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# High sensitivity troponin assays for the early rule-out or diagnosis of acute myocardial infarction in people with acute chest pain: a systematic review and cost-effectiveness analysis

A Diagnostic Assessment Report commissioned by the NIHR HTA Programme on behalf of the National Institute for Health and Care Excellence



Kleijnen Systematic Reviews Ltd in collaboration with Erasmus University Rotterdam and Maastricht University

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## **Contributions of authors**

Marie Westwood and Penny Whiting planned and performed the systematic review and interpretation of evidence. Manuela Joore, Thea van Asselt, Bram Ramaekers and Praveen Thokala

3

planned and performed the cost-effectiveness analyses and interpreted results. Nigel Armstrong contributed to planning and interpretation of cost-effectiveness analyses, acquisition of input data and conducted model peer review. Janine Ross devised and performed the literature searches and provided information support to the project. Jos Kleijnen and Johan Severens provided senior advice and support to the systematic review and cost-effectiveness analyses, respectively. All parties were involved in drafting and/or commenting on the report.

Academic in confidence information is marked

# TABLE OF CONTENTS

Table of Contents5				
List of Tables7				
List of Figures	8			
List of Abbreviations	9			
Glossary	11			
Executive Summary (2,043 words)	.12			
Plain English summary (232 words)	17			
1. Objective	18			
2. Background and definition of the decision problem(s)	.19			
2.1 Population	.19			
2.2 Intervention technologies	20			
2.2.1 Abbott ARCHITECT high-sensitivity troponin I assay	21			
2.2.2 AccuTNI+3 troponin I assay (Beckman-Coulter)	21			
2.2.3 Roche Elecsys high-sensitive troponin T assay	21			
2.3 Comparator	24			
2.4 Care pathway	24			
2.4.1 Diagnostic assessment	24			
2.4.2 Management/treatment	25			
3. Assessment of clinical effectiveness	27			
3.1 Systematic review methods	27			
3.1.1 Search strategy	27			
3.1.2 Inclusion and exclusion criteria	29			
3.1.3 Inclusion screening and data extraction	31			
3.1.4 Quality assessment	31			
3.1.5 Methods of analysis/synthesis	31			
3.2 Results of the assessment of clinical effectiveness assessment	33			
3.2.1 Overview of included studies	33			
3.2.2 Study quality	34			
3.2.3 Diagnostic accuracy of the Roche Elecsys hs-cTnT assay	38			
3.2.4 Diagnostic accuracy of the Abbott ARCHITECT hs-cTnI assay for the rule-out and diagno	osis			
of AMI	51			
3.2.5 Diagnostic accuracy of the Beckman Coulter Access hs-cTnl assay	56			
3.2.6 Comparative diagnostic accuracy of the Roche Elecsys hs-TnT assay, the Abb	ott			
ARCHITECT hs-TnI assay and the Beckman Coulter Access hs-TnI assay	59			
3.2.7 Selection of diagnostic strategies for inclusion in cost-effectiveness modeling	59			
4. Assessment of cost-effectiveness	60			
4.1 Review of economic analyses of hs-cTn assays	60			
4.1.1 Search strategy	60			
4.1.2 Inclusion criteria	60			
4.1.3 Quality assessment	61			
4.1.4 Results	61			
4.2 Model structure and methodology	76			
4.2.1 Troponin tests considered in the model	76			

4.2.2	Model structure	77		
4.2.3	Model parameters	80		
4.2.4	Overview of main model assumptions	85		
4.3	Model analyses	86		
4.3.1	Secondary analysis	86		
4.3.2	Sensitivity analysis	87		
4.3.3	Subgroup analysis	88		
4.4	Results of cost-effectiveness analyses	89		
4.4.1	Base case analysis	89		
4.4.2	Secondary analysis	93		
4.4.3	Sensitivity analysis	96		
4.4.4	Subgroup analysis	99		
5.	Discussion	102		
5.1	Statement of principal findings	102		
5.1.1	Clinical effectiveness			
5.1.2	Cost-effectiveness	104		
5.2	Strengths and limitations of assessment	107		
5.2.1	Clinical effectiveness			
5.2.2	Cost-effectiveness	110		
5.3	Uncertainties	112		
5.3.1	Clinical effectiveness	112		
5.3.2	Cost-effectiveness	114		
6.	Conclusions	116		
6.1	Implications for service provision	116		
6.2	Suggested research priorities	116		
7. References	S	118		
Appendix 1: L	iterature search strategies	146		
Clinical effectiveness search strategies				
Cost-effect	iveness searches	155		
Appendix 2: D	Data extraction tables	162		
a. Baseline study details				
b. Index test and reference standard details171				
c. Study results178				
Appendix 3: QUADAS-2 Assessments				
Appendix 4: Table of excluded studies with rationale201				
Appendix 5: Sensitivity analyses (base case)				
Appendix 6: Sensitivity analyses (secondary analysis)				
Appendix 7: Subgroup analyses (base case)				
Appendix 8: Subgroup analyses (secondary analysis)				
Appendix 9: Subgroup analyses based on accuracy and AMI prevalence (only available for the Roche				
99th centile test)245				
Appendix 10:	NICE guidance relevant to the management of supected ACS	247		
Appendix 11: PRISMA check list248				

# LIST OF TABLES

Table 1: Overview of cardiac biomarkers    23
Table 2: Inclusion criteria    30
Table 3: QUADAS-2 results for studies of hs-cTn assays
Table 4: Accuracy of the Roche hs-cTnT assay: Summary estimates (95% confidence intervals)47
Table 5: Accuracy of the Abbott ARCHITECT hs-cTnl assay: Summary estimates (95% confidence
intervals)
Table 6: Accuracy of the Beckman Coulter Access hs-cTnl assay: Summary estimates (95% confidence
intervals)
Table 7: Comparison between assays (Presentation samples, 99 <sup>th</sup> centile threshold): Sensitivity and
specificity (95% CI)
Table 8: Comparison between assays (Presentation samples, 99 <sup>th</sup> centile threshold): Likelihood
ratios (95% CI)
Table 9: Summary of included full papers69
Table 10: Checklist of study quality for full papers included74
Table 11: Transition probabilities    80
Table 12: Test accuracy81
Table 13: Test outcomes
Table 14: Utility scores
Table 15: Resource use (test specific)83
Table 16: Health state costs, event costs and unit prices
Table 17: Probabilistic results for base case analysis: life years         90
Table 18: Probabilistic results for base case analysis: costs and QALYs91
Table 19: Probabilistic results for secondary analysis: life years
Table 20: Probabilistic results for secondary analysis: costs and QALYs

# LIST OF FIGURES

Figure 1: Flow of studies through the review process
Figure 2: Summary of QUADAS-2 results for studies of hs-cTn assays
Figure 3: SROC for the Roche Elecsys hs-cTnT assay using the 99 <sup>th</sup> centile threshold and a
presentation sample (13 studies)
Figure 4: SROC for the Roche Elecsys hs-cTnT assay using the 99 <sup>th</sup> centile threshold and a
presentation sample (6 studies which excluded participants with STEMI)
Figure 5: SROC for the Roche Elecsys hs-cTnT assay using the 99 <sup>th</sup> centile threshold and a
presentation sample (8 studies with a mixed population, target condition any AMI)40
Figure 6: ROC space plot for the Roche Elecsys hs-cTnT assay using the 99 <sup>th</sup> centile threshold and a
presentation sample in different clinical subgroups42
Figure 7: ROC space plot for the Roche Elecsys hs-cTnT assay using the 99 <sup>th</sup> centile threshold and a
presentation sample in people presenting at different times after symptom onset
Figure 8: ROC space plot of the Roche Elecsys hs-cTnT assay using multiple sampling strategies 46
Figure 9: Testing pathway for the Roche Elecsys hs-cTnT assay used in cost-effectiveness modeling 50
Figure 10: ROC space plot of the Abbott ARCHITECT hs-cTnT assay53
Figure 11: Testing pathway for the Abbott ARCHITECT hs-cTnI assay used in cost-effectiveness
modeling
Figure 12: ROC space plot of the Beckman Coulter Access hs-cTnT assay
Figure 13: Flow of studies through the review process62
Figure 14: Decision tree structure79
Figure 15: Markov model structure79
Figure 16: Cost-effectiveness acceptability curve and incremental cost-effectiveness plane
(incremental costs and QALYs compared to standard troponin) for base case analysis
Figure 17: Cost-effectiveness acceptability frontier for base case analysis
Figure 18: Cost-effectiveness acceptability curve and incremental cost-effectiveness plane
(incremental costs and QALYs compared to standard troponin) for secondary analysis
Figure 19: Cost-effectiveness acceptability frontier for secondary analysis

# LIST OF ABBREVIATIONS

Technical terms and abbreviations are used throughout this report. The meaning is usually clear from the context, but a glossary is provided for the non-specialist reader.

AACC	American Association for Clinical Chemistry
ACC	American College of Cardiology
ACE	angiotensin converting enzyme
ACS	acute coronary syndrome
AHA	American Health Association
AMI	acute myocardial infarction
ARIF	aggressive Research Intelligence Facility
CAD	coronary artery disease
CADTH	Canadian Agency for Drugs and Technologies in Health
ССТ	controlled clinical trial
CDSR	Cochrane Database of Systematic Reviews
CEAC	cost-effectiveness acceptability curve
CEAF	cost-effectiveness acceptability frontier
CENTRAL	Cochrane Central Register of Controlled Trials
CHD	coronary heart disease
CI	confidence interval
CTCA	computed tomography coronary angiography
CV	co-efficient of variation
DARE	Database of Abstracts of Reviews of Effects
DTA	diagnostic test accuracy
ECG	electrocardiography/electrocardiogram
ECLIA	electrochemiluminescence immunoassay
ED	emergency department
EED	Economic Evaluations Database
ESC	European Society of Cardiology
ED	Emergency Department
FN	false negative
FP	false positive
HES	Hospital Episode Statistics
HF	heart failure
H-FABP	heart fatty acid binding protein
HES	hospital episode statistics
HRQoL	Health-Related Quality of Life
hs-cTn	high sensitivity cardiac troponin
hs-cTnl	high sensitivity cardiac troponin I
hs-cTnT	high sensitivity cardiac troponin T
HSROC	hierarchical summary receiver operating characteristic
HTA	Health technology Assessment
ICER	incremental cost-effectiveness ratio
INAHTA	International Network of Agencies for Health Technology Assessment
IQR	interquartile range

LILACS	Latin American and Caribbean Health Sciences Literature				
LoB	limit of blank				
LoD	limit of detection				
LR+	Positive likelihood ratio				
LR-	Negative likelihood ratio				
LY	life year				
MACE	major adverse cardiac event				
MI	myocardial infarction				
NA	not applicable				
NHS	National Health Service				
NICE	National Institute for Health and Care Excellence				
NIH	National Institutes of Health				
NIHR	National Institute for Health Research				
NPV	negative predictive value				
NR	not reported				
NSTE-ACS	non-ST-segment-elevation ACS				
NSTEMI	non-ST segment elevation myocardial infarction				
ONS	Office for National Statistics				
PSA	probabilistic sensitivity analysis				
QALY	Quality-Adjusted Life Year				
RCT	randomised controlled trial				
ROC	receiver operating characteristic				
SCI	Science Citation Index				
SIGN	Scottish Intercollegiate Guidelines Network				
SROC	summary receiver operating characteristic				
STEMI	ST segment elevation myocardial infarction				
Tn	troponin				
TN	true negative				
ТР	true positive				
UA	unstable angina				
WHF	World Heart Federation				

Cost-effectiveness	An economic analysis that converts effects into health terms and describes the		
analysis	costs for additional health gain.		
Decision modelling	A mathematical construct that allows the comparison of the relationship		
	between costs and outcomes of alternative healthcare interventions.		
False negative	Incorrect negative test result – number of diseased persons with a negative test		
	result.		
False positive	Incorrect positive test result – number of non-diseased persons with a positive		
	test result.		
Incremental cost-	The difference in the mean costs of two interventions in the population of		
effectiveness ratio	interest divided by the difference in the mean outcomes in the population of		
(ICER)	interest.		
Index test	The test whose performance is being evaluated.		
Likelihood Ratio	Likelihood ratios describe how many times more likely it is that a person with the		
(LR)	target condition will receive a particular test result than a person without the		
	target condition.		
Markov model	An analytic method particularly suited to modelling repeated events, or the		
	progression of a chronic disease over time.		
Meta-analysis	Statistical techniques used to combine the results of two or more studies and		
-	obtain a combined estimate of effect.		
Meta-regression	Statistical technique used to explore the relationship between study		
	characteristics and study results.		
Opportunity costs	The cost of forgone outcomes that could have been achieved through alternative		
	investments.		
Publication bias	Bias arising from the preferential publication of studies with statistically		
	significant results.		
Quality of life	An individual's emotional, social and physical well-being and their ability to		
	perform the ordinary tasks of living.		
Quality-adjusted	A measure of health gain, used in economic evaluations, in which survival		
life year (QALY)	duration is weighted or adjusted by the patient's quality of life during the survival		
	period.		
Receiver Operating	A graph which illustrates the trade-offs between sensitivity and specificity which		
Characteristic	result from varying the diagnostic threshold.		
(ROC) curve			
Reference standard	The best currently available method for diagnosing the target condition. The		
	index test is compared against this to allow calculation of estimates of accuracy.		
Sensitivity	Proportion of people with the target disorder who have a positive test result.		
Specificity	Proportion of people without the target disorder who have a negative test result.		
True negative	Correct negative test result – number of non-diseases persons with a negative		
0	test result.		
True positive	Correct positive test result – number of diseased persons with a positive test		
	result.		

# GLOSSARY

# **EXECUTIVE SUMMARY (2,043 WORDS)**

# Background

The primary indication for this assessment is the early rule-out of acute myocardial infarction (AMI) in people presenting with acute chest pain and suspected, but not confirmed, non-ST segment elevation myocardial infarction (NSTEMI).

Cardiac troponins (Tn) I and T are used as markers of AMI. They are intended for use in conjunction with clinical history taking and electrocardiography monitoring. Elevated troponin levels are associated with an increased risk of adverse cardiac outcomes. However, the optimal sensitivity of standard troponin assays for AMI occurs several (10-12) hours after the onset of symptoms. Two high-sensitivity troponin (hs-cTn) assays are currently available for use in the NHS in England and Wales, ARCHITECT high-sensitivity troponin I assay (Abbott Diagnostics) and the Elecsys troponin T high-sensitive assay (Roche).One additional assay, AccuTNI+3 troponin I assay (Beckman-Coulter), was included in the scope for this assessment pending CE marking; CE marking has now been confirmed. These are able to detect lower levels of troponin in the blood with analytical sensitivities up to 100 times greater than conventional troponin assays. Use of high sensitivity assays enables the detection of small changes in troponin levels and may enable AMI to be ruled out at an earlier time after the onset of acute chest pain.

This assessment considers hs-cTn assays used singly or in series, up to four hours after the onset of chest pain or up to four hours after presentation; for serial troponin measurements, both data on change in troponin levels and peak troponin are considered.

# Objectives

To assess the clinical- and cost-effectiveness of high sensitivity troponin assays for the management of adults presenting with acute chest pain, in particular for the early (within four hours of presentation) rule-out of AMI.

#### Methods

# Assessment of clinical effectiveness

Sixteen databases, including MEDLINE and EMBASE, research registers and conference proceedings were searched to October 2013. Search results were screened for relevance independently by two reviewers. Full text inclusion assessment, data extraction, and quality assessment were conducted by one reviewer and checked by a second. Study quality was assessed using QUADAS-2. The bivariate/hierarchical summary receiver operating characteristic (HSROC) model was used to estimate summary sensitivity and specificity with 95% confidence intervals (CIs) and prediction

regions around the summary points, and to derive hierarchical summary receiver operating characteristic curves for meta-analyses involving four or more studies. For meta-analyses with fewer than four studies we estimated separate pooled estimates of sensitivity and specificity, using random-effects logistic regression. Summary positive and negative likelihood ratios were derived from the summary estimates of sensitivity and specificity. Analyses were conducted separately for each of the three hs-cTn assays and were stratified according to whether the study evaluated the prediction of AMI or major adverse cardiac event (MACE), test timing, and the threshold used to define a positive hs-cTn result. Stratified analyses were used to investigate heterogeneity and the influence of risk of bias on summary estimates.

## Assessment of cost-effectiveness

We considered the long-term costs and quality adjusted life years (QALYs) associated with different troponin testing methods, to diagnose or rule-out NSTEMI, for patients presenting at the emergency department (ED) with suspected non-ST-segment-elevation acute coronary syndrome (NSTE-ACS). The de novo model consisted of a decision tree and a Markov model. The decision tree was used to model the 30 day outcomes after presentation, based on test results and the accompanying treatment decision. The long-term consequences in terms of costs and QALYs were estimated using a Markov cohort model with a lifetime time horizon (60 years). The following strategies were included in the main economic analysis:

- Standard troponin at presentation and at 10-12 hours (reference standard)
- Roche Elecsys hs-cTnT at presentation: 99<sup>th</sup> centile threshold
- Roche Elecsys hs-cTnT (optimal strategy): LoB threshold at presentation followed by 99<sup>th</sup> centile threshold peak within three hours and/or Δ20% (compared to presentation test) at 1-3 hours
- Abbott ARCHITECT hs-cTnI at presentation: 99<sup>th</sup> centile threshold
- Abbott ARCHITECT hs-cTnI (optimal strategy): LoD threshold at presentation, followed by 99<sup>th</sup> centile threshold at three hours
- Beckman Coulter hs-cTnI at presentation: 99<sup>th</sup> centile threshold

In the base case, it was assumed that standard troponin testing had perfect sensitivity and specificity (reference case) for diagnosing AMI and that only patients testing positive on the reference standard (standard troponin), were at increased risk for adverse events and would benefit from immediate treatment. In a secondary analysis, a proportion of patients testing positive on an hs-cTn test were treated accordingly. These patients were assumed to be treated for the hs-cTn assays and left untreated for the standard troponin test and at increased risk for adverse events. In addition, a

number of sensitivity and subgroup analyses were performed.

#### Results

#### Assessment of clinical effectiveness

Eighteen studies (38 publications) were included in the review. The main potential sources of bias in the included studies related to patient spectrum and patient flow. There were also concerns regarding the applicability of the patient population and the reference standard in some of the included studies.

# Diagnostic accuracy of the Roche Elecsys hs-cTnT assay (15 studies)

The most commonly evaluated testing strategy was the 99<sup>th</sup> centile threshold in a blood sample taken on presentation. Studies (n=six) that excluded patients with ST segment elevation myocardial infarction (STEMI) gave a summary positive likelihood ratio (LR+) of 5.41 (95% CI: 3.40 to 8.63) and summary negative likelihood ratio (LR-) of 0.15 (0.08, 0.26) for this strategy. Estimates were similar when derived from all studies (n=13) that evaluated this strategy. The optimum strategy based on this assay appeared to be one based on the combination of a limit of blank (LoB) threshold in a presentation sample which could be used to rule out AMI (LR- 0.10, 95% CI: 0.05 to 0.18) but has limited potential to rule in an AMI (LR+ 1.83, 95% CI: 1.70, 1.97). Patients testing positive could then have a further sample taken at two hours, a result above the 99<sup>th</sup> centile on either the presentation or two hour sample and a delta of at least 20% has some potential for ruling in an AMI (LR+ 8.42, 95% CI: 6.11 to 11.60) while a result below the 99<sup>th</sup> centile on both samples and a delta less than 20% can be used to rule out an AMI (LR- 0.04, 95% CI: 0.02 to 0.10).

## Diagnostic accuracy of the Abbott ARCHITECT hs-cTnl assay (four studies)

Three studies, all conducted in populations that included patients with STEMI, evaluated this assay at the 99<sup>th</sup> centile threshold in a blood sample taken on presentation. The summary LR+ was 11.47 (95% CI: 9.04 to 16.19) and the summary LR- was 0.22 (95% CI: 0.16 to 0.27). The optimum strategy appeared to be one based on the combination of a limit of detection (LoD) threshold in a presentation sample which could be used to rule out AMI (LR- 0.01, 95% CI: 0.00 to 0.08) but has limited potential to rule in an AMI (LR+ 1.54, 95% CI: 1.47 to 1.62). Patients testing positive could then have a further sample taken at three hours, a result above the 99<sup>th</sup> centile on this sample has some potential for ruling in an AMI (LR+ 10.16, 95% CI: 8.38 to 12.31) while a result below the 99<sup>th</sup> centile can be used to rule out an AMI (LR- 0.02, 95% CI: 0.01 to 0.05).

#### Diagnostic accuracy of the Beckman Coulter Access hs-cTnl (two studies)

One study, conducted in a population that included patients with STEMI, evaluated this assay at the

99<sup>th</sup> centile threshold in a blood sample taken on presentation. The summary LR+ was 3.67 (95% CI: 3.26 to 4.13) and the summary LR- was 0.11 (95% CI: 0.07 to 0.17). Data were not reported for the LoB/LoD threshold. There were insufficient data to determine the optimum testing strategy for this assay.

#### Assessment of cost-effectiveness

#### Base case analysis

In the base case analysis, standard troponin testing was both most effective and most costly. Strategies considered cost-effective depending upon ICER thresholds were Abbott ARCHITECT hscTnI 99<sup>th</sup> centile (thresholds below £6,597), Beckman Coulter hs-cTnI 99<sup>th</sup> centile (thresholds between £6,597 and £30,042), Abbott ARCHITECT hs-cTnI optimal strategy (LoD threshold at presentation, followed by 99<sup>th</sup> centile threshold at three hours) (thresholds between £30,042 and £103,194), and the standard troponin test (thresholds over £103,194). The Roche Elecsys hs-cTnT 99<sup>th</sup> centile and the Roche Elecsys hs-cTnT optimal strategy (LoB threshold at presentation followed by 99<sup>th</sup> centile threshold and/or  $\Delta$ 20% (compared to presentation test) at 1-3 hours) were extendedly dominated in this analysis (one of the more effective strategies was better value in that the ICER was lower).

#### Secondary analysis

In the secondary analysis, which assumed that a proportion of false positives in the hs-cTn testing strategies had an increased risk of adverse events, standard troponin was least effective and most costly, and therefore a dominated strategy. The most effective strategy here was the Abbott ARCHITECT hs-cTnI optimal strategy. The Roche Elecsys hs-cTnT optimal strategy was extendedly dominated (one of the more effective strategies was better value in that the ICER was lower), as was the Beckman Coulter hs-cTnI 99<sup>th</sup> centile in this analysis. Strategies considered cost-effective were Abbott ARCHITECT hs-cTnI 99<sup>th</sup> centile (thresholds below £12,217), Roche Elecsys hs-cTnT 99<sup>th</sup> centile (thresholds below £12,217).

# Sensitivity and subgroup analyses

Sensitivity analyses showed that assumptions regarding the difference between treated and untreated patients (e.g. mortality rate, risk of re-infarction) had the largest impact on relative cost-effectiveness, as well as whether or not patients testing false positive were assigned treatment costs. In general, the base case analysis was affected more by varying these assumptions than the secondary analysis. Results from the subgroup analyses led to the conclusion that hs-cTn testing is likely to be more cost-effective in younger population, in populations with pre-existing coronary

artery disease (CAD), and for patients whose symptom onset was less than three hours ago. A no testing strategy can only be considered cost-effective in populations with a prevalence as low as 1%.

#### Conclusions

#### Implications for service provision

There is evidence to suggest that undetectable levels of Tns (below the LoB/LoD of the assay) on presentation, measured using the Roche Elecsys hs-cTnT assay or the Abbott ARCHITECT hs-cTnI assay, may be sufficient to rule out NSTEMI in people presenting with symptoms suggestive of ACS. There is also evidence to suggest that, for the Roche Elecsys hs-cTnT assay and the Abbott ARCHITECT hs-cTnI assay, a further rule-out step may be possible within the four hour NHS emergency department target. There is insufficient evidence to determine an optimum testing strategy for the Beckman Coulter hs-cTnI assay. There is some limited evidence to suggest that a Tn level below the 99<sup>th</sup> centile on presentation, measured using the Roche Elecsys hs-cTnT assay, may be sufficient to rule out NSTEMI in some groups (people over 70 years old, people without pre-existing CAD and people with a clinically determined high pre-test probability).

The economic model does not provide strong evidence to prefer one hs-cTn testing strategy over another. Results do, however, indicate that hs-cTn testing in general may be cost-effective compared to standard troponin testing given that hs-cTn testing leads to cost-saving at a QALY loss. This becomes more likely if one assumes that hs-cTn testing detects some patients who require treatment despite their testing negative with standard troponin, as shown in the secondary analysis hs-cTn testing. In particular, the Abbott ARCHITECT hs-cTnI optimal strategy, which involves multiple testing and varying cutoff levels, may be promising. The main issue, with regard to service provision, if implementation of an hs-cTn testing strategy is considered, is the balance between the likely reduction in cost and the risk of a reduction in effectiveness, albeit possibly small.

## Suggested research priorities

New studies are needed to fully evaluate the performance of our proposed optimal testing strategies in a clinical setting. Further research (diagnostic cohort studies or multivariable prediction modelling studies) is needed to fully explore possible variation in the performance of hs-cTn assays and the optimal testing strategies for these assays in relevant demographic and clinical subgroups (sex, age, ethnicity, renal function, previous CAD, previous AMI) and to investigate the effects of clinical judgement (assessment of pre-test probability) on test performance. As most of the uncertainties in the economic model were caused by assumptions relating to clinical effectiveness, this type of research would also facilitate economic analyses of hs-cTn testing.

# PLAIN ENGLISH SUMMARY (232 WORDS)

Heart disease is a leading cause of death in the UK, with myocardial infarction (MI) (heart attack) accounting for approximately 5% of all deaths recorded in 2011. Many people attend hospital with chest pain and suspected MI; chest pain has been reported as the most common cause of hospital admissions in the UK and 2011-2012 statistics showed that it accounted for approximately 5% of all emergency admissions. It is important to diagnose people who are suspected of having an MI as early as possible in order to ensure quick and effective treatment. However, only around 20% of emergency admissions for chest pain will actually have an MI and there are many other possible causes of chest pain (e.g. gastro-oesophageal disorders, muscle pain, anxiety, or stable ischaemic heart disease). Tests which can quickly tell which patients do not have MI could therefore avoid unnecessary hospital admissions and anxiety for many people.

This assessment aimed to determine the clinical- and cost-effectiveness of high sensitivity troponin tests, used as single tests or repeated over a short time, for diagnosing or ruling-out MI in people who present to hospital with chest pain. We found that high sensitivity troponin tests may be able to rule-out MI within the four hour NHS emergency department target. Health economic analyses indicated that high sensitivity tests may be cost-effective compared to standard troponin tests, which require repeat testing at 10-12 hours.

## 1. OBJECTIVE

The overall objective of this project is to assess the clinical- and cost-effectiveness of high sensitivity troponin (Tn) assays for the management of adults presenting with acute chest pain, in particular for the early (within four hours of presentation) rule-out of acute myocardial infarction (AMI). The following research questions were defined to address the review objectives:

- What is the clinical effectiveness of new, high sensitivity troponin (hs-cTn) assays (used singly or in series) compared with conventional diagnostic assessment, for achieving early discharge within four hours of presentation, where AMI is excluded without increase in adverse outcomes?
- What is the accuracy of new, hs-cTn assays (used singly or in series, such that results are available within three hours of presentation) for the diagnosis of AMI in adults with acute chest pain?
- What is the accuracy of new, hs-cTn assays (used singly or in series, such that results are available within three hours of presentation), for the prediction of major adverse cardiac events (MACE) (cardiac death, non-fatal AMI, revascularisation, or hospitalisation for myocardial ischemia) during 30 day follow-up in adults with acute chest pain?
- What is the cost-effectiveness of using new, hs-cTn assays (used singly or in series, such that results are available within three hours of presentation), compared with the current standard of serial Tn T and/or I testing on admission and at 10-12 hours post-admission?

# 2. BACKGROUND AND DEFINITION OF THE DECISION PROBLEM(S)

## 2.1 Population

The primary indication for this assessment is the early rule-out of AMI and consequent early discharge in people presenting with acute chest pain and suspected, but not confirmed, non-ST segment elevation myocardial infarction (NSTEMI). The assessment will also consider the potential effects of early diagnosis of AMI and of reduced specificity of testing.

Acute coronary syndrome (ACS) is the term used to describe a spectrum of conditions caused by coronary artery disease (CAD) (also known as coronary heart disease or ischaemic heart disease). ACS arises when atheromatous plaque ruptures or erodes leading to vasospasm, thrombus formation and distal embolization, obstructing blood flow through the coronary arteries. It incorporates three distinct conditions: unstable angina, ST segment elevation myocardial infarction (STEMI) and NSTEMI. CAD and AMI are a significant health burden in the UK, with Office for National Statistics (ONS) mortality data for 2011 showing 23,705 deaths from AMI and 64,435 deaths from ischaemic heart disease; AMI accounted for approximately 5% of all deaths recorded in 2011 and ischaemic heart disease accounted for approximately 13%.<sup>1</sup>

People with ACS usually present with chest pain and chest pain has been reported as the most common cause of hospital admissions in the UK;<sup>2</sup> Hospital Episode Statistics (HES) for 2011-2012 show 243,197 emergency admissions for chest pain, accounting for approximately 5% of all emergency admissions.<sup>3</sup> However, many people presenting with acute chest pain will have non-cardiac underlying causes, such as gastro-oesophageal disorders, muscle pain, anxiety, or stable ischaemic heart disease. A 2003 study on the impact of cardiology guidelines on the diagnostic classification of people with ACS in the UK reported that the majority of people admitted to hospital with chest pain have either no ischaemic heart disease or stable ischaemic heart disease.<sup>4</sup> HES for 2011-2012 are consistent with this observation, showing diagnoses of AMI in 47,783 emergency admissions and unstable angina in 32,369 admissions; this represents approximately 20% and 13% of emergency admissions with chest pain, respectively.<sup>3</sup> Accurate and prompt differentiation of ACS (in particular AMI), stable CAD and other causes of chest pain is therefore vital to ensure appropriate and timely intervention where required and to avoid unnecessary hospital admissions.

STEMI can usually be diagnosed on presentation by electrocardiogram (ECG), hence the main diagnostic challenge in the investigation of suspected ACS is the detection or rule-out of NSTEMI. Investigation of ACS can also involve identification of people with unstable angina (CAD with

worsening symptoms, but no evidence of myocardial necrosis).

Since the development of protein biomarkers of myocardial damage in the 1980s, the number of biomarker assays available has proliferated, cardiac specificity has increased, and the role of biomarkers in the diagnostic work-up of acute chest pain has expanded. Cardiac biomarkers are becoming increasingly sensitive and recent European Society of Cardiology (ESC) and American College of Cardiology (ACC) guidelines enable AMI to be diagnosed with any rise and/or fall of Tn to above the laboratory reference range.<sup>5, 6</sup> This has resulted in fewer people being classified as having unstable angina with no myocardial damage and more people being classified as having NSTEMI.<sup>7</sup> The most recent two years of HES show that the number of Emergency Department (ED) attendances where the first recorded investigation was a cardiac biomarker rose from 13,743 in 2010-2011 to 28,379 in 2011-2012.<sup>3</sup> Cardiac troponins I and T (cTnI and cTnT), together with cardiac troponin C, form the troponin-tropomyosin complex which is responsible for regulating cardiac muscle contraction. cTnI and cTnT are used clinically as markers of cardiomyocyte necrosis, indicative of AMI. Troponin assays are intended for use in conjunction with clinical history taking and ECG monitoring as, although specificity is high, troponins may also be elevated in many other conditions including myocarditis, congestive heart failure, severe infections, renal disease and chronic inflammatory conditions of the muscle or skin. Standard biochemical diagnosis of NSTEMI is based on elevation of the cardiac biomarker Tn above the 99<sup>th</sup> percentile of the reference range for the normal population.<sup>8</sup> Elevated Tn levels have been shown to be associated with an increased risk of adverse cardiac outcomes.<sup>9</sup> However, the optimal sensitivity of standard Tn assays for AMI occurs several hours after the onset of symptoms;<sup>10</sup> this is reflected in current clinical guidelines, which recommend cTnI or cTnT testing at initial hospital assessment and again 10-12 hours after the onset of symptoms.<sup>11, 12</sup> Since the majority of people presenting with chest pain do not have NSTEMI, where presentation is within a few hours of symptom onset delayed biomarker measurement may result in unnecessary periods of extended observation or hospitalisation and associated costs. The development of cardiac biomarkers which can be used at an earlier stage without reduction in sensitivity is, therefore, desirable.

# 2.2 Intervention technologies

The development of high-sensitivity cTn (hs-cTn) assays means that it is possible to detect lower levels of Tn in the blood. Current generations of commercially available assays have analytical sensitivities up to 100 times greater than was the case for early Tn assays (1 ng/L versus 100 ng/L).<sup>13</sup> Use of these high-sensitivity assays enable the detection of small changes in cTn levels, and may enable AMI to be ruled out at an earlier time after the onset of acute chest pain. Use of the hs-cTn

assays has the potential to facilitate earlier discharge for people with normal cTn levels and earlier intervention for those with elevated levels of cTn. The recommended definition of an hs-cTn assay uses two criteria:<sup>13, 14</sup>

- The total imprecision, co-efficient of variation (CV), of the assay should be ≤10% at the 99<sup>th</sup> percentile value for the healthy reference population.
- The limit of detection (LoD) of the assay should be such as to allow measurable concentrations to be attainable for at least 50% (ideally >95%) of healthy individuals.

A number of high-sensitivity cTnI and cTnT (hs-cTnI and hs-cTnT) assays are currently available for use in the NHS in England and Wales; all are designed for use in clinical laboratory settings.

# 2.2.1 Abbott ARCHITECT high-sensitivity troponin I assay

The ARCHITECT hs-cTnI STAT assay can be used with the Abbott ARCHITECT i2000SR and i1000SR analysers. The assay is a quantitative, chemiluminescent micro particle immunoassay (CMIA) for serum or plasma samples. Results are available within 16 minutes. The ARCHITECT hs-cTnI STAT assay can detect cTnI in 96% of the reference population, and has a recommended 99<sup>th</sup> percentile cut-off of 26.2ng/L with a CV of 4%.<sup>15</sup> The assay is CE marked and available to the NHS.

## 2.2.2 AccuTNI+3 troponin I assay (Beckman-Coulter)



# 2.2.3 Roche Elecsys high-sensitive troponin T assay

The Elecsys cTnT-hs and Elecsys cTnT-hs STAT assays can be used on the Roche Elecsys 2010 analyser and the cobas Modular Analytics e series immunoassay analysers. The assay is a quantitative, sandwich electrochemiluminescence immunoassay (ECLIA) for serum and plasma samples. Results are available within 18 minutes with the standard assay and within nine minutes if the STAT assay is used. Both versions of the assay can detect cTnT in 61% of the reference population and have a recommended 99<sup>th</sup> percentile cut off of 14ng/L with a CV of <10%.<sup>18</sup> Both versions of the assay are CE marked and available to the NHS.

A summary of the product properties of hs-cTnI and hs-cTnT assays available as in the NHS in England and Wales is provided in Table 1.

hs-cTn assays can be used as single diagnostic tests, or in combination with other cardiac biomarkers, e.g. heart fatty acid binding protein (H-FABP) and copeptin. The use of combinations of cardiac biomarkers may increase sensitivity, where a positive result on either test is considered to be indicative of AMI, although this increase may be achieved at the expense of decreased specificity. Conversely, if a positive result on both tests is required before AMI is diagnosed, increased specificity and reduced sensitivity are likely. It is currently unclear which, if any, of the available cardiac biomarkers could add clinical benefit if used in combination with hs-cTnI and hs-cTnT, compared to hs-cTnI and hs-cTnT alone. A recent systematic review reported some data for combination testing, but none of the identified studies of Tns combined with other biomarkers used high sensitivity methods.<sup>7</sup> Retrospective analysis of data from one arm of a randomised controlled trial by the same authors provided some indication that the use of H-FABP in combination with hscTn, on admission, may increase sensitivity for AMI without decreasing specificity.<sup>19</sup> This increase was equivalent to the sensitivity achieved by serial hs-cTn testing on admission and at 90 minutes.<sup>19</sup> However, these tests are not readily available for analytical platforms in routine use in the NHS and discussions at the scoping stage of this assessment concluded that practical applications of H-FABP and copeptin assays and evidence for their effectiveness are not yet sufficiently developed to justify their inclusion.

This assessment will consider hs-cTn assays used singly or in series, up to four hours after the onset of chest pain or up to four hours after presentation (as reported); for serial Tn measurements, both data on change in Tn levels and peak Tn will be considered (as reported).

# Table 1: Overview of cardiac biomarkers

Manufacturer	System	Assay	LoD (ng/L)	LoB (ng/L)	99 <sup>th</sup> percentile (ng/L) <sup>*</sup>	CV at 99 <sup>th</sup> percentile <sup>*</sup>	Turnaround time (mins) <sup>*</sup>	CE marked
Abbott Diagnostics	ARCHITECT	STAT hs-cTnl	1.1 to 1.9	0.7 to 1.3	26.2	4%	16	~
Beckman Coulter	Access and UniCel Dxl	AccuTnl+3	10	<10	40.0	<10%	13	√
Roche	Elecsys	cTnT-hs	5	3	14	<10%	18	✓
Roche	Elecsys	cTnT-hs STAT	5	3	14	<10%	9	$\checkmark$

\* Information supplied to NICE by the manufacturer

LoD: limit of detection

LoB: limit of blank

#### 2.3 Comparator

The comparator for this technology appraisal is the current UK standard of serial TnT and/or I testing (using any method not defined as a hs-cTn test) on admission and at 10-12 hours after the onset of symptoms.<sup>11</sup>

#### 2.4 Care pathway

# 2.4.1 Diagnostic assessment

The assessment of patients with suspected ACS is described in NICE clinical guideline 95 (CG95) "Chest pain of recent onset: Assessment and diagnosis of recent onset chest pain or discomfort of suspected cardiac origin."<sup>11</sup> The guideline specifies that initial assessment should include a resting 12-lead ECG along with a clinical history, a physical examination and biochemical marker analysis. For people in whom a regional ST-segment elevation or presumed new left branch bundle block is seen on ECG, management should follow NICE clinical guideline 167 (CG167) "The acute management of AMI with ST-segment elevation."<sup>20</sup> People without persistent ST-elevation changes on ECG, i.e. with suspected non-ST-segment-elevation ACS (NSTE-ACS), should receive further investigation using cardiac biomarkers with the aim of distinguishing NSTEMI from unstable angina. NICE CG95 makes the following recommendations on the use of cardiac biomarkers:<sup>11</sup>

- Take a blood sample for cTnI or cTnT on initial assessment in hospital. These are the preferred biochemical markers to diagnose AMI.
- Take a second blood sample for cTnI or cTnT measurement 10-12 hours after the onset of symptoms.
- Do not use biomarkers such as natriuretic peptides and high sensitivity C-reactive protein to diagnose an ACS.
- Do not use biomarkers of myocardial ischemia (such as ischemia modified albumin) as opposed to markers of necrosis when assessing people with acute chest pain.
- Take into account the clinical presentation, from the time of onset of symptoms and the resting 12-lead ECG findings, when interpreting Tn measurements.

CG95 recommends that a diagnosis of NSTEMI should be made using the universal definition of AMI.<sup>8</sup> However, the third universal definition of AMI has been up-dated since the publication of CG95.<sup>21</sup> The most recent version states that AMI is defined as "The detection of a rise and/or fall of cardiac biomarker values (preferably cardiac Tn) with at least one value above the 99<sup>th</sup> percentile upper reference limit and with at least one of the following: symptoms of ischemia, new or presumed new significant ST-segment-T wave changes or new left branch bundle block, development of pathological Q waves in the ECG, imaging evidence of new loss of viable

myocardium or new regional wall motion abnormality, or identification of an intracoronary thrombus by angiography or autopsy."

The Scottish Intercollegiate Guidelines Network guideline 93 (SIGN 93) provides similar recommendations on the diagnostic work-up of people with suspected ACS, stating that:<sup>12</sup>

- Immediate assessment with a 12-lead ECG
- Repeat 12-lead ECG if there is diagnostic uncertainty or change in clinical status, and at discharge
- Serum Tn measurement on arrival at hospital
- Repeat serum Tn measurement 12 hours after the onset of symptoms
- Troponin concentrations should not be interpreted in isolation, but with regard to clinical presentation

Guidelines from the ESC on the diagnostic assessment of people with a suspected NSTE-ACS are consistent with those of NICE and SIGN, but additionally acknowledge the use of high-sensitivity Tn assays and make recommendations on a fast track rule out protocol. The guidelines state that hs-cTn assays have a negative predictive value (NPV) of greater 95% for AMI on admission; including a second sample of hs-cTn at three hours can increase this to 100%.<sup>22</sup>

#### 2.4.2 Management/treatment

NICE clinical guideline 94 (CG94) provides recommendations on the management of people with suspected NSTE-ACS "Unstable angina and NSTEMI: The early management of unstable angina and non-STEMI."<sup>23</sup> The guideline states that initial treatment should include a combination of antiplatelet (aspirin, clopidogrel and glycoprotein IIb/IIIa inhibitors) and antithrombin therapy, and should take into account contraindications, risk factors and the likelihood of percutaneous coronary intervention. SIGN 93 makes similar recommendations.<sup>12</sup> It is recommended that people with a diagnosis of NSTEMI, who are assessed as being at low risk of future complications, receive conservative treatment with aspirin and/or clopidogrel, or aspirin in combination with ticagrelor. People at a higher risk of future complications should be offered coronary angiography (within 96 hours of admission) with subsequent coronary revascularisation by percutaneous coronary intervention or coronary artery bypass grafting where indicated.<sup>23</sup> Additional testing to quantify inducible ischemia may also be used, before discharge, to identify those who may need further intervention<sup>23</sup> and SIGN 93 also recommends functional testing to identify people at higher risk.<sup>12</sup> SIGN 93 states that people in whom an elevated Tn level is not observed may be discharged for further follow up according to clinical judgement and, in some cases, the results of ischemia testing.<sup>12</sup>

Longer term follow-up of people who have had an AMI is described in full in NICE Clinical Guideline 48 (CG48) "Secondary prevention in primary and secondary care for patients following a myocardial infarction". This includes recommendations on lifestyle changes, cardiac rehabilitation programmes, drug therapy (including a combination of angiotensin converting enzyme (ACE) inhibitors, aspirin, beta-blockers and statins), and further cardiological assessment to determine whether coronary revascularisation is required.<sup>24</sup>

# 3. ASSESSMENT OF CLINICAL EFFECTIVENESS

A systematic review was conducted to summarise the evidence on the clinical effectiveness of hscTn assays for the early rule-out or diagnosis of AMI in people with acute chest pain. Systematic review methods followed the principles outlined in the Centre for Reviews and Dissemination (CRD) guidance for undertaking reviews in health care<sup>25</sup> and NICE Diagnostics Assessment Programme manual.<sup>26, 27</sup>

# 3.1 Systematic review methods

# 3.1.1 Search strategy

Search strategies were based on intervention (high-sensitivity Tn assays) and target condition, as recommended in the Centre for Reviews and Dissemination (CRD) guidance for undertaking reviews in health care<sup>25</sup> and the Cochrane Handbook for Diagnostic Test Accuracy Reviews.<sup>27</sup>

Candidate search terms were identified from target references, browsing database thesauri (e.g. Medline MeSH and Embase Emtree), existing reviews identified during the rapid appraisal process and initial scoping searches. These scoping searches were used to generate test sets of target references, which informed text mining analysis of high-frequency subject indexing terms using Endnote reference management software. Strategy development involved an iterative approach testing candidate text and indexing terms across a sample of bibliographic databases, aiming to reach a satisfactory balance of sensitivity and specificity.

The following databases were searched for relevant studies from 2005 to October 2013:

- MEDLINE (OvidSP): 2005-2013/10/wk1
- MEDLINE In-Process Citations and Daily Update (OvidSP): up to 2013/10/1
- EMBASE (OvidSP): 2005-2013/10/10
- Cochrane Database of Systematic Reviews (CDSR) (Wiley): Cochrane Library Issue 10 2005-2013/10/11
- Cochrane Central Register of Controlled Trials (CENTRAL) (Wiley): Cochrane Library Issue 9 2005-2013/10/11
- Database of Abstracts of Reviews of Effects (DARE) (Wiley): Cochrane Library Issue 3 2005-July 2013
- Health Technology Assessment Database (HTA) (Wiley): Cochrane Library Issue 3 2005-July 2013
- Science Citation Index (SCI) (Web of Science): 2005-2013/10/14
- Conference Proceedings Citation Index Science (CPCI) (Web of Science): 2005-2013/10/14

 LILACS (Latin American and Caribbean Health Sciences Literature) (Internet): 2005-2013/10/11

http://regional.bvsalud.org/php/index.php?lang=en

 International Network of Agencies for Health Technology Assessment (INAHTA) Publications (Internet): 2005-2013/10/15

http://www.inahta.org/

- Biosis Previews (Web of Knowledge): 2005-2013/10/11
- NIHR Health Technology Assessment Programme (Internet): 2005-2013/10/14
- Aggressive Research Intelligence Facility (ARIF) database (Internet) : 2005-2013/10/16
   <a href="http://www.birmingham.ac.uk/research/activity/mds/projects/HaPS/PHEB/ARIF/index.aspx">http://www.birmingham.ac.uk/research/activity/mds/projects/HaPS/PHEB/ARIF/index.aspx</a>
- MEDION database (Internet): 2005-2013/10/16
   <a href="http://www.mediondatabase.nl/">http://www.mediondatabase.nl/</a>
- PROSPERO (International Prospective Register of Systematic Reviews) (Internet): up to 2013/10/10 http://www.crd.york.ac.uk/prospero/

Completed and on-going trials were identified by searches of the following resources (2005-present):

- NIH ClinicalTrials.gov (Internet): up to 2013/10/1 http://www.clinicaltrials.gov/
- Current Controlled Trials (Internet): up to 2013/10/10 http://www.controlled-trials.com/
- WHO International Clinical Trials Registry Platform (ICTRP) (Internet): up to 2013/10/10 http://www.who.int/ictrp/en/

No restrictions on language or publication status were applied. Date restrictions were applied based on expert advice on the earliest appearance of literature of high sensitivity Tn assays. Searches took into account generic and other product names for the intervention. The main EMBASE strategy for each set of searches was independently peer reviewed by a second Information Specialist, using the CADTH Peer Review Checklist.<sup>28</sup> Search strategies were developed specifically for each database and the keywords associated with high sensitivity Tn T/I were adapted according to the configuration of each database. Full search strategies are reported in Appendix 1.

Electronic searches were undertaken for the following conference abstracts (selected based on advice from expert committee members):

 American Heart Association (AHA) Scientific Sessions (Internet): 2009-2013 <u>http://my.americanheart.org/professional/Sessions/ScientificSessions/Archive/Archive-Scientific-Sessions\_UCM\_316935\_SubHomePage.jsp</u>

- American Association for Clinical Chemistry (AACC) (Internet): 2009-2013
   <u>http://www.aacc.org/resourcecenters/meet\_abstracts\_archive/abstracts\_archive/annual\_m\_eeting/Pages/default.aspx#</u>
- European Society of Cardiology (ESC) (Internet): 2009-2013 http://spo.escardio.org/abstract-book/search.aspx

Identified references were downloaded in Endnote X4 software for further assessment and handling. References in retrieved articles were checked for additional studies. The final list of included papers were checked on PubMed for retractions, errata and related citations.<sup>29-31</sup>

# 3.1.2 Inclusion and exclusion criteria

Inclusion criteria for each of the clinical effectiveness questions are summarised in Table 2. Studies which fulfilled these criteria were eligible for inclusion in the review.

#### Table 2: Inclusion criteria

Question	What is the accuracy of hs-cTn assays (used singly or in	What is the effectiveness of hs-cTn assays (used singly or in			
	series, such that results are available within 3 hours of	series) compared with conventional diagnostic assessment, for			
	presentation) for the diagnosis of AMI in adults with acute	achieving successful early discharge of adults with acute chest			
	chest pain?	pain within 4 hours of presentation?			
Participants:	Adults (≥18 yrs) presenting with acute 'pain, discomfort or pressure in the chest, epigastrium, neck, jaw, or upper limb without an apparent non-cardiac source' <sup>32</sup> due to a suspected.				
	but not proven, AMI				
Setting:	Seconda	ry or tertiary care			
Interventions (index test):	Any hs-cTnT or hs-cTnI test, listed in Table 1, hs-cTn assays (used singly or in series, such that results were available within 3 hours of				
	presentation)				
Comparators:	Any other hs-cTn test, as specified above, or no comparator	Troponin T or I measurement on presentation and 10-12 hours after			
		the onset of symptoms			
Reference standard:	Universal definition of AMI, including measurement of	Not applicable			
	troponin T or I (using any method not defined as a hs-cTn				
	test) on presentation and 10-12 hours after the onset of				
	symptoms in $\ge$ 80% of the population <sup>\$</sup> or occurrence of MACE				
	(any definition used in identified studies) during 30 day				
	follow-up				
Outcomes <sup>\$\$</sup> :	Test accuracy (the numbers of true positive, false negative,	Early discharge (≤4 hrs after initial presentation) without MACE during			
	false positive and true negative test results)	follow-up, incidence of MACE during follow-up, re-attendance at or re-			
		admission to hospital during follow-up, time to discharge, patient			
		satisfaction or health-related quality of life (HRQoL) measures			
Study design:	Diagnostic cohort studies	Randomised controlled trials (RCTs) (controlled clinical trials (CCTs)			
		will be considered if no RCTs are identified)			

\* A high sensitivity assay is defined as one which has a CV  $\leq 10\%$  at the 99<sup>th</sup> percentile value for the healthy reference population, and where the LoD allows measurable concentrations to be attained for at least 50% of healthy individuals

\*\* For serial hs-cTn assays, both data on change in Tn levels and peak Tn values were be considered

<sup>\$</sup> Studies that used only new diagnostic ECG changes or outcome-based MACE (cardiac death, non-fatal AMI, revascularisation, or hospitalisation for myocardial ischemia) alongside a Tn-based reference standard were eligible for inclusion<sup>7</sup>

<sup>\$\$</sup> Any estimates of the relative accuracy/effectiveness of different hs-cTnT or hs-cTnI tests, were derived from direct, within study comparisons

#### 3.1.3 Inclusion screening and data extraction

Two reviewers (MW and PW) independently screened the titles and abstracts of all reports identified by searches and any discrepancies were discussed and resolved by consensus. Full copies of all studies deemed potentially relevant were obtained and the same two reviewers independently assessed these for inclusion; any disagreements were resolved by consensus. Details of studies excluded at the full paper screening stage are presented in Appendix 4.

Studies cited in materials provided by the manufacturers of hs-cTn assays were first checked against the project reference database, in Endnote X4; any studies not already identified by our searches were screened for inclusion following the process described above.

Data were extracted on the following: study details, inclusion and exclusion criteria, participant characteristics (demographic characteristics and cardiac risk factors), target condition (NSTEMI or AMI), details of the hs-cTnT or hs-cTnI test (manufacturer, timing, and definition of positive diagnostic threshold), details of reference standard (manufacturer, timing, diagnostic threshold for conventional Tn T or I testing, clinical and imaging components of the reference standard, method of adjudication (e.g. two independent clinicians)), and test performance outcome measures (numbers of true positive (TP), false positive (FP), false negative (FN) and true negative (TN) test results). Data were extracted by one reviewer, using a piloted, standard data extraction form and checked by a second (MW and PW); any disagreements were resolved by consensus. Full data extraction tables are provided in Appendix 2.

# 3.1.4 Quality assessment

The methodological quality of included diagnostic test accuracy (DTA) studies was assessed using QUADAS-2.<sup>33</sup> Quality assessments was undertaken by one reviewer and checked by a second (MW and PW); any disagreements were resolved by consensus.

The results of the quality assessments are summarised and presented in tables and graphs in the results of the systematic review and are presented in full, by study, in Appendix 3.

## 3.1.5 Methods of analysis/synthesis

Sensitivity and specificity were calculated for each set of 2×2 data and plotted in receiver operating characteristic space. The bivariate/hierarchical summary receiver operating characteristic (HSROC) model was used to estimate summary sensitivity and specificity with 95% confidence intervals (CIs) and prediction regions around the summary points, and to derive hierarchical summary receiver operating characteristic curves for meta-analyses involving four or more studies.<sup>34-36</sup> This approach

allows for between-study heterogeneity in sensitivity and specificity, and for the trade-off (negative correlation) between sensitivity and specificity commonly seen in diagnostic meta-analyses. For meta-analyses with fewer than four studies we estimated separate pooled estimates of sensitivity and specificity, using random-effects logistic regression.<sup>37</sup> Heterogeneity was assessed visually using summary receiver operating characteristic plots and statistically using the variance of logit (sensitivity) and logit (specificity), where "logit" indicates the logistic function: the smaller these values the less heterogeneity between studies. Summary positive and negative likelihood ratios were derived from the summary estimates of sensitivity and specificity. Analyses were performed in Stata 10 (StataCorp LP, College Station, Texas, USA), mainly using the *metandi* command. For analyses that would not run in Stata we used MetaDisc.<sup>38</sup>

Analyses were conducted separately for each of the three hs-cTn assays. Analyses were stratified according to whether the study evaluated the prediction of AMI or MACE, timing of collection of blood sample for testing, and the threshold used to define a positive hs-cTn result. We investigated possible sources of heterogeneity using stratified analyses based on the following variables:

- Population: studies included mixed populations compared to those that excluded patients with STEMI.
- Age >70 years compared to age ≤70 years
- Patients with pre-existing CAD at baseline compared to patients without pre-existing CAD
- Time from symptom onset to presentation <3 hours compared to >3 hours
- Time from symptom onset to presentation <6 hours compared to >6 hours
- Low to moderate pre-test probability of disease compared to high pre-test probability of disease

Stratified analyses were conducted for all time points and thresholds for which sufficient data were available. To investigate the influence of risk of bias on the studies we restricted analyses to studies conducted in patients at low or unclear risk of bias for the two QUADAS items considered to have the greatest potential to have introduced bias into these studies: the item on patient spectrum (1) and the item on patient flow (4). As the focus of this review was the diagnosis of NSTEMI, we conducted these analyses in studies that excluded patients with STEMI. We used summary receiver operating characteristic (ROC) plots to display summary estimates from the various primary and stratified analyses.

We compared the accuracy of the three different hs-cTn assays by tabulating summary estimates from analyses for common time points and thresholds assessed for all assays. Only one study provided a direct comparison of all three assays. Estimates of sensitivity, specificity, and positive and

32

negative likelihood ratio (LR+ and LR-) for each assay derived from this study were included in the summary tables.

## 3.2 Results of the assessment of clinical effectiveness assessment

The literature searches of bibliographic databases identified 6,766 references. After initial screening of titles and abstracts, 261 were considered to be potentially relevant and ordered for full paper screening; of these 35 were included in the review.<sup>39-73</sup> All potentially relevant studies cited in documents supplied by the test manufacturers had already been identified by bibliographic database searches. One additional study was identified from hand searching of conference abstracts,<sup>74</sup> and two additional studies were identified from information supplied by clinical experts.<sup>75, 76</sup> Figure 1 shows the flow of studies through the review process, and Appendix 4 provides details, with reasons for exclusions, of all publications excluded at the full paper screening stage.

## 3.2.1 Overview of included studies

Based on the searches and inclusion screening described above, 38 publications <sup>39-76</sup> of 18 studies<sup>39, 41, 43, 45, 47, 48, 50, 53, 54, 56, 57, 62, 63, 65, 67, 71, 74, 75</sup> were included in the review; the results section of this report cites studies using the primary publication and, where this is different, the publication in which the referenced data were reported. Fifteen studies reported accuracy data for the Roche Elecsys hs-cTnT assay, <sup>39, 41, 43, 45, 48, 50, 53, 54, 56, 57, 63, 65, 67, 71, 75</sup> four studies reported accuracy data for the Abbott ARCHITECT hs-cTnI assay, <sup>47, 57, 62, 75</sup> and two studies reported accuracy data for the Beckman Coulter Access hs-cTnI assay;<sup>74, 75</sup> two studies reported data for more than one assay.<sup>57, 75</sup> No RCTs or CCTs were identified; no studies provided data on the effects on patient-relevant outcomes of management based on hs-cTn assays at presentation and after 10 to 12 hours. All studies included in the systematic review were diagnostic cohort studies, which reported data on the diagnostic or prognostic accuracy hs-cTn assays.

Thirteen<sup>39, 41, 43, 47, 48, 50, 54, 56, 63, 65, 67, 74, 75</sup> of the 18 included studies were conducted in Europe (two in the UK<sup>65, 67</sup>), four were conducted in Australia and New Zealand,<sup>45, 53, 57, 62</sup> and one was conducted in the USA.<sup>71</sup> Thirteen of the 18 included studies reported receiving some support from test manufacturers, including supply of assay kits; <sup>39, 41, 45, 47, 48, 50, 53, 54, 56, 62, 63, 71, 75</sup> two studies did not report any information on funding.<sup>57, 74</sup>

Full details of the characteristics of study participants, study inclusion and exclusion criteria, and hscTn assay used and reference standard, and detailed results are reported in the data extraction tables presented in Appendix 2 (Tables a, b and c).

# Figure 1: Flow of studies through the review process



#### 3.2.2 Study quality

The main potential sources of bias in the 18 studies included in this assessment relate to patient spectrum and patient flow. There were also concerns regarding the applicability of the patient population and the reference standard in some of the included studies. The results of QUADAS-2 assessments are summarised in Table 3 and Figure 2; full QUADAS-2 assessments for each study are provided in Appendix 3. A summary of the risks of bias and applicability concerns within each QUADAS-2 domain is provided below.

#### 3.2.2.1 Patient spectrum

Three studies<sup>41, 45, 50</sup> were rated as high risk of bias for patient selection and a further six were rated as unclear risk of bias. Most studies rated as unclear risk of bias did not provide sufficient details to make a judgement on whether appropriate steps were taken to minimise bias when enrolling patients into the study.<sup>43, 57, 63, 67, 71</sup> In one study a large number of patients were not enrolled due to 'technical reasons' that were not fully defined and so it was not possible to judge whether these constituted inappropriate exclusions; this study was also judged as unclear risk of bias for this domain.<sup>54</sup> One study only enrolled patients presenting between 05.30 and 20.00 and so patients who presented outside these hours were excluded; as these patients may differ in their presenting characteristics (e.g. time from symptom onset) this was considered to introduce a potential bias into the study.<sup>45</sup> A further study stated that consecutive patients were enrolled except for temporary interruptions of the study due to high work load in the coronary care unit.<sup>50</sup> This was also considered to have the potential to lead to the inclusion of a different spectrum of patients than if consecutive patients had been enrolled. The last study judged at high risk of bias for patient enrolment excluded certain patient groups including those with a Tn elevation in any two serial determinations, a prior diagnosis of ischemic heart disease, structural heart disease, concomitant heart failure or significant bradyarrhythmia.<sup>41</sup>

Although this assessment included studies that enrolled both mixed populations (i.e. when the target condition was any AMI) and studies restricted to populations where patients with STEMI were excluded (i.e. target condition NSTEMI), the primary focus was the population of patients with STEMI excluded. Studies not restricted to this specific patient group were therefore considered to have high concerns regarding applicability. Seven studies were restricted to patients in whom STEMI had been excluded<sup>39, 43, 45, 50, 54, 63, 65</sup>, an additional study enrolled a mixed population but also presented data for patients in whom STEMI had been excluded.<sup>75</sup> Three of these studies<sup>43, 50, 54</sup> were restricted to patients admitted to coronary care/chest patients units and so were considered to represent patients with more severe disease. A further study had strict inclusion criteria which resulted in the

35
inclusion of a very low risk population.<sup>65</sup> These four studies were not considered to be representative of patients with chest pain presenting to the emergency department who are the main focus of this assessment and so were also rated as having high concerns regarding applicability. Therefore only four studies<sup>39, 45, 75, 77</sup> (one only for a subset of data<sup>75</sup>) were considered to have low concerns regarding the applicability of the included patients.

