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Development of type 2 diabetesmellitus in people with intermediate hyperglycaemia (Review)

Richter, B; Hemmingsen, B; Metzendorf, Maria Inti; Takwoingi, Yemisi

DOI: 10.1002/14651858.CD012661.pub2

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Document Version Publisher's PDF, also known as Version of record

Citation for published version (Harvard):

Richter, B, Hemmingsen, B, Netzendorf, MI & Takwoingi, Y 2018, 'Development of type 2 diabetesmellitus in people with intermediate hyperglycaemia (Review)', *Cochrane Database of Systematic Reviews*, vol. 2018, no. 10, CD012661. https://doi.org/10.1002/14651858.CD012661.pub2

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[Prognosis Review]

Development of type 2 diabetes mellitus in people with intermediate hyperglycaemia

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Editorial group: Cochrane Metabolic and Endocrine Disorders Group. **Publication status and date:** New, published in Issue 10, 2018.

Citation: Richter B, Hemmingsen B, Metzendorf MI, Takwoingi Y. Development of type 2 diabetes mellitus in people with intermediate hyperglycaemia. *Cochrane Database of Systematic Reviews* 2018, Issue 10. Art. No.: CD012661. DOI: 10.1002/14651858.CD012661.pub2.

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ABSTRACT

Background

Intermediate hyperglycaemia (IH) is characterised by one or more measurements of elevated blood glucose concentrations, such as impaired fasting glucose (IFG), impaired glucose tolerance (IGT) and elevated glycosylated haemoglobin A1c (HbA1c). These levels are higher than normal but below the diagnostic threshold for type 2 diabetes mellitus (T2DM). The reduced threshold of 5.6 mmol/ L (100 mg/dL) fasting plasma glucose (FPG) for defining IFG, introduced by the American Diabetes Association (ADA) in 2003, substantially increased the prevalence of IFG. Likewise, the lowering of the HbA1c threshold from 6.0% to 5.7% by the ADA in 2010 could potentially have significant medical, public health and socioeconomic impacts.

Objectives

To assess the overall prognosis of people with IH for developing T2DM, regression from IH to normoglycaemia and the difference in T2DM incidence in people with IH versus people with normoglycaemia.

Search methods

We searched MEDLINE, Embase, ClincialTrials.gov and the International Clinical Trials Registry Platform (ICTRP) Search Portal up to December 2016 and updated the MEDLINE search in February 2018. We used several complementary search methods in addition to a Boolean search based on analytical text mining.

Selection criteria

We included prospective cohort studies investigating the development of T2DM in people with IH. We used standard definitions of IH as described by the ADA or World Health Organization (WHO). We excluded intervention trials and studies on cohorts with additional comorbidities at baseline, studies with missing data on the transition from IH to T2DM, and studies where T2DM incidence was evaluated by documents or self-report only.

Data collection and analysis

One review author extracted study characteristics, and a second author checked the extracted data. We used a tailored version of the Quality In Prognosis Studies (QUIPS) tool for assessing risk of bias. We pooled incidence and incidence rate ratios (IRR) using a random-effects model to account for between-study heterogeneity. To meta-analyse incidence data, we used a method for pooling

proportions. For hazard ratios (HR) and odds ratios (OR) of IH versus normoglycaemia, reported with 95% confidence intervals (CI), we obtained standard errors from these CIs and performed random-effects meta-analyses using the generic inverse-variance method. We used multivariable HRs and the model with the greatest number of covariates. We evaluated the certainty of the evidence with an adapted version of the GRADE framework.

Main results

We included 103 prospective cohort studies. The studies mainly defined IH by IFG_{5.6} (FPG mmol/L 5.6 to 6.9 mmol/L or 100 mg/ dL to 125 mg/dL), IFG_{6.1} (FPG 6.1 mmol/L to 6.9 mmol/L or 110 mg/dL to 125 mg/dL), IGT (plasma glucose 7.8 mmol/L to 11.1 mmol/L or 140 mg/dL to 199 mg/dL two hours after a 75 g glucose load on the oral glucose tolerance test, combined IFG and IGT (IFG/IGT), and elevated HbA1c (HbA1c_{5.7}: HbA1c 5.7% to 6.4% or 39 mmol/mol to 46 mmol/mol; HbA1c_{6.0}: HbA1c 6.0% to 6.4% or 42 mmol/mol to 46 mmol/mol). The follow-up period ranged from 1 to 24 years. Ninety-three studies evaluated the overall prognosis of people with IH measured by cumulative T2DM incidence, and 52 studies evaluated glycaemic status as a prognostic factor for T2DM by comparing a cohort with IH to a cohort with normoglycaemia. Participants were of Australian, European or North American origin in 41 studies; Latin American in 7; Asian or Middle Eastern in 50; and Islanders or American Indians in 5. Six studies included children and/or adolescents.

Cumulative incidence of T2DM associated with IFG_{5.6}, IFG_{6.1}, IGT and the combination of IFG/IGT increased with length of followup. Cumulative incidence was highest with IFG/IGT, followed by IGT, IFG_{6.1} and IFG_{5.6}. Limited data showed a higher T2DM incidence associated with HbA1c_{6.0} compared to HbA1c_{5.7}. We rated the evidence for overall prognosis as of moderate certainty because of imprecision (wide CIs in most studies). In the 47 studies reporting restitution of normoglycaemia, regression ranged from 33% to 59% within one to five years follow-up, and from 17% to 42% for 6 to 11 years of follow-up (moderate-certainty evidence).

Studies evaluating the prognostic effect of IH versus normoglycaemia reported different effect measures (HRs, IRRs and ORs). Overall, the effect measures all indicated an elevated risk of T2DM at 1 to 24 years of follow-up. Taking into account the long-term follow-up of cohort studies, estimation of HRs for time-dependent events like T2DM incidence appeared most reliable. The pooled HR and the number of studies and participants for different IH definitions as compared to normoglycaemia were: IFG_{5.6}: HR 4.32 (95% CI 2.61 to 7.12), 8 studies, 9017 participants; IFG_{6.1}: HR 5.47 (95% CI 3.50 to 8.54), 9 studies, 2818 participants; IGT: HR 3.61 (95% CI 2.31 to 5.64), 5 studies, 4010 participants; IFG and IGT: HR 6.90 (95% CI 4.15 to 11.45), 5 studies, 1038 participants; HbA1c_{5.7}: HR 5.55 (95% CI 2.77 to 11.12), 4 studies, 5223 participants; HbA1c_{6.0}: HR 10.10 (95% CI 3.59 to 28.43), 6 studies, 4532 participants. In subgroup analyses, there was no clear pattern of differences between geographic regions. We downgraded the evidence for the prognostic effect of IH versus normoglycaemia to low-certainty evidence due to study limitations because many studies did not adequately adjust for confounders. Imprecision and inconsistency required further downgrading due to wide 95% CIs and wide 95% prediction intervals (sometimes ranging from negative to positive prognostic factor to outcome associations), respectively.

This evidence is up to date as of 26 February 2018.

Authors' conclusions

Overall prognosis of people with IH worsened over time. T2DM cumulative incidence generally increased over the course of followup but varied with IH definition. Regression from IH to normoglycaemia decreased over time but was observed even after 11 years of follow-up. The risk of developing T2DM when comparing IH with normoglycaemia at baseline varied by IH definition. Taking into consideration the uncertainty of the available evidence, as well as the fluctuating stages of normoglycaemia, IH and T2DM, which may transition from one stage to another in both directions even after years of follow-up, practitioners should be careful about the potential implications of any active intervention for people 'diagnosed' with IH.

PLAIN LANGUAGE SUMMARY

Development of type 2 diabetes mellitus in people with intermediate hyperglycaemia ('prediabetes')

Review question

We wanted to find out whether raised blood sugar ('prediabetes') increases the risk of developing type 2 diabetes and how many of these people return to having normal blood sugar levels (normoglycaemia). We also investigated the difference in type 2 diabetes development in people with prediabetes compared to people with normoglycaemia.

Background

Type 2 diabetes is often diagnosed by blood sugar measurements. These include fasting blood glucose, which is a measurement of the sugar in blood after an oral glucose tolerance test (drinking 75 g of glucose on an empty stomach) or by measuring glycosylated haemoglobin A1c (HbA1c), a long-term marker of blood glucose levels. Type 2 diabetes can have bad effects on health in the long term (diabetic complications), like severe eye or kidney disease or diabetic feet, eventually resulting in foot ulcers.

Raised blood glucose levels (hyperglycaemia), which are above normal ranges but below the limit of diagnosing type 2 diabetes, indicate prediabetes, or intermediate hyperglycaemia. The way prediabetes is defined has important effects on public health because some physicians treat people with prediabetes with medications that can be harmful. For example, reducing the threshold for defining impaired fasting glucose (after an overnight fast) from 6.1 mmol/L or 110 mg/dL to 5.6 mmol/L or 100 mg/dL, as done by the American Diabetes Association (ADA), dramatically increased the number of people diagnosed with prediabetes worldwide.

Study characteristics

We searched for observational studies (studies where no intervention takes place but people are observed over prolonged periods of time) that investigated how many people with prediabetes at the beginning of the study developed type 2 diabetes. We also evaluated studies comparing people with prediabetes to people with normoglycaemia. Prediabetes was defined by different blood glucose measurements.

We found 103 studies, monitoring people over 1 to 24 years. More than 250,000 participants began the studies. In 41 studies the participants were of Australian, European or North American origin, in 7 studies participants were primarily of Latin American origin and in 50 studies participants were of Asian or Middle Eastern origin. Three studies had American Indians as participants, and one study each invited people from Mauritius and Nauru. Six studies included children, adolescents or both as participants.

This evidence is up to date as of 26 February 2018.

Key results

Generally, the development of new type 2 diabetes (diabetes incidence) in people with prediabetes increased over time. However, many participants also reverted from prediabetes back to normal blood glucose levels. Compared to people with normoglycaemia, those with prediabetes (any definition) showed an increased risk of developing type 2 diabetes, but results showed wide differences and depended on how prediabetes was measured. There were no clear differences with regard to several regions in the world or different populations. Because people with prediabetes may develop diabetes but may also change back to normoglycaemia almost any time, doctors should be careful about treating prediabetes because we are not sure whether this will result in more benefit than harm, especially when done on a global scale affecting many people worldwide.

Certainty of the evidence

The certainty of the evidence for overall prognosis was moderate because results varied widely. The certainty of evidence for studies comparing prediabetic with normoglycaemic people was low because the results were not precise and varied widely. In our included observational studies the researchers often did not investigate well enough whether factors like physical inactivity, age or increased body weight also influenced the development of type 2 diabetes, thus making the relationship between prediabetes and the development of type 2 diabetes less clear.

SUMMARY OF FINDINGS FOR THE MAIN COMPARISON [Explanation]

Outcome: development of T2DM Prognosis of people with intermediate hyperglycaemia

| Follow-up (years) | Cumulative T2 [no of studies; | Regression from intermediate hy- pergly- caemia to normo- glycaemia % (95% CI) [no of studies; no of participants | | | | | | |
|----------------------|----------------------------------|--|---------------------------------|--------------------------------|--------------------------------|--------------------------------|-----------------------------------|-------------------------------|
| | IFG _{5.6} | IFG _{6.1} | IGT | IFG + IGT | HbA1c _{5.7} | HbA1c _{6.0} | with intermediate hyperglycaemia] | |
| 1 | - | - | 13 (5-23) [3; 671] | 29 (23-36) [1; 207] | - | - | 59 (54-64) [2; 375] | ⊕⊕⊕⊖ Moderate ^b |
| 2 | 2 (1-2) [1; 1335] | 11 (8-14) [2; 549] | 16 (9-26) [9; 1998] | - | - | - | 46 (36-55) [9; 2852] | |
| 3 | 17 (6-32) [3; 1091] | 9 (2-20) [3; 927] | 22 (18-27) [3; 417] | 34 (28-41) [1; 209]- | - | 7 (5-10) [1; 370] | 41 (24-69) [7; 1356] | |
| 4 | 17 (13-22) [3; 800] | 30 (17-44) [2; 1567] | 22 (12-34) [5; 1042] | - | 14 (7-23) [3; 5352] | 44 (40-48) [2; 627] | 33 (26-40) [3; 807] | |
| 5 | 18 (10-27) [7; 3530] | 26 (19-33) [11; 3837] | 39 (25-53) [12; 3444] | 50 (37-63) [5; 478] | 25 (18-32) [4; 3524] | 38 (26-51) [3; 1462] | 34 (27-42) [9; 2603] | |
| 6 | 22 (15-31) [4; 738] | 37 (31-43) [5; 279] | 29 (25-34) [7; 775] | 58 (48-67) [4; 106] | 17 (14-20) [1; 675] | - | 23 (3-53) [5; 1328] | |
| 7 | 18 (8-30) [5; 980] | 15 (0-45) [4; 434] | 19 (13-26) [5; 835] | 32 (20-45) [4; 753] | 21 (16-27) [1; 207] | - | 41 (37-45) [4; 679] | |

| 8 | 34 (27-40) [2; 1887] | 48 (31-66) [1;29] | 43 (37-49) [4; 1021] | 52 (47-57) [1; 356] | - | - | 39 (33-44) [2; 328] |
|----|--------------------------------|--------------------------------|--------------------------------|-------------------------------|--------------------------------|------------------------------|-------------------------------|
| 9 | 38 (10-70) [3; 1356] | - | 53 (45-60) [1; 163] | 84 (74-91) [1; 69] | - | - | 17 (14-22) [1; 299] |
| 10 | 23 (14-33) [6; 1542] | 29 (17-43) [6; 537] | 26 (17-37) [6; 443] | 30 (17-44) [2; 49] | 31 (29-33) [2; 2854] | - | 42 (22-63) [7; 894] |
| 11 | - | 38 (33-43) [1; 402] | 46 (43-49) [1; 1253] | - | - | - | 28 (17-39) [2; 736] |
| 12 | 31 (19-34) [3; 433] | 31 (28-33) [1; 1382] | 41 (38-43) [2; 1552] | 70 (63-76) [2; 207] | - | - | - |
| 15 | | - | - | - | - | 29 (19-40) [1; 70] | - |
| 20 | - | - | 60 (5-68) [1; 114] | - | - | - | |

Cl: confidence interval; HbA1c_{5.7}: glycosylated haemoglobin A1c, 5.7% threshold; HbA1c_{6.0}: glycosylated haemoglobin A1c, 6.0% threshold; IFG_{5.6}: impaired fasting glucose, 5.6 mmol/L threshold; IFG_{6.1}: impaired fasting glucose, 6.1 mmol/L threshold; IGT: impaired glucose tolerance; T2DM: type 2 diabetes mellitus.

GRADE Working Group grades of evidence

High quality: further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Very low quality: we are very uncertain about the estimate.

^aWith phase 2 explanatory studies aiming to confirm independent associations between the prognostic factor and the outcome, GRADE starts with 'high quality' (Huguet 2013). We assumed the GRADE factor publication bias was inherent with this type of research (phase 2 design), so we did not use it as a potential downgrading factor

^bDowngraded by one level because of imprecision (wide CIs for most intermediate hyperglycaemia definitions and the association with T2DM incidence and regression from intermediate hyperglycaemia)

BACKGROUND

For a glossary of terms please see Appendix 1.

'Prediabetes', 'borderline diabetes', 'prediabetic stage', 'high risk of diabetes', 'dysglycaemia' or 'intermediate hyperglycaemia' (IH) are terms used to characterise various measurements of elevated blood glucose concentrations, such as impaired fasting glucose (IFG), impaired glucose tolerance (IGT), elevated glycosylated haemoglobin A1c (HbA1c) or combinations of these conditions (WHO/IDF 2006). Elevated blood glucose levels that indicate hyperglycaemia are too high to be considered normal, but they are below the diagnostic threshold for type 2 diabetes mellitus (T2DM). Therefore, due to the continuous glycaemic spectrum from normal to the diabetic stage, a sound evidence base is needed to define glycaemic thresholds for people at high risk of T2DM, especially because dysglycaemia is commonly an asymptomatic condition, so naturally it often remains undiagnosed (CDC 2015). The various terms used to describe stages of hyperglycaemia may cause people to have marked emotional reactions. For example, the term prediabetes may imply (at least for non-experts) that diabetes is unavoidable, whereas (high) risk of diabetes gives people the impression that they can possibly avoid the disease altogether. In addition to the disputable construct of intermediate health states termed 'predisease' (Viera 2011), many people may associate the label 'prediabetes' with dire consequences. Alternatively, any diagnosis of prediabetes may be an opportunity to reassess, for example, eating habits and physical activity levels, thus enabling affected individuals to actively change their health-related behaviours.

Several institutional bodies like the American Diabetes Association (ADA) and the World Health Organization (WHO) have established commonly used criteria to define people who are at a high risk of developing T2DM.

• In 1979, the National Diabetes Data Group (NDDG) described glucose intolerance as an intermediate metabolic state between normoglycaemia and diabetes (NDDG 1979). NDDG defined this IGT as an elevated plasma glucose concentration (7.8 mmol/L to 11.1 mmol/L or 140 mg/dL to 199 mg/dL) two hours after a 75 g glucose load on the oral glucose tolerance test (OGTT).

• In 1997, the Expert Committe on the Diagnosis and Classification of Diabetes Mellitus and later the WHO defined two intermediate states of glucose regulation existing between regular glucose homeostasis and diabetes: IGT was diagnosed two hours after a 75 g OGTT by a plasma glucose level of 7.8 mmol/L to 11.1 mmol/L (140 mg/dL to 199 mg/dL) or by the concept of IFG (ADA 1997; WHO 1999). The initial definition of IFG was a fasting plasma glucose (FPG) level of 6.1 mmol/L to 6.9 mmol/L (110 mg/dL to 125 mg/dL). In 2003, the ADA reduced the lower threshold to 5.6 mmol/L (100 mg/dL) (ADA 2003). However, the WHO did not endorse this lower cut-off point for IFG (WHO/IDF 2006). • More recently, an elevated HbA1c has been introduced to identify people at high risk of developing T2DM. In 2009, the International Expert Committee (IEC) proposed HbA1c measurements of 6.0% to 6.4% (42 mmol/mol to 46 mmol/mol) to identify people at a high risk of T2DM (IEC 2009). In 2010, the ADA re-defined this HbA1c level as 5.7% to 6.4% (39 mmol/mol to 46 mmol/mol) (ADA 2010), a decision not endorsed by WHO, IEC or other organisations.

The various glycaemic tests do not identify the same people at risk, as there is an imperfect overlap among the glycaemic modalities available to define IH (Cheng 2006; Gosmanov 2014; Morris 2013; Selvin 2011). Unlike IFG and IGT, HbA1c reflects longer-term glycaemic control, that is, how a person's blood glucose concentrations have been during the preceding two to three months (Inzucchi 2012). Compared with IFG and IGT measurements, HbA1c assessments have less intrapersonal variability when repeated. However, haemoglobin variants, genetic haemoglobinopathies, thalassemias and iron deficiency anaemia substantially influence HbA1c measurements (Mostafa 2011). The FPG thresholds of defining IFG and the question whether HbA1c is an adequate tool to diagnose IH are still a subject of debate (Buysschaert 2011; Buysschaert 2016). In studies investigating the risk of IH as measured by HbA1c, the association is probably underestimated if time-dependent effects are not taken into account (Lind 2009). On the other hand, some investigators question whether HbA1c as such is the right outcome measure for studies of diabetes (Lipska 2017).

Also, IFG and IGT differ in their age and sex distribution, and both increase with advancing age (Nathan 2007), as glucose tolerance deteriorates with age (Gale 2013). 'Ethnicity' and geography are additional important features: the prevalence of elevated HbA1c in black people is twice as high as in non-Hispanic white people, but the opposite is true for IGT (Selvin 2011; Ziemer 2010). The number of people with IH identified in South Asian compared with European cohorts and the associated cardiovascular disease (CVD) risk depend on how prediabetes is diagnosed (Eastwood 2016).

The increase in T2DM results from an interaction between genetic and environmental factors, reflecting behavioural changes over time such as decreased physical activity levels and increased body weight (DeFronzo 2011; Nathan 2007). Both IFG and IGT are insulin-resistant states, and insulin resistance is thought to be the core defect in T2DM: people with (isolated) IFG predominantly have β -cell dysfunction with impaired insulin secretion (DeFronzo 1989), plus moderate hepatic insulin resistance, but near-normal muscle insulin sensitivity. The consequence is excessive fasting hepatic glucose production followed by elevated FPG. During an OGTT the early insulin response (0 to 30/60 min) is impaired, resulting in an excessive early rise in postload glucose (PG). The late insulin response (60 min to 120 min) appears in-

tact and the two-hour PG returns to its approximately starting FPG level (DeFronzo 2011; Nathan 2007). People with (isolated) IGT have normal to slightly reduced hepatic insulin sensitivity and moderate to severe muscle insulin resistance (Abdul-Ghani 2006; Jensen 2002). During an OGTT both the early and the late insulin response are impaired. Hyperglycaemia is progressive and prolonged after the glucose load, and the two-hour PG remains above its starting FPG level (DeFronzo 2011; Nathan 2007).

There are some known risk indicators for the development of T2DM, including a positive family history, gestational diabetes mellitus, obesity, 'ethnicity' (e.g. the risk of diabetes is thought to be higher among Asians, Hispanics, and 'black' people), polycystic ovarian syndrome, impaired insulin secretion and insulin resistance, abnormal coagulation factors and endothelial dysfunction. However, the evidence base for the weight of a single risk indicator and the interplay of various factors is still under investigation. Type 2 diabetes mellitus is a rather complex metabolic state and could be described as an asymptomatic risk factor for a future disease (Yudkin 2016), and hence prediabetes a risk factor for another risk factor (Nathan 2007).

Diabetes is a category, whereas IFG and IGT reflect a continuous variable with more or less arbitrarily chosen cut-off points (Yudkin 1990; Yudkin 2014). The reduced lower threshold of 5.6 mmol/L (100 mg/dL) to define IFG by the ADA in 2003 substantially increased the prevalence of IFG with potentially significant public health and socioeconomic implications (Davidson 2003; Yudkin

2014; Yudkin 2016). Some authors have argued that substantial benefits might ensue even if it were only possible to delay the onset of diabetes by detecting and treating prediabetes (Cefalu 2016). Interestingly, some people with IH will not develop T2DM, and some people will return or 'regress' to normoglycaemia. In the Diabetes Prevention Program (DPP), the hazard ratio of developing T2DM was 0.44 (95% confidence interval 0.37 to 0.55) in people having at least one normal OGTT during the DPP compared with people who never regressed to normoglycaemia during the DPP (Perreault 2012; Perreault 2014). The ADA associated regression with remission and defined it as a partial or complete diabetes remission of glycaemic measurements for at least one year without pharmacological or surgical interventions (Buse 2009). This could have significant impact on "the therapeutic strategy from diabetes prevention and lifelong glucose-lowering treatment to induction of regression and monitoring for relapse" (Yakubovich 2012).

OBJECTIVES

Objective 1: to assess the overall prognosis of people with IH for the development of T2DM and to assess how many people with IH revert back to normoglycaemia (regression).

With regard to objective 1 we established the following 'Population, Intervention, Outcome, Timing, Setting' (PICOTS) table (adapted according to the PICOTS system presented in Debray 2017).

| Item | Definition |
|---------------|---|
| Population | People with intermediate hyperglycaemia (defined by IFG, IGT or elevated HbA1c) |
| Intervention | None |
| Comparator | None |
| Outcome | Development of type 2 diabetes Regression to normoglycaemia |
| Timing | At least 1 year follow-up |
| Setting | Outpatients |
| IFG: impaired | l fasting glucose; IGT : impaired glucose tolerance; HbA1c : glycosylated haemoglobin A1c |

Objective 2: to assess the difference in T2DM incidence in people with IH versus people with normoglycaemia. With regard to objective 2 we established the following PICOTS table (adapted according to the PICOTS system presented in Debray 2017).

| Item | Definition |
|--------------|---|
| Population | People with intermediate hyperglycaemia (defined by IFG, IGT or elevated HbA1c) |
| Intervention | Intermediate hyperglycaemia as a prognostic factor |
| Comparator | Normoglycaemia |
| Outcome | Development of type 2 diabetes |
| Timing | At least one year follow-up |
| Setting | Outpatients |
| | |

IFG: impaired fasting glucose; IGT: impaired glucose tolerance; HbA1c: glycosylated haemoglobin A1c

METHODS

Criteria for considering studies for this review

Study design

Prospective cohort studies investigating either the overall prognosis of people with IH for developing T2DM or IH versus normoglycaemia as a prognostic factor for developing T2DM (Altman 2001).

Inclusion criteria

Types of participants

To study the overall prognosis of people with IH and regression from IH to normoglycaemia, we included cohort studies in people with IH at baseline, defined by impaired fasting glucose (IFG), impaired glucose tolerance (IGT), elevated glycosylated haemoglobin A1c (HbA1c) or any combination of these. IH had to be established by standard cut-off values for IFG, IGT or elevated HbA1c, as defined by ADA or WHO (ADA 1997; ADA 2003; ADA 2010; ICH 1997; IEC 2009; WHO 1998; WHO/IDF 2006).

To study whether IH compared to normoglycaemia is a prognostic factor for developing T2DM, we included cohort studies in people with IH and normoglycaemia at baseline.

Definition of IH

We defined IH according to ADA and WHO descriptions.

 IFG_{5.6} threshold, usually defined as a fasting plasma glucose level between 5.6 mmol/L and 6.9 mmol/L at baseline.

• IFG_{6.1} threshold, usually defined as a fasting plasma glucose level between 6.1 mmol/L and 6.9 mmol/L at baseline.

• IGT, usually defined as a plasma glucose level between 7.8 mmol/L and 11.1 mmol/L two hours after a 75 g OGTT at baseline.

• Isolated IFG was defined as IFG_{5.6} or IFG_{6.1} only (without IGT), and isolated IGT was defined as IGT only (without IFG_{5.6} or IFG_{6.1}).

• HbA1c_{5.7} threshold, usually defined as HbA1c measurement between 5.7% and 6.4% at baseline.

• HbA1c_{6.0} threshold, usually defined as HbA1c measurement between 6.0% and 6.4% at baseline.

Types of outcome measures

Our outcome of primary interest was the diagnosis of newly developed T2DM (T2DM incidence). T2DM incidence should have been diagnosed by blood glucose measurements such as fasting plasma glucose (FPG), two-hour postload glucose (PG) or HbA1c. Diagnosis could have been combined with self-reported diabetes, physician-diagnosed diabetes or use of antihyperglycaemic medications such as oral hypoglycaemic drugs, insulin or both.

Exclusion criteria

• Intervention trials and study designs other than prospective cohort studies.

• People with comorbidities at baseline (e.g. people with coronary heart disease and IGT).

• Missing data on transition from IH to T2DM.

• Follow-up period after baseline assessment not specified (not possible to associate T2DM incidence with length of follow-up).

• T2DM incidence evaluated by documents (e.g. hospital records, retrospective use of registers) or self-report only.

Search methods for identification of studies

The fundamental challenge of this review question was to define the population of interest, that is, people with IH. We expected a great number of terms describing this population, such as people with prediabetes, mentions of IFG, IGT or HbA1c somewhere in the title or abstract of relevant publications, and terms like risk factors, predictors, prevalence, incidence and several other concepts which cannot be foreseen when developing a Boolean search strategy in a conceptual way.

One option to address this problem would have been to design a highly sensitive search strategy, which would have resulted in a yield of more than 15,000 references, which was unfeasible for fast human screening but could be addressed in the future with robust automated classification algorithms. Instead, we designed a more specific Boolean search approach based on text analysis and augmented by the following complementary search methods.

1. Identification of systematic reviews addressing our review question.

2. Careful checking of reference lists and Discussion sections of relevant studies.

3. A non-human skill dependent search method based on PubMed's 'similar articles' algorithm.

Boolean search

We developed the search strategy using analytical text mining of 44 relevant publications (range of publication years 2008 to 2015, from 31 journals) already known to review author BR. We used the tools PubReMiner, TerMine and AntConc and applied the prognosis filters by the Hedges Team (Wilczynski 2004; Wilczynski 2005).

We searched the following sources from database inception to the specified date.

• MEDLINE Ovid Epub Ahead of Print, In-Process & Other Non-Indexed Citations, Ovid MEDLINE(R) Daily and Ovid MEDLINE(R) (1946 to 15 December 2016 and then updated to 26 February 2018).

• Embase Ovid (1974 to 2016 Week 50, last searched 15 December 2016).

- ClinicalTrials.gov (searched 15 December 2016).
- WHO International Clinical Trials Registry Platform

(ICTRP) Search Portal (apps.who.int/trialsearch; searched 15 December 2016).

Before publication, we updated the MEDLINE search as reflected above. We restricted the update to MEDLINE because 98% of the publications of included studies identified up to the point of updating (on 26 February 2018) were indexed in MEDLINE. The search strategy consisted of two tiers.

1. Prediabetes as predictor for cardiovascular disease (CVD), mortality, stroke, cancer, micro- and macrovascular complications.

2. Prediabetes as predictor for diabetes incidence.

We combined both strategies with the conjunction 'OR' because it was likely that search results for prediabetes as a predictor for complications also contained data on diabetes incidence. For details of all search strategies see Appendix 2.

Study extraction of relevant systematic reviews

In addition, we extracted relevant publications from 16 identified systematic reviews (Echouffo-Tcheugui 2016; Erqou 2013; Ford 2010; Hope 2016; Huang 2014b; Huang 2014a; Huang 2016; Lee 2012; Morris 2013; Santos-Oliveira 2011; Sarwar 2010; Schottker 2016; Twito 2015; Xu 2015; Zhang 2012a; Zhong 2016).

Reference checking of included studies

We extracted relevant publications after handsearching the full texts of included studies (Methods section, Discussion section, reference lists).

'Similar articles'-based search method

On 15 March 2018 we ran PubMed's 'similar articles' algorithm with the 224 publications of included studies identified by our search methods so far ('seed publications' in Appendix 2). When using the 'similar articles' algorithm, search results in PubMed are retrieved and ranked according to pre-calculated similarities of the seed publications. We downloaded the first 500 results (of 24,124), deduplicated them against the already identified seed publications and screened the resulting set.

Selection of studies

Two review authors (BR and BH) independently scanned the title, abstract, or both, of every record retrieved in the literature searches to determine which studies to assess further. We investigated the full text of all potentially relevant articles, resolving discrepancies through consensus or by recourse to a third review author (MIM). We prepared a flow diagram of the number of studies identified and excluded at each stage in accordance with the PRISMA flow diagram of study selection (Liberati 2009).

Data extraction and management

For studies that fulfilled our inclusion criteria, one review author (BR) extracted key study characteristics, inclusion and exclusion criteria of study participants, stated aim of the study, definitions of prognosis, prognostic factor and outcome (normoglycaemia, intermediate glycaemia and T2DM incidence), baseline characteristics of study participants and data on transition from IH (as defined by IFG, IGT, elevated HbA1c or combinations thereof) to T2DM. Another author (MIM) checked these data extractions, and we resolved any disagreements by discussion or, if required, by consultation with a third review author (BH). We used parts of the checklist for critical appraisal and data extraction for systematic **r**eviews of prediction **m**odelling **s**tudies (CHARMS), which helps to evaluate prediction modelling studies (Moons 2014), and we established our own context-specific data extraction sheets after piloting data extraction for 15 studies.

Dealing with companion publications

In the event of companion publications or multiple reports of a prospective cohort study (e.g. because of different time points investigated) we focused on the analysis of the publication describing the longest follow-up from baseline and extracted data from shorter follow-ups in case some measures were not reported in the publication on the longest follow-up (e.g. the most recent paper might have described the association between elevated HbA1c and T2DM incidence, but an older publication might have described the association between IGT and T2DM incidence). Companion publications or multiple reports of a primary study were listed as secondary references under the primary reference of the included, ongoing or excluded study.

Assessment of risk of bias in included studies

One review author (BR) assessed the risk of bias of each included study and another review author (MIM) checked the accuracy of this assessment. We resolved any disagreements by consensus, or by consultation with a third review author (BH). We used a tailored version of the Quality In Prognosis Studies (QUIPS) tool for assessing risk of bias in studies of the prognostic factor IH versus normoglycaemia (Dretzke 2014; Hayden 2013; see Appendix 3). Our tool consisted of six risk of bias domains: study participation, study attrition, glycaemic status measurement, outcome measurement, study confounding; and statistical analysis and reporting. The study participation domain consisted of five items: description of the source population or population of interest, description of the baseline study sample, adequate description of the sampling frame and recruitment, adequate description of the period and place of recruitment, and adequate description of inclusion and exclusion criteria. The study attrition domain consisted of four items: description of attempts to collect information on participants who dropped out, reasons for loss to follow-up provided, adequate description of participants lost to follow-up, and

no important differences between participants who completed the study and those who did not. The glycaemic status measurement domain consisted of four items: provision of clear definition or description of the glycaemic status, adequately valid and reliable method of measuring glycaemic status, reporting of continuous variables or use of appropriate cut points, and use of same method and setting of measurement of glycaemic status in all study participants. The outcome measurement domain consisted of three items: provision of clear definition of the outcome, use of adequately valid and reliable method of outcome measurement, and use of same method and setting of outcome measurement in all study participants. The study confounding domain consisted of the seven items: measurement of all important confounders, provision of clear definitions of the important confounders measured, adequately valid and reliable measurement of all important confounders, use of same method and setting of confounding measurement in all study participants, appropriate imputation methods used for missing confounders (if applicable), important potential confounders accounted for in the study design, and important potential confounders accounted for in the analysis. The statistical analysis and reporting domain consisted of two items: sufficient presentation of data to assess the adequacy of the analytic strategy, and adequate statistical model for the design of the study. There is no recommended tool for assessing risk of bias in studies of overall prognosis. Therefore, we applied the tailored QUIPS tool to these studies as well but without the domains for study confounding and statistical analysis and reporting because these were not suitable to basic calculations of cumulative incidence. We planned to investigate the influence of low risk of bias (low risk of bias in all domains) versus unclear/high risk of bias (unclear or high risk of bias in at least one of these domains).

Measures of T2DM incidence and unit of analyses issues

If more than one group from the same cohort study was eligible for inclusion in the same meta-analysis, we included the groups only if separate information was available (e.g. data on T2DM incidence for female and male participants). If more than one time point of T2DM was available for a study (e.g. cumulative incidence data) we included data in the appropriate meta-analysis for each time point separately and did not pool data across different follow-up periods.

Data synthesis

Our primary aim for overall prognosis in people with IH was to provide a transparent overview of the whole data matrix describing a wide variety of possible associations between various isolated and combined definitions of IH and incident T2DM in dissimilar populations covering diverse time periods. We also evaluated whether IH compared to normoglycaemia is a prognostic factor for developing T2DM.

First, we grouped studies on IH definitions, i.e. isolated IFG 5.6 mmol/L to 6.9 mmol/L (**IFG**_{5.6} **threshold**), isolated IFG 6.1 mmol/L to 6.9 mmol/L (**IFG**_{6.1} **threshold**), isolated **IGT** (glucose concentration 7.8 mmol/L to 11.1 mmol/L two hours after a 75 g glucose load on the OGTT), **IFG and IGT** combined, HbA1c 6.0% to 6.4% (**HbA1c**_{6.0} **threshold**), and HbA1c 5.7% to 6.4% (**HbA1c**_{5.7} **threshold**). Then we evaluated subgroups of different geographic locations/ ethnicities' for each IH definition. We expected the following outcome measures.

• Cases (cumulative incidence at follow-up; e.g. 20 new diabetes cases out of 400 people with IFG at baseline (5%)) and cumulative incidence rates (cases per 1000 person-years) for overall prognosis of people with IH.

• Odds ratios (ORs), incidence rate ratios (IRRs), and hazard ratios (HRs) for IH versus normoglycaemia as a prognostic factor for developing T2DM.

We pooled incidence and incidence rate ratios (IRR) using a random-effects model to account for between-study heterogeneity. For meta-analysis of incidence data, we used a method for pooling proportions which uses the Freeman-Tukey Double Arcsine Transformation to stabilise the variances (Freeman 1950). The meta-analysis was performed using the Stata software user written programme metaprop (Stata 2015). For the confidence intervals (CI) for individual studies shown on the forest plots for incidence, we used the Wilson approach (Newcombe 1998). For meta-analysis of IRRs, we first computed the log IRRs and their approximate standard errors and then used an inverse variance weighted random-effects model to pool the log IRRs (Hasselblad 1994; Higgins 2011b). We exponentiated the pooled log IRR to obtain the pooled IRR. The meta-analysis of log IRRs was performed using the Stata user written programme metan.

If publications reported HRs with associated 95% CIs, we obtained standard errors from these CIs as described in chapter 7.7.7.3 of the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011a), and we performed meta-analysis using the generic inverse-variance method (RevMan 2014). When possible, we reported both adjusted and unadjusted HRs, but we primarily used adjusted HRs from multivariable models of studies incorporating similar covariates (Dretzke 2014).

Assessment of heterogeneity

We expected substantial clinical heterogeneity between studies because of geographical/'ethnic' and methodological diversity. We did not intend to address statistical heterogeneity (inconsistency) using the I^2 statistic because this statistic does not indicate how much the effect size varies, which is what people want to know when asking about the implications of heterogeneity (Borenstein 2017a). Also, the I^2 statistic is problematic in the context of prognosis studies because individual studies often have large sample sizes resulting in narrow CIs, which can result in high I^2 values even if inconsistency between studies is moderate (Iorio 2015). Instead, when there were at least three studies, we reported the range of the effects of the random-effects meta-analyses using prediction intervals (Borenstein 2017b; Higgins 2009; IntHout 2016; Riley 2011; Riley 2015). In a random-effects meta-analysis, the prediction interval reflects the whole distribution of effects across study populations, including the effect expected in a future study (IntHout 2016; Riley 2015).

Certainty of the evidence

We created a 'Summary of findings' table using Review Manager 5 (RevMan 2014). We used an adapted version of the GRADE framework for prognostic factor research for describing the influence of IFG, IGT, elevated HbA1c and both IFG and IGT on the development of T2DM (Huguet 2013). We justified all decisions to downgrade the certainty of evidence using footnotes, and we made comments to aid the reader's understanding of this Cochrane Review where necessary.

Sensitivity analysis

We planned to perform sensitivity analyses to explore the influence of the following factors (when applicable) on effect sizes by excluding:

• studies at high or unclear risk of bias;

• very long or large studies to establish the extent to which they dominate the results.

Subgroup analysis

Because we stratified the analyses by IH definition and geographical locations/'ethnicity', which we thought were the main sources of heterogeneity, we did not plan to perform subgroup analyses. However, if at least 10 studies specifying diabetes incidence data were included, we would have investigated age and sex by testing for interactions between subgroups.

If T2DM incidence data were available for children and adolescents, we reported the results separately.

RESULTS

Description of studies

Results of the search

We identified a total of 8354 records through database searching and an additional 259 records from 16 systematic reviews. After excluding duplicates and non-relevant records based on title and abstract screening, we assessed 450 full-text records. Of

these we excluded 213 full-text articles; the remaining 237 articles were reports of 110 studies. Of the 110 studies, 4 were potentially relevant ongoing trials (NCT00786890; NCT02838693; NCT02958579; Vilanova 2017), and 3 are awaiting classification (Li 2001; Misnikova 2011; NCT00816608). Therefore, we included 103 studies. We added 86 new publications after handsearching the full texts of included studies, but these were all secondary publications of the included studies.

The complementary 'similar articles' algorithm search using our set of known publications yielded 263 publications for screening after deduplication. This resulted in 24 new publications after excluding irrelevant articles based on title and abstract screening. We did not identify new studies but found 13 secondary publications of studies we had already included.

Altogether, we included 103 prospective cohort studies (329 publications) in the review. After the initial search in four databases (in December 2016), we observed that 98% of all included publications were indexed in Ovid MEDLINE. Therefore, we decided to restrict the pre-publication update search in February 2018 to Ovid MEDLINE.

For full details of search results see Figure 1.

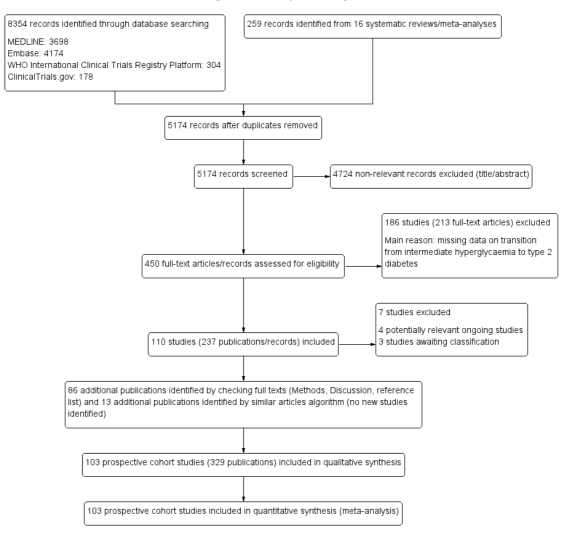


Figure I. Study flow diagram

Included studies

For a detailed description of the characteristics of the included studies, see Characteristics of included studies; Appendix 4; Appendix 5; Appendix 6; Appendix 7; Appendix 8; Appendix 9; Appendix 10; Appendix 11; Appendix 12; Appendix 13; Appendix 14; Appendix 15; Appendix 16; and Appendix 17. The following is a succinct overview.

Source of data

The 103 studies took place in the following regions of the world.

• Australia: 3 studies.

• Latin America: 7 studies (Chile, 1 study; Columbia, 1 study; Mexico, 5 studies (2 studies with primarily Mexican Americans took place in the USA (Garcia 2016; Lorenzo 2003)).

• North America: 12 studies (USA ,12 studies, with 4 studies in particular populations: Pima Indians/Native Americans, 3 studies (Vijayakumar 2017; Wang 2011; Wheelock 2016); and Japanese Americans, 1 study (McNeely 2003)).

• Africa: 1 study (performed in South Africa but with a population consisting of South African Indians (Motala 2003)).

• Middle East: 7 studies (Iran, 5 studies; Israel, 1 study; Jordan, 1 study).

• Asia: 42 studies (China, 11 studies; India, 5 studies; Japan, 8 studies; Korea, 11 studies; Singapore, 2 studies; Taiwan, 2 studies; Thailand, 3 studies).

• Islands: 2 studies (Mauritius, 1 study; Micronesia (Nauru), 1 study).

• Europe: 29 studies (Denmark, 1 study; Finland, 5 studies; France, 3 studies; Germany, 3 studies; Greece, 1 study; Italy, 3 studies; Malta, 1 study; Spain, 3 studies; Sweden, 3 studies; Netherlands, 4 studies; UK, 2 studies). One study in the Netherlands included a mixed population of South-Asian Surinamese participants, African Surinamese participants and "Ethnic Dutch" participants (Admiraal 2014).

Fifty-eight studies contributed most of the data (Appendix 4).

Measurements of overall prognosis of people with IH and of the prognostic factor IH versus normoglycaemia

Of the 103 included studies, 17 evaluated the overall prognosis of people with IH for the development of type 2 diabetes mellitus without a normoglycaemic comparison group. Of these studies, six recruited participants with IFG at baseline (Baena-Diez 2011; Gautier 2010; Lecomte 2007; Leiva 2014; Levitzky 2008; Sharifi 2013), six recruited participants with IGT at baseline (Kleber 2010; Kleber 2011; Ko 1999; Marshall 1994; Rajala 2000; Ramachandran 1986), two recruited a mixed IFG/IGT cohort (Rasmussen 2008; Toshihiro 2008), and three recruited participants with various definitions of IH (Kim 2014; Lee 2016; Song 2016a). In addition, 76 studies with a normoglycaemic comparison group contributed data to evaluate the overall prognosis of people with IH by means of cumulative incidence. Therefore, analysis of overall prognosis is based on 93 studies.

Fifty-two studies assessed the prognostic effect of IH versus normoglycaemia for the development of type 2 diabetes mellitus and provided outcome measures as ratios (hazard ratio (HR), incidence rate ratio (IRR) and/or odds ratio (OR)). Forty-seven studies explicitly defined normoglycaemia, often by a combination of FPG thresholds and two hour post-load glucose thresholds (Anjana 2015; Baena-Diez 2011; Bergman 2016; Chen 2003; Chen 2017; Coronado-Malagon 2009; Den Biggelaar 2016; Derakhshan 2016; Dowse 1991; Forouhi 2007; Guerrero-Romero 2006; Heianza 2012; Janghorbani 2015; Jaruratanasirikul 2016; Kim 2005; Ko 1999; Ko 2001; Larsson 2000; Lecomte 2007; Leiva 2014; Li 2003; Ligthart 2016; Lipska 2013; Liu 2014; Liu 2017; Lyssenko 2005; Magliano 2008; Man 2017; Meigs 2003; Motala 2003; Motta 2010; Mykkänen 1993; Nakanishi 2004; Peterson 2017; Qian 2012; Rajala 2000; Rathmann 2009; Rijkelijkhuizen 2007; Sasaki 1982; Soriguer 2008; Toshihiro 2008; Vaccaro 1999; Valdes 2008; Viswanathan 2007; Wang 2011; Wat 2001; Weiss 2005; Yeboah 2011). In the remaining studies, it was evident that normoglycaemia reflected the population with neither IH nor T2DM at baseline.

IH was commonly defined by the IFG_{5.6} threshold (FPG level 5.6 mmol/L to 6.9 mmol/L or 100 mg/dL to 125 mg/dL), IFG_{6.1} threshold (FPG level 6.1 mmol/L to 6.9 mmol/L or 110 mg/dL to 125 mg/dL), IGT (plasma glucose concentration 7.8 mmol/L to 11.1 mmol/L or 140 mg/dL to 199 mg/dL two hours after a 75 g glucose load on the OGTT), or combinations of these criteria (Appendix 5; Appendix 6). Sixty-six studies used an OGTT at baseline as part of the strategy to assess glycaemic status, and 46 studies used OGTT at baseline and follow-up (Appendix 5).

Twelve studies defined IH by applying the HbA1c_{5.7} threshold (HbA1c 5.7% to 6.4% or 39 mmol/mol to 46 mmol/mol) (Bae 2011; Cederberg 2010; Han 2017; Heianza 2012; Kim 2014; Kim 2016a; Lee 2016; Lipska 2013; Man 2017; Nakagami 2016; Vijayakumar 2017; Warren 2017), and 10 studies used the HbA1c_{6.0} threshold (HbA1c 6.0% to 6.4% or 42 mmol/mol to 46 mmol/mol) (Bae 2011; Bonora 2011; Chamnan 2011; Han 2017; Heianza 2012; Kim 2016a; Nakagami 2016; Sato 2009; Wang 2011; Warren 2017).

Overview of study populations

Sixty-nine studies (67%) started recruitment after 1990 (see Characteristics of included studies), and overall follow-up ranged from 1 year in Bai 1999, Coronado-Malagon 2009 and Kleber 2010 to 24 years in Bergman 2016 (see Characteristics of included studies; Appendix 7).

Depending on the phase of the study, the number of participants differed. The first phase of every study often constituted a large epidemiological investigation of, for example, the importance of various risk factors for cardiovascular health; in total, more than

250,000 participants began the studies (Appendix 8). The number of participants with IH depended on how the studies defined this condition at baseline and the way they measured the development of T2DM.

The overall prognosis of participants with IH at baseline and across all follow-up times (1 to 20 years) was based on the following data (Table 1).

- IFG_{5.6}: 13,692 participants.
- IFG_{6.1}: 9943 participants.
- IGT: 13,728 participants.
- Both IFG and IGT: 2434 participants.
- HbA1c5.7: 9758 participants.
- HbA1c_{6.0}: 2529 participants.

Follow-up time across all measures of IH at baseline had the following number of participants per year of follow-up (in parentheses, number of people with IH who regressed to normoglycaemia); see Table 1.

- 1 year: 878 (375) participants.
- 2 years: 3882 (2852) participants.
- 3 years: 3014 (1356) participants.
- 4 years: 9388 (807) participants.
- 5 years: 16,275 (2603) participants.
- 6 years: 2573 (1328) participants.
- 7 years: 3209 (679) participants.
- 8 years: 3293 (328) participants.
- 9 years: 1588 (299) participants.
- 10 years: 5425 (894) participants.
- 11 years: 1655 (736) participants.
- 12 years: 3574 (no data) participants.
- 15 years: 70 (no data) participants.
- 20 years: 114 (no data) participants.

Data on the prognostic factor IH versus normoglycaemia for the development of T2DM were based on the following number of participants with IH at baseline (Table 2). Data were reported by ratio measures (HR, IRR, OR).

- IFG_{5.6}: 42,694 participants.
- IFG_{6.1}: 12,507 participants.
- IGT: 25,617 participants.
- Both IFG and IGT: 6160 participants.
- HbA1c5.7: 8094 participants.
- HbA1c_{6.0}: 6126 participants.
- Both HbA1c5.7 and IFG5.6: 3761 participants.

The mean age of adult participants at baseline ranged from 30 years to 77 years (Appendix 9). In two studies all the participants were female (De Abreu 2015; Larsson 2000), and in eight studies all the participants were male (Charles 1997; Lecomte 2007; Nakanishi 2004; Park 2006; Sato 2009; Stengard 1992; Toshihiro 2008; Zethelius 2004). The body mass index (BMI) of the participants at baseline ranged from 23.2 kg/m² to 39.1 kg/m². A family history of diabetes was reported in 3% to 100% of the study participants.

At baseline, 60 studies (58%) reported diastolic and systolic blood pressure; 43 studies (22%), smoking status; 66 studies (64%), FPG; 24 studies (23%), HbA1c; 44 studies (43%), two-hour glucose measurements; 7 studies (7%), medications; 26 studies (25%), comorbidities; 20 studies (19%), hypertension; and 5 studies (5%), dyslipidaemia (Appendix 10).

Categorisation of studies

In order to address the complexity of our dataset with regard to factors potentially influencing the definition, detection and development of T2DM, such as genetics, environmental and social conditions, the way risk factors and T2DM incidence were measured, and access to health care (Avilés-Santa 2016; De Rekeneire 2007; Herman 2012; Likhari 2010; Maruthur 2011; Parrinello 2016) - with all of these features interacting to some degree - we choose to provide the reader with a broad overview mainly focusing on geographic regions in the following way.

Groups consisted of participants from studies taking place in Australia, Europe or North America; people from Latin America; individuals from Asia or the Middle East; and American (Pima) Indians and Pacific/Indian Ocean islanders ('American Indians/Islands' group). The logic of grouping participants in the last cohort together resided in the fact that they shared some characteristics relevant to T2DM, including a considerable genetic background risk, historic isolation from outside communities with substantial influence from Western diets, or both (Hanson 2014; Jowett 2009; Nair 2015; Serjeantson 1983).

For 41 studies, we categorised the origin of participants as 'Australia/Europe/North America' (Admiraal 2014; Baena-Diez 2011; Bonora 2011; Cederberg 2010; Chamnan 2011; Charles 1997; Cugati 2007; De Abreu 2015; Den Biggelaar 2016; Filippatos 2016; Forouhi 2007; Gautier 2010; Hanley 2005; Kleber 2010; Kleber 2011; Larsson 2000; Lecomte 2007; Levitzky 2008; Ligthart 2016; Lipska 2013; Lyssenko 2005; Magliano 2008; Marshall 1994; McNeely 2003; Meigs 2003; Motta 2010; Mykkänen 1993; Peterson 2017; Rajala 2000; Rasmussen 2008; Rathmann 2009; Rijkelijkhuizen 2007; Schranz 1989; Soriguer 2008; Stengard 1992; Vaccaro 1999; Valdes 2008; Warren 2017; Weiss 2005; Yeboah 2011; Zethelius 2004).

For seven studies, we categorised the origin of participants as 'Latin America' (Coronado-Malagon 2009; Ferrannini 2009; Garcia 2016; Gomez-Arbelaez 2015; Guerrero-Romero 2006; Leiva 2014; Lorenzo 2003). Although Garcia 2016 and Lorenzo 2003 took place in the USA, they included primarily Mexican Americans, hence the rationale for this categorisation.

We categorised 50 studies as 'Asia/Middle East' (Aekplakorn 2006; Ammari 1998; Anjana 2015; Bae 2011; Bai 1999; Bergman 2016; Chen 2003; Chen 2017; Derakhshan 2016; Han 2017; Heianza 2012; Inoue 1996; Janghorbani 2015; Jaruratanasirikul 2016; Jeong 2010; Jiamjarasrangsi 2008a; Kim 2005; Kim 2008; Kim 2014; Kim 2016a; Ko 1999; Ko 2001; Latifi 2016; Lee 2016; Li 2003; Liu 2008; Liu 2014; Liu 2016; Liu 2017; Man 2017;

Mohan 2008; Motala 2003; Nakagami 2016; Nakanishi 2004; Noda 2010; Park 2006; Qian 2012; Ramachandran 1986; Sadeghi 2015; Sasaki 1982; Sato 2009; Sharifi 2013; Shin 1997; Song 2015; Song 2016a; Toshihiro 2008; Viswanathan 2007; Wang 2007; Wat 2001; Wong 2003). Of these, 37 studies recruited participants from China, Japan, South Korea, Singapore, Taiwan and Thailand (Aekplakorn 2006; Bae 2011; Chen 2003; Chen 2017; Han 2017; Heianza 2012; Inoue 1996; Jaruratanasirikul 2016; Jeong 2010; Jiamjarasrangsi 2008a; Kim 2005; Kim 2008; Kim 2014; Kim 2016a; Ko 1999; Ko 2001; Lee 2016; Li 2003; Liu 2008; Liu 2014; Liu 2016; Liu 2017; Man 2017; Nakagami 2016; Nakanishi 2004; Noda 2010; Park 2006; Qian 2012; Sasaki 1982; Sato 2009; Shin 1997; Song 2015; Song 2016a; Toshihiro 2008; Wang 2007; Wat 2001; Wong 2003), 5 studies recruited participants from India (Anjana 2015; Bai 1999; Mohan 2008; Ramachandran 1986; Viswanathan 2007), 1 study involved Indian-South African participants (Motala 2003), and 7 studies recruited participants from Iran, Israel and Jordan (Ammari 1998; Bergman 2016; Derakhshan 2016; Janghorbani 2015; Latifi 2016; Sadeghi 2015; Sharifi 2013).

We categorised the origin of participants as 'American Indians/ Islands' in five studies. Three of the five studies had American Indians as participants (Vijayakumar 2017; Wang 2011; Wheelock 2016), one included Mauritians (Söderberg 2004), and the remaining study included Nauruans (Dowse 1991).

Six studies included black participants (Admiraal 2014; Bergman 2016; Hanley 2005; Söderberg 2004; Warren 2017; Yeboah 2011), representing 25% to 47% of all participants in these studies.

Six studies included children, adolescents or both as participants (Jaruratanasirikul 2016; Kleber 2010; Kleber 2011; Vijayakumar 2017; Weiss 2005; Wheelock 2016).

Measurement of the development of T2DM

Almost all studies combined criteria to define incident T2DM, using indicators such as FPG of 7.0 mmol/L or more, two-hour postload glucose level of 11.1 mmol/L or more, HbA1c of 6.5% or more, receipt of antidiabetic medication, physician diagnosis or self-report.

Of the 103 included studies, 64 included FPG of 7.0 mmol/L or more, and 52, two-hour postload glucose level of 11.1 mmol/L or more, in their definition of incident T2DM. Eighteen studies used HbA1c as part of the definition of T2DM, typically an HbA1c level of 6.5% or more. One study defined T2DM incidence based only on an HbA1c level of 6.5% or more (Lee 2016). In 34 studies, antidiabetic treatment comprised part of the definition of T2DM, and in 15 studies physician diagnosis or self-report was part of the T2DM incidence definition.

Risk of bias in included studies

For details on the QUIPS tool and the risk of bias of the included studies see Appendix 3 and Characteristics of included studies. The results are summarised below separately for studies that provided data on overall prognosis for people with IH and on IH versus normoglycaemia as a prognostic factor.

a) Overall prognosis of people with IH for the development of T2DM and b) regression from IH to normoglycaemia

There were 93 studies providing data on cumulative incidence. Figure 2 summarises the risk of bias results across all studies, while the results for each study are shown in Figure 3 and Figure 4 (split into two figures because of the large number of studies). We evaluated the first four risk of bias domains (i.e. study participation, study attrition, glycaemic status measurement, outcome measurement) of the QUIPS tool.

Figure 2. Risk of bias graph for studies of overall prognosis of people with intermediate hyperglycaemia for developing type 2 diabetes: review authors' judgements about each risk of bias item presented as percentages across all included studies

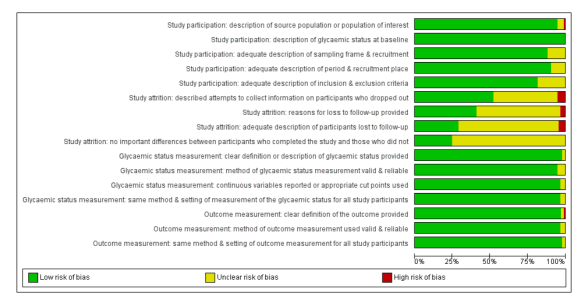


Figure 3. 'Risk of bias' summary for studies of overall prognosis in people with intermediate hyperglycaemia for developing type 2 diabetes: review authors' judgements about each risk of bias item for each included study (part 1). The summary was split into part 1 (Figure 3) and part 2 (Figure 4) for better legibility

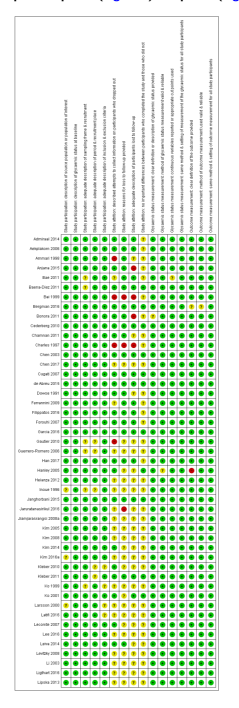
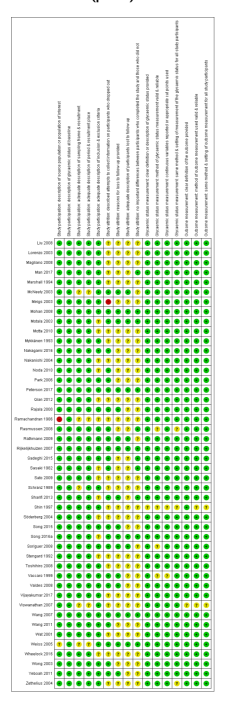


Figure 4. Risk of bias summary for studies of overall prognosis of people with intermediate hyperglycaemia for developing type 2 diabetes: review authors' judgements about each risk of bias item for each included study (part 2)



Study participation

Study authors described the five items in this domain sufficiently in most (65 studies; 70%) included studies. Eleven studies did not adequately characterise the sampling frame and/or recruitment procedures (Bae 2011; Baena-Diez 2011; Gautier 2010; Guerrero-Romero 2006; Inoue 1996; Ko 1999; McNeely 2003; Ramachandran 1986; Schranz 1989; Viswanathan 2007; Weiss 2005). One study was at high risk of bias for the item 'description of the source population or population of interest' (Ramachandran 1986).

Study attrition

Forty-eight studies attempted to collect information on participants who were lost to follow-up, while 40 studies were at unclear risk of bias and five studies were at high risk of bias (Ammari 1998; Bai 1999; Charles 1997; Gautier 2010; Meigs 2003).

In most (61 studies; 66%) of the studies we could not identify the reasons for loss to follow-up or adequate descriptions of these participants. Five studies were at high risk of bias for one or both of the items (Anjana 2015; Bai 1999; Bonora 2011; Charles 1997; Jaruratanasirikul 2016).

Only 23 studies (25%) provided information on potentially im-

portant differences between participants who completed the studies and those who did not.

Glycaemic status measurement

Study authors described these items sufficiently in 85 studies (91%). One study did not describe three of the four items ('clear definition of the outcome provided', 'adequately valid and reliable method of measurement', and 'continuous variables reported or appropriate cut points used') in enough detail (Shin 1997).

Outcome measurement

Study authors described the three items sufficiently in 89 studies (96%). One study was at high risk of bias for the item 'provision of clear definition of the outcome' (Hanley 2005).

c) Development of T2DM in people with IH as compared to people with normoglycaemia

There were 52 studies comparing IH with normoglycaemia as a prognostic factor for T2DM. Figure 5 shows the results for the six domains summarised across studies, and the result for each study is shown in Figure 6.

Figure 5. Risk of bias graph for studies of intermediate hyperglycaemia versus normoglycaemia as a prognostic factor for developing type 2 diabetes: review authors' judgements about each risk of bias item presented as percentages across all included studies

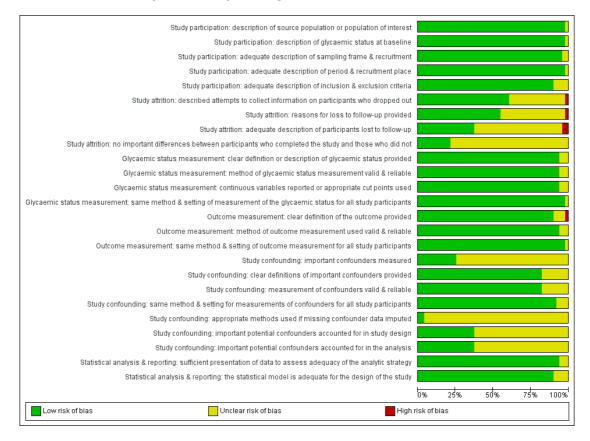
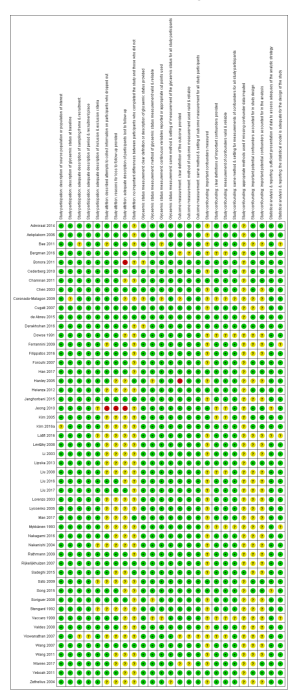


Figure 6. Risk of bias summary for studies of intermediate hyperglycaemia versus normoglycaemia as a prognostic factor for developing type 2 diabetes: review authors' judgements about each risk of bias item for each included study



Fourteen studies provided data on multivariable HRs of T2DM incidence, adjusted for 2 to 13 covariates (Bae 2011; Bonora 2011; Forouhi 2007; Han 2017; Heianza 2012; Janghorbani 2015; Kim 2005; Li 2003; Liu 2016; Lyssenko 2005; Nakagami 2016; Wang 2011; Warren 2017; Yeboah 2011). Whenever possible, we used the reported model with the greatest number of covariates.

Study participation

Study authors described the items of this domain sufficiently in most (42 studies; 82%) of the included studies. Two studies did not adequately characterise the sampling frame and/or recruitment procedures (Bae 2011; Viswanathan 2007).

Study attrition

Study authors usually described these items sufficiently and attempted to collect information on participants who were lost to follow-up. However, in most (32 studies; 63%) of the included studies we could not identify the reasons for losses to follow-up or adequate descriptions of these participants. Only 10 studies (20%) provided information on potentially important differences between participants who completed the studies and those who did not. Two studies were at high risk of bias on one of the four items (Bonora 2011; Jeong 2010).

Glycaemic status measurement

Study authors described the items sufficiently in 40 (78%) studies.

Outcome measurement

Study authors described these items sufficiently in 46 studies (90%). One study had a high risk of bias for the item 'clear definition of the outcome provided' (Hanley 2005).

Study confounding

Only one study described all items sufficiently (Derakhshan 2016). It was difficult to judge study confounding because the number of important covariates measured was limited. If studies analysed data by means of multivariable regression models, they often adjusted these analyses taking into account several covariates: age (43 out of 52 studies), anthropometric measures such as BMI (33 out of 52 studies), sex (31 out of 52 studies), family history of diabetes (24 out of 52 studies), smoking status (24 out of 52 studies), blood pressure/hypertension (19 out of 52 studies), triglycerides (18 out of 52 studies), cholesterol (17 out of 52 studies), physical activity (14 out of 52 studies), drinking status (12 out of 52 studies), socioeconomic status (8 out of 52 studies), 'ethnicity' (5 out 52 studies), medications (3 out of 52 studies) and renal function (1 study); for details see Appendix 16 and Appendix 17.

Twenty studies (39%) adjusted their analyses for age, sex and anthropometric measures (e.g. BMI or waist circumference) (Admiraal 2014; Bergman 2016; Bonora 2011; Chamnan 2011; Chen 2003; Derakhshan 2016; Forouhi 2007; Han 2017; Heianza 2012; Janghorbani 2015; Kim 2005; Kim 2016a; Li 2003; Man 2017; Sadeghi 2015; Soriguer 2008; Valdes 2008; Wang 2011; Warren 2017; Yeboah 2011). Six studies (12%) adjusted for age, sex, anthropometric measures and physical activity (Bonora 2011; Derakhshan 2016; Forouhi 2007; Han 2017; Kim 2016a; Yeboah 2011), and five studies (10%) also included smoking status (Bonora 2011; Derakhshan 2016; Forouhi 2007; Han 2017; Kim 2016a). When used, covariates were usually clearly defined and measured. However, only two studies reported an imputation method for missing confounders (Derakhshan 2016; Sadeghi 2015).

Statistical analysis and reporting

Study authors addressed this domain sufficiently in 44 studies (86%).

Development of T2DM in people with IH

In the following we report the results of the analyses for the overall prognosis of people with IH as well as regression from IH to normoglycaemia, and the effects of glycaemic status (IH versus normoglycaemia) as a prognostic factor for T2DM.

Definition of IH at baseline

Studies defined IH as follows.

• IFG_{5.6} threshold, usually defined as a fasting plasma glucose level of 5.6 mmol/L to 6.9 mmol/L.

• IFG_{6.1} threshold, usually defined as a fasting plasma glucose level of 6.1 mmol/L to 6.9 mmol/L.

- IGT, usually defined as a plasma glucose level of 7.8 mmol/ L to 11.1 mmol/L two hours after a 75 g OGTT.
- Isolated IFG was defined as IFG_{5.6} or IFG_{6.1} alone, without IGT, and isolated IGT was defined as IGT alone, without IFG_{5.6} or IFG_{6.1}.
- HbA1c_{5.7} threshold, usually defined as HbA1c measurement of 5.7% to 6.4%.
 - HbA1c_{6.0} threshold, usually defined as HbA1c

measurement of 6.0% to 6.4%.

Depending on how investigators measured IH, the following number of study cohorts provided information on T2DM incidence associated with glycaemic status at baseline (one study might have investigated several associations between glycaemic status and T2DM incidence within the same study, for example, one cohort

with $IFG_{5.6}$, another cohort with $IFG_{6.1}$ and a third cohort with IGT).

- IFG_{5.6}/isolated IFG_{5.6}: 27/10 study cohorts.
- IFG_{6.1}/isolated IFG_{6.1}: 22/9 study cohorts.
- IGT/isolated IGT: 39/18 study cohorts.
- Combined IFG and IGT: 15 study cohorts.
- HbA1c_{5.7}: 7 study cohorts.
- HbA1c_{6.0}: 10 study cohorts.
- Combined HbA1c5.7 and IFG5.6: 3 study cohorts.

a) Overall prognosis of people with IH for developing T2DM

Irrespective of the definition of IH at baseline, the cumulative incidence of T2DM seemed to increase with length of followup, though there was no obvious linear trend. There was no clear pattern of differences between geographic regions.

IH defined by IFG_{5.6} mmol/L threshold

Diabetes incidence associated with $IFG_{5.6}$ at baseline and followup periods from 2 to 12 years showed pooled cumulative incidences of 2% to 38% (Figure 7; Figure 8).

Figure 7. Impaired fasting glucose 5.6 mmol/L (IFG_{5.6}) threshold: association with cumulative type 2 diabetes mellitus (T2DM) incidence over 2-5 years *Isolated IFG_{5.6}

CI: confidence interval; M: men; n/N: events/number of participants; W: women

| Study, location | Cumulative incidence (95% CI) % Weight n/N |
|---|---|
| 2 years Kim 2008, Asia/Middle East | 0.02 (0.01, 0.02) 100.00 22/1335 |
| 3 years Jiamjarasrangsi 2008a, Asia/Middle East Chen 2017*, Asia/Middle East Rasmussen 2008*, Australia/Europe/North America Subtotal (Tau² = 0.09) | 0.10 (0.07, 0.14) 33.23 33/320 0.12 (0.09, 0.16) 33.25 40/329 0.32 (0.28, 0.36) 33.52 141/442 0.17 (0.06, 0.32) 100.00 |
| 4 years Park 2006, Asia/Middle East Kim 2014*, Asia/Middle East Song 2015_M*, Asia/Middle East Song 2015_W*, Asia/Middle East Subtotal (Tau² = 0.01) | 0.12 (0.09, 0.17) 28.85 40/321 0.15 (0.10, 0.22) 23.64 24/158 0.19 (0.14, 0.26) 23.42 30/154 0.23 (0.17, 0.30) 24.10 38/167 0.17 (0.13, 0.22) 100.00 |
| 5 years Noda 2010_M, Asia/Middle East Noda 2010_W, Asia/Middle East Liu 2008, Asia/Middle East Heianza 2012, Asia/Middle East Nakagani 2016, Asia/Middle East Latifi 2016, Asia/Middle East Wang 2007*, Asia/Middle East Vijayakumar 2017, Australia/Europe/North America Subtotal (Tau² = 0.01) | 0.09 (0.06, 0.14) 12.40 18/202 0.09 (0.06, 0.14) 12.40 19/203 0.11 (0.07, 0.16) 12.29 18/169 0.16 (0.14, 0.17) 12.91 262/1680 0.16 (0.13, 0.20) 12.72 77/467 0.17 (0.11, 0.25) 12.06 21/124 0.20 (0.16, 0.26) 12.53 53/261 0.52 (0.48, 0.57) 12.70 222/424 0.18 (0.10, 0.27) 100.00 |
| 0 | I I I 0.2 0.4 0.6 mulative incidence |

T2DM cumulative incidence associated with IFG 5.6 mmol/L threshold: 2 to 5 years follow-up

Figure 8. Impaired fasting glucose 5.6 mmol/L (IFG_{5.6}) threshold: association with cumulative type 2 diabetes mellitus (T2DM) incidence over 6-12 years

*Isolated IFG_{5.6}

**'Africa': African Surinamese cohort, 'Asia': Asian Surinamese cohort, 'Australia/Europe/North America':

'ethnic Dutch' cohort.

CI: confidence interval; M: men; n/N: events/number of participants; W: women

Cumulative incidence (95% CI) % Weight n/N Study, location 6 years Valdes 2008°, Australia/Europe/North America Rijkelijkhuizen 2007, Australia/Europe/North America McNeely 2003, Australia/Europe/North America 0.12 (0.07, 0.20) 0.21 (0.17, 0.25) 0.22 (0.15, 0.30) 14/114 24.78 29.68 101/488 25.26 27/125 Soriguer 2008*, Australia/Europe/North America Subtotal (Tau² = 0.03) 0.41 (0.29, 0.54) 0.22 (0.15, 0.31) 20.29 23/56 100.00 7 years Janghorbani 2015*, Asia/Middle East 0.10 (0.07, 0.15) 20.49 23/230 0.20 (0.13, 0.27) 0.36 (0.31, 0.41) 0.11 (0.07, 0.16) Sharifi 2013, Asia/Middle East Sadeghi 2015*, Asia/Middle East 19.79 24/123 134/373 20.82 Lipska 2013, Australia/Europe/North America Ferrannini 2009*, Latin America Subtotal (Tau² = 0.10) 20.31 20/189 0.17 (0.10, 0.28) 18.58 11/65 0.18 (0.08, 0.30) 100.00 8 years Wang 2011_M, American Indians/Islands Wang 2011_W, American Indians/Islands Yeboah 2011, Australia/Europe/North America 0.33 (0.28, 0.37) 31.86 137/418 0.39 (0.35, 0.44) 33.02 208/529 35.13 0.29 (0.26, 0.32) 273/940 Subtotal (Tau² = 0.01) 0.34 (0.27, 0.40) 100.00 9 years Anjana 2015*, Asia/Middle East Gautier 2010, Australia/Europe/North America 0.48 (0.36, 0.60) 0.15 (0.12, 0.17) 32.54 32/67 33.84 142/979 Garcia 2016, Australia/Europe/North America Subtotal (Tau² = 0.34) 0.55 (0.49, 0.60) 0.38 (0.10, 0.70) 33.62 169/310 100.00 10 years Admiraal 2014**, Africa Admiraal 2014**, Asia/Middle East Admiraal 2014**, Australia/Europe/North America Forouhi 2007, Australia/Europe/North America 0.35 (0.22, 0.50) 0.42 (0.26, 0.59) 11.27 14/40 13/31 10.67 0.08 (0.03, 0.20) 3/40 53/633 11.27 0.08 (0.06, 0.11) 13.78 de Abreu 2015, Australia/Europe/North America 0.11 (0.07, 0.17) 13.30 21/187 Filippatos 2016, Australia/Europe/North America Cugati 2007, Australia/Europe/North America 0.25 (0.21, 0.31) 0.30 (0.25, 0.36) 13.52 13.42 71/279 69/229 0.38 (0.29, 0.48) 0.23 (0.14, 0.33) McNeely 2003, Australia/Europe/North America 12 78 39/103 Subtotal (Tau² = 0.10) 100.00 12 years Aekplakorn 2006, Asia/Middle East Han 2017*, Asia/Middle East 0.29 (0.24, 0.35) 0.41 (0.34, 0.48) 43.53 65/223 42.91 81/199 Vaccaro 1999*, Australia/Europe/North America Subtotal (Tau² = 0.03) 0.09 (0.02, 0.38) 13.57 1/11 100.00 0.31 (0.19, 0.43) 0 0.2 0.4 0.6 0.8 Cumulative incidence

T2DM cumulative incidence associated with IFG 5.6 mmol/L threshold: 6 to 12 years follow-up

The number of studies and participants, and the cumulative incidence of T2DM (pooled if more than one study) according to follow-up period were as follows.

• 2 years' follow-up: 1 study, 1335 participants, cumulative incidence 2% (95% confidence interval (CI) 1 to 2).

• 3 years' follow-up: 3 studies, 1091 participants, cumulative incidence 17% (95% CI 6 to 32).

• 4 years' follow-up: 3 studies, 800 participants, cumulative incidence 17% (95% CI 13 to 22).

• 5 years' follow-up: 7 studies, 3530 participants, cumulative incidence 18% (95% CI 10 to 27).

• 6 years' follow-up: 4 studies, 783 participants, cumulative incidence 22% (95% CI 15 to 31).

• 7 years' follow-up: 5 studies, 980 participants, cumulative incidence 18% (95% CI 8 to 30).

• 8 years' follow-up: 2 studies, 1887 participants, cumulative incidence 34% (95% CI 27 to 40).

• 9 years' follow-up: 3 studies, 1356 participants, cumulative incidence 38% (95% CI 10 to 70).

• 10 years' follow-up: 6 studies, 1542 participants, cumulative incidence 23% (95% CI 14 to 33).

• 12 years' follow-up: 3 studies, 433 participants, cumulative incidence 31% (95% CI 19 to 34).

IH defined by IFG_{6.1} mmol/L threshold

Diabetes incidence, as associated with $IFG_{6.1}$ at baseline and a follow-up period of 2 to 15 years, showed pooled cumulative incidences of 9% to 48% (Figure 9; Figure 10).

