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Phenotypic and genotypic differences between Indian and Scandinavian women with gestational diabetes mellitus

Short running title: Diabetes during pregnancy: India and Scandinavia

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Abstract

Objective: Gestational diabetes mellitus (GDM) is a transient form of diabetes characterized by impaired insulin secretion and action during pregnancy. Population-based differences in prevalence exist which could be explained by phenotypic and genetic differences. The aim of this study was to examine these differences in pregnant women from Punjab, India and Scandinavia.

Methods: 85 GDM/T2D loci in European and/or Indian populations from previous studies were assessed for association with GDM based on Swedish GDM criteria in 4018 Punjabi Indian and 507 Swedish pregnant women. Selected loci were replicated in Scandinavian cohorts, Radiel (N=398, Finnish), STORK/STORK-G (N=780, Norwegian).

Results: Punjabi Indian women had higher GDM prevalence, lower insulin secretion and better insulin sensitivity than Swedish women. There were significant frequency differences of GDM/T2D risk alleles between both populations. rs7178572 at *HMG20A*, previously associated with GDM in South Indian and European women was replicated in North Indian women. The T2D risk SNP rs11605924 in the *CRY2* gene was associated with increased GDM risk in Scandinavian but decreased risk in Punjabi Indian women. No other overlap was seen between GDM loci in both populations.

Conclusions: GDM is more common in Indian than Swedish women, which partially can be attributed to differences in insulin secretion and action. There was marked heterogeneity in the GDM phenotypes between the populations which could only partially be explained by genetic differences.

Introduction

Gestational diabetes mellitus (GDM) is defined as “carbohydrate intolerance resulting in hyperglycemia of variable severity with onset or first recognition during pregnancy, excluding those with diabetes in pregnancy likely to represent overt diabetes mellitus” (<http://www.who.int/>). GDM develops when women no longer can increase their insulin secretion to meet the increased demands of insulin resistance during the second and third trimester of pregnancy ¹. The risk of GDM is exacerbated by age, obesity, and a family history of GDM and type 2 diabetes (T2D) ²; however, the exact etiology is unknown. Women with GDM are at increased risk of gestational hypertension, pre-eclampsia, and T2D, as well as metabolic syndrome later in life ³. Untreated GDM predisposes to adverse neonatal outcome and predicts future development of T2D in both the mothers and offspring ⁴.

Ethnicity has a great impact on the prevalence of GDM which of GDM differs between 1% and 10-35% in different populations and applying different criteria ^{5,6}. Individuals of Asian descents have 2-7 -fold greater risk of developing GDM than their Caucasian counterparts ⁷. These differences can have several explanations including differences in predisposing risk factors including diet and lifestyle ⁸, but also genetics and diagnostic criteria applied ⁹.

Family history of T2D or previous history of GDM increases risk for developing GDM during pregnancy. Several candidate gene studies have confirmed a role for T2D risk loci in GDM. To our knowledge, only one Asian GWAS study on GDM in Korean women has been published ¹⁰. Few studies have examined the genetic susceptibility to GDM in the Indian population; these included two studies in South Indian women and reported association of variants in the *CDKALI*, *HMG20A* and *HNF4A* genes with GDM ^{11,12}. It is quite possible that the genetic background contributing to GDM differs between women of North Indian and of

Scandinavian origin and could explain some disparities in the prevalence of GDM. To address these questions, we examined phenotypic and genetic differences in pregnant women with GDM from India and Sweden.

Methods

Study population and GDM diagnosis

PunjabiGDM study: A multistage protocol was applied for recruiting study participants between 2009 and 2012 in a representative group of 5100 pregnant women from Punjab. All women between gestational weeks 24-28 weeks visiting selected study sites, both urban and rural for antenatal checkup were screened. Information of demographic factors including diet, age, family history of diabetes, BMI, location (urban / rural), education status and religion was obtained in a standard questionnaire by trained health-care professionals. Written material was provided in three languages (Hindi, English and Punjabi). Consent was obtained from each study participant after full explanation of the purpose and nature of all procedures used.

The project was approved by Independent ethics committee, Ludhiana in 2009 (registered with Office of Drugs Controller General (India) Directorate General of Health Service). All participants underwent a 75-g OGTT as previously described^{5,13}. Based on availability of DNA and clinical data a total of 4018 women were included in the study.

Scandinavian Cohorts

Malmö Study: From a total of 188 women with GDM referred to the Department of Endocrinology in Malmö, Sweden, between 1995 and 1999, 83 women of Swedish ethnicity were included in the present study. A 75-g OGTT was performed at the 28th week of

gestation. The OGTT was then repeated with venous measurements of blood glucose concentrations at 0, 30, 60 and 120 min with simultaneous measurements of insulin concentrations. Blood glucose values were converted to plasma glucose by multiplying by a factor of 1.11 according to the IFCC recommendation ¹⁴.

Mamma study: Pregnant women giving birth in the County of Skåne in southern Sweden between 2003 and 2005 were recruited to the Mamma study. A 75-g OGTT was offered to all the women at 27-28 week of gestation as part of routine antenatal care. From a total of 424 women of Swedish ancestry, 89 women with GDM (2-h capillary plasma glucose concentration ≥ 10.0 mmol/l), and 335 women without GDM (2-h capillary plasma glucose concentration < 9.9 mmol/l) with DNA available were included in the study.

Informed consent was obtained from each study participant after full explanation of the purpose and nature of all procedures used and the studies were approved by the Ethics Committee of Lund University. Glucose concentrations were measured using HemoCue devices (HemoCue, Ängelholm, Sweden).

Diagnosis of GDM

In order to maintain consistency between studies, GDM in the present study was defined as 2-h plasma blood glucose concentration ≥ 10 mmol /l (2-h capillary blood glucose concentration of ≥ 9.0 mmol/l) in accordance with the definition in the Malmö study, where only 2-h glucose values were available.

Biochemical measurements

Serum insulin concentrations were measured with an enzyme-linked immunosorbent assay (ELISA, Dako, Glostrup, Denmark), and homeostasis model assessment (HOMA2-B and

HOMA2-IR) was used for estimation of insulin secretion and action, using the HOMA2 calculator v2.2.3 (<http://www.dtu.ox.ac.uk/homacalculator/>)¹⁵.

