



The Randomised Assessment of Treatment using Panel Assay of Cardiac Markers (RATPAC) trial: a randomised controlled trial of point-of-care cardiac markers in the emergency department

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ABSTRACT

Objectives To determine whether using a point-of-care cardiac biomarker panel would increase the rate of successful discharge home after emergency department assessment, and affect the use of cardiac tests and treatments, subsequent attendance at or admission to hospital and major adverse events.

Design and setting Pragmatic multicentre randomised controlled trial in six acute hospitals in the UK.

Participants Patients attending with acute chest pain due to suspected myocardial infarction (N=2243).

Interventions Diagnostic assessment using a point-of-care biomarker panel consisting of creatine kinase, myocardial type, myoglobin and troponin I measured at baseline and 90 min compared with standard care without the point-of-care panel.

Main outcome measures The primary outcome was successful discharge home, defined as having left hospital or awaiting transport home by 4 h after attendance and no major adverse events up to 3 months. Secondary outcome measures included length of stay, use of coronary care, cardiac interventions and inpatient beds, emergency department attendances, subsequent admissions, outpatient visits and major adverse events.

Results Point-of-care panel assessment was associated with an increased rate of successful discharge (358/1125 (32%) vs 146/1118 (13%); OR 3.81, 95% CI 3.01 to 4.82; $p < 0.001$), reduced median length of initial hospital stay (8.8 vs 14.2 h; $p < 0.001$) and greater use of coronary care (50/1125 (4.0%) vs 31/1118 (3.0%); $p = 0.041$), but no difference in mean length of initial stay (29.6 vs 31.7 h; $p = 0.462$), mean inpatient days over follow-up (1.8 vs 1.7; $p = 0.815$) or major adverse events (36 (3%) vs 26 (2%); OR 1.31, 95% CI 0.78 to 2.20; $p = 0.313$).

Conclusions Point-of-care panel assessment increases successful discharge home and reduces median length of stay, but does not alter overall hospital bed use.

Trial registration Current controlled trials ISRCTN37823923.

The rise in hospital admissions has recently been described as a problem that 'threatens bankruptcy for the NHS'.¹ Chest pain is responsible for approximately a quarter of emergency hospital admissions² and recent developments are adding to the problem. The main reason for hospital attendance is the possibility of acute myocardial infarction. Recent public information campaigns have encouraged

patients to call for an emergency ambulance if they have acute chest pain. However, current recommendations suggest that diagnosis requires testing with a troponin sample taken 10–12 h after their symptom onset.^{3–4} This approach requires many patients to be unnecessarily admitted to hospital until the time delay has elapsed, thus incurring additional health service costs and inconvenience for patients.

It has been suggested that the cardiospecificity of troponin measurement should be combined with other biomarkers that are released earlier to facilitate rapid diagnosis. Combined measurement of cardiac troponin, myoglobin and the MB isoenzyme of creatine kinase (myocardial type; CK-MB) as a cardiac panel has been proposed as a way combining early and later markers. Recently developed sensitive troponin assays may further improve the early sensitivity of diagnostic testing,^{5–6} thus allowing the rapid rule-out of myocardial infarction and avoidance of admission, although this has yet to be tested in practice.

Point-of-care testing can substantially reduce turnaround time and allow rapid provision of results to the emergency department. The combination of cardiac troponin, CK-MB and myoglobin measured by point-of-care testing at baseline and 90 min later has been widely evaluated and used. Diagnostic cohort studies of this combination report high sensitivity for myocardial infarction^{7–14} and before-and-after intervention studies suggest its use can reduce turnaround times¹⁰ and coronary care unit admissions.¹¹ However, there have been no randomised controlled trials of this diagnostic strategy to determine reliably whether it can change patient management and reduce hospital admissions.

Although point-of-care cardiac marker panels have clear theoretical benefits, pragmatic evaluation is required to determine whether the rapid availability of accurate diagnostic information leads to changes in clinical decision-making and patient management. The Randomised Assessment of Treatment using Panel Assay of Cardiac markers (RATPAC) trial aimed to measure the effect of using a point-of-care cardiac marker panel upon discharge home after emergency department assessment, use of coronary and intensive care and cardiac treatments, subsequent re-attendance at and/or re-admission to hospital and major adverse events.

