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Association analyses of 249,796 individuals reveal eighteen new loci associated with body mass index

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Abstract

Obesity is globally prevalent and highly heritable, but the underlying genetic factors remain largely elusive. To identify genetic loci for obesity-susceptibility, we examined associations between body mass index (BMI) and ~2.8 million SNPs in up to 123,865 individuals, with targeted follow-up of 42 SNPs in up to 125,931 additional individuals. We confirmed 14 known obesity-susceptibility loci and identified 18 new loci associated with BMI ($P < 5 \times 10^{-8}$), one of which includes a copy number variant near *GPRC5B*. Some loci (*MC4R*, *POMC*, *SH2B1*, *BDNF*) map near key hypothalamic regulators of energy balance, and one is near *GIPR*, an incretin receptor. Furthermore, genes in other newly-associated loci may provide novel insights into human body weight regulation.

Obesity is a major and increasingly prevalent risk factor for multiple disorders, including type 2 diabetes and cardiovascular disease¹,². While lifestyle changes have driven its prevalence to epidemic proportions, heritability studies provide evidence for a substantial genetic contribution (h²~40–70%) to obesity risk³,⁴. BMI is an inexpensive, non-invasive measure of obesity that predicts the risk of related complications⁵. Identifying genetic determinants of BMI could lead to a better understanding of the biological basis of obesity.

Genome-wide association (GWA) studies of BMI have previously identified ten loci with genome-wide significant ($P < 5 \times 10^{-8}$) associations in or near *FTO*, *MC4R*, *TMEM18*, *GNPDA2*, *BDNF*, *NEGR1*, *SH2B1*, *ETV5*, *MTCH2*, and *KCTD15*⁶–¹⁰. Many of these genes are expressed or known to act in the central nervous system, highlighting a likely neuronal component to the predisposition to obesity⁹. This pattern is consistent with results in animal models and studies of monogenic human obesity, where neuronal genes, particularly those expressed in the hypothalamus and involved in regulation of appetite or energy balance, are known to play a major role in susceptibility to obesity¹¹–¹³.

A full list of author contributions appears in the Supplementary Note.

Competing interests statement

The authors declare competing financial interests. A full list of competing interests appears in the Supplementary Note.

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Author contributions

The ten previously identified loci account for only a small fraction of the variation in BMI. Furthermore, power calculations based on the effect sizes of established variants have suggested that increasing the sample size would likely lead to the discovery of additional variants⁹. To identify more loci associated with BMI, we expanded the GIANT (Genetic Investigation of ANtropometric Traits) consortium GWA meta-analysis to include a total of 249,769 individuals of European ancestry.

Results

Stage 1 GWA studies identify novel loci associated with BMI

We first conducted a meta-analysis of GWA studies of BMI and ~2.8 million imputed or genotyped SNPs using data from 46 studies including up to 123,865 individuals (Online Methods, Supplementary Fig. 1 and Supplementary Note). This stage 1 analysis revealed 19 loci associated with BMI at $P < 5 \times 10^{-8}$ (Table 1, Fig. 1a and Supplementary Table 1). These 19 loci included all ten loci from previous GWA studies of BMI⁶ $^{-10}$, two loci previously associated with body weight¹⁰ (FAIM2 and SEC16B) and one locus previously associated with waist circumference¹⁴ (near TFAP2B). The remaining six loci, near GPRC5B, MAP2K5/LBXCOR1, TNNI3K, LRRN6C, FLJ35779/HMGCR, and PRKD1, have not previously been associated with BMI or other obesity-related traits.

Stage 2 follow-up leads to additional novel loci for BMI

To identify additional BMI-associated loci and to validate the loci that reached genomewide significance in stage 1 analyses, we examined SNPs representing 42 independent loci (including the 19 genome-wide significant loci) with stage 1 $P < 5 \times 10^{-6}$. Variants were considered to be independent if the pair-wise linkage disequilibrium (LD; r^2) was less than 0.1 and if they were separated by at least 1 Mb. In stage 2, we examined these 42 SNPs in up to 125,931 additional individuals (79,561 newly genotyped individuals from 16 different studies and 46,370 individuals from 18 additional studies for which GWA data were available; Table 1, Supplementary Note, and Online Methods). In a joint analysis of stage 1 and stage 2 results, 32 of the 42 SNPs reached $P < 5 \times 10^{-8}$. Even after excluding SNPs within these 32 confirmed BMI loci, we still observed an excess of small P-values compared to the distribution expected under the null hypothesis (Fig. 1b), suggesting that more BMI loci remain to be uncovered.

The 32 confirmed associations included all 19 loci with $P < 5 \times 10^{-8}$ at stage 1, 12 additional novel loci near *RBJ/ADCY3/POMC*, *QPCTL/GIPR*, *SLC39A8*, *TMEM160*, *FANCL*, *CADM2*, *LRP1B*, *PTBP2*, *MTIF3/GTF3A*, *ZNF608*, *RPL27A/TUB*, *NUDT3/HMGA1*, and one locus (*NRXN3*) previously associated with waist circumference¹⁵ (Table 1, Supplementary Table 1, Supplementary Fig. 1 and 2). In all, our study increased the number of loci robustly associated with BMI from 10 to 32. Four of the 22 new loci were previously associated with body weight¹⁰ or waist circumference¹⁴,¹⁵, whereas 18 loci had not previously associated with any obesity-related trait in the general population. Whilst we confirmed all loci previously established by large-scale GWA studies for BMI⁶–¹⁰ and waist circumference¹⁴,¹⁵, four loci identified by GWA studies for early-onset or adult morbid obesity¹⁶,¹⁷ [at *NPC1* (rs1805081; P = 0.0025), *MAF* (rs1424233; P = 0.25), *PTER*

(rs10508503; P = 0.64), and TNKS/MSRA (rs473034; P = 0.23)] showed limited or no evidence of association with BMI in our study.