## 3.2.2.2 Index test

All but one of the studies were rated as low risk of bias for the index test as all reported data for at least one threshold that was pre-specified (generally the 99<sup>th</sup> centile threshold, LoD or LoB threshold). The study that was rated as high risk of bias on this domain assessed the accuracy of the Beckman Coulter Access hs-cTnI assay at a single threshold which was derived from the ROC curve.<sup>74</sup> As the reference standard (diagnosis of AMI or MACE) was interpreted after the high sensitivity Tn test blinding was not considered important for these studies. Inclusion criteria were very tightly defined in terms of the high sensitivity Tn assays that we were interested in and so all studies were considered to have low concerns regarding the applicability of the index test.

#### 3.2.2.3 Reference standard

Six studies were rated as unclear risk of bias for reference standard.<sup>39, 41, 43, 54, 71, 74</sup> In five this was because it was unclear whether the diagnosis of AMI/MACE was made without knowledge of the high sensitivity Tn results.<sup>39, 41, 43, 54, 56</sup> Two studies reported as abstracts provided insufficient details on how the diagnosis of AMI was made, including whether adjudicators were blinded to the high sensitivity Tn results, to judge whether an appropriate reference standard had been used.<sup>71, 74</sup> No studies were rated as high risk of bias for this domain as these would not have fulfilled the inclusion criteria for the review. In our review question we specified that an appropriate reference standard had to include a standard Tn measurement at baseline and at 10-12 hours after the onset of symptoms in 80% of the population.<sup>11</sup> Only five studies<sup>41, 50, 62, 65, 75</sup> met this criteria for standard Tn measurement and were judged to have low concerns regarding the applicability of the reference standard; all but one of the remaining studies were judged at high risk of bias, the other study did not provide exact details on the timing of the standard Tn assay.<sup>39</sup>

#### 3.2.2.4 Patient flow

Six studies were considered at high risk of bias for patient flow<sup>43, 47, 53, 63, 65, 75</sup> and a further three were considered at unclear risk of bias.<sup>57, 71, 74</sup> In all cases this was related to withdrawals from the study; verification bias was not considered to be a problem in any of the studies. The four studies that were rated as unclear risk of bias were only reported as abstracts and did not provide sufficient details to judge whether there were any withdrawals in the study. The studies judged at high risk of

bias on this domain generally excluded patients for whom samples or high sensitive Tn results were not available.

Study	<b>RISK OF BI</b>	AS		-	APPLICABI		ICERNS
	Patient	Index	Reference	Flow	Patient	Index	Reference
	selection	test	standard	and	selection	test	standard
	-		-	timing	2		-
Aldous(2011) <sup>55</sup>	$\odot$	$\odot$	$\odot$	$\overline{\odot}$	$\overline{\mbox{\scriptsize (i)}}$	$\odot$	$\overline{\mathbf{O}}$
Aldous(2012) <sup>45</sup>	$\overline{\mbox{\scriptsize (S)}}$	$\odot$	$\odot$	$\odot$	$\odot$	$\odot$	$\overline{\mbox{\scriptsize (S)}}$
Body(2011) <sup>67</sup>	?	$\odot$	$\odot$	$\odot$	$\overline{\mathbf{i}}$	$\odot$	8
Christ(2010) <sup>56</sup>	$\odot$	$\odot$	?	$\odot$	8	$\odot$	8
Collinson(2013) <sup>65</sup>	$\odot$	$\odot$	$\odot$	$\overline{\mbox{\scriptsize (s)}}$	8	$\odot$	$\odot$
Cullen(2013) <sup>62</sup>		$\odot$	<u></u>	$\odot$	$\overline{\overline{\mathbf{o}}}$	$\odot$	$\odot$
Eggers(2012) <sup>43</sup>	?	$\odot$	?	8	8	$\odot$	$\overline{\mathbf{i}}$
Freund(2011) <sup>48</sup>	$\odot$	$\odot$	$\odot$	$\odot$	8	$\odot$	8
Hoeller(2013) <sup>75</sup>	$\odot$	$\odot$	$\odot$	8	<mark>8/</mark> 3	$\odot$	$\odot$
Keller(2011) <sup>47</sup>	$\odot$	$\odot$	$\odot$	$\overline{\otimes}$	$\overline{\otimes}$	$\odot$	$\overline{\mbox{\scriptsize (S)}}$
Kurz(2011) <sup>54</sup>	?	$\odot$	$\odot$	$\odot$	8	$\odot$	8
Lippi(2012) <sup>74</sup>	$\odot$	8	?	?	$\overline{\overline{\mathbf{o}}}$	$\odot$	$\overline{\mathbf{i}}$
Melki(2011) <sup>50</sup>	1	$\odot$	$\odot$	$\odot$	$\overline{\otimes}$	$\odot$	$\odot$
Parsonage(2013) <sup>57</sup>	?	$\odot$	$\odot$	?	8	$\odot$	$\overline{\mathbf{i}}$
Saenger(2010) <sup>71</sup>	?	$\odot$	?	?	$\overline{\otimes}$	$\odot$	$\overline{\mbox{\scriptsize (S)}}$
Sanchis(2012) <sup>41</sup>	$\overline{\otimes}$	$\odot$	?	$\odot$	$\overline{\otimes}$	$\odot$	$\odot$
Santalo(2013) <sup>39</sup>	$\odot$	$\odot$	?	$\odot$	$\odot$	$\odot$	?
Sebbane(2013) <sup>63</sup>	?	$\odot$	$\odot$	$\overline{\mbox{\scriptsize (c)}}$		$\odot$	$\overline{\mbox{\scriptsize (c)}}$
Cow Risk 😕 H	igh Risk	? Uncle	ear Risk				

Table 3: QUADAS-2 results for studies of hs-cTn assays

Figure 2: Summary of QUADAS-2 results for studies of hs-cTn assays



#### 3.2.3 Diagnostic accuracy of the Roche Elecsys hs-cTnT assay

#### 3.2.3.1 Study details

Fifteen diagnostic cohort studies, <sup>39, 41, 43, 45, 48, 50, 53, 54, 56, 57, 63, 65, 67, 71, 75</sup> reported in 34 publications, <sup>39-46,</sup> <sup>48-61, 63-68, 70-73, 75, 76</sup> provided data on the diagnostic performance of the Roche Elecsys hs-cTnT assay. Fourteen of the 15 studies in this section assessed the accuracy of the Roche Elecsys hs-cTnT assay for the detection of AMI, and the remaining study assessed accuracy for the prediction of MACE within 30 days of the index presentation.<sup>41</sup> Eight studies provided data specific to the population of interest for this assessment; participants with STEMI were excluded, i.e. the target condition was NSTEMI rather than any AMI.<sup>39, 43, 45, 50, 54, 63, 65, 75</sup>

All 14 of the studies which assessed accuracy for the detection of AMI reported data on the diagnostic performance of a single sample taken on presentation. All but one of the studies reported data for the 99<sup>th</sup> centile for the general population the remaining study reported data for a ROC-derived threshold of 9.5 ng/L.<sup>54</sup> Studies additionally assessed the diagnostic performance of a LoD/LoB threshold (5 ng/L or 3 ng/L) in a single sample taken on presentation, <sup>45, 52, 53, 67, 75</sup> of a single sample taken 1 to 3 hours after presentation, <sup>45, 50</sup> and/or the diagnostic performance of a specified change in, or peak value of hs-cTnT level over the initial three hours from presentation.<sup>39, 45, 57, 71, 75</sup> Table 4 provides summary estimates of the diagnostic performance of all combinations of population, diagnostic threshold and hs-cTnT test timing which were assessed by more than one study. For analyses based on NSTEMI patients only, where sufficient data were available sensitivity analyses which excluded studies rated as 'high risk of bias' on one or more QUADAS domains, were also reported. Where combinations were assessed by a single study, diagnostic performance estimates derived from that study alone are provided. Key results used in the cost-effectiveness modelling conducted for this assessment are highlighted in bold. Full results (including numbers of TP, FP, FN and TN test results), for all studies and all datasets, are provided in Appendix 2 (Table c).

## 3.2.3.2 Presentation samples

The summary estimates of sensitivity and specificity, where the diagnostic threshold was defined as the 99<sup>th</sup> centile for the general population, were 89% (95% CI: 85 to 92%) and 82% (95% CI: 77 to 86%), based on data from 13 studies; <sup>39, 43, 45, 48, 50, 53, 56, 57, 63, 65, 67, 71, 75</sup> the SROC curve for this analysis is shown in Figure 3. The LR+ and LR- were 4.96 (95% CI: 3.84 to 6.39) and 0.14 (95% CI: 0.10 to 0.19), respectively. These estimates were similar when the analysis was restricted to studies which excluded participants with STEMI; summary estimates of sensitivity and specificity were 88% (95% CI: 78 to 93%) and 84% (95% CI: 74 to 90%), respectively (SROC curve shown in Figure 4) and the LR+ and LR- were 5.41 (95% CI: 3.40 to 8.63) and 0.15 (95% CI: 0.08 to 0.26), respectively, based on six

studies.<sup>39, 43, 45, 50, 63, 65</sup> The only study, conducted in a population which excluded participants with STEMI, which was rated as 'low or unclear risk of bias' on all QUADAS domains,<sup>39</sup> reported similar sensitivity and negative LR (see Table 4) to the summary estimates, but lower estimates of specificity (71% (95% CI: 66 to 76%)) and LR+ (3.11 (95% CI: 2.55 to 3.79)). Results were also similar when the analysis was restricted to eight studies with a mixed population (i.e. where the target condition was any AMI); summary estimates of sensitivity and specificity were 89% (95% CI: 86 to 91%) and 81% (95% CI: 76 to 85%), respectively (SROC curve shown in Figure 5) and the LR+ and LR- were 4.64 (95% CI: 3.73 to 5.76) and 0.14 (95% CI: 0.11 to 0.17), respectively. 40, 48, 53, 56, 57, 67, 71, 75 Based on these data, it is unlikely that hs-cTnT testing on a single admission sample, using the 99<sup>th</sup> centile diagnostic threshold, would be considered adequate for either rule-out or rule-in of any AMI or NSTEMI. Although there was little apparent variation in the estimates of test performance derived from the three meta-analyses described above, the results of the second analysis (studies which excluded participants with STEMI) was selected to inform our cost-effectiveness analyses, as it best matched the main population of interest for this assessment (i.e. the target condition was NSTEMI rather than any AMI). The approach of, where possible, selecting data based on a population which excluded STEMI rather than a mixed population to inform cost-effectiveness modelling was applied throughout.







Figure 4: SROC for the Roche Elecsys hs-cTnT assay using the 99<sup>th</sup> centile threshold and a presentation sample (6 studies which excluded participants with STEMI)

Figure 5: SROC for the Roche Elecsys hs-cTnT assay using the 99<sup>th</sup> centile threshold and a presentation sample (8 studies with a mixed population, target condition any AMI)



Limited data were identified on additional clinical subgroups (age >70 years versus ≤70 years.<sup>52, 75</sup> without pre-existing CAD versus with pre-existing CAD,<sup>46,75</sup> and high versus low to moderate pre-test probability (determined by clinical judgement based on cardiovascular risk factors, type of chest pain, physical findings, and ECG abnormalities)<sup>48</sup>). None of these studies excluded participants with STEMI. The study which stratified participants by age,<sup>52, 75</sup> reported a higher estimate of sensitivity (97% (92% to 99%)) and a lower estimate of LR- (0.05 (95% CI: 0.02 or 0.18)) in participants >70 years of age than for patients ≤70 years of age (88% (95% CI:78 to 94%) and 0.14 (95% CI: 0.07 to 0.28), respectively); the estimates of sensitivity and LR- for people >70 years of age were also higher and lower, respectively, than the corresponding summary estimates derived from all 15 studies which used the 99<sup>th</sup> centile diagnostic threshold. A similar pattern was apparent for people with a high pre-test probability compared to those with a low to moderate pre-test probability<sup>48</sup> and for participants without pre-existing CAD compared to those with pre-existing CAD,<sup>46, 75</sup> see Table 4. As with the age stratification, the estimates of sensitivity and LR- were higher and lower, respectively, than the corresponding summary estimates derived from all 15 studies which used the 99<sup>th</sup> centile diagnostic threshold, for people with a high pre-test probability and for people without pre-existing CAD. Figure 6 illustrates the variation in performance characteristics of a single admission sample, using the 99<sup>th</sup> centile diagnostic threshold, when used in different clinical subgroups. These data provide some indication that hs-cTnT testing on a single admission sample, using the 99<sup>th</sup> centile diagnostic threshold, may be adequate for rule-out of AMI in certain selected populations (older people (≥70 years), those without pre-existing CAD, and people classified by clinical judgement as having a high pre-test probability.



Figure 6: ROC space plot for the Roche Elecsys hs-cTnT assay using the 99<sup>th</sup> centile threshold and a presentation sample in different clinical subgroups

Time from onset of chest pain to presentation was inconsistently reported across studies; where reported, the median time from onset ranged from 2.7 to 8.25 hours. Full details of all information reported is provided in Appendix 2 (Table a). Two studies specifically investigated variation in test performance according to time from symptom onset to presentation.<sup>67, 75</sup> Both of these studies were conducted in a mixed population, i.e. the target condition was any AMI. Study participants were stratified by presentation before or after three hours,<sup>67, 75</sup> and before or after six hours.<sup>67</sup> Summary estimates for the three hour stratification indicated that a presentation sample, using the 99<sup>th</sup> centile threshold had higher sensitivity (94% (95% CI: 92 to 96%)) and lower specificity (77% (95% CI: 75 to 79%)) for any AMI, when used to assess people presenting more than three hours after the onset of chest pain than when used to assess early presenters (sensitivity 78% (95% CI: 71 to 73%) and specificity 84% (95% CI: 81 to 86%)), see Table 4. The LR- was also lower when the test was used in people presenting after three hours from the onset of chest pain (0.08 (95% CI: 0.05 to 0.11)) than in early presenters (0.26 (95% CI: 0.178 to 0.39)). Test performance in people presenting after three hours, see Table 4. Figure 7 illustrates the variation in performance characteristics of a single

admission sample, using the 99<sup>th</sup> centile diagnostic threshold, when used in people presenting at different times from the onset of chest pain. These data provide some indication that hs-cTnT testing on a single admission sample, using the 99<sup>th</sup> centile diagnostic threshold, may be adequate for ruleout of AMI where people present after three hours from the onset of chest pain, but that longer delays in presentation did not appear to further improve rule-out performance.





Five studies considered the performance of a presentation sample using a threshold equivalent to the LoD (5 ng/L) or LoB (3 ng/L) of the assay for the diagnosis of AMI.<sup>45, 52, 53, 56, 67, 75</sup> Three studies reported data for the 5 ng/L threshold;<sup>45, 52, 53, 75</sup> one of these studies only reported data at this threshold for participants over 70 years of age.<sup>52, 75</sup> When this study was excluded, the summary estimates of sensitivity and specificity were 95% (95% CI: 92 to 97%) and 54% (95% CI: 51 to 58%), respectively, and the LR+ and LR- were 2.06 (95% CI: 1.40 to 2.64) and 0.09 (95% CI: 0.07 to 0.17), respectively (Table 4). Three studies reported data for the 3 ng/L threshold.<sup>41, 45, 67</sup> The summary estimates of sensitivity and specificity derived from these studies were 98% (95% CI: 95 to 99%) and 40% (95% CI: 38 to 43%), respectively, and the LR+ and LR- were 1.63 (95% CI: 1.24 to 1.86) and 0.05

(95% CI: 0.02 to 0.21), respectively (Table 4). Only one study was conducted in a population which excluded people with STEMI,<sup>45</sup> however, estimates of test performance from this study were similar to the summary estimates. For the 3 ng/L threshold, sensitivity and specificity derived from this study were 95% (95% CI: 92 to 98%) and 48% (95% CI: 44 to 51%), respectively, and the LR+ and LR-were 1.83 (95% CI: 1.70 to 1.97) and 0.10 (95% CI: 0.05 to 0.18), respectively, see Table 4.<sup>45</sup> For the 5 ng/L threshold, sensitivity and specificity derived from this study were 93% (95% CI: 89 to 96%) and 58% (95% CI: 55 to 62%), respectively, and the LR+ and LR- were 2.20 (95% CI: 2.00 to 2.50) and 0.11 (95% CI: 0.07 to 0.19), respectively, see Table 4.<sup>45</sup> These data provide some indication that hs-cTnT testing on a single admission sample may be adequate to rule out any AMI or NSTEMI, where a lower diagnostic threshold (5 ng/L or 3 ng/L) is used.

#### 3.2.3.3 Subsequent samples

The summary estimates of sensitivity and specificity, where the diagnostic threshold was defined as the 99<sup>th</sup> centile for the general population but the sample was taken 1 to 3 hours after presentation, were 95% (95% CI: 92 to 97%) and 80% (95% CI: 77 to 82%), based on data from two studies.<sup>45, 50</sup> The LR+ and LR- were 4.75 (95% CI: 3.98 to 5.23) and 0.06 (95% CI: 0.00 to 0.63), respectively, see Table 4. Both of these studies were conducted in populations which excluded people with STEMI. Unsurprisingly, these data indicate a similar improvement in rule-out performance to that seen when the test is used only in people presenting more than three hours after the onset of chest pain.

#### 3.2.3.4 Multiple samples

Six studies (data reported in multiple publications) provided data on the performance of a variety of diagnostic strategies involving multiple sampling,<sup>39, 45, 49, 51, 57, 64, 71, 75</sup> most commonly involving a combination of a peak hs-cTn value above the 99<sup>th</sup> centile diagnostic threshold and a 20% change in hs-cTnT over two or three hours following presentation, see Table 4. Figure 8 shows the results of these studies plotted in ROC space. One study reported data for this combination over two hours, in a population which excluded people with STEMI,<sup>45, 49</sup> and this study was used in cost-effectiveness modelling. It is important to give full consideration to the optimal way of interpreting combination data of this type. As can be seen from the values reported in Table 4, a positive result from the 'AND' combination (defined as both a peak value above the 99<sup>th</sup> centile AND a change of >20% over two hours) provides the optimum rule-in performance (LR+ 8.42 (95% CI: 6.11 to 11.60)); conversely, a negative result from the 'OR' combination (defined as both no value above the 99<sup>th</sup> centile AND a change of <20% over two hours) provides the optimum rule-in performance (LR+ 8.42 (95% CI: 6.11 to 11.60)); conversely, a negative result from the 'OR' combination (defined as both no value above the 99<sup>th</sup> centile AND a change of <20% over two hours) provides the optimum rule-in performance (LR+ 0.04 (95% CI: 0.02 to 0.10)). Where a patient has a negative result from the 'AND' combination/positive result from the 'OR' combination (Defined as both no value above the 99<sup>th</sup> centile OR a change of >20% over

two hours), further investigation is likely to be needed. This optimal interpretation strategy is illustrated in Figure 9, along with a potential initial rule-out step, based on a presentation sample below the LoB threshold (3 ng/L); this strategy is included in cost-effectiveness modelling. Figure 9 shows the application of this two stage approach to a theoretical cohort of 1,000 people presenting with symptoms suggestive of ACS (STEMI excluded); the estimated number of people with AMI and a negative test result who would be erroneously discharged based on this testing strategy is 14 (nine at the first stage and five at the second stage). The prevalence of NSTEMI was estimated to be 17%, based on data from three studies conducted in populations which excluded people with STEMI.<sup>39, 45,</sup> <sup>63</sup> Four studies were excluded from the estimate of prevalence because they were considered to have unrepresentative populations; three were conducted in coronary care unit populations, <sup>43, 50, 54</sup> and one was conducted in a low risk population.<sup>78</sup> It was assumed that the diagnostic performance of 'AND'/'OR' combinations of peak values of hs-cTnT and change over two hours, using the 99<sup>th</sup> centile diagnostic threshold, are the same for people in whom NSTEMI is not ruled out by the initial test (hs-cTnT > LoB) as for the initial population; this was because no test performance data were available for the combination of initial hs-cTnT test using the LoB diagnostic threshold followed by combined peak hs-cTnT and change over two hours using the 99<sup>th</sup> centile threshold.

### 3.2.3.5 Prognostic accuracy

One study assed the performance of a presentation sample at the LoB (3 ng/L) threshold for the prediction of MACE within 30 days of the index presentation.<sup>41</sup> The results of this study indicate that a positive test was a poor predictor of occurrence of MACE and a negative test was not adequate to rule out MACE within 30 days (Table 4).





Grouping	Population	Risk of bias	N	Sensitivity (%)	Specificity (%)	LR+	LR-
Presentation samples							
Any threshold*	All	Mixed	14	88 (84, 91)	82 (77, 86)	4.88 (3.84, 6.21)	0.14 (0.11, 0.19)
	All	Low/unclear risk of bias on patient spectrum	13	86 (83, 89)	82 (77, 87)	4.89 (3.76, 6.35)	0.16 (0.14, 0.20)
	All	Low/unclear risk of bias on patient flow	11	90 (87, 93)	80 (77, 84)	4.69 (3.88, 5.66)	0.12 (0.09, 0.16)
	All	Low/unclear risk of bias on patient spectrum and patient flow	8	89 (85, 92)	80 (74 <i>,</i> 85)	4.49 (3.47, 5.80)	0.14 (0.11, 0.18)
99 <sup>th</sup> centile threshold	All	Mixed	13	89 (84, 91)	82 (77, 86)	4.96 (3.84, 6.69)	0.14 (0.10, 0.19)
	Mixed	Mixed	8	89 (86, 91)	81 (76, 85)	4.64 (3.74, 5.76)	0.14 (0.11, 0.17)
	STEMI excluded	Mixed	6	88 (78, 93)	84 (74, 90)	5.41 (3.40, 8.63)	0.15 (0.08, 0.26)
	STEMI excluded	Low/unclear risk of bias on patient spectrum	4	81 (75, 86)	85 (70, 93)	5.33 (2.65, 10.72)	0.22 (0.17, 0.29)
	STEMI excluded	Low/unclear risk of bias on patient flow	3	92 (88, 94)	79 (76, 82)	4.38 (3.02, 6.11)	0.10 (0.05, 0.22)
	STEMI excluded	Low/unclear risk of bias on patient spectrum and patient flow	1 <sup>39</sup>	89 (81, 94)	71 (66, 76)	3.11 (2.55, 3.79)	0.15 (0.08, 0.28)
	age ≤70 years	High risk for patient flow	1 <sup>52, 75</sup>	88 (78, 94)	86 (83, 89)	6.24 (5.03, 7.74)	0.14 (0.07, 0.28)
	age >70 years	High risk for patient flow	1 <sup>52, 75</sup>	97 (92, 99)	49 (44, 55)	1.91 (1.71, 2.14)	0.05 (0.02, 0.18)
	patients with pre-existing CAD	High risk for patient flow	1 <sup>46, 75</sup>	93 (85, 97)	60 (55, 65)	2.32 (2.02, 2.68)	0.12 (0.05, 0.26)
	patients without pre-existing CAD	High risk for patient flow	1 <sup>46, 75</sup>	94 (88, 97)	82 (79, 85)	5.18 (4.36, 6.16)	0.07 (0.04, 0.16)
	Mixed; Low to moderate pre-test probability	Low	1 <sup>48</sup>	89 (70, 97)	85 (79, 89)	5.79 (4.16, 8.06)	0.13 (0.04, 0.41)
	Mixed; High pre-test probability	Low	1 <sup>48</sup>	94 (77, 99)	66 (50 <i>,</i> 79)	2.78 (1.75, 4.41)	0.09 (0.02 <i>,</i> 0.45)
	Symptom onset <3 hours	1 study high risk for patient flow	2 <sup>67, 75</sup>	78 (71, 83)	84 (81, 86)	4.88 (3.91, 5.74)	0.26 (0.18, 0.39)
	Symptom onset >3 hours	1 study high risk for patient flow	<b>2</b> <sup>67, 75</sup>	94 (92 <i>,</i> 96)	77 (75, 79)	4.09 (3.33, 5.70)	0.08 (0.05, 0.11)
	Symptom onset <6 hours	Low	1 <sup>67</sup>	83 (74 <i>,</i> 89)	83 (79 <i>,</i> 86)	4.80 (3.80, 6.08)	0.21 (0.14, 0.32)

# Table 4: Accuracy of the Roche hs-cTnT assay: Summary estimates (95% confidence intervals)

Grouping	Population	Risk of bias	N	Sensitivity (%)	Specificity (%)	LR+	LR-
	Symptom onset >6 hours	Low	167	94 (78, 99)	81 (75, 86)	4.99 (3.66, 6.81)	0.07 (0.02, 0.34)
LoD (<5ng/L)	All	Mixed	3	96 (94, 98)	41 (39, 44)	1.63 (0.34, 7.07)	0.10 (0.07, 0.17)
	All; Outlying study conducted in	Mixed	2	95 (92, 97)	54 (51, 58)	2.06 (1.40, 2.64)	0.09 (0.07, 0.17)
	patients age>70 years removed						
	Age >70 years	High risk for patient flow	152,75	100 (95, 100)	1 (0, 3)	1.01 (0.99, 1.03)	0.45 (0.02, 8.56)
	STEMI excluded	High risk for patient	1 <sup>45</sup>	93 (89, 96)	58 (55 <i>,</i> 62)	2.20 (2.00, 2.50)	0.11 (0.07, 0.19)
		spectrum					
LoB (<3ng/L)	All	Mixed	3	98 (95 <i>,</i> 99)	40 (38 <i>,</i> 43)	1.63 (1.24, 1.86)	0.05 (0.02, 0.21)
	STEMI excluded	High risk for patient	1 <sup>45</sup>	95 (92, 98)	48 (44 <i>,</i> 51)	1.83 (1.70, 1.97)	0.10 (0.05, 0.18)
		spectrum					
	Mixed; symptom onset <3 hours	Low	1 <sup>67</sup>	99 (94, 100)	64 (57 <i>,</i> 69)	2.73 (2.31, 3.23)	0.01 (0.00, 0.16)
	Mixed; symptom onset >3 hours	Low	1 <sup>67</sup>	99 (91, 100)	33 (28 <i>,</i> 38)	1.47 (1.36, 1.59)	0.03 (0.00, 0.47)
	Mixed; symptom onset <6 hours	Low	1 <sup>67</sup>	100 (96, 100)	34 (30, 39)	1.52 (1.41, 1.64)	0.01 (0.00, 0.22)
	Mixed; symptom onset >6 hours	Low	1 <sup>67</sup>	100 (84, 100)	33 (27, 40)	1.47 (1.31, 1.65)	0.06 (0.00, 0.91)
1-3 hours after							
presentation							
1-3 hours after	STEMI excluded	High risk for patient	2 <sup>45, 50</sup>	95 (92 <i>,</i> 97)	80 (77 <i>,</i> 82)	4.75 (3.98, 5.23)	0.06 (0.00, 0.63)
presentation, 99 <sup>th</sup>		spectrum					
centile threshold							
Multiple samples							
99 <sup>th</sup> centile threshold	All	High risk for patient	1 <sup>45, 49</sup>	50 (43, 56)	94 (92 <i>,</i> 96)	8.40 (6.10, 11.60)	0.54 (0.47, 0.62)
(peak) and ∆20%		spectrum					
(presentation-3hrs)							
99 <sup>th</sup> centile (peak)	All	High risk for patient	1 <sup>45, 49</sup>	97 (94, 99)	65 (61, 68)	2.80 (2.50, 3.10)	0.04 (0.02, 0.10)
threshold or $\Delta 20\%$		spectrum					
(presentation-3hrs)							
99 <sup>th</sup> centile (peak)	STEMI excluded	Low	1 <sup>45, 49</sup>	50 (43, 56)	94 (92, 96)	8.42 (6.11, 11.60)	0.54 (0.47, 0.62)
threshold and $\Delta 20\%$							
(presentation-2hrs)							
99 <sup>th</sup> centile (peak)	STEMI excluded	Low	1 <sup>45, 49</sup>	97 (94, 99)	65 (61 <i>,</i> 68)	2.76 (2.50, 3.05)	0.04 (0.02, 0.10)
threshold or Δ20%							
(presentation-2hrs)							

Grouping	Population	Risk of bias	Ν	Sensitivity (%)	Specificity (%)	LR+	LR-
Peak above 99 <sup>th</sup>	All	Mixed	2 <sup>45, 49, 57</sup>	94 (91 <i>,</i> 97)	84 (82 <i>,</i> 86)	5.88 (3.56, 10.24)	0.07 (0.04, 0.11)
centile							
On presentation (30	STEMI excluded	Low	1 <sup>39</sup>	99 (94 <i>,</i> 100)	66 (61, 72)	2.94 (2.50, 3.47)	0.01 (0.00, 0.15)
minutes after arrival),							
and at 2, 4 and 6-8							
hours or until							
discharge: ∆20%							
On presentation and	STEMI excluded	High risk for patient flow	1 <sup>64, 75</sup>	60 (51 <i>,</i> 69)	72 (69 <i>,</i> 75)	2.15 (1.77, 2.60)	0.55 (0.44, 0.70)
at 1 hour: Δ17%							
On presentation and	STEMI excluded	High risk for patient flow	1 <sup>51, 75</sup>	64(52, 74)	84 (80 <i>,</i> 87)	3.97(3.05, 5.17)	0.43 (0.31, 0.59)
at 2 hours: Δ30%							
On presentation and	Mixed	Low	1 <sup>71</sup>	95(89 <i>,</i> 98)	95(91, 97)	19.19 (10.31, 35.72)	0.05 (0.02, 0.12)
at 3 hours: Δ8 ng/L							
Prediction of MACE							
On presentation, LoB	STEMI excluded	Low	1 <sup>41</sup>	85 (74, 92)	46 (41, 51)	1.58 (1.37, 1.81)	0.33 (0.18, 0.59)
threshold	+h						

\*All but one study used the 99<sup>th</sup> centile as the threshold, the remaining study used at threshold of 9.5ng/L

Key results, used in cost-effectiveness modelling are highlighted in bold

## Figure 9: Testing pathway for the Roche Elecsys hs-cTnT assay used in cost-effectiveness modeling



# 3.2.4 Diagnostic accuracy of the Abbott ARCHITECT hs-cTnl assay for the rule-out and diagnosis of AMI

#### 3.2.4.1 Study details

Four diagnostic cohort studies provided data on the diagnostic performance of the Abbott ARCHITECT hs-cTnI assay.<sup>47, 57, 62, 75</sup> Three of these studies assessed the accuracy of the Abbott ARCHITECT hs-cTnI assay for the detection of AMI,<sup>47, 57, 75</sup> and the remaining study assessed accuracy for the prediction of MACE within 30 days of the index presentation.<sup>62</sup> None of the studies in this section provided data specific to the population of interest for this assessment; participants with STEMI excluded, i.e. the target condition was NSTEMI rather than any AMI. All four studies were conducted in mixed populations. Full details of the baseline characteristics of study populations, including baseline cardiac risk factors are provided in Appendix 2 (Table a).

Where a single diagnostic threshold was used to define a positive test result for AMI, all studies in this section reported data for the 99<sup>th</sup> centile for the general population and a single sample taken at presentation. Table 5 provides summary estimates of diagnostic performance for this testing strategy. All other combinations of diagnostic threshold and hs-cTnI test timing were assessed by only one study. Figure 10 shows the diagnostic performance of all testing strategies assessed plotted in ROC space. Diagnostic performance estimates derived from these studies are also provided. Key results used in the cost-effectiveness modelling conducted for this assessment are highlighted in bold. Full results (including numbers of true positive (TP), false positive (FP), false negative (FN) and true negative (TN) test results), for all studies and all datasets, are provided in Appendix 2 (Table c).

#### 3.2.4.2 Presentation samples

Summary estimates of sensitivity and specificity based on a diagnostic threshold was defined as the 99<sup>th</sup> centile for the general population were 80% (95% CI: 77 to 83%) and 93% (95% CI: 92 to 94%), based on data from three studies.<sup>47, 57, 75</sup> The LR+ and LR- were 11.47 (95% CI: 9.04 to 16.19) and 0.22 (95% CI: 0.16 to 0.27), respectively. All three studies were conducted in a mixed population (i.e. where the target condition was any AMI). Based on these data, it is unlikely that hs-cTnI testing on a single admission sample, using the 99<sup>th</sup> centile diagnostic threshold, would be considered adequate for rule-out of any AMI, but a positive test result may be useful in ruling-in AMI.

No studies reported clinical subgroup data, or data on the performance of the test in people presenting at different times after symptom onset for the Abbott ARCHITECT hs-cTnl assay.

One study also considered the performance of a presentation sample using the LoD of the assay as the threshold for diagnosing AMI.<sup>47</sup> This study provided estimates of sensitivity and specificity of

100% (95% CI: 98 to 100%) and 35% (95% CI: 32 to 38%), respectively, and the LR+ and LR- were 1.54 (95% CI: 1.47 to 1.62) and 0.01 (95% CI: 0.00 to 0.08), respectively, see Table 5. These data provide some indication that hs-cTnI testing on a single admission sample may be adequate to rule out any AMI, where a lower diagnostic threshold (the LoD of the assay) is used.

#### 3.2.4.3 Subsequent samples

One study assessed the performance of hs-cTnI testing on a sample taken three hours after presentation, where the diagnostic threshold was defined as the 99<sup>th</sup> centile for the general population.<sup>57</sup> The summary estimates of sensitivity and specificity, derived from this study, were 98% (95% CI: 96 to 99%) and 90% (95% CI: 88 to 92%). The LR+ and LR- were 10.16 (95% CI: 8.38 to 12.31) and 0.02 (95% CI: 0.01 to 0.08), respectively, see Table 5. These data provide some indication that a sample taken at three hours after presentation may be informative, at the 99<sup>th</sup> centile threshold, for both rule-out and rule-in of AMI.

#### 3.2.4.4 Multiple samples

Two studies provided data on the performance of a variety of diagnostic strategies involving multiple sampling (Table 5).<sup>47, 57</sup> None of these strategies appeared to offer a performance advantage over testing based on a single sample. Figure 11 illustrates our proposed optimal testing pathway for the Abbott ARCHITECT hs-cTnI assay; this strategy is included in cost-effectiveness modelling. As with Figure 9, which presents the Roche Elecsys hs-cTnT optimal strategy, Figure 11 shows the application of this two stage approach to a theoretical cohort of 1,000 people presenting with symptoms suggestive of ACS (STEMI excluded), with a prevalence of NSTEMI of 17%; the estimated number of people with AMI and a negative test result who would be erroneously discharged based on this the diagnostic performance of hs-TnI using the 99<sup>th</sup> centile diagnostic threshold on a sample taken three hours after presentation is the same for people in whom NSTEMI is not ruled out by the initial test (hs-cTnI > LoD) as for the initial population; this was because no test performance data were available for the combination of initial hs-cTnI test using the LoD diagnostic threshold followed by three hour hs-cTnI and using the 99<sup>th</sup> centile threshold.

#### 3.2.4.5 Prognostic accuracy

One study assed the performance of a presentation sample at the 99<sup>th</sup> centile for the prediction of MACE within 30 days of the index presentation.<sup>62, 75</sup> The results of this study indicate that a positive test may be helpful in predicting the occurrence of MACE, whilst a negative test was not adequate to rule out MACE within 30 days, see Table 5.

0



Figure 10: ROC space plot of the Abbott ARCHITECT hs-cTnT assay

Population	Risk of bias	Ν	Sensitivity (%)	Specificity (%)	LR+	LR-
Mixed	Mixed	3 <sup>47, 57, 75</sup>	80 (77, 83)	93 (92, 94)	11.47 (9.04,	0.22 (0.16, 0.27)
					16.19)	
Mixed	High risk for patient	1 <sup>47</sup>	100(98, 100)	35 (32 <i>,</i> 38)	1.54 (1.47, 1.62)	0.01 (0.00, 0.08)
	flow					
Mixed	High risk for patient	1 <sup>47</sup>	98 (96 <i>,</i> 99)	90 (88 <i>,</i> 92)	10.16 (8.38,	0.02 (0.01, 0.05)
	flow				12.31)	
Mixed	Unclear risk for patient	1 <sup>57</sup>	91 (81 <i>,</i> 96)	93 (91 <i>,</i> 95)	12.94 (9.74,	0.09 (0.04, 0.23)
	spectrum and flow				17.19)	
Mixed	High risk for patient	1 <sup>47</sup>	82 (78 <i>,</i> 86)	52 (49 <i>,</i> 55)	1.73 (1.59, 1.88)	0.34 (0.26, 0.43)
	flow					
Mixed	High risk for patient	1 <sup>47</sup>	77 (72, 82)	26 (23 <i>,</i> 29)	1.04 (0.97, 1.12)	0.87 (0.69, 1.11)
	flow					
Mixed	High risk for patient	2 <sup>62, 75</sup>	88 (85, 91)	93 (91, 94)	12.57 (8.88,	0.13 (0.06, 0.28)
	flow for 1 study				15.35)	
	Population Mixed Mixed Mixed Mixed Mixed Mixed	PopulationRisk of biasMixedMixedMixedHigh risk for patient flowMixedHigh risk for patient flowMixedUnclear risk for patient spectrum and flowMixedHigh risk for patient flowMixedHigh risk for patient flow	PopulationRisk of biasNMixedMixed $3^{47, 57, 75}$ MixedHigh risk for patient $1^{47}$ flowflow1MixedHigh risk for patient $1^{47}$ flowUnclear risk for patient $1^{57}$ MixedUnclear risk for patient $1^{57}$ MixedHigh risk for patient $1^{47}$ flow11MixedHigh risk for patient $1^{47}$ flow11MixedHigh risk for patient $1^{47}$ flow111MixedHigh risk for patient $1^{47}$ flow111flow11MixedHigh risk for patient $2^{62, 75}$ flow for 1 study11	PopulationRisk of biasNSensitivity (%)MixedMixed $3^{47, 57, 75}$ 80 (77, 83)MixedHigh risk for patient $1^{47}$ 100(98, 100)flow1198 (96, 99)MixedHigh risk for patient $1^{47}$ 98 (96, 99)flow1191 (81, 96)spectrum and flow1182 (78, 86)MixedHigh risk for patient $1^{47}$ 82 (78, 86)flow11177 (72, 82)MixedHigh risk for patient $1^{47}$ 88 (85, 91)MixedHigh risk for patient $2^{62, 75}$ 88 (85, 91)MixedHigh risk for patient $2^{62, 75}$ 88 (85, 91)	Population         Risk of bias         N         Sensitivity (%)         Specificity (%)           Mixed         Mixed         3 <sup>47, 57, 75</sup> 80 (77, 83)         93 (92, 94)           Mixed         High risk for patient         1 <sup>47</sup> 100(98, 100)         35 (32, 38)           Mixed         High risk for patient         1 <sup>47</sup> 98 (96, 99)         90 (88, 92)           Mixed         High risk for patient         1 <sup>57</sup> 91 (81, 96)         93 (91, 95)           Spectrum and flow         1 <sup>47</sup> 82 (78, 86)         52 (49, 55)           Mixed         High risk for patient         1 <sup>47</sup> 82 (78, 86)         52 (49, 55)           Mixed         High risk for patient         1 <sup>47</sup> 77 (72, 82)         26 (23, 29)           Mixed         High risk for patient         1 <sup>47</sup> 88 (85, 91)         93 (91, 94)           Mixed         High risk for patient         1 <sup>47</sup> 78 (85, 91)         93 (91, 94)	Population         Risk of bias         N         Sensitivity (%)         Specificity (%)         LR+           Mixed         Mixed         3 <sup>47, 57, 75</sup> 80 (77, 83)         93 (92, 94)         11.47 (9.04, 16.19)           Mixed         High risk for patient         1 <sup>47</sup> 100(98, 100)         35 (32, 38)         1.54 (1.47, 1.62)           Mixed         High risk for patient         1 <sup>47</sup> 98 (96, 99)         90 (88, 92)         10.16 (8.38, 12.31)           Mixed         Unclear risk for patient         1 <sup>57</sup> 91 (81, 96)         93 (91, 95)         12.94 (9.74, 17.19)           Mixed         Unclear risk for patient         1 <sup>57</sup> 91 (81, 96)         93 (91, 95)         12.94 (9.74, 17.19)           Mixed         High risk for patient         1 <sup>47</sup> 82 (78, 86)         52 (49, 55)         1.73 (1.59, 1.88)           flow         1 <sup>47</sup> 82 (78, 86)         52 (49, 55)         1.73 (1.59, 1.88)         160w           Mixed         High risk for patient         1 <sup>47</sup> 77 (72, 82)         26 (23, 29)         1.04 (0.97, 1.12)           flow         1 <sup>477</sup> 78 (85, 91)         93 (91, 94)         12.57 (8.88, 15.35)

Table 5: Accuracy of the Abbott ARCHITECT hs-cTnI assay: Summary estimates (95% confidence intervals)

Key results, used in cost-effectiveness modelling are highlighted in bold

Figure 11: Testing pathway for the Abbott ARCHITECT hs-cTnI assay used in cost-effectiveness modeling



#### 3.2.5 Diagnostic accuracy of the Beckman Coulter Access hs-cTnl assay

#### 3.2.5.1 Study details

Two diagnostic cohort studies,<sup>74, 75</sup> reported in three publications,<sup>64, 74, 75</sup> provided data on the diagnostic performance of the Beckman Coulter Access hs-cTnI assay. Both studies assessed a precommercial version of the assay and both reported accuracy data for the diagnosis of AMI (any AMI,<sup>64, 74</sup> or NSTEMI<sup>75</sup>). No study assessed the performance of the Beckman Coulter Access hs-cTnI assay for the prediction of MACE within 30 days of the index admission. The diagnostic performance estimates, for all combinations of diagnostic threshold and test timing assessed by in cluded studies, are summarised in Table 6. Figure 12 shows the diagnostic performance of all testing strategies assessed, plotted in ROC space.

#### 3.2.5.2 Presentation samples

Both studies assessed the diagnostic performance of a single sample taken at presentation. One study used the 99<sup>th</sup> centile for the general population as the diagnostic threshold.<sup>75</sup> This study was considered the most relevant to our assessment and was used to inform cost effectiveness analyses; this was the only testing strategy modelled for the Beckman Coulter Access hs-cTnI assay and, for a theoretical cohort of 1,000 people presenting with symptoms suggestive of ACS (STEMI excluded) with a prevalence of NSTEMI of 17%, the estimated number of people with AMI and a negative test result who would be erroneously discharged based on this testing strategy is 14. However, it should be noted that the Beckman Coulter hs-cTnI assay evaluated in the this study was described as 'an investigational prototype';<sup>75</sup> the 99<sup>th</sup> centile (9 ng/L), described as 'acording to the manufacturer', differs from the 99<sup>th</sup> centile given in the current product information leaflet (40 ng/L).<sup>16</sup> The estimates of sensitivity and specificity derived from this study were 92% (95% CI: 88 to 95%) and 75% (95% CI: 72 to 78%) respectively, and the LR+ and LR- were 3.67 (95% CI: 3.26 to 4.13) and 0.11 (95% CI: 0.07 to 0.17), respectively, see Table 6. The summary estimates, for the two studies combined, were very similar (Table 6).

No studies reported clinical subgroup data, or data on the performance of the test in people presenting at different times after symptom onset, for the Beckman Coulter Access hs-cTnl assay.

#### 3.2.5.3 Subsequent samples

Neither of the studies reported data for single samples taken at time points other than presentation.

#### 3.2.5.4 Multiple samples

One study assessed the diagnostic performance of a >27% change in hsTnI from presentation to one hour.<sup>75</sup> This testing strategy produced results indicating a decline in both rule-in and rule-out

performance compared to the single presentation sample described above (Table 6).



Figure 12: ROC space plot of the Beckman Coulter Access hs-cTnT assay

Grouping	Population	Risk of bias	Ν	Sensitivity (%)	Specificity (%)	LR+	LR-
Prediction of AMI							
Presentation sample, 9ng/L and	All	High risk for	2 <sup>74, 75</sup>	92 (88, 95)	75 (72, 77)	3.68 (2.46, 4.48)	0.11 (0.07, 0.16)
18 ng/L		patient flow					
		on 1 study					
Presentation sample, 99 <sup>th</sup>	Mixed	High risk for	1 <sup>75</sup>	92 (88 <i>,</i> 95)	75 (72, 78)	3.67 (3.26, 4.13)	0.11 (0.07, 0.17)
centile (9ng/L)		patient flow					
On presentation and at 1 hour: $\Delta$	STEMI excluded	High risk for	1 <sup>64, 75</sup>	63 (53, 71)	66 (63, 69)	1.85 (1.55, 2.21)	0.56 (0.44, 0.72)
27%		patient flow					

## Table 6: Accuracy of the Beckman Coulter Access hs-cTnl assay: Summary estimates (95% confidence intervals)

Key results, used in cost-effectiveness modelling are highlighted in bold

# 3.2.6 Comparative diagnostic accuracy of the Roche Elecsys hs-TnT assay, the Abbott ARCHITECT hs-TnI assay and the Beckman Coulter Access hs-TnI assay

Only one study provided data for a direct comparison of the diagnostic performance of all thee hscTn assays in the same polulation.<sup>75</sup> These data were for the use of the 99<sup>th</sup> centile threshold in a sample taken at presentation. This was also the only time point and threshold assessed for each study by individual included studies. As can be seen from Tables 7 and 8 below, the summary estimates of the performance of each test, derived from all studies reporting data for this threshold, were similar to estimates derived from the direct comparison study alone.

 Table 7: Comparison between assays (Presentation samples, 99<sup>th</sup> centile threshold): Sensitivity and specificity (95% CI)

	Indirect comparison			Direct comparison <sup>75</sup>		
Assay	Ν	Sensitivity (%)	Specificity (%)	Sensitivity (%)	Specificity (%)	
Beckman Coulter	2	92 (88 <i>,</i> 95)	75 (72, 77)	92 (88, 98)	75 (72, 78)	
Access hs-cTnl						
Abbott ARCHITECT	3	80 (77, 83)	93 (92, 94)	77 (72, 82	93 (91, 94)	
hs-cTnI						
Roche Elecsys hs-	15	88 (85, 91)	82 (78 <i>,</i> 86)	90 (86, 92)	78 (76, 79)	
cTnT						

 Table 8: Comparison between assays (Presentation samples, 99<sup>th</sup> centile threshold): Likelihood ratios (95% CI)

	Indirect comparison			Direct comparison <sup>75</sup>		
Assay	Ν	LR+	LR-	LR+	LR-	
Beckman Coulter	2	3.32 (2.46, 4.48)	0.11 (0.07, 0.16)	3.68 (3.27, 4.14)	0.11 (0.07, 0.17)	
Access hs-cTnl						
Abbott ARCHITECT	3	12.10 (9.04, 16.19)	0.21 (0.16, 0.27)	10.42 (8.49, 12.79)	0.25 (0.20, 0.30)	
hs-cTnl						
Roche Elecsys hs-	15	5.02 (4.02, 6.28)	0.14 (0.11, 0.18)	4.02 (3.65, 4.43)	0.13 (0.10, 0.18)	
cTnT						

#### 3.2.7 Selection of diagnostic strategies for inclusion in cost-effectiveness modeling

Diagnostic strategies, for each hs-cTn assay, were selected for inclusion in cost-effectiveness modeling based on optimal diagnostic performance as indicated by data from the systematic review. In addition, wherever possible data from studies which excluded patients with STEMI (i.e. where the target condition was NSTEMI) were preferentially selected.

#### 4. ASSESSMENT OF COST-EFFECTIVENESS

This chapter explores the cost-effectiveness of hs-cTn assays (used singly or in series, up to four hours from the onset of chest pain/presentation), compared with the current standard of serial troponin T and/or I testing on admission and at 10-12 hours after the onset of symptoms for the early rule out of AMI in people with acute chest pain.

## 4.1 Review of economic analyses of hs-cTn assays

## 4.1.1 Search strategy

Searches were undertaken to identify cost-effectiveness studies of high sensitivity TnT/I. As with the clinical effectiveness searching, the main EMBASE strategy for each set of searches was independently peer reviewed by a second Information Specialist, using the CADTH Peer Review Checklist.<sup>28</sup> Search strategies were developed specifically for each database and keywords associated with high sensitivity TnT/I were adapted according to the configuration of each database. Full search strategies are reported in Appendix 1.

The following databases were searched for relevant studies from 2005-October 2013:

- MEDLINE (OvidSP): 2005-2013/10/wk1
- MEDLINE In-Process Citations and Daily Update (OvidSP): up to 2013/10/1
- EMBASE (OvidSP): 2005-2013/10/17
- NHS Economic Evaluation Database (NHS EED) (Wiley): Cochrane Library Issue 3 2005-July 2013
- Health Economic Evaluation Database (HEED) (Wiley): 2005-2013/10/18
- EconLit (EBSCO): 2005-2013/09/01
- Science Citation Index (SCI) (Web of Science): 2005-2013/10/21
- Conference Proceedings Citation Index Science (CPCI) (Web of Science): 2005-2013/10/21
- Research Papers in Economics (REPEC) (Internet): up to 2013/10/21
- http://repec.org/

Identified references were downloaded in Endnote X4 software for further assessment and handling.

References in retrieved articles were checked for additional studies.

## 4.1.2 Inclusion criteria

Studies reporting a full economic analysis, which related explicitly to the cost-effectiveness of hs-cTn or standard cTn (with cTn implying either cTnI or cTnT) testing, with survival and/or Quality-Adjusted Life Years (QALYs) as an outcome measure, were eligible for inclusion. Specifically, one of the

strategies had to include cTn testing. Studies that only reported a cost-analysis of cTn testing were not included in the review.

#### 4.1.3 Quality assessment

Full cost-effectiveness studies were appraised using the Drummond checklist.<sup>79</sup>

#### 4.1.4 Results

The literature search identified 152 reports. After initial screening of titles and abstracts, five reports were considered potentially relevant: two full papers, and three HTA reports. Two additional reports were identified provided by a clinical expert: a Canadian optimal use report (comparable to an HTA report) and an abstract which was referred to in this report. All seven identified reports fulfilled inclusion criteria based on full text assessment. The seven publications related to five studies. Figure 13 shows the flow of studies through the review process, Table 9 lists the study details and the results of the quality assessment are shown in Table 10.

# 4.1.4.1 Goodacre (2011)<sup>80</sup> and Fitzgerald (2011)<sup>81</sup>

This study was based on the multicentre pragmatic controlled trial 'Randomised Assessment of Treatment using Panel Assay of Cardiac Markers' (RATPAC).<sup>80</sup> An economic evaluation was undertaken to assess the cost-effectiveness of management based on testing with a panel of point-of-care cardiac markers compared with management without point-of-care panel assessment. The included population consisted of patients presenting to hospital with chest pain due to suspected, but not proven, AMI and no other potentially serious alternative pathology or co-morbidity. The analysis was performed from an NHS perspective using trial data to estimate the mean costs per patient of chest pain-related care and the mean number of QALYs accrued by patients in each arm of the trial, with a time horizon of three months. In addition, a decision-analytic model was constructed to duplicate (validate) trial results and extrapolate results to a longer time horizon.

## Figure 13: Flow of studies through the review process



CA=conference abstract; JA=journal article; HTA=health technology assessment

Resource use data were collected for all patients. Cost and outcome data were collected using patient notes and self-completed questionnaires. Unit prices were based partly on a micro-costing study on a sample of patients, partly on a study previously undertaken by the investigators, and partly on purchase price and national unit costs. QALYs were calculated based on EQ-5D measurements. In a sensitivity analysis, productivity costs were included as reported by the patients.

As it was anticipated that the trial would have limited power to detect a difference in major adverse events, the decision-analytic model was intended to explore whether uncertainty around the effect of the intervention upon the major adverse event rate could influence the potential cost-effectiveness of the intervention. The model used trial data to estimate costs and QALYs up to three months. Beyond this, lifetime cost and QALYs were estimated from a previous study.<sup>82</sup> It was assumed that patients who had died at three months would accrue no further costs or QALYs. Those who had survived non-fatal myocardial infarction (MI) would accrue costs and QALYs associated with coronary heart disease (CHD) (estimated at £10,079 and 6.829, respectively). Those without CHD were assigned zero costs and 20 QALYs.

Empirical results showed that the point-of-care test strategy was dominated by standard care, which delivered slightly more QALYs at a lower cost. The probability that point-of-care testing would be more cost-effective than standard care at a willingness to pay threshold of £20,000 per QALY was less than 1%. The decision-analytic model again resulted in higher costs and less effect for the point-of-care panel assay compared to standard care, also when extrapolated to lifetime survival. The probability of the point-of-care panel assay being cost-effective for the three month and lifetime model was 22.3% and 33.6%, respectively.

The main conclusion was that point-of-care panel assay testing is unlikely to be considered costeffective in the NHS, with an 89% probability that standard care was dominant. Cost-effectiveness was mainly driven by differences in mean cost, with point estimates suggesting that, per patient, point-of-care panel assessment was £211 more expensive than standard care.

## 4.1.4.2 Vaidya (2012)<sup>83</sup>

This study aimed to assess the cost-effectiveness of an hs-TnT assay, alone or in combination with the H-FABP assay in comparison with the conventional cTnT assay for the diagnosis of AMI in patients presenting to hospital with chest pain. A decision analytic model was developed to perform both a cost-utility analysis (cost per QALY gained) and a cost-effectiveness analysis (cost per life year (LY) gained and cost per AMI averted), using a health care perspective and a lifetime time horizon. One way and probabilistic sensitivity analyses were conducted.

63

The incremental cost-effectiveness ratio (ICER) for hs-TnT compared to conventional cTnT was  $\notin$ 3,748 per QALY gained. For hs-cTnT in combination with H-FABP compared to conventional cTnT the ICER was  $\notin$ 5,717 per QALY gained. For LY and AMI averted, no ICERs were reported in the abstract. The probabilistic sensitivity analysis showed the hs-TnT assay to be the preferable strategy with a probability of over 90%, at a ceiling ratio of  $\notin$ 4,800 per QALY. This led to the conclusion that the hs-TnT assay is very cost-effective relative to the conventional cTnT assay. Combining hs-TnT with H-FABP did not seem to offer any additional economic or health benefit over the hs-TnT test alone.

## 4.1.4.3 Goodacre (2013)<sup>7</sup> and Thokala(2012)<sup>84</sup>

This study aimed to estimate the cost-effectiveness of using alternative biomarker strategies to diagnose MI, and using biomarkers, computed tomography coronary angiography (CTCA) and exercise ECG to risk-stratify troponin-negative patients. As the second aim was outside the scope of this review, we have only summarised the analysis which compares the biomarker strategies for diagnosing MI, referred to in the HTA report as 'the diagnostic phase model'. The different diagnostic strategies were applied to a hypothetical cohort of patients attending the ED with suspected, but not proven, ACS. Patient characteristics were defined using data from the RATPAC trial, <sup>85</sup> as well as patients' arrival times during the day at the ED. The model assigned each patient a probability of re-infarction or death depending on their characteristics and whether or not they had treatment. The model took a lifetime time horizon. The economic perspective was that of the NHS in England and Wales.

The following strategies were applied to each patient:

- No testing: discharge all patients without treatment (hypothetical)
- Standard troponin assay measured at presentation using the 10% coefficient of variation as the threshold for positivity
- Standard troponin assay measured at presentation using the 99<sup>th</sup> percentile threshold
- High-sensitivity troponin assay measured at presentation using the 99<sup>th</sup> percentile threshold
- Standard troponin assay measured at presentation and 10 hours after symptom onset using the 99<sup>th</sup> percentile threshold

Blood tests at presentation were assumed to be taken in the ED and so a decision could be made within one hour of the test results becoming available. For the 10-12 hours troponin measurement, three different scenarios were tested:

- 'doctor-on-demand' scenario, with medical staff available 24 hours a day to make a disposition decision within one hour of the results being available
- twice-daily ward round scenario, with medical staff only available at twice daily ward rounds to make disposition decisions
- once-daily ward round scenario, with medical staff only available at a once daily ward round to make disposition decisions

Sensitivity and specificity estimates for the presentation troponin tests were obtained by performing meta-analysis of estimates from individual primary studies included in the accompanying review. The 10 hour troponin test was assumed to have perfect sensitivity and specificity as it was the reference standard for the review. This implies that false-positives of the hs-Tn testing at presentation will still be discharged home after the 10-12 hour troponin test, but false negatives will be discharged home without treatment. The 'discharge without testing or treatment' by definition has perfect specificity, but a sensitivity of 0%.

The risk of re-infarction and death for patients with MI was based on a study by Mills et al.<sup>86</sup> Life expectancy of patients with MI and MI with re-infarction was estimated from Polanczyk et al,<sup>87</sup> while the utility of patients with MI was based on Ward et al.<sup>88</sup> The utility of patients with re-infarction was estimated by using a multiplicative factor of 0.8 for patients with MI (expert opinion). Patients without MI were assigned the life expectancy and utility scores of the general population. Lifetime costs for patients with MI were based on Ward et al.<sup>88</sup> One-way sensitivity analyses were performed, as well as a probabilistic sensitivity analysis. In a secondary analysis, a strategy was added that involved alternative biomarkers in combination with the presentation troponin testing.

The results showed that measuring a 10 hour troponin level in all patients was the most effective strategy (ICER £27,546-103,560). However, at a threshold of £30,000 per QALY, the optimal strategy in all but one scenario was measurement of high-sensitivity troponin at presentation, with a 10 hour troponin test if positive and discharge home if negative (ICER £7,487–£17,191 per QALY). The exception was a scenario involving patients without known CAD and doctor available on demand to discharge the patient, where, using the £30,000 per QALY threshold, the strategy of measuring a 10 hour troponin level in all patients was optimal (ICER of £27,546 per QALY). Sensitivity analyses showed the optimal strategy to vary with different levels of sensitivity and timing of the tests.

The report concluded that the additional costs that are likely to be incurred by measuring a 10 hour troponin level, compared with a presentation high-sensitivity troponin level, are unlikely to represent a cost-effective use of NHS resources in most of the scenarios tested.

## 4.1.4.4 CADTH optimal use report<sup>89</sup>

This report aimed to determine the cost-effectiveness of hs-cTnT and hs-cTnI assays compared with each other as well as with cTnI assays in patients with suspected ACS symptoms in the ED. For this purpose, three comparators were considered: hs-cTnT, hs-cTnI, and cTnI. As cTnT is no longer available in Canada, it was not taken into account in the analysis. The target population consisted of 65-year old patients presenting to the ED, without ST-segment elevation, who required cTn testing for diagnosis of NSTEMI. For the economic evaluation, a decision tree was constructed which calculated lifetime cost per QALY from the perspective of a publicly funded health care system.

The model consisted of a short-term part, which had a time horizon of one year, and a long-term part. The short-term part incorporated the testing and treatment procedures and short-term outcomes. Patients were tested at presentation at the ED and, if they were not admitted to hospital after the first test, they were tested again after six hours. When the patient was admitted after the first test, treatment was said to be initiated early, and when a patient was admitted after the second test, treatment was late. One year mortality depended on whether a patient had NSTEMI and whether they were treated early, treated late, or untreated (in the case of false negative test results). Those not suffering from NSTEMI were further stratified into unstable angina (UA) or not having acute coronary syndrome (non-ACS). The annual probability of death in the long-term part of the model was dependent on patient age, gender, and whether they had suffered an NSTEMI, UA, or did not have any type of ACS in the short-term part of the model.

The sensitivity and specificity for each cTn test at presentation to the ED was derived from the systematic review which was also part of this study. In the model, patients with a negative cTn test at presentation were assumed to be observed and have a second cTn test six hours later. After the second cTn test, 90% of these false negatives were assumed to become true positives.

Short-term mortality rates and relative risks for treated/non-treated were taken from published clinical studies and one non-referenced study. The relative risk for late versus early treatment was derived from expert opinion. Long-term mortality rates were taken from published clinical studies, and one non-referenced study. QALYs were calculated by incorporating an age-specific utility decrement for patients with NSTEMI. A number of one-way sensitivity analyses were performed, as well as a probabilistic sensitivity analysis.

The base-case results indicated that hs-cTnI was dominated by hs-cTnT, when compared to cTnI, at an ICER of \$119,377 per QALY. The probabilistic sensitivity analysis showed that, for willingness-topay thresholds up to \$124,000, cTnI had the highest probability of being cost-effective. For thresholds over \$124,000, hs-cTnT had the highest probability of being cost-effective. The hs-cTnI test was not likely to be cost-effective for any value of the threshold.

The authors concluded that hs-cTnT would be considered the most cost-effective testing strategy if willingness to pay for a QALY is \$119,377 or more, otherwise cTnI would be the most cost-effective test. However, there was a lot of uncertainty in results when model assumptions were changed.

