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Defining the role of common variation in the genomic and biological architecture of adult human height

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Abstract

Using genome-wide data from 253,288 individuals, we identified 697 variants at genome-wide significance that together explain one-fifth of heritability for adult height. By testing different numbers of variants in independent studies, we show that the most strongly associated ~2,000, ~3,700 and ~9,500 SNPs explained ~21%, ~24% and ~29% of phenotypic variance. Furthermore, all common variants together captured the majority (60%) of heritability. The 697 variants clustered in 423 loci enriched for genes, pathways, and tissue-types known to be involved in growth and together implicated genes and pathways not highlighted in earlier efforts, such as signaling by fibroblast growth factors, WNT/beta-catenin, and chondroitin sulfate-related genes. We identified several genes and pathways not previously connected with human skeletal growth, including mTOR, osteoglycin and binding of hyaluronic acid. Our results indicate a genetic architecture for human height that is characterized by a very large but finite number (thousands) of causal variants.

Height is a classical polygenic trait that has provided general insights into the genetic architecture of common human traits and diseases, and into the prospects and challenges of different methods used to identify genetic risk factors. Studies consistently estimate that the additive genetic contribution to normal variation in adult height ("narrow sense heritability") is approximately $80\%^{1-3}$. Previous analysis of genome-wide association studies (GWAS) of adult height showed that common variants together account for 50% of this heritable contribution to height variation^{4,5}. The most recent GWAS of adult height identified 180 loci, which together highlighted many genes relevant to human skeletal growth that had not been implicated in previous studies⁶. Common variants in these loci, however, only accounted for 10% of the phenotypic variation (~12% of heritability). Here, we report results from a GWAS meta-analysis of adult height in 253,288 individuals of European ancestry. We show that additive contributions of fewer than 10,000 SNPs (at $P < 5 \times 10^{-3}$) can account for 36% of the heritability of adult height. Variants reaching genome-wide significance ($P < 5 \times 10^{-8}$) in this larger study (697 SNPs) clustered in loci, were substantially enriched for regulatory variants, and implicated multiple known and previously unknown

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genes and pathways relevant to growth. More broadly, our results provide evidence that increasing GWAS sample sizes to the order of 100,000s, now plausible for many common traits, will likely continue to identify the variants and loci that close the "missing heritability" gap, whilst improving knowledge of the biology of those traits.

Results

The overall analysis strategy is illustrated in Supplementary Figure 1. We first performed a GWAS meta-analysis of adult height using summary statistics from 79 studies consisting of 253,288 individuals of European ancestry (Online Methods). We identified 697 SNPs that reached genome-wide significance ($P < 5 \times 10^{-8}$) using an approximate conditional and joint multiple-SNP (COJO) analysis⁷ in GCTA⁸ (Online Methods) which takes linkage disequilibrium (LD) between SNPs into account (Supplementary Table 1; Supplementary Figs. 2-3). The 697 SNPs clustered in 423 loci, with a locus defined as one or multiple jointly associated SNPs located within ±1Mb of each other. Most of these 697 SNPs are uncorrelated although those in close physical proximity (e.g. < 1Mb) may be in partial LD (see Supplementary Table 1 for LD between adjacent pairs of the 697 SNPs). The clustering of signals was non-random (empirical enrichment of 1.4 fold, $P<1\times10^{-4}$) with 90, 26 and 31 loci containing 2, 3 and 4 signals respectively, (Supplementary Note and Supplementary Tables 1 and 2). We observed strong evidence of clustering of association signals within loci across a range of locus sizes, from 100kb to 1.25Mb, but the clustering was almost entirely driven by variants within 250kb of index SNPs (Supplementary Note and Supplementary Table 2). As shown in Figure 1 and Supplementary Figure 4, in some loci, multiple signals cluster tightly around a single gene, whereas in other cases, the clustering of associated variants is likely due to multiple different height-related genes being in close proximity.

Of the 697 SNPs, 403 were represented on the Metabochip array⁹. Using data from 80,067 individuals genotyped on the Metabochip array from 37 independent studies, we observed very strong evidence of concordance of effect sizes between the Metabochip and GWAS samples ($P = 1.9 \times 10^{-160}$); and >99% of variants were directionally consistent between Metabochip and GWAS (Online Methods, Supplementary Note, and Supplementary Table 3).

We observed a large genome-wide 'inflation' factor of the test statistic for association even after we corrected each study's test statistics by its individual inflation factor (single $\lambda_{GC}=1.94$). At least two phenomena could have contributed to this observation. First, as described previously ¹⁰, highly polygenic models of inheritance are expected to increase the genomic inflation factor to levels comparable to what we observe. Second, height is particularly susceptible to confounding by population ancestry (stratification), which can also lead to inflation of the test statistics. We addressed these possibilities by comparing our results with those obtained using more stringent corrections for stratification (linear mixed models), and with results obtained in subsets of studies in which a purely family-based analysis was feasible, and by performing a within-family prediction analysis which partitioned the variance in the genetic predictor into the contributions of true associations and population stratification.

Our linear mixed model (LMM) analyses, performed in a subset of 15 individual studies comprising 59,380 individuals, provided strong evidence that the inflated statistics were driven predominantly by the highly polygenic nature of the trait. This approach utilizes a genomic relationship matrix (GRM) calculated through genome-wide SNP data to correct for distant relatedness between all pairs of individuals within a study. This resulted in a single λ_{GC} of 1.20. This value was entirely consistent with the single λ_{GC} of 1.20 obtained from the standard GWAS analysis of the same individuals and a single λ_{GC} of 1.94 obtained from the full 253,288 individuals (Supplementary Table 4). Because this approach may be overly conservative for a strongly genetic and highly polygenic trait, each study additionally repeated the analyses for each chromosome using a GRM generated from the remaining 21 chromosomes, or in the case of the largest study (WGHS) repeating the analysis for all odd numbered chromosomes using a GRM generated from the even numbered chromosomes and vice versa. The single λ_{GC} inflation factor for this analysis, 1.23, was also entirely consistent with the standard GWAS results (Online Methods, Supplementary Note, and Supplementary Table 4).

Our family based analyses also provided strong evidence that the inflated statistics are driven predominantly by the highly polygenic nature of height. We assessed whether variants that reached genome-wide significance after single GC correction replicated in family-based analyses of up to 25,849 samples (effective sample size 14,963, using methods that are immune to stratification (Online Methods, Supplementary Note, and Supplementary Tables 5 and 6). We identified genome-wide significant associations from a meta-analysis that excluded the family-based samples, and tested these associations for replication in the family-based samples; a lower rate of replication than expected could be due to inflation of effect sizes in the discovery sample from the "winner's curse" and/or stratification. Of 416 genome-wide significant SNPs representing multiple signals selected after exclusion of family-based studies, 371 SNPs had a consistent direction of effect (compared with 208 expected by chance, and 400 expected in the absence of any inflation of estimated effect sizes), and 142 replicated with P<0.05 (compared with 21 expected by chance, and 210 expected in the absence of effect size inflation; Supplementary Table 5). These analyses (particularly the directional consistency) shows that most of the loci represent true associations, but also shows that there is a modest inflation in the effect size estimates, due to stratification and/or the winner's curse. To distinguish between these possibilities, we repeated this analysis, substituting for the family-based samples a random set of studies with similar total effective sample size. The number of replicating loci was only slightly lower in the family-based cohorts than in the random samples (Supplementary Table 5, 12-17 fewer replications attributable to stratification at different P-value thresholds). This indicates that most of the modest inflation in effect estimates is due to the winner's curse, that a small amount of inflation is due to residual stratification, and that few (upper limit ~15-25; Supplementary Note and Supplementary Table 5) if any of the loci that reach genome-wide significance after single GC correction are likely to be complete false positives due to stratification (that is, no real association whatsoever with height).