As expected, the effect sizes of the 18 newly discovered loci are slightly smaller, for a given minor allele frequency, than those of the previously identified variants (Table 1 and Fig. 1c). The increased sample size also brought out more signals with low minor allele frequency. The BMI-increasing allele frequencies for the 18 newly identified variants ranged from 4% to 87%, covering more of the allele frequency spectrum than previous, smaller GWA studies of BMI (24%–83%)⁹, ¹⁰ (Table 1 and Fig. 1c).

We tested for evidence of non-additive (dominant or recessive) effects, SNP×SNP interaction effects and heterogeneity by sex or study among the 32 BMI-associated SNPs (Online Methods). We found no evidence for any such effects (P > 0.001, no significant results after correcting for multiple testing) (Supplementary Tables 1 and Supplementary Note).

Impact of 32 confirmed loci on BMI, obesity, body size, and other metabolic traits

Together, the 32 confirmed BMI loci explained 1.45% of the inter-individual variation in BMI of the stage 2 samples, with the FTO SNP accounting for the largest proportion of the variance (0.34%) (Table 1). To estimate the cumulative effect of the 32 variants on BMI, we constructed a genetic-susceptibility score that sums the number of BMI-increasing alleles weighted by the overall stage 2 effect sizes in the ARIC study (N = 8,120), one of our largest population-based studies (Online Methods). For each unit increase in the geneticsusceptibility score, approximately equivalent to one additional risk allele, BMI increased by 0.17 kg/m², equivalent to a 435–551 g gain in body weight in adults of 160–180 cm in height. The difference in average BMI between individuals with a high geneticsusceptibility score (38 BMI-increasing alleles, 1.5% (n=124) of the ARIC sample) and those with a low genetic-susceptibility score (21 BMI-increasing alleles, 2.2% (n=175) of the ARIC sample) was 2.73 kg/m², equivalent to a 6.99 to 8.85 kg body weight difference in adults 160-180 cm in height (Fig. 2a). Still, we note that the predictive value for obesity risk and BMI of the 32 variants combined was modest, although statistically significant (Fig. 2b, Supplementary Fig. 4). The area under the receiver operating characteristic (ROC) curve for prediction of risk of obesity (BMI 30 kg/m^2) using age, age² and sex only was 0.515 (P =0.023 compared to AUC of 0.50), which increased to 0.575 ($P < 10^{-5}$) when also the 32 confirmed SNPs were included in the model (Fig. 2b). The area under the ROC for the 32 SNPs only was 0.574 ($P < 10^{-5}$).

All 32 confirmed BMI-increasing alleles showed directionally consistent effects on risk of being overweight (BMI 25 kg/m²) or obese (30 kg/m²) in stage 2 samples, with 30 of 32 variants achieving at least nominally significant associations. The BMI-increasing alleles increased the odds of overweight by 1.013 to 1.138-fold, and the odds for being obese by 1.016- to 1.203-fold (Supplementary Table 2). In addition, 30 of the 32 loci also showed directionally consistent effects on the risk of extreme and early-onset obesity in a meta-analysis of seven case-control studies of adults and children (binomial sign test $P = 1.3 \times 10^{-7}$) (Supplementary Table 3). The BMI-increasing allele observed in adults also increased the BMI in children and adolescents with directionally consistent effects observed

for 23 of the 32 SNPs (binomial sign test P = 0.01). Furthermore, in family-based studies, the BMI-increasing allele was over-transmitted to the obese offspring for 24 of the 32 SNPs (binomial sign test P = 0.004) (Supplementary Table 3). As these studies in extreme obesity cases, children and families were relatively small ($N_{range} = 354 - 15,251$) compared to the overall meta-analyses, their power was likely insufficient to confirm association for all 32 loci. Nevertheless, these results show that the effects are unlikely to reflect population stratification and that they extend to BMI differences throughout the life course.

All BMI-increasing alleles were associated with increased body weight, as expected from the correlation between BMI and body weight (Supplementary Table 2). To confirm an effect of the loci on adiposity rather than general body size, we tested association with body fat percentage, which was available in a subset of the stage 2 replication samples (n = 5,359-28,425) (Supplementary Table 2). The BMI-increasing allele showed directionally consistent effects on body fat percentage at 31 of the 32 confirmed loci (binomial sign test $P = 1.54 \times 10^{-8}$) (Supplementary Table 2).

We also examined the association of the BMI loci with metabolic traits (type 2 diabetes 18, fasting glucose, fasting insulin, indices of beta-cell function (HOMA-B) and insulin resistance (HOMA-IR)¹⁹, and blood lipid levels²⁰) and with height (Supplementary Tables 2 and 4). Although many nominal associations are expected because of known correlations between BMI and most of these traits and because of overlap in samples, several associations stand out as possible examples of pleiotropic effects of the BMI-associated variants. Particularly interesting is the variant in the GIPR locus where the BMI-increasing allele is also associated with increased fasting glucose levels and lower 2-hour glucose levels (Supplementary Table 4)¹⁹, ²¹. The direction of the effect is opposite to what would be expected due to the correlation between obesity and glucose intolerance, but is consistent with the suggested roles of GIPR in glucose and energy metabolism (see below)²². Three loci show strong associations ($P < 10^{-4}$) with height (MC4R, RBJ/ADCY3/POMC and MTCH2/NDUFS3). Because BMI is weakly correlated with height (and indeed, the BMIassociated variants as a group show no consistent effect on height), these associations are also suggestive of pleiotropy. Interestingly, analogous to the effects of severe mutations in *POMC* and *MC4R* on height and weight²³, ²⁴, the BMI-increasing alleles of the variants near these genes were associated with decreased (POMC) and increased (MC4R) height, respectively (Supplementary Table 2).