METHODS

We undertook a pragmatic randomised controlled trial of a point-of-care cardiac marker panel in the management of patients with suspected, but not confirmed, myocardial infarction in six emergency departments in the UK: Barnsley District General Hospital, Derriford Hospital (Plymouth), Edinburgh Royal Infirmary, Frenchay Hospital (Bristol), Leeds General Infirmary and Leicester Royal Infirmary. We selected hospitals that would provide a diverse range of settings, were able to support point-of-care testing in the emergency department, did not already use a point-of-care cardiac marker panel and were able to support emergency care research. Research nurses or emergency department staff screened all patients with chest pain and excluded those with electrocardiogram (ECG) changes for myocardial infarction or high-risk acute coronary syndrome (>1 mm ST deviation or >3 mm inverted T waves), known coronary heart disease (CHD) presenting with prolonged (>1 h) or recurrent episodes of cardiac-type pain, confirmed or suspected serious non-coronary pathology (eg, pulmonary embolus), co-morbidity or social problems that require hospital admission, an obvious non-cardiac cause (eg, pneumothorax or muscular pain), more than 12 h since their most significant episode of pain, previous participants, those unable to understand the trial information and those unwilling to consent. Research nurses recorded details of all excluded patients during every fourth week of the trial.

Research nurses or emergency department staff provided trial information and obtained written consent. Participants were then randomly allocated to receive either: (1) diagnostic assessment using the point-of-care biomarker panel; or (2) standard care without the panel, according to a simple randomisation sequence, stratified by centre, generated by the Nottingham Clinical Trials Unit and accessed through a secure website. The allocated treatment group was only revealed after the participant's details were entered, written consent was confirmed and the participant irrevocably entered into the trial.

Patients in the intervention arm received testing with the point-of-care panel, while all other tests and treatments were available as normal. RATPAC was a pragmatic trial so the point-of-care test was provided with a recommended protocol for use but management decisions were ultimately at the discretion of the clinical staff and all other diagnostic tests and the use of laboratory blood tests in the control group were at the discretion of the clinical staff. Awareness of the intervention was essential to its implementation so there was no attempt to blind clinical staff, patients or carers to the allocated treatment group.

The point-of-care cardiac marker panel comprised CK-MB (mass), myoglobin and troponin I, measured at presentation and 90 min later, using the Siemens Stratus CS analyser (Siemens Healthcare Diagnostics, Frimley, UK).¹⁵ The analytical characteristics of the assays were as follows: troponin I detection limit 0.02 µg/l, analytical range 0.02–50 µg/l, interassay coefficient of variation (CV) 4.3–5.1% (0.03–0.22 µg/l). The 99th centile of the assay is 0.07 µg/l. Myoglobin: detection limit 1 µg/l; analytical range 1–900 µg/l; interassay CV 1.9–12.7% (56–308 µg/l); 95% reference interval, men 21 to 98 µg/l, women 19 to 56 µg/l, combined 20 to 82 µg/l. CK-MB: detection limit 0.3 µg/l; analytical range 0.3–150 µg/l; interassay CV 0.15–1.27% (3.7–39.3 µg/l); 95% reference interval 0.6 to 3.5 µg/l.

Clinical staff were trained to use the test and given guidance in interpretation of the results. We provided guidance that advised a first panel test immediately after initial emergency

department assessment and a second panel test 90 min later. Initially the guidance recommended hospital admission if any troponin level exceeded 0.02 µg/l, if any CK-MB level exceeded 5 µg/l, if the CK-MB gradient exceeded 1.6 µg/l or if the myoglobin gradient exceeded 25% of the initial level. The troponin threshold was amended during the trial in the light of recommendations that the 99th percentile should be used as a diagnostic threshold for myocardial infarction to recommend admission if either sample level exceeded 0.07 µg/l or if the first sample was less than 0.03 µg/l and the second was between 0.03 and 0.07 µg/l.

The standard care group was managed without point-of-care testing, according to existing guidance at each hospital. One hospital used a troponin T assay with a diagnostic threshold of less than 0.01 µg/l measured 6 h after worst symptoms, whereas the other hospitals used troponin I assays with a diagnostic threshold ranging from less than 0.05 to less than 0.2 µg/l measured at 12 h after worst symptoms. Two hospitals admitted patients to an inpatient ward, one to a medical assessment unit and three to a clinical decision unit.