## 4.1.4.5 Collinson (2013)<sup>65</sup>

This study used the decision tree developed in the related HTA by Goodacre et al<sup>7</sup> to compare the cost-effectiveness of five diagnostic strategies to a hypothetical cohort of patients presenting to hospital with symptoms suggestive of myocardial infarction but with no diagnostic ECG changes, no known history of coronary heart disease and no major co-morbidities requiring inpatient treatment. Essentially, this was a sub-study of the point-of-care arm of the RATPAC trial. All methods and model inputs were identical to the study by Thokala et al<sup>84</sup> and the HTA report by Goodacre et al,<sup>7</sup> but with slightly different strategies applied to the cohort of patients:

- No testing: discharge all patients without treatment (theoretical 'zero' option)
- High-sensitivity cTnT at presentation: discharge home if test is negative or admit to hospital for troponin-testing at 10-12 hours if positive
- High-sensitivity cTnT and H-FABP at presentation: discharge home if both tests are negative or admit to hospital for troponin testing at 10-12 hours if either test is positive
- High-sensitivity cTnT at presentation and at 90 minutes as in the RATPAC protocol: discharge home if both tests are negative or admit to hospital testing at 10-12 hours if either test is positive
- Standard troponin testing at 10-12 hours (current standard as per NICE guidelines)

The difference with the other studies is in the addition of H-FABP in the 3<sup>rd</sup> strategy and in the second high-sensitive troponin test at 90 minutes in the 4<sup>th</sup> strategy. In a secondary analysis, cTnT was replaced by cTnI. Sensitivity and specificity of presentation biochemical testing were estimated using data from within the study (RATPAC). Standard troponin testing at 10-12 hours was assumed to have perfect sensitivity and specificity as this was again the reference standard.

At the £20,000 per QALY threshold, 10 hour troponin testing was cost-effective (£12,090 per QALY) in the doctor-on demand scenario, but not in the other scenarios (once-daily ward round and twice-daily ward rounds), when high-sensitivity cTnT and H-FABP measurement at presentation was cost-effective. At the £30,000 per QALY threshold, 10 hour troponin testing was cost-effective in the doctor-on-demand scenario and twice-daily ward rounds scenario (£24,600 per QALY), whereas the

troponin T and H-FABP measurement at presentation strategy was cost-effective (£14,806 per QALY) in the once-daily ward round scenario. Secondary analysis using cTnI instead of cTnT showed that cTnI testing at presentation and at 90 minutes was cost-effective in all three scenarios at the £20,000 per QALY threshold and in two of the scenarios at the £30,000 per QALY threshold, with 10 hour troponin being cost-effective only in the doctor-on-demand scenario (£24,327 per QALY). The overall conclusion was that 10 hour troponin testing is likely to be cost-effective compared with rapid rule-out strategies only if patients can be discharged as soon as a negative result is available and a £30,000 per QALY threshold is used.

#### 4.1.4.6 Summary of studies included in the cost-effectiveness review

Most of the studies identified in this review have found that the question whether hs-Tn testing is cost-effective cannot be answered unequivocally. In favour of hs-Tn testing, the abstract by Vaidya et al<sup>83</sup> concluded that hsTnT testing is 'very cost effective' and the study by Goodacre<sup>7</sup> concluded that 'the optimal strategy in all but one scenario was high-sensitivity troponin at presentation, with a 10 hour troponin test if positive and discharge home if negative' (p.xv). The other papers reported ICERs that were considerably higher and with substantial uncertainty. The accuracy of high-sensitive tests and the efficiency of decision-making based on test results were important drivers of cost-effectiveness.

Study details	Goodacre et al (2011) <sup>80</sup>	Vaidya et al <sup>83</sup>	Thokala et al <sup>84</sup> Goodacre et al	CADTH report <sup>89</sup>	Collinson et al <sup>65</sup>
	Fitzgerald et al		(2013)		
Population	People presenting to hospital	Patients presenting to the	Patients attending hospital	65-year-old patients	Patients presenting to hospital with
	with chest pain due to	hospital with chest pain	with symptoms suggesting MI,	presenting to an ED with	symptoms suggestive of myocardial
	suspected but not proven		but a normal or non-diagnostic	ischemic chest pain,	infarction but with no diagnostic
	AMI, and no other potentially		ECG, and no major	without ST-segment	ECG changes (ST deviation >1 mm
	serious alternative pathology		comorbidities requiring	elevation ECG who	or T-wave inversion > 3mm), no
	or comorbidity		hospital treatment	require cTn testing for	known history of coronary heart
				diagnosis of NSTEMI	disease and no major
					comorbidities requiring inpatient
					treatment
Time horizon	Lifetime	Lifetime	Lifetime	Lifetime	Lifetime
Objective	Estimate the cost-	Assess the cost-	Estimate the incremental cost	To investigate the cost-	Assess the cost-effectiveness of
-	effectiveness of the point-of-	effectiveness of a high-	per QALY of delayed troponin	effectiveness of hs-cTnT	measuring a combination of
	care panel in terms of mean	sensitive troponin T assav	testing compared with	and hs-cTnI assavs	biomarkers compared with
	costs and OALYs accrued	(hs-cTnT), alone or	presentation testing and no	compared with each	measurement of cardiac troponin
	compared with standard care	combined with the H-	testing to determine which	other as well as with cTnI	alone
		EABP assay in comparison	diagnostic strategy should be	assays in natients with	uione
		with the conventional	recommended	suspected ACS symptoms	
		cardiac tropopin (cTpT)	recommended	in the ED	
				In the ED	
		assay for the diagnosis of			
Course of	Data frame within the trial we	AIVII			Consitivity and enacificity data
Source of	Data from within the trial up	No information	Sensitivity and specificity were	Sensitivity and specificity	Sensitivity and specificity data
effectiveness	to 3 months, and beyond this,		taken from the meta-analysis	from review performed in	derived from data from the HIA
information	lifetime costs and QALY		as reported in the 2013	same report. Proportion	(RAIPAC) itself, short-term survival
	estimates were used from a		Goodacre report , the RATPAC	UA and mortality	and probability of re-infarction
	previous economic		trial <sup>®</sup> was used for sampling	estimated based on	based on Mills et al <sup>66</sup> . Source for
	evaluation.		patient characteristics, Mills <sup>89</sup>	published studies, and	long-term survival and QALYs not
			for risk of re-infarction and	one unpublished study.	specified
			death, Polanczyk <sup>90</sup> for life	Utility decrements based	
			expectancy of patients with MI	on published study	
			and re-MI		

# Table 9: Summary of included full papers

Study details	Goodacre et al (2011) <sup>80</sup>	Vaidya et al <sup>83</sup>	Thokala et al <sup>84</sup> Goodacre et al	CADTH report <sup>89</sup>	Collinson et al <sup>65</sup>
Commentana	Fitzgeraid et al	Conventional sTaT		he eTeT	ne testing, discharge all patients
Comparators	Diagnostic assessment using	Conventional cini	no biochemical testing:	ns-cini	no testing: discharge all patients
	the point-or-care biochemical	he eTeT	discharge all patients without	he cal	without treatment
	marker panel	IIS-CITT	treatment (hypothetical)	ns-chi	he alle at an an an training discharge
		he etc.		-7-1	ns-cin at presentation: discharge
	Conventional diagnostic	ns-cini combined with H-	standard troponin assay	cini	nome if test is negative or admit to
	assessment without the panel	ГАВР	measured at presentation		nospital for troponin testing at 10-
			using the 10% coefficient of		12 hours if positive
			variation as the threshold for		
			positivity		ns-cin and a combination of
					cytopiasmic or neuronormone
			standard troponin assay		biomarkers at presentation:
			measured at presentation		discharge nome if both tests are
			using the 99 percentile		negative or admit to hospital for
			threshold		troponin testing at 10-12 hours if
					either test is positive
			high-sensitivity troponin assay		
			measured at presentation		hs-cTn at presentation and at 90
			using the 99 <sup>th</sup> percentile		minutes as in the RATPAC protocol:
			threshold		discharge home if both tests are
					negative or admit to hospital for
			standard troponin assay		troponin testing at 10-12 hours if
			measured at presentation and		either test is positive
			10h after symptom onset using		
			the 99 <sup>th</sup> percentile threshold		standard troponin testing at 10-12
					hours
Unit costs	Microcosting study within	No information	Admission and treatment were	Costs of hospital	Hospital stay and treatment for MI
	RATPAC; PSSRU unit costs		based on the national tariff.	admission were based on	based on NHS reference cost,
			Lifetime costs for MI patients	the Ontario Case Costing	biochemical testing based on
			were taken from Ward <sup>88</sup> . The	Initiative database and	Goodacre et al <sup>°°</sup>
			price of a troponin test was	the Ontario Schedule of	
			taken from the 2011 Goodacre	Benefits for Physician	

Study details	Goodacre et al (2011) <sup>80</sup>	Vaidya et al <sup>83</sup>	Thokala et al <sup>84</sup> Goodacre et al	CADTH report <sup>89</sup>	Collinson et al <sup>65</sup>
	Fitzgerald et al <sup>81</sup>		(2013)'		
			report <sup>80</sup>	Services. Costs of ED visits	
				were based on a hospital	
				in Soutwestern Ontario	
				and the Ontario Schedule	
				of Benefits. Unit prices of	
				cTn tests were based on	
				information provided by	
				the manufacturers.	
Measure of	QALY	AMI survivor	QALY	QALY	QALYs
benefit					
Study type	Trial-based economic	Model-based cost-	Model-based cost-utility	Model-based cost-utility	Model-based cost-utility study
	evaluation up to 3 months,	effectiveness and cost-	analysis	analysis	
	decision tree lifetime. Cost-	utility study			
	utility analysis.				
Model	2-hour delay between	No information	10h troponin testing has	non-NSTEMI patients are	<ul> <li>10h troponin testing has</li> </ul>
assumptions	sampling and results available		perfect sensitivity and	further classified into	perfect sensitivity and
			specificity (since it is the	Unstable Angina (UA) or	specificity (since it is the
	4 hours after presentation at		reference standard)	non-ACS, with	reference standard)
	ED patients moves to			consequences for costs	
	inpatient dept		2h delay from the time at	and outcome	<ul> <li>presentation blood tests</li> </ul>
	1 hour dolou botwoon		which sampling could be		taken in ED and results
	I noul delay between		performed to results available	there is a small survival	available and decision
	presentation and start		For presentation testing	benefit (RR 1.01) of	made within 2h of
	Diomarker sampling		stratogios: dosision made	treating early compared	sampling
	After short term (test-		within th of results available	to treating late	for tosting at 10,12h
	treatment-outcome), progress		within thor results available	(presentation testing vs.	- IOI testing at 10-1211
	only depends on whether or		For 10h testing strategies:	standard testing)	scenario used
	not patient had MI, and		decision made according to		
	whether or not this was		scenario applied		
	treated				
			Diagnostic strategy only		
Study details	Goodacre et al (2011) <sup>80</sup> Fitzgerald et al <sup>81</sup>	Vaidya et al <sup>83</sup>	Thokala et al <sup>84</sup> Goodacre et al (2013) <sup>7</sup>	CADTH report <sup>89</sup>	Collinson et al <sup>65</sup>
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			influences outcomes among patients with MI		
Perspective	NHS	Healthcare	NHS	Publicly funded health care system	NHS in England and Wales
Discount rate	Not mentioned	No information	Nothing mentioned	5% discount rate applied to costs and QALYs	Nothing mentioned
Uncertainty around cost- effectiveness ratio expressed	iCE plane, probability of strategy being dominated/cost-effective	Cost-effectiveness acceptability curves (not shown in abstract)	Cost-effectiveness acceptability curves for probabilistic sensitivity analysis (PSA) results, per scenario	as reported in outcomes of one-way sensitivity analyses, and also (for PSA) In cost-effectiveness acceptability curves	Cost-effectiveness acceptability curves
Sensitivity analysis	Probabilistic sensitivity analysis	One way and probabilistic	One-way sensitivity analyses, scenario analyses (doctor on demand, twice-daily ward round, and once daily ward- round), and PSA		Secondary analysis using cTnl instead of cTnT, scenario analysis (doctor-on-demand, once-daily ward round, twice-daily ward round), and PSA
Outcome (cost and Lys/QALYs) per comparator	Empirical 3 months PoC £ 1217 QALY 0.158 SC £ 1006 QALY 0.161 For the model, no outcomes per comparator were reported	No information	For doctor-on-demand scenario, per 1000 patients without known CAD: No testing £ 965,994 QALY 26,227 Pres standard trop, 10% CV £ 1,560,361 QALY 26,345 Pres standard trop, 99th perc £ 1,609,760 QALY 26,352 Pres hs-trop, 99th perc £ 1,806,910 QALY 26,279 10h troponin £ 2,016,540 QALY 26,286	cTnI \$ 2,018 QALY 8.1385 hs-cTnI \$ 2,082 QALY 3.1389 hs-cTnT \$ 2,186 QALY 8.1399	For doctor-on-demand scenario, per 1000 patients: No testing £ 965,994 QALY 26,227 hs-cTnT at presentation £ 1,581,263 QALY 26,349 hs-cTnT at presentation and 90 min £ 1,715,526 QALY 26,354 hs-cTnT and H-FABP at presentation £ 1,682,362 QALY 26,359 10-hour troponin £ 2,016,540 QALY 26,386
Summary of	Empirical 3 months:	hsTnT vs cTnT: incr 111	For doctor-on-demand	cTnl reference	No testing – reference strategy

Study details	Goodacre et al (2011) <sup>80</sup> Fitzgerald et al <sup>81</sup>	Vaidya et al <sup>83</sup>	Thokala et al <sup>84</sup> Goodacre et al (2013) <sup>7</sup>	CADTH report <sup>89</sup>	Collinson et al <sup>65</sup>
incremental analysis	Fitzgerald et al <sup>81</sup> Increment PoC vs SC £211 QALY -0.00282 Probability PoC cost-effective at £20,000/QALY = 0.4% Decision model 3 months: Increment PoC vs SC £169 QALY -0.002 Probability PoC cost-effective at £20,000/QALY = 22.3% Decision model lifetime: Increment PoC vs SC £329 QALY -0.087 Probability PoC cost-effective at £20,000/QALY = 33.6%	Euros and 16-17 lives per 1,000 AMI ICER 3,748 Euro/QALY hsTnT + H-FABP vs cTnT: incr 178 Euros ICER 5,717 Euro /QALY	(2013)' scenario: Pres standard trop. 10% CV vs no testing: £ 5030/QALY Pres standard trop 99 <sup>th</sup> perc vs pres standard trop 10% CV: £ 6518/QALY Pres hs-trop 99 <sup>th</sup> perc vs pres standard trop 99 <sup>th</sup> perc: £ 7487/QALY 10h trop vs pres hs-trop 99 <sup>th</sup> perc: £ 27,546/QALY	hs-cTnl incr costs \$64 incr QALYS 0.000352 dominated (by extension) hs-cTnT incr costs \$168 incr QALYS 0.001408 ICER \$119,377/QALY	hs-cTnT compared to no testing ICER £ 5012/QALY hs-cTnT at presentation and at 90 minutes: dominated hs-cTnT and H-FABP compared to hs-cTnT at presentation: ICER £11,026/QALY (as reported bu t correct number should be 10,871) 10-hour troponin compared to Hs- cTnT and H-FABP: ICER £12,090/QALY Conclusion: if a rapid-rule out strategy with a sensitivity of 95% (and specificity of around 90%) would be available, then a 10-hour troponin strategy does not seem cost-effective

## Table 10: Checklist of study quality for full papers included

	Goodacre et al. 2011 <sup>80</sup> & Fitzgerald et al	Vaidya et al <sup>83</sup>	Thokala et al <sup>84</sup> & Goodacre et al 2013 <sup>7</sup>	CADTH report <sup>89</sup>	Collinson et al <sup>65</sup>
Study design					
The research question is stated	V	V	V	V	V
The economic importance of the research question is stated	V	Х	V	V	V
The viewpoint(s) of the analysis are clearly stated and justified	V	V	V	V	V
The rationale for choosing alternative programmes or interventions compared is stated	V	х	v	V	v
The alternatives being compared are clearly described	V	V	V	V	٧
The form of economic evaluation used is stated	V	V	V	V	V
The choice of form of economic evaluation is justified in relation to the	v	V	v	v	v
questions addressed					
	,	X	,	,	· · · ·
The source(s) of effectiveness estimates used are stated	ν	X	V	V	ν
Details of the design and results of effectiveness study are given (if based on a single study)	v	х	v	V	V
Details of the methods of synthesis or meta-analysis of estimates are given (if based on a synthesis of a number of effectiveness studies)	V	х	V	V	v
The primary outcome measure(s) for the economic evaluation are clearly stated	V	V	V	V	V
Methods to value benefits are stated	V	Х	V	V	х
Details of the subjects from whom valuations were obtained were given	V	Х	х	х	х
Productivity changes (if included) are reported separately	NA	Х	NA	NA	NA
The relevance of productivity changes to the study question is discussed	NA	Х	NA	NA	NA
Quantities of resource use are reported separately from their unit costs	V	Х	x	х	х
Methods for the estimation of quantities and unit costs are described	V	Х	V	V	٧
Currency and price data are recorded	V	Х	V	V	٧
Details of currency of price adjustments for inflation or currency conversion are given	V	Х	x	x	x

	Goodacre et al. 2011 <sup>80</sup> & Fitzgerald et al	Vaidya et al <sup>83</sup>	Thokala et al <sup>84</sup> & Goodacre et al 2013 <sup>7</sup>	CADTH report <sup>89</sup>	Collinson et al <sup>65</sup>
Details of any model used are given	V	Х	V	V	V
The choice of model used and the key parameters on which it is based are justified	V	х	V	V	V
Analysis and interpretation of results					
Time horizon of costs and benefits is stated	V	V	V	V	V
The discount rate(s) is stated	х	Х	х	V	х
The choice of discount rate(s) is justified	NA	Х	NA	V	NA
An explanation is given if costs and benefits are not discounted	х	Х	х	NA	х
Details of statistical tests and confidence intervals are given for stochastic data	V	Х	V	V	V
The approach to sensitivity analysis is given	V	Х	V	V	V
The choice of variables for sensitivity analysis is justified	V	Х	V	V	V
The ranges over which the variables are varied are justified	V	Х	V	V	V
Relevant alternatives are compared	V	V	V	V	V
Incremental analysis is reported	V	Х	V	V	V
Major outcomes are presented in a disaggregated as well as aggregated form	V	Х	V	V	V
The answer to the study question is given	V	V	V	V	V
Conclusions follow from the data reported	V	V	V	V	V
Conclusions are accompanied by the appropriate caveats	V	X	V	V	V

#### 4.2 Model structure and methodology

#### 4.2.1 Troponin tests considered in the model

The health economic analysis will estimate the cost-effectiveness of different troponin testing methods for diagnosing or ruling-out NSTEMI, in patients presenting at the ED with suspected NSTE-ACS, who have no major comorbidities requiring hospitalisation (e.g. as heart failure (HF) or arrhythmia) and in whom STEMI has been ruled out. Those diagnosed with NSTEMI will then be admitted to the hospital for AMI treatment and those diagnosed as without NSTEMI can be discharged without AMI treatment and further hospital stay. AMI treatment might include aspirin, statins and angiotensin converting enzyme inhibitors and consideration of coronary revascularisation for high-risk cases.<sup>7</sup> Initiating AMI treatment for NSTEMI will reduce the probability of major adverse cardiac events, particularly cardiac death and re-infarction.

Standard serial troponin testing, for patients with acute chest pain due to possible ACS, does not achieve optimal sensitivity in detecting AMI until 10-12 hours after onset of symptoms. Waiting for 10-12 hours after symptoms onset is burdensome for patients and induces additional health care costs. Therefore, various alternatives have been proposed, using more sensitive troponin tests, for the early rule-out of NSTEMI (within the four hour NHS emergency department target).<sup>91</sup>

Two hs-cTn assays (Roche Elecsys hs-cTnT and Abbott ARCHITECT hs-cTnI) are currently used in NHS laboratories in England and Wales. One additional assay (Beckman Coulter hs-cTnI) was listed in the scope for this assessment, pending CE marking. However, each of these tests can be used at different time points and with different diagnostic thresholds, resulting in multiple possible strategies for each test. Whether or not a test strategy was included in the economic model was decided based on optimal diagnostic performance given the available evidence on accuracy for a population with STEMI ruled out, and on applicability in clinical practice (see section 3.2). The test strategies evaluated in the model are:

- Standard troponin at presentation and at 10-12 hours (reference standard)
- Roche Elecsys hs-cTnT at presentation: 99<sup>th</sup> centile threshold
- Roche Elecsys hs-cTnT (optimal strategy): LoB threshold at presentation followed by 99<sup>th</sup> centile threshold peak within three hours and/or Δ20% (compared to presentation test) at 1-3 hours (Figure 9)
- Abbott ARCHITECT hs-cTnI at presentation: 99<sup>th</sup> centile threshold
- Abbott ARCHITECT hs-cTnI (optimal strategy): LoD threshold at presentation, followed by 99<sup>th</sup> centile threshold at three hours (Figure 11)
- Beckman Coulter hs-cTnI at presentation: 99<sup>th</sup> centile threshold

 No testing, discharge all patients without testing or treatment (only in sensitivity analyses). A troponin test may not be indicated when clinical judgment assesses the probability that a patient is experiencing an AMI as low. Therefore, consistent with the protocol, this hypothetical strategy, is included in sensitivity analyses wherein the AMI prevalence is varied.

In the base case, it was assumed that standard troponin had perfect sensitivity and specificity (reference case) for diagnosing AMI. Using this assumption, all patients testing positive on an hs-cTn test but negative on the standard troponin would be classified as false positives. This implies that their risk for adverse events would be the same as for those patients testing negative on both the hs-cTn test and the standard troponin and that they ought to be discharged home without further immediate treatment. However, recent evidence has shown that patients with a negative standard troponin, but a positive hs-cTn, may be at higher risk for adverse events than patients who test negative on both the standard and the high-sensitive troponin.<sup>92</sup> A secondary analysis was therefore performed, which attributed a higher risk of adverse events to a proportion of patients testing false positive with the hs-cTn test.

Based on the available evidence, two analyses were performed:

- Base case analysis
- Secondary analysis, assuming that false positives in the hs-cTn testing strategies do not have the same risk for adverse events as true negatives. Instead, these patients were assigned a higher risk for (re-)infarction and death, to reflect the idea that when the hs-cTn test gives a positive result, in some cases this must be caused by a disease process, whether or not the strict definition of AMI is met. The risk of adverse events in patients with positive hs-cTn but a negative standard troponin is higher than the patients testing negative on both the hs-cTn test and the standard troponin, but lower than risk of adverse events in patients come case with NSTEMI (i.e. both positive hs-cTn and standard troponin).

#### 4.2.2 Model structure

This assessment uses the HTA report by Goodacre et al<sup>7</sup> as a starting point for cost-effectiveness modelling. The Goodacre report compared the cost-effectiveness of several diagnostic strategies for ACS. The assessment group received the health economic model (in Simul8; SIMUL8 Corporation) that this HTA was based on and this model was used as a starting point to develop a de novo model (in Microsoft Excel) adapted to better fit the scope of the current assessment. In the health economic model the mean expected costs and quality adjusted life years (QALYs) were calculated for

each alternative strategy. These long-term consequences were estimated based on the accuracy of the different testing strategies followed by AMI treatment or discharge from the hospital without AMI treatment for patients presenting at the emergency department with suspected NSTE-ACS, including patients with NSTEMI and patients without NSTEMI, who are further subdivided into 'no ACS, no UA' and 'UA'. For this purpose a decision tree and a Markov model were developed. The decision tree was used to model the 30-day outcomes after presentation, based on test results and the accompanying treatment decision. These outcomes consisted of 'no ACS, no UA', 'UA', 'Non-fatal AMI (untreated)', 'Non-fatal AMI (treated)' and 'Death'. The decision tree is shown in Figure 14.

The long-term consequences in terms of costs and QALYs were estimated using a Markov cohort model (Figure 15) with a lifetime time horizon (60 years). The cycle time was one year, except for the first cycle which was adjusted to 335.25 days (365.25-30) to ensure that the decision tree period (30 days) and the first cycle combined summed to one year. The following health states were included:

- No acute coronary syndrome and no unstable angina (no ACS, no UA)
- Unstable angina
- Post AMI (treated and untreated)
- Post AMI with re-infarction
- Death

#### Figure 14: Decision tree structure



<sup>a</sup> During the first year post-AMI a distinction is made between treated and untreated AMI.

#### 4.2.3 Model parameters

Estimates for the model input parameters were retrieved from the literature and by consulting experts for unpublished data. Accuracy estimates were derived from the systematic review component of this assessment (see section 3.2).

#### 4.2.3.1 Transition probabilities

An overview of transition probabilities is provided in Table 11.

### Table 11: Transition probabilities

	Estimate	Se / 95% Cl	Distribution	Source
Decision tree (short term)				
NSTEMI prevalence <sup>a</sup>	0.170	0.028	Beta	Santalo (2013), <sup>39</sup> Aldous (2012) <sup>45</sup>
				Sebbane
				(2013). <sup>63</sup>
				APACE <sup>75</sup>
Proportion of UA (of all non-NSTEMI patients)	0.160	0.038	Beta	CADTH (2013) <sup>89</sup>
Decision tree (30-day) probabilities				
Mortality (30-day) treated AMI	0.097	0.012	Beta	Pope (2000) <sup>93</sup>
Mortality (30-day) untreated AMI	0.105	0.069	Beta	Pope (2000) <sup>93</sup>
Mortality (30-day) treated UA	0.021	0.005	Beta	Pope (2000) <sup>93</sup>
Mortality (30-day) no ACS	b	-	Fixed	ONS <sup>94</sup> )
Markov model (long term)	с		Fixed	Duitich Llocut
AMI Incidence		-	Fixed	Foundation <sup>95</sup>
				roundation
Annual re-infarction (treated) <sup>d</sup>	0.023	0.001	Beta	Smolina (2012) <sup>96</sup>
RR re-infarction (untreated versus treated) <sup>e</sup>	2.568	1.366 - 5.604	LogNormal	Mills (2011) <sup>86</sup>
Annual mortality no ACS	b	-	Fixed	ONS <sup>94</sup>
Annual mortality post-MI <sup>d</sup>	0.066	0.000	Beta	Smolina (2012) <sup>96</sup>
Annual mortality post re-infarction <sup>d</sup>	0.142	0.002	Beta	Smolina (2012) <sup>96</sup>
HR mortality (UA versus NSTEMI)	0.781	0.581 - 1.053	LogNormal	Allen (2006) <sup>97</sup>
RR mortality (untreated versus treated) <sup>d</sup>	1.877	0.951 - 4.239	LogNormal	Mills (2011) <sup>86</sup>
for patients tested false positive)				
OR AMI <sup>†</sup>			LogNormal	personal
4				communication <sup>92</sup>
OR Death'			LogNormal	personal
			_	communication
Proportion of AMI <sup>®</sup>			Beta	personal
Dreparties of Deeth <sup>g</sup>			Data	communication
Proportion of Death-			вета	personal
BR AMI <sup>f, h</sup>			LogNormal	nersonal
			LOGINOITINA	communication <sup>92</sup>
RR Death <sup>f, h</sup>			LogNormal	personal
				communication <sup>92</sup>

<sup>a</sup> Prevalence was used to calculate the proportions of true/false positives/negatives based on test accuracy. Prevalence was calculated using identified studies that included NSTEMI data (see section 3.2.3.4).

<sup>b</sup> Based on age dependent mortality from the general population.

<sup>c</sup> Age dependent incidence from the general population.

<sup>d</sup> Weighted average based on gender (58.1% males <sup>7</sup>).

<sup>e</sup> Increased re-infarction and mortality risk for untreated (versus treated) was assumed for the 1<sup>st</sup> year after presentation at ED, after which no increased risk was assumed (RR = 1.0).

<sup>f</sup> For patients with both positive high sensitivity and standard troponin tests versus patients with positive high sensitivity and negative standard troponin tests.

<sup>g</sup> Proportion for patients with both positive high sensitivity and standard troponin tests. This proportion is only used to covert odds ratios to relative risks.

 $^{
m n}$  ORs were converted to RRs using the method described by Zhang and Yu. $^{
m 98}$ 

#### 4.2.3.2 Decision tree

The proportions of patients testing positive or negative (and thus commencing AMI treatment or being discharged from the hospital) were based on the estimated accuracy of the testing strategies considered (Table 12) and the estimated prevalence of NSTEMI in the UK (17.0% with standard error 2.8%; Table 11).<sup>39,, 45,, 63,, 75</sup> This prevalence was higher than that derived from the RATPAC trial<sup>99</sup> and used in the Goodacre model,<sup>7</sup> because the RATPAC study population was a low risk population.<sup>81,, 85</sup> The proportion of true positives (TP), false positive (FP), false negative (FN) and true negatives (TN) were calculated as follows:

- TP = NSTEMI prevalence × sensitivity
- FP = (1 NSTEMI prevalence) × (1 specificity)
- FN = NSTEMI prevalence × (1 sensitivity)
- TN = (1 NSTEMI prevalence) × specificity

Subsequently, the proportions of patients who receive AMI treatment (TP + FP), and who are discharged without AMI treatment (TN + FN) were calculated. These results are listed in Table 13.

	Sensitivity (Se) <sup>a</sup>	Specificity (Se) <sup>a</sup>	Distribution	Source
Serial standard troponin	1.00 (-)	1.00 (-)	Fixed	Assumption
testing				
Roche Elecsys hs-cTnT (99th			Multivariate	Chapter 3
centile at presentation)	0.88 (0.04)	0.84 (0.04)	normal	
Roche Elecsys hs-cTnT			Multivariate	Chapter 3
(optimal strategy) <sup>b</sup>	0.93 (0.02) <sup>c</sup>	0.82 (0.01) <sup>C</sup>	normal	
Abbott ARCHITECT hs-cTnI			Multivariate	Chapter 3
(99th centile at			normal	
presentation)	0.80 (0.02)	0.93 (0.00)	normai	
Abbott ARCHITECT hs-cTnI			Multivariate	Chapter 3
(optimal strategy) <sup>d</sup>	0.98 (0.01) <sup>c</sup>	0.94 (0.01) <sup>c</sup>	normal	
Beckmann Coulter hs-cTnI			Multivariate	Chapter 3
(99th centile)	0.92 (0.02)	0.75 (0.01)	normal	
No troponin test <sup>e</sup>	0.00 (-)	1.00 (-)	Fixed	Assumption

#### Table 12: Test accuracy

<sup>a</sup> Correlation between sensitivity and specificity was calculated to be -0.262 based on the covariance matrix from the *metandi* output for the Roche Elecsys hs-cTnT (99th centile at presentation) test (see also Chapter 3). This correlation was assumed to be equal for other tests as it was not possible to obtain the covariance matrix for the other tests included in the economic analyses (a minimum of 4 studies is required).

<sup>b</sup> Calculated based on accuracy data for the Roche Elecsys hs-cTnT optimal testing strategy

<sup>c</sup> Standard error based on probabilistic sensitivity analysis

<sup>d</sup> Calculated based on accuracy data for the Abbott ARCHITECT optimal testing strategy

<sup>e</sup> The no testing strategy is only considered in sensitivity analyses.

	ТР	FP	FN	ΤN	PPV	NPV	LR+	LR-
Serial standard troponin testing	0.17	0.00	0.00	0.83	1.00	1.00	1.00	0.00
Roche Elecsys hs-cTnT (99th								
centile at presentation)	0.15	0.13	0.02	0.70	0.53	0.97	5.41	0.15
Roche Elecsys hs-cTnT (optimal								
strategy)	0.16	0.15	0.01	0.68	0.51	0.98	5.05	0.09
Abbott ARCHITECT hs-cTnl (99th								
centile at presentation)	0.14	0.06	0.03	0.77	0.70	0.96	11.47	0.21
Abbott ARCHITECT hs-cTnI								
(optimal strategy)	0.17	0.05	0.00	0.78	0.76	1.00	15.67	0.02
Beckman Coulter hs-cTnI (99th								
centile)	0.16	0.21	0.01	0.62	0.43	0.98	3.67	0.11
No troponin test <sup>a</sup>	0.00	0.00	0.17	0.83	0.00	0.83	0.00	1.00

#### Table 13: Test outcomes

The no testing strategy is only considered in sensitivity analyses, the FN rate represents the prevalence of NSTEMI

After treatment, TP patients in the decision tree were allocated to 'Non-fatal AMI (treated)' and FP patients were further subdivided between 'no ACS, no UA' and 'UA' (based on the proportion of UA among non-NSTEMI patients; Table 11). After being discharged, TN patients were also subdivided between 'no ACS, no UA' and 'UA', whereas FN patients were allocated to 'Non-fatal AMI (untreated)'. The proportions of FN's, reported in Table 13, can be considered as the proportions of AMIs that would have been missed when assuming that standard troponin testing has perfect accuracy. Finally, to calculate the total number of deaths in the decision tree, the probability of 30 day mortality was assigned based on above mentioned subdivision (Table 11). It was assumed that UA is always correctly diagnosed, hence the mortality probability for treated UA was used.

#### 4.2.3.3 Markov model

The age-dependent AMI incidence in the UK<sup>95</sup> was used to model the occurrence of AMI for patients in the health states 'no ACS,' and 'UA'. It was assumed that all AMIs in the Markov trace are diagnosed correctly and thus receive treatment. For patients in the 'Post-MI' health state, the probability of re-infarction after treated AMI was retrieved from a UK record linkage study, (n=387,452) which assessed long-term survival and recurrence after AMI.<sup>96</sup> For the current assessment the probabilities for females and males were weighted according to the estimated proportion of females and males in the population (males = 58.1%<sup>7</sup>). The re-infarction probability for the 'Post-MI with re-infarction' health state is equal to the re-infarction probability for the 'Post-MI' health state. The re-infarction RR for people with untreated versus treated AMI was calculated from a recent study by Mills et al<sup>86</sup> based on patients with a troponin concentration of 5 to 19 ng/L. This RR was assumed only for the first year after presentation at ED, after which no increased risk was assumed (i.e. RR = 1.0 for untreated versus treated AMI after year 1).

Age-dependent mortality from the general population was used for patients in the 'no ACS, no UA' health state.<sup>94</sup> For the 'Post-MI' and 'Post-MI with re-infarction' health states, mortality was extracted from the record linkage study.<sup>96</sup> Again the study by Mills et al<sup>86</sup> was used to calculate the mortality RR for untreated versus treated AMI for the first year, after which an RR of 1.0 was used. Finally, a multivariate adjusted mortality hazard ratio for UA versus NSTEMI was retrieved from a study by Allen et al<sup>97</sup> to calculate mortality after UA.

All input parameters for the Markov model are reported in Table 11.

#### 4.2.3.4 Health state utilities

Age-dependent utility scores, from the UK general population, were calculated for patients in the 'no ACS, no UA' health state based on a linear regression model.<sup>88</sup> These age-dependent utility scores from the general population, were combined with age-dependent disutilities for AMI<sup>89</sup> to calculate utilities for the 'Post-MI' health states (with or without re-infarction). Utility scores for the 'UA' health state were calculated based on Post-MI utility scores and a utility increment of 0.010 (Table 14).<sup>88</sup>

	Estimate	Se	Distribution	Source
No ACS, no UA				
Intercept	1.060	0.029	Normal	88
Disutility for age	0.004	0.001	Normal	88
Post-MI (disutility compared to no				
ACS by age)				
Age = 45	0.060	0.001	Normal	89
Age = 55	0.051	0.001	Normal	89
Age = 65	0.025	0.001	Normal	89
Age = 75	0.007	0.001	Normal	89
UA				
Utility increment compared to AMI	0.010	0.042	Normal	88

#### **Table 14: Utility scores**

#### 4.2.3.5 Resource use and costs

Test specific resource use consisted of the number of tests performed and the duration of hospital stay (hours) before discharge / AMI treatment (see Table 15).

	Estimate	Se / Range	Distribution	Source
Number of tests				
Serial standard troponin testing	2.00	-	Fixed	Assumption
Roche Elecsys hs-cTnT (99th centile at presentation)	1.00	-	Fixed	Assumption
Roche Elecsys hs-cTnT (optimal	1.60	0.02	Beta <sup>a</sup>	Chapter 3

#### Table 15: Resource use (test specific)

strategy)				
Abbott ARCHITECT hs-cTnl (99th	1.00	-	Fixed	Assumption
centile at presentation)				
Abbott ARCHITECT hs-cTnl (optimal	1.71	0.02	Beta <sup>ª</sup>	Chapter 3
strategy)				
Beckman Coulter hs-cTnI (99th	1.00	-	Fixed	Assumption
centile)				
No troponin test <sup>b</sup>	0.00	-	Fixed	Assumption
Hospital stay (hours) before				
discharge / AMI treatment <sup>b</sup>				
Serial standard troponin testing	14	13 - 15	Beta PERT	Assumption
Roche Elecsys hs-cTnT (99th centile at	3	-	Fixed	Assumption
presentation)				
Roche Elecsys hs-cTnT optimal	3	-	Fixed	Assumption
strategy (patients with AMI ruled-out				
on first test)				
Roche Elecsys hs-cTnT optimal	5	4 - 6		Assumption
strategy (patients receiving both tests)				
Abbott ARCHITECT hs-cTnl (99th	3	-	Fixed	Assumption
centile at presentation)				
Abbott ARCHITECT hs-cTnl optimal	3	-	Fixed	Assumption
strategy (patients with AMI ruled-out				
on first test)				
Abbott ARCHITECT hs-cTnl optimal	6	-	Fixed	Assumption
strategy (patients receiving both tests)				
Beckman Coulter hs-cTnI (99th centile	3	-	Fixed	Assumption
at presentation)				
No troponin test <sup>b</sup>	0	-	Fixed	Assumption

<sup>a</sup> Beta distribution is used to estimate the probability of patients receiving a second test (all patients receive the presentation test).

<sup>b</sup> The no testing strategy is only considered in sensitivity analyses.

<sup>c</sup> Includes delay from the time at which sampling could be performed to the time at which results became available (2 hours) and delay between arrival at hospital and troponin assessment commencing (1 hour).

Health state costs (Table 16) were mainly retrieved from previous economic evaluations conducted in the UK.<sup>88, 100</sup> Health state costs for the 'UA', 'Post-MI' and 'Post-MI with re-infarction' consisted of costs for three 15 minute GP consultations and medication costs.<sup>88</sup> For the first year in the 'UA' health state, costs for clopidogrel (for 60%) and hospitalisation (for 50%) were added to this. The first year costs for both 'Post-MI' health states were based on resource data from the Nottingham Heart Attack Register. <sup>100</sup>

Additionally, costs of fatal events, retrieved from a UK economic evaluation,<sup>88</sup> were accumulated for all fatal AMI's. For this purpose, it was assumed that all 30 day deaths after 'true' NSTEMI were due to a fatal AMI event. In addition, AMI treatment costs were calculated based on the national tariff for non-elective AMI without complications (HRG code: EB10Z).<sup>101</sup> To calculate the hospital stay costs for patients, based on the number of hours before the test results become available, non-elective NHS reference costs for the general medical ward were used (HRG code: EB01Z).<sup>101</sup> For this purpose, it was assumed that doctors were available on demand and the time to discharge was delayed due to time between arrival at the emergency department and start of first sampling (one hour) and the time between sampling and the results being available (two hours). In the case of multiple testing, the one hour delay between arrival at the emergency department and start of sampling was only applied to the first test, however, this also affected the timing of the second test if applicable. The two hour delay before test results become available applies to all tests performed. Incorporating these time delays effectively implies that only tests at presentation and tests performed one hour after presentation could inform decisions within the NHS four hour emergency department target. All other multiple testing strategies, as well as standard troponin testing at 10-12 hours, would require a transfer from emergency department to the general ward (patients are transferred to the general ward four hours after presentation at the emergency department). Finally, the test costs includes panel (including reagent, machine and maintenance), calibration and quality control costs. Depending on the annual number of panels, the test costs varied between £16.18 and £21.33, for annual rates of testing of 1,500 and 3,000 respectively.<sup>99</sup> Based on clinical expert input, the average test costs were estimated to be £20 (2011 price level).<sup>7,84</sup>

	Estimate (£)	Se / range (£)	Distribution	Source
Health state costs				
No ACS, no UA first year	0	-	Fixed	Assumption
No ACS, no UA subsequent year	0	-	Fixed	Assumption
UA first year <sup>a</sup>	548	-	Fixed	88
UA subsequent year <sup>a</sup>	213	-	Fixed	88
Post-MI first year <sup>a, b</sup>	5,835	488	Gamma	100
Post-MI subsequent years <sup>a, b</sup>	213	-	Fixed	88
Event costs				
Costs of fatal AMI <sup>a</sup>	1,451	-	Fixed	88
AMI treatment costs	3,436	-	Fixed	101
Unit prices				
Hospital stay costs (per hour) <sup>c</sup>	27	-	Fixed	101
Test costs <sup>a</sup>	20	18 - 26	Beta PERT	7, 84

Table 16: Health state costs, event costs and unit prices

<sup>a</sup> Price inflated to the 2012-2013 price level based on price indices from The Hospital & Community Health Services index.<sup>102</sup>

<sup>b</sup> Post-MI with or without re-infarction.

<sup>c</sup> NHS reference costs was divided by 24 to obtain the hourly costs

#### 4.2.4 Overview of main model assumptions

The main assumptions in the health economic analyses were:

- Serial troponin testing (comparator) has perfect accuracy (sensitivity = 1.0 and specificity = 1.0).
- For the Roche Elecsys hs-cTnT and Abbott ARCHITECT hs-cTnI optimal strategies it was assumed that the sensitivity and specificity for the subpopulation not discharged after the

presentation test is equal to the sensitivity and specificity for the initial group (presenting at the emergency department).

- The life expectancy, quality of life and costs for false positive patients is, in the base case analysis, equal to the life expectancy, quality of life and costs of true negative patients. This assumption was amended in the secondary and sensitivity analyses.
- In contrast with AMIs occurring during the decision tree period, all AMIs (either first or reinfarction) occurring in the Markov trace are diagnosed correctly and thus treated.
- UA is always correctly diagnosed and thus treated.
- The re-infarction probability for the 'Post-MI with re-infarction' health state is equal to the re-infarction probability for the 'Post-MI' health state.
- The increased Post-MI re-infarction and mortality probabilities for untreated AMI were assumed to last one year: afterwards a RR of 1.0 was applied (for untreated versus treated AMI).
- There is no additional benefit of starting treatment early, so treatment effect for highsensitive strategies is equal to treatment effect for standard troponin strategy.
- All 30 day deaths (after presentation at the emergency department) are due to fatal AMI events and will receive the associated costs.

#### 4.3 Model analyses

Expected costs, life years (LYs) and QALYs were estimated for all troponin testing methods. Discount rates of 3.5% and a half-cycle correction were applied for both costs and effects. Incremental cost and QALYs for each strategy versus standard troponin and versus the next best alternative were calculated. The ICER was then calculated by dividing the incremental costs by the incremental QALYs. Probabilistic sensitivity analyses (10,000 simulations) were performed, and cost-effectiveness acceptability curves (CEACs) and cost-effectiveness acceptability frontiers (CEAFs) were constructed. Although CEACs can be used to illustrate decision uncertainty, the option with the highest probability of being cost-effective may not necessarily be the most cost-effective option according to the expected values. Moreover, CEAFs can be used to illustrate the decision uncertainty surrounding the most cost-effective option <sup>103</sup>.

#### 4.3.1 Secondary analysis

For the base case it was assumed that patients who tested negative on standard troponin and positive on hs-cTn tests would experience life expectancy and quality of life equal to true negative patients. This assumption is, however, debatable. As unpublished data<sup>92</sup> show that patients with a negative standard troponin test and positive hs-cTn test have an increased risk of (re-)infarction and

mortality compared to those who test negative on both standard troponin and hs-cTn tests. Although this risk was not as high as in patients with both positive standard troponin and positive hscTn tests, it could still be considered prognostically important. Therefore, in this secondary analysis the risk of re-infarction and mortality was adjusted for patients who tested false positive (Table 11). It was assumed that for this proportion of patients, the relative treatment benefit would be equal to that for true positive patients. As the prevalence of this 'higher risk subgroup' is likely to be the same for all comparators, it was assumed that this proportion was equal to the lowest proportion of FP patients for all hs-cTn tests (Table 13). This 'higher risk subgroup' was assumed to be treated for all hs-cTn tests (since they tested positive with these tests) and untreated for the standard troponin test (since they tested negative with this test), thus affecting the probability of adverse outcomes and treatment costs. In addition, the post-MI utility and health state costs were used for this 'higher risk subgroup'.

#### 4.3.2 Sensitivity analysis

For both the base case and the secondary analysis, the following one-way sensitivity analyses were performed to assess the impact of model assumptions and input parameters on the estimated outcomes:

Model assumptions:

- The assumption that the increased post AMI re-infarction and mortality probabilities for untreated AMI only lasts for one year was replaced by the assumption that these probabilities would remain elevated for a lifetime.
- The assumption that a doctor will be available on demand and thus that a decision could be made immediately (as in the base case) was replaced with an assumed delay (one, two or three hours) before a doctor is available and a decision could be made.
- As for the previous sensitivity analysis except that the delay (one, two or three hours) only applies once patients are transferred to the general ward four hours after presentation; (no delay in the emergency department).
- A total delay of 1.5 hours is assumed (includes delay from the time at which sampling could be performed to the time at which results became available and delay between arrival at hospital and troponin assessment commencing) rather than assuming a total delay of three hours (base case).
- AMI treatment costs are applied for patients who tested false positive rather than using no treatment costs, as assumed in the base case analysis.
- In addition to the health state costs of UA during the first year, the AMI treatment costs

are also applied for patients with UA (during the first year), rather than assuming no additional treatment costs.

Model input parameters (varied to lower and upper boundary of the 95% CI unless stated otherwise):

- Test costs (test costs was varied over a wider range (£5-£40) than the 95% confidence interval)
- AMI treatment costs (- / + 25%)
- Post-MI first year health state costs
- Utility increment for UA compared to AMI
- Post-MI disutility compared to no ACS
- Mortality (30 day) treated AMI (decision tree)
- Mortality (30 day) untreated AMI (decision tree)
- Annual re-infarction (after initial AMI)
- RR re-infarction (untreated versus treated AMI)
- Annual Post-MI mortality
- Annual Post-MI mortality after re-infarction
- HR mortality (UA versus NSTEMI)
- RR mortality (untreated versus treated AMI)

#### 4.3.3 Subgroup analysis

For both the base case and the secondary analysis, a number of subgroup analyses were performed. The main subgroup analyses were based on age- and gender-dependent re-infarction probabilities, mortality probabilities (for all health states), AMI incidence and quality of life, and could be applied to all test strategies. Accuracy was thus assumed to be subgroup independent (equal to the base case values). The following subgroups were identified:

- Gender
- Age (45, 55, 65, 75 and 85)
- People with a history of previous NSTEMI. For this purpose, a proportion of 0% UA was assumed and the probabilities for the initial 'Post-MI' health state were used for the 'no ACS, no UA' health state and the probabilities for 'Post-MI with re-infarction' were used for the 'Post-MI' and 'Post-MI with re-infarction' health states. This subgroup analysis was only performed for the base case as for the secondary analysis this would lead to lower mortality probabilities for false positive patients compared with true negative patients (which seems implausible).

Subgroups with varying AMI prevalence (1%, 5%, 10%, 20%, 30%). In these analyses the no testing strategy was included as a comparator since a troponin test may not be indicated when clinical judgment assesses that the probability that a patient is experiencing an AMI is low. For the no testing strategy it is assumed that patients will be discharged immediately.

It should be noted that the main subgroup analyses (described above) differ from the subgroups described in the systematic review component of this assessment (see section 3.2.3.2), for which specific accuracy and prevalence data were available. Additional subgroup analyses were performed based on these subgroup-specific accuracy data. However, these analyses could only be performed for the Roche Elecsys hs-cTnT assayat presentation sample, using the 99<sup>th</sup> centile diagnostic threshold, compared with standard troponin testing; no subgroup-specific accuracy data were available for the other two hs-cTn assays. The following subgroups were considered:

- Age  $\leq$  70 and age >70
- Patients with pre-existing CAD and patients without pre-existing CAD
- Symptom onset <3 hours before presentation and symptom onset ≥3 hours before presentation

The subgroups with high pre-test probability and low to moderate pre-test probability were not considered as the prevalence data for these subgroups was unknown.

#### 4.4 Results of cost-effectiveness analyses

This section describes the results using probabilistic analyses for the base case analysis and the secondary analysis. In addition the sensitivity analyses (deterministic) and subgroup analyses are described (these deterministic analyses are also presented in tabulated form in Appendices 5 to 9.

#### 4.4.1 Base case analysis

The base case analysis includes six test strategies. Tables 17 and 18 show the probabilistic results of this analysis. Standard troponin testing was both most effective (15.101 life years, 11.730 QALYs) and most expensive (£2,697). The Abbott ARCHITECT hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was least effective (15.076 life years, 11.712 QALYs) and least expensive (£2,253). Compared to standard troponin testing, hs-cTn testing resulted in ICERs ranging between £90,725 and £24,019 savings per QALY lost.

Comparisons based on the next best alternative showed that for willingness to pay values below £6,600 per QALY, the Abbott ARCHITECT hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, would be cost-effective. For thresholds between £6,600 and £30,631 per QALY,

the Beckmann Coulter hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was cost-effective; above £30,631 per QALY the Abbott ARCHITECT hs-cTnI optimal strategy was cost-effective. Standard troponin becomes cost-effective at a threshold of £90,725 or higher (Table 18).

		Compared to Standard
Strategy	Life years	troponin
	15.076	
Abbott ARCHITECT hs-cTnl (99th centile at presentation)	(95% CI: 14.321 - 15.764)	-0.024
	15.085	
Roche Elecsys hs-cTnT (99th centile at presentation)	(95% CI: 14.332 - 15.770)	-0.016
	15.090	
Beckman Coulter hs-cTnI (99th centile at presentation)	(95% CI: 14.338 - 15.774)	-0.010
	15.091	
Roche Elecsys hs-cTnT optimal strategy	(95% CI: 14.340 - 15.776)	-0.009
	15.098	
Abbott ARCHITECT hs-cTnl optimal strategy	(95% CI: 14.351 - 15.780)	-0.003
	15.101	
Standard troponin	(95% CI: 14.356 - 15.781)	

Table 17: Probabilistic results for base case analysis: life years

At a willingness to pay threshold of £20,000 and £30,000 per QALY, the Beckmann Coulter hs-cTnl assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, had a probability of being cost-effective of 47% and 35% respectively. Although the Beckmann Coulter hs-cTnl assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was cost-effective at a willingness to pay threshold of £30,000 per QALY, the Abbott ARCHITECT hs-cTnl optimal strategy had the highest probability of being cost-effective (35%) at this threshold (Figure 16 and 17).

Charles			Compared to Standard		tandard	Compared to next best strategy			
Strategy			troponin		n				
	Co. etc. (05%( Cl))	0 41 1/2 (059/ 01)			ΔCosts /	C			ΔCosts /
	Costs (95% CI)	QALYS (95% CI)		ΔQALYS	ΔQALYS	Comparator	ΔCosts	ΔQALYS	ΔQALYS
	£2,253	11.712							
Abbott ARCHITECT hs-cTnl (99th	(95% CI: £1,702 -	(95% CI: 10.312 -							
centile at presentation)	£2,877)	13.157)	-£444	-0.018	£24,019				
	£2,296	11.718							
Roche Elecsys hs-cTnT (99th	(95% CI: £1,731 -	(95% CI: 10.319 -				Abbott ARCHITECT hs-cTnl (99th			Extendedly
centile at presentation)	£2,936)	13.165)	-£401	-0.012	£33,247	centile at presentation)	£42	0.006	dominated
	£2,324	11.723							
Beckmann Coulter hs-cTnl (99th	(95% CI: £1,755 -	(95% CI: 10.323 -				Abbott ARCHITECT hs-cTnl (99th			
centile at presentation)	£2,971)	13.172)	-£373	-0.008	£48,337	centile at presentation)	£71	0.011	£6,600
	£2,422	11.723							
Roche Elecsys hs-cTnT (optimal	(95% CI: £1,846 -	(95% CI: 10.326 -				Beckmann Coulter hs-cTnI (99th			Extendedly
strategy)	£3,077)	13.171)	-£275	-0.007	£38,528	centile at presentation)	£98	0.001	dominated
	£2,491	11.728							
Abbott ARCHITECT hs-cTnI	(95% CI: £1,908 -	(95% CI: 10.328 -				Beckmann Coulter hs-cTnI (99th			
(optimal strategy)	£3,148)	13.177)	-£206	-0.002	£90,725	centile at presentation)	£167	0.005	£30,631
	£2,697	11.730							
	(95% CI: £2,113 -	(95% CI: 10.334 -				Abbott ARCHITECT hs-cTnl			
Standard troponin	£3,359)	13.179)				(optimal strategy)	£206	0.002	£90,725

### Table 18: Probabilistic results for base case analysis: costs and QALYs

# Figure 16: Cost-effectiveness acceptability curve and incremental cost-effectiveness plane (incremental costs and QALYs compared to standard troponin) for base case analysis





Figure 17: Cost-effectiveness acceptability frontier for base case analysis

#### 4.4.2 Secondary analysis

The secondary analysis includes the same six test strategies. This analysis assumed that in a proportion of patients with a false positive hs-cTn test (i.e. positive hs-cTn test and a negative standard troponin test), there is prognostic significance (i.e. it is associated with an increased risk of adverse events (mortality and re-infarction)).

Standard troponin testing was least effective (14.785 life years, 11.464 QALYs) and most expensive (£3,058). The Abbott ARCHITECT hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold was the least effective hs-cTn test strategy (14.833 life years, 11.501 QALYs) and overall the least expensive strategy (£2,781). The Abbott ARCHITECT hs-cTnI optimal strategy was most effective (14.855 life years, 11.518 QALYs). Standard troponin testing was dominated by all hs-cTn testing strategies.

Comparisons based on the next best alternative showed that for willingness to pay values below £13,623 per QALY, the Abbott ARCHITECT hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was cost-effective. For thresholds between £13,623 and £14,562 per QALY, the Roche Elecsys hs-cTnT assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was cost-

effective; above £14,562 per QALY the Abbott ARCHITECT hs-cTnI optimal strategy was cost-effective (Tables 19 and 20).

Strategy	Life years	Compared to Standard troponin
	14.833	
Abbott ARCHITECT hs-cTnl (99th centile at presentation)	(95% CI: 14.104 - 15.487)	0.048
	14.837	
Roche Elecsys hs-cTnI (99th centile at presentation)	(95% CI: 14.111 - 15.491)	0.052
	14.839	
Beckman Coulter hs-cTnI (99th centile at presentation)	(95% CI: 14.114 - 15.488)	0.054
	14.843	
Roche Elecsys hs-cTnT (optimal strategy)	(95% CI: 14.119 - 15.494)	0.058
	14.855	
Abbott ARCHITECT hs-cTnl (optimal strategy)	(95% CI: 14.129 - 15.502)	0.070
	14.785	
Standard troponin	(95% CI: 14.061 - 15.436)	

Table 19: Probabilistic results	for secondary	analysis: life years
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At a willingness to pay threshold of £20,000 and £30,000 per QALY, the Abbott ARCHITECT hs-cTnI optimal strategy had the highest probability of being cost-effective (53% and 67% respectively; Figures 18 and 19).

#### **Compared to Standard** Compared to next best strategy Strategy troponin ∆Costs / ∆Costs / ΔQALYs **∆Costs** ΔQALYs **∆Costs** Costs (95% CI) QALYs (95% CI) ΔQALYs ΔQALYs Comparator £2,781 11.501 Abbott ARCHITECT hs-cTnI (99th (95% CI: £2,247 (95% CI: 10.087 Dominant centile at presentation) - 12.918) 0.037 - £3,388) -£277 £2,823 11.504 Roche Elecsys hs-cTnT (99th Abbott ARCHITECT hs-cTnI (99th (95% CI: £2,271 (95% CI: 10.092 Dominant centile at presentation) centile at presentation) - £3,442) - 12.920) -£235 0.040 £42 0.003 £13,623 £2,851 11.506 Beckmann Coulter hs-cTnI (99th Roche Elecsys hs-cTnT (99th (95% CI: £2,299 (95% CI: 10.093 Dominant Extendedly centile at presentation) centile at presentation) 0.042 £28 dominated - £3,477) - 12.923) -£207 0.001 £2,949 11.509 Roche Elecsys hs-cTnT (99th Roche Elecsys hs-cTnT (optimal (95% CI: £2,390 (95% CI: 10.095 Dominant Extendedly centile at presentation) strategy) - £3,579) - 12.926) £126 -£109 0.045 0.004 dominated £3,018 11.518 Abbott ARCHITECT hs-cTnl Roche Elecsys hs-cTnT (99th (95% CI: 10.103 (95% CI: £2,446 Dominant (optimal strategy) centile at presentation) - £3,659) - 12.936) -£39 0.054 £196 0.013 £14,562 11.464 £3,058 Abbott ARCHITECT hs-cTnl Standard troponin (95% CI: £2,485 (95% CI: 10.053 (optimal strategy) - £3,708) - 12.869) £39 -0.054 Dominated

#### Table 20: Probabilistic results for secondary analysis: costs and QALYs

# Figure 18: Cost-effectiveness acceptability curve and incremental cost-effectiveness plane (incremental costs and QALYs compared to standard troponin) for secondary analysis





Figure 19: Cost-effectiveness acceptability frontier for secondary analysis

#### 4.4.3 Sensitivity analysis

The deterministic analysis for the base case analysis is presented in Appendix 5. When it was assumed that the Post-MI re-infarction and mortality probabilities would remain elevated for untreated AMI for a life-time period, the Abbott ARCHITECT hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was cost-effective for thresholds below £1,642 per QALY, at which point the Beckman Coulter hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, became cost-effective up to a threshold of £7,602 per QALY. The Abbott ARCHITECT hs-cTnl optimal strategy was cost-effective for thresholds between £7,602 and £26,532 per QALY. Standard troponin testing was cost-effective for thresholds above £26,532 per QALY. Consistent with the base case analysis, all 'no doctor on demand' sensitivity analyses (one, two or three hours) showed that the Beckman Coulter hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was costeffective for thresholds between approximately £8,000 and £40,000 per QALY. Similarly, where the total delay decreased to 1.5 hours (and assuming availability of a doctor on demand), the Beckman Coulter hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was cost-effective for thresholds between £7,778 and £29,653 per QALY at which point the ARCHITECT hs-cTnl optimal strategy became cost-effective. Adding AMI treatment costs for the patients with a false positive test substantially impacted the results: standard troponin testing was cost-effective for all threshold values above £16,050 per QALY. Adding AMI treatment costs to the UA health state for the first year had a negligible impact on the incremental outcomes.

The following input parameters had a noticeable impact on the estimated cost-effectiveness: 30 day mortality for treated and untreated AMI (decision tree) and the mortality RR for treated versus untreated AMI (Markov trace). Varying the remaining parameters did not have a substantial impact on the results (i.e. the Beckman Coulter hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was cost-effective for thresholds between approximately £10,000 and £35,000 per QALY).

The deterministic analysis for the secondary analysis is presented in Appendix 6. When assuming that the post AMI re-infarction and mortality probabilities would remain elevated for untreated AMI for a life-time period, the Abbott ARCHITECT hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was cost-effective for thresholds below £1,853 per QALY, at which point the Roche Elecsys hs-cTnT assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, became cost-effective up to a threshold of £2,017 per QALY. The Beckman Coulter hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was cost-effective for thresholds between £2,017 and £5,889 per QALY. The Abbott ARCHITECT hs-cTnI optimal strategy was cost-effective for thresholds above £5,889 per QALY. For all 'no doctor on demand' sensitivity analyses, the Abbott ARCHITECT hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was costeffective for thresholds below £18,000 per QALY for one, two and three hours delay. The Roche Elecsys hs-cTnT assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was cost-effective for thresholds between £18,000 and £19,000, £20,000 and £22,000 per QALY in case of one, two and three hours delay respectively. The Abbott ARCHITECT hs-cTnI optimal strategy was cost-effective for higher thresholds. Similarly to the deterministic base case, where the total delay decreased to 1.5 hours (assuming availability of a doctor on demand), the Abbott ARCHITECT hs-cTnl assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was cost-effective for thresholds below £14,956 at which point the ARCHITECT hs-cTnl optimal strategy became cost-effective. Adding AMI treatment costs for all patients with a false positive test gave comparable results to the deterministic analysis: the Abbott ARCHITECT hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was cost-effective for all threshold values below £15,508 per QALY at which point the Abbott hs-cTnI optimal strategy became the preferred option. Adding AMI treatment costs to the UA health state for the first year had a negligible impact on the incremental outcomes.

