# **ARTICLE IN PRESS**

**ORIGINAL ARTICLE** 

# Long Term Prognostic Value of a Negative Work-Up for Acute Coronary Disease in Emergency Department Chest Pain Patients Without Known Coronary Artery Disease: A Cohort Study

## **Q1** Anne-Maree Kelly, MD FACEM FFCP<sup>\*</sup>, Sharon Klim, BN

Joseph Epstein Centre for Emergency Medicine Research at Western Health, Melbourne, Vic, Australia

Received 16 March 2016; received in revised form 12 July 2016; accepted 22 July 2016; online published-ahead-of-print xxx

Background	To determine the rate of all cause and cardiac death, new myocardial infarction (MI) or coronary revascu- larisation at over three years from index visit in emergency department chest pain patients without known coronary artery disease (CAD) at index presentation who had a negative electrocardiogram (ECG) and biomarker workup for acute coronary syndrome (ACS).
Methods	An unplanned sub-study of a prospective observational study of consecutive adult patients presenting to the ED with atraumatic chest pain (or equivalents). The primary outcome of interest was the predictive performance of a negative ECG and biomarker work-up for ACS for all cause and cardiac mortality over more than three years' follow-up in patients not known to have pre-existing CAD presenting to the ED with chest pain. Secondary outcomes were rate of new MI or revascularisation not related to the index visit.
Results	237 patients were studied. Median age was 52 (IQR 42 – 62) and 55.3% were male. Median follow-up was 48 months. There were seven deaths (3%, 95% CI 1.4 – 6%), one of which was potentially cardiac in origin with cause of death given as pulmonary hypertension and cardiac failure (0.4%, 95% CI 0.02 – 2.3%). There was one confirmed MI (0.6%, 95% CI 0.03 – 3.8%). The rate of revascularisation not related to the index visit was 3.1% (95% CI 1.1 – 7.4%).
Conclusion	Patients who present to ED with potentially cardiac chest pain but do not have known CAD, have non- ischaemic ECGs and troponin assays below the 99 <sup>th</sup> percentile are at low risk of cardiac death or MI in long- term follow-up. This challenges the recommendation for routine functional or anatomic testing.
Keywords	Prognosis • Chest pain • Coronary artery disease • ED

#### 11 12 13

14

15

16

17

18

7

8

9

## Introduction

**Q2** Emergency department (ED) based processes to identify patients presenting with chest pain who are at low risk of acute coronary syndrome (ACS) or adverse cardiac events have been shown to have a low rate of, and high negative predictive value for, major adverse cardiac events (MACE) in

short-term follow-up [1-4]. Less is known about the predic-19tive performance of these processes in longer term follow-up,20particularly in the subgroup of patients without known pre-21existing coronary artery disease (CAD). Current guidelines22suggest that patients with a negative ACS work-up should23have provocative functional testing (e.g. exercise stress test,24myocardial perfusion imaging or stress echocardiography)25

Please cite this article in press as: Kelly A-M, Klim S. Long Term Prognostic Value of a Negative Work-Up for Acute Coronary Disease in Emergency Department Chest Pain Patients Without Known Coronary Artery Disease: A Cohort Study. Heart, Lung and Circulation (2016), http://dx.doi.org/10.1016/j.hlc.2016.07.015

<sup>\*</sup>Corresponding author at: JECEMR, Sunshine Hospital, Furlong Road, St Albans 3021, Vic, Australia, Email: anne-maree.kelly@wh.org.au © 2016 Australian and New Zealand Society of Cardiac and Thoracic Surgeons (ANZSCTS) and the Cardiac Society of Australia and New Zealand (CSANZ). Published by Elsevier B.V. All rights reserved.

# **ARTICLE IN PRESS**

or anatomical imaging (e.g. CT coronary angiography (CTCA)) to identify 'silent' CAD [5]. However if the rate of events in long-term follow-up is low and not cardiac-related, there may be a case for testing of selected patients only.

The aim of this study was to determine the rate of all cause and cardiac death, new myocardial infarction (MI) or coronary revascularisation, at over three years from index visit in patients without known CAD at index presentation who had a negative ECG and biomarker work-up for ACS in the ED.

## 35 Methods

#### 36 **Design and Setting**

This is an unplanned sub-study of a prospective observational
study of consecutive adult patients (aged over 18 years) presenting to the ED of two community teaching hospitals
between 19 January 2009 and 30 June 2009 with chest pain.