Figure 9. Impaired fasting glucose 6.1 mmol/L (IFG_{6.1}) threshold: association with cumulative type 2 diabetes mellitus (T2DM) incidence over 2-5 years *Isolated IFG_{6.1}

CI: confidence interval; M: men; n/N: events/number of participants; W: women

.

| Study, location | Cumulative incidence (95% CI) % Weight n/N |
|--|--|
| 2 years Kim 2008, Asia/Middle East Ko 2001, Asia/Middle East Subtotal (Tau² = 0) ♢ | • 0.10 (0.07, 0.13) 89.91 48/49 0.25 (0.16, 0.38) 10.09 14/55 0.11 (0.08, 0.14) 100.00 |
| 3 years Chen 2003, Asia/Middle East Charles 1997*, Australia/Europe/North America Motta 2010, Australia/Europe/North America Subtotal (Tau ² = 0.08) | 0.10 (0.06, 0.15)32.3315/150.03 (0.02, 0.05)34.1115/470.17 (0.13, 0.22)33.5650/290.09 (0.02, 0.20)100.00 |
| 4 years Sato 2009, Asia/Middle East Levitzky 2008_M, Australia/Europe/North America Levitzky 2008_W, Australia/Europe/North America Subtotal (Tau ² = 0.07) | → 0.42 (0.39, 0.46) 33.76 334/7 → 0.20 (0.17, 0.24) 33.35 92/46 → 0.28 (0.23, 0.33) 32.89 87/31 → 0.30 (0.17, 0.44) 100.00 |
| 5 years Söderberg 2004, American Indians/Islands Kim 2005, Asia/Middle East Kim 2016a, Asia/Middle East Wang 2007, Asia/Middle East Noda 2010_M, Asia/Middle East Qian 2012*, Asia/Middle East Nakagami 2016, Asia/Middle East Li 2003*, Asia/Middle East Heianza 2012, Asia/Middle East Noda 2010_W, Asia/Middle East Li 2003*, Asia/Middle East Lecomte 2007, Australia/Europe/North America Lecomte 2007, Australia/Europe/North America Subtotal (Tau² = 0.07) | 0.22 (0.16, 0.29) 8.48 32/14 0.05 (0.03, 0.09) 8.84 15/27 0.25 (0.23, 0.27) 9.21 357/1 0.25 (0.18, 0.34) 8.25 28/11 0.32 (0.22, 0.43) 7.88 25/79 0.37 (0.25, 0.51) 7.12 17/46 0.37 (0.30, 0.46) 8.41 50/13 0.38 (0.25, 0.53) 6.96 16/42 0.41 (0.36, 0.46) 8.96 155/3 0.45 (0.34, 0.56) 7.80 33/74 0.12 (0.09, 0.16) 8.95 44/37 0.17 (0.15, 0.20) 9.12 127/7 0.26 (0.19, 0.33) 100.00 |
| | |

Figure 10. Impaired fasting glucose 6.1 mmol/L (IFG_{6.1}) threshold: association with cumulative type 2 diabetes mellitus (T2DM) incidence over 6-15 years *Isolated IFG_{6.1}

CI: confidence interval; n/N: events/number of participants

| Study, location | Cumulative incidence (95% CI) % Weight n/N |
|---|---|
| 6 years McNeely 2003, Australia/Europe/North America Meigs 2003*, Australia/Europe/North America Valdes 2008, Australia/Europe/North America Rijkelijkhuizen 2007, Australia/Europe/North America Leiva 2014, Latin America Subtotal (Tau ² = 0) | 0.23 (0.12, 0.41) 11.23 7/30 0.30 (0.15, 0.52) 7.59 6/20 0.37 (0.25, 0.50) 19.09 19/52 0.42 (0.34, 0.50) 51.59 62/149 0.39 (0.24, 0.58) 10.50 11/28 0.37 (0.31, 0.43) 100.00 |
| 7 years Nakanishi 2004, Asia/Middle East Rathmann 2009, Australia/Europe/North America Lipska 2013, Australia/Europe/North America Ferrannini 2009*, Latin America Subtotal (Tau ² = 0.41) | 0.02 (0.01, 0.05) 25.97 5/246 0.17 (0.10, 0.27) 25.37 12/71 0.48 (0.38, 0.58) 25.61 48/100 0.06 (0.01, 0.27) 23.05 1/17 0.15 (0.00, 0.45) 100.00 |
| 8 years Lorenzo 2003, Latin America | • 0.48 (0.31, 0.66) 100.00 14/29 |
| 10 years Larsson 2000*, Australia/Europe/North America Forouhi 2007, Australia/Europe/North America Baena-Diez 2011, Australia/Europe/North America Bonora 2011, Australia/Europe/North America Cederberg 2010, Australia/Europe/North America McNeely 2003, Australia/Europe/North America Subtotal (Tau ² = 0.12) | 0.12 (0.05, 0.25) 16.03 5/42 0.13 (0.10, 0.18) 18.66 34/257 0.29 (0.21, 0.38) 17.94 33/115 0.33 (0.22, 0.46) 16.69 18/55 0.38 (0.24, 0.53) 15.89 15/40 0.64 (0.46, 0.79) 14.80 18/28 0.29 (0.17, 0.43) 100.00 |
| 11 years Söderberg 2004*, American Indians/Islands | 0.38 (0.33, 0.43) 100.00 153/402 |
| 15 years Ligthart 2016, Australia/Europe/North America | • 0.31 (0.28, 0.33) 100.00 425/1382 |
| 0 0.2 | 0.4 0.6 0.8 |
| | ulative incidence |

The number of studies and participants, and the cumulative incidence of T2DM (pooled if more than one study) according to follow-up period were as follows.

• 2 years' follow-up: 2 studies, 549 participants, cumulative incidence 11% (95% CI 8 to 14).

• 3 years' follow-up: 3 studies, 927 participants, cumulative incidence 9% (95% CI 2 to 20).

• 4 years' follow-up: 2 studies, 1567 participants, cumulative incidence 30% (95% CI 17 to 44).

• 5 years' follow-up: 11 studies, 3837 participants,

cumulative incidence 26% (95% CI 19 to 33).

• 6 years' follow-up: 5 studies, 279 participants, cumulative incidence 37% (95% CI 31 to 43).

• 7 years' follow-up: 4 studies, 434 participants, cumulative incidence 15% (95% CI 0 to 45).

• 8 years' follow-up: 1 study, 29 participants, cumulative incidence 48% (95% CI 31 to 66).

• 10 years' follow-up: 6 studies, 537 participants, cumulative incidence 29% (95% CI 17 to 43).

• 11 years' follow-up: 1 study, 402 participants, cumulative

incidence 38% (95% CI 33 to 43).
15 years' follow-up: 1 study, 1382 participants, cumulative incidence 31% (95% CI 28 to 33).

IH defined by IGT

Diabetes incidence associated with IGT at baseline showed pooled cumulative incidences of 13% to 60% after a follow-up period of 1 to 20 years (Figure 11; Figure 12).

Figure 11. Impaired glucose tolerance (IGT): association with cumulative type 2 diabetes mellitus (T2DM) incidence over 1-5 years *Isolated IGT

CI: confidence interval; n/N: events/number of participants

| 0.06 (0.03, 0.09) 0.24 (0.17, 0.32) 0.12 (0.09, 0.16) 0.13 (0.05, 0.23) | 33.99 14/252 31.66 29/123 34.35 35/296 100.00 |
|---|--|
| $\begin{array}{c} 0.10 \; (0.07, 0.13) \\ 0.13 \; (0.09, 0.19) \\ 0.15 \; (0.08, 0.25) \\ 0.18 \; (0.12, 0.25) \\ 0.08 \; (0.05, 0.12) \\ 0.08 \; (0.05, 0.11) \\ 0.16 \; (0.11, 0.24) \\ 0.28 \; (0.22, 0.36) \\ 0.41 \; (0.36, 0.46) \\ 0.16 \; (0.09, 0.26) \end{array}$ | 11.40 31/322 11.08 20/153 10.39 10/88 10.98 23/131 11.18 14/183 11.46 32/418 10.94 20/123 11.10 45/158 11.48 181/44 100.00 |
| 0.14 (0.06, 0.28) 0.23 (0.18, 0.30) 0.23 (0.18, 0.30) 0.22 (0.18, 0.27) | 8.96 5/37 46.00 45/192 45.04 44/188 100.00 |
| $\begin{array}{c} 0.37 \ (0.33, 0.41) \\ 0.18 \ (0.11, 0.30) \\ 0.10 \ (0.06, 0.15) \\ 0.22 \ (0.14, 0.33) \\ 0.24 \ (0.18, 0.30) \\ 0.22 \ (0.12, 0.34) \end{array}$ | 21.42 198/53 18.56 12/65 20.48 17/170 18.84 16/72 20.70 48/203 100.00 |
| 0.21 (0.18, 0.25) 0.28 (0.21, 0.37) 0.36 (0.28, 0.46) 0.41 (0.32, 0.50) 0.64 (0.59, 0.69) 0.72 (0.66, 0.78) 0.89 (0.83, 0.93) 0.07 (0.05, 0.11) 0.16 (0.14, 0.19) 0.19 (0.14, 0.25) 0.32 (0.27, 0.38) 0.56 (0.51, 0.62) 0.39 (0.25, 0.53) | 7.79 103/4 7.62 33/118 7.60 39/107 7.62 49/12 7.78 251/38 7.73 165/22 7.66 126/14 7.73 17/23 7.81 122/75 7.69 32/17 7.75 88/27 7.77 196/3 7.46 20/67 100.00 |
| | - • • • • • • • • • • • • • • • • • • • |

T2DM cumulative incidence associated with IGT threshold: 1 to 5 years follow-up

Figure 12. Impaired glucose tolerance (IGT): association with cumulative type 2 diabetes mellitus (T2DM) incidence over 6-20 years *Isolated IGT

CI: confidence interval; M: men; n/N: events/number of participants; W: women

T2DM cumulative incidence associated with IGT threshold: 6 to 20 years follow-up

| Study, location | Cumulative incidence (95% CI) % Weig | jht n/N | | |
|--|---|------------------------------------|--|--|
| 6 years Dowse 1991, American Indians/Islands Valdes 2008, Australia/Europe/North America Soriguer 2008, Australia/Europe/North America Schranz 1989, Australia/Europe/North America Rijkelijkhuizen 2007, Australia/Europe/North America Subtotal (Tau ² =0.01) | 0.25 (0.16, 0.39) 8.93 0.24 (0.16, 0.34) 13.17 0.25 (0.19, 0.32) 19.77 0.26 (0.16, 0.39) 9.32 0.31 (0.21, 0.42) 11.82 0.32 (0.24, 0.42) 15.25 0.37 (0.31, 0.44) 21.73 0.29 (0.25, 0.34) 100.00 | 14/54 23/75 36/111 81/218 | | |
| 7 years Sadeghi 2015*, Asia/Middle East Janghorbani 2015*, Asia/Middle East Sasaki 1982, Asia/Middle East Rathmann 2009*, Australia/Europe/North America Ferrannini 2009*, Latin America Subtotal (Tau ² = 0.02) | 0.13 (0.10, 0.17) 26.15 0.17 (0.12, 0.24) 22.51 0.38 (0.18, 0.64) 6.68 0.28 (0.21, 0.37) 21.27 0.17 (0.12, 0.24) 23.39 0.19 (0.13, 0.26) 100.00 | | | |
| 8 years Wang 2011_W, American Indians/Islands Wang 2011_M, American Indians/Islands Wong 2003, Asia/Middle East Mohan 2008, Asia/Middle East Lorenzo 2003, Latin America Subtotal (Tau ² = 0.01) | 0.47 (0.42, 0.52) 25.09 0.49 (0.41, 0.57) 19.84 0.35 (0.30, 0.41) 24.24 0.41 (0.26, 0.57) 8.99 0.44 (0.37, 0.50) 21.84 0.43 (0.37, 0.49) 100.00 | | | |
| 9 years Anjana 2015*, Asia/Middle East - | • 0.53 (0.45, 0.60) 100.00 | 86/163 | | |
| 10 years Motala 2003, Asia/Middle East Larsson 2000*, Australia/Europe/North America Bonora 2011*, Australia/Europe/North America Peterson 2017, Australia/Europe/North America Cederberg 2010*, Australia/Europe/North America McNeely 2003, Australia/Europe/North America Subtotal (Tau ² = 0.06) | 0.37 (0.23, 0.54) 14.74 0.12 (0.06, 0.22) 17.19 0.15 (0.08, 0.27) 16.43 0.21 (0.10, 0.38) 13.87 0.37 (0.28, 0.47) 18.45 0.38 (0.30, 0.45) 19.32 0.26 (0.17, 0.37) 100.00 | 8/53 6/29 38/103 59/157 | | |
| 11 years Söderberg 2004, American Indians/Islands | 0.46 (0.43, 0.49) 100.00 | 575/1253 | | |
| 12 years Han 2017*, Asia/Middle East Vaccaro 1999*, Australia/Europe/North America Subtotal (Tsu ² = 0) | 0.41 (0.39, 0.44) 97.39 0.32 (0.20, 0.48) 2.61 0.41 (0.38, 0.43) 100.00 | 13/40 | | |
| 20 years Bergman 2016, Asia/Middle East | 0.60 (0.50, 0.68) 100.00 | 68/114 | | |
| I I I 0 0.2 0.4 | I I 0.6 0.8 | | | |
| Cumulative incidence | | | | |

The number of studies and participants, and the cumulative incidence of T2DM (pooled if more than one study) according to follow-up period were as follows.

• 1 year's follow-up: 3 studies, 671 participants, cumulative incidence 13% (95% CI 5 to 23).

• 2 years' follow-up: 9 studies, 1998 participants, cumulative incidence 16% (95% CI 9 to 26).

• 3 years' follow-up: 3 studies, 417 participants, cumulative incidence 22% (95% CI 18 to 27).

• 4 years' follow-up: 5 studies, 1042 participants, cumulative incidence 22% (95% CI 12 to 34).

• 5 years' follow-up: 12 studies, 3444 participants, cumulative incidence 39% (95% CI 25 to 53).

• 6 years' follow-up: 7 studies, 775 participants, cumulative incidence 29% (95% CI 25 to 34).

• 7 years' follow-up: 5 studies, 835 participants, cumulative incidence 19% (95% CI 13 to 26).

• 8 years' follow-up: 4 studies, 1021 participants, cumulative incidence 43% (95% CI 37 to 49).

• 9 years' follow-up: 1 study, 163 participants, cumulative

incidence 53% (95% CI 45 to 60).

• 10 years' follow-up: 6 studies, 443 participants, cumulative incidence 26% (95% CI 17 to 37).

• 11 years' follow-up: 1 study, 1253 participants, cumulative incidence 46% (95% CI 43 to 49).

• 12 years' follow-up: 2 studies, 1552 participants,

cumulative incidence 41% (95% CI 38 to 43).

• 20 years' follow-up: 1 study, 114 participants, cumulative incidence 60% (95% CI 50 to 68).

IH defined by combined IFG and IGT

Diabetes incidence associated with the combination of both IFG and IGT at baseline showed pooled cumulative incidences of 29% to 84% at 1 to 12 years (Figure 13).

Figure 13. Combined impaired glucose tolerance (IGT) and impaired fasting glucose (IFG): association with cumulative type 2 diabetes mellitus (T2DM) incidence over 1-12 years CI: confidence interval; M: men; n/N: events/number of participants; W: women

| tudy, location Cumulative incidence (95% CI) % Weightn/N | | |
|--|--|--|
| 1 year Rasmussen 2008, Australia/Europe/North America — | 0.29 (0.23, 0.36) 100.00 60/207 | |
| 3 years Chen 2017, Asia/Middle East —●— | 0.34 (0.28, 0.41) 100.00 71/205 | |
| 5 years Söderberg 2004, American Indians/Islands Li 2003, Asia/Middle East Oian 2012, Asia/Middle East Vijayakumar 2017, Australia/Europe/North America Subtotal (Tau ² = 0.08) | − 0.38 (0.30, 0.47) 21.29 45/118 0.41 (0.28, 0.55) 18.68 20/49 ● 0.50 (0.40, 0.59) 21.11 54/100 ● 0.52 (0.35, 0.67) 16.97 17/33 ● 0.69 (0.61, 0.75) 21.95 116/16 ● 0.50 (0.37, 0.63) 100.00 100.00 | |
| 6 years Soriguer 2008, Australia/Europe/North America — Meigs 2003, Australia/Europe/North America — Valdes 2008, Australia/Europe/North America — Rijkelijkhuizen 2007, Australia/Europe/North America Subtotal (Tau ² = 0) | 0.50 (0.33, 0.67) 26.39 14/28 0.56 (0.37, 0.72) 25.46 15/27 0.60 (0.39, 0.78) 18.98 12/20 0.65 (0.47, 0.79) 29.17 20/31 0.58 (0.48, 0.67) 100.00 | |
| 7 years Sadeghi 2015, Asia/Middle East Kim 2014, Asia/Middle East Janghorbani 2015, Asia/Middle East Rathmann 2009, Australia/Europe/North America Subtotal (Tau ² = 0.07) | 0.17 (0.14, 0.22) 26.98 65/373 0.32 (0.24, 0.41) 25.07 38/115 0.36 (0.30, 0.43) 26.28 78/214 0.47 (0.33, 0.61) 21.67 22/47 0.32 (0.20, 0.45) 100.00 | |
| 8 years Wang 2011_W, American Indians/Islands Wang 2011_M, American Indians/Islands Subtotal (Tau²=0) | 0.52 (0.45, 0.58) 64.85 119/23 0.53 (0.44, 0.61) 35.15 66/125 0.52 (0.47, 0.57) 100.00 | |
| 9 years Anjana 2015, Asia/Middle East | 0.84 (0.74, 0.91) 100.00 58/69 | |
| 10 years Larsson 2000, Australia/Europe/North America Bonora 2011, Australia/Europe/North America Subtotal (Tsu²=0) | • 0.20 (0.10, 0.37) 61.00 6/30 0.47 (0.27, 0.68) 39.00 9/19 0.30 (0.17, 0.44) 100.00 | |
| 12 years Han 2017, Asia/Middle East Vaccaro 1999, Australia/Europe/North America Subtotal (Tau²=0) | • • • • • • • • • • • • • • • • • • • | |
| 0 0.2 0 | 4 0.6 0.8 1 | |
| Cumula | tive incidence | |

T2DM cumulative incidence associated with IFG and IGT thresholds

The number of studies and participants, and the cumulative incidence of T2DM (pooled if more than one study) according to follow-up period were as follows.

• 1 year's follow-up: 1 study, 207 participants, cumulative incidence 29% (95% CI 23 to 36).

• 3 years' follow-up: 1 study, 209 participants, cumulative incidence 34% (95% CI 28 to 41).

• 5 years' follow-up: 5 studies, 478 participants, cumulative incidence 50% (95% CI 37 to 63).

• 6 years' follow-up: 4 studies, 106 participants, cumulative incidence 58% (95% CI 48 to 67).

• 7 years' follow-up: 4 studies, 753 participants, cumulative incidence 32% (95% CI 20 to 45).

• 8 years' follow-up: 1 study, 356 participants, cumulative

incidence 52% (95% CI 47 to 57).

• 9 years' follow-up: 1 study, 69 participants, cumulative incidence 84% (95% CI 74 to 91).

• 10 years' follow-up: 2 studies, 49 participants, cumulative incidence 30% (95% CI 17 to 44).

• 12 years' follow-up: 2 studies, 207 participants, cumulative incidence 70% (95% CI 63 to 76).

IH defined by HbA1c5.7 threshold

Diabetes incidence associated with HbA1c_{5.7} at baseline and a follow-up period of 4 to 10 years showed pooled cumulative incidences of 14% to 31% (Figure 14).

Figure 14. Elevated glycosylated haemoglobin A1c (HbA1c) 5.7% threshold: association with cumulative type 2 diabetes mellitus (T2DM) incidence over 4-10 years CI: confidence interval; n/N: events/number of participants

| Study, location | Cumulative incidence (95% Cl) | % Weight n/N |
|--|---|----------------|
| 4 years | | |
| Kim 2014*, Asia/Middle East | 0.11 (0.05, 0.21) | 25.72 7/64 |
| Lee 2016, Asia/Middle East | 0.11 (0.10, 0.12) | 37.29 390/3497 |
| Bae 2011, Asia/Middle East | 0.21 (0.19, 0.23) | 36.99 373/1791 |
| Subtotal (Tau ² = 0.03) | 0.14 (0.07, 0.23) | 100.00 |
| 5 years | | |
| Nakagami 2016, Asia/Middle East - | 0.15 (0.12, 0.18) | 25.43 87/583 |
| Kim 2016a, Asia/Middle East | 0.22 (0.21, 0.24) | 26.57 435/1951 |
| Heianza 2012, Asia/Middle East | 0.22 (0.20, 0.25) | 25.89 184/822 |
| Vijayakumar 2017, Australia/Europe/North America | 0.45 (0.37, 0.52) | 22.10 75/168 |
| Subtotal (Tau ² = 0.03) | > 0.25 (0.18, 0.32) | 100.00 |
| 6 years | | |
| Man 2017, Asia/Middle East | 0.17 (0.14, 0.20) | 100.00 112/675 |
| 7 years | | |
| Lipska 2013*, Australia/Europe/North America | 0.21 (0.16, 0.27) | 100.00 44/207 |
| 10 years | | |
| Han 2017. Asia/Middle East | • 0.31 (0.29, 0.33) | 99.14 881/2830 |
| Cederberg 2010*, Australia/Europe/North America | • 0.38 (0.21, 0.57) | 0.86 9/24 |
| Subtotal (Tau ² = 0) | Image: Construction of the second s | 100.00 |
| 1 1 | | |
| 0 0.2 | 0.4 0.6 | |
| Cumulativ | <i>i</i> e incidence | |

T2DM cumulative incidence associated with HbA1c 5.7% threshold

The number of studies and participants, and the cumulative incidence of T2DM (pooled if more than one study) according to follow-up period were as follows.

• 4 years' follow-up: 3 studies, 5352 participants, cumulative incidence 14% (95% CI 7 to 23).

• 5 years' follow-up: 4 studies, 3524 participants, cumulative incidence 25% (95% CI 18 to 32).

• 6 years' follow-up: 1 study, 675 participants, cumulative incidence 17% (95% CI 14 to 20).

• 7 years' follow-up: 1 study, 207 participants, cumulative

incidence 21% (95% CI 16 to 27).

• 10 years' follow-up: 2 studies, 2854 participants, cumulative incidence 31% (95% CI 29 to 33).

IH defined by HbA1c_{6.0} threshold

Most studies were undertaken in Asia. Diabetes incidence associated with HbA1c_{6.0} at baseline and a follow-up period of 3 to 15 years showed pooled cumulative incidences of 7% to 44% (Figure 15).

Figure 15. Elevated glycosylated haemoglobin A1c (HbA1c) 6.0% threshold: association with cumulative type 2 diabetes mellitus (T2DM) incidence over 3-15 years CI: confidence interval; n/N: events/number of participants

| Study, location | Cumulativ | e incidence (95% CI) | % Weig | ht n/N |
|--|-------------------|----------------------|--------|---------|
| 3 years | | | | |
| Chamnan 2011, Australia/Europe/North America 🔸 | | 0.07 (0.05, 0.10) | 100.00 | 26/370 |
| 4 years | | | | |
| Sato 2009, Asia/Middle East | _ •_ | 0.42 (0.35, 0.49) | 34.32 | 90/215 |
| Bae 2011, Asia/Middle East | _ | 0.45 (0.41, 0.50) | 65.68 | 187/412 |
| Subtotal (Tau ² = 0) | \diamond | 0.44 (0.40, 0.48) | 100.00 | |
| 5 years | | | | |
| Kim 2016a, Asia/Middle East | -•- | 0.29 (0.27, 0.32) | 35.36 | 322/110 |
| Nakagami 2016, Asia/Middle East | | 0.37 (0.30, 0.45) | 31.89 | 58/156 |
| Heianza 2012, Asia/Middle East | _ - • | - 0.49 (0.42, 0.56) | 32.75 | 100/203 |
| Subtotal (Tau ² = 0.05) | $\langle \rangle$ | 0.38 (0.26, 0.51) | 100.00 | |
| 15 years | | | | |
| Bonora 2011, Australia/Europe/North America | - _ | 0.29 (0.19, 0.40) | 100.00 | 20/70 |
| 1 | 0.2 0.4 | 0.6 | | |

T2DM cumulative incidence associated with HbA1c 6.0% threshold

The number of studies and participants, and the cumulative incidence of T2DM (pooled if more than one study) according to follow-up period were as follows.

• 3 years' follow-up: 1 study, 370 participants, cumulative

incidence 7% (95% CI 5 to 10).

- 4 years' follow-up: 2 studies, 627 participants, cumulative incidence 44% (95% CI 40 to 48).
 - 5 years' follow-up: 3 studies, 1462 participants, cumulative

33

incidence 38% (95% CI 26 to 51).
15 years' follow-up: 1 study, 70 participants, cumulative incidence 29% (95% CI 19 to 40).

Children and adolescents with IH (mostly IGT)

Diabetes incidence in children and adolescents, usually associated with IGT at baseline and with follow-up of 1 to 10 years, showed pooled cumulative incidences of 1% to 56% (Figure 16). We did not observe any distinct pattern between T2DM incidence and geography.

Figure 16. Cumulative type 2 diabetes mellitus (T2DM) incidence in children/adolescents over 1-10 years CI: confidence interval; HbA1c 5.7: glycosylated haemoglobin A1c 5.7% threshold; (i-)IGT: (isolated) impaired glucose tolerance; n/N: events/number of participants; NO: non-overweight; OV: overweight

| Study, location, intermediate hyperglycaemia origin | | Cumulative incidenc | e (95% CI) % | Weight | n/N |
|--|----------------------|---------------------|--------------|--------|--------|
| 1 year | | | | | |
| Kleber 2010, Australia/Europe/North America, IGT 🛛 🗕 🗕 | | 0.01 (0. | 00, 0.07) | 100.00 | 1/79 |
| 2 years | | | | | |
| Weiss 2005, Australia/Europe/North America, i-IGT | - | 0.24 (0. | 13, 0.41) | 100.00 | 8/33 |
| 4 years | | | | | |
| Kleber 2011, Australia/Europe/North America, IGT 🔸 | | 0.03 (0. | 01, 0.07) | 100.00 | 3/119 |
| 5 years | | | | | |
| Wheelock 2016_NO, American Indians/Islands, | • | 0.24 (0. | 13, 0.40) | 14.10 | 9/37 |
| Wheelock 2016_OV, American Indians/Islands, | - _ | 0.37 (0. | 29, 0.46) | 49.81 | 49/13 |
| Vijayakumar 2017, American Indians/Islands, HbA1c 5.7 | • | 0.29 (0. | 19, 0.41) | 23.50 | 18/62 |
| Jaruratanasirikul 2016, Asia/Middle East, i-IGT | • | 0.27 (0. | 15, 0.44) | 12.59 | 9/33 |
| Subtotal (Tau ² = 0) | \diamond | 0.32 (0. | 26, 0.38) | 100.00 | |
| 10 years | | | | | |
| Wheelock 2016_NO, American Indians/Islands, | • | 0.30 (0. | 17, 0.46) | 22.06 | 11/37 |
| Wheelock 2016_OV, American Indians/Islands, | | • 0.64 (0. | 55, 0.71) | 77.94 | 84/132 |
| Subtotal (Tau ² = 0) | < | 0.56 (0. | 49, 0.64) | 100.00 | |
| 1 | 0.2 0.4 | 0.6 0.8 | | | |
| 0 | Cumulative incidence | 0.0 0.0 | | | |

T2DM cumulative incidence in children or adolescents

The number of studies and participants, and the cumulative incidence of T2DM (pooled if more than one study) according to follow-up period were as follows.

• 1 year's follow-up: 1 study, 79 participants, cumulative incidence 1% (95% CI 0 to 7).

Development of type 2 diabetes mellitus in people with intermediate hyperglycaemia (Review) Copyright © 2018 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.

Figure 17. Regression from intermediate hyperglycaemia to normoglycaemia in adults over 1-5 years CI: confidence interval; HbA1c5.7: glycosylated haemoglobin A1c 5.7%; i-IFG5.6/6.1: (isolated) impaired fasting glucose 5.6/6.1 mmol/L threshold;IGT: impaired glucose tolerance; n/N: events/number of participants Regression to normoglycaemia: adults only, 1 to 5 years follow-up Cumulative incidence (95% CI) % Weight n/N Study, location, prediabetic origin

Special populations with IH

incidence 24% (95% CI 13 to 41).

incidence 3% (95% CI 1 to 7).

Studies involving black populations were scarce: one study reported a cumulative T2DM incidence of 35% in African Surinamese after 10 years of follow-up in association with IFG_{5.6} at baseline (Admiraal 2014). Another study, which used IFG_{5.6} at baseline, reported a T2DM cumulative incidence of 33% in

• 2 years' follow-up: 1 study, 33 participants, cumulative

• 4 years' follow-up: 1 study, 119 participants, cumulative

• 5 years' follow-up: 3 studies, 264 participants, pooled

• 10 years' follow-up: 1 study (2 subpopulations), 169

cumulative incidence 32% (95% CI 26 to 38).

African Americans after 7.5 years of follow-up (Yeboah 2011).

b) Regression from IH to normoglycaemia

Adults

In the 47 studies reporting data on regression from IH to normoglycaemia in adults within a follow-up period of 1 to 11 years, pooled percentages ranged from 17% to 59% (Figure 17; Figure 18). Regression to normoglycaemia varied widely and showed neither a clear linear reduction or increase nor a distinct pattern associated with geography. Regression rates were often reported in association with IGT at baseline; however, there were no distinct differences in regression rates when compared with IFG5.6, IFG6.1 or HbA1c5.7 as IH risk factors.

participants, cumulative incidence 56% (95% CI 49 to 64).

| study, location, prediabelic origin | Cumulative incidence (95% CI) | A Weight him |
|---|--|----------------------------|
| 1 year | | |
| Ko 1999, Asia/Middle East, IGT | 0.49 (0.40, 0.58) | 32.85 60/12 |
| Bai 1999, Asia/Middle East, IGT Subtotal (Tau ² = 0) | 0.64 (0.58, 0.70) 0.59 (0.54, 0.64) | 67.15 162/2 100.00 |
| Subtotal (Tau ² = 0) | 0.59 (0.54, 0.64) | 100.00 |
| 2 years | | |
| Kim 2008, Asia/Middle East, IFG 5.6 | 0.56 (0.53, 0.59) | 12.01 747/1 |
| Ko 2001, Asia/Middle East, IFG 6.1 | 0.31 (0.20, 0.44) | 9.77 17/55 |
| Li 2003, Asia/Middle East, IGT | 0.17 (0.11, 0.24) | 11.01 22/13 |
| Ammari 1998, Asia/Middle East, IGT | 0.40 (0.29, 0.52) | 10.14 27/68 11.65 174/3 |
| Wat 2001, Asia/Middle East, IGT Marshall 1994, Australia/Europe/North America, IGT | 0.54 (0.49, 0.59) 0.49 (0.40, 0.58) | 11.65 174/3 10.94 60/12 |
| Rajala 2000, Australia/Europe/North America, IGT | ► 0.63 (0.56, 0.70) | 11.30 115/1 |
| Charles 1997. Australia/Europe/North America, IGT | 0.65 (0.50, 0.70) 0.65 (0.61, 0.70) | 11.76 273/4 |
| Coronado-Malagon 2009, Latin America, Prediabetes | 0.35 (0.29, 0.42) | 11.43 76/21 |
| Subtotal (Tau ² = 0.07) | 0.46 (0.36, 0.55) | 100.00 |
| | | |
| 3 years Jiamjarasrangsi 2008a, Asia/Middle East, IFG 5.6 —● | - 0.62 (0.56, 0.67) | 14.63 197/3 |
| Toshihiro 2008. Asia/Middle East, IFG 5.6 and/or IGT | 0.30 (0.23, 0.39) | 14.34 39/12 |
| Chen 2003, Asia/Middle East, IFG 6.1 | 0.83 (0.76, 0.88) | 14.42 129/1 |
| Liu 2014. Asia/Middle East, IFG or IGT | 0.29 (0.25, 0.33) | 14.69 130/4 |
| Inoue 1996. Asia/Middle East. IGT | 0.30 (0.17, 0.46) | 13.24 11/37 |
| Ramachandran 1986, Asia/Middle East, IGT | 0.32 (0.24, 0.41) | 14.24 34/10 |
| Rijkelijkhuizen 2007, Australia/Europe/North America, IGT — | 0.22 (0.16, 0.29) | 14.43 35/15 |
| Subtotal (Tau ² = 0.22) | 0.41 (0.24, 0.59) | 100.00 |
| 4 years | | |
| Wang 2011, American Indians/Islands, IGT - | 0.28 (0.24, 0.32) | 42.48 147/5 |
| Motala 2003, Asia/Middle East, IGT | 0.39 (0.28, 0.50) | 22.74 28/72 |
| Mykkänen 1993, Australia/Europe/North America, IGT | 0.35 (0.29, 0.42) | 34.78 72/20 |
| Subtotal (Tau ² = 0.01) | 0.33 (0.26, 0.40) | 100.00 |
| 5 years | | |
| Wheelock 2016, American Indians/Islands, IGT | 0.34 (0.27, 0.42) | 11.03 49/14 |
| Heianza 2012, Asia/Middle East, HbA1c 5.7 - | 0.32 (0.29, 0.35) | 12.24 263/8 |
| Latifi 2016, Asia/Middle East, IFG 5.6 | 0.50 (0.41, 0.59) | 10.81 62/12 |
| Qian 2012_iIFG6.1, Asia/Middle East, i-IFG 6.1 | 0.23 (0.14, 0.35) | 9.44 14/60 |
| Cugati 2007, Australia/Europe/North America, IFG 5.6 | 0.41 (0.35, 0.48) | 11.54 94/22 |
| Lecomte 2007, Australia/Europe/North America, IFG 6.1 | 0.40 (0.37, 0.44) 0.34 (0.28, 0.40) | 12.21 297/7 11.56 79/23 |
| Rajala 2000, Australia/Europe/North America, IGT | 0.56 (0.49, 0.63) | 11.24 96/17 |
| Guerrero-Romero 2006, Latin America, IGT | 0.04 (0.01, 0.11) | 9.93 3/75 |
| Subtotal (Tau ² = 0.05) | 0.34 (0.27, 0.42) | 100.00 |
| | 1 1 | |
| | | |
| 0 0.2 0.4 0.6 | 5 0.8 1 | |

Figure 18. Regression from intermediate hyperglycaemia to normoglycaemia in adults over 6-11 years CI: confidence interval; i-IFG_{5.6/6.1}: (isolated) impaired fasting glucose 5.6/6.1 mmol/L threshold; i-IGT: (isolated) impaired glucose tolerance; n/N: events/number of participants

| Study, location, prediabetic origin | | Cumulative incide | nce(95% CI) | % Weigh | nt n/N |
|---|---------------------|-------------------|--------------|---------|---------|
| 6 years | | | | | |
| Dowse 1991, American Indians/Islands, IGT - | | | (0.27, 0.53) | 19.79 | 20/51 |
| Rijkelijkhuizen 2007, Australia/Europe/North America, IFG 5.6 🔸 | | 0.07 | (0.05, 0.09) | 20.47 | 33/488 |
| Lyssenko 2005, Australia/Europe/North America, IFG or IGT | -+ | | (0.52, 0.59) | 20.49 | 379/686 |
| Schranz 1989, Australia/Europe/North America, IGT | • | 0.33 | (0.24, 0.45) | 20.03 | 25/75 |
| Leiva 2014, Latin America, IFG 6.1 | | 0.00 | (0.00, 0.12) | 19.22 | 0/28 |
| Subtotal (Tau ² = 0.51) | | 0.23 | (0.03, 0.53) | 100.00 | |
| 7 years | | | | | |
| Sharifi 2013, Asia/Middle East, IFG 5.6 | - _ | 0.43 | (0.35, 0.52) | 18.14 | 53/123 |
| Sadeghi 2015, Asia/Middle East, IFG 5.6 and/or IGT | | 0.40 | (0.35, 0.45) | 54.85 | 148/373 |
| Sasaki 1982, Asia/Middle East, IGT | • | 0.38 | (0.18, 0.64) | 1.98 | 5/13 |
| Ferrannini 2009, Latin America, IGT | | | (0.36, 0.50) | 25.04 | 73/170 |
| Subtotal (Tau ² = 0) | \diamond | | (0.37, 0.45) | 100.00 | |
| 8 vears | | | | | |
| Mohan 2008. Asia/Middle East, IGT | - | 0.16 | (0.08, 0.31) | 11.40 | 6/37 |
| Wong 2003, Asia/Middle East, IGT | | | (0.36, 0.48) | 88.60 | 122/29 |
| Subtotal (Tau ² =0) | \diamond | | (0.33, 0.44) | 100.00 | |
| 9 years | | | | | |
| Anjana 2015, Asia/Middle East, i-IFG 5.6 or iIGT | | 0.17 | (0.14, 0.22) | 100.00 | 52/299 |
| 10 years | | | | | |
| Motala 2003. Asia/Middle East. IGT | . | - 0.46 | (0.30, 0.62) | 13.73 | 16/35 |
| Cugati 2007, Australia/Europe/North America, IFG 5.6 - | | | (0.04, 0.11) | 14.77 | 15/229 |
| de Abreu 2015, Australia/Europe/North America, IFG 5.6 | | | (0.48, 0.63) | 14.73 | 104/18 |
| Baena-Diez 2011. Australia/Europe/North America. IFG 6.1 | | | (0.48, 0.63) | 14.75 | 57/115 |
| Forouhi 2007, Australia/Europe/North America, IFG 6.1 | | | (0.41, 0.53) | 14.79 | 143/25 |
| | | | | | |
| Peterson 2017, Australia/Europe/North America, IGT | | | (0.15, 0.46) | 13.50 | 8/29 |
| Larsson 2000_ilFG6.1, Australia/Europe/North America, i-IFG 6.1 | | | (0.49, 0.77) | 13.92 | 27/42 |
| Subtotal (Tau ² = 0.31) | | 0.42 | (0.22, 0.63) | 100.00 | |
| 11 years | | | | | |
| Söderberg 2004_iIFG6.1, American Indians/Islands, i-IFG 6.1 | | | (0.33, 0.43) | 35.14 | 153/40 |
| Song 2016a_M, Asia/Middle East, IFG 5.6 and/or IGT | _ | | (0.16, 0.30) | 31.39 | 28/125 |
| Song 2016a_W, Asia/Middle East, IFG 5.6 and/or IGT | | | (0.17, 0.29) | 33.47 | 47/209 |
| Subtotal (Tau ² = 0.04) | | 0.28 | (0.17, 0.39) | 100.00 | |
| | 1 | 1 | | | |
| 0 0.2 | 0.4 | 0.6 0.8 | | | |
| c | umulative incidence | | | | |

Regression to normoglycaemia: adults only, 6 to 11 years follow-up

The number of studies and participants, and the proportion regressing from IH to normoglycaemia (pooled if more than one study) according to follow-up period were as follows.

• 1 year's follow-up: 2 studies, 375 participants, regression to normoglycaemia 59% (95% CI 54 to 64).

- 2 years' follow-up: 9 studies, 2852 participants, regression to normoglycaemia 46% (95% CI 36 to 55).
- 3 years' follow-up: 7 studies, 1356 participants, regression to normoglycaemia 41% (95% CI 24 to 59).
- 4 years' follow-up: 3 studies, 807 participants, regression to normoglycaemia 33% (95% CI 26 to 40).
 - 5 years' follow-up: nine studies, 2603 participants,

regression to normoglycaemia 34% (95% CI 27 to 42).

- 6 years' follow-up: 5 studies, 1328 participants, regression to normoglycaemia 23% (95% CI 3 to 53).
- 7 years' follow-up: 4 studies, 679 participants, regression to normoglycaemia 41% (95% CI 37 to 45).
- 8 years' follow-up: 2 studies, 328 participants, regression to normoglycaemia 39% (95% CI 33 to 44).
- 9 years' follow-up: 1 study, 299 participants, regression to normoglycaemia 17% (95% CI 14 to 22)
- 10 years' follow-up: 7 studies, 894 participants, regression to normoglycaemia 42% (95% CI 22 to 63).

• 11 years' follow-up: 2 studies, 736 participants, regression to normoglycaemia 28% (95% CI 17 to 39).

Children and adolescents

Regression from IH to normoglycaemia in children and adolescents within a follow-up period of one to four years showed percentages from 45% to 81% (Figure 19). There were no distinct patterns with regard to geography. IGT at baseline was often investigated as the IH risk factor.

Figure 19. Regression from intermediate hyperglycaemia to normoglycaemia in children/adolescents over I-4 years

CI: confidence interval; IGT: impaired glucose tolerance; n/N: events/number of participants

| Study, location, prediabetic origin | | | | | Cumulativ | ve incidence (95% Cl) | n/N |
|--|---|-----|-----|-----|-----------|-----------------------|--------|
| 1 year | | | | | | | |
| Kleber 2010, Australia/Europe/North America, IGT | | | | -• | _ | 0.66 (0.55, 0.75) | 52/79 |
| 2 years | | | | | | | |
| Weiss 2005, Australia/Europe/North America, IGT | | | • | | | 0.45 (0.30, 0.62) | 15/33 |
| 4 years | | | | | | | |
| Kleber 2011, Australia/Europe/North America, IGT | | | | | -•- | 0.81 (0.73, 0.87) | 96/119 |
| | 0 | 0.2 | 0.4 | 0.6 | 0.8 | 1 | |

Regression to normoglycaemia: children or adolescents

The number of studies and participants, and the proportion regressing from IH to normoglycaemia according to follow-up period were as follows.

• 1 year's follow-up: 1 study, 79 participants, regression to normoglycaemia 66% (95% CI 55 to 75).

• 2 years' follow-up: 1 study, 33 participants, regression to normoglycaemia 45% (95% CI 30 to 62).

• 4 years' follow-up: 1 study, 119 participants, regression to normoglycaemia 81% (95% CI 73 to 87).

Prognostic factor studies used various definitions for IH and different effect measures (IRR, OR and HR) to express the effect of glycaemic status on development of T2DM. The findings are presented below according to IH definition and effect measure. No data were available on the prognostic factor IH versus normoglycaemia for children or adolescents.

HR as the effect measure

c) IH versus normoglycaemia as a prognostic factor for developing T2DM

IFG 5.6 mmol/L threshold

Eight studies reported HRs and the IFG_{5.6} threshold for IH at baseline (Analysis 1.1). The length of follow-up ranged from 4 to 22 years (studies are ordered with ascending length of follow-up in Analysis 1.1). The studies included 9017 participants with IH and 25,850 participants with normoglycaemia. The overall HR was 4.32 (95% CI 2.61 to 7.12). The 95% prediction interval ranged from 0.75 to 25.01

The comparison of geographic regions showed the following results (Analysis 1.1).

• Asia/Middle East (4 studies, 2385 participants with IH and 12,418 participants with normoglycaemia, 5 to 12 years' followup): the pooled HR was 5.07 (95% CI 3.41 to 7.53). The 95% prediction interval ranged from 1.07 to 24.02.

• Australia/Europe/North America (3 studies, 5685 participants with IH and 12,837 participants with normoglycaemia, 8 to 22 years' follow-up): the pooled HR was 4.15 (95% CI 1.24 to 13.87). Calculation of the 95% prediction interval did not provide a meaningful estimate.

• American Indians/Islands (1 study, 947 participants with IH and 595 participants with normoglycaemia, 4 years' follow-up): the HR was 2.38 (95% CI 1.85 to 3.06).

IFG 6.1 mmol/L threshold

Nine studies reported HRs and the IFG_{6.1} threshold for IH at baseline (Analysis 1.2). The length of follow-up ranged from 5 to 22 years (studies are ordered by ascending length of follow-up in Analysis 1.2). The studies included 2818 participants with IH and 18,591 participants with normoglycaemia. The overall HR was 5.47 (95% CI 3.50 to 8.54). The 95% prediction interval ranged from 1.09 to 27.56

The comparison of geographic regions showed the following results (Analysis 1.2).

• Asia/Middle East (5 studies, 1054 participants with IH and 9756 participants with normoglycaemia, 5 to 11 years' followup): the pooled HR was 10.55 (95% CI 3.61 to 30.81). Calculation of the 95% prediction interval did not provide a meaningful estimate.

• Australia/Europe/North America (4 studies, 1736 participants with IH and 8835 participants with normoglycaemia, 6 to 22 years' follow-up): the pooled HR was 3.30 (95% CI 2.32 to 4.67). The 95% prediction interval ranged from 0.84 to 12.99.

• Latin America (1 study, 28 participants with IH and 66 participants with normoglycaemia, 6 years' follow-up): the HR was 2.06 (95% CI 1.76 to 2.41).

IGT

Five studies reported HRs and IGT for IH at baseline (Analysis 1.3). The length of follow-up ranged from 5 to 16 years (studies

are ordered by ascending length of follow-up in Analysis 1.3). These studies included 4010 participants with IH and 12,566 participants with normoglycaemia. The overall HR was 3.61 (95% CI 2.31 to 5.64). The 95% prediction interval ranged from 0.69 to 18.97.

The comparison of geographic regions showed the following results (Analysis 1.3).

• Asia/Middle East (3 studies, 1780 participants with IH and 6695 participants with normoglycaemia, 5 to 12 years' followup): the pooled HR was 4.48 (95% CI 2.81 to 7.15). Calculation of the 95% prediction interval did not provide a meaningful estimate.

• Australia/Europe/North America (2 studies, 2230 participants with IH and 5871 participants with normoglycaemia, 6 to 16 years' follow-up): the pooled HR was 2.53 (95% CI 1.52 to 4.19).

Combined IFG and IGT

Five studies reported HRs and used both IFG and IGT for defining IH at baseline (Analysis 1.4). The length of follow-up ranged from 4 to 12 years (studies are ordered by ascending length of follow-up in Analysis 1.4). These studies included 1038 participants with IH and 8719 participants with normoglycaemia. The overall HR was 6.90 (95% CI 4.15 to 11.45). The 95% prediction interval ranged from 1.06 to 44.95.

The comparison of geographic regions showed the following results (Analysis 1.4).

• Asia/Middle East (3 studies, 461 participants with IH and 6695 participants with normoglycaemia, 5 to 12 years' followup): the pooled HR was 10.20 (95% CI 5.45 to 19.09). Calculation of the 95% prediction interval did not provide a meaningful estimate.

• Australia/Europe/North America (1 study, 221 participants with IH and 1429 participants with normoglycaemia, 6 years' follow-up): the HR was 3.80 (95% CI 2.30 to 6.28).

• American Indians/Islands (1 study, 356 participants with both IFG and IGT and 595 participants with normoglycaemia, 4 years' follow-up): the HR was 4.06 (95% CI 3.05 to 5.40).

HbA1c 5.7% threshold

Four studies reported HRs and the HbA1c_{5.7} threshold for IH at baseline (Analysis 1.5). The length of follow-up ranged from 4 to 22 years (studies are ordered by ascending length of follow-up in Analysis 1.5). The studies included 5223 participants with IH and 19,824 participants with normoglycaemia. The overall HR was 5.55 (95% CI 2.77 to 11.12). The 95% prediction interval ranged from 0.23 to 141.18.

The comparison of geographic regions showed the following results (Analysis 1.5).

• Asia/Middle East (3 studies, 3196 participants with IH and 13,609 participants with normoglycaemia, 4 to 5 years' followup): the pooled HR was 7.21 (95% CI 5.14 to 10.11). The 95% prediction interval ranged from 0.81 to 64.52.

• Australia/Europe/North America (1 study, 2027 participants with IH and 6215 participants with normoglycaemia, 22 years' follow-up): the HR was 2.71 (95% CI 2.48 to 2.96).

HbA1c 6.0% threshold

Six studies reported HRs and the HbA1c_{6.0} threshold for IH at baseline (Analysis 1.6). The length of follow-up ranged from 4 to 22 years (studies are ordered by ascending length of follow-up in Analysis 1.6). The studies included 4532 participants with IH and 26,167 participants with normoglycaemia. The overall HR was 10.10 (95% CI 3.59 to 28.43). Calculation of the 95% prediction interval did not provide a meaningful estimate.

The comparison of geographic regions showed the following results (Analysis 1.6).

• Asia/Middle East (4 studies, 3492 participants with IH and 19,242 participants with normoglycaemia, 4 to 12 years' followup): the pooled HR was 13.12 (95% CI 4.10 to 41.96). Calculation of the 95% prediction interval did not provide a meaningful estimate.

Australia/Europe/North America (2 studies, 1040

participants with IH and 6925 participants with normoglycaemia, 15 to 22 years' follow-up): the pooled HR was

5.09 (95% CI 1.69 to 15.37).

Both elevated HbA1c and IFG

One study in Japanese participants provided data on elevated HbA1c and IFG for defining IH at baseline and estimated the effect of IH versus normoglycaemia using the HR (Analysis 1.7). The combination of HbA1c_{5.7} plus IFG_{5.6} (410 participants) when compared with normoglycaemia (4149 participants) showed an HR of 32.50 (95% CI 23.00 to 45.92). The combination of

HbA1c_{5.7} plus IFG_{6.1} (159 participants) when compared with normoglycaemia (5198 participants) showed an HR of 37.90 (95% CI 28.10 to 51.12). The combination of HbA1c_{6.0} plus IFG_{5.6} (135 participants) when compared with normoglycaemia (4493 participants) showed an HR of 53.70 (95% CI 38.40 to 75.09). The combination of HbA1c_{6.0} plus IFG_{6.1} (72 participants) when compared with normoglycaemia (5730 participants) showed an HR of 52.30 (95% CI 37.80 to 72.37).

IH in special populations

Data on black populations were scarce: one study in African Surinamese reported an adjusted OR of 5.1 (95% CI 2.0 to 13.3) for the association between IFG_{5.6} at baseline and T2DM incidence at 7.5 years' follow-up (Admiraal 2014). Another study including a subgroup of African Americans reported the association of various measures of IH at baseline with the development of T2DM using HRs (Warren 2017): after 16 years of follow-up the HR for IFG_{5.6} was 2.65 (95% CI 2.11 to 3.32); for IFG_{6.1}, the HR was 3.16 (95% CI 2.47 to 4.06); and for IGT, the HR was 2.55 (95% CI 2.01 to 3.22). After 22 years' follow-up, the HR for IFG_{5.6} was 2.05 (95% CI 1.75 to 2.40); for IFG_{6.1}, the HR was 2.66 (95% CI 2.26 to 3.13); for HbA1c_{5.7}, the HR was 2.24 (95% CI 1.92 to 2.61); and for HbA1c_{6.0}, the HR was 2.60 (95% CI 2.21 to 3.05).

Incidence rate ratio as the effect measure

IFG 5.6 mmol/L threshold

Ten studies reported incidence rate ratios (IRRs) and used the IFG_{5.6} threshold for IH. The studies included 24,357 participants with IH and 155,272 participants with normoglycaemia (Figure 20). Of those with IH, 661 (2.7%) developed T2DM compared with 1270 (0.8%) in participants with normoglycaemia. The overall IRR was 4.81 (95% CI 3.67 to 6.30) with a 95% prediction interval ranging from 1.95 to 11.83.

Figure 20. IFG: impaired fasting glucose; IRR: incidence rate ratio; n: number of cases; T: person-time in years

| | | | | | | | Prediabetic | Normoglycaemi |
|------------------------------------|-------|-------------|------------|-------------|----------------|----------|-------------|---------------|
| Study | | | | IRR (95% | CI) | % Weight | n/T | n/T |
| Asia/Middle East | | | | | | | | |
| Anjana 2015* | | | - | 2.74 (1.8 | 9, 3.98) | 11.26 | 32/525 | 209/9398 |
| Han 2017* | | - | -∎÷ | 4.17 (3.3 | 1, 5.26) | 12.95 | 81/1579 | 657/53461 |
| Janghorbani 2015* | | _ | | 5.34 (2.7 | 5, 10.37) | 7.68 | 23/1409 | 14/4578 |
| Park 2006 | | | | 5.48 (3.8) | 2, 7.85) | 11.42 | 40/1278 | 116/20298 |
| Derakhshan 2016 | | | | 7.46 (5.9) | 8, 9.32) | 13.04 | 150/4950 | 162/39901 |
| Heianza 2012 | | | | 7.92 (5.6 | 1, 11.18) | 11.60 | 108/5920 | 46/19961 |
| Subtotal (Tau ² = 0.13) | | | \sim | - 5.23 (3.7 | 7, 7.25) | 67.94 | | |
| with.estimated prediction interval | | | | (1.7 | 2, 15.89) |) | | |
| Australia/Europe/North America | | | | | | | | |
| Forouhi 2007 | | | - | 4.42 (2.10 | 0, 9.29) | 6.88 | 53/5000 | 8/3333 |
| Valdes 2008 | | | - | 5.18 (2.3 | , 5, 11.41) | 6.45 | 14/718 | 11/2923 |
| Soriguer 2008 | | _ | | 5.29 (2.6) | , 3, 10.44) | 7.52 | 23/604 | 13/1806 |
| Subtotal (Tau ² = 0) | | | \diamond | → 4.96 (3.2 | 5, 7.57) | 20.85 | | |
| with estimated prediction interval | | | | (0.3 | 2, 77.24) |) | | |
| American Indians/Islands | | | | | | | | |
| Wang 2011 | | | - | 2.74 (1.8 | 8, 3.99) | 11.21 | 137/2374 | 34/1613 |
| | | | | | | | | |
| Overall (Tau ² = 0.13) | | | \diamond | 4.81 (3.6 | 7, 6.30) | 100.00 | | |
| with.estimated prediction interval | | | | (1.9 | 5, 11.83) |) | | |
| | | | | | | | | |
| | 0.5 1 | 2 | 4 8 | 16 | | | | |
| | Incid | ence rate r | atio (IRR) | | | | | |

T2DM incidence rate ratio associated with IFG 5.6 mmol/L threshold

The results for the geographic regions were as follows.

• Asia/Middle East (6 studies): T2DM developed in 434/ 15,661 (2.8%) participants with IH and in 1204/145,597 (0.8%) participants with normoglycaemia. The pooled IRR was 5.23 (95% CI 3.77 to 7.25) with a 95% prediction interval ranging from 1.72 to 15.89.

• Australia/Europe/North America (3 studies): T2DM developed in 90/6322 (1.4%) participants with IH and in 32/8062 (0.4%) participants with normoglycaemia. The pooled IRR was 4.96 (95% CI 3.25 to 7.57) with a 95% prediction interval ranging from 0.32 to 77.24.

• American Indians/Islands (1 study): T2DM developed in 137/2374 (5.8%) participants with IH and in 34/1613 (2.1%)

participants with normoglycaemia. The IRR was 2.74 (95% CI 1.88 to 3.99).

IFG 6.1 mmol/L threshold

Six studies reported IRRs and used an IFG_{6.1} threshold for IH. Thee studies included 5115 participants with IH, of whom 127 (2.5%) developed T2DM, plus 56,580 participants with normoglycaemia, of whom 188 (0.3%) developed T2DM (Figure 21). The overall IRR was 6.82 (95% CI 4.53 to 10.25) with a 95% prediction interval ranging from 2.03 to 22.87.

Figure 21. IFG: impaired fasting glucose; IRR: incidence rate ratio; n: number of cases; T: person-time in years

| | | | Pr | ediabetic l | Normoglycaemi |
|---|-------------------|-----------------------|----------|-------------|---------------|
| Study | | IRR (95% CI) | % Weight | n/T | n/T |
| Australia/Europe/North America | | | | | |
| Forouhi 2007 | _ | 7.29 (3.37, 15.75) | 14.38 | 34/1943 | 8/3333 |
| Rijkelijkhuizen 2007* | _ _ | 7.34 (4.78, 11.29) | 22.21 | 35/681 | 51/7286 |
| Bonora 2011 | _ | 8.56 (4.76, 15.42) | 18.29 | 18/486 | 29/6704 |
| Valdes 2008 | | - 15.39 (7.33, 32.35) | 14.92 | 19/328 | 11/2923 |
| Subtotal (Tau ² = 0.002) | ↔ | 8.55 (6.37, 11.48) | 69.80 | | |
| with estimated prediction interval | | (4.37, 16.73) | | | |
| Asia/Middle East | | | | | |
| Nakanishi 2004 - | | 2.23 (0.89, 5.60) | 11.82 | 5/1506 | 51/34308 |
| Li 2003* | | 4.99 (2.78, 8.95) | 18.38 | 16/171 | 38/2026 |
| Subtotal (Tau ² = 0.17) | $\langle \rangle$ | 3.62 (1.67, 7.83) | 30.20 | | |
| Inestimable prediction distribution with <3 studies | | (- , -) | | | |
| Overall (Tau ² = 0.15) | | 6.82 (4.53, 10.25) | 100.00 | | |
| with estimated prediction interval | | (2.03, 22.87) | | | |
| I I 0.25 0.5 | | | | | |

T2DM incidence rate ratio associated with IFG 6.1 mmol/L threshold

Incidence rate ratio (IRR)

The comparison of geographic regions showed a lower IRR for Asia/Middle East as follows.

• Asia/Middle East (2 studies): T2DM developed in 21/1677 (1.3%) participants with IH and in 89/36,334 (0.2%) participants with normoglycaemia. The pooled IRR was 3.62 (95% CI 1.67 to 7.83).

• Australia/Europe/North America (4 studies): T2DM developed in 106/3438 (3.1%) participants with IH and in 99/20,246 (0.5%) participants with normoglycaemia. The pooled IRR was 8.55 (95% CI 6.37 to 11.48) with a 95% prediction interval ranging from 4.37 to 16.73.

IGT threshold

Twelve studies reported IRRs and defined IH using IGT. The studies included 18,468 participants with IH and 98,409 participants with normoglycaemia (Figure 22). T2DM developed in 947 (5.1%) participants with IH compared to 1147 (1.2%) in participants with normoglycaemia. The overall IRR was 4.48 (95% CI 3.69 to 5.44) with a 95% prediction interval ranging from 2.60 to 7.70.

Figure 22. IGT: impaired glucose tolerance; IRR: incidence rate ratio; n: number of cases; T: person-time in years

| Study | | | | | IRR (95% CI) | % Weight | Prediabetic n/T | Normoglycaemi n/T |
|---|--------|-------------|------------|------|-------------------|-------------|--------------------|----------------------|
| Asia/Middle East | | | | | | | | |
| Anjana 2015* | _ | - | | | 3.05 (2.37, 3.92) |) 14.97 | 86/1269 | 209/9398 |
| Li 2003* | _ | - | | | 3.23 (2.03, 5.16) | | 33/544 | 38/2026 |
| Mohan 2008 | | | _ | | 3.48 (1.98, 6.10) | | 15/247 | 64/3665 |
| Han 2017* | | | | | 4.32 (3.87, 4.82) | | 624/11744 | 657/53461 |
| Janghorbani 2015* | | | - | | 8.46 (4.42, 16.2) | | 26/1005 | 14/4578 |
| Subtotal (Tau ² = 0.05) | | $ \diamond$ | | | 3.93 (3.03, 5.10) | -, | | |
| with estimated prediction interval | | Ŭ | | | . (1.71, 9.02 | , | | |
| Australia/Europe/North America | | | | | | | | |
| Bonora 2011 | | _ _ | | | 3.93 (1.80, 8.59) |) 4.72 | 8/471 | 29/6704 |
| Valdes 2008* | | - | | | 4.20 (1.85, 9.49) | , 4.42 | 9/429 | 16/3200 |
| Soriguer 2008 | | | | | 4.32 (2.03, 9.19) | ,) 4.98 | 14/450 | 13/1806 |
| Rijkelijkhuizen 2007* | | | | _ | 8.28 (5.19, 13.2) | 0) 9.29 | 27/466 | 51/7286 |
| Forouhi 2007 | | + | - | | 9.37 (4.04, 21.7 | 1) 4.23 | 17/756 | 8/3333 |
| Subtotal (Tau ² = 0.05) | - | | > | _ | 5.93 (4.11, 8.57) | 27.64 | | |
| with estimated prediction interval | | | | | . (2.38, 14.8 | 1) | | |
| American Indians/Islands | | | | | | | | |
| Dowse 1991 | | | | | 3.86 (1.82, 8.21) |) 4.98 | 13/322 | 14/1339 |
| Wang 2011 | | -+ | | | 4.65 (3.10, 6.97) |) 10.69 | 75/765 | 34/1613 |
| Subtotal (Tau ² = 0) | | \langle | > | | 4.46 (3.12, 6.38) |) 15.68 | | |
| Inestimable prediction distribution with <3 studies | | | | | . (-,-) | | | |
| Overall (Tau ² =0.05) | | \$ | | | 4.48 (3.69, 5.44) |) 100.00 | | |
| with estimated prediction interval | | | | | . (2.60, 7.70 |) | | |
| | | | - | - | | | | |
| 1 | 2 | 4 | 8 | 16 | 32 | | | |
| | Incide | ence ra | te ratio (| IRR) | | | | |

T2DM incidence rate ratio associated with IGT threshold

The findings according to geographic regions were as follows.

• Asia/Middle East (5 studies): T2DM developed in 766/ 14,809 (5.2%) participants with IH and in 390/73,128 (0.5%) participants with normoglycaemia. The pooled IRR was 3.93 (95% CI 3.03 to 5.10) with a 95% prediction interval ranging from 1.71 to 9.02.

• Australia/Europe/North America (5 studies): T2DM developed in 75/2572 participants with IH and in 117/22,329 (0.5%) participants with normoglycaemia. The pooled IRR was 5.93 (95% CI 4.11 to 8.57) with a 95% prediction interval ranging from 2.38 to 14.81.

• American Indians/Islands (2 studies): T2DM developed in 88/1087 (8.1%) participants with IH and in 48/2952 (1.6%)

participants with normoglycaemia. The pooled IRR was 4.46 (95% CI 3.12 to 6.38).

Combined IFG and IGT

Nine studies used both IFG and IGT to define IH and reported IRRs. Of the 4470 participants with IH included in the studies, 551 (12.3%) developed T2DM compared with 1091 of the 90,072 (1.2%) participants with normoglycaemia (Figure 23). The overall IRR was 10.94 (95% CI 7.22 to 16.58) with 95% prediction interval ranging from 2.58 to 46.46.

Figure 23. IFG: impaired fasting glucose; IGT: impaired glucose tolerance; IRR: incidence rate ratio; n: number of cases; T: person-time in years

| | | | | | | | Prediabetic | Normoglycae |
|---|-----------|----------|---------------|---------------|--------------------------------------|---------|-------------|-------------|
| Study | | | | | IRR (95% CI) | % Weigh | t n/T | n/T |
| Asia/Middle East | | | 1 | | | | | |
| Li 2003 | | | - | | 5.96 (3.47, 10.24) | 11.09 | 20/179 | 38/2026 |
| Anjana 2015 | | | | | 6.01 (4.49, 8.04) | 12.81 | 58/434 | 209/9398 |
| Han 2017 | | -8 | H | | 9.31 (7.75, 11.19) | 13.31 | 138/1206 | 657/53461 |
| Janghorbani 2015 | | | | → | 51.95 (30.25, 89.21) | 11.10 | 214/1347 | 14/4578 |
| Subtotal (Tau ² = 0.46) | _ | $-\!\!<$ | \rightarrow | \rightarrow | 11.20 (5.59, 22.43) | 48.32 | | |
| with estimated prediction interval | | | | | (0.42, 300.27) | | | |
| Australia/Europe/North America | | | | | | | | |
| Soriguer 2008 | | | <u> </u> | | 9.17 (4.31, 19.52) | 9.42 | 14/212 | 13/1806 |
| Bonora 2011 | | | • | | 11.37 (5.38, 24.02) | 9.47 | 9/183 | 29/6704 |
| Rijkelijkhuizen 2007 | | | | | 16.05 (9.57, 26.92) | 11.28 | 20/178 | 51/7286 |
| Valdes 2008 | | - | | - | 19.05 (9.01, 40.26) | 9.47 | 12/126 | 16/3200 |
| Subtotal (Tau ² = 0) | | | \sim | | 13.92 (9.99, 19.40) | 39.65 | | |
| with estimated prediction interval | | | | | (6.71, 28.85) | | | |
| American Indians/Islands | | | | | | | | |
| Wang 2011 | | | | | 5.18 (3.42, 7.83) | 12.04 | 66/605 | 34/1613 |
| | | | | | | | | |
| Overall (Tau ² = 0.33) with estimated prediction interval | | | | _ | 10.94 (7.22, 16.58) (2.58, 46.46) | 100.00 | | |
| with estimated prediction merval | | | | | (2.00, 40.46) | | | |
| 0.25 | 1 | 4 | 16 | 64 | | | | |
| | Incidence | | | 04 | | | | |

T2DM incidence rate ratio associated with IFG and IGT thresholds

A lower pooled IRR was observed for the American Indians/Islands cohort compared to other cohorts as shown below.

• Asia/Middle East (4 studies): T2DM developed in 430/ 3166 (13.6%) participants with IH and in 918/69,463 (1.3%) participants with normoglycaemia. The pooled IRR was 11.20 (95% CI 5.59 to 22.43). Calculation of the 95% prediction interval did not provide a meaningful estimate.

• Australia/Europe/North America (4 studies): T2DM developed in 55/699 (7.9%) participants with IH and in 109/ 18,966 (0.6%) participants with normoglycaemia. The pooled IRR was 13.92 (95% CI 9.99 to 19.40) with a 95% prediction interval ranging from 6.71 to 28.85.

• American Indians/Islands (1 study): T2DM developed in 66/605 (10.9%) participants with IH and in 34/1613 (2.1%) participants with normoglycaemia. The pooled IRR was 5.18 (95% CI 3.42 to 7.83).

HbA1c 5.7% threshold only and the combination of HbA1c 5.7% threshold with IFG 5.6 mmol/L threshold

One study, Heianza 2012, reported using HbA1c_{5.7} only or the combination of IFG_{5.6} plus HbA1c_{5.7} to define IH at baseline (Figure 24).

Figure 24. IFG: impaired fasting glucose; HbA1c: glycosylated haemoglobin A1c; IRR: incidence rate ratio; n: number of cases; T: person-time in years

| | | | | | | | P | rediabetic | Normoglycaemic |
|-------------------------------------|------|----------|---------|--------|------|----------------------|----------|------------|----------------|
| Study, prediabetic origin | | 1 | | | | IRR (95% CI) | % Weight | n/T | n/T |
| Asia/Middle East | | | | | | | | | |
| Heianza 2012, HbA1c 5.7 | | | | - | | 6.62 (4.18, 10.49) | 33.16 | 30/1965 | 46/19961 |
| Heianza 2012, HbA1c 5.7 and IFG 5.6 | | | | | | 40.72 (29.30, 56.61) | 33.54 | 154/1641 | 46/19961 |
| | | | | | | 1 | | | |
| | 0.25 | 1 | 4 | 16 | 6 | 64 | | | |
| | Inc | idence i | rate ra | atio (| IRR) | | | | |

T2DM incidence rate ratio: HbA1c 5.7% threshold ± IFG 5.6 mmol/L threshold

T2DM developed in 30/1965 (1.5%) participants with IH defined using HbA1c_{5.7} only compared with 46/19,961 (0.2%) in participants with normoglycaemia. The IRR was 6.62 (4.18 to 10.49).

In the cohort with both HbA1c_{5.7} and IFG_{5.6}, T2DM developed in 154/1641 (9.4%) participants compared with 46/19961 (0.2%) in participants with normoglycaemia. The IRR was 40.72 (95% CI 29.30 to 56.61).

Odds ratio as the effect measure

IFG 5.6 mmol/L threshold

Twenty-one studies reported ORs and the IFG_{5.6} threshold for IH (Analysis 2.1). The length of follow-up ranged from 4 to 24 years (studies are ordered by ascending length of follow-up in Analysis 2.1). The studies included 9320 participants with IH and 38,327 participants with normoglycaemia. The overall OR was 4.15 (95% CI 2.75 to 6.28). The 95% prediction interval ranged from 0.54 to 32.00.

The comparison of geographic regions showed the following results (Analysis 2.1).

• Asia/Middle East (10 studies, 6359 participants with IH and 28,218 participants with normoglycaemia, 4 to 24 years' follow-up): the pooled OR was 2.94 (95% CI 1.77 to 4.86). The 95% prediction interval ranged from 0.43 to 19.93.

• Australia/Europe/North America (9 studies, 1949 participants with IH and 7920 participants with normoglycaemia, 4 to 12 years' follow-up): the pooled OR was 6.47 (95% CI 3.81 to 11.00). The 95% prediction interval ranged from 0.99 to 42.32.

• Latin America (1 study, 65 participants with IH and 1594 participants with normoglycaemia, 7 years' follow-up): the OR was 4.28 (95% CI 3.21 to 5.71).

• American Indians/Islands (1 study, 947 participants with IH and 595 participants with normoglycaemia, 4 years' follow-up): the OR was 3.12 (95% CI 2.31 to 4.21).

The test for subgroup differences did not indicate a significant subgroup effect (P = 0.07). However, two of the four subgroups had only one study each, so the validity of the analysis is uncertain. Furthermore, there is substantial heterogeneity between studies (Tau² = 0.65 and 0.59) within each of the other two subgroups, and the subgroup analysis does not appear to have explained heterogeneity.

IFG 6.1 mmol/L threshold

Fifteen studies reported ORs and the IFG_{6.1} threshold for IH at baseline (Analysis 2.2). The length of follow-up ranged from 3 to 24 years (studies are ordered by ascending length of follow-up in Analysis 2.2). The studies included 4574 participants with threshold for IH and 32,292 participants with normoglycaemia. The overall OR was 6.60 (95% CI 4.18 to 10.43). The 95% prediction interval ranged from 0.93 to 46.82.

The comparison between geographic regions showed the following results (Analysis 2.2).

• Asia/Middle East (7 studies, 3317 participants with IH and 25,604 participants with normoglycaemia, 3 to 24 years' followup): the pooled OR was 5.18 (95% CI 2.32 to 11.53). The 95% prediction interval ranged from 0.29 to 91.37.

• Australia/Europe/North America (7 studies, 1240 participants with IH and 5094 participants with normoglycaemia, 4 to 15 years' follow-up): the pooled OR was 8.69 (95% CI 4.95 to 15.24). The 95% prediction interval ranged from 1.20 to 62.69.

• Latin America (1 study, 17 participants with IH and 1594 participants with normoglycaemia, 7 years' follow-up): the OR was 3.73 (95% CI 2.18 to 6.38).

The test for subgroup differences did not indicate a significant subgroup effect (P = 0.10). However, one of the three subgroups had only one study, and there is substantial heterogeneity between studies (Tau² = 1.08 and 0.57) within each of the other two subgroups.

IGT

Twenty studies reported adjusted ORs and IGT for IH at baseline (Analysis 2.3). The length of follow-up ranged from 5 to 24 years (studies are ordered by ascending length of follow-up in Analysis 2.3). The studies included 3139 participants with IH and 18,413 participants with normoglycaemia. The overall OR was 4.61 (95% CI 3.76 to 5.64). The 95% prediction interval ranged from 2.10 to 10.13.

The comparison of geographic regions showed the following results (Analysis 2.3).

• Asia/Middle East (6 studies, 1226 participants with IH and 7417 participants with normoglycaemia, 5 to 24 years' followup): the pooled OR was 3.74 (95% CI 2.83 to 4.94). The 95% prediction interval ranged from 1.70 to 8.21.

• Australia/Europe/North America (11 studies, 1481 participants with IH and 7684 participants with normoglycaemia, 4 to 12 years' follow-up): the pooled OR was 5.20 (95% CI 3.62 to 7.45). The 95% prediction interval ranged from 1.50 to 18.09.

• Latin America (2 studies, 381 participants with IH and 3097 participants with normoglycaemia, 7 to 8 years' follow-up): the pooled OR was 4.94 (95% CI 3.15 to 7.76).

• American Indians/Islands (1 study, 51 participants with IH and 215 participants with normoglycaemia, 5 to 8 years' followup): the OR was 3.60 (95% CI 1.40 to 9.26).

The test for subgroup differences did not indicate a significant subgroup effect (P = 0.47). However, two of the four subgroups had only one or two studies, so the validity of the analysis is uncertain.

Combined IFG and IGT

Nine studies reported ORs and used both IFG and IGT for defining IH at baseline (Analysis 2.4). The length of follow-up ranged from 5 to 24 years (studies are ordered by ascending length of follow-up in Analysis 2.4). The studies included 652 participants with IH and 9004 participants with normoglycaemia. The overall OR was 13.14 (95% CI 7.41 to 23.30). The 95% prediction interval ranged from 1.84 to 93.66. The comparison of geographic regions showed the following results (Analysis 2.4).

• Asia/Middle East (3 studies, 498 participants with IHT and 3704 participants with normoglycaemia, 5 to 24 years' followup): the pooled OR was 6.99 (95% CI 3.09 to 15.83). Calculation of the 95% prediction interval did not provide a meaningful estimate.

• Australia/Europe/North America (6 studies, 154 participants with IH and 5300 participants with normoglycaemia, 6 to 12 years' follow-up): the pooled OR was 20.95 (95% CI 12.40 to 35.40). The 95% prediction interval ranged from 4.93 to 89.05.

The OR for the Australia/Europe/North America cohort of studies appeared to be higher compared with the Asia/Middle East cohort.

HbA1c 5.7% threshold

Three studies reported ORs and HbA1c_{5.7} threshold for IH at baseline (Analysis 2.5). The length of follow-up ranged from 6 to 10 years (studies are ordered with ascending length of follow-up in Analysis 2.5). The studies included 906 participants with IH and 2562 participants with normoglycaemia. The overall OR was 4.43 (95% CI 2.20 to 8.88). Calculation of the 95% prediction interval did not provide a meaningful estimate.

The results by geographic region are as follows (Analysis 2.5).

• Asia/Middle East (1 study, 675 participants with IH and 462 participants with normoglycaemia, 6 years' follow-up): the OR was 4.54 (95% CI 2.65 to 7.78).

• Australia/Europe/North America (2 studies, 231 participants with IH and 2100 participants with normoglycaemia, 7 to 10 years' follow-up): the pooled OR was 4.38 (95% CI 1.36 to 14.15).

HbA1c 6.0% threshold

Three studies reported ORs and the HbA1 $c_{6.0}$ threshold for IH at baseline (Analysis 2.6). The length of follow-up ranged from three to five years. The studies included 1594 participants with IH and 16,723 participants with normoglycaemia. The overall OR was 12.79 (95% CI 4.56 to 35.85). Calculation of the 95% prediction interval did not provide a meaningful estimate.

The comparison of geographic regions showed the following results (Analysis 2.6).

• Asia/Middle East (1 study, 1103 participants with IH and 10,763 participants with normoglycaemia, 5 years' follow-up): the OR was 23.20 (95% CI 18.70 to 28.78).

• Australia/Europe/North America (1 study, 370 participants with IH and 5365 participants with normoglycaemia, 3 years' follow-up): the OR was 15.60 (95% CI 6.90 to 35.27).

• American Indians/Islands (1 study, 121 participants with IH and 595 participants with normoglycaemia, 4 years' follow-up): the OR was 5.89 (95% CI 4.23 to 8.20).

The OR for the Asia/Middle East and Australia/Europe/North America studies appeared higher compared with the American Indians/Islands study.

Combination of HbA1c 5.7% threshold with IFG 5.6 mmol/L threshold

Two studies defined IH using a combination of HbA1c_{5.7} and IFG_{5.6} at baseline and reported ORs (Analysis 2.7). The length of follow-up ranged from five to seven years (studies are ordered by ascending length of follow-up in Analysis 2.7). The studies included 2120 participants with IH and 11,886 participants with normoglycaemia. The pooled OR was 35.91 (95% CI 20.43 to 63.12).

The findings for each geographic region are as follows (Analysis 2.7).

• Asia/Middle East (1 study, 1951 participants with IH and 10,761 participants with normoglycaemia, 5 years' follow-up): the OR was 46.70 (95% CI 33.60 to 64.91).

• Australia/Europe/North America (1 study, 169 participants with IH and 1125 participants with normoglycaemia, 7 years' follow-up): the OR was 26.20 (95% CI 16.30 to 42.11).

Subgroup and sensitivity analyses

There were not enough data to perform subgroup analyses by age or sex. The special group of children and adolescents is reported under the headings corresponding to the association between IH and T2DM incidence and regression to normoglycaemia.

Sensitivity analyses for risk of bias were not meaningful because of the diversity in measurement of T2DM incidence, definitions of IH, and follow-up periods. The analysis of adequate adjustment for confounding factors in studies reporting HRs may have provided interesting information, but there were not enough data to analyse the impact of at least four or five well-known covariates influencing the relationship between prognostic factor and T2DM incidence. There were no very large studies including participants with IH at baseline.

