

Multi-ancestry genetic study of type 2 diabetes highlights the power of diverse populations for discovery and translation

SUPPLEMENTARY INFORMATION

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Summary of loci identified through recent ancestry-specific and multi-ancestry meta-analyses incorporating GWAS from the DIAMANTE Consortium. Three recently published meta-analyses of T2D GWAS together account for 77.8% of the total effective sample size contributing to the DIAMANTE multi-ancestry meta-regression (**Supplementary Figure 1**). First, the European ancestry-specific DIAMANTE study¹, which includes 74,124 T2D cases and 824,006 controls, accounting for 47.0% of the effective sample size of the multi-ancestry meta-regression. The multi-ancestry meta-regression includes an additional 6,030 T2D cases and 29,810 controls from EGCUT, NEO and MGB, which were not part of the European ancestry-specific DIAMANTE study. Second, the East Asian ancestry-specific DIAMANTE study², which includes 77,418 T2D cases and 356,122 controls, accounting for 28.4% of the effective sample size of the trans-ethnic meta-regression. The multi-ancestry meta-regression does not include 21,151 T2D cases and 128,967 controls from BBJ that were part of the East Asian ancestry-specific DIAMANTE study. Third, the meta-analysis of GWAS from the Million Veteran Program (MVP), DIAMANTE and other cohorts³, which includes 228,499 T2D cases and 1,178,783 controls, accounting for 57.1% of the effective sample size of the multi-ancestry meta-regression. GWAS contributing to the multi-ancestry meta-regression account for 45.4% of the effective sample size of the MVP meta-analysis. Both multi-ancestry and ancestry-specific (European, African, Hispanic/Latino and East/South Asian) meta-analyses were undertaken. Making comparisons of the numbers of (novel) loci reported by each of these investigations is not an appropriate evaluation of their relative power because of differences in thresholds of genome-wide significance, corrections for residual population structure after meta-analysis, reference panels used for imputation and SNV filtering criteria (**Supplementary Note Table 1**). Instead, we sought to present an all-inclusive summary of loci reported in studies to which DIAMANTE GWAS have contributed to provide the most comprehensive overview of the genetic contribution to T2D susceptibility to date.

We began by considering loci reported in each of the published ancestry-specific and multi-ancestry meta-analyses incorporating GWAS from the DIAMANTE Consortium (without adjustment for BMI). For each of these efforts, loci were defined as mapping 500kb up- and downstream of a lead SNV attaining genome-wide significance ($p < 5 \times 10^{-8}$). We also considered loci reported in the multi-ancestry meta-regression, which used a more conservative definition (**Methods**) that: (i) considered the flanking genomic interval mapping 500kb up- and downstream of a lead SNV attaining stringent multi-ancestry genome-wide significance ($p < 5 \times 10^{-9}$); and (ii) merged loci where lead SNVs were separated by less than 1Mb. We then aggregated loci across the four studies, ensuring no overlap between adjacent loci (**Supplementary Figure 2**). Taken together, the four studies report 520 non-overlapping loci spanning 624.5Mb, including 405 (77.9%) attaining stringent multi-ancestry genome-wide significance (**Supplementary Note Table 2**). Of the 520 loci, 35 (6.7%) were reported only in ancestry-specific meta-analyses: 21 European ancestry-specific, 12 East Asian ancestry-specific and 2 African ancestry-specific.

Comparison of ancestry-specific meta-analyses and multi-ancestry meta-regression. To gain insight into the power offered by aggregating GWAS from diverse populations, we compared the number of loci identified in multi-ancestry meta-regression with that detected in ancestry-specific meta-analyses. As there are minor differences in the GWAS

contributing to the multi-ancestry meta-regression and the previously reported European and East Asian ancestry components of DIAMANTE^{1,2}, we restricted comparisons to those that contributed to both. Of the 100 and 193 loci attaining genome-wide significance in the East Asian and European ancestry-specific meta-analyses, respectively, lead SNVs at 94 (94.0%) and 164 (85.0%) demonstrated stronger evidence for association (i.e. smaller p -value) in the multi-ancestry meta-regression (**Extended Data Figure 4**), in line with differences in sample size. In contrast, eleven (5.7%) of the 193 loci identified in the European ancestry-specific meta-analysis did not attain genome-wide significance in the multi-ancestry meta-regression (**Supplementary Note Table 3**). None of these eleven SNVs demonstrated significant evidence of T2D association in a meta-analysis of non-European ancestry GWAS. Such signals could arise when the lead SNV is in strong linkage disequilibrium (LD) with an ancestry-specific causal variant that has not been interrogated in the multi-ancestry meta-regression, or because of haplotype/epistatic effects across variants with differing allele frequency between ancestry groups. Taken together, these results demonstrate the power of multi-ancestry meta-analyses for locus discovery and replication that is afforded by increased sample size, but also emphasize the importance of complementary ancestry-specific GWAS for optimal identification of associations driven by causal variants that are not shared across diverse populations.

Dissection of distinct T2D association signals. Through approximate conditional analyses, conducted using ancestry-matched LD reference panels for each GWAS, we partitioned associations at the 237 T2D loci into 338 distinct signals that were each represented by an index SNV at the same multi-ancestry genome-wide significance threshold (**Methods, Supplementary Tables 6 and 7**). We observed multiple distinct association signals at 52 (21.9%) loci, of which 50 were represented by between two and five index SNVs. The most complex genetic architecture was observed across a 1Mb region flanking the lead SNV at the *TCF7L2* locus, where the T2D association was delineated to 16 distinct signals (**Supplementary Figure 3**), and a 1.7Mb imprinted region encompassing the previously reported loci *INS-IGF2* and *KCNQ1*, which was delineated into 14 distinct signals (**Supplementary Figure 4**).

Assessment of the impact of reference panel choice on approximate conditional analyses undertaken in admixed ancestry groups. We used haplotypes from the 1000 Genomes Project reference panel (phase 3, October 2014 release)⁴ that were specific to each ancestry group (**Supplementary Table 22**) as a reference for LD between SNVs across loci in the approximate conditional analysis. African ancestry GWAS, including admixed African American studies, were matched to African haplotypes, derived from 661 individuals from: African Caribbean in Barbados; African Ancestry in Southwest USA; Esan in Nigeria; Gambian in Western Division, The Gambia; Luhya in Webuye, Kenya; Mende in Sierra Leone; and Yoruba in Ibadan, Nigeria. Hispanic GWAS were matched to American haplotypes, derived from 347 individuals from: Colombian in Medellin, Colombia; Mexican Ancestry in Los Angeles, California; Peruvian in Lima, Peru; and Puerto Rican in Puerto Rico. The 1000 Genomes Project reference panel has the advantage that it includes individuals from diverse populations across each ancestry group, with haplotypes derived from high-quality whole genome sequence data that includes all variants tested in the multi-ancestry meta-analysis. However, the disadvantage of this reference panel is that it includes only of the order of 500

individuals per ancestry group, and approximate conditional analyses may therefore be susceptible to unstable effect size estimates at lower frequency variants.

An alternative approach is to make use of individual-level genotype data from GWAS contributing to the multi-ancestry meta-regression as a reference for LD in approximate conditional analyses. These studies typically include larger numbers of individuals than are available in the 1000 Genomes Project reference panel. However, these GWAS will usually have been imputed, such that many variants present in the reference panel will fail imputation quality control. Furthermore, the approximate conditional analyses implemented in GCTA require that imputed genotypes be converted to hard calls, which can lead to over-confidence in downstream analyses by ignoring imputation uncertainty. A single study may also not be representative of the genetic diversity amongst GWAS from an ancestry group, particularly those with variable levels of admixture.

To gain insight into the robustness of our approximate conditional analyses to the choice of LD reference panel in admixed ancestry groups, we considered subsets of 1,000 African American and 1,000 Hispanic individuals from the Resource for Genetic Epidemiology on Adult Health and Aging (GERA), a large multi-ancestry population-based cohort, created for investigating the genetic and environmental basis of age-related diseases [database of Genotypes and Phenotypes (dbGaP) phs000674.p1]. GERA participants have previously been genotyped using one of four custom arrays, which have been designed to maximise coverage of common and low-frequency variants in non-Hispanic white, East Asian, African American and Hispanic individuals^{5,6}. We undertook quality control of these genotype data, removing individuals from known pedigrees and/or with call rate (<97%), and excluding SNVs with call rate (<95%) and extreme deviation from Hardy-Weinberg equilibrium (autosomes only, exact $p < 10^{-6}$). We constructed a genetic relationship matrix (GRM) from pair-wise identity by descent metrics estimated from LD pruned ($r^2 < 0.01$ across individuals) autosomal SNVs shared across the four genotyping arrays, and with MAF $\geq 1\%$, after exclusion of those in high-LD and complex regions, and those mapping to established T2D loci. We defined related individuals with pair-wise π -hat > 0.2 and removed those with the lowest call rate from each related set.

We applied multi-dimensional scaling, implemented in PLINK⁷, to the GRM to obtain principal components to represent axes of genetic variation that separate the major ancestry groups. Clusters of African American and Hispanic individuals were identified in principal component space, and subsets of 1,000 randomly selected individuals from each cluster for use as a reference for LD in approximate conditional analyses.

For each subset of individuals, we constructed a scaffold for imputation after excluding SNVs with MAF $< 1\%$. The scaffold was then pre-phased using SHAPEITv2.5⁸, based on estimates of recombination rate from the International HapMap Project⁹. The resulting haplotypes were imputed up to the 1000 Genomes Project reference panel (phase 3, October 2014 release) using minimac4 via the Michigan Imputation Server¹⁰. SNVs with $r^2 \geq 0.4$ were retained for downstream analyses. African and Hispanic LD reference panels were then obtained by converting imputed genotype dosages to hard calls using PLINKv1.9¹¹.

For each locus with more than one distinct association signal in the multi-ancestry meta-regression, we used GCTA in each African and Hispanic ancestry GWAS, removing each SNV, in turn, from the conditional set, and adjusting for the remainder, using the "--cojo-cond" option. If any SNV from the conditional set failed imputation quality control in the LD reference panel, the locus was excluded from downstream analyses. We aggregated allelic log-ORs from the approximate conditional analyses across African and Hispanic ancestry

GWAS via fixed-effects meta-analysis using METAL¹² based on inverse-variance weighting. We corrected association p -values and standard errors of allelic effects from each ancestry group for residual inflation due to structure between GWAS using the same genomic control adjustments as in the unconditional analyses (**Methods**).

We compared allelic effect estimates (log-OR) and p -values (on a $-\log_{10}$ scale) in African and Hispanic ancestry-specific meta-analyses using ancestry-matched LD reference panels from the 1000 Genomes Project and GERA (**Supplementary Figure 5, Supplementary Note Tables 4 and 5**). There was a strong correlation in allelic effect estimates from the different LD reference panels: African ancestry $r=0.997$; Hispanic ancestry $r=0.988$. The strength of evidence in favour of association (as measured by the conditional p -value) of each index SNV was mostly within one order of magnitude between the LD reference panels. We conclude, therefore, that our approximate conditional analyses undertaken in admixed ancestry groups are robust to the choice of reference panel.

Impact of obesity on multi-ancestry heterogeneity. We were interested to determine whether ancestry-correlated heterogeneous association signals could be explained by an interaction with obesity, given the leftwards shift in the distribution of body mass index (BMI) in individuals of East Asian ancestry. To do this, we considered the 136 association signals with nominal evidence of ancestry-correlated heterogeneity in allelic effects (**Supplementary Note Table 6**). For each index SNV, we modelled allelic log-ORs across GWAS in a linear regression framework, weighted by the inverse of the variance of the effect estimates. For index SNVs at loci with a single distinct association signal, log-ORs and variances were obtained from unconditional analysis. For index SNVs at loci with multiple distinct association signals, log-ORs and variances were obtained from approximate conditional analysis. We excluded GWAS for which BMI was not reported: BIOME (HIS), GERA (AFR), GERA (EUR), GODARTS, KORA and WTCCC. For each GWAS, we included as covariates: (i) mean BMI; and (ii) the three axes of genetic variation representing ancestry. In this modelling framework, we tested for: (i) heterogeneity in allelic effects on T2D between GWAS that is correlated with BMI, after adjusting for ancestry; (ii) heterogeneity in allelic effects on T2D between GWAS that is correlated with ancestry, after adjusting for BMI; and (iii) residual allelic effect heterogeneity between GWAS due to unmeasured confounders.

The strongest evidence for heterogeneity in allelic effects that was correlated with BMI (after accounting for ancestry) was observed for the T2D association signal at the *CDKAL1* locus (rs9348441, $p_{\text{HET}}=3.0 \times 10^{-6}$). At this signal, the effect of the risk allele on T2D was greatest in East Asian ancestry populations, and there was a negative correlation between BMI and log-OR across GWAS (**Supplementary Figure 6**). This relationship is consistent with a model of “favourable adiposity”, whereby a subset of BMI-increasing alleles are associated with higher subcutaneous-to-visceral adipose tissue ratio and a paradoxical reduction in insulin levels, protecting against T2D through higher adipose storage capacity¹³. A protective interaction with obesity at this locus is supported by evidence of: (i) stronger association at the index SNV after adjustment for BMI in the European and East Asian ancestry components of DIAMANTE^{1,2}; and (ii) significant association of the T2D-risk allele with decreased BMI in European and East Asian ancestry GWAS meta-analyses of obesity in the general population^{14,15}. However, confirmation of the impact of obesity on the heterogeneity of allelic effects at the *CDKAL1* locus requires formal testing of SNV x BMI interaction within GWAS across ancestry groups.

Impact of allele frequency, allelic effect size and LD on fine-mapping resolution. Compared to the European ancestry-specific meta-analysis, some of the most dramatic improvements in fine-mapping resolution after multi-ancestry meta-regression included signals where the index SNV was of lower frequency and/or of smaller effect in European ancestry populations. For these signals, including those at *GCC1-PAX4-LEP*, *SGCG*, *RGMA*, *DSTYK-MDM4* and *MYO3A*, the evidence for association was weak in the European ancestry-specific meta-analysis, resulting in large credible sets compared to other ancestry groups. However, we also observed examples of T2D signals with strong associations across all five ancestry groups, for which the credible sets were smaller in the multi-ancestry meta-regression. The most noticeable improvements in fine-mapping resolution were seen at *TMEM154*, *HMGA2*, *GRP-MC4R*, *IGF2BP2*, *SPRY2* and *FTO* (**Supplementary Table 9**). At *FTO*, for example, the 18 variants in the European ancestry-specific 99% credible set were in strong LD with the index SNV (rs55872725) in European ancestry populations ($r^2 > 0.8$). However, the 99% credible set after multi-ancestry meta-regression included just six of these variants that were in strong LD with the index SNV in all five ancestry groups (**Supplementary Figure 7**).

Improved fine-mapping of T2D coding variant associations. The multi-ancestry meta-regression highlighted two examples of previously reported T2D coding variant associations that were better resolved by fine-mapping across diverse populations (**Supplementary Figure 8**). A set of five coding variants in *SLC16A11* has been associated with T2D in Hispanic populations^{16,17}, but causality could not be ascribed because of strong LD between them. However, after multi-ancestry fine-mapping, *SLC16A11* p.Val113Ile (rs117767867, $p = 6.5 \times 10^{-24}$, $\pi = 59.8\%$) emerged as the variant most likely driving this association signal (the other four coding variants together account for just 14.0% of the posterior probability). Similarly, strong LD at the *KCNJ11-ABCC8* locus has frustrated efforts in European ancestry studies to distinguish the impact on T2D of three missense variants: *KCNJ11* p.Val250Ile (rs5215), *KCNJ11* p.Lys23Glu (rs5219) and *ABCC8* p.Ala1369Ser (rs757110). *ABCC8* and *KCNJ11* code for the two elements of the hetero-octameric beta-cell K_{ATP} channel and both represent strong biological candidates. Whilst multi-ancestry fine-mapping cannot equivocally distinguish between *KCNJ11* p.Val250Ile ($p = 1.3 \times 10^{-54}$, $\pi = 67.1\%$) and *KCNJ11* p.Lys23Glu ($p = 2.6 \times 10^{-54}$, $\pi = 32.5\%$), it is less likely that the association signal is mediated via *ABCC8* p.Ala1369Ser ($p = 1.2 \times 10^{-51}$, $\pi = 0.1\%$).

Multi-ancestry fine-mapping provided a more detailed view of the role of missense variants in driving three distinct T2D association signals at the *ZFAND3-KCNK16-GLP1R* locus (**Supplementary Figure 9**). Previous East Asian ancestry GWAS and exome-array meta-analyses^{18,19} reported T2D association with *GLP1R* p.Arg131Gln (rs3765467). Whilst this variant is included in the 99% credible set of the signal indexed by rs742762, a non-coding SNV, in the multi-ancestry meta-regression, it has a relatively low posterior probability of association ($\pi = 2.0\%$, compared with $\pi = 75.0\%$ for the index SNV). However, we identified a different *GLP1R* missense variant, p.Pro7Leu (rs10305420, $p = 1.1 \times 10^{-9}$, $\pi = 94.1\%$), not in LD with p.Arg131Gln, which seems likely to be causal for the second association signal at the locus. At the third signal, 61.4% of the posterior probability of association could be attributed to three different missense variants: p.Ser21Gly (rs10947804, $\pi = 39.2\%$) in *KCNK17*; and p.Pro254His (rs11756091, $\pi = 14.8\%$) and p.Ala277Glu (rs1535500, $\pi = 13.7\%$) in *KCNK16*. Both genes encode members of the TWIK-related alkaline pH-activated K₂P family, TALK-1 and TALK-2, and are expressed in islets with high specificity. The missense variants

are in strong LD with each other across ancestry groups, and the T2D-risk haplotype is associated with increased *KCNK17* expression in pancreatic islets²⁰. These results highlight the mechanistic complexity at this locus, with evidence that missense variants in *GLP1R*, *KCNK16* and *KCNK17* may each be contributing to T2D susceptibility.

Integration of fine-mapping and chromatin interaction data in diverse tissues. We intersected 99% credible set variants for distinct T2D association signals with genome-wide promoter-focussed chromatin conformation capture data (pcHi-C) from pancreatic islets, subcutaneous adipose and liver (equivalent data are not available in hypothalamus and visceral adipose)²¹⁻²³. Across the three tissues, we observed contacts between credible set variants and putative target gene promoters for 214 (63.3%) of the association signals (**Supplementary Table 18**). The contacts at 119 of these signals were observed in only one tissue: 51 in islets, 45 in liver, and 23 in subcutaneous adipose. Some targets were expected based on their proximity to the index SNV for the T2D association (including *TCF7L2*, *PROX1*, *PTEN*, *DLEU1*, *GLIS3*, *CCND2*, *CMIP* and *BCL2*), but for 143 (66.8%) of the 214 signals, we identified more distant candidate effector genes (including *AQP5* and *AQP6* at the *FAIM2* locus, *P2RX1* at the *ZZEF1* locus, *STX16* at the *GNAS* locus, and *ISL1* at the *ITGA1* locus). Several of these targets provide complementary support for candidate effector genes identified via colocalization with *cis*-eQTLs in the same tissue: *PLEKHA1* in islets and subcutaneous adipose; *ST6GAL1*, *CARD9*, *DNLZ*, *CAMK1D*, *TCF7L2*, *TH*, *DLK1* and *AP3S2* in islets; *DCAF16*, *STEAP2* and *MAN2C1* in subcutaneous adipose; and *CEP68* and *SLC22A3* in liver.

Summary of candidate causal genes at T2D loci identified from functional annotation and colocalization with molecular QTLs. We identified a total of 117 candidate causal genes at T2D loci through integration of multi-ancestry fine-mapping, functional annotation, and molecular QTL data resources (**Supplementary Note Table 7**). First, we identified missense variants accounting for more than 50% posterior probability of driving distinct T2D association signals after annotation-informed fine-mapping. Second, we identified distinct T2D association signals that colocalized with more than 80% posterior probability with: (i) circulating plasma proteins (pQTLs)²⁴; or (ii) gene expression (eQTLs) in diabetes-relevant tissues (pancreatic islets, subcutaneous and visceral adipose, liver, skeletal muscle, and hypothalamus)^{25,26}.

We sought to evaluate the support for these candidate causal genes from complementary analyses undertaken in three recently published meta-analyses of T2D GWAS together account for 77.8% of the total effective sample size contributing to the DIAMANTE multi-ancestry meta-regression. First, the European ancestry-specific DIAMANTE study¹ reported missense variants accounting for more than 50% posterior probability of driving distinct T2D association signals after annotation-informed fine-mapping. Second, the East Asian ancestry-specific DIAMANTE study² reported index SNVs for T2D that are in strong LD ($r^2 > 0.8$) with significant eQTLs in: (i) pancreas²⁵ and pancreatic islets²⁰; (ii) subcutaneous adipose^{25,27}; (iii) skeletal muscle²⁵; and (iv) blood^{25,28}. Here, we focussed only on those significant eQTLs in diabetes-relevant tissues (pancreas, pancreatic islets, subcutaneous adipose and skeletal muscle). Third, the meta-analysis of GWAS from the Million Veteran Program (MVP), DIAMANTE and other cohorts³ reported missense variants attaining genome-wide significance at T2D loci. Here, we considered only those missense variants in strong LD ($r^2 > 0.8$) with lead SNVs for T2D. The study also reported the results of

transcriptome-wide association studies across the diverse range of tissues available in GTEx²⁵. Here, we focussed on those genes with significant association ($p < 1.93 \times 10^{-7}$) of T2D with genetically-regulated expression, which also colocalized with more than 80% posterior probability in diabetes-relevant tissues (pancreas, subcutaneous and visceral adipose, liver, skeletal muscle, and hypothalamus).

Of the 117 candidate causal genes identified in the DIAMANTE multi-ancestry study, 40 were not reported in these complementary analyses (**Supplementary Note Table 7**). These include genes previously implicated in T2D through detailed experimental studies (such as *TCF7L2*, *MTNR1B*, *ARAP1*, *STARD10* and *CAMK1D*), but also novel candidates that provide new leads for functional follow-up. These findings highlight the importance of: (i) diverse populations to enable high-resolution fine-mapping of T2D association signals; (ii) gene expression profiling in diabetes-relevant tissues to understand cell-type specific contexts through which association signals are mediated; and (iii) dissection of both T2D associations and molecular QTLs through conditional analysis to allow colocalization at the level of a signal (and not a locus).

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Supplementary Methods

Exemplar power calculations. Assuming homogeneous effects on T2D across ancestry groups, we estimated power to detect association with an SNV under an additive model from the non-centrality parameter of a chi-squared distribution with one degree of freedom, given by $N_{\text{eff}}\psi^2q(1-q)$, where q denotes the mean effect allele frequency across populations and ψ denotes the allelic log-OR, and N_{eff} is the total effective sample size across studies¹. At our multi-ancestry genome-wide significance threshold, $p < 5 \times 10^{-9}$, under an additive genetic model, we had $\geq 80\%$ power to detect association of SNVs with MAF $\geq 5\%$ and OR ≥ 1.045 or MAF $\geq 0.5\%$ and OR ≥ 1.145 .

Dissection of distinct multi-ancestry association signals. We used iterative approximate conditioning, implemented in GCTA², making use of forward selection and backward elimination, to identify index SNVs at multi-ancestry genome-wide significance ($p < 5 \times 10^{-9}$). We used haplotypes from the 1000 Genomes Project reference panel (phase 3, October 2014 release)³ that were specific to each ancestry group (**Supplementary Table 22**) as a reference for LD between SNVs across loci in the approximate conditional analysis.

For each locus, we first used GCTA in each ancestry-specific GWAS, using the study-level association summary statistics and ancestry-matched LD reference, implementing a forward selection scheme. At each iteration, we adjusted for the “conditional set” of variants at the locus using the “--cojo-cond” option. In the first iteration, the conditional set included only the lead SNV at the locus. Allelic log-ORs from the approximate conditional analyses across GWAS were modelled in the multi-ancestry meta-regression framework, incorporating the three axes of genetic variation as covariates, and weighted by the inverse of the variance of the effect estimates. The meta-regression association p -values were corrected for inflation due to residual structure between GWAS by using the same genomic control adjustment as in the unconditional analysis ($\lambda_{TA} = 1.052$). If no SNVs attained genome-wide significant ($p < 5 \times 10^{-9}$) evidence of residual T2D association in the meta-regression, the iterative approximate conditional analysis for the locus was stopped. Otherwise, the SNV with the strongest residual association signal was added to the conditional set. This process continued, at each iteration adding the SNV with the strongest residual T2D association from the meta-regression to the conditional set, until no remaining SNVs attained genome-wide significance. Note, that at each iteration, GWAS with missing association summary statistics for any variant in the conditional set were excluded from the meta-regression.

For each locus with more than one SNV in the conditional set, we then checked that all variants in the conditional set attained genome-wide significant evidence of association in a joint model after meta-regression. To do this, we used GCTA in each GWAS, but this time using the “--cojo-joint” option, including all SNVs in the conditional set in the joint model. Allelic log-ORs from the approximate conditional analyses across GWAS were modelled in the multi-ancestry meta-regression framework, incorporating the three axes of genetic variation as covariates, and weighted by the inverse of the variance of the effect estimates. The meta-regression association p -values were corrected for inflation due to residual structure between GWAS by using the same genomic control adjustment as in the unconditional analysis ($\lambda_{TA} = 1.052$). If any SNV in the conditional set did not attain genome-wide significant evidence of association, the SNV with the least significant p -value was removed (backward elimination). The procedure then iterated between forward selection

and backward elimination steps until: (i) no SNVs outside the conditional set attained genome-wide significant evidence of residual association in the meta-regression; and (ii) all SNVs in the conditional set attained genome-wide significant evidence of association in the joint model after meta-regression.

For each locus including more than one SNV in the conditional set, we next dissected each distinct association signal. We again used GCTA in each GWAS, but this time removing each SNV, in turn, from the conditional set, and adjusting for the remainder, using the “--cojo-cond” option. Allelic log-ORs from the approximate conditional analyses across GWAS were modelled in the multi-ancestry meta-regression framework, incorporating the three axes of genetic variation as covariates, and weighted by the inverse of the variance of the effect estimates. The meta-regression association p -values were corrected for inflation due to residual structure between GWAS by using the same genomic control adjustment as in the unconditional analysis ($\lambda_{TA} = 1.052$). The variant with the strongest residual association was defined as the “index SNV” for the signal. We also aggregated allelic log-ORs from the approximate conditional analyses across GWAS via fixed-effects meta-analysis using METAL⁴ based on inverse-variance weighting. Standard errors were corrected for residual inflation due to structure between GWAS using the same genomic control adjustment as in the unconditional analysis ($\lambda_{TA}^{FE} = 1.253$).

Ancestry-specific meta-analyses. We aggregated association summary statistics across GWAS from the same ancestry group via fixed-effects meta-analysis using METAL⁴ based on inverse-variance weighting of allelic log-OR to obtain effect size estimates. We corrected association p -values and standard errors of allelic effects from each ancestry group for residual inflation due to structure between GWAS by genomic control adjustment: African $\lambda_{AFR} = 1.056$; East Asian $\lambda_{EAS} = 1.111$; European $\lambda_{EUR} = 1.096$; Hispanic $\lambda_{HIS} = 1.008$; South Asian $\lambda_{SAS} = 0.973$ (no correction made). We estimated the mean effect allele frequency across GWAS from each ancestry group, weighted by the effective sample size of the study. We also aggregated association summary statistics across GWAS from non-European ancestry groups via fixed-effects meta-analysis using METAL based on inverse-variance weighting of allelic log-OR to obtain effect size estimates. We corrected association p -values and standard errors of allelic effects for residual inflation due to structure between GWAS by genomic control adjustment: $\lambda_{nonEUR} = 1.133$. Finally, we aggregated association summary statistics across GWAS from East Asian and European ancestry groups via fixed-effects meta-analysis using METAL based on inverse-variance weighting of allelic log-OR to obtain effect size estimates. We corrected association p -values and standard errors of allelic effects for residual inflation due to structure between GWAS by genomic control adjustment: $\lambda_{EASEUR} = 1.147$.

For each locus with more than one distinct association signal in the multi-ancestry meta-regression, we used GCTA in each GWAS, removing each SNV, in turn, from the conditional set, and adjusting for the remainder, using the “--cojo-cond” option. We aggregated allelic log-ORs from the approximate conditional analyses across GWAS within the same ancestry group (and combined across East Asian and European ancestry groups) via fixed-effects meta-analysis using METAL based on inverse-variance weighting. We corrected association p -values and standard errors of allelic effects from each ancestry group for residual inflation due to structure between GWAS using the same genomic control adjustments as in the unconditional analyses.

Derivation of approximate Bayes' factors in favour of association. For the multi-ancestry meta-regression, we approximated the Bayes' factor for the j th SNV at the i th distinct association signal by

$$\Lambda_{ij} = \exp\left[\frac{D_{ij} - 4\ln K_{ij}}{2}\right],$$

where D_{ij} is the deviance across K_{ij} contributing GWAS⁵.

For the European ancestry-specific meta-analysis and combined East Asian and European ancestry meta-analysis, we approximated the Bayes' factor for the j th SNV at the i th distinct association signal by

$$\Lambda_{ij} = \exp\left[\frac{D_{ij} - \ln K_{ij}}{2}\right],$$

where $D_{ij} = b_{ij}^2/v_{ij}$, and b_{ij} and v_{ij} are the allelic log-OR and corresponding variance, respectively, across K_{ij} contributing GWAS.

Assessment of fine-mapping resolution in “down-sampled” multi-ancestry meta-regression. We selected GWAS contributing to the multi-ancestry meta-regression to approximate the effective sample size of the European ancestry-specific meta-analysis and maintain the distribution of effective sample size across ancestry groups (**Supplementary Table 10**). The selected GWAS were: African ancestry BIOME (AFR 2), CARDIA, CHS, EMERGE, GENOA (AFR), GERA (AFR), REGARDS, WHI (AFR); East Asian ancestry BBJ (1), CAGE-KING (2), CKB-12, CKB-16, CKB-58, CKB-68, CKB-78, CKB-88, KBA (2), SCES (2), SCHS; European ancestry BIOME (EUR), DECODE, DGI, EPIC-INTERACT (2), FHS, FUSION, GCKD, GENOA (EUR), GERA (EUR), GODARTS, GOMAP, KORA, METSIM, MGI, NUGENE, PIVUS, PROSPER, RS (1), UCPH, ULSAM; Hispanic ancestry BIOME (HIS), HCHS/SOL, MACAD, MC (1), MC (2), MESA (HIS); South Asian EPIDREAM, GRCCDS, INDICO, INTERHEART (2), LOLIPOP (1), LOLIPOP (2), LOLIPOP (4), PROMIS (1).

We conducted a “down-sampled” multi-ancestry meta-regression, implemented in the MR-MEGA software⁵, for the selected studies. For each SNV, we modelled allelic log-ORs across GWAS in a linear regression framework, weighted by the inverse of the variance of the effect estimates, incorporating the same three axes of genetic variation as covariates (**Extended Data Figure 2**). We corrected the meta-regression association p -values for inflation due to residual structure between the selected GWAS using genomic control adjustment (allowing for four degrees of freedom): $\lambda_{TA*} = 1.012$.

For each locus with more than one distinct association signal in the complete multi-ancestry meta-regression, we used GCTA in each GWAS, removing each SNV, in turn, from the conditional set, and adjusting for the remainder, using the “--cojo-cond” option (as described above). Allelic log-ORs from the approximate conditional analyses across GWAS were modelled in the multi-ancestry meta-regression framework, incorporating the three axes of genetic variation as covariates, and weighted by the inverse of the variance of the effect estimates. The meta-regression association p -values were corrected for inflation due to residual structure between GWAS by using the same genomic control adjustment as in the unconditional analysis ($\lambda_{TA*} = 1.012$).

Within each locus, we approximated the Bayes' factor⁶, Λ_{ij} , in favour of T2D association of the j th SNV at the i th distinct association signal on the basis of summary statistics from the down-sampled multi-ancestry meta-regression. For loci with a single association signal, the association summary statistics were obtained from unconditional analysis. For loci with multiple distinct association signals, the association summary statistics were obtained from the approximate conditional analyses. The posterior probability for the j th SNV at the i th distinct signal, was then given by $\pi_{ij} \propto \Lambda_{ij}$, where

$$\Lambda_{ij} = \exp \left[\frac{D_{ij} - 4 \ln K_{ij}}{2} \right],$$

and D_{ij} is the deviance across K_{ij} contributing GWAS⁵. We derived a 99% credible set⁷ for the i th distinct association signal by: (i) ranking all SNVs according to their posterior probability π_{ij} ; and (ii) including ranked SNVs until their cumulative posterior probability attains or exceeds 0.99.

Enrichment of distinct T2D association signals in genomic annotations. We tested for enrichment of distinct T2D association signals from the multi-ancestry meta-regression (as measured by the approximate Bayes' factor) that map to genomic annotations using fGWAS⁸ with the region-based input format (-fine). We first considered each annotation separately and identified those with significant enrichment ($p < 0.00023$, Bonferroni correction for 220 annotations), which we refer to as the "enriched set". We then used an iterative approach to identify a joint model of enriched annotations. At each iteration, we added the annotation from the enriched set to the joint model that maximised the improvement in the penalised likelihood. We continued until no additional annotations improved the fit of the joint model at nominal significance ($p < 0.05$). We next used the cross-validation likelihood because the significance of parameter estimates from the penalised likelihood cannot be assessed using standard statistical approaches. For the selected joint model, we identified the penalty that maximised the cross-validation likelihood. Finally, we dropped any annotations from the joint model that resulted in a decrease in the cross-validation likelihood.

Transferability of multi-ancestry GRS across ancestry groups. We selected two studies per ancestry group as test GWAS, prioritising those with larger effective sample sizes and greater genetic diversity: DDS/DCC, WHI (AFR), KBA, SIMES, EPIC-INTERACT (2), UKBB, HCHS/SOL, MC, PROMIS and RHS. We repeated the multi-ancestry meta-regression, after excluding the ten test GWAS, incorporating the same three axes of genetic variation as covariates to account for ancestry. The association p -values from this "reduced" meta-regression were then corrected for inflation due to residual structure between GWAS by means of genomic control adjustment (allowing for four degrees of freedom): $\lambda_{TA} = 1.037$. SNVs reported in $\geq 50\%$ of the total effective sample size of the "reduced" meta-regression ($N_{TE} \geq 179,074$) were included in downstream analyses. We identified loci attaining genome-wide significant evidence of association ($p < 5 \times 10^{-9}$) in the "reduced" meta-regression, and the lead SNV for each locus was selected as the variant with minimum association p -value.

For each test GWAS, we next estimated population-specific "predicted" allelic effects for each lead SNV to be used as weights in the GRS. For the i th study, we estimated the allelic effect of the j th SNV by

$$\hat{b}_{TAij} = \alpha_{TA0j} + \sum_k \alpha_{TAkj} x_{ki},$$

where x_{ki} is the position of the i th study on the k th axis of genetic variation from the “complete” multi-ancestry meta-regression, and α_{TA0j} and α_{TAkj} denote the intercept and effect of the k th axis of genetic variation for the SNV from the “reduced” multi-ancestry meta-regression. For each test GWAS, we then regressed the observed allelic effect estimates at lead SNVs, weighted by their corresponding variances, on the “predicted” allelic effect estimates, as implemented in `grs.summary` function⁹ of the `gtx` package in R. We estimated the OR per unit of the weighted GRS and the corresponding percentage of T2D variance explained, measured by pseudo R^2 .

