
Downloaded from: http://researchonline.lshtm.ac.uk/1495/

DOI: https://doi.org/10.1136/jech.2005.037515

Usage Guidelines:

Please refer to usage guidelines at https://researchonline.lshtm.ac.uk/policies.html or alternatively contact researchonline@lshtm.ac.uk.

Available under license: http://creativecommons.org/licenses/by-nc-nd/2.5/
Ethnic differences in cause specific mortality among hospitalised patients with diabetes: a linkage study in New Zealand

Mona Jeffreys, Craig Wright, Andrea ‘t Mannetje, Ken Huang, Neil Pearce

Study objective: To describe patterns of excess mortality among patients with diabetes in three ethnic groups.

Design: A linkage study of national hospital discharge records to death records.

Setting: New Zealand.

Participants: The study included 74 847 patients (11 268 Māori, 5730 Pacific, and 57 849 non-Māori/non-Pacific) aged over 25 years with a hospital discharge diagnosis of diabetes between 1988 and 2001. By the end of 2001, 29 295 (39%) of the cohort had died. Based on the underlying cause of death, standardised mortality ratios (SMRs) [95% confidence intervals] were calculated for each ethnic group and sex.

Main results: Comparing the mortality patterns of patients with diabetes to the general population of the same ethnic group, adjusting for age and calendar period, all cause SMRs were higher for Māori women and men: 3.80 (95% CI: 3.64 to 3.97) and 3.44 (95%CI: 3.30 to 3.58) than for Pacific (men: 2.41 (95%CI: 2.21 to 2.61); women: 2.23 (95%CI: 2.06 to 2.41)) and non-Māori/non-Pacific (men: 2.98 (95%CI: 2.93 to 3.04); women: 2.99 (95%CI: 2.93 to 3.04)) people. SMRs were significantly raised for several causes of death, including cardiovascular disease and many site specific cancers.

Conclusions: The pattern of excess mortality among Māori with diabetes may relate to severity of disease. This needs further investigation, as the excess mortality may be amenable to intervention.

M ortality rates among people with type 2 diabetes are higher than the general population. Estimates of the magnitude of this excess mortality vary widely, and although the differences are partially dependent on the sources of the diabetes cases, such selection factors do not seem to explain all of the reported differences in mortality. Studies that identify diabetes patients through primary and secondary care sources show slightly increased (25% to 50%) standardised mortality ratios (SMR), although other population based studies report SMRs or relative risks (RR) of about twice that of the general population.3–5 Studies based on patients with diabetes identified through hospital records report higher SMRs,5 presumably because of the selective inclusion of more severe cases of diabetes or patients with comorbidities. Basing diagnosis on insulin use in type 2 diabetes also results in high all cause SMRs.5

Most previous reports examine broad categories of causes of death, and there is general consensus that excess mortality is evident for cardiovascular and renal disease.1,6–7 Cancer mortality may also be increased,6 although not all studies have found this.4,9 and some studies have reported lower SMRs for cancer mortality in patients with diabetes.9 There are few reports that investigate site specific cancer mortality in patients with diabetes.

The prevalence of diabetes in New Zealand differs by ethnicity. A national survey of self reported doctor diagnosed diabetes found lowest rates in New Zealand Europeans (2.5% in women and 3.1% in men), and higher rates in Māori (9.8% in women and 8.0% in men) and Pacific people (8.2% in women and 6.8% in men).10 These differing rates across ethnic groups are only partly explained by differing levels of obesity.17 The aim of this study was to describe cause specific mortality among patients admitted to a public hospital in New Zealand, and to investigate whether these patterns differed by ethnicity.

METHODS

The study entailed linkage between electronic hospital discharge data and mortality data. Patients included in the study were New Zealand residents who were discharged from a public hospital with a code for diabetes mellitus, as recorded on the National Minimum Dataset (NMDS) between 1988 and 2001 and were aged 25 or over at the time of discharge. Note that the included patients may or may not have been diagnosed with diabetes before admission, but all had been diagnosed with diabetes at the time of discharge. The International Classification of Disease (ICD-9) codes used to define diabetes mellitus were 2500 to 2509. People were identified using the National Health Index (NHI) identifier, a number assigned to each person using health and disability support services. For people with multiple diabetes discharges, the first admission during the period 1988 to 2001 was used.