The primary outcome was the proportion of patients successfully discharged home after emergency department assessment, ie, discharged and having no adverse event (as defined below) during the following 3 months. To be defined as discharged the patient had to have left the hospital by 4 h after arrival or be awaiting transport home from hospital with a discharge decision made. This outcome was chosen because hospital admission is inconvenient for the patient and costly for the health service, so reducing the probability of admission would represent a meaningful benefit from rapid diagnostic testing. The 4-h time limit was chosen because the participating hospitals were subject to a national target of discharging patients from the emergency department within 4 h of arrival. Secondary outcomes included use of the coronary care unit, cardiac medications and cardiac interventions, re-attendance at and/or re-admission to hospital over the following 3 months and adverse events (death, non-fatal myocardial infarction according to the universal definition,¹⁶ emergency revascularisation, life-threatening arrhythmia or hospitalisation for myocardial ischaemia). We also measured patient satisfaction, health utility, healthcare costs and the proportion of patients admitted with myocardial infarction, although these will be reported elsewhere (S Goodacre, M Bradburn, P Fitzgerald, *et al*, submitted January 2010).

Recruiting staff recorded baseline data, the results of initial assessment and emergency department disposition. Research nurses then used emergency department and hospital inpatient notes to record management decisions at initial attendance and admission, extract resource use data and identify subsequent hospital attendances or admissions and adverse events up to 3 months. Participants were mailed a questionnaire at 1 and 3 months to identify adverse events and hospital attendances, health and social care resource use, and measure EQ-5D and satisfaction with care.

Initial plans were to recruit 3130 participants to the trial. Previous data suggested that 50% of the routine care group would be successfully discharged.¹⁷ With 1565 evaluable subjects in each arm the trial would have 80% power to detect a 5% improvement (to 55% of patients successfully discharged) at the two-sided significance level of 5%. The same sample size provided 80% power to detect a reduction from 4% to 2% in major adverse events, again at the two-sided 5% level of significance.

Recruitment was terminated early after the funders declined a request for additional resources to continue the trial, having taken into account a conditional power calculation from the Data Monitoring Committee. This estimated the conditional power to detect a significant difference in the primary outcome was greater than 99.9% and the conditional power to detect a difference in the adverse event rate was less than 10%, thus suggesting grounds for termination on the basis of futility (ie, further enrolment would not significantly improve the statistical power).

We analysed the primary outcome through logistic regression, fitting concurrently with intervention group the effects of centre, age, gender and past history of CHD to present adjusted OR along with their corresponding 95% CI. Analysis was on an intention to treat basis.

RESULTS

We recruited 2263 participants between 30 January 2007 and 2 June 2008. There were a total 2658 hospital-days of recruitment with a mean of 0.9 patients recruited per hospital per day. Over 667 hospital-days of screening we identified 9109 patients with chest pain, of whom 1295 (14%) had ECG changes, 1378 (15%) had known CHD with prolonged or recurrent pain, 724 (8%) had suspected serious non-CHD pathology, 414 (5%) had comorbidities or social problems mandating hospital admission, 2506 (28%) had obvious non-cardiac pain, 465 (5%) had over 12 h since their worst pain, 21 (<1%) had previously participated, 109 (1%) were unable to understand trial information, 198 (2%) had other exclusion criteria, 240 (3%) had an unknown reason for exclusion, 40 (<1%) declined consent, 1115 (12%) were eligible but recruitment was not sought and 604 (7%) were recruited.

Table 1 shows the baseline characteristics of the study groups and figure 1 shows the flow of recruited patients through the trial. Some 140 patients (12.4%) in the point-of-care group did not receive point-of-care testing as recommended: 34 did not receive any point-of-care testing, mostly due to problems using the technology, while the remainder received point-of-care in combinations other than those recommended. Two patients (0.2%) in the standard care group received point-of-care testing.

In the point-of-care group 362 patients were initially discharged (319 had left the emergency department and 43 were awaiting transport by 4 h), but four re-attended with a major adverse event, so overall 358/1125 (32%) were successfully discharged. In the standard care group 147 patients were initially discharged (134 had left and 13 were awaiting transport), but one re-attended with a major adverse event, so overall 146/1118 (13%) were successfully discharged. Point-of-care testing was associated with a significant increase in successful discharge (OR 3.81, 95% CI 3.01 to 4.82; $p < 0.001$). Figure 2 shows that the difference in the proportion in hospital lasted until approximately 24 h after attendance.

