RESEARCH ARTICLE

Equity in healthcare for coronary heart disease, Wales (UK) 2004–2010: A population-based electronic cohort study

William King¹, Arron Lacey², James White³, Daniel Farewell⁴, Frank Dunstan⁴, David Fone⁴*

¹ Aneurin Bevan Gwent Local Public Health Team, Public Health Wales, Newport, Wales, United Kingdom, ² College of Medicine, Swansea University, Swansea, Wales, United Kingdom, ³ Centre for the Development and Evaluation of Complex Public Health Interventions and South East Wales Trials Unit, Cardiff University, Wales, United Kingdom, ⁴ Division of Population Medicine, Cardiff University, Cardiff, Wales, United Kingdom

* foned@cardiff.ac.uk

Abstract

Background

Despite substantial falls in coronary heart disease (CHD) mortality in the United Kingdom (UK), marked socioeconomic inequalities in CHD risk factors and CHD mortality persist. We investigated whether inequity in CHD healthcare in Wales (UK) could contribute to the observed social gradient in CHD mortality.

Methods and findings

Linking data from primary and secondary care we constructed an electronic cohort of individuals (n = 1199342) with six year follow-up, 2004–2010. We identified indications for recommended CHD interventions, measured time to their delivery, and estimated risk of receiving the interventions for each of five ordered deprivation groups using a time-to-event approach with Cox regression frailty models. Interventions in primary and secondary prevention included risk-factor measurement, smoking management, statins and antihypertensive therapy, and in established CHD included medication and revascularization. For primary prevention, five of the 11 models favoured the more deprived and one favoured the less deprived. For medication in secondary prevention and established CHD, one of the 15 models favoured the more deprived and one the less deprived. In relation to revascularization, six of the 12 models favoured the less deprived and none favoured the more deprived—this evidence of inequity exemplified by a hazard ratio for revascularization in stable angina of 0.79 (95% confidence interval 0.68, 0.92). The main study limitation is the possibility of under-ascertainment or misclassification of clinical indications and treatment from variability in coding.

Conclusions

Primary care components of CHD healthcare were equitably delivered. Evidence of inequity was found for revascularization procedures, although this inequity is likely to have only a modest effect on social gradients in CHD mortality. Policymakers should focus on reducing
established an application process to be followed by anyone who would like to access data via SAIL
https://www.saildatabank.com/application-process.

Funding: The authors received no specific funding for this work. WK was funded as a Welsh Clinical Academic Track Fellow. AL was employed by the Cardiovascular Research Group Cymru, funded by a grant from Health and Care Research Wales (formerly National Institute for Social Care and Health Research). Part of DFa’s work was supported by a Medical Research Council Methodology Fellowship.

Competing interests: The authors have declared that no competing interests exist.

Abbreviations: BMI, body mass index; CABG, coronary artery bypass graft; CHD, coronary heart disease; CI, confidence interval; DPP, deaths prevented or postponed; HDL, high-density lipoprotein; HR, hazard ratio; LSOA, lower super output area; MI, myocardial infarction; MICE, multiple imputations with chained equations; NICE, National Institute for Clinical Excellence (before 1 April 2005), National Institute for Health and Clinical Excellence (1st April 2005-31st March 2013), National Institute for Health and Care excellence (from 1st April 2013); NSF, National Service Framework; ONS, Office for National Statistics; OPCS, Office of Population Censuses and Surveys; PCI, percutaneous coronary intervention; PEDW, Patient Episode Data for Wales; PSALF, Project specific anonymised linking field; QOF, Quality and Outcomes Framework; SAIL, Secure Anonymized Data Linkage; SBP, systolic blood pressure; UK, United Kingdom; WIMD, Welsh Index of Multiple Deprivation.

inequalities in CHD risk factors, particularly smoking, as these, rather than inequity in healthcare, are likely to be key drivers of inequalities in CHD mortality.