Ethnicity was assigned using the recorded ethnicity in the NHI database, prioritised using the Māori, Pacific, non-Māori/non-Pacific groups.12 The prioritised system minimises the possibility of undercounting Māori and Pacific people. The non-Māori/non-Pacific group is predominantly of European origin, although does include a significant proportion (about 7%) of Asians.

The cohort of patients with a discharge summary including diabetes was linked to the national death registration data held by the New Zealand Health Information Service.
Follow up was from the date of the diabetes admission until the earlier of either date of death or 31 December 2001, the date for which cause of death data were available at the time of linkage. The underlying cause of death was used for all analyses.

Statistical methods
SMRs were calculated by comparing the mortality rate among patients with diabetes to the sex and ethnic specific mortality rates of the New Zealand population. The rates used for comparison were split by five year age bands and two year calendar periods. Analysis was performed using the PC life table analysis system. All reported p values are two sided.

RESULTS
Between February 1988 and December 2001, 76 684 people were discharged from hospital with a code for type 2 diabetes on their discharge summary (see above for clarification). The following exclusions were made: not New Zealand residents on their discharge summary (see above for clarification). The people were discharged from hospital with a code for type 2 diabetes between February 1988 and December 2001, 76 684 people (37 118 women and 37 729 men). These people were followed up for a maximum of 13 years. By the end of 2001, 29 295 (39.1%) cohort members had died.

Table 1 shows the characteristics of the study participants. The median age at index discharge was about 13 years lower for Māori and Pacific people compared with non-Māori/non-Pacific people. The gap in median age at death between the ethnic groups was a correspondingly wide, although Pacific people, who were, on average, slightly younger than Māori at index diagnosis, were older at death than Māori.

The all cause SMR for all patients combined was 3.02 (95% confidence intervals (CI): 2.99 to 3.06). Significantly raised SMRs were found for all major causes of death, including all cancers combined (2.40, 95% CI: 2.34 to 2.47), diseases of the circulatory (2.82, 95% CI: 2.77 to 2.87), respiratory (1.97, 95% CI: 1.89 to 2.06), digestive (3.05, 95% CI: 2.85 to 3.26), and genitourinary (3.31, 95% CI: 3.04 to 3.60) systems, as well as external causes (1.99, 95% CI: 1.81 to 2.19). Site specific cancer analyses showed that the sites at which diabetic patients were most at risk, with an SMR exceeding 3.0, were liver, gallbladder, pancreas, oro-pharynx, brain and nervous system, and non-thyroid endocrine glands. However, significantly raised SMRs (p<0.01) were evident for almost all sites, including important sites of non-smoking related

| Table 1 | Description of all patients aged 25 years or more discharged from New Zealand public hospital with diabetes, 1988 to 2001 |  |
|---------|-------------------------------------------------|---|---|
| Women | | | |
| Age at diabetes discharge | Māori | 5654 | 58.0 | 25.0 to 95.5 | <0.001 | 5614 | 57.7 | 25.2 to 93.9 | <0.001 |
| | Pacific | 3220 | 57.7 | 25.0 to 99.3 | <0.001 | 2510 | 57.5 | 25.2 to 97.1 | <0.001 |
| | Non-Māori/non-Pacific | 28244 | 72.1 | 25.0 to 105.9 | | 29605 | 69.3 | 25.0 to 102.9 | |
| Age at death | Māori | 2003 | 66.3 | 29.1 to 96.7 | <0.001 | 2220 | 64.5 | 25.3 to 95.4 | <0.001 |
| | Pacific | 578 | 69.4 | 32.6 to 99.5 | <0.001 | 621 | 66.5 | 25.9 to 97.2 | <0.001 |
| | Non-Māori/non-Pacific | 11400 | 79.8 | 27.9 to 108.5 | | 12473 | 76.3 | 26.0 to 105.1 | |