Overview of complete data set and certainty of the evidence

Table 1 provides a succinct overview of the overall prognosis of people with IH as well as regression from IH to normoglycaemia over 1 to 20 years of follow-up.

Table 2 provides a succinct overview of IH compared with normoglycaemia as a prognostic factor for developing T2DM according to geographic regions/special populations and type of outcome measurement.

Figure 25 shows the overall prognosis of IH as measured by cumulative incidence over different follow-up periods and across all populations, as well as regression from IH to normoglycaemia.

Figure 25. Overall prognosis of people with intermediate hyperglycaemia (cumulative type 2 diabetes incidence and regression to normoglycaemia) associated with measures of intermediate hyperglycaemia HbA1c5.7/HbA1c6.0: glycosylated haemoglobin A1c 5.7%/6.0% threshold; IFG5.6/6.1: impaired fasting glucose 5.6/6.1 mmol/L threshold; IGT: impaired glucose tolerance

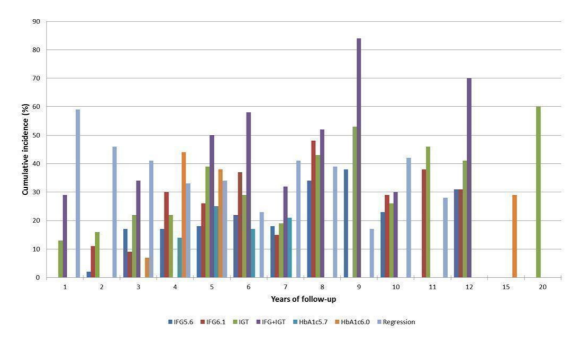
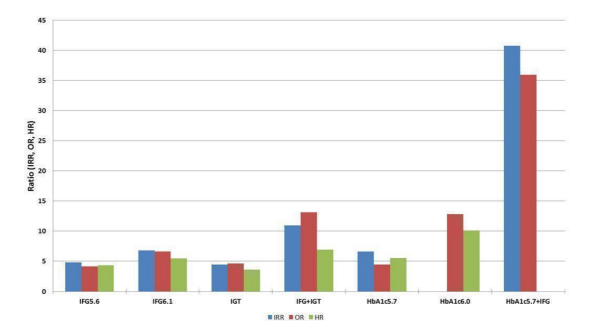


Figure 26 shows IH versus normoglycaemia as a prognostic factor for developing T2DM measured by IRR, OR or HR across all populations.

Figure 26. Intermediate hyperglycaemia versus normoglycaemia as a prognostic factor for developing type 2 diabetes (associated with different measures and relative risks of intermediate hyperglycaemia)
 HbA1c5.7/HbA1c6.0: glycosylated haemoglobin A1c 5.7%/6.0% threshold; IFG5.6/6.1: impaired fasting glucose
 5.6/6.1 mmol/L threshold; IGT: impaired glucose tolerance; IRR: incidence rate ratio; OR: odds ratio; HR: hazard ratio



Taking into account all follow-up times and all populations, the percentages of people with IH *not* developing T2DM over time (i.e. either regressing to normoglycaemia or remaining 'prediabetic') were as follows (see Appendix 11): IFG_{5.6} cohorts, 79.2%; IFG_{6.1} cohorts, 75.4%; IGT cohorts, 66.7%; combined IFG and IGT cohorts, 57.2%; HbA1c_{5.7} cohorts, 79.7%; and HbA1c_{6.0} cohorts, 69.0%.

For overall prognosis, we started with high-certainty evidence because prospective cohort studies represent an adequate study design to investigate overall prognosis. However, we downgraded the certainty of the evidence to moderate because of imprecise results for most definitions of IH (Summary of findings for the main comparison).

We considered the overall certainty of the evidence for the prognostic factor IH versus normoglycaemia as low (Summary of findings 2; Summary of findings 3; Summary of findings 4; Summary of findings 5; Summary of findings 6; Summary of findings 7). We started with a high level of evidence because most included studies were phase 2 explanatory studies, defined as studies that aimed to confirm independent associations between the prognostic factor and the outcome (Huguet 2013). We downgraded the evidence for all IH measurements to low, first one level due to study limitations because many studies did not adequately adjust for confounders (only six studies used the covariate core set of age, sex, anthropometric measures and physical activity for adjustments in multivariable regression analyses - Bonora 2011; Derakhshan 2016; Forouhi 2007; Han 2017; Kim 2016a; Yeboah 2011). Furthermore, we downgraded one level for imprecision/inconsistency (wide 95% CIs/wide 95% prediction intervals, sometimes ranging from negative to positive prognostic factor to outcome associations).

ADDITIONAL SUMMARY OF FINDINGS [Explanation]

Outcome: development of T2DM

Prognostic factor: intermediate hyperglycaemia versus normoglycaemia as measured by IFG_{5.6}

| | ······································ | | | | | | | |
|----------------------------|--|---|--|--|--|--|--|--|
| No of studies | No of participants with interme- diate hyperglycaemia | Geographic region/special popu- lation | Estimated effect (95% CI) [95% prediction interval] | Overall certainty of the evidence (GRADE) ^a | | | | |
| HR: 4 IRR: 6 OR: 10 | HR: 2385 IRR: 15,661 OR: 6359 | Asia/Middle East | HR: 5.07 (3.41-4.86) [1.07-24.02] IRR: 5.23 (3.77-7.25) [1.72-15. 89] OR: 2.94 (1.77-4.86) [0.43-19.93] | | | | | |
| HR: 3 IRR: 3 OR: 9 | HR: 5685 IRR: 6322 OR: 1949 | Australia/Europe/North America | HR: 4.15 (1.24-13.9) [N/M] IRR: 4.96 (3.25-7.57) [0.32-77. 24] OR: 6.47 (3.81-11.00) [0.99-42. 32] | _ | | | | |
| HR: 0 IRR: 0 OR: 1 | HR: 0 IRR: 0 OR: 65 | Latin America | HR: NA IRR: NA OR: 4.28 (3.21-5.71) | | | | | |
| HR: 1 IRR: 1 OR: 1 | HR: 947 IRR: 2374 OR: 947 | American Indians/Islands | HR: 2.38 (1.85-3.06) IRR: 2.74 (1.88-3.99) OR: 3.12 (2.31-4.21) | | | | | |
| HR: 8 IRR: 10 OR: 21 | HR: 9017 IRR: 24,357 OR: 9320 | Overall | HR: 4.32 (2.61-7.12) [0.75-25.0] IRR: 4.81 (3.67-6.30) [1.95-11. 83] OR: 4.15 (2.75-6.28) [0.53-32.4] | | | | | |

CI: confidence interval; HR: hazard ratio; IFG_{5.6}: impaired fasting glucose 5.6 mmol/L threshold; IRR: incidence rate ratio; NA: not applicable; N/M: fewer than 3 studies or calculation of the 95% prediction interval did not provide a meaningful estimate; OR: odds ratio; T2DM: type 2 diabetes mellitus.

GRADE Working Group grades of evidence

High quality: further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate. Very low quality: we are very uncertain about the estimate.

^aWith phase 2 explanatory studies aiming to confirm independent associations between the prognostic factor and the outcome, GRADE starts with 'high quality' (Huguet 2013). We assumed the GRADE factor publication bias was inherent with this type of research (phase 2 design), so we did not use it as a potential downgrading factor

^bDowngraded by one level because of study limitations (many studies did not adequately adjust for confounders, if at all) and by one level because of imprecision (CIs were wide) and inconsistency (wide 95% prediction intervals sometimes ranging from negative to positive prognostic factor to outcome associations)

| rognostic factor: inter | nediate hyperglycaemia as measured by IFC | 6.1 | | |
|---------------------------|--|---|--|--|
| No of studies | No of participants with interme- diate hyperglycaemia | Geographic region/special popu- lation | Estimated effect (95% CI) [95% prediction interval] | Overall certainty of the evidence (GRADE) a |
| HR: 5 IRR: 2 OR: 7 | HR: 1054 IRR: 1677 OR: 3317 | Asia/Middle East | HR: 10.55 (3.61-30.81) [N/M] IRR: 3.62 (1.67-7.83) [N/M] OR: 5.18 (2.32-11.53) [0.29-91. 37] | ⊕⊕⊖⊖ Low ^b |
| HR: 4 IRR: 4 OR: 7 | HR: 1736 IRR: 3438 OR: 1240 | Australia/Europe/North America | HR: 3.30 (2.32-4.67) [0.84-12.99] IRR: 8.55 (6.37-11.48) [4.37-16. 73] OR: 8.69 (4.95-15.24) [1.20-62. 69] | |
| HR: 0 IRR: 0 OR: 1 | HR: 0 IRR: 0 OR: 17 | Latin America | HR: NA IRR: NA OR: 3.73 (2.18-6.38) | |
| HR: 0 IRR: 0 OR: 0 | HR: 0 IRR: 0 OR: 0 | American Indians/Islands | HR: NA IRR: NA OR: NA | |
| HR: 9 IRR: 6 OR: 15 | HR: 2818 IRR: 5115 OR: 4574 | Overall | HR: 5.47 (3.50-8.54) [1.09-27.56] IRR: 6.82 (4.53-10.25) [2.03-22. 87] OR: 6.60 (4.18-10.43) [0.93-46. 82] | |

CI: confidence interval; HR: hazard ratio; IFG_{6.1}: impaired fasting glucose 6.1 mmol/L threshold; IRR: incidence rate ratio; NA: not applicable; N/M: fewer than 3 studies or calculation of the 95% prediction interval did not provide a meaningful estimate; OR: odds ratio; T2DM: type 2 diabetes mellitus.

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Moderate quality: further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate. Very low quality: we are very uncertain about the estimate.

^aWith phase 2 explanatory studies aiming to confirm independent associations between the prognostic factor and the outcome, GRADE starts with 'high quality' (Huguet 2013). We assumed the GRADE factor publication bias was inherent with this type of research (phase 2 design), so we did not use it as a potential downgrading factor

^bDowngraded by one level because of study limitations (many studies did not adequately adjust for confounders, if at all) and by one level because of imprecision (CIs were wide) and inconsistency (wide 95% prediction intervals sometimes ranging from negative to positive prognostic factor to outcome associations)

| No of studies | No of participants with interme- diate hyperglycaemia | Geographic region/special popu- lation | Estimated effect (95% CI) [95% prediction interval] | Overall certainty of the eviden $(GRADE)^a$ |
|----------------------------|--|---|---|---|
| HR: 3 IRR: 5 OR: 6 | HR: 1780 IRR: 14,809 OR: 1226 | Asia/Middle East | HR: 4.48 (2.81-7.15) [N/M] IRR: 3.93 (3.03-5.10) [1.71-9.02] OR: 3.74 (2.83-4.94) [1.70-8.21] | ⊕⊕⊖⊖ Low ^b |
| HR: 2 IRR: 5 OR: 11 | HR: 2230 IRR: 2572 OR: 1481 | Australia/Europe/North America | HR: 2.53 (1.52-4.19) [N/M] IRR: 5.93 (4.11-8.57) [2.38-14. 81] OR: 5.20 (3.62-7.45) [1.50-18.09] | |
| HR: 0 IRR: 0 OR: 2 | HR: 0 IRR: 0 OR: 381 | Latin America | HR: NA IRR: NA OR: 4.94 (3.15-7.76) [N/M] | _ |
| IRR: 2 OR: 1 HR: 0 | IRR: 1087 OR: 51 HR: 0 | American Indians/Islands | IRR: 4.46 (3.12-6.38) [N/M] OR: 3.60 (1.40-9.26) HR: NA | |
| HR: 5 IRR: 12 OR: 20 | HR: 4010 IRR: 18,468 OR: 3139 | Overall | HR: 3.61 (2.31-5.64) [0.69-18.97] IRR: 4.48 (3.59-5.44) [2.60-7.70] OR: 4.61 (3.76-5.64) [2.10-10.13] | |

CI: confidence interval; HR: hazard ratio; IGT: impaired glucose tolerance; IRR: incidence rate ratio; NA: not applicable; N/M: fewer than 3 studies or calculation of the 95% prediction interval did not provide a meaningful estimate; T2DM: type 2 diabetes mellitus.

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Moderate quality: further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Very low quality: we are very uncertain about the estimate.

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^aWith phase 2 explanatory studies aiming to confirm independent associations between the prognostic factor and the outcome, GRADE starts with 'high quality' (Huguet 2013). We assumed the GRADE factor publication bias was inherent with this type of research (phase 2 design), so we did not use it as a potential downgrading factor

^bDowngraded by one level because of study limitations (many studies did not adequately adjust for confounders, if at all) and by one level because of imprecision (CIs were wide) and inconsistency (wide 95% prediction intervals sometimes ranging from negative to positive prognostic factor to outcome associations)

| No of studies | No of participants with interme- diate hyperglycaemia | Geographic region/special popu- lation | Estimated effect (95% CI) [95% prediction interval] | Overall certainty of the evidence (GRADE) ^a |
|--------------------------|--|---|---|--|
| HR: 3 IRR: 4 OR: 3 | HR: 461 IRR: 3166 OR: 498 | Asia/Middle East | HR: 10.20 (5.45-19.09) [N/M] IRR: 11.20 (5.59-22.43) [N/M] OR: 6.99 (3.09-15.83) [N/M] | ⊕⊕⊖⊖ Low ^b |
| HR: 1 IRR: 4 OR: 6 | HR: 221 IRR: 699 OR: 154 | Australia/Europe/North America | HR: 3.80 (2.30-6.28) [N/M] IRR: 13.92 (9.99-19.40) [6.71-28. 85] OR: 20.95 (12.40-35.40) [4.93- 89.05] | _ |
| HR: 0 IRR: 0 OR: 0 | HR: 0 IRR: 0 OR: 0 | Latin America | HR: NA IRR: NA OR: NA | _ |
| HR: 1 IRR: 1 OR: 0 | HR: 356 IRR: 605 OR: 0 | American Indians/Islands | HR: 4.06 (3.05-5.40) IRR: 5.18 (3.42-7.83) OR: NA | _ |
| HR: 5 IRR: 9 OR: 9 | HR: 1038 IRR: 4470 OR: 652 | Overall | HR: 6.90 (4.15-11.45) [1.06-44. 95] IRR: 10.94 (7.22-16.58) [2.58-46. 46] OR: 13.14 (7.41-23.30) [1.84-93. 66] | |

CI: confidence interval; HR: hazard ratio; IFG: impaired fasting glucose; IGT: impaired glucose tolerance; IRR: incidence rate ratio; NA: not applicable; N/M: fewer than 3 studies or calculation of the 95% prediction interval did not provide a meaningful estimate; OR: odds ratio; T2DM: type 2 diabetes mellitus.

GRADE Working Group grades of evidence

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Moderate quality: further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Very low quality: we are very uncertain about the estimate.

^aWith phase 2 explanatory studies aiming to confirm independent associations between the prognostic factor and the outcome, GRADE starts with 'high quality' (Huguet 2013). We assumed the GRADE factor publication bias was inherent with this type of research (phase 2 design), so we did not use it as a potential downgrading factor

^bDowngraded by one level because of study limitations (many studies did not adequately adjust for confounders, if at all) and by one level because of imprecision (CIs were wide) and inconsistency (wide 95% prediction intervals)

| Outcome: development of T2DM Prognostic factor: intermediate hyperglycaemia as measured by HbA1c _{5.7} | | | | |
|--|--|---|---|--|
| No of studies | No of participants with interme- diate hyperglycaemia | Geographic region/special popu- lation | Estimated effect (95% CI) [95% prediction interval] | Overall certainty of the evidence (GRADE) ^a |
| HR: 3 IRR: 1 OR: 1 | HR: 3196 IRR: 1965 OR: 675 | Asia/Middle East | HR: 7.21 (5.14-10.11) [0.81-64. 52] IRR: 6.62 (4.18-10.49) [N/M] OR: 4.54 (2.65-7.78) [N/M] | ⊕⊕⊖⊖ Low ^b |
| HR: 1 IRR: 0 OR: 2 | HR: 2027 IRR: 0 OR: 231 | Australia/Europe/North America | HR: 2.71 (2.48-2.96) [N/M] IRR: NA OR: 4.38 (1.36-14.15) [N/M] | |
| HR: 0 IRR: 0 OR: 0 | HR: 0 IRR: 0 OR: 0 | Latin America | HR: NA IRR: NA OR: NA | _ |
| HR: 0 IRR: 0 OR: 0 | HR: 0 IRR: 0 OR: 0 | American Indians/Islands | HR: NA IRR: NA OR: NA | |
| HR: 4 IRR: 1 OR: 3 | HR: 5223 IRR: 1965 OR: 906 | Overall | HR: 5.55 (2.77-11.12) [0.23-141. 18] IRR: 6.62 (4.18-10.49) [N/M] OR: 4.43 (2.20-8.88) [N/M] | |

Cl: confidence interval; HbA1c_{5.7}: glycosylated haemoglobin A1c 5.7% threshold; HR: hazard ratio; IRR: incidence rate ratio; NA: not applicable; N/M: fewer than 3 studies or calculation of the 95% prediction interval did not provide a meaningful estimate; OR: odds ratio; T2DM: type 2 diabetes mellitus.

GRADE Working Group grades of evidence

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Moderate quality: further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Very low quality: we are very uncertain about the estimate.

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^aWith phase 2 explanatory studies aiming to confirm independent associations between the prognostic factor and the outcome, GRADE starts with 'high quality' (Huguet 2013). We assumed the GRADE factor publication bias was inherent with this type of research (phase 2 design), so we did not use it as a potential downgrading factor

^bDowngraded by one level because of study limitations (many studies did not adequately adjust for confounders, if at all) and by one level because of imprecision (CIs were wide) and inconsistency (95% prediction intervals sometimes ranging from negative to positive prognostic factor to outcome associations)

| Outcome: development of T2DM Prognostic factor: intermediate hyperglycaemia as measured by HbA1c _{6.0} | | | | |
|--|--|---|--|--|
| No of studies | No of participants with interme- diate hyperglycaemia | Geographic region/special popu- lation | Estimated effect (95% CI) [95% prediction interval] | Overall certainty of the evidence (GRADE) ^a |
| HR: 2 IRR: 0 OR: 1 | HR: 1040 IRR: 0 OR: 370 | Australia/Europe/North America | HR: 5.09 (1.69-15.37) [N/M] IRR: NA OR: 15.60 (6.90-35.27) [N/M] | ⊕⊕⊖⊖ Low ^b |
| HR: 4 IRR: 0 OR: 1 | HR: 3492 IRR: 0 OR: 1103 | Asia/Middle East | HR: 13.12 (4.10-41.96) [N/M] IRR: NA OR: 23.20 (18.70-28.78) [N/M] | |
| HR: 0 IRR: 0 OR: 0 | HR: 0 IRR: 0 OR: 0 | Latin America | HR: NA IRR: NA OR: NA | _ |
| IRR: 0 OR: 1 HR: 0 | IRR: 0 OR: 121 HR: 0 | American Indians/Islands | IRR: NA OR: 5.89 (4.23-8.20) [N/M] HR: NA | _ |
| HR: 6 IRR: 0 OR: 3 | HR: 4532 IRR: 0 OR: 1594 | Overall | HR: 10.10 (3.59-28.43) [N/M] IRR: NA OR: 12.79 [4.56-35.85] [N/M] | |

Cl: confidence interval; HbA1c_{6.0}: glycosylated haemoglobin A1c 6.0% threshold; HR: hazard ratio; IRR: incidence rate ratio; NA: not applicable; N/M: fewer than 3 studies or calculation of the 95% prediction interval did not provide a meaningful estimate; OR: odds ratio; T2DM: type 2 diabetes mellitus.

GRADE Working Group grades of evidence

High quality: further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Very low quality: we are very uncertain about the estimate.

Development of type 2 diabetes mellitus in people with intermediate hyperglycaemia (Review) Copyright © 2018 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.

^aWith phase 2 explanatory studies aiming to confirm independent associations between the prognostic factor and the outcome, GRADE starts with 'high quality' (Huguet 2013). We assumed the GRADE factor publication bias was inherent with this type of research (phase 2 design), so we did not use it as a potential downgrading factor

^bDowngraded by one level because of study limitations (many studies did not adequately adjust for confounders, if at all) and by one level because of imprecision (most CIs were wide)

DISCUSSION

Summary of main results

We included 103 prospective cohort studies from many parts of the world evaluating people with IH, usually defined using the IFG_{5.6} or IFG_{6.1} threshold, IGT, combined IFG/IGT or elevated HbA1c. However, we did not identify studies involving black Africans or Eastern Europeans. Participants were of Australian, European or North American origin in 41 studies; primarily of Latin American origin in 7 studies; Asian or Middle Eastern origin in 50 studies; American Indians in 3 studies; Mauritians in 1 study; and Nauruans in 1 study. Six studies included children, adolescents or both. Ninety-three studies contributed data to estimate the overall prognosis of people with IH, and 52 studies evaluated baseline glycaemic status as a prognostic factor by comparing an IH cohort with a normoglycaemic cohort.

Cumulative incidence of T2DM for the IFG_{5.6} threshold, the IFG_{6.1} threshold, IGT, combined IFG/IGT and elevated HbA1c, showed increasing percentages over follow-up time; however, there was no clear linear increase over time. Regression rates to normoglycaemia, though decreasing over follow-up, showed fluctuations and no clear linear decrease over time. The estimates of the prognostic effect of IH versus normoglycaemia were comparable when using HR, IRR or OR across the different definitions of IH. There was no clear pattern of risk differences between geographic regions.

Overall completeness and applicability of evidence

A limiting factor of our review was that most studies took place in Asia, the Middle East, Australia, Western Europe and North America, affecting the generalisability of findings to other populations residing in Africa and Eastern Europe. We are also aware that categorising the included studies based on region or 'ethnicity' has deficiencies with regard to clearly delineating study participants. The complicated interplay of factors like genetics, diets, and changing environmental and social conditions, among others, makes it virtually impossible to achieve a generally accepted categorisation. We chose an approach based primarily on geographic location because we thought that most readers would be interested in having a broad overview of any potential differences in T2DM incidence based on this characteristic. At the same time, we tried not to overload the reader with too much information by fragmenting our dataset into all the different countries or into more precisely defined 'ethnicities', since some investigators even reported several 'ethnic' subgroups within a single study cohort. However, we do provide detailed information, when available, in our appendices to enable the interested reader to identify studies according to whatever combination of factors seems of value to generate hypotheses of potential differences between the diverse study groups.

Only six studies addressed the overall prognosis of IH in 495 children or adolescents, with approximately 50% originating from high-risk American Indian cohorts, also affecting the applicability of findings to other populations. No data were available on the prognostic factor of IH versus normoglycaemia for children or adolescents. Most studies determined the glycaemic status of participants at baseline and follow-up on the basis of a single FPG, glucose tolerance test or HbA1c. Therefore, participants may have been misclassified at baseline, follow-up or both in either direction. Interestingly, 93 studies provided data on overall prognosis of IH, but only 49 studies published information on regression from IH to normoglycaemia.

Certainty of the evidence

To our knowledge there is no validated risk of bias tool for studies addressing overall prognosis. Moreover, information on some applicable risk of bias domains of the QUIPS tools were limited. However, as illustrated in Figure 25, there was a wide fluctuation between the various definitions of IH as well as no linear increase in T2DM incidence over time of follow-up. Of note, regression rates to normoglycaemia were also high, even after more than five years of follow-up, emphasising that transition from IH to T2DM might be an intermediate state (Taylor 2017).

The certainty of the evidence for the overall prognosis of IH was moderate due to imprecise results for most IH definitions. The certainty of the evidence for the prognostic factor of IH versus normoglycaemia was low, mainly because most studies did not adjust for confounders known to be independently associated with T2DM incidence and due to substantial imprecision (wide 95% CIs) and inconsistency (wide 95% prediction intervals). However, the results of the six studies that adjusted for sex, anthropometric measures and physical activity were similar to the rest of the prospective cohort studies.

Limitations in the review process

As described in the Methods section, it was difficult to define a reliable search strategy, which probably holds true for many systematic reviews of prognostic studies. We noted that when checking other systematic reviews on the topic and the references of the included studies, around one third of our included studies were identified through reference checking. However, using PubMed's 'similar articles' algorithm did not yield new studies but did help us identify 13 secondary publications of studies we had already included. The 103 prospective cohort studies included in this review represent by far the largest amount of data synthesised on the overall prognosis of IH and the impact of IH versus normoglycaemia as a prognostic factor for T2DM development. We did not contact study authors for additional information, mainly for logistical reasons but also because we anticipated poor response, since many studies were published long ago. Moreover, retrieval of additional information, often demanding recalculations, would

have imposed a considerable burden on study authors.

During the review process, the need to establish a database of cohort studies specifying details on prognostic factors and outcomes, amongst other things, became clear. Many large cohort studies investigate the association of a great number of prognostic factors with yet another large number of outcomes. These data may only be detected through a detailed analysis of the full text (especially tables and figures). It is evident that screening titles and abstracts will miss this information.

We did not include participants of randomised controlled trials. Though potentially some trials with longer time of follow-up could provide additional data, we decided not to include information from intervention trials at this stage on theoretical grounds, as any intervention will interfere with peoples' lives, as opposed to demonstrating the natural progression of a disorder. In addition, we are conducting a series of Cochrane Reviews on interventions for people with IH and may integrate these data in a later update of this review (Hemmingsen 2016a; Hemmingsen 2016b; Hemmingsen 2016c).

Agreements and disagreements with other reviews

Gerstein 2007 is a widely cited review including 21 cohort studies and nine randomised controlled trials published between 1979 and 2004. The review authors annualised T2DM incidence rates, which varied from 5% to 10%. Their relative risks for T2DM incidence of 6.35 in people with IGT, 4.66 in people with IFG and 12.1 with both IFG and IGT were higher but comparable to our HR data. We did not annualise incidence rates because with pronounced fluctuations between regression and development of T2DM, assumptions to establish a model for annualising incidence data over prolonged period of times appeared too strong. Zhang 2010 examined ranges of HbA1c and also associated these with annualised diabetes incidences. The results of seven included studies reporting HbA1c categories showed an increase in T2DM incidence across an HbA1c range from 5.0% to 6.5%. No metaanalysis was performed. Our results also showed increased T2DM incidence when the threshold of the HbA1c value at baseline was raised from 5.7% to 6.0%. Morris et al. performed a meta-analysis of prospective observational studies in which participants had IH at baseline (Morris 2013). The review included 70 studies and estimated pooled incidence rates using IFG (35.5 incident cases per 1000 person-years as defined by ADA and 47.4 incident cases per 1000 person-years as defined by WHO, 11 and 34 studies, respectively), IGT (45.5 incident cases per 1000 person-years, 46 studies) and IFG/IGT (70.4 incident cases per 1000 person-years, 15 studies) definitions for IH. Elevated HbA1c was associated with a pooled incidence rate of 35.6 per 1000 person-years. Similar to our results, the review found that progression rates to T2DM differed by definition of IH.

AUTHORS' CONCLUSIONS

Implications for practice

Our systematic review on the development of type 2 diabetes mellitus (T2DM) in people with intermediate hyperglycaemia (IH) or 'prediabetes' identified several uncertainties: glycaemic status can be measured in various ways, with IH usually defined by impaired fasting glucose (IFG) with cut-off levels of 5.6 mmol/L or 6.1 mmol/L, by impaired glucose tolerance (IGT) or by elevated HbA1c levels with thresholds of 5.7% or 6.0%. These definitions imply specific settings and demands on resources. It is likely that the accuracy of information provided by the tests will need to be balanced against the time, effort and cost required to capture them. IFG measurement is cumbersome because of the need for overnight fasting. HbA1c measurement is resource intensive and must be standardised, taking into account potential interference factors like anaemia, haemoglobinopathy or renal insufficiency. IGT measurement is cumbersome and also resource intensive. Overall, the certainty of the evidence was low for IH versus normoglycaemia, mainly because many of the prospective cohort studies did not adequately investigate other factors or covariates which could have confounded or modified the prognostic effect of glycaemic status on T2DM incidence. Moreover, results varied widely, making it difficult to specify the best definition for IH. The certainty of the evidence for the overall prognosis of people with IH as well as regression from IH to normoglycaemia was moderate because of imprecise results for most intermediate hyperglycaemia definitions. With increasing years of follow-up, T2DM incidence increased, but regression from IH to normoglycaemia was also high. There was no clear pattern of geographical differences; again, studies showed wide variation depending on the definition of IH, mode of measurement and length of follow-up. Due to the fluctuating stages of normoglycaemia, IH and T2DM, which might show transition from one stage to another in both directions and even after years of follow-up, practitioners should be careful about the potential implications of any active intervention for people 'diagnosed' with IH.

Implications for research

Future prospective cohort studies should address the consequences of IH to minimise secondary analyses of cohort studies where investigators synthetically form a subgroup of people with prediabetes, as such analyses are suboptimal. There is an urgent need for data from Eastern Europe and Africa to enable assessment of the prognostic value of IH in these regions, and for prospective cohort studies designed to examine the relationship between IH and normoglycaemia, T2DM incidence and the development of diabetic complications. The studies should adjust for confounding using important, well-defined factors such as age, sex, 'ethnicity', anthropometric measures and physical activity. Also, studies

should be adequately powered and analysed using suitable statistical techniques such as time-dependent regression methods. There is a need for a database of cohort studies with details on all analysed prognostic factor to outcome associations because many cohort studies start with general questions like the influence of various risk factors on cardiovascular disease, and specific factors may only be identified by investigating the full text. The nature of these investigations means that search strategies basing their retrieval on titles and abstracts only will not be sufficient to identify these studies.

ACKNOWLEDGEMENTS

The World Health Organization (WHO) funded this review.

We thank Megan Harris for the excellent copy-editing of our review. We thank Nuala Livingstone, Kerry Dwan, Toby Lasserson, Alex Sutton and especially Carl Moons for their distinguished peer-reviewing which definitely raised the quality of our review.

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* Indicates the major publication for the study

CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

Admiraal 2014

| Name of study | Surinamese in the Netherlands: study on health and ethnicity/healthy life in an urban setting (SUNSET/HELIUS) |
|--------------------|--|
| Inclusion criteria | Participants of 2 studies (SUNSET and HELIUS), Surinamese and ethnic Dutch, south- east Amsterdam, aged 35-60 years with completed interviews and medical examinations at baseline and follow-up |
| Exclusion criteria | Missing FPG data, diabetes |
| Notes | Baseline data for total cohort included in the analyses (N = 456): South-Asian Surinamese (N = 90), African Surinamese (N = 190), ethnic Dutch (N = 176) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|---|
| Study participation: description of source population or population of interest | Low risk | Surinamese in the Netherlands study |
| Study participation: description of gly- caemic status at baseline | Low risk | 456 participants available for analysis; table 1 specifies people with IFG _{5.7} |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Random sample of 2975 Surinamese and ethnic Dutch individuals, aged 35-60, drawn from the population register of 2 neighbourhoods in southeast Amsterdam |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria specified |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Those who were lost to follow-up were younger, had a higher BMI and greater waist circumference, a higher FPG and more often had baseline IFG than those with follow-up data available after 10 years |
| Study attrition: reasons for loss to follow- up provided | Low risk | 777/1444 lost to follow-up (moved out- side of Amsterdam, declined to participate, died, non-response); figure S1 |

Admiraal 2014 (Continued)

| Study attrition: adequate description of participants lost to follow-up | Low risk | Reported in Table S2 |
|---|--------------|--|
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | See above |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | IFG |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | FPG measurement by G6PD test |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG 5.7-6.9 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; HbA1c \geq 6.5; self-reported T2DM |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Reliable measurement |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Limited number of confounders measured |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |

Admiraal 2014 (Continued)

| Study confounding: important potential confounders accounted for in study design | Low risk | Adjustment for sex, age, BMI and change in BMI after 10 years |
|---|----------|--|
| Study confounding: important potential confounders accounted for in the analysis | Low risk | Unadjusted and adjusted analyses |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, odds ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multivariate logistic regression |
| Aekplakorn 2006 | | |

| Name of study | None |
|--------------------|--|
| Inclusion criteria | Eymployees of the Electric Generation Authority Bangkok, Thailand aged ≥ 35 years ('exploratory cohort'); middle-income social class |
| Exclusion criteria | Diabetes at baseline |
| Notes | Baseline data for cohort becoming diabetic (N = 361) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|---|
| Study participation: description of source population or population of interest | Low risk | Cohort study of employees of the Electric Generation Authority of Bangkok, Thailand |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | 3499 employees aged \geq 35 years; mostly urban dwellers of mid- dle-income social class |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria specified |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Of 3254 participants without diabetes at baseline, 2667 took part in the 1997 survey |

Aekplakorn 2006 (Continued)

| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
|---|--------------|---|
| Study attrition: adequate description of participants lost to follow-up | Low risk | Individuals lost to follow-up were slightly older |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Unclear, limited data only |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | 2-h OGTT after 75-g glucose load |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Glucose oxidase method |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG \geq 5.6 to < 7.0; IGT: 2-h PG \geq 7.8 to < 11.1 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | $FPG \ge 7.0$ or 2-h glucose ≥ 11.1 ; development of T2DM during the follow-up period until 1997 according to FPG or diagnosis and/or receipt of diabetes medication during follow-up |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Limited number of confounders |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |

Aekplakorn 2006 (Continued)

| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
|---|--------------|---|
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Low risk | Age, sex, BMI, waist circumference, smoking status, drinking status, family history, hypertension |
| Study confounding: important potential confounders accounted for in the analysis | Low risk | Yes; IFG status (model 2) and IGT status (model 3) |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Odds ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multivariable logistic regression |

Ammari 1998

| Name of study | None |
|--------------------|---|
| Inclusion criteria | Community-based survey of cardiovascular risk factors in 4 Jordanian towns, individuals aged ≥ 25 years; follow-up on one of the town (Sikhra) and matched control group with non-IGT (normal) individuals from initial survey |
| Exclusion criteria | Diabetes |
| Notes | Few baseline data reported for total study population (N = 212) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|---|
| Study participation: description of source population or population of interest | Low risk | 4 community-based survey of cardiovascular risk factors in 4 Jordanian towns |
| Study participation: description of gly- caemic status at baseline | Low risk | Community-based survey of cardiovascular risk factors in 4 Jor- danian towns |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |

Ammari 1998 (Continued)

| Study participation: adequate description of period & recruitment place | Low risk | Yes |
|---|--------------|---|
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | High risk | Scarce data |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not described (some comparison of participants with non-par- ticipants) |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not described |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | IGT |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | FPG and 2-h 75 g OGTT |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IGT: 2-h PG 7.8 to < 11.1 (WHO 1985) |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | 2-h PG ≥ 11.1 (WHO 1985) |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes (probably FPG and 2-h OGTT was also measured at follow-up) |
| Study confounding: important confounders measured | Unclear risk | Some baseline parameters were investigated (hypercholestero- laemia, hypertriglyceridaemia, obesity, hypertension, family his- |

Ammari 1998 (Continued)

| | | tory of diabetes) |
|---|--------------|----------------------|
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Scarce data |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Scarce data |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Cumulative incidence |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Not reported |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Unclear risk | Not reported |

Anjana 2015

| Name of study | Chennai Urban Rural Epidemiology Study (CURES) | |
|---|---|--|
| Inclusion criteria | Representative sample from Chennai, ≥ 20 years of age | |
| Exclusion criteria | Diabetes at baseline, unknown glycaemic status | |
| Notes | Baseline data for cohort becoming diabetic at follow-up (N = 176) | |
| Risk of bias | | |
| Bias | Authors' judgement | Support for judgement |
| Study participation: description of source population or population of interest | Low risk | Chennai Urban Rural Epidemiology Study |

Anjana 2015 (Continued)

| Study participation: description of gly- caemic status at baseline | Low risk | 299 with 'prediabetes' |
|---|--------------|--|
| Study participation: adequate description of sampling frame & recruitment | Low risk | Representative sample from Chennai, ≥ 20 years |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria specified |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | High risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | i-IFG, i-IGT, IFG/IGT |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | i-IGT: 2-h PG 7.8-11.0 and FPG > 5.6; i- IFG: FPG 5.6-6.9 and 2-h PG < 7.8; pre- diabetes: FPG 5.6-6.9 or 2-h PG 7.8-11.0 (i-IGT or i-IFG or IFG/IGT) |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | $FPG \ge 7.0$; 2-h PG ≥ 11.1 ; diagnosed; antihyperglycaemic medication |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |

Anjana 2015 (Continued)

| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
|---|--|--|
| Study confounding: important confounders measured | Unclear risk | For IFG/IGT, several confounders mea- sured as predictors for incident diabetes |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cox proportional hazards model for vari- ous single factors |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Univariate analyses |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, incidence rate |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Unclear risk | Cox proportional hazards model, univari- ate analyses for single variables |
| Bae 2011 | | |
| Name of study | None | |
| Inclusion criteria | Individuals who participated in comprehensive health check-ups annually for 5 years | |
| Exclusion criteria | Anaemia with a haemoglobin level < 7.4 mmol/L; self-reported diabetes and undiagnosed diabetes (FPG concentration 7.0 mmol/l or HbA1c 6.5%; absence of HbA1c data at any visit | |
| Notes | Baseline data for total cohort | |

Risk of bias

Bae 2011 (Continued)

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Participants partially undergoing annual or biannual health check-ups (Kangbuk Samsung Hospital Total,Healthcare Cen- ter) |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Unclear risk | Scarce data |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Not reported |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Low risk | Yes |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | HbA1c _{5.7} and HbA1c _{6.0} |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Unclear risk | Normal reference for HbA1c: < 5 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |

Bae 2011 (Continued)

| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; HbA1c \geq 6.5; history of diabetes; antihypergly- caemic medication |
|---|--------------|---|
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | 2 covariates measured: age and sex |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | 2 covariates included: age and sex |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | 2 covariates analysed: age and sex |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, incidence rate, hazard ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Unclear risk | Kaplan-Meier method, Cox proportional hazard analysis (2 co- variates), ROC analysis |

Baena-Diez 2011

| Name of study | None |
|--------------------|--|
| Inclusion criteria | Participants aged > 18 years visiting a healthcare centre with impaired fasting glucose measured twice |
| Exclusion criteria | Corticosteroid therapy, terminal illness, life expectancy of 1 year or less, diabetes |

Baena-Diez 2011 (Continued)

Notes Baseline data for cohort with intermediate hyperglycaemia (N = 115) Risk of bias Bias Authors' judgement Support for judgement Study participation: description of source Low risk Healthcare centre in Barcelona, Spain, "Cohorta Zona Franca" population or population of interest Study participation: description of gly- Low risk Yes caemic status at baseline Study participation: adequate description Unclear risk Scarce data of sampling frame & recruitment Study participation: adequate description Low risk Yes of period & recruitment place Study participation: adequate description Low risk Inclusion and exclusion criteria specified of inclusion & exclusion criteria Yes Study attrition: description of attempts to Low risk collect information on participants who dropped out Yes Study attrition: reasons for loss to follow- Low risk up provided Study attrition: adequate description of Low risk Yes participants lost to follow-up Quote: "no significant differences" Study attrition: no important differences Low risk between participants who completed the study and those who did not Glycaemic status measurement: provision Low risk IFG of clear definition or description of glycaemic status Glycaemic status measurement: valid and Low risk FPG measured twice reliable method of glycaemic status measurement IFG: 6.1-6.9 Glycaemic status measurement: continu- Low risk ous variables reported or appropriate cut points used

Baena-Diez 2011 (Continued)

| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
|---|--------------|--|
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0 (measured twice) |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | FPG |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Some variables (univariate analyses) associated with progression to diabetes |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some confounders measured |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Univariate analyses for single variables |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Unclear risk | Cox regression for other risk factors (e.g. obesity) associated with progression to diabetes |

Rai 1999

| Bai 1999 | | |
|--|---|--|
| Name of study | None | |
| Inclusion criteria | Staff of the Indian Institute of Technology of Chennai, along with their family members, aged 20 years and over | |
| Exclusion criteria | Treatment for diabetes | |
| Notes | Baseline data for the I | GT cohort (N = 252) |
| Risk of bias | | |
| Bias | Authors' judgement | Support for judgement |
| Study participation: description of source population or population of interest | Low risk | Staff of the Indian Institute of Technology of Chennai, along with their family members, aged 20 years and over |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | High risk | Not reported |
| Study attrition: reasons for loss to follow- up provided | High risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | High risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | IGT |
| Glycaemic status measurement: valid and | Low risk | Yes |

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reliable method of glycaemic status mea-

surement

Bai 1999 (Continued)

| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IGT: 7.8 to < 11.1 (WHO 1985) |
|---|--------------|------------------------------------|
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | 2-h PG ≥ 11.1 (WHO 1985) |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Not reported, cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Not reported |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Not reported |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Not reported |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Not reported |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Unclear risk | Not reported |

Bergman 2016

| Name of study | Israel study of glucose intolerance, obesity and hypertension (Israel GOH study) |
|--------------------|--|
| Inclusion criteria | Survival until follow-up with fasting blood glucose < 126 mg/dL (7.0 mmol/L) and 1- and 2-h postload glucose values available at baseline |
| Exclusion criteria | Individuals with diabetes |
| Notes | Baseline data for IGT cohort (N = 24) |
| | |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|---|
| Study participation: description of source population or population of interest | Low risk | Israeli general population registry sample |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Low risk | Yes |
| Study attrition: no important differences between participants who completed the study and those who did not | Low risk | Comment: "no differences" between non- participants and participants |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Comment: IGT |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |

Bergman 2016 (Continued)

| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | Comment: FPG 5.6-7.8; 2-h BG 7.8-11.0 |
|---|--------------|---|
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Unclear risk | Comment: FPG \geq 7.8, 2-h BG \geq 11.1; reported diabetes |
| Outcome measurement: method of out- come measurement used valid & reliable | Unclear risk | Non-standard FPG thresholds |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Comment: some confounders were mea- sured |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Comment: scarce data |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Comment: scarce data |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Low risk | Yes |
| Study confounding: important potential confounders accounted for in the analysis | Low risk | Yes |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, odds ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multiple multinomial logistic regression |

Bonora 2011

| Name of study | Bruneck Study |
|--------------------|---|
| Inclusion criteria | White men and women, aged 40-79 years |
| Exclusion criteria | Not reported |
| Notes | No baseline data (except white participants aged > 40 years, N = 919) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|---|
| Study participation: description of source population or population of interest | Low risk | Bruneck study, a long-term prospective population-based study of atherosclerosis and its risk factors |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | High risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Unclear risk | HbA1c categories, IFG (additional analyses) |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |

Bonora 2011 (Continued)

| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | HbA1: 6.0-6.49; IFG: not defined, probably FPG 5.6-6.9 |
|---|--------------|---|
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; HbA1c \geq 6.5; diabetes treatment |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Low risk | Yes |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Low risk | Yes |
| Study confounding: important potential confounders accounted for in the analysis | Low risk | Yes |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, incidence rate, hazard ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Cox proportional hazards models; additional models were run with updates variables (HbA1c and other variables were assessed every 5 years during follow-up) |

Cederberg 2010

| Name of study | None |
|--------------------|--|
| Inclusion criteria | All inhabitants of the city of Oulo, Finland, born in 1935 |
| Exclusion criteria | Diabetes at baseline |
| Notes | Baseline data for the total cohort (N = 553), men (N = 223), women (N = 330) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Part of a longer follow-up study assessing type 2 diabetes and IGT |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Low risk | Yes |
| Study attrition: no important differences between participants who completed the study and those who did not | Low risk | Yes |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | IFG, IGT, IFG/IGT |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |

Cederberg 2010 (Continued)

| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: 6.1-6.9; 2-h PG < 7.8; IGT: FPG > 7.0; 2-h PG 7.8 to < 11.1; elevated HbA1c: 5.7-6.4 |
|---|--------------|---|
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | Confirmed by 2 diabetic 75 g OGTTs (2-h PG \geq 11.1) and/or fasting values |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Some confounders measured |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Low risk | Yes |
| Study confounding: important potential confounders accounted for in the analysis | Low risk | Yes |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, risk ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Log-binomial regression |

Chamnan 2011

| Name of study | European Prospective Investigation of Cancer (EPIC)-Norfolk cohort |
|--------------------|---|
| Inclusion criteria | Participants aged 40-74 years from the Norfolk region, UK; individuals with HbA1c measurements at baseline and the second health assessment |
| Exclusion criteria | Diabetes at baseline, missing data |
| Notes | Baseline data for HbA1c 6.0-6.4 cohort (N = 370) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|---|
| Study participation: description of source population or population of interest | Low risk | Population-based study monitoring indi- viduals recruited from general practice in the Norfolk region, UK |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Scarce data |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | HbA1c (50% of all participants had infor- mation on this measure at baseline); analy- ses were limited to these individuals |

Chamnan 2011 (Continued)

| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
|---|--------------|--|
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | HbA1c 6.0-6.4 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | HbA1c \geq 6.5; reported physician-diagnosed diabetes or diabetes medications; antihyperglycaemic medication; diagnosis through medical records, registers or death certificates; results for clinically and/or biochemically diagnosed diabetes were used |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Low risk | Yes |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Low risk | Yes |
| Study confounding: important potential confounders accounted for in the analysis | Low risk | Yes |

Chamnan 2011 (Continued)

| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | | Cumulative incidence, odds ratio |
|---|----------|---|
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Logistic regression (for every 0.5% increase in HbA1c as well as for different categories of HbA1c) |

Charles 1997

| Name of study | Paris Prospective Study |
|--------------------|--|
| Inclusion criteria | Longitudinal epidemiologic study of cardiovascular risk factors in male employees of the Paris police, born in France between 1917-28 |
| Exclusion criteria | No diabetes or cardiovascular disease |
| Notes | Baseline data for individuals with IGT converting to T2DM (N = 32) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|---|
| Study participation: description of source population or population of interest | Low risk | Longitudinal epidemiologic study of cardiovascular risk fac- tors in male employees of the Paris |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | High risk | Not reported |
| Study attrition: reasons for loss to follow- up provided | High risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | High risk | Not reported |

Charles 1997 (Continued)

| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
|---|--------------|---|
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | IGT |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IGT: 2-h PG ≥ 7.8 to < 11.1 (WHO 1985) |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | 2-h PG \geq 11.1 (WHO 1985); physician diagnosed diabetes |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Low risk | Yes (see below) |

Charles 1997 (Continued)

| Study confounding: important potential confounders accounted for in the analysis | Low risk | Yes |
|---|----------|--|
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multivariate logistic regression (odds ratio for an increase of 1 SD in the population of participants with NGT or IGT) |

Chen 2003

| Name of study | None |
|--------------------|---|
| Inclusion criteria | Residents of Penghu, Taiwan aged 40-79 years were selected for a baseline diabetes prevalence study |
| Exclusion criteria | Diabetes at baseline |
| Notes | Baseline data for cohort converting to T2DM (N = 26) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Random sample of residents of Penghu, Taipei were selected for a baseline diabetes prevalence survey |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria reported |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |

Chen 2003 (Continued)

| Study attrition: adequate description of participants lost to follow-up | Low risk | Yes |
|---|--------------|---|
| Study attrition: no important differences between participants who completed the study and those who did not | Low risk | Quote: "the 600 persons who were re-examined did not signif- icantly differ from the others" |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | IFG |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG 6.1-7.0 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | $FPG \ge 7.0$ |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Some confounders measured |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |

Chen 2003 (Continued)

| Study confounding: important potential confounders accounted for in study design | Low risk | Yes |
|---|--------------|--|
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Age-sex adjusted odds ratio |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, odds ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multiple logistic regression (selected risk factors) |

Chen 2017

| Name of study | None |
|--------------------|--|
| Inclusion criteria | Participants with complete 3 year follow-up and non-pharmacological interventions |
| Exclusion criteria | Participants aged 0-60 years, incomplete baseline data, diabetes at baseline |
| Notes | Baseline data for i-IFG/i-IGTand IFG/IGT across age groups < 40 years + > 60 years (data indicate range across groups) (i-IFG < 40 years: N = 51 and > 60 years: N = 278; i-IGT < 40 years: N = 41 and > 60 years: N = 151; IFG/IGT: < 40 years: N = 34 and > 60 years: N = 175) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Permanent participants of Fujian province (China), part of the baseline survey from the REACTION study investigating the association between diabetes and cancer |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |

Chen 2017 (Continued)

| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Not reported |
|---|--------------|--|
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | IFG, IGT, IFG/IGT |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG 5.6-6.9 + 2-h PG \leq 7.8; IGT: FPG < 5.6 + 2-h PG 7.8 to \leq 11.0; IFG/IGT: FPG 5.6-6.9 + 2-h PG 7.8 to \leq 11.0 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; 2-h PG \geq 11.1; previously diagnosed diabetes |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Low risk | Confounder adjustment for HOMA-IR and HOMA-B |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |

Chen 2017 (Continued)

| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
|---|--------------|---|
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Low risk | Yes |
| Study confounding: important potential confounders accounted for in the analysis | Low risk | Yes (HOMA-IR, HOMA-B) |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Stepwise multiple regression analysis (for HOMA-IR or HOMA-B) |

Coronado-Malagon 2009

| Name of study | None |
|--------------------|--|
| Inclusion criteria | Healthy Mexicans |
| Exclusion criteria | Previous diabetes diagnosis, various diseases and medications affecting glucose metabolism |
| Notes | Baseline characteristics for the prediabetic cohort (N = 217) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Personnel working for Petróleos Mexicanos with annual health- checkups living in the metropolitan area of Mexico City |
| Study participation: description of gly- caemic status at baseline | Unclear risk | Quote: "prediabetes" |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |

Coronado-Malagon 2009 (Continued)

| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
|---|--------------|---|
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Unclear risk | IFG and IGT (ADA 2007), vague definition |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Unclear risk | IFG and IGT: 5.6-6.9 and 7.8 to < 11.1 (ADA 2007), vague definition |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Unclear risk | FPG \geq 7.0 or 2-h PG \geq 11.1 (ADA 2007), vague definition |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Scarce data |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Scarce data |

Coronado-Malagon 2009 (Continued)

| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Scarce data |
|---|--------------|-------------------------------------|
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Scarce data |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Scarce data |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Not reported |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, relative risk |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Logistic regression |

Cugati 2007

| Name of study | Blue Mountains Eye Study (BMES) |
|--------------------|---|
| Inclusion criteria | Survey of vision and common eye diseases in 2 postcode areas west of Sydney; all per- manent non-institutionalised residents with birth date prior to January 1943 (aged 49+ at baseline) were invited to attend a detailed eye examination at a local clinic |
| Exclusion criteria | Nursing home residents, diabetes at baseline, missing data |
| Notes | Baseline data for BMES I study, people without diabetes (N = 3437/3654) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Older community within the geographically de- fined area west of Sydney, Australia; population- based survey of vision and common eye diseases |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |

Cugati 2007 (Continued)

| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
|---|----------|--|
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Low risk | Yes |
| Study attrition: no important differences between participants who completed the study and those who did not | Low risk | Yes, for most variables |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | IFG |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG 5.6 -6.9 (originally FPG ≥ 6.1 to < 7. 0) |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; self-reported diabetes history; anti- hyperglycaemic medication |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |

Cugati 2007 (Continued)

| Study confounding: important confounders measured | Unclear risk | Few variables (adjustment for age and sex) |
|---|--------------|---|
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Few variables |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Few variables |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, odds ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multivariate-adjusted discrete logistic models, few variables |

De Abreu 2015

| Name of study | Geelong Osteoporosis Study (GOS) | |
|---|---|---|
| Inclusion criteria | Female arm of the GOS | |
| Exclusion criteria | No FPG level or self-report of antihyperglycaemic medication or diabetes status | |
| Notes | Baseline data for IFG cohort at baseline (N = 187) | |
| Risk of bias | | |
| Bias | Authors' judgement Support for judgement | |
| Study participation: description of source population or population of interest | Low risk | Utilised data from the female arm of the Geelong Osteoporosis Study, Australia |

De Abreu 2015 (Continued)

| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
|---|----------|--|
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Low risk | Yes |
| Study attrition: no important differences between participants who completed the study and those who did not | Low risk | Yes |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | IFG |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: 5.5-6.9 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; self-reported; antihyperglycaemic medication |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |

De Abreu 2015 (Continued)

| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
|---|--------------|---|
| Study confounding: important confounders measured | Low risk | Yes, also age-standardised incidence rate and additional covariates reported (metabolic syn- drome, fasting glucose at baseline) (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Low risk | Yes |
| Study confounding: important potential confounders accounted for in the analysis | Low risk | Yes |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, incidence rate, odds ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Logistic regression |

Den Biggelaar 2016

| Name of study | Cohort on Diabetes and Atherosclerosis Maastricht (CODAM) |
|--------------------|--|
| Inclusion criteria | Individuals with an elevated risk of type 2 diabetes and cardiovascular disease |
| Exclusion criteria | Previously diagnosed type 2 diabetes at baseline, who did not undergo an OGTT and incomplete OGTT data |
| Notes | Baseline data for prediabetic group (N = 122) |

Den Biggelaar 2016 (Continued)

Risk of bias

| Risk of bias | | |
|--|--------------------|--|
| Bias | Authors' judgement | Support for judgement |
| Study participation: description of source population or population of interest | Low risk | Participants of the Cohort on Diabetes and Atherosclerosis Masstricht (CODAM) study on natural progression of glucose tol- erance |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Low risk | Analyses restricted individuals without T2DM who participated in the follow-up measurements |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Scarce data |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | IFG and IGT |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | FPG 6.1-6.9; 2-h PG 7.8-11.1 |

Den Biggelaar 2016 (Continued)

| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
|---|--------------|--|
| Outcome measurement: clear definition of the outcome provided | Low risk | $FPG \ge 7.0; 2-h PG \ge 11.1$ |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Not reported, cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Not reported |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Not reported |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Not reported |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Not reported |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Discriminatory ability of beta-cell func- tions indices to predict 'prediabetes' and T2DM by means of ROC curves |

Derakhshan 2016

| Name of study | Tehran Lipid and Glucose Study (TLGS) |
|--------------------|--|
| Inclusion criteria | 3 separate analyses to investigate incidence of type 2 diabetes, hypertension and chronic kidney disease |
| Exclusion criteria | Individuals aged < 20 years, type 2 diabetes at baseline, missing data, no follow-ups |
| Notes | Baseline data for 'prediabetes' group with normal blood pressure (IFG and/or IGT, N = 523) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Population-based study on a representative sample of the population of Tehran to deter- mine the prevalence and incidence of non- communicable diseases and their risk fac- tors |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Low risk | Yes |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Unclear risk | Quote: "prediabetes" (IFG and IGT) |

Derakhshan 2016 (Continued)

| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
|---|----------|---|
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | $5.55 \le FPG < 7.0; 7.77 \le 2-h PG \le 11.1;$ no antihyperglycaemic medication |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; 2-h PG \geq 11.1; antihypergly- caemic medication |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Low risk | Yes |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Low risk | Multiple imputation |
| Study confounding: important potential confounders accounted for in study design | Low risk | Yes |
| Study confounding: important potential confounders accounted for in the analysis | Low risk | Yes |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Incidence rate, hazard ratio |

Derakhshan 2016 (Continued)

| Statistical analysis & reporting: the statisti- | Low risk | Cox proportional hazard models |
|---|----------|--------------------------------|
| cal model is adequate for the design of the | | |
| study | | |

Dowse 1991

| Name of study | Nauru Study |
|--------------------|---|
| Inclusion criteria | All Nauruans aged 20 years and over; this survey included 266 individuals who were not diabetic in the combined 1975/76 survey; individuals who had previously attended either or both the 1975/76 and 1982 surveys; individuals with at least one parent identified as being of Nauruan heritage |
| Exclusion criteria | Diabetes |
| Notes | No baseline data |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|---|
| Study participation: description of source population or population of interest | Low risk | Nauruan population, persons of Micronesian ancestry |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Description of inclusion and exclusion criteria |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Some reasons provided |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |

Dowse 1991 (Continued)

| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Scarce data |
|---|--------------|---|
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | IGT |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IGT: FPG < 7.8 and 2-h PG ≥ 7.8 - < 11.1 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | 2-h PG \ge 11.1 (WHO 1985); FPG \ge 7.8 |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Some confounders were measured |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Yes |

Dowse 1991 (Continued)

| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Yes |
|---|--------------|--|
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, incidence rate, odds ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multiple logistic regression models |

Ferrannini 2009

| Name of study | Mexico City Diabetes Study |
|--------------------|--|
| Inclusion criteria | Population-based study of diabetes and cardiovascular risk factors in low-income neigh- bourhoods in Mexico City, participants aged 35-64 years |
| Exclusion criteria | Type 2 diabetes, type 1 diabetes, pregnant women |
| Notes | Baseline characteristics provided for a range across different definitions of 'prediabetes' |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Data were collected as part of the Mexico City Diabetes Study |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Description of inclusion and exclusion criteria |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Not reported |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |

Ferrannini 2009 (Continued)

| Study attrition: adequate description of participants lost to follow-up | Low risk | Yes |
|---|--------------|---|
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Unclear, limited data |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | (i)IFG, (i)IGT |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG 6.1-6.9; IGT: FPG < 7.0 and 2-h PG 7.8-11. 1; i-IFG _{6.1} /i-IFG _{5.6} ; 2-h PG < 7.8 and FPG 6.1-6.9/5. 6-6.1; i-IGT/i-IGT _{6.1} /i-IGT _{5.6} |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG ≥ 7.0; 2-h PG ≥ 11.1 |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Not for transition data (intermediate hyperglycaemia to T2DM) |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |

Ferrannini 2009 (Continued)

| Study confounding: important potential confounders accounted for in study design | Unclear risk | Scarce data |
|---|--------------|---|
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Scarce data |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, relative risk (multiple model odds ratios were calculated for 1 SD of the population value of that variable, in order to compare the relative importance of the variables (sex, familial diabetes, age, BMI, FPG, 2-h PG) |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Unclear risk | Logistic regression (for calculation of odds ratios, see above) |

Filippatos 2016

| Name of study | ATTICA (province of Attica, Greece) |
|--------------------|---|
| Inclusion criteria | 1 participant per household, inhabitants from the Attica province |
| Exclusion criteria | People living in institutions; people with CVD and of those with chronic viral infections |
| Notes | Baseline data for IFG _{5.6} cohort |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|---|
| Study participation: description of source population or population of interest | Low risk | ATTICA study |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described (par- ticipants with no diabetes and no CVD at base- line) |

Filippatos 2016 (Continued)

| Study attrition: description of attempts to | Low risk | Yes (85% participation rate) |
|---|--------------|---|
| collect information on participants who dropped out | | |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Low risk | Yes |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | IFG _{5.6} |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | FBG 5.6-6.9 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FBG > 6.9; use of antidiabetic medication |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Some confounders measured |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |

Filippatos 2016 (Continued)

| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
|---|--------------|-------------------------------------|
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some confounders included |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Some confounders analysed |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, odds ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multiple logistic regression models |

Forouhi 2007

| Name of study | Ely Study (Cambridgeshire, UK) |
|--------------------|---|
| Inclusion criteria | All adults free of known diabetes registered with a single practice serving Ely, adults aged 40-69 years |
| Exclusion criteria | Diabetes |
| Notes | Baseline data for the IFG _{6.1} cohort (N = 257) Cumulative incidence increased across increasing age groups and was higher in men than in women |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | The Ely Study (Cambridgeshire, UK) was a prospec- tive study of the aetiology of T2DM |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |

Forouhi 2007 (Continued)

| Study participation: adequate description of period & recruitment place | Low risk | Yes |
|---|--------------|---|
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Low risk | Yes |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Scarce data |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | IFG |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG _{6.1} : FPG 6.1-6.9 (FPG < 7.0 and 2-h PG < 11. 1) and IFG _{5.6} : FPG 5.6-6.0 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; 2-h PG \geq 11.1; physician diagnosis or treatment for diabetes |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Some confounders measured |

Forouhi 2007 (Continued)

| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
|---|--------------|---|
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Low risk | Yes |
| Study confounding: important potential confounders accounted for in the analysis | Low risk | Yes |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, incidence rate, hazard ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Cox regression (cumulative hazard curves by glucose status) |

Garcia 2016

| Name of study | Sacramento Area Latino Study on Aging (SALSA) |
|--------------------|--|
| Inclusion criteria | Older Mexican Americans residing in the Sacramento metropolitan statistical area |
| Exclusion criteria | Missing baseline diabetes status, certain neighbourhoods |
| Notes | Baseline data for the IFG cohort (N = 310) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Participants were from the Sacramento Area Latino Study on Aging (SALSA), a longitudinal cohort study of physical and cognitive impairment and cardiovas- cular diseases in community-dwelling older Mexican Americans residing in the Sacra- |

Garcia 2016 (Continued)

| | | mento Metropolltan Statistical Area |
|---|----------|---|
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria reported |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Low risk | Yes |
| Study attrition: no important differences between participants who completed the study and those who did not | Low risk | Not reported but only 12/1789 participants were excluded |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | IFG |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | FBG 5.6-6.9 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | $FPG \geq 7.0$; self-reported; antihypergly- caemic medication; diabetes comedication at death |

Garcia 2016 (Continued)

| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
|---|--------------|---|
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Some confounders measured |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Low risk | Multistate Markov models |
| Study confounding: important potential confounders accounted for in the analysis | Low risk | Multistate Markov models |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence (hazard ratio was calculated for the association between neighbourhood scocioeconomic position (NSEP) scores and transitions between var- ious (pre)diabetic stages) |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multistate Markov models |

Gautier 2010

| Name of study | Data from an Epidemiological Study on the Insulin Resistance Syndrome (DESIR) cohort |
|--------------------|--|
| Inclusion criteria | Men and women aged 30-64 years recruited from volunteers who were offered periodic health examinations free of charge by the French Social Security at 10 health centres in western France |

Gautier 2010 (Continued)

| Exclusion criteria | Diabetes at baseline, individuals with unknown diabetes status at the 9-year examination |
|--------------------|--|
| Notes | No baseline data for cohort with intermediate hyperglycaemia |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|---|
| Study participation: description of source population or population of interest | Low risk | Participants of the Data from an Epidemio- logical Study on the Insulin Resistance Syn- drome (DESIR) cohort who had IFG at baseline |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Unclear risk | Key characteristics unclear |
| Study participation: adequate description of period & recruitment place | Unclear risk | Time frame unclear |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | High risk | Not reported |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Scarce data |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | IFG |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |

Gautier 2010 (Continued)

| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG 5.6-6.9 |
|---|--------------|---|
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; treatment for diabetes (at 1 of the 3-yearly examinations) |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Low risk | Some confounders measured |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Low risk | Yes (see below) |
| Study confounding: important potential confounders accounted for in the analysis | Low risk | Yes (see below) |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence (odds ratios for 9- year incident diabetes per 1 SD change in waist circumference and weight in IFG) |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Logistic models (for increases in waist cir- cumference and weight) |

Gomez-Arbelaez 2015

| Name of study | None |
|--------------------|--|
| Inclusion criteria | Adults \geq 35 years attending a general practitioner for any reason |
| Exclusion criteria | Known diabetes, acute illness, pregnancy, use of antihyperglycaemic medication |
| Notes | Baseline data for the total cohort (N = 772) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|---|
| Study participation: description of source population or population of interest | Low risk | Longitudinal observational study conducted in a healthcare cen- tre in Floridablanca, Colombia |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | The sub-sample of people with intermediate hyperglycaemia was followed for diabetes incidence |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | High risk | Not reported |
| Study attrition: reasons for loss to follow- up provided | High risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | High risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Intermediate hyperglycaemia as measured by FPG, OGTT, HbA1c; FINDRISC score |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |

Gomez-Arbelaez 2015 (Continued)

| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: \geq 5.6 to < 7.0; IGT: \geq 7.8 to < 11.1; HbA1c \geq 5.7 to \leq 6.4 |
|---|--------------|---|
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; OGTT \geq 11.1; HbA1c \geq 6.5 |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Age and sex-adjusted odds ratios for FINDRISC score |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Low risk | For FINDRISC score |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Age and sex-adjusted odds ratios |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, incidence rate |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multivariate logistic regression for the association between the FINDRISC score and incident T2DM |

Guerrero-Romero 2006

| Name of study | None |
|--------------------|---|
| Inclusion criteria | Men and non-pregnant women, aged 20-64 years, were recruited from the city of Du- rango, northern Mexico; with NGT or IGT |
| Exclusion criteria | Participants who failed to attend 2 or more visits |
| Notes | Baseline data for IGT cohort at baseline progressing to T2DM (N = 20); all individuals were counselled on the importance of diet and physical exercise (standard care for the whole cohort) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Cohort study in healthy Mexicans to determine predictors for the development of metabolic disorders |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Unclear risk | Time frame unclear |
| Study participation: adequate description of period & recruitment place | Unclear risk | Period of recruitment unclear |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Not reported |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | IGT |

Guerrero-Romero 2006 (Continued)

| Low risk | Yes |
|--------------|--|
| Low risk | IGT: 2-h PG ≥ 7.8 to < 11.1 |
| Low risk | Yes |
| Low risk | 2-h PG: ≥ 11.1 |
| Low risk | Yes |
| Low risk | Yes |
| Unclear risk | Some covariates measured (for association between beta-cell function and IGT/T2DM) (see Appendix 16 and Appendix 17) |
| Unclear risk | Not reported |
| Unclear risk | Not reported |
| Low risk | Yes |
| Unclear risk | Not reported |
| Low risk | For beta-cell function and IGT/T2DM |
| Unclear risk | Some confounders measured |
| Low risk | Cumulative incidence, incidence rate |
| | Low risk Low risk Low risk Low risk Unclear risk Unclear risk Unclear risk Low risk Low risk Low risk Low risk |

Guerrero-Romero 2006 (Continued)

| Statistical analysis & reporting: the statisti- | Low risk | Multivariate logistic regression on relative risk of IGT or T2DM |
|---|----------|--|
| cal model is adequate for the design of the | | associated with beta-cell function |
| study | | |

Han 2017

| Name of study | Ansung-Ansan cohort study, part of the Korean Genome and Epidemiology Study (Ko-GES), to investigate the trends in diabetes and associated risk factors |
|--------------------|--|
| Inclusion criteria | Urban (Ansan) and rural (Ansung) communities (within 60 km of Seoul) |
| Exclusion criteria | Unknown glucose status, individuals with known diabetes, participants who were newly diagnosed with type 2 diabetes at baseline examination; persons with a history of malig- nant diseases, liver failure, end-stage renal disease, rheumatological diseases and acute or chronic infec- tious diseases, individuals who had taken steroids in the previous 3 months; individuals who did not undergo any follow-up examination after the baseline examination |
| Notes | Baseline data for i-IFG, i-IGT and IFG/IGT |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|---|
| Study participation: description of source population or population of interest | Low risk | Ansung-Ansan Cohort Study, part of the Korean Genome and Epidemiology Study (KoGES) |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes (follow-up rate at 12 years 60.6%) |

Han 2017 (Continued)

| Study attrition: adequate description of participants lost to follow-up | Low risk | Yes |
|---|--------------|---|
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG 5.6-6.9 and no diagnosis of dia- betes; IGT: 2-h PG 7.8 to < 11.1; i-IFG _{5.6} : IFG without IGT; i-IGT: IGT without IFG; IGT/IGT: IFG+IGT; 'prediabetes': IFG or IGT |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; 2-h PG \geq 11.1; HbA1c \geq 6. 5; current antihyperglycaemic treatment |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Low risk | Yes |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |

Han 2017 (Continued)

| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
|---|--------------|---|
| Study confounding: important potential confounders accounted for in study design | Low risk | Yes |
| Study confounding: important potential confounders accounted for in the analysis | Low risk | Yes |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, incidence rate, haz- ard ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multivariate Cox proportional hazard model |

Hanley 2005

| Name of study | Insulin Resistance Atherosclerosis Study (IRAS) |
|--------------------|---|
| Inclusion criteria | 4 clinical centres (Oakland, Los Angeles - non-Hispanic whites and blacks recruited from Kaiser Permanente) and San Antonio, San Luis Valley (non-Hispanic whites and Hispanics): from 2 population-based studies (San Antonio Heart Study and the San Luis Valley Diabetes study) |
| Exclusion criteria | Participants with inflammatory diseases; diabetes; no information on metabolic variables of interest and follow-up glucose tolerance status |
| Notes | Baseline data for diabetic cohort at follow-up (N = 131); participants were recruited from 2 population-based studies: the San Antonio Heart Study and the San Luis Valley diabetes study |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Observational study of the relationship be- tween insulin resistance, cardiovascular dis- ease and its known risk factors in differ- ent ethnic groups and varying states of glu- cose tolerance; the study was conducted at 4 clinical centres; report on individu- als who were nondiabetic at baseline and for whom information was available on metabolic variables of interest and follow- up glucose tolerance status |

Hanley 2005 (Continued)

| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
|---|--------------|--|
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Response rate 81% |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Scarce data |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Low risk | Yes |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Unclear risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG, IGT (WHO 1999) |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | High risk | Not specified |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |

Hanley 2005 (Continued)

| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
|---|--------------|--|
| Study confounding: important confounders measured | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Some covariates (age, sex, clinical centre, ethnicity) (see Appendix 16 and Appendix 17) |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, odds ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Logistic regression |

| Name of study | Toranomon Hospital Health Management Center Study (TOPICS) |
|--------------------|---|
| Inclusion criteria | Participants from the TOPICS: apparently healthy Japanese government employees who underwent annual multiphasic health screening examinations; the study attempted to elucidate the incidence of and risk factors for various diseases among the Japanese pop- ulation |
| Exclusion criteria | Diabetes at baseline, missing data at baseline |
| Notes | Baseline data for the total cohort (N = 6241) |

Heianza 2012 (Continued)

Risk of bias

| Risk of bias | | |
|--|--------------------|--|
| Bias | Authors' judgement | Support for judgement |
| Study participation: description of source population or population of interest | Low risk | Healthy Japanese government employees who underwent annual examinations for health screening |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Scarce data |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG 5.6-6.9 or FPG 6.1-6.9; HbA1c 5.7 -6.4 or 6.0-6.4; IFG/HbA1c = 'predia- betes' |
| Glycaemic status measurement: same method and setting of measurement of the | Low risk | Yes |

Heianza 2012 (Continued)

| glycaemic status for all study participants | | |
|---|--------------|---|
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; HbA1c \geq 6.5%; self-reported clinician-diagnosed diabetes |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Low risk | Yes |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Low risk | Yes |
| Study confounding: important potential confounders accounted for in the analysis | Low risk | Yes |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, incidence rate, haz- ard ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Cox regression, multivariate model |

Inoue 1996

| Name of study | None |
|--------------------|---|
| Inclusion criteria | Non-obese participants with IGT and 22 normal control persons were selected from the participants of a health screening programme |
| Exclusion criteria | People with liver or kidney diseases |
| Notes | Baseline data for the IGT cohort (N = 37) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|--|
| Study participation: description of source population or population of interest | Unclear risk | Participants of a health screening programme |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Unclear risk | Scarce data |
| Study participation: adequate description of period & recruitment place | Unclear risk | Scarce data |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Not reported |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | IGT |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |

Inoue 1996 (Continued)

| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IGT: \geq 7.8 to < 11.1 (presumed WHO 1985) |
|---|--------------|---|
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | IGT: \geq 11.1 (presumed WHO 1985) |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Not reported, cumulative incidence data |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Not reported |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Not reported |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Not reported |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Not reported, cumulative incidence data |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Not reported |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Kruskal-Wallis test |

Janghorbani 2015

| Name of study | Isfahan Diabetes Prevention Study (IDPS) |
|--------------------|--|
| Inclusion criteria | Participants with a family history of type 2 diabetes, being non-diabetic |
| Exclusion criteria | Type 1 diabetes, pregnancy |
| Notes | Baseline data for i-IFG, i-IGT and IFG/IGT cohort (N = 770); first-degree relatives of people with T2DM; data on the cohort without hypertension at baseline |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Ongoing cohort in central Iran to assess the various potential risk factors for diabetes in people with a family history of T2DM |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Description of inclusion and exclusion cri- teria |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Low risk | Yes |
| Study attrition: no important differences between participants who completed the study and those who did not | Low risk | Yes |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |

Janghorbani 2015 (Continued)

| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
|---|--------------|--|
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | i-IGT: FPG < 5.6 and 2-h PG 7.8-11.1; i- IFG: 5.6-6.9 and 2-h PG < 7.8; IFG/IGT: 5.6-6.9 and 2-h PG 7.8-11.1 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 11.1; antihyperglycaemic medica- tion; 2nd FPG \geq 7.0; 2-h PG \geq 11.1 |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some covariates measured (age, sex, BMI, triglycerides, total cholesterol) (see Appendix 16 and Appendix 17) |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Some covariates measured (age, sex, BMI, triglycerides, total cholesterol) (see Appendix 16 and Appendix 17) |

Janghorbani 2015 (Continued)

| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | | Cumulative incidence, incidence rate, haz- ard ratio |
|---|----------|---|
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Cox proportional hazards model |

Jaruratanasirikul 2016

| Name of study | None |
|--------------------|---|
| Inclusion criteria | Obese Thai children and adolescents aged 8-15 years, Pediatric Endocrine Clinic at Songklanagarind Hospital (Hat Yai, Songkhia Thailand) |
| Exclusion criteria | No clinical findings of secondary obesity, not on corticosteroids |
| Notes | Baseline data for IGT cohort (N = 27) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Scarce data |
| Study attrition: reasons for loss to follow- up provided | High risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |

Jaruratanasirikul 2016 (Continued)

| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
|---|--------------|---|
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | (i)-IGT: FPG < 5.6 and 2-h PG 7.8 to < 11.1 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG > 7.0; 2-h PG ≥ 11.1 |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Not reported |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Not reported |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Not reported |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Not reported |

Jaruratanasirikul 2016 (Continued)

| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Not reported |
|---|--------------|---|
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Cox regression analysis for ROC curves (cut-off FPG levels) |

Jeong 2010

| Name of study | None |
|--------------------|---|
| Inclusion criteria | People older 20 years living in the rural area of Dalseong County near Daegu visiting community health centres |
| Exclusion criteria | Not reported |
| Notes | 1287 participants were re-evaluated in 2008 and 187 new participants "added to the study"; baseline data for participants with incident diabetes (N = 135) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Population-based survey to determine the prevalence and inci- dence of 'prediabetes' and diabetes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Unclear risk | Only inclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | High risk | Several surveys plus new recruitment; follow-up rate 80.5%; no description of dropouts |

Jeong 2010 (Continued)

| Study attrition: reasons for loss to follow- up provided | High risk | Not reported |
|---|--------------|--|
| Study attrition: adequate description of participants lost to follow-up | High risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG \geq 5.6 to < 7.0; IGT: 2-h PG \geq 7.8 to < 11.1; 'pre-diabetes': IFG or IGT |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG ≥ 7.0; 2-h PG ≥ 11.1 |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Low risk | Several covariates were measured (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Not reported |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Not reported |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |

Jeong 2010 (Continued)

| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
|---|--------------|----------------------------|
| Study confounding: important potential confounders accounted for in study design | Low risk | Yes |
| Study confounding: important potential confounders accounted for in the analysis | Low risk | Yes |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Unclear risk | Odds ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Logistic regression models |

Jiamjarasrangsi 2008a

| Name of study | None |
|--------------------|--|
| Inclusion criteria | Individuals 35 years or older participating in the annual physical checkup at least twice during the years 2001-2005 |
| Exclusion criteria | People with diabetes |
| Notes | Baseline data for total cohort becoming diabetic at follow-up (N = 48) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | University hospital employees |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |

Jiamjarasrangsi 2008a (Continued)

| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Not reported |
|---|--------------|---|
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG \geq 5.6 to < 7.0 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | $FPG \ge 7.0$ |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Logistic regression on hepatic enzymes; incidence rate: few co- variates (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Logistic regression on hepatic enzymes |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Logistic regression on hepatic enzymes |

Jiamjarasrangsi 2008a (Continued)

| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Logistic regression on hepatic enzymes |
|---|--------------|---|
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Logistic regression on hepatic enzymes |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | lLogistic regression on hepatic enzymes |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Logistic regression on hepatic enzymes |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, incidence rate |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Logistic regression (independent variables: hepatic enzymes) and Poisson regression analyses |

