Plasma Glucose Regulation and Mortality in Pima Indians

Nan Hee Kim, MD, PhD; Meda E. Pavkov, MD, PhD; Helen C. Looker, MBBS*; Robert G. Nelson, MD, PhD; Peter H. Bennett, MB, FRCP; Robert L. Hanson, MD, MPH; Jeffrey M. Curtis, MD, MPH; Maurice L. Sievers, MD; William C. Knowler, MD, DrPH

Diabetes Epidemiology and Clinical Research Section, National Institute of Diabetes and Digestive and Kidney Diseases, National Institutes of Health, Phoenix, Arizona.
* Division of Endocrinology, Mount Sinai School of Medicine, New York, New York

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Correspondence:
Dr Nan-Hee Kim
National Institutes of Health
1550 East Indian School Road
Phoenix, AZ 85014-4972 USA
kimnanhee@niddk.nih.gov

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ABSTRACT

Objective: To evaluate whether impaired fasting glucose (IFG) and/or impaired glucose tolerance (IGT) are associated with increased risk of mortality and prevalent ischemic heart disease (IHD); to analyze if the increased risk of death is dependent on subsequent development of diabetes in Pima Indians.

Research Design and Methods: 2,993 Pima Indians aged 35 years and older were included. Prevalent IHD, defined by major ischemic ECG changes, was evaluated according to the following glucose/diabetes categories: normal glucose regulation (NGR), IFG and/or IGT, and diabetic groups by duration. During a median follow-up of 10.4 years, 780 subjects died from natural causes and 156 of these from IHD. Mortality was analyzed according to the same glucose/diabetes categories at baseline and then as time-dependent variables.

Results: Only subjects with diabetes ≥15 years of duration have a higher prevalence of IHD (odds ratio=1.9, 95% CI=1.4-2.5) relative to NGR. In baseline and time-dependent models, age-sex-adjusted death rates from natural causes and from IHD were similar among the nondiabetic groups. Among diabetic subjects, natural mortality was higher in those with ≥15 years diabetes duration (death rate ratio (DRR) relative to NGR=2.6, 95% CI=2.1-3.3). IHD mortality was higher in subjects with long diabetes duration (DRR for diabetes 10-15 yrs=3.8, 95% CI=1.5-9.5; DRR for diabetes ≥15 years=8.6, 95% CI=3.8-19.4) in the time-dependent model.

Conclusions: Natural and IHD mortality are not increased in Pima Indians with IFG and/or IGT. Only after the onset of diabetes do the rates of these events increase relative to NGR.
Considerable evidence indicates that chronic hyperglycemia is a risk factor for natural cause and IHD mortality and for incident IHD (1-4). Low glucose concentration has also been implicated as a risk factor for increased overall and cardiovascular disease (CVD) mortality, including IHD (3). There is some controversy, however, concerning the relative pathophysiological roles of fasting and post-load hyperglycemia in nondiabetic persons. Some epidemiologic studies support the importance of one or the other (1,2) and a few indicate that neither IFG nor IGT confer increased risk for mortality or incident IHD (5), or do not confer increased risk after accounting for associated cardiovascular risk factors (6,7). However, most previous studies consider only the baseline glucose levels and do not account for the change of glucose tolerance status and the development of diabetes during follow-up, so that the risk of IFG or IGT on mortality might be overestimated in these studies. Development of diabetes is reported to confound the relationship between baseline IFG and mortality (8), but not the relationship between baseline IGT and mortality (9).

In the Pima Indians, diabetes has a major impact on mortality and causes of death, and the death rate from IHD has increased in recent years in those with diabetes (10). Sufficient numbers of follow-up examinations permit us to account precisely for the future development of diabetes. The specific purposes of this study are to assess the association between prevalent IHD based on major ischemic ECG changes and plasma glucose and diabetes duration categories, to compare death rates from natural causes and from IHD among these plasma glucose and diabetes duration categories, and to determine if risk of mortality in IFG and/or IGT groups is dependent on development of diabetes during follow-up.

**RESEARCH DESIGN AND METHODS**

Pima Indians and the closely related Tohono O’odham Indians who live in the Gila River Indian Community in central Arizona participate in a longitudinal study of diabetes. Since 1965, members of the community ≥5 years old have been invited to participate in research examinations approximately every 2 years. These examinations included measurements of venous plasma glucose obtained 2h after a 75-g oral glucose load (2hPG); measurement of fasting plasma glucose (FPG) was added in 1975. The present analysis included persons who resided in the community at any time between January 1, 1975 and December 31, 2003 and had one or more research examinations after 35 years of age during this period with a diagnosis of diabetes or measurement of both FPG and 2hPG. Each subject’s vital status as of December 31, 2003 was determined. Causes of death were determined by review of clinical records, autopsy reports and death certificates. Terminology and codes of the International Classification of Disease, Ninth Revision (ICD-9), were used to classify the underlying cause of death as due to natural causes (ICD-9 codes 001.0-799.9) and IHD (ICD-9 codes 410.0-414.9). Death rates and causes of death were computed according to glucose/diabetes categories.

