

Person—diabetes mellitus status, code NN

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Person—diabetes mellitus status, code NN

Identifying and definitional attributes

Metadata item type:	Data Element
Short name:	Diabetes status
METEOR identifier:	270194
Registration status:	Health! , Standard 01/03/2005 Indigenous , Standard 13/03/2015
Definition:	Whether a person has or is at risk of diabetes, as represented by a code.
Data Element Concept:	Person—diabetes mellitus status
Value Domain:	Diabetes mellitus status code NN

Value domain attributes

Representational attributes

Representation class:	Code
Data type:	String
Format:	NN
Maximum character length:	2

	Value	Meaning
Permissible values:	01	Type 1 diabetes
	02	Type 2 diabetes
	03	Gestational diabetes mellitus (GDM)
	04	Other (secondary diabetes)
	05	Previous gestational diabetes mellitus (GDM)
	06	Impaired fasting glucose (IFG)
	07	Impaired glucose tolerance (IGT)
	08	Not diagnosed with diabetes
	09	Not assessed
Supplementary values:	99	Not stated/inadequately described

Collection and usage attributes

Guide for use: Note that where there is a Gestational diabetes mellitus (GDM) or Previous GDM (i.e. permissible values 3 & 5) and a current history of Type 2 diabetes then record 'Code 2' Type 2 diabetes.

This same principle applies where a history of either Impaired fasting glycaemia (IFG) or Impaired glucose tolerance (IGT) and a current history and Type 2 diabetes, then record 'Code 2' Type 2 diabetes.

CODE 01 Type 1 diabetes

Beta-cell destruction, usually leading to absolute insulin deficiency. Includes those cases attributed to an autoimmune process, as well as those with beta-cell destruction and who are prone to ketoacidosis for which neither an aetiology nor pathogenesis is known (idiopathic). It does not include those forms of beta-cell destruction or failure to which specific causes can be assigned (e.g. cystic fibrosis, mitochondrial defects). Some subjects with Type 1 diabetes can be identified at

earlier clinical stages than 'diabetes mellitus'.

CODE 02 Type 2 diabetes

Type 2 includes the common major form of diabetes, which results from defect(s) in insulin secretion, almost always with a major contribution from insulin resistance.

CODE 03 Gestational diabetes mellitus (GDM)

GDM is a carbohydrate intolerance resulting in hyperglycaemia of variable severity with onset or first recognition during pregnancy. The definition applies irrespective of whether or not insulin is used for treatment or the condition persists after pregnancy. Diagnosis is to be based on the Australian Diabetes in Pregnancy Society (ADIPS) Guidelines.

CODE 04 Other (secondary diabetes)

This categorisation includes less common causes of diabetes mellitus, but are those in which the underlying defect or disease process can be identified in a relatively specific manner. They include, for example, genetic defects of beta-cell function, genetic defects in insulin action, diseases of the exocrine pancreas, endocrinopathies, drug or chemical-induced, infections, uncommon forms of immune-mediated diabetes, other genetic syndromes sometimes associated with diabetes.

CODE 05 Previous GDM

Where the person has a history of GDM.

CODE 06 Impaired fasting glycaemia (IFG)

IFG or 'non-diabetic fasting hyperglycaemia' refers to fasting glucose concentrations, which are lower than those required to diagnose diabetes mellitus but higher than the normal reference range. An individual is considered to have IFG if they have a fasting plasma glucose of 6.1 or greater and less than 7.0 mmol/L if challenged with an oral glucose load, they have a fasting plasma glucose concentration of 6.1 mmol/L or greater, but less than 7.0 mmol/L, AND the 2 hour value in the Oral Glucose Tolerance Test (OGTT) is less than 7.8 mmol/L.

CODE 07 Impaired glucose tolerance (IGT)

IGT is categorised as a stage in the natural history of disordered carbohydrate metabolism; subjects with IGT have an increased risk of progressing to diabetes. IGT refers to a metabolic state intermediate between normal glucose homeostasis and diabetes. Those individuals with IGT manifest glucose intolerance only when challenged with an oral glucose load. IGT is diagnosed if the 2 hour value in the OGTT is greater than 7.8 mmol/L. and less than 11.1 mmol/L AND the fasting plasma glucose concentration is less than 7.0 mmol/L.