Variance explained by SNPs at different significance levels

Having established that single GC correction is sufficient to identify SNPs that are likely to be truly associated with height, we next performed a series of analyses using GWAS data from five independent validation studies to quantify the fraction of phenotypic variance explained by SNPs selected from the GCTA-COJO analyses⁷ of the meta-analysis data, which excluded data from the validation studies, at a range of statistical thresholds, and to quantify the accuracy of predicting height using these selected SNPs (Online Methods). We first developed a new method that uses within-family prediction to partition the variance of the SNP-based predictor into components due to real SNP effects, errors in estimating SNP effects, and population stratification (Online Methods), and applied the method to data on full-sib pairs from three of the five validation studies (Online Methods). Consistently across the three studies, all the partitioned variance components increased as a less stringent significance level was used for SNP selection in the discovery sample and the error variance increased more dramatically than the genetic variance when more SNPs selected at a less significance level were included in the predictor (Fig. 2a-c). We demonstrated the partitioning of variance due to population stratification by the within-family prediction analyses with and without adjusting for principal components (PCs) (Supplementary Fig. 5). The results again confirmed that the impact of population stratification on the top associated SNPs was minor and demonstrated that the variation in the predictor due to true SNP effect, estimation error and population stratification was quantifiable. We next inferred, using these partitioned variance components from the within-family prediction analysis, how well different selected sets of SNPs would predict height in independent samples. We showed that the observed prediction accuracy (squared correlation between phenotype and predictor, R^2) in five different population-based cohorts was highly consistent with the values inferred from the within-family based analyses, with prediction accuracy peaking at ~17% using the ~1,900 SNPs reaching $P < 5 \times 10^{-5}$ (Fig. 2d). Finally we estimated variance explained by the selected SNPs in population-based studies using the GCTA-GREML method^{4,8} (Fig. 2e). The results showed that ~670 SNPs at $P < 5 \times 10^{-8}$ and ~9.500 SNPs at $P < 5 \times 10^{-3}$ captured ~16% and ~29% of phenotypic variance respectively (Table 1), which was also consistent with the estimates inferred from the within-family prediction analysis. As shown in equation [19], prediction R^2 is not equal to the variance explained but a function of the variance of true SNP effects and the error variance in estimating SNP effects, in the absence of population structure. This is demonstrated in Figure 2, where at thresholds below genomewide significance, variance explained is higher than the prediction accuracy, because the latter is deflated both by imprecise estimates of effect sizes (estimation errors) and by inclusion of SNPs that are not associated with height. The estimate of variance explained by all the HapMap3 (ref. 11) SNPs without SNP selection was ~50% (Table 1), consistent with previous estimates^{4,5}. Thus, a group of ~9,500 SNPs (representing <1% of common SNPs) selected at $P < 5 \times 10^{-3}$, explained ~29% of phenotypic variance. Since ~50% of phenotypic variance is explained by all common SNPs, the selected set of SNPs, despite being limited to <1% of common SNPs, accounts for the majority of variance attributable to all common SNPs (29/50 ~ 60%). This set of ~9,500 SNPs strongly clustered with the newly established height loci: 1,704 (19%) variants were located within 250kb of one of the 697 genome-wide associated SNPs, suggesting that a substantial fraction of "missing heritability" is within already identified loci. This clustering of additional variants within identified loci was

confirmed in a parallel analysis based on two left-out studies where we observed that SNPs in closer physical proximity with the top associated SNPs explained disproportionally more variance (Online Methods and Supplementary Fig. 6).

Larger GWAS identifies new biologically relevant genes and pathways

Having shown that ~1% of variants can account for the majority of heritability attributable to common variation, we next considered whether the expanded set of height-associated variants could be used to identify the genomic features and biological pathways of most relevance to normal variation in adult height. To test whether our GWAS could implicate new biology, we used established and novel approaches to test whether the height-associated loci were enriched for functionally relevant variants, genes, pathways, and tissues.

As with the 180 variants identified in our previous analysis, the 697 variants were non-randomly distributed with respect to functional and putatively functional regions of the genome (Online Methods). We observed that height associated variants were enriched for non-synonymous SNPs (nsSNPs) (empirical enrichment of 1.2 fold, P=0.02), cis-regulatory effects in blood (empirical enrichment of 1.5 fold, P=0.03), a curated list of genes that underlie monogenic syndromes of abnormal skeletal growth¹² (empirical enrichment 1.4 fold, P=0.013), associations with apparently unrelated complex traits in the NHGRI GWAS catalog (empirical enrichment 2.6 fold, P<1×10⁻⁴) and functional chromatin annotations in multiple tissues and cell types (empirical enrichment 1.8 fold, P<1×10⁻³) (Supplementary Note and Supplementary Tables 7-11).

The greater resolution of height associated variants provided by increased sample size, combined with improved gene prioritization and gene set enrichment approaches, identified multiple new tissues, gene sets and specific genes that are highly likely to be involved in the biology of skeletal growth. Specifically, using a variety of established and novel pathway methods, we identified ~3 times as many enriched pathways and prioritized ~5 times as many genes (including genes newly prioritized in previously identified loci) compared to results derived from identical pathway methods to the previous GWAS of 133,000 individuals (Table 2).

We first focused on existing pathway and gene prioritization methods: (1) MAGENTA¹³, a method designed to identify gene sets enriched in GWAS data, and (2) GRAIL¹⁴, which uses published literature to highlight connections between likely relevant genes within GWAS loci. As expected, the GRAIL and MAGENTA analyses confirmed several previously identified gene sets and pathways clearly relevant to skeletal growth, but in the larger sample they also provided evidence for additional known and novel genes, gene sets and protein complexes not identified in our previous smaller study (for example, FGF signaling, WNT signaling, osteoglycin, and other genes related to bone or cartilage development) (Supplementary Tables 12-13 and Supplementary Fig. 7).

To obtain more detailed insight into height biology, we applied DEPICT, a novel datadriven integrative method that uses gene sets reconstituted based on large scale expression data to prioritize genes and gene sets, and also to identify tissues enriched in highly expressed genes from associated loci (Pers *et al. in preparation;* Online Methods and

Supplementary Note). The DEPICT analysis highlighted 2,330 reconstituted gene sets (after pruning for high levels of redundancy). These gene sets both confirmed and extended the MAGENTA and GRAIL findings, and identified novel pathways not identified in our previous height GWAS (for example regulation of beta-catenin, biology related to glycosaminoglycans such as chondroitin sulfate and hyaluronic acid, and mTOR signaling) (Supplementary Table 14). Gene sets identified based on 327 strictly novel height variants (>1Mb from the 180 known variants loci) highly resembled gene sets highlighted by the already known 180 loci (Spearman's rank correlation coefficient between gene set enrichment Z-scores r=0.91, P=2×10⁻¹⁶). Thus, the variants discovered through increased sample size continued to highlight specific and relevant growth-associated gene sets, while the combined analysis of both old and new loci provided the additional power needed to identify new gene sets (Table 3 and Supplementary Table 14).

The DEPICT analysis also prioritized tissues and individual genes. We found that genes within associated height loci were enriched for expression in tissues related to chondrocytes (cartilage, joint capsule, synovial membrane, and joints; $P<5.5\times10^{-9}$, FDR<0.001), and other musculoskeletal, cardiovascular, and endocrine tissue-types (FDR<0.05) (Fig. 3; Supplementary Fig. 8; Supplementary Table 15). We also showed that a subset of the 697 height associated SNPs that represented lead cis-eQTLs in blood defined 75 genes that were collectively enriched for expression in cartilage (P=0.008) (Supplementary Note and Supplementary Table 8).