Diabetes and nondiabetic categories of glucose regulation were defined by the 2003 ADA criteria (11), i.e., a person was classified as diabetic based on fasting plasma glucose (FPG) ≥7.0 mmol/l, 2h plasma glucose (2hPG) ≥11.1 mmol/l, or a previous clinical diagnosis. In the absence of diabetes, they were classified as having normal glucose regulation (NGR), isolated IFG, isolated IGT, or combined IFG and IGT (IFG+IGT), using glucose limits specified in Table 1. In addition, diabetes was classified as <5 years duration, 5 to 10 years duration, 10 to 15 years duration, and ≥ 15 years duration.
Nondiabetic subjects were also classified by only FPG criteria as NF G (FPG < 5.6 mmol/l) or IFG (5.6 ≤ FPG < 7.0 mmol/l) with or without IGT and similarly, by only 2hPG criteria, NGT (2hPG < 7.8 mmol/l) or IGT (7.8 ≤ 2hPG < 11.1 mmol/l) with or without IFG, to make groups comparable with previous publications (1,3,6). To determine the association between low FPG and mortality, subjects with normal FPG were also categorized into two groups according to their FPG [FPG < 4.5 mmol/l (n=59), 4.5 ≤ FPG < 5.6 mmol/l (n=928)].

A standard 12-lead ECG was obtained at each examination, after the carbohydrate load, with the subject resting in the supine position. ECGs performed before 1992 were coded by a cardiologist who had no knowledge of the clinical data. After 1992, ECGs were coded at the EPICARE Center at Wake Forest University by readers trained in standard ECG measurement techniques who were also blinded to the clinical data and previous tracings. All ECGs were graded using Minnesota Code (MC) classifications. Major ischemic ECG changes such as MC I1-2 (major Q wave abnormalities), MC I3 with IV1-2 or V1-2 (minor Q, QS waves with ST or T abnormalities), IV1,2 (significant ST segment depression), V1,2 (deep or moderate T wave inversion) or VII1,2,4,8 (left bundle branch block, right bundle branch block, intraventricular block) were considered to reflect IHD, as described previously (12).

**Statistical Analysis.** Generalized estimating equations were used to calculate the odds ratio for prevalent IHD according to glucose/diabetes categories using all examinations after controlling for age and sex and accounting for the dependence among multiple examinations in the same individual. Confidence intervals were computed from standard errors which were calculated with the empirical (or robust) estimator (13).

Death rates were calculated as the number of subjects who died per 1,000 person-years of follow-up. For the mortality analysis, the period at risk extended from the first research examination after the age of 35 years to death or the end of 2003, whichever came earlier. Mortality was analyzed according to glucose/diabetes categories at baseline and as time-dependent variables. Although the former analysis ignored changes in glucose tolerance status during follow-up, including development of diabetes, it was intended to replicate most reports in the literature. In the time-dependent analysis, person-time accumulated in the corresponding glucose tolerance categories, thereby accounting for changes over time. Deaths were counted in the most recent glucose category rather than by status at baseline. Nondiabetic glucose categories could change at each subsequent examination, whereas categories of duration of diabetes changed only according to years since diagnosis of diabetes. Age- and sex-adjusted death rates and death rate ratios (DRRs) relative to NGR were calculated with direct standardization to the 1980 Pima Indian population. Confidence intervals were computed from the logarithms of the death rates and rate ratios (14). Tests for general association were computed by the Mantel-Haenszel test (15) and for linear association by the Mantel extension test (16) modified for person-time denominators (17).

**RESULTS**

Baseline characteristics of the study population are presented in Table 1. Controlled for age and sex, prevalent IHD was significantly higher relative to NGR in subjects with ≥15 years duration of diabetes only (Odds ratio=1.9, 95% CI=1.4-2.5) (Figure 1A).

