Neighborhood walkability and risk of gestational diabetes

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ABSTRACT

Objective Higher neighborhood walkability has been associated with a lower risk of type 2 diabetes mellitus (T2DM) by promoting greater physical activity (thereby reducing weight and lowering insulin resistance). However, it is not known if walkability may similarly reduce maternal risk of gestational diabetes mellitus (GDM), which arises in the setting of the severe physiologic insulin resistance of pregnancy. Indeed, the insulin resistance of pregnancy is primarily driven by placental hormones and not maternal weight gain. Thus, we sought to evaluate the impact of neighborhood walkability on maternal risk of GDM and the pathophysiologic determinants thereof (insulin sensitivity and pancreatic beta-cell function).

Methods In this study, 1318 women reported their pregravid physical activity (Baekcke questionnaire) while undergoing an oral glucose tolerance test (OGTT) at mean 29.3 weeks’ gestation. The OGTT identified 290 women with GDM and enabled assessment of insulin sensitivity and beta-cell function. Based on their residential Walk Score, the women were stratified into the following four established categories of neighborhood walkability: car dependent (n=328), somewhat walkable (n=315), very walkable (n=406), and walker’s paradise (n=269).

Results There was a progressive increase in pregravid total physical activity (p=0.002), non-sport leisure-time activity (p=0.009) and sport activity (p=0.01) across the walkability groups (from car dependent to somewhat walkable to very walkable to walker’s paradise), coupled with a concomitant decline in pre-pregnancy body mass index (p=0.007). However, in pregnancy, the groups did not differ in gestational weight gain (p=0.80). Moreover, the walkability groups also did not differ in mean adjusted insulin sensitivity, beta-cell function, or glycermia on the antepartum OGTT. On logistic regression analysis, Walk Score did not predict GDM (OR=1.001, 95% CI 0.995 to 1.007).

Conclusion Neighborhood walkability is not a significant determinant of maternal risk of GDM. Thus, in contrast to T2DM, the effect of neighborhood design on incidence of GDM will be comparatively modest.

INTRODUCTION

Type 2 diabetes mellitus (T2DM) is a global epidemic affecting more than 425 million people worldwide, as per latest estimates from the International Diabetes Federation (IDF).1 As a health problem of this magnitude arguably requires a societal solution(s), there is growing interest in urban planning as a means of promoting healthy lifestyle practices and thereby addressing this epidemic.2 In particular, higher neighborhood walkability has been associated with a lower prevalence of overweight/obesity and lower risk of diabetes.2–5 The presumed mechanism underlying these associations is that neighborhood walkability encourages a more active lifestyle that reduces the likelihood of excessive weight gain over time and thereby limits obesity-associated insulin resistance that would otherwise contribute to the development of T2DM in those individuals in whom pancreatic beta-cell compensation would be insufficient to fully mitigate such a challenge.

In parallel with the global burden of T2DM, there is also a rising incidence of hyperglycemia in pregnancy. Indeed, according to the IDF, the proportion of pregnancies affected...
by maternal hyperglycemia worldwide increased from one in seven in 2011 to one in six pregnancies in 2015.1 Moreover, this rising prevalence potentially may portend a transgenerational future burden of disease in those individuals in whom intrauterine exposure to maternal hyperglycemia may have adverse consequences for developmental programming of metabolic pathways (as per the Developmental Origins of Health and Disease paradigm).6 Thus, if enhanced neighborhood walkability could lower the risk of gestational diabetes mellitus (GDM), then the beneficial effects could be relevant to both mother and child, and thereby provide a transgenerational approach to the diabetes epidemic. However, the impact of neighborhood walkability on risk of GDM is not known.

Although both GDM and T2DM are characterized by the same general pathophysiologic basis (insufficient beta-cell compensation for ambient insulin resistance),7 there are important differences in their respective settings that may relate to the extent to which neighborhood walkability could be beneficial. Specifically, in contrast to the non-gravid state, pregnancy is a discrete 9-month interval during which some degree of weight gain is both physiologic and necessary.8 Second, the progressive insulin resistance that characterizes the second half of pregnancy is physiologic and is primarily mediated by placental hormones (ie, rather than weight gain as in the non-gravid state). Third, compared with the impact of weight gain on insulin sensitivity outside of pregnancy, the normal physiologic insulin resistance of the latter half of gestation is of such a profound magnitude that its capacity for modification by urban design potentially might be limited. Thus, it is quite unclear whether enhanced walkability could have a beneficial effect on GDM, in the way that it does for T2DM. In this context, our objective in this study was to evaluate the impact of neighborhood walkability on maternal risk of GDM and the pathophysiologic determinants thereof (insulin sensitivity/resistance and beta-cell function).

