An additional dimension to health inequalities: disease severity and socioeconomic position

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Abstract

Objective—To investigate the association between the severity of hip pain and disability, and a number of measures of socioeconomic position, using a range of individual and ecological socioeconomic indicators.

Design—Interviewer administered and self-completed questionnaires on symptoms of pain and disability, general health and socioeconomic indicators, completed by people reporting hip pain in a cross-sectional, postal, screening questionnaire.

Setting—40 general practices from inner city, suburban and rural areas of south west England.

Participants—954 study participants who had reported hip pain in a postal questionnaire survey of 26,046 people aged 35 and over, selected using an age-sex stratified random probability sample.

Data—Individual indicators of socioeconomic position: social class based on occupation, maximum educational attainment, car ownership, gross household income, manual or non-manual occupation and living alone. Area level measures of socioeconomic position: Townsend scores for material deprivation at enumeration district level; urban or rural location based on the postcode of residence. Severity of hip disease, measured by the pain, disability and independence components of the New Zealand score for major joint replacement. Self reported comorbidity validated using general practice case notes and summary measures of general health.

Main results—Increasing disease severity was strongly associated with increasing age and a variety of measures of general health, including comorbidity. The data provide considerable evidence for the systematic association of increased severity of hip disease with decreasing socioeconomic position. Measures of socioeconomic position that were systematically associated with increasing disease severity, standardised for age and sex, included educational attainment (relative index of inequality 1.95 (95% confidence intervals 1.29 to 2.62)) and income (relative index of inequality 4.03 (95% confidence intervals 3.49 to 4.64)). Those with access to a car (mean disease severity 15.5) had statistically significant lower severity of hip disease than those without (mean 17.5, p<0.01). Similar results were found for access to higher or further education and living with others. For a given level of income, people with greater comorbidity had more severe hip pain and disability. The gradient in disease severity between rich and poor was steepest among those with the most comorbidity.

Conclusions—People with lower socioeconomic position experience a greater severity of hip disease. The poorest sector of the population are more likely to be in civil jeopardy: they not only experience a greater burden of chronic morbidity but also a greater severity of hip disease. This study has implications for health care provision, if the National Health Service is to live up to its principle of equal treatment for equal medical need.

Over the past few years the issue of inequalities in health has been raised from being politically taboo in Britain,1 to being embraced within the key political goal of reducing the “health gap” between rich and poor.1 One of the founding principles of the National Health Service was that there should be equal treatment for equal medical need, but a compromise had to be built in, which allowed private practice to coexist alongside statutory health care provision, resulting in an inevitable compromise of the equity goal.2 Today, after half a century of comprehensive services that are universally available and free at the point of delivery, there is clear evidence of inequalities in health that are widening in certain groups of society.3 The recent independent inquiry,4 into inequalities in health has provided 39 recommendations of methods to reduce reversible inequalities. Its publication has been welcomed for highlighting the problems of inequalities but the recommendations have been criticised for lack of prioritisation, vagueness and for not being costed.5 The challenge of how to direct health care resources to where they are most needed remains unmet.

In investigating the extent of inequalities in morbidity across the socioeconomic hierarchy, some investigators have used summary measures of health,6 while others have used disease specific measures. A clear social class gradient has been demonstrated for global self assessed health,7 myocardial infarction, stroke, diabetes and chronic bronchitis.8 Limiting longstanding illness,9 and disability,10 increase as household income decreases, as do the prevalences of angina, diabetes, bronchitis,11 and arthritis.12 Psychiatric disorders are more common in civil servants with lower pay and employment grades.13 Increased educational attainment has...
been associated with lower prevalences of self reported diabetes, heart disease and chronic respiratory disease, and arthritis. Ecological measures have demonstrated inequalities at area level, with the prevalence of many forms of self reported chronic morbidity being higher in more deprived areas. Evidence about urban and rural health is mixed, with a review concluding that people from urban areas have poorer health in general, but those from certain rural areas have poorer access to high quality care. Recent evidence asserts that to understand determinants of inequalities in health, studies must use more than one measure of socioeconomic position. A largely ignored dimension of the issue of inequalities in health, is the degree to which disease severity is associated with socioeconomic position. Despite warnings in the literature, about possible differential reporting of disease by educational attainment, income, and the inadequacy of studies relying on just one or two simple questions to pick up the range within musculoskeletal, respiratory and mental diseases, there still remain a dearth of studies that have investigated the association between disease severity and socioeconomic position. One study that has gone some way to investigate this, comprised 1044 men aged under 55 with ankylosing spondylitis. Mean scores for pain, physical activity and disease impact were statistically significantly higher for unemployed men compared with employed men. To try and resolve this gap in the research evidence, this study focuses on severity of hip disease rather than prevalence and examines the association between severity and a number of measures of socioeconomic position, using a wide range of socioeconomic indicators.

Methods

STUDY DESIGN

A multistage sampling strategy, was adopted to take an age/sex stratified random sample of 28 080 people aged 30 and over from 40 general practices in the counties of Avon and Somerset in south west England. The last British Census (1991) revealed that Avon had a resident population of almost 1 million (932 674), whereas Somerset, which is geographically much larger and more rural, had a population of almost half a million (460 368). The percentage of the population that was of pensionable age was 22% for Somerset and 19% for Avon.

After checking the names and addresses with the general practices, those who had died, moved out of the area or had a terminal illness were removed from the sample. A postal screening questionnaire comprising questions on general health, musculoskeletal disease of any joint, symptoms of hip and knee disease, back problems and sociodemographic indicators, was sent to 26 046 people. A question derived from the American Health and Nutrition Examination Survey, was used as the key screening question for hip problems. Details of the screening phase of the project including follow up of non-responders and validation of the questionnaire are published elsewhere. Essentially, 22 204 people provided usable responses to the hip screening question. A total of 3169 (14.0%) reported hip pain on screening, equating to a prevalence of 143 per 1000 people aged 55 and over (95% confidence intervals: 136 to 150 per 1000). An analysis of the effect of the cluster design across 40 practices on the prevalence of self reported pain revealed only minimal effects on the width of the confidence intervals. To adjust for this design effect, the standard errors around the prevalence of people reporting hip pain would need to be inflated by a design factor of just 1.12 for all men and 1.32 for all women. Those reporting hip pain were invited to attend a study clinic for a detailed orthopaedic and general assessment. Study clinics were held at general practices and hospitals throughout the region. People who attended a clinic were offered an assessment at home. All those who attended a clinic had their travel expenses reimbursed and were able to bring dependants to the clinics if necessary. They completed a detailed, interviewer administered questionnaire about symptoms of pain and function in the hip, knee and lower back. They were also asked to self report details of their domestic circumstances and their general health.