Kim 2005

| Name of study | None |
|--------------------|--|
| Inclusion criteria | People visiting the Health Promotion Centre of Samsung Medical Center for a physical health check-up |
| Exclusion criteria | Diabetes |
| Notes | Baseline data for FPG group 4 (6.1-7.0) with baseline and follow-up (N = 276) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|-----------------------|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes (FPG categories) |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |

Kim 2005 (Continued)

| Study participation: adequate description | I ow risk | Inclusion and exclusion criteria described |
|---|--------------|---|
| of inclusion & exclusion criteria | LOW HOR | |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Participation rate 20.9% in group 4; scarce data |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG 6.1 to < 7.0 (group 4) |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; antihyperglycaemic treatment |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Low risk | Several covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Scarce data |

Kim 2005 (Continued)

| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Scarce data |
|---|--------------|------------------------------------|
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Low risk | Yes |
| Study confounding: important potential confounders accounted for in the analysis | Low risk | Yes |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, hazard ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Cox regression analysis |

Kim 2008

| Name of study | None |
|--------------------|---|
| Inclusion criteria | Individuals undergoing a medical examination at Inha University Hospital with a follow- up medical examination 2 years later |
| Exclusion criteria | Individuals diagnosed with diabetes at baseline |
| Notes | Baseline data for IFG _{5.6} /IFG _{6.1} cohort (N = 1335/N = 494) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|---|
| Study participation: description of source population or population of interest | Low risk | Participants who underwent a medical examination at Inha Uni- versity Hospital and had either NGT or IFG |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |

Kim 2008 (Continued)

| Study participation: adequate description of period & recruitment place | Low risk | Yes |
|---|--------------|--|
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Participants diagnosed with diabetes in 2002 were excluded |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Scarce data |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG _{5.6} : FPG 5.6-7.0; IFG _{6.1} : FPG 6.1-7.0 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | $FPG \ge 7.0$ |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Measurement of cumulative incidence |

Kim 2008 (Continued)

| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Not reported |
|---|--------------|--|
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Not reported |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Not reported |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Measurement of cumulative incidence |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | ROC curves for predicting the future onset of diabetes |

Kim 2014

| Name of study | None | |
|--------------------|--|-----------------------|
| Inclusion criteria | Pre-screened individuals with 'prediabetes' visiting the diabetes clinic at Seoul National University Bundang Hospital (SNUB) in 2005/06 after they were diagnosed with pre- diabetes at their health check-up or primary clinic | |
| Exclusion criteria | Taking oral hypoglycaemic agents or insulin | |
| Notes | Baseline data for i-IFG (N = 158)/i-IGT (N = 65)/IFG/IGT (N = 119)/i-HbA1c (N = 64); total: N = 406 | |
| Risk of bias | | |
| Bias | Authors' judgement | Support for judgement |

| Study participation: description of source population or population of interest | Low risk | Pres-screened individuals with 'prediabetes' |
|---|----------|--|

Kim 2014 (Continued)

| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
|---|--------------|---|
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Pre-defined participants with intermediate hyperglycaemia |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | i-IFG: FPG 5.6-6.9 and 2-h PG < 7.8; i-IGT: 2-h PG 7.8- 11.1 and FPG < 5.6; IFG/IGT: combined glucose intolerance; HbA1c: 5.7-6.4 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | $FPG \ge 7.0; 2-h PG \ge 11.1; HbA1c \ge 6.5$ |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |

Kim 2014 (Continued)

| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
|---|--------------|--|
| Study confounding: important confounders measured | Unclear risk | For C-peptide |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | For C-peptide |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | For C-peptide |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | For C-peptide |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | For C-peptide |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | For C-peptide |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multiple logistic regression for association of T2DM develop- ment and C-peptide levels |

Kim 2016a

| Name of study | None |
|--------------------|---|
| Inclusion criteria | Medical examinations at the Health Screening and Promotion Center at Asan Medical Center (Seoul, Korea) |
| Exclusion criteria | History of diabetes mellitus, taking antihyperglycaemic medications, FPG ≥ 7.0 mmol/ L or HbA1c $\geq 6.5\%$ at baseline |
| Notes | 2 baseline data cohorts: 'prediabetes' by FPG and HbA1c (N = 3544 and N = 1713) |
| Risk of bias | |

Kim 2016a (Continued)

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|--|
| Study participation: description of source population or population of interest | Unclear risk | Participants who underwent medical examinations in a health screening and promotion centre |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Not reported |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | FPG 5.6-6.9; HbA1c 5.7-6.4 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | FPG \geq 7.0; HbA1c \geq 6.5; antihyperglycaemic medications |

Kim 2016a (Continued)

| Outcome measurement: clear definition of the outcome provided | Low risk | Yes |
|---|--------------|---|
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Low risk | Several covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Low risk | Yes |
| Study confounding: important potential confounders accounted for in the analysis | Low risk | Yes |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, odds ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multivariate logistic regression |

Kleber 2010

| Name of study | None |
|--------------------|---|
| Inclusion criteria | Obese children and adolescents aged 10-17 years with IGT attending the outpatient centre (Department of Paediatric Nutrition Medicine, Witten/Herdecke Germany) |

Kleber 2010 (Continued)

| Exclusion criteria | Not reported | |
|--|---------------------------------------|--|
| Notes | Baseline data for IGT cohort (N = 79) | |
| Risk of bias | | |
| Bias | Authors' judgement | Support for judgement |
| Study participation: description of source population or population of interest | Low risk | Obese white children and adolescents with IGT attending an outpatient centre |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Unclear risk | Time of recruitment unclear |
| Study participation: adequate description of inclusion & exclusion criteria | Unclear risk | No exclusion criteria reported |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Probably no dropouts |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IGT: 2-h PG > 7.7: IFG: FPG ≥ 5.5 |

Kleber 2010 (Continued)

| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
|---|--------------|-----------------------------|
| Outcome measurement: clear definition of the outcome provided | Low risk | T2DM by ADA 2000 guidelines |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Not reported |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Not reported |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Not reported |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Not reported |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multiple linear regression |

Kleber 2011

| Name of study | None |
|--------------------|---|
| Inclusion criteria | Obese white children with IGT without medication or endocrine/syndromal disorders, aged 10-17 years not participating in the intervention part of the study |
| Exclusion criteria | Children in the intervention part of the study |
| Notes | Baseline data for IFG cohort (N = 128) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Obese children and adolescents with IGT not attending an in- tervention trial |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Unclear risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Low risk | Yes |
| Study attrition: no important differences between participants who completed the study and those who did not | Low risk | Yes |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |

Kleber 2011 (Continued)

| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IGT: not reported (presumed 7.8-11.1) |
|---|--------------|---------------------------------------|
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | "ADA" (2000 criteria - 2-h PG ≥ 11.1) |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Measurement of cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Not reported |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Npt reported |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Not reported |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Not reported |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multiple linear regression |

Ko 1999

| Name of study | None |
|--------------------|-------------------------------|
| Inclusion criteria | Chinese participants with IGT |
| Exclusion criteria | Not reported |
| Notes | Letter to the editor |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|-----------------------------------|
| Study participation: description of source population or population of interest | Low risk | Chinese participants with IGT |
| Study participation: description of gly- caemic status at baseline | Low risk | WHO/NDGG 1979 |
| Study participation: adequate description of sampling frame & recruitment | Unclear risk | Scarce data |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Unclear risk | Only inclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Not reported (IGT cohort) |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported (IGT cohort) |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported (IGT cohort) |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not applicable (IGT cohort) |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | IGT (WHO/NDDG 1979 definition) |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |

Ko 1999 (Continued)

| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | Yes |
|---|--------------|---|
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | Assumed WHO/NDDG 1979 definition |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Cumulative incidence |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Cumulative incidence |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Cumulative incidence |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Cumulative incidence |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Unclear risk | Cox regression analysis (to predict the progression to diabetes with age, sex, BMI, blood pressure, HbA1c, FPG, 1-h PG and 2-h PG as predictor variables) |

Ko 2001

| Ko 2001 | | |
|--|---|---|
| Name of study | None | |
| Inclusion criteria | The Diabetes and Endocrine Centre of the prince of Wales Hospital in Hong Kong screened individuals with risk factors for glucose intolerance (family history of diabetes, history of gestational diabetes, overweight, hypertension) by OGTT | |
| Exclusion criteria | Diabetes at baseline | |
| Notes | Baseline data for IFG | cohort (N = 55) |
| Risk of bias | | |
| Bias | Authors' judgement | Support for judgement |
| Study participation: description of source population or population of interest | Low risk | Individuals with risk factors for glucose intolerance undergoing screening for diabetes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Low risk | Yes |
| Study attrition: no important differences between participants who completed the study and those who did not | Low risk | Yes |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| | | |

Ko 2001 (Continued)

| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
|---|--------------|-------------------------------------|
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG 6.1-6.9 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | $FPG \ge 7.0$ |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Measurement of cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | No ratios reported |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | No ratios reported |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |

Ko 2001 (Continued)

| Statistical analysis & reporting: the statisti- | Low risk | Kaplan-Meier analysis, Cox regression analysis (to predict the |
|---|----------|--|
| cal model is adequate for the design of the | | progression to diabetes with age, sex, BMI, blood pressure, FPG, |
| study | | gestational diabetes, HbA1c, smoking habit and IFG status be- |
| | | ing independent variables - no hazard ratios provided) |

Larsson 2000

| Name of study | None |
|--------------------|--|
| Inclusion criteria | Postmenopausal women aged 55-57 years in a health screening programme; random sample of 265/1843 invited for follow-up (new OGTT); 1843 women were grouped according to WHO and ADA glucose tolerance criteria |
| Exclusion criteria | Not reported |
| Notes | Baseline data for (i)-IGT (N = 66)/(i)-IFG (N = 42)/IFG/IGT (N = 30); 265 follow-up participants were randomly sampled from each glucose tolerance group of the original cohort and invited for follow-up; NGT at baseline vs follow-up: FPG < 5.3 vs < 6.1; FPG 5.3: 15% conversion factor as recommended by the WHO (blood glucose > plasma glucose) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|--|
| Study participation: description of source population or population of interest | Unclear risk | Postmenopausal women participating in a health screening pro- gramme; follow-up: a random sample of the original cohort |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Unclear risk | No exclusion criteria reported |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Not reported |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |

Larsson 2000 (Continued)

| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
|---|--------------|---|
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | (i)-IFG: BG 5.3-5.9 and 2-h BG < 7.8; (i)-IGT: FPG < 5.3 and 2-h BG 7.8-11.0; IFG/IGT: BG 5.3-5.9 and 2-h BG 7.8-11.0 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG ≥ 7.0; 2-h PG ≥ 11.1 |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Measurement of cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Not reported |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Not reported |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Not reported |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |

Larsson 2000 (Continued)

| Study confounding: important potential confounders accounted for in study design | Unclear risk | Not reported | |
|---|--|--|--|
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Not reported | |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence | |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Chi-squared test | |
| Latifi 2016 | | | |
| Name of study | None | | |
| Inclusion criteria | Residents aged over 20 years | | |
| Exclusion criteria | Not reported | | |
| Notes | Baseline for prediabetic cohort becoming diabetic at follow-up | | |
| Risk of bias | | | |
| Bias | Authors' judgement | Support for judgement | |
| Study participation: description of source population or population of interest | Low risk | First phase of prevalence study of the metabolic syndrome and its related factors in Ahvaz Diabetes Research Centre, Iran | |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes | |

Yes

Yes

Not reported

No exclusion criteria reported

Development of type 2 diabetes mellitus in people with intermediate hyperglycaemia (Review) Copyright © 2018 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.

Study participation: adequate description Low risk

Study participation: adequate description Low risk

Study participation: adequate description Unclear risk

Study attrition: description of attempts to Unclear risk

of sampling frame & recruitment

of period & recruitment place

of inclusion & exclusion criteria

dropped out

collect information on participants who

Latifi 2016 (Continued)

| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
|---|--------------|---|
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | $5.6 \leq FPG < 7.0$ |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; antihyperglycaemic medication |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Several covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |

Latifi 2016 (Continued)

| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
|---|--------------|--|
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some covariates (see Appendix 16 and Appendix 17) |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Some covariates (see Appendix 16 and Appendix 17) |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Unclear risk | Cumulative incidence, incidence rate |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Unclear risk | Multiple logistic regression of factors affecting the incidence of diabetes and prediabetes among healthy people in phase 1 (baseline) |

Lecomte 2007

| Name of study | None |
|--------------------|---|
| Inclusion criteria | People with IFG recruited from medical check-ups by the French social security system in the 9 preventive health centres of IRSA (Institut Interrégional pur la Santé) |
| Exclusion criteria | No personal history of diabetes, no hypoglycaemic drug treatment |
| Notes | Baseline data for IFG cohort attending both examinations (N = 743) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|-----------------------|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Yes |

Lecomte 2007 (Continued)

| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
|---|--------------|--|
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG 6.1-6.9; no personal history of diabetes; no hypogly- caemic treatment |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; personal history of diabetes; antihyperglycaemic treatment |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Measurement of cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |

Lecomte 2007 (Continued)

| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
|---|--------------|---|
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Univariate analyses, some covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Some covariates, univariate analyses (see Appendix 16 and Appendix 17) |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Logistic regression, univariate analyses on risk factors for devel- oping diabetes |

Lee 2016

| Name of study | None |
|--------------------|--|
| Inclusion criteria | Individuals undergoing health checkups at a single medical institution (Gangneung Asian Hospital) |
| Exclusion criteria | Previously diagnosed with diabetes, history of diabetes medication use, only 1 measure- ment |
| Notes | Baseline data for the total cohort (N = 3497) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|-----------------------|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |

Lee 2016 (Continued)

| Study participation: adequate description of period & recruitment place | Low risk | Yes |
|---|--------------|--|
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Not reported |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | HbA1c 5.7-6.4 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | HbA1c \geq 6.5 |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Measurement of cumulative incidence |

Lee 2016 (Continued)

| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes for coffee consumption |
|---|--------------|---|
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | 1 covariate |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | No ratios reported |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Unclear risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Kaplan-Meier survival analysis for progression to diabetes ac- cording to coffee consumption |

Leiva 2014

population or population of interest

| Name of study | Programa de Investigación de Factores de Riesgo de Enfermedad Cardiovascular (PIFRECV) | |
|--|---|-----------------------|
| Inclusion criteria | Study participants were recruited in 2005 by the 'Programa de Investigación de Factores de Riesgo de Enfermedad Cardiovascular' (PIFRECV); participants had to have an FPG 5.6-6.9 mmol/L | |
| Exclusion criteria | Diabetes, individuals on corticosteroid treatment, pregnant women, individuals with cardiovascular complications | |
| Notes | Most baseline data for cohort becoming diabetic at follow-up (N = 94 with IFG) | |
| Risk of bias | | |
| Bias | Authors' judgement | Support for judgement |
| Study participation: description of source | Low risk | Yes |

Leiva 2014 (Continued)

| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
|---|--------------|--|
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: 5.6-7.0 (low range: 5.6-6.1; high range: 6.1-6.9) |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0 (on 2 consecutive days); HbA1c \geq 6.5 |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |

Leiva 2014 (Continued)

| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
|---|--------------|--|
| Study confounding: important confounders measured | Unclear risk | Some covariates were measured (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some covariates planned (see Appendix 16 and Appendix 17) |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Some covariates analysed (see Appendix 16 and Appendix 17) |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, odds ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Cox regression analysis (comparing 'high range' glycaemia (> 6.1 mmol/L) with 'low range' glycaemia (< 6.1 mmol/L) |

Levitzky 2008

| Name of study | Framingham Heart Study |
|--------------------|---|
| Inclusion criteria | Participants were drawn from the Framingham Offspring cohort; participants who at- tended examinations (referred to as index examinations) |
| Exclusion criteria | Participants with CHD or diabetes |
| Notes | Baseline data for individuals on first exam, free of CVD (N = 4058) |

Risk of bias

Levitzky 2008 (Continued)

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria reported |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Not reported |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG _{5.6} : FPG 5.6-6.9; IFG _{6.1} : FPG 6.1-6.9 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |

Levitzky 2008 (Continued)

| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; antihyperglycaemic medication |
|---|--------------|--|
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some covariates included (see Appendix 16 and Appendix 17) |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Some covariates analysed (see Appendix 16 and Appendix 17) |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, odds ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Pooled logistic regression, multivariable models |

Li 2003

| Name of study | Kinmen Study (study in Kin-Chen, Kinmen, Taiwan) |
|--------------------|---|
| Inclusion criteria | Individuals aged \geq 30 years in Kin-Chen; FPG 5.6-7.0 and 2-h PG < 11.1 |
| Exclusion criteria | Diabetes |

Li 2003 (Continued)

Baseline data for i-IGT (N = 118)/i-IFG (N = 42)/IFG/IGT (N = 49) cohorts

Risk of bias

Notes

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|---|
| Study participation: description of source population or population of interest | Low risk | Yes, series of community-based epidemio- logical surveys of diabetes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Not reported |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | i-iFG: FPG 6.1-7.0 and 2-h PG < 7.8; i- IGT: FPG < 6.1 and 2-h PG 7.8-11.1; IFG/ IGT: FPG 6.1-7.0 and 2-h PG 7.8-11.1 |

Li 2003 (Continued)

| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
|---|--------------|--|
| Outcome measurement: clear definition of the outcome provided | Low risk | $FPG \ge 7.0$; 2-h PG ≥ 11.0 ; antihypergly- caemic medication |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some covariates included (see Appendix 16 and Appendix 17) |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Some covariates analysed (see Appendix 16 and Appendix 17) |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, incidence rate, haz- ard ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Cox proportional hazard model (hazard ra- tios of T2DM for relative insulin resistance, beta-cell dysfunction and varying degrees of glucose intolerance) |

Ligthart 2016

| Name of study | Rotterdam study, targeting cardiovascular, endocrine, hepatic, neurological, ophthalmic, psychiatric, dermatological, oncological and respiratory diseases |
|--------------------|--|
| Inclusion criteria | Community dwelling population aged 45/55 years and older in Rotterdam, no diabetes at baseline |
| Exclusion criteria | No valid baseline fasting glucose measurement, no informed consent |
| Notes | Baseline data for prediabetic cohort (N = 1382) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Not reported |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |

Ligthart 2016 (Continued)

| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
|---|--------------|--|
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | FBG > 6.0 and < 7.0; non-fasting BG > 7. 7 and < 11.1 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | $FBG \ge 7.0$; non-fasting $BG \ge 11.1$; anti- hyperglycaemic medication |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Some covariates for lifetime risk of diabetes (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | For lifetime risk of diabetes |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | For lifetime risk of diabetes |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Incidence rate |

Ligthart 2016 (Continued)

| Statistical analysis & reporting: the statisti- | Unclear risk | Modified version of survival analysis to cal- |
|---|--------------|---|
| cal model is adequate for the design of the | | culate the lifetime risk of diabetes |
| study | | |

Lipska 2013

| Name of study | Health, Aging, and Body Composition study (Health ABC) |
|--------------------|--|
| Inclusion criteria | Aged 70-79 years from Pittsburgh (PA) and Memphis (TN); no difficulty performing activities of daily living, walking 0.25 mile (402 m) or climbing 10 steps without resting; no reported need of assistive devices (e.g. cane, walker); no active treatment for cancer in the prior 3 years; no life-threatening illness; and no plans to leave the area for 3 years |
| Exclusion criteria | Not surviving baseline, diagnosed diabetes, missing HbA1c or FPG values at baseline, without adequate follow-up after baseline |
| Notes | Baseline data for i-IFG (N = 189)/i-HbA1c _{5.7} (N = 207)/IFG/HbA1c (N = 169) cohorts |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|---|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria reported |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Not reported |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |

Lipska 2013 (Continued)

| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
|---|--------------|--|
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | i-IFG: FPG 5.6-6.9 and HbA1c < 5.7; i- HbA1c: 5.7-6.4 and FPG > 5.6; IFG and HbA1c: FPG 5.6-6.9 and HbA1c 5.7-6.4 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | Single HbA1c \geq 6.5 (years 2,6,7); self-report of physician diagnosis (annually); antihyperglycaemic medication (years 1,2,4, 6,7) |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Low risk | Multiple covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |

Lipska 2013 (Continued)

| Study confounding: important potential confounders accounted for in study design | Low risk | Yes |
|---|----------|-----------------------------------|
| Study confounding: important potential confounders accounted for in the analysis | Low risk | Yes |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, odds ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multivariable logistic regression |

Liu 2008

| Name of study | None |
|--------------------|--|
| Inclusion criteria | Individuals from the JiangSu province of China, aged 35-74 years, to trace the incidence of CVD and diabetes; individuals participating twice in the study |
| Exclusion criteria | Individuals suffering from cancer, severe disability, severe psychiatric disturbances; indi- viduals with diabetes, missing data |
| Notes | Baseline data for non-diabetic participants (N = 1844); men (N = 788)/women (N = 1056) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |

Liu 2008 (Continued)

| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Not reported |
|---|--------------|---|
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG 5.6-6.9 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; 2-h PG \geq 11.0; antihyperglycaemic medication |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Not reported |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Not reported |

Liu 2008 (Continued)

| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
|---|--------------------|--|
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some covariates included (see Appendix 16 and Appendix 17) |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Some covariates analysed (see Appendix 16 and Appendix 17) |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, incidence rate, relative risk |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Cox proportional hazards regression |
| Liu 2014 | | |
| Name of study | None | |
| Inclusion criteria | Shanghai residents | |
| Exclusion criteria | Not reported | |

Notes Baseline data for the prediabetic cohort converting to T2DM (N = 78)

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|-----------------------|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |

Liu 2014 (Continued)

| Study participation: adequate description of inclusion & exclusion criteria | Unclear risk | Only inclusion criteria reported |
|---|--------------|----------------------------------|
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Unclear risk | "WHO criteria" |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Unclear risk | Scarce data |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Unclear risk | Scarce data; IFG or GT |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Unclear risk | "WHO criteria" |
| Outcome measurement: method of out- come measurement used valid & reliable | Unclear risk | Scarce data |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Not reported |

Liu 2014 (Continued)

| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Not reported |
|---|--------------|----------------------|
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Not reported |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Not reported |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Analysis of variance |

Liu 2016

| Name of study | Beijing Longitudinal Study on Aging (BLSA) |
|--------------------|--|
| Inclusion criteria | Chinese elders free of diabetes at baseline |
| Exclusion criteria | Diabetes at baseline |
| Notes | Baseline data for participants without diabetes at baseline (N = 1857) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|-----------------------|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |

Liu 2016 (Continued)

| Study participation: adequate description of period & recruitment place | Low risk | Yes |
|---|--------------|---|
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | FPG 6.1-6.9 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; self-reported; antihypergly- caemic medication |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |

Liu 2016 (Continued)

| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
|---|--------------|--|
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some covariates included (see Appendix 16 and Appendix 17) |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Some covariates analysed (see Appendix 16 and Appendix 17) |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Hazard ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Subdistribution hazards model |

Liu 2017

| Name of study | China Multicenter Collaborative Study of Cardiovascular Epidemiology (ChinaMUCA) | | |
|---|--|-----------------------|--|
| Inclusion criteria | 2 studies: China Multicenter Collaborative Study of Cardiovascular Epidemiology (Chi- naMUCA) study and the China Cardiovascular Health Study | | |
| Exclusion criteria | Individuals with missing baseline glucose information, individuals from Deyang, Sichuan (earthquake) and individuals with ASCVD at baseline | | |
| Notes | Baseline data for IFG cohort at baseline (N = 3607) | | |
| Risk of bias | | | |
| Bias | Authors' judgement | Support for judgement | |
| Study participation: description of source population or population of interest | Low risk | Yes | |

Liu 2017 (Continued)

| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
|---|--------------|--|
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Low risk | Yes |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Participants lost to follow-up e.g. were younger, had lower BMI levels and higher physical activity levels |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | FBG 5.6-6.9 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FBG \geq 7.0; using insulin/antihypergly- caemic medications; self-reported |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |

Liu 2017 (Continued)

| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes | |
|---|--|--|--|
| Study confounding: important confounders measured | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) | |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes | |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes | |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes | |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported | |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some covariates included (see Appendix 16 and Appendix 17) | |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Some covariates analysed (see Appendix 16 and Appendix 17) | |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Odds ratio | |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Cox proportional hazard regression | |
| Lorenzo 2003 | | | |
| Name of study | San Antonio Heart Study (SAHS) | | |
| Inclusion criteria | Mexican-Americans and non-Hispanic whites participating in a study of type 2 diabetes and cardiovascular disease | | |
| Exclusion criteria | Phase 1 participants (waist circumference was not measured), and those in phase 2 with diabetes at baseline | | |
| Notes | Baseline data for cohort converting to T2DM (N = 195) | | |
| Risk of bias | | | |

Lorenzo 2003 (Continued)

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|---|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Scarce data |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Scarce data |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG 6.1-6.9; IGT: 2-h PG 7.8 to < 11.1 (WHO 1999) |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |

Lorenzo 2003 (Continued)

| Outcome measurement: clear definition of the outcome provided | Low risk | $FPG: \ge 7.0; 2-h PHG: \ge 11.1 (WHO 1999/1985)$ |
|---|--------------|---|
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some covariates included (see Appendix 16 and Appendix 17) |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Some covariates analysed (see Appendix 16 and Appendix 17) |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, odds ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multiple logistic regression (diabetes risk of the metabolic syndrome and components of the metabolic syndrome) |

Lyssenko 2005

| Name of study | Botnia Study |
|--------------------|---|
| Inclusion criteria | People with type 2 diabetes in western Finland were invited to participate together with their family members; nondiabetic individuals were invited (family members or 'controls' (spouses), aged 18-73 years; prospective visits every 2-3 years; at least 2 OGTTs |

Lyssenko 2005 (Continued)

| Exclusion criteria | MODY, individuals with missing data |
|--------------------|--|
| Notes | Baseline data for IFG-IGT individuals who converted to T2DM (N = 86) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|---|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Description of inclusion and exclusion criteria |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Not reported |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG ≥ 6.1 (WHO 1999 criteria) |

Lyssenko 2005 (Continued)

| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
|---|--------------|--|
| Outcome measurement: clear definition of the outcome provided | Low risk | WHO 1999 criteria |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Univariate analyses |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Univariate analyses |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Univariate analyses |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, hazard ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Univariate Cox proportional hazards model (adjusted for BMI) |

Magliano 2008

| Name of study | Australian Diabetes, Obesity and Lifestyle Study (AusDiab) |
|--------------------|---|
| Inclusion criteria | National population-based survey in adults aged ≥ 25 years |
| Exclusion criteria | Participants refusing further contact, deceased, moved overseas or into a nursing facility classified for high care, had a terminal illness |
| Notes | Baseline data for cohort becoming diabetic at follow-up (N = 224/5842) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Scarce data |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Scarce data |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |

Magliano 2008 (Continued)

| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG 6.1-6.9 and 2-h PG < 7.8; IGT: FPG < 7.0 and 2-h PG \leq 7.8 to < 11.1 |
|---|--------------|---|
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | $FPG \ge 7.0$; 2-h PG ≥ 11.1 ; current anti- hyperglycaemic medication |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Low risk | Multiple covariates included (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Low risk | Yes |
| Study confounding: important potential confounders accounted for in the analysis | Low risk | ORs per SD changes in FPG and HbA1c |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, incidence rate per year, odds ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multivariate logistic regression (logFRPG and logHbA1c) |

Man 2017

| Name of study | Singapore Malay Eye Study (SIMES) |
|--------------------|---|
| Inclusion criteria | Malay adults in Singapore aged 40-80 years; SIMES aims to assess the prevalence, inci- dence, progression, associated factors and impact of major eye disease as well as access to eye care by Asian Malays |
| Exclusion criteria | Diabetes, missing data |
| Notes | Baseline data for incident diabetes cohort (N = 127) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Scarce data |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Scarce data |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Low risk | Yes |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |

Man 2017 (Continued)

| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
|---|--------------|--|
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | HbA1c 5.7-6.4; no self-reported diabetes or an- tihyperglycaemic medication |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | Random glucose \geq 11.1 or HbA1c > 6.4; self- reported history or antihyperglycaemic medica- tion |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some covariates included (see Appendix 16 and Appendix 17) |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Not reported |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, incidence rate, risk ratio |

Man 2017 (Continued)

| Statistical analysis & reporting: the statisti- | Low risk | Multivariate analyses using modified Poission |
|---|----------|--|
| cal model is adequate for the design of the | | regression models to estimate adjusted risk ratios |
| study | | |

Marshall 1994

| Name of study | San Luis Valley Diabetes Study |
|--------------------|--|
| Inclusion criteria | The San Luis Valley Diabetes Study determined the prevalence and incidence of NIDDM among Hispanic and non-Hispanic white adults; sample without prior diabetes diagnosis aged 30-74 years; IGT at the initial visit |
| Exclusion criteria | Unavailability of complete data |
| Notes | Baseline data for IGT cohort converting to T2DM (N = 20) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Scarce data |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Scarce data |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |

Marshall 1994 (Continued)

| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
|---|--------------|--|
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IGT: 2-h PG \ge 7.8 to < 11.1 (WHO 1985) |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | 2-h PG ≥ 11.1 (WHO 1985) |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Cumulative incidence |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Cumulative incidence |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Cumulative incidence |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Cumulative incidence |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence |

Marshall 1994 (Continued)

| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence |
|---|--------------|---|
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multiple logistic regression (baseline dietary risk fac- tors to predict the development of diabetes; glucose levels as continuous variables) |

McNeely 2003

| Name of study | Japanese American Community Diabetes Study |
|--------------------|--|
| Inclusion criteria | Second-generation (Nisei) and third-generation (Sansei) Japanese-American participants residing in Kong County, Washington |
| Exclusion criteria | Individuals with diabetes at baseline |
| Notes | Baseline data for cohort converting to T2DM at 5-6 years (N = 50)/10 years (N = 74) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Unclear risk | Scarce data |
| Study participation: adequate description of period & recruitment place | Unclear risk | Scarce data |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |

McNeely 2003 (Continued)

| Study attrition: adequate description of participants lost to follow-up | Low risk | Yes |
|---|--------------|---|
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Some difference reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG ≥ 6.1 to < 7.0; IGT: 2-h PG ≥ 7.8 to < 11.1 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; 2-h PG \geq 11.1; antihypergly- caemic medication prescribed by a physi- cian |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Cumulative incidence |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Cumulative incidence |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Cumulative incidence |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Cumulative incidence |

McNeely 2003 (Continued)

| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence |
|---|--------------|--|
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Logistic regression (ROC-curves, clinical model) |

Meigs 2003

| Name of study | Baltimore Longitudinal Study of Aging (BLSA) |
|--------------------|--|
| Inclusion criteria | Community dwelling volunteers, largely from the Baltimore (MD) and Washington, D.C. areas; primarily white middle- and upper-middle socioeconomic class aged 21-96 years, being examined approximately every 2 years; open cohort design with dropouts replaced (around 1000 persons at each study cycle); attending at least 3 examinations and an OGTT within an 8-year period |
| Exclusion criteria | 2 or fewer OGTTs or > 4 years elapsed between any 2 OGTTs |
| Notes | Baseline data for the IFG-IGT cohort (N = 265); follow-up time: at least 6 years 77%, at least 10 years 44%, at least 16 years 16%, at least 20 years 4.5% |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |

Meigs 2003 (Continued)

| Study attrition: description of attempts to collect information on participants who dropped out | High risk | Scarce data |
|---|--------------|---|
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Scarce data |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG 6.1-6.9 and 2-h PG ≤ 7.8; IGT: FPG < 6.1 and 2-h PG 7.8-11.0; IFG/IGT |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; 2-h PG \geq 11.1 (IFG-IGT: diabetes defined by OGTT) |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence, incidence rates |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Cumulative incidence, incidence rates |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Cumulative incidence, incidence rates |

Meigs 2003 (Continued)

| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Cumulative incidence, incidence rates |
|--|---|---|
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Cumulative incidence, incidence rates |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence, incidence rates |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence, incidence rates |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, incidence rate |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Kaplan-Meier product limit estimates |
| Mohan 2008 | | |
| Name of study | Chennai Urban Population Study-19 (CUF | PS-19) |
| Inclusion criteria | Participants of 2 residential colonies in Chennai, India, representing the middle and lower income groups ≥ 20 years of age | |
| | - | nennai, India, representing the middle and |
| Exclusion criteria | - | nennai, India, representing the middle and |
| | lower income groups ≥ 20 years of age | |
| Exclusion criteria | lower income groups ≥ 20 years of age Individuals with diabetes | |
| Exclusion criteria Notes | lower income groups ≥ 20 years of age Individuals with diabetes | |
| Exclusion criteria Notes Risk of bias | lower income groups ≥ 20 years of age Individuals with diabetes Baseline data for cohort becoming diabetic Authors' judgement | at follow-up (N = 64/476) |
| Exclusion criteria Notes Risk of bias Bias Study participation: description of source | lower income groups ≥ 20 years of age Individuals with diabetes Baseline data for cohort becoming diabetic Authors' judgement Low risk | at follow-up (N = 64/476) Support for judgement |
| Exclusion criteria Notes Risk of bias Bias Study participation: description of source population or population of interest | lower income groups ≥ 20 years of age Individuals with diabetes Baseline data for cohort becoming diabetic Authors' judgement Low risk Low risk | at follow-up (N = 64/476) Support for judgement Yes |

Mohan 2008 (Continued)

| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
|---|--------------|--|
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Low risk | Yes |
| Study attrition: no important differences between participants who completed the study and those who did not | Low risk | Yes |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG ≥ 6.1 to < 7; IGT: 2-h PG ≥ 7. 8 to < 11.1 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | $FPG \ge 7$; 2-h $PG \ge 11.1$ |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence, incidence rate |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Cumulative incidence, incidence rate |
| | | |

Mohan 2008 (Continued)

| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Cumulative incidence, incidence rate |
|---|--------------|---|
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Cumulative incidence, incidence rate |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Cumulative incidence, incidence rate |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence, incidence rate |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence, incidence rate |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, incidence rate |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Cox regression analysis (effects of various risk factors but not intermediate hypergly- caemia on diabetes) |

Motala 2003

| Name of study | None |
|--------------------|---|
| Inclusion criteria | South African Indians, mainly living in Durban (1984); survey to determine the preva- lence of NIDDM among South African Indians; non-pregnant participants > 15 years of age |
| Exclusion criteria | Not reported |
| Notes | Baseline data for responders (both baseline and follow-up examination) (N = 563) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|-----------------------|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |

Motala 2003 (Continued)

| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
|---|--------------|--|
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Unclear risk | Only inclusion criteria reported |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Low risk | Yes |
| Study attrition: no important differences between participants who completed the study and those who did not | Low risk | Yes |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IGT: FPG < 7.8 and 2-h PG 7.8 to < 11.1 (WHO 1985) |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \ge 7.8; 2-h PG \ge 11.1 (WHO 1985) |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |

Motala 2003 (Continued)

| Study confounding: important confounders measured | Unclear risk | Cumulative incidence |
|---|--------------|---|
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Cumulative incidence |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Cumulative incidence |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Cumulative incidence |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Cumulative incidence |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multiple logistic regression (to evaluate the effect of various pre- dictor variables for type 2 diabetes) |

Motta 2010

| Name of study | Italian Longitudinal Study on Aging (ILSA) | |
|---|--|-----------------------|
| Inclusion criteria | Elderly participants aged 65-84 years involved in ILSA studies | |
| Exclusion criteria | Not reported | |
| Notes | No baseline characteristics provided | |
| Risk of bias | | |
| Bias | Authors' judgement | Support for judgement |
| Study participation: description of source population or population of interest | Low risk | Yes |

Motta 2010 (Continued)

| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
|---|--------------|-----------------------------------|
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Unclear risk | Only inclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Not reported |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: 6.1 to < 7.0 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | $FPG \ge 7.0$ |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |

Motta 2010 (Continued)

| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
|---|--------------|----------------------|
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Cumulative incidence |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Cumulative incidence |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Cumulative incidence |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Cumulative incidence |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | t-test |

Mykkänen 1993

| Name of study | None | |
|--------------------|--|--|
| Inclusion criteria | Participants from Kuopio, Finland | |
| Exclusion criteria | Diabetes at baseline, incomplete OGTT at the follow-up examination | |
| Notes | Baseline data for cohort developing T2DM (N = 69) | |
| Risk of bias | | |
| Bias | Authors' judgement Support for judgement | |

Mykkänen 1993 (Continued)

| Study participation: description of source population or population of interest | Low risk | Yes |
|---|--------------|---|
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Scarce data |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Scarce data |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IGT: FPG < 7.8 and 2-h PG 7.8-11.1 (WHO 1985) |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | $FPG \ge 7.8$; 2-h PG ≥ 11.1 (WHO 1985) |

Mykkänen 1993 (Continued)

| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
|---|--------------|--|
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Cumulative incidence |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Cumulative incidence |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Cumulative incidence |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, odds ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Unclear risk | ANCOVA, odds ratios (risk of developing diabetes associated with various risk factors) |

Nakagami 2016

| Name of study | Kurihashi Lifestyle Cohort Study |
|--------------------|--|
| Inclusion criteria | Baseline health check-ups at Kurihashi Hospital |
| Exclusion criteria | People < 30 years or \geq 80 years, diabetes at baseline, people with chronic diseases, missing covariate data |
| Notes | Baseline data for cohort converting to T2DM (N = 99) |

Nakagami 2016 (Continued)

Risk of bias

| Risk of bias | | |
|---|--------------------|--|
| Bias | Authors' judgement | Support for judgement |
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Scarce data |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Scarce data |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | FPG 5.5-6.9; HbA1c 5.7-6.4 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |

Nakagami 2016 (Continued)

| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0, HbA1c \geq 6.5; physician diagnosis of diabetes |
|---|--------------|---|
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, hazard ratio (associated with a 1 SD increase in the levels of FPG or HbA1c) |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Cox proportional hazards models |

Nakanishi 2004

| Name of study | None |
|--------------------|--|
| Inclusion criteria | Employees of Company A, one of the largest building contractors in Japan (in major cities around Japan); Japanese men aged 35-59 years with no prior history of coronary heart disease or stroke |

Nakanishi 2004 (Continued)

| Exclusion criteria | Not participating in all the consecutive annual health examinations |
|--------------------|---|
| Notes | Baseline characteristics for IFG cohort (N = 246) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Unclear risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Scarce data |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Scarce data |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG 6.1-6.9 |

Nakanishi 2004 (Continued)

| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
|---|--------------|---|
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; antihyperglycaemic medication |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, incidence rate, relative risk (adjusted for all other components and clustering of components of the metabolic syndrome at study entry) |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Cox proportional hazards model |

Noda 2010

| Name of study | Japanese Public-Health Center-based prospective (Diabetes) Study (JPHC Study) |
|--------------------|--|
| Inclusion criteria | All registered Japanese inhabitants in 11 public health center areas aged 40-59 years old in cohort I and 40-69 years old in cohort II; inhabitants who received annual health- checkups; authors included those who were 51-70 years of age at the time of the baseline survey of diabetes |
| Exclusion criteria | Missing data, casual blood samples in any of the 2 health check-ups; known diabetes or an FPG of 125 mg/dL or more at baseline |
| Notes | Baseline characteristics for the total cohort (N = 2207) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Unclear risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Scarce data |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Scarce data |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |

Noda 2010 (Continued)

| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
|---|--------------|--|
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | Taken from table 2: FPG levels: IFG 5.6 and 6.1 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; HbA1c \geq 6.1%; self-reported |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Cumulative incidence |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Cumulative incidence |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Cumulative incidence |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Cumulative incidence |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |

Noda 2010 (Continued)

| Statistical analysis & reporting: the statisti- | Low risk | Crude incidence, ROC curves |
|---|----------|-----------------------------|
| cal model is adequate for the design of the | | |
| study | | |

Park 2006

| Name of study | None |
|--------------------|---|
| Inclusion criteria | Korean men employed at a semiconductor manufacturing facility in Korea participating in an annual health examination at a university hospital |
| Exclusion criteria | Diabetes, failing to undergo subsequent examinations within 2 years; missing data |
| Notes | Baseline data for incident diabetic participants with IFG at baseline (N = 40) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Scarce data |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Scarce data |

Park 2006 (Continued)

| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
|---|--------------|--------------------------------------|
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: $FPG \ge 5.6$ |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | $FPG \ge 7.0$ |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence, incidence rate |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence, incidence rate |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence, incidence rate |

Park 2006 (Continued)

| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | | Cumulative incidence, incidence rate |
|---|----------|--|
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Cox proportional hazards models (for sequential changes in FPG levels) |

Peterson 2017

| Name of study | Follow-up of a cohort originally from the population-based Västerbotten Intervention Program (VIP), a strategy to reach all middle-aged persons individually at ages 40, 50 and 60 years, by inviting them to participate in systematic risk factor screening and individual counselling about healthy lifestyle habits; neuropathy study part of the VIP |
|--------------------|--|
| Inclusion criteria | All individuals who became 40, 50 or 60 years and who belonged to the list for a specific primary care centre or lived within the area for that centre |
| Exclusion criteria | People not participating in the neuropathy study |
| Notes | Baseline data for IGT cohort (N = 29) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |

Peterson 2017 (Continued)

| Study attrition: adequate description of participants lost to follow-up | Low risk | Yes |
|---|--------------|---|
| Study attrition: no important differences between participants who completed the study and those who did not | Low risk | Yes |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IGT: FPG < 7.0 and 2-h PG \geq 7.8 to < 11. 1 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | Yes |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | $FPG \ge 7.0; 2-h PG \ge 11.1$ |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Cumulative incidence |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Cumulative incidence |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Cumulative incidence |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Cumulative incidence |

Peterson 2017 (Continued)

| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence |
|---|--------------|----------------------------|
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | ANOVA, regression analyses |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Unclear risk | Cumulative incidence |

Qian 2012

| Name of study | None |
|--------------------|--|
| Inclusion criteria | Shanghai residents |
| Exclusion criteria | Not reported |
| Notes | Baseline data for cohort progressing to T2DM (N = 377) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|----------------------------------|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Unclear risk | Only inclusion criteria reported |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Not reported |

Qian 2012 (Continued)

| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
|---|--------------|---|
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | i-IFG: 6.1-6.9 and 2-h PG < 7.8; i-IGT: < 6.1 and 2-h PG 7.8- 11.0; IFG/IGT: 6.1-6.9 and 2-h PG 7.8-11.0 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG ≥ 7.0; 2-h PG ≥ 11.1 |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Cumulative incidence |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |

Qian 2012 (Continued)

| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
|---|--------------|---|
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Logistic regression (to assess the potential contributing factors to diabetes incidence) |

Rajala 2000

| Name of study | None | |
|--------------------|---|--|
| Inclusion criteria | Inhabitants in Oulu (northern Finland) recruited from the official population register to investigate the prevalence of diabetes and IGT, reasons for early retirement and the prevalence of depression | |
| Exclusion criteria | Previoulsy diagnosed diabetic people | |
| Notes | Only few baseline data for IGT cohort (N = 171); new cases identified by OGTTs in 1994 and 1996-8 | |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|-----------------------|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |

Rajala 2000 (Continued)

| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
|---|--------------|--|
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Prevalence of hypertension was higher among people lost to follow-up |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IGT: 2-h PG 7.8 to < 11.1 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | 2-h PG \ge 11.1; 2 × FPG \ge 6.7 |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence, incidence rate |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |

Rajala 2000 (Continued)

| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
|---|--------------|--|
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Cumulative incidence, incidence rate |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence, incidence rate |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence, incidence rate |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, incidence rate |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multiple logistic regression (for effects of hypertension and an- tihypertensive medications) |

Ramachandran 1986

| ndian individuals with IGT |
|--|
| lot reported |
| aseline data for the diabetic cohort at follow-up (N = 39) |
| Jo |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|-----------------------|
| Study participation: description of source population or population of interest | High risk | Not reported |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Unclear risk | Scarce data |

Ramachandran 1986 (Continued)

| Study participation: adequate description of period & recruitment place | Unclear risk | Scarce data |
|---|--------------|------------------------------------|
| Study participation: adequate description of inclusion & exclusion criteria | Unclear risk | Only inclusion criteria reported |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Not reported |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IGT: 7.8-11.0 (presumed NDDG 1979) |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | 2-h PG > 11.0 (presumed NDDG 1979) |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence |

Ramachandran 1986 (Continued)

| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Cumulative incidence |
|---|--------------|----------------------|
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Cumulative incidence |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Cumulative incidence |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Cumulative incidence |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Unclear risk | Not reported |

Rasmussen 2008

| Bias | Authors' judgement | Support for judgement |
|--------------------|---|-----------------------|
| Risk of bias | | |
| Notes | Baseline data for IFG (N = 607)/IGT cohort (N = 903) | |
| Exclusion criteria | Severe concurrent illness, alcohol abuse or subsequently treated by general practitioners not in the addition study; individuals with diabetes | |
| Inclusion criteria | Population-based high-risk screening and intervention study for type 2 diabetes; persons aged 40-69 years registered with the participating practices in 5 counties in Denmark with a risk score of 5 points or more; measurement of fasting capillary blood glucose and OGTT; annual glucose measurement recommended for individuals with IFG and IGT; individuals with 2 diabetic glucose values on separate days were included in the intervention programme | |
| Name of study | Anglo-Danish-Dutch study of Intensive Treatment in People with Screen Detected Di- abetes in Primary Care (ADDITION) | |

Rasmussen 2008 (Continued)

| Study participation: description of source population or population of interest | Low risk | Yes |
|---|--------------|--|
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Scarce data |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Low risk | Yes |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Unclear risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG (i-IFG): FBG 5.6 to < 6.1 and 2-h BG < 7.8; IGT (i-IGT): FBG < 6.1 and 2-h BG 7.8 to < 11.1; IFG/IGT |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Unclear risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | $FBG \ge 6.1 \text{ or } 2\text{-h } BG \ge 11.1$ |

Rasmussen 2008 (Continued)

| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
|---|--|--|
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence, incidence rate |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence, incidence rate |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence, incidence rate |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, incidence rate |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Regression models (for sequential changes in some covariates) |
| Rathmann 2009 | | |
| Name of study | Kooperative Gesundheitsforschung in der Region Augsburg (KORA S4/F4) | |
| Inclusion criteria | People living in Augsburg and surroundings; KORA was follow-up of MONICA WHO- Project (Monitoring Trends and determinants in Cardiovascular Disease); S1: 25-64 years, S2/S3/S4: 25-74 years | |

| | years, 52/55/54: 2)-/4 years |
|--------------------|------------------------------|
| Exclusion criteria | People with known diabetes |

Rathmann 2009 (Continued)

Notes Baseline characteristics for total cohort (participants of the follow-up; age-group 55-74 years; N = 887)

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|---|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Low risk | Yes |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Some differences reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG 6.1-6.9; IGT: 2-h PG 7.8 to < 11.1; 'prediabetes': i-IFG, i-IGT and IFG/ IGT |

Rathmann 2009 (Continued)

| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
|---|--------------|---|
| Outcome measurement: clear definition of the outcome provided | Low risk | $FPG \ge 7.0$; 2-h PG ≥ 11.1 ; validated physician diagnosis |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some covariates included (see Appendix 16 and Appendix 17) |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Some covariates analyses (see Appendix 16 and Appendix 17) |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, incidence rate, odds ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Logistic regression models |

Rijkelijkhuizen 2007

| Name of study | Hoorn Study |
|--------------------|---|
| Inclusion criteria | General Dutch population (Hoorn) aged 50-75 years at baseline; participants completing both measurements in 1989 and 1996 |
| Exclusion criteria | People using antihyperglycaemic medications or diet for diabetes were marked as known diabetes mellitus; missing information of plasma glucose values |
| Notes | Baseline data for IFG _{6.1} (N = 149)/IFG _{5.6} (N = 488) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Low risk | Yes |
| Study attrition: no important differences between participants who completed the study and those who did not | Low risk | No substantial differences |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |

Rijkelijkhuizen 2007 (Continued)

| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
|---|--------------|---|
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG _{5.6} : FPG 5.6-7.0; IFG _{6.1} : FPG 6.1-7.0; IGT: 2-h PG 7.8 to < 11.1 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | $FPG \ge 7.0; 2-h PG: \ge 11.1$ |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some covariates included (see Appendix 16 and Appendix 17) |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Some covariates analysed (see Appendix 16 and Appendix 17) |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, incidence rate, odds ratio |

Rijkelijkhuizen 2007 (Continued)

| Statistical analysis & reporting: the statisti- | Low risk | Cox proportional hazards models |
|---|----------|---------------------------------|
| cal model is adequate for the design of the | | |
| study | | |

Sadeghi 2015

| Name of study | Isfahan Cohort Study (ICS), baseline survey of the Isfahan Healthy Heart Program (IHHP) |
|--------------------|---|
| Inclusion criteria | Participants of the baseline survey of the Isfahan Healthy Heart Program, a community trial for prevention and control of CVD |
| Exclusion criteria | Diabetes at baseline |
| Notes | Baseline data for prediabetic cohort at baseline becoming diabetic at follow-up (N = 131) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Scarce data |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the | Low risk | Yes |

Sadeghi 2015 (Continued)

| study and those who did not | | |
|---|--------------|---|
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG \geq 5.5 and < 7.0; IGT: 2-h OGTT \geq 7.8 and < 11.1 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG > 7.0; 2-h OGTT > 11.1; IFG/IGT; antihyperglycaemic medication |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Low risk | Stochastic regression methods |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some covariates included (see Appendix 16 and Appendix 17) |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Some covariates analysed (see Appendix 16 and Appendix 17) |

Sadeghi 2015 (Continued)

| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | | Cumulative incidence, incidence rate, odds ratio |
|---|----------|---|
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multivariate logistic regression |

Sasaki 1982

| Name of study | None |
|--------------------|---|
| Inclusion criteria | Epidemiological survey on diabetes mellitus in Osaka, Japan and follow-up study |
| Exclusion criteria | Not reported |
| Notes | Baseline data for the IGT cohort (N = 13) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|-----------------------------------|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Unclear risk | Only inclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Scarce data |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |

Sasaki 1982 (Continued)

| Study attrition: no important differences between participants who completed the study and those who did not | Low risk | Yes |
|---|--------------|---|
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IGT: FPG < 7.8 and 2-h PG 7.8-11.1 (WHO 1980) |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | $FPG \ge 7.8 \text{ or } 2\text{-h } PG \ge 11.1 \text{ (WHO } 1980)$ |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Scarce data |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Scarce data |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Scarce data |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence |

Sasaki 1982 (Continued)

| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence, incidence rate |
|---|--------------|---|
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Unclear risk | Multiple logistic regression (standardised regression coefficients for single covariates) |

Sato 2009

| Name of study | Kansai Healthcare Study |
|--------------------|---|
| Inclusion criteria | Japanese male employees of a company in the area of Kansai, aged 40-55 years, not taking an oral antihyperglycaemic or insulin at study entry and considered to be involved in sedentary jobs |
| Exclusion criteria | Not reported |
| Notes | Baseline data for cohort becoming diabetic at follow-up (N = 659/6804); non-standard categories for elevated HbA1c values were used (Table 1, p 645 of the publication) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|----------------------------------|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Unclear risk | Only inclusion criteria reported |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Scarce data |

Sato 2009 (Continued)

| Study attrition: reasons for loss to follow- up provided | Unclear risk | Scarce data |
|---|--------------|---|
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | Table 1: IFG: FPG group 6.1-6.9; HbA1c-group: 6.0-6.4 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; antihyperglycaemic medication |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Low risk | Yes |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |

Sato 2009 (Continued)

| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
|---|--------------|--|
| Study confounding: important potential confounders accounted for in study design | Low risk | Yes |
| Study confounding: important potential confounders accounted for in the analysis | Low risk | Yes |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Odds ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multiple logistic regression (FPG, HbA1c categories) |

Schranz 1989

| Name of study | Study within the WHO-assisted National Diabetes Programme |
|--------------------|--|
| Inclusion criteria | Within the framework of the WHO-assisted National Diabetes Programme a cohort of Maltese people was investigated |
| Exclusion criteria | Known diabetic persons |
| Notes | Baseline data for diabetic cohort at follow-up (N = 166) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|-----------------------|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Unclear risk | Scarce data |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Yes |

Schranz 1989 (Continued)

| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Not reported |
|---|--------------|---|
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IGT: 2-h PG \geq 7.8 to < 11.1 (WHO 1985) |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | 2-h PG ≥ 11.1 (WHO 1985) |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Cumulative incidence |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Cumulative incidence |

Schranz 1989 (Continued)

| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Cumulative incidence |
|---|--------------|----------------------|
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Cumulative incidence |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Unclear risk | Not reported |

Sharifi 2013

| Name of study | Zanjan Healthy Heart Study |
|--------------------|--|
| Inclusion criteria | Participants from the Zanjan Healthy Heart Study, aged 21-75 years, individuals with IFG |
| Exclusion criteria | Not reported |
| Notes | Baseline data for active participants (N = 123) of the IFG cohort |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|-----------------------|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |

Sharifi 2013 (Continued)

| Study participation: adequate description of inclusion & exclusion criteria | Unclear risk | Only inclusion criteria reported |
|---|--------------|---|
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | High attrition rate (> 50%) |
| Study attrition: no important differences between participants who completed the study and those who did not | Low risk | Yes |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | FPG 5.6-7.0 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG > 7.0 (2 measurements); diabetes diagnosis based on documents |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |

Sharifi 2013 (Continued)

| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
|---|--------------|---|
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Unclear risk | Logistic regression (BMI and physical activity for pre- diction of diabetes) |

Shin 1997

| Name of study | Yonchon study |
|--------------------|---|
| Inclusion criteria | Individuals living in Yonchon County (South Korea), free of diabetes aged \geq 30 years |
| Exclusion criteria | Diabetes |
| Notes | Baseline data for individuals converting to T2DM (N = 67) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|-----------------------|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |

Shin 1997 (Continued)

| Study participation: adequate description of period & recruitment place | Low risk | Yes |
|---|--------------|---|
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Scarce data |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Scarce data |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Unclear risk | Scarce data |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Unclear risk | Scarce data |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Unclear risk | Assumed WHO 1985 criteria |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Unclear risk | Scarce data |
| Outcome measurement: clear definition of the outcome provided | Low risk | "WHO criteria"; antihyperglycaemic medication |
| Outcome measurement: method of out- come measurement used valid & reliable | Unclear risk | Scarce data |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Unclear risk | Scarce data |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence |

Shin 1997 (Continued)

| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Yes |
|---|--------------|---|
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multiple logistic regression (1 mmol/L difference for FPG and 2-h plasma glucose) |

Song 2015

| Name of study | Korean Genome Epidemiology Study-Kangwha Study (KoGES) |
|--------------------|---|
| Inclusion criteria | People aged ≥ 40 years |
| Exclusion criteria | Missing key variables, history of stroke, angina pectoris or myocardial infarction, diabetes |
| Notes | Baseline data for prediabetic cohort (men: N = 154; women: N = 167; total: N = 321); ranges for men - women |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|-----------------------|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |

Song 2015 (Continued)

| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
|---|--------------|--|
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Responders had relatively low FPG and HbA1c at baseline compared to non-re- sponders |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG 5.6-6.9 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; HbA1c \geq 6.5; antihypergly- caemic medication |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |

Song 2015 (Continued)

| Study confounding: important confounders measured | Low risk | Yes |
|---|--------------|-------------------------------------|
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Low risk | Yes |
| Study confounding: important potential confounders accounted for in the analysis | Low risk | Yes |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Unclear risk | Cumulative incidence, relative risk |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Generalised linear models |

Song 2016a

| Name of study | None | | |
|---|---|-----|--|
| Inclusion criteria | Survey of the prevalence of T2DM in an urban community; eligible permanent inhabi- tants 15-74 years | | |
| Exclusion criteria | Not reported | | |
| Notes | Baseline data for prediabetic cohort (N = 334) | | |
| Risk of bias | | | |
| Bias | Authors' judgement Support for judgement | | |
| Study participation: description of source population or population of interest | Low risk | Yes | |

Song 2016a (Continued)

| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
|---|--------------|--|
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Unclear risk | Only inclusion criteria reported |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Low risk | Yes |
| Study attrition: no important differences between participants who completed the study and those who did not | Low risk | Yes |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FG 5.6-6.9; IGT: 2-h G 7.8-11.0 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | IFG \geq 7.0; 2-h G \geq 11.0; HbA1c \geq 6.5; self-reported |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |

Song 2016a (Continued)

| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
|---|--|---|
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Logistic regression models (sex-related risk factors associated with the development of diabetes) |
| Soriguer 2008 | | |
| Name of study | Pizarra study, evaluating the prevalence of latent autoimmune diabetes of adults (LADA) in the context of the overall prevalence of diabetes in Southern Spain | |
| Inclusion criteria | People aged 18-65 years from Pizarra, Malaga | |
| Exclusion criteria | Institutionalised persons, pregnant women, severe clinical or psychological disorder | |
| Notes | Baseline data for final sample of follow-up (N = 714); diabetes diagnosis according to capillary blood glucose levels > 6.1 mmol/L or post OGTT BG > 11.1 mmol/L | |
| Risk of bias | | |
| | | |

Soriguer 2008 (Continued)

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Low risk | Yes |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Scarce data |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Unclear risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: BG 5.6-6.1 and 2-h BG < 7.8; IGT: BG < 5.6 and 2-h BG 7.8-11.1 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |

Soriguer 2008 (Continued)

| Outcome measurement: clear definition of the outcome provided | Low risk | BG > 6.1 or 2-h BG > 11.1 |
|---|--------------|---|
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some covariates included (see Appendix 16 and Appendix 17) |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Some covariates analysed (see Appendix 16 and Appendix 17) |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, incidence rate, rel- ative risk |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multivariate logistic regression |

Stengard 1992

| Name of study | Finnish Cohorts of the Seven Countries Study |
|--------------------|---|
| Inclusion criteria | Elderly Finnish men, survivors of the Finnish cohorts of the Seven-Countries Study (studying mortality, morbidity and risk factor levels of cardiovascular diseases in different countries), aged 65-84 years at baseline |

Stengard 1992 (Continued)

| Exclusion criteria | Not reported | | |
|--|--|----------------------------------|--|
| Notes | Baseline data for IGT cohort converting to T2DM (N = 17) | | |
| Risk of bias | | | |
| Bias | Authors' judgement | Support for judgement | |
| Study participation: description of source population or population of interest | Low risk | Yes | |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes | |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes | |
| Study participation: adequate description of period & recruitment place | Low risk | Yes | |
| Study participation: adequate description of inclusion & exclusion criteria | Unclear risk | Only inclusion criteria reported | |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Scarce data | |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Scarce data | |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported | |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported | |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes | |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes | |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IGT: 2-h PG 7.8-11.1 | |

Stengard 1992 (Continued)

| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
|---|--------------|--|
| Outcome measurement: clear definition of the outcome provided | Low risk | 2-h PG \geq 11.1 (WHO 1985); antihyper- glycaemic medications |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some covariates included (see Appendix 16 and Appendix 17) |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Some covariates analysed (see Appendix 16 and Appendix 17) |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, odds ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multiple logistic regression |

Söderberg 2004

| Name of study | None |
|--------------------|---|
| Inclusion criteria | Population based survey in Mauritius, 3 cohorts of nonpregnant participants aged 25- 79 years with classifiable data from 2 separate surveys |
| Exclusion criteria | Not reported |
| Notes | Baseline data for cohort 1987-1998 (N = 2631), 10 years follow-up; 3 cohorts 1987- 1992 (N = 3680), 1992-1998 (N = 4178), 1987-1998 (N = 2631) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|----------------------------------|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Unclear risk | Only inclusion criteria reported |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Scarce data |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Scarce data |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |

Söderberg 2004 (Continued)

| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
|---|--------------|---|
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG \geq 6.1 to < 7.0 and 2-h PG < 7.8; IGT: FPF < 7.0 and 2-h PG \geq 7.8 to < 11.1 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG ≥ 7.0; 2-h PG ≥ 11.1 |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence, incidence rate |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Cumulative incidence, incidence rate |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Cumulative incidence, incidence rate |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Cumulative incidence, incidence rate |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Cumulative incidence, incidence rate |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence, incidence rate |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence, incidence rate |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, incidence rate |

Söderberg 2004 (Continued)

| Statistical analysis & reporting: the statisti- | Low risk | Calculation of incidence rate ratios, Poisson regression analysis |
|---|----------|---|
| cal model is adequate for the design of the | | to estimate sex effects between 1987 and 1998 allowing for ad- |
| study | | justments |

Toshihiro 2008

| Name of study | None |
|--------------------|--|
| Inclusion criteria | Japanese mal workers of a railroad company receiving a health-check at Nishimatsuzono Clinic, IFG and/or IGT cohort |
| Exclusion criteria | People with type B or C hepatitis virus infections |
| Notes | Baseline data for cohort becoming diabetic at follow-up (N = 36/128);participants with IFG and/or IGT were given advice about lifestyle modifications once or twice a year |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Not reported |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the | Unclear risk | Not reported |

Toshihiro 2008 (Continued)

| study and those who did not | | |
|---|--------------|---|
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG 6.1-6.9 and 2-h PG < 7.8; IGT: FPG < 7.0 and 2-h PG 7.8-11.1 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | $FPG \ge 7.0$; 2-h PG > 11.1; non-fasting PG > 11.1 |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence |

Toshihiro 2008 (Continued)

| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | | Cumulative incidence |
|---|----------|--|
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Cox proportional hazards model (multivariate analysis of inde- pendent risk factors and recovery factors) |

Vaccaro 1999

| Name of study | None |
|--------------------|--|
| Inclusion criteria | Telephone company employees in the age range 40-59 years were screened in the province of Naples for major cardiovascular risk factors |
| Exclusion criteria | Taking antihyperglycaemic medication, previous diabetes diagnosis |
| Notes | Baseline data for total cohort (follow-up examination; N = 560) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |

Vaccaro 1999 (Continued)

| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Those lost to follow-up were older and more frequently women |
|---|--------------|--|
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Unclear risk | Unusual thresholds |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Unclear risk | IFG: FPG 5.6-6.0; IGT: 2-h PG 6.7-9.9 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; antihyperglycaemic medication |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Not reported |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Not reported |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Not reported |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Not reported |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Not reported |

Vaccaro 1999 (Continued)

| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Not reported |
|---|--------------|--|
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, odds ratio (probably unadjusted) |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Unclear risk | Quote: "standard methods" |

Valdes 2008

| Name of study | Asturias Study (Asturias) |
|--------------------|--|
| Inclusion criteria | Survey of diabetes and cardiovascular risk factors in the principality of Asturias, northern Spain; participants from basic health area |
| Exclusion criteria | Type 1 diabetes, pregnancy, severe disease, hospitalisation, use of diabetogenic drugs, missing data; diabetes |
| Notes | Baseline data for IFG 5.6-6.1 (N = 114)/IFG 6.1-6.9 (N = 52) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |

Valdes 2008 (Continued)

| Study attrition: reasons for loss to follow- | Low risk | Yes |
|---|--------------|---|
| up provided | | |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG _{5.6} : 5.6-6.1; IFG _{6.1} : 6.1-6.9 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; 2-h PG \geq 11.1; clinical diabetes diagnosis; an- tihyperglycaemic medication, diet |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |

Valdes 2008 (Continued)

| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
|---|--------------|--|
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some covariates included (see Appendix 16 and Appendix 17) |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Some covariates analysed (see Appendix 16 and Appendix 17) |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, incidence rate, odds ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multivariate logistic regression |

Vijayakumar 2017

| Name of study | None |
|--------------------|---|
| Inclusion criteria | Participants were 10-19 years of age at first examination without diabetes, and at least 1 follow-up examination before the 40th birthday |
| Exclusion criteria | History of possibly taking metformin at baseline |
| Notes | Baseline data for adults (A)/children (C) with HbA1c 5.7-6.4 (children: N = 62, adults: N = 168) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |

Vijayakumar 2017 (Continued)

| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Not reported |
|---|--------------|---|
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | FPG 5.6-6.9; 2-h PG 7.8-11.9; HbA1c 5.7-6.4 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; 2-h PG \geq 11.1; previous clinical diagnosis |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence, incidence rate |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Cumulative incidence, incidence rate |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Cumulative incidence, incidence rate |

Vijayakumar 2017 (Continued)

| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Cumulative incidence, incidence rate |
|---|--------------|--|
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Cumulative incidence, incidence rate |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence, incidence rate |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence, incidence rate |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, incidence rate |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | ROC curves, increments in HbA1c and FPG or 2-h PG to cal- culate 10-year cumulative incidence |

Viswanathan 2007

| Name of study | None |
|--------------------|--|
| Inclusion criteria | Programme on primary prevention of diabetes in the population and in high risk people (positive family history of diabetes); individuals with at least 2 follow-up visits; partici- pants were given advice on preventive measures such as dietary modifications and regular exercise |
| Exclusion criteria | Known history of diabetes, newly diagnosed diabetes during screening |
| Notes | Baseline data for IGT group (N = 619); participants were given advice on preventive measures such as dietary modifications and regular exercise |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|-----------------------|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Unclear risk | Scarce data |

Viswanathan 2007 (Continued)

| Study participation: adequate description of period & recruitment place | Unclear risk | Scarce data |
|---|--------------|--|
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Not reported |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IGT: 2-h PG 7.8 to < 11.1 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Unclear risk | Not defined, presumably by OGTT |
| Outcome measurement: method of out- come measurement used valid & reliable | Unclear risk | Scarce data |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Unclear risk | Scarce data |
| Study confounding: important confounders measured | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |

Viswanathan 2007 (Continued)

| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Not reported |
|---|--------------|--|
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Not reported |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some covariates included (see Appendix 16 and Appendix 17) |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Some covariates analysed (see Appendix 16 and Appendix 17) |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, odds ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multiple logistic regression, Cox regression analysis |

Wang 2007

| Name of study | Beijing Project as part of the National Diabetes Survey |
|--------------------|---|
| Inclusion criteria | Inhabitants of Beijing aged 25 years or older |
| Exclusion criteria | Newly diagnosed diabetes or CHD at baseline, known diabetes |
| Notes | Baseline data for cohort with incident diabetes and no CHD (N = 67) |
| | |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|-----------------------|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |

Wang 2007 (Continued)

| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
|---|----------|--|
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Low risk | Yes |
| Study attrition: no important differences between participants who completed the study and those who did not | Low risk | Yes |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG 6.1-6.9; IGT: 2-h PG 7.8-11.0 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | $FPG \ge 7.0; 2-h PG \ge 11.1$ |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |

Wang 2007 (Continued)

| Study confounding: important confounders measured | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |
|---|--------------|--|
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some covariates included (see Appendix 16 and Appendix 17) |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Some covariates analysed (see Appendix 16 and Appendix 17) |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, risk ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Multiple logistic regression |

Wang 2011

| Name of study | Strong Heart Study (SHS) | |
|---|--|-----------------------|
| Inclusion criteria | Data collected from American Indians at the baseline and second exams from those participants who had HbA1c and FPG measured | |
| Exclusion criteria | Antihyperglycaemic medications, renal dialysis, kidney transplant | |
| Notes | No baseline data reported | |
| Risk of bias | | |
| Bias | Authors' judgement | Support for judgement |
| Study participation: description of source population or population of interest | Low risk | Yes |

Wang 2011 (Continued)

| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
|---|--------------|--|
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Scarce data |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Those lost to follow-up had lower BMI |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: 5.6 to < 7.0; HbA1c 6.0 to < 6.5 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | $FPG \ge 7.0$; HbA1c ≥ 6.5 ; FPG/HbA1c: ≥ 6.5 or FPG \ge 7.0; antihyperglycaemic medication |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |

Wang 2011 (Continued)

| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
|---|--------------|--|
| Study confounding: important confounders measured | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some covariates included (see Appendix 16 and Appendix 17) |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Some covariates analysed (see Appendix 16 and Appendix 17) |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Odds ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Logistic regression |

Warren 2017

| Name of study | Atherosclerosis Risk in Communities study (ARIC) |
|--------------------|--|
| Inclusion criteria | Adults aged 45-64 years from the communities of Jackson, MS; Forsyth County, NC; suburban Minneapolis, MN; and Washington County, MD, USA |
| Exclusion criteria | Participants with prevalent diabetes, chronic kidney disease, atherosclerotic cardiovascu- lar disease, or peripheral arterial disease, those who were missing variables of interest, or those who fasted for < 10 h |
| Notes | 2 different baseline cohorts; 4 prediabetes definitions (visit 2: IFG 5.6-6.9: N = 4112; HbA1c 5.7-6.4: N = 2027; visit 4: IFG 5.6-6.9: N = 2142; IGT: N = 2009) |

Warren 2017 (Continued)

Risk of bias

| Risk of bias | | |
|---|--------------------|--|
| Bias | Authors' judgement | Support for judgement |
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Scarce data |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | FPG 5.6-6.9 (ADA); FG 6.1-6.9 (WHO) ; 2-h 7.8-11.0 (ADA); HbA1c 5.7-6.4 (ADA); 6.0-6.4 (IEC) |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |

Warren 2017 (Continued)

| Outcome measurement: clear definition of the outcome provided | Unclear risk | Self-report of physician diagnosis; antihy- perglycaemic medication reported during a study visit or annual telephone call |
|---|--------------|--|
| Outcome measurement: method of out- come measurement used valid & reliable | Unclear risk | Missing lab measurements |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some covariates included (see Appendix 16 and Appendix 17) |
| Study confounding: important potential confounders accounted for in the analysis | Low risk | Some covariates analysed (see Appendix 16 and Appendix 17) |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Hazard ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Cox proportional hazards models |

Wat 2001

| Name of study | Hong Kong Cardiovascular Risk Factor Prevalence Study |
|--------------------|--|
| Inclusion criteria | Follow-up of the Hong Kong Cardiovascular Risk Factor Prevalence Study in Hong Kong Chinese aged 25-74 years; persons with IGT (matched controls from the same population with normal glucose tolerance), investigation of the development of appropriate population-wide coronary heart disease prevention strategies and monitoring their long-term impact |
| Exclusion criteria | Diabetes at baseline |
| Notes | Baseline data for IGT cohort (N = 322) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Scarce data |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Scarce data |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |

Wat 2001 (Continued)

| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
|---|--------------|---|
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IGT: FPG < 7.8 and 2-h PG 7.8 to < 11.1 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | $FPG \ge 7.8; 2-h PG \ge 11.1$ |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |

Wat 2001 (Continued)

| Statistical analysis & reporting: the statisti- | Low risk | Logistic regression (per unit increase for |
|---|----------|--|
| cal model is adequate for the design of the | | some covariates) |
| study | | |

Weiss 2005

| Name of study | None |
|--------------------|--|
| Inclusion criteria | Obese children and adolescents aged 4-18 years were recruited from the Yale Pediatric Obesity Clinic (New Haven, Conneticut, USA) |
| Exclusion criteria | Participants with medical conditions, using medications that may affect glucose metabolism before their first OGTT |
| Notes | Baseline data for IGT cohort (N = 33) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|--|--------------------|---|
| Study participation: description of source population or population of interest | Unclear risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Unclear risk | Scarce data |
| Study participation: adequate description of period & recruitment place | Unclear risk | Scarce data |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria reported |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | No dropouts |
| Study attrition: reasons for loss to follow- up provided | Low risk | No dropouts |
| Study attrition: adequate description of participants lost to follow-up | Low risk | No dropouts |
| Study attrition: no important differences between participants who completed the study and those who did not | Low risk | No dropouts |

Weiss 2005 (Continued)

| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
|---|--------------|--|
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IGT: FPG < 5.6 and 2-h PG 7.8-11.1 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; 2-h PG > 11.1; presentation of hyperglycaemia (more than 2 random glucose measurements > 11.1), glucosuria, polydipsia, and polyuria |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Cumulative incidence |
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Cumulative incidence |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Cumulative incidence |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence |

Weiss 2005 (Continued)

| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence |
|---|--------------|---|
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Mann-Whitney U test and linear regression (to identify predic- tors of 2-h glucose on the second OGTT) |

Wheelock 2016

| Name of study | Pima Indian Study (Gila River Indian Community - near Phoenix, Arizona) |
|--------------------|---|
| Inclusion criteria | Gila River Indian Community in Arizona (mostly Pima or Tohono Indians); children and adolescents 5-19 years who were nondiabetic at baseline and had at least 1 follow- up examination |
| Exclusion criteria | Not reported |
| Notes | Baseline data for the full cohort (N = 5532); prediabetic cohort = non-overweight (N = 37) + IGT group and overweight + IGT group (N = 132); 5-11 years/12-19 years); age-stratified incidence data on overweight participants + IGT <i>or</i> overweight and either hypertension or hypercholesterolaemia + IGT (metabolic set (MSet)) |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|-----------------------------------|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Unclear risk | Only inclusion criteria described |
| Study attrition: description of attempts to collect information on participants who | Unclear risk | Scarce data |

Wheelock 2016 (Continued)

| dropped out | | |
|---|--------------|--|
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Scarce data |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IGT: 2-h PG ≥ 7.8 to < 11.1 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | $FPG \ge 7.0$; 2-h PG ≥ 11.1 ; previous diagnosis |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |

Wheelock 2016 (Continued)

| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
|---|--------------|--|
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Cumulative incidence |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Cox regression model using each metabolic risk factor as a continuous variable; viola- tion of the proportionality assumption was noted, therefore cumulative incidence rates were calculated from a Poisson regression model |

Wong 2003

| Name of study | Singapore Impaired Glucose Tolerance Follow-up Study | |
|--------------------|---|--|
| Inclusion criteria | Representative sample of the Singapore population aged 18-69 years; persons with IGT and matched controls | |
| Exclusion criteria | Antihyperglycaemic medication, venepuncture failure; persons with IFG | |
| Notes | Baseline data for IGT group (N = 291) | |

Risk of bias

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|-----------------------|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |

Wong 2003 (Continued)

| Study participation: adequate description of period & recruitment place | Low risk | Yes |
|---|--------------|---|
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Scarce data |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Scarce data |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IGT: 2-h PG \ge 7.8 to < 11.1 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG ≥ 7.0; 2-h PG ≥ 11.1; physician di- agnosed diabetes |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Cumulative incidence |

Wong 2003 (Continued)

| Study confounding: clear definitions of im- portant confounders provided | Unclear risk | Cumulative incidence |
|---|--------------|---|
| Study confounding: measurement of con- founders valid & reliable | Unclear risk | Cumulative incidence |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Unclear risk | Cumulative incidence |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Cumulative incidence |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Cumulative incidence |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Not reported |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | ANCOVA using general linear models (comparisons between continuous vari- ables) |

Yeboah 2011

| á years |
|---|
| art failure, cedure for pacemaker |
| |
| |
| - |

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|-----------------------|
| Study participation: description of source population or population of interest | Low risk | Yes |

Yeboah 2011 (Continued)

| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
|---|--------------|--|
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Low risk | Yes |
| Study attrition: reasons for loss to follow- up provided | Low risk | Yes |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Scarce data |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Scarce data |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IFG: FPG 5.6-6.9 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Low risk | Yes |
| Outcome measurement: clear definition of the outcome provided | Low risk | FPG > 6.9; antihyperglycaemic medication during examinations 2,3,4 |
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |

Yeboah 2011 (Continued)

| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
|---|--------------|------------------------------------|
| Study confounding: important confounders measured | Low risk | Yes |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Low risk | Yes |
| Study confounding: important potential confounders accounted for in the analysis | Low risk | Yes |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Cumulative incidence, hazard ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Cox proportional hazards model |

Zethelius 2004

| Name of study | None |
|--------------------|---|
| Inclusion criteria | All men residing in Uppsala were invited to a health survey in 1970; reinvestigation 20 years later (= baseline) at 70 years of age |
| Exclusion criteria | Diabetes, antihyperglycaemic medications |
| Notes | Baseline data for cohort converting to T2DM (N = 26) |

Risk of bias

Zethelius 2004 (Continued)

| Bias | Authors' judgement | Support for judgement |
|---|--------------------|--|
| Study participation: description of source population or population of interest | Low risk | Yes |
| Study participation: description of gly- caemic status at baseline | Low risk | Yes |
| Study participation: adequate description of sampling frame & recruitment | Low risk | Yes |
| Study participation: adequate description of period & recruitment place | Low risk | Yes |
| Study participation: adequate description of inclusion & exclusion criteria | Low risk | Inclusion and exclusion criteria described |
| Study attrition: description of attempts to collect information on participants who dropped out | Unclear risk | Not reported |
| Study attrition: reasons for loss to follow- up provided | Unclear risk | Not reported |
| Study attrition: adequate description of participants lost to follow-up | Unclear risk | Not reported |
| Study attrition: no important differences between participants who completed the study and those who did not | Unclear risk | Not reported |
| Glycaemic status measurement: provision of clear definition or description of gly- caemic status | Low risk | Yes |
| Glycaemic status measurement: valid and reliable method of glycaemic status mea- surement | Low risk | Yes |
| Glycaemic status measurement: continu- ous variables reported or appropriate cut points used | Low risk | IGT: 2-h PG 7.8 to < 11.1 |
| Glycaemic status measurement: same method and setting of measurement of the glycaemic status for all study participants | Unclear risk | Yes |

Zethelius 2004 (Continued)

| Outcome measurement: clear definition of the outcome provided | Low risk | FPG \geq 7.0; antihyperglycaemic medications |
|---|--------------|---|
| Outcome measurement: method of out- come measurement used valid & reliable | Low risk | Yes |
| Outcome measurement: same method & setting of outcome measurement for all study participants | Low risk | Yes |
| Study confounding: important confounders measured | Unclear risk | Some covariates measured (see Appendix 16 and Appendix 17) |
| Study confounding: clear definitions of im- portant confounders provided | Low risk | Yes |
| Study confounding: measurement of con- founders valid & reliable | Low risk | Yes |
| Study confounding: same method & set- ting for measurements of confounders for all study participants | Low risk | Yes |
| Study confounding: appropriate methods used if missing confounder data imputed | Unclear risk | Not reported |
| Study confounding: important potential confounders accounted for in study design | Unclear risk | Some covariates included (see Appendix 16 and Appendix 17) |
| Study confounding: important potential confounders accounted for in the analysis | Unclear risk | Some covariates analysed (see Appendix 16 and Appendix 17) |
| Statistical analysis & reporting: sufficient presentation of data to assess adequacy of the analytic strategy | Low risk | Odds ratio |
| Statistical analysis & reporting: the statisti- cal model is adequate for the design of the study | Low risk | Logistic regression, multivariate models (adjusted for BMI, age at baseline and length of follow-up) |

Note: for better readability all IFG/IGT and HbA1c measurements are reported in numerical format only (IFG and IGT were measured in mmol/L, HbA1c was measured in %)

ADA: American Diabetes Association; **ANOVA**: analysis of variance; **BG**: blood glucose; **BMI**: body mass index; **CHD**: coronary heart disease; **CI**: confidence interval; **CVD**: cardiovascular disease; **FG**: fasting glucose; **FBG**: fasting blood glucose; **FINDRISC**: Finnish Diabetes Risk Score; **FPG**: fasting plasma glucose; **G6PD**: glucose-6-P-dehydrogenase test; **HbA1c**: glycosylated haemoglobin A1c; **HbA1c**_{5.7}: intermediate hyperglycaemia with HbA1c 5.7% as lower threshold (usually reflecting 5.7%-6.4%); **HbA1c**_{6.0}: intermediate hyperglycaemia with HbA1c 6.0% as lower threshold (usually reflecting 6.0%-6.4%); **HOMA-B**: homeostatic model assessment

beta-cell function; HOMA-IR: homeostatic model assessment for insulin resistance; HR: hazard ratio; IEC: International Expert Committee; IFG: impaired fasting glucose; IFG_{5.6}: impaired fasting glucose with 5.6 mmol/L as lower threshold; IFG_{6.1}: impaired fasting glucose with 6.1 mmol/L as lower threshold; IFG/IGT: both IFG and IGT; i-IFG: isolated IFG; IGT: impaired glucose tolerance; i-IGT: isolated IGT; JDS: Japanese Diabetes Society; MSet: metabolic set; NDDG: National Diabetes Data Group; NGSP: National Glycohemoglobin Standardization Program; NGT: normal glucose tolerance; OGTT: oral glucose tolerance test; OR: odds ratio; PG: postload glucose; ROC: receiver operating characteristics; RR: risk ratio, relative risk; T2DM: type 2 diabetes mellitus; WHO: World Health Organization.