CODE 08 Not diagnosed with diabetes

The subject has no known diagnosis of Type 1, Type 2, GDM, Previous GDM, IFG, IGT or Other (secondary diabetes).

CODE 09 Not assessed

The subject has not had their diabetes status assessed.

CODE 99 Not stated/inadequately described

This code is for unknown or information unavailable.

Collection methods:

The diagnosis is derived from and must be substantiated by clinical documentation.

Source and reference attributes

Origin:

Developed based on Definition, Diagnosis and Classification of Diabetes Mellitus and its Complications Part 1: Diagnosis and Classifications of Diabetes Mellitus Provisional Report of a World Health Organization Consultation (Alberti & Zimmet 1998).

Data element attributes

Collection and usage attributes

Collection methods: Diabetes (clinical):

A type of diabetes should be recorded and coded for each episode of patient care.


Source and reference attributes

Submitting organisation: Cardiovascular Data Working Group

National Diabetes Data Working Group

Relational attributes

Related metadata references:

Is re-engineered from  [Diabetes status, version 1, DE, NHDD, NHMG, Superseded 01/03/2005.pdf](#) (27.3 KB)

No registration status

See also [Primary Health Network—number of regular clients with diabetes mellitus, total number N\[NNNNNN\]](#)

[Health!](#), Recorded 05/01/2021

Implementation in Data Set Specifications:

[Acute coronary syndrome \(clinical\) DSS](#)

[Health!](#), Superseded 07/12/2005

[Acute coronary syndrome \(clinical\) DSS](#)

[Health!](#), Superseded 01/10/2008

[Acute coronary syndrome \(clinical\) DSS](#)

[Health!](#), Superseded 01/09/2012

[Acute coronary syndrome \(clinical\) DSS](#)

[Health!](#), Superseded 02/05/2013

[Acute coronary syndrome \(clinical\) NBPDS 2013-](#)

[Health!](#), Standard 02/05/2013

Implementation start date: 01/07/2013

[Cardiovascular disease \(clinical\) DSS](#)

[Health!](#), Superseded 15/02/2006

DSS specific information:

People with diabetes have two to five times increased risk of developing heart, stroke and vascular disease (Zimmet & Alberti 1997). Cardiovascular disease is the most common cause of death in people with diabetes.

Diabetes is also an important cause of stroke, and people with diabetes may have a worse prognosis after stroke.

Heart, stroke and vascular disease and diabetes share common risk factors, but also diabetes is an independent risk factor for heart, stroke and vascular disease.

During the 1995 National Health Survey, about 15 per cent of those with diabetes reported having heart disease, at almost six times the rate noted among people without diabetes. In 1996-97, almost one in six hospital separations, with coronary heart disease as any listed diagnosis, also had diabetes recorded as an associated diagnosis. Heart disease appears earlier in life and is more often fatal among those with diabetes.

Diabetes may accentuate the role of elevated blood pressure in stroke. The incidence and prevalence of peripheral vascular disease in those with diabetes increase with the duration of the peripheral vascular disease.

Mortality is increased among patients with peripheral vascular disease and diabetes, in particular if foot ulcerations, infection or gangrene occur. There is limited information on whether the presence of heart, stroke and vascular disease promotes diabetes in some way.

High blood pressure, high cholesterol and obesity are often present along with

diabetes. As well as all being independent cardiovascular risk factors, when they are in combination with glucose intolerance (a feature of diabetes) and other risk factors such as physical inactivity and smoking, these factors present a greater risk for heart, stroke and vascular disease.

Evidence is accumulating that high cholesterol and glucose intolerance, which often occur together, may have a common aetiological factor. Despite these similarities, trends in cardiovascular mortality and diabetes incidence and mortality are moving in opposite directions.

