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Predictors of hyperglycaemic individuals who do not follow up with physicians after screening in Japan: A cohort study.

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Predictors of Hyperglycaemic Individuals Who do not Follow up with Physicians after Screening in Japan: a Cohort Study

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Aims: Although people screened as being hyperglycaemic often fail to follow up with physicians for clinical assessment, epidemiologic findings on the frequency and predictors of not following up (hereafter, “no follow-up”) are lacking. The purpose of this study was to examine the no follow-up rate with physicians after screening for diabetes and its predictors.

Methods: We assessed cases of no follow-ups with physicians within six months after screening based on medical claims data from employee-based social health insurance programs in Japan, for people aged 20 to 68 years from 2005 to 2010.

Results: Among 3,878 screened participants with hyperglycemia, 2,527 (65%) did not follow up with their physicians within six month after screening. Multiple logistic regression analysis revealed that younger age and lower blood glucose level predicted no follow-ups among both men and women, while lower body mass index and negative proteinuria also predicted no follow-ups among men. Treatment for dyslipidaemia facilitated follow-ups among both genders, and treatment for hypertension or depression facilitated follow-ups among men.

Conclusions: Approximately two thirds of individuals screened as having hyperglycemia did not follow up with their physicians within six months after screening. Predictors of no follow-ups
were younger age and milder hyperglycemia. Treatment of comorbidities tended to facilitate follow-ups.

**Keywords**

diabetes; screening; no follow-up; Japan
Introduction

Diabetes mellitus is associated not only with the occurrence of vascular complications, but also with impaired quality of life and increased health care costs\textsuperscript{1-3}. In 2013, 382 million people had diabetes in the world; this number was expected to rise to 592 million by 2035\textsuperscript{4}. The total number of excess deaths attributable to diabetes in 2010 was estimated to be 3.96 million in the age group of 20-79 years\textsuperscript{5}. Approximately 17\% of the Japanese population is estimated to have diabetes, and this number increases every year\textsuperscript{6}.

The effectiveness of screening for diabetes remains controversial\textsuperscript{7-14}. In Japan, a unique system of general health screening that includes glucose testing was established by the Health and Medical Service Act for the Aged in 1983. Screenings have been conducted primarily at worksites or local community facilities, and patients who screen positively are advised to follow up with a physician. Thus, following up with a physician is a requisite for this screening system to work. A Japanese national survey based on self-report revealed that 39\% of people with diabetes had not been treated\textsuperscript{6}, indicating that many individuals who are potentially diabetic neither visited nor followed up with physicians.

A previous cross-sectional study based on self-report suggested that younger people and those with lower risk were more likely to drop out from screenings\textsuperscript{15}. Another cross-sectional study reported that individuals with type 2 diabetes who suffer from depression poorly adhere to
self-care\textsuperscript{16}. Yet, no study has reported longitudinal findings using objective data to address this issue. To this end, we conducted a cohort study to examine the characteristics of people who did not follow up with physicians for clinical assessment after they were screened as being hyperglycaemic, with a focus on age, severity of hyperglycemia and comorbidities.

**Materials and Methods**

**Setting**

In Japan, all people have been insured under universal health coverage since 1961, which primarily consists of employee-based and community-based social health insurance programs\textsuperscript{17}. The Industrial Safety and Health Law enacted in 1972 requires that public screening services be provided at worksites or local community facilities, rather than taking place at a family or attending physician’s office as in the United States and other countries.

In order to obtain a confirmed diagnosis or to initiate treatment after screening, screened individuals with a positive test result must follow up with physicians (Figure 1). In Japan, standard screening for adults was implemented by the Law of Health and Medical Services for the Elderly from 1983 to 2008, although it is now under the control of the Act on Assurance of Medical Care for Elderly People, which was enacted in 2008. Our data include both periods. Both laws instruct those whose examined values exceed cut-off values to visit physicians, and most health insurance programs advise them to do so simply by mail. Disease management, including
education, monitoring or feedback, for screened people is not proactively carried out by the
government. Rather, disease management differs by the particular health insurance system.
Screened people generally receive instructions to consult with a physician within about two
months of the screening, although a medical institution or timing of the visit is not specified. That
is, in Japan, although health insurance plans provide screening to insured people, the plans fall
short of designating a hospital. This reflects the disconnect between health insurance programs
and hospitals, although regular health check-ups are the norm and mass screening is provided for
everyone at school and work or in the community. In 2007, the Ministry of Health, Labour and Welfare proposed a standard program for screening
and health guidance, and screenings are currently conducted according to this program.
Screenings are held by each health insurance program, although the frequency differs by program.
Most screenings are held during the day, and blood and/or urine samples are collected early in the
morning. Screened participants are instructed to fast from 9 p.m. the night before to the end of
screening, although the degree of compliance with these instructions is unclear. In general, there
are two approaches to the screening. Either people visit a specifically-designated facility to
receive the check-up, or screeners visit the worksite. For worksite-based screenings in our study,
vehicles carrying medical testing equipment visited each worksite and carried out the screenings.
Depending on the insurance program, some participants visited a facility designated by the
program to undergo screening. For community-based screenings, non-employed people typically
visit local facilities.

