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BMJ Open  Clinical value of chest pain presentation and prodromes on the assessment of cardiovascular disease: a cohort study

John Robson,1 Luis Ayerbe,1 Rohini Mathur,1 Juliet Addo,2 Andrew Wragg3

ABSTRACT

Objectives: The recognition of coronary artery disease (CAD) among patients who report chest pain remains difficult in primary care. This study investigates the association between chest pain (specified, unspecified or musculoskeletal) and prodromes (dyspepsia, fatigue or dyspnoea), with first-ever acute CAD, and increased longer term cardiovascular risk.

Design: Cohort study.

Setting: Anonymised clinical data recorded electronically by general practitioners from 140 primary care surgeries in London (UK) between April 2008 and April 2013.

Participants: Data were extracted for all patients aged 30 years and over at the beginning of the study period, registered in the surgeries.

Main outcome measures: Clinical data included chest pain, dyspepsia, dyspnoea and fatigue, first-ever CAD and long-term cardiovascular risk (QRisk2).

Regression models were used to analyse the association between chest pain together with prodromes and CAD and QRisk2 ≥20%.

Results: 354 052 patients were included in the study.

4842 patients had first-ever CAD of which 270 reported chest pain in the year before the acute event. 257 019 patients had QRisk2 estimations. Chest pain was associated with acute CAD and longer term cardiovascular risk.

Conclusions: All patients with chest pain, including those with atypical symptoms, require careful assessment for acute and longer term cardiovascular risk. Prodromes may have independent diagnostic value in the estimation of cardiovascular disease risk.

INTRODUCTION

Coronary artery disease (CAD) is the leading cause of global mortality, accounting for 13% of deaths across the world.1 Chest pain is a common presenting symptom of CAD. However, the recognition of CAD among those who report chest pain remains difficult in primary care as chest pain only represents CAD in 8%2 to 11%3–5 of the patients. The guidelines of both the UK National Institute for Health and Care Excellence (NICE) and the American Heart Association acknowledge that clinical assessment may be sufficient to confirm or exclude the diagnosis of CAD.6,7 The NICE guidelines recommend the estimation of risk of CAD based on the typicality of the pain, age, gender and cardiovascular (CV) risk factors, and suggest the consideration of an alternative diagnosis if the estimated risk is below 10%.6 Many patients referred to secondary care are now offered investigations which have high costs and some involve exposure to ionising radiation.6–10 This care pathway emphasises the relevance of the initial clinical assessment in...
primary care. Both unnecessary referral and failure to diagnose CAD are important issues from a resource and clinical outcome perspective.11

The initial clinical assessment in primary care should include the consideration of the typicality of chest pain.6 Clinicians appropriately refer central chest pain with shoulder radiation, or avoid referring typical musculoskeletal chest pain. However, there is a large grey area of unspecified chest pain that fits neither of these categories, particularly in older people at higher risk. The current recommendations for estimating the risk of CAD in patients with chest pain, acknowledging the typicality of symptoms, are based on a single secondary care study published in 1993.12 However, it is reported in the guidelines that this study may overestimate the risk of CAD in primary care.6 12 In addition, about half of the patients with CAD also report other prodromal symptoms to their general practitioner in the months before the acute event3 15-16 with dyspepsia, dyspnoea and fatigue among the most frequently reported.14 16 There is little information on the diagnostic and prognostic value of such symptoms.

There are a number of validated tools, such as QRisk2, for the assessment of long-term CV risk in asymptomatic primary care patients.17 However, it is unclear whether these tools could also be used in patients with chest pain to estimate the risk of acute CAD. We wished to undertake preliminary work to explore the association between type of chest pain, together with prodromal symptoms, and CV risk. This will inform subsequent research, using a larger data set, in which a tool for prediction of acute CAD in primary care patients with chest pain might be derived.

This paper investigates the association between chest pain (specified, unspecified and musculoskeletal) and prodromal symptoms (dyspepsia, dyspnoea and fatigue), with first-ever acute CAD. The association between chest pain and prodromal symptoms with longer term CV risk (QRisk2 scores) is also examined.17

### METHODS

The study conformed to the STROBE (Strengthening the Reporting of Observational Studies in Epidemiology) study design recommendations.18

The study was based on patients living in three inner boroughs of east London (UK), registered with local general practitioners (GPs). This area had a population of 908,096 in April 2013 and over half of the patients were of non-white ethnicity. Anonymised clinical and demographic data recorded in GPs’ electronic health records between 1 April 2008 and 1 April 2013 were extracted using EMIS web software for all patients aged 30 years and over at the beginning of the study period. Patients from 140 of the 144 GP surgeries in the boroughs were used for the analysis (four surgeries used a different computer system and therefore were not included). Sociodemographic variables extracted included age, gender and self-reported ethnic group. The Townsend deprivation score, a measure of material deprivation derived from the UK Census at a small area level, that strongly correlates with standardised mortality ratios, was also recorded.19 Age was grouped into five categories of 35–44 years, 45–54, 55–64, 65–74, and 75 and older. Ethnicity was grouped into four categories: white, south Asian, black African/Caribbean and other. Individuals of mixed ethnicity were grouped with the relevant ethnic minority group. Clinical data included routinely recorded chest pain, prodromal symptoms and a report of a first-ever CAD in the electronic medical record. Chest pain was grouped into three categories of specified, unspecified and musculoskeletal. Table 1 shows the symptoms coded by the GPs that were used to define each of these categories in this study.