During a median follow-up of 10.4 years (range=0.04-29.0 years), 780 deaths from natural causes occurred among the 2,993 subjects (1,268 men, 1,725 women), 156 of these deaths being attributed to IHD. The age- and sex-adjusted death rates from natural
Plasma glucose and mortality

causes were similar within the nondiabetic groups (NGR, isolated IFG, isolated IGT and IFG+IGT) in both the baseline and time-dependent models. When only baseline glucose/diabetes categories were analyzed, death rates were higher in people with diabetes and were associated with diabetes duration. However, in the time-dependent analysis, natural mortality did not increase significantly until the duration of diabetes reached 15 years (Table 2, Figure 1B).

Diabetes at baseline was associated with higher IHD mortality, which increased with baseline duration of diabetes. In the time-dependent model, IHD death rates were lower than those in the corresponding baseline categories, but the associations did not change (Table 2, Figure 1C). IFG with or without IGT was not associated with higher natural or IHD mortality compared to NFG in both baseline and time-dependent models. The same was found for IGT with or without IFG (Figure 2). The major finding that only longer duration of diabetes was associated with increased natural cause and IHD mortality was consistent in both men and women. The associations of diabetes and duration of diabetes with mortality were not changed after adjusting for BMI, total cholesterol, mean arterial pressure, and smoking.

Subjects in the lowest FPG group (FPG <4.5 mmol/l) had a slightly higher death rate than those in the normal FPG group (4.5≤ FPG <5.6 mmol/l) after adjustment for age and sex (DRR compared to normal FPG group=1.5, 95% CI=0.6-3.4). The small number of people in this group, however, leads to a wide confidence interval, making this finding inconclusive.

CONCLUSIONS

Previously, we reported that natural and CVD mortality were not higher in Pima Indians with IFG or IGT than in those with NGR (18). That study, however, had a relatively short follow-up time so that there were only 285 deaths among 1,370 study subjects. Therefore, death rates within the sub-categories of impaired glucose regulation could not be examined. This study extends previous observations in Pima Indians on the relationship between glucose concentration and mortality (18), to encompass a 29-year period.

Although diabetes has a major impact on IHD and mortality in Pima Indians, the prevalence of IHD was not higher in the categories of IFG and/or IGT, but only in those with over 15 years of diabetes. The associations of incident IHD with glucose/diabetes categories (data not shown) were similar to those of prevalent IHD, but the confidence intervals for prevalent IHD were narrower because multiple examinations could be included in the generalized estimating equations and increased the power of the prevalent IHD analysis. Furthermore, mortality from natural causes or from IHD was not higher in individuals with IFG and/or IGT than in those with NGR in this study. IHD and diabetic nephropathy share many risk factors and are the leading causes of death among diabetic Pima Indians. Moreover, persons who develop diabetic nephropathy die primarily of IHD (10). When death rates from diabetic nephropathy and IHD were computed as a composite endpoint, the associations with glucose/diabetes categories were similar as with IHD alone (data not shown). These findings are consistent with the Atherosclerosis Risk in Communities (ARIC) study (5), which included 6,888 white and black nondiabetic persons and showed that neither IFG nor IGT increased the risk for all-cause mortality or incident IHD. Similarly in the Hoorn study (6), increased risk associated with IFG or IGT was mostly, but not completely, attributable to cardiovascular risk factors in a population cohort of 2,363 subjects, without known diabetes. By contrast, in the DECODE (Diabetes Epidemiology: Collaborative
analysis Of Diagnostic criteria in Europe) study (1), which included 22,514 subjects from a number of European population-based studies, IGT but not IFG, was associated with increased mortality from all causes and from CVD, defined as IHD and stroke. This relationship was present even after accounting for conventional cardiovascular risk factors, whereas in our study, IFG and/or IGT did not increase the risk of death after controlling for age, sex, body mass index, total cholesterol, mean blood pressure, and smoking. We computed mortality in persons with IFG (with or without IGT) relative to NFG and in persons with IGT (with or without IFG) relative to NGT, categories that facilitated comparison with the Hoorn (6) and DECODE (1) studies. Although neither of these categories were associated with statistically significantly higher natural-cause or IHD mortality rates, the point estimates of effects were consistent with these previous studies. For example, in the current study, the DRR for IHD mortality in IGT was 1.5 (95% CI=0.6 to 3.5) compared with 1.3 (95% CI=1.0 to 1.6) in DECODE. By contrast, neither the Hoorn study nor the present study provided evidence for an effect of IFG on mortality, although the two studies used different definitions of IFG and classification of causes of death.

