Type 2 diabetes after gestational diabetes: greater than fourfold risk among Indigenous compared with non-Indigenous Australian women†

Abstract

Background Gestational diabetes is associated with a high risk of type 2 diabetes. However, progression rates among Indigenous women in Australia who experience high prevalence of gestational diabetes are unknown.

Methods This retrospective cohort study includes all births to women at a regional hospital in Far North Queensland, Australia, coded as having 'gestational diabetes' from 1 January 2004 to 31 December 2010 (1098 births) and receiving laboratory postpartum screening from 1 January 2004 to 31 December 2011 (n = 483 births). Women who did not receive postpartum screening were excluded from the denominator. Data were linked between hospital electronic records, routinely collected birth data and laboratories, with sample validation by reviews of medical records. Analysis was conducted using Cox-proportional regression models.

Results Indigenous women had a greater than fourfold risk of developing type 2 diabetes within 8 years of having gestational diabetes, compared with non-Indigenous women (hazards ratio 4.55, 95% confidence interval 2.63–7.88, p < 0.0001). Among women receiving postpartum screening tests, by 3, 5 and 7 years postpartum, 21.9% (15.8–30.0%), 25.5% (18.6–34.3%) and 42.4% (29.6–58.0%) Indigenous women were diagnosed with type 2 diabetes after gestational diabetes, respectively, compared with 4.2% (2.5–7.2%), 5.7% (3.3–9.5%) and 13.5% (7.3–24.2%) non-Indigenous women. Multivariate analysis showed significant associations between an increased rate of type 2 diabetes with an early pregnancy body mass index >25 kg/m², partial breastfeeding only at hospital discharge and gestational diabetes diagnosis prior to 17 weeks gestation.

Conclusions This study demonstrates that, compared with non-Indigenous women, Indigenous Australian women have a greater than fourfold risk of developing type 2 diabetes after gestational diabetes. Strategies are urgently needed to reduce rates of type 2 diabetes by supporting a healthy weight and breastfeeding and to improve postpartum screening among Indigenous women with gestational diabetes. Copyright © 2015 John Wiley & Sons, Ltd.

Keywords gestational diabetes mellitus; type 2 diabetes mellitus; diabetes; pregnancy; aboriginal; Indigenous
Introduction

Gestational diabetes mellitus (gestational diabetes), defined as diabetes diagnosed during pregnancy [1], is increasing in prevalence, with particularly high rates reported among Indigenous women worldwide [2]. In addition to causing serious complications in pregnancy and birth [3], women diagnosed with gestational diabetes have a very high risk of developing type 2 diabetes mellitus (type 2 diabetes) postpartum [4]. If left undetected and untreated in the longer term, unmanaged hyperglycaemia associated with type 2 diabetes can cause serious complications in subsequent pregnancies [5] and lead to multiple comorbidities for the mother [6].

Particularly high rates of progression from gestational to type 2 diabetes have been reported among Indigenous women internationally [7]. Progression rates range from 30% to 70% within 4 years postpartum, among Indigenous women in Canada [8], New Zealand [9] and the United States [10,11]. However, there are currently few studies investigating rates of progression from gestational to type 2 diabetes among Aboriginal and Torres Strait Islander (Indigenous) women in Australia. One study reported that two out of seven Indigenous Australian women with gestational diabetes receiving postpartum screening had developed type 2 diabetes, but this sample was too small for substantive analysis [12].

The aim of this article is to investigate the rates and associations with progression to type 2 diabetes among Indigenous and non-Indigenous women, diagnosed with gestational diabetes in Far North Queensland (Australia). Comparisons between Indigenous and non-Indigenous women are also needed to determine whether existing services currently provided for all women are adequate for Indigenous women or whether tailoring of services and additional strategies are required for Indigenous women.

Materials and methods

Study setting and sample

This study includes all women who gave birth at Cairns Hospital and were coded as having gestational diabetes from 1 January 2004 to 31 December 2010. Cairns Hospital is the only secondary referral hospital for Far North Queensland, a vast region on the northeast tip of Australia covering almost 300 000 km². The region has a population of over 230 000, and approximately 40 000 (17%) are Aboriginal and Torres Strait Islander people [13]. About half of the population lives in the main regional centre of Cairns, and the remainder lives in areas classified as rural and remote, in a sparsely populated tropical region that has limited sealed road access and is subject to extreme weather events. More than 80% of women in the region give birth at Cairns Hospital, which includes almost all women with gestational diabetes. The study setting and design details are reported in detail elsewhere [14].

