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Large-scale association analyses identify new loci influencing glycemic traits and provide insight into the underlying biological pathways

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Abstract

Through genome-wide association meta-analyses of up to 133,010 individuals of European ancestry without diabetes, including individuals newly genotyped using the MetaboChip, we have raised the number of confirmed loci influencing glycemic traits to 53, of which 33 also increase type 2 diabetes risk ($q < 0.05$). Loci influencing fasting insulin showed association with lipid levels and fat distribution, suggesting impact on insulin resistance. Gene-based analyses identified further biologically plausible loci, suggesting that additional loci beyond those reaching genome-wide significance are likely to represent real associations. This conclusion is supported by an excess of directionally consistent and nominally significant signals between discovery and follow-up studies. Functional follow-up of these newly discovered loci will further improve our understanding of glycemic control.

The Meta-Analyses of Glucose and Insulin-related traits Consortium (MAGIC) previously undertook meta-analyses of genome-wide association studies (GWAS) of glycemic traits in

non-diabetic individuals, leading to the discovery of multiple associated loci: 16 for fasting glucose concentrations, two for fasting insulin concentrations and five for post-challenge glucose concentrations (2hGlu)¹⁻³. These and subsequent studies highlighted important biological pathways implicated in glucose and insulin regulation^{4,5}. They also demonstrated that some, but not all, loci associated with glycemic traits in non-diabetic individuals also affect the risk of type 2 diabetes (T2D)^{1,6}. Despite the success of these efforts, the identification of new loci was limited by *de novo* genotyping capacity and cost, such that only a limited number of promising loci from discovery analyses were taken forward to follow-up analyses (often those reaching a threshold of $\sim P < 10^{-5}$ in discovery). Therefore, it is likely that many additional associations with common, low penetrance variants remain to be found among SNPs not previously selected for replication^{7,8}.

The Illumina CardioMetaboChip (MetaboChip) is a custom Illumina iSELECT array of 196,725 SNPs developed to support cost-effective large-scale follow-up studies of putative association signals for a range cardiovascular and metabolic traits ($\sim 66,000$ SNPs) and to fine-map established loci ($\sim 120,000$ SNPs) (Supplementary Fig. 1)⁹. The $\sim 66,000$ follow-up SNPs were selected to enable genotyping of the most significant association signals for each of 23 metabolic traits contributed by a range of consortia. MAGIC contributed $\sim 5,000$ top ranking SNPs for fasting glucose, and $\sim 1,000$ each for fasting insulin and 2hGlu that had shown nominal association in discovery analyses ($P_{\text{discovery}} < 0.02$)^{1,2}.

In the present study, we combined newly available samples with genotype data for these 66,000 follow-up SNPs with previous discovery meta-analyses to discover new association signals with glycemic traits. This approach identified 41 glycemic associations not previously described^{1,2}: 20 for fasting glucose, 17 for fasting insulin and four for 2hGlu. This raises the number of associated loci to 36 for fasting glucose, 19 for fasting insulin and 9 for 2hGlu, explaining 4.8%, 1.2% and 1.7% of the variance in these traits, respectively. Of these 53 non-overlapping loci, 33 were also associated with T2D ($P < 0.05$), which while supporting the previous assertion of an imperfect correlation between these traits, also implicates new loci in the etiology of T2D and increases the overlap between glycemic and T2D loci.

RESULTS

Approaches to identify loci associated with glycemic traits

To follow up loci showing evidence of association ($P_{\text{stage one}} < 0.02$) in discovery GWAS, we investigated the 66,000 MetaboChip follow-up SNPs for association with fasting glucose, fasting insulin and 2hGlu. We combined in meta-analysis data from up to 133,010 (fasting glucose), 108,557 (fasting insulin) and 42,854 (2hGlu) non-diabetic individuals of European ancestry, including individuals from the previous meta-analyses^{1,2}, individuals from new GWAS and individuals newly genotyped on the MetaboChip array (Supplementary Fig. 2). All study characteristics are shown in Supplementary Table 1. Genome-wide association data for Filipino women were available (Supplementary Table 1), for which we report the effect directions and allele frequencies in Supplementary Tables 2a,b. Genome-wide significant ($P < 5 \times 10^{-8}$) association signals located more than 500 kb from, and not in LD (Hapmap CEU: $r^2 < 0.05$) with, any variant already known to be associated with the trait were considered novel. Associated loci are referred to by the name of the nearest gene, unless a more biologically plausible gene was nearby, or a nearby gene was previously associated with another trait. In such cases, we list the nearest genes in Supplementary Tables 2a-d. As body mass index (BMI) is a major risk factor for T2D and is correlated with glycemic traits, we also performed analyses adjusted for BMI.

Though not the main focus of this effort, given the increased variant density available on the Metabochip for established glycaemic loci, we investigated whether these data would enable fine-mapping of underlying functional variants¹⁻³. In these analyses, we included data from up to 53,622 individuals for fasting glucose, 42,384 for fasting insulin and 27,602 for 2hGlu from studies with Metabochip genotypes only. However, given the lack of samples from different ancestries and the absence of full conditional analyses, for the most part these analyses did not improve the resolution of association signals.

Beyond individual SNP investigations for each glycaemic trait, we also tested the hypothesis that gene-based analyses using VEGAS¹⁰ would identify genes that harbor multiple association signals, which individually did not reach genome-wide significance. Among the ~66,000 SNPs, we used VEGAS to pool the results for all SNPs within each gene (± 50 kb) to identify genes with more evidence of association than expected by chance (given gene size and linkage disequilibrium structure) by simulation, and significant after Bonferroni-correction for multiple testing ($P < 5 \times 10^{-6}$).

Fasting glucose

In analyses of up to 133,010 individuals, we identified 20 loci with genome-wide significant associations to fasting glucose ($P < 5 \times 10^{-8}$) (Table 1 and Supplementary Figs. 3 and 4) and confirmed previously established loci¹ (Supplementary Table 2e). Of these 20 loci, nine (in or near *IBKAP*, *LOC728489*, *WARS*, *KL*, *TOPI*, *P2RX2*, *AMT*, *RREB1* and *GLS2*) had not previously been associated with other metabolic traits (Box 1). Among these, *KL* (Klotho) is of particular interest. In addition to being associated with fasting glucose (but not fasting insulin), the glucose-raising allele is also associated with an increased risk of T2D (OR = 1.08 (1.04-1.11), $P = 1.1 \times 10^{-5}$) (Fig. 1). *KL* was first identified as a gene related to suppression of aging: its reduced expression was associated with reduced lifespan, as well as hypoglycemia¹¹. Despite further animal studies supporting a role for *KL* in glucose metabolism¹² and insulin sensitivity¹³, human studies have generally been small and inconclusive^{14,15}.

We also identified new associations with fasting glucose in regions previously associated with other metabolic traits or disease outcomes, including T2D^{6,16} (*ARAPI*, *CDKN2B*, *GRB10*, *CDKAL1*, *IGF2BP2* and *ZBED3*, which was identified in BMI-adjusted models) and 2hGlu² (*GIPR*), as well as confirming the recently identified signals for fasting glucose¹⁷⁻¹⁹ at *FOXA2*, *PPP1R3B*, *PCSK1* and *PDX1*. *FOXA2* is a forkhead transcription factor that regulates *PDX1* expression, while *PDX1* encodes a transcription factor critical for pancreatic development²⁰. *PDX1* mutations have been linked to MODY4 (ref. 21), pancreatic agenesis²² and permanent neonatal diabetes²³, although we observed no significant association with T2D in DIAGRAM Metabochip analyses²⁴ (Fig. 1).

Given the overlap between genetic loci for fasting glucose and other metabolic traits, we performed a systematic look-up of all glycaemic loci and their associations with other metabolic traits using data available through other consortia²⁵⁻²⁷. In DIAGRAM Metabochip analyses²⁴, 22 (>60%) of the now 36 genome-wide significant fasting glucose loci showed association ($P < 0.05$; FDR $q < 0.05$) with T2D (Fig. 1). In all cases, the glucose-raising allele was associated with increased risk of T2D, yet the fasting glucose effect size and T2D odds ratio were weakly correlated (Fig. 2a).

Gene-based analyses confirmed many of the loci identified in individual SNP analyses (Supplementary Table 3a) and identified another nine genomic regions (containing 14 genes) with significant association signals ($P < 5 \times 10^{-6}$), including some with biological candidacy, such as the *HKDC1* gene, encoding a putative hexokinase²⁸.

Fasting insulin

In 108,557 individuals, we identified 17 additional loci with genome-wide significant associations to fasting insulin and confirmed known associations¹. These newly identified loci include variants in or near *HIP1*, *TET2*, *YSK4*, *PEPD* and *FAM13A1* (Table 1, Box 1 and Supplementary Figs. 3 and 4), as well as SNPs near loci previously associated with other metabolic traits, including T2D⁶ (*TCF7L2*, *PPARG*), BMI²⁹ (*FTO*), waist-hip ratio²⁶ (WHR) (*LYPLAL1*, *RSPO3*, *GRB14*), triglycerides²⁷ (*ANKRD55-MAP3K1*) and adiponectin³⁰ (*ARL15*). We also confirmed the recent associations with fasting insulin at *GRB14*, *PPP1R3B*, *LYPLAL1*, *IRS1*, *UHRF1BP1* and *PDGFC*¹⁹. The *ANKRD55-MAP3K1* association is of interest as MAP3K1 regulates expression of IRS1 (ref. 31) as well as activation of NF- κ B^{32,33} and the JNK pathway³⁴, both centrally implicated in insulin resistance^{35,36}. Furthermore, data from DIAGRAM Metabochip analyses show that the insulin-raising allele at this SNP is strongly associated with increased risk of T2D²⁴.