MEASURES OF SOCIOECONOMIC POSITION

Several indicators of socioeconomic position were measured at individual and ecological level. The ecological measures were enumeration district and ward level Townsend scores for material deprivation, and urban and rural status derived from the postcode of each study participant. The following individual level measures of socioeconomic position were collected: social class based on each respondent’s most recent full time occupation (housewives with no other occupation were assigned their partner’s occupation, where given); manual/non-manual occupation, gross household income; maximum educational attainment (derived from a list of academic and trade qualifications and age at which the respondent left school); access to any higher or further education; living alone and access to a car.

MEASURES OF ILL HEALTH

Severity of hip disease, indicated by pain and disability in or around the hip joint, was assessed by a continuous score from 0 to 80 using three of the four components of a score for assessing global disease severity, the New Zealand score for major joint replacement (the New Zealand Score). Measures of pain (40 points), disability (20 points), involvement of other joints in the disease process and the degree to which independence was threatened (20 points) are all taken into account by this scoring system. For the purpose of these analyses, the fourth component of the score about clinical and radiological examination findings and which carried a maximum of 20 points, was not included as not all the clinic attendees were able to be examined or have radiology. Prevalence and incidence estimates of hip
disease that may require primary hip replacement surgery, based on data from all four components of the score, are reported elsewhere. Comorbidity was self reported on the screening questionnaire. The variables were grouped into respiratory, cardiovascular or eye diseases, depression and cancers. A summary score was devised to indicate the burden of chronic comorbidity experienced by each study member. Reporting any one from each major disease group gave a score of 1 and an additional 0.5 was given for each extra disease from within a disease group. The sum across all disease groups formed the comorbidity score, which ranged from 0 to 5.5. Because of the small numbers in some of the groups, the comorbidity score has been collapsed into three groups, for ease of interpretation and presentation. A score greater than 1 but less than 2.5 is referred to as “considerable comorbidity” and a score of 2.5 or more is referred to as “multiple comorbidity”. In addition to these details from the postal screening questionnaire, clinic attendees completed Lakert scales to assess their general health, how their health compared with that of a year ago and the extent to which their physical and/or mental health interfered with their social life.

Full ethical committee approval was granted for all stages of the data collection.

VALIDATING THE ILLNESS MEASURE
To test the validity of using the New Zealand score for this purpose, Spearman’s rank correlation was used to correlate the score with a battery of variables: age, four measures of general health and comorbidity.

SUBJECTS INCLUDED IN THE STUDY
Full details of the clinic invitation criteria have been described elsewhere. Essentially, 2027 of the 2149 people who had reported hip pain on screening were invited to a study clinic. A total of 1405 (69%) people attended a study clinic or were given a home visit. There were no statistically significant differences between clinic attendees and non-attendees in terms of comorbidity, with the exception that women who attended a clinic had a statistically significant higher prevalence of cataract than women who did not attend. For all three measures of service utilisation asked on the postal questionnaire (GP consultation in the past 12 months for hips, currently waiting to see a hospital doctor and currently on a waiting list for a hip replacement), there were significant differences for both sexes, with attenders more likely to have accessed care for hip disease.

A total of 1382 people completed sufficient questions to be assigned a hip disease severity score. The mean age of the 23 people who were not able to be assigned a hip need score was 70, with a standard deviation of 18. This compares with a mean of 65 and standard deviation of 13 for those who were assigned a score.

As data on deprivation and urban/rural status were derived from postcodes, these data were available for a larger subset of respondents with a hip disease severity score (1381) than the details on social class (1366), income (1018), education (1155), car ownership (1144), living alone (1155), self assessed general health measures (1276) and affect of health on social life (1250), which were obtained using self completed questionnaires at clinic and home visits. For this reason, the data set for the main analysis comprises the set of people who completed all relevant questionnaires and questions (954). This is referred to as the “main set” throughout. For comparison, and to assess possible bias, the analyses were also carried out on the largest set available for each variable, the “full set” for each variable.

STATISTICAL ANALYSIS
Those with a hip prosthesis in situ were treated for the purposes of these analyses in exactly the same way as those without. The scores for each hip are not additive, and as there is no reason to suppose that one hip might have a different association with socioeconomic position than another, the analysis has been presented for the left hip only. The analysis was also carried out for the right hip and at person level, taking the score of each person’s most severe hip to indicate their maximum need. Any differences between findings using left hip, right hip and most severe hip are reported. The analysis was also separately carried out for the 354 men and 600 women in the main set and similar results were found.

A comparison of the main set with those excluded because of incomplete data was carried out using a two sample test for proportion (“`Prtest`” command in the statistical package STATX) for each of the sex, three age groups, socioeconomic position measured by enumeration district level Townsend score for material deprivation, and level of comorbidity.

The association of the age and sex standardised New Zealand score with measures of general health, and with individual and area measures of socioeconomic position, was assessed using Spearman’s rank correlation. Pearson’s product moment correlation coefficient was also used for continuous variables (age, enumeration district and ward level Townsend scores). t Tests assuming equal variances were used for two factor, categorical variables (car ownership, any further or higher education since leaving school, urban or rural status, manual/non-manual occupation and any comorbid, chronic conditions). The non-parametric Mann-Whitney U test was also carried out, and produced similar findings to the other methods.

To look at gradients in mean disease severity across the socioeconomic hierarchy, the continuous measure, enumeration level Townsend score for deprivation (deprivation category) was quantised. Social class was broken into five major groups according to the Registrar General’s classification of occupations, and education was presented as five groups, depending on qualifications and school leaving age.

