

Genetic architecture of early childhood growth phenotypes gives insights into their link with later obesity

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Abstract

Early childhood growth patterns are associated with adult metabolic health, but the underlying mechanisms are unclear. We performed genome-wide meta-analyses and follow-up in up to 22,769 European children for six early growth phenotypes derived from longitudinal data: peak height and weight velocities, age and body mass index (BMI) at adiposity peak (AP ~9 months) and rebound (AR ~5-6 years). We identified four associated loci ($P < 5 \times 10^{-8}$): *LEPR/LEPROT* with BMI at AP, *FTO* and *TFAP2B* with Age at AR and *GNPDA2* with BMI at AR. The observed AR-associated SNPs at *FTO*, *TFAP2B* and *GNPDA2* represent known adult BMI-associated variants. The common BMI at AP associated variant at *LEPR/LEPROT* was not associated with adult BMI but was associated with *LEPROT* gene expression levels, especially in subcutaneous fat ($P < 2 \times 10^{-51}$). We identify strong positive genetic correlations between early growth and later adiposity traits, and analysis of the full discovery stage results for Age at AR revealed enrichment for insulin-like growth factor 1 (IGF-1) signaling and apolipoprotein pathways. This genome-wide association study suggests mechanistic links between early childhood growth and adiposity in later childhood and adulthood, highlighting these early growth phenotypes as potential targets for the prevention of obesity.

Introduction

Complex developmental processes regulate changes in body mass and growth in early life. Multiple genetic variants are robustly associated with adult body mass index (BMI)¹ and metabolic phenotypes²⁻⁴ but much fewer data exist for childhood traits^{5,6}. Discrete genetic variants may regulate developmental patterns during rapid growth in infancy compared to later growth periods in childhood⁷ as demonstrated by the age-dependent impact of variation at the *FTO* locus on BMI⁸ but there still remains a paucity of relevant data. On the other hand, robust observational associations between early growth traits and adult cardiometabolic risk⁹⁻¹² may, in part, be explained by shared genetic factors. In support of this hypothesis, studies from the Early Growth Genetics (EGG) consortium show that some of the genetic component contributing to size at birth also impacts type 2 diabetes, cardiovascular disease and blood pressure¹³. However, the extent of the genetic correlation between postnatal growth patterns and the regulation of adult body weight, energy metabolism and later metabolic health is unclear.

In infancy and childhood, individuals follow well-characterized and predictable height, weight and BMI trajectories (**Figure 1**)¹⁴. For example, BMI trajectory encompasses three periods characterized by i) a rapid increase in BMI up to the age of 9 months (adiposity peak, AP); ii) a rapid decline in BMI up to the age of 5-6 years (adiposity rebound, AR); and iii) a steady increase in BMI after 5-6 years until early adulthood¹⁰. Here, we set out to model sex-specific individual weight, height and BMI curves in children, using unique data collected from primary health care or clinical research visits, to extract six early growth traits: peak height velocity (PHV; cm/month), peak weight velocity (PWV; kg/month), age at adiposity peak (Age-AP; year), BMI at adiposity peak (BMI-AP, kg/m²), age at adiposity rebound (Age-AR; year), BMI at adiposity rebound (BMI-AR, kg/m²) (**Figure 1**) and to test for genetic associations using genome-wide genotype data. Although a few previous studies have investigated the genetic basis of these well-established measures of longitudinal growth trajectory^{7,9,15,16}, no systematic genome-wide analyses have been carried out to date. Understanding the genetic architecture of early childhood growth phenotypes will provide insight into their regulation and into the mechanisms underlying the observed associations with adult cardiometabolic disease, which will be fundamental for early promotion of metabolic health.