Potential functional roles and pathways analyses

Although associated variants typically implicate genomic regions rather than individual genes, we note that some of the 32 loci include candidate genes with established connections to obesity. Several of the 10 previously identified loci are located in or near genes that encode neuronal regulators of appetite or energy balance, including $MC4R^{12}$, 25 , $BDNF^{26}$, and $SH2BI^{11}$, 27 . Each of these genes has been tied to obesity, not only in animal models, but also by rare human variants that disrupt each of these genes and lead to severe obesity 24 , 28 , 29 . Using the automated literature search programme, Snipper (Online Methods), we identified various genes within the novel loci with potential biological links to obesity-susceptibility (Supplementary Note). Among the novel loci, the location of rs713586

near *POMC* provides further support for a role of neuroendocrine circuits that regulate energy balance in susceptibility to obesity. *POMC* encodes several polypeptides including α -MSH, a ligand of the *MC4R* gene product³⁰, and rare mutations in *POMC* also cause human obesity²³,²⁹,³¹.

In contrast, the locus near GIPR, which encodes a receptor of gastric inhibitory polypeptide (GIP), suggests a role for peripheral biology in obesity. GIP, which is expressed in the K cell of the duodenum and intestine, is an incretin hormone that mediates incremental insulin secretion in response to oral intake of glucose. The variant associated with BMI is in strong LD ($r^2 = 0.83$) with a missense SNP in GIPR (rs1800437, Glu354Gln) that has recently been shown to influence the glucose and insulin response to an oral glucose challenge 21 . Although no human phenotype is known to be caused by mutations in GIPR, mice with disruption of Gipr are resistant to diet-induced obesity 32 . The association of a variant in GIPR with BMI suggests that there may be a link between incretins/insulin secretion and body weight regulation in humans as well.

To systematically identify biological connections among the genes located near the 32 confirmed SNPs, and to potentially identify new pathways associated with BMI, we performed pathway-based analyses using MAGENTA³³. Specifically, we tested for enrichment of BMI genetic associations in biological processes or molecular functions that contain at least one gene from the 32 confirmed BMI loci (Online Methods). Using annotations from the KEGG, Ingenuity, PANTHER, and Gene Ontology databases, we found evidence of enrichment for pathways involved in the platelet-derived growth factor (PDGF) signaling (PANTHER, P = 0.0008, P = 0.0061), translation elongation (PANTHER, P = 0.0008, P = 0.0008, hormone or nuclear hormone receptor binding (Gene Ontology, P < 0.0005, P = 0.0008, homeobox transcription (PANTHER, P = 0.0001, P = 0.001), regulation of cellular metabolism (Gene Ontology, P < 0.0002, P = 0.0002, P = 0.0031), neurogenesis and neuron differentiation (Gene Ontology, P < 0.0002, P = 0.0002, P = 0.0034), protein phosphorylation (PANTHER, P = 0.0001, P = 0.0045) and numerous other pathways related to growth, metabolism, immune and neuronal processes (Gene Ontology, P < 0.002, P = 0.0002, P = 0.0

Identifying possible functional variants

We used data from the 1000 Genomes Project and the HapMap Consortium to explore whether the 32 confirmed BMI SNPs were in LD (r^2 0.75) with common missense SNPs or copy number variants (CNVs) (Online Methods). Non-synonymous variants in LD with our signals were present in the *BDNF*, *SLC39A8*, *FLJ35779/HMGCR*, *QPCTL/GIPR*, *MTCH2*, *ADCY3*, and *LBXCOR1* genes. In addition, the rs7359397 signal was in LD with coding variants in several genes including *SH2B1*, *ATNX2L*, *APOB48R*, *SULT1A2*, and *AC138894.2* (Table 1, Fig. 3, Supplementary Table 6 and Supplementary Fig. 2). Furthermore, two SNPs tagged common CNVs. The first CNV was previously identified and is a 45-kb deletion near *NEGR1*⁹. The second CNV is a 21-kb deletion that lies 50kb upstream of *GPRC5B*; the deletion allele is tagged by the T-allele of rs12444979 ($r^2 = 1$) (Fig. 3). Although the correlations with potentially functional variants does not prove that

these variants are indeed causal, these provide first clues as to which genes and variants at these loci might be prioritized for fine-mapping and functional follow-up.

As many of the 32 BMI loci harbor multiple genes, we examined whether gene expression (eQTL) analyses could also direct us to positional candidates. Gene expression data were available for human brain, lymphocytes, blood, subcutaneous and visceral adipose tissue, and liver³⁴–³⁶ (Online Methods, Table 1 and Supplementary Table 7). Significant *cis*-associations, defined at the tissue-specific level, were observed between 14 BMI-associated alleles and expression levels (Table 1 and Supplementary Table 7). In several cases, the BMI-associated SNP was the most significant SNP or explained a substantial proportion of the association with the most significant SNP for the gene transcript in conditional analyses (*P*_{adj}>0.05). These significant associations included *NEGR1*, *ZC3H4*, *TMEM160*, *MTCH2*, *NDUFS3*, *GTF3A*, *ADCY3*, *APOB48R*, *SH2B1*, *TUFM*, *GPRC5B*, *IQCK*, *SLC39A8*, *SULT1A1*, and *SULT1A2* (Table 1 and Supplementary Table 7), making these genes higher priority candidates within the associated loci. However, we note that some BMI-associated variants were correlated with the expression of multiple nearby genes, making it difficult to determine the most relevant gene.