Table 2 | Cause specific standardised mortality ratios among 37118 women hospitalised for diabetes, New Zealand, 1988 to 2001 |  |
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Cause of death (ICD-9)</td>
<td>Māori (n = 5654)</td>
<td>Pacific (n = 3220)</td>
<td>Other (n = 28244)</td>
</tr>
<tr>
<td>Deaths</td>
<td>SMR</td>
<td>95% CI</td>
<td>Deaths</td>
</tr>
<tr>
<td>All deaths</td>
<td>2003</td>
<td>3.80</td>
<td>3.64, 3.97</td>
</tr>
<tr>
<td>Infectious and parasitic diseases</td>
<td>15</td>
<td>3.17</td>
<td>1.77, 5.23</td>
</tr>
<tr>
<td>All cancers (150-208)</td>
<td>433</td>
<td>2.85</td>
<td>2.59, 3.14</td>
</tr>
<tr>
<td>Endocrine, nutritional, and metabolic diseases (240-259, 270-279)</td>
<td>524</td>
<td>11.36</td>
<td>10.41, 12.37</td>
</tr>
<tr>
<td>Mental disorders (290-319)</td>
<td>6</td>
<td>1.32</td>
<td>0.48, 2.88</td>
</tr>
<tr>
<td>Diseases of nervous system (320-359)</td>
<td>11</td>
<td>3.58</td>
<td>1.79, 6.41</td>
</tr>
<tr>
<td>Diseases of circulatory system (320-399)</td>
<td>665</td>
<td>3.03</td>
<td>2.81, 3.27</td>
</tr>
<tr>
<td>Diseases the respiratory system (460-519)</td>
<td>126</td>
<td>2.17</td>
<td>1.81, 2.59</td>
</tr>
<tr>
<td>Diseases of digestive system (520-579)</td>
<td>52</td>
<td>4.39</td>
<td>3.28, 5.75</td>
</tr>
<tr>
<td>Diseases of genitourinary system (580-629)</td>
<td>55</td>
<td>5.87</td>
<td>4.42, 7.64</td>
</tr>
<tr>
<td>Diseases of skin and subcutaneous tissue (680-709)</td>
<td>6</td>
<td>3.71</td>
<td>1.36, 8.08</td>
</tr>
<tr>
<td>Diseases of musculoskeletal system and connective tissue (710-739)</td>
<td>9</td>
<td>4.57</td>
<td>2.09, 8.68</td>
</tr>
<tr>
<td>External causes (E800-E999)</td>
<td>17</td>
<td>1.96</td>
<td>1.14, 3.14</td>
</tr>
</tbody>
</table>

SMR, standardised mortality ratio; CI, confidence interval.
cancers such as colorectal, breast, prostate, urinary, and haematopoietic system.

Sex and ethnic specific SMRs for the major categories of cause of death and also for specific subcategories of particular interest are shown in table 2 (women) and table 3 (men). SMRs, comparing the mortality experience among people hospitalised for diabetes to the general population of the same ethnicity, differed across the three ethnic groups. The lowest excess mortality was in Pacific people, intermediate in Māori/non-Pacific group, the greatest contribution to the excess mortality in patients with diabetes. Because of the completeness of the data in the NMDS, we restricted our analyses to discharges from public hospitals, at which access to care is free. We estimate that about 2% of all diagnoses of diabetes may have been omitted from our analyses through the omission of private hospital diagnoses. It is unlikely that this will have had a large impact on our results. We chose to use a cut off of age 25 years at hospital discharge to define adult onset disease. This is likely to be reasonably reliable in the later years of the data, because someone who was discharged from hospital for the first time and had not been discharged in the previous 10 years is likely to have been comparatively recently diagnosed. However, for the earlier years of the data, we are likely to have included a proportion of patients with type 1 diabetes, if they had been incorrectly coded as type 2 instead of type 1. This could lead to diabetes. Nor could we exclude the possibility of detection bias, as patients could have been admitted to hospital for a reason related to their cause of death, and had diabetes diagnosed for the first time in hospital, thus creating a spurious association between diabetes and mortality.

**DISCUSSION**

The findings from our study are in agreement with a number of previous studies, showing increased risks of all cause and cardiovascular disease mortality in a cohort of patients discharged from hospital with diabetes. Because of the large size of the cohort, we were also able to explore cause specific mortality, which was found to be raised in comparison with the general population for most causes of death. Importantly, we found that the pattern of excess mortality, although broadly similar across ethnic groups, was substantially higher in Māori and lower in Pacific people compared with non-Māori/non-Pacific people for cardiovascular disease, the single greatest contributor to the excess mortality in patients with diabetes.

The study was based only a cohort of diabetes patients who were hospitalised, thus including those patients with more severe disease than a community based sample. This raises the potential for reverse causality, as we are unable to determine either the temporality of the onset of diabetes and other comorbidities that could underlie the cause of death. This issue may be particularly important for the results for cancer mortality, as cancer treatments could lead to diabetes. Nor could we exclude the possibility of detection bias, as patients could have been admitted to hospital for a reason related to their cause of death, and had diabetes diagnosed for the first time in hospital, thus creating a spurious association between diabetes and mortality.