The study is a retrospective cohort design that used linked electronic data validated by medical record reviews. The following data sources were linked: (1) the Cairns Hospital Clinical Coding system (CHCCS) entries for all women who gave birth at Cairns Hospital between 1 January 2004 and 31 December 2010 and were coded as having gestational diabetes (International Classification Diseases codes 024.41, 024.42, 024.42, 024.43 and 024.44); (2) pregnancy and birth details from the Midwives Perinatal Data Collection (MPDC); and (3) postpartum glucose test details from the three local laboratories. Additionally, available medical records were reviewed for all births among Indigenous women (n = 578) and a random sample of births among non-Indigenous women (n = 332) to validate gestational diabetes coding and extract additional data not reliably available in electronic data [antepartum and postpartum care providers, body mass index (BMI) and English proficiency]. This sample size was assessed as adequate to detect a 10% difference in postpartum screening between Indigenous and non-Indigenous women.

Gestational diabetes diagnostic criteria

During the study period, the diagnostic criteria for gestational diabetes were consistent with the Australian Diabetes in Pregnancy Society guidelines [15], which required fasting plasma glucose (FPG) ≥5.5 mmol/L or 2-h glucose ≥8 mmol/L following a 75-g oral glucose tolerance test (OGTT). However, the standard procedure for gestational diabetes screening varied markedly across Australia during this period [16]. In 2005, local primary care guidelines were developed, which recommended a random blood glucose level at each pregnancy visit, and if the level was ≥5.0 mmol/L, a fasting blood glucose level was offered [17]. At 24 weeks, a random blood glucose level was offered again, and if ≥5.0 mmol/L or the woman was classified as ‘at risk’ of gestational diabetes, a 75-g OGTT was offered. ‘At risk’ included a history of
unexplained miscarriage or stillbirth, large baby (>4000 g or ‘large for dates’), age >30 years, obesity (BMI >30 kg/m²), previous neonatal hyperglycaemia, family history of diabetes, past history of gestational diabetes or polycystic ovary syndrome (indigeneity was not a risk factor). If normal, the 75-g OGTT was to be offered again at 32 weeks. Glycosylated haemoglobin (HbA1c) was recommended around 28 weeks or at first presentation. In 2007, the primary care guidelines changed to include all Indigenous women as ‘high risk’, along with ‘other high risk populations’, such as Pacific Islander, Chinese and Mediterranean people, and recommended a 75-g OGTT at 26–30 weeks gestation [18]. In 2009, the guidelines were further revised to recommend a 75-g OGTT at 24–28 weeks gestation [19], as per the Australian Diabetes in Pregnancy Society guidelines [15].

Type 2 diabetes diagnostic criteria

The diagnostic criteria for type 2 diabetes at the time of the study are outlined in Table 1, which was clarified in consultation with local endocrinology experts. The recommended diagnosis was based on glucose readings following a 75-g OGTT equal to or more than 7 mmol (fasting), 10 mmol/L (after 1 h) or 11 mmol/L (after 2 h) [20]. In this study, women with a diagnosis of type 2 diabetes were identified by the presence of one of these readings, a notation on the laboratory results identifying the woman as having type 2 diabetes, subsequent coding of a pregnancy as type 2 diabetes or documentation of type 2 diabetes in the pregnancy or medical records.

Indigenous status criteria

Indigenous status is a measure of whether a person self-identifies as being of Aboriginal, Torres Strait Islander or Aboriginal and Torres Strait Islander descent, and this is recorded in the CHCCS.

Data analysis

De-identified data were exported from Microsoft Access for analysis in Stata 13 [21]. All analyses were stratified by Indigenous status. Primary analyses were restricted to women who received a laboratory-based postpartum screening test (OGTT, HbA1c, FPG or random plasma glucose) as women who did not receive a test had no chance of being diagnosed with type 2 diabetes. Analyses included all births to mothers coded as having gestational diabetes, who received a laboratory-based postpartum screening test, where routinely collected data on potential moderating factors were consistently available in electronic data, including country of birth, remoteness, maternal age, number of antenatal care visits, parity, smoking during pregnancy, medical complications, induction of labour, mode of birth and confinement date (Table 1). Analyses were restricted only to those records where the medical records were reviewed when data were not consistently recorded in routine electronic data, including BMI at first antenatal visit, location of antenatal or postnatal care (hospital, private general practitioner clinic, government clinic, community-controlled health service or other), breastfeeding at discharge and diagnosis of gestational diabetes prior to 17 weeks gestation (probable type 2 diabetes) (Table 2). Sensitivity analyses were also conducted including all eligible women with gestational diabetes, with and without postpartum screening, in the denominator.