Transferability of ancestry-specific GRS across ancestry groups. We selected two studies per ancestry group as test GWAS. We repeated each of the ancestry-specific fixed-effects meta-analyses after excluding the ten test GWAS. We aggregated association summary statistics across GWAS from the same ancestry group via fixed-effects meta-analysis using METAL⁴ based on inverse-variance weighting of allelic log-OR to obtain effect size estimates. We corrected association p -values for residual inflation due to structure between GWAS by genomic control adjustment: African $\lambda_{AFR} = 1.049$; East Asian $\lambda_{EAS} = 1.092$; European $\lambda_{EUR} = 1.180$; Hispanic $\lambda_{HIS} = 1.004$; South Asian $\lambda_{SAS} = 0.974$ (no correction made). SNVs reported in $\geq 50\%$ of the total effective sample size of the “reduced” ancestry-specific meta-analyses were included in downstream analyses: African $N_{AFR} \geq 11,613$; East Asian $N_{EAS} \geq 57,129$; European $N_{EUR} \geq 85,062$; Hispanic $N_{HIS} \geq 9,480$; South Asian $N_{SAS} \geq 15,789$. We identified loci attaining genome-wide significant evidence of association ($p < 5 \times 10^{-8}$) in each of the “reduced” ancestry-specific meta-analyses, and the lead SNV for each locus was selected as the variant with minimum association p -value. For each test GWAS, we then regressed the observed allelic effect estimates at lead SNVs, weighted by their corresponding variances, on the allelic effect estimates from the each of the ancestry-specific meta-analyses, as implemented in `grs.summary` function⁹ of the `gtx` package in R, and estimated the OR per unit of the weighted GRS and the corresponding percentage of T2D variance explained, measured by pseudo R^2 .

Predictive power of GRS in FinnGen. Individuals from FinnGen were genotyped with Illumina and Affymetrix arrays. After quality control, individuals were imputed with Beagle⁴¹⁰ up to the Finnish population-specific reference panel (SISu version 3), comprising 3,775 whole genome sequences (www.sisuproject.fi). We excluded individuals due to non-Finnish ancestry, relatedness, or missing age and/or sex. We estimated the positions of FinnGen on the three axes of genetic variation from the multi-ancestry meta-regression as the mean of the two Finnish studies, FUSION and METSIM (**Supplementary Tables 1 and 2**), which we denoted x_1 , x_2 and x_3 . We derived Finnish-specific “predicted” allelic effect estimates for each lead SNV from the multi-ancestry meta-regression to be used as weights in the GRS. For the j th SNV, the “predicted” effect was given by

$$\hat{b}_{TAj} = \alpha_{TA0j} + \sum_k \alpha_{TAkj} x_k,$$

where α_{TA0j} and α_{TAkj} denote the intercept and effect of the k th axis of genetic variation for the SNV from the multi-ancestry meta-regression.

For each individual, we calculated the centred GRS from the multi-ancestry meta-regression given by

$$GRS_{TAi} = \sum_j (G_{ij} - 2q_j) \hat{b}_{TAj}$$

for the i th individual. In this expression, q_j denotes the frequency of the effect allele at the j th SNV, and G_{ij} is the effect allele dosage for the i th individual, which we replaced by $2q_j$ if the genotype was missing. We excluded lead SNVs from the GRS that were not reported in FinnGen. T2D status was defined using two variables: T2D-I and T2D-II. T2D-I included individuals with ICD-10 E11 and/or ICD-9 250*A, but excluded individuals with pancreatitis. T2D-II included individuals with ICD-10 E11, T2D complications, or medicine purchases of Anatomical Therapeutic Chemical (ATC) class A10B (blood glucose lowering drugs, excluding insulins). We defined controls as non-diabetic individuals (type 1, type 2, or undefined). We excluded individuals with missing T2D status or BMI from subsequent analyses, resulting in a total of 18,111 affected individuals and 111,119 unaffected individuals.

In a logistic regression framework, we first fitted a “null” model that included age, sex, genotyping batch, and ten axes of genetic variation to account for population structure. We then fitted models that additionally added BMI only, GRS only, and both BMI and GRS. For each model, we calculated the variance in T2D status explained (pseudo R^2) and the AUROC (calculated with a 10-fold cross-validation). We also conducted age-stratified analyses, after excluding age from the “null” model, within five age groups: under 50 years; 50-60 years; 60-70 years; 70-80 years; and over 80 years. We next considered the subset of individuals in the highest and lowest deciles of the GRS, testing for association of T2D status with an indicator variable of high/low decile in a logistic regression framework after adjustment for age, sex, BMI, genotyping batch, and ten axes of genetic variation. Finally, we considered T2D cases only, and tested for association of age of diagnosis of the disease with GRS in a linear regression framework after adjustment for sex, BMI, genotyping batch, and ten axes of genetic variation.

Selection analyses. We tested for evidence of selection for index SNVs for distinct T2D association signals, which were partitioned into two groups, risk and protective, according to the direction of the allelic effect when aligned to the derived allele. For each population, we excluded index SNVs that were not segregating in that population. We also excluded SNVs with T2D association p -value >0.5 for the ancestry group to which the population belongs because allelic effect estimates are close to zero and are imprecise. To test for selection, we sampled 20 variants for each index SNV, selected at random from those with the same derived allele frequency for the population in the 1000 Genomes Project reference panel. We conducted a one-sided Wilcoxon rank-sum test of whether the index SNVs have smaller than expected selection p -values, derived by *Relate*¹¹, when compared to the rest of the genome. We repeated this test 20 times and reported the mean p -value across replicates.

We next tested for selection on a range of traits available in the UK Biobank¹² at the subset of index SNVs for which the derived allele increased risk of T2D. We downloaded association summary statistics from <http://www.nealelab.is/uk-biobank>, which were derived from 361,194 white British individuals using PHEASANT¹³. We considered traits for which the number of significantly associated T2D index SNVs ($p < 0.00015$, Bonferroni

correction for 338 variants) exceeded ten (**Extended Data Figure 10**). For each population, we then evaluated the evidence for selection for the subset of associated index SNVs for the trait, using the same approach as described above. We also conducted a one-side Wilcoxon rank-sum test of whether the subset of index SNVs had younger age, conditional on derived allele frequency, when compared with the rest of the genome.

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Inserted DNA sequences and primers used for functional experimentation at the *PROX1* locus.

Signal 1

Credible set SNP: rs340874 (chr1:206600992, G)

Primers used for amplification:

Primer_F: **TTGTGGGCTAAAGTGCAAGC**

Primer_R: **GGGTGTATTGAGCGGGGAAA**

DNA fragment used for constructs: (>hg19_dna range=chr1:214159081-214159431)

CTATGTGCAATTGACACAAAC**TTGTGGGCTAAAGTGCAAGC**CATTTTTTCGCGTTTGAATCTTTTCTCTGTCCTGAC
TCCTTTCTCCCTACTCCCTCCTCTCTGCTCTCCGCCCTTTAAATGCAAACTGAGCAGATGGTTTTAAGGT
GTGGAAAGGTATATAGCCCTACTCCTACCAGTTATTTGTGGCTGGCGCTAACTTATATGTACAAACCAAGATTCCTA
AAGAAAAGTGTAGGACGAAAATAAGAAAGAAAGTAGCTTTGATCCATTCTCAGATCCCAAG**TTTCCCGCTCAATAC**
ACCGGCTTACCTCGAAGGACCCAACCAAT

Signal 2

Credible set SNP 1: rs17712208 (chr1:214150445, T)

Credible set SNP 2: rs79687284 (chr1:214150821, G)

Primers used for amplification:

Primer_F: **GGACTTCACTGGCAGACACA**

Primer_R: **TACTCATTCCCTGGCTTTGC**

DNA fragment used for constructs: (>hg19_dna range=chr1:214149600-214151600)

GGAAACACTTAAACCACACTGATTATACAGATTTTTCTCATCTAATGGTA TAATATCTATAACTACATGCATCTGGTAAC
ATACTAAATGCTGTCTAGAACAAAGCAAAGCAAACCAAAAGCCCAATTTAAATTAATAAAAAAAAAAAGATCTA
AGCCACCTTCGTAAACATGTGCTCTGTAAAGAAAGTAAAAAAAAAAAAAAAAAGAAAGAAAAGAAAAAGAT
CCACCAGAAGTACCAAAATCAAATATTTAGTCTTTAATTTCTACTTATTTGAAATCAAAAATATTTTCATGCTTT
GATAGGAATGTGCGTTGTTATCATTTCAAAAGCACTTCTCCCTTTATCACGAATCGAAGAAGAACTAACATTGAGAA
ACAAGGAACCAGAAATTTAGAGATGCTGGGAATAACTACAACCTAACATGGTCAAGGGAGAGAAAATATGATCCTC
TCAGAAGAATATGTAACAACAATCAGAGCACATCTGGGATTTGATTCAAACCAACCTGGAAACCAGATTGGATCTCAA
GCTGTTCTGTGTATAC
AGAAAGTTATCCACAGATTAAACCCAGAAA**GGACTTCACTGGCAGACACA**GGCATAACTTTACTCCTTTTGTGATGAC
CCATGAGTGGGGTCTATGGCAGTCTGAATAGATGGGCCTTCTGTTGAAAGATTCTGCCTAATCCTTCCCACCAAGCA
GGGTCTAAAGGTGTCAGCAGGATTTGGCTGACTGGATCCTAATGGAGCTATGGTTAATTATTGACTGATTAGGGATT
TACCTTATCTTTCGTGAGGAGCTGGCTCAAGACTTAACGTAAGCAATTTAGAGCCAGGGTGAACCTACACACATGCC
TTTTTCTTCTTCTTGGGTCACTTTAGCTTGCCCTCCTCCATAATTCACATTCAGGACAGAAATGGCCAGTCTTACAA
GGCGTGGAGTCTCAAGAGCACCGAAAATGAGAGGGGCCAGGTCCACGTGACAAGTGTCCAGAGACAGAGGCTTA
GAGAAATGTGCTTTTGAACACAGTGTATGTGTAAGGTTTTCCAGTTAAGTCCCTGGAGAAAAAAAAAAAAAAAA
GCACTTGTCTTTGTCCTAAAAGGTCTGTGATGCCCGTGGGTGAGAAATCCACCCGCACTCCCCAAGGCCCTTG**GC**
AAAGCCAGGAATGAGTACAGGCAGCTCAGGCCAGCTGCCAGATAGAGGTGGCCCGTAAATGACAGGCTT
CCTCTGCACCTCAGCAGGGCCTTCTTTCTAACAGTCTCCCTTAAATGTTGGCGAATGTTGTTTTCCATTGACTCAAC
ATCTCGCCTGGTGAAGCCAGTGAAGAAAGTTGAGCGGGAGGGGAAAGTGGGAGAGAGTGATGCCAAAGCA
AAAGAGCGGGACGGTCAAGCCAGTTTCCAAACAAGCTAGACACCTGCTTTGAAAAGACAGTACCAAGCCTAGACTTC
TGGCTTCTTCTTCACTTTGATCAGCCTTTGTTCCCTGCGGCTCTGATGAGGCTCCCTGCCCTCCCTCACCACTGCC
CCTTCACTGGGAGCTACTTCACTTAAACCATCAAAAATTCATAGCTTTCTCTATGAATGTAAGTACTGTCTTATCTGAA
GAAAAGGGAAAACAGTTATTGGAAATGCATGAAAGAAAGAAAGGAATTTAAGGAAGAGAAAGTA
AGTGAGAAGAGAAAATTTGGAGAGAAAATAAAGAAAGG
GGAAAAGAGAATAGAGAAATAGAAATTAAGGCAAAAAGAAAAGAAATAAAAGGATAGAACAAAAATAAAGAAAAG
GGAGGAGGGCAAAGGAGAAAGACGGCAGAAAAGTAAATCCAAAAGGGAGCTTTTCTCCAGAAAGTCAAGTTTC

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Ethics statements

Anti-aging study cohort (AASC). The ethics committees of Ehime University Graduate School of Medicine approved all study procedures. Written informed consent was obtained from all participants.

Atherosclerosis Risk in Communities (ARIC). Institutional Review Board approvals were obtained at all study sites: National Heart, Lung, and Blood Institute, University of North Carolina at Chapel Hill, Wake Forest Baptist Medical Center, University of Mississippi Medical Center, University of Minnesota and Johns Hopkins University. All participants provided written informed consent.

Biobank Japan (BBJ). All participants provided written informed consent as approved by the ethical committees of the RIKEN Yokohama Institute and the Institute of Medical Science, University of Tokyo.

Beijing Eye Study (BES). Approval was obtained from the Medical Ethics Committee of the Beijing Tongren Hospital. All participants gave written informed consent.

BioMe Biobank (BIOME). Approval was obtained from the Institutional Review Board at the Icahn School of Medicine at Mount Sinai. All participants provided written informed consent for genomic data sharing.

Bangladesh Population Cohort (BPC). The conduct of the BPC was reviewed and approved by Ethical Committees of the Bangladesh Medical Research Council and Institutional Review Boards of the University of Chicago.

Cardiometabolic Genome Epidemiology (CAGE-AMAGASKI and CAKE-GWAS). Approval was obtained from the Institutional Review Boards at the National Center for Global Health and Medicine. All participants provided written informed consent.

Cardiometabolic Genome Epidemiology (CAGE-KING). Approval was obtained from the ethics committees of Aichi Gakuin University, Jichi Medical University, Nagoya University and Kyushu University. All participants provided written informed consent.

Coronary Artery Risk Development in Young Adults (CARDIA). Participating centers (Northwestern University, University of Alabama Birmingham, University of Minnesota, and Kaiser Foundation Research Institute) provided ethics approval for the CARDIA study, and all participants provided written informed consent to participate.

Cleveland Family Study (CFS). Approval was obtained from the Institutional Review Board of Mass General Brigham (formerly Partners HealthCare). Written informed consent was obtained from all participants.

China Health and Nutrition Survey (CHNS). Approval was obtained from the Institutional review Boards at the University of North Carolina at Chapel Hill, the Chinese National

Human Genome Center at Shanghai, and the Institute of Nutrition and Food Safety at the China Centers for Disease Control. All participants provided written informed consent.

Cardiovascular Health Study (CHS). Approval was obtained from the Institutional Review Boards at Wake Forest University, University of California, Davis, Johns Hopkins, University of Pittsburgh, and the University of Washington, Seattle. All participants provided written informed consent.

China Kadoorie Biobank (CKB). All participants provided written informed consent. Ethical approval was obtained from Oxford Tropical Research Ethics Committee (OxTREC) and from the Ethical Review Committees of the Chinese Centre for Disease Control and Prevention and the Chinese Academy of Medical Sciences/Peking Union Medical College.

Cebu Longitudinal Health and Nutrition Survey (CLHNS). Written informed consent was obtained from all participants. Study protocols were approved by the University of North Carolina Institutional review Board for the Protection of Human Subjects.

Diabetic Cohort and Singapore Prospective Study Program (DC/SP2). Study protocols were approved by the Singapore General Hospital Ethics Committee, and National University of Singapore Institutional Review Board. All participants provided written informed consent.

Durban Diabetes Study and Durban Diabetes Case Control (DDS/DCC). Approvals were granted by the Biomedical Research Ethics Committee at the University of KwaZulu-Natal and the UK National Research Ethics Service. All participants provided written informed consent.

deCODE genetics (DECODE). The study was approved by the Icelandic National Bioethics Committee (approval no. VSN-16-112) after evaluation by the Icelandic Data Protection Authority. We obtained written informed consent for all participants in this study who donated samples. All data processing complies with the Icelandic Data Protection Authority (no. PV_2017060950þS).

Diabetes Gene Discovery Group (DGDG). All participants signed informed consent, and the protocol was approved by the French ethics committee.

Diabetes Genetics Initiative (DGI). The study was approved by the Ethics Committees of the Helsinki University Hospital, Helsinki, Finland, and Lund University, Sweden.

Estonian Genome Center of the University of Tartu (EGCUT). All analyses were approved by the Ethics Review Committee of the University of Tartu. All participants provided written informed consent.

Electronic Medical Records and Genomics Network (EMERGE). Approval was obtained from the Institutional Review Boards at Boston Children's Hospital, Children's Hospital of Philadelphia, Cincinnati Children's Hospital Medical Center, Essentia Institute of Rural Health, Geisinger Clinic, Group Health Cooperative, Marshfield Clinic Research Foundation, Mayo Clinic, Icahn School of Medicine at Mount Sinai, Northwestern University,

Pennsylvania State University, Vanderbilt University Medical Center, and University of Washington. All participants provided written informed consent.

European Prospective Investigation into Cancer and Nutrition (EPIC-INTERACT). The EPIC-InterAct study was approved by the local ethics committee in the participating countries and the Internal Review Board of the International Agency for Research on Cancer. All participants gave written informed consent. The study was coordinated by the Medical Research Council Epidemiology Unit at the University of Cambridge.

Epidemiologic Study of the Screenees for Diabetes Reduction Assessment with Ramipril and Rosiglitazone Medication (EPIDREAM). All study participants consented to analysis of blood samples. Approval was granted by the Hamilton Integrated Research Ethics Board, at McMaster University, Hamilton, Canada.

Family Heart Study (FAMHS). Approval was obtained from the Institutional Review Board at Washington University, St. Louis. Written informed consent, including consent to participate in genetic studies, was obtained from all participants.

Framingham Heart Study (FHS). Approval was obtained from the Institutional review Board of Boston University Medical Campus. All study participants provided written informed consent.

Finland-United States Investigation of NIDDM Genetics (FUSION). Approval was obtained from the coordinating Ethics Committee of the Hospital District of Helsinki and Uusimaa. All participants provided written informed consent.

German Chronic Kidney Disease (GCKD). All participants provided written informed consent. The study was registered in the national registry for clinical studies (DRKS 00003971) and was approved by local ethics committees.

Genetic Study of Atherosclerosis Risk (GENESTAR). Approval was obtained from the Johns Hopkins Medicine Institutional Review Board. All participants gave written informed consent.

Genetic Epidemiology Network of Arteriosclerosis (GENOA). Approval was granted by Institutional Review Boards of the University of Michigan, University of Mississippi Medical Center and Mayo Clinic. Written informed consent was obtained from all participants.

Resource for Genetic Epidemiology on Adult Health and Aging (GERA). The Institutional Review Boards for Human Subjects Research of both Kaiser Permanente Medical Care Plan (Northern California Region) and the University of California at San Francisco approved the project.

Genetics of Diabetes and Audit Research in Tayside Scotland (GODARTS). Approval was obtained from the Tayside Medical Ethics Committee. Informed consent was obtained for all participants.

Genetics of Latinos Diabetic Retinopathy (GOLDR). Approval was granted by the Institutional Review Board of the Lundquist Institute for Biomedical Innovation at Harbor-UCLA Medical Center.

Genetic Overlap Between Metabolic and Psychiatric Traits and Teens of Attica: Genes and Environment (GOMAP-TEENAGE). Ethical permission for TEENAGE was obtained from the Bioethics Committee of Harokopio University, Athens. Ethical permission for GOMAP was obtained from the Dromokaiteio Scientific Committee, Dromokaiteio Management Committee, Dafni Scientific Committee, Eginitio Scientific Committee and Harokopio Ethics Committee. All participants of GOMAP-TEENAGE gave written informed consent.

Genomic Research Cohort for CCMB Diabetes Study (GRCCDS). Ethics committees of CSIR-Centre for Cellular and Molecular Biology and KEM Hospital and Research Centre approved the project.

Health, Aging and Body Composition Study (HABC). The Institutional Review Boards at the University of Memphis and the University of Pittsburgh granted approval to conduct the Health ABC Study, and all participants provided written informed consent.

Healthy Aging in Neighborhoods of Diversity Across the Life Span Study (HANDLS). Approval was granted by the National Institutes of Health Institutional Review Board (study number 09AGN248). All participants provided written informed consent.

Hispanic Community Health Study/Study of Latinos (HCHS/SOL). Approval was obtained from Institutional Review Boards at the University of North Carolina at Chapel Hill, Albert Einstein College of Medicine, University of Illinois at Chicago, University of Miami, and San Diego State University. All participants provided written informed consent.

Hong Kong Diabetes Registry (HKDR). Approval was obtained from the Chinese University of Hong Kong Clinical Research Ethics Committee.

Health Professionals' Follow-Up Study (HPFS). Approval was obtained from the Human Research Committee at the Brigham and Women's Hospital. All participants provided written informed consent.

Mexican American Hypertension and Insulin Resistance (HTNIR). Approval was granted by Human Subjects Protection Institutional Review Boards at the University of California at Los Angeles, University of Southern California, Lundquist/LABioMed/Harbor-UCLA and Cedars-Sinai Medical Center.

Howard University Family Study (HUFS). All human participants from the HUFS included in the analyses of this manuscript provided written informed consent prior to enrollment. The HUFS study was approved by the Institutional Review Board at Howard University.

Indian Diabetes Consortium (INDICO). Approval was obtained by the Human Ethics Committees of All India Institute of Medical Sciences, New Delhi and CSIR-Institute of Genomics and Integrative Biology, New Delhi, India, and was conducted in accordance with

the principles of Helsinki Declarations. Informed written consent was obtained from all of participants.

INTERHEART (INTERHEART). All study participants consented to analysis of blood samples. Approval was granted by the Hamilton Integrated Research Ethics Board, at McMaster University, Hamilton, Canada.

Jackson Heart Study (JHS). Approval was obtained from Institutional Review Boards at Jackson State University, Tougaloo College and the University of Mississippi Medical Center. All participants provided written informed consent.

Korean Association Resource (KARE). Approval was granted by the Institutional review Board at the Korean National Institute of Health. All participants provided written informed consent.

Korean Biobank Array from the Korean Genome and Epidemiology (KoGES) Consortium (KBA). Approval was granted by the Institutional Review Board of the Korean National Institute of Health. All participants provided written informed consent.

Collaborative Health Research in the Region of Augsburg (KORA). Approval was granted by the Ethics Committee of the Medical Association of Bavaria (number 06068). All participants provided informed consent.

Los Angeles Latino Eye Study (LALES). Approval was obtained from the Los Angeles County/University of Southern California Institutional Review Board, and Western Institutional Review Board at Southern California Eye Institute. All participants provided written informed consent.

London Life Sciences Prospective Population (LOLIPOP). Approval was obtained from the London-Fulham Research Ethics Committee (ref 07/H0712/150). All participants gave an written informed consent.

Mexican American Study of Coronary Artery Disease (MACAD). Approval was granted by Human Subjects Protection Institutional Review Boards at the University of California at Los Angeles, University of Southern California, Lundquist/LABioMed/Harbor-UCLA and Cedars-Sinai Medical Center.

Mexico City (MC). Approval was obtained from Institutional Review Boards at the Ethics and Scientific Commission members and the AUTHORIZATION is issued with registration number R-2011-785-018 and the Conacyt SALUD-2010-02-150352. In Canada, approval was obtained from the Research Ethics Board from the University of Toronto (Protocol 15770).

Multi-Ethnic Study of Atherosclerosis (MESA). Approval was obtained from Institutional Review Boards at the University of Washington, Wake Forest School of Medicine, Northwestern University, University of Minnesota, Columbia University, Johns Hopkins University, Cedars-Sinai Medical Center, and the University of California at Los Angeles.

Metabolic Syndrome in Men (METSIM). Approval was granted by the Ethics Committee of the University of Kuopio and the Kuopio University Hospital. All participants gave written informed consent.

Mass General Brigham Biobank (MGB). The MGB Biobank protocol and informed consent documents are reviewed annually by the Partners-MGB Institutional Review Board (#2009P002312). All patients who participate in the MGB Biobank are consented for their samples to be linked to their identified clinical information. They have also consented for their information to be used for a broad range of research and for their deidentified information to be shared outside of MGB.

Michigan Genomics Initiative (MGI). Approval was granted by the IRBMED Institutional Review Board of the University of Michigan. All participants gave written informed consent.

Nagahama Study (NAGAHAMA). Approval was granted by the ethics committees of Kyoto University Graduate School of Medicine. Written informed consent was obtained from all participants.

Netherlands Epidemiology of Obesity (NEO). Approval was obtained from the Medical Ethics Committee of Leiden University Medical Center. All participants gave written informed consent.

Nurses Health Study (NHS). Approval was obtained from the Human Research Committee at the Brigham and Women's Hospital. All participants provided written informed consent.

NIDDM-Atherosclerosis Study Hispanic Cohorts (NIDDM). Approval was granted by Human Subjects Protection Institutional Review Boards at the University of California at Los Angeles, University of Southern California, City of Hope, Lundquist/LABioMed/Harbor-UCLA and Cedars-Sinai Medical Center.

Northwestern University Genetics (NUGENE). Approval was obtained from Institutional Review Boards at Northwestern University and Vanderbilt University.

Prospective Investigation of the Vasculature in Uppsala Seniors (PIVUS). Approval was granted by the Ethics Committee of Uppsala University. All participants provided written informed consent.

Pakistan Risk of Myocardial Infarction Study (PROMIS). The study was approved by the Institutional Review Board of the Center for Non-Communicable Diseases Pakistan and by regional Ethical Review Committees in the different centres across Pakistan involved in the study. Institutional Review Boards at the National Institute of Cardiovascular Disorders, Karachi, Punjab Institute of Cardiology, Lahore, and Tabba Heart Institute, Karachi approved the study. All participants provided written informed consent.

Prospective Study of Pravastatin in the Elderly at Risk (PROSPER). Approval was obtained from the Institutional Ethics Review Boards of Cork University (Ireland), Glasgow University

(UK) and Leiden University Medical Center (The Netherlands). All participants gave written informed consent.

Sea Islands Genetic Network Reasons for Geographic and Racial Differences in Stroke (REGARDS). The REGARDS study protocol was approved by the institutional review boards of each participating institution, and written informed consents were obtained from all participants.

Ragama Health Study (RHS). Approval was obtained from Institutional Review Boards at the National Center for Global Health and the University of Kelaniya (P38/09/2006). All participants provided written informed consent.

Rotterdam Study (RS). Approval was granted by the Institutional review Board at Erasmus University Medical Center. All participants provided written informed consent.

Shanghai Breast Cancer Study and Shanghai Women's Health Study (SBCS/SWHS). Approval was obtained from Institutional review Boards at Vanderbilt University Medical Center and Shanghai Cancer Institute. A written informed consent form was obtained from all study participants.

Singapore Chinese Eye Study (SCES). The study adhered to the Declaration of Helsinki. Ethical approval was obtained from the SingHealth Institutional Review Board and National University of Singapore Institutional Review Board. Written informed consent was obtained from all participants.

Starr County Health (SCH). All protocols were reviewed and approved by the Institutional Committee for the Protection of Human Subjects (HSC-SPH-02-042). All participants provided written informed consent permitting the collection and sharing of data.

Singapore Chinese Health Study (SCHS). Approval was obtained from the Institutional Review Board at the National University of Singapore. All participants provided written informed consent.

Slim Initiative for Genomic Medicine in the Americas (SIGMA). Approval was obtained from the Institutional Review Board of the Instituto Nacional de Ciencias Medicas y Nutricion Salvador Zubiran. All participants provided written informed consent.

Singapore Malay Eye Study (SIMES). The study adhered to the Declaration of Helsinki. Ethical approval was obtained from the SingHealth Institutional Review Board and National University of Singapore Institutional Review Board. Written informed consent was obtained from all participants.

Singapore Indian Eye Study (SINDI). The study adhered to the Declaration of Helsinki. Ethical approval was obtained from the SingHealth Institutional Review Board and National University of Singapore Institutional Review Board. Written informed consent was obtained from all participants.

Samsung Medical Center (SMC). Approval was obtained from the Institutional Review Board of the Samsung Medical Center (No. 2004-12-005). All participants provided written informed consent.

Seoul National University Hospital (SNUH). The Institutional Review Board of the Biomedical Research Institute at Seoul National University Hospital approved the study protocol (1205–130–411). Written informed consent was obtained from each participant.

Taiwan MetaboChip Consortium Zhonghua (TAICHI-G). Approval was granted by Institutional Review Boards at Stanford University School of Medicine, Hudson-Alpha Biotechnology Institute, Lundquist/LABioMed/Harbor-UCLA, Cedars-Sinai Medical Center, Taichung Veterans General Hospital, Taipei Veterans General Hospital, National Health Research Institute, Tri-Service General Hospital, and National Taiwan University Hospital.

Taiwan Type 2 Diabetes (TWT2D). Approval was obtained from Institutional Review Boards at China Medical University Hospital, Chia-Yi Christian Hospital, and National Taiwan University Hospital.

Danish T2D Case-Control Study (UCPH). The studies included in the Danish T2D Case-Control Study (UCPH) were conducted in accordance with the Declaration of Helsinki II and were approved by the local Ethical Committees of Copenhagen County, the Capital Region of Denmark, or the Region of Southern Denmark.

UK Biobank (UKBB). Approval was obtained from the North West Centre for Research Ethics Committee (11/NW/0382).

Uppsala Longitudinal Study of Adult Men (ULSAM). Approval was granted by the Ethics Committee of Uppsala University. All participants provided written informed consent.

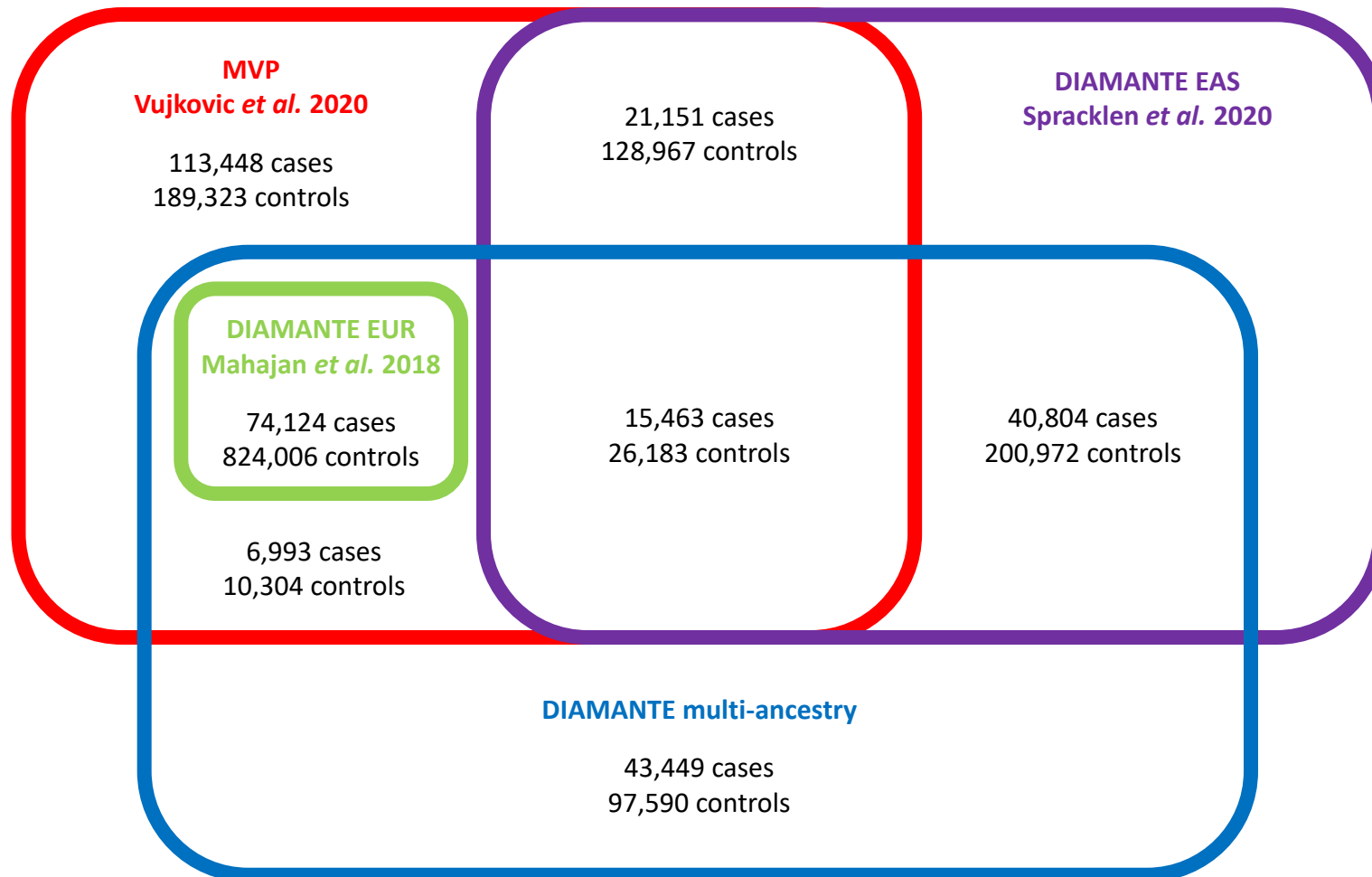
Wake Forest School of Medicine (WFSM). Approval was granted by the Institutional Review Board at Wake Forest School of Medicine. All participants provided written informed consent.

Women's Health Initiative (WHI). Approval was granted by the Institutional review Board at the Fred Hutchinson Cancer Research Centre in accordance with the US Department of Health and Human Services regulations at 45 CFR 46 (approval number IR# 3467-EXT). All participants provided written informed consent. Additional written consent to review medical records was obtained. The Fred Hutchinson Cancer Research Centre has an approved FWA on file with the Office for Human Research Protections under assurance number 0001920.

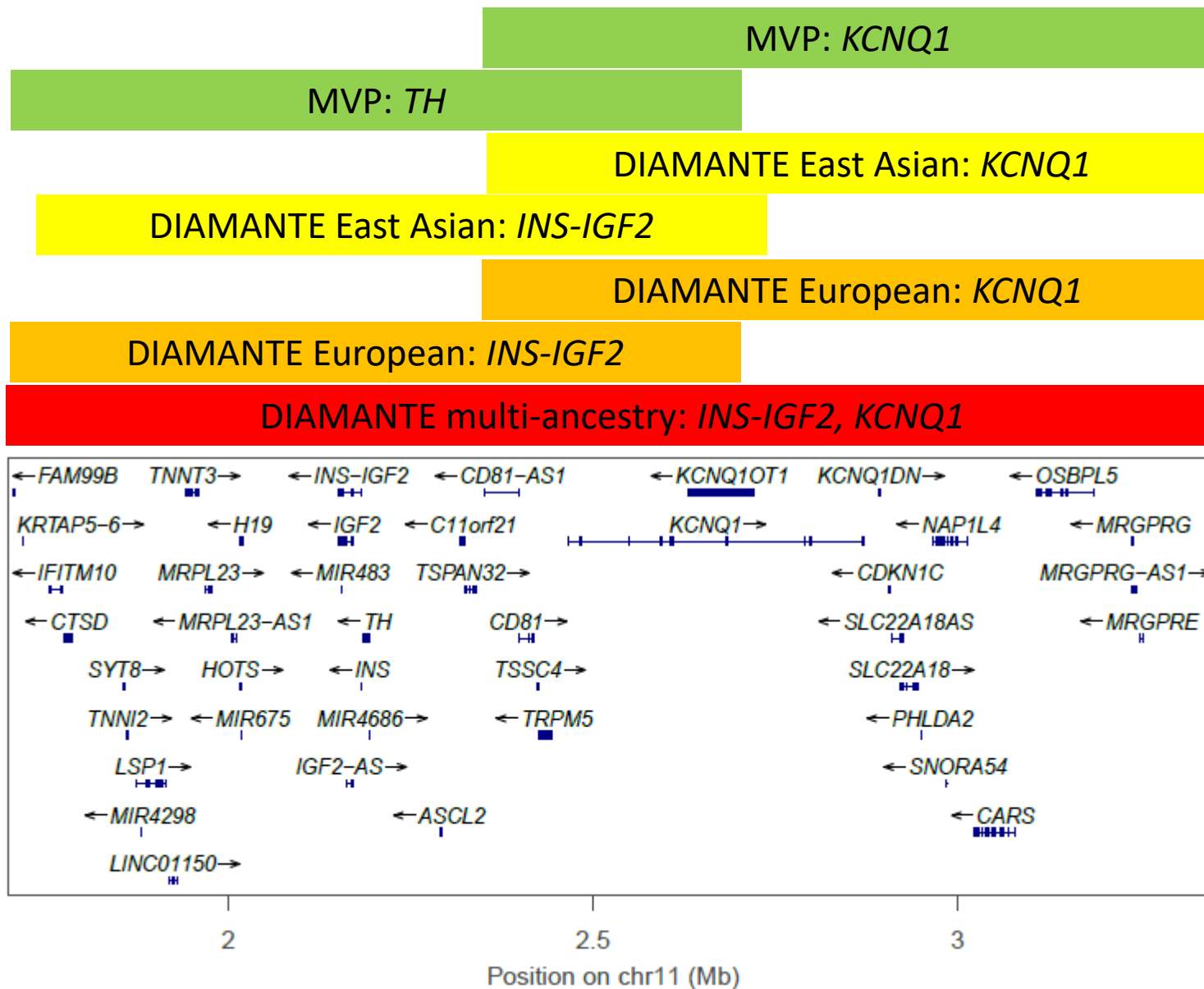
Wellcome Trust Case Control Consortium (WTCCC). Approval for the study was obtained from Peterborough & Fenland Local Research Ethics Committee, National Research Ethics Service, Leeds (East) Research Ethics Committee, South West Multicentre Research Ethics Committee, Tayside Committee on Medical Research Ethics and Oxford Tropical Research Ethics Committee.

