### Novel DNA methylation sites of glucose and insulin homeostasis: an integrative cross-omics analysis

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### Abstract

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Despite existing reports on differential DNA methylation in type 2 diabetes (T2D) and obesity, our understanding of the functional relevance of the phenomenon remains limited. Because obesity is the main risk factor for T2D and a driver of methylation from previous study, we aimed to explore the effect of DNA methylation in the early phases of T2D pathology while accounting for body mass index (BMI). We performed a blood-based epigenome-wide association study (EWAS) of fasting glucose and insulin among 4,808 nondiabetic European individuals and replicated the findings in an independent sample consisting of 11,750 nondiabetic subjects. We integrated blood-based in silico cross-omics databases comprising genomics, epigenomics and transcriptomics collected by BIOS project of the Biobanking and BioMolecular resources Research Infrastructure of the Netherlands (BBMRI-NL), the Meta-Analyses of Glucose and Insulin-related traits Consortium (MAGIC), the DIAbetes Genetics Replication And Meta-analysis (DIAGRAM) consortium, and the tissue-specific Genotype-Tissue Expression (GTEx) project. We identified and replicated nine novel differentially methylated sites in whole blood (P-value  $< 1.27 \times 10^{-7}$ ): sites in LETM1, RBM20, IRS2, MAN2A2 genes and 1q25.3 region were associated with fasting insulin; sites in FCRL6, SLAMF1, APOBEC3H genes and 15q26.1 region were associated with fasting glucose. The association between SLAMF1, APOBEC3H and 15q26.1 methylation sites and glucose emerged only when accounted for BMI. Follow-up in silico cross-omics analyses indicate that the cis-acting meQTLs near SLAMF1 and SLAMF1 expression are involved in glucose level regulation. Moreover, our data suggest that differential methylation in FCRL6 may affect glucose level and the risk of T2D by regulating FCLR6 expression in the liver. In conclusion, the present study provided nine new DNA methylation sites associated with glycemia homeostasis and also provided new insights of glycemia related loci into the genetics, epigenetics and transcriptomics pathways based on the integration of cross-omics data in silico.

### Background

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Type 2 diabetes (T2D) is a common metabolic disease, characterized by disturbances in glucose and insulin metabolism, that are in part genetically driven 1-10 with the heritability ranging from 20% to 80% 11. DNA methylation has been associated with T2D as well as with fasting glucose and insulin<sup>12</sup>. Methylation-based risk scores of T2D predicted incident T2D cases that go beyond traditional risk factors such as obesity and waist-hip ratio<sup>13</sup>. Further, obesity, which is the most important determinant of insulin resistance and glucose levels in the population, <sup>14,15</sup> has also been associated with differential DNA methylation <sup>13</sup>. This raises the possibility that differential methylation associated with glucose and insulin levels could be counfounded by obesity. DNA methylation, mainly depending on the region, results in gene silencing and thus regulates gene expression and subsequent cellular functions<sup>16</sup>. It is very well possible that the epigenetic modifications occur in early phases of the pathology of T2D, requiring research focusing on the early process of the disease, e.g. in subjects free of diabetes. We aimed to determine the association of DNA methylation with fasting glucose and insulin accounting for the effect of obesity in the non-diabetic subjects and to evaluate the impact of DNA methylation on cross-omics level. We followed the hypothesis that genetic variants drive DNA methylation which subsequently regulates gene expression and then glycemic traits, changes of which mark the early phases of diabetes pathology (Figure 1a). First, we performed a blood-based epigenome-wide association study (EWAS) meta-analysis of 4,808 diabetes-free individuals of European descent and replicated our findings among 11 cohorts summing up to 11,750 trans-ethnic non-diabetic individuals, mainly from European ancestry. Subsequently, we explored the role of genetics in determining the regulation of methylation associated with glycemic traits and the effects of the differential methylation on the human transcriptome in silico (Figure 1a).

## Results

### 1. Blood-based epigenome-wide association analysis of glycemic traits

The discovery phase was based on four European cohorts (**Supplementary Table 1**). The meta-analysis revealed DNA methylation in 28 unique CpG sites associated with fasting glucose (11 CpG sites, n = 4,808) and/or insulin (20 CpG sites, n = 4,740) at epigenome-wide significance (P-value <  $1.27 \times 10^{-7}$ ) in either the baseline model without body mass index (BMI) adjustment or in the second model with BMI adjustment. Of these 28 CpG sites, 15 were novel (**Table 1**) while 13 were identified by earlier EWAS studies of either T2D or related traits, including glucose, insulin, hemoglobin A1c (HbA1c), homeostatic model assessment-insulin

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resistance (HOMA-IR) and BMI<sup>12,13,17-25</sup> (Supplementary Table 2). Of the known CpG sites, three located in SLC7A11, CPT1A, and SREBF1 associated with both glucose and insulin. The remaining ten CpG sites associated with insulin and were located in genes ASAM, DHCR24, RNF145, KDM2B, MYO5C, TMEM49, CPT1A, two in ABCG1 and one in the 4p15.33 region. The 15 novel CpGs were tested using the same models in meta-analysis of 11 independent cohorts including 11,750 non-diabetic subjects from the Cohorts for Heart and Aging Research in Genomic Epidemiology (CHARGE) consortium (Supplementary Table 1). As a result, nine unique CpG-trait associations were replicated: four passing the epigenome-wide significance threshold (P-value  $< 1.27 \times 10^{-7}$ ) and five passing the Bonferroni significance threshold after correcting for 15 tests (P-value  $< 3.3 \times 10^{-3}$ ) (Table 1). These included five sites associated with fasting insulin in the baseline model (LETM1, RBM20, IRS2, MAN2A2 and 1q25.3 region), one associated with fasting glucose in the baseline model (FCRL6) and additional three emerged to be associated with fasting glucose in the BMI-adjusted model (SLAMF1, APOBEC3H and 15q26.1 region). Because the replication cohorts also included other ethnic groups than the main European ancestry (European: n = 7,254, African: n = 3,744, and Hispanic: n = 543), we also performed meta-analysis stratified by ancestry. Seven out of nine new CpG sites (FCRL6, LETM1, RBM20, IRS2, MAN2A2, APOBEC3H and the 15q26.1 region) confirmed consistent directions of effect across the three ethnicities. (Supplementary Table 3) 2. Integrated in silico cross-omics studies To evaluate the functional relevance of differential methylation findings, we integrated our EWAS findings with genomics, epigenomics and transcriptomics data obtained from public resources. These included blood-based cis and trans methylation quantitative trait loci (meQTLs), expression quantitative trait methylations (eQTMs), expression quantitative trait loci (eQTLs) from the European BIOS database<sup>26</sup> from the Biobanking and BioMolecular resources Research Infrastructure of the Netherlands (BBMRI-NL), the genome-wide association studies (GWAS) of glycemic traits or T2D from the Meta-Analyses of Glucose and Insulin-related traits Consortium (MAGIC) and the DIAbetes Genetics Replication And Meta-analysis (DIAGRAM) consortium<sup>4-7,27</sup>, and tissue-specific eQTL-phenotype associations from MetaXcan database<sup>28,29</sup> based on Genotype-Tissue Expression (GTEx) project (See resources of these database in URLs). The hypotheses tested were outlined in Figure 1a: DNA methylation and gene expression are partly genetically driven and heritable; genetic variants determine in part methylation, which subsequently influences expression and further fasting glucose and/or insulin. While doing so, we centered on the 11 top independent DNA methylation sites previously identified

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(cg00574958 in CPT1A and cg06500161 in ABCG1 were used) and 9 novel sites from our current study (total n = 20 methylation sites). 2.1 Genomics of the differentially methylated sites involved in glycemic traits Using BIOS database (blood-based)<sup>26</sup>, we found that 2,991 single-nucleotide polymorphisms (SNPs) in 29 unique genetic loci were associated with methylation in either cis or trans across 18 unique CpG sites among the tested 20 target methylation sites. For two CpG sites located in SLC7A11 and LETM1, we did not find any significant meQTLs. Results are shown in Figure 2 and given in detail in Supplementary Table 4. Seven of the 29 meQTLs, 5 cis-acting and 2 trans-acting were found significantly associated with T2D, fasting glucose or HbA1c (shown in Figure 2 and given in detail Supplementary Table 5). Based on our leading hypothesis, we examined whether DNA methylation may influence fasting glucose and insulin in the circulation. To this end, we performed a two-sample-based Mendelian Randomization (MR) analyses<sup>30</sup> to examine the causal effect of the differential DNA methylation sites in blood on fasting glucose or insulin using the summary GWAS results from BIOS<sup>26</sup> and MAGIC databases<sup>5</sup> (Supplementary Table 6). Up to eight independent genetic variants were included in the genetic risk score as the instrumental variable of each methylation site to check the association with the observationally associated traits, either fasting glucose or fasting insulin. Thirteen CpG sites out of the initial 20 met the present MR criteria and were testes by MR. No significant associations were detected when adjusting for multiple testing involving 13 independent tests (Pvalue  $< 3.8 \times 10^{-3}$ ) except for two marginal significant associations between methylation site in RBM20 with fasting insulin (P-value = 0.04) and methylation site in SLAMF1 with fasting glucose (P-value = 0.05). 2.2 Transcriptome associated with the differentially methylated sites of glycemic traits 2.2.1 Association of gene expression with differentially methylated sites in blood To understand if the methylation is also eQTM, we investigated the association between gene expression and the 20 key glycemic methylation sites from the European blood-based BIOS database<sup>26</sup> (intergrated in **Figure** 2). We found that methylation in five CpG sites, including two novel sites (in FCRL6 and SLAMF1) and three known sites (in CPT1A, SREBF1 and ABCG1), was significantly negatively associated with the expression of their respective genes: FCRL6 (P-value =  $4.0 \times 10^{-39}$ ), SLAMF1 (P-value =  $4.1 \times 10^{-5}$ ), CPT1A (P-value =  $3.1 \times 10^{-20}$ ), SREBF1 (P-value =  $4.5 \times 10^{-15}$ ) and ABCG1 (P-value =  $2.2 \times 10^{-37}$ ). The methylation site in SLAMF1 was positively associated with expression of two other genes near *SLAMF1* (*CD244*: P-value =  $2.9 \times 10^{-6}$  and *SLAMF7*: P-value =  $5.4 \times 10^{-9}$ ). (Supplementary Table 7)