Prodromal symptoms included dyspepsia, dyspnoea and fatigue. The terms coded by the GPs that were used in this study to define CAD and each prodromal symptom are presented in online supplement 1. Additional clinical data used in the analyses included long-term CV risk, assessed with QRisk2 10-year CV risk scores.20 The QRisk2 score for prediction of CV disease within 10 years was stratified into lower (0–9%), medium (10–19%) and high risk (≥20%).17

The associations between chest pain, dyspepsia, dyspnoea or fatigue and first-ever reported CAD in the year succeeding the onset of symptoms were investigated. CAD was reported in the year after the start of symptoms because it was considered both observations within that period of time could be assumed to be clinically related. When prodromes (dyspepsia, dyspnoea or fatigue) had been reported in the 12 months before chest pain, and they were independently associated with CAD, they were combined with chest pain. The associations between the combination of chest pain and prodromes with first-ever CAD were also estimated.

Within the 5 years of the study period, the associations of chest pain, dyspepsia, dyspnoea or fatigue with high

| Table 1 | General practitioner notes used to define each category of chest pain |
|---------|----------------------------------------------------------|
| Chest pain category | Read codes from clinical notes included in each variable |
| Specified chest pain | Ischaemic chest pain, Central chest pain, Precordial pain, Parasternal pain, Chest pain on exertion, Retrosternal pain |
| Unspecified chest pain | Anterior chest wall pain, Chest wall pain, Chest pain not otherwise specified, Atypical chest pain, Chest pain unspecified |
| Musculoskeletal chest pain | Rib pain, Pleuritic pain, Painful breathing-pleurodynia |
CV risk (QRisk2 ≥ 20%) were investigated. When prodromes had been reported in the 12 months before chest pain, and they were independently associated with QRisk2 > 20%, they were combined with chest pain. The associations between the combination of chest pain and prodromes with QRisk2 > 20% were also estimated.

For patients with chest pain or prodromes recorded more than once, only the most recent episode of chest pain or the most recent prodrome, before the first-ever CAD or QRisk2 measurement, was included in the analysis. For those who had more than one QRisk2 estimation during the study period, only the first one was included in the analyses. Cox regression was used when CAD was the outcome. Cox regression describes how the risk of CAD changes over time in response to explanatory covariates (chest pain or prodromes). Results of Cox regression are presented in HRs, which are the expression of the risk of CAD occurring in patients with chest pain or prodromes as a ratio of the risk of CAD in patients without these symptoms. Models for the analysis of CAD were adjusted for age, gender, ethnicity and Townsend deprivation score. These variables were included in the models as they were considered potential confounders. Models for the analyses of QRisk2 were not adjusted as the QRisk2 algorithm already includes age, gender, ethnicity and Townsend score. SEs were adjusted for clustering by family practice.

RESULTS
In total, 354,052 patients aged 30 years or older were included in the study. During the 5-year study period, 14,222 (4.0%) patients reported chest pain to their general practitioner, 70,110 (19.8%) patients had reported dyspnoea, dyspepsia or fatigue and 257,019 (72.6%) patients had a QRisk2 estimation. Table 2 shows the sociodemographic and clinical characteristics of patients with chest pain and prodromes.

Among the 4842 patients with first-ever CAD, 270 (5.6%) reported chest pain to the general practitioner in the year before the acute event. Within the 12 months prior to the CAD event, 70 patients had a diagnosis of specified chest pain (25.9%), 187 had a diagnosis for unspecified chest pain (69.2%) and 17 had a diagnosis for musculoskeletal pain (6.3%). Some patients had more than one chest pain event coded in the preceding 12 months. For the purposes of calculating the risk of CAD after experiencing multiple chest pain events, the date of the chest pain closest to the CAD event was used. In the year preceding the CAD, 495 (10.2%) patients had reported prodromes. Dyspepsia was reported by 232 (46.9%) patients, dyspnoea by 235 (47.5%) and fatigue by 54 (10.9%) patients, with some patients having more than one prodrome recorded prior to CAD. Chest pain was independently (after adjusting for confounders) associated with a higher risk of CAD within the following year as shown in Table 3. This association was significant for all types of chest pain but strongest for patients reporting specified chest pain. Dyspepsia, dyspnoea or fatigue was also independently associated with a higher risk of CAD within the following year. The presence of chest pain of all subtypes in combination with dyspepsia, dyspnoea or fatigue was associated with an increased risk of CAD in the following year (Table 3).