Evidence for the existence of additional associated variants

Because the variants identified by this large study explain only 1.45% of the variance in BMI (2–4% of genetic variance based on an estimated heritability of 40–70%), we considered how much the explained phenotypic variance could be increased by including more SNPs at various degrees of significance in a polygene model using an independent validation set (Online Methods)³⁷. We found that including SNPs associated with BMI at lower significance levels (up to P > 0.05) increased the explained phenotypic variance in BMI to 2.5%, or 4% to 6% of genetic variance (Fig. 4a). In a separate analysis, we estimated the total number of independent BMI-associated variants that are likely to exist with similar effect sizes to the 32 confirmed here (Online Methods)³⁸. Based on the effect size and allele frequencies of the 32 replicated loci observed in stage 2 and the power to detect association in the combined stage 1 and stage 2, we estimated that there are 284 (95% CI: 132-510) loci with similar effect sizes as the currently observed ones, which together would account for 4.5% (95% CI: 3.1-6.8%) of the variation in BMI or 6-11% of the genetic variation (based on an estimated heritability of 40–70%) (Supplementary Table 8). In order to detect 95% of these loci, a sample size of approximately 730,000 subjects would be needed (Fig. 4b). This method does not account for the potential of loci of smaller effect than those identified here to explain even more of the variance and thus provides an estimated lower bound of explained variance. These two analyses strongly suggest that larger GWA studies will continue to identify additional novel associated loci, but also indicate that even extremely large studies focusing on variants with allele frequencies above 5% will not account for a large fraction of the genetic contribution to BMI.

We examined whether selecting only a single variant from each locus for follow-up led us to underestimate the fraction of phenotypic variation explained by the associated loci. To search for additional independent loci at each of the 32 associated BMI loci, we repeated our GWA meta-analysis, conditioning on the 32 confirmed SNPs. Using a significance threshold

of 5×10^{-6} for SNPs at known loci, we identified one apparently independent signal at the MC4R locus; rs7227255 was associated with BMI ($P = 6.56 \times 10^{-7}$) even after conditioning for the most strongly associated variant near MC4R (rs571312) (Fig. 5). Interestingly, rs7227255 is in perfect LD ($r^2 = 1$) with a relatively rare MC4R missense variant (rs2229616, V103I, minor allele frequency = 1.7%) that has been associated with BMI in two independent meta-analyses³⁹, ⁴⁰. Furthermore, mutations at the MC4R locus are known to influence early-onset obesity²⁴, ⁴¹, supporting the notion that allelic heterogeneity may be a frequent phenomenon in the genetic architecture of obesity.

Discussion

Using a two-stage genome-wide association meta-analysis of up to 249,796 individuals of European descent, we have identified 18 additional loci that are associated with BMI at genome-wide significance, bringing the total number of such loci to 32. We estimate that more than 250 (i.e. 284 predicted loci – 32 confirmed loci) common variant loci with effects on BMI similar to those described here remain to be discovered, and even larger numbers of loci with smaller effects. A substantial proportion of these loci should be identifiable through larger GWA studies and/or by targeted follow-up of top signals selected from our stage 1 analysis. The latter approach is already being implemented through large-scale genotyping of samples informative for BMI using a custom array (the Metabochip) designed to support follow-up of thousands of promising variants in hundreds of thousands of individuals.

The combined effect on BMI of the associated variants at the 32 loci is modest, and even when we try to account for as-yet-undiscovered variants with similar properties, we estimate that these common variant signals account for only 6–11% of the genetic variation in BMI. There is a strong expectation that additional variance and biology will be explained using complementary approaches that capture variants not examined in the current study, such as lower frequency variants and short insertion-deletion polymorphisms. There is good reason to believe (based on our findings at *MC4R* and other loci – *POMC*, *BDNF*, *SH2B1* – which feature both common and rare variant associations) that a proportion of such low-frequency and rare causal variation will map to the loci already identified by GWA studies.

A primary goal of human genetic discovery is to improve understanding of the biology of conditions such as obesity⁴². One particularly interesting finding in this regard is the association between BMI and common variants near *GIPR*, which may indicate a causal contribution of variation in postprandial insulin secretion to the development of obesity. In most cases, the loci identified by the present study harbor few, if any, annotated genes with clear connections to the biology of weight regulation. This reflects our still limited understanding of the biology of BMI and obesity-related traits and is in striking contrast with the results from equivalent studies of certain other traits (such as autoimmune diseases or lipid levels). Thus, these results suggest that much novel biology remains to be uncovered, and that GWA studies may provide an important entry point. In particular, further examination of the associated loci through a combination of resequencing and finemapping to find causal variants, and genomic and experimental studies designed to assign function, could uncover novel insights into the biology of obesity.

In conclusion, we have performed GWA studies in large samples to identify numerous genetic loci associated with variation in BMI, a common measure of obesity. Because current lifestyle interventions are largely ineffective in addressing the challenges of growing obesity⁴³,⁴⁴, new insights into biology are critically needed to guide the development and application of future therapies and interventions.

Supplementary Material

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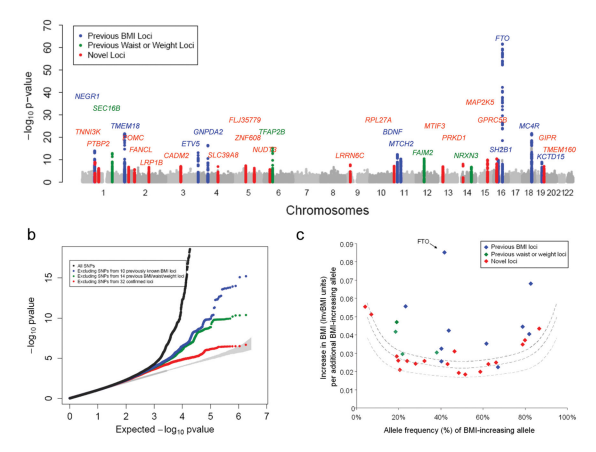