Screening services follow several general steps that represent the analytical framework for diabetes screening\(^8\). We reworked this framework to adapt it for diabetes screening in Japan (Figure 1). As mentioned above, Japanese screenings are held mainly at community facilities or worksites (Step 1 in Figure 1), followed by visiting physicians (Step 2 in Figure 1). For the screening program to work effectively, people need to adhere to all steps.

**Study design and data source**

We carried out a retrospective cohort study using administrative data from both health insurance claims and screenings from Japan Medical Data Center Co., Ltd. (JMDC, Tokyo, Japan). JMDC obtained health insurance claims and screening data from several employee-based social health insurance plans and constructed the JMDC medical database (JMDC-MDB). This database includes 530,000 insured people and their dependents, mainly company employees and their family members, from January 2004 to December 2010. Health insurance claims were anonymously linked with screening data in JMDC-MDB\(^20\). The database provided patient demographics, screening results, test orders, treatments, prescribed drugs and hospital diagnoses. The locality of the health insurance plans or the identities of the insured were not available to the researchers. Under these circumstances, all employees and family members were automatically eligible for the screening and were encouraged to attend it.
The Institutional Review Board of the Kyoto University Graduate School and Faculty of Medicine Ethics Committee approved the study protocols (E1017).

**Study participants**

Study participants were screened individuals with untreated hyperglycemia. The study cohort included 220,409 employees and their dependent family members, from 2005 to 2010. Of the 220,409 participants, we excluded 118,765 who did not attend a screening (screening attendance rate: 46%) and 9,749 who did not undergo glycated haemoglobin (HbA₁c) or fasting glucose tests. Of the remaining 91,895 participants, 5,834 participants (6.3%) were considered hyperglycaemic (HbA₁c ≥ 6.5% [48 mmol/mol], or fasting glucose ≥ 126 mg/dl). Of these, 1,927 were being treated with antihyperglycaemic agents. When participants were identified as hyperglycaemic twice or more, we defined “hyperglycaemic at screening” as the last time of screening in the study period. We excluded people who were born before 1940 (n=19) because of missing birth dates, and people who were under 20 years at the time of screening (n=10). Data from the remaining 3,878 participants (3,420 men and 458 women) were analysed (Figure 2).

**Outcome measurement**

“Follow-up” refers to the initial visit to a physician within six months after the screening, accompanied by an examination of HbA₁c and/or glucose. When a participant who screened
positive did not follow up with a physician, this was designated as “no follow-up”, which was regarded as the primary outcome measurement. We determined the indicated visit as being for treatment of diabetes or not according to whether further glycaemia testing, an examination of HbA$_{1c}$ and/or glucose was done at the visit. Those who visited a physician for the other diseases (e.g., common cold) were excluded. Whether patients were on track for treatment or not is beyond the scope of the present study, and thus “follow-up” also included those who made even a single visit to a physician. For the most part, screenings are performed once (although in some cases, twice a year). Accordingly, the primary outcome measure for this study was “no follow-up with a physician within six month after screening.” For sensitivity analysis, we also examined no follow-up with a physician within three and 12 months of screening.

**Baseline variables**

Demographic characteristics and screening data for participants were obtained directly from JMDC-MDB. Body mass index (BMI [kg/m$^2$]) was calculated from weight and height measured during screening as a measure for obesity. HbA$_{1c}$ values determined by the Japan Diabetes Society (JDS) method was 0.4% lower than NGSP values$^{21}$, so we converted them to NGSP values.

We identified comorbidities using information from health insurance claims on hypertension treatment, dyslipidaemia, proteinuria and depression. Hypertension and dyslipidaemia are chronic
metabolic diseases that, like hyperglycemia, require drug therapy and lifestyle modification. We hypothesized that participants being treated for these diseases may be more likely to follow up with physicians if they were hyperglycaemic at screening. Proteinuria was chosen as a marker of possible chronic kidney disease. A previous study reported that depression is associated with non-adherence to diabetes self-care among patients with type 2 diabetes16, implying that depression precludes engaging in desirable health behaviours. Therefore, treatment for depression was included as a potential predictor of non-adherence to follow-ups. We defined “hypertension” as prescription of hypertension medication within three months prior to screening: C02, 03, 04, 07, 08, or 09 in the Anatomical Therapeutic Chemical Classification System22. Similarly, we defined “dyslipidaemia” as prescription of medication for dyslipidaemia (C10), and “depression” as prescription of medication for depression (N06A or N06BA). Participants who were diagnosed with hypertension, dyslipidaemia and/or depression were excluded if their comorbidities were left untreated. “Glycosuria” was classified as either negative (urine dipstick, negative) or positive (urine dipstick, trace or above). Proteinuria was also classified as negative (urine dipstick, negative or trace) or positive (urine dipstick, 1+ or above). For patients who already had hypertension or dyslipidaemia, we examined whether they visited the same clinics for hyperglycemia as for these other diseases. Men and women were analysed separately.