Results

Genome-wide association meta-analyses of early growth phenotypes

We conducted genome-wide association study (GWAS) meta-analyses to identify genetic loci influencing each of the six early growth traits in up to 7,215 term-born children of European ancestry from up to four population-based studies that had both genetic and early growth trait data (stage 1; **Online Methods, Supplementary Table 1, Supplementary Figure 1**). From the stage 1 inverse variance meta-analyses, we selected a total of 8 loci with either $P < 1 \times 10^{-7}$ or with $P < 1 \times 10^{-5}$ in/near genes known to be associated with obesity and metabolic traits in published GWAS or candidate gene studies (**Supplementary Table 2**), and sought confirmation in up to 16,550 term-born children from up to 11 additional studies (stage 2; **Online Methods, Supplementary Table 3, Supplementary Figure 1**). Our study design is summarized in **Figure 2**, while participant characteristics, genotyping arrays, imputation and quality control for the discovery and follow-up studies are summarized in **Supplementary Tables 1 and 3**. In combined meta-analyses of the discovery and follow-up studies (including up to 22,769 children), we identified common variants at four independent loci, each associated with any of the early growth phenotypes, at genome-wide levels of significance (i.e. $P < 5 \times 10^{-8}$) (**Table 1, Figure 3, Supplementary Figure 2**). The lead signals at each of the four loci were: rs9436303 at the *LEPR/LEPROT* (encoding the leptin receptor and an overlapping transcript) locus associated with BMI at AP, rs1421085 at the *FTO* (encoding a 2-oxoglutarate-dependent demethylase) and rs2817419 at the *TFAP2B* (encoding transcription factor AP-2 beta) loci associated with Age at AR, and rs10938397 near the *GNPDA2* (encoding adiposity regulating glucosamine-6-phosphate deaminase) locus associated with BMI at AR. The lead signals at each of the four loci were located within non-coding regions of the relevant genes except for rs10938397, which was located in an intergenic region proximal to the *GNPDA2* gene (**Figure 3**). The common AR-associated SNPs at *FTO*, *TFAP2B* and *GNPDA2* have previously been associated with other anthropometric measures, especially in adulthood¹. However, the three loci *LEPR/LEPROT*, *TFAP2B* and *GNPDA2* described here are novel with respect to their association with early growth phenotypes, while the association between SNPs at *FTO* and Age at AR in the current study replicates the findings of a previous gene-centric study⁸ at genome-wide levels of significance.

At the *LEPR/LEPROT* locus, the minor G allele of rs9436303 was associated with higher BMI at AP, 0.07SD per allele (95% CI= 0.04, 0.09), $P_{\text{combined}} = 5.1 \times 10^{-9}$, equivalent to a difference of 0.07 kg/m² (95% CI= 0.05, 0.09) per effect allele. The same allele is also associated with earlier age at menarche ($P=0.00019$ in the ReproGen consortium's publicly available age at menarche GWAS data)¹⁷ and with increased plasma soluble leptin receptor levels ($P=1.19 \times 10^{-9}$)¹⁸ but the SNP was not associated with sex combined adult BMI ($P=0.96$ in the GIANT consortium's publicly available BMI GWAS data)¹ amidst a marginal association with BMI in females of ≤ 50 years of age ($P=0.029$) in GIANT. Furthermore, this SNP is in low LD with other common variants at the *LEPR/LEPROT* locus that are also associated with severe early-onset obesity (rs11208659, $r^2=0.008$; rs1137100, $r^2=0.128$)^{19,20}; plasma soluble leptin receptor levels (rs1137100, $r^2=0.128$; rs1137101, $r^2=0.151$)¹⁸ and age at menarche (rs10789181, $r^2=0.006$)¹⁷. It is also in low LD with the CRP-associated variant identified in our own previous study (rs12753193, $r^2=0.015$)²¹ (**Supplementary Table 4**). Using data from the Northern Finland Birth Cohort (NFBC) 1966 study (N=2585) we also show that the previously published genome-wide significant SNPs

associated with other traits in the *LEPR/LEPROT* locus are not associated with BMI at AP, and that our index SNP rs9436303 remains associated with BMI at AP ($P < 1 \times 10^{-6}$) after conditioning on those other trait-associated SNPs (**Online Methods and Supplementary Table 5**). Therefore, to our knowledge, rs9436303 represents a novel, distinct, signal for BMI at AP at this locus.

At the *FTO* locus, the C allele of the index SNP rs1421085 was associated with earlier Age at AR, -0.12 SD per allele (-1.44 months, 95% CI= -1.42, -1.46), $P_{\text{combined}} = 3.12 \times 10^{-30}$, and is directionally consistent with a previous gene-centric study⁸. The *FTO* locus has the largest effect on BMI and obesity risk reported in GWAS carried out to date, and the same allele that is associated with earlier Age at AR is associated with higher childhood and adult BMI²². Close proxies for this index SNP have also been associated with adult BMI²³, type 2 diabetes²⁴, metabolic syndrome²⁵, waist circumference²⁶ and extreme obesity²⁷ (**Supplementary Table 4**).