2.2.2 Common genetic determinants of glycemia related to DNA methylation and gene expression in blood We next explored if the genetic variants associated with the differential expression above were the same as the meQTLs using the eQTL data from the European blood-based BIOS database<sup>26</sup> (integrated in Figure 2 and detailed in Supplementary Table 8). We found three genetic determinants associated with both differential DNA methylation, including two novel methylation sites (in FCRL6 and SLAMF1) and one known (in SREBF1), and gene expression in blood. Rs1577544 near SLAMF1 associated with decreased methylation of the cg18881723 (Z = -5.45, P-value =  $5.1 \times 10^{-8}$ ) as well as *SLAMF1* expession (Z = -6.40, P-value =  $1.6 \times 10^{-10}$ ). Rs11265282 in FCRL6 associated with increased methylation of cg00936728 (Z = 4.17, P-value =  $3.0 \times 10^{-5}$ ) but decreased FCRL6 expession (Z = -6.73, P-value =  $1.7 \times 10^{-11}$ ). Rs6502629 in TOM1L2 associated with increased methylation of cg11024682 (Z = 9.97, P-value =  $2.1 \times 10^{-23}$ ) but decreased SREBF1 expession (Z = -17.93, Pvalue =  $7.2 \times 10^{-72}$ ). 2.2.3 Tissue-specific differential expression associated with T2D and related traits We then explored the tissue-specific differential expression associated with T2D and related traits by datamining from MetaXcan database from the GTEx project<sup>28,29</sup>. This analyses targeted on the eQTM-related expression of seven genes as shown in 2.2.1 in six glucose-metabolism-related tissues including blood, adipose subcutaneous, adipose visceral omentum, liver, pancreas, and muscle skeletal (Supplementary Table 9). The effect direction consistency was checked between methylation sites, gene expression and T2D or related traits. That meant the direction of the association between methylation and T2D or related traits should be a combination of the directions of methylation with gene expression and gene expression with T2D or related traits. The expression of SREBF1 in blood was significantly associated with decreased levels of HbA1c (Z = -3.26, P-value =  $1.1 \times 10^{-3}$ ) and also with decreased risk of T2D (Z = -2.40, P-value = 0.016). Meanwhile, the known cg11024682 in SREBF1 was positively associated with fasting glucose (Z = 6.45, P-value =  $2.7 \times 10^{-8}$ ) and fasting insulin (Z = 6.27, P-value =  $6.7 \times 10^{-9}$ ) and negatively associated with expression of SREBF1 in blood (Z = -7.84,

 $4.5 \times 10^{-15}$ ) (shown in **Figure 1c**). Higher liver gene expression of *FCRL6*, a novel locus, was associated with increased risk of T2D (Z = 2.14, P-value = 0.032) based on MetaXcan<sup>28,29,31</sup> results generated by integrating

functional data in liver<sup>32,33</sup> and the GWAS of T2D<sup>9</sup>. The novel cg00936728 in *FCRL6* was negatively associated

with fasting glucose (Z = -6.17, P-value =  $9.1 \times 10^{-8}$ ) and expression of FCRL6 in blood (Z = -13.09,  $4.0 \times 10^{-39}$ )

(shown in Figure 1b).

## Discussion

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The current large-scale EWAS identified and replicated nine new methylation sites associated with fasting glucose or insulin, including three additionally uncovered sites (in SLAMF1, APOBEC3H and the 15q26.1 region) associated with fasting glucose only after adjustment for BMI. We further validated 13 previous reported CpG sites in 11 independent loci. Based on the cross-omics analyses, our report complements earlier studies 12,13,17-<sup>25,34</sup> for multiple DNA methylation sites related to the pathology early in the development of T2D through genetics and/or gene expression. We also present in silico evidence supporting the potential involvement of the nine new methylation sites. The novel methylation sites annotated to genes that play roles in glucose and energy metabolism (IRS2), metabolism of proteins (MAN2A2 and EDEM3, the nearest gene of cg13222915), RNA and splicing regulation (RBM20), RNA metabolism (APOBEC3H), small molecule transport (LETM1) and immune system process (SLAMF1, FCRL6 and SV2B, the nearest gene of cg18247172). Some of these genes are also involved in other diseases or biomarkers, including inflammatory phenotypes (EDEM3 with systemic lupus erythematosus<sup>35</sup>, SLAMF1 with inflammatory bowel disease<sup>36</sup> and FCRL6 with C-reactive protein (CRP)<sup>37,38</sup>), cardiovascular phenotypes (RBM20 with electrocardiographic traits<sup>39</sup>), cancer (IRS2 with prostate cancer<sup>40</sup>) and schizophrenia (MAN2A2)<sup>41</sup>. Thus, observations provided insight into the pathways that might link T2D to inflammation, cardiovascular disease, cancer and schizophrenia, all disorders associated epidemiologically or clinically with T2D. This phenomenon may point at genetic pleiotropy of the genes, i.e. a gene codes the same products in various cells or have cascade-like signaling function that affects various targets. In this paper, we used the assumption that genetic variants drive DNA methylation which subsequently regulates gene expression and then glycemic traits<sup>42</sup>. Two pathways (on SREBF1 and FCRL6) related to genetics-epigenetics-transcriptomics-phenotype were observed in the present study (Figure 1b and Figure1c). We validated the differential methylation of SREBF1 in insulin metabolism<sup>43</sup> and extended the findings building a pathway based on the cascading cross-omics analysis in the assumption of genetics-epigeneticstranscriptomics-phenotype. We also discovered a new pathway of FCRL6 in glucose metabolism, which still needs further research for its role to be fully understood. From the present study, with the integration of all the significant associations, the effect allele (C) of the genetic variant rs11265282 in FCRL6 increases the methylation level which associates with lower expression of FCRL6 in the blood. The decreased FCRL6 expression in liver was also associates with decreased risk of T2D. This presumably is mediated by a decrease in fasting glucose.

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The present study provides new genomic targets for further work on the pathology of T2D through large-scale EWAS and replication. However, the main findings are based on data from blood which was the only accessible tissue and may not be representative of more glucose-relevant tissues, although concordance of differential methylation between blood and adipose is high for certain pathways<sup>44</sup>. Our present MR analyses yielded no evidence for causality between methylation sites and fasting glucose or insulin. One limitation we faced here was the limited data to perform MR in all the association steps, e.g. the association of methylation with gene expression, the gene expression with phenotypes, some CpG sites with phenotypes, as well as the inverse causal effect of glucose or insulin on DNA methylation, thus we can not exclude entirely the influence of glycemia homeostasis on methylation levels. On the other hand, some of the MR tests performed had low explained variance of the instrumental variables, i.e. seven of the 13 performed CpGs have instrumental variables explained variance less than 5%. This might partly explain the insignificant findings in MR in the current study. Further studies are needed to include additional biologically relevant tissues and perform MR based on the tissue specific meQLTs. In conclusion, our large-scale EWAS and replication have identified nine new differentially methylated sites associated with fasting glucose or insulin. The integrative in silico cross-omics analysis provided new insights of both known and new glycemia related loci into the genetics, epigenetics and transcriptomics pathway. Our study suggests that the expression of seven genes associated with either glycemia related DNA methylation is altered. Two of these seven expressed genes are also associated with T2D or related traits through the tissuespecific differentical expression association analysis: one known loci in SREBF1 and one new loci in FCRL6. Further biological functional experiments are required in more directly glucose-related tissues, e.g. pancreatic cells and liver, to unravel the mechanisms.

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Online methods Study population The discovery samples consisted of 4,808 European individuals without diabetes from four non-overlapped cohorts, recruited by Rotterdam Study III-1 (RS III-1, n = 626), Rotterdam Study II-3 and Rotterdam Study III-2<sup>45</sup> (called as RS-BIOS, n = 705), Netherlands Twin Register (NTR, n = 2,753) and UK adult Twin registry (18) (TwinsUK, n = 724). The replication sets contained up to 11,750 individuals from 11 independent cohorts from the Cohorts for Heart and Aging Research in Genomic Epidemiology (CHARGE), including up to 6,818 individuals from European ancestry, 4,355 from African ancestry and 577 from Hispanic ancestry. We excluded individuals with known diabetes, those on anti-diabetic treatment or fasting glucose ≥ 7mmol/l. Local research ethics committees approved each study, and all participants gave informed consent to each original study. The details of the cohorts and the study design are shown in Supplementary Note. **Glycemic traits and covariates** Venous blood samples were obtained after an overnight fast in all discovery and replication cohorts. Details of fasting glucose and insulin measurements are shown in Supplementary Note. Body mass index (BMI) was calculated as weight over height squared (kg/m²) based on clinical examinations. Smoking status was divided into current, former and never, based on questionnaires. White blood cell counts were quantified using standard laboratory techniques or predicted from methylation data using the standard Houseman method<sup>49</sup> (see **Supplementary Note** for each cohort). **DNA** methylation quantification The Illumina Human Methylation450 array was used in all discovery and replication cohorts to quantify genome-wide DNA methylation in blood samples. We obtained DNA methylation levels reported as β values, which represents the cellular average methylation level ranging from 0 (fully unmethylated) to 1 (fully methylated). Study-specific details regarding DNA methylation quantification, normalization and quality control procedures are provided in the Supplementary Note and Supplementary Table 1. Epigenome-wide association analysis and replication All statistical analyses were performed using R statistical software. Insulin was natural log transformed. In the discovery analysis, we first performed epigenome-wide association studies (EWAS) in each cohort separately. Linear regression analysis was used to test the association between glucose and insulin with each methylation site in the Rotterdam Study samples. Linear mixed models were used in NTR and TwinsUK accounting the