Characteristics of excluded studies [ordered by study ID]

| Study | Reason for exclusion | |
|---|---|--|
| Abdul-Ghani 2011 | No data on transition from intermediate hyperglycaemia to type 2 diabetes | |
| Alvarsson 2009 | Intervention study | |
| Alyass 2015 | No data on transition from intermediate hyperglycaemia to type 2 diabetes | |
| Amoah 2002 | Not a prospective cohort study | |
| Andreou 2017 | No data on transition from intermediate hyperglycaemia to type 2 diabetes (prevalence data) | |
| Bancks 2015 | Only self-reported diabetes, frequency matched population | |
| Birmingham Diabetes Survey Working Party 1976 | Non-standard thresholds for intermediate hyperglycaemia | |
| Bjornholt 2000 | No data on transition from intermediate hyperglycaemia to type 2 diabetes | |
| Bodicoat 2017 | Long-term follow-up of an interventional study | |
| Boned 2016 | Hypertensive cohort | |
| Boucher 2015 | No data on transition from intermediate hyperglycaemia to type 2 diabetes | |
| Brantsma 2005 | No data on transition from intermediate hyperglycaemia to type 2 diabetes | |
| Brateanu 2017 | Retrospective cohort study | |
| Braun 1996 | No data on transition from intermediate hyperglycaemia to type 2 diabetes | |
| Burchfiel 1995 | No cohort with intermediate hyperglycaemia | |
| Chamukuttan 2016 | Intervention trial | |
| Chang 2017 | Investigation of the association between thyroid function and the development of intermediate hyperglycaemia/diabetes | |

| Chen 1995 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
|------------------|--|
| Cheng 2011 | Not a prospective cohort study |
| Cheung 2007 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Choi 2002 | Not a prospective cohort study |
| Cicero 2005 | No valid data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Cosson 2011 | Not a prospective cohort study |
| Costa 2005 | Study design paper |
| Cree-Green 2013 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Cropano 2017 | Investigation of the association between gene variants and development of in- termediate hyperglycaemia/diabetes |
| Dagogo-Jack 2011 | Evaluation of the transition from normoglycaemia to intermediate hypergly- caemia |
| Daniel 1999 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Decode 2003 | Aggregate data of 22 cohorts; no data on transition from intermediate hyper- glycaemia to type 2 diabetes |
| Deedwania 2013 | No data on diabetes incidence |
| DeFina 2012 | Not a prospective cohort study |
| DeJesus 2016 | Not a prospective cohort study |
| Deschenes 2016 | Cohort with depressive symptoms |
| Dinneen 1998 | Not a prospective cohort study |
| Doi 2007 | No cohort with intermediate hyperglycaemia |
| Du 2016 | Cross-sectional study, no cohort with intermediate hyperglycaemia |
| Edelman 2004 | Non-standard thresholds for intermediate hyperglycaemia |
| Edelstein 1997 | Aggregated data on 6 prospective studies, no reliable additional data on transi- tion from intermediate hyperglycaemia to type 2 diabetes |
| Engberg 2010 | Intervention trial |

| Eskesen 2013 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
|---------------------------|--|
| Feizi 2017 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Feskens 1989 | No cohort with intermediate hyperglycaemia |
| Festa 2003 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Folsom 2000 | No cohort with intermediate hyperglycaemia |
| Gil-Montalban 2015 | Diagnosis of type 2 diabetes incidence by database only |
| Giraldez-Garcia 2015 | No data on type 2 diabetes incidence |
| Glauber 2018 | Incidence established by register data |
| Gonzalez-Villalpando 2014 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Gopinath 2013 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Gu 2015 | No data on transition from intermediate hyperglycaemia to type 2 diabetes (database) |
| Gupta 2011 | Intervention trial, hypertensive cohort |
| Hackett 2014 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Haffner 1997 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Haffner 2000 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Hajat 2012 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Hanai 2005 | No data on transition from intermediate hyperglycaemia to type 2 diabetes, OGTTs were unit of analysis |
| He 2018 | Investigation of the association of glycaemic index diets and glycaemic load diets with development of type 2 diabetes |
| Helmrich 1991 | No cohort with intermediate hyperglycaemia |
| Henninger 2015 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Holbrook 1990 | No cohort with intermediate hyperglycaemia |
| | |

| Huang 2014c | Not a prospective cohort study (database) |
|-----------------------|---|
| Hulman 2017 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Inoue 2008 | Retrospective cohort study |
| Invitti 2006 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Jallut 1990 | Not a prospective cohort study |
| James 1998 | No cohort with intermediate hyperglycaemia |
| Jansson 2015 | No cohort with intermediate hyperglycaemia |
| Jarrett 1979 | Intervention trial |
| Jarrett 1982 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Jeanne 2018 | No cohort with intermediate hyperglycaemia, investigation of the association between birth weight and physical activity and cardiometabolic health |
| Jiamjarasrangsi 2008b | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Joshipura 2017 | Diabetes incidence data for 'prediabetes' group only |
| Kadowaki 1984 | Non-standard thresholds for intermediate hyperglycaemia |
| Kametani 2002 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Kanauchi 2003 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Kanaya 2005 | Investigation of a prediction model for development of diabetes |
| Kawahara 2015 | Not a prospective cohort study |
| Khan 2017 | Diabetes incidence defined by register data |
| Khang 2010 | Not a prospective cohort study |
| Kieboom 2017 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Kim 2012a | Not a prospective cohort study |
| Kim 2012b | Not a prospective cohort study |
| Kim 2013 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| | |

| Kim 2016b | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
|-----------------|--|
| Kim 2017a | Investigation of the association between sleep duration and development of type 2 diabetes |
| Kim 2017b | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Ko 2000 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Kosaka 1996 | Non-standard thresholds, no numerical data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Kowall 2013 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Krabbe 2017 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Le Boudec 2016 | Withdrawn publication |
| Lee 2014 | No cohort with intermediate hyperglycaemia |
| Lee 2017 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Leite 2009 | Intervention trial |
| Li 2011 | Evaluation of a diabetes risk tool |
| Liatis 2014 | Participants of a diabetes prevention programme |
| Libman 2008 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Liu 2017a | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Liu 2017b | Investigation of the association between the bone resorption marker CTX and incident intermediate hyperglycaemia/diabetes |
| Malmstrom 2018 | Type 2 diabetes incidence measured mainly through registers; nested case-con- trol study; no transition data |
| Manson 1992 | No cohort with intermediate hyperglycaemia |
| McNeill 2006 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| McPhillips 1990 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Medalie 1975 | No data on transition from intermediate hyperglycaemia to type 2 diabetes; no common thresholds for diagnosis of intermediate hyperglycaemia and type 2 diabetes |

| Metcalf 2017 | No cohort with intermediate hyperglycaemia |
|----------------|--|
| Miranda 2017 | Investigation of the association between advanced glycation end products (AGE) and their receptor (RAGE) and type 2 diabetes incidence |
| Mirbolouk 2016 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Monesi 2012 | No cohort with intermediate hyperglycaemia |
| Morrison 2012 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Nakagami 2017 | No cohort with intermediate hyperglycaemia |
| Nakasone 2017 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Nano 2017 | Investigation of the association between liver transaminases and development of intermediate hyperglycaemia/type 2 diabetes |
| Nguyen 2014 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Nichols 2007 | Not a prospective cohort study |
| Nichols 2010 | Not a prospective cohort study |
| Nichols 2015 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Njolstad 1998 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Norberg 2006 | Not a prospective cohort study |
| Nowicka 2011 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Ohlson 1987 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Oizumi 2011 | Non-standard thresholds for intermediate hyperglycaemia |
| Okada 2017 | Diabetes incidence data for prediabetic cohort only (FPG 5.6-6.9 or HbA1c 5. 7%-6.4%) |
| Onat 2007 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Onat 2013a | Non-standard IFG/IGT definition |
| Onat 2013b | Non-standard IFG/IGT definition |
| Osei 2004 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |

| Paddock 2017 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
|-------------------|---|
| Perry 1995 | Type 2 diabetes mellitus incidence not established by glucose measurements (questionnaires, reviews of primary care records, reviews of death certificates) |
| Pinelli 2011 | Cross-sectional study |
| Polakowska 2011 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Pradhan 2007 | Intervention trial (Women's Health Study) |
| Priya 2013 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Qiao 2003 | Not a prospective cohort study |
| Qiu 2015 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Ramachandran 2012 | Not a prospective cohort study |
| Rauh 2017 | Development of a prediction model for HbA1c levels after 6 years in the non- diabetic general population |
| Reynolds 2006 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Rimm 1995 | No cohort with intermediate hyperglycaemia |
| Sacks 2017 | Investigation of patient activation to predict the course of type 2 diabetes |
| Sai 2017 | No cohort with intermediate hyperglycaemia |
| Samaras 2015 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Schmitz 2016 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Schottker 2011 | Diabetes incidence by self-report only |
| Schulze 2008 | Evaluation of a diabetes risk score |
| Schwarz 2007 | No individuals with intermediate hyperglycaemia at baseline |
| Serrano 2013 | Study design paper |
| Shimazaki 2007 | Not a prospective cohort study |
| Song 2007 | Mix of old an new participants in 2 study phases, participants with with both IFG and IGT were combined into an IFG group |

| Song 2016b | Not a prospective cohort study |
|----------------------|--|
| Sorgjerd 2015 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Soria 2009 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Stampfer 1988 | No cohort with intermediate hyperglycaemia |
| Strauss 1974 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Suvitaival 2018 | Evaluation of a new biomarker ('plasma lipidome') model |
| Tabak 2009 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Tai 2004 | Aggregated data from several prevalence and incidence studies |
| Takkunen 2016 | Cohort from intervention trial, no data on cohort with intermediate hypergly- caemia |
| Tanabe 2009 | Not a prospective cohort study |
| Vaccaro 2005 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Vaidya 2016 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Vazquez 2000 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Vega-Vázquez 2017 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Von Eckardstein 2000 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Wang 2010 | New diabetes cases were identified through hospital records only |
| Warram 1996 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Wei 1999 | Investigation of the association between cardiorespiratory fitness and interme- diate hyperglycaemia/type 2 diabetes mellitus |
| Welborn 1979 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Wheeler 2017 | Investigation of genetic determinants of HbA1c on the development of type 2 diabetes |
| Wingard 1993 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Woo 2015 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| | |

| Wu 2017a | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
|----------------|--|
| Wu 2017b | Intermediate hyperglycaemia determined through register data, retrospective study |
| Wu 2018 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Xu 2014 | Investigation of a prediction model for development of diabetes |
| Yang 2016 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Ye 2014 | No data on people with intermediate hyperglycaemia |
| Yi 2017 | No data on type 2 diabetes incidence |
| Yokota 2017 | Retrospective cohort study |
| Yoshinaga 1996 | Non-standard thresholds for intermediate hyperglycaemia |
| Yoshinaga 1999 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Zargar 2001 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Zethelius 2008 | No data on transition from intermediate hyperglycaemia to type 2 diabetes, establishment of a predictive model |
| Zhang 2012b | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Zhang 2016 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |
| Zimmet 1992 | No data on transition from intermediate hyperglycaemia to type 2 diabetes |

FPG: fasting plasma glucose; HbA1c: glycosylated haemoglobin A1c; IFG: impaired fasting glucose; IGT: impaired glucose tolerance.

Characteristics of studies awaiting assessment [ordered by study ID]

| Li 2001 | |
|---------------------|---|
| Study name | Model development of diabetes in adult Chinese |
| Starting date | 1986, follow-up 6 years |
| Contact information | Guangwei Li, Department of Endocrinology, China-Japan Friendship Hospital, Beijing 100029 China |

Li 2001 (Continued)

| Notes | Establishment of a model for type 2 diabetes and the roles of insulin resistance and insulin secretion impair- ment; needs translation | | | | | | |
|---------------------|---|--|--|--|--|--|--|
| Misnikova 2011 | | | | | | | |
| Study name | Risk of diabetes and cardiovascular events in persons with early glucose metabolism impairments | | | | | | |
| Starting date | 2006, follow-up 3 years | | | | | | |
| Contact information | Misnikova IV, Endocrinology, Moscow Regional Research Clinical Institute, Russian Federation | | | | | | |
| Notes | Conference abstract, no publication available | | | | | | |
| NCT00816608 | | | | | | | |
| Study name | The effect of maximum body weight in lifetime on the development of type 2 diabetes (MAXWEL) | | | | | | |
| Starting date | August 2006 | | | | | | |
| Contact information | Professor Soo Lim, Seoul National University Bundang Hospital | | | | | | |
| Notes | Study completion date: September 2013; no publication available | | | | | | |

Characteristics of ongoing studies [ordered by study ID]

NCT00786890

| Trial name or title | A survey to evaluate the cardiovascular risk status of subjects with pre-diabetes in Hong Kong (JADE-HK2) |
|---------------------|---|
| Starting date | November 2008 |
| Contact information | Juliana Chan, Professor, Chinese University of Hong Kong |
| Notes | Estimated study completion date: December 2018 |

NCT02838693

| Trial name or title | Assessing progression to type-2 diabetes (APT-2D): a prospective cohort study expanded from BRITE-SPOT (Bio-bank and Registry for StratIfication and Targeted intErventions in the Spectrum Of Type 2 Diabetes) (APT-2D) |
|---------------------|--|
| Starting date | March 2016 |
| Contact information | Sue-Anne Toh, MBBChir, MSc, MA; +65 67722195; mdcsates@nus.edu.sg |
| | Sue-Anne Toh, MBBChir, MSc, MA; +65 67722195; mdcsates@nus.edu.sg |

NCT02838693 (Continued)

| Notes | Estimated study completion date: December 2021 |
|---------------------|---|
| NCT02958579 | |
| Trial name or title | A population based study on metabolic syndrome complications, and mortality (MetSCoM) |
| Starting date | January 2005 |
| Contact information | Alireza Esteghamati, MD (esteghamati@tums.ac.ir); Zahra Aryan, MD, MPH (aryanzahra@yahoo.com) |
| Notes | Estimated study completion date: January 2020 |
| Vilanova 2017 | |
| Trial name or title | Prevalence, clinical features and risk assessment of pre-diabetes in Spain: the prospective Mollerussa cohort study |
| Starting date | August 2011 |
| Contact information | Dr Didac Mauricio, MD; didacmauricio@gmail.com |

| Notes | The Mollerussa study completed its recruitment phase in July 2014 and the 12 month follow-up in July 2015. |
|-------|--|
| | Participants will be followed up long-term through annual extraction of data included in the individual's |
| | electronic medical records |

DATA AND ANALYSES

| Outcome or subgroup title | No. of studies | No. of participants | Statistical method | Effect size |
|---|-------------------|------------------------|-------------------------------|----------------------|
| 1 T2DM incidence (IFG _{5.6}) | 8 | 34867 | Hazard Ratio (Random, 95% CI) | 4.32 [2.61, 7.12] |
| 1.1 Asia/Middle East | 4 | 14803 | Hazard Ratio (Random, 95% CI) | 5.07 [3.41, 7.53] |
| 1.2 Australia/Europe/North | 3 | 18522 | Hazard Ratio (Random, 95% CI) | 4.15 [1.24, 13.87] |
| America | | | | |
| 1.3 American Indians/Islands | 1 | 1542 | Hazard Ratio (Random, 95% CI) | 2.38 [1.85, 3.06] |
| 2 T2DM incidence (IFG _{6.1}) | 10 | 21475 | Hazard Ratio (Random, 95% CI) | 5.47 [3.50, 8.54] |
| 2.1 Asia/Middle East | 5 | 10810 | Hazard Ratio (Random, 95% CI) | 10.55 [3.61, 30.81] |
| 2.2 Australia/Europe/North | 4 | 10571 | Hazard Ratio (Random, 95% CI) | 3.30 [2.32, 4.67] |
| America | | | | |
| 2.3 Latin America | 1 | 94 | Hazard Ratio (Random, 95% CI) | 2.06 [1.76, 2.41] |
| 3 T2DM incidence (IGT) | 5 | 16576 | Hazard Ratio (Random, 95% CI) | 3.61 [2.31, 5.64] |
| 3.1 Asia/Middle East | 3 | 8475 | Hazard Ratio (Random, 95% CI) | 4.48 [2.81, 7.15] |
| 3.2 Australia/Europe/North | 2 | 8101 | Hazard Ratio (Random, 95% CI) | 2.53 [1.52, 4.19] |
| America | | | | |
| 4 T2DM incidence (IFG + IGT) | 5 | 9757 | Hazard Ratio (Random, 95% CI) | 6.90 [4.15, 11.45] |
| 4.1 Asia/Middle East | 3 | 7156 | Hazard Ratio (Random, 95% CI) | 10.20 [5.45, 19.09] |
| 4.2 Australia/Europe/North | 1 | 1650 | Hazard Ratio (Random, 95% CI) | 3.80 [2.30, 6.28] |
| America | | | | |
| 4.3 American Indians/Islands | 1 | 951 | Hazard Ratio (Random, 95% CI) | 4.06 [3.05, 5.40] |
| 5 T2DM incidence (HbA1c _{5.7}) | 4 | 25047 | Hazard Ratio (Random, 95% CI) | 5.55 [2.77, 11.12] |
| 5.1 Asia | 3 | 16805 | Hazard Ratio (Random, 95% CI) | 7.21 [5.14, 10.11] |
| 5.2 Australia/Europe/North | 1 | 8242 | Hazard Ratio (Random, 95% CI) | 2.71 [2.48, 2.96] |
| America | | | | |
| 6 T2DM incidence (HbA1c _{6.0}) | 6 | 30699 | Hazard Ratio (Random, 95% CI) | 10.10 [3.59, 28.43] |
| 6.1 Asia/Middle East | 4 | 22734 | Hazard Ratio (Random, 95% CI) | 13.12 [4.10, 41.96] |
| 6.2 Australia/Europe/North | 2 | 7965 | Hazard Ratio (Random, 95% CI) | 5.09 [1.69, 15.37] |
| America | | | | |
| 7 T2DM incidence (HbA1c + | 1 | | Hazard Ratio (Fixed, 95% CI) | Subtotals only |
| IFG) | | | | 2 |
| 7.1 HbA1c _{5.7} + IFG _{5.6} | 1 | 4559 | Hazard Ratio (Fixed, 95% CI) | 32.50 [23.00, 45.92] |
| 7.2 HbA1 $c_{5.7}$ + IFG _{6.1} | 1 | 5357 | Hazard Ratio (Fixed, 95% CI) | 37.90 [28.10, 51.12] |
| 7.3 HbA1 $c_{6.0}$ + IFG _{5.6} | 1 | 4628 | Hazard Ratio (Fixed, 95% CI) | 53.70 [38.40, 75.09] |
| 7.4 HbA1 $c_{6.0}$ + IFG _{6.1} | 1 | 5802 | Hazard Ratio (Fixed, 95% CI) | 52.30 [37.80, 72.37] |

Comparison 1. Hazard ratio as the effect measure for the development of T2DM

| Comparison 2. Ouus ratio as the chect measure for the development of 12D | Comparison 2. | Odds ratio as the effect measure for the developm | ent of T2DM |
|--|---------------|---|-------------|
|--|---------------|---|-------------|

| Outcome or subgroup title | No. of studies | No. of participants | Statistical method | Effect size |
|--|-------------------|------------------------|-----------------------------|----------------------|
| 1 T2DM incidence (IFG _{5.6}) | 21 | 47647 | Odds Ratio (Random, 95% CI) | 4.15 [2.75, 6.28] |
| 1.1 Asia/Middle East | 10 | 34577 | Odds Ratio (Random, 95% CI) | 2.94 [1.77, 4.86] |
| 1.2 Australia/Europe/North | 9 | 9869 | Odds Ratio (Random, 95% CI) | 6.47 [3.81, 11.00] |
| America | | | | |
| 1.3 Latin America | 1 | 1659 | Odds Ratio (Random, 95% CI) | 4.28 [3.21, 5.71] |
| 1.4 American Indians/Islands | 1 | 1542 | Odds Ratio (Random, 95% CI) | 3.12 [2.31, 4.21] |
| 2 T2DM incidence (IFG _{6.1}) | 15 | 36866 | Odds Ratio (Random, 95% CI) | 6.60 [4.18, 10.43] |
| 2.1 Asia/Middle East | 7 | 28921 | Odds Ratio (Random, 95% CI) | 5.18 [2.32, 11.53] |
| 2.2 Australia/Europe/North | 7 | 6334 | Odds Ratio (Random, 95% CI) | 8.69 [4.95, 15.24] |
| America | | 1(11 | | |
| 2.3 Latin America | 1 | 1611 | Odds Ratio (Random, 95% CI) | 3.73 [2.18, 6.38] |
| 3 T2DM incidence (IGT) | 20 | 21552 | Odds Ratio (Random, 95% CI) | 4.61 [3.76, 5.64] |
| 3.1 Asia/Middle East | 6 | 8643 | Odds Ratio (Random, 95% CI) | 3.74 [2.83, 4.94] |
| 3.2 Australia/Europe/North America | 11 | 9165 | Odds Ratio (Random, 95% CI) | 5.20 [3.62, 7.45] |
| 3.3 Latin America | 2 | 3478 | Odds Ratio (Random, 95% CI) | 4.94 [3.15, 7.76] |
| 3.4 American Indians/Islands | 1 | 266 | Odds Ratio (Random, 95% CI) | 3.60 [1.40, 9.26] |
| 4 T2DM incidence (IFG + IGT) | 9 | 9656 | Odds Ratio (Random, 95% CI) | 13.14 [7.41, 23.30] |
| 4.1 Asia/Middle East | 3 | 4202 | Odds Ratio (Random, 95% CI) | 6.99 [3.09, 15.83] |
| 4.2 Australia/Europe/North | 6 | 5454 | Odds Ratio (Random, 95% CI) | 20.95 [12.40, 35.40] |
| America | | | | |
| 5 T2DM incidence (HbA1c _{5.7}) | 3 | 3468 | Odds Ratio (Random, 95% CI) | 4.43 [2.20, 8.88] |
| 5.1 Asia/Middle East | 1 | 1137 | Odds Ratio (Random, 95% CI) | 4.54 [2.65, 7.78] |
| 5.2 Europe/North America | 2 | 2331 | Odds Ratio (Random, 95% CI) | 4.38 [1.36, 14.15] |
| 6 T2DM incidence (HbA1c _{6.0}) | 3 | 18317 | Odds Ratio (Random, 95% CI) | 12.79 [4.56, 35.85] |
| 6.1 Asia/Middle East | 1 | 11866 | Odds Ratio (Random, 95% CI) | 23.20 [18.70, 28.78] |
| 6.2 Australia/Europe/North America | 1 | 5735 | Odds Ratio (Random, 95% CI) | 15.60 [6.90, 35.27] |
| 6.3 American Indians/Islands | 1 | 716 | Odds Ratio (Random, 95% CI) | 5.89 [4.23, 8.20] |
| 7 T2DM incidence (HbA1c _{5.7} + | 2 | 14006 | Odds Ratio (Random, 95% CI) | 35.91 [20.43, 63.12] |
| IFG _{5.6}) | | | | |
| 7.1 Australia/Europe/North America | 1 | 1294 | Odds Ratio (Random, 95% CI) | 26.20 [16.30, 42.11] |
| 7.2 Asia/Middle East | 1 | 12712 | Odds Ratio (Random, 95% CI) | 46.70 [33.60, 64.91] |

Analysis I.I. Comparison I Hazard ratio as the effect measure for the development of T2DM, Outcome I T2DM incidence (IFG_{5.6}).

Review: Development of type 2 diabetes mellitus in people with intermediate hyperglycaemia

Comparison: I Hazard ratio as the effect measure for the development of T2DM

Outcome: I T2DM incidence (IFG_{5.6})

| Study or subgroup | Intermediate hypergly- caemia N | IFG _{5.6} N | log [Hazard Ratio] (SE) | | izard Ratio om,95% Cl | Weight | Hazard Ratio IV,Random,95% Cl |
|---|--|-------------------------|---|----------------|--------------------------|---------------|----------------------------------|
| I Asia/Middle East | | | | | | | |
| Heianza 2012 (1) | 1680 | 4149 | 1.8213 (0.1803) | | | 13.4 % | 6.18 [4.34, 8.80] |
| Kim 2005 (2) | 276 | 2009 | 1.5623 (0.5573) | | _ | 8.6 % | 4.77 [1.60, 14.22] |
| Janghorbani 2015 (3) | 230 | 627 | 2.0015 (0.3537) | | | 11.3 % | 7.40 [3.70, 14.80] |
| Han 2017 (4) | 199 | 5633 | 1.2837 (0.1206) | | - | 14.0 % | 3.61 [2.85, 4.57] |
| Subtotal (95% CI) | 2385 | 12418 | | | • | 47.2 % | 5.07 [3.41, 7.53] |
| Heterogeneity: Tau ² = 0.09; C Test for overall effect: Z = 8.0 2 Australia/Europe/North Am Yeboah 2011 (5) | 4 (P < 0.00001) | 6215 | 2.3514 (0.1139) | | -4 | 14.0 % | 10.50 [8.40, 13.13] |
| Forouhi 2007 (6) | 633 | 407 | 1.0647 (0.4094) | | _ | 10.5 % | 2.90 [1.30, 6.47] |
| Warren 2017 (7) | 4112 | 6215 | 0.8154 (0.0423) | | - | 14.3 % | 2.26 [2.08, 2.46] |
| Subtotal (95% CI) | 5685 | 12837 | | | | 38.9 % | 4.15 [1.24, 13.87] |
| Heterogeneity: Tau ² = 1.08; C Test for overall effect: Z = 2.3 3 American Indians/Islands Wang 2011 (8) | | = 2 (P<0.000 | 01); 1 ² =99% 0.8671 (0.1285) | | - | 13.9 % | 2.38 [1.85, 3.06] |
| Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: Z = 6.7 | 947 5 (P < 0.00001) | 595 | | | * | 13.9 % | 2.38 [1.85, 3.06] |
| Total (95% CI) | 9017 | 25850 | | | • | 100.0 % | 4.32 [2.61, 7.12] |
| Heterogeneity: $Tau^2 = 0.45$; C Test for overall effect: $Z = 5.7$ Test for subgroup differences: | 2 (P < 0.00001) | | | | | | |
| | | | | 0.1 0.2 0.5 | 1 2 5 10 |) | |
| | | | | Normoglycaemia | IFG5.6 | | |

- (1) 5 years follow-up
- (2) 5 years follow-up
- (3) 7 years follow-up
- (4) 12 years follow-up
- (5) 8 years follow-up
- (6) 10 years follow-up
- (7) 22 years follow-up
- (8) 4 years follow-up

Analysis I.2. Comparison I Hazard ratio as the effect measure for the development of T2DM, Outcome 2 T2DM incidence (IFG_{6.1}).

Review: Development of type 2 diabetes mellitus in people with intermediate hyperglycaemia

Comparison: I Hazard ratio as the effect measure for the development of T2DM

Outcome: 2 T2DM incidence (IFG_{6.1})

| Study or subgroup | IFG _{6.1} N | Normoglycaemia N | log [Hazard Ratio] (SE) | Hazard Ratio IV,Random,95% Cl | Weight | Hazard Ratio IV,Random,95% Cl |
|--------------------------------|--------------------------|-----------------------|----------------------------|--|--------|----------------------------------|
| I Asia/Middle East | | | | | | |
| Heianza 2012 (1) | 380 | 4149 | 2.4336 (0.175) | + | 11.0 % | .40 [8.09, 6.06] |
| Kim 2005 (2) | 276 | 2009 | 3.543 (0.5323) | | 7.1 % | 34.57 [12.18, 98.13] |
| Li 2003 (3) | 42 | 435 | 1.7544 (0.3017) | | 9.7 % | 5.78 [3.20, 10.44] |
| Nakagami 2016 (4) | 134 | 1528 | 3.5522 (0.2929) | - | 9.8 % | 34.89 [19.65, 61.95] |
| Liu 2016 (5) | 222 | 1635 | 0.6881 (0.1905) | + | 10.8 % | 1.99 [1.37, 2.89] |
| Subtotal (95% CI) | 1054 | 9756 | | • | 48.5 % | 10.55 [3.61, 30.81] |
| Heterogeneity: $Tau^2 = 1.4$ | 0; Chi ² = 90 | .70, df = 4 (P<0.0000 | I); I ² =96% | | | |
| Test for overall effect: $Z =$ | 4.31 (P = 0. | 000017) | | | | |
| 2 Australia/Europe/North | America | | | | | |
| Lyssenko 2005 (6) | 211 | 1503 | 0.8329 (0.2533) | | 10.2 % | 2.30 [1.40, 3.78] |
| Forouhi 2007 (7) | 257 | 407 | 1.4816 (0.4285) | | 8.3 % | 4.40 [1.90, 10.19] |
| Bonora 2011 (8) | 55 | 710 | 1.763 (0.3013) | | 9.7 % | 5.83 [3.23, 10.52] |
| Warren 2017 (9) | 1213 | 6215 | 1.0473 (0.0468) | - | 11.7 % | 2.85 [2.60, 3.12] |
| | | | ſ | 0.01 0.1 1 10 Normoglycaemia IFG _{6.1} | 100 | |

(Continued . . .)

| | | | | | | (Continued) |
|--|---------------------------|---------------------------------------|--------------------------|------------------|---------------|---------------------|
| Study or subgroup | IFG _{6.1} | Normoglycaemia | log [Hazard Ratio] | Hazard Ratio | Weight | Hazard Ratio |
| | Ν | Ν | (SE) | IV,Random,95% CI | | IV,Random,95% CI |
| Subtotal (95% CI) | 1736 | 8835 | | • | 39.9 % | 3.30 [2.32, 4.67] |
| Heterogeneity: $Tau^2 = 0.07$ | 7; Chi ² = 7.2 | 29, df = 3 (P = 0.06); l ² | =59% | | | |
| Test for overall effect: Z = | 6.69 (P < 0. | 00001) | | | | |
| 3 Latin America | | | | | | |
| Leiva 2014 (10) | 28 | 66 | 0.7227 (0.0803) | - | 11.6 % | 2.06 [1.76, 2.41] |
| Subtotal (95% CI) | 28 | 66 | | • | 11.6 % | 2.06 [1.76, 2.41] |
| Heterogeneity: not applicat | ble | | | | | |
| Test for overall effect: Z = | 9.00 (P < 0. | 00001) | | | | |
| Total (95% CI) | 2818 | 18657 | | • | 100.0 % | 5.47 [3.50, 8.54] |
| Heterogeneity: Tau ² = 0.44 | 1; Chi ² = 18 | 8.70, df = 9 (P<0.0000 |)); ² =95% | | | |
| Test for overall effect: $Z =$ | 7.48 (P < 0. | 00001) | | | | |
| Test for subgroup difference | es: $Chi^2 = I$ | 3.70, df = 2 (P = 0.00) |), l ² =85% | | | |
| | | | | | | |

0.01 0.1 1 10 100

Normoglycaemia IFG_{6.1}

(1) 5 years follow-up

(2) 5 years follow-up

(3) 5 years follow-up

(4) 5 years follow-up

(5) II years follow-up

(6) 6 years follow-up; isolated IFG $_{6.1}$; univariate analysis

(7) 10 years follow-up

(8) 15 years follow-up

(9) 22 years follow-up

(10) 6 years follow-up

Analysis I.3. Comparison I Hazard ratio as the effect measure for the development of T2DM, Outcome 3 T2DM incidence (IGT).

Review: Development of type 2 diabetes mellitus in people with intermediate hyperglycaemia

Comparison: I Hazard ratio as the effect measure for the development of T2DM

Outcome: 3 T2DM incidence (IGT)

| Study or subgroup | IGT N | Normoglycaemia N | log [Hazard Ratio] (SE) | Hazard Ratio IV,Random,95% Cl | Weight | Hazard Ratio IV,Random,95% Cl | | | |
|--|--|---------------------------------------|----------------------------|----------------------------------|---------|----------------------------------|--|--|--|
| Asia/Middle East | | | | | | | | | |
| Li 2003 (I) | 118 | 435 | 1.0784 (0.2475) | - | 18.7 % | 2.94 [1.81, 4.78] | | | |
| Janghorbani 2015 (2) | 150 | 627 | 2.2407 (0.3429) | | 15.6 % | 9.40 [4.80, 18.41] | | | |
| Han 2017 (3) | 1512 | 5633 | 1.4012 (0.0585) | - | 23.7 % | 4.06 [3.62, 4.55] | | | |
| Subtotal (95% CI) | 1780 | 6695 | | * | 58.0 % | 4.48 [2.81, 7.15] | | | |
| Heterogeneity: Tau ² = 0.12 | Heterogeneity: Tau ² = 0.12; Chi ² = 7.68, df = 2 (P = 0.02); I ² = 74% | | | | | | | | |
| Test for overall effect: $Z = $ | 6.30 (P < 0.0 | 0001) | | | | | | | |
| 2 Australia/Europe/North A | America | | | | | | | | |
| Lyssenko 2005 (4) | 221 | 1429 | 1.2528 (0.2606) | + | 18.3 % | 3.50 [2.10, 5.83] | | | |
| Warren 2017 (5) | 2009 | 4442 | 0.7227 (0.0576) | - | 23.7 % | 2.06 [1.84, 2.31] | | | |
| Subtotal (95% CI) | 2230 | 5871 | | • | 42.0 % | 2.53 [1.52, 4.19] | | | |
| Heterogeneity: $Tau^2 = 0.10$ |); Chi ² = 3.95 | 5, df = 1 (P = 0.05); I^2 | =75% | | | | | | |
| Test for overall effect: $Z = 2$ | 3.59 (P = 0.0 | 0033) | | | | | | | |
| Total (95% CI) | 4010 | 12566 | | • | 100.0 % | 3.61 [2.31, 5.64] | | | |
| Heterogeneity: Tau ² = 0.22 | 2; Chi ² = 80.5 | 52, df = 4 (P<0.00001) | ; I ² =95% | | | | | | |
| Test for overall effect: $Z = $ | 5.63 (P < 0.0 | 0001) | | | | | | | |
| Test for subgroup difference | es: Chi ² = 2. | 67, df = 1 (P = 0.10), l ⁴ | 2 =62% | | | | | | |
| | | | | | | | | | |

0.01 0.1 1 10 100

Normoglycaemia IGT

(1) 5 years follow-up; isolated IGT

(2) 7 years follow-up

(3) 12 years follow-up; isolated IGT

(4) 6 years follow-up

(5) 16 years follow-up

Analysis I.4. Comparison I Hazard ratio as the effect measure for the development of T2DM, Outcome 4 T2DM incidence (IFG + IGT).

Review: Development of type 2 diabetes mellitus in people with intermediate hyperglycaemia

Comparison: I Hazard ratio as the effect measure for the development of T2DM

Outcome: 4 T2DM incidence (IFG + IGT)

| Study or subgroup | IFG+IGT N | Normoglycaemia N | log [Hazard Ratio] (SE) | | zard Ratio m,95% Cl | Weight | Hazard Ratio IV,Random,95% Cl |
|--|----------------------------|-------------------------|----------------------------|------------|------------------------|----------------|----------------------------------|
| I Asia/Middle East | | | | | | | |
| Li 2003 (I) | 49 | 435 | 1.8197 (0.3019) | | | 17.9 % | 6.17 [3.41, 11.15] |
| Janghorbani 2015 (2) | 214 | 627 | 3.1135 (0.304) | | | 17.9 % | 22.50 [12.40, 40.83] |
| Han 2017 (3) | 198 | 5633 | 2.1054 (0.0969) | | - | 22.9 % | 8.21 [6.79, 9.93] |
| Subtotal (95% CI) | 461 | 6695 | | | • | 58. 7 % | 10.20 [5.45, 19.09] |
| Heterogeneity: Tau ² = 0.25 Test for overall effect: Z = 2 Australia/Europe/North | 7.26 (P < 0.00 | | 12 =82% | | | | |
| Lyssenko 2005 (4) | 221 | 1429 | 1.335 (0.2562) | | | 19.2 % | 3.80 [2.30, 6.28] |
| Subtotal (95% CI) | 221 | 1429 | | | • | 19.2 % | 3.80 [2.30, 6.28] |
| Heterogeneity: not applica | ble | | | | | | |
| Test for overall effect: $Z =$ | 5.21 (P < 0.00 | (1000 | | | | | |
| 3 American Indians/Islands | | | | | | | |
| Wang 2011 (5) | 356 | 595 | 1.4012 (0.1459) | | - | 22.0 % | 4.06 [3.05, 5.40] |
| Subtotal (95% CI) | 356 | 595 | | | • | 22.0 % | 4.06 [3.05, 5.40] |
| Heterogeneity: not applica | ble | | | | | | |
| Test for overall effect: Z = | 9.60 (P < 0.00 | (1000 | | | | | |
| Total (95% CI) | 1038 | 8719 | | | + | 100.0 % | 6.90 [4.15, 11.45] |
| Heterogeneity: $Tau^2 = 0.28$ | 3; Chi ² = 37.0 | 3, df = 4 (P<0.00001) | ; I ² =89% | | | | |
| Test for overall effect: Z = | 7.46 (P < 0.00 | (1000 | | | | | |
| Test for subgroup difference | es: $Chi^2 = 7.4$ | 7, df = 2 (P = 0.02), l | 2 =73% | | | | |
| | | | | | | | |
| | | | C | 0.02 0.1 1 | 10 5 | 0 | |

Normoglycaemia IFG+IGT

- (1) 5 years follow-up
- (2) 7 years follow-up
- (3) 12 years follow-up
- (4) 6 years follow-up
- (5) 4 years follow-up

Analysis 1.5. Comparison I Hazard ratio as the effect measure for the development of T2DM, Outcome 5 T2DM incidence (HbA1c_{5.7}).

Review: Development of type 2 diabetes mellitus in people with intermediate hyperglycaemia

Comparison: I Hazard ratio as the effect measure for the development of T2DM

Outcome: 5 T2DM incidence (HbA1c_{5.7})

| Study or subgroup | HbAlc _{5.7} | Normoglycaemia | log [Hazard Ratio] | Ha | zard Ratio | Weight | Hazard Ratio | |
|--|-----------------------------|--------------------------------------|--------------------|------------------|------------|---------|----------------------|--|
| | Ν | Ν | (SE) | IV,Random,95% Cl | | | IV,Random,95% Cl | |
| l Asia | | | | | | | | |
| Bae 2011 (1) | 1791 | 7932 | 1.8718 (0.2875) | | | 24.2 % | 6.50 [3.70, 11.42] | |
| Heianza 2012 (2) | 822 | 4149 | 1.8764 (0.2776) | | | 24.5 % | 6.53 [3.79, 11.25] | |
| Nakagami 2016 (3) | 583 | 1528 | 2.2742 (0.3433) | | | 22.7 % | 9.72 [4.96, 19.05] | |
| Subtotal (95% CI) | 3196 | 13609 | | | • | 71.4 % | 7.21 [5.14, 10.11] | |
| Heterogeneity: $Tau^2 = 0.0$; | Chi ² = 1.01, d | $f = 2 (P = 0.60); ^2 = 0$ | .0% | | | | | |
| Test for overall effect: Z = | .44 (P < 0.00 | 0001) | | | | | | |
| 2 Australia/Europe/North | America | | | | | | | |
| Warren 2017 (4) | 2027 | 6215 | 0.9969 (0.0453) | | | 28.6 % | 2.71 [2.48, 2.96] | |
| Subtotal (95% CI) | 2027 | 6215 | | | • | 28.6 % | 2.71 [2.48, 2.96] | |
| Heterogeneity: not applical | ble | | | | | | | |
| Test for overall effect: Z = | 22.01 (P < 0.00 | 0001) | | | | | | |
| Total (95% CI) | 5223 | 19824 | | | - | 100.0 % | 5.55 [2.77, 11.12] | |
| Heterogeneity: Tau ² = 0.44 | 4; Chi ² = 31.07 | df = 3 (P<0.00001); | 12 =90% | | | | | |
| Test for overall effect: Z = | 4.83 (P < 0.000 | 01) | | | | | | |
| Test for subgroup difference | es: Chi ² = 30.0 | 6, df = 1 (P = 0.00), l ² | 2 =97% | | | | | |
| | | | | | | | | |
| | | | 0.0 | 5 0.2 | 5 2 | D | | |
| | | | Norm | noglycaemia | HbAlc5.7 | | | |

(1) 4 years follow-up

(2) 5 years follow-up

(3) 5 years follow-up

(4) 22 years follow-up

Analysis 1.6. Comparison I Hazard ratio as the effect measure for the development of T2DM, Outcome 6 T2DM incidence (HbAlc_{6.0}).

Review: Development of type 2 diabetes mellitus in people with intermediate hyperglycaemia

Comparison: I Hazard ratio as the effect measure for the development of T2DM

Outcome: 6 T2DM incidence (HbAlc_{6.0})

| Study or subgroup | HbAlc <u>6.0</u> N | Normoglycaemia N | log [Hazard Ratio] (SE) | Hazard Ratio IV,Random,95% Cl | Weight | Hazard Ratio IV,Random,95% Cl |
|--|-----------------------------|-----------------------------|----------------------------|----------------------------------|----------------|----------------------------------|
| I Asia/Middle East | | | | | | |
| Bae 2011 (1) | 412 | 7932 | 3.7209 (0.2623) | + | 14.7 % | 41.30 [24.70, 69.06] |
| Heianza 2012 (2) | 203 | 4149 | 2.0042 (0.3592) | - | 14.2 % | 7.42 [3.67, 15.00] |
| Nakagami 2016 (3) | 156 | 1528 | 4.1457 (0.3169) | - | 14.4 % | 63.16 [33.94, 117.54] |
| Han 2017 (4) | 1306 | 2715 | 1.454 (0.293) | - | 14.6 % | 4.28 [2.41, 7.60] |
| Han 2017 (5) | 1415 | 2918 | 1.3987 (0.5568) | | 13.0 % | 4.05 [1.36, 12.06] |
| Subtotal (95% CI) | 3492 | 19242 | | • | 7 0.9 % | 13.12 [4.10, 41.96] |
| Heterogeneity: $Tau^2 = 1.62$ | 2; Chi ² = 62.70 |), df = 4 (P<0.00001); | l ² =94% | | | |
| Test for overall effect: Z = | 4.34 (P = 0.00 | 0014) | | | | |
| 2 Australia/Europe/North | America | | | | | |
| Bonora 2011 (6) | 70 | 710 | 2.2762 (0.428) | | 13.8 % | 9.74 [4.21, 22.53] |
| Warren 2017 (7) | 970 | 6215 | 1.1378 (0.0534) | • | 15.2 % | 3.12 [2.81, 3.46] |
| Subtotal (95% CI) | 1040 | 6925 | | • | 29.1 % | 5.09 [1.69, 15.37] |
| Heterogeneity: $Tau^2 = 0.5$ | 5; Chi ² = 6.97, | df = (P = 0.01); $I^2 =$ | =86% | | | |
| Test for overall effect: Z = | 2.89 (P = 0.00 | 39) | | | | |
| Total (95% CI) | 4532 | 26167 | | • | 100.0 % | 10.10 [3.59, 28.43] |
| Heterogeneity: Tau ² = 1.83 | 3; Chi ² = 183.4 | 5, df = 6 (P<0.00001 |); l ² =97% | | | |
| Test for overall effect: Z = | 4.38 (P = 0.00 | 0012) | | | | |
| Test for subgroup difference | es: Chi ² = 1.34 | $H, df = I (P = 0.25), I^2$ | =25% | | | |
| | | | | | 1 | |
| | | | 0.0 | 02 0.1 1 10 50 | 00 | |

0.002 0.1 1 10 5 Normoglycaemia HbA1c6.0

(1) 4 years follow-up

(2) 5 years follow-up

- (3) 5 years follow-up
- (4) 12 years follow-up; HR for male participants
- (5) 12 years follow-up; HR for female participants
- (6) 15 years follow-up

(7) 22 years follow-up

Analysis 1.7. Comparison I Hazard ratio as the effect measure for the development of T2DM, Outcome 7 T2DM incidence (HbAlc + IFG).

Review: Development of type 2 diabetes mellitus in people with intermediate hyperglycaemia

Comparison: I Hazard ratio as the effect measure for the development of T2DM

Outcome: 7 T2DM incidence (HbAIc + IFG)

| Study or subgroup | HbA1c+IFG N | Normoglycaemia N | log [Hazard Ratio] (SE) | Hazard Ratio IV,Fixed,95% Cl | Weight | Hazard Ratio IV,Fixed,95% Cl |
|--|------------------|---------------------|----------------------------|---------------------------------|---------|---------------------------------|
| HbA1c _{5.7} + IFG _{5.6} Heianza 2012 (1) | 410 | 4149 | 3.4812 (0.1764) | - | 100.0 % | 32.50 [23.00, 45.92] |
| Subtotal (95% CI) | 410 | 4149 | | • | 100.0 % | 32.50 [23.00, 45.92] |
| Heterogeneity: not applical Test for overall effect: $Z =$ 2 HbA1c5.7 + IFG6.1 | | 001) | | | | |
| Heianza 2012 (2) | 159 | 5198 | 3.635 (0.1526) | | 100.0 % | 37.90 [28.10, 51.12] |
| Subtotal (95% CI) | 159 | 5198 | | • | 100.0 % | 37.90 [28.10, 51.12] |
| Heterogeneity: not applical Test for overall effect: Z = 3 HbA1c6.0 + IFG5.6 | | 001) | | | | |
| Heianza 2012 (3) | 135 | 4493 | 3.9834 (0.1711) | -+- | 100.0 % | 53.70 [38.40, 75.09] |
| Subtotal (95% CI) Heterogeneity: not applical | 135 | 4493 | | • | 100.0 % | 53.70 [38.40, 75.09] |
| Test for overall effect: $Z =$ | 23.28 (P < 0.000 | 001) | | | | |
| 4 HbA1c _{6.0} + IFG _{6.1} Heianza 2012 (4) | 72 | 5730 | 3.957 (0.1657) | | 100.0 % | 52.30 [37.80, 72.37] |
| Subtotal (95% CI) | 72 | 5730 | | • | 100.0 % | 52.30 [37.80, 72.37] |
| Heterogeneity: not applical Test for overall effect: Z = | | 2012 | | | | |
| Test for subgroup difference | | , | =52% | | | |
| | | | | | | |
| | | | 0.0 | 005 0.1 1 10 20 | 00 | |

Normoglycaemia HbA1c+IFG

(1) 5 years follow-up

(2) 5 years follow-up

(3) 5 years follow-up

(4) 5 years follow-up

Analysis 2.1. Comparison 2 Odds ratio as the effect measure for the development of T2DM, Outcome I T2DM incidence (IFG_{5.6}).

Review: Development of type 2 diabetes mellitus in people with intermediate hyperglycaemia

Comparison: 2 Odds ratio as the effect measure for the development of T2DM

Outcome: I T2DM incidence (IFG_{5.6})

| Study or subgroup | Intermediate hypergly- caemia N | IFG _{5.6} N | log [Odds Ratio] (SE) | Odds Ratio IV,Random,95% Cl | Weight | Odds Ratio IV,Random,95% Cl |
|--|--|-----------------------------|--------------------------|--|---------------|--------------------------------|
| I Asia/Middle East | | | | | | |
| Song 2015 (1) | 167 | 1092 | 1.4516 (0.527) | | 3.7 % | 4.27 [1.52, 12.00] |
| Song 2015 (2) | 154 | 666 | 2.0149 (0.51) | | 3.8 % | 7.50 [2.76, 20.38] |
| Jeong 2010 (3) | 495 | 792 | 1.7334 (0.2541) | | 4.5 % | 5.66 [3.44, 9.31] |
| Liu 2008 (4) | 169 | 470 | 1.5041 (0.4137) | | 4.1 % | 4.50 [2.00, 10.12] |
| Latifi 2016 (5) | 124 | 394 | 0.0392 (0.02) | - | 4.8 % | 1.04 [1.00, 1.08] |
| Wang 2007 (6) | 261 | 400 | 0.9969 (0.3262) | | 4.3 % | 2.71 [1.43, 5.14] |
| Sadeghi 2015 (7) | 373 | 2607 | 1.1939 (0.2162) | | 4.6 % | 3.30 [2.16, 5.04] |
| Liu 2017 (8) | 3607 | 15003 | 1.3002 (0.0699) | + | 4.8 % | 3.67 [3.20, 4.21] |
| Aekplakorn 2006 (9) | 223 | 2444 | 0.8796 (0.1546) | | 4.7 % | 2.41 [1.78, 3.26] |
| Derakhshan 2016 (10) | 523 | 3611 | 1.0986 (0.1356) | | 4.7 % | 3.00 [2.30, 3.91] |
| Bergman 2016 (11) | 263 | 739 | 0.1044 (0.1933) | _ <u></u> | 4.7 % | 1.11 [0.76, 1.62] |
| Subtotal (95% CI) Heterogeneity: Tau ² = 0.65; C Test for overall effect: Z = 4.1 | 9 (P = 0.000029) | 28218 10 (P<0.000 | 01); I ² =98% | • | 48.8 % | 2.94 [1.77, 4.86] |
| 2 Australia/Europe/North Am Levitzky 2008 (12) | nerica 460 | 0 | 2.5416 (0.2295) | | 4.6 % | 2.70 [8.10, 19.91] |
| Levitzky 2008 (13) | 313 | 0 | 3.1046 (0.2753) | → | 4.5 % | 22.30 [13.00, 38.25] |
| Soriguer 2008 (14) | 56 | 1806 | 1.6677 (0.3441) | | 4.3 % | 5.30 [2.70, 10.40] |
| Valdes 2008 (15) | 4 | 510 | 1.361 (0.4546) | | 4.0 % | 3.90 [1.60, 9.51] |
| Lipska 2013 (16) | 189 | 1690 | 1.2528 (0.3117) | | 4.4 % | 3.50 [1.90, 6.45] |
| Admiraal 2014 (17) | 111 | 354 | 1.8083 (0.3454) | | 4.3 % | 6.10 [3.10, 12.00] |
| Cugati 2007 (18) | 229 | 1512 | 2.9513 (0.2557) | | 4.5 % | 19.13 [11.59, 31.58] |
| De Abreu 2015 (19) | 187 | 342 | 1.7492 (0.5758) | | 3.6 % | 5.75 [1.86, 17.77] |
| Filippatos 2016 (20) | 279 | 1206 | 1.2326 (0.2336) | | 4.6 % | 3.43 [2.17, 5.42] |
| | | | Ν | 0.05 0.2 I 5 20 Iormoglycaemia IFG <u>5.6</u> | | Continued |

(Continued ...)

| Study or subgroup | Intermediate hypergly- caemia N | IFG _{5.6} N | log [Odds Ratio] (SE) | Odds Ratio IV,Random,95% Cl | Weight | Odds Ratio IV,Random,95% CI |
|--|--|-------------------------|--------------------------|--------------------------------|---------|--------------------------------|
| Vaccaro 1999 (21) | 11 | 500 | 0.1823 (0.7073) | | 3.1 % | 1.20 [0.30, 4.80] |
| Subtotal (95% CI) | 1949 | 7920 | | • | 41.7 % | 6.47 [3.81, 11.00] |
| Heterogeneity: Tau ² = 0.59; Cł | $hi^2 = 61.74, df = 1$ | 9 (P<0.00001 |); I ² =85% | | | |
| Test for overall effect: $Z = 6.90$ | (P < 0.00001) | | | | | |
| 3 Latin America | | | | | | |
| Ferrannini 2009 (22) | 65 | 1594 | 1.454 (0.1468) | - | 4.7 % | 4.28 [3.21, 5.71] |
| Subtotal (95% CI) | 65 | 1594 | | • | 4.7 % | 4.28 [3.21, 5.71] |
| Heterogeneity: not applicable | | | | | | |
| Test for overall effect: $Z = 9.90$ | (P < 0.00001) | | | | | |
| 4 American Indians/Islands | | | | | | |
| Wang 2011 (23) | 947 | 595 | 1.1378 (0.1534) | | 4.7 % | 3.12 [2.31, 4.21] |
| Subtotal (95% CI) | 947 | 595 | | • | 4.7 % | 3.12 [2.31, 4.21] |
| Heterogeneity: not applicable | | | | | | |
| Test for overall effect: $Z = 7.42$ | (P < 0.00001) | | | | | |
| Total (95% CI) | 9320 | 38327 | | • | 100.0 % | 4.15 [2.75, 6.28] |
| Heterogeneity: Tau ² = 0.92; Ch | ni² = 972.07, df = | 22 (P<0.000 | 01); 12 =98% | | | |
| Test for overall effect: $Z = 6.74$ | (P < 0.0001) | | | | | |
| Test for subgroup differences: C | Chi ² = 7.19, df = | 3 (P = 0.07), | l ² =58% | | | |
| | | | | | | |
| | | | | 0.05 0.2 I 5 2 | 20 | |
| | | | No | ormoglycaemia IFG5.6 | | |

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- (1) 4 years follow-up; female participants
- (2) 4 years follow-up; male participants
- (3) 5 years follow-up; unclear whether IFG or IGT cohort
- (4) 5 years follow-up
- (5) 5 years follow-up
- (6) 5 years follow-up
- (7) 7 years follow-up; isolated $\mathsf{IFG}_{5.6}$
- (8) 8 years follow-up
- (9) 12 years follow-up
- (10) 12 years follow-up; unclear if IFG_{5.6} or IFG_{6.1}
- (11) 24 years follow-up; isolated IFG5.6
- (12) 4 years follow-up; male IFG cohort, total numbers from $\mathsf{IFG}_{6.1}$ cohort
- (13) 4 years follow-up; female IFG cohort, total numbers from $\mathsf{IFG}_{6.1}$ cohort
- (14) 6 years follow-up; univariate analysis
- (15) 6 years follow-up
- (16) 7 years follow-up
- (17) 10 years follow-up
- (18) 10 years follow-up
- (19) 10 years follow-up
- (20) 10 years follow-up
- (21) 12 years follow-up; upper confidence limit in publication: 10.2
- (22) 7 years follow-up; univariate analysis
- (23) 4 years follow-up; univariate analysis

Analysis 2.2. Comparison 2 Odds ratio as the effect measure for the development of T2DM, Outcome 2 T2DM incidence (IFG_{6.1}).

Review: Development of type 2 diabetes mellitus in people with intermediate hyperglycaemia

Comparison: 2 Odds ratio as the effect measure for the development of T2DM

Outcome: 2 T2DM incidence (IFG_{6.1})

| Study or subgroup | IFG _{6.1} N | Normoglycaemia N | log [Odds Ratio] (SE) | Odds Ratio IV,Random,95% Cl | Weight | Odds Ratio IV,Random,95% Cl |
|--|-------------------------|---------------------|-------------------------------------|--|---------------|--------------------------------|
| I Asia/Middle East | | | | | | |
| Chen 2003 (I) | 156 | 444 | 1.4816 (0.4285) | | 5.6 % | 4.40 [1.90, 10.19] |
| Sato 2009 (2) | 794 | 4147 | 3.1144 (0.122) | - | 6.8 % | 22.52 [17.73, 28.60] |
| Wang 2007 (3) | 112 | 400 | 0.5878 (0.3207) | | 6.2 % | 1.80 [0.96, 3.37] |
| Kim 2016a (4) | 1433 | 10763 | 3.0493 (0.1163) | - | 6.8 % | 21.10 [16.80, 26.50] |
| Nakanishi 2004 (5) | 246 | 5500 | 0.27 (0.4813) | - | 5.4 % | 1.31 [0.51, 3.36] |
| Derakhshan 2016 (6) | 523 | 3611 | 1.411 (0.1767) | - | 6.7 % | 4.10 [2.90, 5.80] |
| Bergman 2016 (7) | 53 | 739 | 1.2326 (0.3068) | - | 6.2 % | 3.43 [1.88, 6.26] |
| Subtotal (95% CI) Heterogeneity: Tau ² = 1.08; C Test for overall effect: $Z = 4.0$ | | . , | l ² =96% | • | 43.8 % | 5.18 [2.32, 11.53] |
| 2 Australia/Europe/North An | | , | | | | |
| Levitzky 2008 (8) | 313 | 0 | 3.2696 (0.2108) | • | 6.6 % | 26.30 [17.40, 39.76] |
| Levitzky 2008 (9) | 460 | 0 | 2.5572 (0.1669) | • | 6.7 % | 12.90 [9.30, 17.89] |
| Valdes 2008 (10) | 52 | 510 | 2.4932 (0.4935) | | 5.3 % | 12.10 [4.60, 31.83] |
| Rijkelijkhuizen 2007 (TT) | 149 | 1125 | 2.3026 (0.2522) | + | 6.4 % | 10.00 [6.10, 16.39] |
| Lipska 2013 (12) | 100 | 1690 | 2.4336 (0.2416) | + | 6.5 % | .40 [7.10, 8.30] |
| Rathmann 2009 (13) | 71 | 649 | 1.5476 (0.3873) | | 5.8 % | 4.70 [2.20, 10.04] |
| Cederberg 2010 (14) | 40 | 410 | 0.8629 (0.2368) | + | 6.5 % | 2.37 [1.49, 3.77] |
| Bonora 2011 (15) | 55 | 710 | 1.7405 (0.3627) | - | 6.0 % | 5.70 [2.80, 11.60] |
| Subtotal (95% CI) | 1240 | 5094 | | • | 49.9 % | 8.69 [4.95, 15.24] |
| Heterogeneity: Tau ² = 0.57; C Test for overall effect: Z = 7.5 3 Latin America Ferrannini 2009 (16) | | . , | ² =90% 1.3164 (0.274) | | 6.4 % | 3.73 [2.18, 6.38] |
| Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: Z = 4.8 | | 1594 | | • | 6.4 % | 3.73 [2.18, 6.38] |
| | | | | 0.002 0.1 I IO 50 Normoglycaemia IFG _{6.1} | 0 | (Continued) |

| Study or subgroup | IFG _{6.1} N | Normoglycaemia N | log [Odds Ratio] (SE) | | Odds Ratio 1dom,95% Cl | Weight | (Continued Odds Ratio IV,Random,95% CI |
|----------------------------------|-----------------------------|------------------------------|--------------------------|----------------|---------------------------|---------|---|
| Total (95% CI) | 4574 | 32292 | | | • | 100.0 % | 6.60 [4.18, 10.43] |
| Heterogeneity: $Tau^2 = 0.78;$ | $Chi^2 = 251.22$ | 3, df = 15 (P<0.00001 |); I ² =94% | | | | |
| Test for overall effect: $Z = 8$ | | , | | | | | |
| Test for subgroup differences | s: Chi² = 4.57 | , df = 2 (P = 0.10), I^2 : | =56% | | | | |
| | | | | 0.002 0.1 | I 10 50 | 0 | |
| | | | | Normoglycaemia | IFG _{6.1} | 0 | |
| (1) 3 years follow-up | | | | | | | |
| (2) 4 years follow-up | | | | | | | |
| (3) 5 years follow-up | | | | | | | |
| (4) 5 years follow-up | | | | | | | |
| (5) 7 years follow-up | | | | | | | |
| (6) 12 years follow-up, uncl | ear if IFG _{5.6} c | or IFG _{6.1} | | | | | |
| (7) 24 years follow-up | | | | | | | |
| (8) 4 years follow-up; femal | e IFG cohort | | | | | | |
| (9) 4 years follow-up; male | IFG cohort | | | | | | |
| (10) 6 years follow-up | | | | | | | |
| (11) 6 years follow-up | | | | | | | |
| (12) 7 years follow-up | | | | | | | |
| (13) 7 years follow-up | | | | | | | |
| (14) 10 years follow-up | | | | | | | |
| (15) 15 years follow-up | | | | | | | |

(16) 7 years follow-up

Analysis 2.3. Comparison 2 Odds ratio as the effect measure for the development of T2DM, Outcome 3 T2DM incidence (IGT).

Review: Development of type 2 diabetes mellitus in people with intermediate hyperglycaemia

Comparison: 2 Odds ratio as the effect measure for the development of T2DM

Outcome: 3 T2DM incidence (IGT)

| Study or subgroup | IGT N | Normoglycaemia N | log [Odds Ratio] (SE) | Odds Ratio IV,Random,95% Cl | Weight | Odds Ratio IV,Random,95% CI |
|--|---------------|---------------------------------------|--------------------------|-----------------------------------|--------|--------------------------------|
| I Asia/Middle East | | | | | | |
| Wang 2007 (1) | 126 | 400 | 1.1474 (0.3456) | | 4.3 % | 3.15 [1.60, 6.20] |
| Jeong 2010 (2) | 495 | 792 | 1.7934 (0.3168) | | 4.6 % | 6.01 [3.23, 11.18] |
| Li 2003 (3) | 118 | 435 | 1.0784 (0.2475) | | 5.6 % | 2.94 [1.81, 4.78] |
| Sadeghi 2015 (4) | 373 | 2607 | 0.9243 (0.1919) | - | 6.4 % | 2.52 [1.73, 3.67] |
| Aekplakorn 2006 (5) | 0 | 2444 | 1.4725 (0.1254) | + | 7.3 % | 4.36 [3.41, 5.57] |
| Bergman 2016 (6) | 4 | 739 | 1.7299 (0.3683) | | 4.0 % | 5.64 [2.74, 11.61] |
| Subtotal (95% CI) Heterogeneity: Tau ² = 0.06; C | 1226 | 7417 | -57% | • | 32.2 % | 3.74 [2.83, 4.94] |
| Test for overall effect: $Z = 9.3$ | | . , | 5270 | | | |
| 2 Australia/Europe/North An Mykkänen 1993 (7) | nerica 203 | 689 | 2.2875 (0.2412) | - | 5.7 % | 9.85 [6.14, 15.80] |
| Stengard 1992 (8) | 234 | 216 | 1.1314 (0.4842) | | 2.9 % | 3.10 [1.20, 8.01] |
| Hanley 2005 (9) | 274 | 603 | 1.6901 (0.2088) | - | 6.1 % | 5.42 [3.60, 8.16] |
| Soriguer 2008 (10) | 54 | 1806 | 1.4586 (0.3906) | | 3.8 % | 4.30 [2.00, 9.25] |
| Valdes 2008 (11) | 88 | 510 | 1.9021 (0.3461) | | 4.3 % | 6.70 [3.40, 13.20] |
| Rijkelijkhuizen 2007 (12) | 111 | 1125 | 2.3888 (0.3046) | | 4.8 % | 10.90 [6.00, 19.80] |
| Rathmann 2009 (13) | 120 | 649 | 2.1748 (0.2884) | | 5.0 % | 8.80 [5.00, 15.49] |
| Zethelius 2004 (14) | 201 | 466 | 0.7793 (0.2151) | | 6.0 % | 2.18 [1.43, 3.32] |
| Bonora 2011 (15) | 53 | 710 | 1.361 (0.4546) | | 3.2 % | 3.90 [1.60, 9.51] |
| Cederberg 2010 (16) | 103 | 410 | 1.0647 (0.2157) | | 6.0 % | 2.90 [1.90, 4.43] |
| Vaccaro 1999 (17) | 40 | 500 | 1.8245 (0.4241) | | 3.4 % | 6.20 [2.70, 14.24] |
| Subtotal (95% CI) | 1481 | 7684 | | • | 51.2 % | 5.20 [3.62, 7.45] |
| Heterogeneity: $Tau^2 = 0.27$; C Test for overall effect: $Z = 8.9$ 3 Latin America | | · · · · · · · · · · · · · · · · · · · | l ² =76% | | | |
| Ferrannini 2009 (18) | 179 | 1594 | 1.3888 (0.128) | - | 7.3 % | 4.01 [3.12, 5.15] |
| ıını 2009 (18) | 179 | 1594 | .0 | D2 0.1 1 10 50 noglycaemia IGT | | 4.01 [3.12, 5.15] |

(Continued . . .)

(... Continued)

| Study or subgroup | IGT N | Normoglycaemia N | log [Odds Ratio] (SE) | | Odds Ratio dom,95% Cl | Weight | Odds Ratio IV,Random,95% Cl |
|------------------------------------|----------------------------|-----------------------------|--------------------------|-------------|--------------------------|---------|--------------------------------|
| Lorenzo 2003 (19) | 202 | 1503 | 1.8516 (0.1923) | | - | 6.4 % | 6.37 [4.37, 9.29] |
| Subtotal (95% CI) | 381 | 3097 | | | • | 13.6 % | 4.94 [3.15, 7.76] |
| Heterogeneity: $Tau^2 = 0.08;$ | Chi ² = 4.01, d | $f = (P = 0.05); ^2 = 7$ | 75% | | | | |
| Test for overall effect: $Z = 6.9$ | 94 (P < 0.000 |))) | | | | | |
| 4 American Indians/Islands | | | | | | | |
| Dowse 1991 (20) | 51 | 215 | 1.2809 (0.4819) | | | 2.9 % | 3.60 [1.40, 9.26] |
| Subtotal (95% CI) | 51 | 215 | | | • | 2.9 % | 3.60 [1.40, 9.26] |
| Heterogeneity: not applicable | е | | | | | | |
| Test for overall effect: $Z = 2.6$ | 66 (P = 0.0079 | 9) | | | | | |
| Total (95% CI) | 3139 | 18413 | | | • | 100.0 % | 4.61 [3.76, 5.64] |
| Heterogeneity: $Tau^2 = 0.13$; | Chi ² = 61.78, | df = 19 (P<0.00001); | I ² =69% | | | | |
| Test for overall effect: $Z = 14$ | 1.78 (P < 0.000 | 001) | | | | | |
| Test for subgroup differences | s: $Chi^2 = 2.5I$, | df = 3 (P = 0.47), $I^2 =$ | =0.0% | | | | |
| | | | | | | | |
| | | | 0. | 02 0.1 | I IO 50 |) | |
| | | | Norr | moglycaemia | IGT | | |

Normoglycaemia

(1) 5 years follow-up

(2) 5 years follow-up; unclear whether IFG or IGT cohort

(3) 5 years follow-up; isolated IGT

(4) 7 years follow-up; univariate analysis

(5) 12 years follow-up; number of participants with IGT not reported, univariate analysis

(6) 24 years follow-up

(7) 4 years follow-up; univariate analysis

(8) 5 years follow-up

(9) 5 years follow-up

(10) 6 years follow-up

(11) 6 years follow-up; univariate analysis

(12) 6 years follow-up; isolated IGT

(13) 7 years follow-up; univariate analysis

(14) 7 years follow-up

(15) 10 years follow-up; univariate analysis

(16) 10 years follow-up

(17) 12 years follow-up; isolated IGT

(18) 7 years follow-up; univariate analysis

(19) 8 years follow-up

(20) 5 years follow-up

Analysis 2.4. Comparison 2 Odds ratio as the effect measure for the development of T2DM, Outcome 4 T2DM incidence (IFG + IGT).

Review: Development of type 2 diabetes mellitus in people with intermediate hyperglycaemia

Comparison: 2 Odds ratio as the effect measure for the development of T2DM

Outcome: 4 T2DM incidence (IFG + IGT)

| Study or subgroup | IFG+IGT N | Normoglycaemia N | log [Odds Ratio] (SE) | | Odds Ratio om,95% Cl | Weight | Odds Ratio IV,Random,95% Cl |
|----------------------------------|----------------------------|-------------------------------------|--------------------------|----------------|-------------------------|---------|---|
| I Asia/Middle East | | | | | | | |
| Wang 2007 (1) | 26 | 187 | 1.9615 (0.5212) | | | 9.3 % | 7.11 [2.56, 19.75] |
| Wang 2007 (2) | 36 | 171 | 2.3253 (0.4999) | | | 9.6 % | 10.23 [3.84, 27.25] |
| Sadeghi 2015 (3) | 373 | 2607 | 2.5337 (0.2722) | | - | 11.9 % | 12.60 [7.39, 21.48] |
| Bergman 2016 (4) | 63 | 739 | 1.026 (0.2966) | | - | 11.7 % | 2.79 [1.56, 4.99] |
| Subtotal (95% CI) | 498 | 3704 | | | • | 42.5 % | 6.99 [3.09, 15.83] |
| Heterogeneity: $Tau^2 = 0.54$ | ; Chi ² = 14.83 | s, df = 3 (P = 0.002); I | 2 =80% | | | | |
| Test for overall effect: $Z = 4$ | 4.66 (P < 0.00 | 001) | | | | | |
| 2 Australia/Europe/North A | America | | | | | | |
| Soriguer 2008 (5) | 28 | 1806 | 2.2192 (0.3881) | | | 10.8 % | 9.20 [4.30, 19.69] |
| Valdes 2008 (6) | 20 | 510 | 3.8199 (0.5408) | | | 9.1 % | 45.60 [5.80, 3 .6] |
| Rijkelijkhuizen 2007 (7) | 31 | 1125 | 3.6763 (0.4302) | | | 10.3 % | 39.50 [17.00, 91.79] |
| Rathmann 2009 (8) | 47 | 649 | 3.054 (0.3634) | | | 11.0 % | 21.20 [10.40, 43.22] |
| Bonora 2011 (9) | 19 | 710 | 3.0204 (0.5063) | | | 9.5 % | 20.50 [7.60, 55.30] |
| Vaccaro 1999 (10) | 9 | 500 | 2.3321 (0.7876) | | | 6.8 % | 10.30 [2.20, 48.22] |
| Subtotal (95% CI) | 154 | 5300 | | | • | 57.5 % | 20.95 [12.40, 35.40] |
| Heterogeneity: $Tau^2 = 0.20$ | ; Chi ² = 9.54, | $df = 5 (P = 0.09); I^2 =$ | =48% | | | | , ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,, |
| Test for overall effect: Z = | | · · · · · | | | | | |
| Total (95% CI) | 652 | 9004 | | | • | 100.0 % | 13.14 [7.41, 23.30] |
| Heterogeneity: $Tau^2 = 0.64$ | ; Chi ² = 43.02 | , df = 9 (P<0.00001); | l ² =79% | | | | |
| Test for overall effect: $Z = 8$ | | | | | | | |
| Test for subgroup difference | es: Chi² = 4.91 | , df = 1 (P = 0.03), 1 ² | =80% | | | | |
| | | . , | | | | L | |
| | | | | 0.005 0.1 | 1 10 20 | 0 | |
| | | | | Normoglycaemia | IFG+IGT | | |

(1) 5 years follow-up; female participants (IFG_{6.1}+IGT); IFG_{5.6}+IGT: 4.67 (1.87-11.62)

(2) 5 years follow-up; male participants (IFG_{6.1}+IGT); IFG_{5.6}+IGT: 9.81 (3.5-27.21)

(3) 7 years follow-up

(4) 24 years follow-up; IFG_{5.6}+IGT (IFG_{6.1} + IGT: 3.85 (1.73-8.54))

(5) 6 years follow-up

(6) 6 years follow-up; univariate analysis

(7) 6 years follow-up

(8) 7 years follow-up; univariate analysis

(9) 10 years follow-up; IFG_{6.1}+IGT, univariate analysis

(10) 12 years follow-up; univariate analysis

Analysis 2.5. Comparison 2 Odds ratio as the effect measure for the development of T2DM, Outcome 5 T2DM incidence (HbA1c_{5.7}).