Table 2 shows initial care provided in the first 24 h. Patients receiving point-of-care were more likely to be admitted to the coronary care unit and receive clopidogrel, whereas patients receiving standard care were more likely to receive aspirin.

The mean length of the initial hospital stay was 29.6 h in the point-of-care group and 31.8 h in the standard care group (mean difference 2.2 h, 95% CI -3.7 to 8.0; $p = 0.462$ (t-test)). The median length of initial hospital stay was markedly shorter than the mean and was shorter in the point-of-care arm: 8.8 h for the point-of-care group and 14.2 h for the standard care group ($p < 0.001$, equality-of-medians test).

Table 1 Patient demographics and characteristics

	Point-of-care (N = 1125)	Standard care (N = 1118)
Mean age, years (SD)	54.5 (13.8)	54.6 (14.4)
Median age, years (IQR)	53.4 (44–64)	53.1 (44–64)
Min–max age, years	22, 93	23, 96
Male	683 (61%)	624 (56%)
Female	442 (39%)	494 (44%)
Centre		
Barnsley District General Hospital	162 (14%)	164 (15%)
Derriford Hospital, Plymouth	164 (15%)	164 (15%)
Edinburgh Royal Infirmary	228 (20%)	224 (20%)
Frenchay Hospital, Bristol	233 (21%)	231 (21%)
Leeds General Infirmary	173 (15%)	171 (15%)
Leicester Royal Infirmary	165 (15%)	164 (15%)
Past history of CHD		
No	985 (88%)	973 (88%)
Yes	132 (12%)	137 (12%)
Previous MI	60 (5%)	65 (6%)
Angina plus positive diagnostic test	46 (4%)	53 (5%)
CABG surgery	12 (1%)	15 (1%)
Angioplasty	37 (3%)	34 (3%)
Stenosis >50% on coronary angiography	14 (1%)	12 (1%)
Unproved clinical label of CHD	36 (3%)	31 (3%)
Other	12 (1%)	10 (1%)
Risk factors		
Diabetes	86 (8%)	92 (8%)
Hypertension	376 (34%)	361 (33%)
Hyperlipidaemia	271 (26%)	282 (27%)
Present smoker	310 (28%)	316 (29%)
Ex-smoker—last 10 years	144 (13%)	129 (12%)
Cocaine abuse	6 (1%)	10 (1%)
First-degree relative with angina or MI, onset age <60 years	344 (33%)	352 (34%)
Source of referral		
General practitioner	188 (17%)	189 (17%)
Emergency ambulance	481 (43%)	510 (46%)
Self-referred	419 (37%)	375 (34%)
Other	35 (3%)	41 (4%)

CABG, coronary artery bypass grafting; CHD, coronary heart disease; IQR, interquartile range; MI, myocardial infarction.

Table 3 shows the total number of days spent in hospital over the 3-month follow-up and the number of days spent on the coronary care or intensive care unit, including the initial hospital admission. We only recorded an inpatient day if the patient stayed overnight, so a proportion of patients were recorded as having no inpatient days if their initial hospital visit did not result in an overnight stay and they were not admitted on a subsequent occasion. More patients in the point-of-care group had no inpatient days (54% vs 40%; $p < 0.001$) but there was no difference in mean inpatient days. Figure 3 shows that these data are highly skewed with a few patients having many inpatient days. The point-of-care group had more patients with very long stays. This attenuated the effect on mean inpatient days of having a greater proportion with no hospital stay. Point-of-care testing was also associated with a higher use of coronary care (4% vs 3%), although only a small minority of patients received any coronary or intensive care.

Figure 1 Flow of patients through the trial.