The time to type 2 diabetes diagnosis from confinement date, among births to mothers coded as having gestational diabetes, was summarized using Kaplan–Meier survival estimates and analysed using Cox-proportional hazards regression models. Women were censored from the analysis if they became pregnant and once type 2 diabetes was diagnosed. The censorship was calculated from time of onset of subsequent pregnancy, calculated as 273 days prior to subsequent confinement or 20 weeks prior to date of test if the test was coded as ‘during pregnancy’ and no pregnancy was recorded (including

Table 1. Glucose parameters used during study period in Far North Queensland* [20]

<table>
<thead>
<tr>
<th>Test</th>
<th>Glucose load (g)</th>
<th>Normal reference range (mmol/L)</th>
<th>Impaired glucose tolerance (mmol/L)</th>
<th>Abnormal glucose tolerance (mmol/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Pregnant</td>
<td>Not pregnant</td>
<td>Pregnant</td>
</tr>
<tr>
<td>OGTT fasting/FPG</td>
<td>75</td>
<td>3.6–5.4</td>
<td>3.6–5.4</td>
<td>5.5–6.9</td>
</tr>
<tr>
<td>OGTT 1 h</td>
<td>75</td>
<td>&lt;5.5</td>
<td>&lt;7.8</td>
<td>≥10.0</td>
</tr>
<tr>
<td>OGTT 2 h</td>
<td>75</td>
<td></td>
<td></td>
<td>≥11.0</td>
</tr>
</tbody>
</table>

FPG, fasting plasma glucose; OGTT, oral glucose tolerance test.

*Modified in consultation with local endocrinology experts about criteria used during study period.
Table 2. Associations with progression time from gestational diabetes to type 2 diabetes, by Indigenous status [among births coded as gestational diabetes with any post-partum screening test (OGTT, HbA1c, FPG and RPG) recorded]

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Combined Indigenous and non-Indigenous (n = 483)</th>
<th>Indigenous (n = 155)</th>
<th>Non-Indigenous (n = 328)</th>
<th>Indigenous versus non-Indigenous difference</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n (%)&lt;sup&gt;c&lt;/sup&gt;</td>
<td>HR</td>
<td>CI</td>
<td>p</td>
</tr>
<tr>
<td>Indigenous status</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Non-Indigenous</td>
<td>328 (68)</td>
<td>4.55</td>
<td>2.63–7.88</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Indigenous</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aboriginal</td>
<td>328 (68)</td>
<td>Ref</td>
<td>NA</td>
<td></td>
</tr>
<tr>
<td>Torres Strait</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Islander</td>
<td>12 (2)</td>
<td>8.18</td>
<td>2.78–24.09</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Remoteness&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Remote or very remote</td>
<td>88 (18)</td>
<td>Ref</td>
<td>NA</td>
<td></td>
</tr>
<tr>
<td>Cairns</td>
<td>394 (82)</td>
<td>0.73</td>
<td>0.40–1.32</td>
<td>0.30</td>
</tr>
<tr>
<td>Country of birth</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Outside Australia</td>
<td>88 (18)</td>
<td>Ref</td>
<td>NA</td>
<td></td>
</tr>
<tr>
<td>Australia</td>
<td>394 (82)</td>
<td>0.73</td>
<td>0.40–1.32</td>
<td>0.30</td>
</tr>
<tr>
<td>Maternal age (years)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;25</td>
<td>42 (9)</td>
<td>Ref</td>
<td>NA</td>
<td></td>
</tr>
<tr>
<td>25–29</td>
<td>96 (20)</td>
<td>0.79</td>
<td>0.32–1.94</td>
<td>0.61</td>
</tr>
<tr>
<td>30–34</td>
<td>134 (28)</td>
<td>0.69</td>
<td>0.27–1.65</td>
<td>0.38</td>
</tr>
<tr>
<td>35+</td>
<td>211 (44)</td>
<td>0.84</td>
<td>0.37–1.86</td>
<td>0.66</td>
</tr>
<tr>
<td>Antenatal visits</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;8</td>
<td>82 (17)</td>
<td>Ref</td>
<td>NA</td>
<td></td>
</tr>
<tr>
<td>8+</td>
<td>391 (81)</td>
<td>1.37</td>
<td>0.65–2.90</td>
<td>0.41</td>
</tr>
<tr>
<td>Parity</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>0</td>
<td>153 (32)</td>
<td>Ref</td>
<td>NA</td>
<td></td>
</tr>
<tr>
<td>1-4</td>
<td>276 (57)</td>
<td>1.15</td>
<td>0.59–2.26</td>
<td>0.68</td>
</tr>
<tr>
<td>5+</td>
<td>52 (11)</td>
<td>1.31</td>
<td>0.57–3.05</td>
<td>0.52</td>
</tr>
<tr>
<td>Smoking at 20 weeks of gestation</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>No</td>
<td>348 (72)</td>
<td>Ref</td>
<td>NA</td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>98 (20)</td>
<td>1.27</td>
<td>0.70–2.29</td>
<td>0.43</td>
</tr>
<tr>
<td>Medical complications&lt;sup&gt;b&lt;/sup&gt;</td>
<td></td>
<td></td>
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<tr>
<td>No</td>
<td>290 (60)</td>
<td>Ref</td>
<td>NA</td>
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<tr>
<td>Yes</td>
<td>190 (39)</td>
<td>0.80</td>
<td>0.46–1.38</td>
<td>0.43</td>
</tr>
<tr>
<td>Induction of labour</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>No</td>
<td>278 (58)</td>
<td>Ref</td>
<td>NA</td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>202 (42)</td>
<td>1.28</td>
<td>0.76–2.13</td>
<td>0.36</td>
</tr>
<tr>
<td>Mode of birth</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unassisted vaginal</td>
<td>236 (49)</td>
<td>Ref</td>
<td>NA</td>
<td></td>
</tr>
<tr>
<td>Assisted vaginal</td>
<td>25 (52)</td>
<td>1.40</td>
<td>0.32–6.05</td>
<td>0.65</td>
</tr>
<tr>
<td>Caesarean</td>
<td>220 (46)</td>
<td>1.43</td>
<td>0.84–2.42</td>
<td>0.84</td>
</tr>
</tbody>
</table>