We used DEPICT to prioritize 649 genes (at FDR<0.05) within height-associated loci (Table 3 and Supplementary Table 16). Of these 649 genes, 202 genes (31%) were either significant in the GRAIL analysis (Supplementary Tables 13 and 16) and/or overlapped with a list of abnormal skeletal growth syndromes that we assembled from the OMIM database¹² (n=40; Supplementary Tables 9 and 16). Many other newly prioritized genes had additional supporting evidence (Supplementary Table 16), including specific expression in the growth plate¹², and/or connections to relevant pathways (for example: *GLI2* and *LAMA5* [hedgehog signaling]; *FRS2* [FGF signaling]; *AXIN2*, *NFATC1*, *CTNNB1*, *FBXW11*, *WNT4*, *WNT5A* and *VANGL2* [WNT/beta-catenin signaling]; *SMAD3* and *MTOR* [TGF-beta and/or mTOR signaling]; *WWP2/miR140*, *IBSP*, *SHOX2* and *SP3* [required in mice for proper bone and cartilage formation]; *CHYS1*, *DSE* and *PCOLCE2* [glycosaminoglycan/collagen metabolism]; *SCARA3*, *COPZ2*, *TBX18*, *CRISPLD1* and *SLIT3* [differential expression in growth plate and predicted to be in highly relevant pathways]).

DEPICT also prioritizes genes that are new candidates for playing a role in skeletal growth. The genes newly and strongly implicated in this study included not only genes with obvious relationships to skeletal biology, such as *SOX5* and collagen genes, but also genes that have no clear published connection to skeletal growth, and likely represent as yet unknown biology (Table 3 and Supplementary Table 16). DEPICT strongly prioritized genes that do not have published annotations related to growth-related pathways but are predicted to be in gene sets that are both enriched in the associated loci and clearly connected to growth. These include genes newly predicted to be in pathways related to cartilage or bone development (*FAM101A*, *CRISPLD1* and the noncoding RNA *LINC00476*), collagen or extracellular matrix (*GLT8D2*, *CCDC3*, and *ZCCHC24*), histone demethylation (*ATAD2B* and *TSTD2*)

and other genes predicted to have skeletal phenotypes but not currently annotated as belonging to relevant pathways (*ARSJ*, *PSKH1*, *COPZ2*, *ADAMTS17* and the microRNA cluster *MIR17HG*). Of note, mutations in both *ADAMTS17* and *MIR17HG* have been identified as causes of syndromic short stature in humans ^{15,16}.

As suggested by the prioritization of *ADAMTS17* and *MIR17HG*, it is possible that some of the newly highlighted genes may also underlie new syndromes of abnormal skeletal growth. As a further proof of principle, the second entry on our list of prioritized genes (Table 3 and Supplementary Table 16), *CHSY1*, was not a known monogenic gene in the OMIM database¹² when we assembled our list, but mutations in this gene have since been shown to cause a syndrome including brachydactyly and short stature^{17,18}. Thus, the novel DEPICT method, applied to the larger GWAS data set, not only identified similar biology to GRAIL and MAGENTA but also implicated a large number of additional genes, gene sets and pathways that that are likely important in skeletal biology and human growth.

Discussion

By performing a large GWAS study on adult height, a highly heritable polygenic trait, we have provided answers to several current questions of relevance to the genetic study of polygenic diseases and traits. First, we showed that by conducting larger GWAS, we can identify SNPs that explain a substantial proportion of the heritability attributable to common variants. As hypothesized by Yang et al. (2010), the heritability directly accounted for by variants identified by GWAS and inferred by whole-genome estimation approaches are converging with increasing sample size. The variance explained by genome-wide significant SNPs has increased from 3-5% with discovery samples of ~25,000 (ref. 19) to 10% with a discovery sample size of ~130,000 (ref. 6) to 16% with a discovery sample size of 250,000 (this study), and the variance explained from all captured common SNPs is ~50% 4.5. The variance explained by genome-wide significant SNPs on a chromosome is also proportional to its length, consistent with the conclusion made by Yang et al.⁵ using all SNPs (Supplementary Fig. 9). Our new results show that ~21%, ~24% and ~29% of phenotypic variance in independent validation samples is captured by the best ~2,000, ~3,700 and ~9,500 SNPs respectively selected in the discovery samples (Table 1), and that the correlation between actual and predicted height in independent samples from the same population has increased to 0.41 (maximum prediction $R^2 = 0.41^2 = 0.17$, Fig. 2d). The results are consistent with a genetic architecture for human height that is characterized by a very large but finite number (thousands) of causal variants, located throughout the genome but clustered in both a biological and genomic manner. Such a genetic architecture may be described as pseudo-infinitesimal, and may characterize many other polygenic traits and diseases. There is also strong evidence of multiple alleles at the same locus segregating in the population and for associated loci to overlap with Mendelian forms, suggesting a large but finite genomic mutational target for height, and effect sizes ranging from minute (<1mm; ~0.01 SDs) to gigantic (>300mm; >3 SDs, in the case of monogenic mutations).

It has been argued that the biological information emerging from GWA studies will become less relevant as sample sizes increase, because as thousands of associated variants are discovered, the range of implicated genes and pathways will lose specificity and cover

essentially the entire genome²⁰. If this were the case, then increasing sample sizes would not help to prioritize follow up studies aimed at identifying and understanding new biology, and the associated loci would blanket the entire genome. Our study provides strong evidence to the contrary: the identification of many 100's and even 1000's of associated variants can continue to provide biologically relevant information. In other words, the variants identified in larger sample sizes both display a stronger enrichment of pathways clearly relevant to skeletal growth and prioritize many additional new and relevant genes. Furthermore, the associated variants are often non-randomly and tightly clustered (typically separated by <250 kb), resulting in the frequent presence of multiple associated variants in a locus. The observations that genes and especially pathways are now beginning to be implicated by multiple variants suggests that the larger set of results retain biological specificity but that at some point, a new set of associated variants will largely highlight the same genes, pathways and biological mechanisms as have already been seen. This endpoint (which we have not clearly reached for height) could be considered analogous to reaching "saturation" in model organism mutagenesis screens, where new alleles typically map to previously identified genes²¹.

We have identified a large number of gene sets and pathways that are enriched for associations with height. Although the number of gene sets and pathways is large, many are overlapping and likely represent multiple annotations of a much smaller set of core biological mechanisms. We also highlight individual genes within associated loci as being relevant to skeletal growth, including candidates for contributing to syndromes of abnormal skeletal growth; for example, we strongly implicated *CHSY1*, recently identified as an underlying cause of a monogenic syndrome with short stature and brachydactyly^{17,18}. The lists of prioritized genes and pathways should therefore provide a rich trove of data for future studies of skeletal growth; to facilitate such studies, we have made our results (including genome-wide association results and complete list of highlighted genes and pathways) publicly available. Based on the results of large genetic studies of height, we anticipate that increasing the number of associated loci for other traits and diseases could yield similarly rich lists that would generate new biological hypotheses and motivate future research into the basis of human biology and disease.

ONLINE METHODS

Genome-wide association study meta-analysis

We combined height summary association statistics from 79 genome-wide association (GWA) studies in a meta-analysis of 253,288 individuals using the same methods and studies as previously described and additional studies as described in Supplementary Tables 17-19. A total of 2,550,858 autosomal SNPs were meta-analyzed using inverse-variance fixed effects method using METAL 22 .