Smoking (yes/no) and frequency of alcohol consumption (everyday/sometimes/rarely) were self-reported and missing in a considerable proportion of the participants (23% and 36%,
respectively); participants without these data were excluded from the analysis. Given the lack of
data regarding education level, marital status, race or ethnicity, type of employee-based social
health insurance plan and insured/dependent individual were used as surrogates of these
socio-economic variables.

**Statistical analysis**

We identified mean or median values, and frequencies of explanatory variables that were potential
predictors or confounders, stratified by gender. We excluded variables if there were more than
20% missing data. For the remaining variables, we used mean values for missing data of
continuous variables, and “no” for dichotomous variables.

We calculated the frequency of no follow-ups after screening, and used logistic regression models
to estimate odds ratios (ORs) and 95% confidence intervals (CIs) to determine predictors. As
explanatory variables, we used baseline variables for which there were less than 20% missing
data (age, gender, HbA1c, fasting glucose, BMI, glycosuria, proteinuria, haemoglobin,
hypertension, dyslipidaemia, depression, type of health insurance plans and insured/dependent
status at the health insurance plans). Multiple logistic regression analyses were performed
following analyses using age-adjusted models. We also analysed data for two additional
follow-up periods: within three months and within 12 months. The statistical models were
examined using the area under the receiver operating characteristic (ROC) curve (C statistic).
Tests of statistical significance were two-tailed, with an $\alpha$-level of 0.05. All analyses were performed with STATA, version 11.1 (Stata Corp., College Station, TX, USA).

Results

Screening attendance rate and baseline characteristics of participants with hyperglycaemia

Among 101,644 of 220,409 cohort members who participated in a screening (attendance rate: 46%), 91,895 underwent glucose testing, of whom 5,834 (6.3%) were hyperglycaemic and 3,907 (4.3%) were hyperglycaemic without treatment. There were 3,878 study participants remaining after excluding those under 20 years of age or born in 1940 or before (Figure 2). HbA1c and fasting glucose levels were used to identify participants with suspected diabetes (i.e., if they had values higher than the cut-off). Among the 3,878 study participants, HbA1c and fasting glucose were examined in 3,784 (97.6%) and 3,363 (86.7%) participants, respectively. For 94 (2.4%) participants, only data for fasting glucose were available.

We used age, gender, HbA1c, fasting glucose, BMI, glycosuria, proteinuria, haemoglobin, hypertension, dyslipidaemia, depression, type of health insurance plan, and insured/dependent status in the health insurance plan as variables; these data were complete or missing less than 20% (Table 1).

Table 1 shows the baseline characteristics of 3,878 participants screened as hyperglycaemic. More
men than women had higher haemoglobin levels (15.2 g/dl vs. 13.1 g/dl), were more likely to smoke (40.4% vs. 7.6%), consumed alcohol more frequently (14.4% vs. 2.0% for everyday drinking), had depression more often (2.3% vs. 6.6% for depression), had hypertension more often (16.9% vs. 13.8%) and were insured (99.97% vs. 49.1%).

**Frequency of no follow-up**

Of the 3,878 hyperglycaemic participants, 2,527 (65.2%) did not follow up with physicians within six months (Figure 2, Table 1). Men were less likely to follow up with physicians than women during the three, six, and 12-month follow-up periods (74.9% vs. 67.0%, 65.9% vs. 59.4% and 58.2% vs. 53.7% for no follow-ups, respectively). When stratified by HbA1c levels, >40% of participants with an HbA1c of 8.0% (64 mmol/mol) or over, and 35% of participants with HbA1c of 10.0% (86 mmol/mol) or over, did not follow up with physicians within six months (Table 2).

**Frequency of follow-up with physicians treating pre-existing conditions**

For participants being treated for hypertension (n=640) or dyslipidaemia (n=410), 367 (57%) of those with hypertension and 270 (66%) of those with dyslipidaemia followed up with physicians after screening for diabetes. Most of these participants followed up with the physicians who were already treating them (80%: hypertension, 90%: dyslipidaemia).

**Predictors of no follow-up**
Table 3 shows age-adjusted analyses for predicting no follow-ups. Lower BMI, lower glucose level and negative proteinuria predicted no follow-ups in men, while treatment for comorbidities (hypertension, dyslipidaemia and depression) facilitated follow-ups. Lower glucose level, negative proteinuria and insured status predicted no follow-ups in women, while treatment for comorbidities (hypertension and dyslipidaemia) facilitated follow-ups.

Table 4 shows the adjusted ORs and 95% CIs for no follow-ups by each variable. This multiple logistic regression analysis revealed that younger age, lower BMI, lower glucose level and negative proteinuria significantly predicted no follow-ups in men, while treatment for comorbidities (hypertension, dyslipidaemia and depression) facilitated follow-ups. For women, younger age and lower glucose level predicted no follow-ups, while treatment for comorbidities (dyslipidaemia) facilitated follow-ups. Areas under the ROC curve for the multiple logistic regression models were 0.72, 0.73 and 0.74 for men and 0.74, 0.76 and 0.77 for women at three, six and 12 months, respectively.