Genotyping

DNA was extracted from buffy coats using the QIAGEN Autopure W kit. Six SNPs previously associated with GDM and /or T2D in Indian people in either GWAS or candidate studies^{11,16-18} and 79 SNPs associated with T2D/GDM from previous GWAS studies with replication, with overall p-values $<5 \times 10^{-8}$ ¹⁹ (supplementary table 1) were genotyped in the current study using a Sequenom mass ARRAY platform or Taqman. All SNPs passed the Bonferroni threshold of < 0.0006 for Hardy-Weinberg equilibrium test.

Statistical analyses

Anthropomorphic and glycemic measures are presented as means \pm SEM. Significance of differences between group means was tested by the Mann-Whitney U test or analysis of variance or covariance (ANCOVA) with BMI and age as covariates. Inverse normal transformation was used to normalize data with skewed distributions.

Allele and genotype frequencies were compared between groups by chi-square or Fisher's exact test. Association of selected SNPs with GDM was assessed by logistic regression analysis adjusted for maternal age and results presented as ORs with their 95% confidence intervals (CI) in plink (plink v1.09). Alleles were also analyzed for association with glucose, insulin and HOMA2-B and HOMA2-IR) using linear regression model adjusted for age.

Two-sided p values of less than 0.05 were considered statistically significant. For the Indian study population, power to detect association with GDM (125 cases and 3893 controls) for 79 markers at a significance level of 0.05, was 0.04 under an additive model and 0.12 under a

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multiplicative model at 0.50 MAF and OR of 1.5. For the Swedish cohort with 245 cases and 335 controls, the corresponding figures were 0.06 and 0.17 respectively. For association with quantitative traits, power to detect an association was 1 at alpha 0.05 for an allele frequency of 0.3²⁰. Polygenic risk scores which is the sum of trait-associated alleles across many genetic loci, typically equally weighted for effect sizes, were calculated using PLINK and regressed against the phenotype. In order to construct genetic risk scores (GRS), we used SNPs from previous GWAS studies on T2D risk and previously published candidate studies for GDM. We constructed unweighted polygenic genetic risk scores using the Plink 1.9 software package. We constructed scores using (1) known GDM variants only (2) GDM variants reported in India, (3) T2D loci from previous GWAS study (i.e. genome-wide significant SNPs), (4) T2D loci showing genome wide significance in Indian populations alone, (5) All reported T2D and GDM loci and (6) T2D and GDM loci from Indian populations studies alone (supplementary table 1). Twelve SNPs previously associated with insulin secretion indices and 5 SNPs with insulin resistance were here used to construct genetic risk scores (GRS) for insulin secretion and action respectively using PLINK in women with GDM²¹ (supplementary table 1).

To compare the relationship between (i) HOMA2-B and BMI, (ii) HOMA2-B and HOMA2-IR and (iii) HOMA2-IR and BMI in non-diabetic women from Indian and European studies, we selected non-diabetic women from the Prevalence, Prediction and Prevention of diabetes (PPP)–Botnia Study, a population-based study in Western Finland started in 2004²², due to lack of data on normal glucose tolerance (NGT) women from the Malmö and Mamma studies. For the present analyses, non-diabetic women (age : mean \pm SD = 49.59 \pm 15.75) were selected and age adjusted residuals of HOMA2-B, HOMA2-IR and BMI were calculated and compared for (i) and (iii) and age and BMI adjusted residuals were calculated for (ii) .

F_{ST} analysis for estimating genetic differentiation between Indian and Scandinavian study populations overall as well as for affected women only was computed using the Weir and Cockerham method²³ implemented in vcfTools²⁴.

Replication in European study populations

The Radiel study comprising 398 women (age = 32.7 ± 4.4 years, mean BMI = 32.8 kg/m^2 SD, 8 with GDM) were included for replication of 18 selected loci showing nominal significance in either Indian or Swedish studies or both. Two SNPs including rs4402960 and rs10010131 had HWE < 0.05 in controls and were excluded.

The STORK study is a prospective cohort of 1031 healthy pregnant women of Scandinavian heritage who registered for obstetric care at the Oslo University Hospital Rikshospitalet from 2001 to 2008²⁵. Exclusion criteria were multiple pregnancies, known history of type 1 or type 2 diabetes mellitus, and severe chronic diseases (pulmonary, cardiac, gastrointestinal, or renal). Results of a 75 g OGTT, age, height and weight were recorded at inclusion at gestational weeks 14–16. The OGTT test was repeated at the third visit in gestational week 30–32. The study was approved by the Norwegian “Regional Committee for Medical Health Research Ethics South East”. Written consents were obtained for all participants. In STORK 512 women with European ancestry were selected for the present study (8 cases and 504 controls, age: 31.6 ± 3.78 years).

The STORK Groruddalen study (STORK-G) is a population-based cohort which included 823 healthy women attending three public mother child health clinics for antenatal care in the multi-ethnic area of Groruddalen, Oslo, Norway²⁶. Women were eligible if they: 1) lived in the study districts; 2) planned to give birth at one of two study hospitals; 3) were < 20 weeks

pregnant; 4) could communicate in Norwegian or any of the eight translated languages; 5) were able to give an informed consent. Women with pre-gestational diabetes or in need of intensive hospital follow-up during pregnancy were excluded. The participation rate was 74 %, varying from 63.9 % to 82.6 % across ethnic groups. The study was approved by the Norwegian “Regional Committee for Medical Health Research Ethics South East”. Written consents were obtained for all participants. In STORK-G, 268 women with European ancestry were included (4 cases and 264 controls, age: 30.7 ± 4.51 years).

Meta-analysis: of the selected SNPs in the European studies was performed using METAL (<http://csg.sph.umich.edu/abecasis/metal/>) with beta coefficients for directionality and weighted on the study population.

eQTL lookups: The association of SNPs with gene expression in human pancreatic islets was looked up in data from RNAseq data from 191 donors from a previously published study²⁷.

Results

Clinical characteristics

Applying Swedish GDM criteria cut-offs, the prevalence of GDM in the PunjabiGDM study was 3.11% (125 out of 4018 women). Swedish women with GDM were >10 years older ($p=1.21 \times 10^{-40}$) and had higher BMI (28.09 ± 0.64 vs 24.08 ± 0.42 , $p = 3.76 \times 10^{-07}$) than the Punjabi women (table 1).