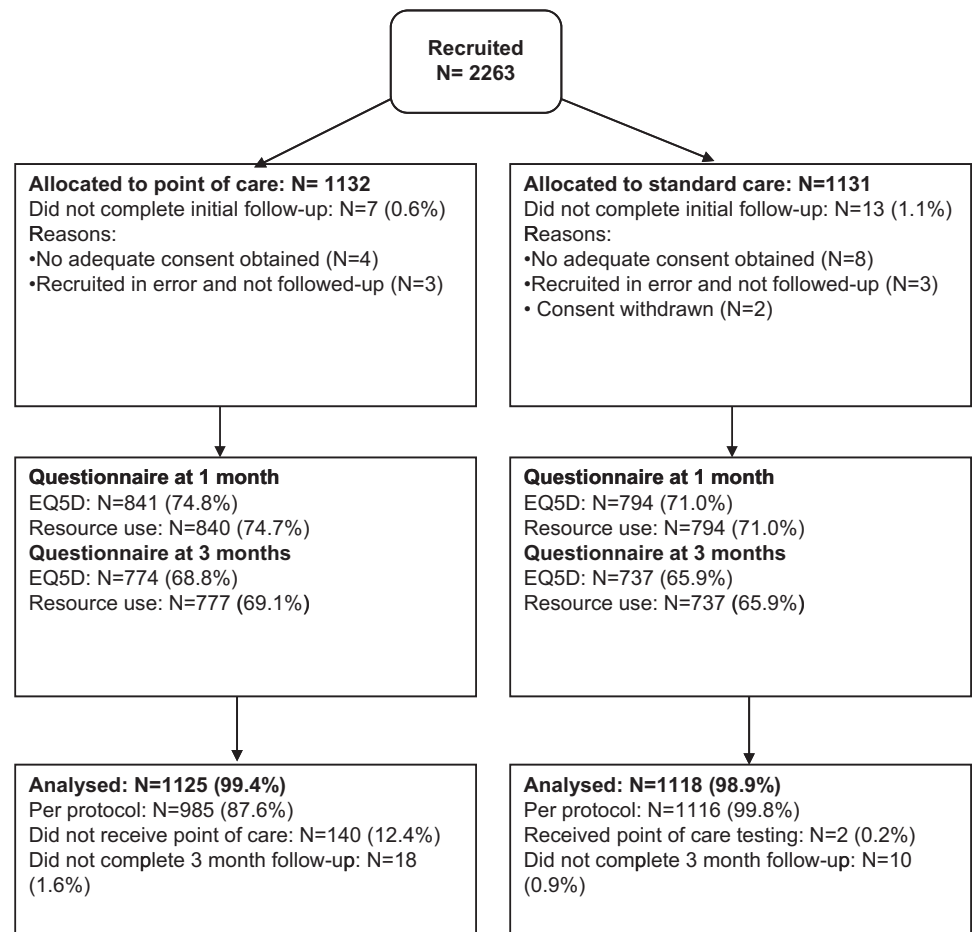


Table 4 shows cardiac interventions, emergency department attendances, hospital admissions (excluding admission at initial attendance) and outpatient reviews over the 3-month follow-up. The only difference between the groups was a borderline significant difference in chest pain-related outpatient attendances.

Overall 62 of 2243 patients (3%) experienced an adverse event, including eight deaths and 10 non-fatal myocardial infarctions. Table 5 shows that there were no significant differences in major adverse events between the two groups. Five adverse events

occurred in patients who were discharged home after emergency department assessment. Details are provided in table 6.

DISCUSSION

The use of the point-of-care cardiac marker panel resulted in a greater proportion of patients being successfully discharged after emergency department assessment and a reduction in the median, but not the mean length of initial hospital stay. It was associated with more patients avoiding any inpatient stay over the 3-month follow-up but did not lead to any difference in the total or mean number of inpatient hospital days. This was

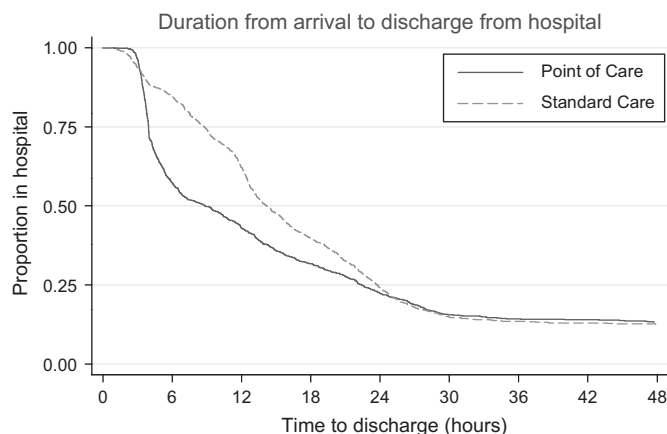


Figure 2 Variation in the proportion of patients in hospital over time from initial attendance.