Introduction

Coronary heart disease (CHD) mortality rates have declined rapidly in recent decades in most middle- to high-income countries [1,2,3]. However a steep social gradient in age-adjusted CHD mortality persists [1,4]. In the UK the rate ratio for premature CHD death in men was 1.84 comparing residents in the most and least deprived quintiles in 2008 [1] and the decline in CHD mortality in England (1982–2006) was faster in the least deprived areas [5]. Marked socioeconomic inequalities in major risk factors for CHD have been found in the UK [6,7,8] but it is not clear whether these inequalities fully explain the mortality gradient, as inequity (inequality to the disadvantage of more deprived groups) in provision of or access to healthcare might contribute to the gradient.

Modelling studies of UK populations have estimated that the decline in CHD mortality has been largely due to population-level reduction in risk factors, particularly rates of smoking and levels of blood pressure and cholesterol. [9–12]. The IMPACT studies, which estimated the proportions of the fall in CHD mortality attributable to changes in risk factors or treatments (effectiveness and provision) suggest that in England and Wales, between 1981 and 2000, 58% of the fall in CHD mortality could be attributed to population-level reduction in major risk factors and 42% to treatments [9]. An IMPACT study of the period 2000–2007 in England, during which CHD mortality fell by 36%, estimated that improved uptake of treatments accounted for approximately 50% of the fall [13].

A number of UK-based ecological studies have reported inequity in use of antihypertensive medication [14–16] and lipid-lowering medication [17,18] using analysis of practice-level data. A large individual-level UK study of secondary prevention of CHD found no evidence of inequity, and some findings suggested that more deprived groups were more likely to receive treatment [19]. UK-based individual-level studies of the management of diabetes found no evidence of inequity in the prescribing of antihypertensive medication and lipid-lowering therapy [20–22]. Other studies have reported clear evidence of inequity in the use of percutaneous coronary intervention (PCI) and coronary artery bypass grafting (CABG) in the UK [23–29].

A number of studies have examined different components of the CHD healthcare pathway [13,14,19]. We are not aware of any that examined whether there are inequities across the entire CHD pathway, from risk assessment, to primary and secondary prevention, medication in established CHD, and revascularization procedures. This is an important gap in the literature as the existing studies of different parts of the CHD pathway do not permit strong inferences to be made about the cumulative effect of inequity in one part of the pathway on inequities that become apparent at a later stage, and do not investigate inequity as a systematic, whole-pathway phenomenon.

We examined socioeconomic inequalities across a recommended CHD healthcare pathway in a population-level record-linked cohort study based on primary care, secondary care and demographic and mortality data from over one million adults, 2004–2010.

The study period followed introduction of the National Service Frameworks (NSF) for Coronary Heart Disease (introduced in 2000 in England and in 2001 in Wales) [30, 31] which set standards for all aspects of management of CHD, and the period coincided with the Quality Outcomes Framework (QOF) [32] introduced in 2004 to improve primary care including CHD healthcare in the UK.
Methods

All analyses were performed within the Secure Anonymized Information Linkage databank (SAIL) at Swansea University [33,34]. This system allows researchers to link anonymised data, including routine primary care, hospital activity, mortality and demographic data in a secure environment. SAIL implements rigorous information governance arrangements involving systematic data anonymisation, access limitations and disclosure controls. Permission to undertake the analysis was obtained from the Information Governance Review Panel at SAIL in line with the Collaborative Review System (project reference number 0156).

Datasets

We defined an electronic cohort of individuals aged 20 or over, resident in Wales and registered with SAIL-submitting general practices between 1 January 2004 and 31 December 2010. Routine data from the Welsh Demographic Service, Office of National Statistics (ONS) mortality files (ICD10 codes for cause of death), Patient Episode Data for Wales (PEDW) hospital admission data (ICD 10 and Office of Population and Censuses (OPCS) codes for CHD-related hospital episodes and procedures), and primary care data (Read codes for diagnosis, investigation and treatment of CHD and for the prescribing of antihypertensive, lipid-lowering and anti-platelet therapy) was extracted to form a linked dataset (for codes see S1 File).