The study was based only a cohort of diabetes patients who were hospitalised, thus including those patients with more severe disease than a community based sample.

**Table 3** Cause specific standardised mortality ratios among 37729 men hospitalised for diabetes, New Zealand, 1988 to 2001

<table>
<thead>
<tr>
<th>Cause of death (ICD-9)</th>
<th>Māori (n = 5614)</th>
<th>Pacific (n = 2510)</th>
<th>Other (n = 29605)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Deaths</td>
<td>SMR</td>
<td>95% CI</td>
<td>Deaths</td>
</tr>
<tr>
<td>All deaths</td>
<td>2220</td>
<td>3.44</td>
<td>3.30, 3.58</td>
</tr>
<tr>
<td>Infectious and parasitic diseases (001–199)</td>
<td>22</td>
<td>3.76</td>
<td>2.56, 5.70</td>
</tr>
<tr>
<td>All cancers (150–208)</td>
<td>414</td>
<td>2.26</td>
<td>2.05, 2.49</td>
</tr>
<tr>
<td>Mental disorders (290–319)</td>
<td>6</td>
<td>1.32</td>
<td>0.48, 2.87</td>
</tr>
<tr>
<td>Diseases of nervous system (320–399)</td>
<td>11</td>
<td>2.80</td>
<td>1.40, 5.01</td>
</tr>
<tr>
<td>Diseases of circulatory system (400–499)</td>
<td>860</td>
<td>3.08</td>
<td>2.87, 3.39</td>
</tr>
<tr>
<td>Diseases of respiratory system (460–519)</td>
<td>113</td>
<td>1.91</td>
<td>1.57, 2.29</td>
</tr>
<tr>
<td>Diseases of digestive system (520–579)</td>
<td>55</td>
<td>3.78</td>
<td>2.85, 4.92</td>
</tr>
<tr>
<td>Diseases of genitourinary system (580–629)</td>
<td>40</td>
<td>3.51</td>
<td>2.51, 4.78</td>
</tr>
<tr>
<td>Diseases of skin and subcutaneous tissue (680–709)</td>
<td>7</td>
<td>6.56</td>
<td>2.63, 13.52</td>
</tr>
<tr>
<td>Diseases of musculoskeletal system and connective tissue (710–739)</td>
<td>5</td>
<td>2.58</td>
<td>0.84, 6.03</td>
</tr>
<tr>
<td>External causes (E800–E999)</td>
<td>38</td>
<td>1.92</td>
<td>1.36, 2.63</td>
</tr>
</tbody>
</table>

SMR, standardised mortality ratio; CI, confidence interval.
explain the comparatively high all cause SMR that we report in comparison with other studies, as SMRs tend to be higher for type 1 than type 2 diabetes.4

Although the recording of diabetes on death certificates of people with diabetes is notoriously low, both in New Zealand14 and internationally, this will not affect the results that we present for cardiovascular disease or cancer, because there is no reason to think that these causes of disease are differentially underreported in patients with and without diabetes. Similarly, although the high prevalence of undiagnosed patients with suspected diabetes in New Zealand is high,15 this should not affect the results from the sample on which the analysis is based.

Previous studies of mortality among diabetic patients have found similar results to ours, with excess mortality from all causes, cardiovascular disease, and renal disease evident in most reports.1-8 The evidence for the presence of a positive association between diabetes and cancer mortality is less consistent. Many studies have shown that patients with diabetes are at a higher risk of liver6 and pancreatic cancer,16-17 with SMRs of over threefold commonly reported. There is less consistency in the reported associations between cancer mortality in populations with type 2 diabetes compared with the general population. Several of the cohort studies that have failed to find an association between diabetes and all cancer mortality include a very small number of cases, such as a follow up of the NHANES-1 survey (486 deaths among patients with diabetes)18 the Whitehall cohort of men (188 deaths)17 compared with over 2500 in our study. A Swedish study of similar design to ours reported an SMR of 1.46 (95% CI: 1.40, 1.52) in their group of hospitalised patients who had “probably NIDDM”. A SMR of similar magnitude was found in the probable IDDM group.

Associations between diabetes and cause specific mortality may be evident because of shared common risk factors (for example, smoking) or an influence of lack of glycaemic control on mortality.
control on vascular or other tissues. It is probable that associations between smoking related cancers and diabetes may be attributable to the common cause of smoking in these cancers and the development of diabetes.\textsuperscript{15, 16} However, such obvious confounding is less clear for non-smoking related cancers.