Table 2 Initial care in the first 24 h

	Point-of-care N = 1125	Standard care N = 1118	p Value
Admitted to coronary care unit	50/1121 (4%)	21/1114 (2%)	0.001
Received GTN	446/1113 (40%)	460/1092 (42%)	0.327
Received heparin	206/1114 (18%)	186/1090 (17%)	0.381
Received glycoprotein inhibitors	8/1111 (1%)	7/1092 (1%)	0.822
Received antacid	90/1110 (8%)	108/1083 (10%)	0.128
Received angiography	16/1119 (1%)	8/1109 (1%)	0.105
Received aspirin	618/1118 (55%)	663/1109 (60%)	0.031
Received beta-blocker	108/1109 (10%)	105/1084 (10%)	0.967
Received clopidogrel	237/1110 (21%)	176/1091 (16%)	0.002
Received analgesic	373/1110 (34%)	407/1086 (37%)	0.058
Received any other drugs	316/1108 (29%)	340/1085 (31%)	0.150
Received coronary angioplasty or stent	11/1120 (1%)	4/1108 (<1%)	0.073

GTN, glyceryl trinitrate.

Table 3 Inpatient care at 3 months

	Point-of-care N = 1125	Standard care N = 1118	p Value
Days in hospital at any location			
N (%) with no days	607 (54%)	448 (40%)	<0.001
Mean days per patient	1.8	1.7	0.815
Total days (all patients)	1961	1908	
Days in coronary care			
N (%) with days on coronary care	50 (4%)	31 (3%)	0.041
Mean days per patient	0.17	0.09	0.033
Total days (all patients)	190	105	
Days in intensive care			
N (%) with days on intensive care	8 (1%)	3 (<1%)	0.225
Mean days per patient	0.08	0.01	0.259
Total days (all patients)	76	14	

because patients in the point-of-care arm who were admitted to hospital tended to accrue more inpatient days. Point-of-care assessment was also associated with a small increase in coronary care admission and chest pain-related outpatient follow-up. These findings suggest that point-of-care assessment changes the emergency department disposition of patients with undiagnosed chest pain and may reduce inpatient bed turnover, but does not reduce inpatient bed occupancy.