Assessment of socioeconomic inequalities

The Welsh Index of Multiple Deprivation (WIMD) (2008) for the area of residence of the individual assessed at the Lower Super Output Area (LSOA) was used as a measure of socioeconomic deprivation. An LSOA is a unit of small-area geography used in the UK with a mean population of 1500. WIMD 2008 at LSOA level is based on residents’ income, employment status, education, housing, health and geographical access to services [35].

Pathway of CHD care

The pathway consisted of a sequence of evidence-based interventions recommended in NSFs [30, 31] and National Institute for Health and Clinical Excellence (NICE) guidelines [36–40]. These guidelines include CHD risk-assessment, primary prevention, secondary prevention, and medical and surgical treatment of established disease. The pathway of care investigated is shown in Fig 1.

Because of the complexity of the clinical algorithms for treating hypertension, the analysis of this area of the pathway was simplified by considering initiation of treatment as the prescription of any antihypertensive medication.

We identified ‘clinical trigger states’, defined as arising when an individual had an indication for an intervention according to NICE guidelines. Fig 1 shows the points at which clinical triggers (along the top of the figure) would be expected to prompt clinical actions (left hand side of the figure). In our comparisons for revascularization procedures we included a composite measure that included both PCI and CABG in order to avoid the possibility that increased use of either procedure might lead to a reduced need for the other procedure.

Covariates. For each clinical trigger identified in an individual, we determined covariates at the first appearance of the clinical trigger. Covariates available included demographic factors (age, sex, WIMD 2008); risk-factors (systolic blood pressure (SBP), BMI, smoking status, cholesterol: HDL ratio); co-morbidities based on the Charlson co-morbidity index [41] collapsed to a binary variable because some components (CHD, cerebrovascular disease and diabetes) were already considered as covariates; and the Framingham non-laboratory risk assessment
score (comprising sex, age, SBP, BMI, smoking status, reported diabetic status, and current treatment for hypertension) [42]. To take account of an individual’s previous progress in the pathway we included covariates for the timing of successive indications for the same intervention (for example high cardiovascular disease risk and subsequent angina both being indications for lipid-lowering therapy). Thus our models were able to represent an individual’s cumulative experience in the pathway.

### Statistical methods

We used a Cox model with random effects to examine associations with time-to-healthcare provision, measuring from the initiation of the clinical trigger state to the delivery of the indicated clinical action. We adjusted for important covariates, with the individual’s general practice or admitting hospital modelled as random effects, to allow for unobserved hospital or GP specific factors. Modelling was performed using the `coxme` package in R [43]. For each point on the pathway we selected covariates on the basis of relevance. Information on the covariates used in the models is provided in S2 File.

Absolute inequalities were examined by comparing WIMD quintile 5 (most deprived) to quintile 1 (least deprived). Deprivation quintile (Welsh Index of Multiple Deprivation 2008) of

---

**Fig 1. Clinical triggers and clinical actions investigated in the healthcare pathway for coronary heart disease.** Top row shows the clinical triggers in the healthcare pathway. The left-hand side shows the clinical actions identified in the pathway of care. Where boxes corresponding to a clinical trigger and clinical action are ticked, equity in the provision of care for that combination of clinical trigger and clinical action was investigated.