There is accumulating evidence that colorectal cancer risk is higher in patients with diabetes, and in particular this increased risk may be related to glycaemic control.\textsuperscript{17} Such findings shed important light on the role of the insulin/insulin-like growth factor (IGF) axis in the development of or survival from these cancers. One of the symptoms of insulin resistance is high levels of circulating insulin, which are related to increased bioactivity of IGF-I, as well as to its bioavailability through the association with IGF binding proteins.\textsuperscript{18, 19} Hyperinsulinaemia has been related to several hormone responsive cancers,\textsuperscript{20} particularly of the colorectum, endometrium, and breast.\textsuperscript{21} Several previous studies (reviewed in Rodriguez et al\textsuperscript{12}) have shown lower risks of prostate cancer in patients with diabetes compared with the general population. We have found moderately raised SMRs for prostate cancer, consistent with the suggestion that high insulin levels present during insulin resistance and the early phase of diabetes are associated with prostate cancer risk. It has been proposed that the relation between diabetes and prostate cancer is modified by duration of diabetes, with higher risks of prostate cancer in the early years after a diabetes diagnosis reducing to lower risks in later years. Given the decades during which prostate cancer develops, these data suggest that raised insulin levels may act as a promoter of prostate cancer, rather than having a carcinogenic effect in itself.

Some studies have reported sex differences in the mortality experiences of diabetic patients. However, most larger studies, such as NHANES\textsuperscript{22} and the Swedish linkage study\textsuperscript{23} found no evidence to suggest sex differences in mortality among diabetics. In this study, most off the site specific SMRs were similar in women and men. An important finding among diabetics. In this study, most off the site specific SMRs were particularly high for Māori, the indigenous population of New Zealand, but not for Pacific people living in New Zealand.

The results suggest that Māori admitted to hospital with diabetes may have more severe disease that non-Māori, which could be attributable to differential access to primary care for Māori with diabetes.

What this paper adds

\begin{itemize}
  \item A pattern of high mortality rates from cardiovascular disease, renal disease, and cancer was found in hospitalised patients with diabetes compared with the general population.
  \item These SMRs were particularly high for Māori, the indigenous population of New Zealand, but not for Pacific people living in New Zealand.
  \item The results suggest that Māori admitted to hospital with diabetes may have more severe disease that non-Māori, which could be attributable to differential access to primary care for Māori with diabetes.
\end{itemize}

In summary, we have reported high SMRs for several causes of death in a cohort of patients admitted to hospital for diabetes. The findings with respect to cancer add to our understanding of the aetiology of non-smoking related cancers. The pattern of excess mortality by ethnicity should be further investigated, as this may be amenable to intervention to improve outcomes for Māori.

Authors’ affiliations

M Jeffreys, A ’t Mannetje, K Huang, N Pearce, Centre for Public Health Research, Massey University, Wellington, New Zealand

C Wright, K Huang, Public Health Intelligence, Ministry of Health, Wellington, New Zealand

Funding: the Centre for Public Health Research is supported by a program grant from the Health Research Council of New Zealand.

Competing interests: none declared.

REFERENCES


THE JECH GALLERY

Sir John Pringle’s Observations on the Diseases of the Army—an early scientific account of epidemiology and the prevention of cross infection

Born in 1707, Pringle studied medicine at Leiden and Paris before practising in Edinburgh. In 1744 he became physician general to the British army and he was present at the battle of Culloden. In 1752 the first of the many editions of Pringle’s Diseases of the Army was published. In the volume Pringle identified hospitals as a major cause of sickness and began to think in terms of septic and antiseptic. Pringle felt that fevers, dysentery, and jail fever were the three most prevalent and fatal diseases affecting armies and although he was wrong about the way they were transmitted (he blamed putrid air) his recommendations for prevention helped to control epidemics. After his retiral from the army Pringle started a successful London practice. He was a president of the Royal Society and physician-in-ordinary to the king. Sir John died in 1782.

FURTHER READING

- Pringle’s historic manuscripts on display http://news.bbc.co.uk/1/hi/scotland/4007407.stm

I Milne
Library and Information Services, Royal College of Physicians of Edinburgh, 9 Queen Street, Edinburgh EH2 2JQ, UK
i.milne@rcpe.ac.uk

Illustration courtesy of the Royal College of Physicians of Edinburgh.