



The genetic architecture of type 2 diabetes

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The genetic architecture of type 2 diabetes

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For more than a century, people have debated the genetic architecture of common traits: the number, frequency, and effect sizes of inherited variants that contribute to individual risk. Genome-wide association studies have identified scores of common variants associated with type 2 diabetes, but in aggregate, these explain only a fraction of heritability. To test the hypothesis that lower-frequency variants explain much of the remainder, we performed whole genome and exome sequencing in large numbers of individuals with and without type 2 diabetes and, to increase statistical power, expanded sample size via genotyping and imputation. Variants associated with type 2 diabetes after sequencing were overwhelmingly common and most fell within regions previously identified by genome-wide association studies. Comprehensive enumeration of sequence variation is necessary to discover functional alleles that provide important clues to disease pathophysiology, but large-scale sequencing does not support a major role for lower-frequency variants in predisposition to type 2 diabetes.

Whole genome sequence analysis of 2,657 individuals

The Genetics of Type 2 Diabetes (GoT2D) study selected for whole genome sequencing cases of type 2 diabetes (T2D) and ancestry-matched normoglycemic controls from northern and central Europe (**Methods; Supplementary 1**). To increase power to identify low-frequency ($0.5\% < \text{MAF} < 5\%$) and rare ($\text{MAF} < 0.5\%$) T2D variants of large effect, GoT2D preferentially ascertained individuals from the extremes of genetic risk (**Methods**). The genome sequence of 1,326 cases and 1,331 control individuals was determined through joint statistical analysis of low-coverage whole-genome sequence ($\sim 5x$), deep-coverage exome sequence ($\sim 82x$), and array-based genotypes at 2.5M single nucleotide variants (SNVs) (**Extended Data Fig. 1, Table 2**).

We detected, genotyped, and estimated haplotype phase for 26.7M genetic variants (**Extended Data Fig. 1, Table 3**), including 1.5M short insertion-deletion variants (indels) and 8.9K large deletions. Individual diploid genomes carried a mean of 3.30M variants (range: 3.20M-3.35M), including 271K indels (262K-327K), and 669 (579-747) large deletions. These data include many variants not directly studied by previous genome-wide association studies (GWAS), including all of the indels as well as 420K common

and 2.4M low-frequency SNVs poorly tagged ($r^2 \leq 0.30$)^{1,2} by arrays. We estimate near-complete ascertainment (98.2%) of SNVs with minor allele count >5 (MAF>0.1%), and high accuracy (>99.1%) at heterozygous genotypes (**Methods; Fig. 1a**). As half the sequenced individuals were T2D cases, ascertainment was enhanced for any rare or low-frequency variants that substantially increase T2D risk (**Fig. 1a**). Specifically, we estimate $\geq 80\%$ power to detect (at genome-wide significance, $\alpha = 5 \times 10^{-8}$) T2D risk variants with MAF $\geq 5\%$ and OR ≥ 1.87 , or MAF $\geq 0.5\%$ and OR ≥ 4.70 (**Extended Data Fig. 4**).

We tested all 26.7M variants for T2D association by logistic regression assuming an additive genetic model (**Supplementary 2**). Analyses using a mixed-model framework to account for population structure and relatedness generated almost identical results. At genome-wide significance, 126 variants at four loci were associated with T2D (**Fig. 1b**). This included two previously-reported common-variant loci (*TCF7L2*, *ADCY5*), a previously-reported low-frequency variant in *CCND2*³ (rs76895963, MAF=2.6%, $p_{\text{seq}} = 4.2 \times 10^{-9}$), and a novel common-variant association near *EML4* (MAF=34.8%, $p_{\text{seq}} = 1.0 \times 10^{-8}$). There was no significant evidence of T2D association for sets of low-frequency or rare variants within coding regions, nor within specified non-coding regulatory elements (**Methods**).

Increasing sample size across the genome by statistical imputation

Power to detect association with low-frequency and rare variants of modest effect is limited in 2,657 individuals. To increase power for variants discovered via genome sequencing, we imputed sequence-based genotypes into 44,414 additional European-origin individuals (11,645 T2D cases, 32,769 controls; **Methods**) from 13 studies (**Supplementary 3**). We estimated power in the combined sequence plus imputed data, adjusting for imputation quality, to be $\geq 80\%$ for variants with MAF $\geq 5\%$ and OR ≥ 1.23 , or MAF $\geq 0.5\%$ and OR ≥ 1.92 (**Extended Data Fig. 4**). Meta-analysis combining results for the sequence and imputed data identified 674 variants across 14 loci associated with T2D at genome-wide significance (**Fig. 1c**). All were common except the *CCND2* variant described above. We observed a novel association with a common variant near *CENPW* (rs11759026, MAF=23.2%, $p_{\text{meta}} = 3.5 \times 10^{-8}$; **Fig. 1c**) and replicated this association in an additional 14,201 cases and 100,964 controls from the DIAGRAM consortium ($p = 2.5 \times 10^{-4}$; $p_{\text{combined}} = 1.1 \times 10^{-11}$; **Methods**). The *EML4* signal detected in the sequence data did not replicate in the imputed data ($p = 0.59$; $p_{\text{meta}} = 0.26$; **Fig. 1c**).

To test for additional association signals we performed conditional analysis at loci previously associated with risk of T2D (**Methods**). We identified two novel association signals, both involving low-frequency variants, at a corrected significance threshold ($\alpha < 1.8 \times 10^{-6}$; **Methods**): one at the *IRS1* locus (rs78124264, MAF=2.2%, $p_{\text{conditional}} = 2.5 \times 10^{-7}$) and one upstream of *PPARG* (rs79856023, MAF=2.2%, $p_{\text{conditional}} = 9.2 \times 10^{-7}$) (**Extended Data Table 5**). The *PPARG* signal overlaps regulatory elements in hASC pre-adipose and HepG2 cells, consistent with evidence that altered adipose regulation drives the primary *PPARG* signal⁴.

Exome sequence analysis in 12,940 individuals

To further define the contribution of low-frequency alleles, we next focused on variants in protein-coding sequence, seeking to improve power to detect rare-variant association by exploiting the more robust functional annotation of coding variation and the potential to aggregate multiple alleles of similar presumed impact in the same gene⁵. Specifically, the T2D-GENES study implemented exome sequencing in large samples of diverse ancestry ($n=10,437$). We combined data from GoT2D and T2D-GENES, generating exome sequence data from 12,940 individuals (6,504 cases; 6,436 controls) drawn from five ancestry groups: 4,541 of European origin, and ~2,000 [range: 1,943-2,217] each of South Asian, East Asian, Hispanic, and African American origin (**Extended Data Fig. 1, Table 2; Supplementary 4**). Mean coverage was 82x across the coding sequence of 18,281 genes, identifying 3.04M variants (1.19M protein-altering) (**Extended Data Table 3; Supplementary 5,6**). Each diploid genome carried a mean of 9,243 (range: 8,423-11,487) synonymous, 7,636 (6,935-9,271) missense, and 250 (183-358) protein-truncating alleles (**Extended Data Table 3; Supplementary 7**).

We tested for T2D association within the five ancestral groups, assuming an additive genetic model, using mixed-model approaches that account for population structure and relatedness⁶, and combined ancestry-specific results via trans-ethnic meta-analysis (**Methods**). We estimate $\geq 80\%$ power to detect (at genome-wide significance) T2D risk variants with MAF $\geq 5\%$ and OR ≥ 1.36 , or MAF $\geq 0.5\%$ and OR ≥ 2.29 (**Methods; Extended Data Fig. 4**). Only one variant reached genome-wide significance (*PAX4* Arg192His, rs2233580, $p = 9.3 \times 10^{-9}$) (**Table 1; Extended Data Figs. 6,7; Supplementary 8**). This association was exclusive to East Asians, in whom the 192His allele is, in fact, common (MAF $\sim 10\%$) with a substantial effect size (allelic OR=1.79[1.47-2.19]); 192His is virtually absent in other ancestries (MAF=0.014%). The rs2233580 association replicated in independent East Asian case-control data

($n=3,301$; $p=5.9 \times 10^{-7}$; **Supplementary 9**) and was distinct ($r^2 < 0.05$) from previously-reported GWAS SNVs at the *GCC1-PAX4* locus^{7,8}. *PAX4* encodes a transcription factor involved in islet differentiation and function⁹ (**Supplementary 10**), and *PAX4* variants have been implicated in early-onset monogenic diabetes¹⁰. However, in East Asian cases, 192His was not associated with age of diabetes diagnosis ($p=0.64$), indicating this variant influences risk of type 2 rather than early-onset monogenic diabetes¹⁰ (**Supplementary 9**).

To increase power to detect association of rare variants that cluster in individual genes, we deployed gene-level variant aggregation tests¹¹ across the exome sequence data (**Methods; Supplementary 11**). We observed no deviation from the null distribution of association statistics, and no single gene reached exome-wide significance ($\alpha=2.5 \times 10^{-6}$) (**Methods; Supplementary 12,13**). When we focused on 634 genes mapping to known GWAS regions, only *FES* exceeded a reduced significance threshold of $\alpha=7.9 \times 10^{-5}$ ($p_{\text{SouthAsian}}=7.2 \times 10^{-6}$, $p_{\text{multiethnic}}=1.9 \times 10^{-5}$) (**Method; Supplementary 14**). This aggregate signal was driven entirely by the South Asian-specific Pro536Ser variant (MAF=0.9%, OR=6.7 [2.6-17.3], $p=7.5 \times 10^{-6}$), indicating that *FES* is likely to be the effector gene at the *PRC1* GWAS locus².

Increasing sample size across the exome by array-based genotyping

To increase power to detect coding variant associations (**Extended Data Fig. 4**), we contributed early T2D-GENES exome data to the design of Illumina exome array¹², and then collected genotypes from an additional 28,305 T2D cases and 51,549 controls of European-ancestry from 13 studies (**Extended Data Fig. 1, Table 2; Supplementary 15**). Of 27,904 protein-altering variants with MAF>0.5% detected in exome sequence data from $n=4,541$ European individuals, variation at 81.6% was captured on the array (**Supplementary 16**).

Association analysis in the combined sequence and array data from >90,000 individuals identified 18 coding variants (17 nonsynonymous), at 13 loci, which exceeded genome-wide significance ($\alpha=5 \times 10^{-8}$) (**Table 1; Extended Data Figs. 6,7**). All of these were common (MAF>5%) and all but one mapped within established common-variant GWAS regions^{1,13}. The exception, which we replicated in the INTERACT study¹⁴ ($n=9,292$; $p_{\text{INTERACT}}=2.4 \times 10^{-4}$; $p_{\text{meta}}=2.2 \times 10^{-11}$), involved a common haplotype of four strongly-correlated coding variants in *MTMR3* and *ASCC2* (**Table 1**). Of these, *MTMR3* Asn960Ser

(MAF=8.3%) had the strongest residual association signal on conditional analysis, implicating *MTMR3*, encoding a phosphatidylinositol phosphatase¹⁵, as the probable effector transcript at this locus (**Extended Data Table 5, Figs. 6,7; Supplementary 10,17**).

The remaining coding variant signals provided an opportunity to highlight causal alleles and effector transcripts for known GWAS signals. For five loci (*SLC30A8*, *GCKR*, *PPARG*, *KCNJ11-ABCC8*, *PAM*), the coding variants identified had previously been nominated as causal for their respective GWAS signals^{3,13,16}. For the other seven loci, GWAS meta-analyses had previously highlighted a lead variant in non-coding sequence^{7,13,17}. We (re)evaluated these relationships with conditional and credible set analyses, finding that, at most, the evidence supported a direct causal role for the coding variants concerned (**Extended Data Table 5, Figs. 6,7; Supplementary 10,17**).

For example, at the *CILP2* locus¹³, previous GWAS had identified the non-coding variant, rs10401969, as the lead SNV. However, direct genotyping of *TM6SF2* Lys167Glu on the exome array, revealed complete linkage disequilibrium with rs10401969, and reciprocal signal extinction in conditional analyses (**Extended Data Table 5, Figs. 6,7**). In previous GWAS, the association at Lys167Glu had been obscured by incomplete genotyping and poor imputation (**Supplementary 18**). The *TM6SF2* Lys167 allele has been shown to underlie predisposition to hepatic steatosis¹⁸, and was associated with fasting hyperinsulinemia ($p=1.0 \times 10^{-4}$) in 30,824 non-diabetic controls from the present study. This combination of genetic and functional data, consistent with known mechanistic links between insulin resistance, T2D, and fatty liver disease¹⁹, implicates *TM6SF2* Lys167Glu as the likely T2D-risk variant at this locus.

In contrast, the association at *RREB1* Asp1171Asn represented a novel signal, conditionally independent of the adjacent common-variant GWAS signal. This association, together with that involving a second associated coding variant, Ser1554Tyr, which has a marked association with fasting glucose ($p=2.7 \times 10^{-9}$ in levels in 38,338 non-diabetic subjects from the present study) (**Supplementary 19**) establishes *RREB1*²⁰ as the probable effector gene at the *SSR1* locus.