https://doi.org/10.1371/journal.pone.0172618.g001

---

<table>
<thead>
<tr>
<th>CLINICAL TRIGGER</th>
<th>Ascertained of smoking status</th>
<th>Measurement of BMI</th>
<th>Measurement of BP</th>
<th>Measurement of cholesterol</th>
<th>Full cardiovascular risk assessment</th>
<th>Provision of smoking cessation advice</th>
<th>Referral to smoking cessation services</th>
<th>Treatment with antihypertensive medication</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aged 40+ with no previous diagnosis</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td></td>
</tr>
<tr>
<td>First identified as smoker</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td></td>
</tr>
<tr>
<td>BP raised and low-risk</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td></td>
</tr>
<tr>
<td>Risk assessed high-risk</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td></td>
</tr>
<tr>
<td>High-risk diagnosis</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td></td>
</tr>
<tr>
<td>Stable angina</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td></td>
</tr>
<tr>
<td>Stable angina and diabetes</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td></td>
</tr>
<tr>
<td>MI</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>PRIMARY PREVENTION</th>
<th>SECONDARY PREVENTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Statin</td>
<td>✓</td>
</tr>
<tr>
<td>Aspirin</td>
<td>✓</td>
</tr>
<tr>
<td>ACE inhibitor</td>
<td>✓</td>
</tr>
<tr>
<td>Beta-blocker</td>
<td>✓</td>
</tr>
<tr>
<td>Clopidogrel</td>
<td>✓</td>
</tr>
<tr>
<td>PCI</td>
<td>✓</td>
</tr>
<tr>
<td>CABG</td>
<td>✓</td>
</tr>
<tr>
<td>Revascularisation</td>
<td>✓</td>
</tr>
</tbody>
</table>
residence for the individual was included as a term in every model. We used multiple imputations with chained equations (MICE) to create five imputed datasets. We imputed all missing covariates (systolic blood pressure, BMI, cholesterol: HDL ratio, smoking status, admission specialty, and admission type), using the MICE package in R 2.13.2. [44]. The type 1 error probability was set to 0.05 for all analyses.

We performed several sensitivity analyses. We re-ran the analyses using the Framingham 1991 risk-assessment tool [45] rather than the non-laboratory tool, the 2001 Townsend deprivation index [46] rather than WIMD, and 20 imputations rather than five in the chained equations for multiple imputation. We also repeated models using a slope index of inequality across all quintiles, instead of looking at the HR between the most deprived and least deprived quintiles.

Results

The initial cohort totalled 1201399 but after exclusion of individuals with clearly incorrectly coded date of birth (202), absent coding for gender (7) or with discontinuous registration with SAIL (1848) the cohort was reduced to 1199342. The primary care data available for our study was available only from SAIL-submitting practices and covered approximately 40% of the population of Wales, with a disproportionately high level of coverage in south west Wales. There is no available evidence that these practices were unrepresentative, and the distribution of urban and rural residency of the population resembled that of Wales as a whole. Comparing our data with ONS mid-year data for the whole of Wales, small differences were seen in age distribution, our cohort having 1.2% fewer in the proportion aged over 40.

The clinical triggers and related clinical actions are summarized in Table 1. Fig 2 shows the hazard ratio for the most deprived compared with the least deprived quintile (with 95% confidence intervals) for socioeconomic inequalities across the pathway of CHD healthcare.

Management of risk-factors

The clinical trigger, based on NSF and NICE guidance, for CHD risk-factor assessment was that the individual was aged 40 or over and was not previously recognized as being at high CHD risk. Three of the five comparisons showed components of CHD risk-assessment that favoured the most deprived quintile: ascertainment of smoking status, HR 1.20 (95% CI 1.17–1.24), BMI ascertainment, HR 1.12 (95% CI 1.08–1.16), measurement of BP, HR 1.03 (95% CI 1.00–1.06). The recording of a full risk-profile favoured the least deprived quintile, HR 0.97 (95% CI 0.95–1.00).

Provision of smoking-cessation advice favoured the most deprived group, HR 1.1 (95% CI 1.06–1.14).

Use of antihypertensive medication in individuals with raised systolic blood pressure (three readings above 160 mm Hg [38]) but otherwise at low risk, favoured the most deprived group, HR 1.22 (95% CI 1.13–1.31).