**Discussion**

We showed that 2,527 of 3,878 individuals (65.2%; 65.9% of men and 59.4% of women) who were hyperglycaemic without treatment at the time of screening did not follow up with physicians within six months after the screening. Younger age, lower blood glucose level, lower BMI and negative proteinuria predicted no follow-ups, while treatment for comorbidities (hypertension,
dyslipidaemia or depression) facilitated follow-ups. Our study is the first to reveal the frequency
and predictors of no follow-ups in the Japanese diabetes screening system using a large database
that links health claim data with mass screening results.

Direct evidence of the effectiveness of diabetes screening and its clinical implementation is
lacking (Key Question 1 in Figure 1), although there have been several previous studies
addressing this7-14, 23, 24. The U.S. Preventive Services Task Force summarizes screening for type 2
diabetes mellitus in adults as a grade I statement, meaning that current evidence is insufficient to
assess the service7. A modelling study simulated that screening for type 2 diabetes would be
cost-effective when initiated between the ages of 30 and 45 years, and repeated every three to five
years25. Another study suggested that screening for type 2 diabetes and early intensive
multifactorial treatment were feasible in general practice13. Adverse outcomes from diabetes
screenings were not apparent in previous studies26-29, possibly because the studies did not
sufficiently analyse dropout cases. Focus should also be placed on these cases in order to increase
the effectiveness of diabetes screenings.

The screening attendance rate in the present study was 46% (Figure 2), and was 78% among
insured participants and 29% among dependents, which was similar to results of a Japanese
national survey that showed 62% of citizens attended a general health screening (employed: 69%,
unemployed: 49%)30. The screenings involved measurements of HbA1c and fasting or casual
blood glucose to identify individuals who are possibly diabetic. Participants were instructed to fast beginning at the night before the screening, and for those who failed to maintain the fasting requirement, casual blood glucose was measured and the time since the last meal was recorded. In the present study, only HbA1c and fasting glucose were used to identify participants because these measurements are more stable than that of casual blood glucose. Participants were considered to be potentially diabetic if they had values above the cut-off for HbA1c and/or fasting glucose.

The rate of no follow-ups after screening was much higher in the present study (step 2 in Figure 1) than in national data showing that 39% of people with diabetes were not being treated. This discrepancy can be attributed to the difference in research design. The national survey was based on self-report and could be affected by information bias, whereas our study used objective administrative data from health insurance claims and screenings. Moreover, the sampling process differed in that participants of the national survey were those who answered ‘I have never been treated’ or ‘I was treated in the past, but not currently’ to the question ‘have you ever been referred to as a diabetic at a screening or at a medical institution?’ In contrast, we asked the question “how many individuals did not follow up after screening positive for potential diabetes.” Another previous study reported a 50% response rate to a pre-screening questionnaire for diabetes, and 77% of respondents at high risk for diabetes visited clinics. Although the follow-up rate after screening was higher in the previous study than in ours, it was conducted in an interventional
study setting and did not represent actual behaviour after screenings. Also, while it has been suggested that screening programs should follow-up on positive tests, specific goals have not been set. It may be worth examining a concrete follow-up process to screening in the future. Even with moderate to severe hyperglycaemia, considerable proportions of participants did not follow up with physicians. Surprisingly, one third of participants with an HbA1c value of 10% (86 mmol/mol) or over failed to follow up with physicians. This highlights the urgent need for screeners to develop an effective reminder/recall system for such individuals. In addition, potential barriers to following up, e.g., pressure against leave work to visit doctors, need to be further investigated.

Our study identified a number of predictors for no follow-ups. The first was younger age. Younger people may lack awareness regarding the risks of hyperglycaemia and are less health-conscious than older people. Most of the younger people were working, and would not go to clinics if they were experiencing no symptoms or only mild symptoms because of their environment. This finding was consistent with that of the previous study.

Second, lower BMI predicted no follow-ups in men. Some studies reported that obese people tend to visit general practice physicians more often than non-obese people. Although our study was conducted in a screening setting, our results support the generalization that obese men are more likely to follow up with physicians than non-obese men. There are two possible reasons for
this: (1) obesity increases the risk of diseases that require treatment, and (2) obesity increases a
patient’s sense of risk and leads to follow-up with physicians. On the other hand, obese women
tended not to follow up with physicians. This implies that obese women might be less health
conscious than obese men, and the tendency of obese individuals to engage in healthy behaviours,
including follow-ups with physicians, might be influenced by gender. Examination of this
hypothesis is beyond the scope of our study and further study will be needed to address this issue.