Review: Development of type 2 diabetes mellitus in people with intermediate hyperglycaemia

Comparison: 2 Odds ratio as the effect measure for the development of T2DM

Outcome: 5 T2DM incidence (HbA1c5.7)

| Study or subgroup | HbA1c5.7 N | Normoglycaemia N | log [Odds Ratio] (SE) | Odds Ratio IV,Random,95% Cl | Weight | Odds Ratio IV,Random,95% Cl |
|-------------------------------|------------------------------|----------------------------|--------------------------|--------------------------------|---------|--------------------------------|
| I Asia/Middle East | | | | | | |
| Man 2017 (1) | 675 | 462 | 1.5129 (0.2747) | | 32.7 % | 4.54 [2.65, 7.78] |
| Subtotal (95% CI) | 675 | 462 | | • | 32.7 % | 4.54 [2.65, 7.78] |
| Heterogeneity: not applical | ble | | | | | |
| Test for overall effect: Z = | 5.51 (P < 0.000 | 01) | | | | |
| 2 Europe/North America | | | | | | |
| Lipska 2013 (2) | 207 | 1690 | 2.0794 (0.2606) | -#- | 33.3 % | 8.00 [4.80, 13.33] |
| Cederberg 2010 (3) | 24 | 410 | 0.8838 (0.244) | - | 34.1 % | 2.42 [1.50, 3.90] |
| Subtotal (95% CI) | 231 | 2100 | | • | 67.3 % | 4.38 [1.36, 14.15] |
| Heterogeneity: $Tau^2 = 0.65$ | 5; Chi ² = 11.22, | df = I (P = 0.0008I); | ² =9 % | | | |
| Test for overall effect: Z = | 2.47 (P = 0.013 |) | | | | |
| Total (95% CI) | 906 | 2562 | | • | 100.0 % | 4.43 [2.20, 8.88] |
| Heterogeneity: $Tau^2 = 0.3$ | I; Chi ² = 11.26, | df = 2 (P = 0.004); I^2 | =82% | | | |
| Test for overall effect: Z = | 4.18 (P = 0.000 | 029) | | | | |
| Test for subgroup difference | es: $Chi^2 = 0.00$, | df = 1 (P = 0.96), I^2 : | =0.0% | | | |
| | | | | | 1 | |
| | | | 0. | 01 0.1 1 10 10 | 00 | |
| | | | Nor | moglycaemia HbA1c5.7 | | |

(1) 6 years follow-up

(2) 7 years follow-up; isolated HbA1c5.7

(3) 10 years follow-up

Analysis 2.6. Comparison 2 Odds ratio as the effect measure for the development of T2DM, Outcome 6 T2DM incidence (HbA1c_{6.0}).

Review: Development of type 2 diabetes mellitus in people with intermediate hyperglycaemia

Comparison: 2 Odds ratio as the effect measure for the development of T2DM

Outcome: 6 T2DM incidence (HbAlc_{6.0})

| Study or subgroup | HbAlc _{6.0} N | Normoglycaemia N | log [Odds Ratio] (SE) | Odd IV,Random | s Ratio ,95% Cl | Weight | Odds Ratio IV,Random,95% Cl |
|---------------------------------|------------------------------|-----------------------|--------------------------|------------------|--------------------|---------|--------------------------------|
| I Asia/Middle East | | | | | | | |
| Kim 2016a (1) | 1103 | 10763 | 3.1442 (0.11) | | | 35.6 % | 23.20 [18.70, 28.78] |
| Subtotal (95% CI) | 1103 | 10763 | | | * | 35.6 % | 23.20 [18.70, 28.78] |
| Heterogeneity: not applicat | ole | | | | | | |
| Test for overall effect: $Z =$ | 28.58 (P < 0.00 | 0001) | | | | | |
| 2 Australia/Europe/North / | America | | | | | | |
| Chamnan 2011 (2) | 370 | 5365 | 2.7473 (0.4162) | | | 29.5 % | 15.60 [6.90, 35.27] |
| Subtotal (95% CI) | 370 | 5365 | | | • | 29.5 % | 15.60 [6.90, 35.27] |
| Heterogeneity: not applicat | ole | | | | | | |
| Test for overall effect: $Z =$ | 6.60 (P < 0.000 | 001) | | | | | |
| 3 American Indians/Islands | | | | | | | |
| Wang 2011 (3) | 121 | 595 | 1.7733 (0.1689) | | - | 34.9 % | 5.89 [4.23, 8.20] |
| Subtotal (95% CI) | 121 | 595 | | | • | 34.9 % | 5.89 [4.23, 8.20] |
| Heterogeneity: not applicat | ole | | | | | | |
| Test for overall effect: Z = | 10.50 (P < 0.00 | 0001) | | | | | |
| Total (95% CI) | 1594 | 16723 | | | - | 100.0 % | 12.79 [4.56, 35.85] |
| Heterogeneity: $Tau^2 = 0.76$ | ó; Chi ² = 46.26, | df = 2 (P<0.00001); | $ ^2 = 96\%$ | | | | |
| Test for overall effect: $Z = $ | 4.85 (P < 0.000 | 001) | | | | | |
| Test for subgroup difference | es: Chi ² = 46.2 | 6, df = 2 (P = 0.00), | 2 =96% | | | | |
| | | | | | | | |
| | | | | 0.01 0.1 1 | 10 10 | 00 | |
| | | | Ν | lormoglycaemia | HbAlc6.0 | | |

(1) 5 years follow-up

(2) 3 years follow-up

(3) 4 years follow-up

Analysis 2.7. Comparison 2 Odds ratio as the effect measure for the development of T2DM, Outcome 7 T2DM incidence (HbA1c_{5.7} + IFG_{5.6}).

Review: Development of type 2 diabetes mellitus in people with intermediate hyperglycaemia

Comparison: 2 Odds ratio as the effect measure for the development of T2DM

Outcome: 7 T2DM incidence (HbA1c5.7 + IFG5.6)

| Study or subgroup | HbATc5.7+IFG5.6 N | Normoglycaemia N | log [Odds Ratio] (SE) | | lds Ratio m,95% Cl | Weight | Odds Ratio IV,Random,95% Cl |
|------------------------------|-------------------------------------|----------------------------------|--------------------------|--------------|-----------------------|---------|--------------------------------|
| I Australia/Europe/North | America | | | | | | |
| Lipska 2013 (1) | 169 | 1125 | 3.2658 (0.2421) | | - | 45.5 % | 26.20 [16.30, 42.11] |
| Subtotal (95% CI) | 169 | 1125 | | | • | 45.5 % | 26.20 [16.30, 42.11] |
| Heterogeneity: not applica | able | | | | | | |
| Test for overall effect: Z = | = 13.49 (P < 0.00001) | | | | | | |
| 2 Asia/Middle East | | | | | | | |
| Kim 2016a (2) | 1951 | 10761 | 3.8437 (0.168) | | - | 54.5 % | 46.70 [33.60, 64.91] |
| Subtotal (95% CI) | 1951 | 10761 | | | • | 54.5 % | 46.70 [33.60, 64.91] |
| Heterogeneity: not applica | able | | | | | | |
| Test for overall effect: Z = | = 22.88 (P < 0.00001) | | | | | | |
| Total (95% CI) | 2120 | 11886 | | | • | 100.0 % | 35.91 [20.43, 63.12] |
| Heterogeneity: $Tau^2 = 0.1$ | 2; Chi ² = 3.85, df = 1 | (P = 0.05); I ² =74% | 6 | | | | |
| Test for overall effect: Z = | = 12.44 (P < 0.00001) | | | | | | |
| Test for subgroup differen | ices: Chi ² = 3.85, df = | I (P = 0.05), I ² =74 | 1% | | | | |
| | | . (| | | | j. | |
| | | | | 005 0.1 1 | | .00 | |
| | | | No | rmoglycaemia | HbAlc5.7+ | IFG5.6 | |

(1) 7 years follow-up

(2) 5 years follow-up

ADDITIONAL TABLES

Table 1. Overview: overall prognosis of people with intermediate hyperglycaemia and regression from intermediate hyperglycaemia to normoglycaemia

| Follow-up time (years) | % (95% CI) [no of studi | % (95% CI) regression from IH to normogly- caemia [no of studies; no of participants with IH] | | | | | |
|---------------------------|--------------------------------|---|---------------------------------|-------------------------------|--------------------------------|--------------------------------|--------------------------------|
| | IFG _{5.6} | IFG _{6.1} | IGT | IFG + IGT | HbA1c _{5.7} | HbA1c _{6.0} | |
| 1 | - | - | 13 (5-23) [3; 671] | 29 (23-36) [1; 207] | - | - | 59 (54-64) [2; 375] |
| 2 | 2 (1-2) [1; 1335] | 11 (8-14) [2; 549] | 16 (9-26) [9; 1998] | - | - | - | 46 (36-55) [9; 2852] |
| 3 | 17 (6-32) [3; 1091] | 9 (2-20) [3; 927] | 22 (18-27) [3; 417] | 34 (28-41) [1; 209] | - | 7 (5-10) [1; 370] | 41 (24-59) [7; 1356] |
| 4 | 17 (13-22) [3; 800] | 30 (17-44) [2; 1567] | 22 (12-34) [5; 1042] | - | 14 (7-23) [3; 5352] | 44 (40-48) [2; 627] | 33 (26-40) [3; 807] |
| 5 | 18 (10-27) [7; 3530] | 26 (19-33) [11; 3837] | 39 (25-53) [12; 3444] | 50 (37-63) [5; 478] | 25 (18-32) [4; 3524] | 38 (26-51) [3; 1462] | 34 (27-42) [9; 2603] |
| 6 | 22 (15-31) [4; 738] | 37 (31-43) [5; 279] | 29 (25-34) [7; 775] | 58 (48-67) [4; 106] | 17 (14-20) [1; 675] | - | 23 (3-53) [5; 1328] |
| 7 | 18 (8-30) [5; 980] | 15 (0-45) [4; 434] | 19 (13-26) [5; 835] | 32 (20-45) [4; 753] | 21 (16-27) [1; 207] | - | 41 (37-45) [4; 679] |
| 8 | 34 (27-40) [2; 1887] | 48 (31-66) [1;29] | 43 (37-49) [4; 1021] | 52 (47-57) [1; 356] | - | - | 39 (33-44) [2; 328] |
| 9 | 38 (10-70) [3; 1356] | - | 53 (45-60) [1; 163] | 84 (74-91) [1; 69] | - | - | 17 (14-22) [1; 299] |
| 10 | 23 (14-33) [6; 1542] | 29 (17-43) [6; 537] | 26 (17-37) [6; 443] | 30 (17-44) [2; 49] | 31 (29-33) [2; 2854] | - | 42 (22-63) [7; 894] |
| 11 | - | 38 (33-43) [1; 402] | 46 (43-49) [1; 1253] | - | - | - | 28 (17-39) [2; 736] |
| 12 | 31 (19-34) [3; 433] | 31 (28-33) [1; 1382] | 41 (38-43) [2; 1552] | 70 (63-76) [2; 207] | - | - | - |
| 15 | - | - | - | - | - | 29 (19-40) [1; 70] | - |

 Table 1. Overview: overall prognosis of people with intermediate hyperglycaemia and regression from intermediate hyperglycaemia to normoglycaemia (Continued)

| 20 | - | - | 60 (5-68) | - | - | - | - |
|----|---|---|-----------|---|---|---|---|
| | | | [1; 114] | | | | |

CI: confidence interval; HbA1c: glycosylated haemoglobin A1c; HbA1c_{5.7/6.0} (threshold 5.7% or 6.0%); IFG_{5.6/6.1}: impaired fasting glucose (threshold 5.6 mmol/L or 6.1 mmol/L); IGT: impaired glucose tolerance; IFG + IGT: both IFG and IGT; IH: intermediate hyperglycaemia; T2DM: type 2 diabetes mellitus

Table 2. Overview: intermediate hyperglycaemia versus normoglycaemia as a prognostic factor for the development of type 2 diabetes

| Ratio (95% CI) 95% prediction interval ^{a,b} [no of studies; no of participants with IH/no of participants with normoglycaemia] | | | | | | | |
|--|---|--|--|--|---|---|--|
| Hazard ratio | | | | | | | |
| Region | IFG _{5.6} cohort | IFG _{6.1} cohort | IGT cohort | IFG + IGT cohort | HbA1c _{5.7} co- hort | HbA1c _{6.0} co- hort | HbA1c _{5.7} + IFG _{5.6} cohort |
| Asia/Middle East | 5.07 (3.41-7. 53) 1.07-24.02 [4; 2385/12, 837] | 10.55 (3.61- 30.81) NA ^b [5; 1054/ 9756] | 4.48 (2.81-7. 15) NA ^b [3; 1780/ 6695] | 10.20 (5.45- 19.09) NA ^b [3; 461/6695] | 7.21 (5.14- 10.11) 0.81-64.52 [3; 3196/13, 609] | 13.12 (4.10- 41.96) NA ^b [4; 3492/19, 242] | 32.50 (23.00- 45.92) ^c NA ^a [1; 410/4149] |
| Australia/ Europe/North America | 4.15 (1.24- 13.87) NA ^b [3; 5685/12, 837] | 3.30 (2.32-4. 67) 0.84-12.99 [4; 1736/ 8835] | 2.53 (1.52-4. 19) NA ^{<i>a</i>} [2; 2230/ 5871] | 3.80 (2.30-6. 28) NA ^a [1; 221/1429] | 2.71 (2.48-2. 96) NA ^a [1: 2027/ 6215] | 5.09 (1.69- 15.37) NA ^a [2; 1040/ 6925] | - |
| Latin America | - | 2.06 (1.76-2. 41) NA ^b [1; 28/66] | - | - | - | - | - |
| American In- dians/Islands | 2.38 (1.85-3. 06) NA ^{<i>a</i>} [1; 947/595] | - | - | 4.06 (3.05-5. 40) NA ^{<i>a</i>} [1; 356/595] | - | - | - |
| Overall | 4.32 (2.61-7. 12) 0.75-25.01 [8; 9017/25, 850] | 5.47 (3.50-8. 54) 1.09-27.56 [9; 2818/18, 591] | 3.61 (2.31-5. 64) 0.69-18.97 [5; 4010/12, 566] | 6.90(4.15-11.45)1.06-44.95[5;1038/8719] | 11.12) 0.23-141.18 | 10.10 (3.59- 28.43) NA ^b [6; 4532/26, 167] | 45.92) NA ^a |

Table 2. Overview: intermediate hyperglycaemia versus normoglycaemia as a prognostic factor for the development of type 2diabetes(Continued)

| Incidence rate | ratio | | | | | | |
|---------------------------------------|---|---|---|---|--|---|---|
| Region | IFG _{5.6} cohort | IFG _{6.1} cohort | IGT cohort | IFG + IGT cohort | HbA1c _{5.7} co- hort | HbA1c _{6.0} co- hort | HbA1c _{5.7} + IFG _{5.6} cohort |
| Asia/Middle East | 5.23 (3.77-7. 25) 1.72-15.89 [6; 15,661/ 145,597] | 3.62 (1.67-7. 83) NA ^a [2; 1677/36, 334] | 3.93 (3.03-5. 10) 1.71-9.02 [5; 14,809/73, 128] | 11.20 (5.59- 22.43) NA ^b [4; 3166/69, 463] | 6.62 (4.18- 10.49) NA ^a [1; 1965/ 19961] | - | 40.72 (29.30- 56.61) NA ^a [1; 1641/19, 961] |
| Australia/ Europe/North America | 4.96 (3.25-7. 57) 0.32-77.24 [3; 6322/ 8062] | 11.48) 4.37-16.73 | 5.93 (4.11-8. 57) 2.38-14.81 [5; 2572/22, 329] | 13.92 (9.99- 19.40) 6.71-28.85 [4; 699/18, 966] | - | - | - |
| Latin America | - | - | - | - | - | - | - |
| American In- dians/Islands | 2.74 (1.88-3. 99) NA ^a [1; 2374/ 1613] | - | 4.46 (3.12-6. 38) NA ^a [2; 1087/ 2952] | 5.18 (3.42-7. 83) NA ^a [1; 605/1613] | - | - | - |
| Overall | 4.81 (3.67-6. 30) 1.95-11.83 [10; 24,357/ 155,272] | 6.82 (4.53- 10.25) 2.03-22.87 [6; 5115/56, 580] | 4.48 (3.69-5. 44) 2.60-7.70 [12; 18,468/ 98,409] | 10.94 (7.22- 16.58) 2.58-46.46 [9; 4470/90, 072] | 6.62 (4.18- 10.5) NA ^a [1; 1965/ 19961] | - | 40.72 (29.30- 56.61) NA ^{<i>a</i>} [1; 1641/19, 961] |
| Odds ratio | | | | | | | |
| | IFG _{5.6} cohort | IFG _{6.1} cohort | IGT cohort | IFG + IGT cohort | HbA1c _{5.7} co- hort | HbA1c _{6.0} co- hort | HbA1c _{5.7} + IFG _{5.6} cohort |
| Asia/Middle East | 2.94 (1.77-4. 86) 0.43-19.93 [10; 6359/28, 218] | 11.53) 0.29-91.37 | 3.74 (2.83-4. 94) 1.70-8.21 [6; 1226/ 7417] | 6.99 (3.09- 15.83) NA ^b [3; 498/3704] | 4.54 (2.65-7. 78) NA ^a [1; 675/462] | 23.20 (18.70- 28.78) NA ^{<i>a</i>} [1; 1103/10, 763] | 46.70 (33.60- 64.91) NA ^{<i>a</i>} [1; 1951/10, 761] |
| Australia/ Europe/North America | 6.47 (3.81- 11.00) 0.99-42.32 [9; 1949/ 7920] | 8.69 (4.95- 15.24) 1.20-62.69 [7; 1240/ 5094] | 5.20 (3.62-7. 45) 1.50-18.09 [11; 1481/ 7684] | 20.95 (12.40- 35.40) 4.93-89.05 [6; 154/5300] | 4.38 (1.36- 14.15) NA ^{<i>a</i>} [2; 231/2100] | 15.60 (6.90- 35.27) NA ^{<i>a</i>} [1; 370/5365] | 26.20 (16.30- 41.11) NA ^{<i>a</i>} [1; 169/1125] |

| Latin America | 71) NA ^a | 3.73 (2.18-6. 38) NA ^{<i>a</i>} [1; 17/1594] | 7 6) NA ^a | - | - | - | - |
|-------------------------------|--|--|--|-----------------------------|-------------------------------|---|---|
| American In- dians/Islands | 3.12 (2.31-4. 21) NA ^{<i>a</i>} [1; 947/595] | - | 3.60 (1.40-9. 26) NA ^a [1; 51/215] | - | - | 5.89 (4.23-8. 20) NA ^{<i>a</i>} [1; 121/595] | - |
| Overall | 28) 0.54-32.00 | 10.43) 0.93-46.82 | 64) 2.10-10.13 | 23.30) 1.84-93.66 | 88) NA ^b | 12.8 [4.56 - 35.9] NA ^b [3; 1594/16, 723] | 63.12) NA ^{<i>a</i>} |

Table 2.Overview: intermediate hyperglycaemia versus normoglycaemia as a prognostic factor for the development of type 2diabetes(Continued)

CI: confidence interval; HbA1c: glycosylated haemoglobin A1c; HbA1c_{5.7/6.0} (threshold 5.7% or 6.0%); HbA1c_{5.7} + IFG_{5.6}: both HbA1c_{5.7} and IFG_{5.6}; IFG_{5.6/6.1}: impaired fasting glucose (threshold 5.6 mmol/L or 6.1 mmol/L); IGT: impaired glucose tolerance; IFG + IGT: both IFG and IGT; IH: intermediate hyperglycaemia; NA: not applicable; T2DM: type 2 diabetes mellitus; NR: not reported

^{*a*}With fewer than 3 studies a prediction interval could not be calculated

^bCalculation of the 95% prediction interval did not provide a meaningful estimate

^cCombination of HbA1c_{6.0} plus IFG_{5.6} at baseline showed a hazard ratio for T2DM development of 53.7 (95% CI 38.4-75.1)

APPENDICES

Appendix I. Glossary of terms

| Abbreviation | Explanation | | | | |
|--------------|------------------------------|--|--|--|--|
| ADA | merican Diabetes Association | | | | |
| ALAT | anine aminotransferase | | | | |
| ASAT | Aspartate transaminase | | | | |
| BG | Blood glucose | | | | |
| BMI | Body mass index | | | | |
| BW | Body weight | | | | |

| CI | Confidence interval |
|----------------------|---|
| FG | Fasting glucose |
| FBG | Fasting blood glucose |
| FINDRISC | Finnish Diabetes Risk Score |
| FPG | Fasting plasma glucose |
| G6PD | Glucose-6-P-dehydrogenase test |
| HbA1c | Glycosylated haemoglobin A1c |
| HbA1c _{5.7} | Intermediate hyperglycaemia with HbA1c level 5.7%-6.4% at baseline (HbA1c 5.7% threshold) |
| HbA1c _{6.0} | Intermediate hyperglycaemia with HbA1c level 6.0%-6.4% at baseline (HbA1c 6.0% threshold) |
| h-CRP | High-sensitivity C-reactive protein |
| HOMA-B(eta) | Homeostatic model assessment beta-cell function |
| HOMA-IR | Homeostatic model assessment for insulin resistance |
| HR | Hazard ratio |
| ICTRP | International Clinical Trials Registry Platform |
| IEC | International Expert Committee |
| IFG | Impaired fasting glucose |
| IFG _{5.6} | Intermediate hyperglycaemia with impaired fasting plasma glucose level 5.6-6.9 mmol/L at baseline (IFG 5.6 mmol/ L threshold) |
| IFG _{6.1} | Intermediate hyperglycaemia with impaired fasting plasma glucose level 6.1-6.9 mmol/L at baseline (IFG 6.1 mmol/ L threshold) |
| IFG/IGT | Combination of both IFG and IGT |
| i-IFG | Isolated IFG |
| IGT | Impaired glucose tolerance (intermediate hyperglycaemia defined by IGT: plasma glucose 7.8-11.1 mmol/L 2 hours after a 75 g OGTT at baseline) |
| i-IGT | Isolated IGT |
| IQR | Interquartile range |

| IRR | Incidence rate ratio | | | | |
|-------|--|--|--|--|--|
| JDS | Japanese Diabetes Society | | | | |
| М | Men | | | | |
| NCEP | National cholesterol education program | | | | |
| NDDG | National Diabetes Data Group | | | | |
| NGSP | National Glycohemoglobin Standardization Program | | | | |
| NGT | Normal glucose tolerance | | | | |
| OGTT | Oral glucose tolerance test | | | | |
| OR | Odds ratio | | | | |
| PG | Postload glucose | | | | |
| QUIPS | Quality In Prognosis Studies tool | | | | |
| ROC | Receiver operating characteristics | | | | |
| RR | Risk ratio, relative risk | | | | |
| SD | Standard deviation | | | | |
| SE | Standard error | | | | |
| T2DM | Type 2 diabetes mellitus | | | | |
| W | Women | | | | |
| WHO | World Health Organization | | | | |
| γ-GT | Gamma-glutamyl transferase/transpeptidase | | | | |

Appendix 2. Search strategies

Search strategy overview

Tier 1: prediabetes as predictor for CVD, mortality, stroke, cancer, micro/macrovascular complications
(
1. Population block (prediabetes AND prognosis filter)
OR
2. Prediabetes risk factors / diagnostic criteria block ((IFG, IGT, HbA1c) ADJ6 prognosis terms)
)
AND
3. Outcomes block (diabetes complications, micro/macrovascular, mortality)
Tier 2: prediabetes as predictor for diabetes incidence
(
1. Population block (prediabetes AND prognosis filter)
OR
2. Prediabetes risk factors / diagnostic criteria block ((IFG, IGT, HbA1c) ADJ6 prognosis terms)
)
AND
3. Outcomes block (prediabetes AND prognosis filter)
OR
2. Prediabetes risk factors / diagnostic criteria block ((IFG, IGT, HbA1c) ADJ6 prognosis terms)
)
AND
3. Outcomes block (diabetes incidence)

MEDLINE (Ovid SP)

Whole strategy (combining tier 1: 'prediabetes' as predictor for cardiovascular disease, mortality, stroke, cancer, micro/macrovascular complications and tier 2: 'prediabetes' as predictor for diabetes incidence)

- 1. Prediabetic state/
- 2. (prediabet* or pre diabet*).tw.
- 3. intermediate hyperglyc?emi*.tw.
- 4. or/1-3

5. incidence.sh. or exp mortality/ or follow-up studies.sh. or prognos*.tw. or predict*.tw. or course*.tw. [Wilczynski 2004: MEDLINE prognosis filter sensitivity maximizing]

6. prognosis/ or diagnosed.tw. or cohort*.mp. or predictor*.tw. or death.tw. or exp models, statistical/ [Wilczynski 2004: MEDLINE prognosis filter best balance]

7. or/5-6

8. 4 and 7 [population block (prediabetes + prognosis filter)]

9. ((impaired fasting adj2 glucose) or IFG or (impaired adj FPG)).tw

10. (impaired glucose tolerance or IGT).tw.

11. ("HbA(1c)" or HbA1 or HbA1c or "HbA 1c" or ((glycosylated or glycated) adj h?emoglobin)).tw

12. or/9-11

- 13. (predict* or associa* or prognos*).tw.
- 14. ((prognostic or predict*) adj2 model?).tw.
- 15. predictive value?.tw.
- 16. (risk adj (predict* or factor? or score)).tw.
- 17. or/13-16

18. (((impaired fasting adj2 glucose) or IFG or "impaired FPG" or impaired glucose tolerance or IGT or "HbA1cc)" or HbA1 or HbA1c or "HbA1c" or ((glycosylated or glycated) adj h?emoglobin)) adj3 (predict* or associa* or prognos* or ((prognostic or predict*) adj2 model?) or predictive value? or (risk adj (predict* or factor? or score)))).tw. *[12 adj3 17 // risk factor block]*

19. 8 or 18 [block 1 or block 2]

20. complication?.tw.

21. mortality.tw.

22. (CHD or CVD).tw. 23. (coronary adj2 disease).tw. 24. (coronar* adj (event? or syndrome?)).tw. 25. (heart adj (failure or disease? or attack? or infarct*)).tw 26. (myocardial adj (infarct* or isch?emi*)).tw. 27. cardiac failure.tw. 28. angina.tw. 29. revasculari*.tw. 30. (stroke or strokes).tw. 31. cerebrovascular.tw. 32. ((brain* or cerebr*) adj (infarct* or isch?emi*)).tw. 33. apoplexy.tw. 34. ((vascular or peripheral arter*) adj disease?).tw. 35. cardiovascular.tw. 36. (neuropath* or polyneuropath*).tw. 37. (retinopath* or maculopath*).tw. 38. (nephropath* or nephrotic or proteinuri* or albuminuri*).tw 39. ((kidney or renal) adj (disease? or failure or transplant*)).tw 40. ((chronic or endstage or end stage) adj (renal or kidney)).tw 41. (CRD or CRF or CKF or CRF or CKD or ESKD or ESKF or ESRD or ESRF).tw 42. (microvascular or macrovascular or ((micro or macro) adj vascular)).tw 43. (cancer or carcino* or neoplas* or tumo?r?).tw. 44. (amputation? or ulcer* or foot or feet or wound*).tw. 45. or/20-44 [tier 1 strategy outcomes block] 46. 19 and 45 47. ((diabet* or type 2 or type II or T2D*) adj4 (progress* or inciden* or conversion or develop* or future)).tw. [tier 2 strategy outcomes block] 48. 19 and 47 49. 46 or 48 50. exp animals/ not humans/ 51. 49 not 50 52. (gestational or PCOS).tw. 53. 51 not 52 54. (comment or letter or editorial).pt. 55. 53 not 54 56. remove duplicates from 55

Embase (Ovid SP)

Whole strategy (combining tier 1: 'prediabetes' as predictor for cardiovascular disease, mortality, stroke, cancer, micro/macrovascular complications and tier 2: 'prediabetes' as predictor for diabetes incidence)

1. (prediabet* or pre diabet*).tw.

2. intermediate hyperglyc?emi*.tw.

3. or/1-2

4. exp disease course or risk*.mp. or diagnos*.mp. or follow-up.mp. or ep.fs. or outcome.tw. [Wilczynski 2005: Embase prognosis filter sensitivity maximizing]

5. follow-up.mp. or prognos*.tw. or ep.fs. [Wilczynski 2005: Embase prognosis filter best balance]

6. or/4-5

7. 3 and 6 [population block (prediabetes + prognosis filter)]

8. ((impaired fasting adj2 glucose) or IFG or (impaired adj FPG)).tw

9. (impaired glucose tolerance or IGT).tw.

10. ("HbA(1c)" or HbA1 or HbA1c or "HbA 1c" or ((glycosylated or glycated) adj h?emoglobin)).tw

11. or/8-10

12. (predict* or associa* or prognos*).tw.

13. ((prognostic or predict*) adj2 model?).tw.

- 14. predictive value?.tw.
- 15. (risk adj (predict* or factor? or score)).tw.

16. or/12-15

17. (((impaired fasting adj2 glucose) or IFG or "impaired FPG" or impaired glucose tolerance or IGT or "HbA1c)" or HbA1 or HbA1c or "HbA1c" or ((glycosylated or glycated) adj h?emoglobin)) adj3 (predict* or associa* or prognos* or ((prognostic or predict*) adj2 model?) or predictive value? or (risk adj (predict* or factor? or score)))).tw. *[12 adj3 17 // risk factor block]*

18. 7 or 17 [block 1 or block 2]

19. complication?.tw.

20. mortality.tw.

- 21. (CHD or CVD).tw.
- 22. (coronary adj2 disease).tw.
- 23. (coronar* adj (event? or syndrome?)).tw.

24. (heart adj (failure or disease? or attack? or infarct*)).tw

- 25. (myocardial adj (infarct* or isch?emi*)).tw.
- 26. cardiac failure.tw.
- 27. angina.tw.
- 28. revasculari*.tw.
- 29. (stroke or strokes).tw.
- 30. cerebrovascular.tw.
- 31. ((brain* or cerebr*) adj (infarct* or isch?emi*)).tw.
- 32. apoplexy.tw.
- 33. ((vascular or peripheral arter*) adj disease?).tw.
- 34. cardiovascular.tw.
- 35. (neuropath* or polyneuropath*).tw.
- 36. (retinopath* or maculopath*).tw.
- 37. (nephropath* or nephrotic or proteinuri* or albuminuri*).tw
- 38. ((kidney or renal) adj (disease? or failure or transplant*)).tw
- 39. ((chronic or endstage or end stage) adj (renal or kidney)).tw
- 40. (CRD or CRF or CKF or CRF or CKD or ESKD or ESKF or ESRD or ESRF).tw
- 41. (microvascular or macrovascular or ((micro or macro) adj vascular)).tw
- 42. (cancer or carcino* or neoplas* or tumo?r?).tw.
- 43. (amputation? or ulcer* or foot or feet or wound*).tw.
- 44. or/19-43 [tier 1 strategy outcomes block]
- 45. 18 and 44

46. ((diabet* or type 2 or type II or T2D*) adj4 (progress* or inciden* or conversion or develop* or future)).tw. [tier 2 strategy outcomes block]

- 47. 18 and 46
- 48. 45 or 47

[49-53: TSC Portal filter for exclusion of animal references]

49. exp animals/ or exp invertebrate/ or animal experiment/ or animal model/ or animal tissue/ or animal cell/ or nonhuman/

- 50. human/ or normal human/ or human cell/
- 51. 49 and 50
- 52. 49 not 51

53. 48 not 52
54. (gestational or PCOS).tw.
55. 53 not 54
56. (comment or letter or editorial or conference).pt.
57. 55 not 56
58. remove duplicates from 57

ClinicalTrials.gov (Expert search)

(prediabetes OR prediabetic OR "pre diabetes" OR "pre diabetic" OR "intermediate hyperglycemia" OR "intermediate hyperglycemic" OR "intermediate hyperglycemic" OR "impaired fasting glucose") AND (complication OR complications OR mortality OR CHD OR CVD OR coronary OR heart OR myocardial OR infarct OR infarction OR infarcts OR infarctions OR ischemia OR ischemic OR ischaemia OR ischaemic OR failure OR angina OR revascularization OR revascularizations OR neuropathy OR neuropathies OR stroke OR strokes OR cerebrovascular OR apoplexy OR vascular or peripheral OR cardiovascular OR maculopathy OR neuropathies OR nephropathies OR revascular OR adding or cardiovascular OR maculopathies OR cRF OR CKF OR CKF OR CKD OR ESKD OR ESKF OR ESRD OR ESRF OR microvascular OR macrovascular OR "micro vascular" OR "macro vascular" OR cardiovascular OR tumor OR tumors OR tumour OR tumours OR amputation OR amputations OR ulcer OR foot OR feet OR wounds OR (diabetes OR diabetic OR "type 2" OR "type 11" OR T2D OR T2DM) AND (progress OR progression OR progressed OR incident OR incidence OR conversion OR developed OR development OR future)) [OUTCOME]

ICTRP Search Portal (Standard search)

prediabet* AND prognos* OR prediabet* AND predict* OR prediabet* AND inciden* OR prediabet* AND mortality OR prediabet* AND prevent* OR prediabet* AND progress* OR prediabet* AND develop* OR pre diabet* AND prognos* OR pre diabet* AND predict* OR pre diabet* AND inciden* OR pre diabet* AND mortality OR pre diabet* AND prevent* OR pre diabet* AND progress* OR pre diabet* AND develop* OR impaired glucose tolerance AND prognos* OR impaired glucose tolerance AND predict* OR impaired glucose tolerance AND inciden* OR impaired glucose tolerance AND mortality OR impaired glucose tolerance AND prevent* OR impaired glucose tolerance AND progress* OR impaired glucose tolerance AND develop* OR impaired fasting glucose AND prognos* OR impaired fasting glucose AND predict* OR impaired fasting glucose AND inciden* OR impaired fasting glucose AND mortality OR

impaired fasting glucose AND prevent* OR impaired fasting glucose AND progress* OR impaired fasting glucose AND develop* OR HbA* AND prognos* OR HbA* AND predict* OR HbA* AND inciden* OR HbA* AND mortality OR HbA* AND prevent* OR HbA* AND progress* OR HbA* AND develop*

Seed publications (for PubMed's 'similar articles'-algorithm)

| 24355200[PMID] OR 16873795[PMID] OR 9705020[PMID] OR 25906786[PMID] OR 9363520[PMID] OR |
|---|
| 21278140[PMID] OR 21676480[PMID] OR 21300382[PMID] OR 10862313[PMID] OR 18689695[PMID] OR |
| 27596059[PMID] OR 12397006[PMID] OR 18673544[PMID] OR 21307378[PMID] OR 15220202[PMID] OR |
| 22647753[PMID] OR 28258520[PMID] OR 10663216[PMID] OR 20573752[PMID] OR 20622160[PMID] OR |
| 9300248[PMID] OR 2060716[PMID] OR 27459384[PMID] OR 12757990[PMID] OR 10414941[PMID] OR |
| 21335372[PMID] OR 9653617[PMID] OR 20073428[PMID] OR 17309402[PMID] OR 17315136[PMID] OR |
| 14025561[PMID] OR 10466767[PMID] OR 26273669[PMID] OR 28698884[PMID] OR 11311100[PMID] OR |
| 14710970[PMID] OR 27933333[PMID] OR 27543801[PMID] OR 2035513[PMID] OR 12062857[PMID] OR |
| 11978676[PMID] OR 11679461[PMID] OR 19224196[PMID] OR 14693710[PMID] OR 28278309[PMID] OR |
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| 26606421[PMID] OR 18282630[PMID] OR 8635647[PMID] OR 9243105[PMID] OR 8886564[PMID] OR 7589843[PMID] |
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| 11916954[PMID] OR 16344402[PMID] OR 19531260[PMID] OR 19414206[PMID] OR 1216390[PMID] OR |
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| 28004008[PMID] OR 17320447[PMID] OR 11772900[PMID] OR 2260546[PMID] OR 26885316[PMID] OR |
| 25215305[PMID] OR 29074816[PMID] OR 18206734[PMID] OR 12590020[PMID] OR 26575606[PMID] OR |
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| |

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Appendix 3. QUIPS tool signalling questions

| Study ID | | | | | |
|---|--|--|--|--|--|
| Signalling question | Authors' judgement for 'yes' | | | | |
| Study participation: yes/no ^a /unclear ^b /NA ^c | | | | | |
| a. Adequate participation in the study by eligible people | NA: usually participants with information on glycaemic status and follow-up data providing information on development of type 2 diabetes are selected from a greater study cohort (e.g. study evaluating several cardiovascular risk factors) | | | | |
| b. Description of the source population or population of interest | Source population for cohort with intermediate hyperglycaemia is clearly described | | | | |
| c. Description of the baseline study sample | Number of people with intermediate hyperglycaemia at baseline is clearly described | | | | |
| d. Adequate description of the sampling frame and recruitment | Way of establishing the source population, selection criteria and key characteristics of the source population clearly described | | | | |
| e. Adequate description of the period and place of recruitment | Time period and place of recruitment for both baseline and follow- up examinations are clearly described | | | | |
| f. Adequate description of inclusion and exclusion criteria | Definiton of people with normoglycaemia, intermediate hyper- glycaemia or diabetes mellitus and description of other inclusion and exclusion criteria | | | | |
| Study participation: risk of bias rating (high/low/unclear) | High : most items are answered with 'no'; Low : all items answered with 'yes'; Unclear : most items are answered with 'unclear' Note: potentially a single item may introduce a high risk of bias, depending on study specifics | | | | |

Study attrition: yes/no/unclear/NA

| a. Adequate response rate for study participants | NA: usually participants with information on glycaemic status and follow-up data providing information on development of type 2 diabetes are selected from a greater study cohort (e.g. study evaluating several cardiovascular risk factors) |
|---|---|
| b. Attempts to collect information on participants who dropped out described | Attempts to collect information on participants who dropped out are described (e.g. telephone contact, mail, registers) |
| c. Reasons for loss to follow-up provided | Reasons on participants who dropped out are available (e.g. de- ceased participants between baseline and follow-up, participants moving to another location) |
| d. Adequate description of participants lost to follow-up | Key characteristics of participants lost to follow-up are described (age, sex, glucose status at baseline, body mass index) |
| e. No important differences between participants who completed the study and those who did not | Study authors described differences between participants com- pleting the study and those who did not as not important or in- formation provided to judge the differences |
| Study attrition: risk of bias rating (high/low/unclear) | High : most items are answered with 'no'; Low : all items answered with 'yes'; Unclear : most items are answered with 'unclear' Note: potentially a single item may introduce a high risk of bias, depending on study specifics |
| Glycaemic status measurement: yes/no/unclear/NA | |
| a. Clear definition or description provided | Measurements for glycaemic status are provided (e.g. IFG, IGT, elevated HbA1c) |
| b. Adequately valid and reliable method of measurement | Ideally measurements for glycaemic status are repeated to ensure diagnosis, single measurements are accepted as well; technique for glucose measurement or HbA1c measurement described |
| c. Continuous variables reported or appropriate cut points used | Standard categories for intermediate hyperglycaemia (FPG 5.6-6. 9 mmol/L (IFG _{5.6}), FPG 6.1-6.9 mmol/L (IFG _{6.1}), 2-h PG 7.8 to < 11.0 mmol/L (IGT), HbA1c 6.0-6.4% (HbA1c _{6.0}), HbA1c 5.7-6.4% (HbA1c _{5.7})) |
| d. Same method and setting of measurement used in all study participants | Measurements of glycaemic status are the same for all study par- ticipants |
| e. Adequate proportion of the study sample had complete data | NA: usually participants with information on glycaemic status and follow-up data providing information on development of type 2 diabetes are selected from a greater study cohort (e.g. study evaluating several cardiovascular risk factors) |
| f. Appropriate methods of imputation were used for missing data | NA: missing laboratory measurements for glycaemic status cannot be reliably imputed |

| Glycaemic status measurement: risk of bias rating (high/low/ unclear) | High : most items are answered with 'no'; Low : all items answered with 'yes'; Unclear : most items are answered with 'unclear' Note: potentially a single item may introduce a high risk of bias, depending on study specifics |
|--|--|
| Outcome measurement: yes/no/unclear | |
| a. Clear definition of the outcome provided | Measurement of type 2 diabetes mellitus has to be defined |
| b. Use of adequately valid and reliable method of outcome mea- surement | Measurement of type 2 diabetes mellitus: a glucose (FPG, PG) or HbA1c measurement has to be a part of the diagnosis (self-reported diabetes alone will not be accepted) |
| c. Use of same method and setting of outcome measurement in all study participants | Measurements of type 2 diabetes mellitus are the same for all study participants |
| Outcome measurement: risk of bias rating (high/low/unclear) | High : most items are answered with 'no'; Low : all items answered with 'yes'; Unclear : most items are answered with 'unclear' Note: potentially a single item may introduce a high risk of bias, depending on study specifics |
| Study confounding: yes/no/unclear | |
| a. Measurement of all important confounders | Important confounders are: age, sex, family history of diabetes, 'ethnicity', body mass index, blood pressure and hypertension, smoking and drinking status, socioeconomic status, comedica- tions and comorbidities, physical activity |
| b. Provision of clear definitions of the important confounders measured | Measurement of confounders has to be clearly described |
| c. Adequately valid and reliable measurement of all important confounders | Measurement of confounders is valid and reliable |
| d. Use of same method and setting of confounding measurement in all study participants | Measurements of confounders are the same for all study participants |
| e. Appropriate imputation methods used for missing confounders (if applicable) | Strategy to impute missing confounder data is described |
| f. Important potential confounders were accounted for in the study design | Methods section of the publication describes strategy to account for confounders |
| g. Important potential confounders were accounted for in the analysis | Important confounders are accounted for in multivariable logistic regression and Cox proportional hazards models |

| Study confounding measurement: risk of bias rating (high/ low/unclear) | High: most items are answered with 'no'; Low: all items answered with 'yes'; Unclear: most items are answered with 'unclear' Note: potentially a single item may introduce a high risk of bias, depending on study specifics | | | |
|---|---|--|--|--|
| Statistical analysis and reporting: yes/no/unclear/NA | | | | |
| a. Sufficient presentation of data to assess the adequacy of the analytic strategy | Mean or median values, including confidence intervals or standard errors or standard deviations | | | |
| b. Strategy for model building is appropriate and based on a con- ceptual framework or model | NA: we do not anticipate conceptual frameworks or explicit model building strategies for this type of research question (focusing on one prognostic factor only) | | | |
| c. Statistical model is adequate for the study design | Mainly incidence rates, uni- and multivariate logistic regression, Cox proportional hazard model | | | |
| d. No selective reporting of results | NA: development of type 2 diabetes mellitus and potentially re- gression to normoglycaemia from intermediate hyperglycaemia are the only outcomes; if missing the study will be excluded | | | |
| Statistical analysis and reporting: risk of bias rating (high/ low/unclear) | High : most items are answered with 'no'; Low : all items answered with 'yes'; Unclear : most items are answered with 'unclear' Note: potentially a single item may introduce a high risk of bias, depending on study specifics | | | |

 a No: no or no relevant information to answer the signalling question

^bUnclear: not enough information to answer signalling question with yes or no

 c **NA** (not applicable): signalling question not appropriate for this type of prognostic review

FPG: fasting plasma glucose; **HbA1c**: glycosylated haemoglobin A1c; **IFG**: impaired fasting glucose; **IGT**: impaired glucose tolerance; **PG**: postload glucose (after an oral glucose tolerance test)

Appendix 4. Major cohort studies

| Cohort study acronym | Full study name |
|----------------------|--|
| ADDITION | Anglo-Danish-Dutch study of Intensive Treatment in People with Screen Detected Diabetes in Primary Care (Rasmussen 2008) |
| - | Ansung-Ansan Cohort Study (part of the Korean Genome and Epidemiology Study (KoGES)) - (Han 2017) |
| Asturias | Asturias Study (Valdes 2008) |

| ARIC | Atherosclerosis Risk in Communities study (Warren 2017) |
|---------------------|--|
| ATTICA | Province of Attica, Greece Study (Filippatos 2016) |
| AusDiab | Australian Diabetes, Obesity and Lifestyle Study (Magliano 2008) |
| BLSA | Baltimore Longitudinal Study of Aging (Meigs 2003) |
| BLSA | Beijing Longitudinal Study on Aging (Liu 2016) |
| - | Beijing Project as part of the National Diabetes Survey (Wang 2007) |
| BMES | Blue Mountains Eye Study (Cugati 2007) |
| - | Botnia Study (Lyssenko 2005) |
| - | Bruneck Study (Bonora 2011) |
| CUPS-19 | Chennai Urban Population Study-19 (Mohan 2008) |
| CURES | Chennai Urban Rural Epidemiology Study (Anjana 2015) |
| ChinaMUCA | China Multicenter Collaborative Study of Cardiovascular Epidemiology (Liu 2017) |
| CODAM | Cohort on Diabetes and Atherosclerosis Maastricht (Den Biggelaar 2016) |
| DESIR | Data from an Epidemiological Study on the Insulin Resistance Syndrome (Gautier 2010) |
| - | Ely Study (Forouhi 2007) |
| EPIC-Norfolk cohort | European Prospective Investigation of Cancer Norfolk cohort (Chamnan 2011) |
| - | Finnish Cohorts of the Seven Countries Study (Stengard 1992) |
| None | Framingham Heart Study (Levitzky 2008) |
| GOS | Geelong Osteoporosis Study (De Abreu 2015) |
| Health ABC | Health, Aging, and Body Composition Study (Lipska 2013) |
| - | Hoorn Study (Rijkelijkhuizen 2007) |
| None | Hong Kong Cardiovascular Risk Factor Prevalence Study (Wat 2001) |
| IRAS | Insulin Resistance Atherosclerosis Study (Hanley 2005) |

| ICS | Isfahan Cohort Study (baseline survey of the Isfahan Healthy Heart Program) (Sadeghi 2015) |
|------------------|--|
| IDPS | Isfahan Diabetes Prevention Study (Janghorbani 2015) |
| Israel GOH Study | Israel Study of Glucose Intolerance, Obesity and Hypertension (Bergman 2016) |
| ILSA | Italian Longitudinal Study on Aging (Motta 2010) |
| - | Japanese American Community Diabetes Study (McNeely 2003) |
| JPHC Study | Japanese Public-Health Center-based prospective (Diabetes) Study (Noda 2010) |
| - | Kansai Healthcare Study (Sato 2009) |
| - | Kinmen Study (Li 2003) |
| KORA S4/F4 | Kooperative Gesundheitsfroschung in der Region Augsburg (Rathmann 2009) |
| KoGES | Korean Genome Epidemiology Study-Kangwha Study (Song 2015) |
| - | Kurihashi Lifestyle Cohort Study (Nakagami 2016) |
| - | Mexico City Diabetes Study (Ferrannini 2009) |
| MESA | Multi-Ethnic Study of Atherosclerosis (Yeboah 2011) |
| - | Nauru Study (Dowse 1991) |
| - | Paris Prospective Study (Charles 1997) |
| - | Pima Indian Study (Gila River Indian Community) (Wheelock 2016) |
| - | Pizarra study (Soriguer 2008) |
| PIFRECV | Programa de Investigación de Factores de Riesgo de Enfermedad Cardiovascular (Leiva 2014) |
| - | Rotterdam study (Ligthart 2016) |
| SALSA | Sacramento Area Latino Study on Aging (Garcia 2016) |
| SAHS | San Antonio Heart Study (Lorenzo 2003) |
| - | San Luis Valley Diabetes Study (Marshall 1994) |
| - | Singapore Impaired Glucose Tolerance Follow-up Study (Wong 2003) |

| SIMES | Singapore Malay Eye Study (Man 2017) |
|---------------|---|
| SDPP | Stockholm Diabetes Prevention Programme (Alvarsson 2009a) |
| SHS | Strong Heart Study (Wang 2011) |
| - | Study within the WHO-assisted National Diabetes Programme (Schranz 1989) |
| SUNSET/HELIUS | Surinamese in the Netherlands: study on health and ethnicity/Healthy life in an urban setting (Admiraal 2014) |
| TLGS | Tehran Lipid and Glucose Study (Derakhshan 2016) |
| TOPICS | Toranomon Hospital Health Management Center Study (Heianza 2012) |
| - | Yonchon study (Shin 1997) |
| - | Zanjan Healthy Heart Study (Sharifi 2013) |

Appendix 5. Definition of normoglycaemia, intermediate hyperglycaemia and incident type 2 diabetes

| Study ID | Normogly- caemia (mmol/L or %) | Intermedi- ate hypergly- caemia (mmol/L or %) | 2 diabetes (mmol/L or | OGTT mea- surement (glucose load) | OGTT at baseline | OGTT at fol- low-up | Notes |
|--------------------|---|--|---|--|---------------------|------------------------|-------|
| Admiraal 2014 | - | IFG: FPG 5.7- 6.9 | $FPG \ge 7.0;$ $HbA1c \ge 6.5;$ self-reported diabetes | - | - | - | - |
| Aekplakorn 2006 | - | IFG: FPG \geq 5.6 to < 7.0; IGT: 2-h PG \geq 7.8 to < 11. | agnosis and/or | 75 g | Yes | No | - |
| Ammari 1998 | - | IGT: 2-h PG 7.8 to < 11.1 (WHO 1985) | 2-h PG ≥ 11.1 (WHO 1985) | 75 g | Yes | Yes | - |

| Anjana 2015 | FPG < 5.6 and 2-h PG < 7.8 | FPG > 5.6; i- IFG: FPG 5. | h PG ≥ 11.1; diagnosed; an- tihypergly- caemic medi- | 75 g | Yes | Unclear | - |
|--------------------|---|-------------------------------------|---|-------|------|---------|--|
| Bae 2011 | - | | $FPG \ge 7.0;$ $HbA1c \ge 6.5;$ history of di- abetes; antihy- perglycaemic medication | None | None | None | - |
| Baena-Diez 2011 | FPG < 6.1 | IFG: 6.1-6.9 | $\begin{array}{l} \text{FPG} \geq \\ 7.0 \text{ (measured twice)} \end{array}$ | - | - | - | - |
| Bai 1999 | - | IGT: 7.8 to < 11.1 (WHO 1985) | 2-h PG ≥ 11.1 (WHO 1985) | 75 g | Yes | Yes | - |
| Bergman 2016 | FPG < 5.6 + and no antihy- pergly- caemic medi- cation and 2- h BG < 7.8 (if available) | | $FPG \ge 7.8, 2-$ h BG $\ge 11.1;$ self-reported | 100 g | Yes | Unclear | - |
| Bonora 2011 | - | 49; IFG: not | $FPG \ge 7.0;$ $HbA1c \ge 6.5;$ diabetes treat- ment | 75 g | Yes | Unclear | - |
| Cederberg 2010 | - | | 2-h PG: ≥ 11. 1, confirmed by 2 OGTTs | - | - | - | Diabetes inci- dence and IFG/IGT not exactly de- fined |

| | | HbA1c: 5.7-6. 4 | | | | | |
|------------------------------|-------------------------------|---|--|------|-----|---------|----------------------------|
| Chamnan 2011 | - | HbA1c 6.0-6. 4 | $HbA1c \ge 6.5;$ reported physician- diagnosed di- abetes or dia- betes medica- tions; antihy- pergly- caemic med- ication; diag- nosis through registers | - | - | - | - |
| Charles 1997 | - | | 2-h PG ≥ 11.1 (WHO 1985) ; physician diagnosed dia- betes | 75 g | Yes | Yes | 2nd and 4th examination |
| Chen 2003 | FPG < 6.1 | IFG: FPG 6.1- 7.0 | $FPG \ge 7.0$ | - | - | - | - |
| Chen 2017 | FPG < 5.6 and 2-h PG < 7.8 | | h PG \geq 11. 1; previously diagnosed dia- | 75 g | Yes | Unclear | - |
| Coronado- Malagon 2009 | ADA 2007 | ADA 2007 (IFG/IGT: 5. 6-6.9/7.8 to < 11.1) | ADA 2007 (≥ 7.0/≥ 11.1) | - | - | - | - |
| Cugati 2007 | - | IFG: FPG 5.6- 6.9 (originally FPG \geq 6.1 to < 7.0) | FPG ≥ 7.0; self- reported dia- betes history; antihypergly- caemic medi- cation | - | - | - | - |

| De Abreu 2015 | - | IFG: 5.5-6.9 | $FPG \geq 7.0;$ self-reported; antihypergly- caemic medi- cation | - | - | - | - |
|--------------------------|-------------------------------|--|---|--------|------|---------|---|
| Den Biggelaar 2016 | FPG < 6.1 and 2-h PG < 7.8 | FPG 6.1-6.9; 2-h PG 7.8- 11.1 | | 75 g | Yes | Unclear | - |
| Derakhshan 2016 | | $5.55 \leq$ FPG < 7.0; 7.77 \leq 2- h PG \leq 11.1; no antihyper- glycaemic medication | h PG \geq 11.1; | 82.5 g | Yes | Unclear | Glu- cose monohy- drate solution, equivalent to 75 g anhy- drous glucose |
| Dowse 1991 | FPG and 2-h PG < 7.8 | IGT: FPG < 7. 8 and 2-h PG ≥ 7.8 to < 11. 1 | (WHO 1985) | 75 g | Yes | Yes | - |
| Ferrannini 2009 | - | IFG: FPG 6.1- 6.9; IGT: FPG < 7.0 and 2- h PG 7.8-11. 1; i-IFG _{6.1} /i- IFG _{5.6} : 2-h PG < 7.8 and FPG 6.1- 6.9/5.6-6.1; i- IGT/i- IGT _{6.1} /i- IGT _{5.6} | FPG ≥ 7.0; 2- h PG ≥ 11.1 | 75 g | Yes | Yes | - |
| Filippatos 2016 | - | IFG _{5.6} : FBG 5.6-6.9 | FBG > 6.9; antihypergly- caemic medi- cation | None | None | None | - |
| Forouhi 2007 | FPG < 5.6 | 6.1-6.9 (FPG | $FPG \ge 7.0; 2-$ h PG $\ge 11.$ 1; doctor diag- nosis or treat- ment for dia- betes | 75 g | Yes | Yes | - |

| Garcia 2016 | - | Prediabetes: FBG 5.6-6.9 | $FPG \geq 7.0;$ self-reported; antihypergly- caemic medi- cation; di- abetes comed- ication of death | - | - | - | - |
|----------------------------|--|---------------------------------------|---|------|-----|-----|---|
| Gautier 2010 | - | IFG: FPG 5.6- 6.9 | FPG \geq 7.0; treat- ment for dia- betes (at one of the 3-yearly examinations) | - | - | - | - |
| Gomez- Arbelaez 2015 | - | < 7.0; IGT: \geq | $\begin{array}{l} \text{FPG} \geq 7.0;\\ \text{OGTT} \geq 11.\\ 1; \text{HbA1c} \geq 6.\\ 5 \end{array}$ | OGTT | Yes | Yes | OGTTs from hospital's database |
| Guerrero- Romero 2006 | FPG < 6.1 and 2-h PG < 7.8 | IGT: 2-h PG ≥ 7.8 to < 11. 1 | | OGTT | Yes | Yes | OGTT: as baseline and each year dur- ing the 5-year follow-up |
| Han 2017 | FPG < 5.6 and 2-h PG < 7.8 | | h PG \geq 11. 1; HbA1c \geq 6. 5; current an- | 75 g | Yes | Yes | OGTT was performed ev- ery 2 years |
| Hanley 2005 | - | IFG,IGT (WHO 1999) | Unclear | 75 g | Yes | No | - |
| Heianza 2012 | Absence of IFG or ele- vated HbA1c | IFG: FPG 5.6- 6.9 or FPG 6.1-6. | FPG ≥ 7.0; HbA1c ≥ 6.5%; self- | - | - | - | - |

| | | 6.4 or 6.0-6.4; | reported clini- cian-diag- nosed diabetes | | | | |
|--------------------------------|-------------------------------|--|--|--|-----|-----|---------------------------------------|
| Inoue 1996 | - | IGT: ≥ 7.8 to < 11.1 (pre- sumed WHO 1985) | | 75 g | Yes | Yes | OGGT was performed ev- ery year |
| Janghorbani 2015 | FPG < 5.6 and 2-h PG < 7.8 | i-IFG: 5.6-6.9 | antihypergly- caemic medi- cation; 2nd $FPG \ge 7.0; 2$ - | 75 g | Yes | Yes | - |
| Jaru- ratanasirikul 2016 | FPG < 5.6 | i-IGT: FPG < 5.6 and 2-h PG 7.8 to < 11.1 | FPG > 7.0; 2- h PG ≥ 11.1 | 1.75 g/kg (maximum 75 g) glucose so- lution | Yes | No | - |
| Jeong 2010 | - | IFG: FPG \geq 5.6 to < 7.0; IGT: 2-h PG \geq 7.8 to < 11.1: 'prediabetes': IFG or IGT | FPG ≥ 7.0; 2- h PG ≥ 11.1 | 75 g | Yes | Yes | - |
| Jiamjaras- rangsi 2008a | - | IFG: FPG \geq 5.6 to < 7.0 | FPG ≥ 7.0 | - | - | - | - |
| Kim 2005 | FPG < 5.0 | 6.1 to < 7. | FPG ≥ 7.0; antihypergly- caemic treat- ment | - | - | - | - |
| Kim 2008 | - | IFG _{5.6} : FPG 5. 6-7.0; IFG _{6.1} : FPG 6.1-7.0 | FPG ≥ 7.0 | - | - | - | - |

| Kim 2014 | - | | FPG ≥ 7.0; 2- h PG ≥ 11.1; HbA1c ≥ 6.5 | 75 g | Yes | Unclear | - |
|--------------|-------------------------------|--|--|---|-----------------------------------|---------|---|
| Kim 2016a | - | | $FPG \ge 7.0;$ $HbA1c \ge 6.5;$ antihypergly- caemic medi- cations | - | - | - | - |
| Kleber 2010 | - | IGT: 2-h PG > 7.7: IFG: FPG ≥ 5.5 (WHO definition) | ADA 2000 | 1.75 g/kg body mum 75 g) flav | y weight (maxi- roured glucose | Yes | - |
| Kleber 2011 | - | reported (pre- | "ADA" (2000 criteria, 2-h PG ≥ 11. 1) | 1. 75 g/kg body weight (max. 75 g) | Yes | Yes | |
| Ko 1999 | WHO/ NDDG 1979 | WHO/ NDDG 1979 | WHO/ NDDG 1979 | - | Yes | Yes | - |
| Ko 2001 | FPG < 6.1 | IFG: FPG 6.1- 6.9 | $FPG \ge 7.0$ | 75 g | Yes | Yes | Annual OGTTs |
| Larsson 2000 | FPG < 5.3 and 2-h BG < 7.8 | i-IFG: BG 5. 3-5.9 and 2- h BG < 7.8; i-IGT: FPG < 5.3 and 2-h BG 7.8-11.0; IFG/IGT: BG 5.3-5.9 and 2- h BG 7.8-11.0 | | 75 g | Yes | Yes | NGT at base- line vs follow- up: FPG < 5.3 vs < 6.1; FPG 5.3: 15% con- version fac- tor as recom- mended by the WHO (blood glucose > plasma glu- cose) |

| Latifi 2016 | - | 5.6 ≤ FPG < 7.0 | $FPG \geq 7.0;$ antihypergly- caemic medi- cation | - | - | - | - |
|------------------|-------------------------------|--------------------|---|------|-----|-----|---|
| Lecomte 2007 | personal his- tory of dia- | | personal history of di- abetes; hypo- | - | - | - | - |
| Lee 2016 | - | HbA1c 5.7-6. 4 | HbA1c ≥ 6.5 | - | - | - | - |
| Leiva 2014 | - | | $FPG \geq 7.0$ (2 cons. days), HbA1c ≥ 6.5 | - | - | - | - |
| Levitzky 2008 | - | 5. | $FPG \geq 7.0;$ antihypergly- caemic medi- cation | - | - | - | - |
| Li 2003 | FPG < 6.1 and 2-h PG < 7.8 | PG < 7.8; i- | h PG \geq 11.0; | 75 g | Yes | Yes | - |
| Ligthart 2016 | FBG ≤ 6.0 | < 7.0; non- | $FBG \geq 7.$ 0; non-fasting $BG \geq 11.1;$ antihypergly- caemic medi- cation | - | - | - | - |
| Lipska 2013 | FPG < 5.6 and HbA1c < 5.7 | HbA1c < 5.7; | Single HbA1c \geq 6.5 (years 2,6,7); self-re- port of physi- | - | - | - | - |

| | | 6.4 and FPG > 5.6; IFG and HbA1c: FPG 5.6-6.9 and HbA1c 5. 7-6.4 | ; antihypergly- caemic agent | | | | |
|------------------|---|---|--|------|-----|---------|---|
| Liu 2008 | - | IFG 5.6-6.9 | $FPG \ge 7.0; 2-$ h PG $\ge 11.0;$ antihypergly- caemic medi- cation | - | - | - | - |
| Liu 2014 | WHO | IFG; IGT (WHO) | WHO | 75 g | Yes | Unclear | - |
| Liu 2016 | - | FPG 6.1-6.9 | $FPG \ge 7.0;$ self-reported; antihypergly- caemic medi- cation | - | - | - | - |
| Liu 2017 | FPG 3.9-5.5 | FG 5.6-6.9 | FG ≥ 7.0; us- ing insulin/ hypogly- caemic agents; self-reported | - | - | - | - |
| Lorenzo 2003 | - | 1-6.9; IGT: 2- | FPG: ≥ 7.0; 2-h PHG: ≥ 11.1 (WHO 1999/1985) | 75 g | Yes | Yes | - |
| Lyssenko 2005 | FPG < 6.1 | IFG: FPG \geq 6.1; WHO 1999 criteria | WHO 1999 criteria | 75 g | Yes | Yes | - |
| Magliano 2008 | FPG < 6.1 and 2-h PG < 7.8 | h PG < 7.8; IGT: FPG < 7. | h PG \geq 11. 1; current an- tihypergly- caemic medi- | 75 g | Yes | Yes | - |
| Man 2017 | Not 'predia- betes', not dia- betes | HbA1c 5.7-6. 4; no self-re- ported diabetes or an- | Random glu- cose ≥ 11.1 or HbA1c > 6.4; | - | - | - | - |

| | | tihypergly- caemic medi- cation | self-reported history or an- tihypergly- caemic medi- cation | | | | |
|------------------|-----------------|---|--|--|-----|-----|---|
| Marshall 1994 | - | IGT: 2-h PG ≥ 7.8 to < 11.1 (WHO 1985) | 2-h PG ≥ 11.1 (WHO 1985) | 75 g | Yes | Yes | - |
| McNeely 2003 | - | 6.1 to < 7.0; IGT: 2-h PG | $FPG \ge 7.0; 2-$ h PG $\ge 11.1;$ antihypergly- caemic medi- cation prescribed by a physician | 75 g | Yes | Yes | - |
| Meigs 2003 | | IFG: FPG 6. 1-6.9 and 2- h PG ≤ 7. 8; IGT: FPG < 6.1 and 2- h PG 7.8-11. 0; IFG/IGT | $\begin{array}{l} h \ PG \geq 11.1 \\ (IFG-IGT \\ person: \ diabetes \ defined \end{array}$ | 07/1977: 1.75 g glucose/ kg BW, aver- | Yes | Yes | Serial OGTTs over subse- quent biennial examinations |
| Mohan 2008 | - | IFG: FPG \geq 6.1 to < 7; IGT: 2-h PG \geq 7.8 to < 11. 1 | FPG ≥ 7; 2-h PG ≥ 11.1 | 75 g | Yes | Yes | - |
| Motala 2003 | 2-h PG < 7.8 $$ | IGT: FPG < 7. 8 and 2-h PG 7.8 to < 11.1 (WHO 1985) | h PG \geq 11.1 | 75 g glucose monohydrate dissolved in 250 mL of wa- ter (modified OGTT) | Yes | Yes | - |
| Motta 2010 | FPG < 6.1 | IFG: 6.1 to < 7.0 | FPG ≥ 7.0 | - | Yes | | - |

| Mykkänen 1993 | FPG and 2-h PG < 7.8 | IGT: FPG < 7.8 and 2-h PG 7.8-11.1 (WHO 1985) | h PG \geq 11.1 | 75 g | Yes | Yes | - |
|-------------------|-------------------------------|---|---|------|-----|---------|---|
| Nakagami 2016 | - | | $FPG \ge 7.0,$ $HbA1c \ge 6.5;$ physician di- agnosis of dia- betes | - | - | - | - |
| Nakanishi 2004 | FPG < 6.1 | IFG: FPG 6.1- 6.9 | $FPG \geq 7.0;$ antihypergly- caemic medi- cation | - | - | - | - |
| Noda 2010 | - | | FPG \geq 7.0; HbA1c \geq 6.1%; self- reported | - | - | - | - |
| Park 2006 | - | IFG: FPG ≥ 5.6 | $FPG \ge 7.0$ | - | - | - | - |
| Peterson 2017 | FPG < 6.1 and 2-h PG < 7.8 | IGT: FPG < 7. 0 and 2-h PG ≥ 7.8 to < 11. 1 | | - | Yes | Yes | 2 standardised OGTT at baseline with about 1 week's interval to ver- ify glucose sta- tus |
| Qian 2012 | FPG < 6.1 and 2-h PG < 7.8 | i-IFG: 6.1-6.9 and 2-h PG < 7.8; i-IGT: < 6.1 and 2- h PG 7.8-11. 0; IFG/IGT: 6.1-6.9 and 2- h PG 7.8-11.0 | | 75 g | Yes | Unclear | - |
| Rajala 2000 | 2-h PG < 7.8 | IGT: 2-h PG 7.8 to < 11.1 | 2-h PG ≥ 11. 1; 2x FPG ≥ 6.7 | 75 g | Yes | Yes | New cases identified by OGTTs in 1994 and 1996-8 |

| Ramachan- dran 1986 | - | IGT: 7.8-11.0 (presumed NDDG 1979) | 2-h PG > 11.0 (presumed NDDG 1979) | 75 g | Yes | Yes | - |
|------------------------------|---|---|---|------|-----|---------|---|
| Rasmussen 2008 | - | | FBG ≥ 6.1 or 2-h BG ≥ 11. 1 | 75 g | Yes | Unclear | - |
| Rathmann 2009 | WHO 1999 | 1-6.9; IGT: 2- h PG 7.8 to | idated physi- | 75 g | Yes | Yes | - |
| Rijkeli- jkhuizen 2007 | ADA 1997/ 2003 | IFG _{5.6} : FPG 5. 6-7.0; IFG _{6.1} : FPG 6.1-7.0; IGT: 2-h PG 7.8 to < 11.1 | FPG ≥ 7.0; 2- h PG: ≥ 11.1 | 75 g | Yes | Yes | - |
| Sadeghi 2015 | - | 5.5 and < 7. 0; IGT: 2-h | FPG > 7.0; 2- h OGTT > 11.1; IFG/IGT; an- tihypergly- caemic medi- cation | - | Yes | Yes | - |
| Sasaki 1982 | FPG < 7.8 and 2-h PG < 7.8 (WHO 1980) | IGT: FPG < 7.8 and 2-h PG 7.8-11.1 (WHO 1980) | | 50 g | Yes | Yes | - |
| Sato 2009 | - | (Table 1): IFG: FPG group 6.1-6.9; HbA1c- group: 6.0-6.4 | FPG ≥ 7.0; antihypergly- caemic medi- cation | - | - | - | - |

| Schranz 1989 | - | IGT: 2-h PG ≥ 7.8 to < 11.1 (WHO 1985) | 2-h PG ≥ 11.1 (WHO 1985) | OGTT | Yes | Yes | - |
|-------------------|---|---|---|------|-------------|-----|--|
| Sharifi 2013 | - | FPG 5.6-7.0 | FPG > 7.0 (2 mea- surements) ; diabetes diag- nosis based on documents | OGTT | Yes (twice) | - | - |
| Shin 1997 | - | As- sumed WHO 1985 criteria | | 75 g | Yes | Yes | - |
| Söderberg 2004 | - | IFG: FPG \geq 6.1 to < 7.0 and 2-h PG < 7.8; IGT: FPG < 7.0 and 2-h PG \geq 7.8 to < 11.1 | FPG ≥ 7.0; 2- h PG ≥ 11.1 | 75 g | Yes | Yes | - |
| Song 2015 | - | IFG: FPG 5.6- 6.9 | $FPG \ge 7.0;$ $HbA1c \ge 6.5;$ antihypergly- caemic medi- cation | - | - | - | - |
| Song 2016a | - | | $\begin{array}{l} FG \geq 7.0; \ 2-\\ h \ G \geq 11.0;\\ HbA1c \geq 6.5;\\ self-reported \end{array}$ | 75 g | Yes | Yes | 100 g steamed bread at fol- low-up |
| Soriguer 2008 | | IFG: BG 5. 6-6.1 and 2- h BG < 7.8; IGT: BG < 5. 6 and 2-h BG 7.8-11.1 | | 75 g | Yes | Yes | - |
| Stengard 1992 | - | IGT: 2-h PG 7.8-11.1 | 2-h PG ≥ 11.1 (WHO 1985) ; antihypergly- caemic medi- cations | 75 g | Yes | Yes | - |

| Toshihiro 2008 | FPG < 6.1 and 2-h PG < 7.8 | IFG: FPG 6. 1-6.9 and 2- h PG < 7.8; IGT: FPG < 7. 0 and 2-h PG 7.8-11.1 | h PG > 11. 1; non-fasting | 75 g | Yes | Yes | Annual OGTT dur- ing the obser- vation period (3.2 years) |
|---------------------|---|---|---|------|-----|---------|--|
| Vaccaro 1999 | FPG < 5.6; 2- h PG < 6.7; 2- h PG < 6.7 | IFG: FPG 5.6- 6.0; IGT: 2-h PG 6.7-9.9 | FPG> 6.1; an- tihypergly- caemic medi- cations; 2-h PG \geq 10.0 | 75 g | Yes | No | Retro- spective classi- fication; note thresholds (whole blood) |
| Valdes 2008 | FPG < 5.6 | | $FPG \ge 7.0; 2-$ h PG \ge 11. 1; clinical di- abetes diagno- sis; antihyper- gly- caemic medi- cation, diet | 75 g | Yes | Yes | - |
| Vijayakumar 2017 | - | FG 5.6-6.9; 2- h PG 7.8-11. 9; HbA1c 5.7- 6.4 | $FPG \ge 7.0; 2-$ h $PG \ge 11.1;$ previous clini- cal diagnosis | 75 g | Yes | Yes | HbA1c new method = -0.1916 + (0. 9829 × HbA1c old method) |
| Viswanathan 2007 | FPG and 2-h PG < 6.1 and < 7.8 | IGT: 2-h PG 7.8 to < 11.1 | Not defined, presumably by OGTT | 75 g | Yes | Yes | All partic- ipants under- went a second OGTT to confirm the diagno- sis in order to be included in the study; fol- low-up: a re- minder letter was sent ev- ery 6 months to participants to undergo an OGTT |
| Wang 2007 | - | IFG: FPG 6.1- 6.9; IGT: 2-h PG 7.8-11.0 | $FPG \ge 7.0; 2-h PG \ge 11.1$ | 75 g | Yes | Unclear | - |

| Wang 2011 | | IFG: 5.6 to < 7.0; HbA1c 6. 0 to < 6.5 | | - | - | - | - |
|------------------|-------------------------------|--|---|-------------------------------|---------------|---------|--|
| Warren 2017 | - | FPG 5.6-6.9 (ADA); FG 6. 1-6.9 (WHO) ; 2-h 7.8-11.0 (ADA); HbA1c 5.7-6. 4 (ADA); 6.0- 6.4 (IEC) | Self-report of physician di- agno- sis; antihyper- glycaemic medication re- ported during a study visit or annual tele- phone call | 75 g | Yes (visit 4) | Unclear | - |
| Wat 2001 | FPG and 2-h PG < 7.8 | IGT: FPG < 7. 8 and 2-h PG 7.8 to < 11.1 | | 75 g | Yes | Yes | - |
| Weiss 2005 | FPG < 5.6 and 2-h PG < 7.8 | IGT: FPG < 5. 6 and 2-h PG 7.8-11.1 | h PG > 11.1; presenta- tion of hyper- glycaemia | body weight flavoured glu- | Yes | Yes | OGTT was repeated every 18-24 months |
| Wheelock 2016 | - | | $FPG \ge 7.0; 2-$ h PG $\ge 11.1;$ previous diag- nosis | 75 g | Yes | Unclear | Modified OGTT |

| Wong 2003 | - | | $FPG \ge 7.0; 2-$ h $PG \ge 11.1;$ physician diagnosed dia- betes | 75 g | Yes | Yes | - |
|-------------------|-----------|------------------------------|--|------|-----|-----|---|
| Yeboah 2011 | FPG < 5.6 | IFG: FPG 5.6- 6.9 | FPG > 6.9; an- tihypergly- caemic medi- cation during exami- nations 2,3, 4 | - | - | - | - |
| Zethelius 2004 | - | IGT: 2-h PG 7.8 to < 11.1 | $FPG \ge 7.0;$ antihypergly- caemic medi- cations | 75 g | Yes | No | - |

BG: blood glucose; BW: body weight; FPG: fasting plasma glucose; HbA1c: glycosylated haemoglobin A1c; i-IFG: (isolated) impaired fasting glucose; i-IGT: (isolated) impaired glucose tolerance; IFG/IGT: both impaired fasting glucose and impaired glucose tolerance; NDDG: National Diabetes Data Group; NGT: normal glucose tolerance; OGTT: oral glucose tolerance test; PG: postload glucose; WHO: World Health Organization