In contrast to fasting glucose (Supplementary Fig. 5a), in fasting insulin analyses adjusted for BMI, we observed a systematic decrease in the standard errors of the SNP effect estimates (Supplementary Fig. 5b), perhaps because BMI explains more of the variance in fasting insulin ($R^2 = 32.6\%$) than in fasting glucose ($R^2 = 8.6\%$) or 2hGlu ($R^2 = 11.0\%$) (data from the Fenland study). Therefore, BMI adjustment removes more variance in fasting insulin, thereby rendering genetic associations more readily detectable. This is supported by the identification of five additional loci in BMI-adjusted models by this approach (Table 1 and Supplementary Figs. 3 and 4). As expected, BMI-adjustment abolished fasting insulin associations at *FTO* ($P = 0.71$; Supplementary Table 2b), suggesting that the association with fasting insulin is mediated entirely through association with BMI.

In total, 13 of the 19 loci associated with fasting insulin also showed associations with T2D ($P < 0.05$; FDR $q < 0.05$) (Fig. 1), with the insulin-raising allele associated with higher risk of T2D, except for *TCF7L2* (Fig. 2b,c), where the allele associated with lower fasting insulin is associated with higher fasting glucose (Table 1). Notably, the loci associated with fasting insulin showed a pattern of association with lipid traits consistent with insulin resistance, and not observed for either fasting glucose or 2hGlu loci (Fig. 1). Thirteen (~68%) of the 19 loci were associated with HDL-cholesterol ($q < 0.05$): all insulin-raising alleles associated with lower HDL levels, nine of which were also associated with higher triglycerides ($q < 0.05$) (Fig. 1). Further, the insulin-raising alleles of four SNPs were associated with higher WHR (adjusted for BMI) ($q < 0.05$) (Fig. 1), another trait linked to insulin resistance, while five SNPs were also associated with BMI, although with inconsistent direction ($q < 0.05$) (Fig. 1).

In gene-based analyses, we focused on BMI-adjusted results to account for the variance in fasting insulin explained by BMI. Beyond those loci containing genome-wide significant SNPs, we identified 7 distinct regions (containing 22 genes) after Bonferroni-correction ($P < 5 \times 10^{-6}$). Among these genes, we identified many for which prior biological evidence suggests their role in pathways involved in insulin secretion or action (Supplementary Table 3b). While the lead SNP in *PPARD* was not genome-wide significant ($P = 3.9 \times 10^{-6}$), the *PPARD* gene, a regulator of adipose, hepatic and skeletal muscle metabolism³⁷, reached the gene-based significance threshold ($P < 1 \times 10^{-6}$). *PPARD* agonists have also been shown to induce insulin sensitising effects in a mouse model³⁸. In addition, we identified *PTEN* to be associated (Supplementary Table 3b), a gene previously suggested to affect glucose metabolism through regulation of insulin signalling³⁹, and in which a muscle-specific deletion protected mice from insulin resistance and diabetes resulting from high fat feeding⁴⁰.

2-h glucose

In 42,854 individuals, we identified four additional loci to be associated with 2hGlu (Table 1 and Supplementary Figs. 3 and 4), including a signal near *ERAP2* and three signals near loci previously associated with fasting glucose¹ (*GCK*), HDL-cholesterol²⁷ (*PPP1R3B*) and T2D⁶ (*IGF2BP2*), as well as confirming the five previous associations². To determine whether these associations reflected differences in the response to a glucose challenge, or were partly driven by effects on fasting glucose, we also performed analyses adjusted for fasting glucose. No additional loci were identified as genome-wide significant after adjustment for fasting glucose, although the *GCK* association with 2hGlu was severely attenuated ($\beta = 0.04$ (SE = 0.016) mmol/L/allele, $P = 0.005$ vs. $\beta = 0.1$ (0.016) mmol/L/allele, $P = 5.3 \times 10^{-11}$ in the model unadjusted for fasting glucose), suggesting that the association with 2hGlu is driven, at least in part, by a primary association with fasting glucose (Supplementary Table 2d). The association of SNPs near *GCK* with both fasting glucose and 2hGlu suggests a generalized raising of the glucose set point, consistent with inactivating mutations of *GCK* that cause MODY⁴¹. As for fasting glucose, when 2hGlu models were adjusted for BMI, no systematic differences were observed, although again the *IGF2BP2* SNP rs7651090 reached genome-wide significance (Table 1).

Eight of the 9 SNPs associated with 2hGlu at genome-wide levels of significance were also associated with T2D ($q < 0.05$) (Fig. 1), although the 2hGlu-raising alleles at *PPP1R3B*, *GCKR* and *VPS13C-C2CD4A/B* were associated with lower risk of T2D (Fig. 2d), consistent with their association with lower fasting glucose levels (Table 1 and Supplementary Table 2e).

In addition to SNPs that were genome-wide significant in individual SNP analyses, we identified three regions (containing six genes) showing association with 2hGlu in gene-based analyses. These included the *HKDC1* gene, as well as an association signal at *CRHR1* ($P = 2 \times 10^{-6}$) (Supplementary Table 3c), mostly driven by the lead SNP in this gene (rs17762954), which approached genome-wide significance ($P = 7.4 \times 10^{-7}$). *CRHR1*, together with *GIPR*, belongs to the family of class B GPCRs and is highly expressed in pancreatic β -cells, where stimulation of the receptor potentiates insulin secretion in response to glucose⁴².

Fine-mapping of established loci

Analyses at higher SNP density around previously established loci did not generally yield stronger associations or more plausible functional variants (Supplementary Table 4). For fasting glucose, markedly more significant SNPs or larger effect size than the previous lead SNP were observed for four of the 16 loci: *PROX1*, *GCK*, *ADRA2A* and *VPS13C-C2CD4A/B* (Supplementary Table 4). Regional plots for these loci are shown in Supplementary Figure 6. While the new lead SNP near *ADRA2A* was not markedly more significant than the previous lead SNP, the effect size is almost double that of the previous lead SNP (Supplementary Table 4). However, this and other new lead SNPs were without more plausible functionality. The new lead SNP at *VPS13C-C2CD4A/B*, previously associated with proinsulin⁴³, is far more significant and of larger effect size than the previous lead SNP ($\beta = 0.0273$ (SE = 0.0035) mmol/L/allele, $P = 4.8 \times 10^{-15}$ vs. $\beta = 0.0057$ (0.0036) mmol/L/allele, $P = 0.111$; $r^2 = 0.27$). For fasting insulin, another SNP downstream of *IGF1* was found to be more significant and with a larger effect size, although with no known functionality (Supplementary Table 4 and Supplementary Fig. 6). For 2hGlu, again, another SNP at *VPS13C-C2CD4A/B* was more significant than the previous lead SNP (Supplementary Table 4 and Supplementary Fig. 6) and was previously associated with diabetes in Chinese individuals⁴⁴.

Pathway analysis

Next, we explored whether glycemic loci were enriched for connectivity between genes representing particular pathways or processes. To do this, we used GRAIL software⁴⁵ and investigated both an excess of connectivity between the established loci (genome-wide significant) and then between established loci and those loci that did not reach genome-wide significance but that showed a lower level of association ($P < 0.0005$) (Online Methods). We aimed to establish whether there were any biologically relevant genes among this longer list of suggestively-associated loci. This more liberal threshold yielded 218, 155 and 100 regions for fasting glucose, fasting insulin and 2hGlu, respectively. To further assess whether the established loci represented common biological pathways, we used MAGENTA to undertake gene-set enrichment analyses (Online Methods).

We found that genes near the 36 loci associated with fasting glucose had a high degree of connectivity (refer to Online Methods for definition of how genes were selected), with eight genes demonstrating highly significant similarity to genes in other loci at a $P_{\text{grail}} < 0.01$ level, connected by keywords such as “glucose”, “insulin”, “pancreatic” and “diabetes” (Supplementary Table 5a and Supplementary Fig. 7), and more than expected by chance ($P_{\text{permutation}} = 0.003$). We observed less connectivity among the genome-wide significant fasting insulin and 2hGlu loci, with no genes reaching $P_{\text{grail}} < 0.01$ for fasting insulin (Supplementary Table 5b) and only one out of nine genes for 2hGlu ($P_{\text{permutation}} = 0.07$) (Supplementary Table 5c).

Among the list of 218 suggestively-associated fasting glucose loci ($P < 0.0005$), we observed 13 genes to be connected to those in the genome-wide significant loci at $P_{\text{grail}} < 0.01$, more than expected by chance ($P_{\text{permutation}} = 0.003$) (Supplementary Table 6a). These included genes such as *GLP1R* ($P = 3.3 \times 10^{-7}$) (a glucagon receptor that mediates the GLP-1 incretin effect and stimulates insulin release), *IRS2* ($P = 6.9 \times 10^{-5}$; central to development and maintenance of β -cell mass and function^{46,47}) and *INS* ($P = 2.5 \times 10^{-6}$; the insulin gene encoding proinsulin). The presence of these and other genes support our conjecture that many of the SNPs approaching genome-wide significance are likely to represent true associations. Of the 155 suggestively-associated loci for fasting insulin (adjusted for BMI), we observed seven to be connected to the genome-wide significant loci at $P_{\text{grail}} < 0.01$; more than expected by chance ($P_{\text{permutation}} = 0.002$), and these included *INSR* ($P_{\text{grail}} = 1.5 \times 10^{-4}$; encoding insulin receptor precursor), *CD36* ($P_{\text{grail}} = 0.001$; previously implicated in insulin resistance⁴⁸), *GCG* ($P_{\text{grail}} = 0.008$; glucagon gene) and *HNF1A* ($P_{\text{grail}} = 0.005$; mutations in which are associated with MODY3)⁴⁹ (Supplementary Table 6b). Of the 100 suggestively-associated loci for 2hGlu ($P < 0.0005$), we found three to reach $P_{\text{grail}} < 0.01$ ($P_{\text{permutation}} = 0.014$) and the gene highlighted as most biologically connected to the genome-wide significant loci was again *HNF1A* ($P_{\text{grail}} = 3.4 \times 10^{-4}$) (Supplementary Table 6c).