The Indian GDM women had higher fasting glucose (5.72 ± 0.15 vs 4.79 ± 0.10 , 1.60×10^{-5}) and 2-h glucose (12.07 ± 0.20 vs 10.99 ± 0.08 , $p = 3.13 \times 10^{-02}$) which associated with lower fasting insulin (51.8 ± 5.35 vs 78.17 ± 12.67 , $p = 3.74 \times 10^{-06}$) and HOMA2-B (76.6 ± 3.83 vs 123.98 ± 7.54 , $p = 2.99 \times 10^{-9}$) as well as better insulin sensitivity estimated by HOMA2-IR (1.036 ± 0.97 vs 1.26 ± 0.097 , $p = 0.001$) compared with Swedish GDM women adjusted for BMI and age (table 1).

Insulin secretion, resistance and BMI in Indian and Scandinavian women

We examined the relationship of HOMA2-B, HOMA2-IR and BMI in women with NGT in Indian and Swedish women. Since we lacked OGTT data from Swedish pregnant non-diabetic women, we here used OGTT from women with NGT from the Botnia-PPP cohort. There was a stronger relationship between BMI and insulin secretion as well as insulin resistance in Scandinavian women compared to Indian. Here we found significant differences in the relationship between HOMA2-B - HOMA2-IR ($p = 0.0028$, figure 1a), HOMA2-B - BMI ($p = 8.48 \times 10^{-17}$, figure 1b) and HOMA2-IR - BMI ($p = 8.11 \times 10^{-25}$, figure 1c) between Indian and Scandinavian women with NGT.

Frequency differences and population differentiation for GDM and T2D associated SNPs between populations

Significant differences were observed for minor allele frequencies of SNPs at the *CDKALI*, *SLC30A8*, *IGF2BP2*, *ADCY5*, *G6PC2*, *WFS1*, *HHEX/IDE* and *MTNR1B* loci between all pregnant women from the Indian and Swedish studies (supplementary figure 1). F_{ST} analysis

showed clear genetic differentiation at the rs998451 (*TMEM163*), rs8042680 (*PRC1*), and rs1799999 (*PPP1R3A*) loci whereas differences at 14 loci including rs560887 (*G6PC2*) and rs9552911 (*SGCG*) loci were less pronounced (figure 2).

When only focusing on women with GDM, we observed significant differences in frequency of GDM risk alleles rs560887 in *G6PC2* ($p = 0.0008$), rs11708067 in *ADCY5* ($p = 0.005$), rs10010131 in *WFS1* ($p = <0.0001$) and rs10811661 ($p=0.0073$) in *CDKN2B* genes between Indian and Swedish women with GDM were seen (supplementary table 2). F_{ST} values here was identical to that seen when the entire study population was considered with the exception of *DUSP9*, *HHEX*, and *WFS1* which showed moderate differentiation in GDM women whereas very little when all women were considered (figure 2).

Association of genetic loci with GDM in Indian and Swedish women

SNPs previously associated with GDM/T2D in India. The risk allele C of rs7178572 SNP near *HMG20A* was nominally associated with risk of GDM in Indian but not in Swedish women ($p = 0.03$, table 2), thereby replicating previous findings in Indian populations. This SNP also showed nominally significant association with GDM in the STORK-G study, which predominantly comprised multi-ethnic women. rs7178572 is an eQTL for the *PSTPIP1* ($p = 0.003$) and *HMG20A* ($p=0.007$) genes in human pancreatic islets (Supplementary table 4).

The Asp/Tyr missense variant of SNP rs1799999 in the *PPP1R3A* gene, which previously has been shown to associate with T2D risk in Jat Sikhs¹⁸, showed a trend towards association in Indian women ($p = 0.06$) (table 2). The variant was also nominally associated with decreased 2-hour (2-h) insulin in Swedish women ($p = 0.02$, supplementary table 3).

SNPs previously associated with GDM or T2D in Europeans: Of 12 SNPs previously associated with Scandinavian GDM^{28,29} (Supplementary table 1), the rs1111875 SNP near the *HHEX/IDE* genes was nominally associated with GDM in Swedish women ($p = 0.031$, table 3). While the association of the SNP with GDM in the other European cohorts was not significant, the direction of effect was consistent with that in the Swedish study in two out of three studies (table 3). The rs1111875 variant influences expression of *NHP2P1* and *BTAFL* genes in human pancreatic islets (supplementary table 4). The frequency of rs1111875 differed between Indian and Swedish women ($p = <0.0001$, supplementary table 2, supplementary figure 1).

The risk allele rs11708067 in the *ADCY5* gene was associated with increased 2-h glucose ($p = 0.037$), and decreased HOMA2-B ($p = 0.010$) in Swedish GDM women (supplementary table 3). The same SNP was associated with 2-h glucose in all Swedish women (GDM and non-GDM) ($\beta = 0.12 \pm 0.04$, $p = 0.004$).

The rs11605924 SNP in the intron of the *CRY2* gene was nominally associated with GDM in both study populations (table 4). Interestingly, the T2D risk allele A was protective in the Indian population (OR = 0.67, $p = 0.0026$, table 4) whereas it was associated with risk in the Swedish women (OR = 1.44, $p = 0.012$, table 4). A similar direction of effect was observed in the Radiel, STORK and STORK-G studies and the meta-analysis of European studies showed a significant association with GDM ($p = 0.014$, table 4). The same SNP showed differences in frequencies between Indian and Swedish women ($p = 0.0004$, supplementary table 2, supplementary fig 1). The rs11605924 nominally influenced expression of *CRY2* in human pancreatic islets (supplementary table 4).

The rs8090011 SNP in an intron of the *LAMA1* gene was nominally associated with GDM risk in Swedish women (table 4). The same SNP also associated with decreased 2-h insulin concentration (supplementary table 4).

SNPs rs12571751 in the intron of *ZMIZ1*, rs5945326 near *DUSP9*, and rs2237895 in the intron of *KCNQ1* were nominally associated with GDM in Swedish women whereas only the rs7593730 SNP near *RBMS1* was associated with GDM risk in Indian women (table 4).