At the *TFAP2B* locus each additional copy of the A allele of our index SNP rs2817419 was associated with earlier Age at AR, -0.08 SD per allele (~-0.96 months, 95% CI= -0.94, -0.98), $P_{\text{combined}} = 4.4 \times 10^{-11}$, and the same allele is robustly associated with higher adult BMI, 0.03 SD per allele (95% CI= 0.02, 0.04), $P = 3.7 \times 10^{-15}$ in the GIANT consortium's BMI GWAS data¹. The G allele of the adult BMI^{1,28} and waist circumference²⁹ associated index SNP rs2207139 in *TFAP2B* from GIANT results in earlier Age at AR in our data, -0.09 per SD (~-1.1 months, 95% CI: -1.12, 1.08), $P_{\text{discovery}} = 0.0002$ (**Supplementary Table 6**). However, rs2207139 is in very low LD ($r^2 = 0.035$) with our index SNP rs2817491, and these are likely to represent two distinct signals at this locus. In support of this, conditional analyses in the NFBC1986 data adjusting the effect of rs2817419 on Age at AR for rs2207139 and *vice versa* showed that the two SNPs have independent effects on Age at AR (-0.07 SD (95% CI= -0.13, -0.01), $P = 0.03$ and -0.10 SD (95% CI= -0.17, -0.03), $P = 0.007$ respectively) (**Online Methods**).

Overall, the identified associations at *FTO* and *TFAP2B* are directionally consistent with observational associations between the timing of AR and adult BMI⁹ (i.e. reaching AR at an earlier age is associated with higher BMI whereas reaching AR later is associated with lower BMI).

At the *GNPDA2* locus, the G allele of the index SNP rs10938397 was associated with higher BMI at AR, 0.06 SD per allele (0.06 kg/m², 95% CI= 0.04, 0.07), $P_{\text{combined}} = 2.9 \times 10^{-8}$, with the same allele being associated with higher adult BMI in previous reports³⁰.

There was no heterogeneity between the studies, except at *LEPR/LEPROT* (**Supplementary Figure 3**). For example, in the fixed effect meta-analysis of the discovery stage results for the BMI at AP associated SNP rs9436303 in *LEPR/LEPROT* locus showed heterogeneity (**Supplementary Figure 3a**) and random effect meta-analysis did not change the heterogeneity (**Supplementary Figure 3b**). However, excluding the smaller LISA-D study (N=282) that showed directionally inconsistent effects to the other studies, from the discovery stage meta-analysis for this SNP reduced the heterogeneity but did not change the point estimates (**Supplementary Figure 3c**). Similarly, there was heterogeneity in the combined stage fixed meta-analysis for the same SNP in *LEPR/LEPROT* (**Figure 3d**), and random effect meta-analysis

did not change the heterogeneity (**Figure 3e**). Fixed effect meta-analysis excluding the EDEN and NFBC1966 studies that showed inflated results showed no heterogeneity, and the point estimates were similar with and without the two studies (**Supplementary Figure 3f & 3g**). All four SNPs together explain 0.1 %, 0.7% and 0.3 % variance in BMI at AP, Age at AR and BMI at AR, respectively.

Prioritizing candidate genes and pathways in the four early growth trait associated loci for functional characterization

We first considered the genes that harbor, or are nearest to, the index SNP as a potential candidate. The known biological functions and molecular mechanisms of the four proteins encoded by these are given in **Supplementary Table 7**. However, as the four genome-wide significant signals are located in regions harboring multiple genes with dense linkage disequilibrium (LD) structure, we are unable to locate with certainty the causal gene or gene variant. Therefore, the gene nearest to the lead signals may not necessarily be the gene influenced by the underlying causal variant/s. For example, at the *FTO* locus more than one gene, which includes the *FTO*^{31,32}, *RPGRIP1L*³³ and *IRX3/IRX5* in the *IRXB* gene cluster^{34,35} have been implicated as the candidates for mediating the observed associations with adult adiposity. Given our index SNP at *FTO* associated with Age at AR is highly correlated with the BMI associated SNPs at the same locus, it is plausible that the same genes play a role in the regulation of Age at AR and adult adiposity.