Third, lower blood glucose level predicted no follow-ups. This is related to a lack of risk
awareness, similar to that found for age differences. Also, the above-mentioned study suggested
that dropouts at the screening step were more likely to have lower risk scores on the pre-screening
questionnaire.¹⁵

Fourth, having or being treated for comorbidities including hypertension, dyslipidaemia,
proteinuria and depression tended to facilitate follow-ups. Furthermore, over 80% of people who
had been treated for hypertension or dyslipidaemia visited the same physicians after screening as
hyperglycaemic. Treatment for comorbidities, particularly hypertension and dyslipidaemia,
increased the likelihood that people who were hyperglycaemic at screening would follow up with
physicians and receive clinical care. This suggests that many patients tend to attend the same
clinics as part of the free access system in Japan. A previous study reported that patients with type
2 diabetes who are depressed poorly adhere to self-care; however, the definition of depression in
this report was based only on self-reported data, which is different from the definition adopted in our study. Our study participants who were classified as depressed were undergoing treatment and might have been more inclined to comply with instructions after screening. A lack of risk awareness, as discussed above, may be associated with proteinuria as well.

Sensitivity analyses was also conducted to examine differences in follow-up periods. When setting three or 12 months as the follow-up period, all explanatory variables showed similar trends as those for the six-month follow-up. Moreover, multiple logistic regression models and ROC curves were used to assess the ability of these models to predict outcomes. AUCs showed acceptable discrimination, with values ranging from 0.72 to 0.77.

There are some limitations to this study. First, smoking and alcohol consumption were not included in the analysis due to missing data. Consequently, this study may not have considered potentially important information on behavioural factors. Second, as information on socioeconomic status was not available, we used the type of health insurance plan and insured/dependent status as surrogates for socioeconomic status. In Japan, the particular company one works for dictates the health insurance plan one has. While the type of health insurance plan and insured/dependent status have limitations as surrogates of socioeconomic status, the focus of the present study was the middle-aged working class of company employees, rather than a representative population comprising socially diverse sub-populations from around Japan. Given
this design, our study population could be considered more socially homogeneous than a national
representative population. The existence of unadjusted variables (i.e., residual confounding) may
have limited the fitness of the obtained models. Third, the analysed data were only from three
employee-based health insurance plans. As a result, the participants were younger than the
general Japanese population and the number of women was limited. Therefore, extrapolation of
the present findings to other Japanese populations, especially elderly people requires careful
consideration. On the other hand, recent clinical guidelines have reconsidered the strict
glycaemic goals for treating older adults with diabetes, and aggressive screening for older adults
to detect possible diabetes remains controversial. Against this backdrop, our focus on a
younger population (working-age adults) would contribute to maintaining work productivity in
society.

In conclusion, we found that two-thirds of participants who were hyperglycaemic did not follow
up with physicians after screening. Younger age, lower BMI, lower blood glucose level and
negative proteinuria predicted no follow-ups, and treatment for comorbidities including
hypertension, dyslipidaemia and depression facilitated follow-ups. To improve the screening
system and health care cost efficiency, a more effective follow-up process that includes
recommendations to visit physicians must be offered to those at high risk of not following up.
Further studies will be needed to accurately evaluate diabetes screening, including no attendance
or no follow-ups, which will contribute to a more effective health care system.
Conflict of interest

The authors declare that they have no conflict of interest.

Acknowledgments

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References


Figure legends

Fig. 1 - Japanese diabetes screening process.

Note: Screenings are primarily taken place at work sites or local community facilities, and screened patients are advised to follow up with physicians after screening.

KQ, key question(s);

KQ1: Is there direct evidence that systematic screening for diabetes among asymptomatic adults improves health outcomes?

KQ2: Does follow-up with physicians early after screening for diabetes provide an incremental benefit in health outcomes?

KQ3: Does initiating early treatment for type 2 diabetes as a result of screening provide an incremental benefit in health outcomes?

Does initiating early treatment of impaired fasting glucose or impaired glucose tolerance as a result of screening provide an incremental benefit in final health outcomes?

Step 1: Attending diabetes screening

Step 2: Follow-up with physicians after screening

Step 3: Initiation of appropriate treatment
Fig. 2 - Flowchart of the study participants.

HbA1c, glycated haemoglobin
Fig. 1
Fig. 2

220,409 employees and their family

191,644 attended screening
(Attendance rate 46%)

91,885 were tested fasting glucose or HbA1c

5334 were hyperglycemic at screening

3907 were hyperglycemic without treatment

3878 were hyperglycemic without treatment aged 20-65

118,765 did not attend screening
101,422 visited clinics
17345 did not visit clinic

9749 did not tested fasting glucose or HbA1c
not tested

56,051 were not hyperglycemic at screening

1927 were being treated for diabetes

18 were under 20 years of age at time of screening
19 were born 1940 or before

1351 followed up with physicians (35%)

2527 did not follow up with physicians (65%)
### Table 1 - Characteristics of 3878 participants with hyperglycemia, stratified by gender