Given the concentration of coding-variant associations within established GWAS loci, we sought to

nominate additional single-variant signals in 634 genes mapping to established T2D GWAS regions using a Bonferroni-corrected $\alpha=1.6 \times 10^{-5}$ (**Methods; Supplementary 14,20**). At *HNF4A*, we confirmed a T2D association at Thr139Ile (European MAF range 0.7-3.8%, OR=1.15 [1.08-1.22], $p=2.9 \times 10^{-6}$)²¹ distinct both from the common non-coding lead GWAS SNV^{1,13,17}, and multiple rare *HNF4A* variants implicated in monogenic diabetes²². Additional coding variant associations in *TSPAN8* and *THADA* highlighted these two genes as probable effector transcripts in their respective GWAS regions (**Supplementary 10,21**).

Rare coding variant associations in Mendelian genes

We extended gene-based tests for rare-variant associations to gene-sets implicated in monogenic or syndromic diabetes and in altered glucose metabolism²². Across 81 genes harboring rare alleles causal for monogenic or syndromic diabetes or related glycemic traits ('Monogenic All'; **Supplementary 22**), the only variant or gene reaching genome-wide significance was the previously-mentioned *PAX4* Arg192His. However, across the entire gene-set, we observed a weak aggregate association with T2D-risk ($p=0.023$; **Fig. 2a**). The association was considerably stronger in two subsets of genes more directly implicated in monogenic and syndromic diabetes: a manually-curated set of 28 genes for which diabetes was the primary phenotype ('Monogenic Primary') and a partially-overlapping set of 13 genes reported in OMIM as causal for MODY or neonatal diabetes ('Monogenic OMIM') (**Supplementary 22**).

The 'Monogenic OMIM' gene-set had a statistically robust signal of association ($p=2.8 \times 10^{-5}$, OR=1.51 [1.25-1.83]) driven by allelic burden of MAF<1% alleles. Effect size estimates tracked with increasing stringency of variant annotation and gene-set definition, consistent with progressive enrichment for functional over neutral alleles (**Fig. 2b**). This signal does not reflect inclusion among T2D cases of individuals who, in reality, had monogenic diabetes: the association was not concentrated among genes most frequently responsible for monogenic diabetes²² (**Fig. 2c**), and age of diabetes diagnosis was no younger in variant carriers than non-carriers (**Supplementary 23**). The association signal remained after all alleles listed as 'disease-causing' within the Human Genetic Mutation Database were excluded ($p=2.9 \times 10^{-4}$, OR=1.50 [1.21-1.86]).

These analyses point to widespread enrichment for T2D association among rare coding alleles in genes causal for monogenic diabetes: in these genes, alleles of penetrance sufficient to drive familial segregation of early-onset diabetes coexist alongside those of more modest effect predisposing to later-onset T2D. No other compelling signals of rare-variant enrichment were detected using gene-set enrichment or protein-protein interaction analysis in other pre-defined gene-sets (**Supplementary 24-26**).

No evidence for synthetic association

In 2010, Goldstein and colleagues proposed that common-variant GWAS signals may be the consequence of low-frequency and rare variants that by chance cluster on common haplotypes²³. While this hypothesis has been debated^{24,25} and assessed indirectly^{1,26}, we used the near-complete ascertainment of genetic variation in 2,657 genome-sequenced individuals to directly test the importance of ‘synthetic’ associations²⁷. We focused on the ten T2D GWAS loci at which our sample provided the strongest statistical evidence for association ($p < 0.001$), implementing a conditional analysis procedure to assess whether combinations of SNVs within a 5Mb window could explain the common-variant signal (**Extended Data Table 8; Methods**).

We first focused on missense variants, finding that none of the ten signals could be explained by low-frequency and rare variants within 2.5Mb of the common index SNV (**Fig. 3a**). For example, at the *IRS1* locus, including the five observed missense *IRS1* alleles in the model did not meaningfully diminish the index SNV association ($p_{\text{unconditional}} = 2.8 \times 10^{-6}$, $p_{\text{conditional}} = 4.3 \times 10^{-6}$). With 99.7% ascertainment of low-frequency coding variants (**Methods**), these results rule out synthetic associations produced by missense variants at these ten loci.

We expanded the search to include all low-frequency and rare variants, non-coding and coding, within 2.5Mb of index SNVs. At no locus was a single low-frequency or rare variant sufficient to explain the GWAS signal (**Fig. 3a**). At 8 of the 10 loci, ≥ 10 low-frequency and rare variants were needed to reverse the direction of effect at the common index SNV; at *TCF7L2*, even 50 were insufficient (**Fig. 3b**). We note that the statistical procedure we developed and deployed is biased *in favor* of the synthetic association hypothesis, since it is highly prone to over-fitting. Nonetheless, at 8 of the 10 loci the data were

indistinguishable from a null model of no synthetic association (**Extended Data Table 8; Supplementary 27**).

Nominating candidate functional alleles

Using the GoT2D whole genome sequence data, we constructed 99% ‘credible sets’ for each T2D GWAS locus on the assumption of one causal variant per locus (**Methods**)²⁸. Across 78 published autosomal loci at which the reported index SNV had MAF>1%, 99% credible set sizes ranged from 2 (*CDKN2AB*) to ~1,000 (*POU5F1*) variants; at 71 loci, the credible set contained >10 variants (**Fig. 3c; Supplementary 28**). The GoT2D dataset provides near-complete ascertainment of common and low-frequency variants to support more comprehensive credible set analysis than studies based on genotyping or imputation alone^{1,29}: of the credible set variants identified from whole genome sequence data, ~60% are absent from HapMap and ~5% from 1000G Phase 1 (**Fig. 3c**).

Genomic maps of chromatin state or transcription factor binding³⁰⁻³³ have been used to prioritize causal variants within credible sets^{34,35}. We jointly modeled genetic association and genomic annotation data at T2D GWAS loci using fgwas³⁶. Consistent with previous reports^{32,33}, associated variants were enriched in coding exons, transcription factor binding sites, and enhancers active in pancreatic islets and adipose tissue (**Extended Data Fig. 9**). Overall, including the functional annotation data reduced credible set size by 35%. At several loci, access to complete sequence data prioritized variants that overlap relevant regulatory annotations and were previously overlooked. For example, at the *CCND2* locus, three variants not present in HapMap Phase 2 have combined probability of 90.0% of explaining the common-variant signal¹³ (**Extended Data Fig. 9**); one of these (rs3217801) is a 2bp indel overlapping an islet enhancer element.

Modelling disease architecture

To evaluate the overall contribution of low-frequency coding variation to T2D risk, we estimated the proportion of variance in T2D-liability attributable to each such variant³⁷ (**Methods; Extended Data Fig. 10**). We focused on exome array data to maximize sample size, and on variants with MAF>0.1% (sensitivity of variant ascertainment and accuracy of OR estimation decline below this threshold). Among the 31,701 variants on the exome array with 0.1%<MAF<5% there was a progressive increase in

the maximum OR estimates with decreasing frequency. However, the liability variance explained (LVE) for these variants rarely exceeded 0.05%, limiting power to detect association in the sample size available (**Extended Data Fig. 10**). We estimated (**Methods**) that the LVE collectively attributable to coding variants in the $0.1\% < \text{MAF} < 5\%$ range was 2.9%, compared to 6.3% for common variants.

Finally, we compared our T2D association results with predictions from population genetic simulations³⁸ under twelve models that vary widely with respect to the proportion of heritability explained by common, low-frequency, and rare variants. We mirrored the GoT2D study design (with imputation) and performed in parallel the same association analysis on empirical and simulated data, focusing on variants with $\text{MAF} > 0.1\%$ and allowing for power loss due to imperfect imputation (**Methods**).

Figure 4 displays results for three representative models: a ‘purifying selection’ model in which low-frequency and rare variants explain ~75% of T2D heritability, an intermediate model in which low-frequency/rare and common variants both contribute substantially, and a ‘neutral’ model in which common variants explain ~75% of T2D heritability. Predictions of the first two models differ markedly in the numbers of low-frequency and rare risk variants that are associated with T2D. Specifically, these two models predict a larger number and greater effect size of low-frequency variants found in our whole genome sequencing study as compared to those observed in the empirical data. In contrast, empirical data are consistent with predictions under the ‘neutral’ common-variant model.

Discussion

The century-old Mendelian-biometrician debate pitted those who attributed trait variation to rare variants of large effect against those who argued that trait variation is largely due to many common variants of small effect. The debate today is about whether the ‘missing heritability’ after GWAS is due largely to individually rare, highly-penetrant variants³⁹ or to a large universe of common alleles of modest effect⁴⁰. The results are of more than academic interest, since genetic architecture plays out powerfully in relation to the power of genetic diagnosis and the application of precision medicine.

Our data and analysis indicate that nearly all common-variant associations detectable by whole genome sequencing were previously found by GWAS based on genotyping arrays and imputation: concerns

about incomplete coverage due to ‘holes’ in HapMap⁴¹ coverage were, we show, unfounded. Of more lasting interest, the combination of genome and exome sequencing in large samples provides limited evidence to document the role of lower-frequency variants — both coding and genome wide — in T2D predisposition. Of course, rare risk alleles have long been known to contribute in families with early-onset forms of diabetes, and sequencing of Mendelian and GWAS genes has identified rare variants that influence disease risk^{42,43}. Sequencing of T2D cases in much larger samples will undoubtedly uncover additional low-frequency and rare variants that provide biological and potentially clinical value. Nonetheless, our empirical and simulated data argue that these lower-frequency variants contribute much less to T2D heritability than do common variants. Moreover, the frequency spectrum of variant association signals is consistent with a model whereby limited selective pressure distributes most the genetic variance influencing T2D risk among common alleles³⁸, consistent with the frequency distribution of inter-individual sequence variation.

Our results further strengthen the case for sequencing of diverse samples: the population-enriched T2D risk variant in *PAX4* dovetails with similar findings involving *SLC16A11*⁴⁴ in East Asian and Native American populations and *TBC1D4*⁴⁵ in Greenland Inuits. Populations subject to bottlenecks and/or extreme selective pressures^{42,45,46} may be particularly fruitful.

Understanding the inherited basis of T2D will require much greater progress in uncovering the mechanisms whereby common, mostly non-coding, variants influence disease risk. The combination of global epigenetic measurements, genome editing⁴⁷, and high-throughput functional assays⁴⁸ make it increasingly practical to characterize large numbers of non-coding variants and the processes they impact. Genome sequencing in much larger sets of individuals will no doubt provide foundational information to guide such experimentation and connect the results to human population variation, physiology, and disease. Integration of biological insights gleaned from common and rare variants into a unified picture of disease pathophysiology will be required to fully understand the basis of this common but challenging disease.

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ENDNOTES

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Data deposition: Whole genome sequence data from the GoT2D project are available by application to the European Genotype Archive (EGAS00001001459) and from dbGAP (phs000840.v1.p1). Whole exome sequence data from the T2D-GENES project are available from the European Genotype Archive (EGAS00001001460), and will also be available from dbGAP during 2016. Summary level data from the exome array component of this project (and from the exome and genome sequence) can be freely accessed at the Accelerating Medicines Partnership T2D portal (www.type2diabetesgenetics.org), and similar data from the GoT2D-imputed data at www.diagram-consortium.org.