METHODS

The study population consisted of pregnant women in Toronto, Canada, who were recruited to undergo metabolic characterization at the time of antepartum screening for GDM. The study protocol has been previously described in detail.9 At our institution, all pregnant women undergo universal GDM screening at 24–28 weeks’ gestation with a 50 g glucose challenge test (GCT), which is then followed by a diagnostic oral glucose tolerance test (OGTT) if the GCT result is abnormal (blood glucose ≥7.8 mmol/L at 1 hour after challenge). For this study, healthy pregnant women are recruited either prior to or just after their GCT. Regardless of the GCT result (ie, even if normal), all study participants undergo a 3-hour 100 g OGTT for determination of GDM status. As previously described,9 the recruitment of women after an abnormal GCT serves to enrich the study population for those with GDM. The current analysis was performed in women living in Toronto for whom complete home address information was provided (n=1318).

Assessment of study participants

On the morning of the 3-hour 100 g OGTT in pregnancy, data regarding medical, obstetrical, and family history were collected by interviewer-administered questionnaire. Pregravid physical activity in the year before pregnancy was assessed using the Baecke questionnaire, an established instrument that has been extensively validated in several populations, including women of childbearing age.10 11 The Baecke questionnaire measures total physical activity and its three component domains: (1) occupation-associated activity (work index), (2) sport-related physical activity (sport index), and (3) non-sport leisure-time activity (leisure-time index). The work index quantifies the exertion related to occupational activities, such as sitting, standing, lifting, and walking, as well as associated effects on the individual (eg, fatigue, perspiration). The sport index characterizes vigorous/sports activity with respect to intensity (using the compendium of physical activities),12 duration, and frequency. The leisure-time index quantifies exertion associated with non-sport recreational activities (eg, walking and television viewing). Participants completed the Baecke questionnaire during their OGTT (ie, prior to knowledge of GDM status). GDM was diagnosed according to National Diabetes Data Group (NDDG) criteria13 which require at least two of the following on the OGTT: fasting blood glucose ≥5.8 mmol/L, 1-hour blood glucose ≥10.6 mmol/L, 2-hour blood glucose ≥9.2 mmol/L, or 3-hour blood glucose ≥8.1 mmol/L.

Metabolic characterization on OGTT

All OGTTs were performed in the morning after overnight fast, with venous blood samples drawn for the measurement of glucose and specific insulin at fasting and at 30, 60, 120 and 180 min following the ingestion of the 100 g glucose load. Specific insulin was measured with the Roche Elecsys 1010 immunoassay analyzer and electrochemiluminescence immunoassay kit (Roche Diagnostics, Laval, Canada). Area-under-the-glucose curve on the OGTT (AUCglc) was calculated by trapezoidal rule. Insulin sensitivity was measured with the Matsuda index, an established measure of whole-body insulin sensitivity that has been validated against the euglycemic-hyperinsulinemic clamp.14 Hepatic insulin resistance was assessed with the homeostasis model assessment of insulin resistance (HOMA-IR).15 Beta-cell function was assessed with the insulin secretion-sensitivity index-2 (ISSI-2), an OGTT-derived measure that is analogous to the disposition index from the intravenous glucose tolerance test against which it has been validated.16 17 ISSI-2 is defined as the product of (1) insulin secretion measured from the intravenous glucose tolerance test, against which it has been validated.16 17 ISSI-2 is defined as the product of (1) insulin secretion measured from the intravenous glucose tolerance test, against which it has been validated.16 17 ISSI-2 is defined as the product of (1) insulin secretion measured from the intravenous glucose tolerance test, against which it has been validated.16 17 ISSI-2 is defined as the product of (1) insulin secretion measured from the intravenous glucose tolerance test, against which it has been validated.16 17 ISSI-2 is defined as the product of (1) insulin secretion measured from the intravenous glucose tolerance test, against which it has been validated.16 17 ISSI-2 is defined as the product of (1) insulin secretion measured from the intravenous glucose tolerance test, against which it has been validated.16 17 ISSI-2 is defined as the product of (1) insulin secretion measured from the intravenous glucose tolerance test, against which it has been validated.16 17
(insulinogenic index/HOMA-IR) provided a second established measure of beta-cell function (with insulinogenic index defined as the ratio of the incremental change in insulin during the first 30 min of the OGTT to the incremental change in glucose over the same time period).9