family structure. We fitted two models for each cohort: 1) the baseline model adjusting for age, sex, technical covariates (chip array number and position on the array), white blood cell counts (lymphocytes, monocytes, and granulocytes) and smoking status, and 2) a second model additionally adjusting for BMI. We removed probes that have evidence of multiple mapping or contain a genetic variant in the methylation site<sup>50</sup>. All cohort-specific EWAS results for each model were then meta-analysed using inverse variance-weighted fixed effects meta-analysis as implemented in the "metafor" R package<sup>51</sup>. In total, we meta-analysed 403,011 CpGs that passed quality control in all four discovery cohorts. The detail of the quality control for each cohort could be found in the **Supplementary Note**. The association was later corrected by the genomic control factor ( $\lambda$ ) in each meta-EWAS<sup>52</sup>. We produced quantile-quantile (QQ) plots of the -log<sub>10</sub> (P) to evaluate inflation in the test statistic (Supplementary Figure 1). A Bonferroni correction was used to correct for multiple testing and identify epigenome-wide significant results ( $P < 1.27 \times 10^{-7}$ ). We did not correct the number of glycemic traits and models, as they are highly correlated and not independent. The genome coordinates were provided by Illumina (GRCh37/hg19). The correlation of the CpG sites located in the same gene was further checked in the overall RS III-1 and RS-BIOS samples by Pearson's correlation test (n = 1,544) to find the independent top CpGs. For the associations discovered in the meta-EWAS that have not been reported previously, we attempted replication in independent samples using the same traits and models as in the discovery analyses. Studyspecific details of replication cohorts are provided in Supplementary Table 1 and Supplementary Note. Results from each replication cohort were meta-analysed using the same methods as in the discovery analyses. Bonferroni P-value  $< 3.3 \times 10^{-3}$  (0.05 corrected by 15 loci tested for associations) was considered significant.

### Genomics of the differentially methylated sites and glycemic traits

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We identified the genetic determinants of the significant CpG sites known or replicated through the current EWAS using the results of the *cis* and *trans* methylation quantitative trait loci (meQTLs) from European bloodbased BIOS database<sup>26</sup> from the Biobanking and BioMolecular resources Research Infrastructure of the Netherlands (BBMRI-NL) which captured meQTLs, expression quantitative trait loci (eQTLs) and expression quantitative trait methylations (eQTMs) from genome-wide database of 3,841 Dutch blood samples (See resources of the database in **URLs**). All the reported single-nucleotide polymorphisms (SNPs) with P-value adjusted for false discovery rate (FDR) less than 0.05 in the database were treated as the target genetic variants in the present study. The SNPs were annotated based on the information in the previous study<sup>26</sup> or the nearest protein-coding gene list from SNPnexus<sup>53,54</sup> on GRCh37/hg19.

We explored the associations of these DNA methylation-related SNPs with type 2 diabetes (T2D) or related traits, i.e. fasting glucose, insulin, hemoglobin A1c (HbA1c), based on public genome-wide association study (GWAS) datasets in European ancestry<sup>4-7,27</sup>. We checked the effect direction consistency of the association between the SNPs, methylation sites and T2D or related traits. That is the direction of the association between SNP and T2D or related traits should be a combination of the directions of SNP with methylation sites and methylation sites with T2D or related traits. A multiple-testing correction was performed by Bonferroni adjustment (P-value <  $1.8 \times 10^{-3}$ , 0.05 corrected by the 29 genetic loci shown in **Supplementary Table 4**). For the significant CpG sites known or replicated through EWAS, we attempted to evaluate the causality effect of CpGs on their significant traits, either fasting glucose or fasting insulin, using two-sample Mendelian Randomization (MR) approach as described in detail before by Dastani *et al*<sup>30,55,56</sup> based on the summary statistic GWAS results from BIOS database and the Meta-Analyses of Glucose and Insulin-related traits Consortium (MAGIC) database <sup>5,26</sup> (**Supplementary Figure 2**). Briefly, we constructed a weighted genetic risk score for individual CpG on phenotype using independent SNPs as the instrument variables of the CpG, implemented in the R-package "gtx". The effect of each score on phenotype was calculated as

$$ahat = \frac{\sum (\omega_i \beta_i / s_i^2)}{\sum (\omega_i^2 / s_i^2)}$$

, where  $\beta_i$  is the effect of the CpG-increasing alleles on phenotype,  $s_i$  its corresponding standard error and  $\omega_i$  the SNP effect on the respective CpG. Because the genetic variants might be close (cis) or far (trans) from the methylated site, we also performed MR test in the cis only SNPs if the CpG site has both cis and trans genetic markers. All SNPs were mapped to human genome build hg19. For each test (one CpG site with one trait), we extracted all the genetic markers of the CpG site in the fasting glucose or insulin GWAS from the MAGIC dataset (n = 96,496)<sup>5</sup> with their effect estimate and standard error on fasting glucose or insulin. Within the overlapped SNPs, we removed SNPs in potential linkage disequilibrium (LD, pairwise  $R^2 \ge 0.05$ ) in 1-Mbp window based on the 1000 Genome imputed genotype dataset from the general population: Rotterdam Study I (RSI, n = 6,291)<sup>45</sup>. We managed to exclude the genetic loci which were genome-wide significantly associated with glycemic traits, but none of the genetic loci meet this exclusion criteria. The instrumental variables that explain more than 1% of variance in exposure (DNA methylation) were taken forward for MR test. The Bonferroni P-value threshold was used to correct for the 13 CpG sites available for MR (P-value <  $3.8 \times 10^{-3}$ ).

## Gene expression analyses

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To explore whether the differential methylation sites were associated with differential expression in blood, we explored the European blood-based BIOS database for eQTMs<sup>26</sup>. The significantly associated gene expression probes were searched in the eQTL data in BIOS database<sup>26</sup>. We also investigated if the genetic variants associated with these gene expression probes in blood were also related to the DNA methylation sites with glycemia. Finally, we tested whether the expression of the genes that harbor the eQTMs was associated with T2D and related traits in glucose metabolism-related tissues (adipose subcutaneous, adipose visceral omentum, liver, whole blood, pancreas, and muscle skeletal) using MetaXcan <sup>28,29,31</sup> package. MetaXcan associates the expression of the genes with the phenotype by integrating functional data generated by largescale efforts, e.g Genotype-Tissue Expression (GTEx)<sup>32,33</sup> with that of the GWAS of the trait. MetaXcan is trained on transcriptome models in 44 human tissues from GTEx and is able to estimate their tissue-specific effect on phenotypes from GWAS. For this study we used the GWAS studies of T2D<sup>9</sup>, fasting glucose traits<sup>5,6</sup>, fasting insulin<sup>6</sup>, hemoglobin A1c (HbA1c)<sup>57</sup> and homeostatic model assessment-insulin resistance (HOMA-IR)<sup>4</sup>. We used the nominal P-value threshold (P-value = 0.05) as we had separate assumptions for each terminal pathway between gene expressions and phenotype. Further, we checked the effect direction consistency of the association between the methylation sites and fasting glucose or insulin with the combination of the associations between the methylation sites and gene expression and between the gene expression and T2D or related traits. URLs. BIOS database, https://genenetwork.nl/biosqtlbrowser/; SNPnexus, http://snp-nexus.org/index.html; GWAS database of glycemic traits, https://www.magicinvestigators.org/; GWAS database of T2D, http://diagram-consortium.org/; MetaXcan, https://s3.amazonaws.com/imlabopen/Data/MetaXcan/results/metaxcan\_results\_database\_v0.1.tar.gz. (available: 1st Jan, 2018)

Table1 Epigenome-wide association study (EWAS) results: novel differentially methylated sites associated with fasting glucose or insulin at an epigenome-wide significance level