Genetic risk scores (GRS) based on all T2D / GDM loci predicted GDM risk in Swedish but not in Indian women ($p = 0.036$, table 5) whereas GRS based on previous GDM associations in Indian populations predicted GDM risk in India ($p = 0.042$, table 5). GRS for insulin resistance was $0.11 (\pm 0.1, p = 0.49)$ for Swedish whereas $0.13 (\pm 0.07, p = 0.059)$ for Indian women. GRS for insulin secretion was $-0.034 (\pm 0.06, p = 0.53)$ for Swedish and $-0.078 (\pm 0.039, p = 0.42)$ for Indian women.

Discussion

The key findings in the current study were that Indian and Swedish women with GDM showed clear differences in measures of insulin secretion and action (i.e. HOMA2-B and HOMA2-IR), which cannot fully be accounted for by genetic effects alone. Despite being on average 10 years younger, North Indian women had a higher prevalence (3.11%) of GDM than previously reported in Swedish women (2.6%) from comparable time periods⁶. Of note, the prevalence figure of 2.6% during 2009-2012 was based on a study population of mixed ethnicity residing in Sweden, and a lower prevalence could be expected if only Swedish women were included (estimated 1.2-1.5%)³⁰. This is consistent with previous

reports showing higher GDM frequency in populations with a high frequency of T2D³¹. The prevalence of T2D from previous studies was higher in India than in Sweden (8.3-9.4% vs 6.8%)^{32,33}.

Indian women had lower HOMA2-B, which was associated with lower BMI and better insulin sensitivity than Swedish GDM women. The better insulin sensitivity could at least partially be a corollary of the lower BMI (supplementary figure 2). As Indian women seem to develop GDM at lower BMI and with better insulin sensitivity, this could point at a more severe defect in insulin secretion, which was also supported by lower HOMA2-B. However, we need to acknowledge that HOMAs are only surrogate markers for insulin secretion and action.

Previously, 6 loci have been associated with T2D or GDM in India^{11,16-18}. Of them, the rs7178572 SNP near the *HMG20A* gene was associated with GDM in Indian but not in Swedish women. Considering the effect sizes observed in both populations, this is likely an effect of low power in the Swedish studies and to a lesser extent heterogeneity, given that this SNP has also been associated with T2D in other European populations³⁴. The power to detect association of rs7178572 (*HMG20A*) with GDM at a significance level of 0.05 was 0.76. Notably, only when we used the older WHO1999 criteria with higher cut-off values for glucose, we could observe this association. This is consistent with our previous reports based on the same study population (PunjabiGDM), rs7178572 did not associate with GDM diagnosed either using WHO1999 or WHO2013 criteria³⁵. Interestingly, a GRS constructed of previous GDM associations in Indian populations including this SNP significantly predicted GDM in the Indian but not Swedish women. Also of interest is that moderate genetic differentiation observed at this locus ($F_{ST} = 0.07$). It should be also noted that the Indian population has complex genetic origins, with high genetic diversity between the North

and South. The Punjabi-Indian population belongs to the “Ancestral North Indians” group and shares genetic similarities with those from Middle East, Central Asia and to some extent, Europe whereas the South Indian population belongs to the genetically distinct “Ancestral South Indian” group³⁶. Due to differences in haplotype structures, variants associated with GDM in the South Indian population might not necessarily associate with GDM in the North India. Earlier studies have shown a weak association of rs7178572 with *PSTPIP1* gene expression in lymphoblastoid cell lines³⁴. Here we showed that this SNP also was an eQTL in human pancreatic islets influencing expression of both *PSTPIP1* and *HMG20A*. The Proline-Serine-Threonine Phosphatase Interacting Protein 1 (*PSTPIP1*) gene is a tyrosine phosphatase that inhibits T-cell activation upon T-cell receptor (TCR) and *CD28* engagement, regardless of CD2 co-stimulation³⁷. The *HMG20A* gene had higher expression in islets than in muscle and adipose tissue³⁸ and a transient increase in expression levels were observed upon glucose stimulation³⁸. *HMG20A* has been reported to be down-regulated in T2D and T1D islets, and knockdown of *HMG20A* decreased expression of *NEUROD*, *INS* and *GK* with an accompanying impairment in GSIS³⁸. Therefore, this SNP could through its eQTL effect on *HMG20A* expression in islets be a plausible candidate gene for GDM.

The early GWAS SNP rs1111875 near the *HHEX/IDE* genes was associated with GDM risk in Swedish women but not Indian. Notably, the T2D risk SNP rs11605924 in *CRY2* showed a protective effect against GDM in Indian but conferred risk in Swedish women. The power to detect association of rs11605924 (*CRY2*) with GDM at a significance level of 0.05 was 0.85. This protective effect was consistent even when WHO1999 criteria was applied in our previous study on the same population³⁵. *CRY2* encodes the circadian rhythm gene cryptochrome 2, and is a target for the *CLOCK-BMAL1*, which are core components of the endogenous clock. The *CRY2* variant is also associated with fasting glucose and reduced liver fat content in human liver³⁹. *CRY2* mRNA expression has been associated with hepatic

triglyceride content³⁹ suggesting that *CRY2* could serve as a switch between fat and glucose metabolism in the liver³⁹. Interestingly, as the same allele had effects in opposite directions in Indian and Swedish populations, the question rises whether risk seen in the Swedish population could be related to marked differences in circadian rhythm during seasons in Sweden, which is lacking in India. However, this finding needs to be replicated in other Indian studies.

In our previous study, we observed that the T2D risk SNP in the *TCF7L2* gene did not associate with GDM either using WHO1999 or WHO2013 criteria³⁵. However, using the Swedish GDM diagnostic criteria (2-h capillary blood glucose concentration ≥ 10 mmol /l), we see a trend towards an association. Increasing fasting glucose cut-offs could further capture the extreme GDM cases and a stronger signal could be obtained. Alternately, this could also be due to population-based differences³⁵.