Medication in established disease

Use of statins in individuals with stable angina favoured the least deprived, HR 0.87 (95% CI 0.78–0.97). Use of aspirin in individuals with unstable angina favoured the most deprived, HR 1.24 (95% CI 1.05–1.46). The other 13 comparisons showed no statistically significant differences.
<table>
<thead>
<tr>
<th>Pathway position</th>
<th>Clinical trigger</th>
<th>Clinical action</th>
<th>Number of clinical triggers</th>
<th>Number of clinical actions</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Aged 40+ with no high risk diagnosis</td>
<td>Ascertainment of smoking status</td>
<td>122486</td>
<td>72291</td>
</tr>
<tr>
<td>2</td>
<td>Aged 40+ with no high risk diagnosis</td>
<td>Measurement of BMI</td>
<td>122486</td>
<td>46235</td>
</tr>
<tr>
<td>3</td>
<td>Aged 40+ with no high risk diagnosis</td>
<td>Measurement of BP</td>
<td>122486</td>
<td>64312</td>
</tr>
<tr>
<td>4</td>
<td>Aged 40+ with no high risk diagnosis</td>
<td>Measurement of cholesterol</td>
<td>122486</td>
<td>28652</td>
</tr>
<tr>
<td>5</td>
<td>Aged 40+ with no high risk diagnosis</td>
<td>Full cardiovascular risk assessment</td>
<td>122486</td>
<td>84969</td>
</tr>
<tr>
<td>6</td>
<td>First identified as smoker</td>
<td>Referral to smoking-cessation services</td>
<td>55161</td>
<td>2514</td>
</tr>
<tr>
<td>7</td>
<td>First identified as smoker</td>
<td>Provision of smoking-cessation advice</td>
<td>55161</td>
<td>45926</td>
</tr>
<tr>
<td>8</td>
<td>BP raised and low-risk</td>
<td>Treatment with antihypertensive medication</td>
<td>13814</td>
<td>9899</td>
</tr>
<tr>
<td>9</td>
<td>BP raised and high-risk</td>
<td>Treatment with antihypertensive medication</td>
<td>106079</td>
<td>75797</td>
</tr>
<tr>
<td>10</td>
<td>Risk assessed high</td>
<td>Statin</td>
<td>105301</td>
<td>20661</td>
</tr>
<tr>
<td>11</td>
<td>High-risk diagnosis</td>
<td>Statin</td>
<td>34387</td>
<td>19389</td>
</tr>
<tr>
<td>12</td>
<td>Stable angina</td>
<td>Statin</td>
<td>11104</td>
<td>4660</td>
</tr>
<tr>
<td>13</td>
<td>Stable angina and diabetes</td>
<td>Statin</td>
<td>2457</td>
<td>968</td>
</tr>
<tr>
<td>14</td>
<td>Unstable angina</td>
<td>Statin</td>
<td>4462</td>
<td>2178</td>
</tr>
<tr>
<td>15</td>
<td>MI</td>
<td>Statin</td>
<td>10442</td>
<td>5372</td>
</tr>
<tr>
<td>16</td>
<td>Stable angina</td>
<td>Aspirin</td>
<td>9433</td>
<td>3923</td>
</tr>
<tr>
<td>17</td>
<td>Stable angina and diabetes</td>
<td>Aspirin</td>
<td>2736</td>
<td>919</td>
</tr>
<tr>
<td>18</td>
<td>Unstable angina</td>
<td>Aspirin</td>
<td>4172</td>
<td>2041</td>
</tr>
<tr>
<td>19</td>
<td>Stable angina</td>
<td>Statin</td>
<td>11104</td>
<td>4660</td>
</tr>
<tr>
<td>20</td>
<td>Stable angina and diabetes</td>
<td>ACE inhibitor</td>
<td>3361</td>
<td>1092</td>
</tr>
<tr>
<td>21</td>
<td>Unstable angina</td>
<td>ACE inhibitor</td>
<td>5287</td>
<td>1967</td>
</tr>
<tr>
<td>22</td>
<td>MI</td>
<td>ACE inhibitor</td>
<td>10595</td>
<td>5270</td>
</tr>
<tr>
<td>23</td>
<td>Unstable angina</td>
<td>Beta-blocker</td>
<td>10405</td>
<td>285</td>
</tr>
<tr>
<td>24</td>
<td>MI</td>
<td>Beta-blocker</td>
<td>16639</td>
<td>363</td>
</tr>
<tr>
<td>25</td>
<td>Unstable angina</td>
<td>Clopidogrel</td>
<td>13907</td>
<td>5783</td>
</tr>
<tr>
<td>26</td>
<td>MI</td>
<td>Clopidogrel</td>
<td>20467</td>
<td>10132</td>
</tr>
<tr>
<td>27</td>
<td>Stable angina</td>
<td>PCI</td>
<td>18934</td>
<td>1172</td>
</tr>
<tr>
<td>28</td>
<td>Stable angina and diabetes</td>
<td>PCI</td>
<td>8956</td>
<td>300</td>
</tr>
<tr>
<td>29</td>
<td>Unstable angina</td>
<td>PCI</td>
<td>13907</td>
<td>2130</td>
</tr>
<tr>
<td>30</td>
<td>MI</td>
<td>PCI</td>
<td>20467</td>
<td>5118</td>
</tr>
<tr>
<td>31</td>
<td>Stable angina</td>
<td>CABG</td>
<td>18934</td>
<td>1150</td>
</tr>
<tr>
<td>32</td>
<td>Stable angina and diabetes</td>
<td>CABG</td>
<td>8956</td>
<td>385</td>
</tr>
<tr>
<td>33</td>
<td>Unstable angina</td>
<td>CABG</td>
<td>13907</td>
<td>1155</td>
</tr>
<tr>
<td>34</td>
<td>MI</td>
<td>CABG</td>
<td>20467</td>
<td>1645</td>
</tr>
<tr>
<td>35</td>
<td>Stable angina</td>
<td>Revascularisation</td>
<td>18934</td>
<td>2298</td>
</tr>
<tr>
<td>36</td>
<td>Stable angina and diabetes</td>
<td>Revascularisation</td>
<td>8956</td>
<td>676</td>
</tr>
<tr>
<td>37</td>
<td>Unstable angina</td>
<td>Revascularisation</td>
<td>13907</td>
<td>3230</td>
</tr>
<tr>
<td>38</td>
<td>MI</td>
<td>Revascularisation</td>
<td>20467</td>
<td>6649</td>
</tr>
</tbody>
</table>
Fig 2. Hazard ratios (HRs) and 95% confidence intervals (CIs) for the association between absolute socioeconomic inequalities and provision of healthcare for coronary heart disease. Where the association is not statistically significant at the p<0.05 level the box is empty.
Revascularization