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TABLES AND FIGURES

Locus	Gene	Variant	RAF range	Eur MAF	Alleles	Exomes (N=12,940)		Exome-chip (N=79,854)		Combined (N=92,794)	
						p-value	OR (95% CI)	p-value	OR (95% CI)	p-value	OR (95% CI)
Established common causal coding variant signals											
<i>GCKR</i>	<i>GCKR</i>	rs1260326 Pro446Leu	0.49- 0.86	0.37	C, T	0.075	1.05 (0.99-1.11)	4.8x10 ⁻⁹	1.07 (1.04-1.11)	1.2x10 ⁻⁹	1.07 (1.04-1.10)
<i>PPARG</i>	<i>PPARG</i>	rs1801282 Pro12Ala	0.86- 0.99	0.14	C, G	0.0030	1.16 (1.06-1.27)	1.8x10 ⁻⁷	1.10 (1.06-1.14)	4.2x10 ⁻⁸	1.11 (1.07-1.15)
<i>PAM1</i> <i>PPIP5K</i> <i>2</i>	<i>PAM</i> <i>PPIP5K2</i>	rs35658696 Asp563Gly rs36046591 Ser1207Gly	0.00- 0.05 0.00- 0.05	0.054	G, A G, A	0.00045 0.0099	1.36 1.34 (1.12-1.61)	1.7x10 ⁻⁷ 1.0x10 ⁻⁶	1.15 1.17 (1.10-1.25)	5.7x10 ⁻¹⁰ 3.3x10 ⁻⁸	1.17 1.19 (1.12-1.26)
<i>SLC30A</i> <i>8</i>	<i>SLC30A8</i>	rs13266634 Asp325Trp	0.58- 0.91	0.33	C, T	2.9x10 ⁻⁵	1.15 (1.09-1.22)	2.7x10 ⁻¹⁸	1.14 (1.11-1.17)	4.8x10 ⁻²³	1.14 (1.11-1.17)
<i>KCNJ11</i> <i>/</i> <i>ABCC8</i>	<i>KCNJ11</i> <i>ABCC8</i>	rs5215 Val337Ile rs5219 Lys23Glu rs757110 Ala1369Ser	0.08- 0.40 0.06- 0.40 0.06- 0.40	0.40	C, T T, C C, A	0.11 0.056 0.20	1.07 1.08 1.06 (1.00-1.12)	3.4x10 ⁻⁹ 5.1x10 ⁻⁹ 2.3x10 ⁻⁸	1.07 1.07 1.07 (1.04-1.11)	1.3x10 ⁻⁹ 9.0x10 ⁻¹⁰ 1.7x10 ⁻⁸	1.07 1.07 1.07 (1.05-1.10) (1.05-1.10) (1.04-1.10)
Other coding variant associations within established common variant GWAS regions											
<i>THADA</i>	<i>THADA</i>	rs35720761 Cys1605Tyr	0.85- 1.00	0.10	C, T	0.0021	1.12 (1.01-1.23)	3.5x10 ⁻⁸	1.11 (1.07-1.16)	3.3x10 ⁻¹⁰	1.12 (1.07-1.16)
<i>COBLL1</i>	<i>COBLL1</i>	rs7607980 Asn939Asp	0.84- 1.00	0.12	T, C	1.4x10 ⁻⁵	1.21 (1.11-1.33)	4.7x10 ⁻¹¹	1.14 (1.10-1.19)	8.3x10 ⁻¹⁵	1.15 (1.11-1.19)
<i>WFS1</i>	<i>WFS1</i>	rs1801212 Val333Ile rs1801214 Asn500Asn rs734312 Arg611His	0.70- 1.00 0.59- 0.96 0.11- 0.85	0.30 0.41	A, G T, C A, G	0.0026 0.0019 0.12	1.14 1.08 1.05 (0.99-1.11)	9.3x10 ⁻¹² 2.0x10 ⁻¹² 1.3x10 ⁻¹⁰	1.08 1.08 1.07 (1.03-1.10)	9.0x10 ⁻¹⁴ 1.5x10 ⁻¹⁴ 6.9x10 ⁻¹¹	1.09 1.08 1.06 (1.04-1.09)
<i>RREB1</i>	<i>RREB1</i>	rs9379084 Asp1171Asn	0.87- 0.98	0.11	G, A	2.2x10 ⁻⁵	1.19 (1.09-1.30)	1.1x10 ⁻⁵	1.12 (1.06-1.17)	4.0x10 ⁻⁹	1.13 (1.09-1.18)
<i>PAX4</i>	<i>PAX4</i>	rs2233580 Arg192His	0.00- 0.10	0.00	T, C	9.3x10 ⁻⁹	1.79 (1.47-2.19)	NA	NA	9.3x10 ⁻⁹	1.79 (1.47-2.19)
<i>GPSM1*</i>	<i>GPSM1*</i>	rs60980157 Ser391Leu	0.26	0.26	C, T	NA	NA	1.7x10 ⁻⁹	1.09 (1.06-1.12)	1.7x10 ⁻⁹	1.09 (1.06-1.12)
<i>CILP2</i>	<i>TM6SF2</i>	rs58542926 Glu167Lys	0.03- 0.10	0.082	T, C	0.00015	1.22 (1.10-1.36)	1.9x10 ⁻⁷	1.13 (1.08-1.18)	3.2x10 ⁻¹⁰	1.14 (1.10-1.19)
Coding variant associations outside established common variant GWAS regions											
<i>MTMR3/</i> <i>ASCC2</i>	<i>MTMR3</i> <i>ASCC2</i>	rs41278853 Asn960Ser rs11549795 Val123Ile rs28265 Asp407His rs36571 Pro423Ser	0.92- 1.00 0.92- 1.00 0.92- 1.00 0.92- 1.00	0.083	A, G C, T C, G G, A	9.2x10 ⁻⁵ 0.00040 0.00050 0.0023	1.26 (1.12-1.42) 1.23 (1.10-1.38) 1.21 (1.08-1.36) 1.23 (1.08-1.40)	3.2x10 ⁻⁶ 2.0x10 ⁻⁵ 1.9x10 ⁻⁵ 2.0x10 ⁻⁵	1.12 (1.07-1.17) 1.11 (1.06-1.16) 1.11 (1.06-1.16) 1.11 (1.06-1.16)	5.6x10 ⁻⁹ 1.0x10 ⁻⁷ 1.1x10 ⁻⁷ 3.0x10 ⁻⁷	1.14 (1.09-1.19) 1.13 (1.08-1.18) 1.12 (1.08-1.17) 1.12 (1.08-1.17)

Table 1 | Nonsynonymous coding variants achieving genome-wide significance. These loci were identified through single-variant analyses of exome sequence data in 6,504 cases and 6,436 controls and exome-array in 28,305 cases and 51,549 controls. RAF: Risk allele frequency. Eur MAF: Minor allele frequency in Europeans. OR: odds-ratio. CI: confidence interval. N: Total number of individuals analysed. N: Total number of individuals analysed. Genome-wide significance defined as $p < 5 \times 10^{-8}$. **GPSM1* variant failed quality control in exome sequence: association p-values derive only from exome-array analysis. The synonymous variant Thr515Thr (rs55834942) in *HNF1A* also reached genome-wide significance ($p=1.0 \times 10^{-8}$) in the combined analysis. Alleles are aligned to the forward strand of NCBI Build 37 and represented as risk and other allele.

FIGURE LEGENDS

Figure 1 | Ascertainment of variants and single-variant results.

a, Sensitivity of low-coverage genome sequence data to detect SNVs in the deep exome sequence data, relative to other variant catalogs. Points represent results for a specific minor allele count. All results assume $OR=1$ for all variants, unless stated otherwise. Manhattan plots of single-variant association analyses for: **b**, sequence data alone (1,326 cases and 1,331 controls) and **c**, meta-analysis of sequence and imputed data (total of 14,297 cases and 32,774 controls).

Figure 2 | Association between T2D and variants in genes for Mendelian forms of diabetes. **a**, p -values of aggregate association for variants from 6,504 T2D cases and 6,436 controls in three sets of Mendelian diabetes genes, for five variant “masks” (**Methods**). Dotted line: $p=0.05$. **b**, Estimated T2D odds ratio (OR) for carriers of variants in each gene set and mask. Error bars: one standard error. **c**, Estimated ORs (bars, left axis) and p -values (dots, right axis) for carriers of variants in the PTV+NS_{strict} mask for each gene. Error bars: one standard error. Red: $OR > 1$; blue: $OR < 1$; dotted line: $p=0.05$.

Figure 3 | Exclusion of synthetic associations and construction of credible causal variant sets at T2D GWAS loci. Ten T2D GWAS loci were selected for synthetic association testing ($p<0.001$; **Methods**). **a**, The effect size observed at the GWAS index SNV (sequence data) before (navy blue) and after (light blue, grey) conditioning on candidate rare and low-frequency ($MAF<5\%$) variants which could produce synthetic association. **b**, Example of synthetic association exclusion at the *TCF7L2* locus. **c**, Credible sets for T2D GWAS loci where credible set consisted of <80 variants displaying the proportion of credible set variants present in the HapMap and 1000G catalogs.

Figure 4 | Empirical T2D association results compared to results under different simulated disease models. Observed number of rare and low-frequency ($MAF<5\%$) genetic association signals for T2D detected genome-wide after imputation compared to the numbers seen under three simulated disease models for T2D which were plausible given results (T2D recurrence risks, GWAS, linkage) prior to large-scale sequencing. Simulated models were defined by two parameters: disease target size T and degree of coupling τ between the causal effects of variants and the selective pressure against them³⁸. Simulated

data were generated to match GoT2D imputation quality as a function of MAF (**Methods**).

EXTENDED METHODS

Ethics statement. All human research was approved by the relevant institutional review boards and conducted according to the Declaration of Helsinki. All participants provided written informed consent.

1 Data generation

1.1 GoT2D integrated panel generation

1.1.1. GoT2D sequenced samples

Here we describe how we generated, processed, and carried out quality control (QC) on sequence and genotype data for the 2,891 individuals initially chosen for GoT2D from four studies, and how this resulted in 2,657 individuals (1,326 T2D cases and 1,331 non-diabetic controls) for analysis (**Extended Data Figure 1**). We preferentially sampled early-onset, lean, and/or familial T2D cases and overweight controls with low fasting glucose levels⁴⁹. Specific details of selected samples are provided in **Extended Data Table 2** and **Supplementary 1**.

1.1.2. DNA sample preparation

De-identified DNA samples were sent to the Broad Institute (DGI, FUSION), Wellcome Trust Centre for Human Genetics in Oxford (UKT2D), and Helmholtz Zentrum München (KORA) and prepared for genetic analysis. DNA quantity was measured by Picogreen (all), and samples with sufficient total DNA and minimum concentrations for downstream experiments were genotyped for a set of 24 SNVs using the Sequenom iPLEX assay (DGI, FUSION, UKT2D): one gender assay and 23 SNVs located across the autosomes. The genotypes for these SNVs were used as a quality filter to advance samples and a technical fingerprint for subsequent sequencing and genome-wide array genotypes.

1.1.3. Exome sequencing

Genomic DNA was sheared, end repaired, ligated with barcoded Illumina sequencing adapters, amplified, size selected, and subjected to in-solution hybrid capture using the Agilent SureSelect Human All Exon 44Mb v2.0 (DGI, FUSION, UKT2D) and v3.0 (KORA) bait set (Agilent Technologies, USA). Resulting Illumina exome sequencing libraries were qPCR quantified, pooled, and sequenced with 76bp paired-end reads using Illumina GAII or HiSeq 2000 sequencers to ~82-fold mean coverage.

1.1.4. Genome sequencing

Whole-genome Illumina sequencing library construction was performed as described for exome

capture above, except that genomic DNA was sheared to a larger target size and hybrid capture was not performed. Resulting libraries were size selected to contain fragment insert size of 380bp±20% (DGI, FUSION, KORA) and 420bp±25% (UKT2D) using gel electrophoresis or the SAGE Pippin Prep (Sage Science, USA). Libraries were qPCR quantified, pooled, and sequenced with 101bp paired-end reads using Illumina GAII or HiSeq 2000 sequencers to ~5-fold mean coverage.

1.1.5. HumanOmni2.5 array genotyping

Genotyping was performed by the Broad Genetic Analysis Platform. DNA samples were placed on 96-well plates and genotyped using the Illumina HumanOmni2.5-4v1_B SNV array.

1.1.6. Alignment and processing of exome and genome sequence data

1.1.6.1. Alignment of sequence reads to reference genome

Sequence data were processed and aligned to hg19 using the Picard (broadinstitute.github.io/picard/), BWA⁵⁰, and GATK^{51,52} pipelines. Resulting BAM and VCF files were submitted to NCBI and are available in dbGaP (accession number phs000840.v1.p1, study name NIDDK_GoT2D).

1.1.6.2. Coverage and QC of aligned sequence reads

We excluded 151 exome samples with average coverage ≤20x in >20% of the target bases and 68 genome samples with average coverage ≤5x. After sequence alignment and post-processing, aligned sequence reads were screened based on multiple QC criteria, including number of mapped reads, number of mapped bases with <1% estimated base call error rate (>Q20), fraction of duplicate reads, fraction of properly paired reads, distribution of insert sizes, distribution of mean base quality with respect to sequencing cycles, and GC bias (**Extended Data Figure 1**).

1.1.6.3. Detecting and handling contamination of sequence reads

We assessed possible DNA contamination in the genome and exome sequence data using verifyBamID⁵³ using two methods. First, we estimated the contamination level of sequenced samples using allele frequencies estimated from the HumanOmni2.5 array on a thinned set of 100,000 markers with minor allele frequency (MAF)>5%. Second, for samples with HumanOmni2.5 genotypes, we used these genotypes together with sequence data to estimate contamination and identify possible sample swaps. We excluded exome sequence data for 7 individuals and genome sequence data for 59 individuals with estimated contamination ≥2% using either method. Prior to variant calling, uncontaminated sample swaps were assigned to the correct sample label after searching for the matching pairs using the same method.

1.1.7. GoT2D integrated panel genotype calling

1.1.7.1. SNV identification

We processed whole-genome sequence reads across the remaining 2,764 QC-passed individuals by two SNV calling pipelines: GotCloud (www.gotcloud.org) and GATK UnifiedGenotyper⁵⁴. We merged unfiltered SNV calls across the two call sets and then processed the merged site list through the SVM and VQSR filtering algorithms implemented by those pipelines. SNVs that failed both filtering algorithms were removed before genotyping and haplotype integration. For the 2,733 QC-passed exome sequenced individuals, we used GATK UnifiedGenotyper to call SNVs.

1.1.7.1.1. Illumina HumanOmni2.5 array genotyping

We used Illumina GenomeStudio v2010.3 with default clusters to call HumanOmni2.5 genotypes after comparing different clustering algorithms and observing that the default cluster resulted in highest concordance with sequence-based genotypes. Called genotypes were run through a standard QC pipeline; samples passing a call rate threshold of 95%, and genetic fingerprint (24 marker panel) and gender concordance were passed on to downstream GWAS QC. SNVs with GenTrain score<0.6, cluster separation score<0.4, or call rate<97% were considered technical failures at the genotyping laboratory and deleted before data release. We removed samples with call rate<98%, and SNVs monomorphic across all samples, failed by 1000G Omni 2.5 QC filter, or with Hardy-Weinberg equilibrium $p < 10^{-6}$ (**Extended Data Figure 1**). 85 samples were removed in this process.