To identify other potential candidate genes and provide further insights into the molecular mechanisms underlying the four genome-wide significant loci associated with BMI at AP, BMI at AR or Age at AR, we searched for association of the index SNP with *cis*-acting expression quantitative trait loci (eQTLs) in five different living tissues: liver, skin, whole blood and subcutaneous and omental fat, and repeated these lookups in 44 post-mortem tissues from the Genotype-Tissue Expression (GTEx) consortium³⁶ (**Online Methods**), which mostly reproduced the results of the eQTL association in living tissues. For two out of the four loci analysed, the index SNP was strongly associated (at $P < 0.001$) with the expression of one or more nearby (+/-1 Mb) genes (**Supplementary Table 8**). For example, the BMI at AP associated SNP rs9436303 in the *LEPR/LEPROT* locus was associated with gene expression levels in all five tissues: the strongest association was with *LEPROT* ($P=1.5 \times 10^{-51}$) gene expression levels in subcutaneous fat. The consistency of the results for this locus across five different tissues and in four different datasets (MuTHER, KORA, deCODE and Kaplan) suggests that these findings are robust and imply that, at least for this locus, the underlying causal variant functions through gene expression. Both the *LEPR* and *LEPROT* genes are good biological candidates for having a role in the regulation of BMI at AP in infancy because the *LEPR* gene encodes the receptor for leptin, an adipocyte-specific hormone that plays a major role in the regulation of appetite and energy balance, reproduction, growth and the immune system^{37,38}. Variation in leptin concentrations is also associated with adult body mass³⁸ and rare mutations in the *LEPR* gene cause monogenic obesity³⁹. The *LEPROT* gene negatively regulates *LEPR* cell-surface expression⁴⁰ and growth hormone (GH) receptor expression in the liver⁴¹, thereby decreasing hepatic responses to leptin and growth hormone. This is consistent with the finding by Wu *et al*⁴² in mice showing that *LEPROT* could be central to the nutritional regulation of growth and growth hormone binding in the liver and chondrocytes. This evidence further suggests a role for *LEPROT* in the hepatic synthesis of the insulin-like growth factor 1 (IGF-1)⁴². The BMI at AR associated index SNP rs10938397 in *GNPDA2* was associated

with *GUF1* (*GUF1* Homolog, GTPase) transcription in subcutaneous fat ($P=7.10 \times 10^{-4}$); a gene implicated in mitochondrial protein synthesis (**Supplementary Table 8**).

To explore the biological pathways and networks underlying early growth, we applied a gene set enrichment analysis (MAGENTA)⁴³ to the discovery stage GWAS results (**Online Methods**). We identified enrichment of two gene sets (**Supplementary Tables 9 and 10**), suggesting that genes in Age at AR associated regions are involved in the IGF-1 signaling and apolipoprotein pathways (false discovery rate (*FDR*) < 0.05, $P < 1.6 \times 10^{-3}$) (**Supplementary Figure 6**). The apolipoprotein pathway has an important role in lipid transport and metabolism, while the IGF-1 signaling pathway has a well-established role in neonatal and pubertal growth^{44,45} and in the regulation of energy metabolism through the activation of PI3K/AKT pathway via either the insulin receptor or the IGF-1 receptor⁴⁶. The present finding supporting the overexpression of the IGF-1 pathway with an earlier Age at AR is consistent with the IGF-1 and the high-protein hypothesis exemplified by the work of Rolland-Cachera and colleagues in pediatric nutrition (for a recent review see Rolland-Cachera *et al.* 2016)⁴⁷. This hypothesis suggests that the risk of childhood obesity associated with high-protein intake during lactation could be caused by stimulation of the IGF-1/GH pathway leading to an early age at adiposity rebound. It is therefore possible that higher IGF-1 levels, via genetic and/or nutritional factors, might reduce GH levels and expression via a negative feedback^{48,49}. Subsequent, lower circulating levels of GH might also suppress lipolysis and contribute to fat accumulation^{50,51}, potentially affecting normal BMI trajectories and Age at AR, and thereby risk of adult obesity and metabolic disorders.