The following input parameters had a noticeable impact on the estimated cost-effectiveness of the secondary analysis: increased test cost (of £40 per test), 30 day mortality for treated and untreated

98

AMI (decision tree), and the re-infarction and mortality RR for treated versus untreated AMI (Markov trace). Varying the remaining parameters did not have a substantial impact on the results.

#### 4.4.4 Subgroup analysis

Additional analyses were performed for subgroups based on age, gender, people with a history of previous NSTEMI, and AMI prevalence. These deterministic subgroup analyses (for the base case) analysis are presented in Appendix 7. Consistent with the base case analyses, analyses based on age and gender subgroups indicated that, up to an age of 75 year, the Beckman Coulter hs-cTnl assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was cost-effective for thresholds between approximately £10,000 and £35,000 per QALY. The Abbott ARCHITECT hs-cTnI optimal strategy was cost-effective for higher thresholds up to £115,000-£170,000, at which point standard troponin testing became cost-effective. For females aged over 85 years, the Beckman Coulter hs-cTnl assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was cost-effective for thresholds between £15,793 and £74,597 per QALY; the Abbott ARCHITECT hs-cTnI optimal strategy was cost-effective for thresholds between £74,597 and £259,592 per QALY and standard troponin testing was costeffective for thresholds of £259,592 per QALY and higher. For males aged over 85 years, the Abbott ARCHITECT hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was costeffective for thresholds below £28,711 per QALY; the Beckman Coulter hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was cost-effective for thresholds between £28,711 and £143,225 per QALY and the Abbott ARCHITECT hs-cTnI optimal strategy was cost-effective for thresholds between £143,225 and £503,476 per QALY, at which point standard troponin testing became cost-effective. The results for the subgroup with a history of previous NSTEMI were almost identical to the base case analysis.

For subgroup analyses considering AMI prevalence, no testing was included as additional comparator. For an AMI prevalence of 1%, the no testing strategy was cost-effective up to thresholds of £27,409 per QALY at which the Beckmann Coulter hs-cTnI (99<sup>th</sup> centile) test became cost-effective up to a threshold of £447,934 per QALY. For an AMI prevalence of 5%-20%, the no testing strategy was cost-effective up to thresholds of £8,759-£11,703 per QALY at which point the Beckmann Coulter hs-cTnI (99<sup>th</sup> centile) test became cost-effective up to thresholds of £32,042-£97,709 per QALY. For an AMI prevalence of 30%, the no testing strategy was cost-effective up to a threshold of £24,745 per QALY. The Abbott ARCHITECT hs-cTnI optimal strategy was cost-effective for thresholds between £24,745 and £70,942 per QALY.

In addition, cost-effectiveness estimates for the subgroups, described in section 3.2.3.2, based on subgroup-specific accuracy and prevalence are reported in Appendix 9 (only comparing the Roche Elecsys hs-cTnT assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, and standard troponin testing). The results of these analyses indicated that differences in accuracy and AMI prevalence between subgroups had a substantial impact on the cost-effectiveness of the Roche Elecsys hs-cTnT assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, compared with standard troponin testing (ICER range: £22,111-£355,571; deterministic base case: £41,233).

The deterministic subgroup analyses for the secondary analysis are presented in Appendix 8. For females aged 45 and males aged 45 or 55, the Abbott ARCHITECT hs-cTnl assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was cost-effective for thresholds below £16,023-£17,836 per QALY. The Abbott ARCHITECT hs-cTnI optimal strategy became cost-effective for higher thresholds. For females aged 55 or 65 and males aged 65 or 75, the Abbott ARCHITECT hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was cost-effective for thresholds below £13,064-£16,994 per QALY. From this threshold up to £18,999-£25,149 per QALY the Roche Elecsys hs-cTnT assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was most cost-effective. The Abbott ARCHITECT hs-cTnl optimal strategy was cost-effective for higher thresholds. For females aged 75 or 85, the Abbott ARCHITECT hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was cost-effective up to thresholds of £12,392-£21,140 per QALY, at which point the Roche Elecsys hs-cTnT assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, became cost-effective up to thresholds of £16,407-£26,911 per QALY. The Abbott ARCHITECT hs-cTnI optimal strategy became cost-effective for thresholds higher than £24,020-£45,709 per QALY. For males aged 85, the Abbott ARCHITECT hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was cost-effective for thresholds below £66,418 per QALY. The Abbott ARCHITECT hs-cTnI optimal strategy became cost-effective for higher thresholds.

For subgroup analyses considering AMI prevalence, no testing was included as additional comparator. For an AMI prevalence of 1%, the no-testing strategy was cost-effective up to a threshold of £4,563 per QALY at which point the Abbott ARCHITECT hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, became cost-effective up to a threshold of £109,991 per QALY where the Abbott ARCHITECT hs-cTnI optimal strategy became cost-effective. Similarly, for an AMI prevalence of 5% and 10% the thresholds were £5,209 and £35,574 and £5,820 and £22,684 respectively. For an AMI prevalence of 20% and 30%, the Abbott ARCHITECT hs-cTnI optimal strategy was cost-effective for thresholds above £16,319 and £15,410 respectively.

100

In contrast with the base case analysis (described above), the subgroup–specific accuracy and prevalence (only comparing the Roche Elecsys hs-cTnT assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, and standard troponin testing) did not have an important impact on the cost-effectiveness (Appendix 9). The Roche Elecsys hs-cTnT assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was dominant for all subgroups.

#### 5. DISCUSSION

#### 5.1 Statement of principal findings

#### 5.1.1 Clinical effectiveness

All 18 studies (37 publications) included in the systematic review assessed the accuracy of one or more hs-cTn tests for the diagnosis of any AMI or for NSTEMI. There were no controlled trials comparing clinical outcomes in people assessed using hs-cTn tests to those assessed using conventional Tn assays. The majority (15/18) of the included studies reported data for the Roche Elecsys hs-cTnT assay; four studies reported data for the Abbott ARCHITECT hs-cTnI assay and two reported data for pre-commercial versions of the Beckman Coulter Access hs-cTnI assay. Not all of the included studies reported data on accuracy for the diagnosis of NSTEMI (i.e. for a population which excluded people with STEMI), which was the target population for this assessment. However, where data were available for both any AMI (population with symptoms suggestive of ACS) and NSTEMI (population which excluded people with STEMI), estimates of test performance were generally similar, see Table 4 section 3.2.3 and Table 6 section 3.2.5.

When diagnosis was based on a single sample taken at presentation, using the 99<sup>th</sup> centile for the general population as the diagnostic threshold, positive LRs derived from summary estimates of sensitivity and specificity indicated that neither the Roche Elecsys hs-cTnT assay or the Beckman Coulter Access hs-cTnl would be adequate to rule-in a diagnosis of NSTEMI. The LR+ for the Roche Elecsys hs-cTnT assay was 5.41 (95% CI: 3.40 to 8.63) and the LR+ for the Beckman Coulter Access hscTnI was 3.67 (95% CI: 3.26 to 4.13). By contrast, the LR+ for the Abbott ARCHITECT hs-cTnI assay, in a population which did not exclude STEMI, was 11.47 (95% CI: 9.04 to 16.19), indicating that a positive test using this assay may have some utility in confirming a diagnosis of AMI. The corresponding LR-s indicated that a negative test result on a single sample taken at presentation, using the 99th centile for the general population as the diagnostic threshold, would not be adequate to rule-out NSTEMI using any of the three assays assessed. LR- was 0.15 (95% CI: 0.08 to 0.26) for the Roche Elecsys hs-cTnT, 0.11 (95% CI: 0.07 to 0.17) for the Beckman Coulter Access hs-cTnI, and 0.22 (95% CI: 0.16 to 0.27), for the Abbott ARCHITECT hs-cTnl assay. Although, these LRs are fairly low, the consequences of missing an AMI are so great that a test needs to be able to rule out an AMI with a very high degree of certainty. It should be noted that the Beckman Coulter hs-cTnl assay evaluated in the APACE study was described as 'an investigational prototype';<sup>75</sup> the 99<sup>th</sup> centile (9 ng/L), described as 'acording to the manufacturer', differs from the 99<sup>th</sup> centile given in the current  $ng/L),^{16}$ information leaflet product (40

hypothetical cohort of 1,000 people is considered, assuming a prevalence of NSTEMI of 17% (derived from studies included in our systematic review, see section 3.2.3.2), the estimated number of people with AMI and a negative test result who would be erroneously discharged based on this testing protocol is 20 for the Roche Elecsys hs-cTnT assay, 14 for the Beckman Coulter Access hs-cTnI assay and 34 for the Abbott ARCHITECT hs-cTnI assay.

Some limited data were available on the diagnostic performance of the Roche Elecsys hs-cTnT assay in clinical subgroups, using a single sample taken at presentation and the 99<sup>th</sup> centile diagnostic threshold. These data indicated a lower LR- when the test is used in certain population groups, (e.g. people over 70 years of age LR- 0.05, 95% CI: 0.02 to 0.18, people without pre-existing CAD LR- 0.07, 95% CI: 0.04 to 0.16) and with a high pre-test probability (determined by clinical judgement based on cardiovascular risk factors, type of chest pain, physical findings, and ECG abnormalities; LR- 0.09, 95% CI: 0.02 to 0.45). Using the hypothetical cohort of 1,000 people described above, the estimated number of people with AMI and a negative test result who would be erroneously discharged if the test were used to rule-out AMI in these selected populations is five for people over 70 years of age, 10 for people without pre-existing CAD, and 10 for people with a clinical assessment of high pre-test probability. When the performance of the Roche Elecsys hs-cTnT assay was assessed in a population restricted to people who presented more than three hours after the onset of symptoms, a similar fall in the LR- was observed (LR- 0.08, 95% CI: 0.05 to 0.11); the estimated number of people with AMI and a negative test resouly discharged if the test were used to rule-out AMI in these selected number of people with AMI and a negative test resouly discharged if the test were used to rule-out AMI in the hours after the onset of symptoms, a similar fall in the LR- was observed (LR- 0.08, 95% CI: 0.05 to 0.11); the estimated number of people with AMI and a negative test result who would be erroneously discharged if the test were used to rule-out AMI in this populations is 10.

We constructed optimal testing strategies for the Roche Elecsys hs-cTnT assay (Figure 9, section 3.2.3.4) and for the Abbott ARCHITECT hs-cTnI assay (Figure 11, section 3.2.4.4). Both strategies employ a two step process, which provides two potential oportunities to rule-out AMI and hence to discharge patients within the four hour window specified in the scope for this assessment. This potential is conditional upon the acheivement of short (<1 hour) turnarround times for hs-cTn testing, as recommended by the joint National Academy of Clinical Biochemistry and IFCC guidelines on troponin testing<sup>104</sup> and in line with clinical opinion; a study of 1,355 emergency department physicians in the USA indicated that 75% believed that the results of troponin testing should be available to them within 45 minutes.<sup>105</sup> The initial step for both the Abbott ARCHITECT hs-cTnI optimal strategy and Roche Elecsys hs-cTnT optimal strategy was based on the use of an LoB (3 ng/L) diagnostic threshold in a sample taken at presentation and was selected for optimal rule-out potential (low negative LR), regardless of poor rule-in performance. For the Roche Elecsys hs-cTnT optimal strategy, the second step involves an additional sample taken two to three hours after

admission and was selected to provide the best possible combination of rule-out and rule-in performance. Using the hypothetical cohort of 1,000 people, previously described, the intial step of the prososed Roche Elecsys hs-cTnT optimal strategy would result in discharge of 407 people, nine of whom would have been erroneously discharged with AMI. The second step of this strategy involves a combination of testing on admission and after two hours, where a negative result is defined as both no sample above the 99<sup>th</sup> centile AND a change of <20% over two hours and provides the optimum rule-out performance (LR- 0.04, 95% CI: 0.02 to 0.10); conversely, a positive result is defined as both a peak value above the 99<sup>th</sup> centile AND a change of >20% over two hours and provides the optimum rule-in performance (LR+ 8.42 (95% CI: 6.11 to 11.60)). Application of the ruleout component of the second step would result in discharge of a further 286 people, five of whom would have been erroneously discharged. For the proposed Abbott ARCHITECT hs-cTnl optimal strategy, the initial rule-out step would result in discharge of 291 people all of whom would have been appropriately discharged. The second step of this strategy involves repeat testing on a sample taken three hours after admission, using the 99<sup>th</sup> centile diagnostic threshold. Application of the rule-out component of the second step would result in discharge of a further 489 people, three of whom would have been erroneously discharged. Available data on the Beckman Coulter hs-Tnl assay were insufficient to support construction of an optimal testing strategy.

#### 5.1.2 Cost-effectiveness

The review of economic analyses of hs-cTn (i.e. either hs-cTnI or hs-cTnT) testing for the early ruleout of AMI in people with acute chest pain found four HTA reports, two full papers and one abstract. Based on all of these publications, it can be said that, in general, the question of whether hs-cTn testing is cost-effective cannot yet be answered unequivocally. The majority of papers reported substantial ICERs, with considerable uncertainty. In particular, the accuracy of high-sensitive tests as well as the efficiency of decision-making based on test results were found to be important drivers of cost-effectiveness.

In our health economic analysis, the cost-effectiveness of different testing strategies involving hs-cTn for the early rule-out of AMI in people with acute chest pain presenting to the ED with suspected ACS and STEMI ruled out was assessed. All analyses had the same comparator: standard troponin testing at 10-12 hours, which is considered the reference standard and therefore was assumed to have perfect sensitivity and specificity. In addition to the base case analysis, given some evidence that false positives versus this reference standard also have a poor prognosis, a secondary analysis was conducted which assumed an increased adverse event risk for patients with false positive hs-cTn tests. A number of subgroup and sensitivity analyses were also performed.

104

In the base case analysis, standard troponin testing was both most effective and most costly. Strategies considered cost-effective depending upon ICER thresholds were Abbott ARCHITECT hscTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, (thresholds below £6,597), Beckman Coulter hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, (thresholds between £6,597 and £30,042), Abbott ARCHITECT hs-cTnI optimal strategy (LoD threshold at presentation, followed by 99<sup>th</sup> centile threshold at three hours) (thresholds between £30,042 and £103,194), and the standard troponin test (thresholds over £103,194). The Roche Elecsys hs-cTnT assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold and the Roche Elecsys hs-cTnT optimal strategy (LoB threshold at presentation followed by 99<sup>th</sup> centile threshold and/or  $\Delta 20\%$  (compared to presentation test) at 1-3 hours) were extendedly dominated in this analysis (one of the more effective strategies was better value in that the ICER was lower).

In the secondary analysis, which assumed a proportion of false positives in the hs-cTn testing strategies had an increased risk of adverse events, standard troponin was least effective and most costly, and therefore a dominated strategy. The most effective strategy here was the Abbott ARCHITECT hs-cTnI optimal strategy. The Roche Elecsys hs-cTnT optimal strategy was extendedly dominated (one of the more effective strategies was better value in that the ICER was lower), as was the Beckman Coulter hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, in this analysis. Strategies considered cost-effective were Abbott ARCHITECT hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, (thresholds below £12,217), Roche Elecsys hs-cTnT assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, (thresholds between £12,217 and £14,992) and Abbott ARCHITECT hs-cTnI optimal strategy (thresholds over £14,992).

Sensitivity analyses showed that in general, there were no major changes in the relative costeffectiveness of strategies. That is, dominancy and order of relative cost-effectiveness were comparable, although the ICERs were different. Exceptions included assuming that the increased 30 day mortality for treated versus untreated MI applied to a lifetime (instead of only during the first year after presentation at ED), which meant that standard troponin could be cost-effective from a threshold of £26,352 or higher. The same assumption applied to the secondary analysis meant that the Beckman Coulter hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, strategy was no longer extended dominated but was considered cost-effective at thresholds between £2,017 and £5,889. Another sensitivity analysis that resulted in substantial changes was assigning AMI treatment costs to patients who tested false positive. In the base case, under this assumption, standard troponin became cost-effective at an ICER threshold of £20,000 (ICER £16,050 as compared to the Abbott ARCHITECT hs-cTnI optimal strategy). In the secondary analysis, however, assigning treatment costs to false positive patients did not impact the position of standard troponin; it was still dominated by another strategy i.e. less effective and more costly.

Subgroup analyses (with non-subgroup specific accuracy data) for the base case showed that ICERs compared to the next best strategy were slightly higher for males at all ages. Also, for both females and males, ICERs increased with age. In addition, from ages 55 upwards (base case 53), the Roche Elecsys hs-cTnT assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, became extendedly dominated. In the subgroup with previous NSTEMI, again the Roche Elecsys hs-cTnT assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was extendedly dominated and ICERs are slightly higher as compared to the whole group. Subgroup analysis based on MI prevalence (including a no testing strategy) indicated that only when MI prevalence is as low as 1% (base case 17%) was the no testing strategy considered cost-effective up to an ICER threshold of £27,409 after which the Beckman Coulter hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, strategy takes over. The higher the prevalence, the lower the point at which the Beckman Coulter hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, strategy became cost-effective (i.e. £11,703 for prevalence 5%, £9,740 for prevalence 10% and £6,597 for 17%).

For the secondary analysis, again, the ICERS for males were slightly higher than for females. For the various age categories, results were rather diffuse, but as in the base case ICERs appeared to increase with age. There did not appear to be a substantial difference between the MI prevalence subgroups, that is, the no testing strategy was only cost-effective up to rather modest ICER thresholds (£4,563-£7,109) for all values of prevalence.

The subgroup analyses using subgroup-specific accuracy and prevalence could only be performed for the Roche Elecsys hs-cTnT assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold as there were no subgroup data on Beckman Coulter hs-cTnI and Abbott ARCHITECT hs-cTnI assays. The comparator was the standard troponin at 10-12 hours, which was assumed to have perfect sensitivity and specificity. For the base case, the Roche Elecsys hs-cTnT assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, was always less costly and less effective, but ICERs were more favorable for the following subgroups as compared to their counterparts: Age  $\leq$ 70, with pre-existing CAD, and symptom onset <3 hours. For the secondary analysis, the standard troponin was dominated by the Roche Elecsys hs-cTnT assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, overall, as this test was both less costly and more effective. However, the subgroups which rendered the highest savings per QALY gained were consistent with the base case analysis i.e. Age  $\leq$ 70, with pre-existing CAD, and symptom onset <3 hours. Although data are lacking, it seems likely that these differences between subgroups can be extrapolated, at least partly, to the other tests considered in the base case analysis.

#### 5.2 Strengths and limitations of assessment

#### 5.2.1 Clinical effectiveness

Extensive literature searches were conducted in an attempt to maximise retrieval of relevant studies. These included electronic searches of a variety of bibliographic databases, as well as screening of clinical trials registers and conference abstracts to identify unpublished studies. Because of the known difficulties in identifying test accuracy studies using study design-related search terms,<sup>106</sup> search strategies were developed to maximise sensitivity at the expense of reduced specificity. Thus, large numbers of citations were identified and screened, relatively few of which met the inclusion criteria of the review.

The possibility of publication bias remains a potential problem for all systematic reviews. Considerations may differ for systematic reviews of test accuracy studies. It is relatively simple to define a positive result for studies of treatment, e.g. a significant difference between the treatment and control groups which favours treatment. This is not the case for test accuracy studies, which measure agreement between index test and reference standard. It would seem likely that studies finding greater agreement (high estimates of sensitivity and specificity) will be published more often. In addition, test accuracy data are often collected as part of routine clinical practice, or by retrospective review of records; test accuracy studies are not subject to the formal registration procedures applied to randomised controlled trials and are therefore more easily discarded when results appear unfavourable. The extent to which publication bias occurs in studies of test accuracy remains unclear, however, simulation studies have indicated that the effect of publication bias on meta-analytic estimates of test accuracy is minimal.<sup>107</sup> Formal assessment of publication bias in systematic reviews of test accuracy studies remains problematic and reliability is limited.<sup>27</sup> We did not undertake a statistical assessment of publication bias in this review. However, our search strategy included a variety of routes to identify un-published studies and resulted in the inclusion of a number of conference abstracts.

Clear inclusion criteria were specified in the protocol for this review, a copy of which is provided in Appendix 6. The eligibility of studies for inclusion is therefore transparent. In addition, we have provided specific reasons for exclusion for all of the studies which were considered potentially relevant at initial citation screening and were subsequently excluded on assessment of the full publication (Appendix 4). The review process followed recommended methods to minimise the potential for error and/or bias;<sup>25</sup> studies were independently screened for inclusion by two
reviewers and data extraction and quality assessment were done by one reviewer and checked by a second (MW and PW). Any disagreements were resolved by consensus.

Studies included in this review were assessed for risk of bias and applicability using the QUADAS-2 tool developed by the authors<sup>33</sup> and recommended by the Cochrane Collaboration.<sup>27</sup> QUADAS-2 is structured into four key domains covering participant selection, index test, reference standard, and the flow of patients through the study (including timing of tests). Each domain is rated for risk of bias (low, high, or unclear); the participant selection, index test and reference standard domain are also, separately rated for concerns regarding the applicability of the study to the review question (low, high, or unclear). The results of the QUADAS-2 assessment are reported, in full, for all included studies in Appendix 3 and are summarised in section 3.2.2. The main potential sources of bias in the studies included in this assessment were related to patient spectrum and patient flow (QUADAS domains 1 and 4). Reporting of the participant selection process was frequently unclear; a further study was rated as unclear for this domain as a large number of patients were not enrolled due to 'technical reasons' that were not fully defined and so it was not possible to judge whether these constituted inappropriate exclusions.<sup>54</sup> The most common feature of studies rated as 'high risk of bias' for patient selection was the inclusion of participants based on staffing or work flow considerations, e.g. participants were excluded if they presented at night or during busy periods.<sup>41, 45,</sup> <sup>50</sup> All ratings of 'high risk of bias' for patient flow were due to high proportions of withdrawals. There were also concerns regarding the applicability of the patient population and the reference standard in some of the included studies. The main area of concern, with respect to population, was for studies that enrolled mixed populations (i.e. when the target condition was any AMI); because the primary focus of this assessment was the diagnosis of NSTEMI in populations where patients with STEMI were excluded (i.e. target condition NSTEMI), the primary focus was the population of patients with STEMI excluded, mixed population studies which were not restricted to this specific patient group were considered to have high concerns regarding applicability. However, as noted above (section 5.1.1), where data were available for both any AMI (mixed population) and NSTEMI (population which excluded people with STEMI), estimates of test performance were generally similar. In accordance with current NICE guidance,<sup>11</sup> our review question specified that an appropriate reference standard had to include a standard Tn measurement at baseline and at 10-12 hours after the onset of symptoms in 80% of the population. Although studies generally included a baseline and a second, later standard Tn measurement, only five<sup>41, 50, 62, 65, 75</sup> met the specific timing criterion for the second standard Tn measurement; studies which did not meet this criterion were classified as having high concerns regarding applicability.

We identified one recently published systematic review which included an assessment of the accuracy of hs-cTn assays for the diagnosis of AMI and prediction of MACE.<sup>7</sup> This review, by Goodacre et al, also evaluated standard cTn assays (alone and in combination with other cardiac biomarkers) and the diagnostic accuracy of other cardiac biomarkers, as well as including prediction modelling studies, all of which were outside the scope of this assessment. Our systematic review represents an advance on Goodacre et al as it provides a more up-to-date and comprehensive assessment of the performance of hs-cTn assays. Although the Goodacre review was published in 2013, search dates were reported as 1995 to November 2010; hence it only included two studies, which met the definition of an hs-cTn assay used in our assessment.<sup>56, 73</sup> Both of these studies assessed the diagnostic performance of the Roche Elecsys hs-cTnT assay when applied to a single sample taken at presentation, using the 99<sup>th</sup> centile diagnostic threshold, and neither excluded participants with STEMI. Both studies were also included in our systematic review and one<sup>56</sup> contributed data to our summary estimates (based on a total of 15 studies) of the performance of the Roche Elecsys hs-cTnT assay for the diagnosis of any AMI at this threshold studies; the other was an early publication of the APACE study,<sup>73</sup> the most recent publication from which contributed data to our main analysis (accuracy for the diagnosis of NSTEMI), which included a total of six studies.<sup>75</sup> The summary estimate of sensitivity derived from our systematic review was lower (88% for both any AMI or NSTEMI analyses) than that reported by the Goodacre review (96% for any AMI),<sup>7</sup> and our summary estimate of specificity was higher (82% for any AMI and 84% for NSTEMI) than that reported by the Goodacre review (72% for any AMI).<sup>7</sup> A pre-publication copy of a more recent academic-in-confidence systematic review, provided by the authors as material,



Our assessment represents an advance on both of these systematic reviews in that we provide up-todate estimates of the diagnostic performance of assays meeting a strict definintion for hs-cTn, which are stratified by hs-cTn assay type, diagnostic threshold and timing of the Tn test.

We believe that our assessment provides information of direct relevance to UK clinical practice as we focus on the performance of hs-cTn within the four hour time window corresponding to the target for NHS emergency departments, which specifies that 'no one should be waiting more than four hours in the emergency department from arrival to admission, transfer or discharge.<sup>91</sup> Furthermore, we have used the data from our systematic review to propose strategies for how hscTn assays might be applied and interpreted in order to maximise diagnostic performance. These strategies were devised with consideration to test timing, diagnostic threshold and interpretation of combinations of multiple test results. One limitation of this approach is that our estimates of the effectiveness and cost-effectiveness of the proposed two step strategies require the assumption that the diagnostic performance of the second step is the same when used in people in whom NSTEMI is not ruled out by the first step as it is when used in the whole population (see sections 3.2.3.4 and 3.2.4.4). This assumption was necessary because no combined test performance data were available for the proposed strategies. However, it can be argued that the assumption is reasonable as the first step in both strategies focuses on rule-out performance and thus has a low positive LR. This means that there is a relatively small change in the prevalence of AMI between the first and second steps (17% to 27% for the Roche Elecsys hs-cTnT optimal strategy and 17% to 24% for the Abbott ARCHITECT hs-cTnl optimal strategy).

Our assessment was less comprehensive for the Abbott ARCHITECT hs-cTnI assay and the Beckman Coulter hs-cTnI assay than for the Roche Elecsys hs-cTnT, because available data were limited for these two assays.

#### 5.2.2 Cost-effectiveness

Our cost-effectiveness analysis is the most comprehensive to date in terms of the number of relevant hs-cTn test strategies for the early rule-out of AMI in people presenting to the ED with acute chest pain and suspected ACS. Moreover, the de novo probabilistic model was based on one previously developed for a published and peer reviewed HTA.<sup>84</sup> This model was also used in a later assessments on the cost-effectiveness of biomarkers in patients with suspected ACS.<sup>65</sup> For the present analysis, a number of adjustments were made to the model, but most of the assumptions were maintained.

The model was also informed by a comprehensive, high quality systematic review of diagnostic test accuracy. Additional parameters were either those from the original HTA model, or any of the further assessments, or, where necessary, were based on a pragmatic literature review. Such a review is standard practice in economic modeling given the large number of parameters required and we expect that the review has delivered the most relevant information given that it focused on identifying the most recent large UK based studies.

As in any economic model, a number of major and minor assumptions had to be made. It is important to understand the impact of these assumptions in order to correctly interpret the results of the model. The impact of most assumptions has been explored in sensitivity and secondary analyses. However, one major assumption that was maintained throughout all analyses was the conservative assumption of no health benefit of early treatment in the hs-cTn strategies as compared to 'late' treatment in the standard cTn strategy. Although many experts believe that there must be a benefit, at least to some extent, of treating patients early, there is no evidence to support or quantify a timing effect, as yet. In addition, there may well also be adverse effects associated with early treatment also, (e.g. the risk of bleeding, unnecessary PCIs, etc.). The Canadian HTA report<sup>89</sup> identified in the economic review (section 4.1.4.4) did include an advantage for early versus late treatment, based on one study, which investigated the effect of a 36 hour treatment delay.<sup>109</sup> The RR found in this study was then recalculated, assuming a constant effect of timing on treatment benefit, to a RR of 1.035 of mortality for a treatment delay of six hours versus early treatment, was again adjusted to 1.01 based on expert opinion. Any possible adverse effect of early treatment was not considered in this analysis. A similar approach would have been possible in the present model, but in our view, this would not be informative, given the level of uncertainty underlying this final estimate. Therefore, it was decided to leave out a possible effect of timing of treatment. This could be considered a conservative approach, but even this is uncertain.

The assumption that standard troponin, as the reference standard, has perfect sensitivity and specificity was also maintained throughout all analyses. Although a simplification, given that the actual reference standard is standard troponin plus clinical information, this approach is consistent with previous modeling and incorporation of the effect of clinical information to the hs-cTn test would be very difficult, given the current lack of data. To some extent, clinical judgment might already be incorporated into the modeling because, for the effect of treatment (RR for re-infarction and mortality), the study performed by Mills et al was used.<sup>86</sup> In this study not all patients with negative tests results were left untreated; we might therefore speculate that, where patients who tested negative were treated, this was because of clinical judgment. However, we cannot be certain that the observations from this trial reflect the true contribution of clinical judgment. On the other hand, there is recent evidence that the prognostic performance of standard troponin testing may be imperfect. For example, a negative troponin test might assess correctly that a patient is not experiencing a NSTEMI, but some patients with negative test results may still benefit from

treatment. To take this possibility into account, a secondary analysis was performed, which resulted in the standard troponin strategy being dominated by the hs-cTn testing strategies. In other words, it seems reasonable to conclude that not only might hs-cTn be cost effective, it might also be more effective than standard troponin.

Another assumption, which was varied in sensitivity analysis, with a rather substantial impact on results, was how to attribute costs of treatment to patients testing false positive in the hs-cTn treatment strategies. In the base case analysis, false positive patients were assigned survival, quality of life, and costs of true negative patients, i.e. they were basically assumed not to be treated. However, if hs-cTn assays were incorporated in clinical practice, patients with a positive result would be treated, at least up to the point where it is discovered they were false positive. Therefore, in a sensitivity analysis, false positive patients were assigned treatment costs as if they were true positive, but mortality and quality of life as if they were true negative. For the base case, this would change results quite dramatically, as the hs-cTn strategies would become more expensive but not more effective, whereas for the standard troponin nothing would change. For the secondary analysis (some hs-cTn false positives need and get treatment) things are different, since in this case treatment costs would be incurred for a proportion of patients (5%), but these patients would also receive the benefits of treatment. This approach had a very limited effect on results, in terms of strategies that were cost-effective. In our opinion, the secondary analysis, which assigns treatment costs to all false positives, but also assumes that some of these patients benefit treatment, is the most plausible scenario.

## 5.3 Uncertainties

## 5.3.1 Clinical effectiveness

The performance of any test that uses the 99<sup>th</sup> centile for the general population as the diagnostic threshold will be dependent upon the characteristics of the reference population from which this value was derived. Although the product information leaflet for the Abbott ARCHITECT hs-cTnI assay recommends that 'each laboratory should verify that the 99<sup>th</sup> centile is transferable to its population or establish its own 99<sup>th</sup> centile', <sup>15</sup> test accuracy data included in the assessment are predominantly based on the 99<sup>th</sup> centiles for the three assays (Roche Elecsys hs-cTnT, Abbott ARCHITECT hs-cTnI, Beckman Coulter hs-cTnI) as reported by their respective manufacturers. <sup>15, 16, 18</sup> The 99<sup>th</sup> centile for the Roche Elecsys hs-cTnT was reported as being derived from a study population of 616 apparently healthy volunteers and blood donors, with an age range of 20 to 71 years and equal proportions of males and females; <sup>110</sup> no further details were reported. The 99<sup>th</sup> centile for the Abbott ARCHITECT hs-cTnI assay was described being derived from a study of '1,531 apparently healthy individuals in

a US population with normal levels of BNP, HbA1c, and estimated GFR values'.<sup>15</sup> Although a 2012 'in press' reference for this study was given in the APACE study,<sup>75</sup> we were not able to identify any corresponding publication. It should also be noted that the Beckman Coulter hs-cTnI assay evaluated in the APACE study was described as 'an investigational prototype';<sup>75</sup> the 99<sup>th</sup> centile (9 ng/L), described as 'according to the manufacturer', differs from the 99<sup>th</sup> centile given in the current product information leaflet (40 ng/L).<sup>16</sup> The product information leaflet describes this value as being derived from general practice samples obtained from London, UK, and the surrounding area; samples were from 1,000 people over 40 years of age, with approximately equal numbers of males and females, and samples from people with abnormal urea and electrolytes, liver function tests, glucose, or NT-proBNP, were excluded.<sup>16</sup> Expected values, and hence diagnostic thresholds derived from groups of healthy volunteers may have limited applicability to the population in whom hs-cTn testing would be applied in practice, e.g. with respect to age range. Data provided in the product information leaflets for the Roche Elecsys hs-cTnT assay and the Abbott ARCHITECT hs-cTnI assay both indicated that 99<sup>th</sup> centile values differed between males and females; the Abbott ARCHITECT hs-cTnI assay reported values of 15.6 ng/L and 34.2 ng/L for females and males, respectively,<sup>15</sup> and the Roche Elecsys hs-cTnT assay reported values of 10.0 ng/L and 14.2 ng/L for females and males, respectively.<sup>18</sup> Despite this we were unable to identify any data on whether the diagnostic performance of tests varies according to sex, when a single common diagnostic threshold is used for both males and females; the effectiveness of using sex-specific diagnostic thresholds therefore remains uncertain. Similarly, we were unable to identify any data on the diagnostic performance of hs-cTn assays when used in people with impaired renal function.

Differences in the populations used to derive the 99<sup>th</sup> centile diagnostic threshold, and hence in the Tn level at which this threshold set, may also affect the ability of an assay to acheive the first point of the accepted definition of a hs-cTn assay, i.e. a CV of  $\leq 10\%$  at the 99<sup>th</sup> centile for the general population. A standardised definition of the required reference population would be useful in ensuring a 'level playing field' for classification of assays as 'high sensitive' and would aid comparisons between tests.

We identified some data on the diagnostic performance of hs-cTn testing in clinically important subgroups (older people,<sup>52, 75</sup> and people with and without pre-existing CAD).<sup>46, 75</sup> However, these data were very limited and were only available for the Roche Elecsys hs-cTnT assay. Therefore, there remains some uncertainty about how the diagnostic performance of individual hs-cTn assays may vary in clinically relevant subgroup, as well as what may constitute the optimal testing strategy in these groups.

113

A significant limitation of this assessment follows from the design of the primary studies included in the systematic review. The objective of these studies was to evaluate the diagnostic performance of hs-cTn assays when compared to a reference standard based on the universal definition of AMI endorsed the European Cardiology Society, the American Colledge of Cardiology, the American Health Association and the World Heart Federation.<sup>8, 21, 22</sup> The scope for this assessment did not include studies which evaluated the use of hs-cTn testing in combination with other tests, thus, studies which assessed the combined accuracy of a clinical risk score and a hs-cTn test used together would have been excluded, however, we did not identify any studies which were excluded on this basis. Studies assessing the diagnostic performance of a hs-cTn test alone, where participants were subgrouped by clinical risk, met our inclusion criteria and were included in the systematic review. We identified only one study of this type,<sup>48</sup> which, as described in section 5.1.1, indicated that the rule-out performance of hs-cTnT testing may be improved if the test is used in a population with high clinically determined pre-test probability. There remains uncertainty arround how hs-cTn testing would perform if used, as it would be in clinical practice, in combination with a clinical assessment of pre-test probaility (with or without formal risk scoring). Full assessment of the independent predictive value of hs-cTn testing requires multivariable prediction modelling.

A final area of uncertainty exists with respect to the clinical significance of a 'false positive' hs-cTn result, i.e. does a positive hs-cTn result imply a clinically important change in cardiac risk, where a diagnosis of AMI is not confirmed (based on standard Tns and the universal definition)? Re-adjudication of the final diagnosis, using later hs-cTn measurements in place of the conventional Tn results, can provide some insight into this issue. The most recent publication from the APACE study reported that when hs-cTnT results (including a six hour time point) were included in the reference standard diagnosis, this resulted in 131 participants being classified as having had a small AMI, which would have been classified as 'no AMI' where adjudication was based on standard Tn results.<sup>75</sup>

## 5.3.2 Cost-effectiveness

The main uncertainties for the cost-effectiveness analysis lie in the model assumptions, particularly regarding the effect of actual clinical practice in terms of both other diagnostic information and treatment given this information. Although many of these assumptions have been varied in one-way sensitivity analysis, the precise implication of false negative test results, where patients are discharged without essential treatment or of false positive test results, where patients stay in hospital and may receive unnecessary interventions, is unknown.

114

It should also be emphasised that the uncertainty resulting from the above mentioned assumptions was not parameterised in the model and is therefore not reflected in the probabilistic sensitivity analyses or in the cost-effectiveness acceptability curves.

### 6. CONCLUSIONS

#### 6.1 Implications for service provision

We propose the use of two step testing strategies to optimise the diagnostic performance of hs-cTn testing. There is evidence to suggest that undetectable levels of Tns (below the LoB/LoD of the assay) on presentation, measured using the Roche Elecsys hs-cTnT assay or the Abbott ARCHITECT hs-cTnI assay, may be sufficient to rule out NSTEMI in people presenting with symptoms suggestive of ACS. There is also evidence to suggest that a further rule-out step may be possible, within the four hour NHS emergency department target. For the Abbott ARCHITECT hs-cTnI assay, this second rule-out step would be based on a Tn level below the 99<sup>th</sup> centile in a sample taken three hours after presentation. For the Roche Elecsys hs-cTnT assay, the second rule-out step would be based on a Tn level below the 99<sup>th</sup> centile in all samples AND a change in Tn level of <20% between presentation and two hours. There is insufficient evidence to determine an optimal testing strategy for the Beckman Coulter hs-cTnI assay. There is some limited evidence to suggest that a Tn level below the 99<sup>th</sup> centile on presentation, measured using the Roche Elecsys hs-cTnT assay, may be sufficient to rule out NSTEMI in some groups (people over 70 years old, people without pre-existing CAD and people with a clinically determined high pre-test probability).

When considering the base case analysis it appears that the Beckman Coulter hs-cTnI assay at presentation, using the 99<sup>th</sup> centile diagnostic threshold, would be the cost-effective strategy, given an ICER threshold of £20,000 - £30,000. However, both cost and QALY differences between the strategies were small. This means that within the hs-cTn testing strategies, ICERs can change substantially especially with small changes in either costs or QALYs. Therefore, it is difficult to be confident that other hs-cTn strategies might not be cost effective.

Overall, the model does not provide strong evidence to prefer one hs-cTn testing strategy over another. Results do however indicate that hs-cTn testing in general may be cost-effective compared to standard troponin testing. This becomes more likely if one assumes that hs-cTn testing detects some patients who require treatment despite their testing negative with standard troponin, as shown in the secondary analysis. In particular, the Abbott ARCHITECT hs-cTnl optimal strategy, which involves multiple testing and varying diagnostic thresholds, may be promising. The main issue, with regard to service provision, if implementation of an hs-cTn testing strategy is considered, is the balance between the likely reduction in cost and the risk of a reduction in effectiveness, albeit possibly small.

### 6.2 Suggested research priorities

Diagnostic cohort studies are needed to fully evaluate the performance of our proposed optimal

testing strategies in a clinical setting.

If adoption of the Beckman Coulter hs-cTnl is to be considered, further studies are needed to fully evaluate the diagnostic accuracy of this test at the thresholds currently recommended by the manufacturer and to inform the development of an optimal testing strategy.

Further diagnostic cohort studies, or subgroup analyses of existing data sets, are needed to fully explore possible variation in the accuracy of hs-cTn assays and the optimal testing strategies for these assays in relevant demographic and clinical subgroups: sex; age; ethnicity; renal function; previous CAD; previous AMI.

It is important to further explore the effects of clinical judgement (assessment of pre-test probability) on the diagnostic performance of hs-cTn testing. This could be achieved by assessing the combined diagnostic accuracy of risk scoring tools, such as TIMI or GRACE, and hs-cTn tests, or by assessing the accuracy of hs-cTn testing in subgroups stratified by pre-test probability.

Multivariable prediction modelling studies may be useful to assess the independent prognostic value of a positive hs-cTn test result, in the context of other clinical risk factors and tests.

As most of the uncertainties in the economic model were caused by assumptions relating to clinical effectiveness, this type of research would also facilitate economic analyses of hs-cTn testing.

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#### **APPENDIX 1: LITERATURE SEARCH STRATEGIES**

#### **Clinical effectiveness search strategies**

# Medline (OvidSP): 1946 to 2013/10/Week 1

# Searched: 11.10.13

1 (Hstnt or hs-tnt or hsctnt or hs-ctnt or tnt-hs or tnths or ctnths or ctnt-hs).ti,ab,ot. (229)

2 (Hstni or hs-tni or hsctni or hs-ctni or tni-hs or tnihs or ctnihs or ctni-hs or ctni-ultra or accutni or accu-tni).ti,ab,ot. (99)

3 ((troponin t or tnt or ctnt or tropt or trop t) adj2 (sensitiv\$ or hs or early or initial or rapid or present\$ or ultra or high performance or ultrasensitive)).ti,ab,ot. (563)

4 ((troponin I or tni or ctni or tropI or tropI) adj2 (sensitiv\$ or hs or early or initial or rapid or present\$ or ultra or high performance or ultrasensitive)).ti,ab,ot. (349)

5 (troponin\$ adj2 (sensitiv\$ or hs or early or initial or rapid or present\$ or ultra or high performance or ultrasensitive)).ti,ab,ot. (769)

6 (troponin\$ adj5 (architect or elecsys or accutni or accu-tni or access or unicel)).ti,ab,hw,ot. (66)

- 7 or/1-6 (1215)
- 8 troponin t/ or troponin I/ or (60304-72-5 or 77108-40-8).rn. (8642)

9 (sensitiv\$ or hs or early or initial or rapid or present\$ or ultra or high performance or ultrasensitive).ti,ab,ot. (4878300)

- 10 8 and 9 (4209)
- 11 7 or 10 (4559)
- 12 chest pain/ (9293)

13 ((chest or thorax or thoracic) adj2 (pain\$ or discomfort or tight\$ or pressure)).ti,ab,ot,hw. (28602)

- 14 exp myocardial ischemia/ (357748)
- 15 (acute adj2 coronary adj2 syndrome\$).ti,ab,ot. (16495)
- 16 (preinfarc\$ Angina\$ or pre infarc\$ Angina\$).ti,ab,ot. (285)
- 17 Unstable angina\$.ti,ab,ot. (10718)

18 ((heart\$ or myocardi\$ or cardiac or coronary) adj2 (preinfarc\$ or infarc\$ or attack\$ or arrest\$ or occlusion\$ or isch?emia\$)).ti,ab,ot. (194088)

19 (MI or ACS or STEMI or NSTE-ACS or NSTEACS or nonSTEMI or NSTEMI or AMI or UAP or

- OMI).ti,ab,ot. (53168)
- 20 or/12-19 (444673)
- 21 11 and 20 (2503)
- 22 animals/ not (animals/ and humans/) (3957888)
- 23 21 not 22 (2336)

# Medline In-Process & Other Non-Indexed Citations (OvidSP): up to 2013/10/01 Medline Daily Update: up to 2013/10/01

Searched: 11.10.13

1 (Hstnt or hs-tnt or hsctnt or hs-ctnt or tnt-hs or tnths or ctnths or ctnt-hs).ti,ab,ot. (32)

2 (Hstni or hs-tni or hsctni or hs-ctni or tni-hs or tnihs or ctnihs or ctni-hs or ctni-ultra or accutni or accu-tni).ti,ab,ot. (9)

3 ((troponin t or tnt or ctnt or tropt or trop t) adj2 (sensitiv\$ or hs or early or initial or rapid or present\$ or ultra or high performance or ultrasensitive)).ti,ab,ot. (62)

4 ((troponin I or tni or ctni or tropI or tropI) adj2 (sensitiv\$ or hs or early or initial or rapid or present\$ or ultra or high performance or ultrasensitive)).ti,ab,ot. (29)

5 (troponin\$ adj2 (sensitiv\$ or hs or early or initial or rapid or present\$ or ultra or high performance or ultrasensitive)).ti,ab,ot. (99)

- 6 (troponin\$ adj5 (architect or elecsys or accutni or accu-tni or access or unicel)).ti,ab,hw,ot. (3)
- 7 or/1-6 (125)
- 8 troponin t/ or troponin I/ or (60304-72-5 or 77108-40-8).rn. (5)
- 9 (sensitiv\$ or hs or early or initial or rapid or present\$ or ultra or high performance or ultrasensitive).ti,ab,ot. (388942)
- 10 8 and 9 (3)
- 11 7 or 10 (127)
- 12 chest pain/ (13)
- 13 ((chest or thorax or thoracic) adj2 (pain\$ or discomfort or tight\$ or pressure)).ti,ab,ot,hw. (1742)
- 14 exp myocardial ischemia/ (170)
- 15 (acute adj2 coronary adj2 syndrome\$).ti,ab,ot. (1544)
- 16 (preinfarc\$ Angina\$ or pre infarc\$ Angina\$).ti,ab,ot. (3)
- 17 Unstable angina\$.ti,ab,ot. (378)
- 18 ((heart\$ or myocardi\$ or cardiac or coronary) adj2 (preinfarc\$ or infarc\$ or attack\$ or arrest\$ or occlusion\$ or isch?emia\$)).ti,ab,ot. (8220)

19 (MI or ACS or STEMI or NSTE-ACS or NSTEACS or nonSTEMI or NSTEMI or AMI or UAP or OMI).ti,ab,ot. (4224)

- 20 or/12-19 (12386)
- 21 11 and 20 (76)
- 22 animals/ not (animals/ and humans/) (1462)
- 23 21 not 22 (76)

### Embase (OvidSP): 1974 to 2013/10/10

#### Searched: 11.10.13

- 1 "high sensitivity cardiac troponin T"/ or high sensitivity troponin t assay/ (12)
- 2 "high sensitivity cardiac troponin I"/ or high sensitivity troponin i assay/ (3)
- 3 (Hstnt or hs-tnt or hsctnt or hs-ctnt or tnt-hs or tnths or ctnths or ctnt-hs).ti,ab,ot. (565)

4 (Hstni or hs-tni or hsctni or hs-ctni or tni-hs or tnihs or ctnihs or ctni-hs or ctni-ultra or accutni or accu-tni).ti,ab,ot. (190)

5 ((troponin t or tnt or ctnt or tropt or trop t) adj2 (sensitiv\$ or hs or early or initial or rapid or present\$ or ultra or high performance or ultrasensitive)).ti,ab,ot,hw. (1052)

- 6 ((troponin I or thi or cthi or tropI or tropI) adj2 (sensitiv\$ or hs or early or initial or rapid or present\$ or ultra or high performance or ultrasensitive)).ti,ab,ot,hw. (598)
- 7 (troponin\$ adj2 (sensitiv\$ or hs or early or initial or rapid or present\$ or ultra or high performance or ultrasensitive)).ti,ab,ot,hw. (1478)
- 8 (troponin\$ adj5 (architect or elecsys or accutni or accu-tni or access or unicel)).ti,ab,hw,ot. (106)
- 9 or/1-8 (2142)
- 10 troponin t/ or troponin I/ or (60304-72-5 or 77108-40-8).rn. (18661)
- 11 (sensitiv\$ or hs or early or initial or rapid or present\$ or ultra or high performance or ultraconsitive) tilab et hw (6501005)
- ultrasensitive).ti,ab,ot,hw. (6591905)
- 12 10 and 11 (9505) 13 9 or 12 (10097)
- 13 9 or 12 (10097)
- 14 thorax pain/ (44504)
- 15 ((chest or thorax or thoracic) adj2 (pain\$ or discomfort or tight\$ or pressure)).ti,ab,ot,hw.(64208)
- 16 acute coronary syndrome/ (24295)
- 17 (acute adj2 coronary adj2 syndrome\$).ti,ab,ot,hw. (34428)
- 18 exp heart muscle ischemia/ (73551)
- 19 exp heart infarction/ (266027)
- 20 exp Unstable-Angina-Pectoris/ (16552)

21 (preinfarc\$ Angina\$ or pre infarc\$ Angina\$).ti,ab,ot,hw. (374)

22 Unstable angina\$.ti,ab,ot. (14593)

23 ((heart\$ or myocardi\$ or cardiac or coronary) adj2 (preinfarc\$ or infarc\$ or attack\$ or arrest\$ or occlusion\$ or isch?emia\$)).ti,ab,ot,hw. (406203)

24 (MI or ACS or STEMI or NSTE-ACS or NSTEACS or nonSTEMI or NSTEMI or AMI or UAP or OMI).ti,ab,ot,hw. (85655)

25 or/14-24 (498902)

26 13 and 25 (6007)

27 animal/ (1890932)

28 animal experiment/ (1720343)

29 (rat or rats or mouse or mice or murine or rodent or rodents or hamster or hamsters or pig or pigs or porcine or rabbit or rabbits or animal or animals or dogs or dog or cats or cow or bovine or sheep or ovine or monkey or monkeys).ti,ab,ot,hw. (5825865)

- 30 or/27-29 (5825865)
- 31 exp human/ (15014990)
- 32 human experiment/ (317206)
- 33 or/31-32 (15016431)
- 34 30 not (30 and 33) (4642837)
- 35 26 not 34 (5642)
- 36 limit 35 to yr="2005 -Current" (4374)

37 remove duplicates from 36 (4282)

Cochrane Database of Systematic Reviews (CDSR) (Wiley). Issue 10/October: up to 2013/10/11 Cochrane Central Register of Controlled Trials (CENTRAL) (Wiley). Issue 9/September: 2013 Database of Abstracts of Reviews of Effects (DARE) (Wiley). Issue 3/July:2013 Health Technology Assessment Database (HTA) (Wiley). Issue 3/July:2013 NHS Economic Evaluation Database (NHS EED) (Wiley). Issue 3/July:2013 Searched 11.10.13

#1 (Hstnt or hs-tnt or hs-ctnt or hs-ctnt or tnt-hs or tnths or ctnths or ctnt-hs):ti,ab,kw 5

#2 (Hstni or hs-tni or hsctni or hs-ctni or tni-hs or tnihs or ctnihs or ctni-hs or ctni-ultra or accutni or accu-tni):ti,ab,kw 5

#3 ((troponin t or tnt or ctnt or tropt or trop t) near/2 (sensitiv\* or hs or early or initial or rapid or present\* or ultra or high performance or ultrasensitive)):ti,ab,kw 12

#4 ((troponin I or thi or cthi or tropI or tropI) near/2 (sensitiv\* or hs or early or initial or rapid or present\* or ultra or high performance or ultrasensitive)):ti,ab,kw
 10

#5 (troponin\* near/2 (sensitiv\* or hs or early or initial or rapid or present\* or ultra or high performance or ultrasensitive)):ti,ab,kw
 27

- #6 (troponin\* near/5 (architect or elecsys or accutni or accu-tni or access or unicel)):ti,ab,kw
  2
- #7 #1 or #2 or #3 or #4 or #5 or #6 42

#8 MeSH descriptor: [Troponin T] this term only 265

#9 MeSH descriptor: [Troponin I] this term only 309

#10 #8 or #9 543

#11 (sensitiv\* or hs or early or initial or rapid or present\* or ultra or high performance or ultrasensitive):ti,ab,kw 170016

#12 #10 and #11 236

- #13 #7 or #12 249
- #14 MeSH descriptor: [Chest Pain] this term only 335
- #15 ((chest or thorax or thoracic) near/2 (pain\* or discomfort or tight\* or pressure)):ti,ab,kw 1793
- #16 (acute near/2 coronary near/2 syndrome\*):ti,ab,kw 1678

#17 MeSH descriptor: [Myocardial Ischemia] explode all trees 20427

#18 (preinfarc\* Angina\* or pre infarc\* Angina\*):ti,ab,kw 90

#19 (Unstable angina\*):ti,ab,kw 1818

#20 ((heart\* or myocardi\* or cardiac or coronary) near/2 (preinfarc\* or infarc\* or attack\* or arrest\* or occlusion\* or isch?emia\*)):ti,ab,kw 16156

#21 (MI or ACS or STEMI or NSTE-ACS or NSTEACS or nonSTEMI or NSTEMI or AMI or UAP or OMI):ti,ab,kw 4740

#22 #14 or #15 or #16 or #17 or #18 or #19 or #20 or #21 28923

#### #23 #13 and #22 from 2005 to 2013 114

CDSR search retrieved 0 references

**CENTRAL search retrieved 108 references** 

DARE search retrieved 2 references

HTA search retrieved 1 references

(NHS EED search retrieved 3 references)

Science Citation Index – Expanded (SCI) (Web of Science): 1970-2013/10/14 Conference Proceedings Citation Index (CPCI-S) (Web of Science): 1990-2013/10/14 Searched 14.10.13

#### Databases=SCI-EXPANDED, CPCI-S, CCR-EXPANDED, IC Timespan=2005-2013

# 1 228 TS=(Hstnt or hs-tnt or hsctnt or hs-ctnt or tnt-hs or tnths or ctnths or ctnt-hs) # 2 90 TS=(Hstni or hs-tni or hsctni or hs-ctni or tni-hs or tnihs or ctnihs or ctni-hs or ctni-ultra or accutni or accu-tni) # 3 1,438 TS=((troponin\* or tnt or ctnt or tropt or tni or ctni or tropI or "trop t" or "trop I") NEAR/2 (sensitiv\* or hs or early or initial or rapid or present\* or ultra or "high performance" or ultrasensitive)) # 4 1,470 #3 OR #2 OR #1

# 5 13,963 TS=((chest or thorax or thoracic) NEAR (pain\* or discomfort or tight\* or pressure))

# 6 19,298 TS=(acute NEAR/2 coronary NEAR/2 syndrome\*)

# 7 393 TS=(preinfarc\* angina\* or pre infarc\* angina)

# 8 5,481 TS=unstable angina\*

# 9 115,395 TS=((heart\* or myocard\* or cardiac or coronary) NEAR/2 (preinfarc\* or infarc\* or attack\* or arrest\* or occlusion\* or isch?emia\*))

# 10 40,133 TS=(MI or ACS or STEMI or NSTE-ACS or NSTEACS or nonSTEMI or NSTEMI or AMI or UAP or OMI)

# 11 155,342 #10 OR #9 OR #8 OR #7 OR #6 OR #5

#### # 12 835 #11 AND #4

#### LILACS (Latin American and Caribbean Health Sciences): 1982-2013/09/24

http://regional.bvsalud.org/php/index.php?lang=en

Searched 14.10.13

Terms	Records
(Troponin\$ or MH:D05.750.078.730.825.925 or MH:D12.776.210.500.910.925 or	247
MH:D12.776.220.525.825.925 or MH:D05.750.078.730.825.962 or	
MH:D12.776.210.500.910.962 or MH:D12.776.220.525.825.962 or	
MH:D05.750.078.730.825 or MH:D12.776.210.500.910 or	
MH:D12.776.220.525.825 or Hstnt or hs-tnt or hsctnt or hs-ctnt or tnt-hs or tnths	
or ctnths or ctnt-hs or Hstni or hs-tni or hsctni or hs-ctni or tni-hs or tnihs or	
ctnihs or ctni-hs or ctni-ultra or accutni or accu-tni)	
Total	247

Spanish and Portuguese translations of MeSH terms identified using the DECS (Health Sciences Descriptors) thesaurus: <u>http://decs.bvs.br/l/homepagei.htm</u>

# INAHTA (International Network of Agencies for Health Technology Assessment): up to 2013/10/15 http://www.inahta.org/Search2/?pub=1

#### Searched 15.10.13

Search Term	Results
Troponin	9
Elecsys	2
Architect	0
Accutni	0/1
unicel	0
Total	11

#### Biosis Previews (Web of Knowledge): 1956-2013/10/11 Searched 14.10.13

#### Databases=BIOSIS Previews Timespan=2005-2013

# 1 266 TS=(Hstnt or hs-tnt or hsctnt or hs-ctnt or tnt-hs or tnths or ctnths or ctnt-hs) # 2 114 TS=(Hstni or hs-tni or hsctni or hs-ctni or tni-hs or tnihs or ctnihs or ctni-hs or ctni-ultra or accutni or accu-tni) # 3 1,055 TS=((troponin\* or tnt or ctnt or tropt or tni or ctni or tropl or "trop t" or "trop I") NEAR/2 (sensitiv\* or hs or early or initial or rapid or present\* or ultra or "high performance" or ultrasensitive)) # 4 1,095 #3 OR #2 OR #1 # 5 7,468 TS=((chest or thorax or thoracic) NEAR (pain\* or discomfort or tight\* or pressure)) # 6 11,149 TS=(acute NEAR/2 coronary NEAR/2 syndrome\*) # 7 196 TS=(preinfarc\* angina\* or pre infarc\* angina) # 8 3,025 TS=unstable angina\* # 9 62,717 TS=((heart\* or myocard\* or cardiac or coronary) NEAR/2 (preinfarc\* or infarc\* or attack\* or arrest\* or occlusion\* or isch?emia\*)) # 10 28,931 TS=(MI or ACS or STEMI or NSTE-ACS or NSTEACS or nonSTEMI or NSTEMI or AMI or UAP or OMI) # 11 83,999 #10 OR #9 OR #8 OR #7 OR #6 OR #5 # 12 628 #11 AND #4 **NIHR HTA (Internet)** 

http://www.hta.ac.uk/ up to 2013/10/14 Searched 14.10.2013

Browsed with Troponin terms - 6 results

ARIF (Internet): 1996-2013/10/16

http://www.birmingham.ac.uk/research/activity/mds/projects/HaPS/PHEB/ARIF/databases/inde

# Searched 16.10.13

Search terms	Quick Search
Troponin*	21
Hstnt or hs-tnt or hsctnt or hs-ctnt or tnt-hs or tnths or ctnths or ctnt-hs	0
Hstni or hs-tni or hsctni or hs-ctni or tni-hs or tnihs or ctnihs or ctni-hs or	0
ctni-ultra or accutni or accu-tni	
Total	21

#### MEDION database: up to 2013/10/16

http://www.mediondatabase.nl/

Searched 16.10.13

Searched in 'Whole Database'

Search Term in 'Topics'	Results
Troponin	0
Troponins	0
Total	0

# PROSPERO (International Prospective Register of Systematic Reviews) (Internet): up to 2013/10/10 http://www.crd.york.ac.uk/prospero/

Searched 10.10.13

Searched in 'All fields'

Terms	Records
Troponin*	8
Total	8

#### Clinicaltrials.gov (Internet) http://clinicaltrials.gov/ct2/search/advanced Searched 14.10.13

Advanced search option – search terms box

Search terms	Condition	Intervention	Records
troponin% AND (sensitiv% OR hs			186
OR early OR initial OR rapid OR			
present% OR ultra OR high			
performance OR ultrasensitive			
OR elecsys OR architect OR			
accutni OR access OR unicel)			
		Troponin%	109

(Hstnt OR hs-tnt OR hsctnt Or		17
hs-ctnt OR tnt-hs OR tnths OR		
ctnths OR ctnt-hs OR Hstni OR		
hs-tni OR hsctni OR hs-ctni OR		
tni-hs OR tnihs OR ctnihs OR		
ctni-hs OR ctni-ultra OR accutni		
OR accu-tni)		
Total		312

#### mRCT – metaRegister of Controlled Trials (Internet) http://www.controlled-trials.com/ Up to 10.10.13 Searched 10.10.13

Search terms	Results
(troponin* AND (sensitiv* or hs or early or initial or rapid or present* or ultra or	333
high performance or ultrasensitive))	
TOTAL	333

#### WHO International Clinical Trials Registry Platform (ICTRP) (Internet) http://www.who.int/ictrp/en/ Searched 10.10.2013

Advanced search option Date of registration limited to 01/01/2005 – 10/10/2013

Title	Condition	Intervention	Records
Troponin OR Troponins			67
		Troponins	2
		Troponin	This search does not work – the results are irrelevant and do not contain the word troponin in the intervention field
Total			69