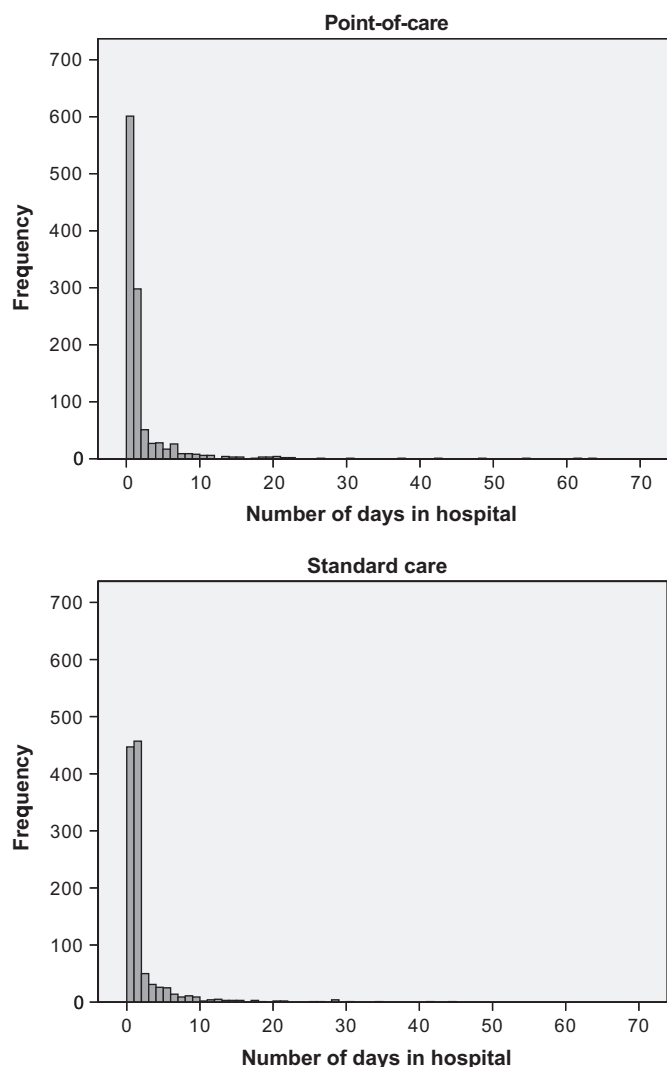


Figure 3 Histograms of number of inpatient days.

Table 4 Cardiac interventions, emergency department attendances, admissions and outpatient reviews

	Point-of-care N = 1125	Standard care N = 1118	p Value
N (%) needing thrombolysis	1 (<1%)	2 (<1%)	0.624
N (%) needing PCI	29 (3%)	29 (3%)	1.000
N (%) needing emergency PCI	7 (1%)	12 (1%)	0.260
N (%) needing CABG	11 (1%)	5 (<1%)	0.209
N (%) of emergency department attendances	140 (12%)	138 (12%)	0.949
N (%) of chest pain related emergency department attendances	107 (10%)	103 (9%)	0.828
N (%) of hospital admissions*	117 (10%)	122 (11%)	0.732
N (%) of chest pain-related hospital admissions*	84 (7%)	99 (9%)	0.247
N (%) with any outpatient attendance	334 (30%)	322 (29%)	0.676
N (%) with a chest pain related outpatient attendance	241 (21%)	202 (18%)	0.05

*Excluding admission at initial attendance.
CABG, coronary artery bypass grafting; PCI, percutaneous coronary intervention.

Interpretation of these findings depends upon one's perspective. For the patient, emergency physician or admitting physician point-of-care testing has the potentially beneficial effect of reducing the need for hospital admission. For the health service manager point-of-care testing may offer some benefit by reducing inpatient bed turnover but does not appear to reduce bed occupancy. An accompanying economic analysis (P Fitzgerald, SW Goodacre, E Cross, S Dixon, on behalf of the RATPAC research team, submitted July 2010) explores whether the potential benefits of point-of-care testing justify the additional costs.

Previous studies of point-of-care cardiac marker panels have focused on estimating accuracy for diagnosing myocardial infarction, but few have evaluated the effect on patient care. Newby *et al*⁹ demonstrated earlier identification of myocardial infarction than laboratory testing and Carragher *et al*¹⁰ showed expedited decision-making with turnaround times reduced by 55%, but it was not clear whether these led to meaningful changes in patient care. Ng *et al*¹¹ compared management with the panel to previous practice and showed a 40% reduction in coronary care unit admissions. Our finding of increased coronary care admissions may reflect differences between the healthcare systems or differences between the methods used. Historically controlled studies tend to overestimate the effects of intervention compared with randomised studies.

Table 5 Major adverse events

	Point-of-care N = 1125	Standard care N = 1118	OR (95% CI)*	p Value*
Any event	36 (3%)	26 (2%)	1.31 (0.78 to 2.20)	0.313
Death	6 (1%)	2 (<1%)	3.4 (0.7 to 17.3)	0.142
Non-fatal myocardial infarction	5 (<1%)	5 (<1%)	0.9 (0.3 to 3.2)	0.903
Hospitalisation for acute coronary syndrome (without myocardial infarction)	18 (2%)	9 (1%)	1.8 (0.8 to 4.1)	0.149
Life threatening arrhythmia	6 (1%)	2 (<1%)	3.2 (0.6 to 15.9)	0.160
Emergency revascularisation	10 (1%)	14 (1%)	0.7 (0.3 to 1.5)	0.324

*Adjusted for age, gender and known coronary heart disease.