A relative index of inequality, was calculated to measure the extent to which ill health is
systematically associated with socioeconomic position. The relative index of inequality can be interpreted as the health differential between the hypothetical person at the bottom of the socioeconomic hierarchy and the hypothetical person at the top of the hierarchy. For this analysis, each category of a socioeconomic indicator was ranked by socioeconomic position, beginning with the richest. The cumulative proportion of the population at the midpoint of each category was calculated to produce a range from 0 (richest) to 1 (poorest). This was regressed against age and sex standardised severity of hip disease to produce a coefficient that is the slope of the regression of a socioeconomic group’s relative morbidity on its relative rank. An F test for trend across the categories of each measure of socioeconomic position and self-assessed health with those who were excluded because of incomplete data is significant. The test reveals the difference across the groups is statistically significant.

Table 2 describes the characteristics in terms of age, gender, deprivation category (measured by Townsend score for material deprivation at enumeration district level) and comorbidity for the 954 people, for whom there are complete data and who comprise the main set for this analysis, and the 451 people who were excluded because of incomplete data. The poorest response rate among the clinic attendees with a hip severity score was to the question asking for self reported details of gross household income (74%). In the main set, there were almost twice as many women as men, 75% were aged 55 or above, 41% of the sample had no comorbidity and only 10% had multiple severe comorbidity. Those excluded from the main set because of incomplete data were similar in terms of gender, socioeconomic position measured by deprivation category and levels of comorbidity. The only statistically significant difference between the groups is that those excluded comprised a greater proportion of people aged 75+. This reflects the fact that many home visits were carried out in this age group and the response to self completed questionnaires on general health and socioeconomic position was not as good as home visits as it was for those who attended a study clinic.

Figure 1 illustrates the distribution of severity of left hip disease in the main set. For the left hip, 35% of the sample had a New Zealand score of below 10, indicating only mild pain or limitation while 9% of the sample had a score of 40 or above, suggesting severe disease.

**KEY POINTS**

- Lower socioeconomic position is associated with increased severity of hip disease.
- The association of socioeconomic position and severity of hip disease is seen with a variety of different indicators of socioeconomic position.
- Those with the lowest incomes experience more comorbidity, as well as greater severity of hip disease, than their richer counterparts.
- This additional dimension of inequalities in health—that of disease severity—reinforces the need to introduce policies to reduce such inequalities.

**Table 1** Age standardised prevalence per 100 of self reported musculoskeletal disorders (any joint) across the socioeconomic hierarchy, measured using the Townsend Score for material deprivation

<table>
<thead>
<tr>
<th>Category</th>
<th>Number</th>
<th>Prevalence</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>1st fifth (affluent)</td>
<td>2597</td>
<td>14.1</td>
<td>(9.6 to 18.7)</td>
</tr>
<tr>
<td>2nd fifth</td>
<td>2367</td>
<td>15.1</td>
<td>(10.6 to 19.6)</td>
</tr>
<tr>
<td>3rd fifth</td>
<td>1624</td>
<td>16.3</td>
<td>(10.9 to 21.8)</td>
</tr>
<tr>
<td>4th fifth</td>
<td>1696</td>
<td>16.1</td>
<td>(10.8 to 21.5)</td>
</tr>
<tr>
<td>5th fifth (deprived)</td>
<td>2024</td>
<td>17.7</td>
<td>(10.7 to 24.7)</td>
</tr>
</tbody>
</table>

Relative index of inequality (95% CI) 1.47 (1.22 to 1.77) **p<0.001**

Figure 1 Distribution of the New Zealand score for severity of hip disease (n=954).

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Table 3: Correlations of the crude score for severity of hip disease with age and the SEP at area level measures with measures of self reported health

<table>
<thead>
<tr>
<th>Measure of ill health</th>
<th>Total mean (SE)</th>
<th>number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Left</td>
<td>Right</td>
<td>55–59</td>
</tr>
<tr>
<td>Pain</td>
<td>165 (0.1)</td>
<td>165 (0.1)</td>
</tr>
<tr>
<td>Correlation coefficient</td>
<td>0.07*</td>
<td>0.07</td>
</tr>
</tbody>
</table>

***(p<0.01)***, ***(p<0.001)***. †Spearman’s rank correlation.

Table 4: Dichotomous indicators of socioeconomic position and comorbidity by severity of hip disease standardised for age

<table>
<thead>
<tr>
<th>Socioeconomic indicator</th>
<th>car available for use</th>
<th>any higher or further education</th>
<th>living alone</th>
<th>manual occupation</th>
<th>living in rural location</th>
<th>comorbidity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean New Zealand score of “no” group (SE)</td>
<td>17.5 (0.1)</td>
<td>17.5 (0.1)</td>
<td>17.5 (0.1)</td>
<td>17.5 (0.1)</td>
<td>17.5 (0.1)</td>
<td>17.5 (0.1)</td>
</tr>
<tr>
<td>Mean New Zealand score of “yes” group (SE)</td>
<td>15.4 (0.2)</td>
<td>15.4 (0.2)</td>
<td>15.4 (0.2)</td>
<td>15.4 (0.2)</td>
<td>15.4 (0.2)</td>
<td>15.4 (0.2)</td>
</tr>
<tr>
<td>Correlation coefficient</td>
<td>0.23**</td>
<td>0.23</td>
<td>0.23</td>
<td>0.23</td>
<td>0.23</td>
<td>0.23</td>
</tr>
</tbody>
</table>

***(p<0.01)***. ***(p<0.001)***. †Spearman’s rank correlation.

Table 5: Association of hip disease severity, standardised for age and sex, with socioeconomic position using individual and area level measures

<table>
<thead>
<tr>
<th>Socioeconomic position</th>
<th>Educational attainment</th>
<th>Income category</th>
<th>BDS level Townsend score</th>
</tr>
</thead>
<tbody>
<tr>
<td>Social class</td>
<td>mean (SE)</td>
<td>mean (SE)</td>
<td>mean (SE)</td>
</tr>
<tr>
<td>1 (most deprived)</td>
<td>18.0 (0.2)</td>
<td>18.0 (0.2)</td>
<td>18.0 (0.2)</td>
</tr>
<tr>
<td>Correlation coefficient (p value)</td>
<td>0.21 (&lt;0.05)</td>
<td>1.45** (1.29-2.62)</td>
<td>0.43 (0.43-1.94)</td>
</tr>
<tr>
<td>test for trend p value</td>
<td>0.51</td>
<td>&lt;0.01</td>
<td>&lt;0.01</td>
</tr>
</tbody>
</table>

***p<0.001***, ***(p<0.01)***. †Spearman’s rank correlation.
A comorbidity score of greater than 2.5.

A comorbidity score of greater than 1 but less than 2.5.