FinnGen. Patients and control subjects in FinnGen provided informed consent for biobank research, based on the Finnish Biobank Act. Alternatively, separate research cohorts, collected prior the start of FinnGen (August 2017), were collected based on study-specific consents and later transferred to the Finnish biobanks after approval by Fimea, the National Supervisory Authority for Welfare and Health. Recruitment protocols followed the biobank protocols approved by Fimea. The Coordinating Ethics Committee of the Hospital District of Helsinki and Uusimaa (HUS) approved the FinnGen study protocol Nr HUS/990/2017. The FinnGen project is approved by Finnish Institute for Health and Welfare (THL), approval number THL/2031/6.02.00/2017, amendments THL/1101/5.05.00/2017, THL/341/6.02.00/2018, THL/2222/6.02.00/2018, THL/283/6.02.00/2019), Digital and population data service agency VRK43431/2017-3, VRK/6909/2018-3, the Social Insurance Institution (KELA) KELA 58/522/2017, KELA 131/522/2018, KELA 70/522/2019 and Statistics Finland TK-53-1041-17. The Biobank Access Decisions for FinnGen samples and data utilized in FinnGen Data Freeze 4 include: THL Biobank BB2017_55, BB2017_111, BB2018_19, BB_2018_34, BB_2018_67, BB2018_71, BB2019_7 Finnish Red Cross Blood Service Biobank 7.12.2017, Helsinki Biobank HUS/359/2017, Auria Biobank AB17-5154, Biobank Borealis of Northern Finland_2017_1013, Biobank of Eastern Finland 1186/2018, Finnish Clinical Biobank Tampere MH0004, Central Finland Biobank 1-2017, and Terveystalo Biobank STB 2018001.

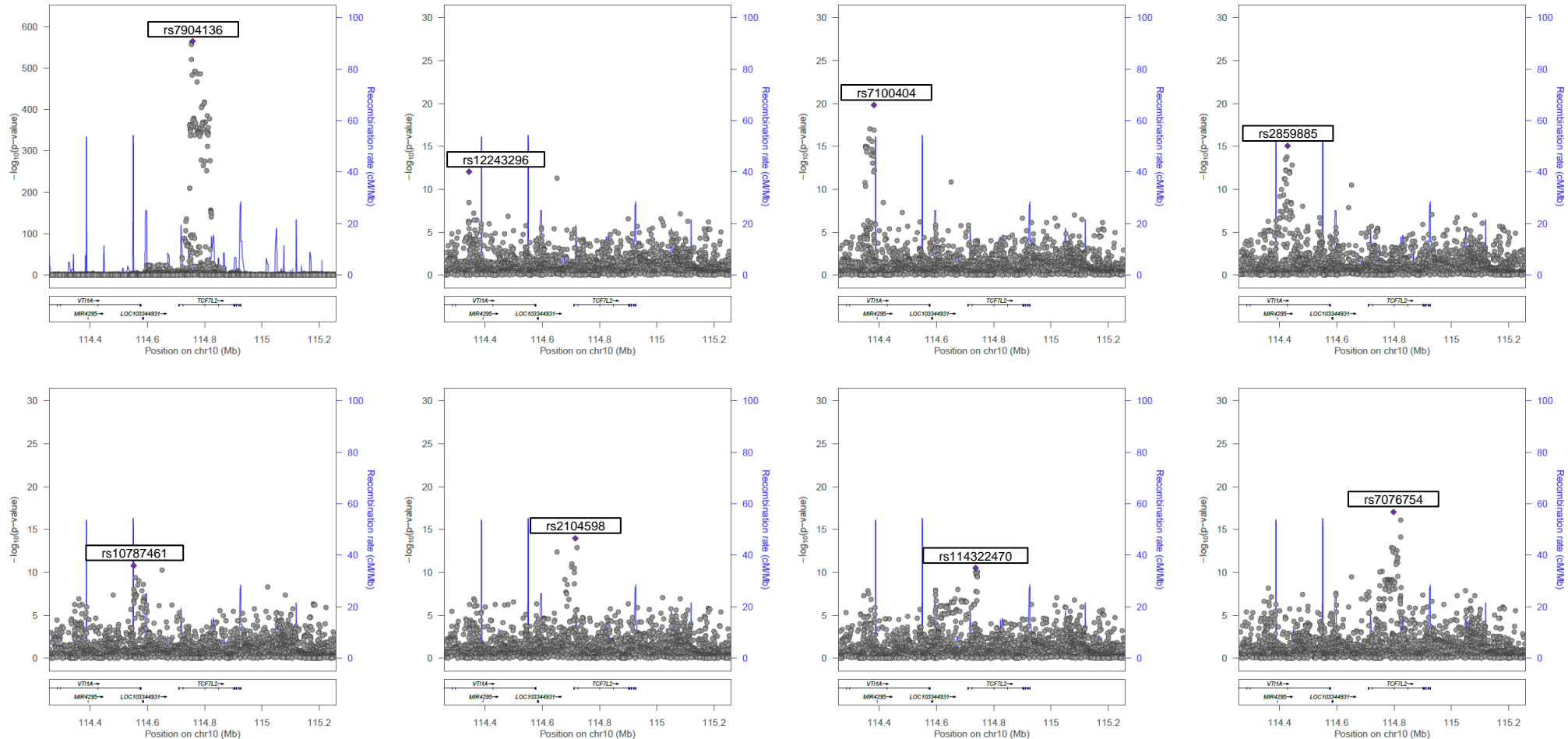
Supplementary Figure 1. Overlap of samples from the DIAMANTE multi-ancestry meta-analysis with recent investigations incorporating T2D GWAS from the DIAMANTE Consortium. The DIAMANTE multi-ancestry meta-analysis includes 180,834 cases and 1,159,055 controls of diverse ancestry, of which 137,385 cases and 1,061,465 controls (77.8% of total effective sample size) have contributed to previous investigation of the genetic contribution to T2D.

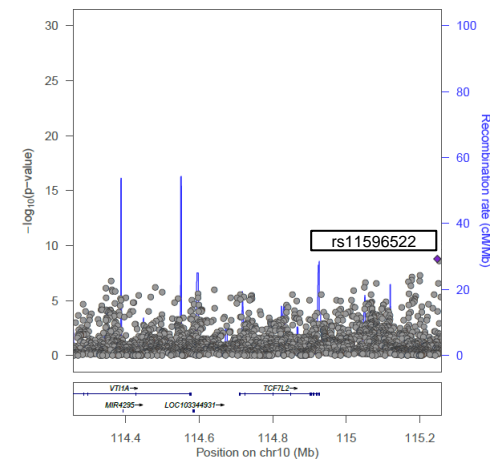
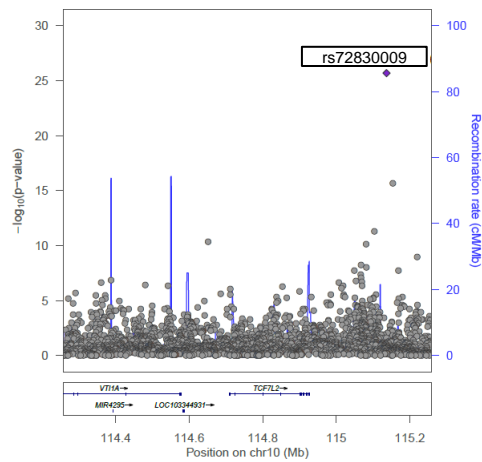
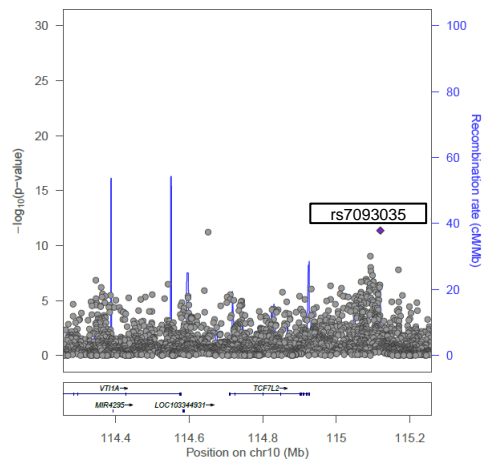
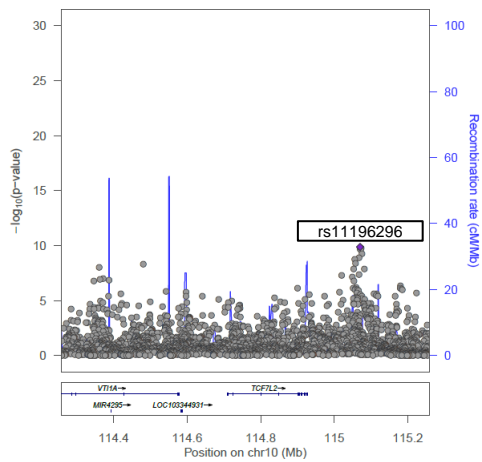
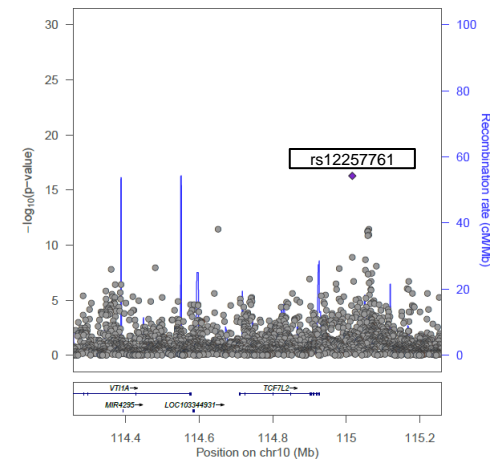
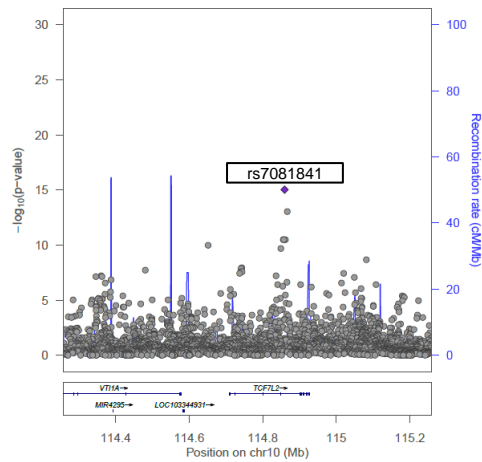
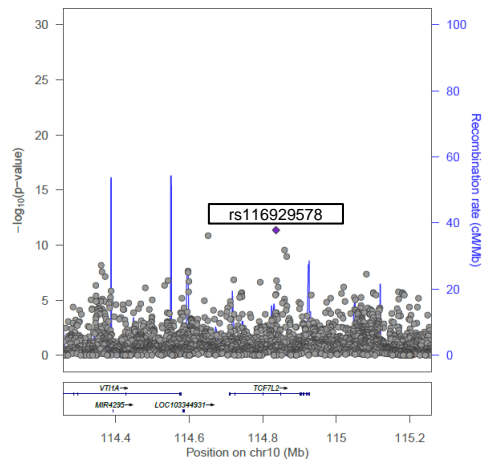
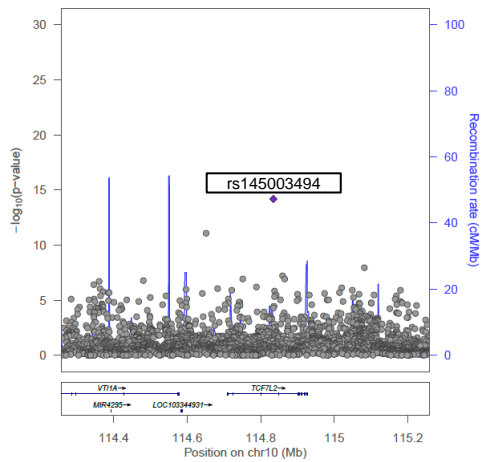


Supplementary Figure 2. Construction of loci across studies incorporating GWAS from the DIAMANTE Consortium. The locus encompassing T2D association signals at *INS-IGF2* and *KCNQ1* was defined by combining overlapping loci across studies and included the region spanning chromosome 11 from 1,697,132bp to 3,358,546bp (build 37).

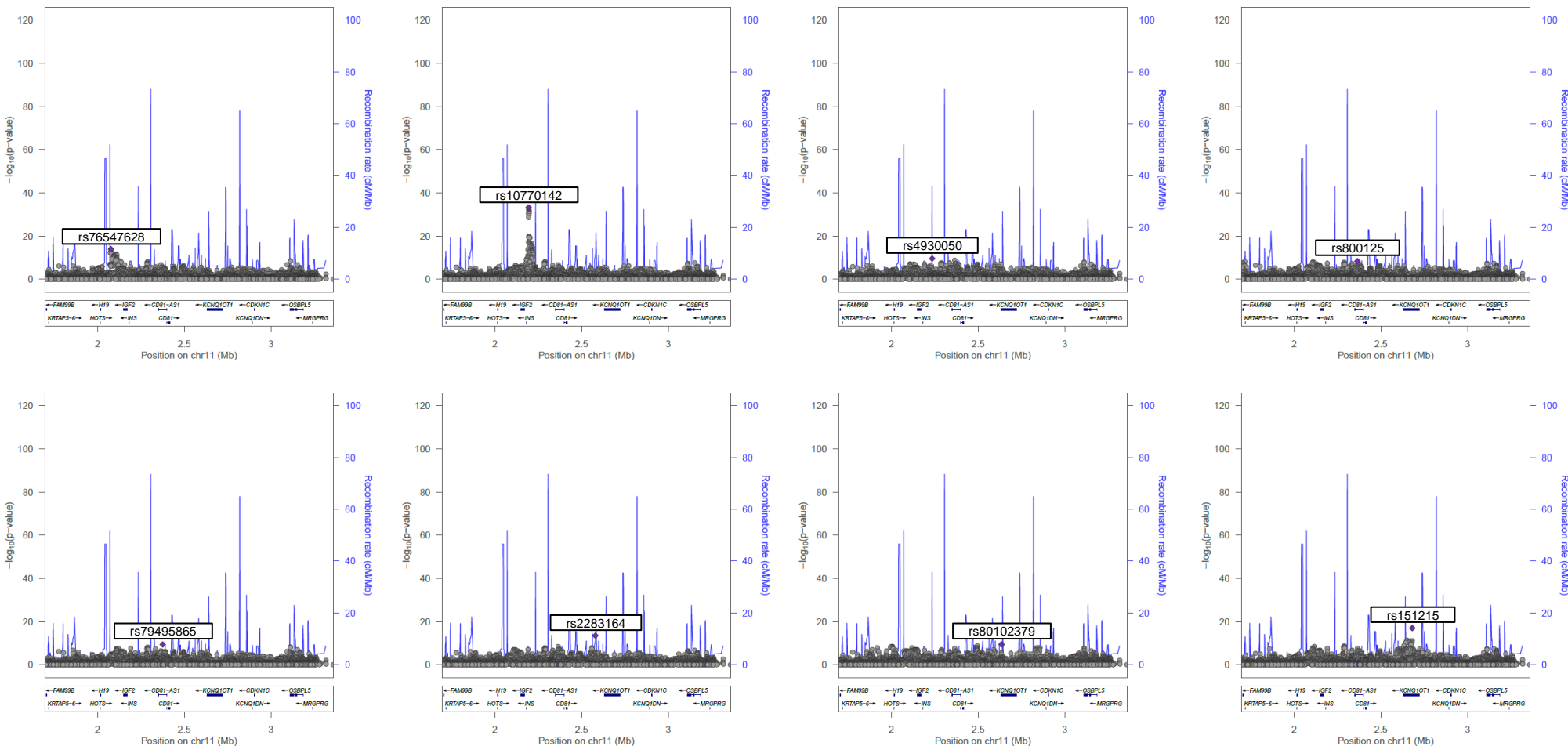


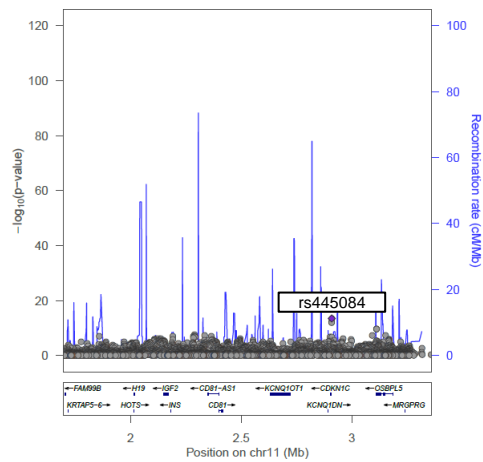
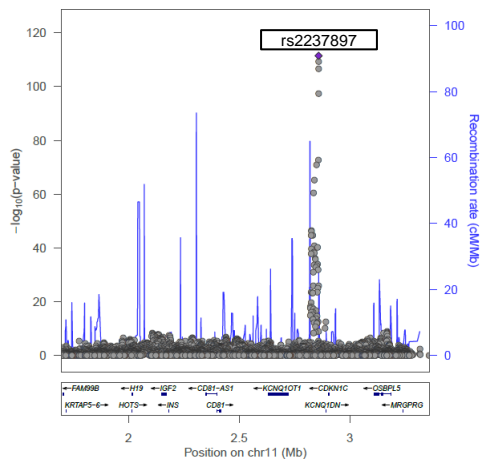
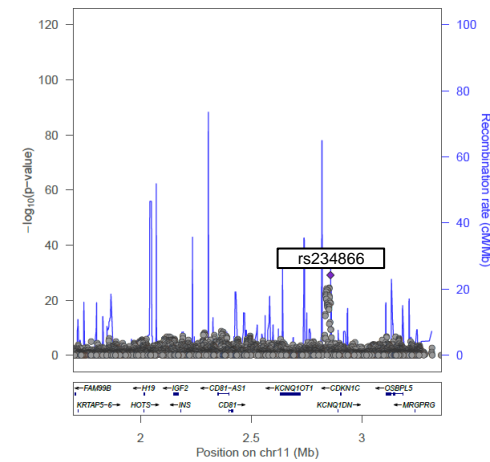
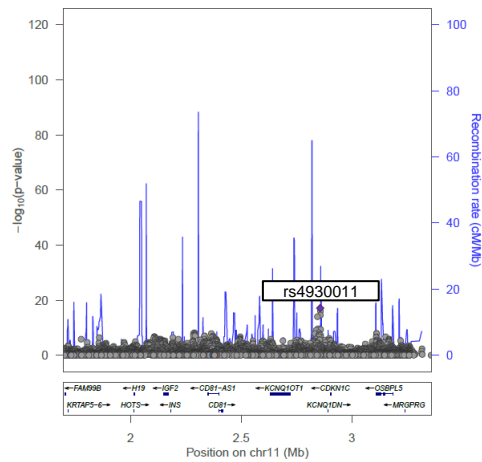
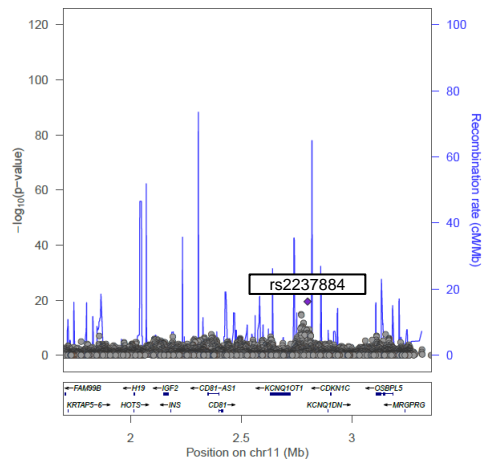
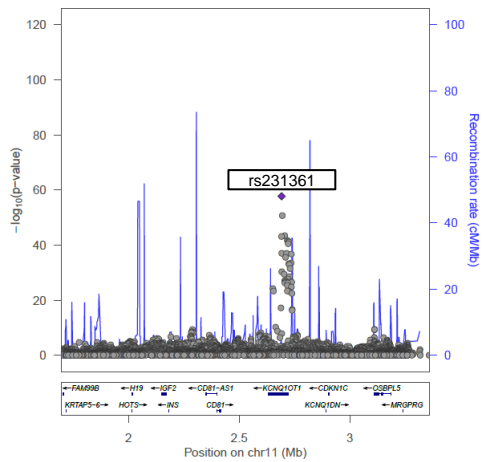
Supplementary Figure 3. Signal plots for distinct T2D association signals at the *TCF7L2* locus from multi-ancestry meta-regression (MR-MEGA) of up to 180,834 cases and 1,159,055 controls. Association summary statistics for each signal are obtained from approximate conditioning after adjusting for all other index SNVs at the locus. Each point represents a SNV passing quality control in the multi-ancestry meta-regression, plotted with their conditional p -value (on a $-\log_{10}$ scale) as a function of genomic position (NCBI build 37). In each plot, the index variant is represented by the purple diamond. Gene annotations are taken from the University of California Santa Cruz genome browser. Recombination rates are estimated from the Phase II HapMap.



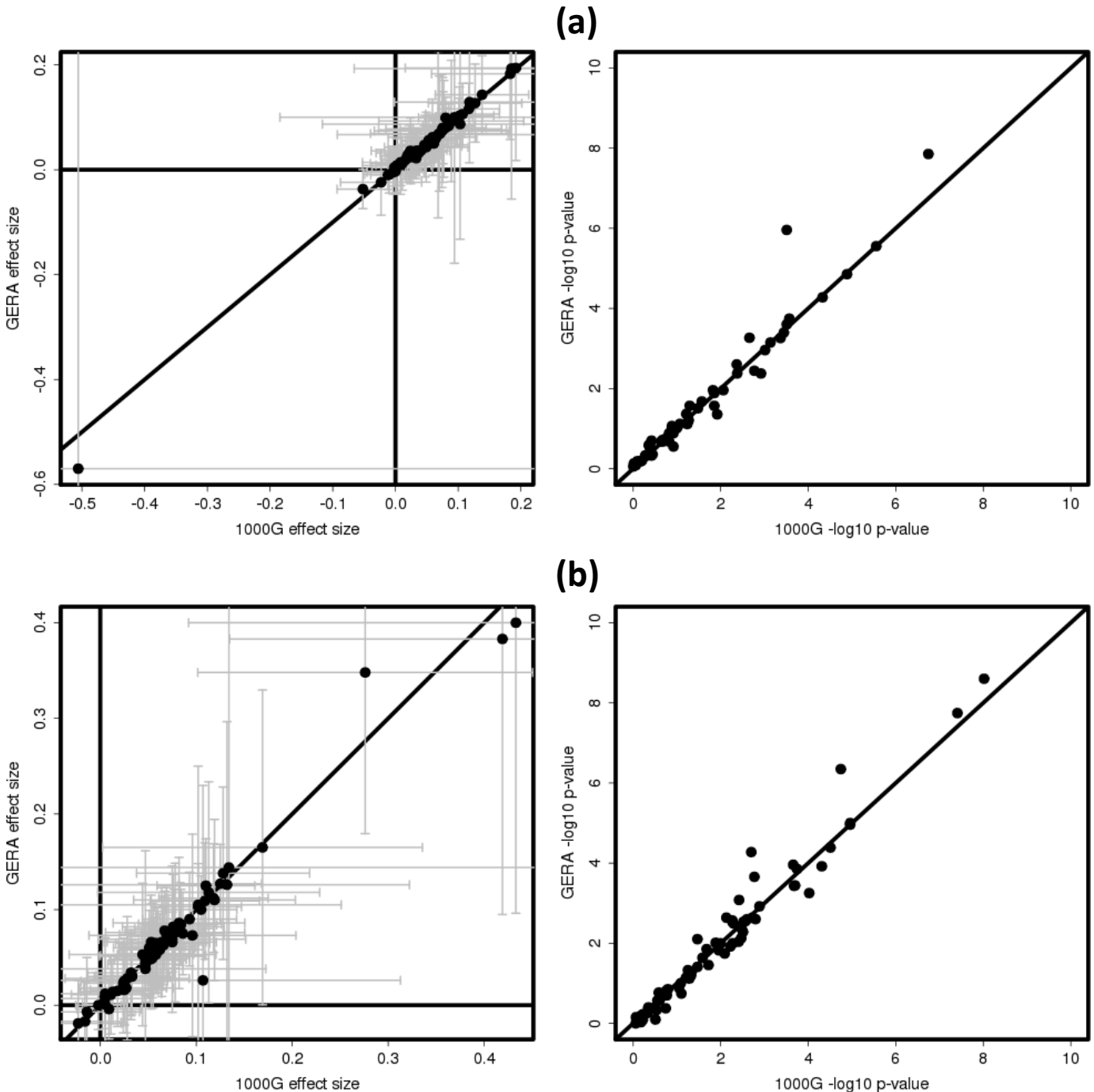


Supplementary Figure 4. Signal plots for distinct T2D association signals at the *INS-IGF2-KCNQ1* locus from multi-ancestry meta-regression (MR-MEGA) of up to 180,834 cases and 1,159,055 controls. Association summary statistics for each signal are obtained from approximate conditioning after adjusting for all other index SNVs at the locus. Each point represents a SNV passing quality control in the multi-ancestry meta-regression, plotted with their conditional p -value (on a $-\log_{10}$ scale) as a function of genomic position (NCBI build 37). In each plot, the index variant is represented by the purple diamond. Gene annotations are taken from the University of California Santa Cruz genome browser. Recombination rates are estimated from the Phase II HapMap.



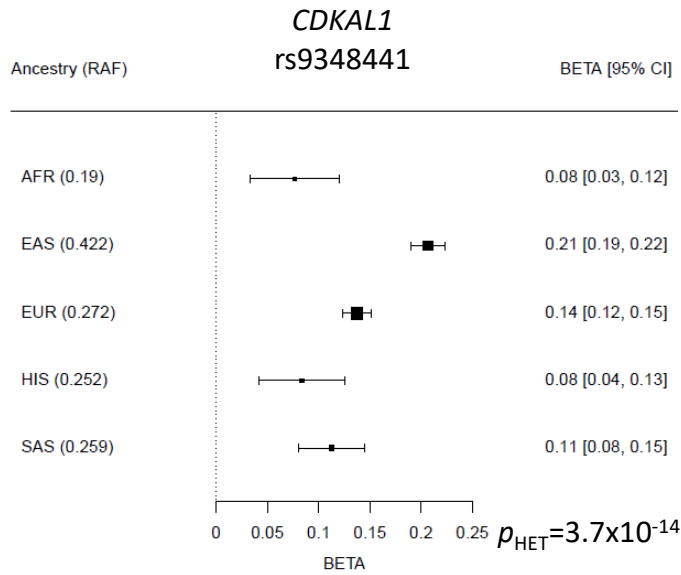


Supplementary Figure 5. Comparison of ancestry-specific association summary statistics obtained from approximate conditional analysis undertaken in loci with multiple distinct signals using LD reference panels from the 1000 Genomes Project and GERA. (a) African ancestry-specific association summary statistics derived from 661 individuals of African ancestry from the 1000 Genomes Project and 1,000 African American individuals from GERA. Association summary statistics were derived from a meta-analysis of 15,487 T2D cases and 23,709 controls. In the left panel, each point represents the log-odds ratio from the approximate conditional analysis and the error bars represent 95% confidence limits. (b) Hispanic ancestry-specific association summary statistics derived from 347 individuals of Hispanic ancestry from the 1000 Genomes Project and 1,000 Hispanic individuals from GERA. Association summary statistics were derived from a meta-analysis of 12,385 T2D cases and 21,423 controls. In the left panel, each point represents the log-odds ratio from the approximate conditional analysis and the error bars represent 95% confidence limits.

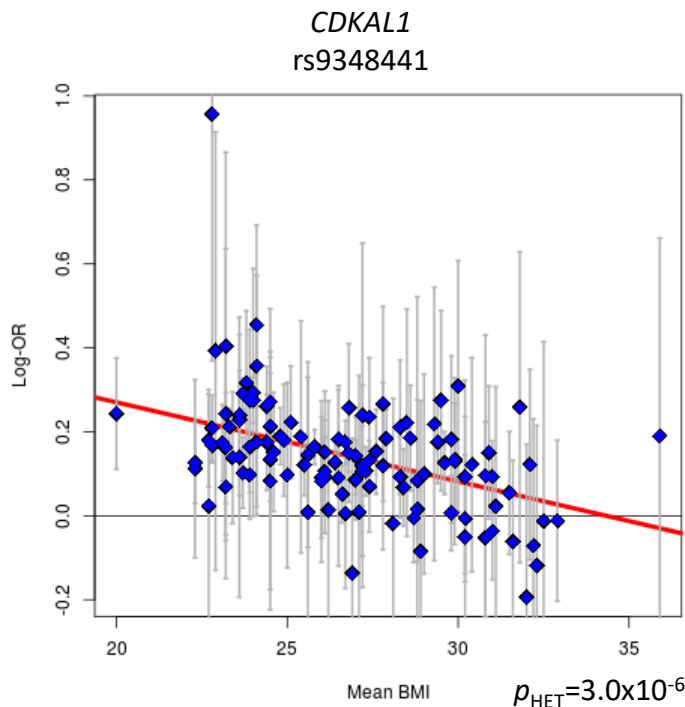


Supplementary Figure 6. Source of heterogeneity in allelic effects on T2D at the *CDKAL1* locus. (a) Forest plot presenting ancestry-specific allelic effects. The plot presents the risk allele frequency (RAF), the point represents the log-OR (BETA) for the risk allele, and the bars represent the corresponding 95% confidence interval (CI), from ancestry-specific fixed-effects meta-analysis. The size of each point represents the relative inverse-variance of the log-OR. The sample size contributing to each ancestry: African 15,043 cases and 22,318 controls; East Asian 56,268 cases and 227,155 controls; European 67,192 cases and 831,463 controls; Hispanic 11,027 cases and 18,885 controls; and South Asian 16,540 cases and 32,952 controls. (b) Correlation between study-level allelic effects and mean BMI. In the plot, each point represents a study contributing to the multi-ancestry meta-regression, plotted according to the mean BMI on the x-axis and the log-OR for the risk allele on the y-axis. The bars represent the 95% confidence interval for the log-OR. The red line is the line of best fit from linear regression of mean BMI on log-OR.

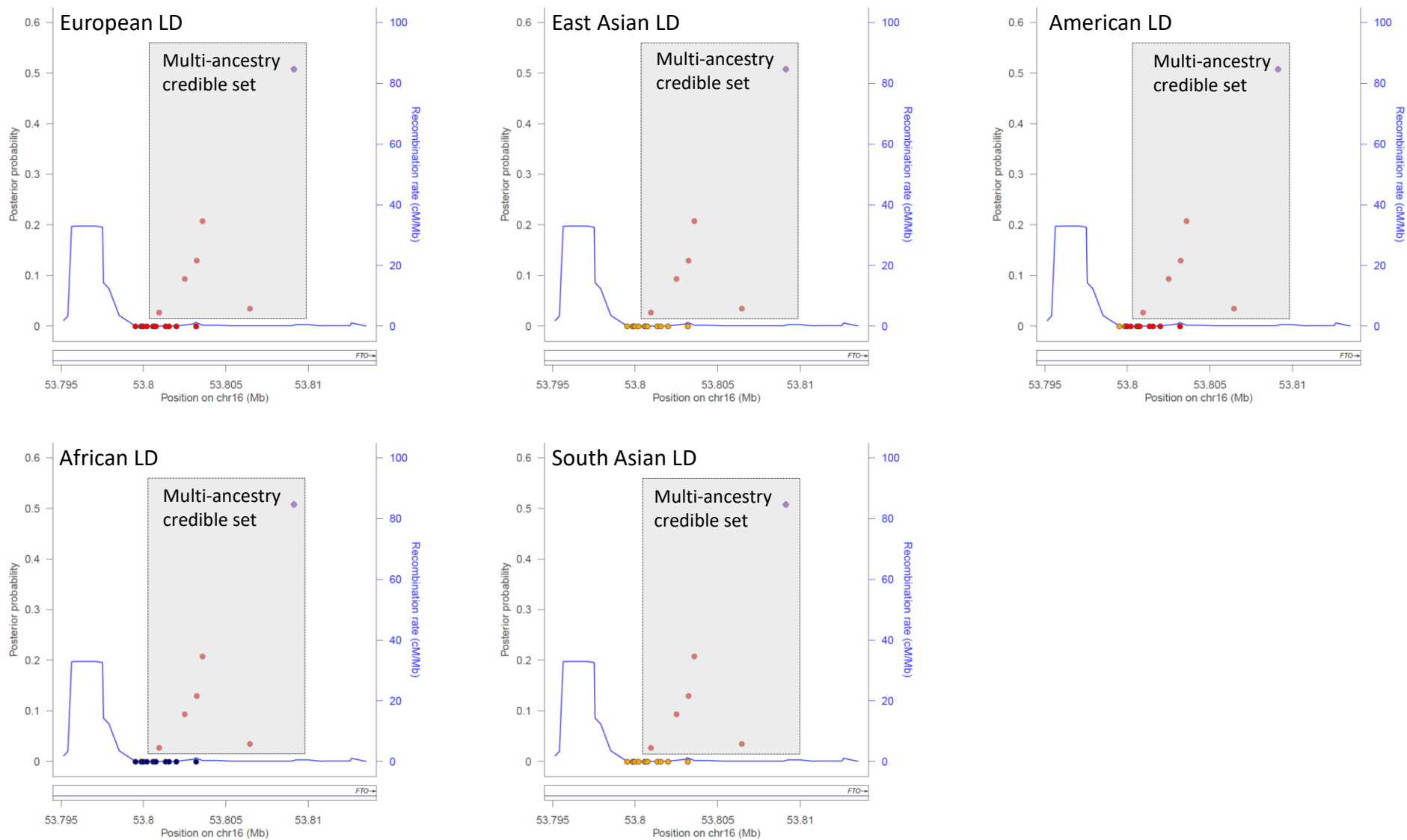
(a)



(b)

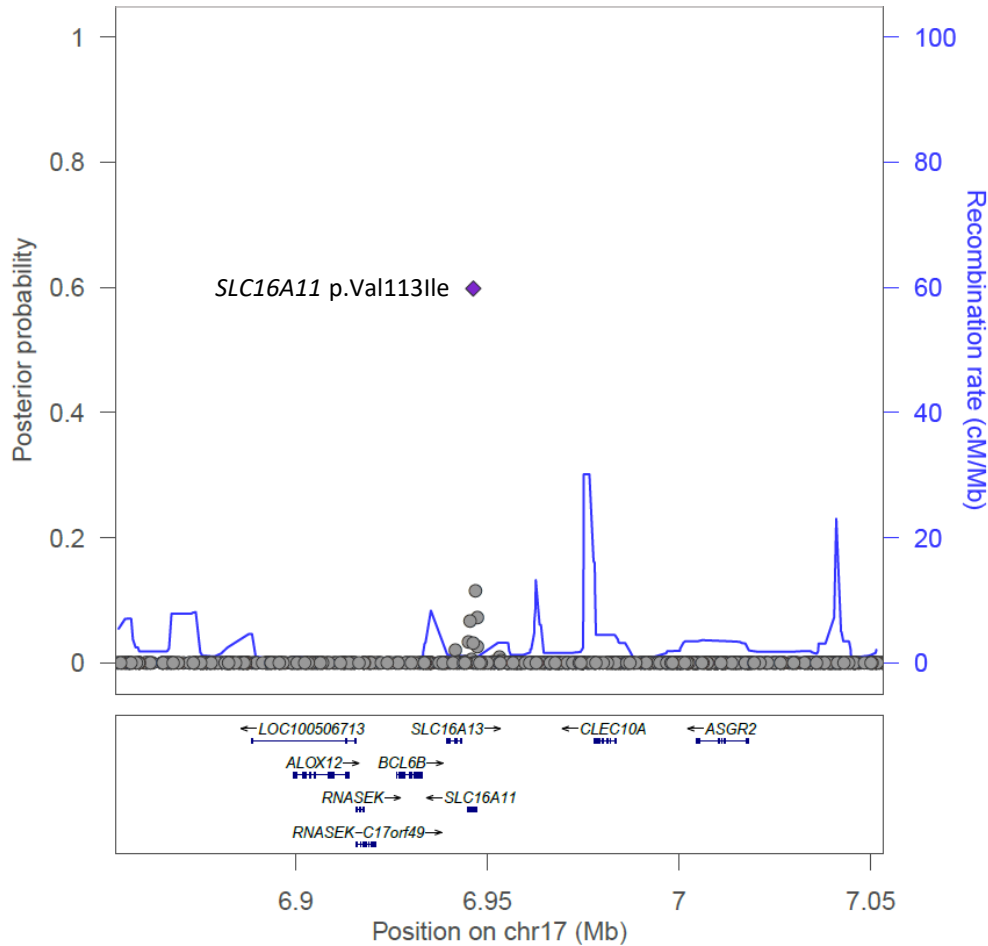


Supplementary Figure 7. Refinement of European ancestry-specific 99% credible set (under uniform prior model of causality) for T2D association at the *FTO* locus after multi-ancestry meta-regression (MR-MEGA) of up to 180,840 cases and 1,159,185 controls. Each point represents a credible set variant, plotted with their posterior probability of association as a function of genomic position (NCBI build 37). The index SNV (rs55872725) is represented by the purple symbol. The colour coding of all other SNVs indicates LD with the index variant in ancestry-specific haplotypes from the 1000 Genomes Project reference panel: red $r^2 \geq 0.8$; gold $0.6 \leq r^2 < 0.8$; green $0.4 \leq r^2 < 0.6$; cyan $0.2 \leq r^2 < 0.4$; blue $r^2 < 0.2$; grey r^2 unknown. Recombination rates are estimated from Phase II HapMap and gene annotations are taken from the University of California Santa Cruz genome browser.