Interestingly, significant frequency differences were observed for 6 out of 12 GDM risk alleles, two of which showed a reversal of major and minor alleles. High to moderate genetic differentiation was observed at 17 loci. Of particular interest was the highest differentiation observed at the *TMEM163* locus, which had previously only shown an association with T2D in Indian but not European studies. A previous study of 12 T2D risk alleles showed decreasing frequencies going from Africa to Europe to East Asia. These decreasing frequencies were associated with different risk of T2D, with the highest in Africa and lowest in Asia. The authors hypothesized that these differences might be caused by optimizing energy storage and usage in environments with inconsistent energy intake⁴⁰. *CRY2* could also potentially represent such an example.

There were significant differences in the relationship between insulin secretion and insulin resistance or between insulin resistance and BMI between glucose-tolerant Indian and Swedish women, which could not be explained by differences in age. A GRS comprised of all previously reported GDM and T2D loci from European studies predicted GDM in the Swedish but not Indian women. These data support the view that differences between these ethnic groups could be partially explained by genetic differences. Since most T2D loci were identified in European ancestry cohorts, this could reflect differences in tagging SNPs due to differences haplotypes between populations. This could also be partly attributed to the differences in anthropometry between Indian and Scandinavian populations, with the former manifesting the distinct “thin-fat” phenotype from birth^{34,35}.

The criteria applied to diagnose GDM markedly influence risks. WHO1999 clearly identifies a more severe dysregulation of glucose metabolism than the other criteria. On the other hand, WHO 2013 is designed to identify risk of adverse pregnancy outcomes in the offspring. The PRS derived from T2D loci identified a shared genetic background between GDM and T2D in India, but not in Sweden probably due to too low power. This does not exclude the possibility that a GWAS could identify shared genetic background also for the other criteria and thereby risk for offspring.

A limitation of the study was the low power in some of the studies to allow multiple testing; we hope that the meta-analysis of Scandinavian studies including Radiel, STORK and STORK-G compensated for this to some extent. Additionally, a proportion of women diagnosed with GDM based on WHO1999 or WHO2013 criteria were excluded due to application of Swedish GDM criteria. Despite differences in study settings, the stringent

GDM diagnostic criteria applied (based on Sweden criteria) should identify the most extreme cases. Nevertheless this is largest comparative study comparing GDM between European and non-European populations and further, the first study comparing GDM in India with that of Europe.

Hypertension during pregnancy is an important aspect, however this was unfortunately not measured for all participants due to limitations in the screening process in many rural areas of screening and is one of the limitations of the study.

Taken together Indian women develop GDM at lower BMI and better insulin sensitivity than Scandinavian women. This points at problems to increase insulin secretion to meet the increased demands imposed by even small increases in insulin resistance during the third trimester. The genetic contribution to GDM seems to be shared with T2D.

Author Contributions

GPA, MA, CB, NW and RPB researched data and reviewed/edited the manuscript. RPB, MA, NW, GHM, and PA analyzed the data. RPB, GPA, AAV, KB, LG contributed to study design and reviewed/edited the manuscript. RT, MOM, NW, GHM, CS, EQ, AKJ, JE, KB, KIB, AV reviewed/edited the manuscript. KB and LG contributed to the discussion and extensively reviewed/edited the manuscript. RPB wrote the manuscript.

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Conflict of interest statement

Dr. Prasad reports other from Novo Nordisk, other from AstraZeneca, outside the submitted work.

CB is a stockholder in Novo Nordisk A/S and AV is employed by AstraZeneca. On behalf of all the other authors, Dr. Prasad B has nothing to disclose. RPB and LG take responsibility for the contents of the article.

Data availability statement

The datasets generated during and/or analysed during the current study are available from the corresponding author on reasonable request.

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Table 1. Clinical characteristics of Indian and Swedish women with GDM (diagnosed based on 2-h glucose cut-offs ≥ 10 mmol/l). Mean \pm SEM are represented. P-values are calculated based on inverse normal transformed data.

Variable	Swedish	N (Swedish)	Indian	N (Indian)	P value
Age (years)	31.78 \pm 0.36	149	20.97 \pm 0.33	125	1.21 x 10 ⁻⁴⁰
BMI (kg/m ²)	28.09 \pm 0.64	56	24.08 \pm 0.42	125	3.76 x 10 ⁻⁰⁷
Fasting glucose ^a (mmol/l)	4.79 \pm 0.10	49	5.72 \pm 0.15	125	1.60 x 10 ⁻⁰⁵
2-h glucose ^a (mmol/l)	10.99 \pm 0.08	149	12.07 \pm 0.20	125	3.13 x 10 ⁻⁰²
Fasting insulin ^a (pmol/l)	78.17 \pm 12.67	51	51.8 \pm 5.35	125	3.74 x 10 ⁻⁰⁶
HOMA2-B ^a	123.99 \pm 7.55	45	76.61 \pm 3.83	109	3.00 x 10 ⁻⁰⁹
HOMA2-IR ^a	1.26 \pm 0.10	45	1.04 \pm 0.10	109	1.11 x 10 ⁻⁰³

^a adjusted for age and BMI

Table 2. Association of T2D and GDM risk loci previously reported in the Indian population based studies with GDM risk in Indian and Sweden pregnant women. CHR = chromosome, BP = base pair coordinates, A1 = effect allele, n = study population size, OR = odds ratio (CI = 95% confidence intervals), P = two-tailed p-value for single test.

CHR	SNP	locus	A1	Study	n	GDM_N	OR (CI)	P	
15	rs7178572	<i>HMG20A</i> <i>/DUSP9</i>	G/A	PunjabiGDM	3346	122	1.34 (1.01 – 1.76)	0.03	
				Malmö-Mamma	476	144	1.22 (0.87 – 1.69)	0.25	
	rs7177055 *	<i>HMG20A</i> <i>/DUSP9</i>	G/A				8		
				Radiel	398		0.35 (0.08 - 1.57)	0.15	
				Stork	512		1.69 (0.47 - 6.25)	0.41	
				Stork-G	268		0.22 (.05 - 0.96)	0.02	
				meta-analysis EU				0.60	
meta-analysis ALL				0.15					
7	rs1799999	<i>PPP1G.G</i>	C/A	PunjabiGDM	3664	124	1.30 (0.99 – 1.70)	0.06	
				Malmö-Mamma	465	144	1.20 (0.70 – 2.08)	0.50	
				meta-analysis				0.04	
20	rs4812829	<i>HNF4A</i>	G/A	PunjabiGDM	3756	124	1.16 (0.87 – 1.57)	0.30	
				Malmö-Mamma meta-analysis					
11	rs689	<i>INS</i>	A/T	PunjabiGDM	3676	122	1.09 (0.78 – 1.52)	0.60	
						489	144	1.101 (0.80 – 1.51)	0.55
				Malmö-Mamma meta-analysis				0.49	
13	rs9552911	<i>SGCG</i>	A/G	PunjabiGDM	3665	121	0.85 (0.54 – 1.32)	0.47	
				Malmö-Mamma meta-analysis	486	144	1.23 (0.11 – 13.76)	0.86	
								0.54	
2	rs998451	<i>TMEM163,</i> <i>RAB3GAP1</i>	G/A	PunjabiGDM	3656	122	1.18 (0.76 – 1.82)	0.47	
				Malmö-Mamma	482	144	0.93 (0.70 – 1.25)	0.65	
				meta-analysis				0.16	