While the ageing of the population following reductions in cardiovascular mortality may have contributed to these contrasting trends, the role of other factors also needs to be clearly understood if common risk factor prevention strategies are to be considered. (From Commonwealth Department of Health & Aged Care and Australian Institute of Health and Welfare (1999) National Health Priority Areas Report: Cardiovascular Health).

In settings such as general practice where the monitoring of a person's health is ongoing and where diabetes status can change over time, the service contact date should be recorded.

[Cardiovascular disease \(clinical\) DSS](#)
Health!, Superseded 04/07/2007

DSS specific information:

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[Cardiovascular disease \(clinical\) DSS](#)

[Health!](#), Superseded 22/12/2009

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[Cardiovascular disease \(clinical\) DSS](#)

[Health!](#), Superseded 01/09/2012

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In settings such as general practice where the monitoring of a person's health is ongoing and where diabetes status can change over time, the service contact date should be recorded.

[Cardiovascular disease \(clinical\) NBPDS Health!](#), Superseded 17/10/2018

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[Cardiovascular disease \(clinical\) NBPDS](#)

[Health!](#), Standard 17/10/2018

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[Diabetes \(clinical\) DSS](#)

[Health!](#), Superseded 21/09/2005

DSS specific information:

Uncontrolled diabetes leads to a variety of complications, often resulting in limitation of activity, disability, illness and premature mortality. Therefore ongoing assessment is required to identify people at risk of developing complications so that early preventive strategies can be applied. Although there is no cure for diabetes, with modern treatment most people can lead a full and active life and avoid long-term complications.

Aetiological classifications contained in the scientific paper 'Definition, Diagnosis and Classification of Diabetes Mellitus and its Complications Part 1: Diagnosis and Classifications of Diabetes Mellitus Provisional Report of a WHO Consultation (Alberti & Zimmet 1998)'.

[Diabetes \(clinical\) NBPDS](#)

[Health!](#), Standard 21/09/2005

DSS specific information:

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[Indigenous primary health care DSS 2012-14](#)

[Health!](#), Superseded 21/11/2013

[Indigenous](#), Superseded 21/11/2013

Implementation start date: 01/07/2012

Implementation end date: 30/06/2014

[Indigenous primary health care DSS 2014-15](#)

[Health!](#), Superseded 13/03/2015

[Indigenous](#), Superseded 13/03/2015

Implementation start date: 01/07/2014

Implementation end date: 30/06/2015

[Indigenous primary health care DSS 2015-17](#)

[Health!](#), Superseded 25/01/2018

[Indigenous](#), Superseded 27/02/2018

Implementation start date: 01/07/2015

Implementation end date: 30/06/2017

[Indigenous primary health care NBEDS 2017–18](#)

[Health!](#), Superseded 06/09/2018

[Indigenous](#), Superseded 22/10/2018

Implementation start date: 01/07/2017

Implementation end date: 30/06/2018

[Indigenous primary health care NBEDS 2018–19](#)

[Health!](#), Superseded 12/12/2018

[Indigenous](#), Superseded 02/04/2019

Implementation start date: 01/07/2018

Implementation end date: 30/06/2019

[Indigenous primary health care NBEDS 2019–20](#)

[Health!](#), Superseded 16/01/2020

[Indigenous](#), Superseded 14/07/2021

Implementation start date: 01/07/2019

Implementation end date: 30/06/2020

[Indigenous primary health care NBEDS 2020–21](#)

[Health!](#), Retired 13/10/2021

Implementation in Indicators:

Implementation start date: 01/07/2020

Implementation end date: 30/06/2021

[Indigenous-specific primary health care NBEDS December 2020](#)

[Indigenous, Standard 14/07/2021](#)

Use as a National Indicator start date: 01/07/2020

Implementation end date: 31/12/2020
[Indigenous primary health care: PI05a-Number of regular clients with Type II diabetes who have had an HbA1c measurement result recorded, 2012](#)

[Health!, Superseded 23/02/2012](#)