Table 6 Mean severity of hip disease, standardised for age and sex, within self assessed comorbidity categories by income (n=954)

<table>
<thead>
<tr>
<th>Comorbidity measure</th>
<th>None</th>
<th>One comorbid condition</th>
<th>Considerable comorbidity*</th>
<th>Multiple comorbidity†</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>mean (SE)</td>
<td>number</td>
<td>mean (SE)</td>
<td>number</td>
<td>mean (SE)</td>
</tr>
<tr>
<td>Income</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>£20 000+</td>
<td>13.9 (2.1)</td>
<td>97</td>
<td>15.6 (1.8)</td>
<td>55</td>
<td>14.9 (2.6)</td>
</tr>
<tr>
<td>£15 000-19 999</td>
<td>14.1 (2.2)</td>
<td>76</td>
<td>15.1 (2.1)</td>
<td>26</td>
<td>14.2 (2.1)</td>
</tr>
<tr>
<td>£10 000-14 999</td>
<td>15.7 (2.8)</td>
<td>93</td>
<td>15.8 (2.7)</td>
<td>64</td>
<td>16.6 (3.0)</td>
</tr>
<tr>
<td>£5000-9999</td>
<td>17.3 (2.4)</td>
<td>94</td>
<td>17.2 (2.6)</td>
<td>99</td>
<td>17.5 (2.7)</td>
</tr>
<tr>
<td>£1 500-2 999</td>
<td>16.3 (2.0)</td>
<td>79</td>
<td>16.8 (3.1)</td>
<td>50</td>
<td>17.7 (3.1)</td>
</tr>
<tr>
<td>£500-999</td>
<td>15.4 (2.9)</td>
<td>80</td>
<td>12.9 (3.0)</td>
<td>294</td>
<td>16.6 (3.0)</td>
</tr>
</tbody>
</table>

* A comorbidity score of greater than 1 but less than 2.5. † A comorbidity score of greater than 2.5.

Discussion

The study has replicated the finding that there is an association between socioeconomic position and morbidity and it has also allowed for the examination of how the severity of hip disease is distributed across the socioeconomic hierarchy, measuring socioeconomic position by a battery of different indicators. Our study adds to the growing body of work demonstrating the need to use multiple measures of socioeconomic position in health studies.14 15

INDIVIDUAL LEVEL INDICATORS OF SOCIOECONOMIC POSITION

The results show significant difference between mean hip disease levels using dichotomous individual level measures of socioeconomic position (any further or higher education, access to a car and living alone). Categorical variables (income and educational attainment) are also significantly associated with increasing severity of hip disease.

The systematic association of income and educational attainment with disease severity complements the literature on income inequalities in health measured at individual level and extends the findings of two large surveys in the United States that revealed that the prevalence of arthritis increases in those with fewer years of formal education.16 17

All individual level socioeconomic measures revealed an association of disease severity with ability to command resources at an individual level. After standardising for age and sex, however, the gradient in the means for social class disappeared. This may be explained by the fact that the majority of the main set are older women and many of these will not have had full time occupations. In such cases, a woman’s social class will have been ascertained from her husband’s last main occupation or, if this was not given, from her full time occupation that she held in her early adulthood or a part time position. In such cases, the social class assigned can only be a proxy for a woman’s socioeconomic position in later life.

ECOLOGICAL INDICATORS OF SOCIOECONOMIC POSITION

Despite previous work on this cohort (table 1) finding a systematic association between self reported musculoskeletal disease and area deprivation, this survey did not reveal any statistically significant association of the severity of mean hip disease with ecological measures of socioeconomic position (material deprivation and urban or rural location). In addition, there

The table also reports age and sex standardised relative indices of inequality. Both education and income are systematically associated with disease severity, with income having the strongest association, revealing that for each decrease in socioeconomic position (defined by education or income) there is an increase in disease severity. Analysis of the full set available for each variable produced similar means, medians and relative indices of inequality, with the association for income and education again proving highly statistically significant. (Tables are available from the first author on request).

Table 6 reveals that, using income as an indicator of socioeconomic position, poorer people not only experience a higher mean severity of hip disease but also greater comorbidity. Within comorbidity categories, disease severity increases as income decreases. In addition, the gradient between rich and poor is steepest among those with the greatest burden of comorbidity. Also within income bands, there is a general pattern of mean severity of hip disease increasing as comorbidity increases.

Despite the increase in the means for social class position (any further or higher education, access to a car and living alone). Categorical variables (income and educational attainment) are also significantly associated with increasing severity of hip disease.

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Individual level indicators of socioeconomic position

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Ecological indicators of socioeconomic position

Despite previous work on this cohort (table 1) finding a systematic association between self reported musculoskeletal disease and area deprivation, this survey did not reveal any statistically significant association of the severity of mean hip disease with ecological measures of socioeconomic position (material deprivation and urban or rural location). In addition, there
was also no gradient in median hip disease severity by area level measures. Although the differences between the self reported, binary variable and the severity score could be partially explained by the different types of measures being used, there may be other factors that explain why area level effects are weak in identifying inequalities in disease severity that are clearly observable at individual level. This could have important implications as the majority of health care resources are allocated at area level and underestimating the burden of disease among the poorest people may serve to exacerbate inequalities in provision.

While there is some evidence of people living in rural areas having better general health, and less chronic disease, another study found that those living in rural parishes in East Anglia had a higher proportion of acute and chronic morbidity. In contrast with these findings, this study found no difference in the severity of hip disease between those living in urban areas and those living in rural areas. Defining urban and rural status just as a binary variable based on postcode of residence may be too crude and mask any real differences as postcode groups cover large areas in rural parts of the country and a binary classification makes no distinction between inner city and suburban areas, and small market towns, classifying all of these as urban.