Table 6 Major adverse events

Age and gender	Study group	Testing at initial presentation	Working diagnosis	Adverse event
43, female	Point-of-care	Point-of-care tests negative, taken 505 and 600 min after worst pain	Musculoskeletal pain	Hospitalisation for ACS 50 days later
63, male	Point-of-care	Point-of-care tests negative, taken 721 and 824 min after worst pain	Musculoskeletal pain	Hospitalisation for ACS 1 day later
64, female	Standard care	Troponin negative 1606 min after worst pain	Angina no ACS	Non-fatal MI 32 days later
78, female	Point-of-care	Point-of-care tests negative, taken 800 and 900 min after worst pain	Gastro-oesophageal pain	Died from metastatic pancreatic carcinoma 75 days later
78, male	Point-of-care	CK-MB and myoglobin negative, troponin 0.06 and 0.07 at 909 and 1005 min after worst pain	Angina no ACS	Life-threatening arrhythmia (SVT) 48 days later

ACS, acute coronary syndrome; CK-MB, creatine kinase, myocardial type; MI, myocardial infarction; SVT, supraventricular tachycardia.

Randomised trials have compared the use of point-of-care and laboratory troponin assays. Renaud *et al*¹⁸ showed that point of care troponin testing in an emergency department reduced time to anti-ischaemic therapy and physician notification of troponin results, but did not change emergency department length of stay or patient outcomes. Ryan *et al*¹⁹ evaluated point-of-care troponin testing in four emergency departments and found that the effect varied between settings, with length of stay in the emergency department being increased in one hospital and decreased in another. These studies only evaluated the impact of point-of-care technology, whereas RATPAC evaluated the effect of point-of-care testing in a rapid rule-out protocol.

Our study has a number of strengths and weaknesses. Individual patient randomisation ensured allocation concealment and allowed an unbiased comparison of the intervention with standard care. Participation of six varied hospitals ensured that findings are generalisable across a range of settings. An important limitation is that the trial was not powered to detect potentially important differences in the adverse event rate. The overall adverse event rate was very low and most events occurred in patients admitted after initial assessment. Only one of the five adverse events in patients who were initially discharged home occurred within 1 month of recruitment, so there is little evidence of significant missed pathology. Furthermore, point-of-care testing was performed relatively late in this case so admission for

a 12-h troponin is unlikely to have yielded a positive result. Another limitation is that clinician behaviour may have been influenced by participation in the trial, with perhaps more cautious management of patients in either, or both, study groups. This may be reflected in the lower than anticipated rate of discharge in the control group. This could have been addressed by using cluster randomisation, albeit with associated problems of reduced statistical power and loss of allocation concealment. Finally, it should be emphasised that this is a pragmatic evaluation of point-of-care panel assessment in which patients were selected on the basis of clinical and ECG characteristics and decision-making was ultimately at the discretion of the clinician. The biomarker panel was used to augment, and not replace, clinical assessment.

In summary, we have shown that use of the point-of-care cardiac marker panel safely reduced hospital admissions but did not alter the mean length of hospital stay or inpatient days. The implications for practice will depend upon whether avoiding hospital admission is important to patients and the hospital, and the findings of the economic analysis undertaken alongside this trial (P Fitzgerald, SW Goodacre, E Cross, S Dixon, on behalf of the RATPAC research team, submitted July 2010)

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What is already known on this subject

- ▶ Chest pain due to suspected myocardial infarction is responsible for a substantial and increasing number of emergency department attendances and hospital admissions.
- ▶ A point-of-care cardiac biomarker panel consisting of CK-MB, myoglobin and troponin can safely rule out myocardial infarction in patients with acute chest pain within 90 min of presentation to hospital.

What this study adds

- ▶ Using the point-of-care panel increases the proportion of patients successfully discharged after emergency department assessment and reduces the median length of stay.
- ▶ Using the point-of-care panel does not alter the mean length of stay or hospital days, and increases the use of coronary care.

(RATPAC research nurse, Frenchay Hospital, Bristol); Sue Mackness (RATPAC research nurse, Leicester Royal Infirmary); Yvonne Meades (RATPAC research nurse, Leeds General Infirmary); David Newby (professor of cardiology, University of Edinburgh/co-applicant); Dawn Newell (RATPAC research nurse, Leicester Royal Infirmary); Doris Quartey (RATPAC research nurse, Leeds General Infirmary); Karen Robinson (RATPAC research nurse, Leicester Royal Infirmary); Glen Sibbick (RATPAC research nurse, Leicester Royal Infirmary); Jason Smith (consultant in emergency medicine, Derriford Hospital, Plymouth); Roz Squire (RATPAC research nurse, Derriford Hospital, Plymouth).

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