1.1.7.2. Short insertion and deletion (indel) identification

For the whole-genome sequence data, we used the GATK UnifiedGenotyper to call short indels (<50bp). Because short indels are known to have high false positive rates due to systematic sequencing and alignment errors⁵⁴, we used stringent filtering criteria in SVM and VQSR and excluded indels that failed either algorithm. For exome sequencing, we used GATK UnifiedGenotyper to call short indels, following best practices described elsewhere⁵¹.

1.1.7.3. Large deletion identification

We used GenomeSTRiP⁵⁵ to call large (>100bp) deletions in the whole-genome sequence data. After initial discovery of large deletions in 2,764 QC-passed individuals, we merged the discovered sites with deletions identified in 1,092 sequenced individuals from the 1000G Project to increase sensitivity and then genotyped the merged site lists across the 2,764 individuals. After applying the default filtering implemented in GenomeSTRiP, pass-filtered sites variable in any of the samples were identified as candidate variant sites. Among these candidate sites, we excluded variants in

known immunoglobulin loci to reduce the impact of possible cell-line artifacts. We then excluded 136 more individuals owing to an unusually large number of variants per sample ($>\text{median}+3\times\text{mean}$ absolute deviation). Variants present only in these excluded individuals were removed from further analysis.

1.1.8. GoT2D integrated panel haplotype integration

1.1.8.1. Genotype likelihood calculation

We merged SNVs discovered from the three experimental platforms into one site list and calculated genotype likelihoods across all sites separately by platform. Because exome sequence data have substantial off-target coverage, we calculated likelihoods across the genome combining data from the genome and exome sequence experiments. For genome sequence, we calculated likelihoods using GotCloud; for exomes, we used GATK UnifiedGenotyper; for HumanOmni2.5 genotypes, we converted hard genotype calls into genotype likelihoods assuming a genotype error rate of 10^{-6} . For indels, we calculated likelihoods in a similar way except the HumanOmni2.5 data could not be used. For structural variants (SVs), genotype likelihoods were calculated from GenomeSTRiP using the whole-genome sequence data.

1.1.8.2. Integration of genotype and sequence data

We calculated combined genotype likelihoods across each of the 2,874 individuals as the product of the corresponding genome, exome, and HumanOmni2.5 likelihoods assuming independent data across platforms (**Extended Data Figure 1**). We then phased the genotype data using the strategy developed for 1000G Phase 1⁵⁴. Specifically, we phased the integrated likelihoods using Beagle⁵⁶ with 10,000 SNVs per chunk and 1,000 overlapping SNVs between consecutive chunks. We refined phased sequences using Thunder⁵⁷ as implemented in GotCloud (genome.sph.umich.edu/wiki/GotCloud) with 400 states to improve genotype and haplotype quality.

1.1.9. GoT2D integrated panel QC

2,874 individuals were available in the integrated haplotype panel. To identify population outliers, we carried out principal components analysis (PCA). We computed PCs for each of the three variant types (SNVs, short indels, large deletions) using EPACTS on an LD-pruned ($r^2 < 0.20$) set of autosomal variants obtained by removing large high-LD regions^{58,59}, variants with $\text{MAF} < 0.01$, and variants with Hardy-Weinberg equilibrium $p < 10^{-6}$. Inspecting the first ten PCs for each variant type, we identified 43 population outliers and 136 additional outliers for large deletions only; we excluded these 179 individuals. We excluded an additional 38 individuals based on close relationships (estimated genome-wide identity-by-descent proportion of alleles shared > 0.20) with other study members.

2,657 individuals remained available for downstream analyses (**Extended Data Figure 1**).

1.1.10. GoT2D integrated panel evaluation of variant detection sensitivity

Since we had no external data to evaluate SNV and indel variant detection sensitivity and genotype accuracy for our integrated haplotype panel, we evaluated accuracy for the low-pass whole-genome sequence data using the exome sequence data as gold standard for variants at which exome sequence depth was ≥ 10 . We consider the resulting sensitivity and accuracy estimates as lower bounds for the integrated panel, which combined information from the genome, exome, and HumanOmni2.5 data.

We estimated the sensitivity of low-pass genome sequence data to detect true SNVs by calculating the proportion of exome-sequencing-detected SNVs detected by low-pass genome sequencing in the 2,538 individuals with data for all three experimental platforms. For exome sequence allele counts $< 1,000$, we merged adjacent allele count bins until the number of alleles was $> 1,000$. We estimated the sensitivity of low-pass genome sequencing to detect common, low-frequency, and rare SNVs as 99.8%, 99.0%, and 48.2%, respectively. Similarly, we estimated the sensitivity of low-pass genome sequence to detect true short indels by calculating the proportion of exome sequencing-detected short indels detected by low-pass genome sequencing. Sensitivity estimates were $> 99.9\%$, 93.8%, and 17.9% for common, low-frequency, and rare short indels, respectively.

To estimate the sensitivity of the combined low-pass genome and exome sequence data, we focused on coding SNVs and calculated the proportion of HumanOmni2.5 SNVs detected by either sequencing platform. Because HumanOmni2.5 SNVs are enriched for common variants, we calculated a weighted averaged sensitivity at each allele count, weighted by the number of exome-detected variants given the allele count. Sensitivity estimates were 99.9%, 99.7%, and 83.9% for common, low-frequency, and rare variants.

1.1.11. GoT2D integrated panel evaluation of genotype accuracy

To evaluate genotype accuracy for SNVs, we focused on chromosome 20, and compared the concordance of low-pass whole-genome-sequence-based genotypes with those based on exome sequence. Overall genotype concordance was 99.86%. Homozygous reference, heterozygous, and homozygous non-reference concordances were 99.97%, 98.34%, and 99.72%. We also compared genotype concordance between exome sequence and HumanOmni2.5 genotypes. Overall concordance was 99.4%. When the HumanOmni2.5 genotypes were homozygous reference, heterozygous, and homozygous non-reference, concordances were 99.97%, 99.69%, and 99.88%. We evaluated genotype accuracy of indels for the 210 chromosome 20 indels that overlapped

between those discovered by exome and genome sequencing. Overall genotype concordance was 99.4%. When the exome genotypes were homozygous reference, heterozygous, and homozygous non-reference, concordances were 99.8%, 95.8%, and 98.6%.

To evaluate the genotype accuracy of our low-pass genome sequence data to detect true structural variants, we took advantage of the 181 individuals in our study previously included in the WTCCC array-CGH based structural variant detection experiment⁶⁰. Taking the WTCCC data as gold standard, we estimated genotype accuracy across 1,047 overlapping structural variants (with reciprocal overlap>0.8) genome-wide. The overall genotype concordance was 99.8%. When the WTCCC genotypes were homozygous reference, heterozygous, and homozygous non-reference, concordances were 99.9%, 99.6%, and 99.7%.

1.2. GoT2D+T2D-GENES multiethnic exome panel generation and QC

1.2.1. Samples

We considered 6,504 T2D cases and 6,436 controls from 14 studies of African American, East Asian, South Asian, Hispanic, and European ancestry. In contrast to the GoT2D whole-genome integrated panel, this data set also includes GoT2D individuals for whom whole genome data were not available. Sample characteristics are provided in **Extended Data Table 2** and **Supplementary 4**.

Sequence reads were processed and aligned to the reference genome (hg19) with Picard (<http://picard.sourceforge.net>). Polymorphic sites and genotypes were called with GATK, with filtering of sites performed using Variant Quality Score Recalibration (VQSQR) for SNVs, and hard filters for indels. Genotype likelihoods were computed controlling for contamination.

Hard calls (the GATK-called genotypes but set as missing at a genotype quality (GQ)<20 threshold⁵¹) and dosages (the expected value of the genotype, defined as $\text{Pr}(RX|\text{data})+2\text{Pr}(XX|\text{data})$, where X is the alternative allele) were computed for each sample at each variant site. Hard calls were used only for quality control, while dosages were used in all downstream association analyses. Multi-allelic SNVs and indels were dichotomized by collapsing alternate alleles into one category because downstream association analyses required bi-allelic variants.

Individuals were excluded from analysis if they were outliers on one of multiple metrics: poor array genotype concordance (where available), high number of variant alleles or singletons, high or low allele balance (average proportion of non-reference alleles at heterozygous sites), or excess mean heterozygosity or ratio of heterozygous to homozygous genotypes.

Within this reduced set of individuals, we then performed extended QC using ethnicity and T2D

status to provide high-quality genotype data for downstream association analyses. Within each ethnicity, we excluded variants based on hard call rate (<90% in any cohort), deviation from Hardy-Weinberg equilibrium ($p < 10^{-6}$ in any ancestry group), or differential call rate between T2D cases and controls ($p < 10^{-4}$ in any ancestry group). We then considered autosomal variants that passed extended QC and with MAF > 1% in all ancestry groups for trans-ethnic kinship analyses. We calculated identity-by-state (IBS) between each pair of samples based on independent variants (trans-ethnic $r^2 < 0.05$) and constructed axes of genetic variation through PCA implemented in EIGENSTRAT⁶¹ to identify ethnic outliers (**Supplementary 29**). We also identified duplicates based on IBS, and excluded the sample from each pair with lowest call rate and/or mismatch with external information. The extended QC excluded 68 individuals, and 9.9% of SNVs and 90.8% of indels from the clean dataset.

2. Association analysis

2.1.1. Power calculation

We used the genetic power calculator (<http://pngu.mgh.harvard.edu/~purcell/gpc/>) to estimate power to detect T2D association assuming 8% prevalence. For the T2D-GENES+GoT2D exome sequence data set we assumed: (i) a fixed-effect across all five ancestry groups (12,940 individuals); and (ii) an effect specific to one group (2,000 individuals) (**Extended Data Figure 4**). We repeated our calculations for combined exome sequence and exome array data, assuming a fixed effect across all ethnicities, for an effective total sample size of 82,758 individuals (**Extended Data Figure 4**).

For the GoT2D integrated panel we allowed for incomplete variant detection by multiplying power by the estimated sensitivity to detect the variant as a function of MAF. For imputed variants, we first multiplied the sample size by the median imputation quality (rsq_hat) obtained from MaCH/Thunder or minimac⁶² for the corresponding MAF bin across the analyzed cohorts, and then multiplied the estimated power by the fraction of variants that passed the imputation quality cutoff for that MAF bin.

For gene-based tests in the T2D-GENES+GoT2D data, we made use of a Bonferroni correction for 20,000 genes, corresponding to $p < 2.5 \times 10^{-6}$. We used a simulated haplotype dataset from the SKAT package (<http://cran.r-project.org/web/packages/SKAT/vignettes/SKAT.pdf>) and estimated the power of SKAT-O to detect association of variants within a gene at this threshold as a function of the phenotypic variance (1%) in a liability scale explained by additive genetic effects and the percentage of variants that were causal (50% and 100%). As for single-variant power calculations, we

considered: (i) a fixed-effect across all ethnicities (12,940 individuals); and (ii) an effect specific to one ancestry group (2,000 individuals) (**Extended Data Figure 4**).

2.2. GoT2D integrated panel association analysis

2.2.1. Single-variant association analysis

We tested for T2D association in a logistic regression framework assuming an additive genetic model. We used the Firth bias-corrected likelihood ratio test^{63,64} as our primary analysis strategy; we repeated association analysis using the score test for inclusion in sample-size-weighted meta-analysis (**Supplementary 2**). Tests were adjusted for sex, the first two genotype-based PCs to account for population stratification, and an indicator function for observed temporal stratification based on sequencing date and center. PCs were calculated using linkage-disequilibrium (LD) pruned ($r^2 < 0.20$) HumanOmni2.5M array variants with MAF > 1% after removing large high-LD regions^{58,59}.

2.2.2. Aggregate association analysis

To test for aggregate association within coding regions of the genome, we used the approach described in **2.3.6**. For every gene and mask tested, p-values were greater than 2.5×10^{-4} .

We also tested for aggregate association among variants in non-coding regions of the genome. We aggregated variants in individual pancreatic islet enhancer elements (see **6.1**), as these elements collectively demonstrated strongest genome-wide enrichment of T2D association. We performed both the burden and SKAT tests using genotypes from the integrated panel on variants with MAF < 5% in each islet enhancer element. We used a Bonferroni threshold $p < 1.68 \times 10^{-7}$ based on a nominal significance level of $\alpha = 0.05$ corrected for 298,240 elements with at least one variant. All elements tested in this manner had p-value greater than 2.5×10^{-6} .