Appendix 6. Number of participants with and without intermediate hyperglycaemia at baseline

| Study ID | N partic- ipants with/ without IH | Definitions of | Definitions of IH at baseline | | | | | | | |
|--------------------|---|---------------------------|-------------------------------|--|------------|------------------|----------------|--|--|--|
| | | 'Prediabetes' a (%) | Elevated HbA1c (%) | IFG (%) | IGT (%) | IFG/HbA1c (%) | IFG/IGT (%) | | | |
| Admiraal 2014 | IFG _{5.6} total: 111/456 | - | - | IFG5.6: Total 24.3 South-Asian Surinamese 34.4 African Suri- namese 21.1 "Ethnic Dutch" 22.7 | - | - | - | | | |
| Aekplakorn 2006 | IFG _{5.6} : 223/ 2667 | - | - | IFG _{5.6} : 8.4 | - | - | - | | | |

| Ammari 1998 | IGT: 68 | - | - | - | 100 | - | - |
|--------------------|--|------|---|-----------------------------|-------------|------|-----|
| Anjana 2015 | 'Predi- abetes' (i-IFG, i-IGT or both) : 299/1376 | 21.7 | - | i-IFG _{5.6} : 4.9 | i-IGT: 11.8 | - | 5.0 |
| Bae 2011 | HbA1c _{5.7} : 1791/9723; HbA1c _{6.0} : 412/1791 | - | HbA1c _{5.7} : 18. 4 HbA1c _{6.0} : 4.2 | - | - | - | - |
| Baena-Diez 2011 | IFG _{6.1} : 115 | - | - | IFG _{6.1} : 100 | - | - | - |
| Bai 1999 | IGT: 252/696 | - | - | - | 36.2 | - | - |
| Bergman 2016 | IGT: 68/853 | - | - | - | 8 | - | - |
| Bonora 2011 | HbA1c _{6.0} :70/ 842 | - | 8.3 | - | - | - | - |
| Cederberg 2010 | IFG _{6.1} : 40/ 553 IGT: 103/553 IFG/IGT: 15/ 553 | - | - | IFG _{6.1} : 7.2 | 18.7 | - | 2.7 |
| Chamnan 2011 | HbA1c _{6.0} : 370/5735 | - | HbA1c _{6.0} : 6.5 | - | - | - | - |
| Charles 1997 | IGT: 418/4089; i- IFG _{6.1} : 476/ 5042 | - | - | i-IFG _{6.1} : 9.4 | 10.2 | - | - |
| Chen 2003 | IFG _{6.1} : 156/ 600 | - | - | IFG _{6.1} : 26 | - | - | - |
| Chen 2017 | i-IFG _{5.6} : 329/ 1347 i-IGT: 192/ 1347 IFG/IGT: 209/1347 | - | - | i-IFG _{5.6} : 24.4 | i-IGT: 14.2 | 15.5 | - |

| Coronado- Malagon 2009 | 'Prediabetes': 217/656 | 33.1 | - | - | - | - | - |
|------------------------------|---|---------------------------------------|---|--|--|---|---|
| Cugati 2007 | IFG _{5.6} : 244/ 2123 | - | - | IFG _{5.6} : 11.5 | - | - | - |
| De Abreu 2015 | IFG _{5.6} : 187/ 1167 | - | - | IFG _{5.6} : 16 | - | - | - |
| Den Biggelaar 2016 | IFG _{6.1} and/or IGT: 122/476 | 25.6 | - | - | - | - | - |
| Derakhshan 2016 | | IFG _{5.6} and/or IGT: 6.4 | - | - | - | - | - |
| Dowse 1991 | IGT: 105/ 1201 | - | - | - | 8.7 | - | - |
| Ferrannini 2009 | $\begin{array}{cccc} i-IFG_{5.6}:& 65'\\ 1941\\ i-IFG_{6.1}:& 17'\\ 1941\\ IGT:& 179'\\ 1941\\ i-IGT\\ (IFG_{5.6}):& 57'\\ 1941\\ i-IGT\\ (IFG_{6.1}):& 29'\\ 1941\\ \end{array}$ | - | - | i-IFG _{5.6} : 3.3 i-IFG _{6.1} : 0.9 | IGT: 9.2 i-IGT _{5.6} : 2.9 i-IGT _{6.1} : 1.5 | - | - |
| Filippatos 2016 | IFG _{5.6} : 279/ 1485 | - | - | IFG _{5.6} : 18.8 | - | - | - |
| Forouhi 2007 | IFG _{6.1} : 257/ 1040 IFG _{5.6} : 633/ 1040 | - | - | IFG _{5.6} : 60.9 IFG _{6.1} : 24.7 | - | - | - |
| Garcia 2016 | IFG _{5.6} : 310/ 1777 | - | - | IFG5.5: 17.5 | - | - | - |
| Gautier 2010 | IFG _{5.6} : 979 | - | - | IFG _{5.6} : 100 | - | - | - |
| Gomez- Arbelaez 2015 | 'Prediabetes': 186/772 | 24.1 | - | - | - | - | - |

| | (Men: 61/ 772, women: 125/772) | | | | | | |
|--------------------------------|--|---|---|---|-------------|------|------|
| Guerrero- Romero 2006 | IGT: 75/375 | - | - | - | 20 | - | - |
| Han 2017 | i-IFG _{5.6} : 199/ 7542 i-IGT: 1512/ 7542 IFG/IGT: 198/7542 | - | - | i-IFG _{5.6} : 2.6 | i-IGT: 20.0 | - | 2.6 |
| Hanley 2005 | IGT: 274/882 | - | - | - | 31.6 | - | - |
| Heianza 2012 | IFG _{5.6} : 1680/ 6241 IFG _{6.1} : 380/ 6241 HbA1c _{5.7} : 822/6241 HbA1c _{6.0} : 203/6241 IFG _{5.6} / HbA1c _{5.7} : 2092/6241 | - | HbA1c _{5.7} : 13. 2 HbA1c _{6.0} : 3.3 | IFG _{5.6} : 26.9 IFG _{6.1} : 6.1 | - | 33.5 | - |
| Inoue 1996 | IGT: 37 | - | - | - | 100 | - | - |
| Janghorbani 2015 | i-IFG _{5.6} : 304/ 1530 i-IGT: 198/ 1530 IFG/IGT: 268/1530 | - | - | i-IFG _{5.6} : 19.9 | i-IGT: 12.9 | - | 17.5 |
| Jaru- ratanasirikul 2016 | i-IGT: 27/177 | - | - | - | i-IGT: 15.3 | - | - |
| Jeong 2010 | IFG _{5.6} : 16% IGT: 5.3% | - | - | IFG _{5.6} : 16 | 5.3 | - | - |
| Jiamjaras- rangsi 2008a | IFG _{5.6} : 320/ 2370 | - | - | IFG _{5.6} : 13.5 | - | - | - |
| Kim 2005 | IFG _{6.1} : 276/ 2964 | - | - | IFG _{6.1} : 9.3 | - | - | - |

| Kim 2008 | IFG total: 1829/7211 IFG _{5.6} : 1335/ 7211 IFG _{6.1} : 494/ 7211 | - | - | IFG total: 25. 4 IFG _{5.6} : 18.5 IFG _{6.1} : 6.9 | - | - | - |
|-----------------|---|---|----------------------------------|--|-------------|------|------|
| Kim 2014 | i-IFG _{5.6} : 158/ 406 i-IGT: 65/406 IFG/IGT: 119/406 i-HbA1c _{5.7} : 64/406 | - | i-HbA1c _{5.7} : 15.8 | i-IFG _{5.6} : 38.9 | i-IGT: 16 | - | 29.3 |
| Kim 2016a | IFG _{5.6} : 3544/ 17971 HbA1c _{5.7} : 1713/17971 IFG _{5.6} / HbA1c _{5.7} : 1951/17971 | - | HbA1c _{5.7} : 9.5 | IFG _{5.6} : 19.7 | - | 10.9 | - |
| Kleber 2010 | IGT: 79 | - | - | - | 100 | - | - |
| Kleber 2011 | IGT: 119 | - | - | - | 100 | - | - |
| Ko 1999 | IGT: 123 | - | - | - | 100 | - | - |
| Ko 2001 | IFG _{6.1} : 55/ 319 | - | - | IFG _{6.1} : 17.2 | - | - | - |
| Larsson 2000 | i-IFG _{6.1} : 42/ 265 i-IGT: 66/265 IFG/IGT: 30/ 265 | - | - | i-IFG _{6.1} : 15.8 | i-IGT: 24.9 | - | 11.3 |
| Latifi 2016 | IFG _{5.6} : 124/ 593 | - | - | IFG _{5.6} : 20.9 | - | - | - |
| Lecomte 2007 | IFG _{6.1} : 743 | - | - | IFG _{6.1} : 100 | - | - | - |
| Lee 2016 | HbA1c _{5.7} : 3497 | - | HbA1c _{5.7} : 100 | - | - | - | - |
| Leiva 2014 | IFG _{6.1} : 28/94 | - | - | IFG _{6.1} : 29.8 | - | - | - |

| Levitzky 2008 | Not reported | - | - | - | - | - | - |
|------------------|---|------|---------------------------------|-----------------------------|-------------|------|-----|
| Li 2003 | i-IFG _{6.1} : 42/ 644 i-IGT: 118/ 644 IFG/IGT: 49/ 644 | - | - | i-IFG _{6.1} : 6.5 | i-IGT: 18.3 | - | 7.6 |
| Ligthart 2016 | IFG _{6.1} : 1382/ 10,050 | - | - | IFG _{6.1} : 13.8 | - | - | - |
| Lipska 2013 | IFG _{5.6} : 189/ 1690 i-HbA1c _{5.7} : 207/1690 IFG/HbA1c: 169/1690 | - | i-HbA1c: 12.2 | IFG _{5.6} : 11.2 | - | 10.0 | - |
| Liu 2008 | IFG _{5.6} : 169/ 1844 | - | - | IFG _{5.6} : 9.2 | - | - | - |
| Liu 2014 | 'Prediabetes' (IFG or IGT): 450/2271 | 19.8 | - | - | - | - | - |
| Liu 2016 | IFG _{6.1} : 222/ 1857 | - | - | IFG _{6.1} : 12.0 | - | - | - |
| Liu 2017 | IFG _{5.6} : 3607/ 18610 | - | - | IFG _{5.6} : 19.4 | - | - | - |
| Lorenzo 2003 | IFG _{6.1} : 29/ 1734 IGT: 202/ 1734 | - | - | IFG _{6.1} : 1.7 | 11.6 | - | - |
| Lyssenko 2005 | i-IFG _{6.1} : 263/ 2115 i-IGT: 250/ 2115 IFG/IGT: 173/2115 | - | - | i-IFG _{6.1} : 12.4 | i-IGT: 11.8 | - | 8.2 |
| Magliano 2008 | Not reported | - | - | - | - | - | - |
| Man 2017 | HbA1c _{5.7} : 675/1137 | - | HbA1c _{5.7} : 59. 4 | - | - | - | - |

| Marshall 1994 | IGT: 123 | - | - | - | 100 | - | - |
|-------------------|--|---|---|---|--|---|---|
| McNeely 2003 | 5-6 years: IFG _{6.1} : 30/ 465 IGT: 178/465 10 years: IFG _{6.1} : 28/ 412 IGT: 157/412 | - | - | 5-6 years: IFG _{6.1} : 6.5 10 years: IFG _{6.1} : 6.8 | 5-6 years: 38.3 10 years: 38.1 | - | - |
| Meigs 2003 | i-IFG _{5.6} : 126/ 753 i-IGT (IFG _{5.6}): 115/ 753 IFG _{5.6} /IGT: 103/753 i-IFG _{6.1} : 20/ 753 i-IGT (IFG _{6.1}): 218/ 753 IFG _{6.1} /IGT: 27/753 | - | - | i-IFG _{5.6} : 16.7 i-IFG _{6.1} : 2.7 | i-IGT _{5.6} : 15.3 i-IGT _{6.1} : 29 | - | IFG _{5.6} /IGT: 13.7 IFG _{6.1} /IGT: 3.6 |
| Mohan 2008 | IGT: 37/513 | - | - | - | 7.2 | - | - |
| Motala 2003 | IGT: 35/563 | - | - | - | 6.2 | - | - |
| Motta 2010 | IFG _{6.1} : 295/ 2603 | - | - | IFG _{6.1} : 11.3 | - | - | - |
| Mykkänen 1993 | IGT: 203/892 | - | - | - | 22.8 | - | - |
| Nakagami 2016 | IFG _{5.6} : 467/ 2267 IFG _{6.1} : 134/ 2267 HbA1c _{5.7} : 583/2267 HbA1c _{6.0} : 156/2267 | - | HbA1c _{5.7} : 25. 7 HbA1c _{6.0} : 6.9 | IFG _{5.6} : 20.6 IFG _{6.1} : 5.9 | - | - | - |
| Nakanishi 2004 | IFG _{6.1} : 246/ 5588 | - | - | IFG _{6.1} : 4.4 | - | - | - |

| Noda 2010 | IGF _{5.6} : 558/ 2207 IFG _{6.1} : 153/ 2207 | - | - | IFG _{5.6} : 25.3 IFG _{6.1} : 6.9 | - | - | - |
|------------------------------|---|------|---|--|-------------|---|-----|
| Park 2006 | IFG _{5.6} : 321/ 5296 | - | - | IFG5.6: 6.1 | - | - | - |
| Peterson 2017 | IGT: 29/74 | - | - | - | 39.2 | - | - |
| Qian 2012 | i-IFG _{6.1} : 46/ 1042 i-IGT: 120/ 1042 IFG/IGT: 33/ 1042 | - | - | i-IFG _{6.1} : 4.4 | i-IGT:11.5 | - | 3.2 |
| Rajala 2000 | IGT: 100 | - | - | - | 100 | - | - |
| Ramachan- dran 1986 | IGT: 107 | - | - | - | 100 | - | - |
| Rasmussen 2008 | i-IFG _{5.6} : 607/ 1510 i-IGT 903/ 1510 | - | - | i-IFG _{5.6} : 40.2 | i-IGT: 59.8 | - | - |
| Rathmann 2009 | i-IFG _{6.1} : 71/ 887 i-IGT: 120/ 887 IFG/IGT: 47/ 887 | - | - | i-IFG _{6.1} : 8 | i-IGT: 13.5 | - | 5.3 |
| Rijkeli- jkhuizen 2007 | IFG _{5.6} : 488/ 1428 IFG _{6.1} : 149/ 1428 | - | - | IFG _{5.6} : 34.2 IFG _{6.1} : 10.4 | - | - | - |
| Sadeghi 2015 | 'Predia- betes' (IFG _{5.6} and/or IGT): 373/2980 | 12.5 | - | - | - | - | - |
| Sasaki 1982 | IGT: 13/207 | - | - | - | 6.3 | - | - |
| Sato 2009 | Unclear | - | - | - | - | - | - |

| Schranz 1989 | IGT: 75/2128 | - | - | - | 3.5 | - | - |
|-------------------|---|-------------------------|---|--|--|---|-----|
| Sharifi 2013 | IFG _{5.6} : 123 | - | - | IFG _{5.6} : 100 | - | - | - |
| Shin 1997 | IGT: 153/ 1193 | - | - | - | 12.8 | - | - |
| Söderberg 2004 | i-IFG _{6.1} : 87-98: 402/ 6690 87-92: 149/ 3193 92-98: 253/ 3437 IGT: 87-98: 1253/ 6690 87-92: 600/ 3193 92-98: 662/ 3437 | - | - | i-IFG _{6.1} : 87-98: 6 87-92: 4.7 92-98: 7.4 | 87-98: 18. 9 87-92: 18.8 92-98: 19.3 | - | - |
| Song 2015 | IFG _{5.6} : 321/ 2467 | - | - | IFG _{5.6} : 13 | - | - | - |
| Song 2016a | 'Prediabetes': 344 | 100 | - | - | - | - | - |
| Soriguer 2008 | IFG _{5.6} : 56/ 714 IGT: 54/714 IFG/IGT: 28/ 714 | - | - | IFG5.5: 7.8 | 7.6 | - | 3.9 |
| Stengard 1992 | IGT: 234/637 | - | - | - | 36.7 | - | - |
| Toshihiro 2008 | IFG6.1: 14/ 128 IFG and/or IGT: 114/128 | IFG and/or IGT: 89.1 | - | IFG _{6.1} : 10.9 | - | - | - |
| Vaccaro 1999 | i-IFG _{5.6} : 36/ 1141 i-IGT: 861141 IFG/IGT: 11/ 1141 | - | - | i-IFG _{5.6} : 3.1 | i-IGT: 7.5 | - | 1.0 |

| Valdes 2008 | IFG _{5.6} : 114/ 630 IFG _{6.1} : 52/ 630 IGT: 50/630 | - | - | IFG _{5.6} : 18.1 IFG _{6.1} : 8.3 | 7.9 | - | - |
|---------------------|--|---|--|---|--|---|--|
| Vijayakumar 2017 | IFG _{5.6} adults: 423/2005 IFG _{5.6} children: 193/ 2095 HbA1c _{5.7} adults: 168/ 2005 HbA1c _{5.7} children: 62/ 2095 IGT adults: 347/2005 IGT children: 170/2095 IFG/IGT adults: 169/ 2005 IFG/ IGT children: 53/2095 | - | HbA1c _{5.7} adults: 8.4 HbA1c _{5.7} children: 3.0 | IFG _{5.6} adults: 21.1 IFG _{5.6} children: 9.2 | Adults: 17.3 Children: 8.1 | - | IFG/IGT adults: 8.4 IFG/IGT children: 2.5 |
| Viswanathan 2007 | IGT: 619/ 1659 | - | - | - | 37.3 | - | - |
| Wang 2007 | IGT: 141/541 | - | - | - | 26 | - | - |
| Wang 2011 | i-IGT total: 135/10 i-IGT men: 29/447 i-IGT women: 106/ 635 | - | - | - | i-IGT total: 12.5 i-IGT men: 6. 5 i-IGT women: 16.7 | - | - |
| Warren 2017 | IFG _{5.6} : 4112/ 10844 IFG _{6.1} : 1213/ 10844 IGT: 2009/ 7194 HbA1c _{5.7} : 2027/10844 | - | HbA1c _{5.7} : 19 HbA1c _{6.0} : 9 | IFG _{5.6} : 38 IFG _{6.1} : 11 | 28 | - | - |

| | HbA1c _{6.0} : 970/10844 | | | | | | |
|-------------------|--|---|---|---------------------------|-------------|---|---|
| Wat 2001 | IGT: 322 | - | - | - | 100 | - | - |
| Weiss 2005 | i-IGT (IFG _{5.6}): 33/ 117 | - | - | - | i-IGT: 28.2 | - | - |
| Wheelock 2016 | IGT: 169/ 5532 | - | - | - | 3.1 | - | - |
| Wong 2003 | IGT: 291 | - | - | - | 100 | - | - |
| Yeboah 2011 | IFG _{5.6} : 940/ 6753 | - | - | IFG _{5.6} : 13.9 | - | - | - |
| Zethelius 2004 | IGT: 201/667 | - | - | - | 30.1 | - | - |

^aTerm 'prediabetes' as used by study authors (usually defined by various combinations of glycaemic status measurements, e.g. IFG *and/or* IGT)

FG: fasting glucose; **FPG**: fasting plasma glucose; **HbA1c**: glycosylated haemoglobin A1c; **HbA1c**_{5.7/6.0}: HbA1c threshold 5.7% or 6.0% (usually reflecting 5.7% to 6.4% and 6.0% to 6.4%, respectively); **HbA1c/IFG**: both HbA1c and IFG; **i**-: isolated;**IFG**_{5.6/6.1}: impaired fasting glucose (threshold 5.6 mmol/L or 6.1 mmol/L); **IGT**: impaired glucose tolerance; **IFG/IGT**: both IFG and IGT; **PG**: postload glucose;**IH**: intermediate hyperglycaemia; **T2DM**: type 2 diabetes mellitus

Appendix 7. Follow-up time and type of outcome measurement of the development of type 2 diabetes

| Study ID | Length of follow-up | Time-points of mea- surements | Outcome measurement of the development of T2DM | Notes |
|-----------------|----------------------------|----------------------------------|--|---|
| Admiraal 2014 | 10 years | Baseline, follow-up | Incidence, odds ratio | Data for total popu- lation/South-Asian Suri- namese/African Suri- namese/"Ethnic Dutch" |
| Aekplakorn 2006 | 12 years | Baseline, follow-up | Incidence, odds ratio | - |
| Ammari 1998 | 2 years | Baseline, follow-up | Incidence | - |
| Anjana 2015 | Median 9.1 years (IQR 2.6) | Baseline, follow-up | Incidence, incidence rate | - |

| Bae 2011 | 4 years (mean 47.2 months) | Baseline, follow-up (par- tially annually/biannu- ally) | | - |
|--------------------------|----------------------------|---|--|--|
| Baena-Diez 2011 | 10 years | Baseline, follow-up | Incidence | - |
| Bai 1999 | 1 year | Baseline, follow-up | Incidence | - |
| Bergman 2016 | 24 years | Baseline, follow-up | Incidence, odds ratio | Also adjusted for fast- ing blood glucose; 100 g OGTT |
| Bonora 2011 | 15 years | Baseline, follow-up (5, 10, 15 years) | Incidence, incidence rate, hazard ratio | HbA1c category used: 6. 0% to 6.49% |
| Cederberg 2010 | Mean 9.7 years (SD 0.7) | Baseline, follow-up | Incidence, risk ratio | Total incident cases = mixture of isolated and combined intermediate glycaemic conditions |
| Chamnan 2011 | Median 3 years | Baseline, follow-up | Incidence, odds ratio | Data for HbA1c 6.0% to 6.4% group, focus on clinically and/or bio- chemically diagnosed di- abetes |
| Charles 1997 | 2 years | Baseline, follow-up (5 annual clinical examina- tions) | Incidence | - |
| Chen 2003 | 3 years | Baseline, follow-up | Incidence, odds ratio | Also adjusted for apolipoprotein B |
| Chen 2017 | 3 years | Baseline, follow-up | Incidence | - |
| Coronado-Malagon 2009 | 1 and 2 years | Baseline, follow-up | Incidence, relative risk | Results are given for year 1/year 2 of follow-up |
| Cugati 2007 | 10 years | Baseline, follow-up (5 and 10 years) | Incidence, odds ratio | Odds-ratio, age-and sex- adjusted |
| De Abreu 2015 | 10 years | Baseline, follow-up | Incidence, incidence rate, odds ratio | Age-standard- ised incidence rate; addi- tional co- variates: metabolic syn- drome, fasting glucose at baseline |

| Den Biggelaar 2016 | 7 years | Baseline, follow-up | Incidence | - |
|-------------------------|-------------------------------------|---|--|---|
| Derakhshan 2016 | Median 11.7 years (IQR 8.4-13.2) | Baseline, follow-up | Incidence rate, hazard ratio | - |
| Dowse 1991 | Approx. 5 years | Baseline, follow-up | Incidence, incidence rate, odds ratio | Incidence rates for the periods 1975/76- 1982 and 1982-1987 |
| Ferrannini 2009 | 7 years | Baseline, follow-up | Incidence, relative risk | - |
| Filippatos 2016 | 10 years | Baseline, follow-up (in- termediate 5 -year fol- low-up) | Incidence, odds ratio | - |
| Forouhi 2007 | 10 years | Baseline, follow-up | Incidence, incidence rate, hazard ratio | Cumulative incidence increased across increas- ing age groups and was higher in men than in women |
| Garcia 2016 | Approx. 9 years | Baseline, follow-up (ev- ery 12-15 months, max. 6 follow-ups) | Incidence | - |
| Gautier 2010 | 9 years | Baseline, follow-up (3- yearly examinations) | Incidence | - |
| Gomez-Arbelaez 2015 | Approx. 2 years | Baseline, follow-up | Incidence, incidence rate | Rate was given in terms of per 100 person-years (recalculated to 1000 person-years) |
| Guerrero-Romero 2006 | 5 years | Baseline, follow-up | Incidence, incidence rate | - |
| Han 2017 | 12 years | - | Incidence, incidence rate, hazard ratio | - |
| Hanley 2005 | Average 5.2 years (range 4.5-6.6) | Baseline, follow-up | Incidence, odds ratio | - |
| Heianza 2012 | Median 5 years | Baseline, follow-up (an- nual follow-ups) | Incidence, incidence rate, hazard ratio | Adjusted odds ratios: mean age and sex-ad- justed |
| Inoue 1996 | 2.5 years | Baseline, follow-up | Incidence | - |

| Janghorbani 2015 | Mean 6.8 years (SD 1.7) | Baseline, follow-up (OGTT at 3- year intervals) | | Date for cohort without hypertension |
|------------------------|--|---|--|--|
| Jaruratanasirikul 2016 | 3-6 years | Baseline, follow-up | Incidence | - |
| Jeong 2010 | 5 years | Baseline, follow-up | Odds ratio | Also adjusted for ALAT, ASAT, γ-GT, h-CRP |
| Jiamjarasrangsi 2008a | Mean 2.6 years (SD 0. 97) | Baseline, follow-up (an- nual follow-ups, 1-4 years) | Incidence | - |
| Kim 2005 | 5 years | Baseline, follow-up | Incidence, hazard ratio | - |
| Kim 2008 | 2 years | Baseline, follow-up | Incidence | - |
| Kim 2014 | Median 46 months | Baseline, follow-up (ev- ery 3-6 months, up to 9 years) | Incidence | 81 par- ticipants were diagnosed with diabetes with a con- version rate of 20% (81/ 406); conversion rates are given within predi- abetes groups (e.g. 24/ 158 i-IFG converters = 15.2%) |
| Kim 2016a | Mean 5.2 years (range 3. 1-6.7) | Baseline, follow-up | Incidence, odds ratio | - |
| Kleber 2010 | 1 year | Baseline, follow-up | Incidence | - |
| Kleber 2011 | Mean 3.9 years (SD 0.6) | Baseline, follow-up | Incidence | - |
| Ко 1999 | Mean 1.4 years (range 0. 9-7.6) | Baseline, follow-up (an- nual OGTTs) | Incidence | - |
| Ко 2001 | Median 1.7 years | Baseline, follow-up (an- nual OGTTs) | Incidence | - |
| Larsson 2000 | Mean 10 years (SD 1 year 10 months) | Baseline, follow-up | Incidence | - |
| Latifi 2016 | Median 5 years | Baseline, follow-up | Incidence, incidence rate, odds ratio | - |
| Lecomte 2007 | 5 years | Baseline, follow-up | Incidence | - |

| Lee 2016 | Mean 3.7 years (SD 2.3) | Baseline, follow-up | Incidence | - |
|---------------|-------------------------------------|--|---|---|
| Leiva 2014 | 6 years | Baseline, follow-up | Incidence, hazard ratio | - |
| Levitzky 2008 | 4 years | Baseline, follow-up (ap- prox. 4-year intervals) | Incidence, odds ratio | - |
| Li 2003 | 5 years | Baseline, follow-up (ex- amination every 2 years) | | Incidence rates for 5- year cumulative inci- dence; further adjust- ments for HOMA-IR and HOMA beta-cell |
| Ligthart 2016 | 14.7 years | Baseline, follow-up (blood glucose measures approx. every 4 years) | Incidence rate | - |
| Lipska 2013 | 7 years | Baseline (year 4), follow- up (years 5,6,7) | Incidence, odds ratio | IFG _{6.1} : sensitivity anal- ysis, analysis for 'ethnic- ity', sex analysis |
| Liu 2008 | 5 years | Baseline, follow-up | Incidence, incidence rate, relative risk | - |
| Liu 2014 | 3 years | Baseline, follow-up | Incidence, incidence rate | No exact definition of 'prediabetes' and dia- betes incidence |
| Liu 2016 | Median 10.9 years (IQR 8.0-15.3) | Baseline, follow-up | Hazard ratio | Subdistribution hazard ratios; also adjusted for self-rated health |
| Liu 2017 | 7.8 years | Baseline, follow-up | Odds ratio | - |
| Lorenzo 2003 | 7-8 years | Baseline, follow-up | Incidence, odds ratio | Also adjusted for NCEP metabolic syndrome def- inition, fasting insulin |
| Lyssenko 2005 | Median 6 years (range 2- 12) | Baseline, follow-up (ev- ery 2-3 years) | Incidence, hazard ratio | 1372 persons 1 visit, 392 persons 2 visits, 219 per- sons 3 visits, 132 persons 4 visits |
| Magliano 2008 | 5 years | Baseline, follow-up | Incidence, incidence rate, odds ratio | 5-year cu- mulative incidence rate was standardised to the 1998 Australian popula- tion (age and sex-specific |

| | | | | incidence rates) |
|-------------------|---|---|--|---|
| Man 2017 | 6 years | Baseline, follow-up | Incidence, incidence rate, risk ratio | Male: female, age stan- dardised rate |
| Marshall 1994 | Mean 22.6 months (range 11-40) | Baseline, follow-up | Incidence | - |
| McNeely 2003 | 10 years | Baseline, follow-up (5-6 years and 10 years) | Incidence | - |
| Meigs 2003 | 5 years, 10 years | Baseline, follow-up (3 to 10 biennial examina- tions) | Incidence, incidence rate | - |
| Mohan 2008 | Mean 8 years (SD 1.3) | Baseline, follow-up | Incidence, incidence rate | - |
| Motala 2003 | 10 years | Baseline, follow-up | Incidence | - |
| Motta 2010 | 3 years | Baseline, follow-up | Incidence | - |
| Mykkänen 1993 | Mean 3.5 years (42 months (SD 4)) | Baseline, follow-up | Incidence, odds ratio | - |
| Nakagami 2016 | 5 years | Baseline, follow-up | Incidence, hazard ratio | - |
| Nakanishi 2004 | 7 years | Baseline, follow-up (an- nual health examina- tions) | | Also adjusted for all other components of the metabolic syndrome at study entry |
| Noda 2010 | 5 years | Baseline, follow-up | Incidence | - |
| Park 2006 | Mean 4.1 years | Baseline, follow-up (an- nual examinations) | Incidence, incidence rate | - |
| Peterson 2017 | 10 years | Baseline, follow-up | Incidence | - |
| Qian 2012 | 5 years | Baseline, follow-up | Incidence | - |
| Rajala 2000 | 4.6 years (1.9-6.4) | Baseline, follow-up (in- cluding a separate co- hort) | Incidence, incidence rate | - |
| Ramachandran 1986 | Reverters: 3.3 years (SD 2) Converters: 5.1 years (SD 3.5) | Baseline, follow-up ("pe- riodically") | Incidence | All individuals were ad- vised a calorie-restricted high carbohydrate high- fibre diet |

| Rasmussen 2008 | 3.5 years i-IFG _{5.6} : median 2.5 years i-IGT: median 2.1 years | Baseline, follow-up | Incidence, incidence rate | - |
|----------------------|--|--|---|---|
| Rathmann 2009 | 7 years | Baseline, follow-up | Incidence, incidence rate, odds ratio | - |
| Rijkelijkhuizen 2007 | Mean 6.4 years | Baseline, follow-up | Incidence, incidence rate | - |
| Sadeghi 2015 | 7 years | Baseline, follow-up (biannual) | Incidence, incidence rate | - |
| Sasaki 1982 | 7 years | Baseline, follow-up | Incidence,odds ratio | - |
| Sato 2009 | 4 years | Baseline, follow-up | Odds ratio | - |
| Schranz 1989 | 6 years | Baseline, follow-up | Incidence | - |
| Sharifi 2013 | 7 years | Baseline, follow-up | Incidence | - |
| Shin 1997 | 2 years | Baseline, follow-up | Incidence | - |
| Söderberg 2004 | 11 years | Baseline, follow-up | Incidence, incidence rate | Incidence rates are given for periods 1987-1992 and 1992-1998, strati- fied by men:women |
| Song 2015 | Median 3.97 years | Baseline, follow-up | Incidence, relative risk | Also adjusted for glucose |
| Song 2016a | Mean 10.8 years (range 10.5-12) | Baseline, follow-up (ad- ditional follow-up 2014) | Incidence | - |
| Soriguer 2008 | Mean 6 years | Baseline, follow-up | Incidence, incidence rate, relative risk | - |
| Stengard 1992 | 5 years | Baseline, follow-up | Incidence, odds ratio | - |
| Toshihiro 2008 | Mean 3.2 years (SD 0.1) | Baseline, follow-up (an- nual OGTT) | Incidence | - |
| Vaccaro 1999 | 11.5 years | Baseline, follow-up | Incidence, odds ratio | Odds ratios probably un- adjusted |
| Valdes 2008 | Mean 6.3 years (5.9-6.8) | Baseline, follow-up | Incidence, incidence rate, odds ratio | Also adjusted for 2-h PG |

| Vijayakumar 2017 | Adults median 4.6 years (IQR 2.8-7.9) Children: median 5.2 years (IQR 2.7-9.6) | Baseline, follow-up (ex- aminations every 2 years) | Incidence, incidence rate | Data for adults/children; incidence rate taken from figure 2 (boys:men; girls:women) | |
|------------------|---|---|---------------------------|--|--|
| Viswanathan 2007 | Median 5 years | Baseline, follow-up (re- minder to undergo an OGTT every 6 months) | Incidence, odds ratio | Also adjusted for FPG and 2-h PG | |
| Wang 2007 | 5 years | Baseline, follow-up | Incidence, risk ratio | - | |
| Wang 2011 | 4 years | Baseline, follow-up | Odds ratio | Unclear which confounders were used in the multivariate model | |
| Warren 2017 | Cohort 1 (visit 2): 22 years Cohort 2 (visit 4): 16 years | Baseline, follow-up (3 visits every 3 years, 5th visit 2011-13) | Hazard ratio | Data for IFG _{5.6} , IFG _{6.1} , HbA1c _{5.7} , HbA1c _{6.0} , IGT (cohort 2 only) | |
| Wat 2001 | 2 years | Baseline, follow-up | Incidence | - | |
| Weiss 2005 | Mean 20.4 months (SD 10.3) | Baseline, follow-up (biannual) | Incidence | - | |
| Wheelock 2016 | Median 12.4 years (IQR 6.0-22.9) | Baseline, follow-up (ap- prox. annual intervals for repeated OGTTs) | Incidence | Non-overweight partici- pants with IGT cohort and overweight partici- pants with IGT group | |
| Wong 2003 | 8 years | Baseline, follow-up | Incidence | Odds ratios from Tai 2004 | |
| Yeboah 2011 | 7.5 years | Baseline, follow-up (3 examinations) | Incidence, hazard ratio | - | |
| Zethelius 2004 | 7 years | Baseline, follow-up | Odds ratio | Also adjusted for (split) proinsulin, intact insulin | |

ALAT: alanine aminotransferase; ASAT: aspartate transaminase; FG: fasting glucose; FPG: fasting plasma glucose; h-CRP: highsensitivity C-reactive protein; HOMA-beta: homeostatic model assessment of beta-cell function; HOMA-IR: homeostatic model assessment of insulin resistance; HbA1c: glycosylated haemoglobin A1c; HbA1c_{5.7/6.0}: HbA1c threshold 5.7% or 6.0% (usually reflecting 5.7% to 6.4% and 6.0% to 6.4%, respectively); HbA1c/IFG: both HbA1c and IFG; i-: isolated; IFG_{5.6/6.1}: impaired fasting glucose (threshold 5.6 mmol/L or 6.1 mmol/L); IGT: impaired glucose tolerance; IFG/IGT: both IFG and IGT; IQR: interquartile range; NCEP: national cholesterol education program; OGTT: oral glucose tolerance test; PG: postload glucose; SD: standard deviation; T2DM: type 2 diabetes mellitus; γ -GT: gamma-glutamyl transferase/transpeptidase

Appendix 8. Baseline characteristics (I)

| Study ID | Setting | N participants in origi- nal cohort (several phases of the cohort study) | (several phases of the | Notes |
|-----------------|---------------------------------|---|--------------------------------|---|
| Admiraal 2014 | Amsterdam, The Netherlands | 2975 | 2975 456 | |
| Aekplakorn 2006 | Bangkok, Thailand | 3499/3245 | 2667 | Baseline data for cohort becoming diabetic (N = 361) |
| Ammari 1998 | Jordan | Unclear | 121/68-200/144 (con- trols) | Few baseline data re- ported for study popula- tion (N = 212) |
| Anjana 2015 | Chennai, India | 26,001 | 3589/2207 | Baseline data for cohort becoming diabetic at fol- low-up (N = 176) |
| Bae 2011 | South Korea | 10,959 | 9723 | Baseline data for the total cohort (N = 9723) |
| Baena-Diez 2011 | Barcelona, Spain | 2248 | 168 | Baseline data for predia- betic cohort (N = 115) |
| Bai 1999 | Chennai, India | 4885/1082 | 1082/696 | Baseline data for the IGT cohort (N = 252) |
| Bergman 2016 | Israel | 1970 | 1037 | Baseline data for IGT co- hort (N = 24) |
| Bonora 2011 | Bruneck (South Tyrol), Italy | 1000 | 936 | No baseline data (except white participants aged > 40 years, N = 919) |
| Cederberg 2010 | Finland | 593 | 553/499 | Baseline data for the co- hort (total N = 553, men N = 223, women N = 330) |

| Chamnan 2011 | Norfolk (East Anglia), UK | 77,630/25,639 | 6372/5735 | Baseline data for HbA1c _{6.0-6.4} cohort (N = 370) |
|--------------------------|---------------------------------------|---------------|---|---|
| Charles 1997 | Paris, France | Unclear | 7540 (2nd clinical exam- ination)/4089 | Baseline data for individ- uals with IGT convert- ing to T2DM (N = 32) |
| Chen 2003 | Penghu, Taiwan | 1601 | 1306/600 | Baseline data for cohort converting to T2DM (N = 26) |
| Chen 2017 | China | 8845 | 1374 | Baseline data for i-IFG/i- IGTand IFG/IGT across age groups < 40 years + > 60 years (data indicate range across groups) (i- IFG < 40 years N = 51 and > 60 years N = 278; i-IGT < 40 years N = 41 and > 60 years N = 151; IFG/IGT: < 40 years N = 34 and > 60 years N = 175) |
| Coronado-Malagon 2009 | Mexico | 820 | 656 | Baseline characteristics for the prediabetic co- hort (N = 217) |
| Cugati 2007 | Australia, Blue Moun- tains region | 4433/3654 | 2335 (5 years)/1952 (10 years)/2123 complete data (10 years) | Baseline data for people without diabetes (N = 3437) |
| De Abreu 2015 | Australia | Unclear | 1167/395 (IFG _{5.6}) | Baseline data for IFG co- hort at baseline (N = 187) |
| Den Biggelaar 2016 | The Netherlands | 574/491 | 476 | Baseline data for predia- betic group (N = 122) |
| Derakhshan 2016 | Tehran, Iran | 12808 | 8231 | Baseline data for predia- betes group with normal blood pressure |
| Dowse 1991 | Nauru, Micronesia | 1497/1201 | 830 (1982/1987-includ- ing 143 nondiabetic per- son from 1975/76) | No baseline data pro- vided |

| Ferrannini 2009 | Mexico | 3505 | 2282/1963 | Baseline characteristics: range across different definitions of predia- betes | |
|-------------------------|------------------------------|---------------------------------------|---------------|--|--|
| Filippatos 2016 | Attica, Greece | 4056/3042/1875 | 1485 | Baseline data for IFG _{5.6} cohort (N = 343) | |
| Forouhi 2007 | Ely (Cambridgeshire), UK | 1571/1122 (phase 1)/ 912 (phase 2) | 683 (phase 3) | Baseline data for IFG _{6.1} cohort (N = 257) | |
| Garcia 2016 | Sacramento (CA), USA | 1789 | 1777 | Baseline data for predia- betic cohort (N = 310) | |
| Gautier 2010 | France | 3817 | 979 | No baseline data | |
| Gomez-Arbelaez 2015 | Columbia | 2012 | 772 | Baseline data for the total cohort (N = 772) | |
| Guerrero-Romero 2006 | Durango, Mexico | Unclear | 375 | Baseline data for IGT cohort at baseline pro- gressing to T2DM (N = 20); all individuals were counselled on the impor- tance of diet and physical exercise (standard care for the whole cohort) | |
| Han 2017 | Ansung-Ansan, South Korea | 10,030 | 7542 | Baseline data for i-IFG, i-IGT and IFG/IGT co- hort | |
| Hanley 2005 | ley 2005 USA | | 822 | Baseline data for dia- betic cohort at follow- up (N = 131); par- ticipants were recruited from 2 population-based studies: the San Anto- nio Heart Study and the San Luis Valley diabetes study | |
| Heianza 2012 | Japan | 32057 | 6636/6241 | Baseline data for total co- hort (N = 6241) | |
| Inoue 1996 | Gunma (Gyeonggi), Japan | Unclear | Unclear | Baseline data for the IGT cohort (N = 37) | |

| Janghorbani 2015 | Isfahan, Iran | 3370 | 1489 | Baseline data for i-IFG, i-IGT and IFG/IGT co- hort at baseline (N = 770); first-degree rela- tives of people with T2DM | |
|------------------------|---------------------------------|---------------|-----------|--|--|
| Jaruratanasirikul 2016 | Thailand | 181 | 177 (157) | Baseline data for IGT co- hort (N = 27) | |
| Jeong 2010 | Dalseong County, South Korea | 1806/1599 | 1474 | 1287 participants were re-evaluated in 2008 and 187 new participants "added to the study"; baseline data for partici- pants with incident dia- betes (N = 135) | |
| Jiamjarasrangsi 2008a | Bangkok, Thailand | 3989 | 3243/2370 | Baseline data for total co- hort becoming diabetic at follow-up (N = 48) | |
| Kim 2005 | Seoul, South Korea | 20,203/15,936 | 2964 | Baseline data for FPG group 4 (6.1-7.0) with baseline and follow-up $(N = 276)$ | |
| Kim 2008 | Incheon, South Korea | 7510 | 7211 | Baseline data for IFG _{5.6} / IFG _{6.1} cohort (N = 1335/494) | |
| Kim 2014 | Seoul, South Korea | 418 | 418 | Baseline data for i-IFG (N = 158)/i-IGT (N = 65)/IFG/IGT (N = 119) /i-HbA1c (N = 64); total (N = 406) | |
| Kim 2016a | Seoul, South Korea | 19,356 | 17,971 | 2 baseline data cohorts: prediabetes by FPG only and HbA1c only (N = 3544 and N = 1713) | |
| Kleber 2010 | Germany | 79 | 79 | Baseline data for IGT co- hort (N = 79) | |
| Kleber 2011 | Germany | 128 | 128 | Baseline data for IFG co- hort (N = 128) | |

| Ко 1999 | Hong Kong | 123 | 123 | Baseline data for the IGT cohort (N = 123) |
|---------------|---------------------------------|---------------|---------------|--|
| Ко 2001 | Hong Kong | 657 | 319 | Baseline data for IFG co- hort (N = 55) |
| Larsson 2000 | Sweden | 1843 | 265 | Baseline data for i-IGT (N = 66)/i-IFG (N = 42)/IFG/IGT (N = 30) ; 265 follow-up par- ticipants were randomly sampled from each glu- cose tolerance group of the original cohort and invited for follow-up |
| Latifi 2016 | Ahvaz (Khuzestan), Iran | 12,514/6640 | Unclear/593 | Baseline for prediabetic cohort becoming dia- betic at follow-up |
| Lecomte 2007 | France | 56,650 | 4532 | Baseline data for IFG co- hort attending both ex- aminations (N = 743) |
| Lee 2016 | South Korea | 6246 | 5528 | Baseline data for the total cohort (N = 3497) |
| Leiva 2014 | Chile | 1007 | 177 | Most baseline data for cohort becoming dia- betic at follow-up (N = 94 with IFG) |
| Levitzky 2008 | Framingham (MA), USA | Unclear | 3634 | Baseline data for individ- uals on first exam, free of cardiovascular disease (N = 4058) |
| Li 2003 | Kinmen, Taiwan | Unclear | 644 | Baseline data for i-IGT (N = 118)/i-IFG (N = 42)/IFG/IGT (N = 49) |
| Ligthart 2016 | Rotterdam, The Nether- lands | 14,926/11,740 | 11,740/10,050 | Baseline data for predia- betic cohort (N = 1382) |
| Lipska 2013 | USA | 3075 | 1690 | Baseline data for i-IFG (N = 189)/i-HbA1c _{5.7} (N = 207)/IFG/HbA1c (N = 169) |

| Liu 2008 | Jiang Su province, China | 6400/5888 | 1844 | Baseline data for non-di- abetic participants (N = 1844); M (N = 788)/W (N = 1056) | |
|---------------|---|---------------|---------------------------------|---|--|
| Liu 2014 | Shanghai, China | 4556 | 3174 | Baseline data for the pre- diabetic cohort convert- ing to T2DM (N = 78) | |
| Liu 2016 | Beijing, China | 2101 | 1857 | Baseline data for partici- pants without diabetes at baseline (N = 1857) | |
| Liu 2017 | China | 27,020 | 23,626/18,610 | Baseline data for IFG co- hort at baseline (N = 3607) | |
| Lorenzo 2003 | San Antonio (TX), USA | 2941/2569 | 1734 | Baseline data for cohort converting to T2DM (N = 195) | |
| Lyssenko 2005 | Finland | Unclear | 2115 | Baseline data for IFG- IGT individuals who converted to T2DM (N = 86) | |
| Magliano 2008 | Australia | 20,347/11,247 | 6537 | Baseline data for cohort becoming diabetic at fol- low-up (N = 224) | |
| Man 2017 | Singapore | 3280 | 1279/1137 | Baseline data for inci- dent diabetes cohort (N = 127) | |
| Marshall 1994 | Colorado, USA | 1321 | 173/134 | Baseline data for IGT cohort convert- ing to T2DM (N = 20) | |
| McNeely 2003 | Seattle (WA), USA | 518 | 465 (5 years)/412 (10 years) | Baseline data for cohort converting to T2DM at 5-6 years (N = 50) and 10 years (N = 74) | |
| Meigs 2003 | Baltimore (MD) and Washington, D.C., USA | Unclear | 815/753 | Baseline data for the IFG-IGT cohort (N = 265); follow-up time: at least 6 years 77%, at least 10 years 44%, at least | |

| | | | | 16 years 16%, at least 20 years 4.5% |
|-------------------|---|--------------|-------------------------------------|---|
| Mohan 2008 | Chennai, India | 1061 | 513 | Baseline data for cohort becoming diabetic at fol- low-up (N = 64) |
| Motala 2003 | Durban (KwaZulu-Na- tal), South Africa | 2479 | 563 | Baseline data for respon- ders (both baseline and follow-up examination) (N = 563) |
| Motta 2010 | Italy | 2603 | 2603 | No baseline data pro- vided |
| Mykkänen 1993 | Kuopio (Northern Savo- nia), Finland | 1300 | 1054/892 | Baseline data for cohort developing T2DM (N = 69) |
| Nakagami 2016 | Japan | 6012 | 2770/2267 | Baseline data for cohort converting to T2DM (N = 99) |
| Nakanishi 2004 | Japan | Unclear/6812 | 5746 | Baseline characteristics for IFG cohort (N = 246) |
| Noda 2010 | Japan | 22387 | 2207 | Baseline characteristics for the total cohort (N = 2207) |
| Park 2006 | South Korea | 6305 | 5557 | Baseline data for in- cident diabetic partici- pants with IFG at base- line (N = 40) |
| Peterson 2017 | Sweden | 119 | 87/74/29 | Baseline data for IGT co- hort (N = 29) |
| Qian 2012 | Shanghai, China | 1869 | 1042 | Baseline data for cohort progressing to T2DM (N = 377) |
| Rajala 2000 | Oulo (North Ostroboth- nia), Finland | 1008/768 | 183 (1st)/193 (2nd, other group) | Few baseline data for IGT cohort (N = 171) |
| Ramachandran 1986 | Madras, India | Unclear | 107 | Baseline data for the dia- betic cohort at follow-up (N = 39) |

| Rasmussen 2008 | Denmark | 1821 | 1510/1002 | Baseline data for IFG (N = 607)/IGT cohort (N = 903) |
|----------------------|---------------------------------|----------------|-----------|--|
| Rathmann 2009 | Augsburg (Bavaria), Germany | 2656 | 1202 | Baseline data for total co- hort (follow-up partici- pants, age-group 55-74 years, N = 887) |
| Rijkelijkhuizen 2007 | The Netherlands | 2484/1513 | 1428 | Baseline data for IFG _{6.1} (N = 149)/IFG _{5.6} (N = 488) |
| Sadeghi 2015 | Isfahan, Iran | 6323 | 2980 | Baseline data for predi- abetic cohort becoming diabetic at follow-up (N = 131) |
| Sasaki 1982 | Osaka, Japan | 507 | 207 | Baseline data for the IGT cohort (N = 13) |
| Sato 2009 | Japan | 12,647 | 9116/6804 | Baseline data for cohort becoming diabetic at fol- low-up (N = 659) |
| Schranz 1989 | Malta | 2128 | 1422 | Baseline data for diabetic cohort at follow-up (N = 166) |
| Sharifi 2013 | Zanjan, Iran | 2941 | 395 | Baseline data for active participants (N = 123) |
| Shin 1997 | Yonchon County, South Korea | 2520/2293 | 2248/1193 | Baseline data for in- dividuals converting to T2DM (N = 67) |
| Söderberg 2004 | Mauritius | 5083/6616/6291 | Unclear | Baseline data for cohort 1987-1998 (N = 2631) , 10 years follow-up; 3 cohorts 1987-1992 (N = 3680), 1992-1998 (N = 4178), 1987-1998 (N = 2631) |
| Song 2015 | ng 2015 South Korea 4899 | | 2079 | Baseline data for predi- abetic cohort (men N = 154; women N = 167; total N = 321) |

| Song 2016a | Shanghai, China | 2132 | 778/526 | Baseline data for predia- betic cohort (N = 334) |
|------------------|--|-------------|---|---|
| Soriguer 2008 | Pizarra (Andalusia), Spain | 1051 | 824 | Baseline data for final sample of follow-up (N = 714) |
| Stengard 1992 | Finland | 1711 | 716/637 | Baseline data for IGT cohort convert- ing to T2DM (N = 17) |
| Toshihiro 2008 | Japan | 732 | 128 | Baseline data for cohort becoming diabetic at fol- low-up (N = 36); partic- ipants with IFG and/or IGT were given advice about lifestyle modifica- tions once or twice a year |
| Vaccaro 1999 | Naples, Italy | 1285/1245 | 1141/560 | Baseline data for total co- hort (follow-up exami- nation N = 560) |
| Valdes 2008 | Spain | 1626/1034 | 943/630 | Baseline data for IFG _{5.6-6.1} (N = 114) /IFG _{6.1-6.9} (N = 52) |
| Vijayakumar 2017 | Phoenix (AZ), USA | Unclear | 2095 (10-19 years)/ 2005 (20-39 years) | Baseline data for adults/ children with HbA1c 5. 7%-6.4% (children N = 62, adults N = 168) |
| Viswanathan 2007 | India (probably Chen- nai) | 4084 | 1659 | Baseline data for IGT group (N = 619); partic- ipants were given advice on preventive measures such as dietary modifica- tions and regular exercise |
| Wang 2007 | Beijing, China | 20,682/1566 | 902 | Baseline data for cohort with incident diabetes and no coronary heart disease (N = 67) |
| Wang 2011 | Arizona/North/South Dakota/Oklahoma, USA | Unclear | 2849/1670 (2nd exam) | No baseline data |

| Warren 2017 | USA, 4 communities | 15,792 | 10844: 1990-1992 (FG, HbA1c) as baseline | diabetes definitions (visit 2: IFG _{5.6-6.9} N = 4112; |
|----------------|--------------------|----------------|---|---|
| Wat 2001 | Hong Kong | 2900 | 434/322 | Baseline data for IGT co- hort (N = 322) |
| Weiss 2005 | Conneticut, USA | 129 | 117 | Baseline data for IGT co- hort (N = 33) |
| Wheelock 2016 | Arizona, USA | Unclear | 5532 | Baseline data for the full cohort (N = 5532); pre- diabetic cohort = non- overweight (N = 37) + IGT group and over- weight + IGT group (N = 132); 5-11 years/12-19 years |
| Wong 2003 | Singapore | 3568 | 469/291 | Baseline data for IGT group (N = 291) |
| Yeboah 2011 | USA | 6814 | 6814/6753 | Baseline data for IFG co- hort (N = 940) |
| Zethelius 2004 | Uppsala, Sweden | 2322/1221/1010 | 840/667 | Baseline data for cohort converting to T2DM (N = 26) |

FG: fasting glucose; **FPG**: fasting plasma glucose; **HbA1c**: glycosylated haemoglobin A1c; **HbA1c**_{5.7/6.0}: HbA1c threshold 5.7% or 6.0% (usually reflecting 5.7% to 6.4% and 6.0% to 6.4%, respectively); **HbA1c/IFG**: both HbA1c and IFG; **i**-: isolated; **IFG**_{5.6/6.1}: impaired fasting glucose (threshold 5.6 mmol/L or 6.1 mmol/L); **IGT**: impaired glucose tolerance; **IFG/IGT**: both IFG and IGT; **PG**: postload glucose; **T2DM**: type 2 diabetes mellitus

| Study ID | Sex, % women | Age (SD), years | 'Eth- nicity', % white | 'Ethnic- ity', % Arabian/ Asian/ (Pima) In- dians | 'Ethnic- ity', % Hispanic | 'Ethnic- ity', % Black | Family history of diabetes, % | BMI (SD) , kg/m ² | Notes |
|-------------------------|----------------------|----------------------|------------------------------|---|------------------------------------|---------------------------------|--|--|--|
| Admiraal 2014 | 59 57 68 51 | 45 44 44 47 | 39 | 20 | - | 42 | 55 77 59 38 | 26.4 25.7 27.4 25.6 | Total cohort South- Asian Suri- namese African Suri- namese "Ethnic Dutch" (the Nether- lands) |
| Aek- plakorn 2006 | 19 | 43.6 (5.0) | - | 100 | - | - | 53 | 24.8 (3.2) | - |
| Ammari 1998 | - | 63% > 40 | - | 100 | - | - | 99 | - | - |
| Anjana 2015 | 61 | 47 (13.1) | - | 100 | - | - | 47 | 25.8 (4.3) | - |
| Bae 2011 | 25 | 44.7 (5.4) | - | 100 | - | - | - | 23.8 (2.8) | - |
| Baena- Diez 2011 | 52 | 61.2 (11. 8) | - | - | 100 | - | 26 | - | - |
| Bai 1999 | 35 | Mainly 40- 60+ | - | 100 | - | - | - | - | - |
| Bergman 2016 | 38 | 50.5 (8.3) | 42 | 29 | - | 47 | - | Men: 26.5 (3.8) Women: 26.8 (5.2) | - |
| Bonora 2011 | - | - | 100 | - | - | - | - | - | - |

Appendix 9. Baseline characteristics (II)

| Cederberg 2010 | - | - | 100 | - | - | - | - | Men: 27.6 (3.5) Women: 27.9 (4.5) | - |
|-----------------------------------|-------|-------------------------|----------------------------------|-----|-----|---|-------|--|---|
| Chamnan 2011 | 54 | 62.4 (8.2) | 100 | - | - | - | 14 | 26.6 (4.0) | - |
| Charles 1997 | 0 | 48.8 (1.8) | 100 | - | - | - | - | 27 (4) | - |
| Chen 2003 | 49 | 59.6 | - | 100 | - | - | 21 | 25.7 (3.1) | - |
| Chen 2017 | 54-58 | 40-67 | - | 100 | - | - | 9-37 | 23.8-24.8 | - |
| Coron- ado- Malagon 2009 | 10 | 47.9 (8.6) | - | - | 100 | - | - | 26.8 (3.0) | - |
| Cugati 2007 | 57 | 67.4 | 100 | - | - | - | 19 | 26 | - |
| De Abreu 2015 | 100 | 53.8 (IQR 44.0-64.4) | Mostly white Aus- tralians | - | - | - | - | 27.7 (IQR 24.3-31.4) | - |
| Den Biggelaar 2016 | 39 | 60.8 (IQR 55.3-64.9) | 100 | - | - | - | - | 28.0 (IQR 26.5-31.2) | - |
| Der- akhshan 2016 | 56 | 42.8 (11. 7) | - | 100 | - | - | - | 26.9 (4.1) | - |
| Dowse 1991 | - | - | - | 100 | - | - | - | - | - |
| Ferran- nini 2009 | 52-70 | 47-50 | - | - | 100 | - | 27-45 | 29.1-30.5 | - |
| Filippatos 2016 | 35 | 46.4 (12. 4) | 100 | - | - | - | 22 | 27.4 (4.7) | - |
| Forouhi 2007 | 44 | 55.5 (7.9) | 100 | - | - | - | - | 27.8 (4.6) | - |

| Garcia 2016 | - | 69.8 (6.9) | - | - | 49 | - | - | 31.1 (5.6) | - |
|-------------------------------|----------------|--|-----|-------------------|-----|----|----------------|---|--|
| Gautier 2010 | 31 | 30-64 | 100 | - | - | - | - | - | - |
| Gomez- Arbelaez 2015 | 70 | 58 (12) | - | - | 100 | - | - | 27.4 (4.6) | - |
| Guerrero- Romero 2006 | - | 38 | - | - | 100 | - | - | 32.9 (5.6) | - |
| Han 2017 | 28 60 33 | 50.4 (8.3) 53.1 (8.9) 52.4 (8.7) | - | 100 100 100 | - | - | 15 12 15 | 25.5 (3.4) 24.9 (3.2) 25.4 (3.2) | i-IFG _{5.6} i-IGT IFG/IGT |
| Hanley 2005 | 60 | 56.2 (7.9) | 38 | - | 36 | 26 | - | - | - |
| Heianza 2012 | 25 | 49.9 (8.7) | - | 100 | - | - | - | 22.8 (2.8) | - |
| Inoue 1996 | - | - | - | 100 | - | - | - | 23.2 | - |
| Janghor- bani 2015 | - | 44.4 42.9 44.1 | - | 100 | - | - | 100 | 29.2 29.0 30.0 | i-IFG i-IGT IFG/IGT |
| Jaru- ratanasiriku 2016 | 37 | 12.4 (2.3) | - | 100 | - | - | - | 35.3 (5.8) BMI SDS: 3.66 (0. 86) | - |
| Jeong 2010 | - | 61 (9) | - | 100 | - | - | 7 | 24.6 (3.2) | - |
| Jiamjaras- rangsi 2008a | 67 | 49.5 (12) | - | 100 | - | - | 15 | 26.9 (0.6) | - |
| Kim 2005 | 15 | 50.7 (7.2) | - | 100 | - | - | 9 | 24.6 (2.2) | - |
| Kim 2008 | 7 5 | 41 43 | - | 100 | - | - | 9 8 | 24 25 | IFG _{5.6} IFG _{6.1} |

| Kim 2014 | 49 57 48 56 | 60.2 (11. 3) 63.0 (11. 0) 59.1 (10. 1) 59.3 (10. 1) | - | 100 | - | - | 29 14 22 16 | 24.7 (3.0) 23.2 (3.5) 25.1 (3.3) 24.9 (4.7) | i-IFG i-IGT IFG/IGT i-HbA1c |
|------------------|----------------------|--|-----------------|-----|-----|---|----------------------|--|--|
| Kim 2016a | 24 47 | 49.5 51.2 | - | 100 | - | - | 22 22 | 24.4 23.9 | IFG HbA1c |
| Kleber 2010 | 51 | 13.1 (2.1) | 100 | - | - | - | - | 31.8 (6.3) BMI SDS: 2.56 (0. 62) | - |
| Kleber 2011 | 53 | 13.5 (2.1) | 100 | - | - | - | - | 31.7 (6.1) | - |
| Ko 1999 | 88 | 22-26 | - | 100 | - | - | - | - | - |
| Ko 2001 | 84 | 37.4 (9.3) | - | 100 | - | - | 38 | 25.9 (4.0) | - |
| Larsson 2000 | 100 | 66 (2.3) | 100 | - | - | - | - | 24.6 26.2 26.7 | i-IGT i-IFG IFG/IGT (age at fol- low-up) |
| Latifi 2016 | 38 | 46.6 (12. 5) | - | 100 | - | - | 80 | - | - |
| Lecomte 2007 | 0 | 44.5 (7.5) | 100 | - | - | - | 3 | 26.4 (3.6) | - |
| Lee 2016 | 33 | 46.1 (8.5) | - | 100 | - | - | 24 | 24.8 (3.1) | - |
| Leiva 2014 | 57 | 25-80 | - | - | 100 | - | - | 33.1 (4.3) | - |
| Levitzky 2008 | 53 | Women: 48 Men: 49 | Mainly white | - | - | - | - | Men: 27.3 (3.9) Women: 25.6 (5.4) | - |
| Li 2003 | 57 36 53 | 56.1 48.4 58.9 | - | 100 | - | - | - | 24.8 23.8 25.5 | i-IGT i-IFG IFG/IGT |

| Ligthart 2016 | 51 | 66.6 (9.4) | 92 | - | - | - | - | 27.9 (4.2) | - |
|------------------|----------------|-------------------------|----------------|-----|----|---|--------------------|--|---|
| Lipska 2013 | 33 60 47 | 76.6 76.7 76.6 | 82 36 60 | - | - | - | - | 27.9 27.9 29.0 | i-IFG i-HbA1c IFG + HbA1c |
| Liu 2008 | 57 | Men: 52 Women: 50 | - | 100 | - | - | Men: 6 Women: 8 | - | - |
| Liu 2014 | 48 | 68.6 (6.7) | - | 100 | - | - | - | 23.5 (3.0) | - |
| Liu 2016 | - | Men: 70 Women: 69 | - | 100 | - | - | - | - | - |
| Liu 2017 | 50 | 50.9 (9.7) | - | 100 | - | - | - | 24.2 (3.6) | - |
| Lorenzo 2003 | 61 | 47.7 (0.8) | 19 | - | 81 | - | 46 | 31.3 | - |
| Lyssenko 2005 | 50 | 52 (11) | 100 | - | - | - | 100 | - | - |
| Magliano 2008 | 49 | 55.8 (12. 0) | 85 | - | - | - | 31 | Men: 29.3 (0.4) Women: 29.7 (0.6) | - |
| Man 2017 | 57 | 54.4 (9.7) | - | 100 | - | - | 39 | 28.5 (5.3) | - |
| Marshall 1994 | 75 | 58.6 | 40 | - | 60 | - | 53 | 29.2 | - |
| McNeely 2003 | 52 41 | 58.9 57.5 | | 100 | - | - | 60 62 | 24.9 25.1 | 5-6 years follow-up 10 years follow-up |
| Meigs 2003 | 28 | 61.8 (14) | 95 | - | - | - | 29 | ≥ 25: 60% | - |
| Mohan 2008 | - | 43 (14) | - | 100 | - | - | 28 | 24.4 (4.4) | - |
| Motala 2003 | 60 | 36.4 (13. 9) | - | 100 | - | - | 45 | 22.6 (6.0) | - |

| Motta 2010 | - | 65-84 | 100 | - | - | - | - | - | - |
|--------------------------------|----------|-----------------------------|-----|-----|---|---|----|--|--|
| Mykkänen 1993 | 57 | 68.6 | 100 | - | - | - | 29 | 29 | - |
| Nakagami 2016 | 27 | 53 (7) | - | 100 | - | - | 19 | 24.6 (3.5) | - |
| Nakanishi 2004 | 0 | 49 (5.8) | - | 100 | - | - | 16 | 24.6 (3.0) | - |
| Noda 2010 | 63 | Men: 62.4 Women: 61.5 | - | 100 | - | - | - | Men: 24.1 (3.0) Women: 24.2 (3.2) | - |
| Park 2006 | 0 | 36.4 (3.9) | - | 100 | - | - | - | 24.8 (3.0) | - |
| Peterson 2017 | 48 | 61.4 (0.8) | 100 | - | - | - | - | 26.9 (5.4) | - |
| Qian 2012 | - | 60 (13) | - | 100 | - | - | - | 24.9 (3.7) | - |
| Rajala 2000 | 58 | - | 100 | - | - | - | - | - | - |
| Ra- machan- dran 1986 | 31 | 48 | - | 100 | - | - | 49 | 25.2 | - |
| Ras- mussen 2008 | 43 56 | 59.9 61.2 | 100 | - | - | - | - | 29.1 29.6 | IFG IGT |
| Rath- mann 2009 | 49 | 63.2 (5.4) | 100 | - | - | - | 23 | 28.1 (4.0) | - |
| Rijkeli- jkhuizen 2007 | 46 53 | 62.5 61.5 | 100 | - | - | - | - | 27.6 27.0 | IFG _{6.1} IFG _{5.6} |
| Sadeghi 2015 | 59 | 51.3 (9.8) | - | 100 | - | - | 20 | 29.4 (4.5) | - |

| Sasaki 1982 | 54 | 57.4 | - | 100 | - | - | - | - | - |
|--------------------------|----------|-----------------------------|-----|-----|---|----|-------------------------|--|--|
| Sato 2009 | 0 | 48.6 (4.2) | - | 100 | - | - | 20 | 24.7 (3.3) | - |
| Schranz 1989 | 56 | Women: 59.8 Men: 57.7 | 100 | - | - | - | - | - | - |
| Sharifi 2013 | 63 | 40 (14) | - | 100 | - | - | - | 27.5 (4) | - |
| Shin 1997 | 34 | 59.6 | - | 100 | - | - | 6 | 24.5 | - |
| Söderberg 2004 | 56 | 41.2 | - | 70 | - | 30 | - | 23.9 | - |
| Song 2015 | 52 | 56-57 | - | 100 | - | - | Men: 10 Women: 22 | Men: 25.2 (2.7) Women: 25.8 (3.4) | - |
| Song 2016a | 63 | 57.2 (10. 0) | - | 100 | - | - | - | - | - |
| Soriguer 2008 | 65 | 45.0 (13. 4) | 100 | - | - | - | 58 | 28.3 (5.2) | - |
| Stengard 1992 | 0 | 70.8 (4.8) | 100 | - | - | - | - | 26.1 (4.2) | - |
| Toshihiro 2008 | 0 | 50.5 (5.8) | - | 100 | - | - | - | 24.9 (3.3) | - |
| Vaccaro 1999 | 23 | 44.1 (4.0) | 100 | - | - | - | - | 26.9 (4.4) | - |
| Valdes 2008 | - | 54.8 56.7 | 100 | - | - | - | - | 28.2 29.8 | IFG _{5.6} IFG _{6.1} |
| Vijayaku- mar 2017 | 97 79 | 29.9 14 | - | 100 | - | - | - | 39.1 32.0 | Adults Children |
| Viswanatha 2007 | 39 | 42.4 (9.8) | - | 100 | - | - | - | - | - |
| Wang 2007 | 46 | 47.9 (10. 7) | - | 100 | - | - | - | 25.2 (3.5) | - |

| Wang 2011 | - | - | - | 100 | - | - | - | - | - |
|-------------------|----|------------|-----|-----|----|----|----|--|---|
| Warren 2017 | 48 | 57.6 (5.7) | - | - | - | 25 | 25 | 28.9 (5.2) | Data for cohort 1 (IFG _{5.6}) |
| Wat 2001 | 57 | 51 | | 100 | - | - | - | 25.6 | - |
| Weiss 2005 | 73 | 12.5 (2.7) | 45 | 39 | 12 | - | - | 36.6 (8.7) BMI z score: 2.42 (0.41) | - |
| Wheelock 2016 | 53 | 11.4 (3.6) | 100 | 100 | - | - | - | Percentile: 87.6 | - |
| Wong 2003 | 53 | 43.8 | - | 100 | - | - | 28 | 25.2 | - |
| Yeboah 2011 | 44 | 64.2 (9.8) | 31 | 15 | 25 | 30 | - | 30.1 (5.7) | - |
| Zethelius 2004 | 0 | 77 | 100 | - | - | - | - | 26.7 (3.2) | - |

BMI: body mass index; **FG**: fasting glucose; **FPG**: fasting plasma glucose; **i-HbA1c**: (isolated) glycosylated haemoglobin A1c; **HbA1c**_{5.7/6.0}: HbA1c threshold 5.7% or 6.0% (usually reflecting 5.7% to 6.4% and 6.0% to 6.4%, respectively); **HbA1c/IFG**: both HbA1c and IFG; **i**-: isolated; **IFG**_{5.6/6.1}: impaired fasting glucose (threshold 5.6 mmol/L or 6.1 mmol/L); **IGT**: impaired glucose tolerance; **IFG/IGT**: both IFG and IGT; **IQR**: interquartile range; **SD**: standard deviation; **SDS**: standard deviation score

Appendix 10. Baseline characteristics (III)

| Study ID | ••• | Mean (SD)/ median (IQR) /range di- astolic BP (SD), mmHg | Smoking: cur- rent and/ or past, % | Medica- tions, % | Comor- bidities, % | Mean (SD)/ median (IQR)/ range FPG, mmol/L | Mean (SD)/ median (IQR) /range 2- h glucose, mmol/L | - | Notes |
|------------------|-----|---|---|---------------------|--------------------------|--|---|---|-----------------|
| Admiraal 2014 | - | - | 38 26 41 | - | Hyperten- sion: 26 | 5.2 5.3 5.2 | - | - | Total cohort |

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| | | | 41 | | 26 32 19 | 5.3 | | | South- Asian Suri- namese African Suri- namese "Ethnic Dutch" |
|-------------------------|---------------------------|-------------------------|-------------------------|---|---|----------------------------|----------------------------|-----------|--|
| Aek- plakorn 2006 | - | - | 42 | - | Hyperten- sion: 33 | - | - | - | - |
| Ammari 1998 | - | - | - | - | Hyperten- sion: 47 | - | - | - | - |
| Anjana 2015 | 129 (21) | 78 (11) | 13 | - | - | 5.2 (0.6) | 8.7 (1.4) | 6.2 (0.7) | - |
| Bae 2011 | 113 (14) | 76 (10) | - | - | - | 5.3 (0.5) | - | 5.4 (0.3) | - |
| Baena- Diez 2011 | - | - | 33 | - | Hyperc- holestero- laemia: 38 Hyper- triglyceri- daemia: 15 Hyperten- sion: 55 | - | - | - | - |
| Bai 1999 | - | - | - | - | - | - | - | - | - |
| Bergman 2016 | 128 (16) | 84 (10) | 38 | - | - | 5.2 (0.5) | 8.6 (1.0) | - | - |
| Bonora 2011 | - | - | - | - | - | - | - | - | - |
| Cederberg 2010 | Men: 142 Women: 142 | Men: 80 Women: 79 | Men: 18 Women: 15 | - | - | Men: 5.0 Women: 5. 0 | Men: 6.8 Women: 7. 0 | - | - |
| Chamnan 2011 | 139 (17) | 84 (11) | 15 | BP lower- ing: 21 Corticos- teroids: 4 | - | - | - | - | - |
| Charles 1997 | - | - | - | - | - | 6.6 (0.8) | 9.3 (0.9) | - | - |

| Chen 2003 | - | - | 38 | - | Hyperten- sion: 46 | - | - | - | - |
|-----------------------------------|----------------------|-------------------|-------|---|--|----------------------|----------------------|----------------------|---|
| Chen 2017 | - | - | 12-24 | - | Hyperten- sion: 28- 55 | 5.1-6.1 | 5.9-9.2 | - | Range for i-IFG, i-IGT and IFG/IGT co- horts sepa- rated by < 40 years and > 60 years |
| Coron- ado- Malagon 2009 | - | - | - | - | - | 5.9 (0.3) | - | - | - |
| Cugati 2007 | 146 | 83 | - | - | - | 5 | - | - | - |
| De Abreu 2015 | 128 (IQR 114-140) | 79 (IQR 72-86) | 13 | - | Hyperten- sion: 43 | 5.3 (IQR 5.0-5.8) | - | - | - |
| Den Biggelaar 2016 | 141 (IQR 132-155) | 83 (IQR 78-92) | 18 | - | - | 6.0 (IQR 5.5-6.3) | 8.8 (IQR 7.8-9.9) | 5.8 (IQR 5.6-6.1) | - |
| Der- akhshan 2016 | - | - | 26 | - | - | - | - | - | - |
| Dowse 1991 | - | - | - | - | - | - | - | - | - |
| Ferran- nini 2009 | 118-128 | 71-78 | - | - | - | 4.9-6.4 | 6.7-9.5 | - | Range for i-IFG _{5.6} , i- IFG _{6.1} , i- IGT, IGT5.6 and IGT _{6.1} co- horts |
| Filippatos 2016 | 127 (17) | 82 (10) | 62 | - | Hyperten- sion: 36 Hyperc- holestero- | 5.9 (0.3) | - | - | - |

| | | | | | laemia: 54 | | | | |
|-------------------------------|----------------------------------|-------------------------------|----------------|--|---|-------------------------------------|-------------------------------------|-------------------------------------|--|
| Forouhi 2007 | 136 (16) | 82 (10) | 52 | - | - | - | - | - | - |
| Garcia 2016 | - | - | 58 | - | - | - | - | - | - |
| Gautier 2010 | - | - | - | - | - | - | - | - | - |
| Gomez- Arbelaez 2015 | - | - | - | - | - | 5.2 (0.7) | 6.0 (1.8) | 6.5 (1.3) | - |
| Guerrero- Romero 2006 | - | - | - | - | Dyslipi- daemia: 41 Hyperten- sion: 24 | 6.4 (0.6) | - | - | - |
| Han 2017 | 120 (17) 119 (18) 124 (18) | 78 (12) 76 (12) 80 (11) | 64 34 59 | - | Hyperten- sion: 28 27 36 | 5.9 (0.3) 4.8 (0.4) 5.9 (0.3) | 6.1 (1.2) 8.9 (0.9) 9.3 (0.9) | 5.5 (0.4) 5.5 (0.4) 5.7 (0.4) | i-IFG _{5.6} i-IGT IFG/IGT |
| Hanley 2005 | 132 (20) | 79 (10) | - | BP lower- ing: 38 Lipid low- ering: 7 | - | 5.9 (0.7) | 8.5 (1.7) | - | - |
| Heianza 2012 | 125 (16) | 76 (11) | - | - | - | 5.3 (0.5) | | 5.3 (0.3) | - |
| Inoue 1996 | 142 (9) | 73 (7) | - | - | - | - | - | - | - |
| Janghor- bani 2015 | 116-117 | 76-77 | - | - | Hyperten- sion: 20- 23 | 5.1-61 | 5.9-9.2 | 5.1-5.3 | Range for i-IFG, i-IGT and IFG/IGT cohorts |
| Jaru- ratanasiriku 2016 | 124 (15) | 77 (9) | - | - | - | - | - | - | - |
| Jeong 2010 | 139 (21) | 87 (12) | 43 | - | - | - | - | 5.7 (0.5) | - |

| Jiamjaras- rangsi 2008a | - | - | 4 | - | - | - | - | - | - |
|-------------------------------|---------------------------|---------|-------------------------|--|---|-----------------|-----------------|-----------|---|
| Kim 2005 | - | - | - | - | - | 6.4 (0.2) | - | - | - |
| Kim 2008 | 128/132 | 80/83 | - | - | - | 5.8/6.4 | - | - | - |
| Kim 2014 | 127-129 | 78 | 20-31 | - | - | - | - | - | Range for i-IFG, i- IGT, IFG/ IGT and i- HbA1c co- horts |
| Kim 2016a | 116-120 | 72-75 | 24-25 | - | - | 5.1-5.9 | - | 5.3-5.8 | Range for IFG and HbA1c co- horts |
| Kleber 2010 | 120 (16) | 73 (13) | - | - | - | 5.1 (1.1) | 8.5 | 5.6 (0.7) | - |
| Kleber 2011 | 120 (14) | 73 (12) | - | - | - | 4.8 (0.4) | 8.4 (0.6) | - | - |
| Ko 1999 | - | - | - | - | - | - | - | - | - |
| Ko 2001 | 125 (21) | 78 (10) | 2 | - | - | 6.5 (0.3) | 9.1 (2.1) | 6.2 (0.6) | - |
| Larsson 2000 | - | - | - | - | - | 4.7/5.5/5. 5 | 8.6/6.8/8. 7 | - | - |
| Latifi 2016 | - | - | - | - | Hyperten- sion: 40 | - | - | - | - |
| Lecomte 2007 | 135 (13) | 81 (10) | 23 | - | Hyperten- sion: 48 | 6.4 (0.2) | - | - | - |
| Lee 2016 | 125 (15) | 81 (11) | 20 | - | Hyperten- sion: 22 | - | - | 5.9 (0.2) | - |
| Leiva 2014 | 134 (16) | 77 (10) | - | - | - | - | - | - | - |
| Levitzky 2008 | Women: 122 Men: 127 | - | Women: 29 Men: 28 | Antihyper- tensives: Women: 14 Men: 16 | Hyperten- sion: Women: 26 Men: 35 | - | - | - | - |

| Li 2003 | 136-138 | 85-87 | - | - | - | 5.4-6.4 | 6.8-9.1 | - | Range for i-IFG, i-IGT and IFG/IGT cohorts |
|------------------|---------------------------|-------------------------|-------|---|--|----------------------------|------------|-----------|--|
| Ligthart 2016 | 145 (21) | 81 (12) | 50 | BP lower- ing: 33 Lipid low- ering: 18 | Stroke: 3 CHD: 8 Hyperten- sion: 64 | - | - | - | - |
| Lipska 2013 | 140-143 | - | 54-65 | - | - | 5.1-6.1 | - | 5.3-5.9 | Range for i-IFG, i-HbA1c and IFG/ HbA1c co- horts |
| Liu 2008 | Men: 126 Women: 124 | Men: 80 Women: 77 | - | - | - | Men: 5.3 Women: 5. 4 | - | - | - |
| Liu 2014 | 132 (16) | 82 (8) | - | - | - | 5.8 (0.8) | 9.2 (1.2) | - | - |
| Liu 2016 | - | - | - | - | - | - | - | - | - |
| Liu 2017 | 128 (21) | 81 (11) | 37 | - | - | 5.9 (0.4) | - | - | - |
| Lorenzo 2003 | 124 | 75 | - | - | - | 5.3 | 7.6 | - | - |
| Lyssenko 2005 | 140 | 85 (11) | - | - | - | 6.3 (IQR 5.8-6.6) | 8.3 (1.6) | 5.7 (0.4) | - |
| Magliano 2008 | - | - | 48 | - | - | 6 | 8 | 5.5 | - |
| Man 2017 | 145 (20) | 80 (12) | 13 | - | Hyperten- sion: 74 | - | - | - | - |
| Marshall 1994 | - | - | - | - | - | 6.1 | 9.5 | - | - |
| McNeely 2003 | 139 137 | 80 80 | - | - | - | 5.5 5.6 | 9.0 8.8 | - | 5-6 years follow-up 10 years follow-up |

| Meigs 2003 | - | - | - | - | - | - | - | - | - |
|--------------------------------|----------|---------|----|-------------------------------|--|----------------------------|-----------|----------------------------|--|
| Mohan 2008 | 127 (19) | 81 (11) | - | - | - | 4.5 (0.6) | - | - | - |
| Motala 2003 | 119 (19) | 78 (13) | - | - | - | 4.6 (1.8) | 6.2 (3.8) | - | - |
| Motta 2010 | - | - | - | - | - | - | - | - | - |
| Mykkänen 1993 | 159 | 84 | 1 | Antihyper- tensives: 24 | Hyperten- sion: 47 | 6.2 | 8.4 | - | - |
| Nakagami 2016 | 134 (18) | 82 (12) | 35 | - | - | 6.0 (0.6) | - | 6.0 (0.3) | - |
| Nakanishi 2004 | 133 (16) | 81 (11) | 53 | - | Dyslipi- daemia: 40 Protein- uria: 5 Hyperten- sion: 35 | 6.4 (0.2) | - | - | - |
| Noda 2010 | - | - | - | - | - | Men: 5.4 Women: 5. 2 | - | Men: 5.0 Women: 5. 1 | - |
| Park 2006 | - | - | - | - | - | 6.0 (0.3) | - | - | - |
| Peterson 2017 | - | 75 (11) | - | - | - | - | - | 5.5 (0.4) | - |
| Qian 2012 | 126 (21) | 81 (12) | - | - | - | 5.2 (0.7) | 6.1 (1.5) | - | - |
| Rajala 2000 | - | - | - | - | Hyperten- sion: 49 | - | - | - | - |
| Ra- machan- dran 1986 | - | - | - | - | - | - | - | - | - |
| Ras- mussen 2008 | 140-142 | - | - | - | - | - | - | - | Range for IFG and IGT cohorts |

| Rath- mann 2009 | 133 (19) | 80 (10) | 49 | Lipid low- ering: 11 | Hyperten- sion: 49 | 5.5 (0.5) | 6.3 (1.7) | 5.6 (0.4) | - |
|------------------------------|----------|---------|------|-------------------------|--|----------------------------|----------------------------|-----------|--|
| Rijkeli- jkhuizen 2007 | 139-145 | 84-85 | - | - | - | - | - | - | Range for IFG _{5.6} and IFG _{6.1} co- horts |
| Sadeghi 2015 | 127 (21) | 81 (11) | 14 | - | - | 5.7 (0.7) | 8.4 (1.5) | - | - |
| Sasaki 1982 | - | - | - | - | - | 5.6 (0.9) | 9.0 (0.9) | - | - |
| Sato 2009 | - | - | 91 | - | - | 6.0 (0.6) | - | 5.6 (0.6) | - |
| Schranz 1989 | - | - | - | - | - | Women: 7. 2 Men: 6.2 | Women: 10.8 Men: 9.7 | - | - |
| Sharifi 2013 | 130 (12) | 79 (8) | 5 | - | Hyper- triglyceri- daemia: 48 Hyperten- sion: 25 | - | - | - | - |
| Shin 1997 | 130 | 84 | - | - | - | 6.1 | 6.7 | - | - |
| Söderberg 2004 | 125 | 77 | 27 | - | - | 5.5 | 6.5 | - | - |
| Song 2015 | 123-127 | 76-80 | 2-27 | - | Dyslipi- daemia: 64-66 Hyperten- sion: 35- 44 | - | - | 5.7-5.8 | Ranges for male and female cohorts |
| Song 2016a | 134 (20) | 85 (12) | 23 | - | - | 6.0 (0.4) | 5.9 (1.6) | - | - |
| Soriguer 2008 | - | - | - | - | - | - | - | - | - |
| Stengard 1992 | 156 | 88 | - | - | Hyperten- sion: 53 | 5.4 (1.1) | 9.7 (0.8) | - | - |
| Toshihiro 2008 | 126 (12) | 81 (10) | 47 | - | - | 6.1 (0.6) | 8.8 (1.3) | - | - |

| Vaccaro 1999 | - | - | - | - | - | 4.2 (0.8) | 4.5 (1.7) | - | - |
|--------------------------|----------|---------|----|---|-----------------------|------------------|------------------|------------------|---|
| Valdes 2008 | 135-144 | 84-92 | - | - | - | 5.8-6.4 | 6.4-7.3 | 4.9-5.1 | Ranges for IFG _{5.6} and IFG _{6.1} co- horts |
| Vijayaku- mar 2017 | - | - | - | - | - | A: 5.4/C: 5.2 | A: 6.7/C: 6.5 | A: 5.8/C: 5.7 | - |
| Viswanatha 2007 | - | - | - | - | - | 6.1 (0.7) | 8.9 (1.0) | - | - |
| Wang 2007 | 124 (19) | 78 (11) | 28 | - | Hyperten- sion: 36 | 5.8 (0.9) | 7.4 (1.7) | - | - |
| Wang 2011 | - | - | - | - | - | - | - | - | - |
| Warren 2017 | - | - | 22 | - | Hyperten- sion: 38 | 6.0 (0.4) | - | 5.6 (0.4) | Data for cohort 1 (IFG _{5.6}) |
| Wat 2001 | 126 | 78 | - | - | - | 5.4 | 8.9 | - | - |
| Weiss 2005 | - | - | - | - | - | 5.2 | 8.9 | - | - |
| Wheelock 2016 | - | - | - | - | - | - | 5.4 (1.2) | - | - |
| Wong 2003 | 125 | 74 | 24 | - | - | 5.7 | 8.9 | - | - |
| Yeboah 2011 | 132 (21) | 74 (11) | 50 | BP lower- ing: 56 Lipid low- ering (statins): 17 | - | 6.0 (0.4) | - | - | - |
| Zethelius 2004 | - | - | - | - | - | 5.7 (0.7) | 7.9 (1.9) | - | - |

2-h: 2-h measurement after an OGTT; **BP**: blood pressure; **CHD**: coronary heart disease; **FG**: fasting glucose; **FPG**: fasting plasma glucose; **HbA1c**: glycosylated haemoglobin A1c; **HbA1c**_{5.7/6.0}: HbA1c threshold 5.7% or 6.0% (usually reflecting 5.7% to 6.4% and 6.0% to 6.4%, respectively); **HbA1c/IFG**: both HbA1c and IFG; **i**-: isolated; **IFG**_{5.6/6.1}: impaired fasting glucose (threshold 5.