**THE NEW ZEALAND SCORE AND SELF REPORTED MORBIDITY**

The New Zealand priority setting tool was chosen because it incorporates a wide variety of variables collected at study clinics and covers pain, disability, comorbidity and the degree to which independence is threatened. The symptoms of a chronic disease, such as osteoarthritis, may change from day to day and even at different times within a day. In view of this, measuring morbidity using questionnaire data is difficult. Other scoring systems used in surgical practice were largely designed to assess the efficacy of treatment rather than need, and tend to have a narrower focus. The strength of the New Zealand Score for the purposes of this research is that it covers the broader issues around the impact of arthritis, including disability, handicap and degree to which independence is threatened.

As systematic underreporting of illness by people with lower socioeconomic position has been highlighted, the data were examined to ascertain whether there was any systematic difference in mean severity of hip disease across the socioeconomic hierarchy within categories of self assessed comorbidity. The data in table 6 reveal that for a person with a given hip disease score, a poor person is more likely to be also suffering more of the chronic comorbid conditions, or to rate their health as worse across the large number of questions asked, than one who is richer. This needs to be interpreted in the light of a Finnish study, which compared detailed health interview data with simple self reported questions on morbidity. Less educated people underestimated the prevalence of self reported arthritis, mental health problems and chronic respiratory disorders when compared with examination data. Dutch health surveys have shown large variations in prevalence of self reported health problems by education and income, and it has been suggested that if there is a socioeconomic differential in reporting bias, this leads to an underestimate of socioeconomic inequalities in health. As both countries have publically funded health care systems, it is probable that these findings would apply to the United Kingdom.

**LIMIATONS OF THE STUDY**

As the data include several different measures of socioeconomic position and self assessed general health, several caveats need to be made. Firstly, as is common in surveys, the income question had the poorest response despite care being taken to explain in the introduction the scope of the question. It is obviously a sensitive area and, in addition, requires a degree of mental arithmetic. Those excluded from the main analyses (largely because of missing data for the income question) were significantly more likely to be women and aged 55 or over and in view of the fact that arthritis is more common in women and increases with age, it is not surprising that the mean severity of hip disease was higher in this group (19.7) than in the main set (16.0), meaning that our findings, with regard to income in particular, may be an underestimate. In fact, the association of disease severity with socioeconomic position was stronger in the full set for all indicators, with income, maximum educational attainment, any higher education, car ownership and living alone again reaching statistical significance. This suggests that if anything, the associations reported here may be underestimated of the true associations.

Our findings are based on cross sectional data and without longitudinal data it is not possible to disentangle causal relations. In addition, there is increasing evidence to support the hypothesis that the whole of the life course influences current disease status. Ascertaining which came first, the hip disease or the low socioeconomic position, is not easy from these data but the breadth of measures of socioeconomic position can provide clues. While occupation, access to a car or living alone may be situations that change suddenly, lifetime educational attainment and gross household income are more general indicators of lifetime socioeconomic position. It is the latter two indicators that were the most strongly correlated with disease severity, suggesting that instead of disease leading to downward social mobility, other factors may...
predispose poorer people to more severe hip disease. For example, being in a manual occupation, such as farming, may be associated with low income, and may predispose to arthritis of the hip.6

FUTURE RESEARCH

There is now a considerable body of research exploring the factors underlying inequalities in health. However, similar analytical studies do not exist regarding differences in disease severity rather than prevalence. More research, both quantitative and qualitative, is needed to investigate the reasons for the greater severity of hip disease experienced by poorer people. Future research may answer the following questions: is it a reflection of inequities in access to and use of health care services; unmer for treatment; a large prevalence pool of people too ill with comorbidity to be operated on, or a result of broader sociological factors, and is this finding evident for other chronic conditions?

It seems that inequalities in a tax funded health care system occur not simply because of different persons’ ability to command health care resources,7 but because the poorest people experience a higher prevalence of morbidity but also because, for a common chronic disorder, such as disease of the hip, the pain and disability experienced by poorer people is more severe than that of their richer counterparts. This has implications for the modern National Health Service if it is to continue striving to direct care and resources to those in the most medical need.

In conclusion, these data reveal that the amount of pain and disability experienced by people with hip disease is associated with lower socioeconomic position, measured by a variety of indicators. People with lower socioeconomic position experience a greater severity of hip disease. As there is also a clear gradient of indicators. People with lower socioeconomic position experience a greater severity of hip disease but also a greater burden of comorbidity.

This study has implications for health care provision of the National Health Service to live up to its principle of equal treatment for equal medical need.