Using MAGENTA, we identified four pathways enriched for fasting glucose associations: GOTERM pathways lens development in camera-type eye ($P = 0.004$), PANTHER processes gut mesoderm development ($P = 0.009$), other steroid metabolism ($P = 0.02$) and KEGG MODY pathway ($P = 0.03$), although these were no longer significant after removing lead genes ($P > 0.05$), all of which were known fasting glucose loci: *PROX1* for eye and gut, and *G6PC2* and *GCK* for steroid and MODY pathways, respectively.

Directional consistency of associations between discovery and follow-up studies

Given the wealth of biologically plausible genes in loci near genome-wide significance (Supplementary Tables 6a-c) and the deviation of the observed distribution from the expected in QQ plots even after removing all established loci (Supplementary Fig. 8a-d), we

hypothesized that additional loci not reaching genome-wide significance were likely to represent true associations with small effects. To establish the presence of further true associations that did not reach genome-wide significance, we compared SNP associations in discovery studies (those included in the original meta-analyses for 42,078 (fasting glucose), 34,230 (fasting insulin) and 15,252 (2hGlu) individuals^{1,2}) with those in the “follow-up” studies (consisting of 85,710 (fasting glucose), 69,240 (fasting insulin) and 27,602 (2hGlu) individuals). We identified all SNPs which had a nominally significant association ($P < 0.05$) in the follow-up studies alone and, for these SNPs, performed a binomial test of whether more SNPs than expected by chance (50%) had a consistent direction of effect with that observed in the discovery analyses. We were also able to compare among SNPs nominated for follow-up by different consortia (Supplementary Fig. 9a-d).

For each trait, evaluation of the 66,000 MetaboChip follow-up SNPs demonstrated a significant excess of SNPs showing directionally consistent associations ($P < 0.05$) compared to that expected by chance (fasting glucose: $P_{binomial} = 5.01 \times 10^{-12}$; fasting insulin: $P_{binomial} = 7.58 \times 10^{-13}$; fasting insulin (adjusted for BMI): $P_{binomial} = 9.76 \times 10^{-9}$; 2hGlu: $P_{binomial} = 2.37 \times 10^{-6}$; Supplementary Table 7 and Supplementary Fig. 9a-d). FDR analyses suggested that a number of these nominal associations in the follow-up studies are true positives for fasting glucose and fasting insulin in particular (fasting glucose: 23%; fasting insulin: 24%; Supplementary Table 7). Interestingly, when we evaluated consistency of association with fasting insulin (between discovery and follow-up) among SNPs submitted to the MetaboChip by other consortia, SNPs submitted by GIANT (anthropometric traits) ($P_{binomial} = 1.52 \times 10^{-8}$) and GLGC (lipid traits) ($P_{binomial} = 1.15 \times 10^{-6}$) and for BMI and triglycerides in particular also demonstrated a marked excess of directional consistency (Supplementary Table 7 and Supplementary Fig. 9b). When we performed the same test for fasting insulin adjusted for BMI, the observed enrichment among SNPs submitted by GIANT and GLGC was attenuated (Supplementary Table 7 and Supplementary Fig. 9c), although SNPs nominated to follow up on triglyceride associations remained the most significant ($P = 3.18 \times 10^{-7}$, Supplementary Table 7 and Supplementary Fig. 9c). Of the 3,353 SNPs submitted for follow-up of triglyceride associations, 158 SNPs showed nominal significance ($P < 0.05$) in follow-up studies and consistent direction of association with fasting insulin (adjusted for BMI) in both discovery and follow-up (Supplementary Table 7). In 139 (88%) of these SNPs, the insulin-raising alleles were associated with higher levels of triglycerides, consistent with the positive correlations between fasting insulin and triglyceride associations observed among the genome-wide significant fasting insulin loci (Fig. 1).

DISCUSSION

In the current meta-analysis of ~66,000 MetaboChip follow-up SNPs in up to 133,010 individuals, we identified a large number of loci to be associated with glycemic traits, explaining 4.8%, 1.2% and 1.7% of the variance in fasting glucose, fasting insulin and 2hGlu, respectively. Of the 53 glycemic loci, 33 are also associated with increased T2D risk ($q < 0.05$), extending the overlap between glycemic and T2D loci. Given the current DIAGRAM effective sample size of 106,953 individuals, we can exclude an effect on T2D of 1.04 with 80% power for alleles more frequent than 5%, effectively confirming that the overlap is incomplete and that many loci associated with glycemic traits have no discernible effect on T2D (Figs. 1 and 2).

Previously, we had detected only two loci associated with fasting insulin and had hypothesized that this might be due to a different genetic architecture of this trait compared to fasting glucose, with potentially smaller effect sizes, lower frequency alleles or greater environmental influence on fasting insulin¹. In the current meta-analysis including up to

108,557 individuals (compared to 62,264 individuals previously), we expanded the number of loci associated with this trait to 19. Of note was the effect of BMI-adjustment on our ability to detect additional loci (five non-overlapping with unadjusted results)¹⁹. We also noted that some of the loci influencing fasting insulin uncovered after BMI-adjustment are likely to have been negatively confounded in previous efforts: at some loci, the insulin-raising allele was nominally associated with lower BMI (potentially via insulin resistance attenuating the anabolic effects of insulin). Given the positive correlation between BMI and fasting insulin, it is likely that this association previously masked their effect on fasting insulin. Fasting insulin loci showed directionally consistent association with lipid levels (HDL and triglycerides); that is, the insulin-raising allele was associated with lower HDL and higher triglyceride levels, a hallmark combination in insulin resistant individuals. We also observed some overlap between fasting insulin loci and those associated with abdominal obesity (Fig. 1). Jointly, these data suggest links of these fasting insulin loci to insulin resistance-related phenotypes. Indeed, some of the fasting insulin loci identified, such as *IRSI* and *PPARG*, are classically known to exert effects on insulin action or sensitivity^{50,51}.

There are now 36 established fasting glucose loci, many of which contain compelling biological candidate genes with plausible causality, including those encoding transcription factors with known roles in pancreas development (e.g. *PDX1*, *FOXA2*, *PROX1*, *GLIS3*) and genes involved in β -cell function and insulin secretion pathways (*SLC2A2*, *GCK*, *PCSK1*). For 2hGlu, only nine loci have been established to date, which likely reflects the smaller sample size available and consequently reduced power.

Comparing the consistency of the direction of associations for glycemic traits between “discovery” and “follow-up” studies suggests that we are observing more directionally consistent associations than expected by chance among MetaboChip follow-up SNPs (Supplementary Fig. 9a-d). This, combined with the excess of biologically plausible genes among the borderline loci (Supplementary Table 6a-c), suggests that beyond the genome-wide significant loci there is a more extensive list of loci still likely to contain true associations. Indeed, some of these loci are implicated by gene-based analyses, which identify genes with compelling biological credentials. For fasting insulin, these analyses revealed additional loci with previously suggested links to insulin resistance (*PPARD* and *PTEEN*). These results lend further support to the proposal that a long tail of common variants of small effect size are likely to account for a substantial proportion of the variance of complex traits^{7,8}.

Of note is the number of glycemic loci associated with other metabolic traits ($q < 0.05$; 34 of 53) and also at genome-wide levels of significance ($P < 5 \times 10^{-8}$) (14 of 53) (Fig. 1), potentially implicating pleiotropic effects. Further support for this notion comes from the analysis of loci nominated for the MetaboChip by other consortia and their associations with glycemic traits (Supplementary Fig. 9a-d). Indeed, some of the loci associated with glycemic traits at genome-wide significance levels were not originally nominated to the MetaboChip for follow-up by MAGIC (Table 1). MetaboChip data available across all contributing consortia will facilitate systematic exploration of these correlated phenotypes with more sophisticated statistical methods for joint analysis⁵²⁻⁵⁴, yielding greater insight into the underlying pathways and genetic networks they represent. As data from human genetic networks accrue, we will be better placed to test whether there is support for the notion of “hub” genes; that is, genes highly connected with others in the network and proposed by experiments in *C. elegans* to act as buffers for genetic variation and that could act as modifier genes for many different disorders⁵⁵.

In summary, we present a large number of genome-wide significant loci influencing glycemic traits, many with a compelling biological basis for their association, as well as a number of loci not previously implicated in glycemic regulation, and for which fine-mapping and functional follow-up will expand and improve our understanding. Use of the MetaboChip for deep follow-up has identified additional loci to be involved in glycemic regulation that, due to insufficient sample size and power, did not reach genome-wide significance. Consideration of such loci in future studies will better exploit data from GWAS and complimentary approaches and further improve our biological understanding of glycemic control and the etiology of diabetes.

ONLINE METHODS

Study design

The Illumina CardioMetaboChip (MetaboChip) is a custom Illumina iSELECT array of 196,725 SNPs. It has been designed to support efficient large-scale follow-up of putative associations for glycemic (including fasting glucose, fasting insulin and post-challenge glucose concentrations (2hGlu) and other metabolic and cardiovascular traits (Supplementary Fig. 1)⁹, and to enable the fine-mapping of established loci. Overall, there were 65,435 SNPs genotyped on the MetaboChip for follow-up of previous associations including a total of 23 cardio-metabolic traits. Traits contributing SNPs to the MetaboChip were prioritized into “primary” (including fasting glucose) and “secondary” (including fasting insulin and 2hGlu) contributing ~5K and ~1K SNPs, respectively, from the most significantly associated variants for each phenotype in the discovery meta-analyses from each contributing consortium. This included 5,055 SNPs for follow-up of fasting glucose, 1,046 for fasting insulin and 1,038 for follow up of 2hGlu associations. In the present analysis, we focused our analysis on this set of “follow-up” SNPs available on the MetaboChip to establish variants among these SNP associated with glycemic traits. While we also included newly available studies genotyped on genome-wide platforms, we limited our primary analyses to only these ~66,000 SNPs.