2.3. GoT2D+T2D-GENES multiethnic association analysis

2.3.1. Kinship analysis

Within each ancestry group, we considered autosomal variants that passed QC with MAF > 1% for ethnic-specific kinship analyses. We calculated IBS between each pair of samples in the ancestry group based on independent variants (ethnic-specific $r^2 < 0.05$) and constructed a kinship matrix to account for intra-ethnic population structure and relatedness in downstream mixed-model (EMMAX) based association analyses⁶. We also used IBS to identify pairs of related individuals within each ancestry group (defined by $\pi\text{-hat} > 0.3$). We then defined intra-ethnic related exclusion lists for downstream non-EMMAX association analyses using the following steps: (i) remove the control from

each T2D-status discordant pair; and (ii) remove the sample with lowest call rate from each T2D-status concordant pair. We also constructed intra-ethnic axes of genetic variation through PCA implemented in EIGENSTRAT⁶¹. We identified axes of genetic variation in each ancestry group for inclusion as covariates in downstream non-EMMAX association analyses to account for intra-ethnic population structure that: (i) explain at least 0.5% genotypic variation; and/or (ii) demonstrate nominal association ($p < 0.05$) with T2D in logistic regression analysis.

2.3.2. Single-variant association analysis

Within each ancestry group, we performed a score test of T2D association with each variant passing ethnic-specific QC in a linear regression framework under an additive model in EMMAX⁶. We also performed a Wald test of T2D association with each variant passing ethnic-specific QC in a logistic regression framework under an additive model with adjustment for ethnic-specific axes of genetic variation after exclusion of related samples (**Supplementary 30**). Within each ancestry group, we calculated genomic control inflation factors (score EMMAX and Wald) based on independent variants used for the ethnic-specific kinship analyses and corrected association summary statistics (p-value and SE) to account for residual population structure.

Subsequently, we performed trans-ethnic fixed-effects meta-analysis of ancestry-specific association summary statistics at each variant based on: (i) sample size weighting of score EMMAX directed p-values; and (ii) inverse-variance weighting of Wald beta/SE (to obtain unbiased estimates of allelic odds ratios and confidence intervals that cannot be constructed from EMMAX effect estimates). We also performed trans-ethnic meta-analysis of ancestry-specific association summary statistics (score EMMAX beta/SE) at each variant using MANTRA⁶⁵, using pair-wise mean allele frequency differences at the subset of independent variants used for trans-ethnic kinship analyses as a prior for relatedness between ancestry groups.

2.3.3. Validation of *PAX4* association signal in additional East Asian studies

We validated the *PAX4* Arg192His (rs2233580) association signal in an additional 1,789 T2D cases and 1,509 controls of East Asian ancestry from Hong Kong, Korea, and Singapore (**Supplementary 9**). Within each study, we tested for association with T2D in a logistic regression model, and combined association summary statistics across studies through fixed-effects meta-analysis (**Supplementary 9**). Among T2D cases, we also tested for association with age of diagnosis in a linear regression model, and combined association summary statistics across studies through fixed-effects meta-analysis (**Supplementary 9**).

2.3.4. Admixture analysis

Admixed populations can offer greater statistical power to detect association because diverse ancestry increases genetic variation. However, admixture can also introduce false-positive signals due to population stratification and heterogeneity of effects because of differential LD⁶⁶. To assess the contribution of ancestral background in the two admixed groups (African American and Hispanic), we inferred local ancestry based on SNVs in available GWAS data using two approaches. For African Americans, we ran HAPMIX⁶⁷ using CEU and YRI haplotypes from HapMap as reference, and estimated the proportion of European ancestry at each genomic position. For Hispanics, we ran Multimix⁶⁸ using European, West African, and Native American haplotypes from HapMap as reference, and estimated the proportion of European ancestry at each genomic position, since we observe only a very low West African contribution (1.1-3.2%, **Supplementary 31**). We then repeated our intra-ethnic EMMAX-based analyses within African American and Hispanic ancestry groups, this time adjusting for local ancestry by including the estimated proportion of European ancestry at each variant as a covariate. Adjustment for local ancestry resulted in numerically similar association statistics as those from unadjusted analyses in the African American and Hispanic samples.

2.3.5. Gene-based analysis

We generated four variant lists ('masks') based on MAF and functional annotation. We mapped variants to transcripts in Ensembl 66 (GRCh37.66). Using annotations from CHAOs v0.6.3, SnpEFF v3.1, and VEP v2.7, we identified variants predicted to be protein-truncating (e.g. nonsense, frameshift, essential splice site) denoted PTV-only or 'Mask 1'; or protein-altering (e.g. missense, in-frame indel, non-essential splice site) in at least one mapped transcript (by at least one of the three algorithms) with MAF<1%, denoted PTV+missense or 'Mask 2'. We additionally used the procedure described by Purcell et al.⁶⁹ to identify subsets of missense variants with MAF<1% meeting 'strict' or 'broad' criteria for being deleterious, using annotation predictions from Polyphen2-HumDiv, PolyPhen2-HumVar, LRT, Mutation Taster, and SIFT; variants predicted deleterious by all five algorithms or by at least one algorithm were denoted PTV+NS_{strict} or 'Mask 3' and PTV+NS_{broad} or 'Mask 4', respectively. Indels predicted by CHAOs, SnpEFF, or VEP to introduce frameshifts were included in the 'strict' category. We calculated MAFs for each ancestry using high-quality genotype calls (GQ>20) for all samples passing extended QC. We considered a variant to have MAF<1% if MAF estimates for every ancestry group were <1%.

We used the MetaSKAT R package (v0.32)¹¹ with the SKAT v0.93 library to perform SKAT-O⁷⁰ analysis within each ancestry, and in meta-analysis. Within each ancestry group, we analyzed genotype

dosages with adjustment for ethnic-specific axes of genetic variation after exclusion of 96 related individuals. We assumed homogenous allele frequencies and genetic effects for all studies within an ancestry group. We performed meta-analysis using genotype-level data, allowing for heterogeneity of allele frequencies and genetic effects between (but homogeneity within) ancestry groups. All analyses were completed using the recommended rho vector for SKAT-O: (0, 0.12, 0.22, 0.32, 0.52, 0.5, 1).

2.4. Imputed data

2.4.1. Samples

We carried out genotype imputation into 44,414 individuals (11,645 T2D cases and 32,769 controls) from 13 studies using the GoT2D integrated haplotypes as reference panel. Characteristics of the imputed studies are provided in **Extended Data Table 2** and **Supplementary 3**.

2.4.2. Single-variant association meta-analysis

The one sequenced and thirteen imputed studies totaled 12,971 T2D cases and 34,100 controls. Each study performed its own sample- and variant-based QC. In each study, SNVs with minor allele count (MAC) ≥ 1 passing QC were tested for T2D association assuming an additive genetic model adjusting for study-specific covariates. Association testing was performed using logistic regression Firth bias-corrected, likelihood ratio, or score tests as implemented in EFACTS (genome.sph.umich.edu/wiki/EFACTS) or SNPTTEST⁷¹. To account for related samples in the Framingham Heart Study, generalized estimating equations (GEE) were used, as implemented in R. Residual population stratification for each study was accounted for using genomic control⁷². We then carried out fixed-effects sample-size weighted meta-analysis as implemented in METAL⁷³.

2.4.3. Conditional analyses in established GWAS loci

We compiled a list of 143 previously-reported genome-wide significant SNVs in 81 T2D autosomal loci (a) from Morris et al.¹³ and Voight et al.²; (b) from papers they referenced; and (c) from references in the NHGRI GWAS catalog⁷⁴. We LD pruned these SNVs ($r^2 < 0.95$), yielding a list of 129 SNVs. We deleted the *CILP2* locus (and two SNVs) from subsequent whole-genome analyses owing to large regions in which no variants passed QC, resulting in a list of 127 index SNVs at 80 autosomal loci. To identify additional T2D-associated variants within these 80 T2D autosomal loci in the genome-wide data, we repeated GWA analysis for 12 of the 13 studies (conditional analysis results for FHS were unavailable), conditioning on the 127 index SNVs. We performed fixed-effects inverse-variance meta-analysis to combine conditional analysis results from the studies totaling 12,298 cases

and 26,440 controls. For each known locus, we analyzed all SNVs within 500kb of the known index SNVs; if there were multiple known index SNVs, we analyzed all SNVs within 500kb of the most proximal and distal index SNVs. We imposed a conditional-analysis significance threshold of $\alpha=1.8\times 10^{-6}$ based on a proportional number of multiple tests for ~83Mb of the ~3000Mb genome.

2.5. Exome array data

2.5.1. Samples

We considered 28,305 T2D cases and 51,549 controls from 13 studies of European ancestry, genotyped with the Illumina exome array. Characteristics of the studies are provided in **Extended Data Table 2** and **Supplementary 15**.

2.5.2. Overlap of exome sequence variation with exome array

We assessed overlap of variants present on the exome array with those observed in our trans-ethnic exome-sequence data. Since exome array primarily contains SNVs that are predicted to be protein altering, we focused on nonsense, essential splice site, and missense variants. Only variants passing QC in both sequence and array data were included in our overlap assessment.

2.5.3. Data processing, QC, and kinship analysis

Within each study, exome array genotypes were initially called using GenCall (https://support.illumina.com/downloads/gencall_software.html) and Birdseed⁷⁵. Sample and variant QC was then undertaken within each study based on several quality control filters. Criteria for sample exclusion included low call rate (<99%), mean heterozygosity, high singleton counts, non-European ancestry, sex discrepancy, GWAS discordance (where data were available), genotyping platform fingerprint discordance, and duplicate discordance. Variants were excluded based on call rate (<99%), deviation from Hardy-Weinberg equilibrium ($p<10^{-6}$), duplicate, chromosome or allele mismatch, GenTrain score <0.6, Cluster separation score <0.4, and manual cluster checks. Missing genotypes were subsequently re-called using zCall, with a second round of QC to exclude poor quality samples (call rate <99% and mean heterozygosity) and variants (call rate <99%). Within each study, we considered independent autosomal variants that passed QC with MAF>1% for kinship analyses, and calculated IBS between each pair of samples. We used these statistics to: (i) identify non-European ancestry samples to be excluded from all downstream analyses; (ii) construct a kinship matrix to account for fine-scale population structure and relatedness in downstream EMMAX-based association analyses; (iii) identify related samples to be excluded from downstream non-EMMAX association analyses; and (iv) calculate axes of genetic variation for inclusion as

covariates in downstream non-EMMAX association analyses to account for fine-scale population structure (if required).

2.5.4. Single-variant association analysis

Within each study, we performed a score test of T2D association with each variant passing QC in a mixed-model regression framework under an additive model in EMMAX⁶. We also performed a Wald test of T2D association with each variant in a logistic regression framework under an additive model with adjustment for axes of genetic variation after exclusion of related samples. For each test, we corrected SE and p -value for the genomic control inflation factor (if >1) calculated based on the independent autosomal variants used for kinship analysis.

Across studies, we performed fixed-effects meta-analysis of association summary statistics at each variant based on: (i) inverse-variance weighting of score EMMAX beta/SE; (ii) sample size weighting of score EMMAX directed p -values; and (iii) inverse-variance weighting of Wald beta/SE. For each of these meta-analyses, we applied a second round of correction of SE and p -value by genomic control, again calculated based on the independent autosomal SNVs used for kinship analyses.

2.5.5. Combined exome sequence and exome array single-variant analysis

We considered variants that were represented both in the exome sequence and on the exome chip. We began by performing fixed-effects meta-analysis of association summary statistics (after correction for genomic control, as described above) from the exome-chip meta-analysis and the European ancestry sequenced samples using: (i) inverse-variance weighting of score EMMAX beta/SE; (ii) sample size weighting of score EMMAX directed p -values; and (iii) inverse-variance weighting of Wald beta/SE. Subsequently, we performed trans-ethnic fixed-effects meta-analysis of ancestry-specific association summary statistics (after correction for genomic control, as described above) at each variant based on: (i) sample size weighting of score EMMAX directed p -values; and (ii) inverse-variance weighting of Wald beta/SE.

2.5.6. Gene-based analyses

We made use of the four variant masks defined for exome sequence gene-based analyses, but with MAF calculated across all exome array studies. Within each study, we performed SKAT-O analyses⁷⁰, with adjustment for axes of genetic variation after exclusion of related samples. We combined p -values for association across studies via meta-analysis with Stouffer's method⁷⁶.

2.5.7. Evaluating relationships between association signals for coding variants and previously reported lead SNVs at established GWAS loci

For coding variants mapping to established T2D susceptibility loci and achieving genome-wide

significance in combined exome sequence and/or exome array analysis, we used complementary approaches with a range of available genetic data resources to evaluate their contribution to the association signals of previously reported lead SNVs. If the previously reported lead SNV (or a good proxy, $r^2 \geq 0.8$) was genotyped on the exome array, we performed reciprocal conditional analyses with the available exome array data. Within each study, we repeated EMMAX analyses in GWAS loci, including additively coded genotypes at the previously reported¹³ lead SNV or genome-wide significant coding variant as an additional covariate in the regression model. Across studies, we performed fixed-effects meta-analysis of association summary statistics at each variant based on: (i) inverse-variance weighting of score EMMAX beta/SE; (ii) sample size weighting of score EMMAX directed p -values. If the previously reported lead SNV (or a good proxy) was not genotyped on the exome array, we performed approximate reciprocal conditional analysis, implemented in GCTA⁷⁷, using genome-wide meta-analysis association summary statistics from 12,971 T2D cases and 34,100 controls from the combined GoT2D integrated panel and imputed data. Patterns of LD between variants were estimated using a subset of the GoT2D integrated panel, restricted to 2,389 individuals with pairwise genetic relationship < 0.025 , as defined by the GCTA A statistic⁷⁸. Finally, we interrogated 99% credible sets of variants at each GWAS locus, which together represent $\geq 99\%$ of the probability of driving each association signal. We determined whether the coding variant at each locus was included in the credible set for the association signal for the previously reported lead SNV, and recorded its rank.