Among the 2027 patients with 10-year CV risk of 20% or more as defined by QRisk2 reported after chest pain or prodromes, prior to a QRisk2 value of 20% or greater being recorded on the patient record, 195 (9.6%) had a diagnosis of specified chest pain, 1437 (70.9%) had a diagnosis of unspecified chest pain and 492 had...
a diagnosis of musculoskeletal chest pain (24.3%), with some patients having more than one chest pain event recorded. Of the 10,337 patients who had CV risk of 20% or more after any prodrome, the most recent prodrome recorded was dyspepsia for 7,022 (67.9%), dyspnoea for 2,865 (27.7%) and fatigue for 1,977 (19.1%) cases, with some patients having more than one prodrome recorded. Chest pain of all subtypes was associated with a high CV risk of 20% or more. Dyspepsia and dyspnoea were also associated with a high CV risk of 20% or more. Finally, the presence of dyspepsia or dyspnoea in combination with chest pain of all subtypes was associated with a high long-term CV risk of 20% or more (table 4).

**DISCUSSION**

Our study suggests that all presentations of chest pain were associated with an increased risk of CAD, within the next year, and increased longer term CV risk. These associations were strongest for specified chest pain, but importantly were also present for other categories of chest pain. Dyspepsia and dyspnoea were also associated with CAD and longer term CV risk. Fatigue was associated with increased risk of CAD but not with longer term CV risk.

This study has strengths and weaknesses. The data are derived from an almost complete population and not selected individuals or organisations and are likely to be representative of similar ethnically diverse populations. The inclusion of the entire local population in a large data set provides the least biased sampling frame. In contrast with many previous articles, this study has a high number of cases and improved statistical power. It was also possible to analyse several variables simultaneously, allowing the effect of multiple symptoms to be assessed. CAD may have been over-reported in our study, since we used an extensive list of terms for its definition. However, this broad definition of CAD probably minimised the number of CADs that were missed. The categorisation of chest pain used in this paper included pain location, which has been questioned as a predictor of CAD. Structured data entry templates and clinical facilitation in the east London practices studied enabled routine entry of high-quality data using agreed code sets for recording CV risk.
Clinicians are likely to be aware that patients reporting specified chest pain have a greater risk of CAD than those with atypical symptoms. These patients will require assessment in secondary care in most cases. However, unspecified and musculoskeletal chest pain should not be disregarded as these symptoms are associated with the risk of CAD above 10%. These results are consistent with the guidance for risk stratification included in the NICE guidelines. It has been reported that the association between atypical chest pain and CAD is strongest in older patients, South Asians and those affected by diabetes. Future studies may provide a more accurate estimation of the risk of CAD in primary care patients with chest pain in association with prodromal symptoms and other CV risk factors. It should also be noted that the proportion of patients with CAD who reported chest pain or prodromes to their GP in the year before the acute event was small. The natural history of CAD in the large proportion of patients who were not seen in primary care requires further clinical and epidemiological studies.

The factors that cause the association between chest pain and acute events are likely to explain its association with increased CV risk in the longer term. The typicality of symptoms appears to have less impact on the longer term CV risk. It is possible that some of the prodromal symptoms or episodes of chest pain observed within a year of CAD were not clinically related to it. Future studies may investigate whether the associations between chest pain, prodromes and CAD are similar when a shorter period of time (ie, three months) is observed. It should be noted that, in our sample, patients who reported chest pain were more likely to have QRisk2 estimations; therefore, the association between the symptoms and the long-term CV risk may be overestimated. The association between dyspepsia, dyspnoea or fatigue with CAD could also be partially explained by misdiagnosis in some patients, for example, angina being diagnosed as dyspepsia. Medication that prevents CV events, such as aspirin, may not have been prescribed to patients with dyspepsia, possibly increasing the rate of CAD among them.

A significant number of missed opportunities in the early management of CAD has been observed by other authors in primary care. Scores to estimate CV risk represent an attractive instrument to identify patients at high risk, both acutely and in the long term, and prompt more aggressive interventions. There are a number of reasons why the development of a risk score, using combinations of clinical data, for patients with chest pain in primary care may be appropriate. Patients with suspected CAD who are inadequately managed may have inappropriate and harmful investigations at high cost. Alternatively, those who are missed may have a preventable CV event. There are already validated and widely used scores predicting long-term CV risk in asymptomatic patients. There are also scores for risk assessment in patients with chest pain, which provide useful evidence for ruling out CAD. However, only two of the available scores, routinely used in clinical practice, have been derived and validated in primary care.

Electronic medical records represent clinical data sets of good quality, with larger samples than in most previous studies, and may provide a suitable basis to develop new and accurate scores for the diagnosis of acute and longer term CAD. The typicality of chest pain, together with the presence of dyspepsia and dyspnoea, might be considered in the further development of clinical decision rules based on risk of CAD for patients with chest pain but would require further validation.

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**Contributors** JR conceived the original idea. JR, LA, RM, JA and AW refined the questions and designed the study. RM conducted the analyses. JR, LA, RM, JA and AW co-wrote the paper, gave final approval of the version to be published and agreed to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

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