Studies

In the present effort, collaborating studies within the Meta-Analysis of Glucose and Insulin related traits Consortium (MAGIC) provided results for the 66,000 “follow-up” SNPs genotyped on MetaboChip on a maximum total of 133,010 (fasting glucose), 108,557 (fasting insulin) and 42,854 (2hGlu) individuals. As well as those newly genotyped on the MetaboChip platform, in our overall meta-analysis we were able to include further studies which had genotyped or imputed the same SNPs on other platforms. The largest proportion of our entire sample was directly genotyped on the MetaboChip and comprised 53,622 (fasting glucose), 42,384 (fasting insulin) and 27,602 (2hGlu) individuals from 26/21/12 studies, respectively. We were also able to recruit 11,690 (fasting glucose) and 8,813 (fasting insulin) individuals from up to four additional GWA studies (Prevend, Ascot (FG-only), Prosper and TRAILS) (Supplementary Table 1) not included in the original meta-analysis¹. From another MAGIC study of sex-specific associations with glycemic traits (I. Prokopenko on behalf of the MAGIC authors, personal communication), we were able to recruit another 15/13 independent studies comprising up to 25,618 and 23,130 individuals for fasting glucose and fasting insulin, respectively. The above studies were combined in a single fixed-effects meta-analysis with those studies included in the original GWAS^{1,2}: 20 (fasting glucose), 19 (fasting insulin) and 9 (2hGlu) studies and 42,080 (fasting glucose), 34,230 (fasting insulin) and 15,252 (2hGlu) individuals, as described previously^{1,2}. The study and individual counts from the original GWAS excluded the family-based SardiNIA study where, initially, a large number of the individuals had imputed genotype data only. The entire sample was directly genotyped on MetaboChip, so those data were included in

place of the original GWAS. Some studies had genotyping data available from both MetaboChip and genome-wide arrays but from entirely independent samples within the studies (Supplementary Table 1). Full study characteristics of all MetaboChip studies are shown in Supplementary Table 1, while data from discovery genome-wide studies and those from the sex-specific analyses are reported elsewhere^{1,2} (I. Prokopenko on behalf of the MAGIC authors, personal communication). All participants of the main analysis were of European descent and mostly adults, although data from a total of 7,872 and 7,164 adolescents were also included in the fasting glucose and fasting insulin meta-analyses, respectively (NFBC86, Leipzig-childhood_IFB, TRAILS and ALSPAC studies). All studies were approved by local research ethic committees and all participants gave informed consent. Results from the CLHNS study of Filipino women ($n = 1,682$ and $1,635$ for fasting glucose and fasting insulin, respectively) genotyped on MetaboChip were also available and were included in supplementary analyses to compare effect directions with European-descent studies alone.

Phenotypes

Analyses were undertaken for fasting glucose and fasting insulin measured in mmol/l and pmol/l, respectively. 2hGlu was measured in mmol/l. Similar to the previous MAGIC discovery analysis^{1,2}, individuals were excluded from the analysis if they had a physician diagnosis of diabetes, were on diabetes treatment (oral or insulin), or had a fasting plasma glucose equal to or greater than 7 mmol/l. Individual studies applied further sample exclusions, including pregnancy, non-fasting individuals and type 1 diabetes, as detailed in Supplementary Table 1. Individuals from case-control studies (Supplementary Table 1) were excluded if they had hospitalization or blood transfusion in the 2-3 months before phenotyping took place. 2hGlu measures were done 120 min after a glucose challenge during an oral glucose tolerance test (OGTT). Measures of fasting glucose and 2Glu made in whole blood were corrected to plasma level using the correction factor of 1.13 (ref. 87). Fasting insulin was measured in serum. Detailed descriptions of study-specific glycemic measurements are given in the Supplementary Table 1.

Trait transformations and adjustment

Analyses were performed for untransformed levels of fasting glucose, natural logarithm transformed fasting insulin and untransformed 2hGlu using a linear regression model. All analyses were adjusted for age (if applicable), study site (if applicable) and geographical covariates (if applicable) to evaluate the association using an additive genetic model at each genetic SNP variant.

BMI-adjusted analysis

In the Fenland study (Supplementary Table 1), we investigated the correlation between BMI and natural logarithm transformed fasting insulin, fasting glucose and 2hGlu to establish the variation in each trait explained by BMI. Meta-analyses for each trait were also adjusted for body mass index (BMI). MetaboChip and new GWA studies performed study-level analyses adjusted for BMI. Most studies in the original GWAS (except deCode, GEMs, KORAF4, TwinsUK studies) as well as from the studies analyzed in a sex-specific manner were included in BMI-adjusted meta-analysis. The original discovery 2hGlu meta-analysis adjusted for BMI² was also included in these analyses. We also performed an analysis for 2hGlu adjusted for fasting glucose to investigate if additional variants would be identified with an effect on 2hGlu independent of fasting glucose and also to establish whether identified 2hGlu associations were driven by fasting glucose.

Genotyping and quality control

The MetaboChip or other commercial genome-wide arrays were used by individual studies for genotyping. Details are presented in Supplementary Table 1 or are reported elsewhere^{1,2}. The quality control criteria for both MetaboChip and genome-wide arrays for filtering of poorly genotyped individuals or low quality SNPs prior to imputation included: (i) call rate < 0.95; (ii) sex-discrepancies; (iii) ethnic outliers; (iv) heterozygosity (Supplementary Table 1); (v) SNP minor allele frequency < 0.01; (vi) SNP Hardy-Weinberg equilibrium $P < 10^{-4}$; (vii) SNP effect estimate standard error (SE) > 10 ; (viii) SNPs minor allele count (MAC) < 10 (calculated as total number of observed alleles at each SNP multiplied by MAF).

Studies with genome-wide arrays undertook imputation using the HapMap CEU reference panel using MACH and IMPUTE software (Supplementary Table 1). Parameters used in imputation and filters applied to imputed genotypes are described in Supplementary Table 1 or reported previously^{1,2}. From a total of ~2.5M genome-wide directly genotyped or imputed autosomal SNPs, study-specific results for the ~66,000 MetaboChip follow-up SNPs were considered for the present meta-analyses. SNPs with a meta-analysis result for more than a total 10,000 individuals were included in the analysis.

Statistical analysis

Analyses of previous discovery studies are reported elsewhere^{1,2}, while those studies genotyped on the MetaboChip are described in Supplementary Table 1. SNP effect estimates and their standard errors (for additive genetic model) were combined by inverse-variance weighted fixed effects meta-analysis using METAL⁸⁸ and GWAMA⁸⁹. Two parallel meta-analyses for each trait by different analysts were compared for consistency. Individual cohort results were corrected for residual inflation of the test statistics using lambda of genomic control (GC) estimates. The GC values were estimated for each study using either test statistics from all SNPs for the GWA studies, while for those studies genotyped on the MetaboChip, GC lambda estimates were derived from test statistics for 5,041 SNPs selected for follow-up of QT-interval associations, as we perceived these to have the lowest likelihood of common architecture of associations with glycaemic traits. Individual study-level lambda GC estimates are shown in Supplementary Table 1. Overall QQ plots for the QT follow-up SNPs are shown in Supplementary Figure 10.

Trait-associated signal selection strategy

Meta-analysis results for each trait were considered as genome-wide significant if they achieved $P < 5 \times 10^{-8}$ threshold and were not in LD ($r^2 < 0.05$) or within 500 kb of an established signal. The most significantly associated SNP (lowest P -value) in each region (500 kb) was selected as the lead SNP. Associated loci are referred to by the name of the nearest gene, unless a more biologically plausible gene was nearby, or a nearby gene was previously associated with another trait. In such cases, we maintain consistency with the previous naming, but list the nearest genes in Supplementary Tables 2a-d. To establish the variance in each trait explained by these SNPs, in the Framingham Heart Study, we included all SNPs in a model adjusted for age, sex, BMI and cohort.

Fine-mapping of known glycaemic trait loci

To undertake preliminary fine-mapping analyses, we investigated the patterns of association at 17 known fasting glucose and fasting insulin loci¹ and 5 known 2hGlu loci² using meta-analysis results from 13,644, 1,309 and 1,249 SNPs genotyped on the MetaboChip in 53,622, 42,384 and 27,602 individuals for fasting glucose, fasting insulin and 2hGlu, respectively. Only studies genotyped directly on the MetaboChip were used for fine-mapping purposes in order to have equal sample size and availability of all SNPs. Regional plots for each locus

were created using the previous lead SNP¹ or a suitable proxy ($r^2 > 0.8$) as the index SNP if that marker was not present on MetaboChip. The plots were generated on the LocusZoom web-based plotting software⁹⁰ using LD information from the 1000 Genomes Project (hg19/Nov2010EUR data). Prior to generating the plots, all SNP names and positions from the MetaboChip-only meta-analysis files were aligned to build37 using the Lift Genome annotation tool on the UCSC website (<http://genome.ucsc.edu/cgi-bin/hgLiftOver>) in order to be compatible with the 1000 Genomes SNP naming format (chr:position) and allow more thorough assessment of the pairwise LD patterns around the established SNPs.