In a look up of the index SNP, and the top 10 intragenic SNPs associated with Age at AR (at $P < 0.05$) in genes from the IGF-1 and apolipoprotein pathways (**Supplementary Note 1 and Supplementary Table 11**) in cis eQTL data (**Online Methods**), a total of 29 eQTLs were associated with 16 different transcripts (all with $P < 1 \times 10^{-3}$). The most significant association was observed between *APOL4* RNA abundance in subcutaneous adipose tissue and rs132700 near the same gene in the apolipoprotein pathway ($P = 1.02 \times 10^{-11}$) (**Supplementary Table 12**). One SNP within *PTK2* of the IGF-1 signaling pathway was strongly associated with omental and subcutaneous adipose tissue expression of Argonaute 2 (*EIF2C2*), a nearby gene encoding a protein involved in regulation of microRNA transcription. The SNPs within *HDLBP*, *APOB* and *APOL4* of the apolipoprotein pathway and SNPs in *IGF1R*, *PTK2*, *GRB10* in the IGF-1 signaling pathway are all likely to affect DNase activity and chromatin state and expression, while SNPs within *HDLBP*, *APOB* and *APOL4* are also likely to affect transcription factor binding according to RegulomeDB⁵² and Haploreg⁵³.

Association of the adult BMI associated loci with early growth traits

In a look-up analysis of the 97 adult BMI associated loci from the GIANT consortium¹ in our stage 1 GWAS meta-analysis data, we observed that an excess of adult BMI increasing alleles was associated ($P < 0.05$) with increased BMI at AP (12/97 associations, $P_{\text{binomial}} = 0.003$), BMI at AR (19/97 associations, $P_{\text{binomial}} = 3.05 \times 10^{-7}$) and earlier Age at AR (26/97 associations, $P_{\text{binomial}} = 1.27 \times 10^{-12}$) (**Supplementary table 6**). These were directionally consistent with the reported observational associations between the same early growth traits and adult BMI (**Supplementary Table 13**). We also

created weighted genetic risk scores using the 97 adult BMI associated SNPs from the GIANT consortium¹ in the two NFBC studies (**Online Methods**). When comparing the bottom 20% of the children carrying a maximum of ~84 weighted BMI increasing alleles (i.e. bottom quintile) to the top 20% of the children carrying a maximum of ~111 weighted BMI increasing alleles (i.e. top quintile), there was a mean difference of 0.02 SD (95% CI= 0.01, 0.03) in PWV; 0.16 SD (95% CI= 0.07, 0.24) in Age at AP; 0.14 SD (95% CI= 0.05, 0.22) in BMI at AP; -0.64 SD (95% CI= -0.72, -0.56) in Age at AR and 0.55 SD (95% CI= 0.47, 0.63) in BMI at AR (**Supplementary Figure 4**). Furthermore, we observed correlations (at $P < 0.0001$) between the effects of the 97 adult BMI associated SNPs, and Age at AR and BMI at AR (**Figure 4**) but no correlations were observed with the effects on the other early growth phenotypes (**Supplementary Figure 5**). Taken together our results suggest that the variants involved in the regulation of adult BMI have effects that begin in early childhood, around the age at AR.

Genetic link between early growth and health outcomes

To further explore the genetic links and, thereby, help prioritize the potential causal relationships between early growth and adverse health outcomes, we estimated their genetic correlation using LD score regression (**Online Methods**). In contrast to observational associations, genetic correlations between complex traits can be useful in prioritizing observational associations for subsequent causal analyses, as genetic factors are less likely to be confounded and are not altered by the outcome. Here we summarize the key genetic correlations at 1% FDR, and full results are presented in **Figure 5** and **Supplementary Table 14**. We uncovered several genetic correlations that were directionally consistent with the reported observational associations between the same traits (**Supplementary Note 2** and **Supplementary table 13**).

In these analyses PHV was not genetically correlated with any of the traits tested at 1% FDR. Conversely, PWV showed a strong positive genetic correlation with childhood obesity ($r_g = 0.53$, $P = 7.79 \times 10^{-8}$) and was positively correlated with birth length ($r_g = 0.50$, $P = 0.0003$), adult height ($r_g = 0.21$, $P = 0.0009$) and waist circumference ($r_g = 0.22$, $P = 0.0008$). We found very little evidence that PWV was genetically correlated with cardiometabolic traits. The shared genetic contribution of Age at AP with other traits could not be quantified due to low heritability estimates.

BMI at AP also showed positive genetic correlations with childhood obesity ($r_g = 0.65$, $P = 1.19 \times 10^{-8}$), adult BMI ($r_g = 0.26$, $P = 0.0002$), adult overweight ($r_g = 0.26$, $P = 0.0007$) and adult low-risk obesity class 1 (BMI 30.0-34.9kg/m²) ($r_g = 0.32$, $P = 0.0001$).