Figure 1. Genome-wide association results for the BMI meta-analysis

(a) Manhattan plot showing the significance of association between all SNPs and BMI in the stage 1 meta-analysis, highlighting SNPs previously reported to show genome-wide significant association with BMI (blue), weight or waist circumference (green), and the 18 new regions described here (red). The 19 SNPs that reached genome-wide significance at Stage 1 (13 previously reported and 6 new) are listed in Table 1). (b) Quantile-quantile (Q-Q) plot of SNPs in stage 1 meta-analysis (black) and after removing any SNPs within 1 Mb of the 10 previously reported genome-wide significant hits for BMI (blue), after additionally excluding SNPs from the four loci for waist/weight (green) and after excluding SNPs from all 32 confirmed loci (red). The plot was abridged at the Y-axis (at $P < 10^{-20}$) to better visualise the excess of small P-values after excluding the 32 confirmed loci (Supplementary Fig. 3 shows full-scale Q-Q plot). The shaded region is the 95% concentration band. (c) Plot of effect size (in inverse normally transformed units (invBMI)) versus effect allele frequency of newly identified and previously identified BMI variants after stage 1 + stage 2 analysis; including the 10 previously identified BMI loci (blue), the four previously identified waist and weight loci (green) and the 18 newly identified BMI loci (blue). The dotted lines represent the minimum effect sizes that could be identified for a given effectallele frequency with 80% (upper line), 50% (middle line), and 10% (lower line) power, assuming a sample size of 123,000 individuals and a α -level of 5×10^{-8} .

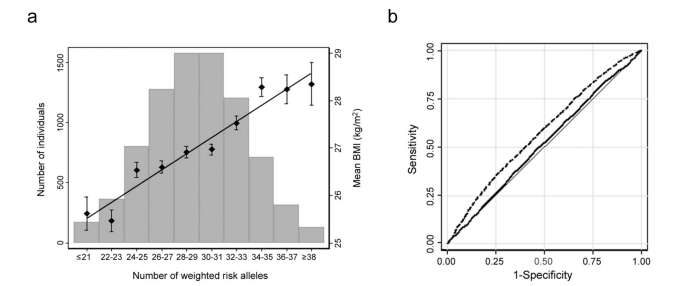


Figure 2. Combined impact of risk alleles on BMI/obesity

(a) Combined effect of risk alleles on average BMI in the population-based Atherosclerosis Risk in Communities (ARIC) study (n = 8,120 individuals of European descent). For each individual, the number of "best guess" replicated (n = 32) risk alleles from imputed data (0,1,2) per SNP was weighted for their relative effect sizes estimated from the stage 2 data. Weighted risk alleles were summed for each individual and the overall individual sum was rounded to the nearest integer to represent the individual's risk allele score (range 16–44). Along the x-axis, individuals in each risk allele category are shown (grouped 21 and 38 at the extremes), and the mean BMI (+/- SEM) is plotted (y axis on right), with the line representing the regression of the mean BMI values across the risk-allele scores. The histogram (y-axis on left) represents the number of individuals in each risk-score category. (b) The area under the ROC curve (AUC) of two different models predicting the risk of obesity (BMI = 30 kg/m^2) in the n = 8,120 genotyped individuals of European descent in the ARIC Study. Model 1, represented by the solid line, includes age, age², and sex (AUC = 0.515, P = 0.023 for difference from AUC_{null} = 0.50). Model 2, represented by the dashed line, includes age, age², sex, and the n = 32 confirmed BMI SNPs (AUC = 0.0575, $P < 10^{-5}$ for difference from AUC_{null} = 0.50). The difference between both AUCs is significant (P < 10^{-4}).

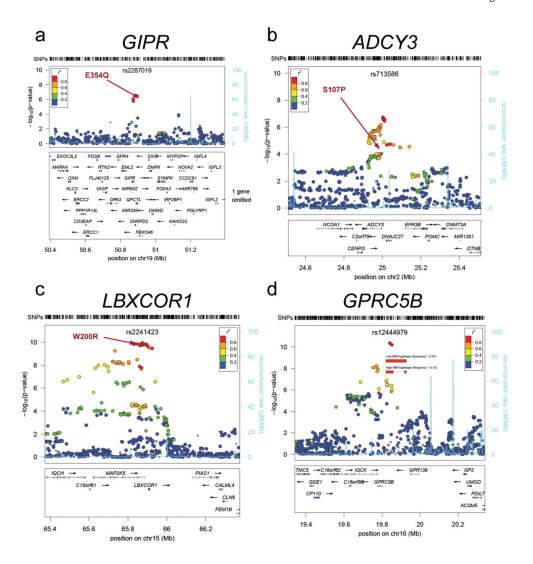


Figure 3. Regional plots of selected replicating BMI loci with missense and CNV variants SNPs are plotted by position on chromosome against association with BMI ($-\log_{10} P$ -value). The SNP name shown on the plot was the most significant SNP after stage 1 meta-analysis. Estimated recombination rates (from HapMap) are plotted in cyan to reflect the local LD structure. The SNPs surrounding the most significant SNP are color-coded to reflect their LD with this SNP (taken from pairwise r^2 values from the HapMap CEU database, www.hapmap.org). Genes, position of exons, and direction of transcription from UCSC genome browser (http://genome.ucsc.edu) are noted. Hashmarks represent SNP positions available in the meta-analysis. (a, b, c) Missense variants noted with their amino acid change for the gene noted above the plot. (d) Structural haplotypes and BMI association signal in the *GPRC5B* region. A 21 kb deletion polymorphism is associated with 4 SNPs (r^2 =1.0) that comprise the best haplogroup associating with BMI. Plots were generated using LocusZoom (http://csg.sph.umich.edu/locuszoom).