Associations of glycemic trait variants with related traits

For those SNPs which we identified to be genome-wide significant, we also investigated their association with other metabolic and disease traits. We exchanged reciprocal data for such SNPs with the latest DIAGRAM MetaboChip analyses²⁴, and checked associations of these SNPs in publicly available data from previous studies of lipid traits from the GLGC²⁷ (triglycerides, HDL- and LDL-cholesterol; <http://www.sph.umich.edu/csg/abecasis/public/lipids2010/>) as well as BMI and waist-hip ratio (WHR) from the GIANT consortium^{25, 26} (http://www.broadinstitute.org/collaboration/giant/index.php/Main_Page). From these data, we were able to establish the presence of any association and the direction of effect for these other traits aligned to our trait-raising alleles. We highlighted associations with other traits at $P < 0.05$, and also performed FDR analyses. We performed FDR analyses for each trait separately (removing duplicate loci that were associated with more than one glycemic trait) and identified those where $q < 0.05$.

eQTL analyses

Liver gene expression data from the Advanced Study of Aortic Pathology (ASAP) study has been described previously⁹¹. In brief, liver biopsies were collected from patients at the Karolinska University Hospital, Stockholm, Sweden undergoing aortic valve surgery alone or combined with surgery for aortic aneurysm, starting from February 13, 2007. All subjects gave their informed consent and the study was approved by the ethics committee of Karolinska Institute, Stockholm, Sweden. After hybridization of extracted RNA to Affymetrix ST 1.0 Exon arrays, data was RMA normalized and log-transformed. DNA was extracted from whole blood and genotyping was carried out using the Illumina 610w-Quad bead array platform. Imputation was carried out on SNPs with a call rate exceeding 95%, using the MACH algorithm. Imputation quality scores of $RSQ < 0.3$ were excluded from analysis. An additive genetic model was used to test for association between SNPs and gene expression.

VEGAS

To identify genes with multiple associated SNPs, we performed gene-based analysis using VEGAS, described in detail previously¹⁰. Briefly, on all available samples and among the ~66,000 follow-up SNPs, VEGAS pooled the information for all SNPs within each gene (± 50 kb) to identify genes with higher evidence of association than expected by chance, while adjusting for gene size and the linkage disequilibrium structure of the SNPs, by simulation (maximum number of simulations used was 10^6). We identified genomic regions (separated by >1 Mb) showing evidence of association and described the genes contained within those regions. While we often identified multiple genes within an associated region, it is probable that some of these are significant via LD. Bonferroni correction was used to adjust for multiple testing, based on the number of independent tests (number of genes tested) (~9,300) and P -values $< 5.0 \times 10^{-6}$ were considered significant. While the number of genes represented was constrained by those SNPs submitted to the MetaboChip, our analyses asked the question: of the genes represented on the MetaboChip, all with a slightly raised prior

likelihood of association, which genes show the most evidence for association with glycemic traits?

GRAIL

We used the GRAIL⁴⁵ to evaluate whether genome-wide loci associated with glycemic traits were enriched for connectivity between genes representing particular pathways or molecular processes. As described in detail previously⁴⁵, to define the genes near each SNP, GRAIL finds the furthest neighboring SNPs in the 3' and 5' direction in LD (Hapmap CEU: $r^2 > 0.5$) and proceeds outwards in each direction to the nearest recombination hotspot⁹². All genes that overlap that interval are considered implicated by the SNP. If there are no genes in that region, the interval is extended by 250 kb in either direction. The method performs a text-based analysis looking at abstracts in PubMed prior to December 2006 (to avoid confounding from GWAS results arising after that date). We performed two analyses for each trait: first, we took all genome-wide signals for each trait as seed and query loci to investigate biological connectivity among those loci (fasting glucose = 35, fasting insulin = 16, 2hGlu = 9). For fasting insulin, we did not include *FTO* as the association with fasting insulin was entirely mediated by BMI. Secondly, we also investigated connectivity between these established signals (as seed regions) and those which did not reach genome-wide significance but were suggestively associated with each trait ($P < 0.0005$) (as query regions) as described previously⁹³. For fasting insulin, we used BMI-adjusted results to define the query regions. Query regions were defined by taking all SNPs more significant than $P < 0.0005$, removing those associated at genome-wide levels of significance and pruning SNPs of $r^2 > 0.05$ in each region using PLINK⁹⁴. As GRAIL tests connectivity of regions, we also removed any duplicates where a region was represented by more than one SNP. For those SNPs not found by the software, we submitted the region as a 500 kb window centered at the location of the SNP. This approach identified 218, 155 and 100 query regions (representing 715, 639 and 298 genes) for fasting glucose, fasting insulin (adjusted for BMI), and 2hGlu, respectively. The number of loci reaching $P_{\text{grail}} < 0.01$ was identified from these analyses, and to establish the level of enrichment, we randomly sampled 1,000 random sets of matched numbers of SNPs and calculated the proportion with as many or more reaching $P_{\text{grail}} < 0.01$ to derive a permutation based P -value ($P_{\text{permutation}}$).

Pathway analyses

Pathway analysis was carried out for fasting glucose, fasting insulin and 2hGlu (uniform/FG-BMI adjusted) using data from previous discovery GWAS studies only¹ to avoid bias towards pathways represented on the MetaboChip (build36, $n > 10,000$ and MAF 1% cutoff used). The software used for this analysis was MAGENTA 2.4 (July 2011, <http://www.broadinstitute.org/mpg/magenta/>). SNPs from the meta-analysis file were assigned to a gene if they mapped within 110 kb upstream and 40 kb downstream of transcript boundaries. The smallest P -value for the set of SNPs assigned to the gene was adjusted for confounders, such as gene length, marker density and LD in a linear regression, creating a gene association score. If a top SNP was assigned to multiple genes, only the gene with the lowest score was kept to avoid positional clustering. The HLA region was removed due to high LD and gene density. Pathway terms from multiple databases (GO, PANTHER, Ingenuity, KEGG) was attached to each gene. The genes were ranked on their association score, and a GSEA test was performed testing all pathway terms using a 5% and 75% cutoff. Initially, 10,000 gene set permutations were performed for GSEA P -value estimation. This number was then increased with GSEA $P < 1 \times 10^{-4}$, and up to 1,000,000 permutations were performed. Results were sorted on FDR (5% cutoff), and $FDR < 0.05$ was considered to be significant.

Analyses of directional consistency of associations between discovery and follow-up studies

We investigated whether the Metabochip follow-up SNPs were likely to contain further true associations in addition to those SNPs which reached genome-wide significance. To do so, we meta-analyzed those studies involved in the original discovery analyses^{1,2} comprising 42,078 individuals for fasting glucose, 34,230 for fasting insulin and 15,252 for 2hGlu, and also then separately meta-analyzed all studies newly available to follow up, comprising 85,710 individuals for fasting glucose, 69,240 for fasting insulin and 27,602 for 2hGlu. For each trait (fasting glucose, fasting insulin, FI-BMIadj and 2hGlu), we then identified all SNPs which had a nominally significant association ($P < 0.05$) in the follow-up studies alone and, for these SNPs, performed a two-sided binomial test of whether more SNPs than expected by chance (50%) had a consistent direction of effect with that observed in the discovery analyses. Before performing these analyses, SNPs were filtered by LD ($r^2 < 0.01$) to identify independent variants, and all SNPs (and those in LD, $r^2 \geq 0.01$) associated with glycemic traits (fasting glucose, fasting insulin, 2hGlu, HbA1c and proinsulin) at genome-wide levels of significance (including those SNPs identified in the present study) were excluded. These analyses were initially performed for all 66,000 SNPs, but we were then also able to compare across SNPs submitted to the Metabochip by different consortia and for SNPs submitted to follow up on particular traits amongst these consortia. The results of each of these tests were plotted overall, within SNPs from each consortia, and within SNPs submitted for follow-up of each trait in Supplementary Figure 9. The numbers of SNPs meeting these criteria are shown are Supplementary Table 7. We supplemented these results with FDR analyses and noted the q -value at $P = 0.05$ in the follow-up studies to identify the likelihood of true positives among these nominally significant SNPs (Supplementary Table 7).

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Footnotes

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Box 1**Fasting glucose**

IKBKAP (inhibitor of kappa light polypeptide gene enhancer in β -cells, kinase complex-associated protein) encodes a scaffold protein that binds IKKs and NF- κ B-inducing kinase (NIK), assembling them into different active complexes. Splicing mutations in this gene lead to familial dysautonomia⁵⁶. Also mapping to this region are **C9orf4**, **C9orf5** and **C9orf6**, **MIR32** (microRNA 32, unknown function), as well as **ACTL7A** (actin-like 7A) and **ACTL7B** (actin-like 7B).

WARS (tryptophanyl-tRNA synthetase) catalyzes the aminoacylation of tRNA(Trp) with tryptophan. The intronic SNP rs3783347 is associated with **WARS** expression in liver: the glucose-raising allele associated with lower mRNA expression (age- and sex-adjusted $P = 4.19 \times 10^{-5}$) and is in perfect LD ($r^2 = 1$, $D' = 1$) with a 3' UTR SNP in **SLC25A47** (rs3736952) and in modest LD ($r^2 = 0.3$, $D' = 1$) with non-synonymous Arg135Leu (qualified as tolerated by SIFT and probably damaging by Polyphen). Nearby **YY1** (YY1 transcription factor) codes for a zinc-finger transcription factor involved in regulating a broad set of promoters. It has been suggested that YY1-regulated transcription is linked to glucose metabolism via O-GlcNAcylation⁵⁷.

KL (klotho) encodes a type-I membrane protein related to β -glucosidases. rs576674 lies ~36 kb upstream of **KL**. Variation in **KL** has been associated with insulin regulation, insulin resistance phenotypes and cardiovascular disease in some studies^{14,15,58,59} but **KL** variants were not associated with diabetes risk⁶⁰. The various SNPs in these studies are all in weak LD with rs576674 ($r^2 < 0.125$). Variation in **KL** is also associated with bone metabolism and may play a role in associations of energy metabolism with bone metabolism^{61,62}.