**Determination of neighborhood walkability**

The walkability of each woman’s residential neighborhood was assessed using Walk Score, which is an established index that reflects the walkability of an address using an algorithm that considers its distance to 13 amenities (grocery stores, coffee shops, restaurants, bars, movie theaters, schools, parks, libraries, bookstores, fitness centers, drugstores, hardware stores, clothing/music stores) and aspects of pedestrian friendliness (such as population density, block length, and intersection density). Walk Score is a validated measure18–20 that has been associated with cardiometabolic risk factors such as physical activity, weight and blood pressure,21–24 including previous studies in Toronto. It yields a numerical score that ranges from 0 to 100, from which categories of walkability have been previously established, ranging from ‘car-dependent’ (Walk Score <50) to ‘somewhat walkable’ (Walk Score 50–69 inclusive) to ‘very walkable’ (Walk Score 70–89 inclusive) to ‘walker’s paradise’ (Walk Score 90–100 inclusive). We determined the walkability for each participant’s home address using the publicly available Walk Score interface (https://www.walkscore.com/).

**Statistical analyses**

All analyses were conducted using SAS V.9.2 (SAS Institute). Walk Score was analyzed as both a continuous and categorical variable. Based on their home address Walk Score, the study participants were first stratified into four previously established Walk Score categories reflecting neighborhood walkability: (1) car-dependent (Walk Score <50), (2) somewhat walkable (Walk Score 50–69 inclusive), (3) very walkable (Walk Score 70–89 inclusive) and (4) walker’s paradise (Walk Score 90–100 inclusive). Continuous variables were tested for normality of distribution, and natural log transformations of skewed variables were used, where necessary, in subsequent analyses. The demographic and clinical characteristics of the four groups were compared by analysis of variance (ANOVA) for continuous variables, or either χ² or Fisher’s exact test for categorical variables. Mean values of non-sport leisure-time index, vigorous/sport index, work index, total physical activity score, pre-pregnancy body mass index (BMI), and weight gain in pregnancy were compared across the four Walk Score groups by ANOVA. In addition, pairwise comparisons with Bonferroni adjustment were performed to determine if significant differences existed between any pairs of groups.

Spearman correlation analysis was performed to evaluate univariate correlations between continuous Walk Score and (1) pregravid features (physical activity, BMI), and (2) metabolic function in pregnancy, respectively. Multiple linear regression analyses were performed to compare the four Walk Score groups with respect to mean adjusted values for the following metabolic outcomes in pregnancy: insulin sensitivity/resistance (Matsuda index, HOMA-IR), beta-cell function (ISSI-2, insulinogenic index/HOMA-IR), and glycemia (fasting glucose, AUCglucose on the OGTT). These comparisons were performed after adjustment for age, ethnicity, family history of diabetes, parity, pre-pregnancy BMI, weight gain in pregnancy up to the OGTT, and weeks’ gestation at the OGTT. For each of these six metabolic outcomes, we conducted sensitivity analyses (1) by further adjusting for fetal sex and (2) by replacing Walk Score groups with continuous Walk Score. In addition, we conducted sensitivity analyses of these comparisons of mean adjusted metabolic outcomes in (1) women of white ethnicity, (2) nulliparous women, and (3) women who had no family history of diabetes (i.e., reflecting variables that differed between the four Walk Score groups). Logistic regression analysis was performed to evaluate whether continuous Walk Score was an independent predictor of GDM after adjustment for the same covariates as in the multiple linear regression analyses.

**RESULTS**

**Comparison of Walk Score groups**

The study population of 1318 women underwent the antepartum OGTT at mean 29.3 weeks’ gestation, which identified 290 women as having GDM. Table 1 shows demographic and clinical characteristics of the study population stratified into four Walk Score groups reflecting different degrees of neighborhood walkability: (1) car dependent (n=328), (2) somewhat walkable (n=315), (3) very walkable (n=406), and (4) walker’s paradise (n=269). These groups did not differ with respect to age or smoking. The lower walkability groups had a higher proportion of women of white ethnicity (p=0.0006) and higher parity (p=0.0004), coupled with a slightly greater prevalence of family history of diabetes (p=0.02).