Compared with analyses using only baseline glucose categories to predict deaths, in the time-dependent models the effect of diabetes on mortality was attenuated. Indeed, in the time-dependent models, only persons with diabetes of longer duration had significantly higher mortality than those with NGR. Considering this difference, reports of higher death rates from natural causes or IHD in persons with IFG and/or IGT that are based solely on baseline glucose overestimate the effect of glucose in the nondiabetic range, since some subjects in these groups could have developed diabetes after the baseline examination and yet remained in the nondiabetic group in the analysis. Two studies examining the impact of development of diabetes on the increased risk for IHD morbidity and mortality in persons with IFG or IGT reported conflicting results. In the Finnish study (9), IGT remained an important risk factor even when controlled for the subsequent development of diabetes. Persons with IGT who did not develop diabetes after 10 years of follow-up had a 49% higher risk of incident IHD and a 65% higher risk of all-cause mortality compared with NGT. Conversely, in the Hoorn study, the excess mortality associated with IFG was present only in those who developed diabetes during follow-up (8). The Hoorn study included follow-up glucose measurements, whereas the Finnish study relied on the registered drug data or ICD codes provided by the national Hospital Discharge Register for identifying new cases of diabetes. Therefore, the Finnish study was more likely to include undiagnosed diabetic subjects in the IGT group. Thus, failure to account for progression to diabetes may result in incorrect conclusions about the pathogenetic importance of IFG and IGT in relation to IHD and mortality.

Several studies have indicated a lower glycemic threshold for macrovascular than for microvascular disease (3,19). This was not evident in the Pima Indians (18,20), i.e., the risk of these complications was increased in subjects with diabetes but not in those with lesser degrees of glucose abnormality. This finding may be due to the much lower rate of IHD in nondiabetic Pimas than in many other populations (20). Low serum concentrations of total and low density lipoprotein cholesterol, and a low rate of heavy smoking (20) may be responsible, in part, for these differences. Indeed, only 22 nondiabetic IHD deaths occurred in over 17,000 person-years of follow-up during the 29-year study period. With low rates of traditional risk factors for macrovascular disease, diabetes, especially when accompanied by renal complications,
Plasma glucose and mortality may represent the predominant risk factor for IHD. This impact of diabetic nephropathy enhances the association of diabetes duration with the incidence of IHD.

Lower glucose concentration has also been previously associated with CVD and natural mortality (3). Although not statistically significant, there was 50% higher natural-cause mortality in those with FPG less than 4.5 mmol/l than those with FPG 4.5-5.5 mmol/l. The pathogenesis of this association remains to be defined. Nevertheless, poor general health and ECG changes such as increased ectopic activity, flattening of the T waves, ST depression, ventricular tachycardia, and atrial fibrillation may play a role (21,22).

The small number of subjects who had or died of IHD in the nondiabetic and the short duration diabetes groups may limit the power of the analysis to reveal potential differences in IHD risk among IFG, IGT, and short-duration diabetes.

In conclusion, the present findings are consistent with the hypothesis that any association of IHD with impaired glucose regulation is due primarily to factors other than hyperglycemia per se. Although impaired glucose regulation is associated with abnormal insulin secretion and action and predicts diabetes, it is not an independent predictor of mortality except in those who subsequently develop diabetes.