CI, confidence interval; FPG, fasting plasma glucose; HbA1c, glycosylated haemoglobin; HR, hazards ratio; NA, not assessable; OGTT, oral glucose tolerance test; RPG, random plasma glucose.

<sup>a</sup> Accessibility/remoteness index of Australia code.

<sup>b</sup> Medical complications existing prior to pregnancy.

<sup>c</sup> Missing/unclear data not included.
all tests after 1 March 2010). The date of type 2 diabetes diagnosis was calculated from the data of any records identifying type 2 diabetes from any data source (e.g. medical records, MPDC data and laboratory tests). The Cox-proportional hazards regression models were extended using interaction terms to investigate possible effect modification of time to type 2 diabetes diagnosis between Indigenous and non-Indigenous women, with likelihood ratio tests used to calculate a single p-value where there are multiple categories of continuous variables. Throughout, two-tailed tests were conducted, and p < 0.05 was considered statistically significant.

**Ethics**

This project was conducted with assistance from Cairns Diabetes Centre and Apunipima Cape York Health Council. Governance was provided by a project advisory group that included Indigenous researchers and representatives of Apunipima Cape York Health Council (community-controlled organization). Ethical approval was granted for this project by the Cairns Hospital and Hinterland Research Ethics Committee, the Monash University Human Research and Ethics Committee, and the Queensland Health Research Ethics and Governance Unit (no. 201101190).

**Results**

From 1 January 2004 to 31 December 2010, 1012 women were identified in the CHCSC as giving birth at Cairns Hospital and coded as having an episode of gestational diabetes, from a total of 16,765 births during the same period. This includes 352 Indigenous women and 660 non-Indigenous women. These women had 1,505 births during the study period, and 1,098 of these births were coded as gestational diabetes after linkage with MPDC data and medical record review of 912 pregnancies (crude gestational diabetes prevalence = 6.5%). Approximately 6% of births where medical records were reviewed were reclassified as gestational diabetes, and hence, gestational diabetes case ascertainment is likely to be underestimated by approximately 37 cases in this sample, detailed elsewhere [22]. Two women who died in the early postpartum period were excluded from analysis. Postpartum screening tests were available following 483/1098 births coded as gestational diabetes, and analysis of type 2 diabetes progression was restricted to this subset with postpartum tests recorded (Figure 1). Compared with women who did not receive postpartum screening, Indigenous women who received postpartum screening were more likely to live in remote areas, and non-Indigenous women who received screening were more likely to be born outside Australia, be older, have more than five previous pregnancies, not smoke or breastfeed [23]. There were no type 2 diabetes diagnoses recorded among births where no postpartum screening test was available. There were 82 type 2 diabetes diagnoses reported following births coded as having gestational diabetes: 76 identified by laboratory test results and six identified in clinical coding or medical records, where the subsequent pregnancy was coded as type 2 diabetes. As previously reported [22], Indigenous women had significantly longer times to postpartum screening compared with non-Indigenous women.