<table>
<thead>
<tr>
<th></th>
<th><strong>Men</strong> (n=3420)</th>
<th><strong>Women</strong> (n=458)</th>
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<tr>
<td><strong>No follow-ups with physicians, n (%)</strong></td>
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<tr>
<td>0-3 months</td>
<td>2563 (74.9)</td>
<td>307 (67.0)</td>
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<tr>
<td>0-6 months</td>
<td>2255 (65.9)</td>
<td>272 (59.4)</td>
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<td>0-12 months</td>
<td>1989 (58.2)</td>
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<td><strong>Age groups, n (%)</strong></td>
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<td>20-24 years</td>
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<tr>
<td>65-68 years</td>
<td>26 (0.76)</td>
<td>9 (2.0)</td>
</tr>
<tr>
<td><strong>Mean body mass index (SD), kg/m(^2)</strong></td>
<td>25.1 (4.4)</td>
<td>25.0 (5.4)</td>
</tr>
<tr>
<td></td>
<td>Group 1 (Mean ± SD)</td>
<td>Group 2 (Mean ± SD)</td>
</tr>
<tr>
<td>--------------------------</td>
<td>---------------------</td>
<td>---------------------</td>
</tr>
<tr>
<td>Mean fasting glucose (SD), mg/dl</td>
<td>141.0 (34.5)</td>
<td>135.7 (38.0)</td>
</tr>
<tr>
<td>Mean HbA1c (SD), %</td>
<td>6.5 (1.3)</td>
<td>6.6 (1.4)</td>
</tr>
<tr>
<td>Glycosuria, n (%)</td>
<td>758 (22.2)</td>
<td>63 (13.8)</td>
</tr>
<tr>
<td>Proteinuria, n (%)</td>
<td>223 (6.5)</td>
<td>26 (5.7)</td>
</tr>
<tr>
<td>Hypertension, n (%)</td>
<td>577 (16.9)</td>
<td>63 (13.8)</td>
</tr>
<tr>
<td>Dyslipidemia, n (%)</td>
<td>357 (10.4)</td>
<td>53 (11.6)</td>
</tr>
<tr>
<td>Depression, n (%)</td>
<td>8 (0.2)</td>
<td>3 (0.7)</td>
</tr>
<tr>
<td>Smoking, n (%)</td>
<td>1383 (40.4)</td>
<td>35 (7.6)</td>
</tr>
<tr>
<td>Alcohol consumption, n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Everyday</td>
<td>493 (14.4)</td>
<td>9 (2.0)</td>
</tr>
<tr>
<td>Sometimes</td>
<td>772 (22.6)</td>
<td>38 (8.3)</td>
</tr>
<tr>
<td>Rarely</td>
<td>897 (26.2)</td>
<td>262 (57.2)</td>
</tr>
<tr>
<td>Missing data</td>
<td>1258 (36.8)</td>
<td>149 (32.5)</td>
</tr>
<tr>
<td>Mean haemoglobin (SD), g/dl</td>
<td>15.2 (1.2)</td>
<td>13.3 (1.6)</td>
</tr>
<tr>
<td>Health insurance plans, n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>A</td>
<td>612 (17.9)</td>
<td>104 (22.7)</td>
</tr>
<tr>
<td>B</td>
<td>2702 (79.0)</td>
<td>343 (74.9)</td>
</tr>
<tr>
<td>C</td>
<td>106 (3.1)</td>
<td>11 (2.4)</td>
</tr>
<tr>
<td>Insured status at the health insurance plans, n (%)</td>
<td>3419 (99.97)</td>
<td>225 (49.1)</td>
</tr>
</tbody>
</table>
\( ^a \) Data missing for 1 man.

\( ^b \) Data missing for 497 men and 18 women.

\( ^c \) Data missing for 89 men and 6 women.

\( ^d \) Data missing for 149 men and 7 women.

\( ^e \) Data missing for 787 men and 113 women.

\( ^f \) Data missing for 1258 men and 149 women.

\( ^g \) Data missing for 109 men and 49 women.
<table>
<thead>
<tr>
<th>HbA1c</th>
<th>3 months</th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>total, N (%)</td>
<td>men, N (%)</td>
<td>women, N (%)</td>
<td>total, N (%)</td>
<td>men, N (%)</td>
<td>women, N (%)</td>
<td>total, N (%)</td>
</tr>
<tr>
<td>&lt; 6%</td>
<td>1471 (84)</td>
<td>1332 (85)</td>
<td>139 (78)</td>
<td>1394 (80)</td>
<td>1255 (80)</td>
<td>139 (78)</td>
<td>1296 (74)</td>
</tr>
<tr>
<td>6% ≤, &lt; 7%</td>
<td>1063 (69)</td>
<td>935 (70)</td>
<td>128 (62)</td>
<td>875 (57)</td>
<td>771 (58)</td>
<td>104 (50)</td>
<td>735 (47)</td>
</tr>
<tr>
<td>7% ≤, &lt; 8%</td>
<td>165 (57)</td>
<td>146 (58)</td>
<td>19 (53)</td>
<td>126 (44)</td>
<td>110 (43)</td>
<td>16 (44)</td>
<td>97 (34)</td>
</tr>
<tr>
<td>8% ≤, &lt;10%</td>
<td>120 (61)</td>
<td>111 (62)</td>
<td>9 (56)</td>
<td>96 (49)</td>
<td>89 (49)</td>
<td>7 (44)</td>
<td>80 (41)</td>
</tr>
<tr>
<td>10% ≤</td>
<td>46 (46)</td>
<td>39 (51)</td>
<td>7 (37)</td>
<td>35 (35)</td>
<td>29 (36)</td>
<td>6 (32)</td>
<td>27 (27)</td>
</tr>
</tbody>
</table>
Table 3 - Age-adjusted Odds Ratios for No Follow-ups, Stratified by Gender