GCTA-COJO: conditional and joint multiple SNPs analysis

We used GCTA-COJO analysis^{7,8} to select the top associated SNPs. This method uses the summary statistics from the meta-analysis and LD correlations between SNPs estimated from a reference sample to perform a conditional association analysis⁷. The method starts

with an initial model of the SNP that shows the strongest evidence of association across the whole genome. It then implements the association analysis conditioning on the selected SNP(s) to search for the top SNPs one-by-one iteratively via a stepwise model selection procedure until no SNP has a conditional P-value that passes the significance level. Finally, all the selected SNPs are fitted jointly in the model for effect size estimation. We used 6,654 unrelated individuals from the ARIC cohort as the reference sample for LD estimation. There were ~3.0M SNPs included in the original meta-analysis. We included in this analysis only the SNPs (~2.48M) on HapMap2 and with sample size > 50,000. We used the genomewide significance level P<5×10⁻⁸ (as reported in Supplementary Table 1).

Metabochip replication

We combined height summary association statistics from 37 independent studies genotyped using Illumina's Metabochip array⁹ in a meta-analysis of 80,067 individuals of European ancestry (Supplementary Tables 20-22). Each study tested association between each genotyped SNP and the same QC procedures, height transformations, adjustment, and inheritance model as described for the GWA analysis. Genomic control correction was applied to results for each study prior to meta-analysis, using a set of 4,427 SNPs associated with QT interval to control study-specific inflation factors. We used the inverse-variance fixed effects meta-analysis method.

Validation - linear mixed model (LMM) based association analysis

Each of 15 studies (59,380 individuals) used genome-wide SNP information to calculate a genomic relationship matrix (GRM) for all pairs of individuals and used this to correct association statistics for cryptic relatedness and population stratification. Each study used a linear mixed model as implemented in the software EMMAX²³. Meta-analysis was performed as described for the standard GWAS and using a single GC correction. Each study additionally repeated the analyses for each chromosome using a GRM generated from the remaining 21 chromosomes, or in the case of the largest study (WGHS) repeating the analysis for all odd numbered chromosomes using a GRM generated from the even numbered chromosomes and vice versa. Each study then combined association results from the 22 or 2 parts of the genome into one set of data and we repeated the single GC meta-analysis.

Validation – within family (transmission) association analyses

A pure transmission based analysis was performed in seven cohorts for SNPs representing 416 signals of association (Supplementary Note), selected after repeating meta-analysis excluding these studies, with single GC correction. Filtering of low imputation quality SNPs in the studies was followed by inverse variance method of meta-analysis of the family based results. Because of the presence of related individuals, family based studies have lower power at a given sample size. For each study, we calculated the effective sample size (the size of a sample of unrelated individuals that would have the equivalent power; see Supplementary Note and Winkler *et al.* ²⁴). Estimation of winner's curse in our data set was performed by repeating the meta-analysis excluding either the family-based studies or excluding random sets of studies from GIANT matched by effective sample size to the

family based studies. Independent genome-wide significant loci were selected from each meta-analysis. Power for replication in the excluded samples was estimated at different *P*-value thresholds and the deficit in replications (number of replications expected minus number observed) was calculated. The contribution of the winner's curse to the deficit in replications was estimated as the average deficit across the three sets of random non-family-based cohorts. By subtracting this from the deficit observed for the family-based cohorts, we estimated the lack of replication that could be attributed to stratification (either inflation of effect size for true associations, or false positive associations).

Variance and heritability explained

We used GCTA-COJO analysis (Online Methods) to select the top associated SNPs at a range of stringent significance levels $(5\times10^{-3}, 5\times10^{-4}, 5\times10^{-5}, \dots, 5\times10^{-8})$ for estimation and prediction analyses. We then quantified the variance explained by those selected SNPs using a three-stage analysis, i.e. within-family prediction, GCTA-GREML analysis and population based prediction, in five validation studies (B-PROOF, FRAM, QIMR, TwinGene and WTCCC-T2D). To avoid sample overlap, we repeated the main GWAS meta-analysis and the multiple-SNP analysis five times, each time excluding one of the five validation studies. This approach ensured complete independence between data used to discover SNPs, and data used to estimate how much variance in height these SNPs explained and how well they predicted height. For the within-family prediction analyses, we selected 1,622, 2,758 and 1,597 pairs of full sibs from the QIMR, TwinGene and FRAM cohorts, respectively, with one sib pair per family. For the whole-genome estimation and prediction analyses, we used GCTA-GRM8 to estimate the genetic relatedness between individuals and selected unrelated individuals with pairwise genetic relatedness < 0.025 in each of the five studies, i.e. B-PROOF (n = 2,555), FRAM (n = 1,145), QIMR (n = 3,627), TwinGene (n = 1,145), QIMR (n = 1,145), QIMR (n = 1,145), TwinGene (n = 1,145), TwinGene (n = 1,145), TwinGene (n = 1,145), TwinGene (n = 1,145), QIMR (n = 1,145), TwinGene (n = 15,668) and WTCCC-T2D (n = 1,914).

Within-family prediction analysis

We used the SNPs selected from GCTA-COJO analysis to create a genetic predictor (also called "genetic profile score") for each of all the full sibs using PLINK²⁵. We then adjusted the genetic predictor by the first 20 principal components (PCs) generated from the principal component analysis (PCA)²⁶. By comparing the predictors within and between families, we partitioned the variance in the predictor analysis into components due to real SNP effects (V_g) , errors in estimating SNP effects (V_e) , and population structure $(C_g + C_e)$, as described in the Online Methods below.

We calculated the weighted average of each of the four (co)variance components over the three cohorts by their sample size, i.e. Σ_i ($V_{g(i)}$ n_i)/ Σ_i (n_i) with the subscript i indicating the cohort and n being the sample size. From the results of these partitioning analyses within families we can infer what the prediction R^2 (Equation 19 in Online Methods below) and what the proportion of variance explained by SNPs (i.e. V_g/V_P with V_P being the phenotypic variance) would be in a sample of unrelated individuals when using the same set of SNPs. We then tested these inferred values in unrelated samples.

GCTA-GREML analysis

We performed the GREML analysis⁴ in GCTA⁸ to estimate the variance explained by the selected SNPs (h^2_g) in each of the five validation studies. This method fits the effects of a set of SNPs simultaneously in a model as random effects and estimates the genetic variance captured by all the fitted SNPs without testing the significance of association of any single SNPs. We combined the estimates of h^2_g from the five studies by the inverse-variance

approach, i.e.
$$\Sigma_i \left(h^2_{\mathrm{g}(i)}/SE^2_i \right)/\Sigma_i \left(1/SE^2_i \right)$$
.

Population-based prediction analysis

We created a genetic predictor using the selected SNPs for the unrelated individuals in each of the five validation studies. We then calculated the squared correlation (R^2) between phenotype and predictor in each validation study, and calculated the weighted average of the prediction R^2 by the sample size across the five studies, i.e. $\Sigma_i \left(R^2_i n_i\right)/\Sigma_i \left(n_i\right)$.

Theory and method to partition the variance in a genetic predictor

Under the assumption of an additive genetic model, the phenotype of a quantitative trait can be written as

$$y=g+\varepsilon=\sum_{i}x_{i}b_{i}+\varepsilon$$
 [1]

where y is the trait phenotype, g is the total genetic effect of all SNPs, x is an indicator variable for SNP genotypes, b is the SNP effect, and ε is the residual.

From this model, the additive genetic variance is

$$var\left(g\right) = \sum_{i} var\left(x_{i}\right) b_{i}^{2} + \sum_{i} \sum_{j\left(i \neq j\right)} cov\left(x_{i}, x_{j}\right) b_{i} b_{j}$$
 [2]

with the first component being the expected value of additive genetic variance under linkage equilibrium (LE) and second component being the deviation from the expected value could be caused by linkage disequilibrium (LD), population structure or selection²⁷.