**Rates of progression from gestational diabetes to type 2 diabetes**

Among women who received a postpartum screening test, Indigenous women with gestational diabetes had a greater than fourfold risk of developing type 2 diabetes [hazards ratio (HR) 4.55, 95% confidence interval 2.63–7.88, p < 0.0001].

By 3 years postpartum, among the 498/1098 gestational diabetes births where postpartum screening tests were recorded, 21.9% (15.8–30.0%) Indigenous women and 4.2% (2.5–7.2%) non-Indigenous women were diagnosed with type 2 diabetes following gestational diabetes. By 5 years postpartum, 25.5% (18.6–34.3%) Indigenous and 5.7% (3.3–9.5%) non-Indigenous women were diagnosed with type 2 diabetes following gestational diabetes. And by 7 years postpartum, 42.4% (29.6–58.0%) Indigenous women and 13.5% (7.3–24.2%) non-Indigenous women were diagnosed with type 2 diabetes following gestational diabetes (Figure 2).

**Sensitivity analysis**

Sensitivity analyses including all eligible women with and without postpartum screening in the denominator continued to show almost a fourfold risk of developing type 2 diabetes among Indigenous women, compared with non-Indigenous women (HR 3.86, 95% confidence interval 2.24–6.78, p < 0.0001). However, as would be expected given that less than 50% women received a postpartum screening test, if all women are included in the denominator, the proportions of women progressing to type 2 diabetes were less than half that reported when only women receiving screening were included. By 3 years postpartum, 9.6% (6.8–13.4%) Indigenous women and 2.0% (1.2–3.4%) non-Indigenous women were diagnosed with type 2 diabetes. By 5 years postpartum, 11.2% (8.0–15.5%) Indigenous women and 2.7% (1.6–4.5%) non-Indigenous women were diagnosed with type 2 diabetes.
And by 7 years postpartum, 18.3% (12.6–26.3%) Indigenous women and 6.4% (3.4–11.7%) non-Indigenous women were diagnosed with type 2 diabetes postpartum (Supporting Information Figure 1).

Factors associated with rates of progression from gestational diabetes to type 2 diabetes

Indigenous status remained strongly associated with a faster time to progression from gestational diabetes to type 2 diabetes in all multivariate regression analyses (p < 0.0001).

In combined analysis, there appeared to be an increased rate of progression from gestational to type 2 diabetes among women with a BMI >25 kg/m² and women partially breastfeeding at discharge from hospital (Table 3). As would be expected, there was also a significantly increased ‘risk of progression’ among women who were first diagnosed with gestational diabetes prior to 17 weeks gestation and were likely to have undiagnosed type 2 diabetes in pregnancy (Table 3).

Among Indigenous women, there was an increased rate of type 2 diabetes progression among women partially breastfeeding, compared with women fully breastfeeding, at discharge from hospital and among women diagnosed with gestational diabetes prior to 17 weeks gestation (‘probable type 2 diabetes’). Despite higher rates in comparison with non-Indigenous women, there were no significant differences in type 2 diabetes progression rates between women who were coded as Aboriginal (HR 3.56, 1.89–6.78), Torres Strait Islander (HR 5.63, 2.92–10.83) or Aboriginal and Torres Strait Islander (HR 8.18, 2.78–24.08) (Table 2; Supporting Information Figure 2).