* r^2 of 0.89 with rs7178572

Table 3. Association of previously reported GDM risk loci discovered in the European population based studies with GDM risk in Indian and Sweden pregnant women. CHR = chromosome, BP = base pair coordinates, A1 = effect allele, n = study population size, OR = odds ratio, P = two-tailed p-value for single test.

CHR	SNP	Locus	RA/OA	Study	n	GDM_N	OR (CI)	P
4	rs10010131	WFS1	G/A	PunjabiGDM	3617	121	1.24 (0.92 - 1.66)	0.15
				Malmö-Mamma	444	144	1,19 (0.88 - 1.64)	0.25
				Radiel	-	8	-	-
				Stork	512	8	2.38 (0.75 - 7.69)	0.13
				Stork-G	268		2.17 (0.43 - 11.11)	0.32
				meta-analysis_EU				0.45
				meta-analysis_ALL				0.10
9	rs10811661	CDKN2B	T/C	PunjabiGDM	3666	122	0.99 (0.68 - 1.44)	0.96
				Malmö-Mamma	428	144	1.31 (0.81 - 2.12)	0.25
				Radiel	398	8	0.66 (0.18 - 2.38)	0.53
				Stork	512	8	1.56 (0.34 - 7.14)	0.56
				Stork-G	-	4	-	-
				meta-analysis_EU				0.93
				meta-analysis_ALL				0.99
10	rs1111875	HHEX/ IDE	G/A	PunjabiGDM	3675	120	1.02 (0.79 - 1.33)	0.86
				Malmö-Mamma	443	144	1.41 (1.03 - 1.92)	0.031
				Radiel	398	8	3.03 (0.84 - 11.11)	0.075
				Stork	512	8	0.69 (0.25 - 1.85)	0.45
				Stork-G	268	4	1.33 (0.31 - 5.88)	0.70
				meta-analysis_EU				0.08
				meta-analysis_ALL				0.26
3	rs11708067	ADCY5	A/G	PunjabiGDM	3648	123	0.78 (0.58 - 1.04)	0.084
				Malmö-Mamma	466	144	1.44 (1.00 - 2.08)	0.054
				Radiel	398	8	0.95 (0.36 - 3.33)	0.94
				Stork	512	8	0.46 (0.16 - 1.31)	0.13
				Stork-G	268	4	0.87 (0.17 - 4.54)	0.86
				meta-analysis_EU				0.93
				meta-analysis_ALL				0.16
8	rs13266634	SLC30A8	T/C	PunjabiGDM	3671	122	0.97 (0.72 - 1.31)	0.84
				Malmö-Mamma	458	144	0.88 (0.64 - 1.20)	0.42
				Radiel	398	8	1.36 (0.50 - 3.69)	0.54
				Stork				
				Stork-G				
				meta-analysis_EU				0.86
				meta-analysis_ALL				0.78

3	rs1801282	PPARG	C/G	PunjabiGDM	3436	116	1.06 (0.71 - 1.58)	0.76
				Malmö-Mamma	421	144	0.75 (0.48 - 1.16)	0.21
				Radiel				
				Stork				
				Stork-G				
				meta-analysis_EU				0.21
				meta-analysis_ALL				0.89
9	rs2796441	TLE1	C/T	PunjabiGDM	3677	122	0.98 (0.75 - 1.27)	0.88
				Malmö-Mamma	457	144	1.00 (0.33 - 2.94)	0.98
				Radiel	398	8	0.64 (0.24 - 1.72)	0.38
				Stork	512	8	1.44 (0.53 - 3.86)	0.46
				Stork-G	268	4	0.48 (0.09 - 2.38)	0.35
				meta-analysis_EU				0.63
				meta-analysis_ALL				0.69
3	rs4402960	IGF2BP2	T/G	PunjabiGDM	3535	121	0.89 (0.69 - 1.15)	0.36
				Malmö-Mamma	352	144	1.06 (0.73 - 1.53)	0.77
				Radiel				
				Stork	512	8	1.48 (0.53 - 4.13)	0.44
				Stork-G	268	4	0.73 (0.14 - 3.70)	0.71
				meta-analysis_EU				0.62
				meta-analysis_ALL				0.59
11	rs5219	KCNJ11	T/C	PunjabiGDM	3382	117	1.04 (0.80 - 1.36)	0.73
				Malmö-Mamma	264	144	1.09 (0.70 - 1.67)	0.71
				Radiel	398	8	1.43 (0.53 - 3.89)	0.47
				Stork				
				Stork-G				
				meta-analysis_EU				0.43
				meta-analysis_ALL				0.53
2	rs560887	G6PC2	G/A	PunjabiGDM	3678	122	0.93 (0.64 - 1.35)	0.70
				Malmö-Mamma	383	144	1.25 (0.85 - 1.81)	0.26
				Radiel	398	8	0.90 (0.30 - 2.63)	0.85
				Stork	512	8	0.96 (0.33 - 2.80)	0.94
				Stork-G	268	4	0.36 (0.04 - 2.97)	0.32
				meta-analysis_EU				0.36
				meta-analysis_ALL				0.86
6	rs7754840	CDKAL1	C/G	PunjabiGDM	3502	116	1.19 (0.90 - 1.57)	0.22
				Malmö-Mamma	426	144	0.97 (0.71 - 1.32)	0.83
				Radiel	398	8	1.42 (0.52 - 3.84)	0.49
				Stork	512	8	0.96 (0.33 - 2.79)	0.93
				Stork-G	268	4	0.70 (0.14 - 3.51)	0.66