							Discovery	phase (EA)		ı	Replication pha	se (EA+AA	\+HS)
Locus	СрG	Chr	Position	Regulatory feature	Trait (s)		Nodel 1	N	lodel 2	N	Model 1	M	lodel 2
						Z	P-value*	Z	P-value*	Z	P-value	Z	P-value
FCRL6	cg00936728	1	159772194	NA	Glucose	-6.17	9.1 × 10 <sup>-8</sup>	-5.71	1.9 × 10 <sup>-7</sup>	-3.90	9.55 × 10 <sup>-5</sup>	NP	NP
SLAMF1	cg18881723	1	160616870	Promoter associated	Glucose	6.11	7.5 × 10 <sup>-8</sup>	6.94	3.4 × 10 <sup>-10</sup>	2.67	7.66 × 10 <sup>-3</sup>	3.23	1.2 × 10 <sup>-3</sup>
1q25.3	cg13222915	1	184598594		Insulin	-6.50	2.6 × 10 <sup>-9</sup>	-4.61	4.1 × 10 <sup>-6</sup>	-8.16	3.33 × 10 <sup>-16</sup>	NP	NP
BRE	cg20657709	2	28509570	NA	Glucose	-5.46	2.7 × 10 <sup>-6</sup>	-5.88	4.1 × 10 <sup>-8</sup>	NP	NP	-2.10	0.036
LRPPRC	cg01913188	2	44223249	Promoter associated	Glucose	5.13	9.4 × 10 <sup>-6</sup>	6.27	5.7 × 10 <sup>-9</sup>	NP	NP	0.12	0.90
IRAK2	cg14527942	3	10276383	NA	Insulin	6.97	3.4 × 10 <sup>-10</sup>	6.69	2.9 × 10 <sup>-11</sup>	-0.70	0.48	-0.75	0.45
LETM1	cg13729116	4	1859262	Promoter associated	Insulin	5.95	4.3 × 10 <sup>-8</sup>	4.56	4.5 × 10 <sup>-6</sup>	4.96	6.98 × 10 <sup>-7</sup>	NP	NP
RBM20	cg15880704	10	112546110	NA	Insulin	6.41	3.8 × 10 <sup>-9</sup>	1.38 4.06	6.7 × 10 <sup>-5</sup>	6.83	8.62 × 10 <sup>-12</sup>	NP	NP
IRS2	cg25924746	13	110432935	Promoter associated	Insulin	6.59	3.0 × 10 <sup>-9</sup>	4.55	4.9 × 10 <sup>-6</sup>	6.65	3.01 × 10 <sup>-11</sup>	NP	NP
SPTB	cg07119168	14	65225253	NA	Glucose	-5.86	4.4 × 10 <sup>-7</sup>	-5.82	4.9 × 10 <sup>-8</sup>	NP	NP	-1.81	0.070
15q26.1	cg18247172	15	91370233	NA	Glucose	-5.25	4.9 × 10 <sup>-6</sup>	-5.90	2.8 × 10 <sup>-8</sup>	NP	NP	-3.47	5.1 × 10 <sup>-4</sup>
MAN2A2	cg20507228	15	91460071	Promoter associated (Cell type specific)	Insulin	5.90	5.5 × 10 <sup>-8</sup>	4.83	9.0 × 10 <sup>-7</sup>	7.93	2.28 × 10 <sup>-15</sup>	NP	NP
FAM92B	cg06709610	16	85143924	NA	Insulin	6.35	6.5 × 10 <sup>-9</sup>	7.24	5.8 × 10 <sup>-13</sup>	0.24	0.81	0.55	0.59
CD300A	cg08087047	17	72461209	NA	Glucose	-5.19	5.9 × 10 <sup>-6</sup>	-5.80	1.1 × 10 <sup>-7</sup>	NP	NP	-1.08	0.28
АРОВЕСЗН	cg06229674	22	39492189	NA	Glucose	-5.40	1.8 × 10 <sup>-6</sup>	-5.86	4.7 × 10 <sup>-8</sup>	NP	NP	-4.82	1.4 × 10 <sup>-6</sup>

Genome-wide DNA methylation sites were tested for association with fasting glucose or fasting insulin in two models. Novel epigenome-wide significant (P-value <  $1.27 \times 10^{-7}$ ) results in the discovery phase and the replication are shown. Model 1 adjusted for age, sex, technical covariates, white blood cell, and smoking status, accounting for family structure if needed in each cohort. Model 2 adjusted for BMI additionally. The significant associations in non-reported CpG sites were promoted for replication of the same models and traits. NP: Replication was not performed in the non-significant associated model or trait. Locus: the cytogenetic location or the gene symbol of the CpGs from Illumina annotation. Regulatory feature: the regulatory feature group of the CpGs from Illumina annotation. Chr: Chromosome. \* Genomic controlled P-value. EA+AA+HA: European ancestry, African ancestry and Hispanic ancestry population. Z: effect estimate per standard error. **Bold print**: Significant results (P-value <  $1.27 \times 10^{-7}$  in the discovery phase, P-value <  $3.3 \times 10^{-3}$  in the replication phase). NA: Not available.

# **Supplementary Table 1 Characteristics of cohorts**

		Discover	y cohorts									Rej	plication cohort	S							
COHORT	RS III-1	RS-BIOS	NTR	Twins UK	ARIC	BL	SA	С	HS	FHS	GENOA	GOLDN	HyperGEN	inCHIANTI	KORA		WHI-BAA23			WHI-EMPC	
Ethnicity	European	European	European	European	African American	European	African American	European	African American	European	African American	European	African American	European	European	European	African American	Hispanic	European	African American	Hispanic
Fasting glucose (N)	626	705	2753	724	1875	402	142	160	147	2117	268	917	469	433	1488	836	503	319	425	951	258
Fasting insulin (N)	626	705	2685	724	NA	402	142	160	147	2157	267	915	466	421	1488	817	495	314	413	917	246
Fasting glucose (mmol/L)	5.3 (0.5)	5.4 (0.6)	5.2 (0.6)	5.0 (0.5)	5.7 (0.6)	5.3 (0.5)	5.3 (0.5)	5.5 (0.5)	5.5 (0.6)	5.6 (0.5)	5.3 (0.6)	5.48 (0.67)	5.37 (1.3)	4.9 ( 0.6)	5.3 (0.5)	5.28 (0.5)	5.2 (0.6)	5.2 (0.5)	5.2 (0.6)	5.2 (0.6)	5.2 (0.6)
Fasting insulin (pmol/L; mlU/L)	89.0 (44.3)	79.4 (44.4)	57.0 (36.1)	57.0 (53.6)	NA	58.8 (38.9)	78.7 (54.2)	76.3 (38.8)	91.4 (63.7)	4.07 (0.6)	62.4 (55.8)	13.7 (7.5)	9.4 (9.0)	72.8 (37.9)	45.8 (82.9)	5.3 (0.8)	5.79 (0.7)	5.7 (0.7)	90.2 (44.3)	75.2 (41.0)	88.3 (46.6)
Age (years)	59.8 (8.1)	67.5 (6.0)	36.3 (12.8)	58.1 (9.3)	56.1 (5.8)	69.0 (14.4)	64.3 (10.9)	76.2 (5.0)	73.2 (5.7)	65.7 (8.9)	60.7 (7.7)	48.0 (15.8)	51.0 (13.7)	62.1 (16.0)	60.2 (8.8)	68.4 (6.2)	62.6 (6.7)	62.4 (6.8)	62.1 (6.9)	64.5 (7.2)	61.4 (6.1)
BMI (kg/m2)	27.4 (4.5)	27.6 (4.1)	24.1 (3.7)	26.4 (4.6)	29.3 (6.0)	26.4 (4.4)	287 (4.9)	26.3 (4.6)	28.4 (4.7)	27.7 (4.9)	30.5 (6.4)	28.5 (5.6)	30.7 (7.0)	27.0 (3.9)	27.7 (4.5)	28.2 (5.4)	30.9 (6.4)	28.7 (4.9)	31.2 (6.0)	28.4 (5.7)	29.1 (4.8)
Female (%)	351 (56)	414 (59)	1817 (66)	684 (100)	1198 (64)	202 (50)	85 (60)	89 (56)	99 (67)	1204 (57)	191 (71)	488 (49)	365 (60)	234 (54)	783 (53)	836 (100)	503 (100)	319 (100)	425 (100)	951 (100)	258 (100)
Never Smoker (%)	187 (30)	244 (35)	1625 (59)	423 (62)	841 (45)	219 (54)	90 (63)	73 (46)	74 (50)	771 (37)	151 (56)	700 (70.3)	84 (17.8)	244 (56)	662 (44)	438 (52)	234 (467)	188 (59)	207 (49)	482 (51)	168 (65)
Former Smoker (%)	270 (43)	389 (55)	633 (23)	197 (30)	558 (30)	173 (43)	48 (34)	64 (40)	51 (35)	1180 (56)	69 (26)	209 (21)	207 (44.9)	105 (24)	607 (41)	381 (46)	258 (51)	123 (59)	176 (41)	402 (42)	73 (28)

NA: Not available. The unit of fasting insulin in FHS, GOLDN, HyperGEN, WHI-BAA23 are mIU/L; the units of fasting insulin in other cohorts are pmol/L.

# Supplementary Table 2 Epigenome-wide association study (EWAS) results: known differentially methylated sites associated with fasting glucose or insulin at epigenome-wide significance level in the discovery phase