Considering a pair of full siblings in a family, the additive genetic covariance between the sibs is

$$cov\left(g_{1},g_{2}\right) = cov\left(\Sigma_{i}x_{1i}b_{i},\Sigma_{i}x_{2i}b_{i}\right) = \Sigma_{i}cov\left(x_{1i},x_{2i}\right)b_{i}^{2} + \sum_{i}\sum_{j\left(i\neq j\right)}cov\left(x_{1i},x_{2j}\right)b_{i}b_{j} \quad \text{[3]}$$

For full sibs,

$$cov(x_{1i}, x_{2i}) = 1/2var(x_i)$$
,

 $cov\left(x_{1i},x_{2j}\right)$ =1/2 $cov\left(x_{i},x_{j}\right)$ for SNPs that are in LD, and $\sum_{i}\sum_{j(i\neq j)}cov\left(x_{1i},x_{2j}\right)b_{i}b_{j}=\sum_{i}\sum_{j(i\neq j)}cov\left(x_{i},x_{j}\right)b_{i}b_{j}$ for SNPs that are not in LD (as shown by both empirical and simulation results).

Let $V_g = \sum_i var\left(x_i\right) b_i^2 + \sum_i \sum_{j(i \neq j)} cov\left(x_i, x_j\right) b_i b_j$ (SNPs are in LD), and $C_g = \sum_i \sum_{j(i \neq j)} cov\left(x_i, x_j\right) b_i b_j$ (SNPs are not in LD but correlated due to population structure)

Therefore, the genetic variance is

$$var(g) = V_q + C_q$$
 [4]

The genetic covariance between a pair of full-sibs is

$$cov(g_1, g_2) = 1/2V_q + C_q$$
 [5]

If we take a set of SNPs with their effects estimated from GCTA-COJO analysis (Online Methods), and create a predictor using these SNPs in an independent validation sample, we can write the predictor as

$$\hat{q} = \sum_{i} x_i \hat{b}_i$$
 [6]

where \hat{b} is the estimate of b with $\hat{b}=b+e$ with e being the error in estimating b.

If we assume b and e are independent and denote $V_e = \sum_i var(x_i) e_i^2$ and

$$C_e = \sum_i \sum_{j(i\neq j)} cov(x_i, x_j) e_i e_j$$
, the variance of the predictor is

$$\begin{aligned} var\left(\hat{g}\right) &= var\left(\Sigma_{i}x_{i}\hat{b}_{i}\right) = \Sigma_{i}var\left(x_{i}\right)\hat{b}_{i}^{2} + \sum_{i}\sum_{j\left(i\neq j\right)}cov\left(x_{i},x_{j}\right)\hat{b}_{i}\hat{b}_{j} \\ &= \Sigma_{i}var\left(x_{i}\right)b_{i}^{2} + \Sigma_{i}var\left(x_{i}\right)e_{i}^{2} + \sum_{i}\sum_{j\left(i\neq j\right)}cov\left(x_{i},x_{j}\right)b_{i}b_{j} + \sum_{i}\sum_{j\left(i\neq j\right)}cov\left(x_{i},x_{j}\right)e_{i}e_{j} \end{aligned} \ \ [7]$$

$$&= V_{g} + V_{e} + C_{g} + C_{e}$$

The covariance between the predictors of a pair of full-sibs is

$$cov(\hat{g}_{1}, \hat{g}_{2}) = cov\left(\sum_{i} x_{1i} \hat{b}_{i}, \sum_{i} x_{2i} \hat{b}_{i}\right) = 1/2 \sum_{i} var(x_{i}) \hat{b}_{i}^{2} + \sum_{i} \sum_{j(i \neq j)} cov(x_{1i}, x_{2j}) \hat{b}_{i} \hat{b}_{j}$$

$$= 1/2 V_{g} + 1/2 V_{e} + C_{g} + C_{e}$$
[8]

The covariance between the true phenotype and the predictor of a same individual is

$$cov(y,\hat{g}) = cov(g+\varepsilon,g+e) = var(g) = V_g + C_g$$
 [9]

The covariance between the true phenotype of one sib and the predictor of the other sib is

$$\begin{array}{ll} cov\left(y_{1},\hat{g}_{2}\right) = & cov\left(g_{1}+\varepsilon_{1},g_{2}+e_{2}\right) \\ & = & 1/2\Sigma_{i}var\left(x_{i}\right)b_{i}^{2}+\sum_{i}\sum_{j\left(i\neq j\right)}cov\left(x_{i},x_{j}\right)b_{i}b_{j}=1/2V_{g}+C_{g} \end{array} \tag{10}$$

If we define $\Delta \hat{g} = \hat{g}_1 - \hat{g}_2$ and $y = y_1 - y_2$,

$$var(\Delta \hat{g}) = var(\hat{g}_1) + var(\hat{g}_2) - 2cov(\hat{g}_1, \hat{g}_2) = V_q + V_{e}$$
 [11]

$$cov(\Delta y, \Delta \hat{g}) = cov(y_1, \hat{g}_1) + cov(y_2, \hat{g}_2) - cov(y_1, \hat{g}_2) - cov(y_2, \hat{g}_1) = V_q$$
 [12]

We therefore can calculate these four parameters as

$$V_{\rm g} = cov(\Delta y, \Delta \hat{g})$$
 [13]

$$V_{\rm e} = var\left(\Delta \hat{g}\right) - V_{\rm g}$$
 [14]

$$C_{\rm g} = cov(y, \hat{g}) - V_{\rm g}$$
 [15]

$$C_{\rm e} = 2cov(\hat{g}_1, \hat{g}_2) - var(\hat{g}) - C_{\rm g}$$
 [16]

where $V_{\rm g}$ can be interpreted as the variance explained by real SNP effects, $C_{\rm g}$ is the covariance between predictors attributed to the real effects of SNPs that are not in LD but correlated due to population stratification, $V_{\rm e}$ is the accumulated variance due to the errors in estimating SNP effects, and $C_{\rm e}$ is the covariance between predictors attributed to errors in estimating the effects of SNPs that are correlated due to population stratification.

To assess the prediction accuracy, we usually perform a regression analysis of the real phenotype against the predictor, i.e.

$$y=\beta_0+\hat{g}\beta_1+\varepsilon$$
 [17]

so that the regression slope is actually

$$\beta = cov(y, \hat{g}) / var(\hat{g}) = (V_g + C_g) / (V_g + V_e + C_g + C_e)$$
 [18]

with the regression R^2 being

$$R^2 = (V_g + C_g)^2 / (V_g + V_e + C_g + C_e)$$
 [19]

In the absence of population structure,

$$R^2 = V_g^2 / (V_g + V_e)$$
 [20]

Variance explained by SNPs in proximity to the top associated SNPs

We performed analyses to quantify the variance explained by SNPs in close physical proximity to the top associated SNPs in 9,500 unrelated individuals (pairwise genetic relatedness < 0.025) from a combined dataset of the QIMR and TwinGene cohorts. As in previous analyses, to avoid sample overlap between discovery and validation studies, we repeated the discovery meta-analysis excluding the QIMR and TwinGene cohorts, and

identified 643 genome-wide significant SNPs from the GCTA-COJO analysis of the summary statistics using ARIC data for LD estimation. We used GCTA-GREML analysis 4,8 to quantify the phenotypic variance explained by all the common SNPs (MAF > 0.01) within 100Kb, 500Kb or 1Mb of the 643 genome-wide significant SNPs. We show in Supplementary Figure 6a that there are 104K, 423K and 745K SNPs within 100Kb, 500Kb and 1Mb of the top associated SNPs, which explain 20.8% (s.e. = 1.3%), 25.7% (s.e. = 1.8%) and 29.5% (s.e. = 2.2%) of phenotypic variance, respectively. We then applied a regression-based approach 28 to adjust for LD between SNPs. The estimates of variance explained after LD-adjustment were slightly higher than those without adjustment, and the ratio of between the estimates with and without LD-adjustment was consistently ~1.05 regardless of the window size (Supplementary Fig. 6a). However, the difference is small.