6 mmol/L or 6.1 mmol/L);**IGT**: impaired glucose tolerance; **IFG/IGT**: both IFG and IGT; **IQR**: interquartile range; **OGTT**: oral glucose tolerance test; **SD**: standard deviation

Appendix 11. Cumulative incidence as the measurement for the development of T2DM

| Study ID (years of follow- up) | Diabetes cumulative incidence | | | | | | | | | | |
|---|-------------------------------|--|--------------------------------|--------------------------------|--------------------------------|-------------------|-------------------|-------------------|---|--|--|
| | NGT co- hort | IFG _{5.6} co- hort | i-IFG _{5.6} cohort | IFG _{6.1} co- hort | i-IFG _{6.1} cohort | IGT cohort | i-IGT co- hort | IFG/IGT cohort | HbA1c cohort | | |
| Admiraal 2014 (10) | Unclear/ 354 | Total cohort: 51/ 111 (45. 9%) Asian 13/31 (41. 9%) African 14/ 40 (35%) "Ethnic Dutch" 3/ 40 (7.5%) | - | - | - | - | - | - | - | | |
| Aek- plakorn 2006 (12) | Unclear/ 2444 | 65/223 (29.1%) | - | - | - | - | - | - | - | | |
| Ammari 1998 (2) | 10/144 (6. 9%) | - | - | - | - | 10/68 (14. 7%) | - | - | - | | |
| Anjana 2015 (9.1) | 209/1077 (19.4%) | - | 32/67 (47. 8%) | - | - | - | 86/163 (52.8%) | 58/69 (84. 1%) | - | | |
| Bae 2011 (4) | 228/7932 (2.9%) | - | - | - | - | - | - | - | HbA1c _{5.7} : 373/1791 (20.8%) HbA1c _{6.0} : 187/412 (45.4%) | | |

| Baena- Diez 2011 (10) | 0 (IFG co- hort) | - | - | 33/115 (28.7%) | - | - | - | - | - |
|--|--|-------------------|-------------------|--------------------------------|-------------------------------|-------------------------------|-------------------------------|-------------------------------|---|
| Bai 1999 (1) | 1/444 (0. 2%) | - | - | - | - | 14/252 (5. 6%) | - | - | - |
| Bergman 2016 (20) | 202/739 (27.3%) | - | - | - | - | 68/114 (59.6%) | - | - | - |
| Bonora 2011 (15) | 29/710 (4. 1%) | - | | 10 years: 18/55 (32. 7%) | - | - | 10 years: 8/53 (15. 1%) | 10 years: 9/19 (47. 4%) | HbA1c _{6.0} : 20/70 (28. 6%) |
| Cederberg 2010 (9.7) | 11/410 (2. 7%) | - | - | 15/40 (37. 8%) | 6.3% | 38/103 (37.1%) | 23.4% | - | HbA1c _{5.7} : 9/24 (37. 5%) |
| Chamnan 2011 (3) | 37/5365 (0.7%) | - | - | - | - | - | - | - | HbA1c _{6.0} : 26/370 (7%) |
| Charles 1997 (2) | 27/3671 (0.7%) | - | - | - | 3 years: 15/476 (3. 2%) | 2 years: 32/418 (7. 7%) | - | - | - |
| Chen 2003 (3) | 11/444 (2. 5%) | - | - | 15/156 (9. 6%) | - | - | - | - | - |
| Chen 2017 (3) | 60/644 (9. 3%) | - | 40/329 (12.2%) | - | - | - | 45/192 (23.4%) | 71/209 (34%) | - |
| Coronado- Malagon 2009 ^{<i>a</i>} (1, 2) | Year 1: 3/ 439 (0. 7%) Year 2: 3/ 439 (0. 6%) | - | - | - | - | - | - | - | - |
| Cugati 2007 (10) | 108/1512 (7.1%) | 69/229 (30%) | - | - | - | - | - | - | - |
| De Abreu 2015 (10) | 11/342 (3. 2%) | 21/187 (11.2%) | - | - | - | - | - | - | - |
| Den Biggelaar 2016 ^b (7) | 17/294 (5. 8%) | - | - | - | - | - | - | - | - |

| Der- akhshan 2016 ^c (11. 7) | 162/3611 (4.5%) | - | - | - | - | - | - | - | - |
|--|---------------------------------|---------------------|-------------------|--------------------|-----------------|---|--|--------------------|--|
| Dowse 1991 (6.2) | 14/215 (6. 5%) | - | - | - | - | 13/51 (25. 5%) | - | - | - |
| Ferran- nini 2009 (7) | 89/1594 (5.6%) | - | 11/65 (16. 9%) | - | 1/17 (5. 9%) | - | 31/179 (17.3%) 3 years: 44/188 (23.4%) | - | - |
| Filippatos 2016 (10) | 120/1206 (10.0%) | 71/279 (25.4%) | - | - | - | - | - | - | - |
| Forouhi 2007 (10) | 8/407 (2%) | 53/633 (8. 3%) | - | 34/257 (24.7%) | - | 4.4 years: 17/170 (10%) | - | - | - |
| Garcia 2016 (9) | 132/881 (15.0%) | 169/310 (54.5%) | - | - | - | - | - | - | - |
| Gautier 2010 (9) | 0 (IFG co- hort) | 142/979 (14.5%) | - | - | - | - | - | - | - |
| Gomez- Arbelaez 2015 ^d (2) | Unclear/ 586 | - | - | - | - | - | - | - | - |
| Guerrero- Romero 2006 (5) | 1/272 (0. 4%) | - | - | - | - | 20/67 (29. 9%) | - | - | - |
| Han 2017 (12) | 657/5633 (11.7%) | - | 81/199 (40.7%) | - | - | - | 624/1512 (41.3%) | 138/198 (69.7%) | 10 years: HbA1c _{5.7} : 881/2830 (31.1%) |
| Hanley 2005 (5.2) | 5 years: 47/603 (7. 8%) | - | - | - | - | 88/274 (32.1%) 5 years: 101/303 (33.3%) | - | - | - |
| Heianza 2012 (5) | 4.7 years: 34/4149 (0.8%) | 262/1680 (15.6%) | - | 155/380 (40.8%) | - | - | - | - | HbA1c _{5.7} : 184/822 |

| | | | | | | | | | (22.4%) HbA1c _{5.7} and IFG _{5.6} : 292/2092 (14%) HbA1c _{6.0} : 100/203 (49.3%) HbA1c _{6.0} and IFG _{5.6} : 271/1748 (15.5%) |
|---|---|-------------------|-------------------|---------------------|---|------------------|-------------------|-------------------|---|
| Inoue 1996 (2.5) | 1/22 (4. 5%) | - | - | - | - | 5/37 (13. 5%) | - | - | - |
| Janghor- bani 2015 (6.8) | 14/627 (2. 2%) | - | 23/230 (10%) | - | - | - | 26/150 (17.3%) | 78/214 (36.4%) | - |
| Jaru- ratanasirikul 2016 (3- 6) | 12/108 (11.1%) | - | - | - | - | - | 9/33 (27. 3%) | - | - |
| Jeong 2010 ^e (5) | 228/792 (28.8%) | - | - | - | - | - | - | - | - |
| Jiamjaras- rangsi 2008a (2. 6) | 15/2050 (0.7%) | 33/320 (10.3%) | - | - | - | - | - | - | - |
| Kim 2005 (5) | Unclear/ 2009 | - | - | 15/276 (5. 5%) | - | - | - | - | - |
| Kim 2008 (2) | 21/5382 (0.4%) | 22/1335 (1.6%) | - | 48/494 (9. 7%) | - | - | - | - | - |
| Kim 2014 (3.8) | 0 (cohort with inter- medi- ate hyper- glycaemia) | - | 24/158 (15.2%) | - | - | - | 12/65 (18. 5%) | 38/119 (31.9%) | i- HbA1c _{5.7} : 7/64 (10. 9%) |
| Kim 2016a (5. 2) | 43/10,763 (0.4%) | - | - | 357/1433 (24.9%) | - | - | - | - | HbA1c _{6.0} : 322/1103 |

| | | | | | | | | | (29.2%) IFG _{5.6} and HbA1c _{5.7} : 435/1951 (22.3%) |
|----------------------|---|-------------------|---|--|-------------------|-------------------------------|------------------|-------------------|--|
| Kleber 2010 (1) | 0 (IGT co- hort) | - | - | - | - | 1/79 (1. 3%) | - | - | - |
| Kleber 2011 (3.9) | 0 (IGT co- hort) | - | - | - | - | 3/119 (2. 5%) | - | - | - |
| Ko 1999 (1.4) | 0 (IGT co- hort) | - | - | - | - | 29/123 (23.6%) | - | - | - |
| Ko 2001 (1.7) | 13/264 (4. 9%) | - | - | 14/55 (25. 5%) | - | - | - | - | - |
| Larsson 2000 (10) | 5/127 (3. 9%) | - | - | - | 5/42 (11. 9%) | - | 8/66 (12. 1%) | 6/30 (20. 0%) | - |
| Latifi 2016 (5) | 25/394 (6. 3%) | 21/124 (16.9%) | - | - | - | - | - | - | - |
| Lecomte 2007 (5) | 0 (IFG co- hort) | - | - | 127/743 (17.1%) | - | - | - | - | - |
| Lee 2016 (3.7) | 0 (cohort with inter- medi- ate hyper- glycaemia) | - | - | - | - | - | - | - | HbA1c _{5.7} : 390/3497 (11.2%) |
| Leiva 2014 (6) | 0 (IFG co- hort) | - | - | 11/28 (39. 3%) | - | - | - | - | - |
| Levitzky 2008 (4) | 0 (IFG co- hort) | - | - | Women: 87/313 (27.8%) Men: 92/ 460 (20. 0%) | - | - | - | - | - |
| Li 2003 (5) | 38/435 (8. 7%) | - | - | - | 16/42 (38. 1%) | 2 years: 23/131 (17.6%) | 33/118 (28%) | 20/49 (40. 8%) | - |

(Continued)

| Ligthart 2016 (14. 7) | Unclear/ 7462 | - | - | 425/1382 (30.8%) | - | - | - | - | - |
|---|--|---|---|--|---|---|---|---|---|
| Lipska 2013 (7) | 38/1690 (2.2%) | 20/189 (10.6%) | - | 48/100 (48%) | - | - | - | - | i- HbA1c5.7: 44/207 (21.3%) IFG and HbA1c5.7: 81/169 (47.9%) |
| Liu 2008 (5) | 9/470 (1. 9%) | 18/169 (10.7%) | - | - | - | - | - | - | - |
| Liu 2014 ^f (3) | 153/1821 (8.4%) | - | - | - | - | - | - | - | - |
| Liu 2016 (10.9) | Unclear/ 1635 | - | - | - | - | - | - | - | - |
| Liu 2017 (7.8) | Unclear/ 15003 | - | - | - | - | - | - | - | - |
| Lorenzo 2003 (7- 8) | Unclear/ 1503 | - | - | 14/29 (48. 3%) | - | 88/202 (43.6%) | - | - | - |
| Lyssenko 2005 ^g (6) | 41/1429 (2.9%) | - | - | - | - | - | - | - | - |
| Magliano 2008 (5) | 58/4715 (1.2%) | - | - | 44/370 (11.9%) | - | 122/757 (16.1%) | - | - | - |
| Man 2017 (6) | 15/462 (3. 2%) | - | - | - | - | - | - | - | HbA1c _{5.7} : 112/675 (16.6%) |
| Marshall 1994 (1.9) | 0 (IGT co- hort) | - | - | - | - | 20/123 (16.3%) | - | - | - |
| McNeely 2003 (10) | 5-6 years: 5/277 (1. 8%) 10 years: 13/277 (4. 5%) | 5-6 years: 27/125 (21.6%) 10 years: 39/103 (37.9%) | - | 5-6 years: 7/30 (23. 3%) 10 years: 18/28 (64. 3%) | - | 5-6 years: 45/178 (25.3%) 10 years: 59/157 (37.6%) | - | - | - |

| Meigs 2003 (5, 10) | 6 (SD 5) years: 55/488 (11.3%) | - | - | - | 6 (SD 5) years: 6/20 (30. 0%) | - | 6 (SD 5) years: 81/218 (37.1%) | 6 (SD 5) years: 15/27 (55. 6%) | - |
|--------------------------|---|--|---|---|--|--|---|---|--|
| Mohan 2008 (8) | 64/476 (13.4%) | - | - | - | - | 15/37 (40. 5%) | - | - | - |
| Motala 2003 (10) | 36/482 (7. 5%) | - | - | - | - | 13/35 (37. 1%) 4 years: 16/72 (22. 2%) | - | - | - |
| Motta 2010 (3) | 52/2018 (2.6%) | - | - | 50/295 (16.9%) | - | - | - | - | - |
| Mykkänen 1993 (3.5) | 21/689 (3. 0%) | - | - | - | - | 48/203 (23.6%) | - | - | - |
| Nakagami 2016 (5) | 1528 | 77/467 (16.5%) | - | 50/134 (37.3) | - | - | - | - | HbA1c _{6.0} : 58/156 (37.2%) HbA1c _{5.7} : 87/583 (14.9%) |
| Nakanishi 2004 (7) | 51/5500 (0.9%) | - | - | 5/246 (2. 0%) | - | - | - | - | - |
| Noda 2010 (5) | Total: 30/ 1649 (1. 8%) Men: 13/540 (2. 4%) Women: 17/1109 (6.4%) | To- tal: 37/405 (9.1%) Men: 18/202 (8. 9%) Women: 19/203 (9. 4%) | - | To- tal: 58/153 (37.9%) Men: 25/79 (31. 6%) Women: 33/74 (44. 6%) | - | - | - | - | - |
| Park 2006 (4.1) | 116/4975 (2.3%) | 40/321 (12.5%) | - | - | - | - | - | - | - |
| Peterson 2017 (10) | 2/39 (5. 1%) | - | - | - | - | 6/29 (20. 7%) | - | - | - |
| Qian 2012 (5) | 59/843 (7. 0%) | - | - | - | 17/46 (37%) | - | 49/120 (41%) | 17/33 (51%) | - |

| Rajala 2000 (4.6) | 0 (IGT co- hort) | - | - | - | - | 32/171 (18.7%) 2.1 years: 14/183 (7. 7%) | - | - | - |
|---|---------------------------|--------------------|--------------------|--------------------|-----------------|--|------------------------------|----------------------------|---|
| Ra- machan- dran 1986 (5.1) | 0 IGT co- hort) | - | - | - | - | 39/107 (36.4%) | - | - | - |
| Rasmussen 2008 (3.5) (i- IFG _{5.6} : 2. 5, IGT: 2. 1) | 0 (IFG, IGT cohort) | - | 141/442 (32%) | - | - | 181/442 (41%) | 1 year: 35/296 (11.8%) | 1 year: 60/207 (29%) | - |
| Rath- mann 2009 (7) | 25/649 (3. 9%) | - | - | 12/71 (16. 9%) | - | - | 34/120 (28.3%) | 22/47 (46. 8%) | - |
| Rijkeli- jkhuizen 2007 (6.4) | 51/1125 (4.5%) | 101/488 (20.7%) | - | 62/149 (41.6%) | 35/106 (33%) | 36/111 (32.4%) 2 years: 45/158 (28.5%) | 27/80 (33. 8%) | 20/31 (64. 5%) | - |
| Sadeghi 2015 (7) | 141/2607 (5.4%) | - | 134/373 (35.9%) | - | - | - | 49/373 (13.1%) | 65/373 (17.4%) | - |
| <mark>Sasaki</mark> 1982 (7) | 7/161/4. 3%) | - | - | - | - | 5/13 (38. 5%) | - | - | - |
| Sato 2009 (4) | 118/4147 (2.9%) | - | - | 334/794 (42.1%) | - | - | - | - | HbA1c _{6.0} : 90/215 (41.9%) |
| Schranz 1989 (6) | 54/1251 (4.3%) | - | - | - | - | 23/75 (30. 7%) | - | - | - |
| Sharifi 2013 (7) | 0 (IFG co- hort) | 24/123 (19.5%) | - | - | - | - | - | - | - |
| Shin 1997 (2) | 47/1040 (4.5%) | - | - | - | - | 20/153 (13.1%) | - | - | - |

| Söderberg 2004 (11) | Unclear/ 2522 | - | - | 5 years: 32/148 (21.6%) | 153/402 (38%) | 575/1253 (45.9%) | 5 years: 103/489 (21.1%) | 5 years: 45/118 (38.1%) | - |
|--|--|---|---|-------------------------------|------------------|---|--------------------------------|---|--|
| Song 2015 (4) | 74/1758 (4.2%) | - | 68/321 (21.2%) Men: 30/ 154 (19. 5%) Women: 38/167 (22.8%) | - | - | - | - | - | - |
| Song 2016a (10.8) | 0 (cohort with inter- medi- ate hyper- glycaemia) | - | - | - | - | - | - | - | - |
| Soriguer 2008 (6) | 13/1806 (0.7%) | - | 23/56 (41. 1%) | - | - | 14/54 (25. 9%) | - | 14/28 (50%) | - |
| Stengard 1992 (5) | 6/216 (2. 8%) | - | - | - | - | 17/234 (7. 3%) | - | - | |
| Toshihiro 2008 (3.2) h | 0 (co- hort with IFG and/ or IGT) | - | - | - | - | - | - | - | - |
| Vaccaro 1999 (11. 5) | 36/500 (7. 2%) | - | 1/11 (9. 1%) | - | - | - | 13/40 (32. 5%) | 4/9 (44. 4%) | - |
| Valdes 2008 (6.3) | 16/510 (3. 1%) | 14/114 (12.3%) | 7/32 (21. 9%) | 19/52 (36. 5%) | - | 21/88 (23. 9%) | 9/68 (13. 2%) | 12/20 (60%) | - |
| Vijayaku- mar 2017 (adults: 4. 6, children: 5.2) | Adults: 58/ 1466 (3.9) Children: 26/1795 (1.4%) [estimated from figure 2] | Adults: 222/424 (52.4%) Children: 52/193 (26.9%) | - | - | - | Adults: 196/347 (56.5%) Children: 55/169 (32.5%) | - | IFG _{5.6} / IGT: Adults: 116/169 (68.7%) Children: 26/53 (49. 1%) | HbA1c _{5.7} : adults: 75/ 168 (44. 6%) HbA1c _{5.7} : chil- dren: 18/ 62 (29%) |
| Viswanatha 2007 (5) | Total: 154/ 465 33. 1%) | - | - | - | - | Total: 416/ 619 (67. | - | - | - |

| | M: 99/265 (37.4%) W: 55/200 (27.5%) | | | | | 2%) M: 251/ 391 (64. 2%) W: 165/ 228 (72. 4%) | | | |
|--|--|--|-----------------|-----------------|---|---|-------------------|--|---|
| Wang 2007 (5) | 51/358 (14.2%) | - | 53/261 (20%) | 28/112 (25%) | - | 126/141 (89.4%) | 31/95 (32. 6%) | IFG _{5.6} / IGT: 54/ 109 (49. 5%) IFG _{6.1} / IGT: 36/52 (69. 2%) | - |
| Wang 2011 (7.8) | 84/595 (14.1%) | Total: 345/947 (36.4%) Men: 137/ 418 (32. 8%) Women: 208/529 (39.3%) | - | - | - | Total: 233/491 (47.5%) Men: 75/ 154 (48. 7%): Women: 158/337 (46.9%) 4 years: Total 198/ 532 (37. 2%) | - | Total: 185/356 (52%%) Men: 66/ 125 (52. 8%) Women: 119/231 (51.5%) | HbA1c _{6.0} : 19/121 (15.7%) |
| Warren 2017 (co- hort 1: 22, cohort 2: 16) | 16 years: | - | - | - | - | - | - | - | - |
| Wat 2001 (2) | 4/333 (0. 1%) | - | - | - | - | 31/322 (9. 6%) | - | - | - |
| Weiss 2005 (1.7) | 8/84 (9. 5%) | - | - | - | - | - | 8/33 (24. 2%) | - | - |
| Wheelock 2016 (4.3) | Unclear/ 5363 | - | - | 5 years: 31% | - | Non-over- weight: 5 years: 9/ 37 (24%) 10 years: 11/37 (29. | - | 5 years: 41.2% | - |

| | | | | | | 7%) Over- weight: 5 years: 49/ 132 (37%) 10 years: 84/132 (63.6%) | | | |
|-----------------------------------|-------------------|--------------------|---|---|---|--|---|---|---|
| Wong 2003 (8) | 12/278 (4. 3%) | - | - | - | - | 102/291 (35.1%) | - | - | - |
| <mark>Yeboah</mark> 2011 (7.5) | Unclear/ 4973 | 273/940 (29.0%) | - | - | - | - | - | - | - |
| Zethelius 2004 (7) | Unclear/ 466 | - | - | - | - | Not reported/ 201 | - | - | - |

^a Development of T2DM from 'prediabetes' (not defined) at year 1: 11/217 (5.1%), at year 2: 16/217 (7.6%)

^bDevelopment of T2DM from 'prediabetes' (IFG_{6.1} and/or IGT): 46/122 (37.7%).

^cDevelopment of T2DM from IFG_{5.6} and/or IGT: 11.7 years150/523 (28.7%); 2.3 years: 121/911 (13.3%).

^d Development of T2DM from IFG_{5.6} or IGT or HbA1c_{5.7}: 20/186 (10.8%).

^eDevelopment of T2DM from IFG or IGT: not reported.

^f Development of T2DM from IFG or IGT: 78/450 (17.3%).

^gDevelopment of T2DM from IFG or IGT:86/686 (12.5%).

^hDevelopment of T2DM from IFG and/or IGT: 36/128 (28.1%).

FPG: fasting plasma glucose; **HbA1c**: glycosylated haemoglobin A1c; **HbA1c**_{5.7/6.0}: HbA1c threshold 5.7% or 6.0% (usually reflecting 5.7% to 6.4% and 6.0% to 6.4%, respectively); **HbA1c/IFG**: both HbA1c and IFG; **i**-: isolated; **IFG**_{5.6/6.1}: impaired fasting glucose (threshold 5.6 mmol/L) or 6.1 mmol/L); **IGT**: impaired glucose tolerance; **IFG/IGT**: both IFG and IGT; **NGT**: normal glucose tolerance; **PG**: postload glucose; **SD**: standard deviation; **T2DM**: type 2 diabetes mellitus

Appendix 12. Diabetes incidence (cases per 1000 person-years)

| Study ID | Rate (diabe | Rate (diabetes cases/1000 person-years (95% CI)) | | | | | | | | | |
|----------------|----------------------|--|--------------------------------|---|--------------------------------|----------------------|-------------------|-----------------------------|-------------------------------------|--|--|
| | Follow-up (years) | NGT co- hort | 'Predi- abetes' co- hort | | IFG _{5.6} co- hort | IGT cohort | IFG/IGT cohort | Elevated HbA1c cohort | Elevated HbA1c/ IFG cohort | | |
| Anjana 2015 | 9.1 | 22.2 (19. 4-25.4) | 78.9 (68. 0-90.9) | - | | 67.8 (54. 6-83.0) | | - | - | | |

| Bae 2011 | 4 | - | - | - | - | - | - | Per 100 person- years: HbA1c _{5.7} : 5.6 HbA1c _{6.0} : 14.0 | - |
|--------------------------|------|--|---------------------|------------------------------------|--|---------------------------------------|-----------------------------------|---|--|
| Bonora 2011 | 15 | 10 years: 4. 3 (2.7-5.9) | - | 10 years: 37.0 (20. 2-53.8) | - | 10 years: 17.0 (5.3-28.7) | 10 years: 49.2 (17. 9-80.5) | HbA1c _{6.0} : 25.8 | - |
| De Abreu 2015 | 10 | - | - | - | 18.1 (10. 7-28.2) | - | - | - | - |
| Der- akhshan 2016 | 11.7 | - | 30.3 | 6.5 years: 69.4 (56. 0-86.1) | 6.5 years: 39.5 (34. 4-45.4) | 6.5 years: 41.6 (36. 1-47.9) | - | - | - |
| Dowse 1991 | 6.2 | 10.5 | - | - | - | 40.4 | - | - | - |
| Forouhi 2007 | 10 | 2.4 (1.2-4. 8) 4 years: 2. 64 (1.23- 4.05) | - | 17.5 (12. 5-24.5) | 10.6 (8.1- 13.9) (IFG _{5.6} : FPG 5.6- 6.9) | 4 years: 22. 5 (20.4- 24.6) | - | - | - |
| Han 2017 | 12 | 12.3 | IFG or IGT: 58.0 | - | i-IFG _{5.6} : 51.3 | i-IGT: 53. 1 | 114.4 | 10 years: HbA1c _{5.7} : 43.2 | - |
| Heianza 2012 | 5 | 2.3 | - | 104 | 34.6 | - | - | HbA1c _{5.7} : 51.0 HbA1c _{6.0} : 129.2 | HbA1c _{5.7} and IFG _{5.6} : 30. 6 HbA1c _{6.0} and IFG _{5.6} : 34. 4 |
| Janghor- bani 2015 | 6.8 | 3.1 (1.5-4. 7) 2.3 years: 4.6 (1.28- 11.7) | - | - | 3-24.4) 2.3 years: i- | 0-37.7) 2.3 years: i- IGT: 99.7 | 57.9 (46. 1-71.7) | - | - |

| Jiamjaras- rangsi 2008a | 2.6 | - | - | - | 31.5 (11. 4-86.8) | - | - | - | - |
|-------------------------------|-------|--|---|--|----------------------|--|--|---|---|
| Latifi 2016 | 5 | 21.9 | - | - | 34.5 | - | - | - | - |
| Li 2003 | 5 | 18.8 | - | 93.7 | - | 60.7 | 117 | - | - |
| Ligthart 2016 | 14.7 | - | - | 43.0 (39. 2-47.2) | - | - | - | - | - |
| Liu 2008 | 5 | 9 | - | - | 22.5 | - | - | - | - |
| Magliano 2008 | 5 | 0.2 (0.2-0. 3) (incidence percent per years) | - | i- IFG _{6.1} : 2.6 (1.8-3.4) (incidence percent per years) | - | i-IGT: 3.5 (2.9-4.2) (incidence percent per years) | - | - | - |
| Meigs 2003 | 5, 10 | Per 100 person- years (an- nu- alised rate) : FPG \geq 7. 0: 0.64 (0. 32-1.13) 2-h PG \geq 11.1: 2.77 (2. 01-3.71) | - | - | - | - | Per 100 person- years (an- nualised rate): IFG or IGT FPG \geq 7. 0: 0.98 (0. 65-1.41) 2-h PG \geq 11.1: 4.61 (3. 77-5.56) | - | - |
| Mohan 2008 | 8 | 17.5 | - | - | - | 64.8 | - | - | - |
| Nakagami 2016 | 5 | - | - | 1 | - | - | - | - | - |
| Nakanishi 2004 | 7 | 1.5 | - | 3.3 | - | - | - | - | - |
| Park 2006 | 4.1 | 5.7 | - | - | 31.3 | - | - | - | - |

| Rajala 2000 | 4.6 | - | - | - | - | 41 (28-57) | - | - | - |
|------------------------------|--|--|---|--|--|--|---|---|---|
| Ras- mussen 2008 | 3.5 (i- IFG _{5.6} : 2.5 , IGT: 2.1) | - | - | - | i-IFG _{5.6} : 11.8 (9.9- 13.8) per 100 person- years | | 27 (22.5- 31.7) per 100 person- years | - | - |
| Rath- mann 2009 | 7 | - | - | i-IFG _{6.1} : 24.2 (12. 5-42.3) | - | i-IGT: 42.0 (29. 0-58.7) | 77.9 (48. 8-117.9) | - | - |
| Rijkeli- jkhuizen 2007 | 6.4 | 7 | - | 66.5 (49. 9-83.0) | 32.7 (26. 3-39.1) | i-IGT: 57. 9 | 112.2 | - | - |
| Sadeghi 2015 | 7 | Total: 14.1 (12.5-15. 9) Men: 12.8 (10.7-15. 3) Women: 15.5 (13. 1-18.3) | - | - | Total: 48.4 (35.0-66. 7) Men: 46.4 (28.9-74. 7) Women: 50.1 (32. 3-77.7) | (25.7-66. 6) Women: | 6 (103.7- 182.5) | - | - |
| Söderberg 2004 | 11 | - | - | 87-92: Men: 54.1 (48.0-60. 1) Women: 35.1 (30. 3-40.0) 92-98: Men: 60.5 (54.1-67. 0) Women: 74-7 (67. 8-81.8) | - | 87-92: Men: 60.7 (54.3-67. 1) Women: 47.9 (42. 2-53.6) 92-98: Men: 119. 6 (110.6- 128.6) Women: 81.0 (73. 6-88.4) | - | - | - |
| Soriguer 2008 | 6 | 7.2 (4.2- 12.4) | - | - | 38.1 (25. 3-57.3) | 31.1 (18. 4-52.5) | 66.0 (39. 1-111.5) | - | - |

| Valdes 2008 | 6.3 | 3.8 (2.1-6. 8) for i-IGT and IFG/ IGT: 5.0 (2.8-8) | - | 58.0 (37- 90.9) | 19.5 (11. 5-32.9) | 37.9 (24. 7-58.1) i-IGT: 21 (10.9-40. 4) | 95.2 (54. 1-167.7) | - | - |
|--------------------------|---------------------------------|---|---|--------------------|---|--|-----------------------|---|---|
| Vijayaku- mar 2017 | Adults: 4.6 Children: 5.2 | - | - | - | Boys: 22 Men: 70 Girls: 55 Women: 101 | Boys: 38 Men: 94 Girls: 60 Women: 118 | - | Boys: 52 Men: 100 Girls: 100 Women: 118 | - |
| Wang 2011 | 7.8 | 21.1 | - | - | | Total: 95.8 Men: 98.1 Women: 94.8 | | - | - |

CI: confidence interval; FPG: fasting plasma glucose; HbA1c: glycosylated haemoglobin A1c; HbA1c_{5.7/6.0}: HbA1c threshold 5. 7% or 6.0% (usually reflecting 5.7% to 6.4% and 6.0% to 6.4%, respectively); HbA1c/IFG: both HbA1c and IFG; i-: isolated; IFG_{5.6/6.1}: impaired fasting glucose (threshold 5.6 mmol/L or 6.1 mmol/L); IGT: impaired glucose tolerance; IFG/IGT: both IFG and IGT;NGT: normal glucose tolerance; T2DM: type 2 diabetes mellitus

| Appendix 13. | T2DM cases and | person-time (fo | or calculation | incidence rate ratios) |
|--------------|----------------|-----------------|----------------|------------------------|
|--------------|----------------|-----------------|----------------|------------------------|

| Study ID | Persons (cases) with | n diabetes with/witho | ut IH at baseline | | |
|-----------------|----------------------|--|---|------------------------------------|--|
| | Follow-up (years) | Cases in IH group | Person-years for IH group | Cases in normogly- caemic group | Person- years for normo- glycaemic group |
| Anjana 2015 | 9.1 | i-IFG _{5.6} : 32 i-IGT: 86 IFG/IGT: 58 | i-IFG _{5.6} : 525 i-IGT: 1269 IFG _{5.6} /IGT: 434 | 209 | 9398 |
| De Abreu 2015 | 10 | IFG _{5.6} : 21 | IFG _{5.6} : 1768 | 11 | - |
| Bae 2011 | 4 | HbA1c _{5.7} : 373 HbA1c _{6.0} : 187 | HbA1c _{5.7} : 6594 HbA1c _{6.0} : 1338 | - | - |
| Bonora 2011 | 10 | IFG _{6.1} : 18 IGT: 8 IFG/IGT: 9 | IFG _{6.1} : 486 IGT: 471 IFG/IGT: 183 | 29 | 6704 |
| Derakhshan 2016 | 11.7 | IFG _{5.6} : 150 | IFG _{5.6} : 4950 | 162 | 39,901 |

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| Dowse 1991 | 6.2 | IGT: 13 | IGT: 322 | 14 | 1339 |
|-------------------------|-------|--|--|-----------------------|-----------------------------|
| Forouhi 2007 | 10 | IFG _{6.1} : 34 IFG _{5.6} : 53 4.44 years: IGT: 17 | IFG _{6.1} : 1943 IFG _{5.6} : 5000 4.44 years: IGT: 756 | 8 4.44 years: 9 | 3333 4.44 years: 3409 |
| Guerrero-Romero 2006 | 5 | IGT: 20 | IGT: 343 | 1 | 1388 |
| Han 2017 | 12 | i-IFG _{5.6} : 81 i-IGT: 624 IFG/IGT: 138 | i-IFG _{5.6} : 1579 i-IGT: 11,744 IFG/IGT: 1206 | 657 | 53,461 |
| Heianza 2012 | 5 | IFG _{5.6} : 108 HbA1c _{5.7} : 30 HbA1c _{5.7} /IFG _{5.6} : 154 | IFG _{5.6} : 5920 HbA1c _{5.7} : 1965 HbA1c _{5.7} /IFG _{5.6} : 1641 | 46 | 19,961 |
| Janghorbani 2015 | 6.8 | i-IFG _{5.6} : 23 i-IGT: 26 IFG/IGT: 214 | i-IFG _{5.6} : 1409 i-IGT: 1005 IFG/IGT: 1347 | 14 | 4578 |
| Li 2003 | 5 | i-IFG _{6.1} : 16 i-IGT: 33 IFG/IGT: 20 | i-IFG _{6.1} : 171 i-IGT: 544 IFG/IGT: 179 | 38 | 2026 |
| Ligthart 2016 | 14.7 | IFG _{6.1} : 425 | iFG _{6.1} : 9884 | - | - |
| Meigs 2003 | 5, 10 | IFG or IGT T2DM measured by: FPG \geq 7.0: 26 2-h PG \geq 11.1: 101 | IFG or IGT T2DM measured by: FPG ≥ 7.0: 2647 2-h PG ≥ 11.1: 2192 | 28 | 1539 |
| Mohan 2008 | 8 | IGT: 15 | IGT: 247 | 64 | 3665 |
| Nakanishi 2004 | 7 | IFG _{6.1} : 5 | IFG _{6.1} : 1506 | 51 | 34,308 |
| Park 2006 | 4.1 | IFG _{5.6} : 40 | IFG _{5.6} : 1278 | 116 | 20,298 |
| Rijkelijkhuizen 2007 | 6.4 | i-IFG _{6.1} : 35 i-IGT: 27 IFG/IGT: 20 | i-IFG _{6.1} : 681 i-IGT: 466 IFG/IGT: 178 | 51 | 7286 |
| Soriguer 2008 | 6 | IFG _{5.6} : 23 IGT: 14 IFG/IGT: 14 | IFG _{5.6} : 604 IGT: 450 IFG/IGT: 212 | 13 | 1806 |

| Valdes 2008 | 6.3 | IFG _{5.6} : 14 IFG _{6.1} : 19 i-IGT: 9 IFG/IGT: 12 | IFG _{5.6} : 718 IFG _{6.1} :328 i-IGT: 429 IFG/IGT: 126 | 11 (16 for i-IGT and IFG/IGT) | 2923 (3200 for i-IGT and IFG/IGT) |
|-------------|-----|---|---|-------------------------------------|---|
| Wang 2011 | 7.8 | IFG _{5.6} : 137 IGT: 75 IFG/IGT: 66 | IFG _{5.6} : 2374 IGT: 765 IFG/IGT: 605 | 34 | 1613 |

FPG: fasting plasma glucose; **HbA1c**: glycosylated haemoglobin A1c; **HbA1c**_{5.7/6.0}: HbA1c threshold 5.7% or 6.0% (usually reflecting 5.7% to 6.4% and 6.0% to 6.4%, respectively); **HbA1c/IFG**: both HbA1c and IFG; **i**-: isolated; **IFG**_{5.6/6.1}: impaired fasting glucose (threshold 5.6 mmol/L or 6.1 mmol/L); **IGT**: impaired glucose tolerance; **IFG/IGT**: both IFG and IGT; **IH**: intermediate hyperglycaemia;**T2DM**: type 2 diabetes mellitus

Appendix 14. Odds ratios and hazard ratios as the effect measures for the development of T2DM

| Study ID | Adjusted [u baseline | Adjusted [unadjusted] ratios (95% CI) for the development of diabetes comparing IH with normoglycaemia at baseline | | | | | | | | | | |
|------------------|-------------------------|--|--|-----|--------------------|---------|-------|---------------|------------|--|--|--|
| | Follow-up (years) | IFG _{6.1} | IFG _{5.6} | IGT | 'Predia- betes' | IFG/IGT | HbA1c | HbA1c/ IFG | Ratio | | | |
| Admiraal 2014 | 10 | - | Total cohort: 6.1 (3.1- 12.1) [5.7 (3.1- 10.5)] South- Asian Suri- namese: 11.1 (3.0- 40.8) [9.9 (2.9- 34.3)] African Suri- namese: 5.1 (2.0- 13.3) [6.2 (2.6- 14.9)] "Ethnic Dutch": | - | - | - | - | - | Odds ratio | | | |

| | | | 2.2 (0.5- 10.2) [2.1 (0.5- 9.3)] | | | | | | |
|-------------------------|-----|--|---|--|---|--|--|---|--|
| Aek- plakorn 2006 | 12 | - | [2.41 (1. 78-3.28)] | [4.36 (3. 41-5.57)] | - | - | - | - | Odds ratio |
| Bae 2011 | 4 | - | - | - | - | - | HbA1c _{5.7} : 6.5 (3.7- 10.2) HbA1c _{6.0} : 41.3 (24. 7-69.2) [com- pared with HbA1c < 5.0] | - | Hazard ra- tio |
| Bergman 2016 | 24 | IFG _{6.1} : 3. | 20 years: i- IFG _{5.6} : 1. 11 (0.76- 1.61) | 74-12.33) | - | $IFG_{5.6} + IGT: 2.$ 79 (1.56- 5.00) $IFG_{6.1} + IGT: 3.$ 85 (1.73- 8.54) | - | - | Odds ratio |
| Bonora 2011 | 15 | 5.83 (3. 23-10.54) 10 years: 5.7 (2.8- 11.4) [3.9 (1.56- 9.3)] | | 10 years: [3.9 (1.6- 9.3)] | - | 10 years: [20.5 (7.6- 55.3)] | HbA1c _{6.0} : 9.74 (4. 21-22.56) | - | Hazard ra- tio, odds ratio (10 years) |
| Cederberg 2010 | 9.7 | 2.37 (1. 49-3.78) [2.56 (1. 57-4.16)] | - | 2.90 (1. 90-4.43) [2.98 (1. 94-4.569] | - | - | HbA1c _{5.7} : 2.42 (1. 50-3.91) [2.78 (1. 80-4.31)] | - | Risk ratio |
| Chamnan 2011 | 3 | - | - | - | - | - | HbA1c _{6.0} : 15.6 (6.9- 35.7) [15.5 (7.2- 33.3)] | - | Odds ratio |

| Chen 2003 | 3 | 4.4 (1.9- 10.6) | - | - | - | - | - | - | Odds ratio |
|-----------------------------------|------|---------------------------------|---|---------------------------------|--|----------------------|--|---|--|
| Coron- ado- Malagon 2009 | 1, 2 | - | - | - | [At 1 year: 7.7 (2.1- 27.9)] | - | - | - | Relative risk |
| Cugati 2007 | 10 | [19. 13 (11.59- 31.66)] | - | - | - | - | - | - | Odds ratio |
| De Abreu 2015 | 10 | 5.75 (1. 86-17.78) | - | - | - | - | - | - | Odds ratio |
| Der- akhshan 2016 | 11.7 | 6.5 years: 4.1 (2.9-5. 6) | 6.5 years: 3.0 (2.3-3. 9) | - | IFG _{5.6} and/or IGT: 4.98 (4. 08-6.07) | - | - | - | Hazard ra- tio, relative risk (6.5 years) |
| Dowse 1991 | 6.2 | - | - | [3.6 (1.4- 9.1)] | - | - | - | - | Odds ratio |
| Ferran- nini 2009 | 7 | [3.73 (2. 18-6.39)] | [4.28 (3. 21-5.71)] | [4.01 (3. 12-5.14)] | - | - | - | - | Relative risk |
| Filippatos 2016 | 10 | - | 3.43 (2. 17-5.44) | - | - | - | - | - | Odds ratio |
| Forouhi 2007 | 10 | 4.4 (1.9- 10.0) | 2.9 (1.3-6. 3) | - | - | - | - | - | Hazard ra- tio |
| Han 2017 | 12 | - | i- IFG _{5.6} : 3. 61 (2.85- 4.57) | i-IGT: 4. 06 (3.62- 4.55) | - | 8.21 (6. 79-9.94) | 6 years: HbA1c _{6.0} : Men: 4.28 (2.41-7. 58) Women: 4. 05 (1.36- 12.07) | - | Hazard ra- tio |
| Hanley 2005 | 5.2 | - | - | 5.42 (3. 60-8.17) | - | - | - | - | Odds ratio |
| Heianza 2012 | 5 | 11.4 (8. 09-16.1) | 6.18 (4. 34-8.80) | - | - | - | HbA1c _{5.7} : 6.53 (3. | | Hazard ra- tio |

| | | | | | | | 79-9.64) HbA1c _{6.0} : 7.42 (3. 67-15.0) | $\begin{array}{l} 32.5 & (23. \\ 0-45.8) \\ HbA1c_{5.7} \\ + IFG_{6.1} \\ 37.9 & (28. \\ 1-51.1) \\ HbA1c_{6.0} \\ + IFG_{5.6} \\ 53.7 & (38. \\ 4-75.1) \\ HbA1c_{6.0} \\ + IFG_{6.1} \\ 52.3 & (37. \\ 8-72.3) \end{array}$ | |
|--------------------------|-----|---|--|---|---|--|--|---|-------------------|
| Janghor- bani 2015 | 6.8 | - | 7.4 (3.7- 14.8) [8.2 (4.2- 16.0)] | 9.4 (4.8- 18.6) [10.0 (5.2- 19.1)] | - | 22.5 (12. 4-41.0) [26.7 (15. 1-47.2)] | - | - | Hazard ra- tio |
| Jeong 2010 | 5 | - | 5.66 (3. 44-9.31) | 6.01 (3. 23-11.2) | - | - | - | - | Odds ratio |
| Kim 2005 | 5 | 57 (12.18- 98.10) Men: 76. 02 (10.42- 544.51) Women: | Total: 4.77 (1.60-14. 15) Men: 9.5 (1.25- 72.24) Women: 1. 91 (0.45- 8.21) | - | - | - | - | - | Hazard ra- tio |
| Kim 2016a | 5.2 | 21.1 (16. 8-26.3) | - | - | - | - | HbA1c _{6.0} : 23.2 (18. 7-28.7) | HbA1c _{5.7} + IFG _{5.6} : 46.7 (33. 5-64.9) | Odds ratio |
| Latifi 2016 | 5 | - | 1.04 (1. 00-1.07) | - | - | - | - | - | Odds ratio |
| Leiva 2014 | 6 | 2.06 (1. 76-5.14) | - | - | - | - | - | - | Odds ratio |
| Levitzky 2008 | 4 | Women: 26.3 (17. 4-39.8) Men: 12.9 (9.3-18.1) | Women: 22.3 (13. 0-38.1) Men: 12.7 (8.1-20.0) | - | - | - | - | - | Odds ratio |

| Li 2003 | 5 | 5.78 (3. 20-10.43) | - | i-IGT: 2. 94 (1.81- 4.76) | - | 6.17 (3. 41-11.15) | - | - | Hazard ra- tio |
|------------------|------|--|--|---------------------------------|---|-----------------------|--|--|-------------------|
| Lipska 2013 | 7 | 11.4 (7.1- 18.4) | IFG _{5.6} : Total: 3.5 (1.9-6.3) Men: 8.6 (3.4-21.9) Women: 1. 5 (0.5-4.6) White: 3.2 (1.5-6. 6) Black: 4.6 (1.6- 13.3) | - | - | - | i- HbA1c _{5.7} : Total: 8.0 (4.8-13.2) Men: 24.2 (9.5-61.8) Women: 4. 6 (2.4-8.7) White: 10.2 (5.0- 20.8) Black: 5.8 (2.9- 11.7) | HbA1c _{5.7} + IFG _{5.6} : Total: 26.2 (16. 3-42.1) Men: 51.1 (21. 2-123.2) Women: 20.4 (10. 9-38.0) White: 34.9 (19. 1-63.8) Black: 14.9 (6.8- 32.6) | Odds ratio |
| Liu 2008 | 5 | - | 4.5 (2.0- 10.1) | - | - | - | - | - | Risk ratio |
| Liu 2016 | 10.9 | 1.99 (1. 37-2.90) [2.12 (1. 46-3.08)] | - | - | - | - | - | - | Hazard ra- tio |
| Liu 2017 | 7.8 | - | 3.67 (3. 20-4.21) [4.36 (3. 83-4.97)] | - | - | - | - | - | Odds ratio |
| Lorenzo 2003 | 7-8 | - | - | 6.37 (4. 37-9.28) | - | - | - | - | Odds ratio |
| Lyssenko 2005 | 6 | [i-IFG _{6.1} : 2.3 (1.4-3. 7)] | - | [i-IGT: 3.5 (2.1-5.8)] | - | [3.8 (2.3- 6.2)] | - | - | Hazard ra- tio |
| Man 2017 | 6 | - | - | - | - | 4.54 (2. 65-7.78) | - | - | Risk ratio |
| Mykkänen 1993 | 3.5 | - | - | [9.85 (6. 14-15.8)] | - | - | - | - | Odds ratio |

| Nakagami 2016 | 5 | 34.89 (19. 65-61.95) [37. 85 (22.73- 63.05)] | - | - | - | - | HbA1c _{6.0} : [63. 16 (33.94- 117.52)] HbA1c _{5.7} : 8.77(4.47- 17.21) [9.72(4. 96-19.05)] | - | Hazard ra- tio |
|------------------------------|------|--|---|---------------------------------|---|------------------------|--|---|-------------------|
| Nakanishi 2004 | 7 | 1.31 (0. 51-3.34) | - | - | - | - | - | - | Risk ratio |
| Rath- mann 2009 | 7 | [4.7 (2.2- 10.0)] | - | [8.8 (5.0- 15.6)] | - | [21.2 (10. 4-43.3)] | - | - | Odds ratio |
| Rijkeli- jkhuizen 2007 | 6.4 | i-IFG _{6.1} : 10.0 (6.1- 16.5) | - | i-IGT: 10. 9 (6.0-19. 9) | - | 39.5 (17. 0-92.1) | - | - | Odds ratio |
| Sadeghi 2015 | 7 | - | i- IFG _{5.6} : 3. 30 (2.16- 5.06) | i-IGT: 2. 52 (1.73- 3.69) | - | 12.6 (7. 39-21.4) | - | - | Odds ratio |
| Sato 2009 | 4 | 22.52 (17. 73-28.60) | | - | - | - | - | - | Odds ratio |
| Song 2015 | 4 | - | Men: 7.50 (2.76-20. 33) Women: 4. 27 (1.52- 12.00) | - | - | - | - | - | Relative risk |
| Soriguer 2008 | 6 | - | [5.3 (2.7- 10.4)] | 4.3 (2.0-9. 2) | - | 9.2 (4.3- 19.5) | - | - | Relative risk |
| Stengard 1992 | 5 | - | - | 3.1 (1.2-8. 2) | - | - | - | - | Odds ratio |
| Vaccaro 1999 | 11.5 | | [i- IFG _{6.1} : 1. 2 (0.3-10. 2)] | [i-IGT: 6. 2 (2.7-13. 8)] | - | [10.3 (2.2- 46.8)] | - | - | Odds ratio |

| Valdes 2008 | 6.3 | 12.1 (4.6- 31.7) [11.5 (5.6- 23.6] | 3.9 (1.6-9. 8) | [6.7 (3.4- 13.3)] [i-IGT: 4. 7 (1.9-11. 7)] | - | [45.6 (15. 8-131.4)] | - | - | Odds ratio |
|--------------------|------|---|---|---|---|---|------------------------------------|---|---|
| Viswanatha 2007 | 5 | - | - | 1.57 | - | - | - | - | Odds ratio |
| Wang 2007 | 5 | 2.71 (1. 43-5.16) Men: 2.29 (0.95-5. 49) Women: 1. 95 (0.83- 4.61) | 1.80 (0. 96-3.40) Men: 1.79 (0.70-4. 57) Women: 2. 08 (0.93- 4.67) | $\begin{array}{c} 3.15 (1. \\ 60-6.19) \\ i-IGT \\ (IFG_{6.1}): \\ Men: 7.33 \\ (2.62-20. \\ 51) \\ Women: 1. \\ 65 (0.76- \\ 3. \\ 60) i-IGT \\ (IFG_{5.}): \\ Men: 7.50 \\ (1.62-34. \\ 63) \\ Women: 2. \\ 21 (0.77- \\ 6.36) \end{array}$ | - | IGT/ IFG _{6.1} : Men: 10. 23 (3.84- 27.30) Women: 7. 11 (2.56- 19.72) IGT/ IFG _{5.6} : Men: 9.81 (3.5-27. 21) Women: 4. 67 (1.87- 11.62) | - | - | Risk ratio, odds ratio |
| Wang 2011 | 7, 8 | - | 25-3.63] Men: 2.10 (1.40-3. 15) [2.78 (1. 91-4.04)] | Total: 3.47 (2.64-4. 55) [4.11 (3. 20-5.27)] Men: 3.82 (2.41-6. 04) [4.72 (3. 15-7.09)] Women: 3. 16 (2.26- 4.43) [3.74 (2. 72-5.14)] | - | Total: 4.06 (3.05-5. 40) [4.68 (3. 62-6.07)] Men: 4.44 (2.75-7. 15) [5.28 (3. 49-7.99)] Women: 3. 80 (2.66- 5.42) [4.30 (3. 09-5.99)] | HbA1c _{6.0} : 5.89 (4. | - | Hazard ra- tio, odds ratio (4 years) |

| Warren 2017 | 22 | 60-3.12) Black: 2.66 (2.26-3. 13) White: 2. 86 (2.57- 3.19) Cohort 2: 3.41 (3. 01-3.85) Black: 3.16 (2.47-4. 06) White: 3. | 2.26 (2. 08-2.45) Black: 2. 05 (1.75- 2.40) White: 2. 30 (2.10- 2.53) | 2.06 (1. 84-2.31) Black: 2.55 (2.01-3. 22) White: 1. | - | - | Cohort 1: HbA1c _{5.7} : 2.71 (2. 48-2.95) Black: 2.24 (1.92-2. 61) White: 2. 91 (2.63- 3.22) HbA1c _{6.0} : 3.12 (2. 81-3.46) Black: 2.60 (2.21-3. 05) White: 3. 64 (3.20- 4.14) 6 years: HbA1c _{6.0} : 9.24 (7. 20-11.86) | - | Hazard ra- tio |
|-------------------|-----|---|--|---|---|---|--|---|-------------------|
| Yeboah 2011 | 7.5 | - | 10.5 (8.4- 13.1) [13.2 (10. 7-16.2)] | - | - | - | - | - | Hazard ra- tio |
| Zethelius 2004 | 7 | - | - | [2.18 (1. 43-3.34)] | - | - | - | - | Odds ratio |

^aUnreliable adjusted HbA1c_{6.0} interval in publication: 105.47 (29.30-101.86)

CI: confidence interval; **FPG**: fasting plasma glucose; **HbA1c**: glycosylated haemoglobin A1c; **HbA1c**_{5.7/6.0}: HbA1c threshold 5. 7% or 6.0% (usually reflecting 5.7% to 6.4% and 6.0% to 6.4%, respectively); **HbA1c/IFG**: both HbA1c and IFG; i-: isolated; **IFG**_{5.6/6.1}: impaired fasting glucose (threshold 5.6 mmol/L or 6.1 mmol/L); **IGT**: impaired glucose tolerance; **IFG/IGT**: both IFG and IGT; **IH**: intermediate hyperglycaemia; **T2DM**: type 2 diabetes mellitus

Appendix 15. Regression from intermediate hyperglycaemia to normoglycaemia

| Study ID | Follow-up (years) | Regression to normoglycaemia from IH at baseline |
|-----------------------|-------------------|--|
| Ammari 1998 | 2 | IGT: 27/68 (39.7%) |
| Anjana 2015 | 9.1 | i-IFG _{5.6} or i-IGT: 52/299 (17.4%) |
| Baena-Diez 2011 | 10 | IFG _{6.1} : 57/115 (49.6%) |
| Bai 1999 | 1 | IGT: 162/252 (64.3%) |
| Charles 1997 | 2 | IGT: 273/418 (65.3%) |
| Chen 2003 | 3 | IFG _{6.1} : 129/156 (82.6%) |
| Coronado-Malagon 2009 | 1, 2 | 'Prediabetes': 76/217 (35%) |
| Cugati 2007 | 10 | IFG _{5.6} : 5 years: 94/229 (27.9%); 10 years: 15/229 (6.6%) IFG _{6.1} : 5 years: 34/50 (68%); 10 years: 2/50 (4%) |
| De Abreu 2015 | 10 | IFG _{5.6} : 104/187 (55.6%) |
| Dowse 1991 | 6.2 | IGT: 20/51 (39%) |
| Ferrannini 2009 | 7 | IGT: 73/170 (42.9%) |
| Forouhi 2007 | 10 | IFG _{6.1} : 143/257 (55.6%) |
| Guerrero-Romero 2006 | 5 | IGT: 3/75 (4%) |
| Heianza 2012 | 5 | IFG _{5.6} : 383/1680 (22.8%) IFG _{6.1} : 101/380 (26.5%) HbA1c _{5.7} : 263/822 (32%) HbA1c _{6.0} : 63/203 (31.0%) HbA1c _{5.7} /IFG _{5.6} : 428/2092 (20.5%) HbA1c _{6.0} /IFG _{5.6} : 392/1748 (22.4%) |
| Inoue 1996 | 2.5 | IGT: 11/37 (29.7%) |
| Jiamjarasrangsi 2008a | 2.6 | IFG _{5.6} : 197/320 (61.6%) |
| Kim 2008 | 2 | IFG total: 908/1829 (49.6%) IFG5.6: 747/1335 (56%) IFG _{6.1} : 161/494 (32.6%) |
| Kleber 2010 | 1 | IGT: 52/79 (65.8%) |
| Kleber 2011 | 3.9 | IGT: 96/119 (80.1%) |
| Ko 1999 | 1.4 | IGT: 60/123 (48.8%) |

| Ко 2001 | 1.7 | IFG _{6.1} : 17/55 (30.9%) |
|----------------------|-----|--|
| Larsson 2000 | 10 | i-IFG _{6.1} : 27/42 (64.3%) i-IGT: 36/66 (54.6%) IFG/IGT: 17/30 (56.7%) |
| Latifi 2016 | 5 | IFG _{5.6} : 62/124 (50%) |
| Lecomte 2007 | 5 | IFG _{6.1} : 297/743 (44%) |
| Leiva 2014 | 6 | IFG _{6.1} : 0/28 (0%) |
| Li 2003 | 2 | IGT: 22/131 (16.8%) |
| Liu 2014 | 3 | IFG or IGT: 130/450 (28.9%) |
| Lyssenko 2005 | 6 | IFG or IGT: 379/686 (55.2%) |
| Marshall 1994 | 1.9 | IGT: 60/123 (48.8%) |
| Mohan 2008 | 8 | IGT: 6/37 (16.2%) |
| Motala 2003 | 10 | IGT: 16/35 (45.7%) 4 years: IGT: 28/72 (38.9%) |
| Mykkänen 1993 | 3.5 | IGT: 72/203 (35.5%) |
| Peterson 2017 | 10 | IGT: 8/29 (27.6%) |
| Qian 2012 | 5 | i-IFG _{6.1} : 14/46 (30.4%) i-IGT: 45/120 (37.5%) IFG/IGT: 8/33 (24.2%) |
| Rajala 2000 | 4.6 | IGT: 96/171 (56.1%) (2.1 years) IGT: 115/183 (62.8%) |
| Ramachandran 1986 | 3.3 | IGT: 34/107 (31.8%) |
| Rijkelijkhuizen 2007 | 6.4 | IFG _{6.1} : 28/149 (18.8%) IFG _{5.6} : 33/488 (6.8%) (3 years) IGT: 35/158 (22.2%) |
| Sadeghi 2015 | 7 | IFG _{5.6} and/or IGT: 148/373 (39.7%) |
| Sasaki 1982 | 7 | IGT: 5/13 (38.5%) |
| Schranz 1989 | 6 | IGT: 25/75 (33.3%) |

| Sharifi 2013 | 7 | IFG _{5.6} : 53/123 (43.1%) |
|----------------|------|---|
| Söderberg 2004 | 11 | i-IFG _{6.1} : 153/402 (38%) IGT: 296/1253 (23.6%) |
| Song 2016a | 10.8 | Total: 75/334 (22.5%) Men: 28/125 (22.4%) Women: 47/209 (22.5%) |
| Stengard 1992 | 5 | IGT: 79/234 (33.8%) |
| Toshihiro 2008 | 3.2 | IFG and/or IGT: 39/128 (30.5%) |
| Wang 2011 | 4 | IGT: 147/532 (27.6%) |
| Wat 2001 | 2 | IGT: 174/322 (54%) |
| Weiss 2005 | 1.7 | i-IGT: 15/33 (45.5%) |
| Wong 2003 | 8 | IGT: 122/291 (41.9%) |

HbA1c: glycosylated haemoglobin A1c; HbA1c_{5.7/6.0}: HbA1c threshold 5.7% or 6.0% (usually reflecting 5.7% to 6.4% and 6.0% to 6.4%, respectively); HbA1c/IFG: both HbA1c and IFG;i-: isolated; IFG_{5.6/6.1}: impaired fasting glucose (threshold 5.6 mmol/L or 6.1 mmol/L); IGT: impaired glucose tolerance; IFG/IGT: both IFG and IGT; IH: intermediate hyperglycaemia; IQR: interquartile range; SD: standard deviation

Appendix 16. Confounder adjustment (I)

| Study ID | Age | Sex | Body mass index, waist circumfer- ence, waist-to-hip ratio | 'Ethnicity' | Site | Smoking sta- tus | Drinking sta- tus | Physical activ- ity | Medications |
|-------------------------|-----|-----|---|-------------|------|---------------------|----------------------|------------------------|-------------|
| Admiraal 2014 | Yes | Yes | Yes | No | No | No | No | No | No |
| Aek- plakorn 2006 | No | No | No | No | No | No | No | No | No |
| Bae 2011 | Yes | Yes | No | No | No | No | No | No | No |

| Bergman 2016 | Yes | Yes | Yes | No | No | Yes | No | No | No |
|-----------------------------------|-----|-----|-----|-----|-----|-----|-----|-----|-----|
| Bonora 2011 | Yes | Yes | Yes | No | No | Yes | Yes | Yes | No |
| Cederberg 2010 | No | Yes | Yes | No | No | Yes | Yes | Yes | No |
| Chamnan 2011 | Yes | Yes | Yes | No | No | Yes | No | No | Yes |
| Chen 2003 | Yes | Yes | Yes | No | No | No | No | No | No |
| Coron- ado- Malagon 2009 | No |
| Cugati 2007 | Yes | Yes | No |
| De Abreu 2015 | Yes | No | Yes | No | No | Yes | Yes | Yes | No |
| Der- akhshan 2016 | Yes | Yes | Yes | No | No | Yes | No | Yes | No |
| Dowse 1991 | No |
| Ferran- nini 2009 | No |
| Filippatos 2016 | Yes | Yes | No | No | No | Yes | No | Yes | No |
| Forouhi 2007 | Yes | Yes | Yes | No | No | Yes | No | Yes | No |
| Han 2017 | Yes | Yes | Yes | No | Yes | Yes | Yes | Yes | No |
| Hanley 2005 | Yes | Yes | No | Yes | Yes | No | No | No | No |
| Heianza 2012 | Yes | Yes | Yes | No | No | Yes | No | No | No |

| Janghor- bani 2015 | Yes | Yes | Yes | No | No | No | No | No | No |
|--------------------------|-----|-----|-----|-----|-----|-----|-----|-----|-----|
| Jeong 2010 | No | No | Yes | No | No | No | No | No | No |
| Kim 2005 | Yes | Yes | Yes | No | No | No | No | No | No |
| Kim 2016a | Yes | Yes | Yes | No | No | Yes | Yes | Yes | No |
| Latifi 2016 | Yes | No | Yes | Yes | No | No | No | No | No |
| Leiva 2014 | No | No | No | No | No | Yes | No | No | Yes |
| Levitzky 2008 | Yes | No | Yes | No | No | Yes | No | No | No |
| Li 2003 | Yes | Yes | Yes | No | No | No | No | No | No |
| Lipska 2013 | Yes | Yes | No | Yes | Yes | Yes | No | Yes | No |
| Liu 2008 | Yes | Yes | No | No | No | Yes | Yes | No | No |
| Liu 2016 | Yes | No | Yes | No | No | No | No | Yes | No |
| Liu 2017 | Yes | No | No | No | No | Yes | Yes | Yes | No |
| Lorenzo 2003 | Yes | Yes | No |
| Lyssenko 2005 | No | No | Yes | No | No | No | No | No | No |
| Man 2017 | Yes | Yes | Yes | No | No | Yes | No | No | No |
| Mykkänen 1993 | No |
| Nakagami 2016 | Yes | No | Yes | No | No | Yes | Yes | No | No |
| Nakanishi 2004 | Yes | No | No | No | No | Yes | Yes | No | No |

| Rath- mann 2009 | Yes | Yes | No | No | Yes | No | No | No | No |
|------------------------------|-----|-----|-----|-----|-----|-----|-----|-----|-----|
| Rijkeli- jkhuizen 2007 | Yes | Yes | No |
| Sadeghi 2015 | Yes | Yes | Yes | No | No | No | No | No | No |
| Sato 2009 | Yes | NA | Yes | No | No | Yes | Yes | Yes | No |
| Song 2015 | Yes | No | Yes | No | No | Yes | Yes | Yes | No |
| Soriguer 2008 | Yes | Yes | Yes | No | No | No | No | No | No |
| Stengard 1992 | Yes | No | Yes | No | No | No | No | No | No |
| Vaccaro 1999 | No |
| Valdes 2008 | Yes | Yes | Yes | No | No | No | No | No | No |
| Viswanatha 2007 | Yes | No | Yes | No | No | No | No | No | No |
| Wang 2007 | Yes | Yes | No | No | No | Yes | No | No | No |
| Wang 2011 | Yes | Yes | Yes | No | No | Yes | No | No | No |
| Warren 2017 | Yes | Yes | Yes | Yes | No | Yes | Yes | No | Yes |
| Yeboah 2011 | Yes | Yes | Yes | Yes | No | No | No | Yes | No |
| Zethelius 2004 | Yes | No | Yes | No | No | No | No | No | No |

'No' denotes possible confounder but statistical analysis did not adjust for this covariate

'Yes' indicates that statistical analysis adjusted for this confounder

NA: not applicable

Appendix 17. Confounder adjustment (II)

| Study ID | Cardio- vascular disease | Glomeru- lar filtra- tion rate, albu- minuria | Blood pressure, hyperten- sion | Family history of diabetes | Socioeco- nomic status | Region | Depression | Triglyc- erides | Choles- terol |
|-----------------------------------|--------------------------------|---|---|-------------------------------------|------------------------------|--------|------------|--------------------|------------------|
| Admiraal 2014 | No | No | No | No | No | No | No | No | No |
| Aek- plakorn 2006 | No | No | No | No | No | No | No | No | No |
| Bae 2011 | No | No | No | No | No | No | No | No | No |
| Bergman 2016 | Yes | No | Yes | No | No | No | No | Yes | Yes |
| Bonora 2011 | No | No | Yes | Yes | Yes | No | No | Yes | Yes |
| Cederberg 2010 | No | No | No | No | No | No | No | No | No |
| Chamnan 2011 | No | No | Yes | Yes | Yes | No | No | Yes | Yes |
| Chen 2003 | No | No | No | Yes | No | No | No | Yes | No |
| Coron- ado- Malagon 2009 | No | No | No | No | No | No | No | No | No |
| Cugati 2007 | No | No | No | No | No | No | No | No | No |
| De Abreu 2015 | No | No | Yes | No | No | No | No | Yes | Yes |
| Der- akhshan 2016 | No | No | No | Yes | Yes | No | No | Yes | Yes |
| Dowse 1991 | No | No | No | No | No | No | No | No | No |

| Ferran- nini 2009 | No | No | No | No | No | No | No | No | No |
|--------------------------|----|----|-----|-----|-----|-----|----|-----|-----|
| Filippatos 2016 | No | No | Yes | No | No | No | No | Yes | Yes |
| Forouhi 2007 | No | No | No | Yes | No | No | No | No | No |
| Han 2017 | No | No | Yes | Yes | No | Yes | No | Yes | Yes |
| Hanley 2005 | No | No | No | No | No | No | No | No | No |
| Heianza 2012 | No | No | Yes | Yes | No | No | No | Yes | Yes |
| Janghor- bani 2015 | No | No | No | No | No | No | No | Yes | Yes |
| Jeong 2010 | No | No | Yes | No | No | No | No | Yes | Yes |
| Kim 2005 | No | No | Yes | Yes | No | No | No | Yes | Yes |
| Kim 2016a | No | No | Yes | Yes | No | No | No | Yes | Yes |
| Latifi 2016 | No | No | Yes | Yes | No | No | No | No | No |
| Leiva 2014 | No | No | No | Yes | No | No | No | No | No |
| Levitzky 2008 | No | No | No | No | No | No | No | No | No |
| Li 2003 | No | No | No | No | No | No | No | No | No |
| Lipska 2013 | No | No | Yes | No | No | No | No | No | No |
| Liu 2008 | No | No | No | Yes | No | No | No | No | No |
| Liu 2016 | No | No | No | No | No | No | No | No | No |
| Liu 2017 | No | No | No | No | Yes | Yes | No | No | No |

| Lorenzo 2003 | No | No | No | Yes | No | No | No | No | No |
|------------------------------|----|----|-----|-----|-----|----|----|-----|-----|
| Lyssenko 2005 | No | No | No | No | No | No | No | No | No |
| Man 2017 | No | No | Yes | Yes | Yes | No | No | No | Yes |
| Mykkänen 1993 | No | No | No | No | No | No | No | No | No |
| Nakagami 2016 | No | No | Yes | Yes | No | No | No | No | Yes |
| Nakanishi 2004 | No | No | No | Yes | No | No | No | No | No |
| Rath- mann 2009 | No | No | Yes | No | No | No | No | No | No |
| Rijkeli- jkhuizen 2007 | No | No | No | No | No | No | No | No | No |
| Sadeghi 2015 | No | No | No | Yes | No | No | No | No | No |
| Sato 2009 | No | No | No | Yes | No | No | No | No | No |
| Song 2015 | No | No | Yes | Yes | No | No | No | Yes | No |
| Soriguer 2008 | No | No | Yes | Yes | No | No | No | Yes | No |
| Stengard 1992 | No | No | No | No | No | No | No | No | No |
| Vaccaro 1999 | No | No | No | No | No | No | No | No | No |
| Valdes 2008 | No | No | No | No | No | No | No | Yes | No |
| Viswanatha 2007 | No | No | No | Yes | No | No | No | No | No |

| Wang 2007 | No | No | No | Yes | Yes | No | No | No | Yes |
|-------------------|----|-----|-----|-----|-----|----|----|-----|-----|
| Wang 2011 | No | No | Yes | Yes | No | No | No | Yes | Yes |
| Warren 2017 | No | Yes | Yes | Yes | Yes | No | No | Yes | Yes |
| Yeboah 2011 | No | No | No | No | Yes | No | No | No | No |
| Zethelius 2004 | No | No | No | No | No | No | No | No | No |

'No' denotes possible confounder but statistical analysis did not adjust for this covariate

'Yes' indicates that statistical analysis adjusted for this confounder

CONTRIBUTIONS OF AUTHORS

All review authors read and approved the final review draft.

Bernd Richter (BR): protocol and review draft, search strategy development, acquisition of trial reports, trial selection, data extraction of all trials, data analysis, data interpretation and writing of drafts.

Maria-Inti Metzendorf (MIM): search strategy development, trial selection, check of data extraction, review of drafts.

Bianca Hemmingsen (BH): protocol and review draft, trial selection, data interpretation and review of drafts.

Yemisi Takwoingi (YT): protocol and review draft, data analysis, data interpretation and review of drafts

DECLARATIONS OF INTEREST

BR: the World Health Organization (WHO) funded this review.

MIM: none known.

BH: none known.

YT: none known.

SOURCES OF SUPPORT

Internal sources

• No sources of support supplied

External sources

• World Health Organization, Other.

This review is part of a series of reviews on predictors for the development of type 2 diabetes mellitus in people with intermediate hyperglycaemia and interventions for the prevention or delay of type 2 diabetes mellitus and its associated complications in persons at increased risk for the development of type 2 diabetes mellitus which is funded by the WHO (Hemmingsen 2016a; Hemmingsen 2016b; Hemmingsen 2016c)

DIFFERENCES BETWEEN PROTOCOL AND REVIEW

We changed the title of the protocol from 'Intermediate hyperglycaemia as a predictor for the development of type 2 diabetes: prognostic factor exemplar review' to 'Development of type 2 diabetes mellitus in people with intermediate hyperglycaemia' to fit the objectives of the review. We also modified the objectives from "to assess whether intermediate hyperglycaemia is a predictor for the development of type 2 diabetes mellitus (T2DM)" to objective 1 "to assess the overall prognosis of people with IH for the development of T2DM and to assess how many people with IH revert back to normoglycaemia (regression), and objective 2 "to assess the difference in T2DM incidence in people with IH versus people with normoglycaemia". Both changes reflect the fact that our review addresses two prognostic questions at the same time. First, if people have intermediate hyperglycaemia at baseline, how many individuals develop type 2 diabetes in the future? This research question investigates the cumulative incidence of type 2 diabetes over time and does not depend on a comparison with a group with normoglycaemia at baseline; it is also important to note how many people change back from intermediate hyperglycaemia compared with normoglycaemia. The second prognostic question is, how does glycaemic status (intermediate hyperglycaemia compared with normoglycaemia) at baseline affect the development of type 2 diabetes? In particular, we were interested in intermediate hyperglycaemia, defined using impaired fasting glucose, impaired glucose tolerance and elevated glycosylated haemoglobin A1c and combinations thereof.

We specified inclusion criteria in more detail to explain the difference between studies evaluating the overall prognosis of people with intermediate hyperglycaemia and studies evaluating intermediate hyperglycaemia versus normoglycaemia as a prognostic factor developing type 2 diabetes mellitus.

Regarding methods, we explained our exclusion criteria in more detail and deleted 'conference abstract' as an exclusion criterion (we moved one formerly excluded study, Misnikova 2011, to 'Studies awaiting classification').