Among non-Indigenous women, significantly higher rates of type 2 diabetes progression were seen among women who were partially breastfeeding at discharge, compared with those who were fully breastfeeding at discharge from hospital, and women diagnosed with gestational diabetes prior to 17 weeks gestation (‘probable type 2 diabetes’) (Tables 2 and 3).
Table 3. Associations with progression time from gestational diabetes to type 2 diabetes, by Indigenous status, among gestational diabetes births with medical records reviews

<table>
<thead>
<tr>
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</thead>
<tbody>
<tr>
<td></td>
<td>n (%)&lt;sup&gt;b&lt;/sup&gt; HR CI p</td>
<td>n (%)&lt;sup&gt;b&lt;/sup&gt; HR CI p</td>
<td>n (%)&lt;sup&gt;b&lt;/sup&gt; HR CI p</td>
<td></td>
</tr>
<tr>
<td>Body mass index (kg/m²)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>18–24 (ref)</td>
<td>61 (21) Ref</td>
<td>26 (17)</td>
<td>35 (26)</td>
<td></td>
</tr>
<tr>
<td>&lt;18</td>
<td>1 (0) NA</td>
<td>1 (1) NA</td>
<td>0 (0)</td>
<td></td>
</tr>
<tr>
<td>25–29</td>
<td>76 (26) 3.16 1.01–9.86 0.05</td>
<td>33 (21) 2.17 0.65–7.26 0.21</td>
<td>43 (32) NA</td>
<td></td>
</tr>
<tr>
<td>30+</td>
<td>136 (47) 2.73 0.95–7.85 0.06</td>
<td>88 (57) 2.19 0.76–6.28 0.15</td>
<td>48 (36) NA</td>
<td>0.32</td>
</tr>
<tr>
<td>Primary antenatal care location</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>GP</td>
<td>11 (4) Ref</td>
<td>5 (3)</td>
<td>6 (4)</td>
<td></td>
</tr>
<tr>
<td>Hospital</td>
<td>242 (84) 1.74 0.24–12.71 0.59</td>
<td>128 (83) 1.32 0.18–9.73 0.78</td>
<td>114 (85) NA</td>
<td>0.28</td>
</tr>
<tr>
<td>Government clinics</td>
<td>1 (0) NA</td>
<td>1 (1) NA</td>
<td>0 (0)</td>
<td></td>
</tr>
<tr>
<td>Community-controlled organization</td>
<td>3 (1) 4.98 0.45–55.55 0.19</td>
<td>2 (1) 4.35 0.39–48.47 0.23</td>
<td>1 (1) NA</td>
<td></td>
</tr>
<tr>
<td>Postnatal care location</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>GP</td>
<td>140 (48) Ref</td>
<td>29 (19)</td>
<td>111 (83)</td>
<td></td>
</tr>
<tr>
<td>Hospital</td>
<td>2 (1) NA</td>
<td>1 (1) NA</td>
<td>0 (0)</td>
<td></td>
</tr>
<tr>
<td>Government clinics</td>
<td>114 (39) 1.63 0.69–3.83 0.26</td>
<td>98 (63) 2.09 0.73–6.04 0.17</td>
<td>16 (12) 1.42 0.17–12.20 0.75</td>
<td>0.71</td>
</tr>
<tr>
<td>Community-controlled organization</td>
<td>19 (7) 1.68 0.55–5.14 0.36</td>
<td>19 (12) 2.16 0.67–7.67 0.61</td>
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</tr>
<tr>
<td>Breastfeeding at discharge</td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Fully</td>
<td>217 (75) Ref</td>
<td>116 (75)</td>
<td>101 (75)</td>
<td></td>
</tr>
<tr>
<td>Partial</td>
<td>51 (18) 2.34 1.23–4.47 0.009</td>
<td>27 (17) 2.01 0.99–4.10 0.05</td>
<td>24 (18) 12.92 1.34–124.30 0.02</td>
<td>0.22</td>
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<tr>
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<td>234 (81) Ref</td>
<td>117 (75) Ref</td>
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<td>21 (7) 5.38 2.56–11.37 &lt;0.0001</td>
<td>13 (8) 4.76 2.09–10.84 &lt;0.0001</td>
<td>8 (6) 7.76 1.42–42.47 0.02</td>
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<td>25 (16) 2.10 0.96–4.56 0.06</td>
<td>9 (7) NA</td>
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</table>

CI, confidence interval; GP, general practitioner; HR, hazards ratio; NA, not assessable.
<sup>a</sup>Assessed as glucose intolerance diagnosed prior to 17 weeks gestation.
<sup>b</sup>Missing/unclear data not included.
Discussion

Among women with gestational diabetes who received a laboratory-based postpartum screening test, we found a greater than fourfold risk of progression to type 2 diabetes among Indigenous women compared with non-Indigenous women. Within 5 years postpartum, over 25% of Indigenous women with gestational diabetes who received a laboratory-based postpartum diabetes screening test had been diagnosed with type 2 diabetes, compared with almost 6% of non-Indigenous women. There were significantly higher rates of type 2 diabetes progression among women who had a pregnancy BMI >25 compared with those with a BMI <25, those who were ‘partially breastfeeding’, compared with those who were ‘fully breastfeeding’ at discharge from hospital, and women diagnosed with gestational diabetes prior to 17 weeks gestation (‘probable type 2 diabetes’).