Age at AR showed a strong positive genetic correlation with age at menarche ($r_g = 0.44$, $P = 1.90 \times 10^{-9}$), and strong inverse genetic correlations with childhood obesity ($r_g = -0.83$, $P = 1.50 \times 10^{-14}$), fasting insulin levels ($r_g = -0.58$, $P = 8.69 \times 10^{-6}$) and several adult anthropometric traits including adult BMI ($r_g = -0.72$, $P = 3.10 \times 10^{-18}$) and waist circumference ($r_g = -0.62$, $P = 8.40 \times 10^{-12}$).

In contrast to Age at AR, BMI at AR revealed an inverse genetic correlation with age at menarche ($r_g = -0.38$, $P = 4.48 \times 10^{-9}$) and positive correlations with childhood obesity ($r_g = 1.00$, $P = 6.51 \times 10^{-25}$), adult BMI ($r_g = 0.64$, $P = 1.60 \times 10^{-15}$) and waist circumference ($r_g = 0.48$, $P = 6.07 \times 10^{-10}$). Together the genetic correlations of Age at AR and BMI at AR with age at

menarche and BMI are directionally consistent given that earlier age at menarche predicts higher adult BMI⁵⁴ and both are associated with adverse cardiometabolic traits in observational studies⁵⁵⁻⁵⁷.

In summary, the results from LD score regression analysis define a robust link between the genetics of early growth and the genetics of later BMI and childhood /adult obesity. In contrast, the LD score regression analysis revealed relatively little evidence for a shared genetic link between early growth traits and lipid metabolism, blood pressure, type 2 diabetes or cardiovascular disease.

Discussion

We report the first GWAS of six early growth phenotypes derived from longitudinal data. In samples of up to 22,769 term-born children of European descent, we exploited a wealth of repeated weight and height data collected from birth until 13 years. These more refined phenotypes better capture childhood growth patterns relevant to later disease risk than single growth measures in childhood such as BMI and height. The four common variants identified here show consistent effects across several studies despite varying study designs, sample collection methods, and containing data obtained from children born at different geographical regions in Europe demonstrating that our results are robust, and are not affected by any differences between studies. One exception to this is the common variant at *LEPR/LEPROT* where we observed some evidence for heterogeneity. However, excluding the studies that showed inflated results from the meta-analyses reduced or showed no heterogeneity, and did not change the point estimates. Our findings contribute to the understanding of genetic influences on early growth in humans, and provide insights into the underlying molecular mechanisms that link early growth phenotypes with increased risk of obesity in later childhood and adulthood. An exception to this is the common variant in the *LEPR/LEPROT* locus associated with BMI at AP that was not associated with later obesity in childhood or adulthood, although other common variants in this gene are associated with childhood obesity and several metabolic traits, and rare mutations in the same gene cause early onset morbid obesity (OMIM ID: 614963)^{39,58}. This raises questions about how the yet-to-be-determined causal variant at *LEPR/LEPROT*, or the leptin/free leptin surge in infancy, impact on early development and health. It is noteworthy that research in animals has identified an early peak in leptin concentration, which is thought to be essential not only in regulating energy balance but also in brain development⁵⁹.

Our study has limitations that should be taken in to consideration for future genetic studies of early growth traits. **First**, dense longitudinal growth data meshed with GWAS data are only available in a few cohorts worldwide, so we had limited power to detect genetic variants with smaller effects and/or low allele frequencies. This affected our ability to identify robust associations of some of the promising variants identified in the discovery stage, in particular at *PCSK1* (Table 1). Variants at this locus, which are known to be associated with severe obesity⁶⁰, showed a suggestive association with BMI at AP in our discovery GWAS, but failed to replicate and were not included in further analyses. However, in our discovery stage GWAS for BMI at AP among ~6,222 infants, we had 85% power to detect a variant with an effect size of 0.14 SD at $P < 5 \times 10^{-8}$ with minor allele frequency of 0.22. **Second**, it is noteworthy that these derived growth phenotypes are likely to be influenced by a degree of measurement error, especially in cohorts with fewer repeated measures around the time points being estimated, which in turn could have further hampered our power to detect true genetic associations. Despite this, we were still able to discover genetic variants showing robust associations with these derived growth phenotypes. **Third**, in the current GWAS we inferred genotypes based on HapMap Phase 2 imputation; imputation using more comprehensive reference panels can provide greater coverage of the genome and can aid in improving power to detect additional genetic variants. **Fourth**, the present results may not be directly applicable to other ethnic groups as growth patterns and disease risk vary by ethnicity. Trans-ethnic studies may help to identify

