the standard ED diagnostic practices, a concept that warrants further large-scale multicentre investigation.

Table 6 Effect of definition of ‘Prolonged’ closure time on specificity and positive predictive value for a diagnosis of non-cardiac symptoms

<table>
<thead>
<tr>
<th>Threshold</th>
<th>Specificity, 95% CI (%)</th>
<th>Positive predictive value, 95% CI (%)</th>
<th>Patients with non-cardiac symptoms identified (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>95th centile (≥160 s)</td>
<td>100 (96.6 to 100)</td>
<td>100 (83.9 to 100)</td>
<td>6.4 (21/330)</td>
</tr>
<tr>
<td>90th centile (≥138 s)</td>
<td>98.1 (93.3 to 99.8)</td>
<td>95.4 (84.2 to 99.4)</td>
<td>12.4 (41/330)</td>
</tr>
<tr>
<td>80th centile (≥117 s)</td>
<td>88.6 (80.9 to 94.0)</td>
<td>86.2 (77.2 to 92.7)</td>
<td>22.7 (75/330)</td>
</tr>
</tbody>
</table>

Author affiliations
1Department of Emergency Medicine, University of Massachusetts Medical School, Worcester, Massachusetts, USA
2Cardiovascular Research Institute, Wayne State University School of Medicine, Detroit, Michigan, USA
3Department of Physiology, Wayne State University School of Medicine, Detroit, Michigan, USA
4Division of Cardiology, Instituto Modelo de Cardiologia Privado SRL, Cordoba, Argentina
5Department of Cardiovascular Medicine, University of Massachusetts Medical School, Worcester, Massachusetts, USA
6Division of Hematology/Oncology, Center for Platelet Research Studies, Boston Children’s Hospital, Dana-Farber Cancer Institute, Harvard Medical School, Boston, Massachusetts, USA
7Department of Emergency Medicine, Wayne State University School of Medicine, Detroit, Michigan, USA

Contributors KP, CED and ADM participated in conception and design of the study; all authors participated in analysis and interpretation of the data and final approval of the manuscript; KP, CED, RDW, PW, JASM, CSS and ADM participated in drafting of the manuscript or revising it critically for important intellectual content; all authors had full access to all data and take responsibility for the integrity of the data and the accuracy of the analysis. KP is the guarantor.

Funding Siemens Inc provided the Emergency Department at the University of Massachusetts Medical School and the Cardiology Department, Instituto Modelo de Cardiologia Privado SRL with PFA-100 devices on a loan basis, as well as partial support for the purchase of test cartridges used in this study. CED was supported in part by a Research Training Grant from the Society for Academic Emergency Medicine (SAEM), Des Plaines, Illinois, USA.

Competing interests None.

Patient consent Obtained.

Ethics approval Institutional Review Boards from the two study sites, University of Massachusetts (UMASS)-Memorial Medical Center, University Campus, Worcester, Massachusetts, USA, and the Cardiology Department, Instituto Modelo de Cardiologia Privado SRL, Cordoba, Argentina.

Provenance and peer review Not commissioned; externally peer reviewed.

Data sharing statement No additional data are available.

Open Access This is an Open Access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 3.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited and the use is non-commercial. See: http://creativecommons.org/licenses/by-nc/3.0/

REFERENCES