3. Enrichment of exome association signals in GWAS

To define T2D-associated intervals, we first identified all SNVs associated with T2D in published genome-wide association studies (GWAS) by searching literature and the NHGRI GWAS catalog (see also **2.4.3**). We identified 143 autosomal SNVs, with some associated in more than one ancestry (167 SNV-ancestry pairs). For each SNV-ancestry pair, we identified the most distant pair of SNVs with $r^2 > 0.5$ in 1000 Genomes Phase I data, using the appropriate continental subset of 1000 Genomes samples (EUR, AMR, or ASN). We used 1000 Genomes data, rather than our own exome sequence data, because most reported associations for T2D are with common, intergenic SNVs. We then extended each region of interest by moving out 0.02 cM from those two SNVs (to encompass nearby recombination hotspots), and added an additional 300kb upstream and downstream. We merged overlapping intervals, yielding 81 unique associated regions, and identified 634 genes completely or partially included within associated regions. In single-variant analyses, we analyzed 3,147 non-

synonymous variants within these genes in the combined exome sequence and exome array datasets, using a Bonferroni corrected significance threshold of $\alpha=0.05/3,147=1.6\times 10^{-5}$. We considered gene-level association statistics from exome sequence for these 634 genes using a Bonferroni-corrected significance threshold of $\alpha=0.05/634=7.9\times 10^{-5}$.

We note that by reducing the stringency of the significance threshold for variants within GWAS loci, we increase the 'experiment-wise' type I error rate across the entire exome. Assuming that 3% of 100,000 coding variants interrogated in this study map to T2D GWAS loci, as defined above, we would need to change the threshold of significance outside of these regions to $p<2.1\times 10^{-8}$ to maintain an 'experiment-wise' type I error rate of 5%.

4. Testing for 'synthetic associations' at T2D loci in GoT2D genome sequence data

To identify low-frequency or rare variants that could potentially define synthetic associations, we analyzed the ten T2D loci at which a previously-reported tag SNV achieved $p<0.001$ in our single-variant analysis of the genome sequence dataset. We defined as candidates at each locus all low-frequency or rare variants (excluding singletons) within a 5Mb window (centered on the prior GWAS signals) and tested for synthetic associations caused by either (1) a single low-frequency or rare variant or (2) multiple low-frequency or rare variants on a common haplotype.

To identify synthetic associations driven by a single low-frequency or rare variant at each of the ten loci, we performed a series of conditional analyses in which we tested for association between gene dosage at the previously reported GWAS index SNV and T2D risk via logistic regression, while including each candidate low-frequency or rare SNV (excluding singletons) as an additional covariate, one-by-one. If inclusion of the low-frequency or rare variant resulted in a conditional association $p>0.05$ for the tag SNV, we considered the common-variant association signal a potential synthetic association.

To identify synthetic associations based on sets of low-frequency or rare variants, we extended this approach. We (1) defined common haplotypes segregating at each T2D locus; (2) identified all low-frequency or rare (excluding singletons) variants occurring on T2D-associated haplotypes (haplotypes on which the T2D-associated GWAS index SNV minor allele is present); and (3) asked whether any combination of these low-frequency or rare variants could explain the effect observed at the T2D GWAS index SNV. We carried out these analyses restricting attention to protein-coding variants within the window and then again for all low-frequency and rare SNVs in the 5Mb window. To define common haplotypes at each locus, we used the phased whole-genome sequence data. We

first employed the phased genotypes for common (MAF>5%) variants segregating in the interval between recombination hotspots at the locus (to minimize the number of recombinant haplotypes identified). We next identified the haplotypes on which the T2D-associated (risk or protective) GWAS index SNV minor allele was present. We then assembled the set of low-frequency and rare variants from across the 5Mb interval which occurred on the background of these T2D-associated common-variant haplotypes. Due to recombination and imperfect phasing, low-frequency or rare (excluding singletons) variants are often observed on more than one haplotype background. We included all low-frequency or rare variants that occurred more frequently on a T2D-associated haplotype than on other haplotypes.

From this pool of low-frequency and rare variants, we considered only variants with the same direction of effect as the common GWAS index SNV minor allele, as required by the synthetic association hypothesis, which posits that low-frequency or rare variants of larger effect than the common SNV could induce a weaker association signal. We then used a greedy algorithm to select the low-frequency or rare variant which, when added to the index GWAS SNV's dosage in a logistic regression, most reduced the residual effect remaining at the index SNV, as measured by estimated conditional odds ratio. We repeated this process, adding variants to the model, until the estimated effect at the index SNV genotype or gene dosage changed sign, representing no residual effect of the index SNV. At each locus, we also counted the number of variants required to increase the association p-value at the GWAS index SNV beyond the nominal $p=0.05$ significance threshold (**Extended Data Table 8**).

5. Credible set analysis of GoT2D genome sequence data

At 78 of the 80 T2D GWAS loci (**2.4.3**), the previously reported index SNV had MAF>1% in our GoT2D genome-sequenced sample. At these 78 loci, we constructed credible sets of common variants that, with some minimum specified probability (e.g. $\geq 99\%$), contain the variant causal for the corresponding association signal. Our analysis assumes a single causal SNV per signal and that the SNV was genotyped^{28,29}. We constructed credible sets for up to two independent association signals at each locus; at 5 loci with multiple independent ($r^2 < 0.10$) GWAS index SNVs, we constructed two distinct credible sets.

For each GWAS index SNV, we identified the set of common variants with $r^2 \geq 0.10$ with the index SNV within a 5Mb window centered on the index SNV. For each variant in this set, we calculated the posterior probability of being causal²⁹. We first calculated an approximate Bayes' factor (ABF) for

each variant as:

$$ABF = \sqrt{1-r} e^{rz^2/2}$$

where $r=0.04/[SE^2+0.04]$, $z=\beta/SE$, and β and SE are the estimated effect size (log odds ratio) and its standard error from logistic regression. We then calculated the posterior probability for each variant as ABF/T , where T is the sum of the ABF values over all candidate variants across the interval. This calculation assumes a Gaussian prior with mean 0 and variance 0.04 for β , the same prior employed in the commonly used single-variant association program SNPTTEST⁷¹.

We based the analysis on the genome-wide meta-analysis results, since most common variants were included in this analysis, and sample sizes were significantly larger than for the genome sequence data alone.

We calculated the effective imputed sample size for each variant in the meta-analysis data as

$N_{eff} = \sum_{j=1}^{13} r_j^2 n_j^{eff}$, where r_j^2 is the imputation quality and n_j^{eff} is the effective sample size for imputation cohort j . To ensure approximately uniform sample size across variants, we considered to be well-imputed only those variants with effective imputed sample size (N_{eff}) $\geq 80\%$ of the maximum observed across all variants in the window.

Indels were not imputed or meta-analyzed in this study, and $<2\%$ of common SNVs were not well-imputed by the above effective sample size criterion. To include these common variants while using the most precise estimates available, we calculated posterior probabilities separately from each genome-wide data source. Where an indel from the sequence dataset had a SNV proxy in high LD ($r^2 \geq 0.80$) in the meta-analysis dataset, we used the proxy's information instead. Where a common SNV that was poorly imputed had high-quality association data from the genome sequence data alone, the posterior probability from the genome sequence dataset was used instead. In each case, the final posterior probabilities for all SNVs were re-scaled such that their sum across a locus equaled one.

We used these final posterior probabilities to rank variants in decreasing order. To define credible sets of a specified level (e.g. 99%), we included variants with highest final posterior probabilities until their sum reached or exceeded that level (**Supplementary 28**).

6. Genome enrichment analyses of the GoT2D genome sequence data

6.1. Genomic annotation

We collected genome annotation data from several sources. First, we obtained gene transcript

information from GENCODEv14⁷⁹. For protein-coding genes, we included transcripts with a protein-coding tag that either were present in the conserved coding DNA sequence (CCDS) database or had experimentally confirmed mRNA start and end; we then included 5' UTR, exon, and 3' UTR regions from the resulting transcripts. For non-coding genes, we included transcripts with a lncRNA, miRNA, snoRNA, or snRNA tag.

Second, we defined regulatory chromatin states in 12 cell types. We collected sequence reads generated for the following assays: H3K4me1, H3K4me3, H3K27ac, H3K27me3, H3K36me3, and CTCF ChIP, in 9 ENCODE cell types (GM12878, K562, HepG2, Hsmm, HUVEC, NHEK, NHLF, hESC, HMEC)³⁰, pancreatic islets³³, and hASC (adipose stromal cell) pre- and mature adipocytes³¹. We mapped reads to hg19 using BWA⁵⁰ and used the resulting mapped reads for all cell types to call regulatory states using ChromHMM⁸⁰, assuming ten states. We then assigned names to the resulting state definitions: (1) H3K4me3, H3K27ac (active promoter); (2) H3K4me3, H3K27ac, H3K4me1 (active enhancer 1); (3) H3K27ac, H3K4me1 (active enhancer 2); (4) H3K4me1 (weak enhancer); (5) H3K27me3, H3K4me3, H3K4me1 (poised promoter); (6) H3K27me3 (repressed); (7) low/no signal 1; (8) CTCF (insulator); (9) low/no signal 2; and (10) H3K36me3 (transcription).

Third, we obtained transcription factor binding ChIP sites from three sources: 141 proteins from ENCODE³⁰, 5 from Pasquali et al.³³, and 1 from Mikkelsen et al.³¹.

From gene transcript data we defined CDS (protein coding transcript exons); ncRNA (non-coding RNA transcripts); and 3' and 5' UTR (UTR regions of coding transcripts). From chromatin state data for each of the 12 cell types we identified active enhancers (pooled active enhancer 1 and 2 elements); weak enhancers; and active promoters. From transcription factor binding sites we defined transcription factor binding sites (TFBS) (sites pooled across all factors). This resulted in a total of 41 annotation categories (**Extended Data Figure 9**).

6.2. Enrichment of genome annotation

We jointly modeled variants in credible sets using T2D association and the functional annotation classes using the method described by Pickrell³⁶. First, we tested each annotation individually and identified the annotation that most improved the model likelihood. We then iteratively added annotations in this manner until the likelihood did not increase further. Using this set of annotations, we tested a range of penalized likelihoods (from 0-1 in .01 increments) using 10-fold cross-validation, and identified the penalty that gave the best cross-validation likelihood. Using this penalty, we then iteratively dropped annotations to identify the model with the maximal cross-validation likelihood. The resulting model included coding exons, TFBS, hASC mature adipose active

enhancers and promoters, pancreatic islet active and weak enhancers and active promoters, hASC pre-adipose active and weak enhancers, NHEK active enhancers, NHLF active enhancers, K562 weak enhancers, HMEC weak enhancers and active promoters, H1-hESC active promoters, ncRNA, and 5' and 3' UTR (**Extended Data Figure 9**). Finally, we used this model to update posterior probabilities for each variant and re-calculate 99% credible sets.

7. Gene enrichment analyses in the GoT2D+T2D-GENES exome sequence data

We first used the SMP (statistics/matrix/permutation) gene-set enrichment procedure implemented in the PLINK/Seq package (<http://atgu.mgh.harvard.edu/plinkseq/>). This approach calculates enrichment statistics for large sets of genes to establish whether case-enrichment of rare variants is preferentially concentrated in a particular set of genes, controlling for any exome-wide/baseline difference in case and control rates. The procedure uses gene-based association statistics, and forms sums of these statistics over all genes in a set, the significance of which is evaluated by permutation. We considered the relative enrichment statistic, SSET/SEXOME, with significance evaluated empirically (10,000 replicates) based on the null distribution of this ratio. The reported effect sizes from the gene-set enrichment analysis are estimates of the unconditional odds ratio that do not take exome-wide differences in case/control rates into account⁶⁹. We selected 18 'premium' sets of genes (**Supplementary 32**) that reflect the current knowledge of pathways (N=15) involved in type 2 diabetes and the three sets of genes involved in monogenic form of diabetes defined above: 'Monogenic All' (N=81); 'Monogenic Primary' (N=28); and 'Monogenic OMIM' (N=13). We restricted these analyses to singleton and ultra-rare (MAF<0.1%) protein-truncating variants.

We then used biological knowledge to test for enrichment of association signal across established sets of genes from Gene Ontology, KEGG, Reactome, and Biocarta collections from MSigDB (version 4.0) as well as a number of hand-curated gene-sets (**Supplementary 32**) that had been generated for the SMP analyses. These analyses calculated measures of gene-set enrichment from gene-level association results (i.e. from SKAT-O) by means of a pre-ranked GSEA⁸¹ method (version 2.0.13), which consists of a weighted Kolmogorov-Smirnov (random bridge) statistic. In our analysis we performed 10,000 permutations on gene-set sizes from 5 to 5,000 genes.