We then sought to investigate whether or not there is an enrichment of additional association signals at the top associated loci. We varied the window size from 20Kb to 50Kb, 100Kb, 150Kb, 200Kb, 300Kb, 400Kb, 500Kb, 750Kb and 1Mb, and fitted a two-component model in GCTA-GREML analysis, with the first component being the top associated SNPs and the second component being the rest of SNPs within the window. We found that the per-SNP variance explained excluding the top SNPs (variance explained by the second component divided by the number of SNPs included in this component) decreased with the size of window (Supplementary Fig. 6b), implying that SNPs in closer physical proximity to the top associated SNPs tend to explain disproportionally more variance.

Enrichment of associated SNPs in ENCODE regions, loci containing OMIM genes, eQTLs and nsSNPs

To identify putative causal variants among the height-associated markers, we explored whether the height-associated SNPs were in strong LD ($r^2>0.8$) with non-synonymous coding variants in 1000 Genomes Project CEU Phase 1 data, showed an effect on whole blood gene expression levels, were located within ENCODE-annotated regions, were within loci harboring monogenic growth genes, or had previously been associated with other complex traits in NHGRI GWAS catalog ($P<5\times10^{-8}$) (Supplementary Tables 7-11). To estimate the empirical assessment of enrichment for listed features we used 10,000 permutations of random sets of SNPs matched to the pruned (LD $r^2>0.1$) 628 height-associated SNPs by the number of nearby genes (within a distance of LD $r^2>0.5$), physical distance to nearest gene, and minor allele frequency.

Enrichment of genes in associated loci in known and novel pathways

Data-Driven Expression-Prioritized Integration for Complex Traits (DEPICT) analysis—The DEPICT method (T.H.P. *et al.*, unpublished data; see Geller *et al.*²⁹ for an earlier application of DEPICT) relies on pre-computed predictions of gene function based on a heterogeneous panel of 77,840 expression arrays (Fehrmann *et al.*, manuscript in review; ref. 30), 5,984 molecular pathways (based on 169,810 high-confidence experimentally derived protein-protein interactions³¹), 2,473 phenotypic gene sets (based on 211,882 genephenotype pairs from the Mouse Genetics Initiative (see URLs)), 737 Reactome pathways³², 5,083 Gene Ontology terms¹⁴, and 184 KEGG pathways³³. The method leverages these predictions to extend the functional annotations of genes, including genes that previously

had only a few or no functional annotations. DEPICT facilitates the analysis of GWAS data by (1) assessing whether genes in associated loci are enriched in tissue-specific expression, (2) identifying reconstituted gene sets that are enriched in genes from associated loci, and (3) systematically identifying the most likely causal gene(s) at a given locus (see Supplementary Note for a more detailed description of DEPICT). In order to run DEPICT, we first clumped the summary statistics from the meta-analysis using 500kb flanking regions, $r^2>0.1$, and excluded SNPs with P 5×10^{-8} , which resulted in 628 SNPs. We then mapped genes to each of the 628 best-associated SNPs. For a given SNP, this was accomplished by including all genes that resided within LD $r^2 > 0.5$ boundaries of that SNP, and always including the nearest gene, to its locus gene set. We used a locus definition that was calibrated using the GWAS data for height levels presented in this paper and optimized capture of known monogenic genes for those traits. We merged overlapping loci, and excluded loci that mapped near or within the major histocompatibility complex locus (chromosome 6, location: 20 to 40 Mb), which resulted in a list of 566 non-overlapping loci that were used as input to DEPICT. HapMap Project Phase II CEU genotype data was used for all LD calculations.

GRAIL and MAGENTA analysis—The GRAIL¹⁴ algorithm was run using the LD pruned ($r^2>0.1$) 628 SNPs without correcting for gene size, and using text-mining data up to December 2006 (default setting). MAGENTA¹³ was run with the single genomic control adjusted summary statistics as input using default settings and excluding the HLA region.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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New GWAS studies

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Metabochip Studies

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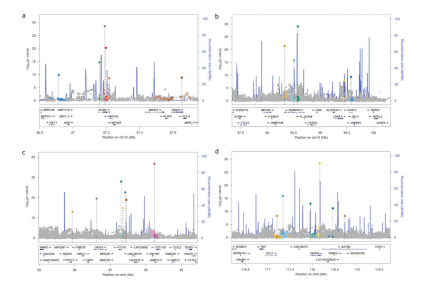


Figure 1. Regional association plots for loci with multiple association signals Panels $\bf a$ to $\bf d$ highlight examples of multiple signals after approximate conditional joint multiple-SNP analysis GCTA-COJO analysis. SNPs are shaded and shaped based on the index SNP with which they are in strongest LD ($r^2>0.4$). Panels $\bf a-c$ show the majority of signals clustering in and around a single gene (ACAN, ADAMTS17, PTCH1, respectively) whereas panel $\bf d$ shows the multiple signals clustering through proximity.

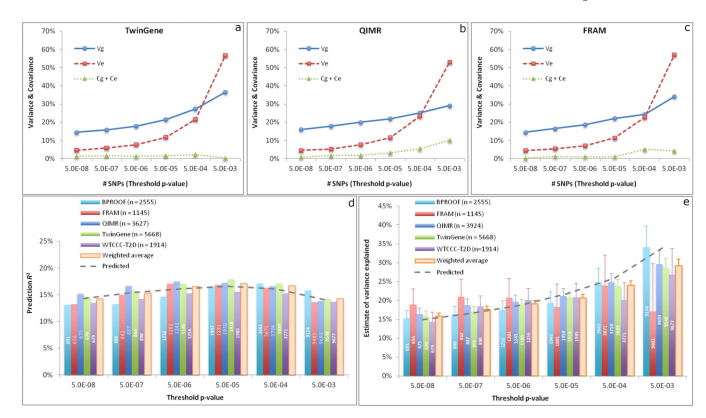


Figure 2. Quantifying the variance explained by height associated SNPs at different levels of significance

The SNPs were selected from the approximate conditional and joint multiple SNPs association analysis using GCTA-COJO analysis with the target cohort being excluded from the meta-analysis. a, b, c, Partitioning the variance in the SNP-derived genetic predictor using a within-family analysis. The SNP-based predictor was adjusted by the first 20 PCs. The four variance-covariance components V_g , V_e , C_g and C_e are defined in Online Methods. d, Accuracy of predicting phenotype by the genetic predictor in unrelated individuals. The prediction R^2 shown on the y-axis is the squared correlation between phenotype and predictor. The SNP-based predictor was adjusted by the first 20 PCs. The solid line is the average prediction R^2 weighted by sample size over the five cohorts. The dashed line is the prediction accuracy inferred from the within-family prediction analysis (Equation 19 in Online Methods). e, The variance explained by the SNPs was estimated by the wholegenome estimation method in GCTA. The phenotype was adjusted by the first 20 PCs. Each error bar represents the standard error of the estimate. The estimates from all the five cohorts (B-PROOF, FRAM, QIMR, TwinGene and WTCCC-T2D) were averaged by the inversevariance approach. The dashed line is the variance explained inferred from the within-family prediction analysis. In panels d and e, the number shown in each column is the number of SNPs used in the analysis.