These findings of increased rates of type 2 diabetes progression among Indigenous Australian women are consistent with previous reports of higher rates of type 2 diabetes among Indigenous women internationally [7–11]. Associations with increased risk of type 2 diabetes and both increased pregnancy BMI and not fully breastfeeding at discharge from hospital are also similar to findings reported among non-Indigenous women [4,24].

Major strengths of this study include the use of linked data validated by medical record review to increase the accuracy of gestational diabetes diagnosis coding and restriction of the sample to women who received postpartum diabetes screening tests. However, there are several limitations. First, we were unable to include ‘point of care’ tests provided by primary healthcare professionals, such as HbA1c, FPG and random plasma glucose, using a capillary sample. We identified six type 2 diabetes diagnoses in coding of subsequent pregnancies and medical records, in addition to the 76 identified by laboratory test records. This suggests that it is possible that some additional cases of type 2 diabetes diagnoses may have been missed (diagnosed in other locations or by ‘point-of-care-tests’), particularly where there was not a subsequent pregnancy, which would increase the observed rate of progression to type 2 diabetes. However, standard practice in this region is that if glucose intolerance is suspected, a venous sample would be sent to one of the three laboratories included in this study, and all women diagnosed with type 2 diabetes in this study had received postpartum diabetes screening. Second, under-enumeration of Indigenous status recording, with data linkage studies suggesting Indigenous mothers in the NSW MPDC, may be under-enumerated by around 40% [25]. Third, we included women who were diagnosed with gestational diabetes prior to 17 weeks gestation, and these women were highly likely to have had undiagnosed type 2 diabetes prior to pregnancy, reflected in the significantly higher risk of ‘progression’, and may be increasing the reported risk of progression in this study. Finally, we do not know the rates of progression among women who did not receive a postpartum screening test, and it is difficult to estimate the direction of the effect of this ‘selection bias’ and whether type 2 diabetes progression rates would be higher or lower among women who did not receive screening. For instance, women in this study who partially breastfed their infants around the time of hospital discharge had higher rates of type 2 diabetes progression than women who fully breastfed, and we had previously reported that ‘fully breastfeeding’ was associated with a higher likelihood of postpartum screening among non-Indigenous women [23]; hence, inclusion of more women partially breastfeeding could result in a higher observed rate of type 2 diabetes progression. On the other hand, one could argue that clinicians may be more strongly encourage screening among women whom they consider ‘at high risk’ and that those women who were not screened may have been more likely to be considered ‘at low risk’; hence, inclusion of more ‘low risk’ women would have decreased the observed rate of type 2 diabetes progression. This highlights the need for prospective research that follows up all women with gestational diabetes.

If the rates of progression to type 2 diabetes among women who received postpartum screening are extrapolated to all women in this study who did not receive postpartum screening, by 7 years postpartum, 20 Indigenous women and 13 non-Indigenous women who did not receive postpartum diabetes screening would have undiagnosed and untreated type 2 diabetes. In this study, we contacted the medical service providers for all women who were identified as being due for a postpartum screening test and advised them to offer their clients a postpartum screening test for diabetes if their records did not already indicate that they had received one. These findings highlight the importance of postpartum screening for type 2 diabetes after gestational diabetes, particularly among Indigenous women. This is of particular concern as we have previously reported very low rates of postpartum screening among all women with gestational diabetes in Far North Queensland, with significantly longer time to and lower rates of early postpartum screening among Indigenous women compared with non-Indigenous women [22] and with lower rates among Indigenous women in Cairns compared with women living in remote areas [23]. While this poses a serious risk for women in the longer term, the risks to women of childbearing age include serious risks to subsequent pregnancies, including congenital anomalies.

We have discussed the barriers to postpartum screening previously [23], including issues for women such as
Type 2 Diabetes after Gestational Diabetes

forgetting, not knowing about the need for a test, inconvenient unpleasant tests, fear of results, time pressures, physician awareness and communication, costs and lack of consistency around postpartum screening guidelines after gestational diabetes for Indigenous women [26]. Recent studies show that some systemic strategies can improve postpartum glucose screening rates among non-Indigenous women [27]. These include case management [28], patient and physician reminders, system changes, proactive postpartum care plans, antenatal education, registers, clinical protocols and electronic records [27]. The effectiveness of these strategies has led to calls for them to be a part of routine postpartum care for women with gestational diabetes [29]. However, given the low rates of postpartum screening among Indigenous women [22], there is a need to assess the specific needs of women and assess whether cultural tailoring or additional strategies are required.