8. Investigation of genes implicated in Mendelian forms of diabetes in the exome data

We first curated a list of 81 genes termed the ‘Monogenic All’ gene-set (**Supplementary 22**), consisting of genes with pathogenic mutations reported to co-segregate with diabetes or a syndrome associated with an increased prevalence of diabetes. Two subsets of the ‘Monogenic All’ gene-set were then additionally defined: the ‘Monogenic Primary’ gene-set (N=28), consisting of genes with mutations leading to diabetes as a primary feature, and the ‘Monogenic OMIM’ gene-set (N=13), consisting of genes linked to Maturity Onset Diabetes of the Young (MODY) or Neonatal Diabetes in the OMIM catalog (entry #606391 and #606176). In addition to examining the significance of single-variant and gene-based tests within these gene-sets, we also performed an aggregate analysis of all variants in the gene-set. For each of the three gene-sets, we constructed five variant lists by applying the same four masks as in the exome-wide gene-level analysis (PTV-only, PTV+missense, PTV+NS_{broad} and PTV+NS_{strict}), as well as an additional mask containing all variants reported as ‘high confidence’ and ‘disease-causing’ in the Human Gene Mutation Database (HGMD), annotated using Biobase ‘GenomeTrax’ software (<http://www.biobase-international.com/product/genome-trax>). We then analyzed each of the fifteen variant lists with the SKAT-O test, using the same meta-analysis procedure and covariates as in the exome-wide gene-based analysis. To obtain effect-size estimates, for each variant list we applied a collapsing burden test, in which logistic regression of T2D status was performed on individual genotypes encoded as 0 (if they carried no variants in the list) or 1 (if they carried at least one variant in the list). Effect size estimates and standard errors were determined using the Firth penalized likelihood method. Analysis in the exome array dataset was performed by first generating fifteen variant lists based on the content of the exome array, computing the collapsing burden test for each cohort, and then combining associations and effect size estimates using an inverse variance weighted meta-analysis. To compare the age of diagnosis of variant carriers to those of non-carriers, we used a two-sided t-test.

9. Protein-protein interaction analyses in the exome data

We performed data-driven extraction of association signal enriched sub-networks (rather than relying on pre-defined gene-sets) from protein-protein interaction (PPI) data. We used two different approaches, both run using the curated PPI database InWeb3⁸².

The first approach consists of two steps. First, the entire human PPI network was searched for protein complexes (clusters) using the algorithm implemented in clusterONE⁸³, which identifies

protein complexes with high cohesiveness. The method was run with default parameter settings (0.3 as density threshold, 0.8 as merging threshold, and 2 as the penalty-value node), and with the --fluff option activated, which allows the addition of highly connected boundary nodes to the cluster.

Second, gene-based association p -values derived from SKAT-O analyses of the 12,844 multiethnic exome sequences were aggregated, using Fisher's method, for the genes encoding each of the proteins within a cluster to generate a 'cluster association' statistic.

An empirical p -value for the significance of these aggregated cluster association statistics was derived by comparing each cluster to a large number of complexes of the same topology, but composed of randomly sampled proteins. Specifically, a background distribution was obtained for each protein complex as follows: each protein in the cluster was randomly substituted by a different protein represented in the InWeb3 database, matched for number of minor allele carriers in the data set. SKAT-O p -values were assigned to each protein from the exome sequencing results, and an aggregated p -value was obtained for each pseudo-complex using Fisher's method, as above. This process was repeated 100,000 times, and the empirical p -value for each complex was calculated as the proportion of the iterations for which the Fisher's p -value of the observed complex was more significant than that of p -values for the pseudo-complexes. This procedure was repeated for all gene-level masks (PTV-only, PTV+missense, PTV+NS_{strict} and PTV+NS_{broad}).

To test the study-wide significance of apparently associated clusters, we used two permutation designs. In the first design, we generated 100,000 pseudo-complexes for each cluster, replacing each protein within each cluster with one protein from InWeb3, matched for the number of minor allele carriers in the data set. We calculated the number of permuted datasets which generated any 'pseudocluster' association p -value more significant than our most enriched cluster. In the second design, we used a Monte-Carlo algorithm to generate 10,000 random PPI networks, with the same degree as observed in the InWeb3 database, ran clusterONE on each, and once again compared the distribution of 'best' cluster association p -value with that observed in the real data.

The second approach uses the dense module searching algorithm (a heuristic 'greedy' method) described in dmGWAS⁸⁴, where a module is defined as a sub-network within the whole network if it contains a locally increased proportion of low p -value genes. This method differs from the earlier method in using the association p -values, in combination with the PPI data, to construct the networks. The module is grown for each protein in the PPI by adding the neighboring nodes within a pre-defined distance ($d=2$) that can yield a maximum increment of the module score $Z_m^{(k)} = \sum Z_i / \sqrt{k}$ for module m , where k is the number of genes in the module and Z_i is calculated from the p -value of

exome gene-based tests using an inverse normal distribution function. The addition of neighborhood nodes is stopped when the increment is less than 10% of $Z_m^{(k)}$ (that is, $Z_m^{(k+1)} < Z_m^{(k)} + Z_m^{(k)} \times 0.1$). As with the clusterONE approach, this procedure was conducted for all four exome gene-based level masks.

To evaluate whether the top ranked-modules are significantly associated with T2D, we permuted case-control status across the 12,844 exomes (maintaining ethnic strata) 10,000 times and generated 10,000 SKAT-O gene-based association tests on all genes in the top 15 modules (once for each gene-based variant mask, 40,000 in total). During each permutation, Z_m was re-calculated for each module, and a set of empirical p-values was obtained by comparing the p-value of the original module to these modules with the SKAT-O results from the swapped labels. Following the above procedure, all 15 top modules were found significantly enriched for the PTV+NS_{strict} and PTV+NS_{broad} gene-based variant masks ($p < 10^{-4}$, after the 10,000 case-control permutations).

10. Modelling disease architecture

10.1. T2D liability risk and architecture bounding in the exome array data

We used a Bayesian framework implemented in R to compute the probability that each variant explains more than a defined amount of the T2D-risk liability-scale variance (LVE). The joint distribution in the MAF-OR space is computed by assuming a T2D prevalence of 8% and beta and normal distributions for the MAF and the odds ratio (OR) respectively. The OR is calculated with reference to the minor allele. The MAF is adjusted to take account of apparent allele frequency heterogeneity between cohorts (subjects from missing cohorts are excluded from calculations). Analyses are restricted to variants with MAF > 0.1% since the representation of variants with MAF below this threshold on the exome array is poor. The probability is obtained by numerically integrating over the joint distribution for MAF-OR combinations that explain more than the defined amount of liability-scale variance. For bounding the maximum number of variants that could contribute to T2D risk variance, we performed a sensitivity analysis on the 88 known T2D index SNVs present on the exome array to define the thresholded variance explained and the probability: this analysis shows that for a probability of > 0.8 to explain 0.01% of the T2D risk variance, we were able to identify 91% of these known T2D SNVs. Ranges of OR and MAF consistent with 80% power to detect single-variant association in this dataset (for exome-wide significance, $p < 5 \times 10^{-7}$) were calculated to reflect the fact that differences in sample size for individual variants (due to differences in allele frequency distribution and genotyping QC) also influence power. The relationship between

power and LVE differs for risk and protective alleles because of unequal numbers of cases and controls.

10.2. Genetic architecture simulations based on GoT2D data and results

10.2.1. Range of simulated disease models

Following our previously published framework³⁸, we conducted population genetic simulations of T2D architecture using the forward simulation program ForSim⁸⁵. We assumed T2D prevalence 8% and heritability ~45%, and chose the mutation rate, recombination rate, a gamma distribution of selection coefficients, and other parameters of demographic history by fitting the simulated site frequency spectrum to empirical high coverage exome sequence data from GoT2D.

We then considered a wide range of disease models by varying two parameters: coupling parameter τ which regulates how strongly selection against a disease-causing allele depends on the per-allele disease risk⁸⁶; and target size T , the summed lengths of the genomic regions within which mutations can influence T2D risk. Specifically, a variant's additive contribution to disease risk g is given by $g = s\tau(1 + \epsilon)$ where s is the selection coefficient under which the variant evolves and ϵ is drawn from a normal distribution³⁸.

By varying τ and T , we generated a wide range of joint distributions for allele frequency and effect size. In total, we evaluated 12 models: $\tau = 0, 0.1, 0.3, \text{ and } 0.5$ crossed with $T = 750\text{kb}, 2.0\text{Mb}, \text{ and } 3.75\text{Mb}$. Under models with higher selection against strongly deleterious alleles (larger τ), rare variants explain the bulk of heritability and can have large effects, while under models with weak dependence (smaller τ), common variants explain the bulk of heritability and rare variants collectively have weaker effects. Although we had previously excluded many models as producing predictions inconsistent with observed sibling relative risk, GWAS, and linkage results, prior work showed that models varying widely in the proportion of total heritability attributable to rare versus common variation were still plausible⁸⁷. In this study, we explored whether the space of plausible disease models could be further constrained using whole genome sequence, imputation, and meta-analysis results.

10.2.2. Simulation procedure

ForSim enables simulation of variants across user-specified loci in large populations. Inputs include a demographic history (trained on European sequence data) and a gamma distribution of selection coefficients for a subset of variants under natural selection. We simulated genotypes for a current population of effective size 500,000 individuals³⁸ and selected potential disease risk variants from

those under selection appropriate to the intended target size. Each risk variant received a disease-specific effect size depending on the selection coefficient under which it evolved and the assumed degree of dependence between selection and effect size. Each individual was then designated as case or control depending on his/her cumulative genetic risk score plus a random environmental risk component chosen to achieve the estimated T2D heritability of ~45%. From this population simulated with both phenotypes and genotypes, we selected appropriate numbers of cases and controls and conducted single-variant association tests in order to compare the distribution of p-values from simulation to that observed in the current study. Results shown are the average of 25 independent simulation replicates for each disease model.

10.2.3. Comparison of simulated outcomes to empirical T2D results

We focused on comparing simulated outcomes under three disease models, each of which were previously found to be consistent with sibling relative risk, GWAS, and linkage results for T2D, but vary widely in causal variant properties (**Fig. 4**): a rare-variant model in which rare variants explain ~75% of T2D heritability (small target size $T=750\text{kb}$ and moderate dependence between effect size and selection $\tau=0.5$), an intermediate model in which rare, low-frequency, and common variants all contribute significantly to T2D heritability ($T=2.0\text{Mb}$ and $\tau=0.3$), and a common polygenic model in which common variants explain ~75% of T2D heritability ($T=3.75\text{Mb}$ and weak dependence $\tau=0.1$). We first compared the simulated outcomes of a whole-genome sequencing study in ~3K samples under each model. All three models predicted similar distributions of variant association test statistics using the sequenced individuals alone (data not shown).

However, the predictions began to diverge when we simulated imputation into GWAS samples and studied the distribution of test statistics after meta-analysis. For each simulated model, we sampled 14,175 cases and 14,175 controls (to match the effective sample size of the actual imputation cohorts used for meta-analysis). Because genotyping accuracy in simulated samples is perfect (unlike in imputation), we calculated average imputation quality as a function of MAC in the empirical data (using the r^2 value reported by the imputation software that was used in each cohort). We then corrected, for each variant, the association test statistic in simulated data by multiplying the chi-squared value by the average imputation r^2 for the variant MAC. We then re-computed association p-values from the corrected chi-squared statistics to compare p-value distributions in simulated versus empirical data. We plotted the distribution of association p-values for variants of different frequency classes in a quantile-quantile (QQ) plot, and compare these curves to the empirical T2D results (**Fig. 4**). Focusing on low-frequency variants, we also asked how many unique low-frequency

signals achieved significant association to T2D risk under each simulated model, and compared these quantities to empirical observation (**Fig. 4**). These analyses demonstrate that the intermediate and rare-variant models produce an excess of association signal among low-frequency variants compared to observation, whereas the common polygenic model is consistent with the genome-wide distribution of association signals observed.

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EXTENDED DATA ITEMS: LEGENDS

Extended Data Figure 1 | Summary of samples and quality control procedures. This figure summarises data generation for whole genome sequencing (GoT2D), exome sequencing (GoT2D and T2D-GENES) and exome array genotyping (DIAGRAM). In addition, GoT2D whole genome sequence data was imputed into GWAS data from 44,414 subjects of European descent.

Extended Data Table 2 | Summary information for samples sets used in the association analyses.

Extended Data Table 3 | Counts and properties of variants identified in sequenced subjects. a.

Variant numbers for the 2,657 individuals with whole genome sequence data passing QC and included in the association analysis data set; **b.** Variant numbers are provided for the 13,008 individuals passing initial rounds of QC from which further QC defined the 12,940 subjects included in the association analysis data set. Private refers to variants seen in only a single ancestral group; cosmopolitan to variants seen in all five major ancestral groups.