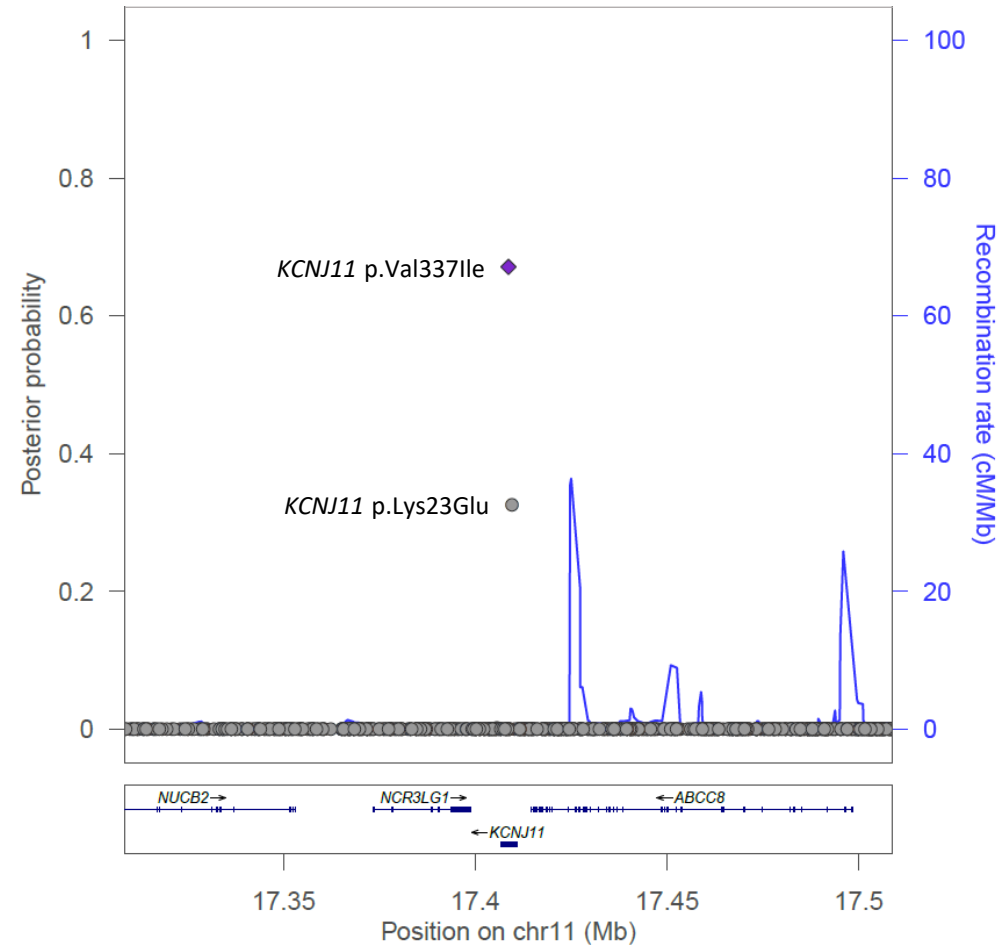


Supplementary Figure 8. Examples of improved fine-mapping of T2D association signals driven by missense variants from multi-ancestry meta-regression (MR-MEGA) of up to 180,834 cases and 1,159,055 controls. Each point represents a SNV passing quality control in the multi-ancestry meta-regression, plotted with their annotation-informed posterior probability of driving T2D association as a function of genomic position (NCBI build 37). In each plot, the index variant is represented by the purple diamond. Gene annotations are taken from the University of California Santa Cruz genome browser. Recombination rates are estimated from the Phase II HapMap.

SLC16A11-SLC16A13 locus

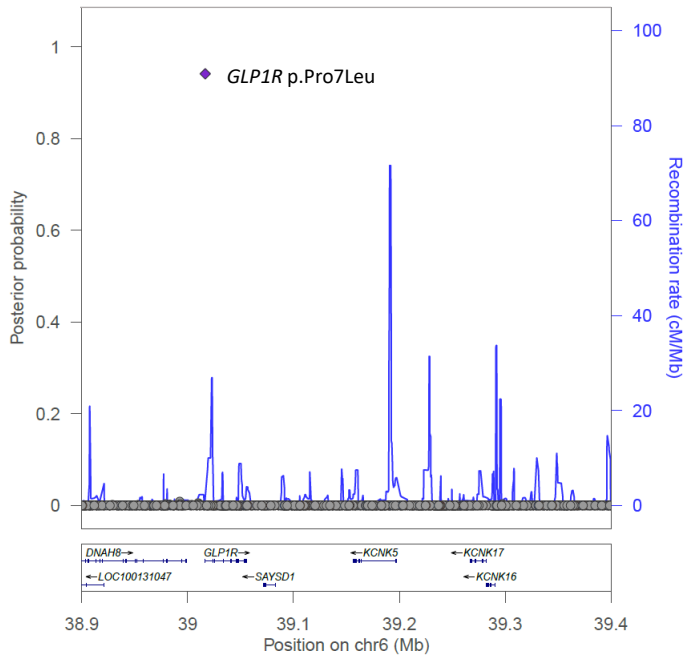


KCNJ11-ABCC8 locus

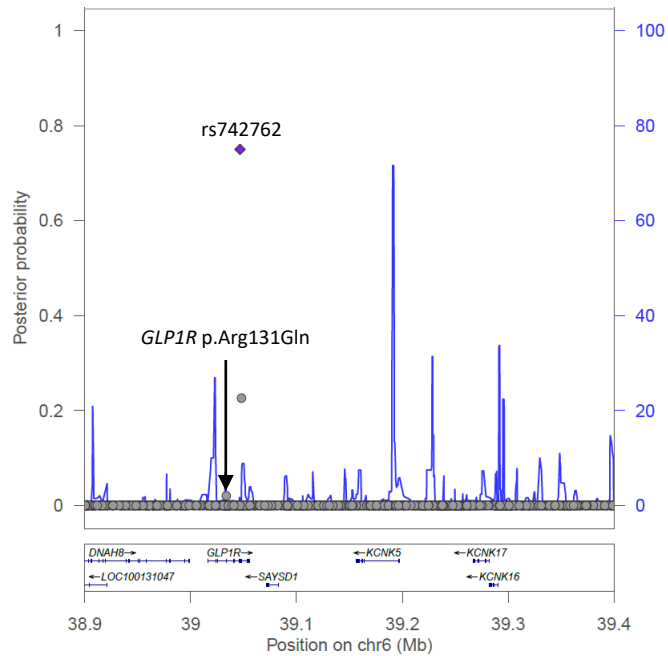


Supplementary Figure 9. The role of coding variation in driving T2D association signals at the *ZFAND3-KCNK16-GLP1R* locus from multi-ancestry meta-regression (MR-MEGA) of up to 180,834 cases and 1,159,055 controls (effective sample size 492,191). Each point represents a SNV passing quality control in the meta-regression, plotted with their annotation-informed posterior probability of driving T2D association as a function of genomic position (NCBI build 37). In each plot, the index variant is represented by the purple diamond. Gene annotations are taken from the University of California Santa Cruz genome browser. Recombination rates are estimated from the Phase II HapMap. (a) Association signal indexed by rs2281342 is driven by novel high-confidence missense variant *GLP1R* p.Pro7Leu (rs10305420). (b) The 99% credible set for the association signal indexed by rs742762, a non-coding SNV, includes *GLP1R* p.Arg131Gln (rs3765467). (c) The 99% credible set for the signal indexed by rs3734618 includes three missense variants that together account for 61.4% of the posterior probability of driving the association: *KCNK17* p.Ser21Gly (rs10947804, $\pi=39.2\%$); *KCNK16* p.Pro254His (rs11756091, $\pi=14.8\%$); and *KCNK16* p.Ala277Glu (rs1535500, $\pi=13.7\%$).

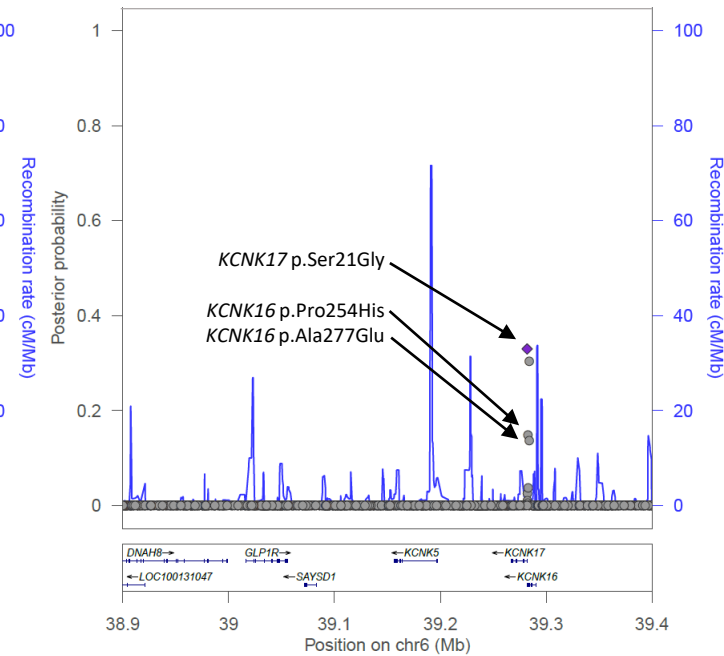
(a) Index SNV rs2281342



(b) Index SNV rs742762



(c) Index SNV rs3734618



Supplementary Note Table 1. Comparison of sample size, distribution of ancestry groups and analytical strategies utilised by four studies that incorporate GWAS from the DIAMANTE Consortium.

Study	Sample size cases/control	Ancestry-specific sample size: cases/controls (% effective sample size)				Genome-wide significance	Meta-analysis strategy	Correction for residual population structure	Variants interrogated
		AFR	ASN	EUR	HIS				
DIAMANTE European ancestry-specific (Mahajan et al. 2018)	74,124/824,006			74,124/824,006 (100%)		$p < 5 \times 10^{-8}$	Fixed-effects (inverse-variance weighted)	Double genomic control	HRC reference panel
DIAMANTE East Asian ancestry-specific (Spracklen et al. 2020)	77,418/356,122		77,418/356,122 (100%)			$p < 5 \times 10^{-8}$	Fixed-effects (inverse-variance weighted)	Double genomic control (LDSC intercept after meta-analysis)	1000G reference panel
MVP (Vujkovic et al. 2020)	228,499/1,178,783	24,646/31,446 (7.5%)	46,511/169,776 (19.8%)	148,726/965,732 (70.0%)	8,616/11,829 (2.7%)	$p < 5 \times 10^{-8}$	Fixed-effects (inverse-variance weighted)	None	1000G or HRC reference panels, MAF >1% per ancestry group
DIAMANTE multi-ancestry	180,834/1,159,005	15,487/23,709 (6.6%)	72,808/260,107 (36.7%)	80,154/853,816 (51.1%)	12,385/21,423 (5.6%)	$p < 5 \times 10^{-9}$	Meta-regression (ancestry correlated heterogeneity)	Double genomic control	Overlap of 1000G and HRC reference panels, MAF >0.5% in at least one ancestry group

AFR: African ancestry. ASN: South and East Asian ancestry. EUR: European ancestry. HIS: Hispanic ancestry. LDSC: LD Score regression. HRC: Haplotype Reference Consortium. 1000G: 1000 Genomes Project.

Supplementary Note Table 2. Overlap of loci reported at genome-wide significance ($p < 5 \times 10^{-8}$) by four studies that incorporate GWAS from the DIAMANTE Consortium: details of sample sizes and analytical approaches summarised in Supplementary Note Table 1.

Locus	Chr	Interval (bp, b37)	DIAMANTE European ancestry			DIAMANTE East Asian ancestry			MVP trans-ancestry/ancestry-specific			DIAMANTE multi-ancestry		
			Lead SNP	Position	p-value	Lead SNP	Position	p-value	Lead SNP	Position	p-value	Lead SNP	Position	p-value
<i>PHF13</i>	1	6,172,729-7,172,729							rs11583755	6,672,729	1.3E-19			
<i>MTOR</i>	1	10,817,932-11,817,932							rs7554251	11,317,932	2.9E-11			
<i>PLEKHM2</i>	1	15,550,470-16,550,470							rs12746673	16,050,470	3.3E-10			
<i>VWA5B1, LINC01141</i>	1	20,188,352-21,229,451				rs60573766	20,688,352	4.3E-10	rs10916780	20,707,153	1.6E-13	rs10916784	20,729,451	1.2E-11
<i>C1orf172, TRIM63</i>	1	25,896,065-27,784,913							rs9438610	26,396,065	6.7E-10			
<i>YTHDF2</i>	1	28,560,898-29,560,898							rs3753693	29,060,898	1.1E-10			
	1	32,696,120-33,696,120							rs59020573	33,196,120	1.3E-8			
<i>EVA1B</i>	1	36,289,546-37,289,546							rs12116935	36,789,546	1.5E-8			
<i>MACF1</i>	1	39,355,177-40,535,928	rs3768321	40,035,928	1.3E-26	rs371894931	39,942,242	2.7E-11	rs61779284	39,855,177	1.4E-48	rs3768301	39,870,793	6.2E-31
<i>MAST2</i>	1	45,744,900-46,858,862				rs562138031	46,244,900	4.0E-12				rs34444543	46,358,862	5.5E-13
<i>FAF1</i>	1	50,691,935-51,756,091	rs58432198	51,256,091	1.8E-10	rs11205766	51,191,935	7.5E-15	rs79090772	51,209,148	1.3E-28	rs12073283	51,219,188	6.7E-18
<i>PATJ, INADL</i>	1	62,079,891-63,079,891	rs12140153	62,579,891	1.2E-8				rs12140153	62,579,891	5.0E-8	rs12140153	62,579,891	1.4E-8
<i>PGM1</i>	1	63,607,284-64,614,429				rs2269245	64,107,893	5.4E-10	rs2269247	64,107,284	3.4E-13	rs11576729	64,114,429	2.5E-17
<i>LEPR</i>	1	65,489,878-66,489,878							rs10889560	65,989,878	2.1E-9			
<i>SGIP1</i>	1	66,510,654-67,510,654							rs4655617	67,010,654	1.7E-10			
<i>NEGR1</i>	1	72,251,552-73,251,552							rs2613499	72,751,552	1.0E-11			
	1	91,548,779-92,548,779							rs4658234	92,048,779	9.1E-9			
<i>RP11-147C23.1</i>	1	95,904,462-96,904,462							rs10159026	96,404,462	2.0E-9			
<i>FAM212B-AS1, ST7L</i>	1	111,789,983-113,606,633							rs197374	112,289,983	7.8E-9	rs12137269	113,106,633	6.1E-9
<i>DENND2C</i>	1	114,644,899-115,644,899	rs184660829	115,144,899	2.5E-8									
<i>PTGFRN, FAM46C</i>	1	117,032,790-118,669,463	rs1127215	117,532,790	2.3E-13				rs1127215	117,532,790	1.8E-26	rs1127215	117,532,790	3.9E-17
<i>NOTCH2</i>	1	119,955,586-121,026,982	rs1493694	120,526,982	2.1E-16				rs2453051	120,499,573	1.1E-23	rs835576	120,455,586	2.8E-17
<i>CHD1L</i>	1	146,214,427-147,621,000							rs79489938 ^a	147,121,000	4.8E-8	rs11588753	146,714,427	4.8E-8
<i>SV2A</i>	1	149,391,028-150,391,028							rs72692804	149,891,028	2.9E-10			
<i>FAM63A, BNIPL</i>	1	150,517,991-151,517,991	rs145904381	151,017,991	2.2E-8				rs145904381	151,017,991	4.0E-13			
<i>ATP8B2, PKLR</i>	1	153,824,384-155,769,776							rs3020781	155,269,776	7.3E-12			
<i>DNM3</i>	1	171,868,310-172,868,310							rs7546252	172,368,310	2.0E-10			
<i>SEC16B</i>	1	177,378,933-178,389,025	rs539515	177,889,025	1.2E-10	rs532504	177,878,933	7.4E-12	rs539515	177,889,025	5.4E-15	rs539515	177,889,025	4.6E-20
	1	178,748,952-179,748,952							rs2816177	179,248,952	6.8E-9			
<i>LAMC1</i>	1	182,504,334-183,504,334							rs4129858	183,004,334	4.4E-9			
<i>TSEN15</i>	1	183,514,593-184,535,116				rs1327123	184,014,593	7.0E-9	rs1327123	184,014,593	6.8E-10	rs1952256	184,035,116	2.6E-8
<i>ZNF281</i>	1	199,697,538-200,916,099							rs12128213	200,197,538	1.5E-9	rs10919928	200,416,099	4.6E-9
<i>IPO9</i>	1	201,349,926-202,349,926							rs41304257	201,849,926	8.5E-12			
<i>CNTN2, MDM4, DSTYK, SRGAP2</i>	1	203,974,581-207,121,028	rs12048743	205,114,873	4.4E-9	rs201297151	204,474,581	3.4E-8	rs61817176	206,621,028	3.6E-11	rs6689629	204,539,291	2.4E-10
<i>PROX1</i>	1	213,655,398-214,659,256	rs340874	214,159,256	5.6E-26	rs12403994	214,155,398	6.1E-12	rs340874	214,159,256	6.5E-45	rs340874	214,159,256	2.5E-33
<i>LYPLAL1</i>	1	219,248,818-220,248,818	rs2820446	219,748,818	3.7E-16				rs2820446	219,748,818	2.3E-22	rs2820446	219,748,818	2.7E-18
<i>ABCB10, NUP133</i>	1	229,142,499-230,172,955	rs348330	229,672,955	3.9E-14	rs238763	229,642,499	5.0E-11	rs348330	229,672,955	2.8E-20	rs348330	229,672,955	3.7E-18
<i>GNG4, TBCE</i>	1	235,042,023-236,190,800	rs291367	235,690,800	6.1E-10				rs10737818	235,542,023	2.0E-11			
<i>TMEM18</i>	2	0-1,153,874	rs62107261	422,144	1.8E-11	rs10634531	632,789	2.4E-17	rs10188334	653,874	2.5E-24	rs6548240	636,929	2.2E-25
<i>FAM49A, AC142119.1</i>	2	15,738,001-17,074,669	rs11680058	16,574,669	1.3E-8				rs28758542	16,238,001	1.7E-9	rs11680058	16,574,669	1.6E-8
	2	18,207,873-19,207,873							rs11096542	18,707,873	3.0E-9			
<i>DTNB, KIF3C</i>	2	25,033,568-26,692,802	rs17802463	25,643,221	3.5E-8				rs34845373	25,635,771	5.3E-12	rs55928417	25,533,568	4.0E-11

GCKR	2	27,230,940-28,230,940	rs1260326	27,730,940	1.3E-24	rs1260326	27,730,940	1.0E-21	rs1260326	27,730,940	2.2E-57	rs1260326	27,730,940	4.6E-38
HEATR5B	2	36,704,168-37,704,168							rs77424687	37,204,168	2.9E-9			
THADA	2	43,111,883-44,198,028	rs80147536	43,698,028	2.7E-30				rs76675804	43,611,883	1.3E-59	rs13414140	43,671,176	5.2E-31
SIX3, SIX2	2	44,692,080-45,692,080				rs12712928	45,192,080	1.8E-14				rs12712928	45,192,080	2.4E-14
EML6	2	54,657,914-55,657,914							rs5010712	55,157,914	2.9E-9			
BNIP1, LINC01122	2	58,461,136-59,807,725	rs10193538	58,981,064	1.7E-8				rs12986742	58,975,143	5.5E-21	rs17049712	58,961,136	2.3E-9
BCL11A, ACO07381.2	2	60,083,665-61,086,707	rs243024	60,583,665	4.4E-20	rs243018	60,586,707	1.5E-15	rs243018	60,586,707	1.2E-39	rs243018	60,586,707	6.7E-35
CEP68	2	64,779,414-66,166,674	rs2249105	65,287,896	1.2E-15				rs2723065	65,279,414	7.2E-28	rs6752053	65,666,674	4.1E-24
ETAA1	2	67,122,243-68,122,243							rs4671799	67,622,243	5.3E-11			
KDM3A	2	86,207,504-87,207,504							rs4832290	86,707,504	3.9E-8			
AFF3	2	100,098,726-101,098,726							rs34506349	100,598,726	1.0E-8			
	2	104,665,674-105,665,674							rs10469860	105,165,674	3.8E-8			
TMEM87B, LOC541471, BCL2L11	2	111,387,754-113,323,114							rs113135335	111,887,754	2.1E-13	rs1345203	112,253,851	2.9E-8
DDX18	2	117,571,061-118,571,061	rs562386202	118,071,061	4.2E-8									
SCTR	2	119,731,070-120,731,070				rs3731600	120,231,070	6.9E-9						
GLI2	2	120,817,747-121,847,612	rs11688682	121,347,612	1.4E-14				rs9308614	121,337,196	4.5E-23	rs11677557	121,317,747	6.5E-15
TEX41	2	145,226,656-146,850,724							rs6716394	146,350,724	1.5E-13			
PABPC1P2	2	147,361,633-148,361,633	rs35999103	147,861,633	8.3E-9									
EPC2	2	148,928,856-150,068,261				rs200576292	149,568,261	1.3E-9	rs66877183	149,428,856	3.0E-8			
	2	151,698,598-152,698,598							rs3845843	152,198,598	2.4E-12			
ACVR1C, CYTIP	2	157,839,550-158,949,081	rs13426680	158,339,550	6.4E-10				rs149447188	158,449,081	2.0E-10	rs7594480	158,390,468	4.0E-12
RBMS1	2	160,635,544-161,833,872	rs3772071	161,135,544	1.6E-11				rs6710938	161,333,872	1.6E-13	rs1020731	161,144,055	1.2E-9
KCNH7	2	163,123,932-164,149,480							rs305686	163,623,932	2.3E-8	rs12614955	163,649,480	8.6E-10
GRB14, COBLL1	2	164,881,518-166,013,091	rs10195252	165,513,091	1.6E-20	rs75536691	165,381,518	1.2E-15	rs10184004	165,508,389	4.4E-54	rs10184004	165,508,389	5.2E-34
GALNT3	2	166,110,827-167,111,006							rs13406280	166,610,827	5.7E-12	rs62174818	166,611,006	3.9E-8
HAT1	2	172,296,774-173,296,774							rs62182438	172,796,774	1.1E-8			
SP9	2	174,697,545-175,697,545							rs12992995	175,197,545	1.1E-8			
TTN	2	179,150,954-180,150,954							rs6715901	179,650,954	1.2E-8			
SCHLAP1	2	181,070,507-182,118,654							rs6741676	181,618,654	6.5E-15	rs12479357	181,570,507	1.8E-8
	2	196,452,010-197,452,010							rs6712905	196,952,010	2.1E-11			
RP11-68606.2	2	202,735,139-203,735,139							rs6714523	203,235,139	6.3E-11			
ACO16903.2	2	204,875,909-205,875,909							rs4482463	205,375,909	1.8E-8			
PLEKHM3	2	208,370,017-209,370,017							rs34329895	208,870,017	1.4E-10			
ERBB4	2	211,774,937-212,774,937							rs3828242	212,274,937	6.5E-9			
ACO79610.1, IKZF2	2	213,187,103-214,329,721				rs75179644	213,687,103	5.4E-10	rs4673712	213,829,721	1.7E-9	rs16849467	213,818,731	4.8E-9
PNKD, CRYBA2	2	218,668,432-220,359,171							rs113414093	219,859,171	1.7E-8			
IRS1	2	226,600,490-227,605,921	rs2972144	227,101,411	7.9E-46				rs2943650	227,105,921	1.9E-81	rs2943648	227,100,490	3.6E-52
SPHKAP	2	228,471,884-229,471,884							rs13415288	228,971,884	9.2E-11			
ATG16L1, DGKD	2	233,691,103-234,803,281				rs117809958	234,191,103	2.0E-15	rs838720	234,303,281	4.3E-18	rs117809958	234,191,103	5.6E-12
	3	3,149,850-4,149,850							rs9842137 ^a	3,649,850	9.3E-9			
SETD5	3	9,014,016-10,014,016							rs3872707	9,514,016	9.8E-12			
PPARG	3	11,829,783-12,885,357	rs11709077	12,336,507	1.6E-27	rs3963364	12,385,357	3.5E-11	rs17036160	12,329,783	1.5E-53	rs17036160	12,329,783	2.9E-38
ANKRD28	3	15,206,124-16,206,124							rs924753	15,706,124	1.2E-11			
UBE2E2	3	22,758,614-23,957,080	rs35352848	23,455,582	9.5E-20	rs11926494	23,258,614	2.7E-37	rs13094957	23,457,080	2.4E-42	rs13094957	23,457,080	5.2E-48
LINC00693	3	28,231,810-29,231,810							rs9869477	28,731,810	4.1E-9			
	3	35,170,150-36,170,150							rs1470560	35,670,150	1.2E-9			
	3	36,370,230-37,370,230							rs11129735 ^a	36,870,230	1.5E-8			
KIF9, SMARCC1	3	46,425,539-48,193,664	rs11926707	46,925,539	1.5E-8				rs62262091	47,693,664	1.5E-10			

RBM6	3	49,480,596-50,674,197	rs4688760	49,980,596	4.5E-10				rs6792892	49,995,518	1.4E-14	rs2624847	50,174,197	6.2E-11
RFT1	3	52,625,429-53,627,677	rs2581787	53,127,677	3.0E-8				rs62255926	53,125,429	4.2E-8	rs2581787	53,127,677	2.0E-8
CACNA2D3	3	54,328,827-55,328,827	rs76263492	54,828,827	6.3E-9				rs76263492	54,828,827	3.2E-9	rs76263492	54,828,827	8.7E-10
PXK	3	57,838,809-58,838,809										rs12629058	58,338,809	3.5E-11
PSMD6, ATXN7, ADAMTS9	3	63,384,800-65,203,394	rs9860730	64,701,146	7.4E-15	rs67114627	63,904,715	9.1E-32	rs13434089	63,948,566	4.5E-31	rs704360	63,884,800	8.0E-33
	3	70,020,917-71,020,917							rs12494424	70,520,917	2.3E-8			
	3	71,148,868-72,148,868							rs853866	71,648,868	2.7E-8			
SHQ1	3	72,303,590-73,365,183	rs13085136	72,865,183	1.4E-8				rs9814945	72,803,590	3.3E-11			
ROBO2	3	77,171,721-78,171,721	rs2272163	77,671,721	1.2E-8				rs2272163	77,671,721	3.6E-9			
	3	86,256,871-87,256,871							rs6549112	86,756,871	1.6E-11			
RP11-159G9.5	3	87,630,136-88,630,136							rs73146095	88,130,136	7.9E-9			
	3	89,486,280-90,486,280							rs11716527 ^a	89,986,280	6.7E-9			
	3	93,481,060-94,481,060							rs978444	93,981,060	3.5E-9			
SIDT1	3	112,788,430-113,788,430							rs11929640	113,288,430	8.5E-9			
ZBTB20	3	114,460,798-115,563,672				rs6806156	114,968,018	1.6E-11	rs7645613	115,063,672	1.5E-12	rs1459513	114,960,798	9.6E-15
CASR	3	121,461,461-122,465,199				rs9859381	121,965,199	2.9E-9	rs13059382	121,961,461	1.4E-9			
ADCY5	3	122,565,778-123,674,832	rs11708067	123,065,778	1.3E-31	rs60054445	123,174,832	5.6E-12	rs11708067	123,065,778	1.6E-57	rs11708067	123,065,778	3.4E-46
SLC12A8	3	124,421,457-125,426,637	rs649961	124,926,637	1.3E-9	rs12497133	124,921,920	1.1E-8	rs9873519	124,921,457	2.5E-22	rs9873519	124,921,457	6.5E-14
TMCC1, PLXND1	3	128,079,324-129,833,182	rs9828772	129,333,182	4.2E-8				rs2255703	129,293,256	3.7E-11			
CPNE4	3	131,250,844-132,250,844							rs9857204	131,750,844	2.2E-10			
STAG1	3	135,569,472-136,569,472							rs667920	136,069,472	5.8E-13			
	3	137,555,136-138,555,136							rs6766859	138,055,136	8.1E-12			
ZBTB38	3	140,601,839-141,601,839							rs56243018	141,101,839	8.9E-16			
TSC22D2, TM4SF4	3	148,721,563-150,566,540	rs62271373	150,066,540	1.0E-9				rs28712435	149,221,563	6.1E-11	rs62271373	150,066,540	2.6E-8
MBNL1	3	151,584,243-152,932,042	rs111729685	152,086,533	3.6E-9	rs1850421	152,382,352	1.4E-9	rs7633673	152,084,243	8.2E-16	rs9877505	152,432,042	1.4E-13
LEKR1, CCNL1	3	156,295,525-157,295,525										rs9854955	156,795,525	1.4E-8
TRIM59	3	159,653,305-160,653,305							rs7629	160,153,305	8.2E-14			
EGFEM1P	3	167,718,841-168,726,052	rs7629630	168,218,841	2.2E-8				rs13099581	168,226,052	1.3E-10			
SLC2A2	3	170,143,788-171,233,076	rs9873618	170,733,076	8.5E-21	rs201018682	170,643,788	1.0E-11	rs8192675	170,724,883	3.9E-27	rs8192675	170,724,883	3.5E-28
NLGN1	3	172,607,443-174,210,695							rs247975	173,107,443	2.0E-11			
CCDC39, FXR1	3	180,045,384-181,045,384										rs4854992	180,545,384	9.3E-9
ABCC5	3	183,238,460-184,238,460	rs2872246	183,738,460	1.8E-8				rs2872246	183,738,460	2.6E-11			
IGF2BP2	3	184,382,015-186,034,482	rs6780171	185,503,456	2.5E-58	rs13092876	185,495,320	1.9E-66	rs9859406	185,534,482	2.0E-169	rs7633675	185,510,613	5.8E-131
ST6GAL1	3	186,149,931-187,165,645	rs3887925	186,665,645	1.4E-17	rs11332772	186,649,931	1.1E-10	rs3887925	186,665,645	1.1E-25	rs3887925	186,665,645	3.8E-22
BCL6, LPP	3	187,198,333-188,241,842	rs4686471	187,740,899	3.1E-20	rs13086331	187,698,333	7.3E-9	rs6777684	187,741,842	1.5E-35	rs4686471	187,740,899	2.9E-21
TFRC	3	195,325,077-196,331,237				rs9866168	195,830,310	1.5E-9	rs9872347	195,831,237	5.4E-14	rs74289356	195,825,077	6.8E-12
CTBP1, PCGF3, MAEA	4	220,681-2,284,605	rs56337234	1,784,403	1.4E-17	rs7656416	1,254,535	9.0E-42	rs730831	1,240,299	4.8E-34	rs730831	1,240,299	7.6E-41
HTT	4	2,741,845-3,741,845	rs362307	3,241,845	1.1E-9				rs362307	3,241,845	3.3E-8			
WFS1	4	5,793,237-6,806,763	rs10937721	6,306,763	1.6E-40	rs147834269	6,303,731	9.1E-12	rs10937721	6,306,763	7.2E-70	rs9998835	6,293,237	3.1E-51
LCORL	4	17,292,869-18,547,401	rs12640250	17,792,869	4.5E-8				rs2169033	18,044,357	1.4E-15	rs6855926	18,047,401	8.4E-12
SLIT2	4	19,765,535-20,765,535							rs7664347	20,265,535	9.5E-9			
GNPDA2	4	44,003,503-45,686,139	rs10938398	45,186,139	4.9E-12	rs10938398	45,186,139	3.8E-10	rs10938398	45,186,139	1.5E-30	rs13130484	45,175,691	1.3E-15
CWH43	4	48,567,323-49,567,323							rs2605281	49,067,323	6.9E-12			
USP46	4	52,298,624-53,318,664	rs2102278	52,818,664	4.5E-8				rs1996617	52,798,624	6.5E-13			
MOB1B	4	71,335,822-72,344,118				rs28599782	71,844,118	4.6E-16	rs7674402	71,835,822	4.0E-16	rs7674402	71,835,822	1.2E-13
ART3, SHROOM3	4	75,996,817-78,033,939							rs6835992	76,496,817	5.6E-11			
SCD5	4	83,078,271-84,087,562	rs79920718	83,578,271	5.7E-10				rs993380	83,584,496	3.7E-12	rs10471048	83,587,562	2.3E-11
NKX6-1, CDS1, RP11-42A4.1	4	84,797,954-85,839,618				rs117624659	85,339,618	2.0E-16	rs117233795	85,297,954	1.4E-12	rs117624659	85,339,618	3.1E-14