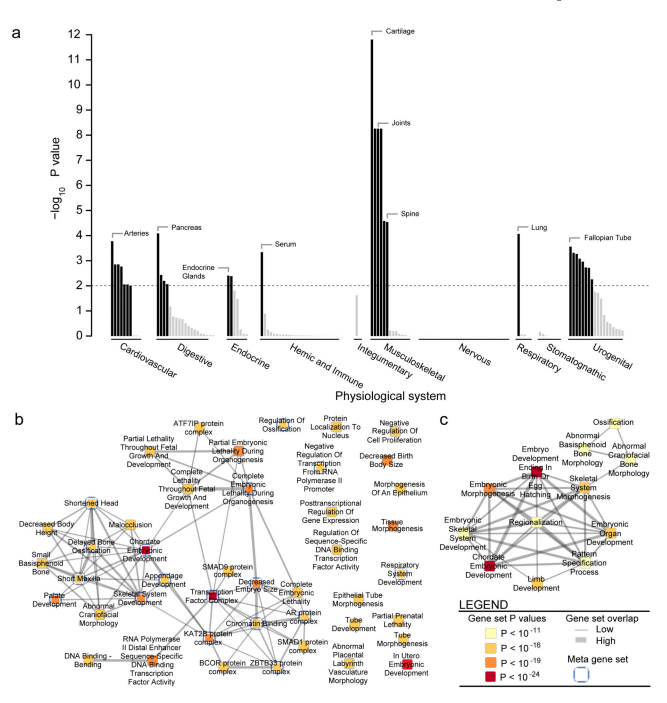


Figure 3. Tissue enrichment combined with pruned gene set network

Genes within genome-wide significant height associated loci enriched for several relevant tissue annotations as well as gene sets. **a**, Genes in associated loci tended to be highly expressed in tissues related to chondrocytes and osteoblasts (cartilage, joints, and spine), and other musculoskeletal, cardiovascular and endocrine tissue-types. The analysis was conducted based on the DEPICT method and 37,427 human microarray samples. Tissue annotations are sorted by physiological system and significance. Significantly enriched (FDR<0.05) tissues are color-coded in black. **b**, Significantly enriched reconstituted gene

sets (P-value< 1×10^{-11} , FDR< 1×10^{-5}) identified by DEPICT. Nodes represent reconstituted gene sets and are color-coded by statistical significance. Edge thickness between nodes is proportional to degree of gene overlap as measured by the *Jaccard* index. Nodes with gene overlap greater than 25% were collapsed into single meta nodes and marked by blue borders. \mathbf{c} , reconstituted gene sets comprised by the Chordate Embryonic Development meta node, which represented several gene sets relevant to human height (e.g. ossification, embryonic skeletal system development, and limb development).

Table 1 Estimates of variance explained by SNPs selected at different significance levels

The SNPs were selected by an approximate conditional and joint multiple-SNP analysis (GCTA-COJO) of the summary statistics from the meta-analysis. The target cohort for variance estimation was excluded from the meta-analysis.

Threehold	QIM	QIMR $(n = 3,924)$	924)	FRA	FRAM $(n = 1,145)$	145)	TwinGe	TwinGene $(n = 5,668)$	(899)	WTCCC	WTCCC-T2D $(n = 1,914)$: 1,914)	B-PRO	$\textbf{B-PROOF}\;(\mathbf{n}=2,555)$	2,555)	^a Weighted average	average	b Pred.
	#SNP	#SNP h^2 g SE	SE	#SNP	h^2 g	SE	#SNP	h^2 g	SE	#SNP	h^2 g	SE	#SNP	h^2 g	SE	h^{2} g	SE	h^{2} g
5E-8	675	0.164	0.164 0.016	959	0.190	0.040	029	0.159	0.013	629	0.143	0.025	691	0.152	0.021	0.159	0.008	0.149
SE-7	887	887 0.187 0.017	0.017	862	0.210	0.045	998	0.170	0.013	890	0.184	0.028	988	0.162	0.022	0.176	0.009	0.166
SE-6	1245	0.196	0.018	1202	0.207	0.050	1186	0.188	0.014	1256	0.201	0.030	1232	0.175	0.024	0.190	0.000	0.186
SE-5	1950	0.212	0.020	1891	0.183	0.060	1918	0.208	0.015	1985	0.208	0.037	1947	0.194	0.029	0.206	0.010	0.218
SE-4	3754	0.248	0.024	3671	0.239	0.080	3689	0.239	0.017	3771	0.201	0.047	3661	0.248	0.037	0.240	0.013	0.259
5E-3	6693	0.297	0.035	9403	0.171	0.126	9548	0.287	0.025	2196	0.267	0.070	9174	0.341	0.055	0.292	0.018	0.339
CHIM3	1.08M	1.08M 0.473	0.086	1.06M	0.313	0.291	1.12M	0.522	090.0	0.97M	0.534	0.170	1.09M	0.463	0.126	0.498	0.044	

 a The estimates from all the five cohorts were averaged by the inverse-variance approach i.e. $\Sigma_{i}\left(h^{2}_{\mathrm{g}(i)}/SE^{2}_{i}
ight)/\Sigma_{i}\left(1/SE^{2}_{i}
ight)$

b the predicted variance explained by the selected SNPs ($V_{\rm g}$) from the within-family prediction analysis;

 c SNPs from HapMap3 project 11 .

Table 2 Comparison of prioritized variants, loci, biology and variance explained from GWASs on human stature with 130,000 individuals (previously published in Lango Allen $\it et~al.$, 2010) and with 250,000 individuals (this paper).

	Height GWAS with 130,000 samples (Lango Allen <i>et al.</i> , Yang et al)*	Height GWAS with 253,288 samples
SNP based comparisons		
GWAS significant SNPS	199	697
Genomic loci [#] (+/- 1Mb)	180	423
Loci [#] with multiple signals	19	147
Secondary associations in loci [#]	19	273
Biological annotation (DEPICT at FDR < 0.05)		
Prioritized genes	92	649
Loci with prioritized gene	74 (43%)	422 (75%)
Pruned gene sets and protein-protein complexes %	813	2,330
Tissues and cell-types	5	43
Variance explained		
GWAS significant SNPs	10%	16%
Deep list of SNPs at 1×10^{-3}	13%	29%
All common SNPs	45% **	50%
Heritability explained		
GWAS significant SNPs	12.5%	20%
Deep list of SNPs at 1×10^{-3}	16%	36%
All common SNPs	56% **	62.5%

^{*} Counts, numbers and estimates for Lango Allen *et al.* are taken from respective publication.

 $^{^{\#}}$ Genomic loci defined by distance: +/- 1Mb from index height SNP

[&]amp; Genomic loci defined by LD: $r^2 > 0.5$ from index height SNP

[%] After clumping of similar gene sets and pathways

^{**} Yang et al. *Nat Genet* 42, 565-9 (2010).

Table 3 Significantly prioritized novel human growth associated genes

The table lists 20 genes prioritized by DEPICT. Genes are ranked by the number of lines of supporting evidence and the DEPICT *P*-value (Supplementary Table 16). Because 20 of the 30 top-ranked genes were in a curated list of genes known to cause syndromes of skeletal¹², these "OMIM genes" are not shown here. The top fifteen genes with prior literature support (based on GRAIL) are shown, followed by the top five novel genes. Each gene is accompanied by the significantly enriched reconstituted gene sets in which it appears in (DEPICT gene set enrichment analysis). Abbreviations; (*GO* – Gene Ontology; *MP* – Mice Phenotypes from Mouse Genome Informatics database; *InWeb* – protein-protein interaction complexes; *KEGG* and *REACTOME* databases).