<table>
<thead>
<tr>
<th></th>
<th>Men</th>
<th></th>
<th></th>
<th>Women</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>3 months</td>
<td>6 months</td>
<td>12months</td>
<td>3 months</td>
<td>6 months</td>
<td>12months</td>
</tr>
<tr>
<td></td>
<td>OR (95% CI)</td>
<td>OR (95% CI)</td>
<td>OR (95% CI)</td>
<td>OR (95% CI)</td>
<td>OR (95% CI)</td>
<td>OR (95% CI)</td>
</tr>
<tr>
<td>Body mass index, 1 kg/m² increase</td>
<td>0.95(0.93-0.96)</td>
<td>0.93(0.91-0.94)</td>
<td>0.92(0.91-0.94)</td>
<td>0.98(0.94-1.02)</td>
<td>0.97(0.93-1.00)</td>
<td>0.96(0.92-0.99)</td>
</tr>
<tr>
<td>Fasting glucose, 10 mg/dl increase</td>
<td>0.93(0.91-0.95)</td>
<td>0.95 (0.93-0.97)</td>
<td>0.95(0.93-0.97)</td>
<td>0.97(0.92-1.02)</td>
<td>0.97(0.92-1.02)</td>
<td>0.99(0.94-1.04)</td>
</tr>
<tr>
<td>HbA₁c, 1% increase</td>
<td>0.69(0.65-0.73)</td>
<td>0.65(0.61-0.69)</td>
<td>0.61(0.57-0.65)</td>
<td>0.74(0.64-0.87)</td>
<td>0.68 (0.58-0.81)</td>
<td>0.70(0.59-0.84)</td>
</tr>
<tr>
<td>Glycosuria</td>
<td>0.59(0.49-0.70)</td>
<td>0.59(0.50-0.69)</td>
<td>0.55(0.46-0.64)</td>
<td>0.29(0.17-0.52)</td>
<td>0.25 (0.14-0.46)</td>
<td>0.28(0.15-0.51)</td>
</tr>
<tr>
<td>Proteinuria</td>
<td>0.54(0.41-0.72)</td>
<td>0.50 (0.38-0.66)</td>
<td>0.46(0.35-0.61)</td>
<td>0.32(0.14-0.73)</td>
<td>0.39 (0.17-0.92)</td>
<td>0.42(0.18-1.02)</td>
</tr>
<tr>
<td>Hypertension</td>
<td>0.43(0.35-0.52)</td>
<td>0.39 (0.32-0.47)</td>
<td>0.35(0.29-0.43)</td>
<td>0.55(0.31-0.96)</td>
<td>0.44 (0.24-0.79)</td>
<td>0.19(0.09-0.39)</td>
</tr>
<tr>
<td>Dyslipidemia</td>
<td>0.28(0.22-0.35)</td>
<td>0.28 (0.22-0.35)</td>
<td>0.25(0.19-0.32)</td>
<td>0.39(0.21-0.72)</td>
<td>0.28 (0.14-0.56)</td>
<td>0.23(0.93-0.97)</td>
</tr>
<tr>
<td>Depression</td>
<td>0.50(0.37-0.67)</td>
<td>0.18 (0.04-0.90)</td>
<td>0.43(0.32-0.58)</td>
<td>0.81(0.38-1.71)</td>
<td>0.29 (0.02-3.47)</td>
<td>0.58(0.27-1.24)</td>
</tr>
<tr>
<td>Smoking</td>
<td>1.02(0.87-1.20)</td>
<td>0.93 (0.80-1.07)</td>
<td>0.93(0.81-1.07)</td>
<td>0.79(0.37-1.68)</td>
<td>1.11 (0.52-2.37)</td>
<td>1.09(0.52-2.28)</td>
</tr>
<tr>
<td>Alcohol consumption</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Everyday</td>
<td>1.00 (Reference)</td>
<td>1.00 (Reference)</td>
<td>1.00 (Reference)</td>
<td>1.00 (Reference)</td>
<td>1.00 (Reference)</td>
<td>1.00 (Reference)</td>
</tr>
<tr>
<td>Sometimes</td>
<td>0.80(0.62-1.03)</td>
<td>0.87 (0.69-1.10)</td>
<td>0.79(0.63-1.00)</td>
<td>2.60(0.58-11.58)</td>
<td>2.91 (0.62-13.7)</td>
<td>2.06(0.44-9.66)</td>
</tr>
<tr>
<td>Rarely</td>
<td>0.86(0.67-1.12)</td>
<td>0.91 (0.72-1.14)</td>
<td>0.83(0.67-1.05)</td>
<td>2.01(0.52-7.73)</td>
<td>2.25 (0.55-9.29)</td>
<td>1.84(0.45-7.63)</td>
</tr>
<tr>
<td>Haemoglobin (g/dl)</td>
<td>1.02(0.95-1.08)</td>
<td>0.98 (0.92-1.04)</td>
<td>0.96(0.90-1.01)</td>
<td>0.98(0.86-1.11)</td>
<td>0.93 (0.82-1.06)</td>
<td>0.96(0.85-1.09)</td>
</tr>
<tr>
<td>Health insurance plan</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>A</td>
<td>1.00 (Reference)</td>
<td>1.00 (Reference)</td>
<td>1.00 (Reference)</td>
<td>1.00 (Reference)</td>
<td>1.00 (Reference)</td>
<td>1.00 (Reference)</td>
</tr>
<tr>
<td>B</td>
<td>1.12(0.91-1.36)</td>
<td>1.21 (1.01-1.46)</td>
<td>1.25(1.04-1.50)</td>
<td>0.80(0.49-1.31)</td>
<td>0.94 (0.59-1.50)</td>
<td>1.15(0.73-1.84)</td>
</tr>
<tr>
<td>C</td>
<td>0.54(0.35-0.84)</td>
<td>0.82 (0.54-1.25)</td>
<td>0.74(0.48-1.12)</td>
<td>1.04(0.26-4.25)</td>
<td>0.71 (0.20-2.55)</td>
<td>1.08(0.30-3.87)</td>
</tr>
<tr>
<td>Dependent status at health insurance plans</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>0.57(0.36-0.90)</td>
<td>0.60 (0.39-0.92)</td>
<td>0.49(0.32-0.75)</td>
</tr>
</tbody>
</table>