FAM13A	4	89,213,121-90,240,894	rs1903002	89,740,894	3.0E-8				rs9991328	89,713,121	2.4E-9			
	4	90,743,865-91,743,865							rs7656001 ^a	91,243,865	1.2E-8			
UNC5C, RP11-363G15.2, SMARCAD1	4	94,591,911-96,614,385	rs6821438	95,091,911	5.4E-11				rs3755879	96,114,385	5.6E-11	rs6821438	95,091,911	2.5E-9
PPP3CA	4	101,635,363-102,635,363										rs2659518	102,135,363	4.5E-9
SLC9B1	4	102,688,709-104,640,848	rs1580278	104,140,848	2.9E-10				rs7659468	103,895,317	1.4E-14	rs223423	103,725,894	1.6E-9
TET2	4	105,548,291-106,548,291							rs17035289	106,048,291	1.5E-13	rs17035289	106,048,291	1.3E-12
PRDM5	4	121,265,788-122,265,788							rs4833687	121,765,788	2.8E-9			
LARP1B	4	128,524,273-129,524,273							rs4834232	129,024,273	9.4E-9			
RP11-422J15.1	4	130,286,346-131,286,346							rs2952858	130,786,346	1.1E-8			
PABPC4L	4	136,583,193-137,583,193	rs1296328	137,083,193	4.3E-8									
MAML3	4	140,406,390-141,406,390							rs12505942	140,906,390	1.8E-12			
HHIP	4	145,112,552-146,112,552							rs12511407	145,612,552	2.1E-10			
TMEM154	4	153,013,369-154,020,475	rs7669833	153,513,369	1.8E-14	rs10011838	153,520,279	1.4E-27	rs6813195	153,520,475	7.7E-31	rs6813195	153,520,475	3.0E-36
GUCY1B3, PDGFC	4	156,197,784-158,225,916	rs28819812	157,652,753	2.7E-8				rs28819812	157,652,753	2.6E-15	rs1425482	157,725,916	2.3E-11
SORBS2, ACSL1	4	185,214,289-187,080,062	rs58730668	185,717,759	1.0E-13				rs55691245	185,716,100	1.8E-18	rs1996546	185,714,289	1.6E-13
ANKH	5	14,251,305-15,268,092	rs146886108	14,751,305	8.7E-16	rs6885132	14,768,092	2.6E-9	rs146886108	14,751,305	1.7E-31	rs6885132	14,768,092	9.5E-23
RANBP3L	5	35,584,426-36,757,018				rs16902871	36,257,018	3.3E-9	rs114136102	36,084,426	2.8E-8			
MRPS30	5	44,144,006-45,182,589	rs6884702	44,682,589	5.8E-9				rs4479849	44,644,006	2.6E-11	rs6884702	44,682,589	6.0E-12
	5	45,704,748-46,704,748							rs8188241	46,204,748	1.5E-8			
PARP8	5	49,579,603-50,645,266				rs74334916	50,079,603	4.3E-8	rs152839	50,145,266	1.1E-12			
ITGA1	5	51,251,574-52,600,489	rs3811978	52,100,489	4.2E-10	rs12109081	51,751,574	1.1E-8	rs12187734	51,763,665	2.2E-13	rs17261179	51,791,225	6.9E-11
ARL15	5	52,771,420-53,797,591	rs702634	53,271,420	2.1E-13				rs4865796	53,272,664	5.4E-30	rs7736354	53,297,591	1.7E-20
ANKRD55, SLC38A9, ACO22431.2	5	54,486,775-56,310,305	rs465002	55,808,475	3.8E-23	rs256904	55,810,305	3.6E-29	rs464605	55,807,370	8.0E-66	rs465002	55,808,475	1.5E-51
RAB3C	5	57,632,702-58,632,702							rs2662390	58,132,702	1.0E-8			
PIK3R1	5	67,214,246-68,216,793							rs4976033	67,714,246	1.0E-10	rs57634870	67,716,793	6.0E-12
POCS, HMGR, ANKDD1B	5	74,074,984-75,503,678	rs2307111	75,003,678	3.3E-16	rs2126736	74,574,984	1.8E-8	rs34341	74,934,009	4.9E-25	rs2307111	75,003,678	1.3E-18
ZBED3	5	75,924,949-76,935,004	rs4457053	76,424,949	1.4E-17				rs7732130	76,435,004	2.2E-32	rs7732130	76,435,004	6.0E-22
DMGDH, JMY	5	77,930,607-79,046,293	rs1316776	78,430,607	3.5E-12				rs2591392	78,546,293	1.9E-15	rs10052346	78,472,599	5.7E-14
RASA1	5	86,018,243-88,197,533	rs7719891	86,577,352	2.9E-8				rs6870983	87,697,533	6.3E-9	rs11953892	86,518,243	1.9E-8
PCSK1, CTD-2337A12.1	5	95,348,503-96,350,250				rs261982	95,843,763	3.1E-9	rs261967	95,850,250	8.1E-10			
SLCO6A1, PAM, CTC-503K11.2	5	100,732,944-103,473,337	rs115505614	102,422,968	1.7E-29				rs75432112	102,586,407	6.4E-37	rs115505614	102,422,968	5.9E-29
CEP120	5	122,150,885-123,204,342							rs144052331	122,650,885	4.0E-13	rs4267865	122,704,342	9.7E-13
JADE2, PHF15	5	133,361,663-134,364,599	rs329122	133,864,599	9.2E-9	rs329122	133,864,599	2.2E-8	rs329118	133,861,663	2.0E-13	rs329122	133,864,599	6.0E-16
WNT8A	5	136,931,501-137,931,501							rs217256	137,431,501	1.4E-8			
CTB-12O2.1	5	150,824,600-151,824,600							rs302395	151,324,600	1.5E-8			
EBF1	5	157,525,983-158,529,734							rs1650505	158,029,734	1.7E-19	rs748510	158,025,983	3.6E-8
RANBP17	5	170,183,134-171,183,134							rs2913873	170,683,134	1.5E-8			
NSD1, FGFR4	5	176,013,896-177,179,407				rs3135911	176,513,896	1.5E-12	rs4343858	176,679,407	5.4E-11	rs244708	176,589,585	3.8E-9
MGAT1	5	179,726,516-180,726,516							rs6885157	180,226,516	4.7E-8			
SSR1, RREB1	6	6,731,843-7,731,843	rs9379084	7,231,843	2.3E-20	rs9379084	7,231,843	2.2E-14	rs9379084	7,231,843	5.3E-31	rs9379084	7,231,843	6.1E-30
JARID2	6	14,975,051-15,999,419							rs727734	15,475,051	6.7E-11	rs7769291	15,499,419	3.0E-8
CDKAL1	6	19,251,516-21,188,121	rs7756992	20,679,709	3.0E-87	rs9350271	20,683,164	5.0E-183	rs10440833	20,688,121	4.5E-215	rs9348441	20,680,678	6.2E-235
HIST1H4E	6	25,711,146-26,711,146							rs9358912	26,211,146	2.3E-8			
MICF	6	28,426,220-30,316,421				rs6915823	30,073,430	1.8E-10	rs9257408	28,926,220	2.0E-10			
MHC region	6	30,526,236-34,736,973	rs601945	32,573,415	2.7E-21	rs76541615	31,026,236	1.1E-17	rs3130931	31,134,888	2.3E-32	rs879882	31,139,452	5.0E-26
ZNF76	6	34,759,397-35,759,397							rs33959228	35,259,397	3.4E-9			
RP1-90K10.4	6	36,411,274-37,411,274							rs72846863	36,911,274	4.6E-8			
ZFAND3, KCNK16, KCNK17, GLP1R	6	38,546,644-39,782,371				rs742762	39,046,644	1.8E-22	rs34247110	39,282,371	2.1E-15	rs34247110	39,282,371	3.4E-21

USP49, LRFN2	6	39,909,243-42,364,441	rs34298980	40,409,243	1.2E-9				rs34298980	40,409,243	1.2E-10	rs34298980	40,409,243	4.1E-9	
VEGFA	6	43,313,711-44,314,190	rs6458354	43,814,190	3.7E-13				rs9472139	43,813,711	1.4E-17	rs6458354	43,814,190	1.3E-13	
SUPT3H	6	44,375,762-45,375,762							rs538801	44,875,762	4.7E-8				
TFAP2B	6	50,287,459-51,913,013	rs3798519	50,788,778	1.1E-12	rs62405419	50,787,459	3.8E-9	rs3798519	50,788,778	5.6E-24	rs3798519	50,788,778	2.4E-22	
RP3-523E19.2	6	53,289,830-54,289,830							rs9370243	53,789,830	1.0E-10				
	6	63,663,807-64,663,807							rs9449295	64,163,807	4.0E-8				
SLC25A51P1	6	66,887,490-67,887,490	rs555402748	67,387,490	4.6E-8										
BEND3	6	106,931,688-107,945,266	rs4946812	107,431,688	1.0E-8				rs7752666	107,445,266	1.7E-12	rs1665901	107,433,400	6.1E-12	
REV3L	6	111,238,793-112,238,793							rs55812705	111,738,793	1.3E-9				
NUS1	6	117,496,631-118,511,723					rs80196932	117,996,631	7.6E-13	rs80196932	117,996,631	9.7E-20	rs72951506	118,011,723	9.9E-11
CENPW, SOGA3, RP11-624M8.1	6	125,561,502-127,916,930	rs11759026	126,792,095	1.3E-18	rs4273712	126,964,510	2.6E-12	rs11759026	126,792,095	1.2E-24	rs11759026	126,792,095	1.0E-36	
	6	129,765,266-130,765,266							rs35164294	130,265,266	1.4E-8				
MED23, ENPP3	6	131,426,334-132,454,797					rs7739842	131,954,797	1.6E-11	rs2608953	131,926,334	3.3E-12	rs7739842	131,954,797	1.8E-13
SLC35D3, RPL35AP3	6	136,791,281-137,800,960	rs1573090	137,302,159	8.4E-15	rs35389258	137,294,771	9.5E-14	rs2876354	137,295,352	1.4E-34	rs6937795	137,291,281	5.3E-15	
NHSL1, REPS1	6	138,355,975-140,337,128					rs9376382	139,205,386	1.5E-8	rs11155073	139,837,128	1.4E-12	rs9376353	138,855,975	2.7E-10
HIVEP2	6	142,556,556-143,558,692					rs9390022	143,056,556	6.4E-9	rs9390022	143,056,556	5.0E-14	rs6570526	143,058,692	8.2E-10
RGS17	6	152,938,573-153,940,770							rs7758002	153,440,770	4.6E-16	rs6932473	153,438,573	5.5E-13	
SLC22A3	6	160,270,312-161,270,918	rs474513	160,770,312	1.0E-9				rs501470	160,770,918	1.6E-17	rs539298	160,770,360	1.4E-15	
QKI, RP1-230L10.1	6	163,633,001-164,633,001	rs4709746	164,133,001	5.0E-9				rs4709746	164,133,001	1.5E-21	rs4709746	164,133,001	7.8E-10	
	7	1,372,921-3,260,750							rs4721089*	1,872,921	7.9E-10				
FOKK1	7	4,183,572-5,191,060							rs62452060	4,683,572	1.7E-9	rs28411900	4,691,060	2.7E-8	
TMEM106B	7	11,769,593-12,769,593							rs13237518	12,269,593	2.0E-13				
EUM1	7	13,386,654-14,387,008					rs7787720	13,886,654	2.3E-15	rs7787720	13,886,654	3.8E-13	rs12154701	13,887,008	1.4E-11
DGKB	7	14,398,282-16,426,228	rs10228066	15,063,569	1.9E-25	rs17168486	14,898,282	8.2E-22	rs10228796	15,064,190	2.0E-71	rs2215383	15,062,983	2.1E-44	
HDAC9	7	17,831,915-18,831,915							rs583769	18,331,915	7.8E-13				
IGF2BP3	7	23,012,896-24,384,697	rs4279506	23,512,896	5.7E-9				rs2188848	23,884,697	1.2E-11				
LOC646588, NFE2L3	7	25,479,338-26,479,338										rs2391174	25,979,338	1.3E-8	
JAZF1	7	27,692,280-28,719,310	rs1708302	28,198,677	4.2E-48	rs3735567	28,219,310	3.1E-12	rs860262	28,194,397	2.9E-82	rs849133	28,192,280	2.3E-69	
CRHR2	7	30,228,452-31,228,452	rs917195	30,728,452	5.6E-11				rs917195	30,728,452	1.5E-20	rs917195	30,728,452	3.6E-16	
AOAH	7	36,242,886-37,242,886							rs6978327	36,742,886	4.8E-9				
SUGCT	7	40,316,653-41,316,653							rs17439448	40,816,653	3.7E-9				
MYL7, CCM2, GCK	7	43,674,857-45,616,468	rs878521	44,255,643	1.6E-14	rs2908279	44,174,857	8.4E-11	rs730497	44,223,721	9.3E-29	rs878521	44,255,643	2.2E-20	
	7	48,339,003-49,339,003							rs12539264	48,839,003	8.6E-11				
DDC, GRB10	7	50,077,968-51,309,085							rs73121277	50,577,968	7.4E-10	rs13236710	50,809,085	2.0E-11	
ZNF713, CICP11	7	55,302,063-56,484,953					rs565050730	55,984,953	4.4E-8	rs6972291	55,802,063	2.5E-9	rs9784904	55,835,078	1.4E-8
AUTS2	7	68,555,951-70,196,905					rs12698877	69,696,905	7.0E-22	rs6975279	69,649,683	4.1E-25	rs2533457	69,055,951	3.3E-19
GTF2I	7	73,576,493-74,576,493							rs13238568	74,076,493	1.7E-9				
MAGI2, RP5-899E9.1	7	76,547,102-78,328,991							rs12669521	77,047,102	2.8E-9				
STEAP1, ACO04969.1	7	89,252,238-90,303,634					rs62469016	89,752,238	1.5E-15	rs6956980	89,803,634	9.6E-14	rs6978118	89,800,241	2.0E-14
CALCR	7	92,607,093-93,618,736					rs2074120	93,107,093	8.4E-9	rs76369672	93,118,736	1.2E-8			
	7	99,813,420-100,813,420							rs506597	100,313,420	1.5E-10				
RASA4, FBXL13, RELN, DNAJC2	7	101,836,979-103,944,978	rs11496066	102,486,254	1.2E-8	rs75990271	102,336,979	3.2E-11	rs187653072	102,976,385	2.2E-12	rs7781557	102,481,891	8.9E-12	
LHFPL3	7	104,016,274-105,016,274							rs73184014	104,516,274	1.2E-8				
CTTNBP2	7	116,995,667-117,995,667	rs6976111	117,495,667	1.5E-8				rs6976111	117,495,667	1.5E-8				
SND1, GCC1, LEP, GRM8, PAX4	7	126,026,991-128,403,272					rs2233580	127,253,550	2.7E-132	rs17866443	127,058,953	8.9E-47	rs12669223	127,250,831	2.4E-39
KLF14	7	129,927,057-130,957,914	rs1562396	130,457,914	7.6E-17				rs3996350	130,427,057	1.2E-18	rs1562396	130,457,914	1.7E-16	
ACO09518.3	7	131,074,608-132,074,608							rs12667919	131,574,608	3.6E-8				
BRAF	7	140,022,073-141,131,823					rs71170768	140,579,350	2.2E-10	rs60251368	140,522,073	2.2E-12	rs11983228	140,631,823	5.0E-10

TRPV5	7	142,107,301-143,107,301							rs4252505	142,607,301	4.5E-9				
CUL1, CNTNAP2	7	147,158,539-149,738,823							rs1922879	147,658,539	2.1E-8				
AOC1	7	150,037,635-151,037,635	rs62492368	150,537,635	1.5E-10				rs62492368	150,537,635	1.5E-14	rs62492368	150,537,635	1.5E-10	
MNX1, UBE3C	7	156,430,550-157,524,510	rs6459733	156,930,550	3.9E-17	rs1182444	157,024,510	1.7E-12	rs6946660	156,948,648	3.3E-33	rs10085650	156,993,413	3.9E-26	
	8	3,686,731-4,686,731							rs117173251 ^a	4,186,731	2.5E-8				
MFHAS1, RP11-115J16.2, XKR6, MSRA	8	8,221,473-11,569,960	rs17689007	9,974,824	1.7E-13				rs60384372	9,974,584	4.3E-17	rs4240673	10,787,612	1.1E-11	
LONRF1, RP11-252C15.1	8	12,118,225-13,143,055							rs12056338	12,643,055	2.2E-10	rs12680692	12,618,225	6.3E-10	
SGCZ	8	13,648,990-14,648,990							rs35753840	14,148,990	9.7E-10				
ASAH1	8	17,427,609-18,427,609					rs34642578	17,927,609	1.6E-9						
LPL	8	19,330,921-20,344,415	rs10096633	19,830,921	8.7E-13				rs10096633	19,830,921	8.1E-10	rs7819706	19,844,415	4.3E-13	
BIN3	8	21,992,103-22,992,103							rs6558173	22,492,103	7.3E-10				
EBF2	8	25,371,721-26,371,721							rs11998023	25,871,721	9.0E-10				
RP11-380I10.3	8	27,595,939-28,595,939							rs11994255	28,095,939	1.8E-8				
PURG	8	30,352,826-31,363,938	rs10954772	30,863,938	2.3E-9				rs2725370	30,852,826	3.3E-12	rs2725370	30,852,826	2.9E-8	
	8	34,002,571-35,002,571							rs4463416	34,502,571	1.1E-8				
ZNF703, RP11-150O12.1, FGFR1, KCNU1	8	36,332,310-38,843,012					rs4739515	37,391,203	1.7E-11	rs13365225	36,858,483	1.6E-12	rs12680217	37,397,803	5.2E-15
ANK1, NKX6-3	8	41,008,577-42,022,991	rs13262861	41,508,577	1.8E-27	rs33981001	41,512,648	5.3E-28	rs13262861	41,508,577	8.2E-79	rs508419	41,522,991	5.7E-47	
PENK, RP11-17A4.1	8	56,996,064-57,998,704							rs3887059	57,496,064	5.1E-12	rs6651357	57,498,704	8.4E-9	
RP11-1102P16.1	8	71,907,374-72,907,374							rs10101067	72,407,374	1.3E-8				
KCNB2	8	73,003,743-74,003,743					rs349359	73,503,743	3.1E-8						
GDAP1, STAU2	8	74,068,099-75,714,398					rs149265787	75,214,398	5.7E-10	rs28792187	74,568,099	4.1E-8	rs3780012	75,147,209	8.0E-10
TP53INP1, RP11-347C18.3	8	95,460,886-96,467,372	rs10097617	95,961,626	1.1E-15	rs896852	95,960,886	6.4E-9	rs10808671	95,967,372	2.4E-21	rs13257021	95,965,695	3.3E-20	
RP11-44N17.2, CPQ	8	96,638,738-98,237,741	rs149364428	97,737,741	1.9E-12				rs546898700	97,724,430	6.0E-12				
AZIN1	8	103,376,325-104,376,325							rs2679745	103,876,325	1.9E-8				
RP11-127H5.1	8	105,162,373-106,162,373							rs112515915	105,662,373	3.2E-11				
TRHR	8	109,623,183-110,623,183	rs12680028	110,123,183	3.1E-8										
TRPS1	8	115,997,173-117,065,365							rs3802219	116,565,365	1.1E-19	rs800909	116,497,173	8.1E-12	
SLC30A8	8	117,684,783-118,685,025	rs3802177	118,185,025	6.3E-55	rs13266634	118,184,783	3.7E-67	rs13266634	118,184,783	4.2E-136	rs13266634	118,184,783	3.2E-115	
TRIB1	8	125,971,274-126,971,274					rs60089934	126,471,274	3.3E-9						
PVT1, CASC11, RP11-89M16.1	8	128,211,742-130,069,999	rs17772814	128,711,742	5.0E-10				rs1561927	129,568,078	5.0E-13	rs4733612	129,569,999	2.4E-12	
EFR3A	8	132,379,795-133,379,795					rs10505581	132,879,777	4.4E-8						
	8	135,275,546-136,275,546							rs4294149	135,775,546	2.3E-8				
BOP1, HSF1	8	145,007,304-146,044,720	rs4977213	145,507,304	4.4E-14				rs13268508	145,525,277	4.1E-20	rs3890400	145,544,720	3.3E-18	
DMRT2	9	532,567-1,533,958					rs1016565	1,032,567	2.2E-8	rs1567353	1,033,773	7.1E-12	rs1509195	1,033,958	1.3E-8
RFX3	9	2,749,708-3,749,708							rs75619936	3,249,708	2.4E-8				
GLIS3	9	3,790,085-4,791,928	rs10974438	4,291,928	1.6E-14	rs4237150	4,290,085	4.5E-27	rs4237150	4,290,085	7.0E-31	rs4237150	4,290,085	1.5E-36	
	9	7,790,816-8,790,816							rs10758950	8,290,816	7.0E-9				
NFIB	9	13,641,703-14,641,703							rs73642097	14,141,703	2.1E-9				
HAUS6	9	18,567,833-19,574,538	rs7022807	19,067,833	3.6E-10				rs12380322	19,074,538	1.6E-13	rs12380322	19,074,538	4.9E-10	
FOCAD	9	19,741,069-21,290,622	rs7867635	20,241,069	4.1E-8				rs2150999	20,790,622	5.5E-10				
CDKN2A, CDKN2B	9	21,632,878-22,634,094	rs10811660	22,134,068	6.6E-79	rs10965248	22,132,878	4.4E-164	rs10811661	22,134,094	9.6E-206	rs10811661	22,134,094	1.1E-201	
	9	22,858,495-23,858,495							rs7029718	23,358,495	1.5E-12				
LINGO2	9	27,910,683-29,589,437	rs1412234	28,410,683	2.5E-10				rs1412234	28,410,683	4.4E-21	rs1412234	28,410,683	1.5E-11	
UBAP2	9	33,574,476-34,574,476	rs12001437	34,074,476	3.7E-10				rs12001437	34,074,476	4.4E-15	rs12001437	34,074,476	5.6E-11	
GBA2	9	35,249,014-36,249,014							rs1570247	35,749,014	2.9E-10				
MTND2P8, TLV4	9	80,844,701-82,417,127	rs17791513	81,905,590	2.9E-14	rs1328412	81,917,111	6.4E-11	rs67269808	81,907,986	1.0E-20	rs13290396	81,914,978	1.4E-26	
TLE1, RP11-154D17.1	9	83,808,948-84,808,948	rs2796441	84,308,948	8.5E-24	rs2796441	84,308,948	1.4E-28	rs2796441	84,308,948	2.9E-55	rs2796441	84,308,948	8.0E-42	
	9	84,812,075-85,812,075							rs654629	85,312,075	5.2E-12				

<i>C9orf3, ZNF169, PTCH1</i>	9	96,415,002-98,778,413	rs55653563	97,001,682	3.2E-9	rs113154802	98,278,413	3.5E-8	rs10993072	96,915,002	2.9E-14	rs113154802	98,278,413	4.3E-12
<i>ABCA1</i>	9	107,097,527-108,097,527				rs201375651	107,597,527	2.6E-8						
<i>EPB41L4B</i>	9	111,438,268-112,438,268							rs10119430	111,938,268	1.8E-9			
<i>COL27A1</i>	9	116,443,357-117,443,357							rs1431819	116,943,357	2.0E-10			
<i>ASTN2</i>	9	118,752,277-119,752,277							rs1885234	119,252,277	2.7E-10			
<i>STRBP, ZBTB26</i>	9	125,189,694-127,086,563							rs10818763 ^a	125,689,694	2.4E-12	rs2416899	126,015,103	3.5E-10
<i>FIBCD1</i>	9	133,286,652-134,286,652							rs6597649	133,786,652	2.3E-9			
<i>MED27</i>	9	134,368,417-135,368,417							rs9411425	134,868,417	4.8E-9			
<i>ABO, LINC00094</i>	9	135,649,229-137,390,704	rs505922	136,149,229	5.4E-12	rs529565	136,149,500	1.7E-10	rs529565	136,149,500	8.0E-27	rs505922	136,149,229	2.0E-21
<i>GP5M1</i>	9	138,741,030-139,748,082	rs28505901	139,241,030	2.6E-21	rs376993806	139,246,588	4.5E-26	rs28642213	139,248,082	5.7E-68	rs28429551	139,243,334	1.4E-39
<i>RN7SL232P, CDC123, CAMK1D</i>	10	11,807,894-12,809,139	rs11257655	12,307,894	3.7E-32	rs11257657	12,309,139	9.8E-62	rs11257655	12,307,894	1.4E-63	rs11257655	12,307,894	1.2E-91
<i>BEND7</i>	10	13,040,869-14,040,869							rs11258422	13,540,869	3.2E-9			
<i>PTF1A</i>	10	22,987,778-23,987,778				rs77065181	23,487,778	1.6E-8						
<i>MYO3A</i>	10	25,997,704-26,997,704										rs7923442	26,497,704	1.2E-9
	10	33,497,227-34,497,227							rs71495046	33,997,227	1.9E-10			
	10	43,527,356-44,527,356							rs3122231	44,027,356	5.8E-10			
<i>ARID5B</i>	10	63,212,602-64,217,113				rs141583966	63,712,602	7.7E-10	rs146716733	63,717,113	1.2E-8			
<i>JMJD1C</i>	10	64,470,928-65,476,133				rs148928116	64,976,133	2.5E-13	rs111765639	64,970,928	5.4E-12	rs41274074	64,974,380	3.1E-11
<i>TFK1, TSPAN15, VPS26A, NEUROG3</i>	10	69,882,179-71,966,578	rs2642588	71,466,578	6.3E-14	rs1955163	71,273,357	1.7E-11	rs177045	71,321,279	2.7E-17	rs177045	71,321,279	5.0E-23
<i>PCBD1</i>	10	72,148,336-73,148,336							rs827237	72,648,336	2.5E-8			
	10	73,335,274-74,335,274							rs12773019 ^a	73,835,274	4.8E-8			
<i>CAMK2G</i>	10	75,098,099-76,098,099							rs2633311	75,598,099	3.1E-9			
<i>ZNF503, LRMDA</i>	10	76,744,336-77,823,643				rs7900112	77,314,617	5.4E-12	rs3012060	77,244,336	5.9E-10	rs3012060	77,244,336	1.6E-12
<i>ZMIZ1</i>	10	80,443,841-81,452,826	rs703972	80,952,826	2.5E-28	rs34204798	80,951,130	5.0E-19	rs697239	80,947,438	4.8E-54	rs703980	80,943,841	8.7E-40
<i>GRID1</i>	10	87,617,318-88,628,637							rs11201992	88,117,318	5.8E-11	rs3814613	88,128,637	1.1E-8
<i>PTEN</i>	10	89,184,214-90,266,368				rs1236816	89,684,214	4.3E-10	rs36062478	89,722,731	6.8E-13	rs10887775	89,766,368	8.9E-11
<i>BTAF1, MYOF, HHEX, IDE</i>	10	93,092,703-95,519,524	rs10882101	94,462,427	1.6E-62	rs35906730	94,435,673	1.3E-71	rs1111875	94,462,882	1.4E-128	rs10882101	94,462,427	1.8E-125
<i>RP11-452K12.4, ARHGAP19, SLIT1</i>	10	98,556,190-99,591,369				rs10748694	99,056,190	9.2E-11	rs945187	99,091,369	8.2E-16	rs10748694	99,056,190	5.1E-17
<i>HPSE2</i>	10	99,921,841-100,921,841							rs524903	100,421,841	2.1E-10			
<i>ERLIN1</i>	10	101,412,194-102,412,194							rs1408579	101,912,194	6.1E-11			
<i>RNU2-43P</i>	10	102,565,789-103,565,789							rs620191	103,065,789	1.7E-8			
<i>RNU6-1231P</i>	10	104,063,743-105,063,743							rs2482506	104,563,743	2.3E-12			
<i>BBIP1</i>	10	112,121,837-113,178,657				rs7895872	112,678,657	1.4E-11	rs7895872	112,678,657	1.3E-13	rs7067540	112,621,837	1.1E-14
<i>TCF7L2</i>	10	114,249,734-115,258,349	rs7903146	114,758,349	<E-300	rs7901695	114,754,088	8.2E-62	rs35011184	114,749,734	<E-300	rs7903146	114,758,349	<E-300
	10	115,321,878-116,321,878							rs10787518	115,821,878	8.8E-9			
	10	118,058,736-119,058,736							rs7912336	118,558,736	1.4E-8			
<i>SEC23IP</i>	10	121,160,400-122,160,400							rs11199116	121,660,400	1.0E-9			
<i>WDR11, FGFR2</i>	10	122,415,345-123,430,568				rs10886863	122,929,493	5.3E-17	rs7071036	122,930,568	2.0E-12	rs72631105	122,915,345	1.9E-18
<i>PLEKHA1</i>	10	123,650,342-124,693,181	rs2280141	124,193,181	2.0E-13	rs112820281	124,150,342	1.4E-10	rs2280141	124,193,181	6.0E-21	rs2421016	124,167,512	9.6E-23
<i>RP11-282I1.1</i>	10	124,726,178-125,726,178							rs705145	125,226,178	3.3E-9			
<i>INS, IGF2, KCNQ1, TH</i>	11	1,697,132-3,358,546	rs2237895	2,857,194	3.6E-44	rs2237897	2,858,546	1.9E-245	rs2237897	2,858,546	2.1E-226	rs2237897	2,858,546	5.5E-233
<i>TRIM66</i>	11	8,154,528-9,177,063							rs7941510	8,677,063	1.9E-17	rs10769936	8,654,528	6.9E-11
<i>SBF2</i>	11	9,356,015-10,356,015							rs76789970	9,856,015	9.2E-11			
<i>ARNTL</i>	11	12,840,710-13,840,710							rs10766076	13,340,710	1.3E-9			
<i>PDE3B, COPB1</i>	11	14,018,419-15,263,828	rs141521721	14,763,828	2.8E-8				rs117316450	14,518,419	9.5E-15	rs141521721	14,763,828	2.3E-8
<i>KCNJ11, ABCC8</i>	11	16,908,404-17,918,477	rs5213	17,408,404	1.9E-26	rs4148646	17,415,190	1.7E-26	rs757110	17,418,477	4.6E-52	rs5215	17,408,630	1.3E-54
<i>NELL1</i>	11	20,452,237-21,452,237							rs16907058	20,952,237	4.6E-8			
<i>BDNF</i>	11	27,183,618-28,229,505				rs988748	27,724,745	1.6E-10	rs10767659	27,686,196	4.6E-10	rs4923464	27,683,618	9.4E-10

MPPED2, RP5-1024C24.1	11	30,108,133-31,120,262								rs10835690	30,620,262	2.5E-8	rs11031140	30,608,133	2.8E-8
QSER1	11	32,427,778-33,456,492	rs145678014	32,927,778	1.1E-11					rs62618693	32,956,492	1.4E-13	rs145678014	32,927,778	5.7E-10
SLC1A2, PDHX, APIP	11	34,408,780-35,933,712	rs2767036	34,982,148	2.5E-8					rs2956092	34,908,780	1.7E-10	rs2985149	34,969,534	1.9E-8
HSD17B12	11	43,316,200-44,378,459	rs1061810	43,877,934	8.5E-13					rs35251247	43,878,459	1.7E-13	rs6485462	43,816,200	4.8E-12
CRY2	11	45,358,584-46,412,013	rs7115753	45,912,013	4.8E-9					rs12419690	45,858,584	1.4E-13	rs12419690	45,858,584	4.2E-11
ORAC9P, FOLH1, CELF1, NUP160	11	46,974,146-50,610,597	rs7124681	47,529,947	6.4E-9					rs3816605	47,857,253	8.9E-14	rs6485981	49,477,266	1.5E-9
OR5D18	11	55,088,216-56,088,216								rs116861182	55,588,216	5.4E-9			
OR5B17	11	57,628,015-58,628,015								rs7483027	58,128,015	5.6E-11			
FEN1	11	61,065,908-62,065,908								rs174541	61,565,908	2.4E-11			
AP003774.1	11	63,600,776-64,600,776								rs1662185	64,100,776	3.5E-8			
MAP3K11, LTBP3	11	64,794,799-65,826,154	rs1783541	65,294,799	1.4E-14					rs12789028	65,326,154	3.9E-20	rs12789028	65,326,154	2.1E-17
TPCN2, CCND1	11	68,335,182-69,963,273	rs11820019	69,448,758	1.0E-11	rs602652	69,462,642	5.3E-9		rs3918298	69,463,273	8.0E-25	rs3918298	69,463,273	3.3E-17
CENTD2, ARAP1	11	71,960,398-72,963,435	rs77464186	72,460,398	2.3E-33	rs7109575	72,463,435	5.5E-21		rs11602873	72,460,762	2.0E-62	rs77464186	72,460,398	3.6E-49
C11orf30	11	74,125,997-76,730,357								rs2513505	76,230,357	4.1E-10	rs61894507	76,156,973	2.2E-10
MTNR1B	11	92,208,710-93,208,710	rs10830963	92,708,710	1.5E-43	rs10830963	92,708,710	4.5E-8		rs10830963	92,708,710	1.3E-66	rs10830963	92,708,710	6.1E-66
MAML2	11	95,210,493-96,210,493								rs7130522	95,710,493	4.8E-8			
FXYD6, FXYD2	11	117,193,255-118,193,255								rs529623	117,693,255	4.3E-9	rs529623	117,693,255	1.5E-8
HMBS	11	118,453,202-119,453,202								rs7127212	118,953,202	2.4E-8			
ETS1	11	127,734,144-128,898,938	rs67232546	128,398,938	1.4E-12					rs10750397	128,234,144	1.7E-21	rs11819995	128,389,391	2.5E-14
CCND2	12	3,881,981-4,884,844	rs76895963	4,384,844	5.3E-70	rs7304270	4,381,981	1.0E-12		rs76895963	4,384,844	1.2E-96	rs76895963	4,384,844	3.7E-71
CHD4	12	6,191,452-7,191,452								rs7316626	6,691,452	3.3E-9			
CDKN1B	12	12,371,099-13,371,099	rs2066827	12,871,099	3.5E-8					rs2066827	12,871,099	2.2E-10	rs2066827	12,871,099	7.1E-11
PDE3A	12	20,079,392-21,091,332								rs7134150	20,591,332	2.7E-10	rs7488780	20,579,392	3.1E-8
LDHB, KCNJ8, RP11-59N23.3	12	21,343,576-22,371,751								rs11046164	21,843,576	4.0E-10	rs10841890	21,871,751	4.2E-8
ITPR2, RP11-283G6.4	12	25,953,283-26,974,867	rs718314	26,453,283	1.1E-10					rs11048457	26,463,174	2.2E-19	rs10842708	26,474,867	5.5E-14
KLHDC5, RN7SKP15	12	27,463,402-28,465,150	rs10842994	27,965,150	2.5E-20	rs3751236	27,963,402	6.6E-21		rs3751239	27,963,676	2.0E-47	rs12578595	27,964,996	7.8E-33
FAM60A, SINHCAF, DENND5B	12	30,917,019-31,941,179				rs80234489	31,441,179	4.3E-32		rs80234489	31,441,179	6.1E-18	rs78345706	31,417,019	1.2E-24
PKP2, SYT10	12	32,870,406-33,910,855								rs10844519	33,410,855	1.4E-12	rs6488140	33,370,406	9.1E-12
	12	38,210,523-39,210,523								rs7315028 ^b	38,710,523	1.5E-8			
PDZRN4	12	41,363,393-42,363,393								rs2730827	41,863,393	2.1E-13			
RP11-25I15.2	12	42,546,449-43,546,449								rs11181613	43,046,449	2.4E-12			
	12	45,368,623-46,368,623								rs2408252	45,868,623	1.1E-9			
	12	48,212,932-49,212,932								rs2732469	48,712,932	4.8E-14			
FAIM2	12	49,763,148-50,769,863				rs77978149	50,269,863	5.7E-9		rs7132908	50,263,148	7.4E-14	rs7132908	50,263,148	6.5E-10
HOXC6	12	53,929,385-54,929,385								rs12422600	54,429,385	3.1E-9			
PRIM1	12	56,646,069-58,468,738								rs2277339	57,146,069	6.9E-15			
	12	60,750,814-61,750,814								rs11173646	61,250,814	1.5E-9			
HMGA2, RPSAP52	12	65,716,162-66,746,181	rs2258238	66,221,060	2.0E-25	rs2583934	66,232,810	5.0E-16		rs2257883	66,216,162	3.9E-47	rs2583930	66,246,181	5.1E-38
TSPAN8, LGR5	12	70,949,521-72,022,953	rs1796330	71,522,953	3.2E-14	rs7313668	71,449,521	4.9E-11		rs10879261	71,520,761	6.8E-19	rs7313668	71,449,521	2.7E-17
LIN7A	12	80,809,262-81,809,262								rs11114650	81,309,262	3.3E-9			
	12	87,838,461-88,838,461								rs10745460 ^b	88,338,461	3.3E-8			
USP44	12	95,428,113-96,428,560	rs2197973	95,928,560	4.4E-8					rs11108094	95,928,113	1.1E-10			
RMST	12	97,348,775-98,351,611	rs77864822	97,848,775	2.2E-8	rs10860209	97,850,215	5.7E-9		rs6538805	97,849,120	1.5E-14	rs7972074	97,851,611	1.1E-10
	12	105,788,445-106,788,445								rs12825669	106,288,445	3.8E-8			
WSCD2	12	108,129,780-109,129,780	rs1426371	108,629,780	1.1E-11	rs1426371	108,629,780	7.8E-12		rs1426371	108,629,780	1.3E-21	rs1426371	108,629,780	9.7E-19
BRAP, SH2B3, ALDH2, PTPN11, HECTD4	12	111,109,727-113,617,897				rs149212747	111,836,771	2.1E-11					rs77753011	113,117,897	4.0E-15
RBM19	12	113,623,722-114,623,722				rs7307263	114,123,722	3.6E-8							
KSR2, NOS1	12	117,223,613-118,912,373	rs34965774	118,412,373	3.5E-9	rs111246699	118,400,856	1.5E-15		rs79310463	118,406,696	7.2E-21	rs34965774	118,412,373	1.5E-21