[Indigenous primary health care: PI05a-Number of regular clients with Type II diabetes who have had an HbA1c measurement result recorded, 2013](#)

[Health!, Superseded 21/11/2013](#)

[Indigenous, Superseded 21/11/2013](#)

[Indigenous primary health care: PI05a-Number of regular clients with Type II diabetes who have had an HbA1c measurement result recorded, 2014](#)

[Health!, Superseded 13/03/2015](#)

[Indigenous, Superseded 13/03/2015](#)

[Indigenous primary health care: PI05a-Number of regular clients with Type II diabetes who have had an HbA1c measurement result recorded, 2015](#)

[Health!, Superseded 05/10/2016](#)

[Indigenous, Superseded 20/01/2017](#)

[Indigenous primary health care: PI05a-Number of regular clients with Type II diabetes who have had an HbA1c measurement result recorded, 2015-2017](#)

[Health!, Superseded 25/01/2018](#)

[Indigenous, Superseded 27/02/2018](#)

[Indigenous primary health care: PI05a-Number of regular clients with Type II diabetes who have had an HbA1c measurement result recorded, 2015-2017](#)

[Health!, Superseded 17/10/2018](#)

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[Indigenous primary health care: PI05a-Number of regular clients with Type II diabetes who have had an HbA1c measurement result recorded, 2018-2019](#)

[Health!, Superseded 16/01/2020](#)

[Indigenous, Superseded 14/07/2021](#)

[Indigenous primary health care: PI05a-Number of regular clients with Type II diabetes who have had an HbA1c measurement result recorded, June 2020](#)

[Health!, Retired 13/10/2021](#)

[Indigenous, Superseded 14/07/2021](#)

[Indigenous primary health care: PI05b-Proportion of regular clients with Type II diabetes who have had an HbA1c measurement result recorded, 2012](#)

[Health!, Superseded 23/02/2012](#)

[Indigenous primary health care: PI05b-Proportion of regular clients with Type II diabetes who have had an HbA1c measurement result recorded, 2013](#)

[Health!, Superseded 21/11/2013](#)

[Indigenous, Superseded 21/11/2013](#)

[Indigenous primary health care: PI05b-Proportion of regular clients with Type II diabetes who have had an HbA1c measurement result recorded, 2014](#)

[Health!, Superseded 13/03/2015](#)

[Indigenous, Superseded 13/03/2015](#)

[Indigenous primary health care: PI05b-Proportion of regular clients with Type II diabetes who have had an HbA1c measurement result recorded, 2015](#)

[Health!, Superseded 05/10/2016](#)

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[Indigenous primary health care: PI05b-Proportion of regular clients with Type II diabetes who have had an HbA1c measurement result recorded, 2015-2017](#)

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[Indigenous, Superseded 27/02/2018](#)

[Indigenous primary health care: PI05b-Proportion of regular clients with Type II diabetes who have had an HbA1c measurement result recorded, 2015-2017](#)

[Health!, Superseded 17/10/2018](#)

[Indigenous, Superseded 17/10/2018](#)

[Indigenous primary health care: PI05b-Proportion of regular clients with Type II](#)

[diabetes who have had an HbA1c measurement result recorded, 2018-2019](#)
[Health!](#), Superseded 16/01/2020
[Indigenous](#), Superseded 14/07/2021

[Indigenous primary health care: PI05b-Proportion of regular clients with Type II diabetes who have had an HbA1c measurement result recorded, June 2020](#)
[Health!](#), Retired 13/10/2021
[Indigenous](#), Superseded 14/07/2021

[Indigenous primary health care: PI06a-Number of regular clients with Type II diabetes whose HbA1c measurement result was within a specified level, 2012](#)
[Health!](#), Superseded 23/02/2012

[Indigenous primary health care: PI06a-Number of regular clients with Type II diabetes whose HbA1c measurement result was within a specified level, 2013](#)
[Health!](#), Superseded 21/11/2013
[Indigenous](#), Superseded 21/11/2013