#### 41 **Participants**

Adult patients presenting with non-traumatic chest pain 42 (or equivalents) and undergoing evaluation for potential 43 ACS were eligible for inclusion in the parent study. For this 44 sub-study, additional inclusion criteria were absence of 45 46 known coronary artery disease at index ED visit, no ischaemic ECG features and all troponin assays during the index ED 47 evaluation being below the 99<sup>th</sup> percentile for the test. Known 48 coronary artery disease was as reported by the patient (con-49 firmed where possible from medical records) and defined as 50 51 any of: previous myocardial infarction (MI), physician-diag-52 nosed angina, percutaneous coronary intervention, coronary 53 artery bypass grafts or coronary angiogram showing stenosis >50%. Patients were not eligible for inclusion if they had 54 clearly ischaemic ECG features identified by the treating 55 56 clinician at initial assessment (including STEMI), they did not have a troponin assay or ECG performed within 24 hours 57 of pain onset, there was a clear non-ACS diagnosis made by 58 the treating clinician at initial assessment, they had a serious 59 arrhythmia pre-hospital or at ED presentation (including 60 cardiac arrest), language barrier or lack of telephone details 61 62 precluded follow-up or they were aged under 18 years. Patients were also excluded if they declined consent to fol-63 low-up or records could not be found. 64

#### Data Collection

65

66 Clinical and investigational data regarding the index presentation were collected on a piloted data collection form. Data 67 collected included demographics, cardiac risk factors, history 68 69 of CAD, cardiac failure, atrial fibrillation or peripheral vas-70 cular disease, clinical features at ED presentation, use of 71 warfarin, aspirin or statins, results of biochemical analyses 72 including cardiac biomarkers, ECG findings, interventions 73 during hospitalisation and in-hospital clinical course. In the parent study, patients were contacted by telephone at 7 and 74 75 30 days after the index ED visit to determine occurrence of 76 defined MACE (defined as all cause and cardiac death, new 77 MI or coronary revascularisation).

Regarding long-term follow-up, we chose to determine patient outcome as at 31 March 2013, representing approximately four years from the index ED visit. The choice of this date was arbitrary. Initially a review of the medical record was undertaken during 2014 to determine if the patient had died (including whether the cause of death was cardiac, noncardiac or unknown) or treatment for a new MI coronary revascularisation had occurred in the study health service during the follow-up period. If complete data was not available covering the relevant period, patients were contacted by telephone. If patients could not be contacted by telephone, a death registry search (Victorian Registry of Births, Deaths and Marriages) was undertaken. For patients who had died and the cause of death was unclear, clarification was made by contact with their family doctor. Deaths of unknown cause were assumed to be cardiac.

The primary outcome of interest was the predictive performance of a negative ECG and biomarker work-up for ACS for all cause and cardiac mortality. Secondary outcomes were the rate of new MI or revascularisation.

The troponin assay used was TnI-Ultra by Siemens Diagnostics performed on an Advia Centaur analyser. The test has a reported range of 0.006 to 50 microg/l. Co-efficient of variation is 10% at TnI 0.03 microg/l, 5.3% at 0.08 microg/l and 4.1% at 0.18 microg/l. The 99<sup>th</sup> percentile is 0.04 microg/l (95%CI 0.03 – 0.05 microg/l) (manufacturer's information). Timing of biomarkers was in accordance with the Australasian guidelines for contemporary troponin assays at the time of the index presentation [5]. Tests are taken at presentation and three to four hours later as long as the latter test is more than six hours from symptom onset. If a patient presented more than six hours from symptom onset, a single assay was deemed sufficient to rule out ACS.

#### Analysis and Sample Size

Data analysis is descriptive. Continuous variables are reported as medians and interquartile ranges (IQR). Categorical data is reported as proportions, with 95% confidence intervals where relevant. Mortality is reported for the whole sample. Myocardial infarction or revascularisations are only reported for patients with full follow-up. No sample size calculation was undertaken as this was an unplanned post-hoc study.

#### **Ethical Approval**

The project was approved by the institution's low risk ethics panel as a quality assurance project under the National Health and Medical Research Council (Australia) guidelines [6]. Patient consent for data collection from medical records was not required. Participants provided verbal consent to telephone follow-up.

## Results

Two hundred and thirty seven patients met inclusion criteria.128Sample derivation is shown in Figure 1. Median age was129

Please cite this article in press as: Kelly A-M, Klim S. Long Term Prognostic Value of a Negative Work-Up for Acute Coronary Disease in Emergency Department Chest Pain Patients Without Known Coronary Artery Disease: A Cohort Study. Heart, Lung and Circulation (2016), http://dx.doi.org/10.1016/j.hlc.2016.07.015

26

27

28

29

30

31

32

33

34

94

95

96

97

98

99

100

101

102

103

104

105

106

107

108

109

110

111

112

113

114

115

116

117

118

119

120

121

122

123

124

125

126

127

78

Download English Version:

# https://daneshyari.com/en/article/5602572

Download Persian Version:

https://daneshyari.com/article/5602572

Daneshyari.com