<i>HNF1A</i>	12	120,863,506-121,932,117	rs56348580	121,432,117	3.8E-19	rs118074491	121,363,506	8.8E-20	rs56348580	121,432,117	1.5E-27	rs1169299	121,429,194	1.9E-21
<i>C12orf65, ZNF664, MPHOSPH9, CCDC92</i>	12	122,950,765-125,045,435	rs4148856	123,450,765	2.2E-10				rs4930726	124,428,331	7.5E-18	rs1790116	123,618,544	2.2E-11
<i>ZNF10, EP400, FBRSL1</i>	12	132,044,643-134,230,500	rs12811407	133,069,698	2.4E-12				rs11614914	133,070,294	2.6E-19	rs12811407	133,069,698	2.0E-15
<i>SGCG, FGF9</i>	13	22,089,883-23,809,382				rs9316706	22,589,883	3.3E-9	rs314879	23,309,382	2.8E-15	rs314879	23,309,382	3.7E-11
<i>RNF6</i>	13	26,276,999-27,281,367	rs34584161	26,776,999	2.9E-10	rs568052023	26,781,367	2.6E-22	rs34584161	26,776,999	1.6E-37	rs34584161	26,776,999	3.5E-19
	13	27,745,127-28,745,127							rs9319382 ^a	28,245,127	2.9E-8			
<i>HMGB1</i>	13	30,517,268-31,542,452	rs11842871	31,042,452	1.5E-8				rs12856169	31,017,268	2.7E-9			
<i>KL</i>	13	33,054,302-34,057,644	rs576674	33,554,302	6.8E-10	rs7983505	33,557,173	3.2E-18	rs57286125	33,557,644	1.8E-33	rs2858980	33,554,587	6.2E-22
	13	41,188,401-42,188,401							rs4397977	41,688,401	1.1E-11			
	13	46,014,492-47,014,492							rs6561273	46,514,492	7.5E-9			
<i>DLEU1, RP11-175B12.2</i>	13	49,931,987-51,596,095	rs963740	51,096,095	2.6E-8	rs123378	51,088,809	2.2E-10	rs9316500	51,094,114	1.0E-22	rs963740	51,096,095	3.8E-11
<i>OLFM4</i>	13	53,607,583-54,607,583							rs9568868	54,107,583	5.8E-11	rs9568868	54,107,583	1.5E-12
<i>PCDH17, SRGAP2D</i>	13	57,866,634-59,577,406	rs9563615	59,077,406	3.9E-9				rs7991679	58,691,107	1.1E-8			
	13	65,704,880-66,704,880							rs9564268	66,204,880	7.7E-10			
<i>SPRY2</i>	13	80,205,315-81,217,156	rs1359790	80,717,156	5.7E-31	rs17072370	80,705,730	1.3E-31	rs11616380	80,705,315	1.0E-69	rs1215468	80,707,429	2.4E-56
<i>MIR17HG</i>	13	91,442,919-92,449,562				rs9523295	91,948,047	7.2E-18	rs9515905	91,949,562	2.1E-27	rs34165267	91,942,919	6.7E-21
<i>HS6ST3</i>	13	96,676,585-97,676,585							rs61967710	97,176,585	2.5E-8			
<i>IRS2</i>	13	109,446,882-110,447,213	rs7987740	109,947,213	4.1E-8				rs9587811	109,946,882	4.9E-11			
	13	111,687,882-112,687,882							rs9560114	112,187,882	3.3E-8			
<i>SLC7A7</i>	14	22,788,935-23,788,935	rs17122772	23,288,935	2.0E-8									
<i>NYNRIN</i>	14	24,378,370-25,378,370				rs12437434	24,878,370	1.0E-9						
	14	25,447,436-26,447,436							rs11159347	25,947,436	1.5E-9			
<i>PRKD1</i>	14	29,586,481-30,586,481							rs12433335	30,086,481	1.6E-8			
<i>AKAP6</i>	14	32,802,882-33,803,540	rs17522122	33,302,882	4.0E-9				rs12883788	33,303,540	2.8E-16	rs12883788	33,303,540	1.9E-11
<i>RP11-85K15.2</i>	14	34,909,701-35,909,701							rs712315	35,409,701	3.9E-8			
<i>CLEC14A</i>	14	38,303,756-39,348,419	rs8017808	38,848,419	2.6E-8	rs61975988	38,809,661	2.0E-9	rs7147483	38,804,675	9.4E-18	rs2183237	38,803,756	5.0E-15
<i>MDGA2</i>	14	46,804,091-47,804,091							rs723355	47,304,091	2.0E-9			
<i>RP11-349A22.5, PSMA3</i>	14	58,212,860-59,232,748							rs12892257	58,732,748	1.0E-8	rs61450169	58,712,860	5.4E-9
<i>MNAT1</i>	14	60,729,411-61,729,411							rs4902002	61,229,411	1.4E-9			
	14	68,959,229-69,959,229							rs242105	69,459,229	3.6E-11			
	14	74,432,641-75,432,641							rs12586772	74,932,641	6.4E-9			
<i>LRRC74A, C14orf166B</i>	14	76,800,863-77,882,503				rs58524310	77,382,503	8.4E-11	rs2056857	77,300,863	5.2E-11	rs72627178	77,372,210	2.1E-8
<i>NRXN3</i>	14	79,432,041-80,444,099	rs17836088	79,932,041	9.7E-14				rs7156625	79,942,647	5.4E-27	rs8008910	79,944,099	3.3E-15
<i>FOXP3</i>	14	89,050,378-90,050,378										rs17714667	89,550,378	4.2E-8
<i>SMEK1</i>	14	91,463,722-92,463,722	rs8010382	91,963,722	8.1E-9				rs8010382	91,963,722	3.0E-12	rs8010382	91,963,722	5.3E-9
<i>UNC79</i>	14	93,539,845-94,539,845							rs11848361	94,039,845	2.2E-8			
<i>DLK1, MEG3</i>	14	100,755,172-101,758,584				rs73347525	101,255,172	7.5E-11	rs112324411	101,258,584	1.4E-12	rs73347525	101,255,172	1.8E-15
<i>MARK3, TRAF3</i>	14	102,737,952-104,460,026	rs62007683	103,894,071	3.8E-8	rs55700915	103,237,952	1.5E-8	rs4906272	103,376,031	5.2E-11	rs11160699	103,252,270	5.9E-12
<i>HERC2</i>	15	28,046,173-29,046,173				rs76704029	28,546,173	3.4E-8						
	15	35,892,562-36,892,562							rs11073147 ^a	36,392,562	4.3E-8			
<i>C15orf52, INFAM2, RP11-624L4.1, RASGRP1</i>	15	38,328,140-41,134,717	rs34715063	38,873,115	3.3E-14	rs12907887	40,615,872	1.7E-20	rs8043085	38,828,140	9.1E-22	rs12912777	38,852,386	2.7E-16
<i>LTK</i>	15	41,301,512-42,318,917	rs11070332	41,809,205	1.3E-13				rs2289739	41,801,512	4.4E-11	rs1473781	41,818,917	2.1E-11
<i>PIIP5K1, STRC</i>	15	43,350,486-44,395,118							rs2447198	43,895,118	5.0E-10	rs475486	43,850,486	3.2E-8
<i>FAM227B</i>	15	49,294,020-50,294,020							rs7169799	49,794,020	3.2E-8			
<i>ONECUT1, WDR72, MYO5C</i>	15	52,017,714-54,247,228	rs2456530	53,091,553	4.7E-9	rs149336329	52,587,740	1.7E-9	rs149336329	52,587,740	2.9E-18	rs3825801	52,517,714	3.8E-11
<i>TCF12</i>	15	56,869,850-58,090,203	rs117483894	57,456,802	3.9E-8				rs28490139	57,369,850	1.5E-12	rs8024992	57,590,203	6.9E-9
<i>ALDH1A2</i>	15	58,176,821-59,176,821							rs11858759	58,676,821	4.3E-8			
	15	60,438,816-61,438,816							rs8033609 ^a	60,938,816	1.1E-8			

C2CD4A, C2CD4B	15	61,891,608-62,894,264	rs8037894	62,394,264	3.7E-13	rs8037894	62,394,264	7.3E-33	rs7163757	62,391,608	6.1E-30	rs7163757	62,391,608	2.4E-37
USP3	15	63,371,292-64,371,292	rs7178762	63,871,292	7.0E-10				rs7178762	63,871,292	1.8E-15	rs7178762	63,871,292	9.2E-12
MAP2K5	15	66,760,238-68,580,886	rs4776970	68,080,886	6.2E-9	rs4776970	68,080,886	3.4E-8	rs4776970	68,080,886	1.1E-12	rs4776970	68,080,886	5.7E-13
PML	15	73,828,576-74,828,576							rs9479	74,328,576	4.0E-13			
PTPN9, SIN3A	15	75,242,095-76,432,129	rs13737	75,932,129	7.3E-10	rs7171507	75,737,287	1.8E-11	rs6495182	75,814,388	2.1E-22	rs11636031	75,815,758	4.9E-19
HMG20A	15	77,276,562-78,318,128	rs1005752	77,818,128	5.7E-29	rs952471	77,776,498	1.6E-26	rs12910361	77,782,335	5.2E-65	rs952472	77,776,562	4.1E-56
FSD2	15	82,961,873-83,961,873							rs36111056	83,461,873	1.4E-8			
ADAMTSL3	15	84,047,222-85,047,222							rs1812707	84,547,222	5.9E-9			
AP3S2	15	89,879,632-90,928,894	rs4932265	90,423,293	7.2E-20	rs10852123	90,428,894	8.4E-13	rs893617	90,381,278	2.1E-38	rs6496609	90,379,632	3.2E-34
PRC1	15	91,011,260-92,022,253	rs12910825	91,511,260	2.4E-15	rs8026714	91,522,253	1.1E-22	rs2290203	91,512,067	2.3E-31	rs2890156	91,513,157	3.8E-34
RP11-26608.1, RGMA	15	93,325,384-94,425,327				rs61021634	93,825,384	1.4E-11	rs4777857	93,925,327	4.8E-11	rs7167984	93,832,067	3.4E-13
IGF1R	15	98,776,521-99,866,409				rs79826452	99,366,409	3.2E-8	rs59646751	99,276,521	3.4E-9			
LMF1, ITFG3	16	0-1,467,241	rs6600191	295,795	7.0E-13				rs55857387	300,388	1.7E-21	rs6600191	295,795	2.4E-15
CLUAP1, SLX4	16	3,083,173-4,156,482	rs3751837	3,583,173	1.7E-8	rs2240885	3,647,098	2.8E-9	rs8061528	3,656,482	2.6E-14	rs12445430	3,613,126	2.5E-11
NTAN1	16	14,653,717-15,653,717							rs9927842	15,153,717	7.0E-10			
GP2	16	19,823,168-20,834,808				rs117267808	20,323,168	4.9E-17	rs4609857	20,334,808	1.1E-10	rs117267808	20,323,168	3.0E-8
ATP2A1	16	28,397,452-29,415,217	rs8046545	28,915,217	2.3E-8				rs8056890	28,897,452	1.5E-12			
FAM57B, TMEM219	16	29,458,216-30,545,789	rs11642430	30,045,789	1.2E-10				rs8054556	29,958,216	1.9E-16	rs11642430	30,045,789	6.6E-10
FTO	16	53,300,954-54,887,084	rs1421085	53,800,954	2.4E-78	rs1421085	53,800,954	1.6E-48	rs1421085	53,800,954	1.3E-189	rs55872725	53,809,123	4.7E-128
AMFR	16	55,959,589-56,959,589							rs111283203	56,459,589	3.4E-8			
PKD1L3, IL34, NFAT5	16	69,151,866-72,522,534	rs862320	69,651,866	5.1E-11	rs12600132	72,022,534	5.9E-9	rs244415	69,666,683	2.0E-22	rs862320	69,651,866	1.5E-10
RP11-346C20.3, ZFH3	16	72,598,091-73,600,308				rs6416749	73,100,308	3.4E-12	rs1075855	73,098,091	3.5E-9	rs6416749	73,100,308	1.5E-13
BCAR1, CTRB2	16	74,734,872-75,746,035	rs72802342	75,234,872	1.3E-27				rs72802365	75,246,035	1.1E-40	rs72802358	75,243,657	2.9E-29
CMIP	16	81,033,789-82,034,790	rs2925979	81,534,790	2.1E-14	rs2925979	81,534,790	1.5E-9	rs56823429	81,533,789	3.3E-20	rs2925979	81,534,790	1.6E-21
GINS2	16	85,216,463-86,216,463							rs11646052	85,716,463	2.7E-11			
ZFPM1	16	87,356,424-89,054,480							rs9937296	88,554,480	2.4E-10	rs9937296	88,554,480	5.3E-10
SPG7, RPL13	16	89,064,055-90,130,630	rs12920022	89,564,055	2.9E-9				rs12932337	89,630,630	8.3E-11	rs12920022	89,564,055	9.9E-10
VPS53	17	0-981,604							rs11870735	481,604	7.6E-9			
ENO3, ZZEF1	17	3,488,451-5,354,480	rs1377807	4,045,440	5.7E-17				rs8071043	3,988,451	5.4E-30	rs8071043	3,988,451	4.3E-15
SAT2, SLC16A11, SLC16A13	17	6,453,155-8,031,965				rs186568031	6,953,781	9.0E-24	rs73239895	6,953,558	3.0E-15	rs113748381	6,953,155	2.3E-24
GLP2R	17	9,285,187-10,287,845	rs7222481	9,785,187	1.7E-8				rs17810376	9,787,845	2.8E-9			
RAI1	17	17,161,802-18,251,478	rs4925109	17,661,802	3.9E-12				rs2297508	17,715,317	2.4E-14	rs1108646	17,751,478	2.8E-13
KCNJ12	17	20,784,910-21,784,910							rs117642733	21,284,910	9.5E-9			
CRYBA1	17	27,070,622-28,070,622							rs9913225	27,570,622	5.4E-11			
NF1	17	28,913,019-30,204,002	rs71372253	29,413,019	4.3E-8	rs7502556	29,642,430	3.8E-11	rs2040792	29,628,549	2.0E-15	rs1048317	29,704,002	1.4E-14
MYO19	17	34,362,220-35,362,220							rs1109442	34,862,220	6.1E-9			
HNF1B, TCF2	17	35,599,840-36,601,586	rs10908278	36,099,952	3.1E-30	rs8064454	36,101,586	6.5E-61	rs11651755	36,099,840	8.6E-67	rs10908278	36,099,952	7.4E-74
	17	37,246,307-38,246,307							rs11078916	37,746,307	1.2E-14			
MLX, LINC00910, RP11-400F19.6	17	40,196,915-41,956,413	rs34855406	40,731,411	3.2E-12				rs676387	40,706,273	3.3E-24	rs684214	40,696,915	3.0E-13
GIP, TTL6, CBX1	17	45,678,674-47,560,322	rs35895680	47,060,322	3.8E-15				rs35895680	47,060,322	8.4E-27	rs35895680	47,060,322	2.3E-14
KIF2B	17	51,640,805-52,640,805	rs569511541	52,140,805	1.5E-8									
ERN1, ACE	17	61,065,025-62,703,304	rs60276348	62,203,304	2.9E-8				rs4335	61,565,025	1.1E-15	rs57676627	62,203,128	1.8E-10
PITPNC1, BPTF	17	65,141,651-66,457,568	rs61676547	65,892,507	1.0E-11	rs2706710	65,641,651	1.7E-8	rs12603589	65,825,248	1.1E-19	rs9899520	65,957,568	1.3E-14
SLC39A11	17	70,145,032-71,145,032							rs61736066	70,645,032	4.9E-11			
SUMO2	17	72,687,031-73,687,031				rs35559984	73,187,031	7.9E-9						
UBE2O	17	73,918,176-75,886,909							rs1656794*	75,386,909	3.6E-9			
CYTH1	17	76,272,288-77,292,179							rs7224711	76,772,288	1.2E-15	rs1044486	76,792,179	3.2E-13
RPTOR	17	77,395,311-79,257,626							rs11150745	78,757,626	5.2E-9			

RP11-172F10.1	18	4,345,027-5,345,027							rs9958640	4,845,027	2.3E-8			
LAMA1	18	6,570,642-7,576,836	rs7240767	7,070,642	2.0E-8	rs9948462	7,076,836	8.7E-10	rs7240767	7,070,642	2.1E-11	rs9948462	7,076,836	3.9E-15
	18	12,771,367-13,771,367							rs11662800 ^a	13,271,367	2.4E-9			
C18orf8	18	20,583,738-21,583,738							rs303760	21,083,738	2.0E-14			
NOL4	18	31,082,890-32,082,890							rs17747955	31,582,890	1.6E-8			
COMMD9	18	35,778,709-37,246,623	rs62080313	36,278,709	9.1E-9				rs7227272	36,746,623	4.7E-13			
LINC00907	18	39,566,006-40,566,006							rs410150	40,066,006	1.2E-9			
TCF4	18	52,550,646-53,550,646	rs72926932	53,050,646	3.6E-13				rs72926932	53,050,646	4.3E-20	rs72926932	53,050,646	1.2E-9
WDR7	18	54,078,482-55,175,384	rs17684074	54,675,384	3.5E-8				rs10048404	54,578,482	6.1E-9			
RNU4-17P, GRP, MC4R	18	56,376,228-58,352,587	rs523288	57,848,369	7.5E-14	rs476828	57,852,587	4.8E-27	rs6567160	57,829,135	7.5E-34	rs6567160	57,829,135	1.1E-37
BCL2A	18	60,345,884-61,345,884	rs12454712	60,845,884	5.1E-13	rs12454712	60,845,884	1.4E-15	rs12454712	60,845,884	2.4E-27	rs12454712	60,845,884	4.1E-20
CDH7	18	62,916,719-63,926,979							rs2032217	63,426,979	2.7E-11	rs1942267	63,416,719	7.3E-9
ZNF236	18	74,055,593-75,058,999							rs6565922	74,558,999	6.2E-13	rs12457906	74,555,593	6.0E-13
TCF3	19	1,146,712-2,146,712							rs4807125	1,646,712	2.6E-8			
UHRF1, PTPRS, KDM4B	19	4,448,862-5,467,739	rs7249758	4,948,862	1.2E-8				rs12185519	4,967,739	2.5E-11	rs262549	4,951,064	2.5E-9
MAP2K7, INSR, ACO10336.1	19	6,740,848-8,486,638	rs4804833	7,970,635	1.1E-12	rs475002	7,986,638	9.8E-10	rs2115107	7,968,168	2.3E-20	rs2115107	7,968,168	1.1E-18
FARSA, ZNF799, GCDH	19	12,005,873-13,538,415	rs3111316	13,038,415	1.6E-12	rs4804181	12,509,536	1.5E-8	rs9384	13,010,643	8.4E-23	rs3111316	13,038,415	1.3E-13
CILP2, CRTCL1, TM6SF2	19	18,334,514-19,888,500	rs8107974	19,388,500	6.3E-15				rs58542926	19,379,549	1.6E-23	rs58542926	19,379,549	1.6E-13
ZNF257, ZNF738	19	21,029,576-22,600,706				rs142395395	22,100,706	6.9E-23				rs142395395	22,100,706	1.2E-15
AC007796.1	19	31,365,946-32,365,946							rs2867570	31,865,946	2.1E-11			
PEPD	19	33,390,838-34,396,432	rs10406327	33,890,838	4.6E-8	rs7250869	33,887,405	2.3E-16	rs4805881	33,896,432	8.4E-22	rs10406327	33,890,838	3.5E-20
RN7SL836P, EML2, TOMM40, APOE, GIPR	19	44,911,941-46,658,417	rs10406431	46,157,019	2.5E-19	rs113036890	46,157,928	1.0E-32	rs8107527	46,158,417	4.2E-41	rs10406431	46,157,019	8.7E-44
ZC3H4	19	47,069,003-48,097,102	rs3810291	47,569,003	1.2E-11				rs10408163	47,597,102	1.1E-13	rs3810291	47,569,003	8.6E-19
FCGRT	19	49,516,759-50,516,759							rs142385484	50,016,759	2.0E-8			
STK35	20	1,600,095-2,600,095							rs6137042	2,100,095	2.3E-8			
CFAP61	20	19,568,635-20,568,635										rs7261425	20,068,635	1.2E-8
FOXA2, NKX2-2	20	20,966,795-22,930,241	rs13041756	21,466,795	1.3E-8	rs73085586	22,430,241	1.7E-9	rs7274134	22,428,284	2.3E-9	rs2181063	22,427,370	5.9E-10
RALY, EIF2S2	20	32,096,704-33,175,727	rs2268078	32,596,704	2.9E-10				rs6059662	32,675,727	2.1E-17	rs4911405	32,674,967	4.4E-12
ZHX3	20	39,332,628-40,332,628							rs17265513	39,832,628	3.0E-9			
IIFT52, HNF4A	20	41,730,695-43,542,364	rs1800961	43,042,364	3.2E-20	rs12625671	42,994,812	2.3E-21	rs12625671	42,994,812	6.1E-28	rs12625671	42,994,812	9.5E-40
EYA2	20	45,094,711-46,098,564	rs6063048	45,598,564	5.8E-11				rs6066138	45,594,711	6.3E-19	rs6063046	45,596,378	1.5E-10
CEBPB	20	48,330,772-49,332,135	rs11699802	48,832,135	2.5E-11	rs13040225	48,830,772	1.6E-14	rs13040225	48,830,772	1.1E-17	rs6091115	48,832,020	9.3E-23
NFATC2, TSHZ2, RP4-723E3.1	20	49,655,386-52,120,857	rs34454109	51,223,594	8.8E-9	rs6021276	50,155,386	6.7E-10	rs4809906	51,033,681	5.1E-18	rs34454109	51,223,594	8.1E-9
GNAS	20	56,887,352-57,977,177	rs6070625	57,394,628	3.2E-12	rs11477757	57,477,177	1.3E-8	rs4810145	57,396,495	1.1E-15	rs736266	57,387,352	6.6E-10
SLCO4A1	20	60,777,014-61,777,014							rs1815591	61,277,014	3.2E-14			
ZBTB46	20	61,950,664-62,950,664							rs6011155	62,450,664	5.9E-13			
	21	47,267,295-48,267,295							rs75756987 ^b	47,767,295	1.6E-9			
ARVCF	22	19,469,696-20,469,696							rs2240716	19,969,696	4.8E-10			
MTMR3, ASCC2, ZNRF3, CTA-85E5.10	22	28,869,398-31,109,554	rs6518681	30,609,554	9.6E-13	rs147413364	29,380,119	3.4E-8	rs56392746	30,451,688	3.5E-14	rs36575	30,205,572	1.9E-13
YWHAH, DEPDC5	22	31,703,334-32,848,841	rs117001013	32,348,841	1.5E-8				rs75307421	32,203,334	2.6E-8	rs75307421	32,203,334	3.1E-11
TOM1	22	35,205,359-36,205,359							rs138771	35,705,359	3.8E-9			
MAFF	22	38,099,767-39,099,767							rs4820323	38,599,767	3.6E-8			
EP300, RP1-85F18.5	22	40,041,838-42,093,581	rs5758223	41,489,920	4.6E-8				rs11913442	41,593,581	8.2E-10	rs738630	41,511,171	5.4E-9
PNPLA3	22	43,824,730-44,824,855	rs738408	44,324,730	1.8E-10				rs3747207	44,324,855	3.7E-21	rs738408	44,324,730	5.5E-10
WNT7B	22	45,813,618-46,813,618				rs28637892	46,313,618	3.7E-9						
PIM3	22	49,856,302-50,856,850	rs1801645	50,356,850	1.5E-10	rs28691713	50,356,302	1.8E-17	rs1801645	50,356,850	4.2E-17	rs28691713	50,356,302	5.9E-22

^aReported only in MVP European ancestry-specific meta-analysis. ^bReported only in MVP African ancestry-specific meta-analysis.

Supplementary Note Table 3. Loci attaining genome-wide significant evidence ($p < 5 \times 10^{-8}$) of association with T2D in European ancestry-specific meta-analysis of up to 80,154 cases and 853,816 controls (effective sample size 251,740) that were not identified at the same threshold in the trans-ancestry meta-regression.

Locus	Lead SNV	Chr	Position (bp, b37)	Alleles		European ancestry-specific meta-analysis				Non-European ancestry meta-analysis		
				Risk	Other	RAF	OR (95% CI)	p -value	Q p -value	RAF	OR (95% CI)	p -value
<i>EGFEM1P</i>	rs7642311	3	168,223,132	A	G	0.872	1.06 (1.04-1.08)	1.5×10^{-8}	0.034	0.823	1.01 (0.98-1.04)	0.67
<i>HTT</i>	rs362307	4	3,241,845	T	C	0.078	1.07 (1.05-1.10)	6.8×10^{-9}	0.47	0.083	1.03 (0.98-1.08)	0.32
<i>FAM13A</i>	rs1903002	4	89,740,894	G	C	0.506	1.04 (1.02-1.05)	1.4×10^{-8}	0.46	0.685	1.01 (0.99-1.02)	0.22
<i>RFX3</i>	rs672271	9	3,273,781	C	T	0.094	1.06 (1.04-1.08)	4.8×10^{-8}	0.38	0.175	0.99 (0.97-1.01)	0.52
<i>USP44</i>	rs61939481	12	95,921,998	C	T	0.068	1.07 (1.05-1.10)	1.7×10^{-8}	0.16	0.043	1.01 (0.96-1.06)	0.72
<i>HMGB1</i>	rs11842871	13	31,042,452	G	T	0.733	1.04 (1.03-1.06)	4.8×10^{-8}	0.097	0.870	1.02 (1.00-1.04)	0.082
<i>PCDH17-SRGAP2D</i>	rs9563615	13	59,077,406	A	T	0.709	1.04 (1.03-1.06)	1.6×10^{-8}	0.12	0.535	1.01 (1.00-1.02)	0.18
<i>LIG4</i>	rs7325671	13	108,797,836	T	C	0.126	1.06 (1.04-1.08)	1.5×10^{-8}	0.74	0.195	1.01 (0.99-1.03)	0.24
<i>SLC7A7</i>	rs17122772	14	23,288,935	G	C	0.226	1.04 (1.03-1.06)	3.3×10^{-8}	0.21	0.145	1.02 (1.00-1.05)	0.044
<i>GLP2R</i>	rs55973554	17	9,793,756	A	G	0.322	1.04 (1.03-1.05)	1.0×10^{-8}	0.57	0.119	1.01 (0.98-1.04)	0.51
<i>PIK3C3-RIT2</i>	rs1431841	18	40,087,098	T	G	0.211	1.04 (1.03-1.06)	3.1×10^{-8}	0.78	0.158	1.02 (1.00-1.03)	0.10

Chr: chromosome. RAF: risk allele frequency. OR: odds-ratio. CI: confidence interval.

Supplementary Note Table 4. Comparison of African ancestry-specific association summary statistics obtained from approximate conditional analysis undertaken in loci with multiple distinct signals using two LD reference panels: 661 individuals of African ancestry from the 1000 Genomes Project; and 1,000 African American individuals from GERA.

Locus	Index SNV	Alleles		RAF	1000 Genomes Project LD reference			GERA African American LD reference		
		Risk	Other		log OR	SE	<i>p</i> -value	log OR	SE	<i>p</i> -value
<i>TMEM18</i>	rs62107261	T	C	0.985	0.103	0.112	0.36	0.087	0.112	0.44
<i>TMEM18</i>	rs10188334	C	T	0.862	0.107	0.030	0.00036	0.106	0.030	0.00040
<i>CEP68</i>	rs2540949	A	T	0.581	0.026	0.019	0.16	0.025	0.019	0.19
<i>CEP68</i>	rs6752053	T	C	0.417	0.036	0.019	0.054	0.035	0.019	0.062
<i>PPARG</i>	rs17036160	C	T	0.976	0.089	0.059	0.13	0.093	0.059	0.11
<i>PPARG</i>	rs4684855	T	C	0.115	0.102	0.028	0.00031	0.103	0.028	0.00025
<i>UBE2E2</i>	rs13094957	T	C	0.642	0.024	0.019	0.21	0.024	0.019	0.19
<i>UBE2E2</i>	rs76435632	G	C	0.011	0.192	0.090	0.033	0.194	0.090	0.031
<i>PSMD6-ADAMTS9</i>	rs2292662	C	T	0.595	0.016	0.019	0.41	0.019	0.019	0.31
<i>PSMD6-ADAMTS9</i>	rs66815886	G	T	0.448	0.046	0.019	0.014	0.047	0.019	0.013
<i>MBNL1</i>	rs1426385	A	G	0.419	0.053	0.019	0.0043	0.056	0.019	0.0025
<i>MBNL1</i>	rs10935897	A	G	0.560	0.035	0.018	0.061	0.036	0.018	0.043
<i>MBNL1</i>	rs75417759	C	T	0.985	0.068	0.082	0.41	0.067	0.081	0.41
<i>ST6GAL1</i>	rs3887925	T	C	0.213	0.033	0.022	0.14	0.033	0.022	0.13
<i>ST6GAL1</i>	rs9799068	A	C	0.575	0.031	0.018	0.085	0.031	0.018	0.076
<i>ANKH</i>	rs147581833	C	T	0.999	-0.506	0.797	0.53	-0.570	0.796	0.47
<i>ANKH</i>	rs30614	A	G	0.637	0.070	0.056	0.21	0.069	0.056	0.21
<i>ARL15</i>	rs702634	A	G	0.751	0.041	0.021	0.058	0.037	0.021	0.077
<i>ARL15</i>	rs6876198	C	T	0.302	0.071	0.020	0.00043	0.069	0.020	0.00055
<i>ANKRD55</i>	rs256904	T	A	0.573	0.101	0.019	1.8x10 ⁻⁷	0.101	0.018	1.4x10 ⁻⁸
<i>ANKRD55</i>	rs42251	A	G	0.206	0.000	0.023	0.99	-0.003	0.022	0.88
<i>ANKRD55</i>	rs3936510	T	G	0.231	0.019	0.022	0.38	0.027	0.021	0.20
<i>SSR1-RREB1</i>	rs77630070	G	T	0.966	0.074	0.055	0.18	0.074	0.055	0.18
<i>SSR1-RREB1</i>	rs9379084	G	A	0.967	0.183	0.064	0.0042	0.183	0.064	0.0042
<i>ZFAND3-KCNK16-GLP1R</i>	rs2281342	T	C	0.920	0.076	0.035	0.027	0.080	0.035	0.021
<i>ZFAND3-KCNK16-GLP1R</i>	rs742762	A	C	0.870	0.022	0.027	0.41	0.032	0.027	0.24
<i>ZFAND3-KCNK16-GLP1R</i>	rs3734618	G	A	0.876	0.047	0.030	0.12	0.045	0.030	0.13
<i>DGKB</i>	rs17168486	T	C	0.109	0.008	0.031	0.80	0.014	0.031	0.65
<i>DGKB</i>	rs2215383	C	T	0.568	0.081	0.019	1.3x10 ⁻⁵	0.081	0.019	1.4x10 ⁻⁵
<i>JAZF1</i>	rs849133	C	T	0.741	0.080	0.022	0.00031	0.099	0.020	1.1x10 ⁻⁶
<i>JAZF1</i>	rs552707	T	C	0.138	0.037	0.015	0.014	0.032	0.015	0.027
<i>JAZF1</i>	rs10226758	C	A	0.798	0.038	0.014	0.0087	0.035	0.014	0.011

<i>KCNU1</i>	rs10092900	G	T	0.403	0.061	0.019	0.00097	0.060	0.019	0.0011
<i>KCNU1</i>	rs12680217	T	C	0.704	0.085	0.021	4.7x10 ⁻⁵	0.084	0.021	5.3x10 ⁻⁵
<i>ANK1</i>	rs12550613	C	G	0.715	0.064	0.021	0.0017	0.059	0.020	0.0036
<i>ANK1</i>	rs508419	G	A	0.724	0.033	0.021	0.12	0.022	0.020	0.28
<i>GLIS3</i>	rs4237150	C	G	0.531	0.008	0.020	0.67	0.009	0.020	0.65
<i>GLIS3</i>	rs4258054	T	C	0.561	-0.004	0.021	0.87	-0.005	0.021	0.82
<i>TLE1</i>	rs9332453	C	T	0.883	-0.023	0.033	0.48	-0.024	0.032	0.46
<i>TLE1</i>	rs2796441	G	A	0.827	0.011	0.026	0.68	0.012	0.026	0.65
<i>VPS26A-NEUROG3</i>	rs190925	A	G	0.438	-0.052	0.021	0.012	-0.037	0.019	0.044
<i>VPS26A-NEUROG3</i>	rs41277236	T	C	0.013	0.185	0.128	0.15	0.193	0.127	0.13
<i>VPS26A-NEUROG3</i>	rs2642588	G	T	0.500	-0.010	0.020	0.63	-0.009	0.018	0.63
<i>HHEX-IDE</i>	rs10882099	T	C	0.767	0.032	0.021	0.13	0.031	0.020	0.12
<i>HHEX-IDE</i>	rs139027698	T	C	0.069	0.059	0.036	0.10	0.062	0.037	0.092
<i>HHEX-IDE</i>	rs1112718	A	G	0.560	-0.012	0.014	0.38	-0.010	0.014	0.47
<i>WDR11</i>	rs11199753	G	T	0.973	0.118	0.061	0.051	0.129	0.059	0.027
<i>WDR11</i>	rs2172073	A	C	0.499	0.061	0.019	0.0012	0.050	0.017	0.0042
<i>WDR11</i>	rs11592107	A	G	0.078	0.049	0.034	0.15	0.043	0.034	0.21
<i>MTNR1B</i>	rs10830963	G	C	0.084	0.138	0.038	0.00027	0.143	0.038	0.00018
<i>MTNR1B</i>	rs11020308	A	C	0.114	0.024	0.032	0.44	0.036	0.032	0.26
<i>ETS1</i>	rs10893827	A	G	0.741	0.033	0.022	0.13	0.036	0.021	0.086
<i>ETS1</i>	rs7104712	C	A	0.202	0.028	0.023	0.23	0.028	0.022	0.21
<i>ETS1</i>	rs11819995	T	C	0.266	0.032	0.021	0.13	0.034	0.020	0.096
<i>HMGA2</i>	rs343093	G	C	0.811	0.117	0.025	2.8x10 ⁻⁶	0.116	0.025	2.8x10 ⁻⁶
<i>HMGA2</i>	rs7970350	T	C	0.377	-0.002	0.018	0.91	0.004	0.018	0.82
<i>RASGRP1</i>	rs28582094	G	A	0.251	0.053	0.022	0.015	0.055	0.022	0.011
<i>RASGRP1</i>	rs34715063	C	T	0.058	0.077	0.047	0.10	0.078	0.047	0.097
<i>GRP-MC4R</i>	rs9957320	G	T	0.866	0.002	0.028	0.96	0.008	0.026	0.74
<i>GRP-MC4R</i>	rs6567160	C	T	0.196	0.075	0.024	0.0022	0.079	0.023	0.00054
<i>GRP-MC4R</i>	rs76227980	C	T	0.956	0.075	0.052	0.15	0.076	0.051	0.14
<i>HNFA4</i>	rs12625671	C	T	0.083	0.127	0.038	0.00073	0.127	0.038	0.00071
<i>HNFA4</i>	rs1800961	T	C	0.008	0.094	0.142	0.51	0.100	0.142	0.48

RAF: risk allele frequency. OR: odds-ratio. SE: standard error.