Of the 12 comparisons made in relation to revascularization procedures six favoured the least deprived, and six showed no significant difference although favouring at a non-significant level the least deprived. In individuals with stable angina the HR for PCI was 0.72 (95% CI 0.59–0.88) and for revascularization (combined) 0.79 (95% CI 0.68–0.92). In unstable angina the HR for PCI was 0.76 (95% CI 0.66–0.88) and for revascularization (combined) 0.83 (95% CI 0.74–0.94). In myocardial infarction (MI) the HR for PCI was 0.83 (95% CI 0.75–0.92) and for revascularization (combined) 0.83 (95% CI 0.77–0.91).

Sensitivity analyses

Sensitivity analyses, using the Framingham 1991 risk-assessment tool rather than the non-laboratory tool, the 2001 Townsend deprivation quintiles rather than WIMD (2005), 20 imputations in the chained equations for multiple imputations, and using a slope-index-of-equality (instead of looking at the HR for the most deprived compared to least deprived deprivation quintile) all had little effect on the overall pattern of our findings.

Discussion

Main findings

For the healthcare pathway (excluding the composite revascularisation outcome), we found six points in the pathway where the most deprived quintile were more likely to receive the clinical action in question and five points where the least deprived were more likely to receive the clinical action; at 23 points there was no significant difference. Our interpretation of these findings was that, in a population-level analysis of the entire CHD healthcare pathway, there was no evidence of systematic inequity in utilisation of healthcare adjusted for need.