additional genetic risk variants due to differences in allele frequencies among different ethnic groups. For example, the type 2 diabetes-associated locus *KCNQ1*, which has an effect across multiple ethnic groups, was first discovered in an East Asian GWAS, due to the allele frequency difference between the East Asian and European populations⁶¹. **Finally**, we did not identify any variants associated with PHV, PWV and Age at AP at genome-wide levels of significance, and this may be due to a combination of smaller genetic effects on growth at this stage of development, reduced statistical power due to smaller sample size or because environmental factors are more influential than genetic influences at this age. The interplay with infant feeding and other environmental factors also warrants additional research.

In conclusion, we have identified three novel loci associated with BMI at AP in infancy at around 9 months, and age and BMI at AR at around 5-6 years, and confirmed the previously reported association of *FTO* with Age at AR⁸, at genome-wide significance levels. The genetic architecture of early growth shares a robust similarity with that of later BMI. For example, the three AR associated loci identified here are strongly associated with adult BMI (**Figure 6**), and the cross-trait genetic correlations based on GWAS data showed the same relationship. The relationship of AR to later adiposity may be due to shared genetic mechanisms or Age at AR and BMI at AR might be on the causal pathway for later adiposity. To test this hypothesis, large independent study populations are required, in which causal analysis methods such as Mendelian randomization can be undertaken using the variants associated with these phenotypes as proxies for early growth phenotypes. However, such studies would currently be underpowered as the variants discovered here only explain a small proportion of the variance in the early growth traits. Taken together, our results suggest that adult obesity has its origins in early childhood, and interventions aiming to promote an optimal growth trajectory in infancy and early childhood could contribute to the prevention of obesity.

Methods

Longitudinal growth modeling and derivation of early growth phenotypes

Early growth phenotypes were derived from sex-specific individual growth curves using mixed effects models of height, weight and BMI measurements from birth to 13 years (**Figure 1**). All height and weight data were collected prospectively via either self-reported data or clinical measurements (**Supplementary Table 1 and 2**). These phenotypes were derived separately in each cohort (**Supplementary Note 4**).

Derivation of peak height velocity (PHV) and peak weight velocity (PWV) - The methods for growth modeling and derivation of growth parameters from the fitted curves is described in detail in a previous publication⁷ Parametric Reed1 growth model was fitted in sex stratified non-linear random-effect model as described previously⁶². Term-born singletons (defined as ≥ 37 completed weeks of gestation) with at least three height or weight measurements from birth to 24 months of age were included in the Reed1 model fitting. Maximum-likelihood method for best fitting curves for each individual was used to estimate the growth parameter, PHV (cm/months) and PWV (kg/months).

Derivation of age and BMI at adiposity peak (AP) and adiposity rebound (AR) - The methods used for growth modeling of age and BMI has been previously described in detail by Sovio et al, 2011⁸. Due to the specificity of longitudinal changes in BMI *i.e.* succession of peak and nadir as described in figure 1, the data was divided into two age windows for modeling i) growth in infancy using height and weight data from 2 weeks to 18 months of age and ii) growth in childhood using growth and weight data from 18 months to 13 years of age. Each cohort contributed most data available within any of these two age windows. In studies where the data available consisted of both height and weight data within a given window, then the data point nearest to the mid time points of that window were used as a proxy for the BMI measurement. Prior to model fitting, age was centered using the median age of the relevant age window. For example, in the infant growth model at 0-1.5 years, the median age was 0.75 years (which was close to the average age at AP), and in the childhood growth model at >1.5-13 years, the median age was 7.25 years (on average shortly after AR). Linear Mixed Effects (LME) models were then fitted for log-transformed BMI. We used sex and its interaction with age as covariates, with random effects for intercepts *i.e.* baseline BMI, and linear slope *i.e.* linear change in BMI over time. In addition to linear age effect, quadratic and cubic terms for age were included in the fixed effects to account for nonlinearity of BMI change over time.