Supplementary Note Table 5. Comparison of Hispanic ancestry-specific association summary statistics obtained from approximate conditional analysis undertaken in loci with multiple distinct signals using two LD reference panels: 347 individuals of American ancestry from the 1000 Genomes Project; and 1,000 Hispanic individuals from GERA.

Locus	Index SNV	Alleles		RAF	1000 Genomes Project LD reference			GERA Hispanic LD reference		
		Risk	Other		log OR	SE	<i>p</i> -value	log OR	SE	<i>p</i> -value
<i>DSTYK-MDM4</i>	rs6689629	A	G	0.910	0.028	0.033	0.41	0.028	0.033	0.41
<i>DSTYK-MDM4</i>	rs12039805	A	G	0.341	0.059	0.020	0.0025	0.059	0.020	0.0025
<i>TMEM18</i>	rs62107261	T	C	0.969	0.113	0.059	0.056	0.118	0.059	0.047
<i>TMEM18</i>	rs10188334	C	T	0.877	0.064	0.029	0.026	0.066	0.029	0.023
<i>CEP68</i>	rs2540949	A	T	0.659	0.033	0.020	0.090	0.030	0.020	0.13
<i>CEP68</i>	rs6752053	T	C	0.649	0.050	0.020	0.011	0.048	0.020	0.015
<i>PPARG</i>	rs17036160	C	T	0.890	0.109	0.029	0.00021	0.109	0.031	0.00037
<i>PPARG</i>	rs4684855	T	C	0.378	0.025	0.018	0.18	0.016	0.019	0.42
<i>UBE2E2</i>	rs13094957	T	C	0.857	0.053	0.028	0.061	0.051	0.028	0.071
<i>UBE2E2</i>	rs76435632	G	C	0.076	0.078	0.037	0.034	0.076	0.037	0.039
<i>PSMD6-ADAMTS9</i>	rs2292662	C	T	0.869	0.005	0.030	0.86	0.012	0.030	0.70
<i>PSMD6-ADAMTS9</i>	rs66815886	G	T	0.758	0.032	0.023	0.16	0.034	0.023	0.14
<i>MBNL1</i>	rs1426385	A	G	0.486	0.018	0.018	0.34	0.015	0.019	0.42
<i>MBNL1</i>	rs10935897	A	G	0.312	0.058	0.020	0.0034	0.053	0.020	0.0075
<i>MBNL1</i>	rs75417759	C	T	0.982	0.102	0.076	0.18	0.105	0.074	0.16
<i>ST6GAL1</i>	rs3887925	T	C	0.376	0.047	0.019	0.011	0.047	0.019	0.011
<i>ST6GAL1</i>	rs9799068	A	C	0.208	0.025	0.023	0.28	0.025	0.023	0.27
<i>ARL15</i>	rs702634	A	G	0.813	0.027	0.025	0.29	0.018	0.024	0.46
<i>ARL15</i>	rs6876198	C	T	0.273	0.066	0.021	0.0019	0.063	0.021	0.0023
<i>ANKRD55</i>	rs256904	T	A	0.743	0.067	0.021	0.0017	0.078	0.021	0.00022
<i>ANKRD55</i>	rs42251	A	G	0.292	0.058	0.020	0.0038	0.065	0.019	0.00083
<i>ANKRD55</i>	rs3936510	T	G	0.218	0.125	0.022	9.6x10 ⁻⁹	0.127	0.021	2.5x10 ⁻⁹
<i>SLCO6A1-PAM</i>	rs78408340	G	C	0.005	0.433	0.174	0.013	0.400	0.155	0.0097
<i>SLCO6A1-PAM</i>	rs115505614	T	C	0.014	0.132	0.097	0.17	0.126	0.087	0.15
<i>SLCO6A1-PAM</i>	rs186327337	G	A	0.998	0.134	0.297	0.65	0.144	0.297	0.63
<i>SSR1-RREB1</i>	rs77630070	G	T	0.934	0.044	0.039	0.26	0.053	0.038	0.17
<i>SSR1-RREB1</i>	rs9379084	G	A	0.913	0.082	0.035	0.021	0.086	0.035	0.014
<i>ZFAND3-KCNK16-GLP1R</i>	rs2281342	T	C	0.766	0.083	0.022	0.00018	0.085	0.022	0.00014
<i>ZFAND3-KCNK16-GLP1R</i>	rs742762	A	C	0.868	0.076	0.027	0.0054	0.082	0.028	0.0029
<i>ZFAND3-KCNK16-GLP1R</i>	rs3734618	G	A	0.485	0.051	0.019	0.0074	0.060	0.020	0.0023
<i>DGKB</i>	rs17168486	T	C	0.428	0.047	0.020	0.019	0.043	0.020	0.035
<i>DGKB</i>	rs2215383	C	T	0.456	0.055	0.019	0.0031	0.053	0.019	0.0052

JAZF1	rs849133	C	T	0.642	0.105	0.019	3.9x10 ⁻⁸	0.100	0.018	1.8x10 ⁻⁸
JAZF1	rs552707	T	C	0.151	0.005	0.012	0.66	0.001	0.013	0.93
JAZF1	rs10226758	C	A	0.812	-0.002	0.012	0.87	0.000	0.013	1.0
MNX1	rs887609	A	G	0.189	0.093	0.025	0.00020	0.090	0.025	0.00036
MNX1	rs2366214	A	G	0.537	0.052	0.019	0.0060	0.048	0.019	0.012
KCNU1	rs10092900	G	T	0.203	0.032	0.023	0.17	0.030	0.023	0.20
KCNU1	rs12680217	T	C	0.760	-0.023	0.023	0.32	-0.019	0.023	0.40
ANK1	rs12550613	C	G	0.598	0.013	0.017	0.45	0.014	0.017	0.43
ANK1	rs508419	G	A	0.776	0.060	0.020	0.0030	0.061	0.021	0.0029
GLIS3	rs4237150	C	G	0.515	0.084	0.019	1.1x10 ⁻⁵	0.084	0.019	1.1x10 ⁻⁵
GLIS3	rs4258054	T	C	0.503	0.025	0.020	0.22	0.025	0.020	0.21
TLE1	rs9332453	C	T	0.708	0.024	0.021	0.25	0.023	0.021	0.27
TLE1	rs2796441	G	A	0.519	0.058	0.019	0.0027	0.058	0.019	0.0028
VPS26A-NEUROG3	rs190925	A	G	0.393	0.033	0.019	0.082	0.031	0.019	0.10
VPS26A-NEUROG3	rs41277236	T	C	0.036	0.096	0.055	0.079	0.073	0.054	0.18
VPS26A-NEUROG3	rs2642588	G	T	0.695	0.011	0.021	0.59	0.011	0.021	0.59
HHEX-IDE	rs10882099	T	C	0.640	0.075	0.018	3.1x10 ⁻⁵	0.066	0.016	4.1x10 ⁻⁵
HHEX-IDE	rs139027698	T	C	0.025	0.047	0.064	0.47	0.038	0.063	0.55
HHEX-IDE	rs1112718	A	G	0.475	0.009	0.017	0.60	-0.004	0.016	0.81
WDR11	rs11199753	G	T	0.875	0.055	0.028	0.048	0.052	0.028	0.065
WDR11	rs2172073	A	C	0.827	0.069	0.025	0.0051	0.074	0.025	0.0032
WDR11	rs11592107	A	G	0.172	0.053	0.025	0.034	0.066	0.025	0.0079
MTNR1B	rs10830963	G	C	0.214	0.102	0.023	1.1x10 ⁻⁵	0.103	0.023	1.0x10 ⁻⁵
MTNR1B	rs11020308	A	C	0.305	-0.016	0.021	0.45	-0.017	0.021	0.40
ETS1	rs10893827	A	G	0.690	0.054	0.020	0.0081	0.049	0.021	0.018
ETS1	rs7104712	C	A	0.392	0.062	0.020	0.0016	0.058	0.019	0.0025
ETS1	rs11819995	T	C	0.205	0.024	0.023	0.31	0.018	0.024	0.45
CCND2	rs10848960	G	C	0.867	-0.014	0.028	0.63	-0.007	0.029	0.81
CCND2	rs3812821	G	C	0.656	0.071	0.022	0.0013	0.072	0.022	0.0012
CCND2	rs3217792	C	T	0.938	0.119	0.043	0.0053	0.110	0.043	0.010
CCND2	rs76895963	T	G	0.991	0.419	0.145	0.0039	0.383	0.147	0.0092
CCND2	rs78470967	T	A	0.985	0.169	0.085	0.046	0.165	0.084	0.051
HNF1A	rs1800574	T	C	0.013	0.276	0.089	0.0020	0.348	0.086	5.3x10 ⁻⁵
HNF1A	rs61953351	G	T	0.820	0.110	0.026	1.8x10 ⁻⁵	0.125	0.025	4.5x10 ⁻⁷
RASGRP1	rs28582094	G	A	0.382	0.056	0.019	0.0033	0.054	0.019	0.0044
RASGRP1	rs34715063	C	T	0.069	0.075	0.039	0.052	0.069	0.039	0.076
INFAM2	rs484943	T	C	0.214	0.005	0.024	0.84	0.007	0.023	0.78
INFAM2	rs3743140	A	G	0.171	0.065	0.025	0.010	0.065	0.025	0.010
GRP-MC4R	rs9957320	G	T	0.751	0.086	0.022	9.5x10 ⁻⁵	0.075	0.022	0.00056

<i>GRP-MC4R</i>	rs6567160	C	T	0.131	0.118	0.029	4.9x10 ⁻⁵	0.112	0.029	0.00012
<i>GRP-MC4R</i>	rs76227980	C	T	0.989	0.107	0.105	0.31	0.026	0.104	0.80
<i>HNF4A</i>	rs12625671	C	T	0.446	0.077	0.021	0.00022	0.080	0.021	0.00011
<i>HNF4A</i>	rs1800961	T	C	0.045	0.128	0.046	0.0054	0.138	0.046	0.0027

RAF: risk allele frequency. OR: odds-ratio. SE: standard error.

Supplementary Note Table 6. Assessment of the impact of BMI on heterogeneity in allelic effects at distinct T2D association signals from multi-ancestry meta-regression (MR-MEGA) of up to 166,070 cases and 1,132,773 controls^a.

Locus	Index SNV	Chr	Position (bp, b37)	Alleles		Heterogeneity <i>p</i> -value				BMI effect (SE) on log-OR of risk allele
				Risk	Other	Ancestry (unadjusted)	Ancestry (BMI adjusted)	BMI (ancestry adjusted)	Residual	
VWA5B1	rs10916784	1	20,729,451	G	C	0.27	0.25	0.55	0.71	0.00267 (0.00428)
MACF1	rs3768301	1	39,870,793	T	C	0.60	0.70	0.91	0.15	-0.00067 (0.00608)
MAST2	rs34444543	1	46,358,862	G	A	0.0063	0.66	0.39	0.89	-0.00399 (0.00420)
FAF1	rs12073283	1	51,219,188	C	G	0.34	0.94	0.47	0.77	-0.00562 (0.00737)
PGM1	rs11576729	1	64,114,429	G	T	0.042	0.042	0.85	0.83	-0.00099 (0.00497)
PTGFRN	rs1127215	1	117,532,790	C	T	0.088	0.19	0.58	0.41	0.00249 (0.00452)
NOTCH2	rs835576	1	120,455,586	C	T	0.037	0.081	0.85	0.18	-0.00127 (0.00707)
SEC16B	rs539515	1	177,889,025	C	A	0.52	0.50	0.66	0.88	-0.00242 (0.00504)
ZNF281	rs10919928	1	200,416,099	A	G	1.8x10 ⁻⁵	1.8x10 ⁻⁵	0.95	0.047	-0.00039 (0.00744)
DSTYK-MDM4	rs6689629	1	204,539,291	A	G	0.77	0.49	0.25	0.68	0.00601 (0.00503)
DSTYK-MDM4	rs12039805	1	205,107,793	A	G	0.19	0.43	0.29	0.14	0.00484 (0.00486)
SRGAP2	rs9429893	1	206,600,992	A	G	0.54	0.097	0.042	0.053	-0.00937 (0.00509)
PROX1	rs79687284	1	214,150,821	C	G	0.073	0.078	0.79	0.39	-0.00522 (0.01953)
PROX1	rs340874	1	214,159,256	C	T	0.40	0.36	0.62	0.72	0.00286 (0.00547)
LYPLAL1	rs2820446	1	219,748,818	C	G	0.029	0.11	0.71	0.20	-0.00179 (0.00515)
ABCB10-NUP133	rs348330	1	229,672,955	G	A	0.50	0.72	0.50	0.79	-0.00311 (0.00438)
TMEM18	rs62107261	2	422,144	T	C	0.29	0.34	0.79	0.47	0.00432 (0.01625)
TMEM18	rs10188334	2	653,874	C	T	0.12	0.094	0.11	0.56	0.01153 (0.00712)
DTNB	rs55928417	2	25,533,568	G	T	0.14	0.11	0.30	0.41	0.00481 (0.00469)
GCKR	rs1260326	2	27,730,940	C	T	0.91	0.93	0.48	0.0030	0.00321 (0.00543)
THADA	rs13414140	2	43,671,176	C	T	3.9x10 ⁻⁶	4.2x10 ⁻⁶	0.92	0.53	0.00076 (0.00734)
SIX3-SIX2	rs12712928	2	45,192,080	C	G	2.1x10 ⁻¹⁴	0.0063	0.45	0.66	-0.00394 (0.00501)
BNIP1	rs17049712	2	58,961,136	T	C	0.75	0.63	0.46	0.58	0.00361 (0.00477)
BCL11A	rs243018	2	60,586,707	G	C	0.83	0.87	0.95	0.80	-0.00027 (0.00414)
CEP68	rs2540949	2	65,284,231	A	T	0.26	0.46	0.97	0.45	-0.00014 (0.00437)
CEP68	rs6752053	2	65,666,674	T	C	0.91	0.92	0.70	0.041	0.00177 (0.00518)
GLI2	rs11688682	2	121,347,612	G	C	0.14	0.17	0.17	0.53	0.00883 (0.00633)
GLI2	rs10864859	2	121,440,218	T	G	0.056	0.044	0.43	0.77	0.00587 (0.00696)
CYTIP	rs7594480	2	158,390,468	T	C	0.92	0.51	0.073	0.58	-0.01747 (0.00953)
RBSM1	rs1020731	2	161,144,055	A	G	0.020	0.20	0.99	0.11	-0.00006 (0.00531)
KCNH7	rs12614955	2	163,649,480	T	C	0.64	0.52	0.41	0.19	-0.00378 (0.00481)
GRB14	rs10184004	2	165,508,389	C	T	0.045	0.19	0.93	0.065	-0.00043 (0.00551)
IKZF2	rs16849467	2	213,818,731	T	C	0.58	0.36	0.19	0.31	-0.00693 (0.00549)
IRS1	rs2943648	2	227,100,490	G	A	0.064	0.14	0.97	0.29	0.00020 (0.00528)
ATG16L1-DGKD	rs117809958	2	234,191,103	A	T	0.21	0.24	0.77	0.94	0.00949 (0.02744)
PPARG	rs17036160	3	12,329,783	C	T	0.89	0.82	0.31	0.99	-0.00742 (0.00606)

PPARG	rs4684855	3	12,490,951	T	C	0.32	0.60	0.97	0.28	0.00018 (0.00494)
UBE2E2	rs13094957	3	23,457,080	T	C	0.0023	0.43	0.96	0.053	-0.00026 (0.00577)
UBE2E2	rs76435632	3	23,632,174	G	C	0.92	0.75	0.39	0.29	-0.00790 (0.00947)
RBM6	rs2624847	3	50,174,197	G	T	0.00043	0.00046	0.42	0.48	0.00445 (0.00548)
CACNA2D3	rs76263492	3	54,828,827	T	G	0.40	0.29	0.32	0.39	0.01415 (0.01436)
PXK	rs12629058	3	58,338,809	T	C	0.074	0.024	0.081	0.74	0.00858 (0.00469)
PSMD6-ADAMTS9	rs2292662	3	63,897,215	C	T	0.0087	0.38	0.80	0.28	-0.00132 (0.00545)
PSMD6-ADAMTS9	rs66815886	3	64,703,394	G	T	0.020	0.060	0.40	0.60	-0.00399 (0.00465)
ZBTB20	rs1459513	3	114,960,798	C	A	0.085	0.073	0.35	0.16	-0.00545 (0.00626)
ADCY5	rs11708067	3	123,065,778	A	G	0.16	0.095	0.26	0.96	-0.00704 (0.00541)
SLC12A8	rs9873519	3	124,921,457	T	C	0.55	0.29	0.10	0.99	-0.00723 (0.00372)
MBNL1	rs1426385	3	151,998,053	A	G	0.18	0.043	0.065	0.32	-0.00785 (0.00438)
MBNL1	rs10935897	3	152,399,693	A	G	0.23	0.20	0.067	0.76	0.00793 (0.00410)
MBNL1	rs75417759	3	152,530,027	C	T	2.7x10 ⁻⁷	0.0097	0.97	0.56	-0.00035 (0.01079)
SLC2A2	rs8192675	3	170,724,883	T	C	9.6x10 ⁻⁵	4.9x10 ⁻⁵	0.056	0.87	0.00927 (0.00447)
IGF2BP2	rs7633675	3	185,510,613	G	T	0.16	0.57	0.53	0.89	-0.00285 (0.00417)
ST6GAL1	rs3887925	3	186,665,645	T	C	0.079	0.25	0.95	0.83	-0.00026 (0.00365)
ST6GAL1	rs9799068	3	186,676,455	A	C	0.099	0.32	0.87	0.081	0.00084 (0.00568)
BCL6-LPP	rs4686471	3	187,740,899	C	T	0.061	0.044	0.35	0.87	0.00516 (0.00507)
TFRC	rs74289356	3	195,825,077	T	C	0.50	0.46	0.59	0.71	-0.00304 (0.00548)
CTBP1-PCGF3-MAEA	rs73221123	4	726,202	T	C	0.85	0.88	0.86	0.88	0.00189 (0.00965)
CTBP1-PCGF3-MAEA	rs730831	4	1,240,299	T	G	0.045	0.051	0.27	0.079	-0.00702 (0.00697)
CTBP1-PCGF3-MAEA	rs6831006	4	1,784,605	G	C	0.66	0.86	0.86	0.41	0.00085 (0.00477)
WFS1	rs9998835	4	6,293,237	G	C	2.5x10 ⁻⁶	0.00017	0.56	0.14	-0.00293 (0.00542)
LCORL	rs6855926	4	18,047,401	A	G	0.17	0.22	0.78	0.39	-0.00132 (0.00479)
GNPDA2	rs13130484	4	45,175,691	T	C	0.31	0.11	0.086	0.75	-0.00770 (0.00427)
MOB1B	rs7674402	4	71,835,822	A	G	0.67	0.84	0.55	0.29	-0.00412 (0.00708)
SCD5	rs10471048	4	83,587,562	G	C	0.0068	0.017	0.62	0.80	0.00221 (0.00416)
NKX6-1-CDS1	rs117624659	4	85,339,618	T	C	0.89	0.80	0.29	0.74	0.03403 (0.03007)
SMARCAD1	rs6821438	4	95,091,911	A	G	8.2x10 ⁻⁵	1.4x10 ⁻⁵	0.056	0.36	-0.00843 (0.00450)
PPP3CA	rs2659518	4	102,135,363	A	G	0.24	0.67	0.11	0.72	0.00927 (0.00557)
SLC9B1	rs223423	4	103,725,894	G	A	3.8x10 ⁻⁵	0.0028	0.53	0.075	-0.00271 (0.00469)
TET2	rs17035289	4	106,048,291	C	T	0.044	0.0096	0.058	0.48	-0.01003 (0.00528)
TMEM154	rs6813195	4	153,520,475	C	T	0.49	0.36	0.23	0.045	0.00544 (0.00504)
PDGFC	rs1425482	4	157,725,916	T	C	5.6x10 ⁻⁵	0.12	0.31	8.8E-01	0.00482 (0.00435)
ACSL1	rs1996546	4	185,714,289	G	T	0.14	0.34	0.026	6.7E-01	-0.01755 (0.00754)
ANKH	rs147581833	5	14,755,919	C	T	0.38	0.39	0.29	2.0E-01	-0.05915 (0.06125)
ANKH	rs30614	5	14,780,521	A	G	0.40	0.27	0.25	6.3E-01	-0.00740 (0.00608)
MRPS30	rs6884702	5	44,682,589	G	A	0.0022	0.0016	0.38	4.2E-02	-0.00389 (0.00491)
ITGA1	rs17261179	5	51,791,225	T	C	0.77	0.66	0.44	8.5E-02	-0.00342 (0.00487)
ARL15	rs702634	5	53,271,420	A	G	0.89	0.90	0.86	5.3E-01	0.00088 (0.00485)
ARL15	rs6876198	5	53,303,595	C	T	0.17	0.61	0.51	5.5E-01	0.00292 (0.00442)
ANKRD55	rs256904	5	55,810,305	T	A	0.78	0.77	0.83	1.3E-01	-0.00095 (0.00479)

ANKRD55	rs42251	5	55,840,633	A	G	0.30	0.33	0.84	8.5E-01	0.00085 (0.00400)
ANKRD55	rs3936510	5	55,860,866	T	G	0.0033	0.00098	0.11	4.4E-01	0.00862 (0.00540)
PIK3R1	rs57634870	5	67,716,793	G	T	0.93	0.93	0.92	8.6E-01	-0.00052 (0.00499)
HMGCR-POC5	rs2307111	5	75,003,678	T	C	9.7x10 ⁻⁷	0.051	0.93	2.1E-01	-0.00038 (0.00464)
ZBED3	rs7732130	5	76,435,004	G	A	0.15	0.073	0.19	1.9E-03	-0.00734 (0.00677)
DMGDH	rs10052346	5	78,472,599	G	T	0.0056	0.36	0.77	5.3E-01	0.00129 (0.00434)
SLCO6A1-PAM	rs78408340	5	102,338,739	G	C	0.28	0.34	0.97	5.9E-01	0.00156 (0.03690)
SLCO6A1-PAM	rs115505614	5	102,422,968	T	C	0.77	0.77	0.92	3.0E-02	-0.00163 (0.02166)
SLCO6A1-PAM	rs186327337	5	103,364,257	G	A	0.012	0.027	0.10	2.8E-01	0.07139 (0.04685)
CEP120	rs4267865	5	122,704,342	G	T	0.18	0.27	0.58	5.0E-01	-0.00475 (0.00865)
PHF15	rs329122	5	133,864,599	A	G	0.30	0.42	0.71	8.0E-01	0.00164 (0.00410)
NSD1	rs244708	5	176,589,585	G	A	0.22	0.24	0.74	5.2E-01	0.00152 (0.00450)
SSR1-RREB1	rs77630070	6	7,196,323	G	T	0.40	0.62	1.0	9.9E-01	0.00003 (0.00609)
SSR1-RREB1	rs9379084	6	7,231,843	G	A	0.0058	0.65	0.43	4.8E-01	0.00565 (0.00716)
CDKAL1	rs9348441	6	20,680,678	A	T	9.4x10 ⁻¹⁴	0.22	3.0x10 ⁻⁶	6.7E-04	-0.02157 (0.00563)
MHC region	rs879882	6	31,139,452	C	T	0.032	0.021	0.34	1.5E-01	0.00519 (0.00597)
MHC region	rs3806155	6	32,373,378	T	A	0.085	0.029	0.10	6.5E-01	0.03172 (0.01850)
MHC region	rs7452864	6	32,439,077	C	T	0.00012	0.055	0.88	5.6E-01	-0.00089 (0.00570)
MHC region	rs62405954	6	33,524,820	T	C	0.34	0.46	0.60	6.9E-01	0.00612 (0.01104)
MHC region	rs4711389	6	34,214,670	A	G	0.00039	0.00067	0.42	6.2E-01	0.00914 (0.01100)
ZFAND3-KCNK16-GLP1R	rs2281342	6	38,992,668	T	C	0.073	0.18	0.72	4.6E-01	-0.00183 (0.00510)
ZFAND3-KCNK16-GLP1R	rs742762	6	39,046,644	A	C	1.1x10 ⁻⁷	7.9x10 ⁻⁵	0.39	3.0E-01	-0.00467 (0.00567)
ZFAND3-KCNK16-GLP1R	rs3734618	6	39,284,184	G	A	0.00042	0.019	0.68	9.8E-01	0.00177 (0.00369)
LRFN2	rs34298980	6	40,409,243	T	C	0.27	0.28	0.12	5.0E-01	0.00699 (0.00454)
VEGFA	rs6905288	6	43,758,873	A	G	0.89	0.89	0.70	5.7E-01	-0.00178 (0.00462)
VEGFA	rs6458354	6	43,814,190	C	T	0.13	0.16	0.87	9.4E-01	0.00087 (0.00469)
TFAP2B	rs3798519	6	50,788,778	C	A	0.78	0.70	0.49	2.3E-01	-0.00358 (0.00542)
BEND3	rs1665901	6	107,433,400	A	T	0.18	0.53	0.25	2.0E-01	-0.00600 (0.00545)
NUS1	rs72951506	6	118,011,723	C	T	0.72	0.77	0.69	2.0E-01	-0.00223 (0.00587)
CENPW-SOGA3	rs11759026	6	126,792,095	G	A	0.12	0.12	0.41	7.6E-01	-0.00441 (0.00512)
CENPW-SOGA3	rs2800733	6	127,416,930	A	G	0.79	0.83	0.16	0.40	-0.00799 (0.00581)
MED23-ENPP3	rs7739842	6	131,954,797	G	T	0.0067	0.063	0.47	0.47	0.00344 (0.00483)
SLC35D3	rs6937795	6	137,291,281	A	C	0.065	0.085	0.27	0.81	-0.00472 (0.00405)
REPS1	rs9376353	6	138,855,975	A	T	0.10	0.11	0.66	0.44	0.00195 (0.00446)
HIVEP2	rs6570526	6	143,058,692	G	C	0.59	0.34	0.23	0.14	-0.00520 (0.00463)
RGS17	rs6932473	6	153,438,573	T	A	0.50	0.53	0.67	0.12	0.00190 (0.00482)
SLC22A3	rs539298	6	160,770,360	A	G	0.22	0.37	0.72	0.86	0.00159 (0.00405)
QKI	rs4709746	6	164,133,001	C	T	0.54	0.49	0.60	0.85	-0.00357 (0.00626)
ETV1	rs12154701	7	13,887,008	A	C	0.020	0.084	0.92	0.32	0.00044 (0.00447)
DGKB	rs17168486	7	14,898,282	T	C	0.27	0.29	0.60	0.20	-0.00252 (0.00509)
DGKB	rs2215383	7	15,062,983	C	T	0.92	0.75	0.35	0.017	-0.00409 (0.00502)
JAZF1	rs849133	7	28,192,280	C	T	0.0015	0.0030	0.76	0.76	-0.00155 (0.00467)
JAZF1	rs552707	7	28,205,303	T	C	6.5x10 ⁻⁵	0.00024	0.73	0.78	-0.00106 (0.00285)

JAZF1	rs10226758	7	28,214,614	C	A	4.4x10 ⁻¹³	3.7x10 ⁻¹²	0.82	0.58	-0.00077 (0.00325)
CRHR2	rs917195	7	30,728,452	C	T	0.61	0.24	0.11	0.69	-0.00826 (0.00503)
GCK	rs882019	7	44,178,829	G	A	0.31	0.22	0.33	0.077	-0.00418 (0.00471)
GCK	rs878521	7	44,255,643	A	G	0.046	0.094	0.63	0.0016	-0.00224 (0.00567)
GRB10	rs13236710	7	50,809,085	G	A	0.10	0.10	1.0	0.93	0.00001 (0.00604)
AUTS2	rs2533457	7	69,055,951	G	A	0.00078	0.0032	0.92	0.85	0.00044 (0.00415)
STEAP1	rs6978118	7	89,800,241	A	T	0.00019	0.0053	0.25	0.56	-0.00511 (0.00441)
FBXL13-RELN-RASA4	rs7781557	7	102,481,891	C	T	0.56	0.38	0.31	0.77	-0.00732 (0.00675)
GCC1-PAX4-LEP	rs12669223	7	127,250,831	A	G	0.040	0.16	0.59	0.39	-0.00898 (0.01696)
KLF14	rs1562396	7	130,457,914	G	A	0.0015	0.0022	0.97	0.80	0.00018 (0.00430)
BRAF	rs11983228	7	140,631,823	C	G	0.42	0.56	0.59	0.60	0.00385 (0.00706)
AOC1	rs62492368	7	150,537,635	A	G	0.064	0.059	0.61	0.91	0.00227 (0.00401)
MNX1	rs887609	7	156,794,983	A	G	7.0x10 ⁻⁵	0.00066	0.90	0.91	0.00071 (0.00508)
MNX1	rs2366214	7	156,992,461	A	G	0.18	0.77	0.64	0.10	0.00207 (0.00481)
MSRA-XKR6	rs4240673	8	10,787,612	T	C	0.14	0.051	0.12	0.38	-0.00769 (0.00499)
LONRF1	rs12680692	8	12,618,225	A	T	0.0088	0.13	0.24	0.044	0.00577 (0.00553)
LPL	rs7819706	8	19,844,415	A	G	0.054	0.051	0.17	0.77	-0.00923 (0.00642)
KCNU1	rs10092900	8	36,854,711	G	T	0.29	0.28	0.72	0.47	0.00181 (0.00504)
KCNU1	rs12680217	8	37,397,803	T	C	0.015	0.013	0.38	0.49	-0.00512 (0.00578)
ANK1	rs12550613	8	41,510,260	C	G	0.25	0.33	0.24	0.26	-0.00474 (0.00421)
ANK1	rs508419	8	41,522,991	G	A	0.52	0.93	0.22	0.24	-0.00618 (0.00522)
GDAP1	rs3780012	8	75,147,209	C	G	0.18	0.54	0.32	0.40	-0.02392 (0.02446)
TP53INP1	rs13257021	8	95,965,695	A	G	0.065	0.098	0.91	0.21	-0.00050 (0.00457)
TRPS1	rs800909	8	116,497,173	T	C	0.014	0.15	0.87	0.39	-0.00077 (0.00477)
SLC30A8	rs13266634	8	118,184,783	C	T	0.23	0.93	0.63	0.040	-0.00226 (0.00529)
PVT1	rs4733612	8	129,569,999	G	A	0.083	0.11	0.056	0.61	-0.01044 (0.00534)
BOP1	rs3890400	8	145,544,720	A	G	0.88	0.91	0.74	0.85	-0.00151 (0.00431)
BOP1	rs7014773	8	145,972,670	T	C	0.83	0.72	0.33	0.074	0.00442 (0.00496)
GLIS3	rs4237150	9	4,290,085	C	G	0.23	0.30	0.40	0.39	0.00352 (0.00424)
GLIS3	rs4258054	9	4,297,892	T	C	0.14	0.12	0.52	0.60	0.00299 (0.00454)
HAUS6	rs12380322	9	19,074,538	G	A	0.24	0.36	0.98	0.54	-0.00009 (0.00450)
CDKN2A-CDKN2B	rs7856455	9	21,840,834	G	T	0.0068	0.0073	0.64	0.44	-0.00347 (0.00757)
CDKN2A-CDKN2B	rs10757282	9	22,133,984	C	T	0.0051	0.00042	0.010	0.28	-0.01006 (0.00407)
CDKN2A-CDKN2B	rs10811661	9	22,134,094	T	C	0.0055	0.00013	0.0048	0.0032	-0.01357 (0.00575)
CDKN2A-CDKN2B	rs1575972	9	22,301,092	T	A	0.090	0.45	0.22	0.75	-0.01324 (0.01018)
LINGO2	rs1412234	9	28,410,683	C	T	0.012	0.092	0.34	0.66	0.00479 (0.00491)
UBAP2	rs12001437	9	34,074,476	C	T	0.058	0.031	0.24	0.19	0.00522 (0.00466)
TLE4	rs13290396	9	81,914,978	C	T	0.77	0.76	0.77	0.039	0.00219 (0.00854)
TLE1	rs9332453	9	83,998,346	C	T	0.0068	0.92	0.12	0.25	-0.00756 (0.00504)
TLE1	rs2796441	9	84,308,948	G	A	0.073	0.18	0.067	0.37	-0.00810 (0.00451)
ZNF169	rs12345069	9	96,971,175	C	T	0.35	0.55	0.012	0.52	0.01388 (0.00549)
PTCH1	rs113154802	9	98,278,413	C	T	0.00061	0.00069	0.62	0.41	0.00390 (0.00795)
STRBP	rs2416899	9	126,015,103	T	G	0.0081	0.017	0.35	0.20	-0.00502 (0.00563)