TOPI (topoisomerase (DNA) I). rs6072275 is intronic in **TOPI** and lies in a large region of high LD in Europeans, which includes the plausible biological candidate **LPIN3** (lipin 3). In mice, a related homolog **Lpin1** is associated with fatty liver dystrophy⁶³, a phenotype similar to human lipodystrophy (loss of body fat, fatty liver, hypertriglyceridemia, and insulin resistance). **Lpin1** mRNA is expressed at high levels in adipose tissue and induced during differentiation of preadipocytes, suggesting that lipin is required for normal adipose tissue development, while **LPIN2** has been suggested to be associated with T2D and glucose metabolism⁶⁴. rs6072275 lies in the middle of a large CNV that extends from within the 3' end of **TOPI** to the 5' end of **PLCG1** (phospholipase C, gamma 1).

P2RX2 (purinergic receptor P2X, ligand-gated ion channel, 2). rs10747083 lies in a small CNV about 150 kb upstream of five protein-coding genes, including **P2RX2**, encoding one of a family of purinoceptors for ATP; **GALNT9** (UDP-N-acetyl-alpha-D-galactosamine:polypeptide N-acetylgalactosaminyltransferase 9 (GalNAc-T9), encoding a member of the UDP-N-acetyl-alpha-D-galactosamine polypeptide N-acetylgalactosaminyltransferase (GalNAc-T) family of enzymes and expressed specifically in the brain; **FBRSL1** (fibrosin-like 1); **PXMP2** (peroxisomal membrane protein 2, 22 kDa); **PGAM5** (phosphoglycerate mutase family member 5), and within 184 kb downstream, **POLE** (polymerase (DNA directed), epsilon) and **LOC100130238** (hypothetical LOC100130238) a miscRNA.

LOC728489. rs3829109 is in low LD with a well-established locus for inflammatory bowel disease. Two recent publications reported **CARD9** SNP rs10781499 ($r^2 = 0.29$) to be associated with ulcerative colitis⁶⁵, and **CARD9-SNAPC4** SNP rs4077515 ($r^2 = 0.27$)

to be associated with Crohn's disease and ulcerative colitis^{66,67}. Several genes are located in the region, but few with high plausibility for a role in glycemia.

AMT encodes the mitochondrial aminomethyltransferase which is a critical component of the glycine cleavage system. Depending upon the *AMT* transcript, rs11715915 is located in 3' UTR or within coding regions, where it causes a synonymous substitution. This SNP is also located downstream of *TCTA* (T-cell leukemia translocation altered), which has no known metabolic function, and upstream of *RHOA* (ras homolog family member A). *RHOA* is a signaling molecule involved actin cytoskeleton stability and reorganization⁶⁸ that binds and activates Rho kinase (ROCK), a regulator of insulin transcription⁶⁹ and action⁷⁰ that is differentially regulated in T2D⁷¹ and hypothesized to play a role in glucose homeostasis⁷⁰.

GLS2 encodes liver-expressed glutaminase 2, which is required for hydrolysis of glutamine. rs2657879 causes a benign (Polyphen) amino acid change (Leu581Pro) in the *GLS2* protein. The *GLS2* protein is highly expressed (human protein atlas) by both liver and pancreas, and it has been shown in liver tumors that alterations in the balance of *GLS2:GLS1* (the kidney-specific homolog) activity are important for regulating glutamate metabolism⁷². The other gene in this region ***SPRYD4*** (*SPRY* domain containing 4) has no known function in metabolism.

RREB1 (ras responsive element binding protein 1) encodes a zinc finger transcription factor, with rs17762454 lying in an intron in the gene. The protein product of *RREB1* binds to RAS-responsive elements (RREs) of gene promoters, including the calcitonin gene promoter. The role of *RREB1* in energy metabolism is not known. An uncorrelated SNP at this locus (rs675209) was associated with serum urate levels ($P = 1.0 \times 10^{-9}$) in a GWAS of serum urate, gout and cardiovascular disease risk factors⁷³. Another gene at this locus, ***SSRI*** (signal sequence receptor, alpha), encodes a glycosylated endoplasmic reticulum membrane receptor associated with protein translocation across the ER membrane. Reactome pathway analysis places this gene in a module with key roles in the synthesis and function of insulin, insulin-like growth factors and ghrelin, making this gene a plausible biological candidate at this locus (REACTOME: REACT_15380). A third gene at this locus, ***CAGE1***, encodes cancer antigen 1. *CAGE1* has no known role in metabolism.

Fasting Insulin

TET2 encodes the tet oncogene family member 2, isoform b, which catalyzes the conversion of methylcytosine to 5-hydroxymethylcytosine. The enzyme is involved in myelopoiesis, and defects in this gene have been associated with several myeloproliferative disorders (NCBI RefSeq). Perhaps more relevant to glycemic regulation is ***PPA2***, which encodes the inorganic pyrophosphatase 2 isoform 1 precursor. Its protein product is localized to mitochondria; it has high homology to members of the inorganic pyrophosphatase family, including the signature sequence essential for its catalytic activity (NCBI RefSeq). Pyrophosphatases catalyze the hydrolysis of pyrophosphate to inorganic phosphate.

HIP1 encodes the huntingtin interacting protein 1, a membrane-associated protein that colocalizes with huntingtin. It is ubiquitously expressed with the highest level in brain. Loss of normal huntingtin-HIP1 interaction in Huntington's disease may contribute to a defect in membrane-cytoskeletal integrity in the brain. Of interest to insulin action, *HIP1* is involved in clathrin-mediated endocytosis and trafficking. Mice transgenic for the mutated form of huntingtin develop diabetes^{74,75}; however, though *Hip1/Hip1r* double-knockout mice have severe vertebral defects, suffer from dwarfism and die in early adulthood, they do not show any fasting glucose abnormalities⁷⁶. The lead SNP

(rs1167800) is only 104 bp away from a missense SNP (rs1167801), encoding a Gln to His amino acid change; however, LD between them is low ($r^2 = 0.196$).

FAM13A (family with sequence similarity 13, member A) encodes a protein with unknown function. Previous GWAS for the study of lung function measures⁷⁷ and chronic obstructive pulmonary disease⁷⁸ described variants in *FAM13A* that affect these traits. **SPP1**, encoding osteopontin, a secreted matrix glycoprotein and pro-inflammatory cytokine involved in cell-mediated immunity, is within 1 Mb. Mice exposed to high fat diet show increased circulating osteopontin and over-expression of Spp1 in the macrophages recruited into adipose tissue improved insulin sensitivity⁷⁹, while SPP1 was highly expressed in obese twins relative to their non-obese siblings⁸⁰. Recent work linked osteopontin to β -cell function through the GIP pathway⁸¹. In carriers of the *GIPR* variant associated with impaired glucose and GIP-stimulated insulin secretion, osteopontin levels were lower compared to non carriers. In addition, both GIP and osteopontin prevented cytokine-induced apoptosis and osteopontin-stimulated cell proliferation of functional β -cell mass.

PEPD (peptidase D) encodes a member of the peptidase family. The protein forms a homodimer that hydrolyzes dipeptides or tripeptides with C-terminal proline or hydroxyproline residues. The enzyme serves an important role in the recycling of proline, and may be rate limiting for collagen production. **CEBPA** (CCAAT/enhancer binding protein (C/EBP) alpha) is ~100 kb downstream of the lead SNP and encodes a transcription factor expressed in adipose tissue that regulates a number of genes involved in lipid and glucose metabolism genes. A SNP in low LD with our lead SNP was previously associated with triglyceride levels⁸². Cells from *Cebpa* (-/-) mice show a complete absence of insulin-stimulated glucose transport, secondary to reduced gene expression and tyrosine phosphorylation of the insulin receptor and IRS1 (ref. 83). CEBPA also modulates expression of leptin by binding to the promoter of the gene⁸⁴, and our lead SNP showed modest association with BMI in previous GIANT meta-analyses ($P = 0.005$).

YSK4 (Sps1/Ste20-related kinase homolog) contains rs1530559 in an intron. This gene has no known function in human energy metabolism. Three other genes at this locus also have no known role in energy metabolism, including **RAB3GAPI** (RAB3 GTPase activating protein subunit 1 (catalytic), encoding the catalytic subunit of a Rab GTPase activating protein and mutated in Warburg micro syndrome; **CCNT2** (cyclin T2), belonging to the highly conserved cyclin family, whose members are characterized by a dramatic periodicity in protein abundance through the cell cycle; and **ACMSD** (aminocarboxymuconate semialdehyde decarboxylase), involved in the *de novo* synthesis pathway of NAD from tryptophan. *ACMSD* has been implicated in the pathogenesis of several neurodegenerative disorders.

2h Glucose

ERAP2 (endoplasmic reticulum aminopeptidase 2) encodes an aminopeptidase that hydrolyzes N-terminal amino acids of proteins or peptide substrates. The lead SNP is strongly associated with *ERAP2* expression in liver ($P = 1.1 \times 10^{-55}$) and in lymphoblastoid cell lines in individuals from the CEU ($P = 8 \times 10^{-21}$) and YRI samples ($P = 2 \times 10^{-15}$). Also near to this lead SNP is **LNPEP** (leucyl/cystinyl aminopeptidase), which is widely expressed and well characterised in muscle and fat cells. In response to insulin, LNPEP translocates to the cell surface and co-localizes with GLUT4 (ref. 85). Although the role it plays in insulin action is unknown, this translocation is impaired in individuals with T2D⁸⁵. **PCSK1** is also within 500 kb of the lead SNP, although on the other side of a recombination hotspot (Supplementary Fig. 4d).