Figure 1 shows the comparisons of the Walk Score groups with respect to physical activity in the year prior to the pregnancy, pre-pregnancy BMI, and gestational weight gain up to the OGTT. As shown in figure 1A, mean non-sport leisure-time activity progressively increased across the four groups as the neighborhoods became more walkable (p=0.009). A similar but milder gradient was seen for the vigorous/sport index (figure 1B, p=0.01). As anticipated, there was no difference across the groups in work index (figure 1C, p=0.49). Overall, there was a clear stepwise increase in pregravid total physical activity from the car-dependent group to somewhat walkable to very walkable to walker’s paradise (figure 1D, p=0.002). This pattern translated to an analogous decrease in pre-pregnancy BMI from the less walkable to more walkable groups (figure 1E, p=0.007), with mean pre-pregnancy
**Table 1** Characteristics of study population, stratified according to Walk Score category as follows: (1) car dependent, (2) somewhat walkable, (3) very walkable, and (4) walker’s paradise

| Characteristics | Car dependent (Walk Score <50) | Somewhat walkable (Walk Score 50–69) | Very walkable (Walk Score 70–89) | Walker’s paradise (Walk Score 90–100) | P value |
|----------------|-------------------------------|--------------------------------------|----------------------------------|--------------------------------------|---------|
| Walk Score mean (SD) | 31.4±13.1 | 60.0±5.9 | 80.6±5.7 | 94.5±3.0 |         |
| Walk Score median (IQR) | 33 (22–43) | 60 (55–65) | 81 (76–86) | 94 (92–97) |         |
| Age (years) | 34±4 | 34±5 | 34±5 | 34±4 | 0.43 |
| Ethnicity | | | | | 0.0006 |
| White, n (%) | 236 (72.0) | 224 (71.1) | 264 (65.0) | 149 (55.4) |         |
| Asian, n (%) | 41 (12.5) | 40 (12.7) | 61 (15.0) | 58 (21.6) |         |
| Other, n (%) | 51 (15.5) | 51 (16.2) | 81 (20.0) | 62 (23.0) |         |
| Family history of DM, n (%) | 221 (69.9) | 198 (65.4) | 232 (59.2) | 156 (61.2) | 0.02 |
| Smoking | | | | | 0.86 |
| Never, n (%) | 224 (68.3) | 226 (72.0) | 283 (69.7) | 190 (70.6) |         |
| Remote, n (%) | 96 (29.3) | 84 (26.7) | 112 (27.6) | 73 (27.2) |         |
| Current, n (%) | 8 (2.4) | 4 (1.3) | 11 (2.7) | 6 (2.2) |         |
| Parity | | | | | 0.0004 |
| Nulliparous, n (%) | 154 (47.0) | 151 (48.0) | 231 (56.9) | 172 (63.9) |         |
| 1, n (%) | 128 (39.0) | 121 (38.7) | 137 (33.7) | 72 (26.8) |         |
| >1, n (%) | 46 (14.0) | 42 (13.3) | 38 (9.4) | 25 (9.3) |         |
| Previous GDM, n (%) | 18 (5.7) | 23 (7.5) | 29 (7.5) | 7 (2.9) | 0.07 |
| Pre-pregnancy BMI class | | | | | 0.14 |
| Obese, n (%) | 61 (18.8) | 55 (17.9) | 52 (13.4) | 36 (13.6) |         |
| Overweight, n (%) | 77 (23.8) | 74 (24.0) | 94 (24.2) | 51 (19.3) |         |
| Normal, n (%) | 186 (57.4) | 179 (58.1) | 242 (62.4) | 177 (67.1) |         |
| Gestational hypertension | 10 (3.1) | 12 (3.8) | 11 (2.7) | 2 (0.7) | 0.09 |

Bold indicates p≤0.001.

Age is presented as mean±SD. Pre-pregnancy BMI classes were defined as follows: (1) obese: BMI ≥30 kg/m²; (2) overweight: 25≤BMI<30 kg/m²; and (3) normal: BMI <25 kg/m².