ABO	rs505922	9	136,149,229	C	T	0.12	0.044	0.14	0.77	0.00660 (0.00418)
GPSM1	rs28429551	9	139,243,334	A	T	0.0051	0.18	0.62	0.080	-0.00290 (0.00636)
GPSM1	rs74604683	9	139,247,229	C	T	0.32	0.13	0.083	0.12	-0.01238 (0.00772)
CDC123-CAMK1D	rs11257655	10	12,307,894	T	C	0.016	0.0035	0.040	0.46	0.00987 (0.00482)
MYO3A	rs7923442	10	26,497,704	A	G	0.37	0.11	0.028	0.99	-0.01114 (0.00432)
JMJD1C	rs41274074	10	64,974,380	G	C	0.23	0.32	0.89	0.21	-0.00118 (0.00864)
VPS26A-NEUROG3	rs190925	10	71,320,943	A	G	1.1x10 ⁻⁵	0.0066	0.70	0.65	-0.00199 (0.00502)
VPS26A-NEUROG3	rs41277236	10	71,332,301	T	C	0.41	0.44	0.83	0.64	-0.00319 (0.01459)
VPS26A-NEUROG3	rs2642588	10	71,466,578	G	T	0.0010	0.40	0.22	0.41	-0.00706 (0.00589)
ZNF503-LRMDA	rs3012060	10	77,244,336	T	A	0.020	0.00021	0.0018	0.84	-0.01880 (0.00559)
ZMIZ1	rs703980	10	80,943,841	G	A	0.29	0.10	0.088	0.32	-0.00745 (0.00450)
PTEN	rs10887775	10	89,766,368	A	G	0.23	0.24	0.92	0.68	0.00054 (0.00522)
HHEX-IDE	rs10882099	10	94,460,650	T	C	1.5x10 ⁻¹⁴	2.2x10 ⁻¹¹	0.55	0.64	-0.00160 (0.00261)
HHEX-IDE	rs139027698	10	94,468,247	T	C	5.8x10 ⁻¹⁰	2.5x10 ⁻⁸	0.20	0.0061	-0.01532 (0.01411)
HHEX-IDE	rs1112718	10	94,479,107	A	G	2.8x10 ⁻⁶	0.00010	0.15	0.17	0.00383 (0.00282)
ARHGAP19-SLIT1	rs10748694	10	99,056,190	A	T	0.0098	0.55	0.11	0.37	-0.00717 (0.00455)
BBIP1	rs7067540	10	112,621,837	C	T	0.097	0.19	0.48	0.15	0.00322 (0.00491)
TCF7L2	rs12243296	10	114,344,288	G	A	0.026	0.35	0.39	0.84	0.00512 (0.00530)
TCF7L2	rs7100404	10	114,381,965	C	T	0.0043	0.054	0.33	0.54	0.00552 (0.00556)
TCF7L2	rs2859885	10	114,428,364	C	T	2.0x10 ⁻⁷	2.4x10 ⁻⁷	0.90	0.38	-0.00096 (0.00748)
TCF7L2	rs10787461	10	114,552,267	G	A	0.0086	0.0088	0.98	0.15	0.00016 (0.00708)
TCF7L2	rs2104598	10	114,715,598	G	A	0.00017	0.00029	0.16	0.071	-0.00843 (0.00701)
TCF7L2	rs114322470	10	114,736,670	T	G	0.80	0.77	0.047	0.50	0.05380 (0.02693)
TCF7L2	rs7903146	10	114,758,349	T	C	0.00043	8.1x10 ⁻⁵	0.053	3.0x10 ⁻⁹	-0.01210 (0.01048)
TCF7L2	rs7076754	10	114,797,893	G	A	0.0048	0.023	0.24	0.13	0.00993 (0.00944)
TCF7L2	rs145003494	10	114,834,411	A	G	0.00018	7.9x10 ⁻⁵	0.044	0.52	-0.06314 (0.03099)
TCF7L2	rs116929578	10	114,836,181	G	A	0.81	0.82	0.84	0.031	-0.00334 (0.02006)
TCF7L2	rs7081841	10	114,859,416	G	C	0.0019	0.0020	0.21	0.41	-0.00817 (0.00667)
TCF7L2	rs12257761	10	115,016,408	T	C	6.2x10 ⁻⁵	0.00018	0.47	0.094	-0.00636 (0.00993)
TCF7L2	rs11196296	10	115,069,951	T	C	1.3x10 ⁻¹¹	4.8x10 ⁻¹¹	0.83	0.50	0.00394 (0.01872)
TCF7L2	rs7093035	10	115,119,864	G	A	0.98	0.91	0.49	0.87	-0.00831 (0.01058)
TCF7L2	rs72830009	10	115,136,540	G	A	0.025	0.015	0.090	0.55	0.03716 (0.02147)
TCF7L2	rs11596522	10	115,247,447	T	G	0.0052	0.0047	0.64	0.12	0.00511 (0.01229)
WDR11	rs11199753	10	122,834,572	G	T	0.037	0.19	0.16	0.64	-0.00938 (0.00653)
WDR11	rs2172073	10	122,909,625	A	C	0.52	0.79	0.88	0.56	0.00089 (0.00574)
WDR11	rs11592107	10	122,968,964	A	G	0.57	0.71	0.76	0.67	-0.00145 (0.00457)
PLEKHA1	rs2421016	10	124,167,512	C	T	0.00092	0.0031	0.26	0.62	-0.00483 (0.00422)
INS-IGF2-KCNQ1	rs76547628	11	2,077,271	T	C	1.4x10 ⁻¹¹	0.00024	0.34	0.14	0.00575 (0.00650)
INS-IGF2-KCNQ1	rs10770142	11	2,194,420	G	C	0.21	0.21	0.61	0.077	-0.00270 (0.00584)
INS-IGF2-KCNQ1	rs4930050	11	2,235,129	G	A	1.9x10 ⁻⁸	0.018	0.82	0.55	0.00257 (0.01104)
INS-IGF2-KCNQ1	rs800125	11	2,364,549	A	C	2.1x10 ⁻⁹	2.2x10 ⁻⁹	0.54	0.13	0.00276 (0.00494)
INS-IGF2-KCNQ1	rs79495865	11	2,375,458	G	A	9.7x10 ⁻⁹	0.0065	0.47	0.54	0.00458 (0.00632)
INS-IGF2-KCNQ1	rs2283164	11	2,579,163	A	G	0.18	0.18	0.35	0.35	0.00981 (0.01077)

<i>INS-IGF2-KCNQ1</i>	rs80102379	11	2,634,177	G	T	0.14	0.23	0.10	0.29	0.02756 (0.01763)
<i>INS-IGF2-KCNQ1</i>	rs151215	11	2,681,072	G	A	1.8x10 ⁻¹¹	1.2x10 ⁻⁵	0.47	0.82	0.00380 (0.00486)
<i>INS-IGF2-KCNQ1</i>	rs231361	11	2,691,500	A	G	1.8x10 ⁻⁵	0.17	0.36	0.15	0.00416 (0.00497)
<i>INS-IGF2-KCNQ1</i>	rs2237884	11	2,799,679	T	C	6.2x10 ⁻⁶	9.0x10 ⁻⁶	0.78	0.24	-0.00136 (0.00515)
<i>INS-IGF2-KCNQ1</i>	rs4930011	11	2,856,658	G	C	4.5x10 ⁻⁵	0.081	0.096	0.14	-0.00641 (0.00418)
<i>INS-IGF2-KCNQ1</i>	rs234866	11	2,857,897	G	A	0.019	0.096	0.60	0.047	0.00228 (0.00489)
<i>INS-IGF2-KCNQ1</i>	rs2237897	11	2,858,546	C	T	7.1x10 ⁻⁹	0.034	0.66	0.081	0.00264 (0.00673)
<i>INS-IGF2-KCNQ1</i>	rs445084	11	2,908,754	G	A	0.047	0.061	0.41	0.62	-0.00420 (0.00500)
<i>TRIM66</i>	rs10769936	11	8,654,528	C	T	0.066	0.061	0.47	0.036	0.00323 (0.00503)
<i>KCNJ11-ABCC8</i>	rs5215	11	17,408,630	C	T	0.24	0.30	0.35	0.15	0.00418 (0.00480)
<i>BDNF</i>	rs4923464	11	27,683,618	C	T	0.0018	0.012	0.99	0.46	0.00004 (0.00501)
<i>QSER1</i>	rs145678014	11	32,927,778	G	T	0.040	0.077	0.85	0.76	0.00300 (0.01524)
<i>HSD17B12</i>	rs6485462	11	43,816,200	C	T	0.00065	0.0051	0.54	0.25	-0.00280 (0.00473)
<i>CRY2</i>	rs12419690	11	45,858,584	G	A	0.52	0.49	0.72	0.41	-0.00162 (0.00454)
<i>FOLH1</i>	rs6485981	11	49,477,266	T	C	0.0019	0.0014	0.42	0.50	0.00502 (0.00619)
<i>MAP3K11</i>	rs12789028	11	65,326,154	A	G	0.35	0.43	0.80	1.0	-0.00150 (0.00501)
<i>TPCN2-CCND1</i>	rs3918298	11	69,463,273	G	A	0.054	0.41	0.18	0.97	-0.01651 (0.01073)
<i>CENTD2</i>	rs77464186	11	72,460,398	A	C	0.076	0.36	0.094	0.10	-0.01083 (0.00701)
<i>C11orf30</i>	rs61894507	11	76,156,973	G	A	0.057	0.0089	0.018	0.49	0.01220 (0.00514)
<i>MTNR1B</i>	rs10830963	11	92,708,710	G	C	7.6x10 ⁻⁶	0.12	0.87	3.6x10 ⁻⁶	0.00081 (0.00638)
<i>MTNR1B</i>	rs11020308	11	93,131,667	A	C	0.18	0.074	0.043	0.036	-0.01039 (0.00577)
<i>ETS1</i>	rs10893827	11	128,040,810	A	G	0.079	0.23	0.36	0.87	0.00484 (0.00487)
<i>ETS1</i>	rs7104712	11	128,235,252	C	A	0.13	0.13	0.99	0.96	0.00005 (0.00430)
<i>ETS1</i>	rs11819995	11	128,389,391	T	C	0.66	0.24	0.075	0.36	-0.00930 (0.00533)
<i>CCND2</i>	rs10848960	12	4,033,222	G	C	0.046	0.072	0.75	0.25	0.00250 (0.00822)
<i>CCND2</i>	rs3812821	12	4,382,324	G	C	0.041	0.033	0.45	0.74	0.00519 (0.00647)
<i>CCND2</i>	rs3217792	12	4,384,696	C	T	0.00070	0.0028	0.85	0.79	-0.00214 (0.01066)
<i>CCND2</i>	rs76895963	12	4,384,844	T	G	0.00034	0.00014	0.16	0.31	-0.04356 (0.03247)
<i>CCND2</i>	rs78470967	12	4,521,511	T	A	0.95	0.85	0.40	0.99	-0.01581 (0.01457)
<i>CDKN1B</i>	rs2066827	12	12,871,099	G	T	0.94	0.98	0.58	0.81	0.00338 (0.00576)
<i>ITPR2</i>	rs10842708	12	26,474,867	G	A	0.026	0.065	0.73	0.81	0.00164 (0.00441)
<i>KLHDC5</i>	rs12578595	12	27,964,996	C	T	0.13	0.17	0.86	0.88	0.00095 (0.00480)
<i>FAM60A</i>	rs78345706	12	31,417,019	A	G	0.57	0.43	0.24	0.47	-0.01136 (0.00972)
<i>PKP2-SYT10</i>	rs6488140	12	33,370,406	A	G	0.0015	0.083	0.36	0.46	-0.00466 (0.00510)
<i>FAIM2</i>	rs7132908	12	50,263,148	A	G	0.15	0.23	0.88	0.16	-0.00072 (0.00497)
<i>HMGA2</i>	rs343093	12	66,255,005	G	C	0.059	0.76	0.53	0.68	0.00312 (0.00481)
<i>HMGA2</i>	rs7970350	12	66,360,164	T	C	0.13	0.18	0.81	0.032	0.00108 (0.00507)
<i>TSPAN8</i>	rs7313668	12	71,449,521	T	G	0.052	0.033	0.27	0.57	0.00508 (0.00451)
<i>RMST</i>	rs7972074	12	97,851,611	C	T	0.030	0.031	0.87	0.63	-0.00086 (0.00499)
<i>WSCD2</i>	rs1426371	12	108,629,780	G	A	0.96	0.72	0.29	0.60	-0.00515 (0.00477)
<i>SH2B3-ALDH2-BRAP</i>	rs3782886	12	112,110,489	T	C	4.0x10 ⁻⁸	3.0x10 ⁻⁸	0.11	2.9x10 ⁻⁶	0.01856 (0.01670)
<i>PTPN11-HECTD4</i>	rs77753011	12	113,117,897	G	T	2.7x10 ⁻⁸	2.3x10 ⁻⁸	0.14	0.00033	0.02023 (0.01862)
<i>KSR2</i>	rs34965774	12	118,412,373	A	G	0.35	0.91	0.20	0.92	-0.00691 (0.00485)

<i>HNF1A</i>	rs1800574	12	121,416,864	T	C	0.00051	0.00087	0.23	0.17	-0.01434 (0.01283)
<i>HNF1A</i>	rs61953351	12	121,456,616	G	T	2.0x10 ⁻⁶	1.7x10 ⁻⁶	0.55	0.00017	0.00342 (0.00758)
<i>MPHOSPH9-ZNF664</i>	rs1790116	12	123,618,544	T	G	0.046	0.10	0.79	0.71	0.00156 (0.00551)
<i>MPHOSPH9-ZNF664</i>	rs2451321	12	124,545,435	C	G	0.28	0.31	0.51	0.48	0.00292 (0.00443)
<i>FBRSL1</i>	rs12811407	12	133,069,698	A	G	0.37	0.39	0.55	0.00010	-0.00304 (0.00648)
<i>SGCG</i>	rs314879	13	23,309,382	C	T	0.32	0.38	0.99	0.63	-0.00009 (0.00539)
<i>RNF6</i>	rs34584161	13	26,776,999	A	G	0.067	0.38	0.22	0.58	-0.00598 (0.00485)
<i>KL</i>	rs2858980	13	33,554,587	G	A	0.12	0.67	0.024	0.20	-0.01224 (0.00570)
<i>DLEU1</i>	rs963740	13	51,096,095	A	T	0.84	0.50	0.20	0.079	0.00603 (0.00514)
<i>OLFM4</i>	rs9568868	13	54,107,583	T	G	0.91	0.99	0.47	0.23	-0.00411 (0.00592)
<i>SPRY2</i>	rs1215468	13	80,707,429	A	G	0.44	0.50	0.27	0.20	-0.00539 (0.00515)
<i>MIR17HG</i>	rs34165267	13	91,942,919	C	T	0.00012	0.012	0.42	0.77	-0.00433 (0.00510)
<i>AKAP6</i>	rs12883788	14	33,303,540	T	C	0.46	0.74	0.69	0.0017	-0.00180 (0.00545)
<i>CLEC14A</i>	rs2183237	14	38,803,756	G	A	0.37	0.27	0.20	0.31	0.00576 (0.00465)
<i>NRXN3</i>	rs8008910	14	79,944,099	A	G	0.62	0.23	0.091	0.34	-0.01089 (0.00662)
<i>DLK1-MEG3</i>	rs12878003	14	101,124,721	G	A	0.022	0.088	0.58	0.030	-0.00287 (0.00586)
<i>DLK1-MEG3</i>	rs73347525	14	101,255,172	A	G	0.66	0.45	0.30	0.58	0.00593 (0.00563)
<i>DLK1-MEG3</i>	rs1053900	14	101,301,866	C	T	0.14	0.064	0.10	0.12	-0.00711 (0.00473)
<i>TRAF3</i>	rs11160699	14	103,252,270	A	G	0.93	0.93	1.0	0.92	0.00001 (0.00482)
<i>RASGRP1</i>	rs28582094	15	38,843,887	G	A	0.20	0.039	0.042	0.054	-0.01042 (0.00576)
<i>RASGRP1</i>	rs34715063	15	38,873,115	C	T	0.86	0.86	0.80	0.11	0.00227 (0.00969)
<i>INFAM2</i>	rs484943	15	40,398,754	T	C	0.26	0.88	0.82	0.033	-0.00114 (0.00549)
<i>INFAM2</i>	rs3743140	15	40,616,742	A	G	0.00015	0.0034	0.66	0.93	0.00251 (0.00520)
<i>LTK</i>	rs1473781	15	41,818,917	A	G	0.0019	0.053	0.63	0.95	-0.00231 (0.00429)
<i>MYO5C</i>	rs3825801	15	52,517,714	C	T	0.058	0.056	0.64	0.30	-0.00299 (0.00661)
<i>C2CD4A-C2CD4B</i>	rs7163757	15	62,391,608	C	T	0.00026	0.18	0.0016	0.11	-0.01378 (0.00472)
<i>USP3</i>	rs7178762	15	63,871,292	C	T	0.63	0.48	0.36	0.45	0.00439 (0.00484)
<i>MAP2K5</i>	rs4776970	15	68,080,886	A	T	0.040	0.19	0.55	0.33	-0.00275 (0.00468)
<i>PTPN9-SIN3A</i>	rs11636031	15	75,815,758	T	C	0.47	0.51	0.90	0.044	0.00060 (0.00524)
<i>HMG20A</i>	rs952472	15	77,776,562	C	A	0.79	0.81	0.97	0.0090	-0.00015 (0.00517)
<i>AP3S2</i>	rs6496609	15	90,379,632	C	A	0.057	0.13	0.62	0.22	-0.00244 (0.00510)
<i>PRC1</i>	rs2890156	15	91,513,157	A	T	0.39	0.41	0.45	0.91	0.00385 (0.00460)
<i>RGMA</i>	rs7167984	15	93,832,067	G	A	0.25	0.51	0.54	0.11	-0.00335 (0.00586)
<i>ITFG3</i>	rs6600191	16	295,795	T	C	0.22	0.33	0.75	0.23	0.00156 (0.00521)
<i>CLUAP1-SLX4</i>	rs12445430	16	3,613,126	T	C	0.89	0.57	0.20	0.38	-0.00665 (0.00529)
<i>FAM57B</i>	rs11642430	16	30,045,789	G	C	0.023	0.040	0.44	0.88	-0.00340 (0.00406)
<i>FTO</i>	rs55872725	16	53,809,123	T	C	0.042	0.055	0.91	0.020	-0.00053 (0.00545)
<i>NFAT5</i>	rs862320	16	69,651,866	C	T	0.11	0.12	0.34	0.28	0.00455 (0.00495)
<i>ZFH3</i>	rs6416749	16	73,100,308	C	T	0.010	0.13	0.99	0.11	0.00004 (0.00553)
<i>BCAR1</i>	rs72802358	16	75,243,657	G	C	1.8x10 ⁻⁶	0.0038	0.047	0.54	-0.01511 (0.00752)
<i>CMIP</i>	rs2925979	16	81,534,790	T	C	0.43	0.37	0.35	0.24	-0.00438 (0.00491)
<i>ZFPM1</i>	rs9937296	16	88,554,480	C	T	0.13	0.058	0.13	0.63	-0.00883 (0.00568)
<i>SPG7</i>	rs12920022	16	89,564,055	A	T	0.044	0.023	0.20	0.22	0.00802 (0.00664)

ZZEF1	rs1043246	17	3,828,086	G	C	0.65	0.67	0.89	0.22	-0.00093 (0.00705)
ZZEF1	rs8071043	17	3,988,451	C	T	0.0021	0.029	0.45	0.90	-0.00407 (0.00481)
SLC16A11-SLC16A13	rs113748381	17	6,953,155	A	G	0.0020	0.00013	0.0024	0.074	-0.03054 (0.01103)
RAI1	rs1108646	17	17,751,478	A	G	0.068	0.0024	0.0065	0.15	0.01319 (0.00519)
NF1	rs1048317	17	29,704,002	T	C	0.41	0.15	0.11	0.43	0.00704 (0.00443)
HNF1B	rs3094515	17	36,043,653	C	T	0.33	0.15	0.13	0.57	-0.00742 (0.00476)
HNF1B	rs12449654	17	36,056,076	C	G	8.7x10 ⁻⁶	1.2x10 ⁻⁵	0.50	0.31	0.00334 (0.00513)
HNF1B	rs10908278	17	36,099,952	T	A	3.1x10 ⁻⁸	0.0083	0.87	0.12	0.00075 (0.00493)
MLX	rs684214	17	40,696,915	T	C	0.18	0.28	0.33	0.52	-0.00476 (0.00490)
GIP-TLL6	rs35895680	17	47,060,322	C	A	0.46	0.46	0.98	0.96	0.00016 (0.00507)
ACE	rs57676627	17	62,203,128	T	C	0.15	0.44	0.63	0.80	-0.00408 (0.00792)
BPTF-PITPNC1	rs80320393	17	65,643,646	T	C	0.0064	0.48	0.72	0.32	-0.00328 (0.00954)
BPTF-PITPNC1	rs9899520	17	65,957,568	A	G	0.0028	0.0099	0.48	0.56	0.00349 (0.00482)
CYTH1	rs1044486	17	76,792,179	G	A	0.32	0.32	0.85	0.50	0.00082 (0.00438)
LAMA1	rs9948462	18	7,076,836	T	C	0.0096	0.21	0.64	0.33	0.00209 (0.00459)
TCF4	rs72926932	18	53,050,646	C	A	0.75	0.65	0.52	0.91	0.00706 (0.00981)
GRP-MC4R	rs9957320	18	56,876,430	G	T	0.0098	0.0087	0.17	0.24	-0.00755 (0.00582)
GRP-MC4R	rs6567160	18	57,829,135	C	T	0.16	0.15	0.49	0.39	0.00354 (0.00517)
GRP-MC4R	rs76227980	18	58,036,384	C	T	0.38	0.45	0.99	0.47	-0.00024 (0.01495)
BCL2A	rs12454712	18	60,845,884	T	C	0.27	0.32	0.76	0.44	-0.00146 (0.00486)
ZNF236	rs12457906	18	74,555,593	G	A	0.82	0.61	0.22	0.38	0.00528 (0.00439)
UHRF1-PTPRS	rs262549	19	4,951,064	G	C	0.78	0.59	0.36	0.48	0.00573 (0.00625)
MAP2K7	rs2115107	19	7,968,168	A	G	0.50	0.78	0.43	1.0	0.00348 (0.00356)
FARSA-ZNF799	rs4804181	19	12,509,536	A	C	0.55	0.72	0.59	0.84	-0.00272 (0.00472)
FARSA-ZNF799	rs3111316	19	13,038,415	A	G	0.34	0.17	0.14	0.29	-0.00688 (0.00480)
CILP2-TM6SF2	rs58542926	19	19,379,549	T	C	0.013	0.030	0.20	0.47	-0.01056 (0.00833)
ZNF257	rs142395395	19	22,100,706	A	G	0.45	0.38	0.38	0.95	0.02193 (0.02071)
PEPD	rs10406327	19	33,890,838	C	G	0.028	0.79	0.59	0.28	-0.00236 (0.00454)
TOMM40-APOE-GIPR	rs1871045	19	45,326,768	T	C	0.70	0.79	0.90	0.62	0.00056 (0.00425)
TOMM40-APOE-GIPR	rs429358	19	45,411,941	T	C	0.071	0.13	0.85	0.67	0.00122 (0.00642)
TOMM40-APOE-GIPR	rs10406431	19	46,157,019	A	G	0.057	0.37	0.69	0.58	0.00168 (0.00419)
TOMM40-APOE-GIPR	rs2238689	19	46,178,661	C	T	0.47	0.52	0.13	0.37	-0.00655 (0.00436)
ZC3H4	rs3810291	19	47,569,003	A	G	0.23	0.26	0.58	0.59	-0.00260 (0.00455)
FOXA2	rs2181063	20	22,427,370	C	G	0.10	0.25	0.029	0.88	-0.01091 (0.00459)
RALY	rs4911405	20	32,674,967	T	C	0.0061	0.22	0.67	0.61	-0.00225 (0.00509)
HNF4A	rs12625671	20	42,994,812	C	T	0.61	0.84	0.70	0.065	0.00201 (0.00585)
HNF4A	rs1800961	20	43,042,364	T	C	0.21	0.27	0.45	0.43	0.01021 (0.01364)
EYA2	rs6063046	20	45,596,378	A	G	0.80	0.16	0.026	0.75	-0.01319 (0.00560)
CEBPB	rs6091115	20	48,832,020	T	C	0.73	0.80	0.64	0.47	-0.00205 (0.00438)
GNAS	rs736266	20	57,387,352	T	A	0.18	0.22	0.55	0.39	-0.00261 (0.00444)
MTMR3-ZNRF3	rs36575	22	30,205,572	C	T	0.67	0.57	0.45	0.98	0.00787 (0.00865)
YWHAH	rs75307421	22	32,203,334	A	G	0.0093	0.014	0.45	0.25	-0.01016 (0.01398)
PNPLA3	rs738408	22	44,324,730	T	C	0.0046	0.23	0.22	0.22	0.00591 (0.00504)

<i>PIM3</i>	rs28691713	22	50,356,302	C	T	0.037	0.44	0.15	0.88	-0.00687 (0.00437)
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Chr: chromosome. SE: standard error. OR: odds-ratio.

^aThe sample size contributing to each ancestry: African 15,043 cases and 22,318 controls; East Asian 56,268 cases and 227,155 controls; European 67,192 cases and 831,463 controls; Hispanic 11,027 cases and 18,885 controls; and South Asian 16,540 cases and 32,952 controls.

Supplementary Note Table 7. Candidate causal genes at T2D loci identified from functional annotation and colocalization with molecular QTLs in the DIAMANTE multi-ancestry study, and support from complementary analyses undertaken by recent T2D GWAS efforts overlapping with DIAMANTE.

Candidate causal gene ^a	Locus	DIAMANTE multi-ancestry			MVP (Vujkovic et al. 2020)		DIAMANTE European (Mahajan et al. 2018)	DIAMANTE East Asian (Spracklen et al. 2020)
		Missense variant	pQTL	eQTL ^b	Missense variant	TWAS ^b	Missense variant	eQTL ^b
ABO	<i>ABO</i>		cis	SM,VAT				
<i>AC012354.6</i>	<i>SIX3-SIX2</i>			I				I
<i>AC122129.1</i>	<i>RAI1</i>			SM		SM,SAT,VAT		
ACVR1C	<i>CYTIP</i>	p.Ile482Val						
<i>ADCY5</i>	<i>ADCY5</i>			I		SM		
<i>ANK1</i>	<i>ANK1</i>			SM,SAT		SM,SAT		SM,SAT
<i>AP3S2</i>	<i>AP3S2</i>			I,SM		L,P,SM,SAT,VAT		
<i>APOE</i>	<i>TOMM40-APOE-GIPR</i>	p.Cys130Arg	cis		p.Cys130Arg		p.Cys130Arg	
ARAP1	<i>CENTD2</i>			I				
<i>ARHGAP19</i>	<i>ARHGAP19-SLIT1</i>			SAT,VAT		SM,SAT,VAT		SAT
<i>ATP2A3</i>	<i>ZZEF1</i>	p.Gly216Arg				SM		
<i>ATP5G1</i>	<i>GIP-TLL6</i>			SM		SM		
<i>C12orf65</i>	<i>MPHOSPH9-ZNF664</i>			SAT		SAT,VAT		
<i>CALR</i>	<i>FARSA-ZNF799</i>			SAT		SAT		
CAMK1D	<i>CDC123-CAMK1D</i>			I				
CARD9	<i>GPSM1</i>			I				
CCDC67	<i>MTNR1B</i>			I				
<i>CCNE2</i>	<i>TP53INP1</i>			VAT		VAT		
CD101	<i>PTGFRN</i>			I				
CDK8	<i>RNF6</i>			I				
<i>CDKN1B</i>	<i>CDKN1B</i>	p.Val109Gly					p.Val109Gly	
<i>CEP68</i>	<i>CEP68</i>			I,L,SM,SAT,VAT		L,P,SM,SAT,VAT		
<i>CLUAP1</i>	<i>CLUAP1-SLX4</i>			H		H,SM,SAT,VAT		
CPB1	<i>BCAR1</i>		trans					
CPLX1	<i>CTBP1-PCGF3-MAEA</i>			I				
CRHR2	<i>CRHR2</i>			I				
CTA-85E5.10	<i>MTMR3-ZNRF3</i>			SAT				
<i>CTD-2021H9.3</i>	<i>TSPAN8</i>			SM		SM		
CTRB1	<i>BCAR1</i>		cis					
<i>DCAF16</i>	<i>LCORL</i>			SAT		P,VAT		
DGKB	<i>DGKB</i>			I				
DLK1	<i>DLK1-MEG3</i>		cis	I				
DNLZ	<i>GPSM1</i>			I				
<i>FAM134C</i>	<i>MLX</i>			SM		SM		
FAM85B	<i>MSRA-XKR6</i>			SAT				
<i>FBXL22</i>	<i>USP3</i>			SM		SM		

GCKR	GCKR	p.Leu446Pro			p.Leu446Pro		p.Leu446Pro	
GLP1R	ZFAND3-KCNK16-GLP1R	p.Pro7Leu			p.Pro7Leu			
GPSM1	GPSM1			I	p.Leu391Ser			
HAUS6	HAUS6			I		SAT,VAT		
HERC1	USP3			VAT		SAT,VAT		
HMG20A	HMG20A			I		VAT		
HNF1A	HNF1A	p.Ala98Val					p.Ala98Val, p.Gly226Ala	
HNF4A	HNF4A	p.Thr139Ile					p.Thr139Ile	
HSD17B12	HSD17B12			H,I,L	p.Leu280Ser	H,L,P,SM,SAT,VAT		
IGF2BP2	IGF2BP2			I				
INHBB	GLI2			VAT				
IRS1	IRS1			SAT,VAT		SAT,VAT		
ITFG3	ITFG3			SAT		SAT,VAT		
ITGB6	RBSM1			H,SAT,VAT		SAT,VAT		
JAZF1	JAZF1			L,SM,SAT,VAT		L,P,SM,SAT,VAT		
KCNJ11	KCNJ11-ABCC8	p.Val337Ile			p.Lys23Glu	SM		
KLF14	KLF14			SAT		SAT		
KLHL42	KLHDC5			I		SM		
MAN2C1	PTPN9-SIN3A			L,SM,SAT		H,P,SM,SAT,VAT		
MED23	MED23-ENPP3			SM		SM		SM
MTNR1B	MTNR1B			I				
MYO5C	MYO5C	p.Glu1075Lys				P		SM
NDUFAF6	TP53INP1			SAT,VAT		SM,SAT,VAT		
NEUROG3	VPS26A-NEUROG3	p.Gly167Arg					p.Gly167Arg	
NKX6-3	ANK1			I				I
NOTCH2	NOTCH2			L		L,P		
NUS1	NUS1			I		P		I,P,SM
PAM	SLCO6A1-PAM	p.Ser539Trp	cis				p.Ser539Trp	
PCGF3	CTBP1-PCGF3-MAEA			SM,SAT,VAT				
PGM1	PGM1		cis					
PLA2G4B	LTK			SM,SAT,VAT		SAT,VAT		
PLEKHA1	PLEKHA1			I,SAT		SM,SAT		
PLRP1	BCAR1		trans					
POC5	HMGCR-POC5	p.His36Arg				SAT,VAT	p.His36Arg	
PRC1-AS1	PRC1			SAT				
PRSS2	BCAR1		trans					
PTGFRN	PTGFRN			I	p.Ile837Val			
PXK	PXK			I				
QSER1	QSER1	p.Arg1101Cys			p.Arg1101Cys		p.Arg1101Cys	
RBM6	RBM6			I,SM,SAT,VAT		H,L,P,SM,SAT,VAT		
RCCD1	PRC1			I,SM,SAT,VAT		H,SM,SAT,VAT		
RNF6	RNF6			I				
RP11-107F6.3	LTK			SAT		SM,SAT		

RP11-282O18.3	<i>MPHOSPH9-ZNF664</i>			SM,VAT			
<i>RP11-395N3.2</i>	<i>IRS1</i>			SAT,VAT		SAT,VAT	
<i>RP11-419C23.1</i>	<i>KCNU1</i>			SAT			SAT
<i>RP11-463M16.4</i>	<i>GIP-TTLL6</i>			SM		SM	
<i>RP11-53O19.3</i>	<i>MRPS30</i>			SAT		H,SAT	
RP11-613D13.5	<i>HSD17B12</i>			SAT			
<i>RP11-817O13.8</i>	<i>PTPN9-SIN3A</i>			SAT,VAT		P,SAT,VAT	
<i>RP11-89K21.1</i>	<i>SIX3-SIX2</i>			I			I
RP1-239B22.5	<i>KCNJ11-ABCC8</i>			SAT			
RP5-1042I8.7	<i>NOTCH2</i>			VAT			
RPL39L	<i>ST6GAL1</i>			I			
<i>RREB1</i>	<i>SSR1-RREB1</i>	p.Asp1171Asn			p.Asp1171Asn		p.Asp1171Asn
<i>SCD5</i>	<i>SCD5</i>	p.Glu197Gln					p.Glu197Gln
<i>SETD8</i>	<i>MPHOSPH9-ZNF664</i>			SM		SM	
<i>SIX2</i>	<i>SIX3-SIX2</i>			I			I
<i>SIX3</i>	<i>SIX3-SIX2</i>			I			I,P
SKOR1	<i>MAP2K5</i>			SM			
SLC12A8	<i>SLC12A8</i>			I			
SLC16A11	<i>SLC16A11-SLC16A13</i>	p.Val113Ile					
<i>SLC22A3</i>	<i>SLC22A3</i>			L		L	
<i>SLC30A8</i>	<i>SLC30A8</i>	p.Arg325Trp			p.Arg325Trp		p.Arg325Trp
<i>SMCO4</i>	<i>MTNR1B</i>			I		P,SM	
<i>ST6GAL1</i>	<i>ST6GAL1</i>			I		P	
STARD10	<i>CENTD2</i>			I			
STEAP2	<i>STEAP1</i>			SAT			
<i>SYCE2</i>	<i>FARSA-ZNF799</i>			L		L	
TCF7L2	<i>TCF7L2</i>			I			
<i>TH</i>	<i>INS-IGF2-KCNQ1</i>			I		P	
<i>TOM1L2</i>	<i>RAI1</i>			SM		SM,SAT,VAT	
<i>TSPAN8</i>	<i>TSPAN8</i>			L		L	
<i>TUBG2</i>	<i>MLX</i>			SM		SM	
UBE2E2	<i>UBE2E2</i>			I			
<i>UBE2Z</i>	<i>GIP-TTLL6</i>			SM		SM	
<i>WFS1</i>	<i>WFS1</i>			SAT		SM,SAT	
<i>WSCD2</i>	<i>WSCD2</i>	p.Thr266Ile			p.Thr266Ile		p.Thr113Ile
<i>ZBTB20</i>	<i>ZBTB20</i>			VAT		SAT	SAT
ZNF236	<i>ZNF236</i>			SAT			
<i>ZNF703</i>	<i>KCNU1</i>			SAT			SAT

pQTL: protein quantitative trait locus. eQTL: expression quantitative trait locus. TWAS: transcriptome-wide association study.

^aGenes highlighted in bold not reported in complementary analyses conducted by DIAMANTE European, DIAMANTE East Asian or MVP.

^bTissues: hypothalamus (H); islet (I); liver (L); skeletal muscle (SM); subcutaneous adipose (SAT); visceral adipose (VAT).