Locus	СрG	Chr	Position	Regulatory feature	Trait (s)	ı	Model 1	N	lodel 2	Previous evidence
Locus	Сро	Cili	rosition	Regulatory reacure	maic (3)	Z	P-value*	Z	P-value*	. Previous evidence
DHCR24	cg17901584	1	55353706	Promoter associated (Cell type specific)	Insulin	-6.19	2.3 × 10 <sup>-8</sup>	-3.58	3.7 × 10 <sup>-4</sup>	BMI <sup>13</sup> ; Insulin <sup>13</sup> ; HbA1c <sup>13</sup> ; Incident T2D <sup>13</sup>
4p15.33	cg10438589	4	14531493	NA	Insulin	6.25	2.4 × 10 <sup>-8</sup>	3.56	5.0 × 10 <sup>-4</sup>	BMI <sup>13</sup> ; Insulin <sup>13</sup> ; Incident T2D <sup>13</sup>
SLC7A11	cg06690548	4	139162808	NA	Glucose	-7.70 -6.21	7.6 × 10 <sup>-10</sup> 2.4 × 10 <sup>-8</sup>	<b>-5.85</b> -3.68	7.8 × 10 <sup>-8</sup> 2.8 × 10 <sup>-4</sup>	BMI <sup>13</sup> ; Glucose <sup>13</sup> ; Insulin <sup>13</sup> ; Incident T2D <sup>13</sup>
RNF145	cg26403843	5	158634085	NA	Insulin	6.29	8.5 × 10 <sup>-9</sup>	4.60	7.7 × 10 <sup>-6</sup>	BMI <sup>13,17,20</sup> ; Insulin <sup>13</sup> ; Incident T2D <sup>13</sup>
СРТ1А	cg00574958	11	68607622	NA	Glucose Insulin	-7.63 -8.00	$2.9 \times 10^{-11}$ $3.5 \times 10^{-13}$	-5.56 -4.11	$2.5 \times 10^{-7}$ $3.5 \times 10^{-5}$	BMI <sup>13,17,18,20</sup> ; Glucose <sup>13,22</sup> ; Insulin <sup>13</sup> ; HbA1c <sup>13</sup> ; Incident  T2D <sup>13</sup> ; Prevalent T2D <sup>17</sup>
CPT1A	cg17058475	11	68607737	NA	Insulin	-6.36	7.4 × 10 <sup>-9</sup>	-4.16	3.4 × 10 <sup>-5</sup>	BMI <sup>13</sup> ; Glucose <sup>13</sup> ; Insulin <sup>13</sup> ; HbA1c <sup>13</sup> ; Incident T2D <sup>13</sup>
ASAM	cg26894079	11	122954435	NA	Insulin	-5.83	7.5 × 10 <sup>-8</sup>	-2.95	3.3 × 10 <sup>-3</sup>	BMI <sup>13</sup> ; Insulin <sup>13</sup> ; Incident T2D <sup>13</sup>
KDM2B	cg13708645	12	121974305	Promoter associated	Insulin	6.00	1.1 × 10 <sup>-7</sup>	3.28	8.8 × 10 <sup>-4</sup>	BMI <sup>17,20</sup>
MYO5C	cg06192883	15	52554171	Unclassified	Insulin	7.96	6.4 × 10 <sup>-13</sup>	4.57	4.4 × 10 <sup>-6</sup>	BMI <sup>13,17,20</sup> ; Insulin <sup>13</sup> ; Incident T2D <sup>13</sup>
SREBF1	cg11024682	17	17730094	Unclassified (Cell type specific)	Glucose Insulin	6.45 6.27	2.7 × 10 <sup>-8</sup> 6.7 × 10 <sup>-9</sup>	4.34 2.69	$6.4 \times 10^{-5}$ $8.6 \times 10^{-3}$	BMI <sup>13,20,24</sup> ; Glucose <sup>13,22</sup> ; Insulin <sup>13</sup> ; HbA1c <sup>13</sup> ; Incident  T2D <sup>13,19</sup> ; Prevalent T2D <sup>17</sup>
TMEM49	cg24174557	17	57903544	NA	Insulin	-7.57	8.8 × 10 <sup>-12</sup>	-4.00	6.7 × 10 <sup>-5</sup>	BMI <sup>13,17</sup> ; Insulin <sup>13</sup> ; Incident T2D <sup>13</sup>
ABCG1	cg27243685	21	43642366	NA	Insulin	7.55	5.9 × 10 <sup>-12</sup>	5.10	4.5 × 10 <sup>-7</sup>	BMI <sup>13,17,20,24,25</sup> ; Glucose <sup>13</sup> ; Insulin <sup>13</sup> ; HbA1c <sup>13</sup> ; Incident  T2D <sup>13</sup>
ABCG1	cg06500161	21	43656587	NA	Insulin	10.16	< 2.2 × 10 <sup>-16</sup>	6.68	5.0 × 10 <sup>-11</sup>	BMI <sup>13,17,20,21</sup> ; Glucose <sup>13,22</sup> ; Insulin <sup>12,13,22</sup> ; HbA1c <sup>13</sup> ; Incident T2D <sup>13,19</sup> ; Prevalent T2D <sup>17,23</sup> ; 2h glucose <sup>22</sup> ; HOMA-IR <sup>12</sup>

Genome-wide DNA methylation sites were tested for association with fasting glucose or fasting insulin in two models. Previously reported epigenome-wide significant (P-value  $< 1.27 \times 10^{-7}$ ) results in the discovery phase and the previous evidence from the EWAS with T2D or related traits in the same CpG sites are shown. Model 1 adjusted for age, sex, technical covariates, white blood cell, and smoking status, accounting for family structure if needed in each cohort. Model 2 adjusted for BMI additionally. Locus: the cytogenetic location or the gene symbol of the CpGs from Illumina annotation. Regulatory feature: the regulatory feature group of the CpGs from Illumina annotation. Chr: Chromosome. \* Genomic controlled P-value. Z: effect estimate per standard error. **Bold print**: Epigenome-wide significant results (P-value  $< 1.27 \times 10^{-7}$ ). NA: Not available.

Supplementary Table 3 Epigenome-wide association study (EWAS) results: replication of newly discovered differentially methylated sites in different ancestry populations

						Replicati	on in EA			Replicati	on in AA			Replication	on in HA	
Locus	СрG	Chr	Position	Trait (s)	ı	Model 1	N	lodel 2	M	odel 1	M	odel 2	М	odel 1	Мо	del 2
					Z	P-value	Z	P-value	Z	P-value	Z	P-value	Z	P-value	Z	P-value
FCRL6	cg00936728	1	159772194	Glucose	-2.48	0.013	NP	NP	-2.90	3.8 × 10 <sup>-3</sup>	NP	NP	-1.84	0.066	NP	NP
SLAMF1	cg18881723	1	160616870	Glucose	2.68	7.3 × 10 <sup>-3</sup>	3.07	2.1 × 10 <sup>-3</sup>	1.14	0.25	1.59	0.11	-0.56	0.58	-0.49	0.63
1q25.3	cg13222915	1	184598594	Insulin	-6.85	7.6 × 10 <sup>-12</sup>	NP	NP	-5.26	1.4 × 10 <sup>-7</sup>	NP	NP	0.10	0.092	NP	NP
BRE	cg20657709	2	28509570	Glucose	NP	NP	-0.93	0.35	NP	NP	-2.43	0.015	NP	NP	0.23	0.82
LRPPRC	cg01913188	2	44223249	Glucose	NP	NP	0.08	0.94	NP	NP	-0.27	0.79	NP	NP	0.99	0.32
IRAK2	cg14527942	3	10276383	Insulin	-1.06	0.29	-1.15	0.25	0.63	0.53	0.45	0.66	0.21	0.83	0.65	0.51
LETM1	cg13729116	4	1859262	Insulin	4.63	3.6 × 10 <sup>-6</sup>	NP	NP	1.69	0.092	NP	NP	NP	NP	0.62	0.54
RBM20	cg15880704	10	112546110	Insulin	6.83	8.6 × 10 <sup>-12</sup>	NP	NP	1.22	0.23	NP	NP	NP	NP	0.55	0.59
IRS2	cg25924746	13	110432935	Insulin	6.20	5.7 × 10 <sup>-10</sup>	NP	NP	1.85	0.064	NP	NP	NP	NP	1.65	0.10
SPTB	cg07119168	14	65225253	Glucose	NP	NP	-0.57	0.57	NP	NP	-2.23	0.026	NP	NP	-0.97	0.33
15q26.1	cg18247172	15	91370233	Glucose	NP	NP	-1.69	0.092	NP	NP	-3.66	2.6 × 10 <sup>-4</sup>	NP	NP	-1.18	0.24
MAN2A2	cg20507228	15	91460071	Insulin	7.59	3.2 × 10 <sup>-14</sup>	NP	NP	1.59	0.11	NP	NP	2.75	6.0 × 10 <sup>-3</sup>	NP	NP
FAM92B	cg06709610	16	85143924	Insulin	0.82	0.41	0.83	0.41	-1.93	0.053	-1.37	0.17	1.13	0.26	1.65	0.10
CD300A	cg08087047	17	72461209	Glucose	NP	NP	-0.95	0.34	NP	NP	-0.21	0.83	NP	NP	-0.96	0.34
APOBEC3H	cg06229674	22	39492189	Glucose	NP	NP	-3.88	1.0 × 10 <sup>-4</sup>	NP	NP	-2.70	6.9 × 10 <sup>-3</sup>	NP	NP	-0.99	0.32

Replication of the epigenome-wide significant (P-value  $< 1.27 \times 10^{-7}$ ) CpGs with fasting glucose or insulin stratified by different ancestry populations. Model 1 adjusted for age, sex, technical covariates, white blood cell, and smoking status, accounting for family structure if needed in each cohort. Model 2 adjusted for BMI additionally. EA:

European ancestry (n = 6,778 for fasting glucose; n = 6,773 for fasting insulin) AA: African ancestry (n = 4,355 for fasting glucose; n = 2,434 for fasting insulin). HA: Hispanic ancestry (n = 577 for fasting glucose; n = 560 for fasting insulin). NP: Replication was not performed in the non-significant associated model or trait from the discovery phase. Locus: the cytogenetic location or the gene symbol of the CpGs from Illumina annotation. Chr: Chromosome. Z: effect estimate per standard error. **Bold print**: Bonferroni significant results (P-value <  $3.3 \times 10^{-3}$ ).