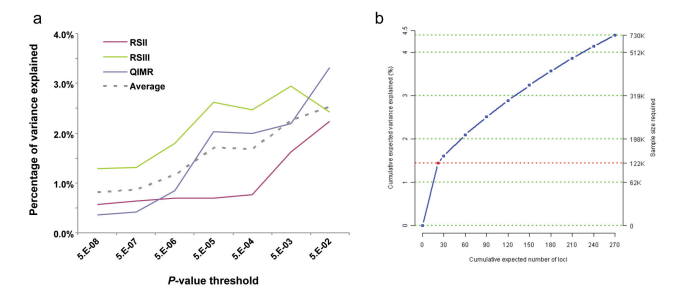


Figure 4. Phenotypic variance explained by common variants

(a) Variance explained is higher when SNPs not reaching genome-wide significance are included in the prediction model. The y-axis represents the proportion of variance explained at different P-value thresholds from stage 1 meta-analysis. Results are given for three studies (RSII, RSIII, QIMR), which were not included in the meta-analysis, after exclusion of all samples from The Netherlands (for RSII and RSIII) and the United Kingdom (for QIMR) from the discovery analysis for this sub-analysis. The dotted line represents the weighted average of the explained variance of three validation sets. (b) Cumulative number of susceptibility loci expected to be discovered, including those we have already identified and others that have yet to be detected, by the expected percentage of phenotypic variation explained and sample size required for a one-stage GWA study assuming a GC correction is utilized. The projections are based on loci that achieved a significance level of $P < 5 \times 10^{-8}$ in the joint analysis of stage 1 and stage 2 and the distribution of their effect sizes in stage 2. The dotted red line corresponds to the expected phenotypic variance explained by the 22 loci that are expected to be discovered in a one-stage GWAS with the sample size of stage 1 of this study.

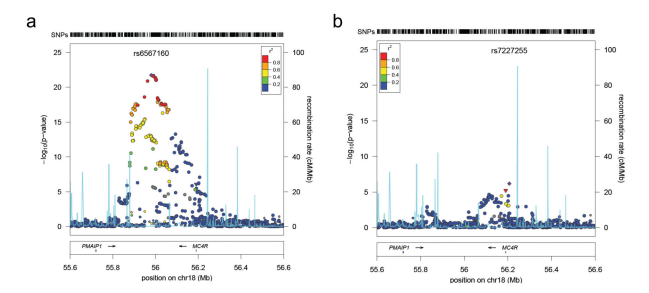


Figure 5. Second signal at the MC4R locus contributing to BMI

SNPs are plotted by position in a 1 Mb window of chromosome 18 against association with BMI (log₁₀ *P*-value). Panel (**a**) highlights the most significant SNP in stage 1 meta-analysis, panel (**b**) the most significant SNP after conditional analysis where the model included the most strongly associated SNP from panel A as a covariate. Estimated recombination rates (from HapMap) are plotted in cyan to reflect the local LD structure. The SNPs surrounding the most significant SNP are color-coded to reflect their LD with this SNP (taken from pairwise r² values from the HapMap CEU database, www.hapmap.org). Genes, exons, and direction of transcription from UCSC genome browser (genome.ucsc.edu) are noted. Hashmarks at the top of the figure represent positions of SNPs in the meta-analysis. Regional plots were generated using LocusZoom (http://csg.sph.umich.edu/locuszoom).

Table 1

Stage 1 and stage 2 results of the 32 SNPs that were associated with BMI at genome-wide significance $(P < 5.10^{-8})$ levels.