#### American Heart Association – Scientific Sessions

http://my.americanheart.org/professional/Sessions/ScientificSessions/Archive/Archive-Scientific-Sessions\_UCM\_316935\_SubHomePage.jsp

Searched: 29.10.13

2013 - Conference not yet taken place at time of searching

2012: http://circ.ahajournals.org/content/vol126/21\_MeetingAbstracts

2011: <u>http://circ.ahajournals.org/content/vol124/21\_MeetingAbstracts</u>

2010: http://circ.ahajournals.org/content/vol122/21\_MeetingAbstracts

2009: http://circ.ahajournals.org/content/120/21/2152.full.pdf

Keyword	2013	2012	2011	2010	2009	Total
Troponin*	N/A	138	131	109	1	379

#### American Association for Clinical Chemistry

http://www.aacc.org/resourcecenters/meet\_abstracts\_archive/abstracts\_archive/annual\_meeting/ Pages/default.aspx#

Searched: 29.10.13

2013 Abstracts from: Clinical Chemistry, 59(S10):A1-295

http://www.aacc.org/events/Annual\_Meeting/abstracts/Documents/AACC\_13\_AbstractBook\_Comp lete.pdf

2012 Abstracts from: Clinical Chemistry, 58(S10):a1-A264 <u>http://www.aacc.org/events/annualmtgdirectory/Documents/AACC\_12\_AbstractBook-Final-</u> Complete.pdf

2011 Abstracts from: Clinical Chemistry, 57 (S10): A1-A235 http://www.aacc.org/events/annualmtgdirectory/documents/AACC 11 FullAbstract.pdf

#### 2010 Abstracts from: Clinical Chemistry, 57 (6 Suppl): A1-276

http://www.aacc.org/events/annualmtgdirectory/Pages/2010PosterAbstracts.aspx#

#### 2009 19-23 July, Chicago, Ill,

http://www.abstractsonline.com/viewer/searchAdvanced.asp?MKey={CA6D749E-BE20-4F85-899B-8A84E2268F72}&AKey={B08F832C-9D23-4F0B-96C3-3FA22F3D94A1}

Keyword	2013	2012	2011	2010	2009	Totals
Troponin	48	21	32	40	29	170

#### **European Society of Cardiology**

http://spo.escardio.org/abstract-book/search.aspx

Searched: 29.10.13

Keyword	2013	2012	2011	2010	2009	Total
Troponin	52	51	61	51	25	240
Troponins	2	1	2	1	2	8
						248

#### Additional searches

Results sorted by Link Ranking http://www.ncbi.nlm.nih.gov/pubmed/ Searched 10.12.13

Nine of the included publications were not indexed on PubMed. Indexed publications were checked for errata and comments. For each reference, the first 20 references were retrieved by carrying out

a Related Citations search using PubMed's similarity matching algorithm. These records were downloaded for screening. All related citations were checked against the Endnote Library to remove duplicates, and only new unique references were imported and screened = 58 records

Reference	PMID	Result retrieved
Santalo <sup>39</sup>	23764266	20/131
Aldous <sup>40</sup>	22109535	20/145
Sanchis <sup>41</sup>	22877804	20/203
Haaf <sup>42</sup>	22623715	20/203
Eggers <sup>43</sup>	22456003	20/145
Reiter <sup>44</sup>	22044927	20/280
Aldous <sup>45</sup>	22291171	20/277
Potocki <sup>46</sup>	22337952	20/304
Keller <sup>47</sup>	22203537	20/300
#403 Meune	22014790	20/252
Freund <sup>48</sup>	21663627	20/142
Aldous <sup>49</sup>	21784766	20/254
Melki <sup>50</sup>	21428843	20/210
Reichlin <sup>51</sup>	21709058	20/162
Reiter <sup>52</sup>	21362702	20/261
Aldous <sup>53</sup>	21441390	20/251
Kurz <sup>54</sup>	20852870	20/207
Hochholzer <sup>55</sup>	21138939	20/138
Christ <sup>56</sup>	20932502	20/201
Parsonage <sup>57</sup>	Not in pubmed	
Collinson <sup>58</sup>	Not in pubmed	
Body <sup>59</sup>	Not in pubmed	
Melki <sup>60</sup>	Not in pubmed	
Aldous <sup>61</sup>	Not in pubmed	
Cullen <sup>62</sup>	23583250	20/133
Sebbane <sup>63</sup>	23816196	20/131
Irfan <sup>64</sup>	23870791	20/134
Collinson <sup>65</sup>	23597479	20/275
Reiter <sup>66</sup>	23514979	20/155
Body <sup>67</sup>	21920261	20/192
Aldous <sup>68</sup>	21441393	20/174
Keller <sup>69</sup>	Not in pubmed	
Collinson <sup>70</sup>	Not in pubmed	
Saenger <sup>71</sup>	Not in pubmed	
Lippi <sup>74</sup>	Not in pubmed	
Hoeller <sup>75</sup>	23604180	20/107
Total		640
Following duplicate removal, number of records screened		58

#### **Cost-effectiveness searches**

#### Medline (OvidSP): 1946 to 2013/10/Week 1 Searched: 18.10.13

- 1 economics/ (27116)
- 2 exp "costs and cost analysis"/ (182544)
- 3 economics, dental/ (1866)
- 4 exp "economics, hospital"/ (19403)
- 5 economics, medical/ (8578)
- 6 economics, nursing/ (3879)
- 7 economics, pharmaceutical/ (2605)
- 8 (economic\$ or cost or costs or costly or costing or price or prices or pricing or
- pharmacoeconomic\$).ti,ab. (427344)
- 9 (expenditure\$ not energy).ti,ab. (17552)
- 10 (value adj1 money).ti,ab. (22)
- 11 budget\$.ti,ab. (17208)
- 12 or/1-11 (551693)
- 13 ((energy or oxygen) adj cost).ti,ab. (2752)
- 14 (metabolic adj cost).ti,ab. (798)
- 15 ((energy or oxygen) adj expenditure).ti,ab. (16662)
- 16 or/13-15 (19503)
- 17 12 not 16 (547348)
- 18 letter.pt. (803396)
- 19 editorial.pt. (334975)
- 20 historical article.pt. (299710)
- 21 or/18-20 (1423597)
- 22 17 not 21 (519320)
- 23 (Hstnt or hs-tnt or hsctnt or hs-ctnt or tnt-hs or tnths or ctnths or ctnt-hs).ti,ab,ot. (229)

24 (Hstni or hs-tni or hsctni or hs-ctni or tni-hs or tnihs or ctnihs or ctni-hs or ctni-ultra or accutni or accu-tni).ti,ab,ot. (99)

25 ((troponin t or tnt or ctnt or tropt or trop t) adj2 (sensitiv\$ or hs or early or initial or rapid or present\$ or ultra or high performance or ultrasensitive)).ti,ab,ot. (563)

- 26 ((troponin I or tni or ctni or tropl or trop I) adj2 (sensitiv\$ or hs or early or initial or rapid or present\$ or ultra or high performance or ultrasensitive)).ti,ab,ot. (349)
- 27 (troponin\$ adj2 (sensitiv\$ or hs or early or initial or rapid or present\$ or ultra or high performance or ultrasensitive)).ti,ab,ot. (769)
- 28 (troponin\$ adj5 (architect or elecsys or accutni or accu-tni or access or unicel)).ti,ab,hw,ot. (66)
- 29 or/23-28 (1215)
- 30 troponin t/ or troponin I/ or (60304-72-5 or 77108-40-8).rn. (8642)
- 31 (sensitiv\$ or hs or early or initial or rapid or present\$ or ultra or high performance or ultrasensitive).ti,ab,ot. (4878300)
- 32 30 and 31 (4209)
- 33 29 or 32 (4559)
- 34 chest pain/ (9293)
- 35 ((chest or thorax or thoracic) adj2 (pain\$ or discomfort or tight\$ or pressure)).ti,ab,ot,hw. (28602)
- 36 exp myocardial ischemia/ (357748)
- 37 (acute adj2 coronary adj2 syndrome\$).ti,ab,ot. (16495)
- 38 (preinfarc\$ Angina\$ or pre infarc\$ Angina\$).ti,ab,ot. (285)
- 39 Unstable angina\$.ti,ab,ot. (10718)

40 ((heart\$ or myocardi\$ or cardiac or coronary) adj2 (preinfarc\$ or infarc\$ or attack\$ or arrest\$ or occlusion\$ or isch?emia\$)).ti,ab,ot. (194088)

41 (MI or ACS or STEMI or NSTE-ACS or NSTEACS or nonSTEMI or NSTEMI or AMI or UAP or OMI).ti,ab,ot. (53168)

- 42 or/34-41 (444673)
- 43 33 and 42 (2503)
- 44 animals/ not (animals/ and humans/) (3957888)
- 45 43 not 44 (2336)
- 46 limit 45 to yr="2005 -Current" (1457)
- 47 22 and 46 (43)

Costs filter:

Centre for Reviews and Dissemination. NHS EED Economics Filter: Medline (Ovid) monthly search York: Centre for Reviews and Dissemination; 2010

# Medline In-Process & Other Non\_Indexed Citations (OvidSP): up to 2013/10/01 Medline Daily Update: up to 2013/10/01

# Searched: 18.10.13

- 1 economics/ (2)
- 2 exp "costs and cost analysis"/ (87)
- 3 economics, dental/(0)
- 4 exp "economics, hospital"/ (8)
- 5 economics, medical/ (0)
- 6 economics, nursing/ (0)
- 7 economics, pharmaceutical/ (1)
- 8 (economic\$ or costs or costly or costing or price or prices or pricing or

pharmacoeconomic\$).ti,ab. (39821)

- 9 (expenditure\$ not energy).ti,ab. (1172)
- 10 (value adj1 money).ti,ab. (4)
- 11 budget\$.ti,ab. (1822)
- 12 or/1-11 (41689)
- 13 ((energy or oxygen) adj cost).ti,ab. (218)
- 14 (metabolic adj cost).ti,ab. (67)
- 15 ((energy or oxygen) adj expenditure).ti,ab. (911)
- 16 or/13-15 (1160)
- 17 12 not 16 (41354)
- 18 letter.pt. (24293)
- 19 editorial.pt. (14525)
- 20 historical article.pt. (68)
- 21 or/18-20 (38878)
- 22 17 not 21 (40906)
- 23 (Hstnt or hs-tnt or hs-ctnt or tnt-hs or tnths or ctnths or ctnt-hs).ti,ab,ot. (32)

24 (Hstni or hs-tni or hsctni or hs-ctni or tni-hs or tnihs or ctnihs or ctni-hs or ctni-ultra or accutni or accu-tni).ti,ab,ot. (9)

25 ((troponin t or tnt or ctnt or tropt or trop t) adj2 (sensitiv\$ or hs or early or initial or rapid or present\$ or ultra or high performance or ultrasensitive)).ti,ab,ot. (62)

26 ((troponin I or thi or cthi or tropI or trop I) adj2 (sensitiv\$ or hs or early or initial or rapid or present\$ or ultra or high performance or ultrasensitive)).ti,ab,ot. (29)

27 (troponin\$ adj2 (sensitiv\$ or hs or early or initial or rapid or present\$ or ultra or high performance or ultrasensitive)).ti,ab,ot. (99)

28 (troponin\$ adj5 (architect or elecsys or accutni or accu-tni or access or unicel)).ti,ab,hw,ot. (3)

- 29 or/23-28 (125)
- 30 troponin t/ or troponin I/ or (60304-72-5 or 77108-40-8).rn. (5)
- 31 (sensitiv\$ or hs or early or initial or rapid or present\$ or ultra or high performance or ultrasensitive).ti,ab,ot. (388942)
- 32 30 and 31 (3)
- 33 29 or 32 (127)
- 34 chest pain/ (13)
- 35 ((chest or thorax or thoracic) adj2 (pain\$ or discomfort or tight\$ or pressure)).ti,ab,ot,hw.

(1742)

- 36 exp myocardial ischemia/ (170)
- 37 (acute adj2 coronary adj2 syndrome\$).ti,ab,ot. (1544)
- 38 (preinfarc\$ Angina\$ or pre infarc\$ Angina\$).ti,ab,ot. (3)
- 39 Unstable angina\$.ti,ab,ot. (378)
- 40 ((heart\$ or myocardi\$ or cardiac or coronary) adj2 (preinfarc\$ or infarc\$ or attack\$ or arrest\$ or occlusion\$ or isch?emia\$)).ti,ab,ot. (8220)
- 41 (MI or ACS or STEMI or NSTE-ACS or NSTEACS or nonSTEMI or NSTEMI or AMI or UAP or
- OMI).ti,ab,ot. (4224)
- 42 or/34-41 (12386)
- 43 33 and 42 (76)
- 44 animals/ not (animals/ and humans/) (1462)
- 45 43 not 44 (76)
- 46 limit 45 to yr="2005 -Current" (75)
- 47 22 and 46 (4)

Costs filter:

Centre for Reviews and Dissemination. NHS EED Economics Filter: Medline (Ovid) monthly search . York: Centre for Reviews and Dissemination; 2010

# Embase (OvidSP): 1974 to 2013/10/17

#### Searched: 18.10.13

- 1 health-economics/ (33273)
- 2 exp economic-evaluation/ (205882)
- 3 exp health-care-cost/ (197503)
- 4 exp pharmacoeconomics/ (169588)
- 5 or/1-4 (471813)
- 6 (econom\$ or cost or costs or costly or costing or price or prices or pricing or
- pharmacoeconomic\$).ti,ab. (590127)
- 7 (expenditure\$ not energy).ti,ab. (23360)
- 8 (value adj2 money).ti,ab. (1320)
- 9 budget\$.ti,ab. (23595)
- 10 or/6-9 (613918)
- 11 5 or 10 (885833)
- 12 letter.pt. (844056)
- 13 editorial.pt. (449323)
- 14 note.pt. (587506)
- 15 or/12-14 (1880885)
- 16 11 not 15 (799169)
- 17 (metabolic adj cost).ti,ab. (876)
- 18 ((energy or oxygen) adj cost).ti,ab. (3163)
- 19 ((energy or oxygen) adj expenditure).ti,ab. (19981)
- 20 or/17-19 (23208)
- 21 16 not 20 (794101)

22 "high sensitivity cardiac troponin T"/ or high sensitivity troponin t assay/ (12)

23 "high sensitivity cardiac troponin I"/ or high sensitivity troponin i assay/ (3)

24 (Hstnt or hs-tnt or hsctnt or hs-ctnt or tnt-hs or tnths or ctnths or ctnt-hs).ti,ab,ot. (571)

25 (Hstni or hs-tni or hsctni or hs-ctni or tni-hs or tnihs or ctnihs or ctni-hs or ctni-ultra or accutni or accu-tni).ti,ab,ot. (193)

26 ((troponin t or tnt or ctnt or tropt or trop t) adj2 (sensitiv\$ or hs or early or initial or rapid or present\$ or ultra or high performance or ultrasensitive)).ti,ab,ot,hw. (1059)

27 ((troponin I or thi or cthi or tropI or trop I) adj2 (sensitiv\$ or hs or early or initial or rapid or present\$ or ultra or high performance or ultrasensitive)).ti,ab,ot,hw. (602)

28 (troponin\$ adj2 (sensitiv\$ or hs or early or initial or rapid or present\$ or ultra or high performance or ultrasensitive)).ti,ab,ot,hw. (1489)

(troponin\$ adj5 (architect or elecsys or accutni or accu-tni or access or unicel)).ti,ab,hw,ot.(106)

- 30 or/22-29 (2155)
- 31 troponin t/ or troponin I/ or (60304-72-5 or 77108-40-8).rn. (18726)
- 32 (sensitiv\$ or hs or early or initial or rapid or present\$ or ultra or high performance or ultrasensitive).ti,ab,ot,hw. (6601404)
- 33 31 and 32 (9548)
- 34 30 or 33 (10144)
- 35 thorax pain/ (44662)

36 ((chest or thorax or thoracic) adj2 (pain\$ or discomfort or tight\$ or pressure)).ti,ab,ot,hw. (64388)

- 37 acute coronary syndrome/ (24412)
- 38 (acute adj2 coronary adj2 syndrome\$).ti,ab,ot,hw. (34558)
- 39 exp heart muscle ischemia/ (73666)
- 40 exp heart infarction/ (266475)
- 41 exp Unstable-Angina-Pectoris/ (16570)
- 42 (preinfarc\$ Angina\$ or pre infarc\$ Angina\$).ti,ab,ot,hw. (374)
- 43 Unstable angina\$.ti,ab,ot. (14604)

44 ((heart\$ or myocardi\$ or cardiac or coronary) adj2 (preinfarc\$ or infarc\$ or attack\$ or arrest\$ or occlusion\$ or isch?emia\$)).ti,ab,ot,hw. (406847)

45 (MI or ACS or STEMI or NSTE-ACS or NSTEACS or nonSTEMI or NSTEMI or AMI or UAP or OMI).ti,ab,ot,hw. (85913)

- 46 or/35-45 (499787)
- 47 34 and 46 (6035)
- 48 animal/ (1890937)
- 49 animal experiment/ (1721607)

50 (rat or rats or mouse or mice or murine or rodent or rodents or hamster or hamsters or pig or pigs or porcine or rabbit or rabbits or animal or animals or dogs or dog or cats or cow or bovine or sheep or ovine or monkey or monkeys).ti,ab,ot,hw. (5828979)

- 51 or/48-50 (5828979)
- 52 exp human/ (15032575)
- 53 human experiment/ (317393)
- 54 or/52-53 (15034016)
- 55 51 not (51 and 54) (4644866)
- 56 47 not 55 (5669)
- 57 limit 56 to yr="2005 -Current" (4401)
- 58 remove duplicates from 57 (4309)
- 59 21 and 58 (129)

Costs filter:

Centre for Reviews and Dissemination. NHS EED Economics Filter: Embase (Ovid) weekly search. York: Centre for Reviews and Dissemination; 2010

#### NHS Economic Evaluation Database (NHS EED) (Wiley) Issue 3/July:2013 Searched 11.10.13

5 #1 (Hstnt or hs-tnt or hsctnt or hs-ctnt or tnt-hs or tnths or ctnths or ctnt-hs):ti,ab,kw #2 (Hstni or hs-tni or hsctni or hs-ctni or tni-hs or tnihs or ctnihs or ctni-hs or ctni-ultra or accutni or accu-tni):ti,ab,kw 5 #3 ((troponin t or tnt or ctnt or tropt or trop t) near/2 (sensitiv\* or hs or early or initial or rapid or present\* or ultra or high performance or ultrasensitive)):ti,ab,kw 12 ((troponin I or tni or ctni or tropI or tropI) near/2 (sensitiv\* or hs or early or initial or rapid #4 or present\* or ultra or high performance or ultrasensitive)):ti,ab,kw 10 #5 (troponin\* near/2 (sensitiv\* or hs or early or initial or rapid or present\* or ultra or high performance or ultrasensitive)):ti,ab,kw 27 #6 (troponin\* near/5 (architect or elecsys or accutni or accu-tni or access or unicel)):ti,ab,kw 2 #7 #1 or #2 or #3 or #4 or #5 or #6 42 #8 MeSH descriptor: [Troponin T] this term only 265 #9 MeSH descriptor: [Troponin I] this term only 309 #10 #8 or #9 543 #11 (sensitiv\* or hs or early or initial or rapid or present\* or ultra or high performance or ultrasensitive):ti,ab,kw 170016 #10 and #11 #12 236 249 #13 #7 or #12 #14 MeSH descriptor: [Chest Pain] this term only 335 ((chest or thorax or thoracic) near/2 (pain\* or discomfort or tight\* or pressure)):ti,ab,kw #15 1793 #16 (acute near/2 coronary near/2 syndrome\*):ti,ab,kw 1678 #17 MeSH descriptor: [Myocardial Ischemia] explode all trees 20427 (preinfarc\* Angina\* or pre infarc\* Angina\*):ti,ab,kw #18 90 (Unstable angina\*):ti,ab,kw #19 1818 ((heart\* or myocardi\* or cardiac or coronary) near/2 (preinfarc\* or infarc\* or attack\* or #20 arrest\* or occlusion\* or isch?emia\*)):ti,ab,kw 16156 (MI or ACS or STEMI or NSTE-ACS or NSTEACS or nonSTEMI or NSTEMI or AMI or UAP or #21 OMI):ti,ab,kw 4740 #14 or #15 or #16 or #17 or #18 or #19 or #20 or #21 28923 #22 #13 and #22 from 2005 to 2013 114 #23 NHS EED search retrieved 3 references

Health Economic Evaluation Database (HEED) (Internet): up to 2013/10/18 http://onlinelibrary.wiley.com/book/10.1002/9780470510933 Searched 18.10.13

Compound search, (all data), unable to limit by date

Troponin\* AND sensitiv\* OR hs OR early OR initial OR rapid OR present OR ultra OR high performance OR ultrasensitive OR elecsys OR architect OR accutni OR access OR unicel N=20 Hstnt or hs-tnt or hsctnt or hs-ctnt or tnt-hs or tnths or ctnths or ctnt-hs or Hstni or hs-tni or hsctni or hs-ctni or tni-hs or ctnihs or ctni-hs or ctni-ultra N=0

Hstni or hs-tni or hsctni or hs-ctni or tni-hs or tnihs or ctnihs or ctni-hs or ctni-ultra or accutni or accu-tni N=0

#### EconLit (EBSCO) 1990-2013/09/01 Searched: 18.10.13

Search modes - Boolean/Phrase

S1 TX Troponin\* (0)

S2 TX Hstnt or hs-tnt or hsctnt or hs-ctnt or tnt-hs or tnths or ctnths or ctnt-hs (0) S3 TX Hstni or hs-tni or hsctni or hs-ctni or tni-hs or tnihs or ctnihs or ctni-hs or ctni-ultra or accutni or accu-tni (0)

# Science Citation Index – Expanded (SCI) (Web of Science): 1970-2013/10/21 Conference Proceedings Citation Index (CPCI-S) (Web of Science): 1990-2013/10/21 Searched 21.10.13

# 1 622,444 TS=(economic\* or cost or costs or costly or costing or price or prices or pricing or pharmacoeconomic\* or budget\*) # 2 10,144 TS=(expenditure\* not energy) # 3 952 TS=(value NEAR money) # 4 626,873 #3 OR #2 OR #1 # 5 22,383 TS=((energy or oxygen) NEAR cost) # 6 1,804 TS=(metabolic NEAR cost) # 7 12,974 TS=((energy or oxygen) NEAR expenditure) # 8 35,684 #7 OR #6 OR #5 # 9 602,398 #4 NOT #8 # 10 230 TS=(Hstnt or hs-tnt or hsctnt or hs-ctnt or tnt-hs or tnths or ctnths or ctnt-hs) # 11 91 TS=(Hstni or hs-tni or hsctni or hs-ctni or tni-hs or tnihs or ctnihs or ctni-hs or ctni-ultra or accutni or accu-tni) # 12 1,442 TS=((troponin\* or tnt or ctnt or tropt or tni or ctni or tropI or "trop t" or "trop I") NEAR/2 (sensitiv\* or hs or early or initial or rapid or present\* or ultra or "high performance" or ultrasensitive)) # 13 1,474 #12 OR #11 OR #10 # 14 14,001 TS=((chest or thorax or thoracic) NEAR (pain\* or discomfort or tight\* or pressure)) # 15 19,324 TS=(acute NEAR/2 coronary NEAR/2 syndrome\*) # 16 393 TS=(preinfarc\* angina\* or pre infarc\* angina) # 17 5,486 TS=unstable angina\* # 18 115,562 TS=((heart\* or myocard\* or cardiac or coronary) NEAR/2 (preinfarc\* or infarc\* or attack\* or arrest\* or occlusion\* or isch?emia\*)) # 19 40,195 TS=(MI or ACS or STEMI or NSTE-ACS or NSTEACS or nonSTEMI or NSTEMI or AMI or UAP or OMI) # 20 155,582 #19 OR #18 OR #17 OR #16 OR #15 OR #14 # 21 839 #20 AND #13 # 22 32 #21 AND #9

#### Databases=SCI-EXPANDED, CPCI-S, CCR-EXPANDED, IC Timespan=2005-2013

#### Research Papers in Economics (REPEC) up to 2013/10/21 http://econpapers.repec.org/scripts/search/search.asp?pg=-1 Searched: 21.10.13 Advanced search

Free text search	Results	Total
Troponin	0/2	0
Troponins	0/1	0

# **APPENDIX 2: DATA EXTRACTION TABLES**

#### a. Baseline study details

Study Details	Selection criteria	Participant details	Test
			Manufacturer
Aldous(2012) <sup>40, 45, 49</sup>	Inclusion criteria:	Median age (IQR): 65( 56, 76)	Roche
	Adults (≥18 years) with symptoms suggestive of cardiac ischemia (acute	Male (%): 60	
Country: New Zealand	chest, epigastric, neck, jaw or arm pain or discomfort or pressure without	White (%): 89	
	an apparent noncardiac source)	Previous CAD (%): 52	
Funding: Funded by the National Heart Foundation		Previous Family History (%): 60	
of New Zealand and assay reagents were provided	Exclusion criteria:	Previous Revascularisation (%): 30	
by the manufacturer (Roche). One author declared	ST-segment elevation on ECG <sup>45</sup> ; unable to provide informed consent;	Diabetes (%): 17	
personal funding from Abbott	would not be available to follow-up	Smoking (%): 61	
		Hypertension (%): 61	
Recruitment: November 2007 - December 2010	Patient category:	Dyslipidaemia (%): 58	
	NSTEMI <sup>45</sup>	Median BMI (IQR): 28(25, 31)	
Number of participants: 939, <sup>45</sup> 385 <sup>40</sup>	Mixed <sup>40</sup>	Median (IQR) time to presentation	
		(hours): 6.3 (3.3, 13.3)	
Aldous(2011) <sup>53, 61, 68</sup>	Inclusion criteria:	Median age (IQR): 64(53,74)	Roche
	Consecutive patients presenting to the emergency department with chest	Male (%): 60	
Country: New Zealand	pain; participants were eligible for inclusion if the attending clinician had	White (%): 85	
	sufficient suspicion of ACS that serial troponins and ECGs were considered	Previous CAD (%): 54	
Funding: Manufacturers (Roche and Abbott)	necessary	Previous Family History (%): 40	
supplied assays. The study was funded by a New		Diabetes (%): 16	
Zealand National Heart Foundation grant	Exclusion criteria:	Smoking (%): 45	
	<18 years; samples not stored for both time points (on admission and at	Hypertension (%): 46	
Recruitment: November 2006 - April 2007	6-24 hours)	Dyslipidaemia (%): 38	
		Median (IQR) time to presentation	
Number of participants: 332	Patient category:	(hours): 4.0 (2.0 to 8.6)	
	Mixed		

Study Details	Selection criteria	Participant details	Test
			Manufacturer
Body(2011) <sup>59, 67, 76</sup>	Inclusion criteria:	Mean age (sd): 59(14)	Roche
	Presenting to ED with chest pain; age >25 years and chest pain within	Male (%): 61	
Country: UK	previous 24h that initial treating physician suspected may be cardiac in	Kidney Disease (%):1	
	nature.	Previous AMI (%): 24	
Funding: Central Manchester NHS Trust		Previous Family History (%): 48	
	Exclusion criteria:	Previous Revascularisation (%): 20	
Recruitment: January 2006 - February 2007	renal failure requiring dialysis, trauma with suspected myocardial	Diabetes (%): 18	
	contusion, or another medical condition mandating hospital admission or	Smoking (%): 31	
Number of participants eligible (enrolled):	if they did not consent to and provide a blood sample for use by the	Dyslipidaemia (%): 48	
1004(703)	research team	Median time to presentation	
		(hours): 3.5 hours	
	Patient category:		
	Mixed		
Christ(2010) <sup>56</sup>	Inclusion criteria:	Mean age (SD): 66(16)	Roche
	Consecutive patients with acute chest pain of possible coronary origin	Male (%): 64	
Country: Germany	presenting to the emergency department	Previous AMI (%): 32	
		Previous CAD (%): 34	
Funding: hsTnT test kits were provided by Roche	Exclusion criteria: NR	Previous Family History (%): 12	
		Previous Revascularisation (%): 24	
Recruitment: 7/9/2009 - 21/9/2009	Patient category:	Diabetes (%): 22	
	Mixed	Smoking (%): 22	
Number of participants: 137		Hypertension (%): 66	
		Dyslipidaemia (%): 35	
		Mean BMI (SD): 28(5)	
		Time to presentation:	
		0-2h 36%; 2-6h 22%; 6-24 h 33%;	
		>24h 20%	

Study Details	Selection criteria	Participant details	Test
			Manufacturer
Collinson(2013) <sup>58, 65, 70</sup>	Inclusion criteria:	Median age (IQR): 54(44,64)	Roche
	Patients presenting to the emergency department with chest pain due to	Male (%): 60	
Country: UK	suspected, but not proven AMI	Previous AMI (%): 40	
		Previous Family History (%):	
Funding: UK Health Technology Assessment	Exclusion criteria:	Previous Revascularisation (%): 1	
Programme	ECG changes diagnostic for AMI or high risk ACS (>1 mm ST deviation, or	Diabetes (%): 8	
	>3 mm inverted T waves); known CAD with prolonged (>1 hr) or recurrent	Smoking (%): 28	
Study Name: Point of care arm of the RATPAC study	typical cardiac-type pain; proven or suspected serious non-cardiac	Hypertension (%): 35	
	pathology (e.g. PE); co-morbidity or social problems requiring hospital	Dyslipidaemia (%): 24	
Recruitment: February 2007 - June 2008	admission even if AMI ruled out; obvious non-cardiac cause of chest pain	Median (IQR) time to presentation	
	(e.g. pneumothorax or muscular pain); presentation >12 hrs after most	(hours): 8.25 (5.17 to 12.30)	
Number of participants: 850	significant episode of pain		
	Patient category: NSTEMI		
Cullen(2013) <sup>62</sup>	Inclusion criteria:	Mean age (SD): 59(13)	Abbott
	Prospectively recruited adults with at least 5 min of	Male (%): 60	
Country: New Zealand and Australia	possible cardiac symptoms in accordance with the American	Previous AMI (%): 24	
	Heart Association case definitions (acute chest, epigastric, neck, jaw, or	Previous Family History (%): 57	
Funding: The manufacturers (Abbott, Roche and	arm pain; or discomfort or pressure without a clear non-cardiac so	Previous Revascularisation (%): 8	
Siemens) provided partial funding		Diabetes (%): 15	
	Exclusion criteria:	Smoking (%): 18	
Study Name: ADAPT study	Pregnancy; unable or unwilling to consent; recruitment inappropriate (e.g.	Hypertension (%): 52	
(ACTRN12611001069943)	terminal illness); transfer from another hospital; follow-up considered	Dyslipidaemia (%): 57	
	impossible (e.g. homeless patients)	Mean (SD) time to presentation	
Recruitment: November 2007 - February 2011		(hours): 22.3 (60.5)	
	Patient category:		
Number of participants: 1635	Mixed		

Study Details	Selection criteria	Participant details	Test
42			Manufacturer
Eggers(2012) <sup>43</sup>	Inclusion criteria:	Median age (IQR): 67(58,76)	Roche
	Chest pain with ≥15 min duration within the last 24h (FAST II-study), or	Male (%): 66	
Country: Sweden	the last 8 h (FASTER I-study). Analysis restricted to patients with symptom	Previous AMI (%): 38	
	onset <8h.	Previous Revascularisation (%): 18	
Funding: Swedish Society of Medicine and the		Diabetes (%): 18	
Selander Foundation	Exclusion criteria:	Smoking (%): 18	
	ST-segment elevation on the admission 12-lead ECG leading to immediate	Hypertension (%): 43	
Study Name: FASTER 1-study and FAST II study	reperfusion therapy or its consideration was used as exclusion criterion.	Dyslipidaemia (%): 38	
		Delay <4 hours (%): 40	
Recruitment: May 2000 (FAST II), October 2002	Patient category:		
(FASTER I) - March 2001 (FAST II), August 2003	NSTEMI		
(FASTER I)			
Number of participants eligible (enrolled):			
495(360)			
Freund(2011) <sup>48, 72</sup>	Inclusion criteria:	Mean (SD): 57(17)	Roche
Country: France	Consecutive adults (>18 years) presenting to the emergency department	Male (%): 65	
	with chest pain suggestive of ACS (onset or peak within the previous 6 hrs)	Previous CAD (%): 26	
Funding: Assay kits for the study were provided by		Previous Family History (%): 32	
the manufacturers (Roche)	Exclusion criteria:	Diabetes (%): 14	
	Patients with acute kidney failure requiring dialysis were excluded	Smoking (%): 40	
Study Name:		Hypertension (%):	
	Patient category:	Dyslipidaemia (%): 36	
Recruitment: August 2005 - January 2007	Mixed (13 were STEMI and 32 NSTEMI)		
Number of participants: 317			

Study Details	Selection criteria	Participant details	Test
			Manufacturer
Hoeller(2011) <sup>42, 44, 46, 51, 52, 55, 62, 64, 66, 73, 75</sup>	Inclusion criteria:	Median age (IQR): 62( 50, 75)	Roche ,
Country: Switzerland, Spain, USA and Germany	Consecutive adults presenting to the ED with symptoms suggestive of AMI	Male (%): 69	Abbott,
	(e.g. acute chest pain, angina pectoris at rest, other thoracic sensations)	Previous AMI (%): 24	Beckman
Funding: Swiss National Science Foundation, Swiss	within an onset or peak within the last 12 hours	Previous CAD (%): 34	Coulter
Heart Foundation, Department of Internal Medicine		Previous Family History (%): 43	
of the University Hospital Basel, Roche, Siemens,	Exclusion criteria:	Previous Revascularisation (%): 24	
Abbott, Brahms, nanosphere, and 8sense	Terminal kidney failure requiring dialysis	Diabetes (%): 18	
		Smoking (%): 61	
Study Name: APACE trial (NCT00470587)	Patient category:	Hypertension (%): 64	
	Mixed	Dyslipidaemia (%): 45	
Recruitment: April 2006 - August 2011		Median BMI (IQR): 27(24, 30)	
		Presenting <3 hours from	
Number of participants: 2245		symptom onset (%): 24	
Keller(2011) <sup>47, 69</sup>	Inclusion criteria:	Mean age (sd): 61(14)	Abbott
	Consecutive adults (18-85 years) presenting to three chest pain units with	Male (%): 66	
Country: Germany	chest pain suggestive of ACS	Previous CAD (%): 36	
		Previous Family History (%): 32	
Funding: Abbott Diagnostics provided study funding	Exclusion criteria:	Diabetes (%): 16	
	Major surgery or trauma within the previous 4 weeks; pregnancy;	Smoking (%): 24	
Recruitment: January 2007 - December 2008	intravenous drug abuse; anaemia (haemoglobin <10 g/dL)	Hypertension (%): 74	
		Dyslipidaemia (%): 73	
Number of participants: 1818	Patient category:	Mean BMI (sd): 28(5)	
	Mixed		

Study Details	Selection criteria	Participant details	Test
			Manufacturer
Kurz(2011) <sup>54</sup>	Inclusion criteria:	Mean age (sd): 66(11)	Roche
	Consecutive patients admitted to a chest pain unit with symptoms	Male (%): 71	
Country: Germany	suggestive of ACS	Previous AMI (%): 37	
		Previous CAD (%): 50	
Funding: Investigators were supported by Roche	Exclusion criteria:	Previous Family History (%): 32	
diagnostics and assay kits were also provided by the	ST-segment elevation; severe kidney dysfunction(GFR <60 mL/min/1.73	Previous Revascularisation (%): 17	
manufacturer	m <sup>2</sup> ); patients undergoing PCI during follow-up sampling	Diabetes (%): 31	
		Smoking (%): 22	
Recruitment: May 2008 - December 2008	Patient category:	Hypertension (%): 78	
	NSTEMI	Dyslipidaemia (%): 65	
Number of participants: 94		Median symptom onset (IQR,	
		minutes): 358 (152, 929)	
		BMI (95% CI/range/IQR): 28(4)	
Lippi(2012) <sup>74</sup>	Inclusion criteria:	No participant details reported	Beckman
	Consecutive patients presenting to the emergency department with chest		
Country: Italy	pain, within 3 hours of the onset of pain		
Funding: NR	Exclusion criteria:		
	None reported		
Recruitment: NR			
	Patient category:		
Conference abstract only	Mixed		
Number of participants: 57			

Study Details	Selection criteria	Participant details	Test
			Manufacturer
Melki(2011) <sup>50, 60</sup>	Inclusion criteria:	Median age (IQR): 65(55,76)	Roche
Country: Sweden	Patients admitted to a coronary care unit with chest pain or other	Male (%): 67	
	symptoms suggestive of ACS within 12 hours of admission	Previous AMI (%): 30	
Funding: Partially supported by a grant from Roche		Previous Revascularisation (%): 21	
Diagnostics, who also provided reagents. Also	Exclusion criteria:	Diabetes (%): 23	
supported by the Swedish Heart and Lung	Patients with persistent ST-segment elevation	Smoking (%): 17	
Foundation and National Board of Health and		Hypertension (%): 50	
Welfare	Patient category:	Mean symptom onset (95%	
	NSTEMI	Cl/range/IQR, hours): 5 (3, 8)	
Recruitment: August 2006 - January 2008			
Number of participants: 233			
Parsonage(2013) <sup>57</sup>	Inclusion criteria:	Mean age (IQR): 54(44,65)	Abbott, Roche
	Patients with symptoms of possible ACS	Male (%): 60	
Country: Australia			
	Exclusion criteria:		
Funding: Not reported	None reported		
Recruitment: NR	Patient category:		
	Mixed		
Conference abstract only			
Number of participants: 737			

Study Details	Selection criteria	Participant details	Test
			Manufacturer
Saenger(2010) <sup>71</sup>	Inclusion criteria:	No further participant details	Roche
Country: USA	Patients presenting to the emergency department with symptoms	reported	
	suggestive of AMI		
Funding: Two authors declared individual funding			
from manufacturers (one from Roche diagnostics	Exclusion criteria:		
and one from Beckman Coulter and Abbott)	None reported		
Study Name:	Patient category:		
	Mixed		
Recruitment: NR - NR			
	Details:		
Conference abstract only	NSTEMI 19%, STEMI 15%		
Number of participants: 288			
Sanchis(2012) <sup>41</sup>	Inclusion criteria:	Mean age (sd): 60(12)	Roche
Country: Spain	Patients presenting to the emergency department with chest pain of	Male (%): 59	
	possible coronary origin and onset of pain within the previous 24 hrs	Previous Family History (%): 14	
Funding: Supported by a grant from Roche		Diabetes (%): 20	
Diagnostics	Exclusion criteria:	Smoking (%): 25	
	Exclusion criteria: persistent ST-segment elevation on ECG; troponin	Hypertension (%): 54	
Study Name: PITAGORAS study	elevation in any of 2 serial determinations (at arrival and 6-8 hours later);	Dyslipidaemia (%): 46	
	prior diagnosis of ischemic heart disease by either the finding of		
Recruitment: NR	significant stenosis in a prior coronary angiogram or previously		
	documented AMI; left bundle-branch block or other non-interpretable		
Number of participants: 446	ECG or inability to performance exercise test; structural heart disease		
	different to ischemic heart disease; concomitant heart failure or		
	significant bradyarrhythmia (<55 beat/min) or tachyarrythmia (>110		
	beat/min) at admission.		
	Patient category:		
	NSTEMI		

Study Details	Selection criteria	Participant details	Test Manufacturer
Santaló(2013) <sup>39</sup>	Inclusion criteria:	Mean age (range): 69( 27, 93)	Roche
	Adult (>18 years) described as presenting with acute coronary syndromes	Male (%): 68	
Country: Spain	and symptom duration ≥5 min; population included 174 people with a	Previous CAD (%): 35	
	final diagnosis of non-acute coronary syndromes.	Diabetes (%): 26	
Funding: Reagents and logistical support were		Hypertension (%): 62	
provided by Roche diagnostics	Exclusion criteria:	Presentation within 3 hours:	
	Exclusion criteria: ST-segment elevation; new left bundle branch block;	46.2%	
Study Name: TUSCA study	pre-admission thrombolytic therapy; defibrillation or cardioversion before		
	sampling; pregnancy; renal failure requiring dialysis; unstable angina		
Recruitment: NR	within 2 months; CABG within 3 month		
Number of participants: 258	Patient category:		
Number of participants. 558	NSTEMI		
Sebbane(2013) <sup>63</sup>	Inclusion criteria:	Median age (IOR): 61(48,75)	Roche
	Adults presenting to the emergency department with chest pain of recent	Male (%): 63	
Country: France	(within 12 hrs of presentation)		
Funding: Study funded by the hospital, with assay	Exclusion criteria:		
reagents supplied by the manufacturers	Traumatic causes of chest pain. STEMI was defined by the persistent		
	elevation of the ST segment of at least 1 mm in 2 contiguous ECG leads or		
Recruitment: December 2009 - November 2011	by the presence of a new left bundle-branch block with positive cardiac		
	enzyme results. Patients with STEMI were excluded from the analysis for		
Number of participants: 248	our review.		
	Detions of a second		
	Patient category:		
	INSTEINIT (Data also reported for mixed Alvii but not extracted)		

D. Index lest and reference standard details	b.	Index test	and reference	standard details
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Study Details	High sensi	tivity tr	oponin de	tails (ng/L)	Reference standard details								
	Manu-	LoD	99 <sup>th</sup>	Coefficient	Target	Time	Reference	Standard troponin	Observer				
	facturer		Centile	of variation	Condition	frame	standard						
Aldous (2012) <sup>40, 45, 49</sup>	Roche Elecsys hs- cTnT	5	14	<10% at 13	NSTEMI	NR	ACC <sup>111</sup>	Conventional troponins were measured using Abbott Diagnostics TnI (LoD 10 ng/L, 99th centile 28 ng/L, CV <10% at 32 ng/L, decision threshold 30 ng/L) <i>Timing:</i> On presentation, and at 2 hours and 6-12 hours	Diagnoses on admission and at follow-up were independently adjudicated by one cardiologist, who was blinded to hs-TnT results				
Aldous (2011) <sup>53, 61, 68</sup>	Roche Elecsys hs- cTnT	5	14	<10% at 13	AMI	NR	Joint ESC, ACC, AHA and WHF <sup>8</sup>	Conventional troponins were measured using Abbott Diagnostics TnI 2 (LoD 10 ng/L, 99th centile 28 ng/L, CV <10% at 32 ng/L) Change (rise or fall) in TnI 2, or no change but no clear alternative cause of troponin elevation, were considered indicative of AMI. <i>Timing:</i> On presentation and at follow- up (6-24 hours)	Final diagnoses were adjudicated independently by cardiologists, blinded to patient history and hs-TnT				

Study Details	High sensi	tivity tı	oponin de	tails (ng/L)	Reference standard details								
	Manu-	LoD	99 <sup>th</sup>	Coefficient	Target	Time	Reference	Standard troponin	Observer				
	facturer		Centile	of variation	Condition	frame	standard						
Body (2011) <sup>59,</sup>	Roche	NR	14	<10% at 9	AMI	12	Joint ESC,	Rise or fall of cTnT, or both, above the	2 independent investigators who had				
67, 76	Elecsys hs-					hours	ACC, AHA	99th percentile (10 ng/l) in the	all clinical, laboratory, and imaging				
	cTnT						and World	appropriate clinical context. For	data available for review, but who				
							Heart	patients with modest elevations of	were blinded to hs-CTnT levels.				
							Federation	cTnT (<0.1 ng/ml) at baseline, an					
							(WHF) <sup>8</sup>	absolute difference of at least 20 ng/l					
								on serial sampling was considered to					
								represent a significant rise, fall, or both					
								based on the analytical performance of					
								the cTnT assay.					
								Timing: at least 12 h after the onset of					
								the most significant symptoms.					

Study Details	High sensit	tivity tr	oponin de	tails (ng/L)	Reference standard details								
	Manu-	LoD	99 <sup>th</sup>	Coefficient	Target	Time	Reference	Standard troponin	Observer				
	facturer		Centile	of variation	Condition	frame	standard						
Christ	Roche	3	14	<10% at 13	AMI	NR	Joint ESC,	Myocardial necrosis was diagnosed on	Two independent consultants				
(2010) <sup>56</sup>	Elecsys hs-						ACC, AHA	the basis of a rising and/or falling cTnT					
	cTnT						and WHF <sup>8</sup>	pattern >20% or <20% compared to the					
								cTnT levels admission) with at least one					
								value above the 99th percentile and an					
								imprecision of <10%. Myocardial					
								necrosis not related to AMI was					
								defined as a typical rise and fall of cTnT					
								levels without clinical evidence of					
								coronary artery disease, and cardiac					
								pain without necrosis was defined as a					
								typical patient history and clinical signs					
								of cardiac pain without increased levels					
								of cTnT. Unstable angina was					
								diagnosed when a patient had normal					
								troponin levels and typical angina at					
								rest or exercise, or a cardiac					
								catheterization result compatible with					
								the diagnosis. cTnT cut-off level of					
								0.04 ug/L,					
								Timing: At presentation and about 6					
								hours at discretion of physician					

Study Details	High sensit	tivity tı	roponin de	etails (ng/L)	Reference standard details								
	Manu-	LoD	99 <sup>th</sup>	Coefficient	Target	Time	Reference	Standard troponin	Observer				
	facturer		Centile	of variation	Condition	frame	standard						
Collinson (2013) <sup>58, 65, 70</sup>	Roche Elecsys hs- cTnT	3	14	<10% at 13	NSTEMI	NR	Joint ESC, ACC, AHA and WHF <sup>8</sup>	Conventional troponins were measured using one of the following methods: Siemens cTnI Ultra (LoD 6 ng/L, 99th centile 40 ng/L, CV 10% at 30 ng/L; Abbott cTnI (LoD 10 ng/L, 99th centile 12 ng/L, CV 10% at 32 ng/L; Beckman AccuTnI (LoD 10 ng/L, 99th centile 40 ng/L, CV 10% at 60 ng/L; Roche cTnT (LoD 10 ng/L, 99th centile 10 ng/L, CV 10% at 30 ng/L Timing: On presentation and at 10 to 12 hours	An initial working diagnosis was recorded by the senior emergency department clinician and reviewed by two independent clinicians; all were blind to hs-TnT results				
Cullen (2013) <sup>62</sup>	Abbott ARCHITECT hs-cTnI <i>STAT</i>	1.2	26.2	<5% at 26.2	MACE	30 days	MACE	NR	Adjudication of all cardiac endpoints was made by two cardiologists, with consultation of a third cardiologist in case of disagreement				
Eggers (2012) <sup>43</sup>	Roche Elecsys hs- cTnT	3	14	<10% at 13	NSTEMI	NR	Joint ESC, ACC, AHA and WHF <sup>8</sup>	cTnI (Stratus CS, Siemens Healthcare Diagnostics, Deerfield, IL, USA). Non- STEMI defined as: cTnI above the 99th percentile of 0.07 μg/L at least at one measurement together with a ≥20% rise and/or fall and an absolute change ≥0.05 μg/L within 24 h. To allow for the calculation of relative changes, cTnI was set to 0.02 μg/L (i.e. a concentration below the lowest level of detection) when reported as 0.00 or 0.01 μg/L. <i>Timing:</i> eight time points during the first 24 h following enrolment	Not reported				

Study Details	High sensi	tivity tr	oponin de	tails (ng/L)	Reference standard details								
	Manu-	LoD	99 <sup>th</sup>	Coefficient	Target	Time	Reference	Standard troponin	Observer				
	facturer		Centile	of variation	Condition	frame	standard						
Freund (2011) <sup>48, 72</sup>	Roche Elecsys hs- cTnT	3	14	<10% at 14	АМІ	30 days	Joint ESC, ACC, AHA and WHF <sup>8</sup>	cTnl (Siemens Healthcare Diagnostica Inc., NewaRK, USA or Access analyser Beckman Coulter Inc., Brea, USA). Threshold for Siemens assay 140 ng/L, CV ≤10% Threshold for Beckman assay 60 ng/L, CV 10% <i>Timing:</i> On presentation and at 3-9 hours if needed	Two independent emergency department physicians, who were blinded to hs-TnT results. Disagreements were adjudicated by a third emergency department physician.				
Hoeller (2011) APACE <sup>46, 51, 52,</sup> 55, 64, 75	Roche Elecsys hs- cTnT	5	14	<10% at 13	AMI	NR	Joint ESC, ACC, AHA and WHF <sup>8</sup>	Conventional troponins were measured using Roche cTnT 4th generation assay (CV <10% at 35 ng/L),	Final diagnoses were adjudicated by two independent cardiologists blind to hsTnT results. Where there was				
APACE <sup>73</sup>		2						Beckman Coulter Accu cTnI (CV <10%	disagreement a third cardiologist was				
APACE <sup>64, 75</sup>	Beckman (pre- commercial assay)	2	9	<10% at 9	AMI and NSTEMI	30 days		at 60 ng/L), or Abbott Axsym cTnI ADV (CV <10% at 160 ng/L). A positive test was defined as change ≥30% of 99th centile or 10% CV level, within 6 to 9	consulted.				
APACE <sup>62</sup>	Abbott ARCHITECT hs-cTnl <i>STAT</i>	1.2	26.2	<5% at 26.2	AMI	30 days		hours. <i>Timing:</i> On presentation and at 6 to 9 hours					
APACE <sup>75</sup>					MACE			NA	Adjudication of all cardiac endpoints was made by two cardiologists, with consultation of a third cardiologist in case of disagreement				
Keller (2011) <sup>47</sup>	Abbott ARCHITECT hs-cTnl <i>STAT</i>	3.4	24-30 for this study popul- ation	10% at 5.2	AMI	30 days	Joint ESC, ACC, AHA and WHF <sup>8</sup>	Conventional serial troponin T or I (no further details) <i>Timing:</i> On presentation and at 3 and 6 hours	Final diagnosis adjudicated by two independent cardiologists, with disagreements referred to a third cardiologist; all three were blinded to hs-Tnl results				

Study Details	High sensit	tivity tı	oponin de	tails (ng/L)	Reference standard details								
	Manu- LoD 99 <sup>th</sup> Coefficient Targe					Time	Reference	Observer					
	facturer		Centile	of variation	Condition	frame	standard						
Kurz (2011) <sup>54</sup>	Roche Elecsys hs- cTnT	3	13.5	8% at 10	NSTEMI	24 hours	Joint ESC, ACC, AHA and WHF <sup>8</sup>	4th generation cTnT (Roche Elecsys, Mannheim, Germany) LoD 10 ng/L, diagnostic threshold 30 ng/L Diagnosis of NSTEMI required elevated cTnT concentration in at least one of the consecutive samples collected within 24 hours of the index event <i>Timing:</i> On presentation, at 6 hours and at least one sample between presentation and 6 hours	NR				
Lippi (2012) <sup>74</sup>	Beckman Coulter prototype hs-cTnI (hs- Accu-TnI)	2.1	8.6	NR	AMI	NR	AMI (unclear method)	NR	NR				
Melki (2011) <sup>50, 60</sup>	Roche Elecsys hs- cTnT	2	14	<10% at 13	NSTEMI	NR	Joint ESC, ACC, AHA and WHF <sup>8</sup>	Conventional troponin Roche 4th generation TnT (LoD 10 ng/L, 10% CV at 35 ng/L), or Beckman Coulter Access AccuTnI (LoD 10 ng/L, 99th centile 40 ng/L, CV <10% at 60 ng/L <i>Timing:</i> On presentation and 9 to 12 hours later	Final diagnosis determined by the individual cardiologist, then adjudicated by two independent evaluators; all three were blinded to hs-TnT results				
Parsonage (2013) <sup>57</sup>	Abbott ARCHITECT hs-cTnI STAT	NR	26.2	NR	AMI	NR	AMI (unclear method)	NR <i>Timing:</i> On admission and >6 hours after presentation	Final diagnosis was adjudicated by two independent cardiologists				
Saenger (2010) <sup>71</sup>	Roche Elecsys hs- cTnT	NR	14	NR	AMI	NR	AMI (unclear method)	NR	NR				

Study Details	High sensi	tivity tı	oponin de	tails (ng/L)	Reference standard details								
	Manu- facturer	LoD	99 <sup>th</sup> Centile	Coefficient of variation	Target Condition	Time frame	Reference standard	Standard troponin	Observer				
Sanchis (2012) <sup>41</sup>	Roche Elecsys hs- cTnT	3	14	<10% at 14	MACE	30 days	MACE	NR	NR				
Santaló (2013) <sup>39</sup>	Roche Elecsys hs- cTnT	NR	14	10% at 9.3	NSTEMI	NR	National Academy of Clinical Biochemistr y and International Federation of Clinical Chemistry Committee <sup>10</sup>	Roche cTnT; NSTEMI was defined as cTnT >10 ng/L and ΔcTnT >20% <i>Timing:</i> 30 minutes after arrival and at 2,4 and 6-8 hours or until discharge	Final diagnosis was made by an adjudication committee				
Sebbane (2013) <sup>63</sup>	Roche Elecsys hs- cTnT	5	14	<10% at 13	NSTEMI	NR	Joint ESC, ACC, AHA and WHF <sup>8</sup>	cTnI measured using the Access2 analyser (Access Immunosystem, Beckman Instruments, France). The LoD was <10 ng/L and the decision threshold was 40 ng/L <i>Timing:</i> Convention cardiac troponin (cTnI) on presentation, 6 hrs later and beyond as needed	Two independent emergency department physicians, blinded to hs- cTnT results				

## c. Study results

Study Details	Troponin	Timing	Threshold	Target	TP	FP	FN	TN	Sens	Spec	LR+ (95% CI)	LR- (95% CI)
	Assay		(ng/L)	Condition					(95% CI)	(95% CI)		
Aldous (2011) <sup>53</sup>	Roche	On presentation	14	AMI	92	36	18	186	83 (75, 89)	84 (78, 88)	5.1 (3.7, 6.9)	0.2 (0.13, 0.3)
			5		106	131	4	91	96 (90, 98)	41 (35, 48)	1.6 (1.4, 1.8)	0.1 (0.04, 0.25)
			13		92	38	18	184	83 (75, 89)	83 (77, 87)	4.8 (3.6, 6.5)	0.2 (0.13, 0.31)
			15		93	29	17	193	84 (76, 90)	87 (82, 91)	6.4 (4.5, 9)	0.18 (0.12, 0.28)
Aldous (2012) <sup>40</sup>	Roche	On presentation	14	AMI	74	54	8	249	90 (81, 95)	82 (77, 86)	5 (3.9, 6.4)	0.12 (0.07, 0.24)
		0-1 hours after	14		77	63	5	240	93 (86 <i>,</i> 97)	79 (74, 83)	4.5 (3.6, 5.6)	0.08 (0.04, 0.19)
		presentation										
		0-2 hours after	14		78	67	4	236	95 (87 <i>,</i> 98)	78 (73, 82)	4.3 (3.4, 5.3)	0.07 (0.03, 0.17)
		presentation										
		On presentation	14 and no		78	74	4	229	95 (87 <i>,</i> 98)	75 (70, 80)	3.9 (3.1, 4.7)	0.07 (0.03, 0.18)
		and at 2 hours	change									
			<14 and $\Delta$ 20%		49	81	33	222	60 (49, 70)	73 (68, 78)	2.2 (1.7, 2.9)	0.55 (0.42, 0.72)
			14 and Δ20%		46	23	36	280	56 (45 <i>,</i> 66)	92 (89, 95)	7.2 (4.7,	0.48 (0.37, 0.61)
											11.2)	
			14 or Δ 20%		81	131	1	172	98 (93, 100)	57 (51, 62)	2.3 (2, 2.6)	0.03 (0.01, 0.16)
Aldous (2012) <sup>45</sup>	Roche	On presentation	14	NSTEMI	181	134	24	600	88 (83 <i>,</i> 92)	82 (79, 84)	4.8 (4.1, 5.7)	0.15 (0.1, 0.21)
		On presentation	5		192	305	13	429	93 (89 <i>,</i> 96)	58 (55, 62)	2.2 (2, 2.5)	0.11 (0.07, 0.19)
		On presentation	3		196	383	9	351	95 (92 <i>,</i> 98)	48 (44, 51)	1.8 (1.7, 2)	0.1 (0.05, 0.18)
		2 hours after	14		189	149	16	585	92 (87, 95)	80 (77, 82)	4.5 (3.9, 5.2)	0.1 (0.06, 0.16)
		presentation	5		196	340	9	394	95 (92, 98)	54 (50, 57)	2.1 (1.9, 2.2)	0.09 (0.05, 0.16)
			3		201	424	4	310	98 (95 <i>,</i> 99)	42 (39, 46)	1.7 (1.6, 1.8)	0.05 (0.02, 0.13)
Data from:		0-2 hours after	Peak 14		189	149	11	590	94 (90, 97)	80 (77, 83)	4.7 (4, 5.4)	0.07 (0.04, 0.13)
Aldous (2011) <sup>49</sup>		presentation	14 and Δ20%		99	43	101	696	50 (43, 56)	94 (92, 96)	8.4 (6.1,	0.54 (0.47, 0.62)
											11.6)	
			14 or ∆20%		195	260	5	479	97 (94, 99)	65 (61, 68)	2.8 (2.5, 3.1)	0.04 (0.02, 0.1)
Body (2011) <sup>67</sup>	Roche	On presentation	3	AMI	130	378	0	195	100 (96, 100)	34 (30, 38)	1.5 (1.4, 1.6)	0.01 (0, 0.18)
		On presentation	14		111	101	19	472	85 (78, 90)	82 (79, 85)	4.8 (4, 5.8)	0.18 (0.12, 0.27)
		On presentation:	3		79	89	0	156	99 (94, 100)	64 (57, 69)	2.7 (2.3, 3.2)	0.01 (0, 0.16)
		Symptom onset										
		<3h										