# Supplementary Table 4 Methylation quantitative trait loci (meQTLs) for known or new replicated CpG sites

Locus (CpG)	CpG	Variant	Chr	Position	Locus (meQTL)	Type (meQTL)	MAF	Alleles	EA	Z	P-value	Cis/Trans
DHCR24	cg17901584	rs7412	19	45412079	APOE	Protein coding	0.08	C/T	Т	6.39	1.7 × 10 <sup>-10</sup>	Trans
DHCR24	cg17901584	rs7701414	5	131585958	P4HA2	Protein coding	0.22	A/G	G	5.26	1.4 × 10 <sup>-7</sup>	Trans
DHCR24	cg17901584	rs174550	11	61571478	FADS1/FADS2	Protein coding	0.30	T/C	С	-6.72	1.8 × 10 <sup>-11</sup>	Trans
DHCR24	cg17901584	rs735665	11	123361397	GRAMD1B (Nearest)	Protein coding	0.10	G/A	А	10.59	3.4 × 10 <sup>-26</sup>	Trans
DHCR24	cg17901584	rs687565	1	55364663	TMEM61 (Nearest)	Protein coding	0.43	C/A	С	11.95	6.2 × 10 <sup>-33</sup>	Cis
FCRL6	cg00936728	rs6657365	1	159782549	FCRL6	Protein coding	0.23	C/G	G	5.04	4.6 × 10 <sup>-7</sup>	Cis
FCRL6	cg00936728	rs2523946	6	29941943	HCG9 (Nearest)	lincRNA	0.49	C/T	Т	5.33	9.8 × 10 <sup>-8</sup>	Trans
SLAMF1	cg18881723	rs3129055	6	29670261	ZFP57 (Nearest)	Protein coding	0.29	A/G	G	6.37	1.9 × 10 <sup>-10</sup>	Trans
SLAMF1	cg18881723	rs11265461	1	160630143	SLAMF1 (Nearest)	Protein coding	0.36	C/T	С	8.39	4.7 × 10 <sup>-17</sup>	Cis
1q25.3	cg13222915	rs72737737	1	184598732	C1orf21	Protein coding	0.06	A/G	G	5.68	1.4 × 10 <sup>-8</sup>	Cis
4p15.33	cg10438589	rs16890352	4	14385522	AC006296.1	lincRNA	0.16	A/G	G	-11.64	2.6 × 10 <sup>-31</sup>	Cis
RNF145	cg26403843	rs7529925	1	199007208	RP11-16L9.4	lincRNA	0.21	T/C	С	5.68	1.3 × 10 <sup>-8</sup>	Trans
RNF145	cg26403843	rs7732603	5	158614357	RNF145	Protein coding	0.48	A/C	С	35.61	$8.6 \times 10^{-278}$	Cis
RBM20	cg15880704	rs7906643	10	112545494	RBM20	Protein coding	0.09	C/T	Т	-29.45	$1.4 \times 10^{-190}$	Cis
CPT1A	cg00574958	rs964184	11	116648917	ZNF259	Protein coding	0.22	C/G	G	-5.54	3.0 × 10 <sup>-8</sup>	Trans
ASAM	cg26894079	rs34817879	11	123023729	CLMP	Protein coding	0.12	T/G	G	-4.67	3.0 × 10 <sup>-6</sup>	Cis
KDM2B	cg13708645	rs60370741	12	121966676	KDM2B	Protein coding	0.30	T/C	С	37.16	$2.6 \times 10^{-302}$	Cis
IRS2	cg25924746	rs9521528	13	110504805	IRS2 (Nearest)	Protein coding	0.44	T/A	Т	-13.64	2.4 × 10 <sup>-42</sup>	Cis

					LINC00396						-4	
IRS2	cg25924746	rs7984800	13	110671951	(Nearest)	lincRNA	0.26	A/G	G	3.88	$1.0 \times 10^{-4}$	Cis
MYO5C	cg06192883	rs1047891	2	211540507	CPS1	Protein coding	0.29	A/C	Α	-6.01	1.8 × 10 <sup>-9</sup>	Trans
MYO5C	cg06192883	rs71472932	15	52541976	MYO5C	Protein coding	0.11	G/A	Α	4.22	2.4 × 10 <sup>-5</sup>	Cis
15q26.1	cg18247172	rs404623	15	91367271	BLM (Nearest)	Protein coding	0.50	G/C	G	-11.70	1.3 × 10 <sup>-31</sup>	Cis
15q26.1	cg18247172	rs3129055	6	29670261	ZFP57 (Nearest)	Protein coding	0.29	A/G	G	-5.22	1.8 × 10 <sup>-7</sup>	Trans
15q26.1	cg18247172	rs4324798	6	28776117	AL662890.3	miRNA	0.04	G/A	Α	-5.48	4.3 × 10 <sup>-8</sup>	Trans
MAN2A2	cg20507228	rs9374080	6	109616420	CCDC162P	pseudogene	0.27	C/T	С	-5.31	1.1 × 10 <sup>-7</sup>	Trans
	20507220	25224262		04466363	MAN2A2		0.17	0/7		0.14	4.0-16	<u> </u>
MAN2A2	cg20507228	rs35831960	15	91466262	(Nearest)	Protein coding	0.17	C/T	Т	8.14	$4.1 \times 10^{-16}$	Cis
SREBF1	cg11024682	rs7701414	5	131585958	P4HA2	Protein coding	0.22	A/G	G	5.21	$1.9 \times 10^{-7}$	Trans
SREBF1	cg11024682	rs7529925	1	199007208	RP11-16L9.4	lincRNA	0.21	T/C	С	-5.15	2.6 × 10 <sup>-7</sup>	Trans
SREBF1	cg11024682	rs6502629	17	17869642	TOM1L2	Protein coding	0.22	G/A	G	9.97	2.1 × 10 <sup>-23</sup>	Cis
TMEM49	cg24174557	rs3774937	4	103434253	NFKB1	Protein coding	0.25	C/T	С	-7.75	9.3 × 10 <sup>-15</sup>	Trans
ABCG1	cg06500161	rs225443	21	43658206	ABCG1	Protein coding	0.40	G/A	Α	-7.16	8.1 × 10 <sup>-13</sup>	Cis
4 D O D C O U	06220674	20F024C4	22	20406502	APOBEC3G	Dontoin andio	0.11	T/C		0.24	9.8 × 10 <sup>-17</sup>	C:-
APOBEC3H	cg06229674	rs28583464	22	39486593	(Nearest)	Protein coding	0.11	T/C	С	8.31	9.8 × 10	Cis

Based on the European blood-based BIOS database (n = 3,841),<sup>26</sup> the meQTL information of known or new replicated CpG sites are shown. Locus (CpG): the cytogenetic location or the gene symbol of the CpGs from Illumina annotation. Locus (meQTL): the located or nearest protein-coding gene of the meQTL from UCSC annotation. Type (meQTL): the gene type of the meQTL. Chr: chromosome. MAF: minor allele frequency. EA: effect allele. Z: effect estimate per standard error.

Supplementary Table 5 Common genetic determinants of glycemia related methylation sites and T2D or related traits in blood

								Association wi	th CnG		Association	n with T20	or related
Variant	Locus	Type (meQTL)	Chr	Position	MAF	EA		Association wi	сро			traits <sup>†</sup>	
variant	(meQTL)	Type (meQTL)	CIII	rosidon	IVIAI	LA	СрG	Locus (CpG)	Z	P-value	Trait	z	P-value
rs6701489	TMEM61 (Nearest)	Protein coding	1	55358459	0.07	Т	cg17901584	DHCR24	4.82	1.4 × 10 <sup>-6</sup>	FG <sup>6</sup>	-3.43	8.5 × 10 <sup>-4</sup>
rs6896438	RNF145 (Nearest)	Protein coding	5	158547876	0.36	С	cg26403843	RNF145	6.15	8.0 × 10 <sup>-10</sup>	FI <sup>5</sup>	3.82	1.4 × 10 <sup>-4</sup>
rs10849885	KDM2B	Protein coding	12	121881848	0.32	Α	cg13708645	KDM2B	29.33	$4.2 \times 10^{-189}$	FG <sup>6</sup>	4.17	2.2 × 10 <sup>-5</sup>
rs9374080	CCDC162P	pseudogene	6	109616420	0.27	С	cg20507228	MAN2A2	-5.31	1.1 × 10 <sup>-7</sup>	HbA1c <sup>58</sup>	-5.11	$2.0 \times 10^{-7}$
rs3818717	RAI1	Protein coding	17	17707105	0.06	Т	cg11024682	SREBF1	8.93	4.1 × 10 <sup>-19</sup>	T2D <sup>27</sup>	1.08	4.9 × 10 <sup>-4</sup>
rs7529925	RP11-16L9.4	lincRNA	1	199007208	0.21	С	cg11024682	SREBF1	-5.15	2.6 × 10 <sup>-7</sup>	HbA1c <sup>58</sup>	-3.60	2.5 × 10 <sup>-4</sup>
rs16960744	TOM1L2	Protein coding	17	17755259	0.37	Α	cg11024682	SREBF1	4.87	1.1 × 10 <sup>-6</sup>	HbA1c <sup>58</sup>	3.11	1.5 × 10 <sup>-3</sup>

The common genetic determinants of glycemia related methylation sites and T2D or related traits are shown. Chr: chromosome. Locus (meQTL): the located or nearest protein-coding gene of the meQTL from UCSC annotation. Type (meQTL): the gene type of the meQTL. MAF: minor allele frequency. EA: effect allele. Locus (CpG): the cytogenetic location or the gene symbol of the CpGs from Illumina annotation. Z: effect estimate per standard error. FG: fasting glucose. FI: fasting insulin. Data sources of associations: 1) association with CpG was from the current discovery phase (n = 4,808), 2) associations with FG (n = 133,010), FI (n = 96,496), T2D (case/control: n = 81,412/370,832) and HbA1c (n = 159,940) were from the MAGIC and DIAGRAM GWAS database<sup>5,6,27,58</sup>