NA, Not applicable.
aAdjusted for age.
<table>
<thead>
<tr>
<th></th>
<th>Men</th>
<th>Women</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>3 months OR (95% CI)</td>
<td>6 months OR (95% CI)</td>
<td>12 months OR (95% CI)</td>
<td>3 months OR (95% CI)</td>
</tr>
<tr>
<td>Age, 10 year increase</td>
<td>0.82 (0.75-0.90)</td>
<td>0.78 (0.71-0.85)</td>
<td>0.76 (0.70-0.83)</td>
<td>0.82 (0.64-1.03)</td>
</tr>
<tr>
<td>Body mass index, 1 kg/m²</td>
<td>0.999 (0.98-1.02)</td>
<td>0.979 (0.96-0.998)</td>
<td>0.98 (0.96-0.99)</td>
<td>1.02 (0.97-1.06)</td>
</tr>
<tr>
<td>HbA₁c, 1 % increase</td>
<td>0.69 (0.65-0.74)</td>
<td>0.67 (0.62-0.71)</td>
<td>0.64 (0.59-0.69)</td>
<td>0.78 (0.65-0.92)</td>
</tr>
<tr>
<td>Proteinuria</td>
<td>0.73 (0.54-0.99)</td>
<td>0.72 (0.54-0.97)</td>
<td>0.69 (0.51-0.94)</td>
<td>0.369 (0.15-0.91)</td>
</tr>
<tr>
<td>Hypertension</td>
<td>0.55 (0.44-0.68)</td>
<td>0.49 (0.40-0.61)</td>
<td>0.45 (0.37-0.56)</td>
<td>0.67 (0.36-1.28)</td>
</tr>
<tr>
<td>Dyslipidemia</td>
<td>0.36 (0.28-0.46)</td>
<td>0.38 (0.30-0.49)</td>
<td>0.35 (0.27-0.46)</td>
<td>0.39 (0.20-0.76)</td>
</tr>
<tr>
<td>Depression</td>
<td>0.17 (0.04-0.78)</td>
<td>0.14 (0.03-0.74)</td>
<td>0.20 (0.04-1.07)</td>
<td>0.19 (0.016-2.31)</td>
</tr>
<tr>
<td>Health insurance plan</td>
<td>A 1.00 (Reference)</td>
<td>1.00 (Reference)</td>
<td>1.00 (Reference)</td>
<td>1.00 (Reference)</td>
</tr>
<tr>
<td></td>
<td>B 0.85 (0.69-1.06)</td>
<td>0.89 (0.73-1.09)</td>
<td>0.89 (0.73-1.09)</td>
<td>0.58 (0.33-1.001)</td>
</tr>
<tr>
<td></td>
<td>C 0.50 (0.32-0.78)</td>
<td>0.77 (0.49-1.20)</td>
<td>0.69 (0.44-1.07)</td>
<td>0.71 (0.15-3.34)</td>
</tr>
<tr>
<td>Dependent status at health insurance plan</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>0.63 (0.36-1.11)</td>
</tr>
</tbody>
</table>

NA, Not applicable.

\(^a\) Adjusted for age, proteinuria, HbA₁c, BMI, hypertension, dyslipidemia, depression, health insurance plan.

\(^b\) Adjusted for age, proteinuria, HbA₁c, BMI, hypertension, dyslipidemia, depression, health insurance plan, and dependent status at the health insurance plan.