Extended Data Figure 4 | Power for single and aggregate variant association. a-g. Power to detect single-variant association ($\alpha=5 \times 10^{-8}$) at varying minor allele frequency (x-axis) and allelic odds-ratio (y-axis) for seven effective sample size (Neff) scenarios relevant to the genomes (**a-c**) and exomes (**d-g**) component of this project. **a.** variant observed in 2,657 samples (the effective size of the GoT2D integrated panel); **b.** variant observed in 28,350 samples (the effective size of the imputed data set); **c.** variant observed in the GoT2D integrated panel and the imputed data set (effective sample size 31,007); **d.** ancestry-specific variant in 2,000 samples (the size of each of the non-European exome sequence data sets); **e.** European specific variant in 5,000 samples (the combined size of the European exome sequence data sets); **f.** variant observed with shared frequency across all ancestry groups in 12,940 samples (the size of the combined exome sequence data set); and **g.** variant observed in the combined exome array and sequencing data set (effective sample size 82,758). **h-i.** Power for gene based test of association (SKAT-O) according to liability variance explained. In **h**, 50% of the variants contribute to disease risk while the remaining 50% have no effect on disease risk; in **i**, 100% of the variants contribute to disease risk. For each, sample sizes considered are 2,000 (ancestry-specific effects; green) and 12,940 (ancestry-shared effects; blue). Power is shown for two levels of significance ($\alpha=2.5 \times 10^{-6}$ and $\alpha=0.001$). From these simulation studies, it is clear that under the optimistic model, where effects are shared across all ethnicities (blue line) and all variants

contribute, power is >60% for 1% variance explained and $\alpha=2.5 \times 10^{-6}$. However, power declines rapidly if either criterion is relaxed.

Extended Data Table 5 | Characterization of variant associations through conditional analysis. For each locus, significantly associated SNVs are presented. Unconditional *p*-values are given in italics, and conditional *p*-values are shown for each pair of SNVs (*p*-values are for SNVs in the “Variant” column, with SNVs listed in header included as covariates in association analysis). The *IRS1* and *PPARG* non-coding associations were characterized using exact conditional analysis in 38,738 samples from the GoT2D genome-wide imputed meta-analysis. Conditional analysis for coding variant associations was, for most loci, restricted to the exome array genotypes (28,305 cases, 51,549 controls). At *THADA* and *RREB1*, neither the non-coding lead GWAS SNVs nor close proxies were typed on the exome array, so approximate conditional analyses were undertaken using GCTA in 44,414 samples from the GoT2D genome-wide imputed meta-analysis (**Methods**). For several of these loci, unconditional association *p*-values for these loci do not reach genome-wide significance as sample sizes are smaller. At the *GPSM1* locus, the previously reported GWAS SNV was not available on exome array and too poorly imputed in the GoT2D meta-analysis to allow meaningful inference. *Conditional analysis was performed once for rs78124264 with all three previously known GWAS variants included as covariates. ¶ Non-coding GWAS lead variant. n.d. indicates “not determined.”

Extended Data Figure 6 | Single variant analyses. Manhattan plot of single-variant analyses generated from **a.** exome sequence data in 6,504 cases and 6,436 controls of African American, East Asian, European, Hispanic, and South Asian ancestry; **b.** exome array genotypes in 28,305 cases and 51,549 controls of European ancestry; and **c.** combined meta-analysis of exome array and exome sequence samples. Coding variants are categorized according to their relationships to the previously reported lead variant from GWAS region. Loci achieving genome-wide significance only in the combined analysis are highlighted in bold. The *HNF1A* variant reaching genome-wide significance in the combined analysis is a synonymous variant (Thr515Thr). The dashed horizontal line in each panel designates the threshold for genome-wide significance ($p < 5 \times 10^{-8}$).

Extended Data Figure 7 | Classification of coding variants according to their relationship to reported lead variants for each GWAS region. The ideogram shows the location of 25 coding variant

associations at 16 loci described in the text. The number in each circle corresponds to the number of associated variants at each locus. Variants are grouped into five categories based on inferred relationship with the GWAS lead variant. For some of these categories, the figure includes representative regional association plots based on exome array meta-analysis data from 28,305 cases and 51,549 controls. The locus displayed for each category is designated in bold. The first plot in each panel shows the unconditional association results; middle plot the association results after conditioning on the non-coding GWAS SNP; and the last plot the results after conditioning on the most significantly associated coding variant. Each point represents a SNP in the exome array meta-analysis, plotted with their p-value (on a $-\log_{10}$ scale) as a function of the genomic position (hg19). In each panel, the lead coding variant is represented by the purple symbol. The color-coding of all other SNPs indicates LD with the lead SNP (estimated by European r^2 from 1000 Genomes March 2012 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). Gene annotations are taken from the University of California Santa Cruz genome browser. GWS: genome-wide significance. *Seven variants, three at *ASCC2*, and one each at *THADA*, *TSPAN8*, *FES* and *HNF4A* did not achieve genome-wide significance themselves, but are included because they fall into genes and/or regions with other significant association signals (see text).

Extended Data Table 8 | Testing for synthetic associations across GWAS-identified T2D loci. Gene names refer to protein-coding transcript(s) closest to the index SNV. Reported index SNVs are the previously-reported GWAS variants (in European populations) with the strongest association signal in the GoT2D sequencing data (n=2657). Relative likelihoods are based on causal models with only the chosen low-frequency and rare missense variants, relative to models with only the GWAS index SNV, assessed using the Akaike Information content (AIC) of each regression model, calculated as $\exp[(AIC_{\text{index}} - AIC_{\text{low-frequency or rare}})/2]$. n_1 provides the number of low-frequency or rare variants required for the residual odds ratio at the GWAS index SNV, after joint conditioning on the low-frequency and rare variants, to switch direction of effect. n_2 provides the number of low-frequency or rare variants required for the association p-value remaining at the GWAS index SNV, after joint conditioning on the low-frequency and rare variants, to exceed 0.05.

Extended Data Figure 9 | Genome enrichment analysis in GoT2D whole genome sequence data (n=2657) **a**, Functional annotation categories were defined using transcription, chromatin state and transcription factor binding data from GENCODE, ENCODE and other studies. **b**, T2D association

statistics for variants at each T2D locus were jointly modelled with functional annotation using fgwas. In the resulting model we identified enrichment of coding exons (CDS), transcription factor binding sites (TFBS), mature adipose active enhancers and promoters (hASC-t4 EnhA, TssA), pancreatic islet active and weak enhancers (HI EnhA, EnhWk), pre-adipose active and weak enhancers (hASC-t1 EnhA, EnhWk), embryonic stem cell active promoters (H1-hESC TssA) and 5' UTR. Dots represent enrichment estimates and horizontal lines the 95% confidence intervals. **c.** At the *CCND2* locus, three variants not present in HapMap2 have a combined 90% posterior probability of being causal (rs4238013, rs3217801, rs73040004). One of these variants, rs3217801, is a 2-bp indel that overlaps an islet enhancer element.

Extended Data Figure 10 | Low frequency variants in exome array data. Results from meta-analysis of 43,045 low-frequency and common coding variants on the exome array (assayed in 79,854 European subjects). **a.** Observed allelic ORs as a property of allele MAF. Variants missing in >8 cohorts or polymorphic in only one cohort were excluded. Colored lines represent contours for liability variance explained. Regions shaded grey denote ranges of OR and MAF consistent with 80% power (in this case, at $\alpha=5 \times 10^{-7}$) to detect single-variant associations in this data set (given the observed range of missing data). Variants with a black collar are those highlighted by a bounding analysis as having a probability >0.8 of having LVE >0.1%; **b.** Distribution of each variant in the MAF/OR space was computed by assuming T2D prevalence of 8% and a beta and normal distribution for MAF and OR respectively. Probability is obtained by integrating the joint MAF-OR distributions over ranges of LVE; **c.** Single variant association, liability and bounding results for the known T2D GWAS variants on the exome array (**Methods**).

FIGURE 1 (low resolution for review purposes)

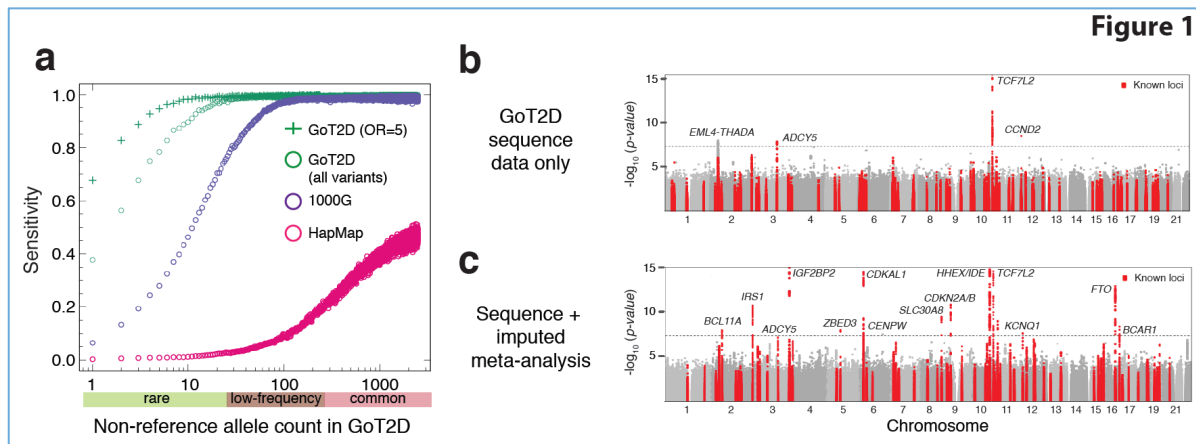


FIGURE 2 (low resolution for review purposes)

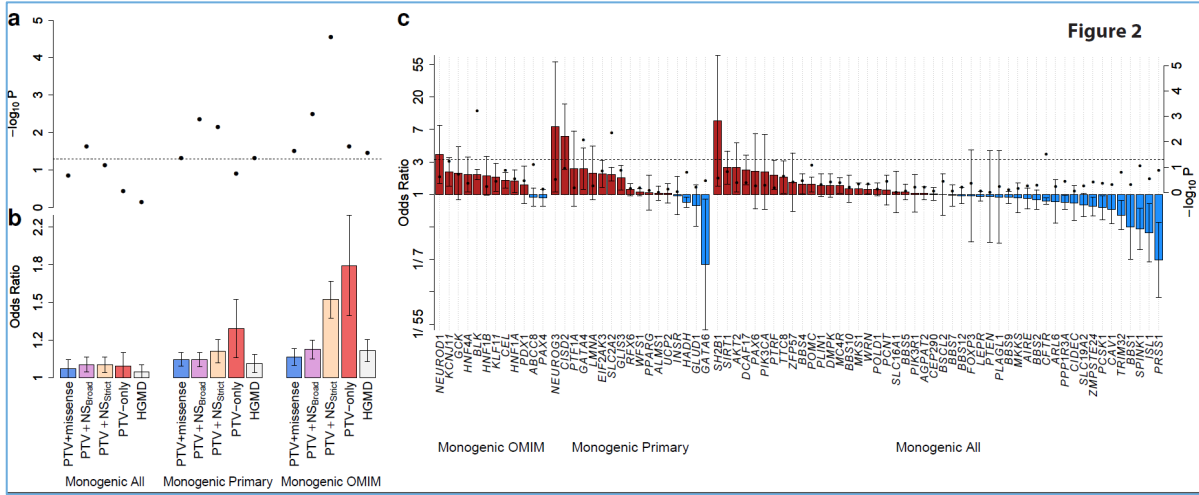


FIGURE 3 low resolution for review purposes)

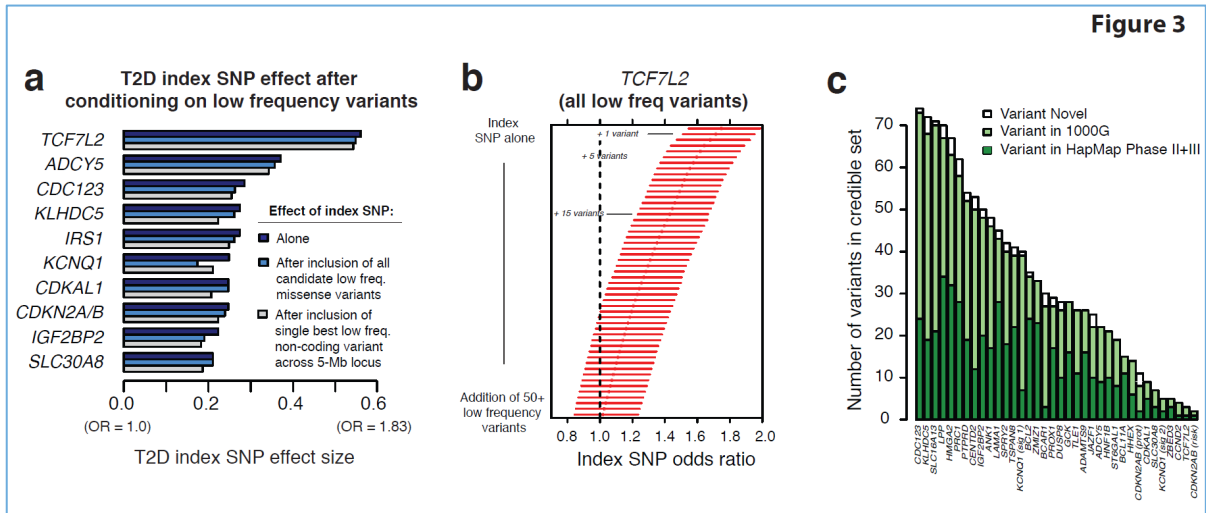


FIGURE 4 (low resolution for review purposes)