SNP	Nearest gene	Other nearby genes*	Chr	Chr Position** (bp)	Alle	Alleles**			Explained variance (%)	Stage 1 P-value	Stage 2 P-value	Stage	Stage 1 + 2
				•	Effect	Other	Frequency effect allele (%)	rer aneie change in bivii beta (se)				u u	P-value
Previous BMI loci	AI loci												
rs1558902	FTO		16	52,361,075	в	t	42%	0.39 (0.02)	0.34%	2.05E-62	1.007E-60	192,344	4.8E-120
rs2867125	TMEM18		2	612,827	၁	t	83%	0.31 (0.03)	0.15%	2.42E-22	4.42E-30	197,806	2.77E-49
rs571312	MC4R B		18	55,990,749	в	၁	24%	0.23 (0.03)	0.10%	1.82E-22	3.19E-21	203,600	6.43E-42
rs10938397	GNPDA2		4	44,877,284	ъъ	в	43%	0.18 (0.02)	0.08%	4.35E-17	1.45E-15	197,008	3.78E-31
rs10767664	$BDNF^{B,M}$		11	27,682,562	в	t.	78%	0.19 (0.03)	0.07%	5.53E-13	1.17E-14	204,158	4.69E-26
rs2815752	NEGRICQ		-	72,585,028	а	ac	61%	0.13 (0.02)	0.04%	1.17E-14	2.29E-09	198,380	1.61E-22
rs7359397	SH2B1Q,B,M	APOB48RQ.M, SULT1A2Q.M, AC138894.2M, ATXN2LM, TUFMQ	16	28,793,160	t t	၁	40%	0.15 (0.02)	0.05%	1.75E-10	7.89E-12	204,309	1.88E-20
rs9816226	ETV5		3	187,317,193	t	а	82%	0.14 (0.03)	0.03%	7.61E-14	1.15E-06	196,221	1.69E-18
rs3817334	MTCH2Q,M	NDUFS3 Q , CUGBP1 Q	11	47,607,569	t	၁	41%	0.06 (0.02)	0.01%	4.79E-11	1.10E-03	191,943	1.59E-12
rs29941	KCTD15		19	39,001,372	ρū	в	%29	0.06 (0.02)	0.00%	1.31E-09	2.40E-02	192,872	3.01E-09
Previous wa	Previous waist & weight loci												
rs543874	SEC16B		-	176,156,103	фD	в	19%	0.22 (0.03)	0.07%	1.66E-13	2.41E-11	179,414	3.56E-23
rs987237	TFAP2B		9	50,911,009	ьø	а	18%	0.13 (0.03)	0.03%	5.97E-16	2.40E-06	195,776	2.90E-20
rs7138803	FAIM2		12	48,533,735	в	50	38%	0.12 (0.02)	0.04%	3.96E-11	7.82E-08	200,064	1.82E-17
rs10150332	NRXN3		14	79,006,717	၁	t	21%	0.13 (0.03)	0.02%	2.03E-07	2.86E-05	183,022	2.75E-11
Newly ident	Newly identified BMI loci												
rs713586	RBJ	ADCY3Q, M , $POMCQ$, B	2	25,011,512	၁	٠	47%	0.14 (0.02)	0.06%	1.80E-07	1.44E-16	230,748	6.17E-22
rs12444979	GPRC5BC,Q	IQCK Q	16	19,841,101	၁	+	87%	0.17 (0.03)	0.04%	4.20E-11	8.13E-12	239,715	2.91E-21
rs2241423	MAP2K5	LBXCOR1 M	15	65,873,892	ad	а	78%	0.13 (0.02)	0.03%	1.15E-10	1.59E-09	227,950	1.19E-18
rs2287019	QPCTL	$GIPR^{B,M}$	19	50,894,012	ပ	t	80%	0.15 (0.03)	0.04%	3.18E-07	1.40E-10	194,564	1.88E-16
rs1514175	TNNI3K		-	74,764,232	æ	ac	43%	0.07 (0.02)	0.02%	1.36E-09	7.04E-06	227,900	8.16E-14
rs13107325	SLC39A8Q,M		4	103,407,732	t	၁	7%	0.19 (0.04)	0.03%	1.37E-07	1.93E-07	245,378	1.50E-13
rs2112347	FLJ35779M	HMGCR B	5	75,050,998	t	5.0	63%	0.10 (0.02)	0.02%	4.76E-08	8.29E-07	231,729	2.17E-13