Locus (height SNP)	Gene symbol	New locus	Prioritization <i>P</i> -value	Levels of biological annotation	Top ranking reconstituted gene sets
		Genes	with prior literature supp	ort (GRAIL)	
rs10748128	FRS2	N	1.0×10 ⁻¹⁶	7	PI 3K cascade (<i>REACTOME</i> , $P=6.2\times10^{-13}$); Chronic Myeloid Leukemia (<i>KEGG</i> , $P=1.6\times10^{-12}$); Response To Fibroblast Growth Factor Stimulus (<i>GO</i> , $P=5.4\times10^{-11}$);
rs2166898	GLI2	Y	4.4×10 ⁻¹⁶	7	Growth Factor Binding (GO , $P=2.6\times10^{-14}$); Regulation Of Osteoblast Differentiation (GO , $P=2.3\times10^{-11}$); WNT-Protein Binding (GO , $P=1.9\times10^{-12}$)
rs526896-rs9327705	TBX4	N	9.9×10 ⁻⁹	7	Short Mandible (MP , $P=3.3\times10^{-19}$); Respiratory System Development (GO , $P=3.1\times10^{-17}$); Abnormal Ulna Morphology (MP , $P=1.9\times10^{-15}$)
rs16860216	SOX8	N	0.016	7	Small Thoracic Cage (<i>MP</i> , P =6.9×10 ⁻¹⁴); Short Ribs (<i>MP</i> , P =2.7×10 ⁻⁸); Short Sternum (<i>MP</i> , P =6.5×10 ⁻⁷)
rs1199734	LATS2	Y	1.0×10 ⁻¹⁶	6	Partial Lethality Throughout Fetal Growth And Development (MP , $P=1.2\times10^{-18}$); Growth Factor Binding (GO , $P=2.6\times10^{-14}$); TGFB1 protein complex ($InWeb$, $P=6.3\times10^{-12}$)
rs12323101	PDS5B	N	1.0×10 ⁻¹⁶	6	Chromatin Binding (<i>GO</i> , <i>P</i> =6.4×10 ⁻¹⁷); Nuclear Hormone Receptor Binding (<i>GO</i> , <i>P</i> =2.4×10 ⁻¹²); RBBP4 protein complex (<i>InWeb</i> , <i>P</i> =1.3×10 ⁻¹¹); WNT16 protein complex (<i>InWeb</i> , <i>P</i> =1.9×10 ⁻⁸)
rs6746356	SP3	Y	1.0×10 ⁻¹⁶	6	BCOR protein complex (InWeb, $P=2.7 \times 10^{-17}$); AFF2 protein complex (InWeb, $P=4.5 \times 10^{-7}$); Intracellular Steroid Hormone Receptor Signaling Pathway (GO, $P=9.0 \times 10^{-6}$)
rs3923086	AXIN2	Y	2.2×10 ⁻¹⁶	6	Signaling By Transforming Growth Factor Beta ($KEGG$, $P=3.8\times10^{-15}$); WNT Receptor Signaling Pathway (GO , $P=6.9\times10^{-14}$); Polydactyly (MP , $P=1.5\times10^{-10}$)
rs3790086	LTBP1	N	1.3×10 ⁻¹³	6	Abnormal Skeleton Morphology (MP , $P=1.1\times10^{-15}$); TGF Beta Signaling Pathway ($KEGG$, $P=3.8\times10^{-15}$); Growth Factor Binding (GO , $P=2.6\times10^{-14}$)

Locus (height SNP)	Gene symbol	New locus	Prioritization P-value	Levels of biological annotation	Top ranking reconstituted gene sets
rs2034172	WNT5A	Y	4.3×10 ⁻¹³	6	Partial Lethality Throughout Fetal Growth And Development (MP, P=1.2×10 ⁻¹⁸); Tissue Morphogenesis (GO, P=4.1×10 ⁻²⁰); Abnormal Skeleton Morphology (MP, P=1.1×10 ⁻¹⁵)
rs3915129	CTNNB1	Y	3.5×10 ⁻¹²	6	AR protein complex (<i>InWeb</i> , $P=8.9\times10^{-17}$); TCEB1 protein complex (<i>InWeb</i> , $P=1.5\times10^{-11}$); GTF21 protein complex (<i>InWeb</i> , $P=4.6\times10^{-11}$)
rs12330322	BMP2	N	5.6×10 ⁻¹⁰	6	Transcription Factor Binding (GO , $P=4.7\times10^{-26}$); Complete Embryonic Lethality During Organogenesis (MP , $P=4.9\times10^{-21}$); Short Mandible (MP , $P=3.3\times10^{-19}$)
rs10958476-rs6999671	ВМР6	N	2.9×10 ⁻⁸	6	Small Basisphenoid Bone (MP , $P=8.9\times10^{-17}$); TGF Beta Signaling Pathway ($KEGG$, $P=3.8\times10^{-15}$); Growth Factor Binding (GO , $P=2.6\times10^{-14}$)
rs564914	SOX5	Y	4.6×10 ⁻⁷	6	Disproportionate Dwarf (<i>MP</i> , <i>P</i> =1.8×10 ⁻¹³); Abnormal Cartilage Morphology (<i>MP</i> , <i>P</i> =1.9×10 ⁻¹³); Short Limbs (<i>MP</i> , <i>P</i> =2.8×10 ⁻¹³)
rs17807185	WNT4	Y	4.6×10 ⁻⁷	6	Morphogenesis Of An Epithelium (GO , $P=2.3\times10^{-17}$); Gland Development (GO , $P=5.4\times10^{-16}$); Basal Cell Carcinoma ($KEGG$, $P=1.5\times10^{-12}$)
		No	ovel genes without prior e	vidence	
rs8042424	CHSYI	N	1.0×10 ⁻¹⁶	7	Abnormal Cartilage Morphology (<i>MP</i> , P=1.9×10 ⁻¹³); Abnormal Bone Ossification (<i>MP</i> , P=2.1×10 ⁻¹²); Signaling by Transforming Growth Factor Beta (<i>REACTOME</i> , P=5.9×10 ⁻⁹)
rs7652177	FNDC3B	N	1.0×10 ⁻¹⁶	5	Abnormal Spongiotrophoblast Layer Morphology (MP , P =3.2×10 ⁻¹⁶); Decreased Length Of Long Bones (MP , P =2.7×10 ⁻¹²); ITGB1 protein complex ($InWeb$, P =5.2×10 ⁻⁸)
rs7284476	TRIOBP	Y	1.0×10 ⁻¹⁶	5	Negative Regulation Of Cell Proliferation (GO , $P=4.3\times10^{-17}$); Abnormal Vitelline Vasculature Morphology (MP , $P=1.7\times10^{-15}$); Beta-Catenin Binding (GO , $P=3.0\times10^{-5}$)
rs2149163-rs3927536	BNC2	N	1.0×10 ⁻¹⁶	5	Short Ulna (MP , P =4.7×10 ⁻¹³); Abnormal Joint Morphology (MP , P =8.6×10 ⁻¹¹); Regulation Of Chondrocyte Differentiation (GO , P =2.9×10 ⁻⁹)
rs3790086	WWP2	Y	1.0×10 ⁻¹⁶	5	Cartilage Development (<i>GO</i> , <i>P</i> =2.0×10 ⁻¹⁹); Chondrocyte Differentiation (<i>GO</i> , <i>P</i> =3.0×10 ⁻¹⁵); Signaling By Platelet-Derived Growth Factor (<i>REACTOME</i> , <i>P</i> =4.8×10 ⁻¹⁰)