Study Details	Troponin	Timing	Threshold	Target Condition	TP	FP	FN	TN	Sens	Spec	LR+ (95% CI)	LR- (95% CI)
	Assay	On procentation:	(IIg/L)	Condition	62	12	12	202			19(26 6 1)	0.21 (0.12, 0.25)
		Sumptom onsot	14		05	42	12	205	02 (72, 09)	05 (70, 07)	4.8 (5.0, 0.4)	0.21 (0.15, 0.55)
		symptom onset										
			2		<b>F</b> 4	224	0	107	00 (01 100)	22 (20, 20)		0.02 (0.0.47)
		On presentation:	3		51	221	0	107	99 (91, 100)	33 (28, 38)	1.5 (1.4, 1.6)	0.03 (0, 0.47)
		Symptom onset										
		>3n							04 (04 05)	00 (77, 00)		
		On presentation:	14		47	59	4	269	91 (81, 96)	82 (77, 86)	5.1 (4, 6.5)	0.11 (0.04, 0.26)
		Symptom onset										
		>3h										
		On presentation:	3		105	253	0	133	100 (96, 100)	34 (30, 39)	1.5 (1.4, 1.6)	0.01 (0, 0.22)
		Symptom onset										
		<6h										
		On presentation:	14		87	66	18	320	83 (74 <i>,</i> 89)	83 (79, 86)	4.8 (3.8, 6.1)	0.21 (0.14, 0.32)
		Symptom onset										
		<6h										
		On presentation:	3		25	125	0	62	98 (84, 100)	33 (27, 40)	1.5 (1.3, 1.6)	0.06 (0, 0.91)
		Symptom onset										
		>6h										
		On presentation:	14		24	35	1	152	94 (78, 99)	81 (75, 86)	5 (3.7 <i>,</i> 6.8)	0.07 (0.02, 0.34)
		Symptom onset										
		>6h										
Christ (2010) <sup>56</sup>	Roche	On presentation	14	AMI	19	45	1	72	93 (74, 98)	61 (52 <i>,</i> 70)	2.4 (1.9, 3.1)	0.12 (0.02, 0.55)
Christ (2010) <sup>56</sup>	Roche	On presentation	14	AMI	20	92	0	25	100 (81, 100)	22 (15, 30)	1.25 (1.11,	0.11 (0.01, 1.74)
											1.40)	
Collinson	Roche	On presentation	14	NSTEMI	53	33	14	733	79 (68, 87)	96 (94, 97)	18 (12.6,	0.22 (0.14, 0.35)
(2013) <sup>65</sup>											25.7)	
		On presentation	Peak 14	NSTEMI	57	43	11	736	83 (73, 90)	94 (93, 96)	14.9 (11,	0.18 (0.1, 0.3)
		and at 1.5 hours									20.3)	
Cullen (2013) <sup>62</sup>	Abbott	On presentation	26.2 on	MACE	227	96	20	129	92 (88, 95)	93 (92, 94)	13.2 (10.9,	0.09 (0.06, 0.13)
. ,		and at 2 hours	admission and					2			16.1)	
			at 2 hours									
Eggers (2012) <sup>43</sup>	Roche	On presentation	14	NSTEMI	101	59	27	173	79 (71, 85)	74 (68, 80)	3.1 (2.4, 3.9)	0.29 (0.2, 0.4)
			45.7	NSTEMI	65	11	63	221	51 (42, 59)	95 (91, 97)	10.3 (5.7.	0.52 (0.43, 0.62)
										(, )	18.5)	
Study Details	Troponin	Timing	Threshold	Target	TP	FP	FN	TN	Sens	Spec	LR+ (95% CI)	LR- (95% CI)
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	Assay		(ng/L)	Condition					(95% CI)	(95% CI)		
Freund (2011) <sup>48</sup>	Roche	On presentation	14	AMI	42	48	3	224	92 (81 <i>,</i> 97)	82 (77 <i>,</i> 86)	5.2 (4, 6.8)	0.09 (0.03, 0.25)
		On presentation:			20	36	2	200	89 (70 <i>,</i> 97)	85 (79 <i>,</i> 89)	5.8 (4.2, 8.1)	0.13 (0.04, 0.41)
		Low/moderate										
		pre-test										
		probability										
		On presentation:			22	12	1	24	94 (77, 99)	66 (50 <i>,</i> 79)	2.8 (1.7, 4.4)	0.09 (0.02, 0.45)
		High pre-test										
11 II (2014) <sup>75</sup>		probability			200	262	10	120	00 (00 00)	70 (76, 00)		0.42 (0.4.0.40)
Hoeller (2011)	Roche	On presentation	14	AMI	398	363	46	126 5	90 (86, 92)	78 (76, 80)	4 (3.6, 4.4)	0.13 (0.1, 0.18)
		On presentation:	14		79	63	28	335	74 (65, 81)	84 (80 <i>,</i> 87)	4.6 (3.6, 6)	0.31 (0.23, 0.43)
		Symptom onset										
		<3h										
		On presentation:	14		318	300	18	931	95 (92 <i>,</i> 96)	76 (73 <i>,</i> 78)	3.9 (3.5, 4.3)	0.07 (0.05, 0.11)
		Symptom onset										
		≥3h										
	Beckman	On presentation	9		209	231	18	693	92 (88, 95)	75 (72, 78)	3.7 (3.3, 4.1)	0.11 (0.07, 0.17)
	Abbott		26.2		240	93	71	116	77 (72, 81)	93 (91, 94)	10.4 (8.4,	0.25 (0.2, 0.3)
							-	3	(00 (00 (00))	/	12.7)	
Data from: $(2000)^{73}$	Roche	On presentation	2		123	512	0	83	100 (97, 100)	14 (11, 17)	1.2 (1.1, 1.2)	0.03 (0.00, 0.46)
Reichlin (2009)	Abbott		10		116	77	7	518	94 (89, 98)	87 (84, 90)	7.3 (5.9, 9.0)	0.07 (0.03, 0.13)
Data from: Reiter (2011) <sup>52</sup>	Roche	On presentation: >70 years only	5		98	305	0	3	99 (95, 100)	1 (0, 3)	1 (1, 1)	0.45 (0.02, 8.56)
		On presentation:	14		96	157	2	151	97 (92, 99)	49 (44, 55)	1.9 (1.7, 2.1)	0.05 (0.02, 0.18)
		>70 years only										
		On presentation:	14		54	87	7	533	88 (78 <i>,</i> 94)	86 (83 <i>,</i> 88)	6.2 (5, 7.7)	0.14 (0.07, 0.28)
		≤70 years										
Data from:		On presentation:	14		73	142	5	213	93 (85 <i>,</i> 97)	60 (55 <i>,</i> 65)	2.3 (2, 2.7)	0.12 (0.05, 0.26)
Potocki (2012) <sup>46</sup>		with pre-existing										
		CAD										
		On presentation:	14		100	114	6	517	94 (88 <i>,</i> 97)	82 (79 <i>,</i> 85)	5.2 (4.4, 6.2)	0.07 (0.04, 0.16)
		without pre-										
-		existing CAD										
Data from:		On presentation	11		129	177	3	454	97 (93 <i>,</i> 99)	72 (68, 75)	3.5 (3.1, 3.9)	0.04 (0.01, 0.1)

Study Details	Troponin	Timing	Threshold	Target	TP	FP	FN	TN	Sens	Spec	LR+ (95% CI)	LR- (95% CI)
	Assay		(ng/L)	Condition					(95% CI)	(95% CI)		
Hochholzer		On presentation	11	NSTEMI	90	177	3	454	96 (90 <i>,</i> 99)	72 (68, 75)	3.4 (3, 3.9)	0.05 (0.02, 0.14)
(2011) <sup>55</sup>												
Data from:		On presentation	Δ 17%		65	202	43	520	60 (51, 69)	72 (69, 75)	2.1 (1.8, 2.6)	0.55 (0.44, 0.7)
Irfan (2013) <sup>64</sup>	Beckman	and at 1 hour	Δ 27%		68	245	40	477	63 (53 <i>,</i> 71)	66 (63 <i>,</i> 69)	1.9 (1.6, 2.2)	0.56 (0.44, 0.72)
Data from:	Roche	On presentation	Δ 30%		43	84	24	439	64 (52 <i>,</i> 74)	84 (80 <i>,</i> 87)	4 (3, 5.2)	0.43 (0.31, 0.59)
Reichlin (2011) <sup>51</sup>		and at 2 hours										
Data from:	Abbott		26.2 on	MACE	129	62	27	691	82 (76 <i>,</i> 88)	92 (90 <i>,</i> 93)	10 (7.8,	0.19 (0.14, 0.27)
Cullen (2013) <sup>62</sup>			admission and								12.8)	
			at 2 hours									
Keller (2011) <sup>47</sup>	Abbott	On presentation	3.4	AMI	282	633	0	345	100 (98, 100)	35 (32, 38)	1.5 (1.5, 1.6)	0.01 (0, 0.08)
			30	AMI	232	77	50	901	82 (77 <i>,</i> 86)	92 (90 <i>,</i> 94)	10.4 (8.3,	0.19 (0.15, 0.25)
											12.9)	
		3 hours after	3.4	AMI	282	959	0	19	100 (98, 100)	2 (1, 3)	1 (1, 1)	0.09 (0.01, 1.46)
		presentation	30	AMI	277	94	5	884	98 (96 <i>,</i> 99)	90 (88, 92)	10.2 (8.4,	0.02 (0.01, 0.05)
											12.3)	
		On presentation	Δ 20%	AMI	218	723	64	255	77 (72, 82)	26 (23, 29)	1 (1, 1.1)	0.87 (0.69, 1.11)
		and at 3 hours	3.4 on	AMI	254	454	54	498	82 (78 <i>,</i> 86)	52 (49 <i>,</i> 55)	1.7 (1.6, 1.9)	0.34 (0.26, 0.43)
			admission and									
			Δ 20%									
			30 after 3 hrs	AMI	187	34	110	929	63 (57 <i>,</i> 68)	96 (95 <i>,</i> 97)	17.6 (12.5,	0.38 (0.33, 0.45)
			and Δ 20%								24.7)	
			30 after 3 hrs	AMI	52	26	4	869	92 (82 <i>,</i> 97)	97 (96 <i>,</i> 98)	31.1 (21.2,	0.08 (0.03, 0.2)
			and ∆ 20%, in								45.7)	
			patients									
			<30ng/L on									
E 4			admission									
Kurz (2011) <sup>34</sup>	Roche	On presentation	9.5	NSTEMI	38	11	8	37	82 (69, 90)	77 (63, 86)	3.5 (2.1, 5.9)	0.24 (0.13, 0.44)
			14	NSTEMI	16	7	10	24	61 (42, 77)	77 (60, 88)	2.6 (1.3, 5.2)	0.51 (0.3, 0.85)
		within 3 hours of	14	NSTEMI	26	7	0	23	98 (84, 100)	76 (58, 87)	4.1 (2.2, 7.6)	0.02 (0, 0.38)
		presentation										
		On presentation	14 and ∆ 20%	NSTEMI	11	27	15	3	43 (26, 61)	11 (4, 27)	0.5 (0.3, 0.8)	5.08 (1.8, 14.37)
		and within 3										
7/		hours										
Lippi (2012)' 🖥	Beckman	On presentation	18	AMI	9	17	0	31	95 (66, 99)	64 (50, 76)	2.7 (1.8, 4)	0.08 (0.01, 1.17)

Study Details	Troponin	Timing	Threshold	Target	TP	FP	FN	TN	Sens	Spec	LR+ (95% CI)	LR- (95% CI)
	Assay		(ng/L)	Condition					(95% CI)	(95% CI)		
Melki (2011)⁵ <sup>0</sup>	Roche	On presentation	14	NSTEMI	112	21	2	98	98 (93 <i>,</i> 99)	82 (74, 88)	5.5 (3.7, 8)	0.03 (0.01, 0.09)
		2 hours after	14	NSTEMI	114	25	0	94	100 (96, 100)	79 (71, 85)	4.7 (3.3, 6.6)	0.01 (0, 0.09)
		presentation										
Parsonage	Abbott	On presentation	26.2	AMI	45	34	6	652	88 (76 <i>,</i> 94)	95 (93 <i>,</i> 96)	17.4 (12.4,	0.13 (0.06, 0.27)
(2013) <sup>57</sup>											24.5)	
		On presentation	26.2 peak	AMI	47	48	4	638	91 (81, 96)	93 (91, 95)	12.9 (9.7,	0.09 (0.04, 0.23)
		and at 2 hours									17.2)	
	Roche	On presentation	14	AMI	44	75	7	611	86 (74, 93)	89 (86, 91)	7.8 (6.1, 9.9)	0.16 (0.08, 0.31)
		On presentation	14 peak	AMI	48	82	3	604	93 (83 <i>,</i> 98)	88 (85, 90)	7.8 (6.3, 9.6)	0.08 (0.03, 0.21)
		and at 2 hours										
Saenger( 2010) <sup>71</sup>	Roche	On presentation	14	AMI	92	38	6	152	93 (87 <i>,</i> 97)	80 (74, 85)	4.6 (3.5, 6.2)	0.08 (0.04, 0.17)
		On presentation	Δ8	AMI	94	9	4	181	95 (89 <i>,</i> 98)	95 (91, 97)	19.2 (10.3,	0.05 (0.02, 0.12)
		and at 3 hours									35.7)	
Sanchis (2012) <sup>41</sup>	Roche	On presentation	3	MACE	53	207	9	177	85 (74, 92)	46 (41, 51)	1.6 (1.4, 1.8)	0.33 (0.18, 0.59)
		On presentation	3	MACE	57	234	5	150	91 (82 <i>,</i> 96)	39 (34, 44)	1.5 (1.3, 1.7)	0.22 (0.1, 0.5)
		and 6-8 hours	14	MACE	21	42	41	342	34 (24, 46)	89 (85, 92)	3.1 (2, 4.8)	0.74 (0.62, 0.89)
Santaló (2013) <sup>39</sup>	Roche	On presentation	14	NSTEMI	71	80	8	199	89 (81, 94)	71 (66, 76)	3.1 (2.5, 3.8)	0.15 (0.08, 0.28)
		On presentation	Δ 20%	NSTEMI	79	94	0	185	99 (94 <i>,</i> 100)	66 (61, 72)	2.9 (2.5, 3.5)	0.01 (0, 0.15)
		and at 2, 4 and 6-										
		8 hours or until										
		discharge										
Sebbane (2013) <sup>63</sup>	Roche	On presentation,	14	NSTEMI	19	25	6	142	75 (56 <i>,</i> 88)	85 (79 <i>,</i> 89)	4.9 (3.2, 7.5)	0.29 (0.15, 0.58)
		or sample taken	18	NSTEMI	19	17	6	150	75 (56 <i>,</i> 88)	90 (84, 93)	7.2 (4.4,	0.28 (0.14, 0.54)
		during pre-									11.8)	
		hospital										
		management										

# **APPENDIX 3: QUADAS-2 ASSESSMENTS**

#### Study: Aldous (2011)<sup>53</sup> DOMAIN 1: PATIENT SELECTION A. RISK OF BIAS

Consecutive adults presenting to the emergency department with chest	pain were eligible for inclusion.	
Was a consecutive or random sample of patients enrolled?	Y	'es
Was a case-control design avoided?	Y	'es
Did the study avoid inappropriate exclusions?	Y	'es
Could the selection of patients have introduced bias?	RISK: Low	
B. APPLICABILITY		
Unselected chest pain population AMI diagnoses may have included bot	th NSTEMI and STEMI	
Do the included patients match the question?	Concerns: High	
DOMAIN 2: INDEX TEST(S)		
A. RISK OF BIAS		
Dealer Flager to TaT an adapticities and after Char. Data are stad for ad		
Roche Elecsys hs-ini on admission and after 6 hrs. Data reported for ad	Imission, for four thresholds	
No details of interpretation reported. One threshold was derived from F	CC analysis; primary analysis based on	
99th centile	like of the state	
were the index test results interpreted without knowledge of the rest	lits of Uncle	ear
the reference standard?		
If a threshold was used, was it pre-specified?	PICK Law	es
Could the conduct or interpretation of the index test have	RISK: LOW	
D. APPLICADILITY	Concernes Louis	
differ from the review question?	Concerns: Low	
differ from the review question?		
DOMAIN 3: REFERENCE STANDARD		
A. RISK OF BIAS		
Reference standard diagnosis of AMI based on joint European Cardiolog	y Society an American College of Cardiolo	gy
criteria and included serial conventional cTnI (10-12 hour time point not	t specified)	07
Determination of diagnosis was made blind to hs-TnT results	, ,	
Is the reference standard likely to correctly classify the target conditio	n? Y	'es
Were the reference standard results interpreted without knowledge of	of the Y	'es
results of the index test?		
Could the reference standard, its conduct, or its interpretation have	RISK: Low	
introduced bias?		
B. APPLICABILITY		
Is there concern that the target condition as defined by the	Concerns: High	
reference standard does not match the review question?	-	

#### DOMAIN 4: FLOW AND TIMING A. RISK OF BIAS

Participants for whom stored samples were not available at both time points were excluded.	
Did all patients receive a reference standard?	Yes
Did patients receive the same reference standard?	Yes
Were all patients included in the analysis?	No
Could the patient flow have introduced bias?	RISK: High

# Study: Aldous (2012)<sup>45</sup> DOMAIN 1: PATIENT SELECTION A. RISK OF BIAS

Patients presenting to the emergency department between 05:30 h and	20:00 h, and with chest pain	
Was a consecutive or random sample of patients enrolled?		No
Was a case-control design avoided?		Yes
Did the study avoid inappropriate exclusions?		Yes
Could the selection of patients have introduced bias?	RISK: High	
B. APPLICABILITY		
Patients with ST-segment elevation excluded		
Do the included patients match the question?	Concerns: Low	
DOMAIN 2: INDEX TEST(S)		
A. RISK OF BIAS		
Roche Elecsys hsTnT		
Data reported for multiple thresholds based on pre-determined propert	ties of the assav	
Frozen samples used, unclear whether interpretation of index test was h	blind to reference standard	
Were the index test results interpreted without knowledge of the resu	ults of	Yes
the reference standard?		
If a threshold was used, was it pre-specified?		Yes
Could the conduct or interpretation of the index test have	RISK: Low	
introduced bias?		
B. APPLICABILITY		
Are there concerns that the index test, its conduct, or interpretation	Concerns: Low	
differ from the review question?		
DOMAIN 3: REFERENCE STANDARD		
A. RISK OF BIAS		
Reference standard final diagnosis of AML based on ACC criteria and inc	luding the results of serial conventiona	1
cTnl (10-12 hour time point not specified) but blinded to hs-TnT results		•
Is the reference standard likely to correctly classify the target condition	n?	Yes
Were the reference standard results interpreted without knowledge o	of the	Yes
results of the index test?		
Could the reference standard, its conduct, or its interpretation have	RISK: Low	
introduced bias?		
B. APPLICABILITY		
Is there concern that the target condition as defined by the	Concerns: High	
reference standard does not match the review question?	C C	
DOMAIN 4: FLOW AND TIMING		
A. RISK OF BIAS		

All participants appear to have been included in the analyses	
Did all patients receive a reference standard?	Yes
Did patients receive the same reference standard?	Yes
Were all patients included in the analysis?	Yes
Could the patient flow have introduced bias?	<b>RISK: Low</b>

#### Study: Body (2011)<sup>67</sup> DOMAIN 1: PATIENT SELECTION A. RISK OF BIAS

Prospective enrolment of patients; unclear if consecutive	
Was a consecutive or random sample of patients enrolled?	Unclear
Was a case-control design avoided?	Yes
Did the study avoid inappropriate exclusions?	Yes
Could the selection of patients have introduced bias?	RISK: Unclear
B. APPLICABILITY	
Mixed chest pain	
Do the included patients match the question?	Concerns: High
DOMAIN 2: INDEX TEST(S)	
A. RISK OF BIAS	
Roche Elecsys HsTnT. Threshold 99th percentile cut point and limit of a	detection. Blinding not reported: objective
test interpreted prior to reference standard so unlikely to have been in	fluenced by knowledge of reference
standard.	
Were the index test results interpreted without knowledge of the res	ults of Yes
the reference standard?	
If a threshold was used, was it pre-specified?	Yes
Could the conduct or interpretation of the index test have	RISK: Low
introduced bias?	
B. APPLICABILITY	
Are there concerns that the index test, its conduct, or interpretation	Concerns: Low
differ from the review question?	
DOMAIN 3: REFERENCE STANDARD	
A. RISK OF BIAS	
Thorgeson criteria; time point not specified. Clinicians were blinded to	Hs-Tn.
Is the reference standard likely to correctly classify the target condition	on? Yes
Were the reference standard results interpreted without knowledge	of the Yes
results of the index test?	
Could the reference standard, its conduct, or its interpretation have	RISK: Low

B. APPLICABILITYConcerns: HighIs there concern that the target condition as defined by the<br/>reference standard does not match the review question?Concerns: High

#### DOMAIN 4: FLOW AND TIMING A. RISK OF BIAS

introduced bias?

301 patients were excluded prior to enrolment; all patients enrolled included in 2x2 table.	
Did all patients receive a reference standard?	Yes
Did patients receive the same reference standard?	Yes
Were all patients included in the analysis?	Yes
Could the patient flow have introduced bias?	<b>RISK: Low</b>

# Study: Christ (2010)<sup>56</sup> DOMAIN 1: PATIENT SELECTION A. RISK OF BIAS

Retrospective analysis of consecutive patients presenting to ED with chest pain	
Was a consecutive or random sample of patients enrolled?	Yes
Was a case-control design avoided?	Yes
Did the study avoid inappropriate exclusions?	Yes
Could the selection of patients have introduced bias? RISK: Low	
B. APPLICABILITY	
Patients with general chest pain symptoms, includes participants with a final diagnosis of STEMI	
Do the included patients match the question? Concerns: High	
DOMAIN 2: INDEX TEST(S)	
A. RISK OF BIAS	
Decks Floorus HoTaT. Threshold 00th percentile out point. Dlinding not reported, retraspective applysic	and co
Roche Elecsys Hstint. Infestiol 99th percentile cut point. Blinding not reported; retrospective analysis	and so
been influenced by knowledge of disease state	/ to have
Were the index tect results interpreted without knowledge of the results of	Uncloar
the reference standard?	Unclear
If a threshold was used, was it pre-specified?	Voc
Could the conduct or interpretation of the index test have <b>PISK:</b> Low	165
introduced bias?	
Are there concerns that the index test its conduct or interpretation Concerns: Low	
differ from the review question?	
A RISK OF BIAS	
Joint European Cardiology Society an American College of Cardiology criteria; time point not specified.	Uncloar
whether clinicians were blinded to Hs-Tn A second consensus diagnosis incorporating Hs-Tn was also r	Unciear
whether enheuris were sinded to its in. A second consensus didghosis meerportering its in was user	nade and so
clinicians may have been aware of the result for the first consensus diagnosis based only on standard tr	nade and so oponin.
clinicians may have been aware of the result for the first consensus diagnosis based only on standard tr Is the reference standard likely to correctly classify the target condition?	nade and so oponin. Yes
clinicians may have been aware of the result for the first consensus diagnosis based only on standard tr Is the reference standard likely to correctly classify the target condition? Were the reference standard results interpreted without knowledge of the	nade and so oponin. Yes Unclear
clinicians may have been aware of the result for the first consensus diagnosis based only on standard tr Is the reference standard likely to correctly classify the target condition? Were the reference standard results interpreted without knowledge of the results of the index test?	nade and so oponin. Yes Unclear
clinicians may have been aware of the result for the first consensus diagnosis based only on standard tr Is the reference standard likely to correctly classify the target condition? Were the reference standard results interpreted without knowledge of the results of the index test? Could the reference standard, its conduct, or its interpretation have RISK: Unclear	nade and so oponin. Yes Unclear
<ul> <li>clinicians may have been aware of the result for the first consensus diagnosis based only on standard tr Is the reference standard likely to correctly classify the target condition?</li> <li>Were the reference standard results interpreted without knowledge of the results of the index test?</li> <li>Could the reference standard, its conduct, or its interpretation have RISK: Unclear introduced bias?</li> </ul>	nade and so oponin. Yes Unclear
<ul> <li>clinicians may have been aware of the result for the first consensus diagnosis based only on standard tr Is the reference standard likely to correctly classify the target condition? Were the reference standard results interpreted without knowledge of the results of the index test?</li> <li>Could the reference standard, its conduct, or its interpretation have introduced bias?</li> <li>B. APPLICABILITY</li> </ul>	nade and so oponin. Yes Unclear
<ul> <li>clinicians may have been aware of the result for the first consensus diagnosis based only on standard tr Is the reference standard likely to correctly classify the target condition?</li> <li>Were the reference standard results interpreted without knowledge of the results of the index test?</li> <li>Could the reference standard, its conduct, or its interpretation have introduced bias?</li> <li>B. APPLICABILITY</li> <li>Is there concern that the target condition as defined by the Concerns: High</li> </ul>	nade and so oponin. Yes Unclear
<ul> <li>clinicians may have been aware of the result for the first consensus diagnosis based only on standard tr Is the reference standard likely to correctly classify the target condition?</li> <li>Were the reference standard results interpreted without knowledge of the results of the index test?</li> <li>Could the reference standard, its conduct, or its interpretation have introduced bias?</li> <li>B. APPLICABILITY</li> <li>Is there concern that the target condition as defined by the reference standard does not match the review question?</li> </ul>	nade and so oponin. Yes Unclear
<ul> <li>clinicians may have been aware of the result for the first consensus diagnosis based only on standard tr Is the reference standard likely to correctly classify the target condition?</li> <li>Were the reference standard results interpreted without knowledge of the results of the index test?</li> <li>Could the reference standard, its conduct, or its interpretation have introduced bias?</li> <li>B. APPLICABILITY</li> <li>Is there concern that the target condition as defined by the reference standard does not match the review question?</li> </ul>	nade and so oponin. Yes Unclear
<ul> <li>clinicians may have been aware of the result for the first consensus diagnosis based only on standard tr clinicians may have been aware of the result for the first consensus diagnosis based only on standard tr Is the reference standard likely to correctly classify the target condition?</li> <li>Were the reference standard results interpreted without knowledge of the results of the index test?</li> <li>Could the reference standard, its conduct, or its interpretation have introduced bias?</li> <li>B. APPLICABILITY</li> <li>Is there concern that the target condition as defined by the reference standard does not match the review question?</li> <li>DOMAIN 4: FLOW AND TIMING</li> </ul>	nade and so oponin. Yes Unclear
<ul> <li>clinicians may have been aware of the result for the first consensus diagnosis based only on standard tr <li>clinicians may have been aware of the result for the first consensus diagnosis based only on standard tr <li>Is the reference standard likely to correctly classify the target condition? <ul> <li>Were the reference standard results interpreted without knowledge of the results of the index test?</li> </ul> </li> <li>Could the reference standard, its conduct, or its interpretation have <ul> <li>RISK: Unclear</li> <li>introduced bias?</li> <li>B. APPLICABILITY</li> <li>Is there concern that the target condition as defined by the reference standard does not match the review question?</li> </ul> </li> <li>DOMAIN 4: FLOW AND TIMING <ul> <li>A. RISK OF BIAS</li> </ul> </li> </li></li></ul>	nade and so oponin. Yes Unclear
<ul> <li>clinicians may have been aware of the result for the first consensus diagnosis based only on standard tr Is the reference standard likely to correctly classify the target condition? Were the reference standard results interpreted without knowledge of the results of the index test?</li> <li>Could the reference standard, its conduct, or its interpretation have introduced bias?</li> <li>B. APPLICABILITY</li> <li>Is there concern that the target condition as defined by the reference standard does not match the review question?</li> <li>DOMAIN 4: FLOW AND TIMING A. RISK OF BIAS</li> <li>No dropouts reported, all included patients accounted for in flow diagram and numbers suggest that tree</li> </ul>	nade and so oponin. Yes Unclear
Could the reference standard results interpreted without knowledge of the results of the index test? Could the reference standard, its conduct, or its interpretation have RISK: Unclear introduced bias? B. APPLICABILITY Is there concern that the target condition as defined by the reference standard does not match the review question? DOMAIN 4: FLOW AND TIMING A. RISK OF BIAS No dropouts reported, all included patients accounted for in flow diagram and numbers suggest that troresults were available for all.	onclear nade and so oponin. Yes Unclear
Could the reference standard results interpreted without knowledge of the results of the index test? Could the reference standard, its conduct, or its interpretation have RISK: Unclear introduced bias? B. APPLICABILITY Is there concern that the target condition as defined by the concerns: High reference standard does not match the review question? DOMAIN 4: FLOW AND TIMING A. RISK OF BIAS No dropouts reported, all included patients accounted for in flow diagram and numbers suggest that troresults were available for all. Did all patients receive a reference standard?	onclear nade and so oponin. Yes Unclear
clinicians may have been aware of the result for the first consensus diagnosis based only on standard tr ls the reference standard likely to correctly classify the target condition? Were the reference standard results interpreted without knowledge of the results of the index test? Could the reference standard, its conduct, or its interpretation have RISK: Unclear introduced bias? B. APPLICABILITY Is there concern that the target condition as defined by the Concerns: High reference standard does not match the review question? DOMAIN 4: FLOW AND TIMING A. RISK OF BIAS No dropouts reported, all included patients accounted for in flow diagram and numbers suggest that tro results were available for all. Did all patients receive a reference standard?	onclear nade and so oponin. Yes Unclear oponin Yes Yes

Could the patient flow have introduced bias?

**RISK: Low** 

# Study: Collinson (2013)<sup>65</sup> DOMAIN 1: PATIENT SELECTION A. RISK OF BIAS

Participants with chest pain and suspected AMI; Study uses subgroup of	f one arm of an RCT. Patients at high risk of
INSTEIVIT excluded	
Was a consecutive or random sample of patients enrolled?	Yes
Was a case-control design avoided?	Yes
Did the study avoid inappropriate exclusions?	Yes
Could the selection of patients have introduced bias?	RISK: Low
B. APPLICABILITY	
Chest pain patients excluding those with diagnostic ECG changes	
Do the included patients match the question?	Concerns: High
DOMAIN 2: INDEX TEST(S)	
A. RISK OF BIAS	
Roche Elecsys hs-TnT on admission and at 90 minutes	
Reference standard (final diagnosis) determined after hs-TnT	
Threshold based on assay characteristics including 99th centile	
Were the index test results interpreted without knowledge of the resu	ults of Yes
the reference standard?	
If a threshold was used, was it pre-specified?	Yes
Could the conduct or interpretation of the index test have	RISK: Low
introduced bias?	
B. APPLICABILITY	
Are there concerns that the index test, its conduct, or interpretation	Concerns: Low

# differ from the review question?

# **DOMAIN 3: REFERENCE STANDARD**

# A. RISK OF BIAS

Reference standard diagnosis of AMI based on joint European Cardiology Society an American College of Cardiology				
criteria and included serial conventional cTnT or cTnI (10-12 hour time point specified)				
Determination of diagnosis was made blind to hs-TnT results				
Is the reference standard likely to correctly classify the target condition?				
Were the reference standard results interpreted without knowledge	of the Yes			
results of the index test?				
Could the reference standard, its conduct, or its interpretation have	RISK: Low			
introduced bias?				
B. APPLICABILITY				
Is there concern that the target condition as defined by the	Concerns: Low			
reference standard does not match the review question?				
DOMAIN 4: FLOW AND TIMING				
A. RISK OF BIAS				
1125 enrolled, 25 no samples collected, 250 samples taken but study sa	amples not collected.			
Did all patients receive a reference standard?	Yes			

Could the patient flow have introduced bias?	RISK: High
Were all patients included in the analysis?	No
Did patients receive the same reference standard?	Yes
Did all patients receive a reference standard?	Yes

# Study: Cullen (2013)<sup>62</sup> DOMAIN 1: PATIENT SELECTION A. RISK OF BIAS

Consecutively recruited adults presenting to the emergency departmer	nt with cardiac symptoms
Was a consecutive or random sample of patients enrolled?	Ye
Was a case-control design avoided?	Ye
Did the study avoid inappropriate exclusions?	Ye
Could the selection of patients have introduced bias?	RISK: Low
B. APPLICABILITY	
Unselected chest pain population AMI diagnoses may have included bo	th NSTEMI and STEMI
Do the included patients match the question?	Concerns: High
DOMAIN 2: INDEX TEST(S)	
A. RISK OF BIAS	
Abbott ARCHITECT hs-STAT Thi: Threshold was 99th centile	
Frozen samples were used, but laboratory technicians were blinded to	natient data
Were the index test results interpreted without knowledge of the res	ults of Ve
the reference standard?	
If a threshold was used, was it pre-specified?	Ye
Could the conduct or interpretation of the index test have	RISK: Low
introduced bias?	
B. APPLICABILITY	
Are there concerns that the index test, its conduct, or interpretation	Concerns: Low
differ from the review question?	
·	
DOMAIN 3: REFERENCE STANDARD	
A. RISK OF BIAS	
30 day MACE, adjudicated blind to index tests, but with access to clinicated	al records, ECG and conventional troponin
results	
Is the reference standard likely to correctly classify the target condition	on? Ye
Were the reference standard results interpreted without knowledge	of the Ye
results of the index test?	<b>-</b> 1011
Could the reference standard, its conduct, or its interpretation have	RISK: LOW
Introduced bias?	
Is there concern that the target condition as defined by the	Concerns: Low
reference standard does not match the review question?	
DOMAIN 4: FLOW AND TIMING	

#### A. RISK OF BIAS

No patients were lost to 30 day follow-up. Procedure for adjudication of 30 day MACE was the same in all	cases,
but investigations undergone by individual patients varied	
Did all patients receive a reference standard?	Yes
Did patients receive the same reference standard?	No
Were all patients included in the analysis?	Yes
Could the patient flow have introduced bias?	RISK: Low

## Study: Eggers (2012)<sup>43</sup> DOMAIN 1: PATIENT SELECTION A. RISK OF BIAS

Unclear whether consecutive or random patients were enrolled.		
Was a consecutive or random sample of patients enrolled?	Une	clear
Was a case-control design avoided?		Yes
Did the study avoid inappropriate exclusions?		Yes
Could the selection of patients have introduced bias?	RISK: Unclear	
B. APPLICABILITY		
Non-STEMI patients with chest pain presenting to coronoary care/ches	t pain unit	
Do the included patients match the question?	Concerns: High	
DOMAIN 2: INDEX TEST(S)		
A. RISK OF BIAS		
· · · · · · · · · · · · · · · · · · ·		
Roche Elecsys HsTnT. Threshold 99th percentile cut point and 95% spe	cificity value. Blinding not reported;	
objective test interpreted prior to reference standard so unlikely to have	e been influenced by knowledge of	
reference standard.		
Were the index test results interpreted without knowledge of the res	ults of Une	clear
the reference standard?		
If a threshold was used, was it pre-specified?		Yes
Could the conduct or interpretation of the index test have	RISK: Low	
introduced bias?		
B. APPLICABILITY		
Are there concerns that the index test, its conduct, or interpretation	Concerns: Low	
differ from the review question?		
DOMAIN 3: REFERENCE STANDARD		
A. RISK OF BIAS		
	·····	
Joint European Cardiology Society an American College of Cardiology ci	iteria; time point not specified. Unclear	
whether clinicians were blinded to Hs-In. A second consensus diagnos	s	
Is the reference standard likely to correctly classify the target condition	on?	Yes
Were the reference standard results interpreted without knowledge	of the Uni	clear
results of the index test?		
Could the reference standard, its conduct, or its interpretation have	RISK: Unclear	
introduced bias?		
B. APPLICABILITY		
	Concerns: High	
is there concern that the target condition as defined by the		
Is there concern that the target condition as defined by the reference standard does not match the review question?		
Is there concern that the target condition as defined by the reference standard does not match the review question?		
Is there concern that the target condition as defined by the reference standard does not match the review question? DOMAIN 4: FLOW AND TIMING		
Is there concern that the target condition as defined by the reference standard does not match the review question? DOMAIN 4: FLOW AND TIMING A. RISK OF BIAS		
Is there concern that the target condition as defined by the reference standard does not match the review question? DOMAIN 4: FLOW AND TIMING A. RISK OF BIAS Only 360 patients out of 495 who fulfilled inclusion criteria had all bioc	nemical tests performed and were includ	led
Is there concern that the target condition as defined by the reference standard does not match the review question? DOMAIN 4: FLOW AND TIMING A. RISK OF BIAS Only 360 patients out of 495 who fulfilled inclusion criteria had all bioch in the analysis: reasons for not performing tests were not reported	nemical tests performed and were includ	led

Could the patient flow have introduced bias?	RISK: High
Were all patients included in the analysis?	No
Did patients receive the same reference standard?	Yes
Did all patients receive a reference standard?	Yes

#### Study: Freund (2011)<sup>48</sup> DOMAIN 1: PATIENT SELECTION A. RISK OF BIAS

Consecutive adults presenting to the emergency department w	ith chest pain (onset or peak within pre-	vious 6 hrs).
Patients with acute kidney failure requiring dialysis were exclude	led	,
Was a consecutive or random sample of patients enrolled?		Yes
Was a case-control design avoided?		Yes
Did the study avoid inappropriate exclusions?		Yes
Could the selection of patients have introduced bias?	RISK: Low	
B. APPLICABILITY		
Unselected emergency department chest pain population, inclu	ides participants with a final diagnosis o	f STEMI;
data also presented for subgroups with low-moderate and with	high pre-test probability	
Do the included patients match the question?	Concerns: High	
DOMAIN 2: INDEX TEST(S) A. RISK OF BIAS		
Roche Elecsys hs-TnT on admission and at 3-9 hours if available	Reference standard (final diagnosis) ac	liudicated by

Roche Elecsys hs-TnT on admission and at 3-9 hours if available. Reference standard (final diagnosis) adjudicated by		
two independent physicians after acute episode. Threshold was 99th ce	ntile	
Were the index test results interpreted without knowledge of the resu	ults of Yes	
the reference standard?		
If a threshold was used, was it pre-specified?	Yes	
Could the conduct or interpretation of the index test have	RISK: Low	
introduced bias?		
B. APPLICABILITY		
Are there concerns that the index test, its conduct, or interpretation	Concerns: Low	
differ from the review question?		

# DOMAIN 3: REFERENCE STANDARD

Α.	RISK	OF	BIAS
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Reference standard final diagnosis, based on joint European Cardiology Society an American College of Cardiology	
criteria and included conventional cTnI on admission and at 3-9 hours i	f needed (10-12 hour time point not
specified). Clinicians adjudicating final diagnosis were blind to hs-TnT re	esults
Is the reference standard likely to correctly classify the target condition	on? Yes
Were the reference standard results interpreted without knowledge	of the Yes
results of the index test?	
Could the reference standard, its conduct, or its interpretation have	RISK: Low
introduced bias?	
B. APPLICABILITY	
Is there concern that the target condition as defined by the	Concerns: High
reference standard does not match the review question?	
DOMAIN 4: FLOW AND TIMING	
A. RISK OF BIAS	
All participants appear to have been included in the analyses	
Did all patients receive a reference standard?	Yes
Did patients receive the same reference standard?	Yes
Were all patients included in the analysis?	Yes

Could the patient flow have introduced bias?

**RISK: Low** 

#### Study: Hoeller (2013)<sup>75</sup> DOMAIN 1: PATIENT SELECTION A. RISK OF BIAS

Patients presenting to the ED with symptoms suggestive of AMI. Consec	cutive patients with hs-TnT measuremen	its
available were included.		
Was a consecutive or random sample of patients enrolled?		Yes
Was a case-control design avoided?		Yes
Did the study avoid inappropriate exclusions?		Yes
Could the selection of patients have introduced bias?	RISK: Low	
B. APPLICABILITY		
Unselected chest pain population AMI diagnoses may have included bot	th NSTEMI and STEMI	
Do the included patients match the question?	Concerns: High	
DOMAIN 2: INDEX TEST(S)		
A. RISK OF BIAS		
Roche Hs-TnT Beckman Coulter Hs-AccuTnl and Abbott ARCHITECT HsT	nl on admission	
Reference standard probably made later than admission 99th Centiles	for assays used as diagnostic thresholds	
(some nublications also reported data for ROC derived thresholds)		
Were the index test results interpreted without knowledge of the resu	ults of	Ves
the reference standard?		105
If a threshold was used was it pre-specified?		Yes
Could the conduct or interpretation of the index test have	BISK: Low	105
introduced bias?		
B. APPLICABILITY		
Are there concerns that the index test, its conduct, or interpretation	Concerns: Low	
differ from the review question?		
DOMAIN 3: REFERENCE STANDARD		
A. RISK OF BIAS		
Reference standard final diagnosis of AMI, European Cardiology Society	criteria and included cTn assays (0 and 6	6
hours). Unclear whether those adjudicating final diagnosis were blind to	hs-Tnl/hsTnT results in all cases, some	
publications reported blinding		
Is the reference standard likely to correctly classify the target conditio	n?	Yes
Were the reference standard results interpreted without knowledge of	of the Yes	s/No
results of the index test?		
Could the reference standard, its conduct, or its interpretation have	RISK: Low	
Could the reference standard, its conduct, or its interpretation have introduced bias?	RISK: Low	
Could the reference standard, its conduct, or its interpretation have introduced bias? B. APPLICABILITY	RISK: Low	
Could the reference standard, its conduct, or its interpretation have introduced bias? B. APPLICABILITY Is there concern that the target condition as defined by the	RISK: Low Concerns: Low	
Could the reference standard, its conduct, or its interpretation have introduced bias? B. APPLICABILITY Is there concern that the target condition as defined by the reference standard does not match the review question?	RISK: Low Concerns: Low	
Could the reference standard, its conduct, or its interpretation have introduced bias? B. APPLICABILITY Is there concern that the target condition as defined by the reference standard does not match the review question?	RISK: Low Concerns: Low	
Could the reference standard, its conduct, or its interpretation have introduced bias? B. APPLICABILITY Is there concern that the target condition as defined by the reference standard does not match the review question? DOMAIN 4: FLOW AND TIMING	RISK: Low Concerns: Low	
Could the reference standard, its conduct, or its interpretation have introduced bias? B. APPLICABILITY Is there concern that the target condition as defined by the reference standard does not match the review question? DOMAIN 4: FLOW AND TIMING A. RISK OF BIAS	RISK: Low Concerns: Low	
Could the reference standard, its conduct, or its interpretation have introduced bias? B. APPLICABILITY Is there concern that the target condition as defined by the reference standard does not match the review question? DOMAIN 4: FLOW AND TIMING A. RISK OF BIAS	RISK: Low Concerns: Low	e

hsTnI (Beckman) analysis, and 1567 were included in the hsTnI (Abbott) analysis Most exclusions were because hsTn measurements were not available

Could the patient flow have introduced bias?	RISK: High
Were all patients included in the analysis?	No
Did patients receive the same reference standard?	Yes
Did all patients receive a reference standard?	Yes

# Study: Keller (2011)<sup>47</sup> DOMAIN 1: PATIENT SELECTION A. RISK OF BIAS

Consecutive patients presenting to chest pain units		
Was a consecutive or random sample of patients enrolled?		Yes
Was a case-control design avoided?		Yes
Did the study avoid inappropriate exclusions?		Yes
Could the selection of patients have introduced bias?	RISK: Low	
B. APPLICABILITY		
General chest pain populations, some participants had a final diagnos	is of STEMI	
Do the included patients match the question?	Concerns: High	
DOMAIN 2: INDEX TEST(S)		
A. RISK OF BIAS		
Abbott Architect STAT hs-Tnl, on admission and at 3 hrs. Reference st	andard (final diagnosis) was adjudica	ted after
hs-Tnl testing. Thresholds based on test properties, appeared to be p	re-specified	
Were the index test results interpreted without knowledge of the re	esults of	Yes
the reference standard?		
If a threshold was used, was it pre-specified?		Yes
Could the conduct or interpretation of the index test have	RISK: Low	
introduced bias?		
B. APPLICABILITY		
Are there concerns that the index test, its conduct, or interpretation	Concerns: Low	
differ from the review question?		
DOMAIN 3: REFERENCE STANDARD		
Reference standard diagnosis of AMI based on joint European Cardiol	ogy Society an American College of C	ardiology
criteria and included serial conventional cTnT (10-12 hour time point	not specified)	
Determination of diagnosis was made blind to hs-TnT results		
Is the reference standard likely to correctly classify the target condition	tion?	Yes
Were the reference standard results interpreted without knowledge	e of the	Yes
results of the index test?		
Could the reference standard, its conduct, or its interpretation have	RISK: Low	
introduced bias?		
B. APPLICABILITY		
Is there concern that the target condition as defined by the	Concerns: High	
reference standard does not match the review question?		
DOMAIN 4: FLOW AND TIMING		
A. KISK OF BIAS		
None of the analyses included all study participants (558 or 867 partic	cipants missing)	
Did all natients receive a reference standard?		Ves

Could the patient flow have introduced bias?	RISK: High
Were all patients included in the analysis?	No
Did patients receive the same reference standard?	Yes
Did all patients receive a reference standard?	Yes

Yes

Unclear

#### Study: Kurz (2011)<sup>54</sup> DOMAIN 1: PATIENT SELECTION A. RISK OF BIAS

Consecutive patients admitted to a chest pain unit. 206 Patients not included due to 'technical reasons' (not fully defined, e.g. venipuncture not possible) Was a consecutive or random sample of patients enrolled? Yes

Was a consecutive or random sample of patients enrolled? Was a case-control design avoided?

Did the study avoid inappropriate exclusions?

Could the selection of patients have introduced bias?

#### **B. APPLICABILITY**

Appears to be an unselected chest pain population, STEMI excluded. Second publication<sup>112</sup> is for a retrospectively selected subgroup of participants with a diagnosis of NSTEMI or unstable angina. Patients were admitted to chest pain units.

Do the included patients match the question?

**Concerns: High** 

**RISK: Unclear** 

#### DOMAIN 2: INDEX TEST(S) A. RISK OF BIAS

Roche Elecsys hs-TnT, data reported for admission, 3 hr and 6 hr sample Reference standard troponin testing occurred after hs-TnT. Threshold w	s (6 hrs data not extracted) as pre-specified for data extracted from <sup>112</sup> ,
but not from <sup>54</sup> (low risk of bias for $112$ data)	
Were the index test results interpreted without knowledge of the resu	Its of Yes
the reference standard?	
If a threshold was used, was it pre-specified?	Yes
Could the conduct or interpretation of the index test have	RISK: Low
introduced bias?	
B. APPLICABILITY	
Are there concerns that the index test, its conduct, or interpretation	Concerns: Low
differ from the review question?	

# DOMAIN 3: REFERENCE STANDARD

#### A. RISK OF BIAS

Reference standard diagnosis of AMI based on joint European Cardiology Society and American College of Cardiology criteria and included serial conventional cTnT (10-12 hour time point not specified) Unclear whether determination of diagnosis was made blind to hs-TnT results	
Is the reference standard likely to correctly classify the target condition?	Yes
Were the reference standard results interpreted without knowledge of the results of the index test?	Unclear
Could the reference standard, its conduct, or its interpretation have RISK: Unclear	
introduced bias?	
B. APPLICABILITY	
Is there concern that the target condition as defined by the Concerns: High	
reference standard does not match the review question?	
DOMAIN 4: FLOW AND TIMING	
A. RISK OF BIAS	
All participants appear to have been included in the analyses	

Could the patient flow have introduced bias?	RISK: Low
Were all patients included in the analysis?	Yes
Did patients receive the same reference standard?	Yes
Did all patients receive a reference standard?	Yes

## Study: Lippi (2012)<sup>74</sup> DOMAIN 1: PATIENT SELECTION A. RISK OF BIAS

Consecutive patients presenting to the emergency department with che	est pain of recent onset (<3 hrs)	
No exclusion criteria reported		
Was a consecutive or random sample of patients enrolled?		Yes
Was a case-control design avoided?		Yes
Did the study avoid inappropriate exclusions?		Unclear
Could the selection of patients have introduced bias?	RISK: Low	
B. APPLICABILITY		
Unselected chest pain population AMI diagnoses may have included bot	th NSTEMI and STEMI	
Do the included patients match the question?	Concerns: High	
DOMAIN 2: INDEX TEST(S)		
A. RISK OF BIAS		
Beckman Coulter HS-AccuTnl on admission. Reference standard final dia admission hs-Tnl. Threshold derived from ROC analysis	agnosis (AMI); probably made later	than
Were the index test results interpreted without knowledge of the resu	Ilts of	Yes
the reference standard?		
If a threshold was used, was it pre-specified?		No
Could the conduct or interpretation of the index test have	RISK: High	
introduced bias?	-	
B. APPLICABILITY		
Are there concerns that the index test, its conduct, or interpretation	Concerns: Low	
differ from the review question?		
DOMAIN 3: REFERENCE STANDARD		
A. RISK OF BIAS		
Reterence standard final diagnosis of AMI, criteria for diagnosis not repo	orted	
Unclear whether those adjudicating final diagnosis were blind to hs-TnI	_	
Is the reference standard likely to correctly classify the target conditio	n?	Unclear
Were the reference standard results interpreted without knowledge of	if the	Unclear
results of the index test?		
Could the reference standard, its conduct, or its interpretation have	RISK: Unclear	
introduced bias?		
B. APPLICABILITY		
Is there concern that the target condition as defined by the	Concerns: High	
reference standard does not match the review question?		
DOMAIN 4: FLOW AND TIMING		

#### DOMAIN 4: FLOW AND TIMING A. RISK OF BIAS

No withdrawals reported	
Did all patients receive a reference standard?	Unclear
Did patients receive the same reference standard?	Unclear
Were all patients included in the analysis?	Unclear
Could the patient flow have introduced bias?	RISK: Unclear

## Study: Melki (2011)<sup>50</sup> DOMAIN 1: PATIENT SELECTION A. RISK OF BIAS

Recruitment described as "consecutive except for temporary interrupt	ions of the study due to high work load i	n
the coronary care unit"		
Was a consecutive or random sample of patients enrolled?		No
Was a case-control design avoided?		Yes
Did the study avoid inappropriate exclusions?		Yes
Could the selection of patients have introduced bias?	RISK: High	
B. APPLICABILITY		
Chest pain patients admitted to chest pain unit, excluding ST-segment	elevation	
Do the included patients match the question?	Concerns: High	
DOMAIN 2: INDEX TEST(S)		
A. RISK OF BIAS		
Roche Elecsys hs-TnT on admission and at 2 hrs. Reference standard (fi	nal diagnosis) determined after hs-TnT	
testing. Threshold based on assay characteristics, appears pre-determi	ned	
Were the index test results interpreted without knowledge of the res	sults of	Yes
the reference standard?		.,
If a threshold was used, was it pre-specified?		Yes
Could the conduct or interpretation of the index test have	RISK: Low	
Introduced bias?		
Are there concerns that the index test, its conduct, or interpretation	Concerns: Low	
differ from the review question?		
DOMAIN 3: REFERENCE STANDARD		
A. RISK OF BIAS		
Reference standard diagnosis of AMI based on joint European Cardiolo	gy Society an American College of Cardio	
criteria and included serial conventional cTnT or cTnI (9-12 hour time n	gy Society an American conege of cardic	лову
Determination of diagnosis was made blind to bs-TnT results	ont specified)	
Is the reference standard likely to correctly classify the target condition	on?	Voc
Were the reference standard results interpreted without knowledge	of the	Voc
results of the index test?		163
Could the reference standard its conduct or its interpretation have	RISK: Low	
introduced hiss?	MON. LOW	
Is there concern that the target condition as defined by the	Concerns: Low	
reference standard does not match the review question?	CONCENTS, LOW	
היכוכויכויכי שנמושמים מסבש וסד וומנטו נווכ ובעוכש קעבשנוטון:		
DOMAIN 4: FLOW AND TIMING		
A. RISK OF BIAS		

All participants appear to have been included in the analyses	
Did all patients receive a reference standard?	Yes
Did patients receive the same reference standard?	Yes
Were all patients included in the analysis?	Yes
Could the patient flow have introduced bias?	RISK: Low

#### Study: Parsonage (2013)<sup>57</sup> DOMAIN 1: PATIENT SELECTION A. RISK OF BIAS

Prospective studies: no further details on recruitment		
Was a consecutive or random sample of patients enrolled?		Jnclear
Was a case-control design avoided?		Yes
Did the study avoid inappropriate exclusions?	l	Jnclear
Could the selection of patients have introduced bias?	RISK: Unclear	
B. APPLICABILITY		
Unselected chest pain population AMI diagnoses may have included bo	th NSTEMI and STEMI	
Do the included patients match the question?	Concerns: High	
DOMAIN 2: INDEX TEST(S)		
A. RISK OF BIAS		
Bacha Elactive betrat and Abbatt ABCHITECT be STAT Tal. Throshold was	00th contile	
Roche Elecsys institut and Abbolic ARCHITECT ins-STAT This. Threshold was	s 99th centile	
More the index test results interpreted without knowledge of the resu	ults of	Voc
the reference standard?		res
If a threshold was used was it pre-specified?		Voc
Could the conduct or interpretation of the index test have	RISK' LOW	103
introduced bias?		
B. APPLICABILITY		
Are there concerns that the index test, its conduct, or interpretation	Concerns: Low	
differ from the review guestion?		
·		
DOMAIN 3: REFERENCE STANDARD		
A. RISK OF BIAS		
Reference standard diagnosis of AMI (criteria unclear) and included seri	al conventional cTnl (10-12 hour time	e point
not specified). Determination of diagnosis was made blind to hs-InI ar	id hs-Inl results	
Is the reference standard likely to correctly classify the target condition	in?	Yes
Were the reference standard results interpreted without knowledge of the finder test?	of the	Yes
results of the index test?	DISK: Low	
could the reference standard, its conduct, or its interpretation have	RISK: LOW	
D. AFFLICADILITT Is there concern that the target condition as defined by the	Concerns: High	
reference standard does not match the review question?	Concerns. nigh	
ירוריבאנים איז		
DOMAIN 4: FLOW AND TIMING		

Patients appear to be missing from the analyses, as 2x2 data (derived from reported sensitivity and specificity		
estimates and total number of AMI) do not match reported number of test positives		
Did all patients receive a reference standard?	Unclear	
Did patients receive the same reference standard?	Yes	
Were all patients included in the analysis?	Unclear	
Could the patient flow have introduced bias? F	RISK: Unclear	

#### Study: Saenger (2010)<sup>71</sup> DOMAIN 1: PATIENT SELECTION A. RISK OF BIAS

No details on how patients were selected. No exclusion criteria reporte	ed.	
Was a consecutive or random sample of patients enrolled?		Unclear
Was a case-control design avoided?		Yes
Did the study avoid inappropriate exclusions?		Unclear
Could the selection of patients have introduced bias?	RISK: Unclear	
B. APPLICABILITY		
No exclusion criteria reported, reference standard was AMI (diagnosis r	nethod not specified),diagnoses incl	uded
STEMI		
Do the included patients match the question?	Concerns: High	
DOMAIN 2: INDEX TEST(S)		
A. RISK OF BIAS		
Roche Elecsys hs-TnT on admission and after 3 hrs. Data reported for ac	Imission and $\Delta$ 0-3 hrs. No details of	
interpretation reported. Threshold for $\Delta$ value derived from ROC analys	is: 99th centile also used	
Were the index test results interpreted without knowledge of the resu	ults of	Unclear
the reference standard?		
If a threshold was used, was it pre-specified?		Yes
Could the conduct or interpretation of the index test have	RISK: Low	
introduced bias?		
B. APPLICABILITY		
Are there concerns that the index test, its conduct, or interpretation	Concerns: Low	
differ from the review question?		
DOMAIN 3: REFERENCE STANDARD		
A. RISK OF BIAS		
Reference standard diagnosis of AMI (no details reported)		
Is the reference standard likely to correctly classify the target condition	200	Unclear
Were the reference standard results interpreted without knowledge of	of the	Unclear
results of the index test?		Uncical
Could the reference standard, its conduct, or its interpretation have	RISK: Unclear	
introduced bias?		
B. APPLICABILITY		
Is there concern that the target condition as defined by the	Concerns: High	
reference standard does not match the review question?		
· · · · · · · · · · · · · · · · · · ·		
DOMAIN 4: FLOW AND TIMING		
A. RISK OF BIAS		
NO WITHORAWAIS REPORTED		1100-0
Did all patients receive a reference standard?		Unclear
Did patients receive the same reference standard?		Unclear
were an patients included in the analysis?	DICK	Unclear
Could the patient flow have introduced blas?	RISK:	unclear

#### Study: Sanchis (2012)<sup>41</sup> DOMAIN 1: PATIENT SELECTION A. RISK OF BIAS

Patients excluded due to troponin elevation in any of 2 serial determinations (at arrival and 6-8 hours later) and		
prior diagnosis of ischemic heart disease. No details on how patients we	ere selected for the study.	
Was a consecutive or random sample of patients enrolled?		Unclear
Was a case-control design avoided?		Yes
Did the study avoid inappropriate exclusions?		No
Could the selection of patients have introduced bias?	RISK: High	
B. APPLICABILITY		
Selected low risk population		
Do the included patients match the question?	Concerns: High	
DOMAIN 2: INDEX TEST(S)		
A. RISK OF BIAS		
Roche Elecsys hs-TnT on admission and at 6-8 hrs (data reported for adr	mission and peak values). Reference	e
standard (30 day composite) occurred after testing. Thresholds were re	ported as pre-specified	
Were the index test results interpreted without knowledge of the resu	ilts of	Yes
the reference standard?		
If a threshold was used, was it pre-specified?		Yes
Could the conduct or interpretation of the index test have	RISK: Low	
introduced bias?		
B. APPLICABILITY		
Are there concerns that the index test, its conduct, or interpretation	Concerns: Low	
differ from the review question?		
DOMAIN 3: REFERENCE STANDARD		
A. RISK OF BIAS		
		1
Composite 30 day end point of AMI, death and revascularisation		
Not clear whether those adjudicating AMI were aware of hs-TnT results		
Is the reference standard likely to correctly classify the target conditio	n?	Yes
Were the reference standard results interpreted without knowledge of	of the	Unclear

Were the reference standard results interpreted without knowledge	or the
results of the index test?	
Could the reference standard, its conduct, or its interpretation have	RISK: Unclear
introduced bias?	
B. APPLICABILITY	
Is there concern that the target condition as defined by the reference standard does not match the review question?	Concerns: Low

#### DOMAIN 4: FLOW AND TIMING A. RISK OF BIAS

All participants appeared to have been included in the analyses	
Did all patients receive a reference standard?	Yes
Did patients receive the same reference standard?	Yes
Were all patients included in the analysis?	Yes
Could the patient flow have introduced bias?	<b>RISK: Low</b>

#### Study: Santalo (2013)<sup>39</sup> DOMAIN 1: PATIENT SELECTION A. RISK OF BIAS

Consecutive adult patients presenting to the emergency department	
Was a consecutive or random sample of patients enrolled?	Ye
Was a case-control design avoided?	Ye
Did the study avoid inappropriate exclusions?	Ye
Could the selection of patients have introduced bias?	RISK: Low
B. APPLICABILITY	
Appears to be an unselected emergency department chest pain popula	ation
Do the included patients match the question?	Concerns: Low
DOMAIN 2: INDEX TEST(S)	
A. RISK OF BIAS	
Roche Elecsys hsTnT on admission and at 2.4. and 6-8 hours or until di	scharge (data reported for admission and A
values) Unclear whether hs. The interpreted blind to CTh	
Were the index test results interpreted without knowledge of the res	sults of Lincle
the reference standard?	
If a threshold was used, was it pre-specified?	V
Could the conduct or interpretation of the index test have	BISK: Low
introduced bias?	
B. APPLICABILITY	
Are there concerns that the index test, its conduct, or interpretation	Concerns: Low
differ from the review question?	
·	
DOMAIN 3: REFERENCE STANDARD	
A. RISK OF BIAS	
Final diagnosis adjudicated by committee, based on Roche cTnT at adm	hission and 2, 4 and 6-8 hours or until
discharge (10-12 hr time point not specified). NSTEMI defined as cinit	$>10$ ng/L and $\Delta c$ in i >20%; also 99th centile
Unclear whether adjudicators were blinded to hs-in i	2
Is the reference standard likely to correctly classify the target condition	ON? Ye
were the reference standard results interpreted without knowledge	of the Unclea
results of the index test?	DICK: Unalgar
Could the reference standard, its conduct, or its interpretation have	RISK: Unclear
B. APPLICABILITY	Concerner Unclear
is there concern that the target condition as defined by the	Concerns: Unclear
reference standard does not match the review question?	
DOMAIN A. FLOW AND TIMING	
DOMAIN 4: FLOW AND THINING	
A. RISK OF BIAS	
All participants appear to have been included in the analyses	
Did all patients receive a reference standard?	Ye
Did all patients receive a reference standard? Did patients receive the same reference standard?	Yı Yı

Could the patient flow have introduced bias?

**RISK: Low** 

#### Study: Sebbane (2013)<sup>63</sup> DOMAIN 1: PATIENT SELECTION A. RISK OF BIAS

No details on how patients were selected for inclusion.		
Was a consecutive or random sample of patients enrolled?		Unclear
Was a case-control design avoided?		Yes
Did the study avoid inappropriate exclusions?		Yes
Could the selection of patients have introduced bias?	RISK: Unclear	
B. APPLICABILITY		
Unselected cohort of adult patients presenting with chest pain of r	recent onset (within 12 hours)	
Do the included patients match the question?	Concerns: Low	
DOMAIN 2: INDEX TEST(S)		
A. RISK OF BIAS		
Roche Elecsys hsTnT on admission or from sample taken during pre	e-hospital management. Final Dia	gnosis
adjudicated one month after acute episode. Optimal diagnostic the	resholds were determined using v	within study
ROC analyses; 99th centile also reported		
Were the index test results interpreted without knowledge of the	e results of	Yes
the reference standard?		
If a threshold was used, was it pre-specified?		Yes
Could the conduct or interpretation of the index test have	RISK: Low	
introduced bias?		
B. APPLICABILITY		
Are there concerns that the index test, its conduct, or interpretat	ion Concerns: Low	
differ from the review question?		
DOMAIN 3: REFERENCE STANDARD		
A. RISK OF BIAS		
Diagnosis determined by two independent emergency department	t physicians, based on Joint Europ	aan Cardiology
Society on American College of Cardiology criteria, Reference stary	dard included cTal taken on admi	csion at 6 brs
and beyond as pooled (10, 12 br time point pot specified). Physicia	and included criticaken on admis	ssion, at o mis
and beyond, as needed (10-12 in time point not specified). Physica		its, but were
billiueu to its-itili results.	adition	Vaa
is the reference standard likely to correctly classify the target cor		Yes
were the reference standard results interpreted without knowle	age of the	Yes
results of the index test?		
Could the reference standard, its conduct, or its interpretation ha	ave RISK: Low	

Diagnosis determined by two independent emergency department phy	sicians, based on Joint Europea	an Cardiology
Society an American College of Cardiology criteria. Reference standard	included cTnI taken on admissi	on, at 6 hrs
and beyond, as needed (10-12 hr time point not specified). Physicians h	ad access to serial cTnI results,	, but were
blinded to hs-TnT results.		
Is the reference standard likely to correctly classify the target condition	on?	Yes
Were the reference standard results interpreted without knowledge of	of the	Yes
results of the index test?		
Could the reference standard, its conduct, or its interpretation have	RISK: Low	
introduced bias?		
B. APPLICABILITY		
Is there concern that the target condition as defined by the	Concerns: High	
reference standard does not match the review question?		