BMI, body mass index; DM, diabetes mellitus; GDM, gestational diabetes mellitus.

BMI in each group as follows: (1) car dependent: 25.3 kg/m²; (2) somewhat walkable: 25.6 kg/m²; (3) very walkable: 24.6 kg/m²; and (4) walker’s paradise: 24.2 kg/m². Despite these pregravid differences, however, gestational weight gain up to the OGTT did not differ across the groups (figure 1F, p=0.80).

**Associations of Walk Score with metabolic outcomes in pregnancy**

We next sought to evaluate the associations of neighborhood walkability with metabolic function in pregnancy. On Spearman correlation analysis (table 2), higher Walk Score (as a continuous measure) was positively associated with pregravid total physical activity (r=0.10, p=0.0003), non-sport leisure-time index (r=0.09, p=0.001), and vigorous/sport index (r=0.09, p=0.0007), and negatively correlated with pre-pregnancy BMI (r=−0.11, p=0.0001). However, there was no association with gestational weight gain up to the OGTT. Moreover, Walk Score showed no evidence of any association with insulin sensitivity or resistance, beta-cell function or glycemia on the OGTT in pregnancy (all |r|<0.05, p non-significant). Thus, despite its associations with pregravid physical activity and BMI, neighborhood walkability did not appear to relate to glucose homeostasis in pregnancy. Accordingly, the prevalence of GDM did not differ between the four Walk Score groups (car dependent: 21.7%; somewhat walkable: 24.1%; very walkable: 23.9%; walker’s paradise: 17.1%; overall p=0.14). Similarly, the prevalence of fasting glucose ≥5.1 mmol/L on the OGTT in pregnancy did not differ between the Walk Score groups (car dependent: 24.9%; somewhat walkable: 23.9%; very walkable: 24.1%; walker’s paradise: 17.8%; overall p=0.28) (data not shown).

We next sought to determine whether neighborhood walkability was independently associated with any of the metabolic outcomes in pregnancy, after adjustment for covariates (age, ethnicity, family history of diabetes, parity, pre-pregnancy BMI, weight gain in pregnancy up to the OGTT, and weeks’ gestation at OGTT). As shown in table 3, the four walkability groups did not differ...
in mean adjusted insulin sensitivity (Matsuda index), insulin resistance (HOMA-IR), beta-cell function (ISSI-2, insulinogenic index/HOMA-IR), or glycemia (fasting glucose, AUC_glucose on the OGTT). Furthermore, these findings were unchanged on sensitivity analyses in which (1) there was further adjustment for fetal sex; (2) further adjustment for GDM in a previous pregnancy, and (3) Walk Score groups were replaced with continuous Walk Score (data not shown). In addition, these findings were unchanged on sensitivity analyses that were restricted to (1) women of white ethnicity, (2) nulliparous women, and (3) women who had no family history of diabetes (data not shown). Finally, on logistic regression analysis, Walk Score did not predict GDM (OR=1.001, 95% CI 0.995 to 1.007) (table 4A). It also did not predict GDM defined by the less stringent Carpenter and Coustan criteria (OR=1.003, 95% CI 0.998 to 1.008) (table 4B).

**DISCUSSION**

In this study of 1318 women, we demonstrate that higher neighborhood walkability is associated with greater pregravid physical activity and lower BMI. However, in pregnancy, it did not have effects on gestational weight gain, insulin sensitivity, beta-cell function or glycemia. Accordingly, neighborhood walkability was not associated with maternal risk of GDM.

In normal human gestation, women experience a physiologic progressive decline in insulin sensitivity from ~20 weeks onwards, resulting in marked insulin resistance by late pregnancy (~50%–70% fall in insulin sensitivity overall). In this setting, the maintenance of glucose homeostasis is dependent on the pancreatic beta-cells appropriately increasing their secretion of insulin to compensate for this degree of insulin resistance. GDM arises in a population of women in whom there exists a chronic defect in beta-cell function such that their beta-cell compensation for this challenge is insufficient (resulting in the hyperglycemia by which GDM is diagnosed). Thus, to lower the risk of GDM, an intervention (such as enhanced walkability) would need to reduce this marked insulin resistance to such an extent that beta-cell compensation in these at-risk women will now be sufficient to keep blood glucose levels in the non-diagnostic range. Accordingly, given the profound insulin resistance of late pregnancy, a modest improvement in insulin sensitivity per se may not lower the incidence of GDM, unless the improvement therein is of a sufficient (ie, prodigious) magnitude.