The study was not designed to identify inequity at specific points in the pathway, and, with the problem of multiple comparison in mind, interpretation of findings at specific stages in the pathway needs to be undertaken with caution. We did identify evidence that pattern of the inequalities related to the stages of the pathway. For primary prevention, significant inequalities favouring the most deprived quintile were found in five of the 11 interventions. Of the 13 interventions for medication in secondary prevention and in established CHD one inequality favouring the least deprived quintile and one the most deprived. There was evidence of inequality favouring the least deprived quintile in six of the 12 interventions relating to revascularization.

Comparison with other studies

An individual-level study comprehensively examined prescribing in secondary prevention of CHD in the UK between 1999 and 2007 and detected no evidence of inequity, its findings suggesting greater prescribing in the most than in the least deprived groups [19]. An IMPACT modelling study to examine the relationship between socioeconomic status and the factors that explain the declining rates of CHD mortality in the UK between 2000 and 2007 found no evidence of inequity in the delivery of a wide range of interventions [13]. Our findings therefore broadly concur with those of these studies and a number of other individual-level studies [20–22] although they conflict with those of several ecological studies [14–18] and studies of...
UK populations during the 1990’s [47–49] that reported evidence of inequity in primary care elements of CHD healthcare.

Our findings replicate those of a number of studies covering the same time period relating to revascularization in UK populations, in which rates of revascularization favoured the less deprived [23–29]. Several studies reporting inequity in revascularization suggest that individuals’ attitudes, expectations and consultation thresholds, differing according to socioeconomic status, may contribute to the inequalities observed [28,50,51]. One qualitative study suggested that attitudes such as low expectations of treatment and fear of hospitals were more likely to be found in more deprived groups and formed barriers to referral for angiography and subsequent revascularization [52].

Implications for policy

We found that the primary care components of CHD healthcare were delivered equitably in the population of Wales during the study period 2004–2010. Despite this finding and the results of comparable studies [13, 19–22], there is evidence that the fall in CHD mortality in the UK has been faster in more affluent groups and that relative inequality in CHD mortality has increased [13,53]. Adverse effects of increasing rates of diabetes and obesity are expected to offset improvements in population risk-profiles [9–12] and, because of higher rates of these conditions in more deprived groups, to increase relative inequality. These studies suggest that even if CHD healthcare is equitably delivered, inequalities in CHD outcomes are likely to persist, and relative inequalities to increase, unless inequalities in major risk-factors such as tobacco use are addressed.

We found clear evidence of inequity in revascularization procedures. Marked increases in the resources for revascularization, and in the volume of procedures carried out, were observed in the UK during our study period. Between 2000 and 2012 the rates of PCI (all types) in Wales increased from 550 per million population to 1363 per million, and between 2004 and 2008 increased from 900 to 1150 per million population. [54]. Our findings suggest that this increase was not applied equitably. It would be valuable to examine reasons for this, in particular deprivation-related attitudes (such as low expectations of healthcare and fear of hospitals) that limit demand, and the distribution of co-morbidities across the social gradient. Comorbidities, potentially acting as contraindications to revascularization procedures, may explain part of the observed inequality, and in our study the evidence of inequity weakened when the model was adjusted for comorbidities.

Evidence from modelling studies helps to clarify the potential effect on CHD mortality of the observed inequity in revascularization. In England and Wales between 1981 and 2000, while 42% of the decline in CHD mortality was due to medical and surgical intervention, only 3.8% of deaths prevented or postponed (DPP) were due to CABG or angioplasty [9]. This estimate is similar to that reported in a comparable USA study [55]. In a study of USA cardiac patients (1980–2000) it was estimated that less than 6% of total life-years gained were attributable to revascularization procedures [56]. In a study of declining CHD mortality in New Zealand 1983–1993 [57] DPP by revascularization procedures were estimated to contribute 5% of the total CHD mortality reduction. A study using IMPACT modelling to explain the decline in CHD mortality in Northern Ireland between 1987 and 2007, calculated that CABG or angioplasty in acute MI or unstable angina accounted for less than 1% of DPP [11]. We assess from such estimates that the degree of inequity that we observed in revascularization procedures would make only a small contribution to the steep social gradient of CHD mortality (hazard ratio 1.72 in our adjusted model). Further work to monitor inequalities in revascularization and to quantify their effects on mortality, would be valuable to policy-makers.
Strengths of the study