ACKNOWLEDGMENTS

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REFERENCES

**TABLE 1.** Baseline characteristics of the study population according to glucose/diabetes categories.

<table>
<thead>
<tr>
<th></th>
<th>NGR</th>
<th>Isolated IFG</th>
<th>Isolated IGT</th>
<th>IFG+IGT</th>
<th>DM</th>
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<tbody>
<tr>
<td>FPG (mmol/l)</td>
<td>&lt; 5.6</td>
<td>5.6-6.9</td>
<td>&lt; 5.6</td>
<td>5.6-6.9</td>
<td>±</td>
</tr>
<tr>
<td>2hPG (mmol/l)</td>
<td>&lt; 7.8</td>
<td>&lt; 7.8</td>
<td>7.8-11.0</td>
<td>7.8-11.0</td>
<td>±</td>
</tr>
<tr>
<td>N</td>
<td>823</td>
<td>299</td>
<td>164</td>
<td>231</td>
<td>1476</td>
</tr>
<tr>
<td>Age (years)</td>
<td>42.1 ± 9.4</td>
<td>43.8 ± 10.2</td>
<td>43.0 ± 9.9</td>
<td>45.6 ± 11.6</td>
<td>47.0 ± 11.2</td>
</tr>
<tr>
<td>Sex (% male)</td>
<td>42.7</td>
<td>46.5</td>
<td>33.5</td>
<td>43.3</td>
<td>42.2</td>
</tr>
<tr>
<td>FPG (mmol/l)</td>
<td>5.0 ± 0.3</td>
<td>5.9 ± 0.3</td>
<td>5.1 ± 0.3</td>
<td>6.0 ± 0.4</td>
<td>11.2 ± 4.4</td>
</tr>
<tr>
<td>2hPG (mmol/l)</td>
<td>5.7 ± 1.2</td>
<td>6.3 ± 1.1</td>
<td>8.7 ± 0.8</td>
<td>9.2 ± 0.9</td>
<td>18.6 ± 6.6</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>32.7 ± 7.4</td>
<td>35.5 ± 7.8</td>
<td>33.6 ± 6.6</td>
<td>36.6 ± 7.7</td>
<td>33.3 ± 7.8</td>
</tr>
<tr>
<td>Waist (cm)</td>
<td>107.6 ± 17.6</td>
<td>114.7 ± 19.2</td>
<td>107.5 ± 14.6</td>
<td>118.0 ± 16.9</td>
<td>113.2 ± 18.9</td>
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<td>SBP (mmHg)</td>
<td>118.7 ± 16.7</td>
<td>123.0 ± 17.7</td>
<td>121.8 ± 18.8</td>
<td>125.6 ± 18.9</td>
<td>130.4 ± 22.4</td>
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<td>DBP (mmHg)</td>
<td>73.8 ± 11.7</td>
<td>76.2 ± 11.7</td>
<td>75.1 ± 11.7</td>
<td>77.4 ± 13.4</td>
<td>79.6 ± 12.3</td>
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<tr>
<td>TC (mmol/l)</td>
<td>4.6 ± 0.9</td>
<td>4.6 ± 0.8</td>
<td>4.6 ± 0.9</td>
<td>4.6 ± 0.9</td>
<td>4.8 ± 1.3</td>
</tr>
<tr>
<td>TG (mmol/l)*</td>
<td>1.2 (0.8,1.7)</td>
<td>1.3 (1.1,2.0)</td>
<td>1.4 (1.0,2.2)</td>
<td>1.5 (1.0,2.1)</td>
<td>1.5 (1.1,2.5)</td>
</tr>
<tr>
<td>HDL-C (mmol/l)</td>
<td>1.3 ± 0.4</td>
<td>1.1 ± 0.4</td>
<td>1.2 ± 0.4</td>
<td>1.1 ± 0.3</td>
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</tr>
<tr>
<td>Smoking (%)</td>
<td>34.8</td>
<td>24.2</td>
<td>27.4</td>
<td>23.4</td>
<td>23.5</td>
</tr>
</tbody>
</table>

Data are mean ± standard deviation (SD) or *median (interquartile range).  
FPG: fasting plasma glucose, 2hPG: 2hr glucose after 75g glucose loading, BMI: body mass index defined as weight divided by the square of height (kg/m²), SBP: systolic blood pressure, DBP: diastolic blood pressure, TC: total cholesterol, TG: triglyceride, HDL-C: HDL cholesterol.  
†Smoking: percent of current smokers  
DM: diabetes.  
* Diagnosis based on FPG ≥ 7.0 mmol/l, 2hPG ≥ 11.1 mmol/l, or previous clinical diagnosis.
<table>
<thead>
<tr>
<th>Group</th>
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<th>IHD mortality</th>
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<tr>
<td></td>
<td>Baseline</td>
<td>Time-dependent</td>
</tr>
<tr>
<td></td>
<td>Pyrs* (N)</td>
<td>Deaths (N)</td>
</tr>
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<td>NGR</td>
<td>8726.1</td>
<td>104</td>
</tr>
<tr>
<td>Isolated IFG</td>
<td>3976.7</td>
<td>44</td>
</tr>
<tr>
<td>Isolated IGT</td>
<td>1687.1</td>
<td>24</td>
</tr>
<tr>
<td>IFG+IGT</td>
<td>2962.2</td>
<td>51</td>
</tr>
<tr>
<td>DM &lt;5</td>
<td>8639.2</td>
<td>165</td>
</tr>
<tr>
<td>5≤ DM &lt;10</td>
<td>3930.9</td>
<td>115</td>
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<tr>
<td>10≤ DM &lt;15</td>
<td>3555.9</td>
<td>163</td>
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<tr>
<td>DM ≥15</td>
<td>1700.4</td>
<td>114</td>
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<td>Total</td>
<td>35178.6</td>
<td>780</td>
</tr>
</tbody>
</table>

* Person-years (pyrs) are shown only for natural mortality as they are the same for IHD mortality.
† Age-sex adjusted and reported as deaths per 1000 pyrs at risk. ‡ 95% CI for each death rate.
FIGURE 1

A. 

B. 

C.
FIGURE 2

A. 

![Graph A](image)

B. 

![Graph B](image)