While there has been considerable debate about gestational diabetes care and treatment pathways during pregnancy, guidelines for postpartum care for women at high risk of developing type 2 diabetes have lagged behind [30], and opportunities for prevention are being lost. A diagnosis of gestational diabetes offers a unique ‘window of opportunity’ to identify women at increased risk of developing type 2 diabetes and offer effective support. The increased insulin resistance occurring as a result of naturally occurring pregnancy hormones offers a ‘natural stress test’ to identify women with impaired glucose tolerance, who may have not been identified had they not become pregnant. Further, frequent scheduled contacts with health services during and after pregnancy provide an additional opportunity to support women to reduce the risk of developing type 2 diabetes for themselves and their infant, by breastfeeding and healthy lifestyle changes, with any effective support having significant benefits for many generations to follow [31]. Finally, studies suggest that women with gestational diabetes are likely to be highly motivated during the postpartum period to improve the health of their infant and family, with a heightened state of ‘change readiness’ [32]. However, while pregnancy and the postpartum period afford opportunities to reduce the risk of type 2 diabetes, there are also a number of challenges including tiredness, maternal attachment and childcare demands in the early postpartum period, work, family and child development in subsequent years [33] and postpartum depression [34]. Several recent studies also suggest that pregnancy care providers lack confidence in talking to women about obesity-related issues [35].

Hence, well-designed strategies to address the needs of women and ensure provision of professional support are essential [36]. Results in this study reinforce calls for supporting a healthy BMI and breastfeeding as important prevention strategies [37]. A recent meta-synthesis of qualitative studies of ‘perceptions of women with gestational diabetes’ suggested that important factors to overcome these challenges included addressing emotional issues, providing clear advice and offering an intervention that works for the whole family [38]. And while technological interventions have been trialled as convenient options [39], it is likely that personal support will be needed to address emotional issues [40,41]. This is likely to be particularly important among Indigenous women with gestational diabetes, as a study in the United States found that Indigenous women with gestational diabetes reported high perceptions of risk coupled with a low sense of self-efficacy [42], a combination associated with avoidance behaviour [43]. This raises questions as to whether traditional risk advice is likely to be helpful and suggests that a strength-based focus that improves self-efficacy may be more effective. This reinforces recommendations in the National Aboriginal and Torres Strait Islander Health Plan [44], which was developed to guide evidence-based strategies to ‘close the gap’ in health inequities, for strength-based family-centred interventions using life-course approaches that focus on social determinants.

Despite the risks of developing type 2 diabetes among women with previous gestational diabetes, there are few studies demonstrating effective strategies to reduce the risks for women after gestational diabetes [30,45]. Importantly, there are no studies investigating strategies to improve postpartum prevention or screening among Indigenous women with gestational diabetes in Australia [46]. This reflects a lack of diabetes intervention research [47] and care [48] among Indigenous peoples more generally. Evidence of effective strategies to prevent gestational diabetes among Indigenous women, and to improve care during and after pregnancy for Indigenous women with gestational diabetes, is urgently needed. This includes collaborative mixed methods research translation strategies to develop strategies to improve postpartum care for Indigenous women with gestational diabetes and reduce the risks of implementation with limited evidence. Strategies should also be contextually and culturally relevant and, where possible, address broader socio-ecological factors [49]. The research must be underpinned by Indigenous research principles, driven by identified needs and include active community engagement.

Conclusions

Evidence of effective strategies to improve postpartum screening and to reduce the risk of developing type 2 diabetes postpartum, including strength-based approaches
for supporting exclusive breastfeeding and healthy lifestyles, are urgently needed. This is particularly important for Indigenous women who have low rates of postpartum screening and more than a fourfold risk of developing type 2 diabetes, compared with non-Indigenous women.

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Author contributions

C.C. proposed the design, developed the methods, prepared and submitted ethics applications, developed data collection tools, linked data and prepared the draft manuscript. B.O. and S.E. contributed to study design. R.W. provided statistical advice and assistance. All authors contributed to and approved the final manuscript.

Conflicts of interest

None declared.

References

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   **How to use it**
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   **How to use it**
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