We based our study on data from a large number of individuals (more than one million) and identified a large number of clinical triggers for each of which we determined the time to clinical action. In contrast to previous studies that had examined different parts of CHD risk-factor assessment, management and treatments, we were able to examine inequalities across an entire recommended CHD pathway at the individual level. We were able to use a hierarchical structure (individual/practice/hospital) in our modelling and this allowed us to take account of supply-side factors as random variables. By using a time-to-event approach we eliminated in our analysis the use of an arbitrary standard for acceptable time-intervals between clinical trigger and action.

We used a set of algorithms for identifying, collecting and classifying relevant information on clinical triggers from a large and unrefined data source. Information on the length of time for which a particular clinical action was indicated and the number of different previous indications that had arisen for that action, enabled us to adjust for elements in an individual’s history as potential confounders in models.

This methodology may be applicable to studies of pathways of care for diseases other than CHD.

Limitations

The main study limitation is that inaccuracies and variability in the use of Read and ICD codes in medical records are known to occur [58,59]. Under-ascertainment or misclassification of clinical indications and interventions would tend to bias the results towards null, potentially reducing the power of our study to identify genuine inequity.

Prescribing data relating to inpatient treatment of acute coronary syndromes was not available in our routine data and we therefore had information on drug treatments for individuals only after they had left hospital. We did not include anti-anginal therapy in the pathway as it is not considered directly to affect CHD mortality, and because our data would not necessarily allow us to distinguish whether some types of medication, including calcium channel blockers and beta blockers, were used in an individual to treat angina or hypertension.

Our study examined a health service in which healthcare is free at the point of delivery and there is no charge for prescriptions. The system of healthcare in the UK, in which interventions such as QOF can operate, contrasts with less integrated systems such as those in the USA, and further work using a similar approach in such health systems without might be revealing.

Conclusions and recommendations

Primary care components of CHD healthcare were equitably delivered in the population of Wales between 2004 and 2011. Clear evidence of inequity was found in relation to revascularization procedures.

Organisations and policymakers should focus on the clear social gradients in risk factors as it is these, rather than inequity in healthcare, that are the key drivers of social gradients in CHD mortality. They should address the increasing rates of obesity and diabetes that are offsetting the benefits of recent reductions in other major CHD risk factors such as smoking.

The time-to-event methodology of this study has been shown to be an effective way of examining evidence of equity in utilization of healthcare and could be similarly used in studies of other disease areas.
Supporting information

S1 File. Clinical codes. Clinical codes used to define clinical conditions from routine data. (PDF)

S2 File. Covariates and outputs from Cox models. (PDF)

Acknowledgments

This study makes use of anonymised data held in the Secure Anonymised Information Linkage (SAIL) system, which is part of the national e-health records research infrastructure for Wales. We would like to acknowledge all the data providers who make anonymised data available for research.

We are grateful to Professor Simon Capewell and to Professor Tim Doran for their helpful feedback on the PhD thesis on which this study is based. We acknowledge the kind support of Professor Julian Halcox for this study.

Author Contributions

Conceptualization: WK D. Fone FD D. Farewell.

Data curation: AL WK.

Formal analysis: WK FD D. Farewell.

Investigation: WK AL.

Methodology: WK D. Fone FD JW D. Farewell AL.

Project administration: WK D. Fone.

Software: WK.

Supervision: D. Fone FD.

Validation: WK.

Visualization: WK D. Fone.

Writing – original draft: WK.

Writing – review & editing: D. Fone FD AL D. Farewell JW.

References


36. National Institute for Clinical Excellence (2001) Prophylaxis for patients who have experienced a myocardial infarction. (CGA)