# Supplementary Table 6 Mendelian randomization (MR) results

Exposure	Locus (CpG)	Trait	No. of SNPs	R <sup>2</sup> (%)	Effect	SE	P- value	Heter ogene ity P- value	Type of SNPs	SNPs list	Z (exposure)	Z (outcome)
cg00574958	CPT1A	NP	1	0.79	NP	NP	NP	NP	All-Trans	rs964184	-5.54	0.42
cg06192883	MYO5C	NP	1	0.91	NP	NP	NP	NP	All-Trans	rs715	-5.94	1.77
cg06229674	АРОВЕСЗН	FG	1	1.71	0.09	0.12	0.48	NA	All-Cis	rs6001423	8.17	0.71
cg06500161	ABCG1	FI	2	2.30	0.09	0.11	0.42	0.27	All-Cis	rs225443;rs225391	-7.16;6.18	-1.33;-0.31
cg10438589	4p15.33	FI	4	7.15	-0.03	0.06	0.60	0.50	All-Cis	rs16890358;rs9291625;rs13131008;rs1 0488977	-11.63;- 7.83;6.91;6.06	-0.3;1.23;0.43;- 0.94
cg11024682	SREBF1	FG	2	4.65	-0.06	0.07	0.44	0.44	All-Cis	rs8070432;rs6502629	9.12;9.97	0.05;-1.09
cg11024682	SREBF1	FI	2	4.65	0.02	0.07	0.81	0.84	All-Cis	rs8070432;rs6502629	9.12;9.97	0.02;0.31
cg13708645	KDM2B	FI	3	30.09	0.04	0.03	0.11	0.22	All-Cis	rs28604990;rs11065536;rs3935332	37.01;-9.78;7.28	1.76;1.13;1.06
cg15880704	RBM20	FI	5	23.38	0.06	0.03	0.04	0.91	All-Cis	rs7906643;rs11195272;rs4918591;rs49 18537;rs10509930	-29.45;-7.86;7.14;- 6.72;5.92	-1.83;-1.05;-0.33;- 0.6;0.56
cg17901584	DHCR24	FI	4	8.33	-0.07	0.06	0.19	0.07	All-CisTrans	rs681123;rs735665;rs174546;rs445925	11.86;10.59;- 6.75;5.5	-0.47;-1.33;2;1.69
cg17901584	DHCR24	FI	1	3.53	-0.04	0.08	0.64	NA	Sub- <i>Cis</i>	rs681123	11.86	-0.47
cg18247172	15q26.1	FG	3	5.82	-0.05	0.07	0.47	0.30	All-CisTrans	rs8038275;rs2518968;rs4324798	11.55;-8.12;-5.48	0.27;0.71;1.53
cg18247172	15q26.1	FG	2	5.04	-0.01	0.07	0.85	0.46	Sub- <i>Cis</i>	rs8038275;rs2518968	11.55;-8.12	0.27;0.71
cg18881723	SLAMF1	FG	2	2.85	0.19	0.09	0.05	0.59	All-CisTrans	rs11265461;rs3129055	8.39;6.37	1.89;0.76
cg18881723	SLAMF1	FG	1	1.80	0.23	0.12	0.06	NA	Sub- <i>Cis</i>	rs11265461	8.39	1.89

cg20507228	MAN2A2	FI	1	1.69	-0.07	0.12	0.59	NA	All-Cis	rs1266482	8.12	-0.55
cg24174557	TMEM49	FI	1	1.54	-0.07	0.13	0.60	NA	All-Trans	rs3774937	-7.75	0.53
cg25924746	IRS2	FI	2	5.89	0.05	0.07	0.48	0.85	All-Cis	rs9521528;rs11842277	-13.64;-7.01	-0.55;-0.49
										rs6556405;rs3846687;rs12188300;rs68	39.82;-16.71;-	-0.3;-
cg26403843	RNF145	FI	8	42.56	0.01	0.02	0.60	0.03	All-CisTrans	90049;rs4244439;rs2043269;rs170567	10.3;8.38;-7.39;-	2.66;0.86;0.31;0.0
										47;rs7529925	7.1;-5.55;5.68	9;1.97;-0.28;2.03
										rs6556405;rs3846687;rs12188300;rs68	39.82;-16.71;-	-0.3;-
cg26403843	RNF145	FI	7	41.73	0.01	0.02	0.78	0.07	Sub- <i>Cis</i>	90049;rs4244439;rs2043269;rs170567	10.3;8.38;-7.39;-	2.66;0.86;0.31;0.0
										47	7.1;-5.55	9;1.97;-0.28

Two-sample MR approach was performed to check the effect of known or replicated CpG sites on their significant traits, either fasting glucose or fasting insulin. We also performed MR test in the *cis*-only SNPs if the CpG site has both *cis* and *trans* genetic markers. Locus (CpG): the cytogenetic location or the gene symbol of the CpGs from Illumina annotation. R2 (%): the percentage of explained variance in the exposure by genetic risk score. Effect/SE/P-value: The effect estimate / standard error / P-value of genetic risk score of the exposure on the outcome (MR results). Heterogeneity P-value: The P-value of the heterogeneity test among the SNPs. Type of SNPs: type of SNPs included in the genetic risk score: 1) All-*CisTrans*: all the genetic markers (included *cis* and *trans* SNPs); 2) All-*Cis*: only *cis* genetic markers available; 3) All-*Trans*: only *trans* genetic markers available; 4) Sub-*Cis*: the sub-analysis with the genetic markers in *cis*-only. Z (exposure): the effect estimate per standard error of the SNP on exposure (CpG) from exposure GWAS result; Z (outcome): the effect estimate per standard error of the SNPs on outcome (fasting glucose or insulin) from outcome GWAS result. NP: the genetic risk score has R<sup>2</sup> less than 1%, and the MR was not performed. FG: fasting glucose. FI: fasting insulin. NA: Not available.

Supplementary Table 7 Blood-based expression quantitative trait methylations (eQTMs): association between gene expression and the glycemia related methylation sites

Locus (CpG)	СрG	Probe	Probe-Chr	Probe-Pos	Gene expression	Z	P-value	Cis/Trans
FCRL6	cg00936728	ENSG00000181036	1	159770301	FCRL6	-13.09	4.0 × 10 <sup>-39</sup>	Cis
SLAMF1	cg18881723	ENSG00000026751	1	160709037	SLAMF7	5.84	5.4 × 10 <sup>-9</sup>	Cis
SLAMF1	cg18881723	ENSG00000122223	1	160832692	CD244	4.68	2.9 × 10 <sup>-6</sup>	Cis
SLAMF1	cg18881723	ENSG00000117090	1	160617085	SLAMF1	-4.10	4.1 × 10 <sup>-5</sup>	Cis
CPT1A	cg00574958	ENSG00000110090	11	68611878	CPT1A	-9.22	$3.1 \times 10^{-20}$	Cis
SREBF1	cg11024682	ENSG00000072310	17	17740325	SREBF1	-7.84	$4.5 \times 10^{-15}$	Cis
ABCG1	cg06500161	ENSG00000160179	21	43619799	ABCG1	-12.78	2.2 × 10 <sup>-37</sup>	Cis

The gene expressions associated with the glycemia related methylation sites are shown based on the European blood-based BIOS database (n = 3,841) <sup>26</sup>. Locus (CpG): the cytogenetic location or the gene symbol of the CpGs from Illumina annotation. Probe: The probe of the gene expression. Chr: chromosome. Pos: position. Z: effect estimate per standard error.

Supplementary Table 8 Common genetic determinants of glycemia related DNA methylation (methylation quantitative trait loci, meQTL) and gene expression (expression quantitative trait loci, eQTL) in blood.

			Locus	Туре				Associ	ation with	СрG		Ass	ociation w	ith gene express	ion
Variant	Chr	Position	(eQTL)	(eQTL)	MAF	EA	СрG	Locus (CpG)	z	P-value	Cis/Tran s	Gene expression	z	P-value	Cis/Trans
rs11265282	1	159774408	FCRL6	Protein coding	0.26	С	cg00936728	FCRL6	4.17	3.0 × 10 <sup>-5</sup>	Cis	FCRL6	-6.73	1.7 × 10 <sup>-11</sup>	Cis
rs1577544	1	160630974	SLAMF1 (Nearest)	Protein coding	0.39	Т	cg18881723	SLAMF1	-5.45	5.1 × 10 <sup>-8</sup>	Cis	SLAMF1	-6.40	1.6 × 10 <sup>-10</sup>	Cis
rs6502629	17	17869642	TOM1L2	Protein coding	0.22	G	cg11024682	SREBF1	9.97	2.1 × 10 <sup>-23</sup>	Cis	SREBF1	-17.93	$7.2 \times 10^{-72}$	Cis

The common genetic determinants of glycemia related methylation sites and gene expression in the European blood-based BIOS database (n = 3,841) <sup>26</sup> are shown. Locus (eQTL): the located or nearest protein-coding gene of the eQTL. Type (eQTL): the gene type of the eQTL. Chr: chromosome. MAF: minor allele frequency. EA: effect allele. Z: effect estimate per standard error. Locus (CpG): the cytogenetic location or the gene symbol of the CpGs from Illumina annotation.

Supplementary Table 9 Association between the gene expression level in the glucose metabolism-related tissue and the T2D or related traits based on the Genotype-Tissue Expression (GTEx) project

Gene expression	Trait	Tissue	Z	P-value
FCRL6	T2D <sup>9</sup>	Liver	2.14	0.032
SREBF1	T2D <sup>9</sup>	Whole blood	-2.40	0.016
SREBF1	HbA1c⁴	Whole blood	-3.26	$1.1 \times 10^{-3}$

The significant associations between the gene expression level in the glucose metabolism-related tissue and the T2D or related traits are shown based on the tissue-specific Genotype-Tissue Expression (GTEx) project<sup>28,29</sup>. It was explored in six glucose related tissues, i.e. adipose subcutaneous, adipose visceral omentum, liver, whole blood, pancreas, and muscle skeletal, and five T2D or related traits, i.e. T2D<sup>9</sup>, fasting glucose<sup>5,6</sup>, fasting insulin<sup>6</sup>, HbA1c<sup>57</sup>, and HOMA-IR<sup>4</sup>. Z: effect estimate per standard error.