Previous studies have established Walk Score as a validated measure of neighborhood walkability that is...
associated with cardiometabolic risk factors (such as physical activity, obesity and blood pressure) in a variety of settings, including Toronto.21-24 In the current study, Walk Score was indeed associated with total physical activity in the year prior to pregnancy. Moreover, this relationship was driven by associations with the component domains of physical activity in the precise pattern that would be anticipated for the impact of neighborhood walkability (ie, primarily affecting non-sport leisure-time activity, with no effect on work index). Furthermore, higher neighborhood walkability was associated with lower pre-pregnancy BMI. Thus, the tracking of Walk Score with these pregravid factors was fully concordant with previous studies in non-pregnant individuals, thereby supporting the fidelity of this measure in capturing walkability in the current study. Despite these pregravid relationships, however, we show that neighborhood walkability did not have any associations with gestational weight gain or glucose homeostasis (insulin sensitivity, beta-cell function, or glycemia) in pregnancy. Indeed, besides not approaching statistical significance, the correlation coefficients between Walk Score and all of the glycemic measures were near 0 (all r<0.02), suggestive of no association at all (Table 2). Similarly, the estimated OR for Walk Score in predicting GDM (OR=1.001, 95% CI 0.995 to 1.007) was also indicative of no effect. Collectively, these data suggest that, in the face of the profound insulin resistance of pregnancy, any impact of neighborhood walkability on insulin sensitivity was too modest to have any discernible effect on glucose homeostasis in over 1300 women. As a reference for perspective, it is worth noting that the modest effect of fetal sex on maternal glucose homeostasis was readily detectable in 1074 women.9

Our finding of an apparent discordance in the metabolic impact of walkability between the pregnant and non-pregnant states is analogous to the difference that has been observed in the relative effects of lifestyle modification on the risks of T2DM and GDM, respectively. Indeed, outside of pregnancy, a series of studies has shown that lifestyle intervention can effectively reduce the incidence of T2DM in at-risk individuals (ie, those with pre-diabetes).26-28 Conversely, studies evaluating lifestyle interventions in early pregnancy to reduce the risk of subsequent GDM have been largely unsuccessful.30-33 Moreover, a recent meta-analysis34 from a consortium of lifestyle intervention trials in pregnancy showed that lifestyle modification can reduce gestational weight gain but not lower the incidence of GDM (possibly by not sufficiently mitigating the marked insulin resistance that characterizes the latter half of gestation). These data serve to underscore the concept that there are inherent differences in the association of weight gain and insulin resistance during pregnancy as compared with the non-gravid state. First, both weight gain and insulin resistance are normal physiologic features of pregnancy. Second, weight gain is not a prime determinant of insulin resistance in pregnancy (which is driven by placental hormones), in contrast to its impact in the non-gravid state. Third, the severity of normal physiologic insulin resistance in late gestation is of an overwhelming magnitude when compared with that which typically occurs with weight gain in non-pregnant individuals. When considered from this perspective, neighborhood walkability can be viewed as a non-targeted lifestyle intervention that can yield metabolic benefit prior to pregnancy but may be insufficient to markedly impact glucose homeostasis in the pregnant state. From the current data, we cannot

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### Table 3

Comparison of mean adjusted metabolic outcomes in pregnancy (Matsuda index, HOMA-IR, ISSI-2, IGI/HOMA-IR, fasting glucose, AUC\textsubscript{glucose} on OGTT) between Walk Score groups, after adjustment for the following covariates: age, ethnicity, family history of diabetes, parity, pre-pregnancy BMI, weight gain in pregnancy up to the OGTT, and weeks’ gestation at OGTT.