#### **DOMAIN 4: FLOW AND TIMING** A. RISK OF BIAS

54 patients were excluded from the analyses because of missing data, including lack of copeptin, hs-cTnT, and cTnI measurements

Could the patient flow have introduced bias?	RISK: High
Were all patients included in the analysis?	No
Did patients receive the same reference standard?	Yes
Did all patients receive a reference standard?	Yes

#### **APPENDIX 4: TABLE OF EXCLUDED STUDIES WITH RATIONALE**

To be included in the review studies had to fulfil the following criteria:

Population:	Adults ( $\geq$ 18 yrs) presenting with acute 'pain, discomfort or pressure in the
	chest, epigastrium, neck, jaw, or upper limb without an apparent non-
	cardiac source' due to a suspected, but not proven, AMI or ACS
Setting:	Secondary or tertiary care
Index Test:	Abbott ARCHITECT (STAT hs-cTnl); Beckman Coulter Access and Unicel Dxl
	(accuTnI+3); Roche Elecsys (cTnT-hs or cTnT-hs STAT); results available within
	3 hours
Reference Standard:	Universal definition of AMI, including measurement of troponin T or I (using
	any method not defined as a hs-cTn test) on presentation and 10-12 hours
	after the onset of symptoms in $\geq$ 80% of the population or occurrence of
	MACE (any definition used in identified studies) during 30 day follow-up
Outcome:	Sufficient data to construct 2x2 table of test performance

The table below summarises studies which were screened for inclusion based on full text publication but did not fulfil one or more of the above criteria. Studies were assessed sequentially against criteria; as soon as a study had failed based on one of the criteria it was not assessed against subsequent criteria. The table shows which of the criteria each study fulfilled ("Yes") and on which item it failed ("No").

Study Details	Primary study	Population	Setting	Index Test	Reference Standard	Outcome
Ahmed(2013) <sup>113</sup>	Yes	Yes	Yes	Yes	No	
Aldous(2010) <sup>114</sup>	Yes	Yes	Yes	Yes	Unclear	No
Aldous(2010) <sup>115</sup>	Yes	Yes	Yes	Yes	Unclear	No
Aldous(2012) <sup>116</sup>	No					
Aldous(2010) <sup>117</sup>	Yes	Yes	Yes	Unclear	No	
Aldous(2012) <sup>118</sup>	Yes	Yes	Yes	Unclear	Yes	No
Aldous(2012) <sup>119</sup>	Yes	Yes	Yes	No		
Aldous(2012) <sup>120</sup>	No					
Aldous(2012) <sup>121</sup>	No					
Alexandra(2013) <sup>122</sup>	Yes	Yes	Yes	No		
Arenja(2010) <sup>123</sup>	Yes	Yes	Yes	Yes	No	
Bahrmann(2012) <sup>124</sup>	Yes	No				
Bahrmann(2013) <sup>125</sup>	Yes	No				
Bahrmann(2013) <sup>126</sup>	Yes	No				
Bahrmann(2012) <sup>127</sup>	Yes	Yes	Yes	Yes	No	
Balmelli(2013) <sup>128</sup>	Yes	Yes	Yes	Yes	Unclear	No
Balmelli(2011) <sup>129</sup>	Yes	Yes	Yes	Yes	Unclear	No
Beyrau(2009) <sup>130</sup>	Yes	No				
Bhardwaj(2011) <sup>131</sup>	Yes	Yes	Yes	No		
Bhardwaj(2011) <sup>132</sup>	Yes	Yes	Yes	No		

Study Details	Primary	Population	Setting	Index Test	Reference	Outcome
$P_{1}^{i} = e^{iH_{2}} (2010)^{133}$	study	No	N/	Unalasi	Standard	
Biasilio(2010)	Yes	Yes	Yes	Unclear	NO	
Biasucci(2010)	Yes	Yes	Yes	Yes	NO	
Biasucci(2010)	Yes	Yes	Yes	Yes	NO	
Biasucci(2010)	Yes	Yes	Yes	Yes	NO	
Biasucci(2010)	Yes	Yes	Yes	Yes	NO	
Biasucci(2011)	Yes	Yes	Yes	Yes	NO	
Biener(2013)	Yes	Unclear	Yes	Unclear	Unclear	NO
Biener(2012)	Yes	Yes	Yes	Unclear	Unclear	NO
Biener(2013)	Yes	NO	N	N -		
$\frac{\text{Biener}(2013)}{\text{Piecite}(2006)^{143}}$	Yes	Yes	Yes	NO		
BIOSITE(2006)	Yes	Yes	Yes	NO	Under	Na
BODY(2012)	Yes	Yes	Yes	Unclear	Unclear	NO
BODY(2012)	Yes	res	res	Yes	res	NO
BO0y(2012)	NO	Vac	Vac	Vac	Vac	No
Braga(2011)	Yes	Yes	Yes	Yes	Yes	NO
Braga(2011)	Yes	Yes	Yes	Yes	Yes	NO
Bronze(2012)	Yes	Yes	Yes	Yes	NO	
Brown(2007)	Yes	Yes	Yes	NO	Na	
$\frac{Buccellettl(2012)}{Buckl(2011)^{152}}$	Yes	Yes	Yes	Yes	NO	
Buni(2011)	Yes	NO	Vaa	Vee	Vaa	Na
Cardino(2012)	Yes	res	res	Yes	res	NO
Carino(2013)	NO					
$\frac{\text{Certain}(2012)}{\text{Charpontiar}(2011)^{156}}$	NO	Vac	Vac	No		
Charpentier(2011)	res	res	res	NO		
$Cobeaux(2013)^{157}$	NO					
Gobeaux(2013)	Voc	Voc	Voc	No		
Collinson(2012)	Yes	Yes	Yes	No		
Collinson(2012)	Yes	Yes	Yes	No		
$Collinson(2006)^{161}$	Yes	Yes	Yes	No		
$Collinson(2010)^{162}$	Ves	Vos	Ves	No		
$Costabel(2013)^{163}$	No	163	163	NO		
Cullen $(2011)^{164}$	Vos	Ves	Vas	No		
$Dawson(2013)^{165}$	Vos	No	163			
Diarcks $(2012)^{166}$	Ves	Vos	Voc	No		
Drevler(2012)	Ves	Vos	Ves	Unclear	Unclear	No
Engel(2007) <sup>168</sup>	Ves	Ves	Ves	No	Unclear	NO
Escabi-Mendoza $(2010)^{169}$	Vos	Ves	Vas	No		
Figiel(2008) <sup>170</sup>	Ves	No	163			
$Fitzgerald(2011)^{81}$	Yes	Yes	Yes	No		
Freund(2011) <sup>171</sup>	Yes	Yes	Yes	Unclear	Unclear	No
Freund(2011) <sup>172</sup>	Yes	Yes	Yes	Unclear	Unclear	No
Giannitsis(2010) <sup>112</sup>	Yes	No				
Giannitsis(2011) <sup>173</sup>	Yes	Yes	Yes	Unclear	Unclear	No
Giavarina(2012) <sup>174</sup>	No					-
Giavarina(2011) <sup>175</sup>	Yes	Yes	Yes	No		
Gimenez(2012) <sup>176</sup>	Yes	Yes	Yes	No		
Gimenez(2012) <sup>177</sup>	Yes	Yes	Yes	No		
Goodacre(2011) <sup>99</sup>	Yes	Yes	Yes	No		
Goodacre(2013) <sup>7</sup>	No					
Goodacre(2011) <sup>85</sup>	Yes	Yes	Yes	No		
Gustapane(2012) <sup>178</sup>	Yes	Yes	Yes	Unclear	No	

Study Details	Primary	Population	Setting	Index Test	Reference	Outcome
Custo non a (2012) <sup>179</sup>	Study	Vaa	Vee	Linglagy	Stalluaru	
Gustapane(2012)	Yes	Yes	Yes	Unclear	INU	Na
Hadi(2011)	Yes	Yes	Yes	Unclear	Unclear	NO
Haaf(2011)	Yes	res	Yes	Unclear	Unclear	NO
Haaf(2013)	NO					
Haaf(2012)	Yes	Yes	Yes	Yes	NO	
Haaf(2012) $(2012)^{185}$	Yes	Yes	Yes	Yes	NO	
Haltern(2010)	Yes	Yes	Yes	NO	11	N -
Heinisch(2010)	Yes	Yes	Yes	Unclear	Unclear	NO
Hochholzer(2011)	Yes	Yes	Yes	Yes	NO	
Hochholzer(2010)	Yes	Yes	Yes	Yes	NO	
Hoeller(2012)	Yes	Yes	Yes	Yes	NO	
Hoeller(2012)	Yes	Yes	Yes	Yes	NO	
11Va(2009)	Yes	Yes	NO	Vee	Na	
Indue(2011)	Yes	Yes	Yes	Yes	INO	Na
rran(2011)	Yes	Yes	Yes	Unclear	Unclear	NO
rran(2011)	Yes	Yes	Yes	Unclear	NO	N -
rran(2013)	Yes	Yes	Yes	Yes	Unclear	NO
Irfan(2013)	Yes	Yes	Yes	Yes	Unclear	NO
Jairam(2011)	Yes	NO	N	NL-		
Januzzi(2010)	Yes	Yes	Yes	NO	N -	
Januzzi(2009)	Yes	Yes	Yes	Yes	NO	
Januzzi(2013)	Yes	Yes	Yes	NO		
Jia(2009)	Yes	Yes	Yes	NO		
KagaWa(2013)	Yes	Yes	Yes	NO	Na	
$\frac{\text{Karakas}(2012)^{204}}{\text{Kaysak}(2012)^{204}}$	Yes	Yes	Yes	res	INU	No
Kavsak(2012)	Ves	Ves	Ves	No	Unclear	NO
$(2007)^{206}$	Vos	Ves	Ves	Unclear	No	
Kavsak(2013)	Vos	Ves	Ves	No	NO	
Kavsak(2003)	Ves	Ves	Ves	Ves	Unclear	No
Kavsak(2012)	Yes	No	103	103	Uncical	110
Kavsak(2000)	Yes	Yes	Yes	Yes	No	
Kavsak(2010) <sup>211</sup>	Yes	Yes	Yes	Yes	Yes	No
Keene(2012) <sup>212</sup>	Yes	Yes	Yes	Yes	No	
Keller(2011) <sup>213</sup>	Yes	Yes	Yes	Yes	Unclear	No
Keller(2011) <sup>214</sup>	Yes	Yes	Yes	Yes	Unclear	No
Keller(2009) <sup>215</sup>	Yes	Yes	Yes	No		
Keller(2010) <sup>216</sup>	Yes	Yes	Yes	No		
Keller(2009) <sup>217</sup>	Yes	Yes	Yes	No		
Kelly(2011) <sup>218</sup>	Yes	Yes	Yes	No		
Khan(2011) <sup>219</sup>	Yes	Yes	Yes	Yes	No	
Khoo(2008) <sup>220</sup>	Yes	Unclear	Yes	No		
Kitamura(2012) <sup>221</sup>	Yes	Yes	Yes	No		
Kobayashi(2011) <sup>222</sup>	Yes	Yes	Yes	Unclear	No	
Kobayashi(2011) <sup>223</sup>	Yes	Yes	Yes	Yes	No	
Koenig(2008)	Yes	Yes	Yes	Yes	NO	
Lachak(2007)	Yes	Yes	Yes	Unclear	NO	
1000000000000000000000000000000000000	Yes	res	res	Unclear	INO	
Linualii(2009)	No	INU				
Linni(2012) <sup>229</sup>	No					
Lippi(2013) <sup>230</sup>	No					
				i		

Study Details	Primary	Population	Setting	Index Test	Reference	Outcome
$10tze(2011)^{231}$	Vos	Voc	Voc	No	Standard	
Lotze(2011)	Yes	Yes	Yos	NO	No	
10(2e(2011))	Yes	Yes	Yes	No	NO	
Maciae(2006)	Yes	Tes No	res	NO		
Mair(2011)	Yes	NO				
Matri(2011)	Yes	NO	Vaa	Linglagy	Unclose	Na
MacSul(2011)	Yes	Yes	Yes	Unclear	Unclear	INO
$\frac{Malancan(2011)}{Malancan(2008)^{238}}$	Yes	Yes	Yes	Unclear	INO	
Malki(2011) <sup>239</sup>	Yes	Yes	Yes	NO	No	
$M_{0}$	Yes	Yes	Yes	No	INU	
$M_{0}$	Yes	Yes	Yes	NO	No	
$M_{enb} = for(2012)^{242}$	Yes	fes	res	Tes	INU	
$M_{2013}$	Yes	NO	Voc	Voc	Voc	No
$M_{2011}$	Yes	Yes	Yes	Yes	Yes	No
Mouro(2012) <sup>245</sup>	Yes	Yes	Yes	Yes	No	NO
$M_{2012}(2013)$	Yes	Yes	Yes	Yes	No	
$Mikkal(2012)^{247}$	Yes	Yes	Yes	Yes	NU	No
Mikkel(2013)	Yes	Yes	Yes	No	Unclear	NO
Mikkel(2013)	Yes	Yes	Yes	NO		
$Milk(2010)^{250}$	Yes	Yes	Yes	INU	No	
$Mills(2010)^{251}$	Yes	Yes	Yes	Unclear	No	
Mills(2010)	Yes	Yes	Yes	Voc	No	
Mins(2012)	Yes	res	res	res	INO	
$\frac{1}{10000000000000000000000000000000000$	Yes	NO	Vac	Vac	No	
$Moehring(2012)^{255}$	Yes	Yes	Yes	Yes	INU	No
$\frac{1}{10000000000000000000000000000000000$	Yes	Yes	Yes	Yes	Unclear	NO
$Morrow(2000)^{257}$	res	res	res	res	res	INO
$N_{2005}^{258}$	Voc	Voc	Voc	No		
Naguriey(2003)	Yes	Yes	Yos	Uncloar	Voc	No
$Narco(2000)^{260}$	Voc	Yes	Yos	No	165	NO
$N_{gap}(2010)^{261}$	Ves	Ves	Vos	Vos	Voc	No
$N_{\text{part}}(2010)^{262}$	Ves	Ves	Ves	Unclear	Unclear	No
Normann $(2012)^{263}$	Ves	Ves	Ves	Ves	No	110
Nusier $(2006)^{264}$	Ves	Ves	Ves	No	110	
$Olivieri(2012)^{265}$	Ves	Ves	Ves	No		
$Orshorne(2012)^{266}$	No	105	105			
Paoloni(2010) <sup>267</sup>	Ves	Ves	Ves	Unclear	No	
$Perego(2011)^{268}$	Yes	105	105	Officieur	110	
Plebani(2009) <sup>269</sup>	Yes	Yes	Yes	No		
Ploner(2011) <sup>270</sup>	Yes	No	No			
Popp(2010) <sup>271</sup>	Yes	Yes	Yes	Yes	No	
Potocki(2011) <sup>272</sup>	Yes	Yes	Yes	No		
Pracon(2012) <sup>273</sup>	Yes	Yes	Yes	No		
Rajdl(2011) <sup>274</sup>	Yes	Yes	Yes	Yes	Unclear	No
Ray(2011) <sup>275</sup>	Yes	Yes	Yes	Unclear	Unclear	No
Reichlin(2012) <sup>276</sup>	No					
Reichlin(2011) <sup>277</sup>	Yes	Yes	Yes	Unclear	Unclear	No
Reichlin(2012) <sup>278</sup>	Yes	Yes	Yes	Yes	No	
Reichlin(2010) <sup>279</sup>	Yes	Yes	Yes	Unclear	No	
Reichlin(2010) <sup>200</sup>	Yes	Yes	Yes	Unclear	No	
Reichlin(2012) <sup>201</sup>	Yes	Yes	Yes	Yes	No	
Rubini Gimenez(2012) <sup>282</sup>	Yes	Yes	Yes	Yes	Unclear	No

Study Details	Primary	Population	Setting	Index Test	Reference Standard	Outcome
Budolph(2011) <sup>283</sup>	Ves	Ves	Ves	Unclear	Unclear	No
$Rudolph(2011)^{284}$	Vas	Ves	Vas	Unclear	Unclear	No
$Rudolph(2012)^{285}$	Vas	Ves	Vas	No	Unclear	NO
Samaraje $(2010)^{286}$	Vas	Ves	Vas	Unclear	No	
Scharphorst(2011) <sup>287</sup>	Vas	Ves	Vas	No	NO	
$Schaub(2012)^{288}$	Vas	Ves	Vas	Ves	Ves	No
Schoos(2013) <sup>289</sup>	Ves	Ves	Ves	Ves	Unclear	No
Schoos(2013) <sup>290</sup>	Yes	Yes	Yes	Yes	Unclear	No
Schreiber(2012) <sup>291</sup>	Yes	Yes	Yes	No	Uncical	110
Sethi(2013) <sup>292</sup>	No	105	105			
Shand(2012) <sup>293</sup>	Yes	Yes	Unclear	Unclear	No	
Shortt(2013) <sup>294</sup>	No	100	Officieur	oncical		
Spanuth(2011) <sup>295</sup>	Yes	Yes	Yes	Unclear	Unclear	No
Spasic-Obradovic $(2011)^{296}$	Yes	Yes	Yes	Yes	No	
Stengaard $(2012)^{297}$	Yes	Yes	Yes	Unclear	Unclear	No
Taisic(2013) <sup>298</sup>	Yes	Yes	Yes	Unclear	Unclear	No
Taisic(2013) <sup>299</sup>	Yes	Yes	Yes	Unclear	Unclear	No
$Taisic(2012)^{300}$	Yes	Yes	Yes	Unclear	No	
Taisic(2013) <sup>301</sup>	Yes	Yes	Yes	Unclear	Unclear	No
Tamimi(2010) <sup>302</sup>	Yes	Yes	Yes	No	Unclear	
Tanaka(2006) <sup>303</sup>	Yes	Yes	Yes	No		
Than(2012) <sup>304</sup>	Yes	Yes	Yes	No		
Thelin(2013) <sup>305</sup>	Yes	Yes	Yes	Yes	No	
Thomas(2007) <sup>306</sup>	Yes	No				
Thomas(2007) <sup>307</sup>	Yes	No				
Truong(2012) <sup>308</sup>	Yes	Yes	Yes	No		
Truong(2011) <sup>309</sup>	Yes	Yes	No	Unclear	Unclear	No
Twerenbold(2010) <sup>310</sup>	Yes	Yes	Yes	Yes	Unclear	No
Twerenbold(2010) <sup>311</sup>	Yes	Yes	Yes	Yes	Unclear	No
Twerenbold(2010) <sup>312</sup>	Yes	Yes	Yes	Yes	Unclear	No
Twerenbold(2011) <sup>313</sup>	Yes	Yes	Yes	No		
Twerenbold(2012) <sup>314</sup>	Yes	Yes	Yes	Yes	No	
University of Edinburgh	Yes	Yes	Yes	Unclear	No	
(2013) <sup>315</sup>						
University of Erlangen	Yes	Yes	Yes	Unclear	Unclear	
(2013) <sup>316</sup>						
van Wijk(2012) <sup>317</sup>	Yes	Yes	Yes	Yes	No	
Vasikaran(2012) <sup>318</sup>	No					
Veljkovic (2012) <sup>319</sup>	Yes	Yes	Yes	Yes	Unclear	No
Venge(2008) <sup>320</sup>	Yes	No				
Venge(2009) <sup>321</sup>	Yes	No				
Venge(2010) <sup>322</sup>	Yes	No				
Weber(2011) <sup>323</sup>	Yes	No				
Weber(2009) <sup>324</sup>	Yes	No				
Wildi(2012) <sup>325</sup>	Yes	Yes	Yes	No		
Wong(2010) <sup>326</sup>	Yes	No	Yes	No		
Worster(2013) <sup>327</sup>	Yes	No	Yes	Yes	No	No
Zahid(2009) <sup>328</sup>	Yes	Yes	Yes	Unclear	Unclear	No
Zahid(2008) <sup>329</sup>	Yes	Yes	Yes	No		
Zellweger(2012) <sup>330</sup>	Yes	Yes	Yes	Yes	No	
Zuily(2011) <sup>331</sup>	Yes	Yes	Yes	Yes	No	

# APPENDIX 5: SENSITIVITY ANALYSES (BASE CASE)

#### Deterministic base case:

Strategy			Com	pared to St	andard troponin	Compared to next best strategy				
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	
Abbott 99th centile	£2,257	11.734	-£440	-0.015	£28,870					
Roche 99th centile	£2,301	11.740	-£396	-0.010	£41,233	Abbott 99th centile	£44	0.006	£7,777	
Beckman 99th centile	£2,327	11.743	-£370	-0.006	£58,988	Roche 99th centile	£26	0.003	£7,777	
Roche strategy	£2,426	11.744	-£271	-0.006	£48,559	Beckman 99th centile	£99	0.001	Extendedly dominated	
Abbott strategy	£2,493	11.748	-£204	-0.002	£124,391	Beckman 99th centile	£167	0.005	£35,904	
Standard troponin	£2,697	11.749				Abbott strategy	£204	0.002	£124,391	

#### Increased re-infarction & mortality risk for no treatment (vs treated) = lifetime (instead of only during the first year after presentation at ED)

Strategy			Comp	ared to Sta	indard troponin	Compared to next best strategy				
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	
Abbott 99th centile	£2,257	11.677	-£440	-0.072	£6,112					
Roche 99th centile	£2,301	11.704	-£396	-0.045	£8,731	Abbott 99th centile	£44	0.027	Extendedly dominated	
Beckman 99th centile	£2,327	11.720	-£370	-0.030	£12,493	Abbott 99th centile	£69	0.042	£1,642	
Roche strategy	£2,426	11.723	-£271	-0.026	£10,284	Beckman 99th centile	£99	0.003	Extendedly dominated	
Abbott strategy	£2,493	11.741	-£204	-0.008	£26,352	Beckman 99th centile	£167	0.022	£7,602	
Standard troponin	£2,697	11.749				Abbott strategy	£204	0.008	£26,352	

#### No doctor on demand, but average waiting time before doctor becomes available is increased with 1, 2 or 3 hours

Waiting time for doctor / decision pending delay = 1 hour(s)													
Strategy			Comp	ared to Sta	andard troponin	Compare	d to next b	est strateg	SY				
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>				
Abbott 99th centile	£2,285	11.734	-£440	-0.015	£28,869								
Roche 99th centile	£2,329	11.740	-£396	-0.010	£41,232	Abbott 99th centile	£44	0.006	Extendedly dominated				
Beckman 99th centile	£2,355	11.743	-£370	-0.006	£58,987	Abbott 99th centile	£70	0.009	£7,776				
Roche strategy	£2,470	11.744	-£255	-0.006	£45,643	Beckman 99th centile	£115	0.001	Extendedly dominated				

Abbott strategy	£2,541	11.748	-£184	-0.002	£112,580	Beckman 99th centile	£186	0.005	£40,072
Standard troponin	£2,725	11.749				Abbott strategy	£184	0.002	£112,580
Waiting time for doctor / decision	pending	delay = 2	hour(s)						
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,313	11.734	-£440	-0.015	£28,868				
Roche 99th centile	£2,357	11.740	-£396	-0.010	£41,231	Abbott 99th centile	£44	0.006	Extendedly dominated
Beckman 99th centile	£2,383	11.743	-£370	-0.006	£58,987	Abbott 99th centile	£70	0.009	£7,776
Roche strategy	£2,515	11.744	-£239	-0.006	£42,727	Beckman 99th centile	£132	0.001	Extendedly dominated
Abbott strategy	£2,588	11.748	-£165	-0.002	£100,769	Beckman 99th centile	£205	0.005	£44,240
Standard troponin	£2,754	11.749				Abbott strategy	£165	0.002	£100,769
Waiting time for doctor / decision	pending	delay = 3	hour(s)						
	Costs	QALYs	ΔCosts	ΔQALYs	$\Delta Costs / \Delta QALYs$	Comparator	∆Costs	ΔQALYs	ΔCosts / ΔQALYs
Abbott 99th centile	£2,342	11.734	-£440	-0.015	£28,868				
Roche 99th centile	£2,386	11.740	-£396	-0.010	£41,231	Abbott 99th centile	£44	0.006	Extendedly dominated
Beckman 99th centile	£2,411	11.743	-£370	-0.006	£58,986	Abbott 99th centile	£70	0.009	£7,775
Roche strategy	£2,559	11.744	-£223	-0.006	£39,811	Beckman 99th centile	£148	0.001	Extendedly dominated
Abbott strategy	£2,636	11.748	-£146	-0.002	£88,957	Beckman 99th centile	£225	0.005	£48,408
Standard troponin	£2,782	11.749				Abbott strategy	£146	0.002	£88,957

# Doctor on demand at ED, but average waiting time before doctor becomes available in the general ward is increased with 1, 2 or 3 hours (discharge to general ward after 4 hours after presenting at ED)

Waiting time for doctor / decision pending delay = 1 hour(s)													
Strategy			Com	pared to St	andard troponin	Compare	d to next b	est strateg	ÿ				
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>				
Abbott 99th centile	£2,258	11.734	-£468	-0.015	£30,665								
Roche 99th centile	£2,302	11.740	-£424	-0.010	£44,080	Abbott 99th centile	£44	0.006	£7,776				
Beckman 99th centile	£2,327	11.743	-£398	-0.006	£63,347	Roche 99th centile	£26	0.003	£7,776				

Roche strategy	£2,443	11.744	-£282	-0.006	£50,541	Beckman 99th centile	£115	0.001	Extendedly dominated
Abbott strategy	£2,513	11.748	-£212	-0.002	£129,290	Beckman 99th centile	£186	0.005	£40,072
Standard troponin	£2,725	11.749				Abbott strategy	£212	0.002	£129,290
Waiting time for doctor / decisi	on pend	ing delay	y = 2 hou	ır(s)					
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,259	11.734	-£495	-0.015	£32,459				
Roche 99th centile	£2,302	11.740	-£451	-0.010	£46,927	Abbott 99th centile	£44	0.006	£7,776
Beckman 99th centile	£2,328	11.743	-£425	-0.006	£67,705	Roche 99th centile	£26	0.003	£7,776
Roche strategy	£2,460	11.744	-£294	-0.006	£52,522	Beckman 99th centile	£132	0.001	Extendedly dominated
Abbott strategy	£2,534	11.748	-£220	-0.002	£134,189	Beckman 99th centile	£205	0.005	£44,240
Standard troponin	£2,754	11.749				Abbott strategy	£220	0.002	£134,189
Waiting time for doctor / decisi	on pend	ing delay	y = 3 hou	ır(s)					
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,260	11.734	-£522	-0.015	£34,254				
Roche 99th centile	£2,303	11.740	-£478	-0.010	£49,774	Abbott 99th centile	£44	0.006	£7,775
Beckman 99th centile	£2,329	11.743	-£453	-0.006	£72,064	Roche 99th centile	£26	0.003	£7,775
Roche strategy	£2,477	11.744	-£305	-0.006	£54,504	Beckman 99th centile	£148	0.001	Extendedly dominated
Abbott strategy	£2,554	11.748	-£228	-0.002	£139,089	Beckman 99th centile	£225	0.005	£48,408
Standard troponin	£2,782	11.749				Abbott strategy	£228	0.002	£139,089

#### Total delay of 1.5 hours

Strategy			Com	pared to S	tandard troponin	Compared to next best strategy				
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	
Abbott 99th centile	£2,214	11.734	-£440	-0.015	£28,871					
Roche 99th centile	£2,258	11.740	-£396	-0.010	£41,234	Abbott 99th centile	£44	0.006	£7,778	
Beckman 99th centile	£2,284	11.743	-£370	-0.006	£58,989	Roche 99th centile	£26	0.003	£7,778	
Roche strategy	£2,359	11.744	-£296	-0.006	£52,933	Beckman 99th centile	£75	0.001	Extendedly dominated	

Abbott strategy	£2,422	11.748	-£233	-0.002	£142,108	Beckman 99th centile	£138	0.005	£29,653
Standard troponin	£2,655	11.749				Abbott strategy	£233	0.002	£142,108

# MI treatment costs added for patients that were tested false positive

Strategy			Comp	ared to St	andard troponin	Compared to next best strategy			
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs
Abbott 99th centile	£2,456	11.734	-£241	-0.015	£15,824				
Abbott strategy	£2,671	11.748	-£26	-0.002	£16,050	Abbott 99th centile	£215	0.014	£15,797
Standard troponin	£2,697	11.749				Abbott strategy	£26	0.002	£16,050
Roche 99th centile	£2,760	11.740	£63	-0.010	Dominated	Standard troponin	£63	-0.010	Dominated
Roche strategy	£2,947	11.744	£251	-0.006	Dominated	Standard troponin	£251	-0.006	Dominated
Beckman 99th centile	£3,038	11.743	£341	-0.006	Dominated	Standard troponin	£341	-0.006	Dominated

#### MI treatment costs added to first year of UA

Strategy			Comp	ared to St	andard troponin	in Compared to next best strategy			
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,703	11.734	-£440	-0.015	£28,870				
Roche 99th centile	£2,747	11.740	-£396	-0.010	£41,233	Abbott 99th centile	£44	0.006	£7,777
Beckman 99th centile	£2,773	11.743	-£370	-0.006	£58,988	Roche 99th centile	£26	0.003	£7,777
Roche strategy	£2,872	11.744	-£271	-0.006	£48,559	Beckman 99th centile	£99	0.001	Extendedly dominated
Abbott strategy	£2,940	11.748	-£204	-0.002	£124,391	Beckman 99th centile	£167	0.005	£35,904
Standard troponin	£3,144	11.749				Abbott strategy	£204	0.002	£124,391

Test costs = £5										
Strategy			Com	pared to S	tandard troponin	Compared to next best strategy				
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	
Abbott 99th centile	£2,240	11.734	-£425	-0.015	£27,856					
Roche 99th centile	£2,284	11.740	-£381	-0.010	£39,624	Abbott 99th centile	£44	0.006	£7,778	
Beckman 99th centile	£2,310	11.743	-£355	-0.006	£56,526	Roche 99th centile	£26	0.003	£7,778	
Roche strategy	£2,400	11.744	-£265	-0.006	£47,439	Beckman 99th centile	£90	0.001	Extendedly dominated	
Abbott strategy	£2,466	11.748	-£199	-0.002	£121,624	Beckman 99th centile	£156	0.005	£33,550	
Standard troponin	£2,665	11.749				Abbott strategy	£199	0.002	£121,624	
Test costs = £40										
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	
Abbott 99th centile	£2,278	11.734	-£460	-0.015	£30,150					
Roche 99th centile	£2,322	11.740	-£416	-0.010	£43,264	Abbott 99th centile	£44	0.006	Extendedly dominated	
Beckman 99th centile	£2,348	11.743	-£390	-0.006	£62,097	Abbott 99th centile	£70	0.009	£7,776	
Roche strategy	£2,458	11.744	-£279	-0.006	£49,972	Beckman 99th centile	£111	0.001	Extendedly dominated	
Abbott strategy	£2,528	11.748	-£210	-0.002	£127,886	Beckman 99th centile	£180	0.005	£38,878	
Standard troponin	£2,737	11.749				Abbott strategy	£210	0.002	£127,886	

#### Test costs

#### AMI treatment costs

AMI treatment costs = £2,577									
Strategy			Comp	pared to St	andard troponin	Compared	d to next k	pest strate	gy
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,119	11.734	-£415	-0.015	£27,188				
Roche 99th centile	£2,154	11.740	-£380	-0.010	£39,551	Abbott 99th centile	£34	0.006	Extendedly dominated
Beckmann 99th centile	£2,174	11.743	-£360	-0.006	£57,307	Abbott 99th centile	£55	0.009	£6,096
Roche strategy	£2,272	11.744	-£262	-0.006	£46,877	Beckmann 99th centile	£98	0.001	Extendedly dominated

Abbott strategy	£2,333	11.748	-£201	-0.002	£122,710	Beckmann 99th centile	£159	0.005	£34,223
Standard troponin	£2,534	11.749				Roche strategy	£201	0.002	£122,710
AMI treatment costs = £4,295									
	Costs	QALYs	∆Costs	ΔQALYs	$\Delta Costs / \Delta QALYs$	Comparator	∆Costs	ΔQALYs	ΔCosts / ΔQALYs
Abbott 99th centile	£2,394	11.734	-£466	-0.015	£30,551	Abbott 99th centile	£53	0.006	£9,458
Roche 99th centile	£2,448	11.740	-£413	-0.010	£42,914	Roche 99th centile	£32	0.003	£9,458
Beckmann 99th centile	£2,479	11.743	-£381	-0.006	£60,669	Beckmann 99th centile	£100	0.001	Extendedly dominated
Roche strategy	£2,579	11.744	-£281	-0.006	£50,240	Beckmann 99th centile	£174	0.005	£37,586
Abbott strategy	£2,654	11.748	-£207	-0.002	£126,073	Roche strategy	£207	0.002	£126,073
Standard troponin	£2,860	11.749							

#### Post-MI health state costs

Post-MI health state costs (1st year) = £6,791												
Strategy			Com	pared to St	andard troponin	Compared to next best strategy						
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>			
Abbott 99th centile	£2,393	11.734	-£443	-0.015	£29,024							
Roche 99th centile	£2,438	11.740	-£398	-0.010	£41,387	Abbott 99th centile	£45	0.006	£7,931			
Beckman 99th centile	£2,464	11.743	-£371	-0.006	£59,142	Roche 99th centile	£26	0.003	£7,931			
Roche strategy	£2,563	11.744	-£272	-0.006	£48,713	Beckman 99th centile	£99	0.001	Extendedly dominated			
Abbott strategy	£2,632	11.748	-£204	-0.002	£124,545	Beckman 99th centile	£167	0.005	£36,059			
Standard troponin	£2,836	11.749				Abbott strategy	£204	0.002	£124,545			
Post-MI health state costs (1st yea	r) = £4,879	9										
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>			
Abbott 99th centile	£2,121	11.734	-£438	-0.015	£28,715							
Roche 99th centile	£2,164	11.740	-£395	-0.010	£41,078	Abbott 99th centile	£43	0.006	£7,623			
Beckman 99th centile	£2,189	11.743	-£369	-0.006	£58,834	Roche 99th centile	£25	0.003	£7,623			
Roche strategy	£2,288	11.744	-£271	-0.006	£48,405	Beckman 99th centile	£99	0.001	Extendedly dominated			

Abbott strategy	£2,355	11.748	-£204	-0.002	£124,237	Beckman 99th centile	£166	0.005	£35,750
Standard troponin	£2,558	11.749				Abbott strategy	£204	0.002	£124,237

#### Utility difference between UA and AMI

Utility difference between UA and AMI = 0.12												
Strategy			Com	pared to St	andard troponin	Compared	to next be	est strategy	1			
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs			
Abbott 99th centile	£2,257	11.779	-£440	-0.015	£28,870							
Roche 99th centile	£2,301	11.785	-£396	-0.010	£41,233	Abbott 99th centile	£44	0.006	£7,777			
Beckman 99th centile	£2,327	11.788	-£370	-0.006	£58,988	Roche 99th centile	£26	0.003	£7,777			
Roche strategy	£2,426	11.789	-£271	-0.006	£48,559	Beckman 99th centile	£99	0.001	Extendedly dominated			
Abbott strategy	£2,493	11.793	-£204	-0.002	£124,391	Beckman 99th centile	£167	0.005	£35,904			
Standard troponin	£2,697	11.794				Abbott strategy	£204	0.002	£124,391			
Utility difference between UA and	AMI = -0.	10										
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>			
Abbott 99th centile	£2,257	11.581	-£440	-0.015	£28,870							
Roche 99th centile	£2,301	11.587	-£396	-0.010	£41,233	Abbott 99th centile	£44	0.006	Extendedly dominated			
Beckman 99th centile	£2,327	11.590	-£370	-0.006	£58,988	Abbott 99th centile	£70	0.009	£7,777			
Roche strategy	£2,426	11.591	-£271	-0.006	£48,559	Beckman 99th centile	£99	0.001	Extendedly dominated			
Abbott strategy	£2,493	11.595	-£204	-0.002	£124,391	Beckman 99th centile	£167	0.005	£35,904			
Standard troponin	£2,697	11.597				Abbott strategy	£204	0.002	£124,391			

#### **MI disutility**

MI disutility = -0.059 (age = 45); -0.050 (age = 55); -0.024 (age = 65); -0.006 (age = 75+)												
Strategy			Compared to Standard troponin Compared to next best strategy									
	Costs	QALYs	ΔCosts	ΔQALYs         ΔCosts / ΔQALYs         Comparator         ΔCosts         ΔQALYs         ΔCosts / ΔQ/								
Abbott 99th centile	£2,257	11.735	-£440	-0.015	£28,832							
Roche 99th centile	£2,301	11.741	11         -£396         -0.010         £41,178         Abbott 99th centile         £44         0.006         £7,767									

Beckman 99th centile	£2,327	11.744	-£370	-0.006	£58,910	Roche 99th centile	£26	0.003	£7,767			
Roche strategy	£2,426	11.745	-£271	-0.006	£48,495	Beckman 99th centile	£99	0.001	Extendedly dominated			
Abbott strategy	£2,493	11.749	-£204	-0.002	£124,227	Beckman 99th centile	£167	0.005	£35,857			
Standard troponin	£2,697	11.751				Abbott strategy	£204	0.002	£124,227			
MI disutility = -0.061(age = 45); -0.052 (age = 55); -0.026 (age = 65); -0.008 (age = 75+)												
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>			
Abbott 99th centile	£2,257	11.733	-£440	-0.015	£28,908							
Roche 99th centile	£2,301	11.738	-£396	-0.010	£41,287	Abbott 99th centile	£44	0.006	Extendedly dominated			
Beckman 99th centile	£2,327	11.742	-£370	-0.006	£59,066	Abbott 99th centile	£70	0.009	£7,787			
Roche strategy	£2,426	11.742	-£271	-0.006	£48,623	Beckman 99th centile	£99	0.001	Extendedly dominated			
Abbott strategy	£2,493	11.746	-£204	-0.002	£124,556	Beckman 99th centile	£167	0.005	£35,952			
Standard troponin	£2,697	11.748				Abbott strategy	£204	0.002	£124,556			

# Mortality (30-day) treated AMI (decision tree)

Mortality (30-day) treated AMI = 0.120												
Strategy			Com	pared to Si	tandard troponin	Compared to next best strategy						
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs			
Abbott 99th centile	£2,219	11.710	-£432	-0.010	£41,819							
Roche 99th centile	£2,260	11.714	-£391	-0.007	£60,062	Abbott 99th centile	£41	0.004	£10,692			
Beckman 99th centile	£2,284	11.716	-£367	-0.004	£86,264	Roche 99th centile	£24	0.002	£10,692			
Roche strategy	£2,383	11.717	-£268	-0.004	£70,874	Beckman 99th centile	£99	0.000	Extendedly dominated			
Abbott strategy	£2,448	11.719	-£203	-0.001	£182,781	Beckman 99th centile	£164	0.003	£52,200			
Standard troponin	£2,651	11.721				Abbott strategy	£203	0.001	£182,781			
Mortality (30-day) treated AMI = 0	0.074	•										
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>			
Abbott 99th centile	£2,295	11.758	-£448	-0.020	£22,206							
Roche 99th centile	£2,342	11.765	-£401	-0.013	£31,543	Abbott 99th centile	£47	0.007	Extendedly dominated			

Beckman 99th centile	£2,369	11.770	-£374	-0.008	£44,952	Abbott 99th centile	£75	0.012	£6,277
Roche strategy	£2,469	11.771	-£274	-0.007	£37,076	Beckman 99th centile	£99	0.001	Extendedly dominated
Abbott strategy	£2,538	11.776	-£205	-0.002	£94,345	Beckman 99th centile	£169	0.006	£27,519
Standard troponin	£2,743	11.778				Abbott strategy	£205	0.002	£94,345

# Mortality (30-day) untreated AMI (decision tree)

Mortality (30-day) untreated AMI = 0.240												
Strategy			Com	pared to St	tandard troponin	Compared	l to next b	est strateg	ÿ			
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>			
Abbott 99th centile	£2,227	11.707	-£470	-0.042	£11,153							
Roche 99th centile	£2,282	11.723	-£415	-0.027	£15,623	Abbott 99th centile	£55	0.016	Extendedly dominated			
Beckman 99th centile	£2,314	11.732	-£383	-0.017	£22,042	Abbott 99th centile	£88	0.025	£3,528			
Roche strategy	£2,414	11.734	-£282	-0.015	£18,271	Beckman 99th centile	£100	0.002	Extendedly dominated			
Abbott strategy	£2,490	11.745	-£207	-0.005	£45,686	Beckman 99th centile	£176	0.013	£13,697			
Standard troponin	£2,697	11.749				Abbott strategy	£207	0.005	£45,686			
Mortality (30-day) untreated AMI	= 0.000											
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>			
Abbott 99th centile	£2,280	11.755	-£417	0.006	Dominant							
Roche 99th centile	£2,316	11.753	-£381	0.004	Dominant	Abbott 99th centile	£35	-0.002	Dominated			
Beckman 99th centile	£2,336	11.752	-£361	0.002	Dominant	Abbott 99th centile	£56	-0.003	Dominated			
Roche strategy	£2,434	11.751	-£263	0.002	Dominant	Abbott 99th centile	£154	-0.004	Dominated			
Abbott strategy	£2,496	11.750	-£201	0.001	Dominant	Abbott 99th centile	£215	-0.005	Dominated			
Standard troponin	£2,697	11.749				Abbott 99th centile	£417	-0.006	Dominated			

Annual re-infarction (after initial AMI) = 0.26												
Strategy			Com	pared to St	tandard troponin	Compare	d to next k	pest strategy	1			
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs			
Abbott 99th centile	£2,286	11.722	-£440	-0.015	£28,543							
Roche 99th centile	£2,330	11.728	-£397	-0.010	£40,757	Abbott 99th centile	£44	0.006	£7,704			
Beckman 99th centile	£2,356	11.731	-£371	-0.006	£58,299	Roche 99th centile	£26	0.003	£7,704			
Roche strategy	£2,455	11.732	-£272	-0.006	£47,995	Beckman 99th centile	£99	0.001	Extendedly dominated			
Abbott strategy	£2,523	11.736	-£204	-0.002	£122,916	Beckman 99th centile	£167	0.005	£35,493			
Standard troponin	£2,727	11.737				Abbott strategy	£204	0.002	£122,916			
Annual re-infarction (after initial A	MI) = 0.19	)										
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>			
Abbott 99th centile	£2,227	11.746	-£440	-0.015	£29,218							
Roche 99th centile	£2,270	11.752	-£396	-0.009	£41,738	Abbott 99th centile	£44	0.006	Extendedly dominated			
Beckman 99th centile	£2,296	11.755	-£370	-0.006	£59,719	Abbott 99th centile	£70	0.009	£7,856			
Roche strategy	£2,395	11.756	-£271	-0.006	£49,157	Beckman 99th centile	£99	0.001	Extendedly dominated			
Abbott strategy	£2,463	11.760	-£204	-0.002	£125,955	Beckman 99th centile	£167	0.005	£36,342			
Standard troponin	£2,666	11.761				Abbott strategy	£204	0.002	£125,955			

#### Annual re-infarction probability (after initial AMI)

# RR re-infarction (untreated versus treated)

RR re-infarction (untreated versus treated) = 5.15												
Strategy			Comp	pared to Sta	andard troponin	Compared to next best strategy						
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>			
Abbott 99th centile	£2,259	11.730	-£438	-0.019	£22,555							
Roche 99th centile	£2,302	11.737	-£395	-0.012	£32,258	Abbott 99th centile	£43	0.007	£5,999			
Beckman 99th centile	£2,327	11.741	-£370	-0.008	£46,195	Roche 99th centile	£25	0.004	£5,999			
Roche strategy	£2,426	11.742	-£271	-0.007	£38,009	Beckman 99th centile	£99	0.001	Extendedly dominated			
Abbott strategy	£2,493	11.747	-£204	-0.002	£97,530	Beckman 99th centile	£166	0.006	£28,076			
Standard troponin	£2,697	11.749				Abbott strategy	£204	0.002	£97,530			
RR re-infarction (untreated versus	treated) =	: 1.28										
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	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>			
Abbott 99th centile	£2,256	11.736	-£441	-0.013	£33,518							
Roche 99th centile	£2,300	11.741	-£397	-0.008	£47,838	Abbott 99th centile	£44	0.005	Extendedly dominated			
Beckman 99th centile	£2,326	11.744	-£371	-0.005	£68,404	Abbott 99th centile	£70	0.008	£9,086			
Roche strategy	£2,425	11.744	-£272	-0.005	£56,324	Beckman 99th centile	£99	0.001	Extendedly dominated			
Abbott strategy	£2,493	11.748	-£204	-0.001	£144,162	Beckman 99th centile	£167	0.004	£41,666			
Standard troponin	£2,697	11.749				Abbott strategy	£204	0.001	£144,162			

### Annual post-MI mortality

Annual post-MI mortality = 0.068										
Strategy			Comp	pared to Sta	andard troponin	Compared to next best strategy				
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	
Abbott 99th centile	£2,248	11.715	-£440	-0.015	£28,843					
Roche 99th centile	£2,292	11.721	-£396	-0.010	£41,191	Abbott 99th centile	£44	0.006	£7,777	
Beckman 99th centile	£2,318	11.724	-£370	-0.006	£58,924	Roche 99th centile	£26	0.003	£7,777	
Roche strategy	£2,417	11.725	-£271	-0.006	£48,508	Beckman 99th centile	£99	0.001	Extendedly dominated	
Abbott strategy	£2,485	11.729	-£204	-0.002	£124,247	Beckman 99th centile	£167	0.005	£35,869	
Standard troponin	£2,688	11.731				Abbott strategy	£204	0.002	£124,247	
Annual post-MI mortality = 0.065										
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	
Abbott 99th centile	£2,266	11.753	-£440	-0.015	£28,897					
Roche 99th centile	£2,309	11.758	-£396	-0.010	£41,275	Abbott 99th centile	£44	0.006	£7,777	
Beckman 99th centile	£2,335	11.762	-£370	-0.006	£59,053	Roche 99th centile	£26	0.003	£7,777	
Roche strategy	£2,434	11.762	-£271	-0.006	£48,610	Beckman 99th centile	£99	0.001	Extendedly dominated	
Abbott strategy	£2,502	11.766	-£204	-0.002	£124,538	Beckman 99th centile	£167	0.005	£35,940	
Standard troponin	£2,706	11.768				Abbott strategy	£204	0.002	£124,538	

Annual mortality post-MI with re-	nnual mortality post-MI with re-infarction = 0.137									
Strategy			Compa	ared to Sta	ndard troponin	Compare	ed to next l	best strategy	1	
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	
Abbott 99th centile	£2,258	11.737	-£440	-0.015	£28,946					
Roche 99th centile	£2,302	11.742	-£396	-0.010	£41,341	Abbott 99th centile	£44	0.006	£7,797	
Beckman 99th centile	£2,328	11.746	-£370	-0.006	£59,144	Roche 99th centile	£26	0.003	£7,797	
Roche strategy	£2,427	11.746	-£271	-0.006	£48,687	Beckman 99th centile	£99	0.001	Extendedly dominated	
Abbott strategy	£2,494	11.750	-£204	-0.002	£124,721	Beckman 99th centile	£167	0.005	£35,999	
Standard troponin	£2,698	11.752				Abbott strategy	£204	0.002	£124,721	
Annual mortality post-MI with re-	infarction	= 0.146								
	Costs	QALYs	<b>∆Costs</b>	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	
Abbott 99th centile	£2,256	11.731	-£440	-0.015	£28,795					
Roche 99th centile	£2,300	11.737	-£396	-0.010	£41,126	Abbott 99th centile	£44	0.006	£7,758	
Beckman 99th centile	£2,325	11.740	-£370	-0.006	£58,835	Roche 99th centile	£26	0.003	£7,758	
Roche strategy	£2,424	11.741	-£271	-0.006	£48,433	Beckman 99th centile	£99	0.001	Extendedly dominated	
Abbott strategy	£2,492	11.745	-£204	-0.002	£124,067	Beckman 99th centile	£167	0.005	£35,812	
Standard troponin	£2,696	11.746				Abbott strategy	£204	0.002	£124,067	

# Annual mortality post-MI after re-infarction

## HR mortality (UA versus NSTEMI)

IR mortality (UA versus NSTEMI) = 1.053									
Strategy			Compa	red to Sta	ndard troponin	Compared to next best strategy			
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,205	11.558	-£440	-0.015	£28,870				
Roche 99th centile	£2,249	11.564	-£396	-0.010	£41,233	Abbott 99th centile	£44	0.006	Extendedly dominated
Beckman 99th centile	£2,274	11.567	-£370	-0.006	£58,988	Abbott 99th centile	£70	0.009	£7,777
Roche strategy	£2,374	11.568	-£271	-0.006	£48,559	Beckman 99th centile	£99	0.001	Extendedly dominated

Abbott strategy	£2,441	11.572	-£204	-0.002	£124,391	Beckman 99th centile	£167	0.005	£35,904
Standard troponin	£2,645	11.573				Abbott strategy	£204	0.002	£124,391
HR mortality (UA versus NSTEMI)	= 0.581								
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,306	11.898	-£440	-0.015	£28,870				
Roche 99th centile	£2,349	11.904	-£396	-0.010	£41,233	Abbott 99th centile	£44	0.006	£7,777
Beckman 99th centile	£2,375	11.907	-£370	-0.006	£58,988	Roche 99th centile	£26	0.003	£7,777
Roche strategy	£2,474	11.908	-£271	-0.006	£48,559	Beckman 99th centile	£99	0.001	Extendedly dominated
Abbott strategy	£2,542	11.912	-£204	-0.002	£124,391	Beckman 99th centile	£167	0.005	£35,904
Standard troponin	£2,746	11.913				Abbott strategy	£204	0.002	£124,391

# RR mortality (untreated versus treated AMI)

RR mortality (untreated versus tre	R mortality (untreated versus treated AMI) = 3.908										
Strategy			Compa	red to Sta	ndard troponin	Compared to next best strategy					
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>		
Abbott 99th centile	£2,224	11.709	-£472	-0.040	£11,771						
Roche 99th centile	£2,280	11.724	-£417	-0.025	£16,467	Abbott 99th centile	£56	0.015	£3,759		
Beckman 99th centile	£2,313	11.733	-£384	-0.017	£23,212	Roche 99th centile	£33	0.009	£3,759		
Roche strategy	£2,414	11.734	-£283	-0.015	£19,250	Beckman 99th centile	£100	0.002	Extendedly dominated		
Abbott strategy	£2,490	11.745	-£207	-0.004	£48,054	Beckman 99th centile	£176	0.012	£14,443		
Standard troponin	£2,697	11.749				Abbott strategy	£207	0.004	£48,054		
RR mortality (untreated versus tre	ated AMI	= 0.901									
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>		
Abbott 99th centile	£2,272	11.746	-£425	-0.003	£128,875						
Roche 99th centile	£2,310	11.747	-£387	-0.002	£186,080	Abbott 99th centile	£38	0.001	Extendedly dominated		
Beckman 99th centile	£2,333	11.748	-£364	-0.001	£268,237	Abbott 99th centile	£61	0.002	£31,275		

Roche strategy	£2,431	11.748	-£266	-0.001	£219,979	Beckman 99th centile	£98	0.000	Extendedly dominated
Abbott strategy	£2,495	11.749	-£202	0.000	£570,869	Beckman 99th centile	£162	0.001	£161,425
Standard troponin	£2,697	11.749				Abbott strategy	£202	0.000	£570,869

### APPENDIX 6: SENSITIVITY ANALYSES (SECONDARY ANALYSIS)

#### Deterministic secondary analysis:

Deterministic secondary analysis									
Strategy			Comp	ared to Sta	ndard troponin	Compared to next best strategy			
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,789	11.530	-£276	0.036	Dominant				
Roche 99th centile	£2,832	11.532	-£232	0.039	Dominant	Abbott 99th centile	£44	0.002	Extendedly dominated
Beckman 99th centile	£2,858	11.532	-£206	0.039	Dominant	Abbott 99th centile	£69	0.003	Extendedly dominated
Roche strategy	£2,957	11.535	-£107	0.042	Dominant	Abbott 99th centile	£168	0.006	Extendedly dominated
Abbott strategy	£3,025	11.543	-£39	0.050	Dominant	Abbott 99th centile	£236	0.014	£17,047
Standard troponin	£3,064	11.493				Abbott strategy	£39	-0.050	Dominated

### Increased re-infarction & mortality risk for no treatment (vs treated) = lifetime (instead of only during the first year after presentation at ED)

Strategy			Compared to Standard troponin			Compared to next best strategy			
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,789	11.473	-£286	0.089	Dominant				
Roche 99th centile	£2,833	11.496	-£242	0.113	Dominant	Abbott 99th centile	£43	0.023	£1,853
Beckman 99th centile	£2,858	11.509	-£217	0.126	Dominant	Roche 99th centile	£26	0.013	£2,017
Roche strategy	£2,957	11.515	-£118	0.131	Dominant	Beckman 99th centile	£99	0.006	Extendedly dominated
Abbott strategy	£3,025	11.537	-£50	0.154	Dominant	Beckman 99th centile	£167	0.028	£5,889
Standard troponin	£3,075	11.383				Abbott strategy	£50	-0.154	Dominated

Waiting time for doctor / decision	pending d	lelay = 1 ł	nour(s)						
Strategy			Comp	ared to Sta	andard troponin	Compare	ed to next l	best strategy	/
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs
Abbott 99th centile	£2,817	11.530	-£275	0.036	Dominant				
Roche 99th centile	£2,861	11.532	-£232	0.039	Dominant	Abbott 99th centile	£44	0.002	£17,587
Beckman 99th centile	£2,887	11.532	-£206	0.039	Dominant	Roche 99th centile	£26	0.000	Extendedly dominated
Roche strategy	£3,002	11.535	-£91	0.042	Dominant	Roche 99th centile	£141	0.003	Extendedly dominated
Abbott strategy	£3,073	11.543	-£20	0.050	Dominant	Roche 99th centile	£212	0.011	£18,628
Standard troponin	£3,093	11.493				Abbott strategy	£20	-0.050	Dominated
Waiting time for doctor / decision	pending d	lelay = 2 ł	nour(s)						
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,846	11.530	-£275	0.036	Dominant				
Roche 99th centile	£2,890	11.532	-£232	0.039	Dominant	Abbott 99th centile	£44	0.002	£17,586
Beckman 99th centile	£2,915	11.532	-£206	0.039	Dominant	Roche 99th centile	£26	0.000	Extendedly dominated
Roche strategy	£3,047	11.535	-£74	0.042	Dominant	Roche 99th centile	£157	0.003	Extendedly dominated
Abbott strategy	£3,121	11.543	£0	0.050	Dominant	Roche 99th centile	£232	0.011	£20,326
Standard troponin	£3,121	11.493				Abbott strategy	£0	-0.050	Dominated
Waiting time for doctor / decision	pending d	lelay = 3 ł	nour(s)						
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,875	11.530	-£275	0.036	Dominant				
Roche 99th centile	£2,918	11.532	-£231	0.039	Dominant	Abbott 99th centile	£44	0.002	£17,584
Beckman 99th centile	£2,944	11.532	-£206	0.039	Dominant	Roche 99th centile	£26	0.000	Extendedly dominated
Roche strategy	£3,092	11.535	-£58	0.042	Dominant	Roche 99th centile	£174	0.003	Extendedly dominated
Standard troponin	£3,149	11.493				Roche 99th centile	£231	-0.039	Dominated
Abbott strategy	£3,169	11.543	£20	0.050	£390	Roche 99th centile	£251	0.011	£22,024

### No doctor on demand, but average waiting time before doctor becomes available is increased with 1, 2 or 3 hours

Doctor on demand at ED, but average waiting time before doctor becomes available in the general ward is increased with 1, 2 or 3 hours (discharge to general ward after 4 hours after presenting at ED)

Waiting time for doctor / decision	pending d	lelay = 1 h	nour(s)						
Strategy			Comp	ared to Sta	indard troponin	Compare	ed to next <b>b</b>	pest strategy	Y
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,790	11.530	-£303	0.036	Dominant				
Roche 99th centile	£2,834	11.532	-£259	0.039	Dominant	Abbott 99th centile	£44	0.002	£17,587
Beckman 99th centile	£2,859	11.532	-£234	0.039	Dominant	Roche 99th centile	£26	0.000	Extendedly dominated
Roche strategy	£2,975	11.535	-£118	0.042	Dominant	Roche 99th centile	£141	0.003	Extendedly dominated
Abbott strategy	£3,046	11.543	-£47	0.050	Dominant	Roche 99th centile	£212	0.011	£18,628
Standard troponin	£3,093	11.493				Abbott strategy	£47	-0.050	Dominated
Waiting time for doctor / decision	pending d	lelay = 2 h	nour(s)						
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,791	11.530	-£330	0.036	Dominant				
Roche 99th centile	£2,835	11.532	-£286	0.039	Dominant	Abbott 99th centile	£44	0.002	£17,586
Beckman 99th centile	£2,860	11.532	-£261	0.039	Dominant	Roche 99th centile	£26	0.000	Extendedly dominated
Roche strategy	£2,992	11.535	-£129	0.042	Dominant	Roche 99th centile	£157	0.003	Extendedly dominated
Abbott strategy	£3,066	11.543	-£55	0.050	Dominant	Roche 99th centile	£232	0.011	£20,326
Standard troponin	£3,121	11.493				Abbott strategy	£55	-0.050	Dominated
Waiting time for doctor / decision	pending d	lelay = 3 h	nour(s)						
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,792	11.530	-£357	0.036	Dominant				
Roche 99th centile	£2,836	11.532	-£313	0.039	Dominant	Abbott 99th centile	£44	0.002	£17,584
Beckman 99th centile	£2,862	11.532	-£288	0.039	Dominant	Roche 99th centile	£26	0.000	Extendedly dominated
Roche strategy	£3,010	11.535	-£140	0.042	Dominant	Roche 99th centile	£174	0.003	Extendedly dominated
Abbott strategy	£3,087	11.543	-£63	0.050	Dominant	Roche 99th centile	£251	0.011	£22,024
Standard troponin	£3,149	11.493				Abbott strategy	£63	-0.050	Dominated

Total delay of	of 1.5 hours
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Strategy			Compared to Standard troponin			Compared to next best strategy				
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	
Abbott 99th centile	£2,746	11.530	-£276	0.036	Dominant					
Roche 99th centile	£2,790	11.532	-£232	0.039	Dominant	Abbott 99th centile	£44	0.002	Extendedly dominated	
Beckman 99th centile	£2,815	11.532	-£207	0.039	Dominant	Abbott 99th centile	£69	0.003	Extendedly dominated	
Roche strategy	£2,890	11.535	-£132	0.042	Dominant	Abbott 99th centile	£144	0.006	Extendedly dominated	
Abbott strategy	£2,953	11.543	-£69	0.050	Dominant	Abbott 99th centile	£207	0.014	£14,956	
Standard troponin	£3,022	11.493				Abbott strategy	£69	-0.050	Dominated	

# MI treatment costs added for patients that were tested false positive

Strategy			Compa	ared to Sta	ndard troponin	Compared to next best strategy				
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	
Abbott 99th centile	£2,841	11.530	-£224	0.036	Dominant					
Abbott strategy	£3,056	11.543	-£9	0.050	Dominant	Abbott 99th centile	£215	0.014	£15,507	
Standard troponin	£3,064	11.493	£0	0.000		Abbott strategy	£9	-0.050	Dominated	
Roche 99th centile	£3,144	11.532	£80	0.039	£2,065	Abbott strategy	£89	-0.011	Dominated	
Roche strategy	£3,331	11.535	£267	0.042	£6,360	Abbott strategy	£275	-0.008	Dominated	
Beckman 99th centile	£3,421	11.532	£356	0.039	£9,142	Abbott strategy	£365	-0.011	Dominated	