а

Abbreviation	Full name	Data source and sample size	Tissue
meQTL	Eethylation quantitative trait loci	Summary statistics from BIOS database (n=3,814)	Blood
eQTL	Expression quantitative trait loci	Summary statistics from BIOS database (n=3,814)	Blood
eQTM	Expression quantitative trait methylation	Summary statistics from BIOS database (n=3,814)	Blood
EWAS	Epigenome-wide association study	Current EWAS results (discovery n=4,808, replicated n=11,750)	Blood
MR	Mendelian randomization	Two-sample MR based on BIOS database (n=3,814) and MAGIC (n=96,496)	Blood
GWAS	Genome-wide association study	Summary statistics from MAGIC and DIAGRAM (n=96,496 ~ 452,244) Blood	
GTEx	Genotype-Tissue Expression project	Summary statistic from GTEx and MAGIC/DIAGRAM (tissues n=153 ~ 491) Glucose related	

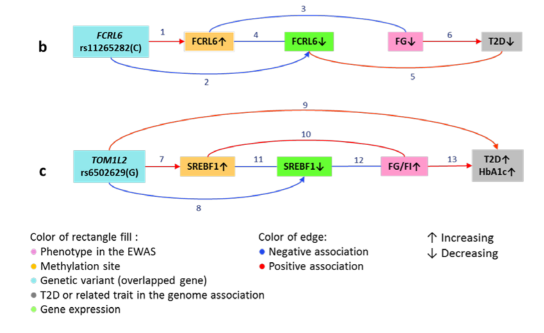
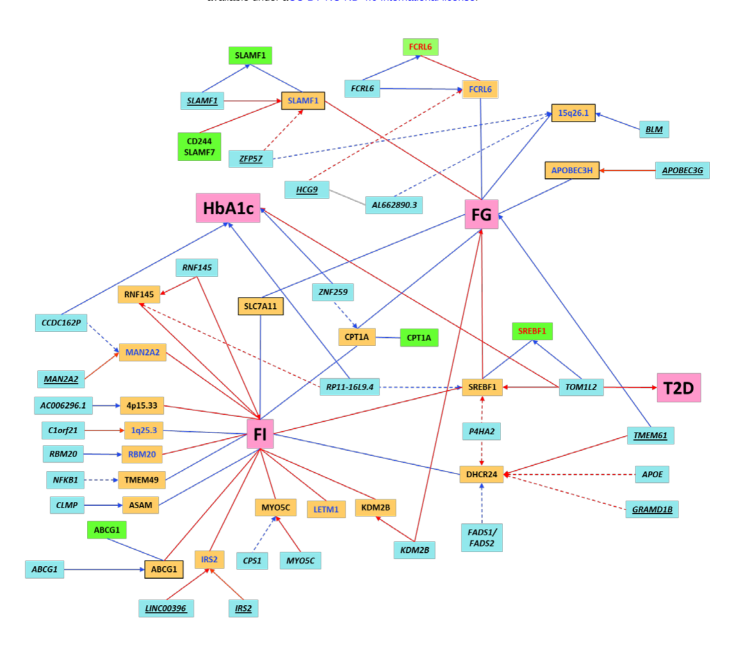
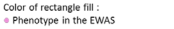


Fig.	NO.	Explanation	Z	P-value
b	1	meQTL	4.17	3.0 × 10 <sup>-5</sup>
	2	eQTL	-6.73	1.7 × 10 <sup>-11</sup>
	3	EWAS	-6.17	9.1 × 10 <sup>-8</sup>
	4	eQTM	-13.09	4.0 × 10 <sup>-39</sup>
	5	GTEx	2.14	0.032
	6	General Knowledge		
С	7	meQTL	9.97	2.1 × 10 <sup>-23</sup>
	8	eQTL	-17.93	7.2 × 10 <sup>-72</sup>
	9	GWAS	3.11	1.5 × 10 <sup>-3</sup>
	10	EWAS with FG	6.45	2.7 × 10 <sup>-8</sup>
	10	EWAS with FI	6.27	6.7 × 10 <sup>-9</sup>
	11	eQTM	-7.84	4.5 × 10 <sup>-15</sup>
	12	GTEx	-3.26	1.1 × 10 <sup>-3</sup>
	13	General Knowledge		

Figure 1 Overview of the cross-omics analysis and examples

Cascading associations cross multiple-omics-based on different data sources were integrated in the network figures. The assumption is genetic variants drive DNA methylation which subsequently regulates gene expression and then glycemic traits. FG: fasting glucose. FI: fasting insulin. T2D: type 2 diabetes





- Methylation site
- Genetic variant (overlapped or <u>nearest</u> gene)
- T2D or related trait in the genome association
- Gene expression

Color of rectangle outline in methylation site:

- BMI-independent methylation site
- No outline: BMI-dependent methylation site

- Color of edge:
- Negative association
- Positive association
- · Association with direction not clear
- Linkage disequilibrium (R<sup>2</sup> > 0.2)
- Color of letter:
- · Novel methylation site and replicated successfully
- Gene expression correlated with T2D or related traits in the glucose metabolism-related tissue

Dash of edge:

-- meQTL or eQTL in trans

-> Casual association

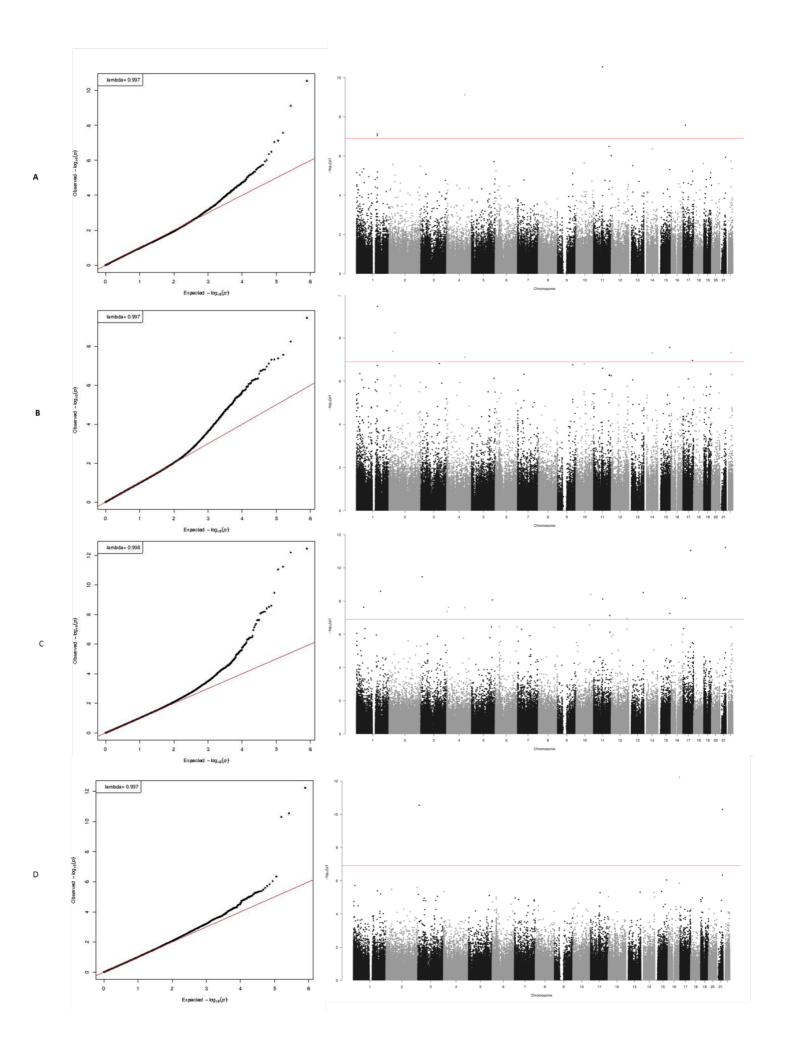
- Others (LD, EWAS)

• Others (gene, phenotype, other methylation site or gene expression)

Figure 2 Associations between genetic variants, DNA methylation sites, gene expressions and fasting glucose,

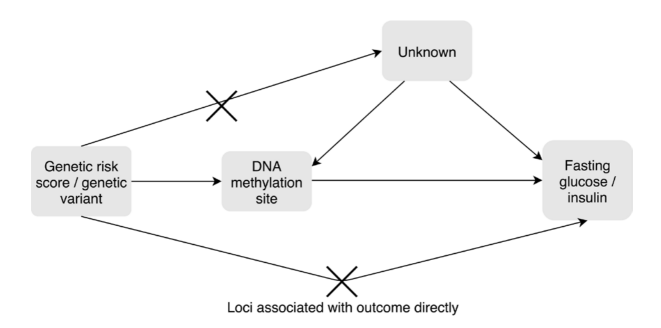
## insulin and related traits based on the integration of cascading associations in Figure 1a

The effect allele is standardized across all associations. Only the significant associations which passed the specific P-value threshold in each association step were shown in the figure. FG: fasting glucose. FI: fasting insulin. T2D: type 2 diabetes.



# Supplementary Figure 1 QQ plots and Manhattan plots of the epigenome-wide association study (EWAS) results

A: EWAS results of fasting glucose in the baseline model; B: EWAS results of fasting glucose in the BMI-adjusted model; C: EWAS results of fasting insulin in the baseline model; D: D EWAS results of fasting insulin in the BMI-adjusted model.



Supplementary Figure 2 Overview of the general Mendelian Randomization process

#### Disclaimer

The views expressed in this manuscript are those of the authors and do not necessarily represent the views of the National Heart, Lung, and Blood Institute; the National Institutes of Health; or the U.S. Department of Health and Human Services.

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