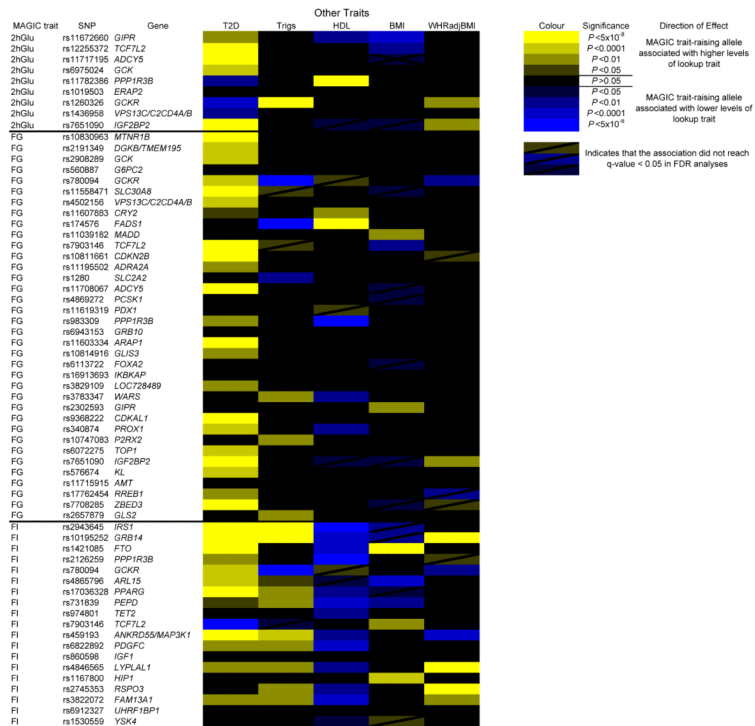
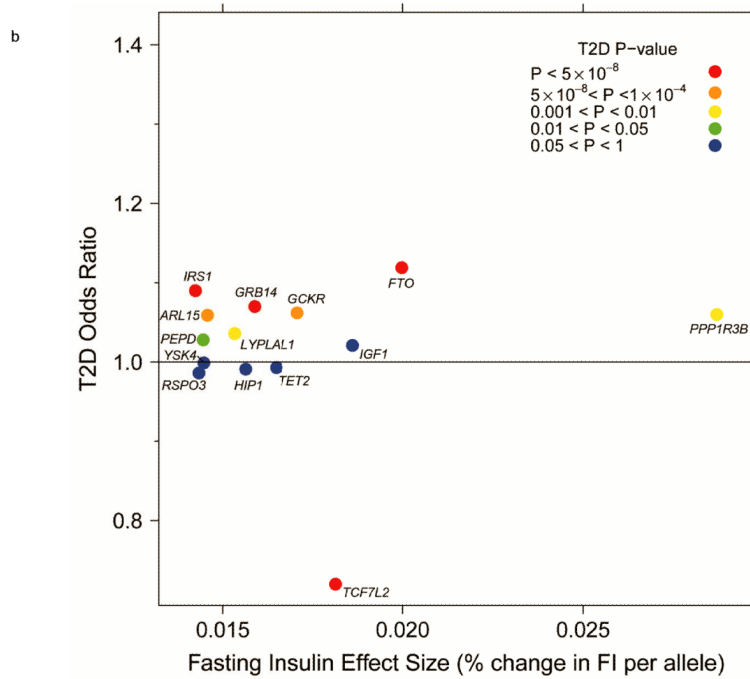
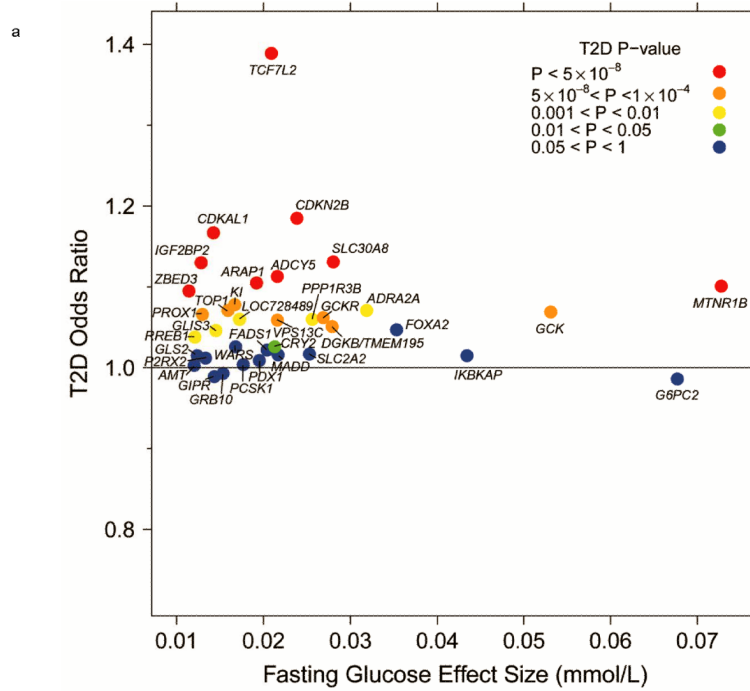


Figure 1. Associations between glycemic loci and T2D, HDL-cholesterol and triglycerides, BMI and WHR. Loci associated with the above traits ($P < 0.05$) are highlighted. Those with positively correlated effect directions are colored yellow, and those with negative correlations are colored blue. Those which did not reach a q -value < 0.05 in FDR analyses are also marked.



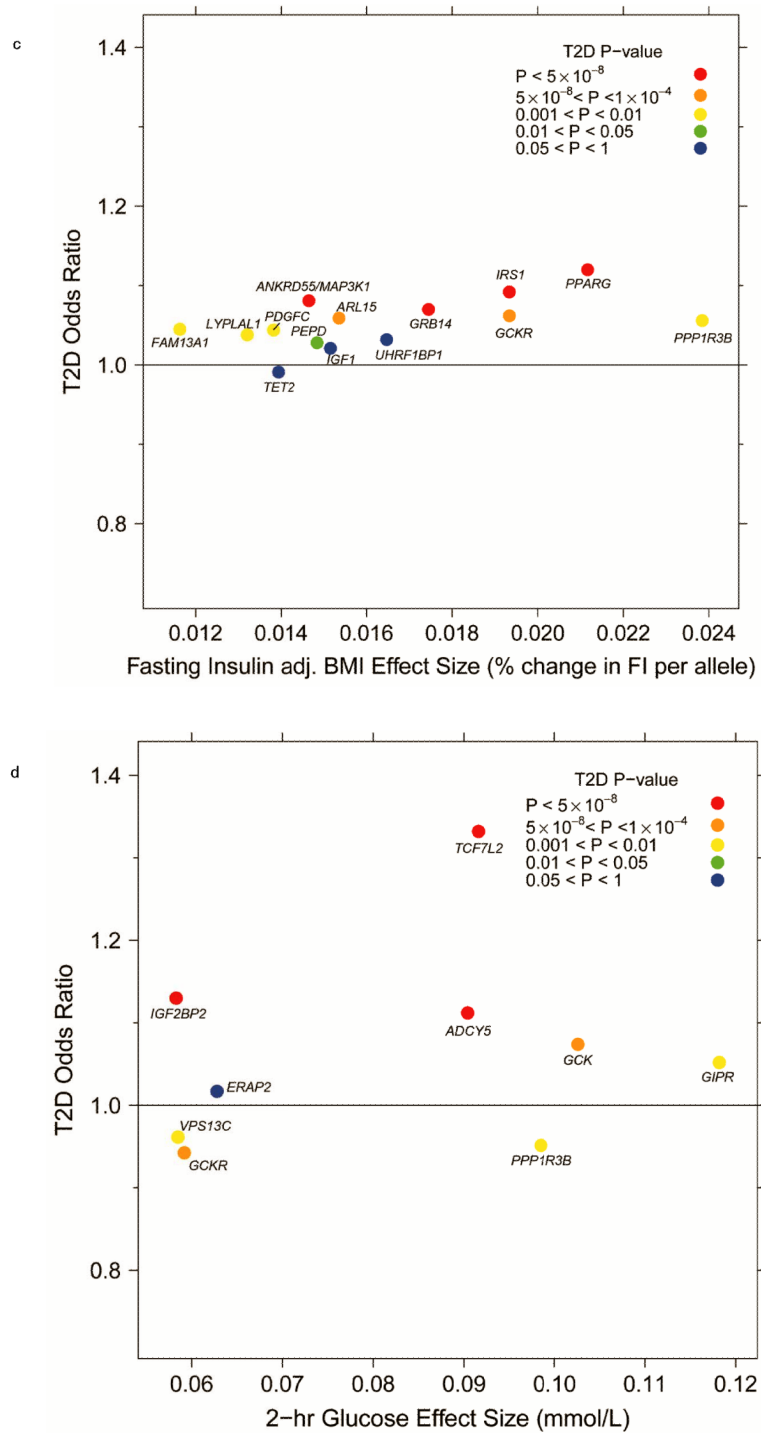


Figure 2. Per-allele beta-coefficients for glucose and insulin concentrations vs. odds ratios for T2D. (a) Fasting glucose vs. T2D. (b) Fasting insulin vs. T2D. (c) Fasting insulin adjusted for BMI vs. T2D. (d) 2-h glucose vs. T2D.