We would like to thank all study participants, the partners and practice staff of participating general practices. We are indebted to the whole of the Somerset and Avon Survey of Health research team and in particular: Kirsty Alchin, Ros Berkeley-Cobbold, Rev. Peter Bally, James Beardsmore, Chris Bannister, Simon Bax, Helen Bedson, Amanda Biggs, Sue Briscoe, Graeme Brown, David Bruce, Helen Butler, Rosemary Campbell, Sarah Cawley, Joanne Charnley, Mark Collins, Chris Denton, Andrew Doherty, Neil Edwards, Chris Evans, Sarah Eveleigh, Jane Flury, Sarah Foona, Sue Goldsmith, Steve Good, Steve Gough, Mike Grant, Glynne Griffiths, John Grimes, Chris Hall, Andrew Hales, Glyn Harper, Andrew Hart, Leslie Hay, Paul Jee, Sue Jones, Linda Jones, Jenny Jordan, Jude Keulemans, Fred Kriel, Louise Knight, Mark Knight, Mike Knights, Claire Krotki, Sue Kure, David Laflin, David Law, Karen Lee, Pete Leary, Mary Lister, Andrew Lomas, Nick Lowes, Margaret Mann, Sarah Marshall, Michael Maskell, cooker Martin, Simon Matthews, Andrew McAndrew, Sue McKie, Mike McRobbie, Andrew Mitchell, Shane Morgan, Robert Moss, Sue Mullen, Tony Neale, Helen Newton, Alison Norris, Andrea O’Connell, David O’Connor, Paul Osborne, Marcus Paick, Simon Paton, Richard Pilkington, Alan Pitt, Joffi Pounds, Shari Pyatt, Karen Quick, Beverley Quinlivan, Sarah Read, John Redfern, Peter Ridgway, Rick Rolfe, Stephen Rowles, Nick Rowlands, Andrew Rule, Sue Sanders, Vivienne Scrivens, Peter Selby, Jenny Scott, Leon Stockwell, Keith Stone, Mike Strange, Clive Thompson, Paul Tait, Paul Taylor, Peter Telford, Mike Thomson, Bridget Thorpe, Paul Torrance, Chris Trigell, Ann UF, Sarah Vickers, Steve Waddingham, Lorna Waters, Sue Williams and Andrea Wilson. We would also like to thank the whole of the Somerset and Avon Survey of Health team and in particular: Kirsty Alchin, Ros Berkeley-Cobbold, Rev. Peter Bally, James Beardsmore, Chris Bannister, Simon Bax, Helen Bedson, Amanda Biggs, Sarah Cawley, Joanne Charnley, Mark Collins, Chris Denton, Andrew Doherty, Neil Edwards, Chris Evans, Sarah Eveleigh, Jane Flury, Sarah Foona, Sue Goldsmith, Steve Good, Steve Gough, Mike Grant, Glynne Griffiths, John Grimes, Chris Hall, Andrew Hales, Glyn Harper, Andrew Hart, Leslie Hay, Paul Jee, Sue Jones, Linda Jones, Jenny Jordan, Jude Keulemans, Fred Kriel, Louise Knight, Mark Knight, Mike Knights, Claire Krotki, Sue Kure, David Laflin, David Law, Karen Lee, Pete Leary, Mary Lister, Andrew Lomas, Nick Lowes, Margaret Mann, Sarah Marshall, Michael Maskell, cooker Martin, Simon Matthews, Andrew McAndrew, Sue McKie, Mike McRobbie, Andrew Mitchell, Shane Morgan, Robert Moss, Sue Mullen, Tony Neale, Helen Newton, Alison Norris, Andrea O’Connell, David O’Connor, Paul Osborne, Marcus Paick, Simon Paton, Richard Pilkington, Alan Pitt, Joffi Pounds, Shari Pyatt, Karen Quick, Beverley Quinlivan, Sarah Read, John Redfern, Peter Ridgway, Rick Rolfe, Stephen Rowles, Nick Rowlands, Andrew Rule, Sue Sanders, Vivienne Scrivens, Peter Selby, Jenny Scott, Leon Stockwell, Keith Stone, Mike Strange, Clive Thompson, Paul Tait, Paul Taylor, Peter Telford, Mike Thomson, Bridget Thorpe, Paul Torrance, Chris Trigell, Ann UF, Sarah Vickers, Steve Waddingham, Lorna Waters, Sue Williams and Andrea Wilson.

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Conflicts of interests: none.
15 Pacca A, Colahan LF, Rushbrooke RV. More chronic diseases are reported more frequently by individuals with fewer than 12 years of formal education in the age 56–84 United States population. J Chronic Dis 1987;40:665–74.
La utilización de las evidencias en las decisiones de política sanitaria (The use of the evidence in health policy decisions)


This book is the result of a seminar held in April 2003 by the University of Alicante (Spain). The publisher and entire support team have made every effort to give the articles written by different authors a homogeneity linked works of a similar origin.

For some years, the health system, both the professionals working in it and the users and those responsible for its management, have been debating whether the decisions taken have a scientific basis.

However, some fields seem to be outside this movement causing some perplexity in the observer. One of these fields is public health and, precisely, health management. As discussed in the first chapter of this book, it is a field in which the evidence is scarce and contradictory and, in that framework, little can be done by those with the task of decision taking to give support to evidence that, very often, is no such thing.

The book has 10 chapters, with the first one dedicated to giving the reader a general view of the matter in question—the use of evidence applied to public health measures. The other nine contributions consider different examples of public health problems with their characteristics and—why not say so?—with their contradictions.

We are given interesting material for a debate as the decisions in this field are influenced not only by the scientific evidence, in many cases arguable as mentioned previously, but also by other factors of a social, cultural, and economic nature that cannot be forgotten.

Antonio Cueto Esparin

When food kills: BSE, E coli and disaster science


Pennington delivers serious messages in this discursive, thought-provoking book by sharing his insight into the failings of food safety (and other) inspectorates. Few have forgotten the hysteria associated with the Escherichia coli 0157 outbreak in central Scotland (linked to Barr’s butchers) or the salmonella in eggs scare when Edwina Currie was Parliamentary Under-Secretary for Health. The situations leading to these and other food poisoning and public health scares are eloquently described throughout, alongside a narrative on the apparent failures of government officials to learn from history through subsequent presentation of new health scares. It is proposed that vCJD cases in humans and BSE cases in animals are a result of these failures. Details regarding the scientific uncertainties over cause, transmission, and scope of these diseases are discussed prior to the UK government’s early assumption that BSE was not likely to be of risk to human health, the future risks to human health from vCJD, and the conclusions of the Phillips inquiry into methods used so far to eradicate BSE and vCJD.

The title indicates that food related incidents will be the principal subject of discussion; however this is not the case. Inferences are drawn throughout from events including the Aberfan tip mining disaster, the Piper Alpha North Sea oil disaster, and even the conditions within lunatic asylums since 1815. Repeated and detailed references to non-food disasters with catastrophic potential through higly-cycling and mobility levels, as compared with the innocuous/unknown long term health problems associated with food scares like BSE somewhat trivialises the importance to public health of food safety scares, and renders the title misleading.

Essentially, this is a non-technical book that describes (with reference to E coli, BSE, and other disasters) the history, the science, the politics, and most significantly, what went wrong. It may leave the reader concerned by the inspectorates’ shortcomings in the mitigation of public health incidents, but delivers an important message: inclusiveness and openness are essential to help avert wide scale disasters in the future.