# MI treatment costs added to first year of UA

Strategy			Compa	ared to Sta	ndard troponin	Compared to next best strategy				
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	
Abbott 99th centile	£3,212	11.530	-£275	0.036	Dominant					
Roche 99th centile	£3,256	11.532	-£231	0.039	Dominant	Abbott 99th centile	£43	0.002	Extendedly dominated	
Beckman 99th centile	£3,281	11.532	-£205	0.039	Dominant	Abbott 99th centile	£69	0.003	Extendedly dominated	
Roche strategy	£3,381	11.535	-£106	0.042	Dominant	Abbott 99th centile	£168	0.006	Extendedly dominated	
Abbott strategy	£3,449	11.543	-£38	0.050	Dominant	Abbott 99th centile	£236	0.014	£17,048	
Standard troponin	£3,487	11.493				Abbott strategy	£38	-0.050	Dominated	

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Test costs = £5									
Strategy			Comp	ared to Sta	andard troponin	Compared to next best strategy			
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,772	11.530	-£260	0.036	Dominant				
Roche 99th centile	£2,816	11.532	-£217	0.039	Dominant	Abbott 99th centile	£44	0.002	Extendedly dominated
Beckman 99th centile	£2,841	11.532	-£191	0.039	Dominant	Abbott 99th centile	£69	0.003	Extendedly dominated
Roche strategy	£2,931	11.535	-£101	0.042	Dominant	Abbott 99th centile	£159	0.006	Extendedly dominated
Abbott strategy	£2,997	11.543	-£35	0.050	Dominant	Abbott 99th centile	£225	0.014	£16,260
Standard troponin	£3,032	11.493				Abbott strategy	£35	-0.050	Dominated
Test costs = £40									
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	<b>∆Costs</b>	ΔQALYs	ΔCosts / ΔQALYs
Abbott 99th centile	£2,810	11.530	-£295	0.036	Dominant				
Roche 99th centile	£2,854	11.532	-£252	0.039	Dominant	Abbott 99th centile	£44	0.002	£17,586
Beckman 99th centile	£2,879	11.532	-£226	0.039	Dominant	Roche 99th centile	£26	0.000	Extendedly dominated
Roche strategy	£2,990	11.535	-£115	0.042	Dominant	Roche 99th centile	£137	0.003	Extendedly dominated
Abbott strategy	£3,060	11.543	-£45	0.050	Dominant	Roche 99th centile	£207	0.011	£18,141
Standard troponin	£3,105	11.493				Abbott strategy	£45	-0.050	Dominated

#### AMI treatment costs

AMI treatment costs = £2,577										
Strategy			Comp	pared to S	tandard troponin	Compared to next best strategy				
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	
Abbott 99th centile	£2,607	11.530	-£286	0.036	Dominant					
Roche 99th centile	£2,641	11.532	-£252	0.039	Dominant	Abbott 99th centile	£34	0.002	£13,770	
Beckmann 99th centile	£2,661	11.532	-£232	0.039	Dominant	Roche 99th centile	£20	0.000	Extendedly dominated	
Roche strategy	£2,759	11.535	-£134	0.042	Dominant	Roche 99th centile	£118	0.003	Extendedly dominated	
Abbott strategy	£2,820	11.543	-£73	0.050	Dominant	Roche 99th centile	£179	0.011	£15,751	

Standard troponin	£2,893	11.493				Abbott strategy	£73	-0.050	Dominated			
AMI treatment costs = £4,295												
	Costs	QALYs	∆Costs	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs			
Abbott 99th centile	£2,971	11.530	-£265	0.036	Dominant							
Roche 99th centile	£3,024	11.532	-£212	0.039	Dominant	Abbott 99th centile	£53	0.002	Extendedly dominated			
Beckmann 99th centile	£3,055	11.532	-£181	0.039	Dominant	Abbott 99th centile	£84	0.003	Extendedly dominated			
Roche strategy	£3,155	11.535	-£81	0.042	Dominant	Abbott 99th centile	£185	0.006	Extendedly dominated			
Abbott strategy	£3,230	11.543	-£6	0.050	Dominant	Abbott 99th centile	£259	0.014	£18,698			
Standard troponin	£3,236	11.493	;			Abbott strategy	£6	-0.050	Dominated			
Post-MI health state costs												
Post-MI health state costs (1st year) = £6,791												
Strategy			Comp	ared to St	andard troponin	Compared to next best strategy						
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>			
Abbott 99th centile	£2,970	11.530	-£276	0.036	Dominant							
Roche 99th centile	£3,014	11.532	-£231	0.039	Dominant	Abbott 99th centile	£44	0.002	Extendedly dominated			
Beckman 99th centile	£3,040	11.532	-£205	0.039	Dominant	Abbott 99th centile	£71	0.003	Extendedly dominated			
Roche strategy	£3,139	11.535	-£106	0.042	Dominant	Abbott 99th centile	£170	0.006	Extendedly dominated			
Abbott strategy	£3,208	11.543	-£37	0.050	Dominant	Abbott 99th centile	£239	0.014	£17,199			
Standard troponin	£3,245	11.493				Abbott strategy	£37	-0.050	Dominated			
Post-MI health state costs (1st ye	ar) = £4,8	79										
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	<b>∆Costs</b>	ΔQALYs	ΔCosts / ΔQALYs			
Abbott 99th centile	£2,608	11.530	-£275	0.036	Dominant							
Roche 99th centile	£2,651	11.532	-£233	0.039	Dominant	Abbott 99th centile	£43	0.002	Extendedly dominated			
Beckman 99th centile	£2,676	11.532	-£207	0.039	Dominant	Abbott 99th centile	£68	0.003	Extendedly dominated			
Roche strategy	£2,775	11.535	-£108	0.042	Dominant	Abbott 99th centile	£167	0.006	Extendedly dominated			
Abbott strategy	£2,842	11.543	-£41	0.050	Dominant	Abbott 99th centile	£234	0.014	£16,896			
Standard troponin	£2,883	11.493				Abbott strategy	£41	-0.050	Dominated			

### Utility difference between UA and AMI

Utility difference between UA and AMI = 0.12												
Strategy			Comp	pared to S	tandard troponin	Compared to next best strategy						
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>			
Abbott 99th centile	£2,789	11.572	-£276	0.036	Dominant							
Roche 99th centile	£2,832	11.575	-£232	0.039	Dominant	Abbott 99th centile	£44	0.002	Extendedly dominated			
Beckman 99th centile	£2,858	11.575	-£206	0.039	Dominant	Abbott 99th centile	£69	0.003	Extendedly dominated			
Roche strategy	£2,957	11.578	-£107	0.042	Dominant	Abbott 99th centile	£168	0.006	Extendedly dominated			
Abbott strategy	£3,025	11.586	-£39	0.050	Dominant	Abbott 99th centile	£236	0.014	£17,046			
Standard troponin	£3,064	11.536				Abbott strategy	£39	-0.050	Dominated			
Utility difference between UA and	nd AMI = -	0.10										
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>			
Abbott 99th centile	£2,789	11.385	-£276	0.036	Dominant							
Roche 99th centile	£2,832	11.387	-£232	0.038	Dominant	Abbott 99th centile	£44	0.003	Extendedly dominated			
Beckman 99th centile	£2,858	11.388	-£206	0.039	Dominant	Abbott 99th centile	£69	0.003	Extendedly dominated			
Roche strategy	£2,957	11.391	-£107	0.042	Dominant	Abbott 99th centile	£168	0.006	Extendedly dominated			
Abbott strategy	£3,025	11.399	-£39	0.050	Dominant	Abbott 99th centile	£236	0.014	£17,051			
Standard troponin	£3,064	11.349				Abbott strategy	£39	-0.050	Dominated			

#### **MI disutility**

MI disutility = -0.059 (age = 45);	MI disutility = -0.059 (age = 45); -0.050 (age = 55); -0.024 (age = 65); -0.006 (age = 75+)													
Strategy			Comp	ared to St	andard troponin	Compared to next best strategy								
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>					
Abbott 99th centile	£2,789	11.531	-£276	0.036	Dominant									
Roche 99th centile	£2,832	11.534	-£232	0.039	Dominant	Abbott 99th centile	£44	0.002	Extendedly dominated					
Beckman 99th centile	£2,858	11.534	-£206	0.039	Dominant	Abbott 99th centile	£69	0.003	Extendedly dominated					
Roche strategy	£2,957	11.537	-£107	0.042	Dominant	Abbott 99th centile	£168	0.006	Extendedly dominated					
Abbott strategy	£3,025	11.545	-£39	0.050	Dominant	Abbott 99th centile	£236	0.014	£17,025					
Standard troponin	£3,064	11.495				Abbott strategy	£39	-0.050	Dominated					

MI disutility = -0.061(age = 45); -0.052 (age = 55); -0.026 (age = 65); -0.008 (age = 75+)												
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs			
Abbott 99th centile	£2,789	11.528	-£276	0.036	Dominant							
Roche 99th centile	£2,832	11.530	-£232	0.039	Dominant	Abbott 99th centile	£44	0.002	Extendedly dominated			
Beckman 99th centile	£2,858	11.531	-£206	0.039	Dominant	Abbott 99th centile	£69	0.003	Extendedly dominated			
Roche strategy	£2,957	11.534	-£107	0.042	Dominant	Abbott 99th centile	£168	0.006	Extendedly dominated			
Abbott strategy	£3,025	11.542	-£39	0.050	Dominant	Abbott 99th centile	£236	0.014	£17,070			
Standard troponin	£3,064	11.492				Abbott strategy	£39	-0.050	Dominated			

# Mortality (30-day) treated AMI (decision tree)

Mortality (30-day) treated AMI = 0.120												
Strategy			Comp	ared to Sta	indard troponin	Compared to next best strategy						
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>			
Abbott 99th centile	£2,750	11.504	-£269	0.039	Dominant							
Roche 99th centile	£2,790	11.504	-£228	0.039	Dominant	Abbott 99th centile	£41	0.000	Dominated			
Beckman 99th centile	£2,814	11.502	-£205	0.038	Dominant	Abbott 99th centile	£64	-0.002	Dominated			
Roche strategy	£2,913	11.506	-£106	0.041	Dominant	Abbott 99th centile	£163	0.002	Extendedly dominated			
Abbott strategy	£2,979	11.514	-£40	0.049	Dominant	Abbott 99th centile	£229	0.010	£24,010			
Standard troponin	£3,019	11.465				Abbott strategy	£40	-0.049	Dominated			
Mortality (30-day) treated AM	l = 0.074											
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>			
Abbott 99th centile	£2,828	11.555	-£282	0.033	Dominant							
Roche 99th centile	£2,875	11.560	-£236	0.038	Dominant	Abbott 99th centile	£47	0.005	£9,175			
Beckman 99th centile	£2,902	11.562	-£208	0.040	Dominant	Roche 99th centile	£28	0.002	£12,967			
Roche strategy	£3,002	11.565	-£109	0.043	Dominant	Beckman 99th centile	£100	0.003	Extendedly dominated			
Abbott strategy	£3,072	11.574	-£39	0.051	Dominant	Beckman 99th centile	£170	0.011	£15,399			
Standard troponin	£3,111	11.522				Abbott strategy	£39	-0.051	Dominated			

Mortality (30-day) untreated AMI = 0.240												
Strategy			Comp	ared to Sta	indard troponin	Compared to next best strategy						
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>			
Abbott 99th centile	£2,759	11.503	-£294	0.066	Dominant							
Roche 99th centile	£2,813	11.515	-£239	0.079	Dominant	Abbott 99th centile	£55	0.012	£4,404			
Beckman 99th centile	£2,846	11.521	-£207	0.085	Dominant	Roche 99th centile	£32	0.006	£5,228			
Roche strategy	£2,946	11.525	-£106	0.089	Dominant	Beckman 99th centile	£101	0.004	Extendedly dominated			
Abbott strategy	£3,022	11.541	-£30	0.104	Dominant	Beckman 99th centile	£176	0.019	£9,139			
Standard troponin	£3,052	11.436				Abbott strategy	£30	-0.104	Dominated			
Mortality (30-day) untreated A	MI = 0.00	0										
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>			
Abbott 99th centile	£2,813	11.551	-£262	0.013	Dominant							
Roche 99th centile	£2,847	11.545	-£227	0.007	Dominant	Abbott 99th centile	£35	-0.005	Dominated			
Beckman 99th centile	£2,868	11.541	-£206	0.003	Dominant	Abbott 99th centile	£55	-0.010	Dominated			
Roche strategy	£2,966	11.543	-£108	0.005	Dominant	Abbott 99th centile	£153	-0.008	Dominated			
Abbott strategy	£3,028	11.546	-£46	0.008	Dominant	Abbott 99th centile	£215	-0.005	Dominated			
Standard troponin	£3,074	11.538				Abbott 99th centile	£262	-0.013	Dominated			

# Mortality (30-day) untreated AMI (decision tree)

#### Annual re-infarction probability (after initial AMI)

Annual re-infarction (after initial AMI) = 0.26													
Strategy			Compa	ared to Sta	indard troponin	Comp	pared to next	best strate	gy				
	Costs	QALYs	ΔCosts	sts ΔQALYs ΔCosts / ΔQALYs Comparator ΔCosts ΔQALYs ΔCosts / ΔQAL									
Abbott 99th centile	£2,830	11.515	-£275	0.036	Dominant								
Roche 99th centile	£2,873	11.517	-£232	0.039	Dominant	Abbott 99th centile	£44	0.003	Extendedly dominated				
Beckman 99th centile	£2,899	11.517	-£206	0.039	Dominant	Abbott 99th centile	£69	0.003	Extendedly dominated				
Roche strategy	£2,998	11.520	-£107	0.042	Dominant	Abbott 99th centile	£169	0.006	Extendedly dominated				
Abbott strategy	£3,066	11.529	-£39	£39 0.050 Dominant Abbott 99th centile £237 0.014 £16,867									
Standard troponin	£3,105	£3,105 11.478 Abbott strategy £39 -0.050 Dominated											

Annual re-infarction (after initial AMI) = 0.19													
Costs QALYs ΔCosts ΔQALYs ΔCosts ΔCosts ΔQALYs ΔCosts ΔCosts ΔQALYs													
Abbott 99th centile	£2,747	11.545	-£276	0.036	Dominant								
Roche 99th centile	£2,791	11.547	-£233	0.038	Dominant	Abbott 99th centile	£43	0.002	Extendedly dominated				
Beckman 99th centile	£2,817	11.548	-£207	0.039	Dominant	Abbott 99th centile	£69	0.003	Extendedly dominated				
Roche strategy	£2,916	11.551	-£108	0.042	Dominant	Abbott 99th centile	£168	0.006	Extendedly dominated				
Abbott strategy	£2,984	11.559	-£40	0.050	Dominant	Abbott 99th centile	£236	0.014	£17,241				
Standard troponin	£3,023	11.509				Abbott strategy	£40	-0.050	Dominated				

# RR re-infarction (untreated versus treated)

R re-infarction (untreated versus treated) = 5.15													
Strategy			Comp	ared to Sta	indard troponin	Con	npared to nex	t best strate	gy				
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs				
Abbott 99th centile	£2,791	11.525	-£277	0.036	Dominant								
Roche 99th centile	£2,834	11.529	-£234	0.041	Dominant	Abbott 99th centile	£43	0.004	£10,647				
Beckman 99th centile	£2,859	11.531	-£209	0.042	Dominant	Roche 99th centile	£25	0.001	Extendedly dominated				
Roche strategy	£2,958	11.534	-£110	0.045	Dominant	Roche 99th centile	£124	0.004	Extendedly dominated				
Abbott strategy	£3,025	11.543	-£43	0.054	Dominant	Roche 99th centile	£192	0.014	£14,126				
Standard troponin	£3,068	11.489				Abbott strategy	£43	-0.054	Dominated				
RR re-infarction (untreated ver	sus treate	d) = 1.28											
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>				
Abbott 99th centile	£2,788	11.532	-£275	0.036	Dominant								
Roche 99th centile	£2,832	11.533	-£231	0.038	Dominant	Abbott 99th centile	£44	0.002	Extendedly dominated				
Beckman 99th centile	£2,858	11.533	-£205	0.038	Dominant	Abbott 99th centile	£70	0.002	Dominated				
Roche strategy	£2,957	11.536	-£106	0.040	Dominant	Abbott 99th centile	£169	0.004	Extendedly dominated				
Abbott strategy	£3,025	11.544	-£38	0.048	Dominant	Abbott 99th centile	£237	0.012	£19,764				
Standard troponin	£3,063	11.496				Abbott strategy	£38	-0.048	Dominated				

### Annual post-MI mortality

Annual post-MI mortality = 0.068												
Strategy			Comp	ared to Sta	andard troponin	Con	npared to nex	t best strate	gy			
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs			
Abbott 99th centile	£2,779	11.509	-£276	0.036	Dominant							
Roche 99th centile	£2,822	11.511	-£232	0.039	Dominant	Abbott 99th centile	£44	0.002	Extendedly dominated			
Beckman 99th centile	£2,848	11.512	-£206	0.039	Dominant	Abbott 99th centile	£69	0.003	Extendedly dominated			
Roche strategy	£2,947	11.514	-£107	0.042	Dominant	Abbott 99th centile	£169	0.006	Extendedly dominated			
Abbott strategy	£3,015	11.523	-£39	0.050	Dominant	Abbott 99th centile	£237	0.014	£17,036			
Standard troponin	£3,054	11.472				Abbott strategy	£39	-0.050	Dominated			
Annual post-MI mortality = 0.0	65				•							
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs			
Abbott 99th centile	£2,799	11.551	-£276	0.036	Dominant							
Roche 99th centile	£2,843	11.553	-£232	0.039	Dominant	Abbott 99th centile	£44	0.002	Extendedly dominated			
Beckman 99th centile	£2,868	11.553	-£206	0.039	Dominant	Abbott 99th centile	£69	0.003	Extendedly dominated			
Roche strategy	£2,968	11.556	-£107	0.042	Dominant	Abbott 99th centile	£168	0.006	Extendedly dominated			
Abbott strategy	£3,035	11.565	-£39	0.050	Dominant	Abbott 99th centile	£236	0.014	£17,059			
Standard troponin	£3,075	11.515				Abbott strategy	£39	-0.050	Dominated			

#### Annual mortality post-MI after re-infarction

Annual mortality post-MI with re-infarction = 0.137													
Strategy			Compa	ared to Sta	indard troponin	Con	npared to nex	t best strate	3 <b>y</b>				
	Costs	QALYs	ΔCosts	osts ΔQALYs ΔCosts / ΔQALYs Comparator ΔCosts ΔQALYs ΔCosts / ΔQAL									
Abbott 99th centile	£2,790	11.532	-£276	0.036	Dominant								
Roche 99th centile	£2,834	11.535	-£232	0.039	Dominant	Abbott 99th centile	£44	0.002	Extendedly dominated				
Beckman 99th centile	£2,859	11.535	-£206	0.039	Dominant	Abbott 99th centile	£69	0.003	Extendedly dominated				
Roche strategy	£2,958	11.538	-£107	0.042	Dominant	Abbott 99th centile	£168	0.006	Extendedly dominated				
Abbott strategy	£3,026	11.546	-£39 0.050 Dominant Abbott 99th centile £236 0.014 £17,091										
Standard troponin	£3,066	3,066 11.496 Abbott strategy £39 -0.050 Dominated											

Annual mortality post-MI with re-infarction = 0.146													
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs				
Abbott 99th centile	£2,788	11.527	-£276	0.036	Dominant								
Roche 99th centile	£2,831	11.529	-£232	0.039	Dominant	Abbott 99th centile	£44	0.002	Extendedly dominated				
Beckman 99th centile	£2,857	11.530	-£206	0.039	Dominant	Abbott 99th centile	£69	0.003	Extendedly dominated				
Roche strategy	£2,956	11.533	-£107	0.042	Dominant	Abbott 99th centile	£168	0.006	Extendedly dominated				
Abbott strategy	£3,024	11.541	-£39	0.050	Dominant	Abbott 99th centile	£236	0.014	£17,005				
Standard troponin	£3,063	11.491				Abbott strategy	£39	-0.050	Dominated				

# HR mortality (UA versus NSTEMI)

HR mortality (UA versus NSTEMI) = 1.053													
Strategy			Compa	ared to Sta	andard troponin	Con	npared to nex	kt best strate	gy				
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	∆Costs	ΔQALYs	<b>ΔCosts / ΔQALYs</b>				
Abbott 99th centile	£2,740	11.363	-£276	0.036	Dominant								
Roche 99th centile	£2,783	11.365	-£232	0.038	Dominant	Abbott 99th centile	£44	0.003	Extendedly dominated				
Beckman 99th centile	£2,809	11.366	-£206	0.039	Dominant	Abbott 99th centile	£69	0.003	Extendedly dominated				
Roche strategy	£2,908	11.369	-£107	0.042	Dominant	Abbott 99th centile	£168	0.006	Extendedly dominated				
Abbott strategy	£2,976	11.377	-£39	0.050	Dominant	Abbott 99th centile	£236	0.014	£17,051				
Standard troponin	£3,015	11.327				Abbott strategy	£39	-0.050	Dominated				
HR mortality (UA versus NSTEN	vII) = 0.581												
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>				
Abbott 99th centile	£2,835	11.685	-£275	0.037	Dominant								
Roche 99th centile	£2,879	11.688	-£232	0.039	Dominant	Abbott 99th centile	£44	0.002	Extendedly dominated				
Beckman 99th centile	£2,904	11.688	-£206	0.039	Dominant	Abbott 99th centile	£69	0.003	Extendedly dominated				
Roche strategy	£3,003	11.691	-£107	0.042	Dominant	Abbott 99th centile	£168	0.006	Extendedly dominated				
Abbott strategy	£3,071	11.699	-£39	0.050	Dominant	Abbott 99th centile	£236	0.014	£17,044				
Standard troponin	£3,111	11.649				Abbott strategy	£39	-0.050	Dominated				

RR mortality (untreated versus treated AMI)

R mortality (untreated versus treated AMI) = 3.908													
Strategy			Comp	ared to Sta	andard troponin	Con	npared to nex	t best strate	gy				
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs				
Abbott 99th centile	£2,757	11.505	-£274	0.042	Dominant								
Roche 99th centile	£2,812	11.516	-£219	0.053	Dominant	Abbott 99th centile	£56	0.012	£4,755				
Beckman 99th centile	£2,845	11.522	-£186	0.059	Dominant	Roche 99th centile	£33	0.006	£5,714				
Roche strategy	£2,945	11.526	-£85	0.063	Dominant	Beckman 99th centile	£101	0.004	Extendedly dominated				
Abbott strategy	£3,022	11.541	-£9	0.078	Dominant	Beckman 99th centile	£177	0.019	£9,476				
Standard troponin	£3,031	11.463				Abbott strategy	£9	-0.078	Dominated				
RR mortality (untreated versus	treated A	MI) = 0.9	01										
Strategy			Comp	ared to Sta	indard troponin	Con	npared to nex	t best strate	gy				
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs				
Abbott 99th centile	£2,804	11.542	-£276	0.034	Dominant								
Roche 99th centile	£2,842	11.540	-£238	0.032	Dominant	Abbott 99th centile	£38	-0.002	Dominated				
Beckman 99th centile	£2,865	11.537	-£216	0.029	Dominant	Abbott 99th centile	£60	-0.004	Dominated				
Roche strategy	£2,963	11.540	-£118	0.032	Dominant	Abbott 99th centile	£159	-0.002	Dominated				
Abbott strategy	£3,027	11.545	-£54	0.037	Dominant	Abbott 99th centile	£223	0.003	£69,543				
Standard troponin	£3,081	11.508				Abbott strategy	£54	-0.037	Dominated				

# APPENDIX 7: SUBGROUP ANALYSES (BASE CASE)

#### Deterministic base case:

Strategy			Comp	ared to Sta	andard troponin	n Compared to next best strategy			
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,257	11.734	-£440	-0.015	£28,870				
Roche 99th centile	£2,301	11.740	-£396	-0.010	£41,233	Abbott 99th centile	£44	0.006	£7,777
Beckman 99th centile	£2,327	11.743	-£370	-0.006	£58,988	Roche 99th centile	£26	0.003	£7,777
Roche strategy	£2,426	11.744	-£271	-0.006	£48,559	Beckman 99th centile	£99	0.001	Extendedly dominated
Abbott strategy	£2,493	11.748	-£204	-0.002	£124,391	Beckman 99th centile	£167	0.005	£35,904
Standard troponin	£2,697	11.749				Abbott strategy	£204	0.002	£124,391

# Age and gender subgroups:

Females									
Age = 45									
Strategy			Comp	ared to Sta	andard troponin	Сон	npared to ne	ext best stra	tegy
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs
Abbott 99th centile	£2,087	12.853	-£443	-0.016	£27,038				
Roche 99th centile	£2,132	12.859	-£398	-0.010	£38,540	Abbott 99th centile	£45	0.006	£7,414
Beckman 99th centile	£2,158	12.863	-£372	-0.007	£55,060	Roche 99th centile	£27	0.004	£7,414
Roche strategy	£2,258	12.864	-£272	-0.006	£45,357	Beckman 99th centile	£99	0.001	Extendedly dominated
Abbott strategy	£2,326	12.868	-£204	-0.002	£115,910	Beckman 99th centile	£168	0.005	£33,583
Standard troponin	£2,530	12.870				Abbott strategy	£204	0.002	£115,910
Age = 55									
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,093	10.615	-£443	-0.016	£28,189				
Roche 99th centile	£2,138	10.620	-£398	-0.010	£40,181	Abbott 99th centile	£45	0.006	Extendedly dominated
Beckman 99th centile	£2,164	10.624	-£372	-0.006	£57,405	Abbott 99th centile	£71	0.009	£7,728

Roche strategy	£2,263	10.624	-£272	-0.006	£47,288	Beckman 99th centile	£99	0.001	Extendedly dominated
Abbott strategy	£2,332	10.629	-£204	-0.002	£120,850	Beckman 99th centile	£168	0.005	£35,013
Standard troponin	£2,536	10.630				Abbott strategy	£204	0.002	£120,850
Age = 65									
	Costs	QALYs	<b>∆Costs</b>	ΔQALYs	ΔCosts / ΔQALYs	Comparator	∆Costs	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,087	8.193	-£443	-0.015	£29,368				
Roche 99th centile	£2,132	8.199	-£398	-0.010	£41,866	Abbott 99th centile	£45	0.006	Extendedly dominated
Beckman 99th centile	£2,158	8.202	-£372	-0.006	£59,816	Abbott 99th centile	£71	0.009	£8,044
Roche strategy	£2,258	8.203	-£272	-0.006	£49,272	Beckman 99th centile	£99	0.001	Extendedly dominated
Abbott strategy	£2,326	8.207	-£204	-0.002	£125,935	Beckman 99th centile	£167	0.005	£36,479
Standard troponin	£2,530	8.208				Abbott strategy	£204	0.002	£125,935
Age = 75									
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,037	5.640	-£442	-0.013	£32,776				
Roche 99th centile	£2,082	5.645	-£398	-0.009	£46,745	Abbott 99th centile	£45	0.005	Extendedly dominated
Beckman 99th centile	£2,108	5.648	-£371	-0.006	£66,808	Abbott 99th centile	£71	0.008	£8,942
Roche strategy	£2,207	5.649	-£272	-0.005	£55,024	Beckman 99th centile	£99	0.001	Extendedly dominated
Abbott strategy	£2,276	5.652	-£204	-0.001	£140,710	Beckman 99th centile	£167	0.004	£40,725
Standard troponin	£2,480	5.654				Abbott strategy	£204	0.001	£140,710
Age = 85									
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£1,826	3.107	-£437	-0.007	£59,890				
Roche 99th centile	£1,869	3.110	-£394	-0.005	£85,736	Abbott 99th centile	£43	0.003	Extendedly dominated
Beckman 99th centile	£1,894	3.112	-£369	-0.003	£122,857	Abbott 99th centile	£68	0.004	£15,793
Roche strategy	£1,993	3.112	-£270	-0.003	£101,053	Beckman 99th centile	£99	0.000	Extendedly dominated
Abbott strategy	£2,059	3.114	-£203	-0.001	£259,592	Beckman 99th centile	£166	0.002	£74,597

Standard troponin	£2,263	3.115				Abbott strategy	£203	0.001	£259,592
Males									
Age = 45									
	Costs	QALYs	∆Costs	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	∆Costs	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,404	14.047	-£438	-0.015	£28,815				
Roche 99th centile	£2,447	14.053	-£395	-0.010	£41,214	Abbott 99th centile	£43	0.006	£7,660
Beckman 99th centile	£2,472	14.056	-£370	-0.006	£59,021	Roche 99th centile	£25	0.003	£7,660
Roche strategy	£2,571	14.057	-£271	-0.006	£48,561	Beckman 99th centile	£99	0.001	Extendedly dominated
Abbott strategy	£2,638	14.061	-£204	-0.002	£124,616	Beckman 99th centile	£166	0.005	£35,870
Standard troponin	£2,842	14.062				Abbott strategy	£204	0.002	£124,616
Age = 55									
	Costs	QALYs	∆Costs	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,407	11.852	-£438	-0.014	£30,338				
Roche 99th centile	£2,450	11.857	-£395	-0.009	£43,396	Abbott 99th centile	£43	0.005	Extendedly dominated
Beckman 99th centile	£2,476	11.860	-£370	-0.006	£62,149	Abbott 99th centile	£68	0.008	£8,059
Roche strategy	£2,575	11.861	-£271	-0.005	£51,134	Beckman 99th centile	£99	0.001	Extendedly dominated
Abbott strategy	£2,642	11.865	-£204	-0.002	£131,231	Beckman 99th centile	£166	0.004	£37,768
Standard troponin	£2,845	11.866				Abbott strategy	£204	0.002	£131,231
Age = 65									
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,371	9.384	-£438	-0.013	£32,627				
Roche 99th centile	£2,413	9.389	-£395	-0.008	£46,682	Abbott 99th centile	£43	0.005	Extendedly dominated
Beckman 99th centile	£2,439	9.392	-£369	-0.006	£66,867	Abbott 99th centile	£68	0.008	£8,647
Roche strategy	£2,538	9.392	-£270	-0.005	£55,011	Beckman 99th centile	£99	0.001	Extendedly dominated
Abbott strategy	£2,605	9.396	-£203	-0.001	£141,222	Beckman 99th centile	£166	0.004	£40,624
Standard troponin	£2,808	9.397				Abbott strategy	£203	0.001	£141,222

Age = 75									
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,253	6.574	-£437	-0.011	£39,186				
Roche 99th centile	£2,295	6.578	-£394	-0.007	£56,106	Abbott 99th centile	£42	0.004	Extendedly dominated
Beckman 99th centile	£2,320	6.581	-£369	-0.005	£80,406	Abbott 99th centile	£68	0.007	£10,317
Roche strategy	£2,419	6.581	-£270	-0.004	£66,133	Beckman 99th centile	£99	0.001	Extendedly dominated
Abbott strategy	£2,486	6.584	-£203	-0.001	£169,919	Beckman 99th centile	£166	0.003	£48,814
Standard troponin	£2,689	6.585				Abbott strategy	£203	0.001	£169,919
Age = 85									
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£1,940	3.634	-£429	-0.004	£114,585				
Roche 99th centile	£1,980	3.635	-£389	-0.002	£164,917	Abbott 99th centile	£40	0.001	Extendedly dominated
Beckman 99th centile	£2,004	3.636	-£366	-0.002	£237,203	Abbott 99th centile	£63	0.002	£28,711
Roche strategy	£2,102	3.636	-£267	-0.001	£194,744	Beckman 99th centile	£99	0.000	Extendedly dominated
Abbott strategy	£2,167	3.637	-£203	0.000	£503,476	Beckman 99th centile	£163	0.001	£143,225
Standard troponin	£2,369	3.638				Abbott strategy	£203	0.000	£503,476

# Subgroup with history of previous NSTEMI<sup>a</sup>

Strategy			Com	pared to St	andard troponin	Compared to next best strategy			
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£4,643	5.764	-£472	-0.019	£25,031				
Roche 99th centile	£4,699	5.771	-£417	-0.012	£35,017	Abbott 99th centile	£56	0.007	Extendedly dominated
Beckman 99th centile	£4,732	5.775	-£384	-0.008	£49,358	Abbott 99th centile	£89	0.011	£7,994
Roche strategy	£4,834	5.776	-£281	-0.007	£40,639	Beckman 99th centile	£103	0.001	Extendedly dominated
Abbott strategy	£4,910	5.781	-£205	-0.002	£101,225	Beckman 99th centile	£178	0.006	£31,052
Standard troponin	£5,115	5.783				Abbott strategy	£205	0.002	£101,225

<sup>a</sup> Based on an AMI prevalence of 20% (see Appendix 9)

# MI prevalence

MI prevalence = 1%									
Strategy			Com	pared to St	andard troponin	Con	npared to n	ext best stra	ategy
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
No testing	£576	12.891	-£439	-0.005	£96,456				
Abbott 99th centile	£687	12.894	-£329	-0.001	£366,354	No testing	£111	0.004	Extendedly dominated
Roche 99th centile	£690	12.895	-£326	-0.001	£576,522	No testing	£113	0.004	Extendedly dominated
Beckman 99th centile	£691	12.895	-£324	0.000	£878,364	No testing	£115	0.004	£27,409
Roche strategy	£774	12.895	-£241	0.000	£734,155	Beckman 99th centile	£83	0.000	Extendedly dominated
Abbott strategy	£813	12.895	-£202	0.000	£2,097,914	Beckman 99th centile	£122	0.000	£447,934
Standard troponin	£1,016	12.895				Abbott strategy	£202	0.000	£2,097,914
MI prevalence = 5%									
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
No testing	£855	12.586	-£581	-0.023	£25,513				
Abbott 99th centile	£1,079	12.604	-£356	-0.004	£79,492	No testing	£224	0.018	Extendedly dominated
Roche 99th centile	£1,092	12.606	-£344	-0.003	£121,526	No testing	£237	0.020	Extendedly dominated
Beckman 99th centile	£1,100	12.607	-£336	-0.002	£181,894	No testing	£245	0.021	£11,703
Roche strategy	£1,187	12.607	-£249	-0.002	£151,398	Beckman 99th centile	£87	0.000	Extendedly dominated
Abbott strategy	£1,233	12.608	-£203	0.000	£420,420	Beckman 99th centile	£133	0.001	£97,709
Standard troponin	£1,436	12.609				Abbott strategy	£203	0.000	£420,420
MI prevalence = 10%									
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	∆Costs	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
No testing	£1,204	12.205	-£758	-0.046	£16,645				
Abbott 99th centile	£1,570	12.242	-£391	-0.009	£43,635	No testing	£366	0.037	Extendedly dominated
Roche 99th centile	£1,596	12.245	-£366	-0.006	£64,651	No testing	£392	0.040	Extendedly dominated
Beckman 99th centile	£1,611	12.247	-£350	-0.004	£94,836	No testing	£407	0.042	£9,740
Roche strategy	£1,703	12.247	-£258	-0.003	£78,554	Beckman 99th centile	£92	0.000	Extendedly dominated

Abbott strategy	£1,758	12.250	-£203	-0.001	£210,733	Beckman 99th centile	£147	0.003	£53,931
Standard troponin	£1,961	12.251				Abbott strategy	£203	0.001	£210,733
MI prevalence = 20%									
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
No testing	£1,900	11.443	-£1,112	-0.091	£12,211				
Abbott 99th centile	£2,551	11.516	-£461	-0.018	£25,706	No testing	£651	0.073	Extendedly dominated
Roche 99th centile	£2,603	11.523	-£410	-0.011	£36,214	No testing	£702	0.080	Extendedly dominated
Beckman 99th centile	£2,633	11.527	-£379	-0.007	£51,306	No testing	£733	0.084	£8,759
Roche strategy	£2,735	11.528	-£277	-0.007	£42,131	Beckman 99th centile	£102	0.001	Extendedly dominated
Abbott strategy	£2,808	11.532	-£204	-0.002	£105,889	Beckman 99th centile	£175	0.005	£32,042
Standard troponin	£3,012	11.534				Abbott strategy	£204	0.002	£105,889
MI prevalence = 30%									
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
No testing	£2,597	10.681	-£1,466	-0.137	£10,733				
Abbott 99th centile	£3,532	10.791	-£531	-0.027	£19,730	No testing	£935	0.110	Extendedly dominated
Roche 99th centile	£3,610	10.801	-£454	-0.017	£26,735	No testing	£1,012	0.120	Extendedly dominated
Beckman 99th centile	£3,655	10.807	-£408	-0.011	£36,797	No testing	£1,058	0.125	£8,431
Roche strategy	£3,767	10.808	-£296	-0.010	£29,991	Beckman 99th centile	£112	0.001	Extendedly dominated
Abbott strategy	£3,858	10.815	-£205	-0.003	£70,942	Beckman 99th centile	£203	0.008	£24,745
Standard troponin	£4,063	10.818				Abbott strategy	£205	0.003	£70,942

# APPENDIX 8: SUBGROUP ANALYSES (SECONDARY ANALYSIS)

### Deterministic secondary analysis:

Strategy			Com	pared to St	tandard troponin	roponin Compared to next best strategy			
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	∆Costs	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,789	11.530	-£276	0.036	Dominant				
Roche 99th centile	£2,832	11.532	-£232	0.039	Dominant	Abbott 99th centile	£44	0.002	Extendedly dominated
Beckman 99th centile	£2,858	11.532	-£206	0.039	Dominant	Abbott 99th centile	£69	0.003	Extendedly dominated
Roche strategy	£2,957	11.535	-£107	0.042	Dominant	Abbott 99th centile	£168	0.006	Extendedly dominated
Abbott strategy	£3,025	11.543	-£39	0.050	Dominant	Abbott 99th centile	£236	0.014	£17,047
Standard troponin	£3,064	11.493				Abbott strategy	£39	-0.050	Dominated

### Age and gender subgroups:

Females											
Age = 45											
Strategy			Con	npared to St	andard troponin	Compared to next best strategy					
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>		
Abbott 99th centile	£2,602	12.547	-£276	0.042	Dominant						
Roche 99th centile	£2,647	12.549	-£231	0.044	Dominant	Abbott 99th centile	£45	0.003	Extendedly dominated		
Beckman 99th centile	£2,673	12.550	-£205	0.044	Dominant	Abbott 99th centile	£71	0.003	Extendedly dominated		
Roche strategy	£2,773	12.553	-£105	0.048	Dominant	Abbott 99th centile	£170	0.006	Extendedly dominated		
Abbott strategy	£2,841	12.562	-£37	0.057	Dominant	Abbott 99th centile	£239	0.015	£16,023		
Standard troponin	£2,878	12.505				Abbott strategy	£37	-0.057	Dominated		
Age = 55											
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>		
Abbott 99th centile	£2,605	10.407	-£276	0.034	Dominant						
Roche 99th centile	£2,650	10.410	-£231	0.037	Dominant	Abbott 99th centile	£45	0.003	£15,224		
Beckman 99th centile	£2,676	10.410	-£205	0.038	Dominant	Roche 99th centile	£26	0.001	Extendedly dominated		
Roche strategy	£2,776	10.413	-£105	0.040	Dominant	Roche 99th centile	£126	0.003	Extendedly dominated		

Abbott strategy	£2,844	10.421	-£37	0.048	Dominant	Roche 99th centile	£194	0.011	£17,150
Standard troponin	£2,881	10.373				Abbott strategy	£37	-0.048	Dominated
Age = 65									
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,592	8.089	-£276	0.025	Dominant				
Roche 99th centile	£2,637	8.092	-£232	0.029	Dominant	Abbott 99th centile	£45	0.003	£13,064
Beckman 99th centile	£2,663	8.094	-£205	0.030	Dominant	Roche 99th centile	£26	0.001	Extendedly dominated
Roche strategy	£2,762	8.096	-£106	0.032	Dominant	Roche 99th centile	£126	0.003	Extendedly dominated
Abbott strategy	£2,831	8.103	-£37	0.039	Dominant	Roche 99th centile	£194	0.010	£18,999
Standard troponin	£2,868	8.064				Abbott strategy	£37	-0.039	Dominated
Age = 75									
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,521	5.618	-£278	0.015	Dominant				
Roche 99th centile	£2,565	5.621	-£234	0.019	Dominant	Abbott 99th centile	£44	0.004	£12,392
Beckman 99th centile	£2,592	5.623	-£207	0.021	Dominant	Roche 99th centile	£26	0.002	£16,407
Roche strategy	£2,691	5.625	-£108	0.022	Dominant	Beckman 99th centile	£99	0.002	Extendedly dominated
Abbott strategy	£2,759	5.630	-£40	0.028	Dominant	Beckman 99th centile	£168	0.007	£24,020
Standard troponin	£2,799	5.602				Abbott strategy	£40	-0.028	Dominated
Age = 85									
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,250	3.104	-£289	0.002	Dominant				
Roche 99th centile	£2,292	3.106	-£247	0.004	Dominant	Abbott 99th centile	£42	0.002	£21,140
Beckman 99th centile	£2,317	3.107	-£222	0.005	Dominant	Roche 99th centile	£25	0.001	£26,911
Roche strategy	£2,416	3.108	-£123	0.006	Dominant	Beckman 99th centile	£99	0.001	Extendedly dominated
Abbott strategy	£2,483	3.111	-£56	0.009	Dominant	Beckman 99th centile	£166	0.004	£45,709
Standard troponin	£2,539	3.102				Abbott strategy	£56	-0.009	Dominated

Males									
Age = 45									
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,958	13.801	-£275	0.042	Dominant				
Roche 99th centile	£3,000	13.803	-£233	0.044	Dominant	Abbott 99th centile	£43	0.002	Extendedly dominated
Beckman 99th centile	£3,026	13.803	-£207	0.044	Dominant	Abbott 99th centile	£68	0.002	Dominated
Roche strategy	£3,125	13.806	-£108	0.047	Dominant	Abbott 99th centile	£167	0.005	Extendedly dominated
Abbott strategy	£3,192	13.815	-£41	0.056	Dominant	Abbott 99th centile	£235	0.014	£16,897
Standard troponin	£3,233	13.759				Abbott strategy	£41	-0.056	Dominated
Age = 55									
	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs	Comparator	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs
Abbott 99th centile	£2,954	11.689	-£276	0.035	Dominant				
Roche 99th centile	£2,997	11.691	-£233	0.037	Dominant	Abbott 99th centile	£43	0.002	Extendedly dominated
Beckman 99th centile	£3,022	11.691	-£208	0.037	Dominant	Abbott 99th centile	£68	0.002	Dominated
Roche strategy	£3,121	11.694	-£109	0.040	Dominant	Abbott 99th centile	£167	0.005	Extendedly dominated
Abbott strategy	£3,188	11.702	-£41	0.048	Dominant	Abbott 99th centile	£234	0.013	£17,836
Standard troponin	£3,230	11.654				Abbott strategy	£41	-0.048	Dominated
Age = 65									
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,902	9.306	-£276	0.026	Dominant				
Roche 99th centile	£2,945	9.309	-£234	0.029	Dominant	Abbott 99th centile	£43	0.003	£16,877
Beckman 99th centile	£2,970	9.310	-£209	0.029	Dominant	Roche 99th centile	£25	0.001	Extendedly dominated
Roche strategy	£3,069	9.312	-£110	0.032	Dominant	Roche 99th centile	£124	0.003	Extendedly dominated
Abbott strategy	£3,136	9.319	-£42	0.039	Dominant	Roche 99th centile	£191	0.010	£19,851
Standard troponin	£3,179	9.280				Abbott strategy	£42	-0.039	Dominated

Age = 75									
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,752	6.560	-£278	0.017	Dominant				
Roche 99th centile	£2,795	6.563	-£236	0.019	Dominant	Abbott 99th centile	£42	0.002	£16,994
Beckman 99th centile	£2,819	6.563	-£211	0.020	Dominant	Roche 99th centile	£25	0.001	Extendedly dominated
Roche strategy	£2,918	6.565	-£112	0.022	Dominant	Roche 99th centile	£124	0.003	Extendedly dominated
Abbott strategy	£2,985	6.570	-£45	0.027	Dominant	Roche 99th centile	£191	0.008	£25,149
Standard troponin	£3,030	6.543				Abbott strategy	£45	-0.027	Dominated
Age = 85									
	Costs	QALYs	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
Abbott 99th centile	£2,374	3.631	-£283	0.006	Dominant				
Roche 99th centile	£2,414	3.631	-£244	0.007	Dominant	Abbott 99th centile	£40	0.001	Extendedly dominated
Beckman 99th centile	£2,437	3.631	-£220	0.007	Dominant	Abbott 99th centile	£63	0.001	Extendedly dominated
Roche strategy	£2,536	3.632	-£122	0.007	Dominant	Abbott 99th centile	£162	0.001	Extendedly dominated
Abbott strategy	£2,601	3.634	-£57	0.010	Dominant	Abbott 99th centile	£227	0.003	£66,418
Standard troponin	£2,657	3.624				Abbott strategy	£57	-0.010	Dominated

## **MI prevalence**

MI prevalence = 1%										
Strategy			Con	npared to St	andard troponin	Compared to next best strategy				
	Costs	QALYs	<b>∆Costs</b>	ΔQALYs	ΔCosts / ΔQALYs	Comparator	<b>∆Costs</b>	ΔQALYs	ΔCosts / ΔQALYs	
No testing	£1,072	12.546	-£439	-0.005	£96,456					
Abbott 99th centile	£1,405	12.619	-£106	0.068	Dominant	No testing	£333	0.073	£4,563	
Roche 99th centile	£1,407	12.615	-£104	0.064	Dominant	Abbott 99th centile	£2	-0.004	Dominated	
Beckman 99th centile	£1,408	12.611	-£103	0.061	Dominant	Abbott 99th centile	£3	-0.008	Dominated	
Roche strategy	£1,492	12.614	-£20	0.064	Dominant	Abbott 99th centile	£87	-0.005	Dominated	
Standard troponin	£1,511	12.550				Abbott 99th centile	£106	-0.068	Dominated	
Abbott strategy	£1,531	12.620	£20	0.070	£290	Abbott 99th centile	£126	0.001	£109,991	

MI prevalence = 5%									
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
No testing	£1,316	12.265	-£581	-0.023	£25,513				
Abbott 99th centile	£1,747	12.348	-£150	0.060	Dominant	No testing	£431	0.083	£5,209
Roche 99th centile	£1,759	12.346	-£137	0.058	Dominant	Abbott 99th centile	£13	-0.002	Dominated
Beckman 99th centile	£1,766	12.343	-£130	0.055	Dominant	Abbott 99th centile	£20	-0.005	Dominated
Roche strategy	£1,854	12.346	-£43	0.058	Dominant	Abbott 99th centile	£107	-0.002	Dominated
Standard troponin	£1,897	12.288				Abbott 99th centile	£150	-0.060	Dominated
Abbott strategy	£1,900	12.352	£4	0.064	£61	Abbott 99th centile	£154	0.004	£35,574
MI prevalence = 10%									
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
No testing	£1,623	11.913	-£758	-0.046	£16,645				
Abbott 99th centile	£2,178	12.008	-£203	0.050	Dominant	No testing	£554	0.095	£5,820
Roche 99th centile	£2,203	12.008	-£178	0.049	Dominant	Abbott 99th centile	£25	0.000	Dominated
Beckman 99th centile	£2,218	12.006	-£163	0.048	Dominant	Abbott 99th centile	£40	-0.002	Dominated
Roche strategy	£2,311	12.009	-£71	0.051	Dominant	Abbott 99th centile	£133	0.001	Extendedly dominated
Abbott strategy	£2,366	12.017	-£15	0.058	Dominant	Abbott 99th centile	£188	0.008	£22,684
Standard troponin	£2,381	11.958				Abbott strategy	£15	-0.058	Dominated
MI prevalence = 20%									
	Costs	QALYs	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	ΔCosts	ΔQALYs	<b>ΔCosts / ΔQALYs</b>
No testing	£2,247	11.202	-£1,112	-0.091	£12,211				
Abbott 99th centile	£3,053	11.324	-£306	0.031	Dominant	No testing	£806	0.122	£6,625
Roche 99th centile	£3,104	11.327	-£255	0.034	Dominant	Abbott 99th centile	£51	0.004	£14,063
Beckman 99th centile	£3,135	11.328	-£224	0.035	Dominant	Roche 99th centile	£30	0.001	Extendedly dominated
Roche strategy	£3,237	11.331	-£122	0.038	Dominant	Roche 99th centile	£132	0.004	Extendedly dominated
Abbott strategy	£3,310	11.340	-£49	0.047	Dominant	Roche 99th centile	£206	0.013	£16,319

Standard troponin	£3,359	11.293				Abbott strategy	£49	-0.047	Dominated
MI prevalence = 30%									
	Costs	QALYs	<b>∆Costs</b>	ΔQALYs	<b>ΔCosts / ΔQALYs</b>	Comparator	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs
No testing	£2,880	10.484	-£1,466	-0.137	£10,733				
Abbott 99th centile	£3,942	10.634	-£404	0.013	Dominant	No testing	£1,062	0.149	£7,109
Roche 99th centile	£4,019	10.641	-£327	0.020	Dominant	Abbott 99th centile	£77	0.008	£10,278
Beckman 99th centile	£4,065	10.645	-£281	0.024	Dominant	Roche 99th centile	£46	0.004	£12,899
Roche strategy	£4,177	10.648	-£169	0.027	Dominant	Beckman 99th centile	£112	0.003	Extendedly dominated
Abbott strategy	£4,268	10.658	-£78	0.037	Dominant	Beckman 99th centile	£203	0.013	£15,410
Standard troponin	£4,346	10.621				Abbott strategy	£78	-0.037	Dominated

Base case	MI prevalence	Roche 99	th centile	Standard troponin		Increments		
		Costs	QALYs	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs
Base case	17%	£2,301	11.740	£2,697	11.749	-£396	-0.010	£41,233
Age ≤70 <sup>b</sup>	28%	£3,411	10.946	£3,853	10.961	-£442	-0.015	£28,633
Age >70 <sup>c</sup>	10%	£1,550	6.274	£1,880	6.275	-£330	-0.001	£355,571
With pre-existing CAD	20%	£2,641	11.528	£3,012	11.534	-£371	-0.006	£58,509
Without pre-existing CAD	16%	£2,236	11.816	£2,592	11.821	-£356	-0.004	£80,454
Symptom onset < 3 hours	22%	£2,726	11.369	£3,222	11.391	-£496	-0.022	£22,111
Symptom onset > 3 hours	13%	£1,929	12.032	£2,277	12.036	-£348	-0.003	£103,107
Symptom onset < 3 hours	17%	£2,241	11.732	£2,697	11.749	-£456	-0.017	£26,327
Symptom onset > 3 hours	17%	£2,341	11.745	£2,697	11.749	-£356	-0.004	£80,677
Secondary analysis	MI prevalence	Roche 99th centile		Standard troponin		Increments		
		Costs	QALYs	Costs	QALYs	ΔCosts	ΔQALYs	ΔCosts / ΔQALYs
Base case	17%	£2,832	11.532	£3,064	11.493	-£232	0.039	Dominant
Age ≤70 <sup>b</sup>	28%	£3,839	10.780	£4,148	10.756	-£310	0.024	Dominant
Age >70 <sup>c</sup>	10%	£2,111	6.245	£2,259	6.222	-£148	0.023	Dominant
With pre-existing CAD	20%	£3,142	11.325	£3,359	11.293	-£217	0.031	Dominant
Without pre-existing CAD	16%	£2,778	11.604	£2,967	11.560	-£189	0.044	Dominant
Symptom onset < 3 hours	22%	£3,209	11.180	£3,556	11.159	-£347	0.021	Dominant
Symptom onset > 3 hours	13%	£2,503	11.806	£2,673	11.760	-£171	0.046	Dominant
Symptom onset < 3 hours	17%	£2,772	11.524	£3,064	11.493	-£292	0.031	Dominant
Symptom onset > 3 hours	17%	£2,873	11.535	£3,064	11.493	-£192	0.042	Dominant

### APPENDIX 9: SUBGROUP ANALYSES BASED ON ACCURACY AND AMI PREVALENCE (ONLY AVAILABLE FOR THE ROCHE 99TH CENTILE TEST)

<sup>a</sup> The two studies presenting data on subgroups <sup>67, 75</sup> were both conducted in patients in whom NSTEMI had not been excluded. They were not at specifically high or low risk of AMI. We calibrated the prevalence (obtained from these studies) in the subgroup to be adapted to a population with a prevalence of 17% (see below).

<sup>b</sup> Average age = 53 (base case value)

<sup>c</sup> Average age = 75

### AMI prevalence in subgroups

Subgroup	Prevalence of AMI (x)	Prevalence of AMI in whole population from subgroups were derived (y)	Prevalence assuming population prevalence of 17% (multiple x <sup>*</sup> y/17)	Source
Age<70 years	24%	15%	28%	APACE <sup>52, 75</sup>
Age>70 years	9%	15%	10%	APACE <sup>52, 75</sup>
Patients with CAD	18%	16%	20%	APACE 46, 75
Patients without CAD	14%	16%	16%	APACE 46, 75
<3 hours from symptoms <sup>67</sup>	24%	18%	22%	APACE, <sup>75</sup> Body (2011) <sup>67</sup>
>3 hours from symptoms <sup>67</sup>	14%	18%	13%	APACE, <sup>75</sup> Body (2011) <sup>67</sup>
<3 hours from symptoms <sup>75</sup>	21%	21%	17%	APACE, <sup>75</sup> Body (2011) <sup>67</sup>
>3 hours from symptoms <sup>75</sup>	21%	21%	17%	APACE, <sup>75</sup> Body (2011) <sup>67</sup>

#### APPENDIX 10: NICE GUIDANCE RELEVANT TO THE MANAGEMENT OF SUPECTED ACS

MI – secondary prevention: secondary prevention in primary and secondary for patients following a myocardial infarction. NICE clinical guideline CG172 (2013). Available from: <a href="http://guidance.nice.org.uk/CG172">http://guidance.nice.org.uk/CG172</a>. Date for review: not stated

Chest pain of recent onset: assessment and diagnosis of recent onset chest pain or discomfort of suspected cardiac origin. NICE clinical Guideline CG95 (2010). Available from: <a href="http://www.nice.org.uk/guidance/CG95">http://www.nice.org.uk/guidance/CG95</a>. Reviewed March 2013, review recommended.

Unstable angina and NSTEMI: the early management of unstable angina and non-ST-segmentelevation myocardial infarction. NICE clinical guideline CG94 (2010). Available from: http://www.nice.org.uk/guidance/CG94. Last modified November 2013.

BRAHMS copeptin assay to rule out myocardial infarction in patients with acute chest pain. NICE medical technology guidance MTG4 (2011). Available from: <u>http://guidance.nice.org.uk/MTG4</u>. Date for review: not stated.

Myocardial Infarction with ST-segment elevation: the acute management of myocardial infarction with ST-segment elevation. NICE clinical guideline CG167 (2013). Available from: <u>http://guidance.nice.org.uk/CG167</u>. Date for review: not stated.

### APPENDIX 11: PRISMA CHECK LIST

Section/topic	#	Checklist item	Reported on page #	
TITLE				
Title	1	Identify the report as a systematic review, meta-analysis, or both.	pg 1	
ABSTRACT				
Structured summary	2	Provide a structured summary including, as applicable: background; objectives; data sources; study eligibility criteria, participants, and interventions; study appraisal and synthesis methods; results; limitations; conclusions and implications of key findings; systematic review registration number.	Executive summary (pg 12-16) PROSPERO registration (pg 2)	
INTRODUCTION				
Rationale	3	Describe the rationale for the review in the context of what is already known.	Background (Section 2, pg 19-26)	
Objectives	4	Provide an explicit statement of questions being addressed with reference to participants, interventions, comparisons, outcomes, and study design (PICOS).	Objectives (Section 1, pg 18)	
METHODS				
Protocol and registration	5	Indicate if a review protocol exists, if and where it can be accessed (e.g., Web address), and, if available, provide registration information including registration number.	PROSPERO: CRD42013005939	
Eligibility criteria	6	Specify study characteristics (e.g., PICOS, length of follow-up) and report characteristics (e.g., years considered, language, publication status) used as criteria for eligibility, giving rationale.	Table 2 (pg 30)	
Information sources	7	Describe all information sources (e.g., databases with dates of coverage, contact with study authors to identify additional studies) in the search and date last searched.	Section 3.1.1 (pg 27-29)	
Search	8	Present full electronic search strategy for at least one database, including any limits used, such that it could be repeated.	Appendix 1	
Study selection	9	State the process for selecting studies (i.e., screening, eligibility, included in systematic review, and, if applicable, included in the meta-analysis).	Section 3.1.3 (pg 31)	
Data collection process	10	Describe method of data extraction from reports (e.g., piloted forms, independently, in duplicate) and any processes for obtaining and confirming data from investigators.	Section 3.1.3 (pg 31)	

Section/topic	#	Checklist item	Reported on page #	
Data items	11	List and define all variables for which data were sought (e.g., PICOS, funding sources) and any assumptions and simplifications made.	Section 3.1.3 (pg 31) Full data extraction tables: Appendix 2	
Risk of bias in individual studies	12	Describe methods used for assessing risk of bias of individual studies (including specification of whether this was done at the study or outcome level), and how this information is to be used in any data synthesis.	Section 3.1.4 (pg 31) Full QUADAS-2 tables: Appendix 3	
Summary measures	13	State the principal summary measures (e.g., risk ratio, difference in means).	Section 3.1.5 (pg 31-33)	
Synthesis of results	14	Describe the methods of handling data and combining results of studies, if done, including measures of consistency (e.g., I <sup>2</sup> ) for each meta-analysis.	Section 3.1.5 (pg 31-33)	
Risk of bias across studies	15	Specify any assessment of risk of bias that may affect the cumulative evidence (e.g., publication bias, selective reporting within studies).	NA	
Additional analyses	16	Describe methods of additional analyses (e.g., sensitivity or subgroup analyses, meta-regression), if done, indicating which were pre-specified.	Section 3.1.5 (pg 31-33)	
RESULTS				
Study selection	17	Give numbers of studies screened, assessed for eligibility, and included in the review, with reasons for exclusions at each stage, ideally with a flow diagram.	Figure 1 (pg 34)	
Study characteristics	18	For each study, present characteristics for which data were extracted (e.g., study size, PICOS, follow-up period) and provide the citations.	Section 3.2.3.1 (pg 38), section 3.2.4.1 (pg 51), section 3.2.5.1 (pg 56). Full data extraction tables:	
Risk of bias within studies	19	Present data on risk of bias of each study and, if available, any outcome level assessment (see item 12).	Section 3.2.2 (pg 35-37) Full QUADAS-2 tables: Appendix 3	
Results of individual studies	20	For all outcomes considered (benefits or harms), present, for each study: (a) simple summary data for each intervention group (b) effect estimates and confidence intervals, ideally with a forest plot.	Appendix 2, Table c	
Synthesis of results	21	Present results of each meta-analysis done, including confidence intervals and measures of consistency.	Figures 3-5 (pg 39-40), Table 4 (pg 47), Table 5 (pg 54), Table 6 (pg 58)	

Section/topic	#	Checklist item	Reported on page #	
Risk of bias across studies	22	Present results of any assessment of risk of bias across studies (see Item 15).	NA	
Additional analysis	23	Give results of additional analyses, if done (e.g., sensitivity or subgroup analyses, meta-regression [see Item 16]).	Table 4 (pg 47), Table 5 (pg 54), Table 6 (pg 58)	
DISCUSSION				
Summary of evidence	24	Summarise the main findings including the strength of evidence for each main outcome; consider their relevance to key groups (e.g., healthcare providers, users, and policy makers).	Section 5.1 (pg 102-107)	
Limitations	25	Discuss limitations at study and outcome level (e.g., risk of bias), and at review-level (e.g., incomplete retrieval of identified research, reporting bias).	Sections 5.2 and 5.3 (pg 107-114)	
Conclusions	26	Provide a general interpretation of the results in the context of other evidence, and implications for future research.	Section 6 (pg 115-116)	
FUNDING				
Funding	27	Describe sources of funding for the systematic review and other support (e.g., supply of data); role of funders for the systematic review.	pg 2	