							meta-analysis_EU	0.99
							meta-analysis_ALL	0.31
6	rs7756992	<i>CDKAL1</i>	G/A	PunjabiGDM	3469	115	1.16 (0.88 - 1.55)	0.29
				Malmö-Mamma	425	144	0.96 (0.70 - 1.32)	0.79
				Radiel				
				Stork	512	8	0.85 (0.27 - 2.79)	0.77
				Stork-G	268	4	0.81 (0.17 - 4.32)	0.85
							meta-analysis_EU	0.65
							meta-analysis_ALL	0.50
10	rs7903146	<i>TCF7L2</i>	T/C	PunjabiGDM	3330	108	1.29 (0.98 - 1.72)	0.07
				Malmö-Mamma	373	144	1.20 (0.84 - 1.70)	0.31
	rs12255372			Radiel	398	8	0.28 (0.04 - 2.15)	0,193
				Stork	512	8	0.63 (0.18 - 2.24)	0.47
				Stork-G	268	4	0.86 (0.17 - 4.32)	0.85
							meta-analysis_EU	0.51
							meta-analysis_ALL	0.27
16	rs9939609	<i>FTO</i>	A/T	PunjabiGDM	2962	88	0.79 (0.57 - 1.10)	0.16
				Malmö-Mamma	0		NA	NA
				Radiel				
				Stork				
				Stork-G				
							meta-analysis_EU	
							meta-analysis_ALL	
11	rs10830963	<i>MTNR1B</i>	G/C	PunjabiGDM	3495	114	0.93 (0.72 - 1.23)	0.65
				Malmö-Mamma	425	144	1.03 (0.75 - 1.41)	0.85
				Radiel	398	8	2.85 (0.95 - 8.6)	0.052
				Stork	512	8	0.88 (0.28 - 2.76)	0.83
				Stork-G	268	4	0.92 (0.18 - 4.63)	0.92
							meta-analysis_EU	0.37
							meta-analysis_ALL	0.89

Table 4. Association of previously reported T2D risk loci discovered in the European population based studies with GDM risk in Indian and Sweden pregnant women. CHR = chromosome, BP = base pair coordinates, A1 = effect allele, n = study population size, OR = odds ratio, P = two-tailed p-value for single test.

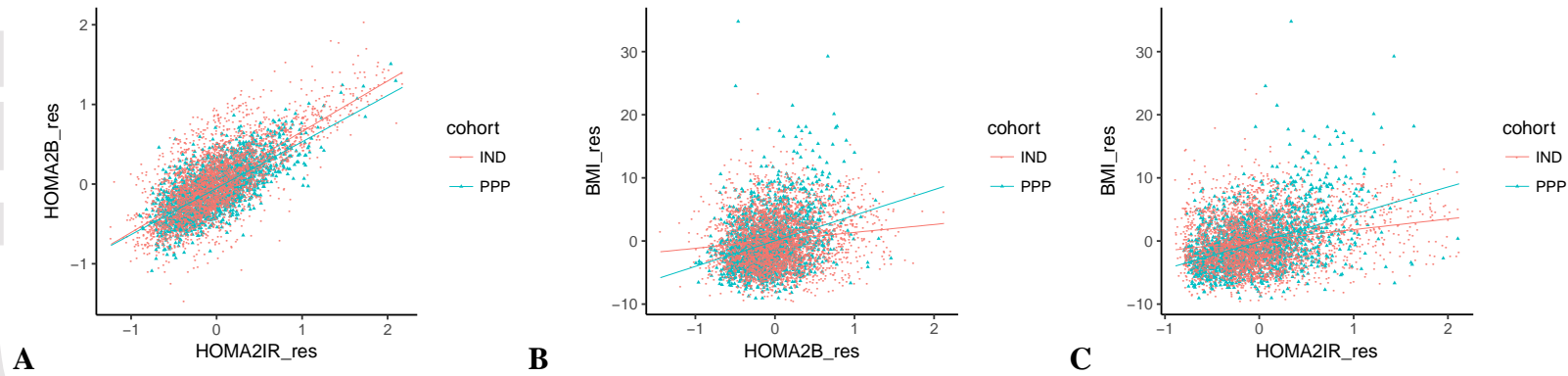
CHR	SNP	Locus	RA/OA	Study	n	GDM_N	OR (CI)	P
10	rs12571751	ZMIZ1	A/G	PunjabiGDM	3390	117	0.80 (0.62 - 1.05)	0.11
				Malmö-Mamma	492	144	0.72 (0.54-0.95)	0.021
				Radiel	398	8	1.68 (0.61 - 4.68)	0.31
				Stork	512	8	0.95 (0.36 - 2.56)	0.92
				Stork-G	268	4	0.30 (0.06 - 1.50)	0.11
				meta-analysis EU				0.15
				meta-analysis ALL				0.03
11	rs11605924	CRY2	A/C	PunjabiGDM	3679	120	0.67 (0.52 - 0.87)	0.002
				Malmö-Mamma	484	144	1.44 (1.08-1.91)	0.0129
				Radiel	398	8	1.39 (0.50 - 3.85)	0.53
				Stork	512	8	1.62 (0.58 - 4.94)	0.35
				Stork-G	268	4	1.69 (0.40 - 7.15)	0.46
				meta-analysis EU				0.014
				meta-analysis ALL				0.26
11	rs2237895	KCNQ1	C/A	PunjabiGDM	3463	113	0.81 (0.62 - 1.06)	0.13
				Malmö-Mamma	410	144	1.43 (1.06-1.94)	0.0204
				Radiel	398	8	1.20 (0.44 - 3.23)	0.72
				Stork	512	8	1.15 (0.43 - 3.10)	0.77
				Stork-G	268	4	0.41 (0.08 - 2.06)	0.26
				meta-analysis EU				0.29
				meta-analysis ALL				0.51
15	rs7177055	HMG20A	A/G	PunjabiGDM	3680	122	1.35 (1.03 - 1.75)	0.024
				Malmö-Mamma	457	144	1.09 (0.79-1.52)	0.58
				Radiel				
				Stork	512	8	1.65 (0.47 - 5.83)	0.43
				Stork-G	268	4	0.21 (0.05 - 0.91)	0.02
				meta-analysis EU				0.82
				meta-analysis ALL				0.06
18	rs8090011	LAMA1	G/C	PunjabiGDM	3683	122	1.02 (0.79 - 1.32)	0.89
				Malmö-Mamma	457	144	1.49 (1.11-2.01)	0.009
				Radiel				
				Stork				
				Stork-G				
				meta-analysis EU				
				meta-analysis ALL				0.31