Claire E Robertson

Health and community design: the impact of the built environment on physical activity


Sedentary behaviour is a major cause of poor health worldwide both through the direct effects of inactivity on health and indirectly via its contribution to obesity. Health and Community Design describes the role of the built environment as a potential contributor to physical inactivity and suggests ways in which communities could be structured to encourage or require physical activity, particularly walking and running.

The “built environment” is defined broadly as “…the form and character of communities” encompassing land use patterns, urban design characteristics, and transportations systems. Frank et al emphasise the influence of the built environment on physical activity and they effectively describe its potential role as a determinant of obesity. However, obesity is likely to be influenced by factors in addition to physical activity, particularly diet. This book highlights the need for a comprehensive assessment of how the built environment influences diverse determinants of energy balance, including diet, and other health behaviours. Occasionally, the authors neglect potential trade offs associated with choices concerning community design. For example, cul de sacs may decrease walking by adults but increase outdoor play of children. Understanding such trade offs is critical to improve planning and prioritisation among design choices.

All in all, we strongly recommend this book as an introduction to connections between urban planning and sedentary behaviour. The authors have done an outstanding job presenting arguments that can be made linking the built environment and physical activity and these arguments should be of great interest to public health, transportation and urban design researchers and planning professionals. The text is also accessible enough for community activists interested in understanding potential consequences of planning decisions and its maps and illustrations are particularly novel and effective for a public health audience.

Brooke Fischer, Sarah Dash, David Berrigan

Health measurement scales. A practical guide to their development and use, 3rd ed


This is the third edition of a successful book whose previous two editions were published in 1989, and 1994. It is a practical guide about health measurement scales called as well “latent outcomes”, such as cognitive abilities, attitudes, quality of life, etc, addressed to clinicians, users, and developers of health measurement scales.

In this third edition the authors have updated most of the chapters, mainly those related to cognitive requirements in answering questions, and include a more in depth chapter on item response theory.

The general content of the book follows the process of development and evaluation of health measurement scales. Chapters are devoted to the process of scale development, which includes basic concepts, how to devise and select items, and building scales. Chapters addressed to analyse attributes of the measures include reliability and validity, as well as measuring change. The chapters about ethical issues that researchers should take into account in their own fieldwork is also of valuable interest. Most chapters are accompanied by practical examples and a considerable number of tables and figures that make easy to understand and to interpret what authors want to explain. Appendices include commented bibliography and recommended reading, sources of developed scales, and a short introduction to exploratory and confirmatory factor analysis.

Through the whole book, authors also try to clarify their point of view on controversial
This book is a very interesting referent document for those who work in the public health research field. In fact it is more appropriate for researchers with some experiences in doing public health research rather than junior students. The readers can find some actual examples with in depth analysis on each case, which is very useful for them. However these examples and illustrations are more focused on American and African countries, so that it weakens the global and international application of the book.

Strengths: one of the strengths of this book is to identify and synthesise the key issues and principles for working with communities. It can be used as a theoretical frame for training courses on community based research. The contribution of this book is to emphasise the importance of community based research, which sometimes is forgotten by traditional epidemiological study. It also brings the sight and attention to the involvement of the community to research, change their role from target group to co-researcher, and from passive to active involvement.

Weaknesses: the authors tried to prepare a comprehensive document on community based research, which included all the issues like introduction, principles, methods, and examples/experiences. But the readers, after looking at the title of the book, are more interested in learning more specific issues and methods for community based research, and in how to distinguish it from public health research in general. It would have been more interesting if the authors had clarified more clearly the differences in methodology applications in epidemiological and community based research.


doi: 10.1136/jech.2003.014611corr1

There were two author errors (one terminological and one relating to data) in this paper by Dr Eachus and others (1999;33:603–11).

Firstly, the authors referred to the index relating socioeconomic position to New Zealand score of severity of hip disease as the relative index of inequality, whereas the statistical presented is actually the slope index of inequality. Secondly, a programming error led to miscalculation of the correlation coefficients and slope indices of inequality presented in table 5. A corrected table is presented here. The direction of associations is the same as for the incorrect results presented in the original paper, but the effect sizes and significance level are both substantially greater when the correct data are seen, in particular for the associations of social class and Townsend deprivation score with hip disease severity. In the light of the correct data the discussion that was included on why the social class association was weak is no longer applicable.

Table 5 (corrected) Association of disease severity, standardised for age and sex, with socioeconomic position using individual and area level measures

<table>
<thead>
<tr>
<th>Socioeconomic position</th>
<th>Social class</th>
<th>Educational attainment</th>
<th>Income category</th>
<th>ED level Townsend score quantile</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n mean (SE)</td>
<td>n mean (SE)</td>
<td>n mean (SE)</td>
<td>n mean (SE)</td>
</tr>
<tr>
<td>Most deprived</td>
<td>42</td>
<td>15.6 (2.5)</td>
<td>430</td>
<td>16.5 (2.8)</td>
</tr>
<tr>
<td>2</td>
<td>175</td>
<td>16.0 (3.0)</td>
<td>69</td>
<td>17.5 (2.8)</td>
</tr>
<tr>
<td>3</td>
<td>436</td>
<td>16.2 (2.9)</td>
<td>90</td>
<td>14.7 (5.6)</td>
</tr>
<tr>
<td>4</td>
<td>261</td>
<td>15.8 (2.9)</td>
<td>177</td>
<td>15.6 (2.7)</td>
</tr>
<tr>
<td>Most affluent</td>
<td>40</td>
<td>15.7 (2.7)</td>
<td>188</td>
<td>15.3 (2.9)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Correlation coefficient (p value)

|                         | SII         |                                    |                             |                               |
|                         | 5.63        | 3.79                               | 5.05                        | 4.04                           |
|                         | (2.23 to 9.04) | (0.41 to 7.18) | (1.78 to 8.32) | (0.04 to 8.13) |

* p<0.05; ** p<0.01. † Spearman’s rank correlation.