[Indigenous primary health care: PI06a-Number of regular clients with Type II diabetes whose HbA1c measurement result was within a specified level, 2014](#)
[Health!](#), Superseded 13/03/2015
[Indigenous](#), Superseded 13/03/2015

[Indigenous primary health care: PI06a-Number of regular clients with Type II diabetes whose HbA1c measurement result was within a specified level, 2015](#)
[Health!](#), Superseded 05/10/2016
[Indigenous](#), Superseded 20/01/2017

[Indigenous primary health care: PI06a-Number of regular clients with Type II diabetes whose HbA1c measurement result was within a specified level, 2015-2017](#)
[Health!](#), Superseded 17/10/2018
[Indigenous](#), Superseded 17/10/2018

[Indigenous primary health care: PI06a-Number of regular clients with Type II diabetes whose HbA1c measurement result was within a specified level, 2015-2017](#)
[Health!](#), Superseded 25/01/2018
[Indigenous](#), Superseded 27/02/2018

[Indigenous primary health care: PI06a-Number of regular clients with Type II diabetes whose HbA1c measurement result was within a specified level, 2018-2019](#)
[Health!](#), Superseded 16/01/2020
[Indigenous](#), Superseded 14/07/2021

[Indigenous primary health care: PI06a-Number of regular clients with Type II diabetes whose HbA1c measurement result was within a specified level, June 2020](#)
[Health!](#), Retired 13/10/2021
[Indigenous](#), Superseded 14/07/2021

[Indigenous primary health care: PI06b-Proportion of regular clients with Type II diabetes whose HbA1c measurement result was within a specified level, 2012](#)
[Health!](#), Superseded 23/02/2012

[Indigenous primary health care: PI06b-Proportion of regular clients with Type II diabetes whose HbA1c measurement result was within a specified level, 2013](#)
[Health!](#), Superseded 21/11/2013
[Indigenous](#), Superseded 21/11/2013

[Indigenous primary health care: PI06b-Proportion of regular clients with Type II diabetes whose HbA1c measurement result was within a specified level, 2014](#)
[Health!](#), Superseded 13/03/2015
[Indigenous](#), Superseded 13/03/2015

[Indigenous primary health care: PI06b-Proportion of regular clients with Type II diabetes whose HbA1c measurement result was within a specified level, 2015](#)
[Health!](#), Superseded 05/10/2016
[Indigenous](#), Superseded 20/01/2017

[Indigenous primary health care: PI06b-Proportion of regular clients with Type II diabetes whose HbA1c measurement result was within a specified level, 2015-2017](#)
[Health!](#), Superseded 17/10/2018

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[Indigenous primary health care: PI06b-Proportion of regular clients with Type II diabetes whose HbA1c measurement result was within a specified level, 2015-2017](#)

[Health!](#), Superseded 25/01/2018

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[Indigenous primary health care: PI06b-Proportion of regular clients with Type II diabetes whose HbA1c measurement result was within a specified level, 2018-2019](#)

[Health!](#), Superseded 16/01/2020

[Indigenous](#), Superseded 14/07/2021

[Indigenous primary health care: PI06b-Proportion of regular clients with Type II diabetes whose HbA1c measurement result was within a specified level, June 2020](#)

[Health!](#), Retired 13/10/2021

[Indigenous](#), Superseded 14/07/2021

[Indigenous primary health care: PI07a-Number of regular clients with a chronic disease for whom a GP Management Plan \(MBS Item 721\) was claimed, 2014](#)

[Health!](#), Superseded 13/03/2015

[Indigenous](#), Superseded 13/03/2015

[Indigenous primary health care: PI07a-Number of regular clients with a chronic disease for whom a GP Management Plan \(MBS Item 721\) was claimed, 2015](#)

[Health!](#), Superseded 05/10/2016

[Indigenous](#), Superseded 20/01/2017

[Indigenous primary health care: PI07a-Number of regular clients with a chronic disease for whom a GP Management Plan \(MBS Item 721\) was claimed, 2015-2017](#)

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