23	rs5945326	DUSP9	A/G	PunjabiGDM	3377	113	0.87 (0.66 - 1.13)	0.29
				Malmö-Mamma	495	144	1.44 (1.03-2.05)	0.035
				Radiel				
				Stork	512	8	0.40 (0.15 - 1.08)	0.06
				Stork-G	268	4	2.31 (0.28 - 18.95)	0.42
				meta-analysis EU				0.62
				meta-analysis ALL				0.52

Table 5. GDM Genetic risk scores for Punjabi GDM and Malmö Mamma studies based on (1) all previously reported GDM SNPs, (2) GDM loci reported in Indian populations, (3) previously reported T2D loci, (4) T2D loci from studies on Indian population, (5) all T2D and GDM loci and (6) T2D and GDM reported on Indian populations. All SNPs for GRS are reported in supplementary table 1.

	B	S.E.	p-value
GDM_All			
PUNJABIGDM study	.507	1.044	.627
MALMÖ-MAMMA study	1.656	1.137	.145
GDM_India			
PUNJABIGDM study	.782	.385	.042
MALMÖ-MAMMA study	.123	.430	.774
T2D			
PUNJABIGDM study	.447	.301	.137
MALMÖ-MAMMA study	-.332	.303	.274
T2D_India			
PUNJABIGDM study	.907	.836	.278
MALMÖ-MAMMA study	-.075	.867	.931
T2DGDM_All			
PUNJABIGDM study	-0.989	2.542	.697
MALMÖ-MAMMA study	6.137	2.923	.036
T2D and GDM India-specific			
PUNJABIGDM study	.921	.608	.130
MALMÖ-MAMMA study	.246	.573	.668

Figures



Compare	PPP_bet		PPP_se	IND_se	z	pvalue
	a	IND_beta				
BMI_res ~ HOMA2B_res	4.055	1.260	0.292	0.165	8.324	8.486e-17
BMI_res ~ HOMA2IR_res	4.339	1.686	0.216	0.141	10.286	8.112e-25
HOMA2B_res ~ HOMA2IR_res	0.582	0.634	0.014	0.010	-2.989	0.0028

Figure 1. Relationship between (A) HOMA2-B-HOMA2-IR (B) HOMA2-B and BMI and (C) HOMA2-IR and BMI in Indian and Scandinavian women from the Botnia-PPP study with normal glucose tolerance (NGT). Z = z scores for the differences between effects in Indian and Swedish women.

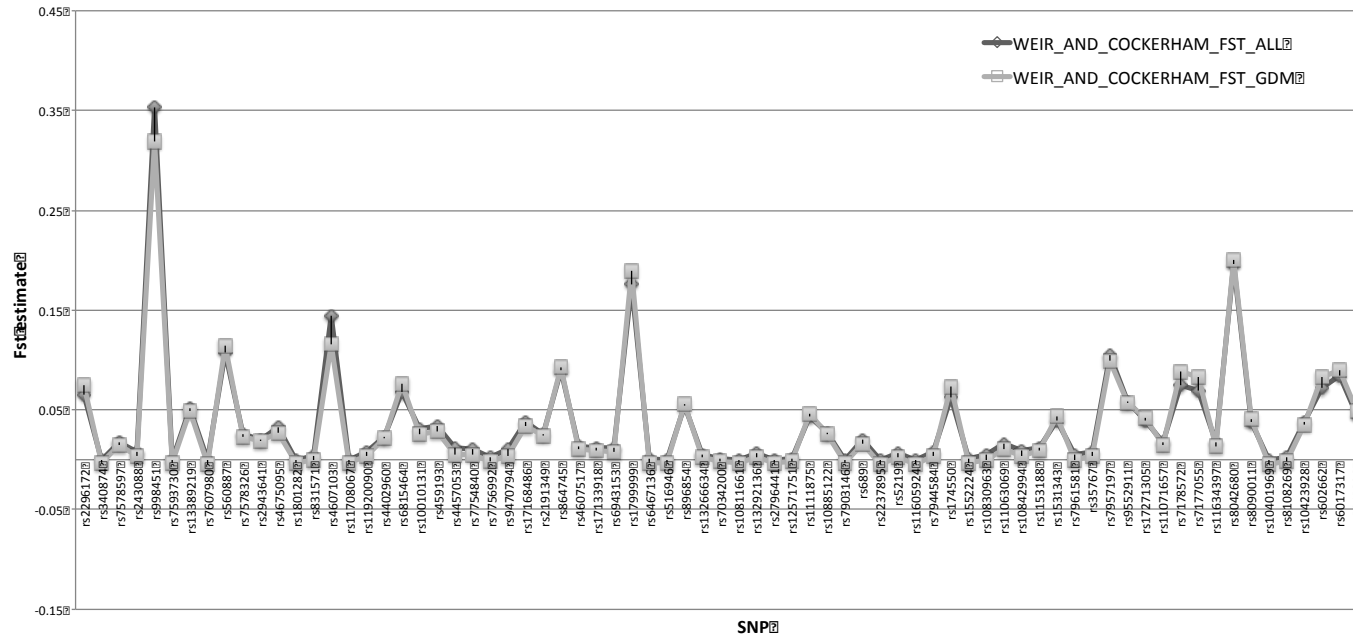


Figure 1. F_{ST} estimates for all (in grey) and GDM (in black). Great differentiation was observed at *TMEM163*, *PRC1* and *PPP1R3A* whereas moderate at *ADAMTS8*, *G6PC2*, *OASL*, *JAZF1*, *HNF4A*, *HMG20A*, *MAEA*, *MACF1*, *FADS1*, *SGCG*, *TP53INP1*, and *GRB14* loci.