Table 1
SNPs associated with fasting glucose, fasting insulin and 2 hour-glucose at genome-wide significance level in Europeans

Primary Trait	SNP	Chr	Position	Gene	Alleles (effect/other)	Freq effect allele	Primary trait		FI (BMI-adjusted)				2hGlu			
							Effect (SE)	Global analysis P value	Global analysis n	I ² estimate (P value)	Effect (SE)	Global analysis P value	Global analysis n	Effect (SE)	Global analysis P value	Global analysis n
	rs10811661	9	22124094	<i>CDKN2B</i>	T/C	0.82	0.0238 (0.003)	5.6 × 10 ⁻¹⁸	128,488	0.00 (1.00)	-0.0065 (0.003)	0.019	98,880	0.0567 (0.014)	8.8 × 10 ⁻⁵	42,801
	rs4869272	5	95565204	<i>PCSK1*</i>	T/C	0.69	0.0177 (0.002)	1.0 × 10 ⁻¹⁵	131,872	0.00 (1.00)	0.0016 (0.002)	0.469	103,493	-0.0322 (0.012)	0.006	42,848
	rs11619319	13	27385599	<i>PDX1</i>	G/A	0.23	0.0195 (0.002)	1.3 × 10 ⁻¹⁵	132,996	0.00 (1.00)	0.0001 (0.002)	0.977	103,492	0.0185 (0.013)	0.156	42,848
	rs983309	8	9215142	<i>PPP1R3B*</i>	T/G	0.12	0.0256 (0.003)	6.3 × 10 ⁻¹⁵	127,470	0.14 (0.32)	0.0223 (0.003)	1.2 × 10 ⁻¹²	99,024	-0.0548 (0.016)	0.001	42,846
FG	rs6943153	7	50759073	<i>GRB10</i>	T/C	0.34	0.0154 (0.002)	1.6 × 10 ⁻¹²	131,795	0.00 (1.00)	0.0091 (0.002)	2.3 × 10 ⁻⁵	103,447	0.0110 (0.011)	0.333	42,794
	rs11603334	11	72110633	<i>ARAP1</i>	G/A	0.83	0.0192 (0.003)	1.1 × 10 ⁻¹¹	128,139	0.00 (1.00)	-0.0046 (0.003)	0.086	99,026	0.0294 (0.014)	0.037	42,839
	rs6113722	20	22505099	<i>FOXA2</i>	G/A	0.96	0.0353 (0.005)	2.5 × 10 ⁻¹¹	123,665	0.04 (0.78)	-0.0095 (0.005)	0.064	103,471	0.0493 (0.030)	0.101	41,416
	rs16913693	9	110720180	<i>IKBKAP</i>	T/G	0.97	0.0434 (0.007)	3.5 × 10 ⁻¹¹	125,115	0.00 (1.00)	-0.0018 (0.007)	0.785	96,357	0.0639 (0.034)	0.062	40,522
	rs3829109	9	138376587	<i>LOC728489</i>	G/A	0.71	0.0172 (0.003)	1.1 × 10 ⁻¹⁰	115,310	0.25 (0.07)	-0.0002 (0.003)	0.948	94,964	0.0343 (0.014)	0.013	36,803
	rs3783347	14	99909014	<i>WARS</i>	G/T	0.79	0.0168 (0.003)	1.3 × 10 ⁻¹⁰	132,544	0.02 (0.89)	0.0017 (0.003)	0.515	103,339	0.0274 (0.014)	0.044	42,850
	rs2302593	19	50888474	<i>GIPR</i>	C/G	0.50	0.0144 (0.002)	9.3 × 10 ⁻¹⁰	116,141	0.27 (0.05)	0.0025 (0.002)	0.265	96,976	-0.0322 (0.012)	0.006	40,781
	rs9368222	6	20794975	<i>CDKALI</i>	A/C	0.28	0.0143 (0.002)	1.0 × 10 ⁻⁹	128,453	0.09 (0.50)	-0.0047 (0.002)	0.037	98,894	0.0279 (0.012)	0.023	42,825
	rs10747083	12	131551691	<i>P2RX2</i>	A/G	0.66	0.0133 (0.002)	7.6 × 10 ⁻⁹	127,111	0.00 (1.00)	-0.0006 (0.002)	0.785	99,895	0.0269 (0.012)	0.026	42,790
	rs6072275	20	39177319	<i>TOP1</i>	A/G	0.16	0.0159 (0.003)	1.7 × 10 ⁻⁸	128,616	0.00 (1.00)	0.0038 (0.003)	0.169	99,018	-0.0110 (0.014)	0.435	42,853
	rs7651090	3	186996086	<i>IGFBP2</i>	G/A	0.31	0.0128 (0.002)	1.75 × 10 ⁻⁸	128,548	0.02 (0.86)	0.0003 (0.002)	0.900	98,924	0.0583 (0.012)	1.05 × 10 ⁻⁶	42,814
	rs576674	13	32452302	<i>KL</i>	G/A	0.15	0.0167 (0.003)	2.3 × 10 ⁻⁸	131,856	0.00 (1.00)	-0.0001 (0.003)	0.983	103,472	0.0308 (0.016)	0.060	42,849

Primary Trait	SNP	Chr	Position	Gene	Alleles (effect/other)	Freq effect allele	Primary trait			FI (BMI-adjusted)			2hGlu			
							Effect (SE)	Global analysis P value	Global analysis n	I ² estimate (P value)	Effect (SE)	Global analysis P value	Global analysis n	Effect (SE)	Global analysis P value	Global analysis n
	rs2943645	2	226807424	<i>IRS1</i>	T/C	0.63	0.0193 (0.002)	2.3×10^{-19}	99,023	0.00 (1.00)	0.0034 (0.002)	0.112	127475	0.0210 (0.011)	0.061	42,846
	rs10195252	2	165221337	<i>GRB14*</i>	T/C	0.60	0.0174 (0.002)	1.3×10^{-16}	98,997	0.00 (1.00)	0.0053 (0.002)	0.014	127005	0.0361 (0.011)	0.001	42,846
	rs2126259	8	9222556	<i>PPP1R3B</i>	T/C	0.11	0.0238 (0.003)	3.3×10^{-13}	99,021	0.14 (0.51)	0.0213 (0.003)	5.4×10^{-10}	127480	-0.0877 (0.017)	1.8×10^{-7}	42,849
	rs4865796	5	53308421	<i>ARL15</i>	A/G	0.67	0.0154 (0.002)	2.2×10^{-12}	98,314	0.48 (0.01)	0.0043 (0.002)	0.052	127784	0.0337 (0.012)	0.004	42,852
	rs17036328	3	12365484	<i>PPARG</i>	T/C	0.86	0.0212 (0.003)	3.6×10^{-12}	98,984	0.21 (0.31)	0.0051 (0.003)	0.103	128567	0.0335 (0.016)	0.031	42,843
	rs731839	19	38590905	<i>PEPD</i>	G/A	0.34	0.0148 (0.002)	5.1×10^{-12}	103,252	0.13 (0.55)	0.0046 (0.002)	0.038	132528	0.0142 (0.012)	0.220	42,847
	rs974801	4	106290513	<i>TET2</i>	G/A	0.38	0.0139 (0.002)	3.3×10^{-11}	103,489	0.09 (0.67)	0.0012 (0.002)	0.582	131866	0.0052 (0.011)	0.643	42,849
FI (BMI-adjusted)	rs459193	5	55842508	<i>ANKRD55/M AP3K1</i>	G/A	0.73	0.0147 (0.002)	1.12×10^{-10}	103,378	0.27 (0.17)	0.0111 (0.002)	1.6×10^{-6}	132989	0.0276 (0.012)	0.023	42,849
	rs6822892	4	157954125	<i>PDGFC</i>	A/G	0.68	0.0138 (0.002)	2.6×10^{-10}	103,432	0.00 (1.00)	0.0010 (0.002)	0.636	132951	0.0256 (0.012)	0.031	42,836
	rs4846565	1	217788727	<i>LYPEL1</i>	G/A	0.67	0.0132 (0.002)	1.8×10^{-9}	99,014	0.00 (1.00)	0.0066 (0.002)	0.003	127468	0.0132 (0.012)	0.254	42,853
	rs3822072	4	89960292	<i>FAM13A1</i>	A/G	0.48	0.0116 (0.002)	1.8×10^{-8}	99,977	0.00 (1.00)	0.0025 (0.002)	0.236	129432	0.0161 (0.011)	0.143	42,850
	rs6912327	6	34872900	<i>UHRF1BP1</i>	T/C	0.80	0.0165 (0.003)	2.3×10^{-8}	80,010	0.04 (0.91)	0.0074 (0.003)	0.011	103826	0.0139 (0.011)	0.391	34,761
											FG			FI (BMI-adjusted)		
	rs6975024	7	44198411	<i>GCK</i>	C/T	0.15	0.1026 (0.016)	5.2×10^{-11}	42,842	0.00 (1.00)	0.0605 (0.003)	2.9×10^{-99}	103,517	0.0063 (0.003)	0.030	98,458
	rs11782386	8	9239197	<i>PPP1R3B*</i>	C/T	0.87	0.0985 (0.017)	2.2×10^{-9}	42,852	0.00 (1.00)	-0.0167 (0.003)	5.5×10^{-7}	100,595	-0.0164 (0.003)	6.9×10^{-7}	95,565
2hGlu	rs1019503	5	96280573	<i>ERAP2</i>	A/G	0.48	0.0628 (0.011)	8.9×10^{-9}	42,851	19.6 (0.42)	-0.0061 (0.002)	0.003	108,113	0.0004 (0.002)	0.851	103,448

Primary Trait	SNP	Chr	Position	Gene	Alleles (effect/other)	Freq effect allele	Primary trait		I ²		FI (BMI-adjusted)		2hGlu			
							Effect (SE)	Global analysis P value	Global analysis n	estimate (P value)	Effect (SE)	Global analysis P value	Global analysis n	Effect (SE)	Global analysis P value	Global analysis n
2hGlu (BMI-adjusted)	rs7651090	3	186996086	<i>IGF2BP2</i>	G/A	0.30	0.064 (0.012)	4.5 × 10 ⁻⁸	42,792	63.4 (0.01)	0.0128 (0.002)	1.8 × 10 ⁻⁸	104,019	0.0003 (0.002)	0.900	98,924

Genome-wide loci for fasting glucose (FG), fasting insulin (FI), FI (adjusted for BMI) and 2hGlu are shown along with results for the other traits aligned to the trait-raising allele for the primary trait. "Non-MAGIC" SNPs (identified in other consortia and selected for the Metabochip to follow-up on other non-MAGIC traits) are indicated in bold. Freq denotes the allele frequency of the primary trait-raising allele. Per allele effect (SE) for FI represents changes of natural-log transformed levels of FI. *N* represents sample size. Heterogeneity was assessed using the I² index⁸⁶. The gene shown is the nearest gene to the lead SNP, other than those loci marked with an asterisk. For these loci, the nearest gene is also listed in Supplementary Tables 2a-d.