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beta (se) \$\frac{3}{2} \text{***}\$ 0.11 (0.02) 0.02% 1.88E-08 3.19E-06 0.09 (0.02) 0.02% 1.04E-07 1.59E-06 0.10 (0.02) 0.03% 2.69E-06 1.72E-07 0.10 (0.02) 0.02% 9.81E-08 5.32E-05 0.17 (0.02) 0.01% 1.11E-08 2.25E-04 0.09 (0.03) 0.01% 7.65E-07 9.47E-05 0.06 (0.02) 0.01% 7.65E-07 8.24E-04 0.07 (0.02) 0.01% 7.64E-07 1.88E-04 0.06 (0.02) 0.01% 7.57E-08 1.00E-03 0.06 (0.02) 0.01% 7.57E-08 1.00E-03	SNP	Nearest gene	Other nearby genes*	Chr	Chr Position** (bp)	Alleles**	es**	Emoraron or officer ellele	Per allele change in BMI	Explained variance (%) Stage 1 P-value Stage 2 P-value	Stage 1 P-value	Stage 2 P-value	Stag	Stage 1 + 2
0 D ZS,404,39 g a 31% 0.11 (0.02) 0.02% 1.88E-08 3.19E-06 0 D ZC3H4 Q 19 22,260,843 a g 67% 0.01 (0.02) 0.02% 1.04E-07 1.59E-06 1 A 25,260,843 1 c 29% 0.10 (0.02) 0.03% 2.69E-06 1.72E-07 2 A 25,568,840 2 a 20% 0.17 (0.02) 0.02% 9.81E-08 5.32E-05 2 A 25,568,840 2 4 20 4 0.17 (0.02) 0.02% 9.81E-08 5.32E-05 2 A 4 25,584,863 4 4 4 4 4 4 4 4 4 4 5.2E-04 5.2E-04 <t< th=""><th></th><th></th><th></th><th></th><th></th><th>Effect</th><th>Other</th><th>riequency effect affete (%)</th><th>beta (se)</th><th></th><th></th><th></th><th>п</th><th>P-value</th></t<>						Effect	Other	riequency effect affete (%)	beta (se)				п	P-value
MANILAS ZASHAQ 152 60.843 a 67% 609 (0.02) 100 (0.02) 10.4E-07 1.59E-06 FANCL ANCL 2 51,56.381 1 c 29% 610 (0.02) 0.03% 1.04E-07 1.59E-06 FANCL ANCADAS 2 51,56.381 1 c 29% 0.10 (0.02) 0.03% 0.03% 0.58E-07 1.72E-07 PANCL ANCADAS 1 c 4% 0.10 (0.02) 0.01% 0.11E-08 5.32E-05 LRPIB ANCADAS 1 25.848.63 1 c 4% 0.04 (0.02) 0.01% 0.01% 0.11E-08 0.25E-04 MITIS ATRAGA 2 3 4 3 4	rs10968576	LRRN6C		6	28,404,339	ρū	g	31%	0.11 (0.02)	0.02%	1.88E-08	3.19E-06	216,916	2.65E-13
FANCL CADM2 EANCL CADM2 CADM2 <th< td=""><td>rs3810291</td><td>TMEM160 Q</td><td>$ZC3H4 \ Q$</td><td>19</td><td>52,260,843</td><td>æ</td><td>50</td><td>%29</td><td>0.09 (0.02)</td><td>0.02%</td><td>1.04E-07</td><td>1.59E-06</td><td>233,512</td><td>1.64E-12</td></th<>	rs3810291	TMEM160 Q	$ZC3H4 \ Q$	19	52,260,843	æ	50	%29	0.09 (0.02)	0.02%	1.04E-07	1.59E-06	233,512	1.64E-12
PRKD1 CADM2 85,966,840 g a 20% 0.10 (0.02) 0.02% 9.81E-08 5.32E-05 PRKD1 LAPIB 2 2,584,863 t c 4% 0.17 (0.05) 0.01% 0.11E-08 5.32E-05 LAPIB LAPIB 2 142,676,401 c t 18% 0.09 (0.03) 0.01% 2.38E-07 9.47E-05 MTIF3 GTF3A Q 13 26,918,180 g a 24% 0.09 (0.03) 0.01% 7.56E-07 9.47E-05 ZNF608 TUB B 13 26,918,180 g a 24% 0.09 (0.03) 0.01% 7.04E-07 1.88E-04 RPLZA TUB B TUB B 1 8,561,169 g t 22% 0.06 (0.02) 0.01% 7.57E-08 7.39E-04 NUDT3 HMGA1 B TMGA1 B 3 34,10,847 g 1 2 2 2 2 2 2 2 2 2 2 2	rs887912	FANCL		2	59,156,381	t	၁	29%	0.10 (0.02)	0.03%	2.69E-06	1.72E-07	242,807	1.79E-12
PRKD1 C PRKD2	rs13078807	CADM2		3	85,966,840	ьa	В	20%	0.10 (0.02)	0.02%	9.81E-08	5.32E-05	237,404	3.94E-11
LRP1B LRP1B c t 18% 0.09 (0.03) 0.02% 2.38E-07 9.4TE-05 PTBP2 PTBP2 a 59% 0.06 (0.02) 0.01% 7.65E-07 4.48E-05 MTIF3 GTF3A Q 13 26,918,180 g a 24% 0.09 (0.03) 0.02% 1.20E-07 8.24E-04 ZNF608 ARD27A TUB B 11 8,561,169 c t 48% 0.06 (0.02) 0.01% 7.54E-08 1.00E-07 1.88E-04 NUDT3 HMGA1 B 6 34,410,847 g a 21% 0.06 (0.02) 0.06 (0.02) 0.01% 0.01% 7.51E-08 7.39E-04	rs11847697	PRKD1		14	29,584,863	t	၁	4%	0.17 (0.05)	0.01%	1.11E-08	2.25E-04	241,667	5.76E-111
PTBP2 ATIF3 C a 59% 0.06 (0.02) 0.01% 7.65E-07 4.48E-05 MTIF3 GTF3A Q 13 26,918,180 g a 24% 0.09 (0.03) 0.02% 1.20E-07 8.24E-04 ZNF608 3 124,360,002 a c 48% 0.07 (0.02) 0.01% 7.57E-08 1.88E-04 NUDT3 HMGA1 B 6 34,410,847 g a 21% 0.06 (0.02) 0.01% 0.01% 7.51E-06 7.39E-04	rs2890652	LRP1B		2	142,676,401	၁	t	18%	0.09 (0.03)	0.02%	2.38E-07	9.47E-05	209,068	1.35E-10
MTIF3 GTF3A Q 13 26,918,180 g a 24% 0.09 (0.03) 0.02% 1.20E-07 8.24E-04 ZNF608 2 124,360,002 a c 48% 0.07 (0.02) 0.01% 7.04E-07 1.88E-04 RPL27A TUB B 11 8,561,169 c t 52% 0.06 (0.02) 0.01% 7.57E-08 1.00E-03 NUDT3 HMGAI B 6 34,410,847 g a 21% 0.06 (0.02) 0.01% 0.01% 2.81E-06 7.39E-04	rs1555543	PTBP2		_	96,717,385	၁	В	29%	0.06 (0.02)	0.01%	7.65E-07	4.48E-05	243,013	3.68E-10
ZNF608 a c 48% 0.07 (0.02) 0.01% 7.04E-07 1.88E-04 RPL27A TUB B 11 8,561,169 c t 52% 0.06 (0.02) 0.01% 7.57E-08 1.00E-03 NUDT3 HMGAI B 6 34,410,847 g a 21% 0.06 (0.02) 0.01% 0.01% 2.81E-06 7.39E-04	rs4771122	MTIF3	GTF3A Q	13	26,918,180	ac	а	24%	0.09 (0.03)	0.02%	1.20E-07	8.24E-04	198,577	9.48E-10
RPL27A TUB B 11 8,561,169 c t 52% 0.06 (0.02) 0.01% 7.57E-08 1.00E-03 NUDT3 HMGA IB 6 34,410,847 g a 21% 0.06 (0.02) 0.01% 2.81E-06 7.39E-04	rs4836133	ZNF608		5	124,360,002	в	၁	48%	0.07 (0.02)	0.01%	7.04E-07	1.88E-04	241,999	1.97E-09
NUDT3 HMGA1B 6 34,410,847 g a 21% 0.06 (0.02) 0.01% 2.81E-06 7.39E-04	rs4929949	RPL27A	TUB B	11	8,561,169	၁	ţ	52%	0.06 (0.02)	0.01%	7.57E-08	1.00E-03	249,791	2.80E-09
	rs206936	NUDT3	HMGA1 B	9	34,410,847	oo	в	21%	0.06 (0.02)	0.01%	2.81E-06	7.39E-04	249,777	3.02E-08

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* Genes within ± -500 kb of the lead SNP

**
Positions according to Build 36 and allele coding based on the positive strand

*** Effect sizes in kg/m2 obtained from Stage 2 cohorts only

 $\mathcal{Q}_{\mathrm{Association}}$ and eQTL data converge to affect gene expression

 $^{\it B}$ Biological candidate

Memory Me

 $c_{
m CNV}$