There were two author errors in this letter by Dr Eachus and others (1999;33:603–11).


doi: 10.1136/jech.2004.022871corr1

There were two author errors in this letter by I D K Dimoliatis (2004;58:1054–5). The unit of measurement in the title should be in lower case (q) [not (Q)]. Also, in line three of the second paragraph it should read (80–50) [not 80–30].
Aetiological heterogeneity of Alzheimer's disease

Editor,—It has regularly been noted that familial cases of Alzheimer's disease (AD) tend to be of relatively early onset. This implies that there must be at least one further aetiological class, comprising sporadic cases that share some common exposure, and that tend to develop later in life than the familial cases. It can now be reported that cases of AD attributable to residence in areas where the public water supply has a relatively high content of aluminum do constitute such a class.

The largest data set so far studied in relation to waterborne aluminum comprises records of 2258 age and sex matched pairs of patients discharged from hospitals in the Province of Ontario during 1985, one member of each pair having a diagnosis of AD. Age at diagnosis was not documented for these cases but it is reasonable to assume that hospital patients with this diagnosis who were older at the time of discharge had also been older, on the average, at the time of onset.

Preliminary analysis of the Ontario data showed that the relative prevalence of hospitalised AD cases increased monotonically through four ranges of aluminum concentration in the public water supply (from <0.1 mg/l to >0.200 mg/l). With the four ranges of aluminum collapsed to two (under and over 0.1 mg/l) these data have now been broken out by age and sex to obtain the results shown in table 1. Although the overall risk of AD in areas with the higher aluminum concentration was only 28%, five of the six estimates for ages beyond 75 were higher than this.

The statistical significance of the apparent age pattern is readily assessed by means of a permutational test procedure. Considering that there are 4+3+2+1 = 10 older-younger comparisons available within each sex, and a further 20 between older and younger age groups of unlike sex, we note that 29 of 40 such comparisons in table 1 show a higher relative risk in the older age group. A computer program was used to generate re-assignments of the estimated relative risk values to the 10 non-overlapping cells of table 1 and to count, for each such permutation, how many of the 40 older-younger comparisons had the higher value for the older group.

After drawing a random sample (but with replacement) of 30 000 from the total of 3 628 800 possible permutations, there were found to have been only 1209 (4.03%) in which 29 or more comparisons had the higher value in the older group. Thus there is conventionally significant evidence against the null hypothesis of no age related aluminum effect and favouring the alternative hypothesis of an aluminum effect more strongly expressed in the age range where it has been judged—on completely separate evidence—that cases of AD are less genetically determinate.

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A survey of smoking among Italian doctors

Editor,—A survey of smoking in a representative sample of Italian doctors was conducted by the Istituto Doxa (the Italian branch of the Gallup Group) between 20 and 22 January 1995, within the framework of a European project coordinated by the British Medical Association. The sample was obtained from the official lists of the Federazione Nazionale degli Ordini dei Medici Chirurghi e degli Odontoiatri, stratified by age, sex and geographical area.

A total of 501 doctors (384 men, 117 women, mean age 45 years) were interviewed; 218 were general practitioners. Table 1 gives their distribution according to smoking status, sex and age group. Of these, 121 (24.2%) were current smokers of >1 cigarettes per day, 17 (3.4%) were occasional smokers, 133 (26.5%) ex-smokers, and 230 (45.9%) never smokers. With reference to number of cigarettes smoked, 12.8% reported smoking <15 cigarettes per day, 9.4% 15 to 24, and 2.0% ≥25 cigarettes per day. The prevalence of current smoking was similar in men (24.5%) and women (23.1%), and in subsequent age groups. Women, however, were more frequently never smokers (37.3% versus 42.4%), and less frequently ex-smokers. Likewise, younger doctors (<40 years) were more frequently never smokers (58.2%).

The overall smoking prevalence among Italian doctors is thus similar to that of the general population of Italian adults (25.3% in 1995). While male doctors smoked less frequently than the male general population (24.5% versus 34.1%), female doctors smoked more frequently than the female general population (23.1% versus 17.2%).

The main message of this survey is that smoking prevalence remains exceedingly high among Italian doctors, and comparable to the general Italian population for both sexes combined, although it has declined since the mid-1980s. Indeed, female Italian doctors smoke more than their general population counterpart.

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Table 1 Estimated relative risk of Alzheimer’s disease associated with residence in areas having a concentration of aluminum in municipal drinking water of 0.1 mg/l or more. Based on subsets of the Ontario data classified by age and sex

<table>
<thead>
<tr>
<th>Age group, years</th>
<th>Number of cases/control pairs</th>
<th>Estimated relative risk: men</th>
<th>Estimated relative risk: women</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;70</td>
<td>325</td>
<td>1.06</td>
<td>1.05</td>
</tr>
<tr>
<td>70–74</td>
<td>381</td>
<td>1.05</td>
<td>1.22</td>
</tr>
<tr>
<td>75–79</td>
<td>507</td>
<td>1.60</td>
<td>1.29</td>
</tr>
<tr>
<td>80–84</td>
<td>547</td>
<td>1.29</td>
<td>1.47</td>
</tr>
<tr>
<td>&gt;85</td>
<td>498</td>
<td>1.24</td>
<td>1.46</td>
</tr>
</tbody>
</table>

Table 1 Smoking habits among a representative sample of 501 Italian doctors, 1999

<table>
<thead>
<tr>
<th>Smoking status</th>
<th>Total</th>
<th>Men</th>
<th>Women</th>
<th>Age group, years</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>%</td>
<td></td>
<td></td>
<td>% Number</td>
</tr>
<tr>
<td>Current smokers</td>
<td>24.2</td>
<td>24.5</td>
<td>23.1</td>
<td>20.0</td>
</tr>
<tr>
<td>Occasional smokers</td>
<td>3.4</td>
<td>3.6</td>
<td>3.4</td>
<td>3.6</td>
</tr>
<tr>
<td>Ex-smokers</td>
<td>26.5</td>
<td>29.4</td>
<td>17.1</td>
<td>29.4</td>
</tr>
<tr>
<td>Never smokers</td>
<td>45.9</td>
<td>42.4</td>
<td>57.3</td>
<td>42.4</td>
</tr>
</tbody>
</table>

*Less than one cigarette per day.

Correction

An authors’ error occurred in the paper by Dr Eachus and others (1999;53:603–11). The index of inequality is incorrectly called the relative index of inequality, while what is reported is the slope index of inequality.