|                      | Car dependent (Walk Score <50) | Somewhat walkable (Walk Score 50–69) | Very walkable (Walk Score 70–89) | Walker’s paradise (Walk Score 90–100) | P value |
|----------------------|---------------------------------|--------------------------------------|----------------------------------|---------------------------------------|---------|
| Insulin sensitivity/resistance |                                |                                      |                                  |                                       |         |
| Matsuda index        | 3.66±0.12                       | 3.71±0.12                            | 3.51±0.11                        | 3.83±0.14                             | 0.19    |
| HOMA-IR              | 1.97±0.07                       | 1.90±0.07                            | 2.07±0.07                        | 1.95±0.07                             | 0.22    |
| Beta-cell function   |                                 |                                      |                                  |                                       |         |
| ISSI-2               | 715.4±15.4                      | 731.1±16.3                           | 681.9±13.8                      | 695.0±16.3                            | 0.04    |
| IGI/HOMA-IR          | 9.6±0.4                         | 10.0±0.5                             | 8.8±0.4                         | 9.2±0.5                              | 0.09    |
| Glycemia             |                                 |                                      |                                  |                                       |         |
| Fasting glucose      | 4.6±0.03                        | 4.6±0.03                             | 4.6±0.03                        | 4.6±0.04                              | 0.12    |
| AUC\textsubscript{glucose} | 23.14±0.24                     | 23.26±0.25                           | 23.51±0.23                      | 23.13±0.26                            | 0.53    |

Data are presented as adjusted mean±SEM.

AUC\textsubscript{glucose}, area-under-the-glucose curve; BMI, body mass index; HOMA-IR, homeostasis model assessment of insulin resistance; IGI, insulinogenic index; ISSI-2, insulin secretion-sensitivity index-2; OGTT, oral glucose tolerance test.

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determine if the lack of metabolic impact in pregnancy is due to gestational changes in behavior or because of physiologic differences between the gravid and non-gravid states (eg, in the severity and determinants of insulin resistance). Either way, however, these data suggest that neighborhood walkability is not a significant determinant of glucose homeostasis in pregnancy.

Strengths of this study include the population of 1318 women reflecting a broad spectrum of neighborhood walkability (ie, ~300 women in each of the four established categories), in whom pregravid physical activity was queried prior to knowledge of the outcome of the OGTT, on which 290 were then diagnosed with GDM. Importantly, neighborhood walkability was assessed with a validated measure in relation to home address and exhibited the anticipated associations with pregravid physical activity. Moreover, all of the women underwent systematic assessment of insulin sensitivity/resistance and beta-cell function, thereby providing pathophysiologic insight into how walkability might affect the risk of GDM. Conversely, a limitation of this study is that the observational design precludes definitive ascertainment of causality in observed associations. Additional limitations are the absence of data on pre-pregnancy clinical risk factors of previous impaired fasting glucose or A1c >5.7% and pre-pregnancy measurements of insulin sensitivity/resistance and beta-cell function. The absence of a measurement of physical activity in pregnancy is also a limitation. Lastly, the absence of an effect of walkability on risk of GDM defined by NDDG criteria does not rule out an impact on GDM defined by less stringent criteria such as those of the International Association of Diabetes in Pregnancy Study Groups (although tables 2 and 3 show no association of Walk Score with continuous measures of glycemia and Walk Score did not predict GDM by Carpenter and Coustan criteria in table 4).

It should be recognized that our data do not rule out any impact of walkability on risk of GDM (however small), particularly since neighborhood design will affect the entire population where even a modest effect can be beneficial for society. Accordingly, these data certainly do not argue against city planning as a societal approach to the T2DM epidemic (as enhanced walkability has been linked to cardiometabolic benefits outside of pregnancy). However, our inability to detect any discernible impact of neighborhood walkability on insulin sensitivity, beta-cell function and glycemia in 1318 pregnant women suggests that the overall effect of urban design on the incidence of GDM may be relatively modest (as compared with its benefits outside of pregnancy). That said, in women who develop GDM, one would anticipate that enhanced neighborhood walkability should still be beneficial in reducing their risk of progression to T2DM in the years after their pregnancy (as weight control is a key modifiable determinant of this risk).

In conclusion, neighborhood walkability is associated with greater physical activity and lower BMI prior to pregnancy but does not have a discernible impact on antepartum weight gain or glucose homeostasis when the same women become pregnant. Accordingly, neighborhood walkability is not a significant determinant of maternal risk of GDM. Taken together, these findings do not argue against urban planning as a means of promoting healthy lifestyle practices and reducing the risk of T2DM in the overall population. Rather, these data suggest that the effect of neighborhood design on the risk of GDM will be comparatively modest, likely reflecting the intrinsic physiologic differences between the gravid and non-gravid states.

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