Growth in Infancy - The following model was used to calculate the age and BMI at adiposity peak (AP), and the analysis was restricted to singletons with BMI measures from two weeks to 18 months of age. The model is as follows:

$$\log(\text{BMI}) = \beta_0 + \beta_1 \text{Age} + \beta_2 \text{Age}^2 + \beta_3 \text{Age}^3 + \beta_4 \text{Sex} + u_0 + u_1(\text{Age}) + \varepsilon$$

where BMI is expressed in kg/m² and age in year. $\beta_0, \beta_1, \beta_2, \beta_3, \beta_4$ are the fixed effects terms, u_0 and u_1 are the individual level random effects and ε is the residual error. The age at AP was calculated from the model as the age at maximum BMI between 0.25 and 1.25 year according to preliminary research^{7,11}.

Growth in Childhood. The model used to measure the age and BMI at adiposity rebound (AR) in childhood is as follows:

$$\log(\text{BMI}) = \beta_0 + \beta_1 \text{Age} + \beta_2 \text{Age}^2 + \beta_3 \text{Age}^3 + \beta_4 \text{Sex} + \beta_5 \text{Age} * \text{Sex} + \beta_6 \text{Age}^2 * \text{Sex} + u_0 + u_1(\text{Age}) + \varepsilon$$

Where BMI is expressed in kg/m^2 and age in year. $\beta_0, \beta_1, \beta_2, \beta_3, \beta_4, \beta_5$ and β_6 are the fixed effects, u_0 and u_1 are the individual level random effects and ϵ is the residual error. Age at AR was calculated as the age at minimum BMI between 2.5 and 8.5 year according to preliminary research^{7,11}.

Stage 1 genome-wide association studies, genotyping and imputation

Stage 1 genome-wide association analyses included up to 7215 children of European descent from five studies (up to four studies for each early growth trait) that had growth data and genome-wide data. These included the Helsinki Birth Cohort Study (HBCS, Finland), Northern Finland Birth Cohort 1966 (NFBC1966, Finland), Lifestyle-Immune System-Allergy Study (LISA, Germany), The Western Australian Pregnancy Cohort Study (Raine, Australia) and Generation R (Netherlands) (**Figure 2**). Informed consent was obtained from all study participants (or parental consent, as appropriate) and the local ethics committees as appropriate approved all study protocols. Study characteristics, genotyping platform, imputation and association test software used, as well as sample and genotyping and imputation quality control steps in each stage 1 study are given in **Supplementary Table 1**. The Stage 1 consisted of a GWAS based on ~2.5 million directly genotyped or imputed SNPs. Imputation of non-genotyped SNPs was undertaken either with MACH or with IMPUTE and were imputed to HapMap Phase 11 CEU reference panel after excluding genotyped SNPs with a minor allele frequency (MAF) < 1%, call rate of at least $\geq 95\%$, and a Hardy-Weinberg Equilibrium (HWE) P -value cut off as given in **Supplementary Table 1**.

Stage 1 genome-wide association analyses and meta-analyses

According to the availability of dense enough data for growth modeling, a total of up to 7215, 6222, 6219 and 6051 children were used to analyse PHV/PWV, Age-AP, BMI-AP and Age-AR /BMI-AR respectively (**Figure 2**). We only included children who were born between 37 and 41 completed weeks of gestation (i.e.: term born) from singleton pregnancies and children who had more than three growth measurements available within the age range in question. Gestational age was either defined from the date of the last menstrual period or ultrasound scans depending on the study. All six early growth traits except for Age-AP and Age-AR were natural log transformed to reduce skewness, and all traits were converted to z-scores prior to association testing to facilitate the comparison of results across the studies. We tested the directly genotyped and imputed variants for association with each of the six early growth traits in a linear regression model assuming an additive genetic effect. The regression models were adjusted for sex and principal components (PC) derived from the genome-wide data to control for potential population substructure (the necessary number of principal components included varied by study). The regression models were also adjusted for gestational age, except for Age-AR and BMI-AR. The genome-wide association analyses (i.e. stage 1) were performed using either SNPTTEST or MACH2QTL in each cohort, and data exchange facilities were provided by the AIMS server⁶³. All stage 1 study beta estimates and their standard errors were meta-analysed using the inverse-variance fixed effects method in the METAL software⁶⁴. SNPs with poor imputation quality (e.g. $r^2 < 0.3$ for MACH and 'proper_info' score < 0.4 for IMPUTE) and/or a HWE $P < 1 \times 10^{-4}$

were excluded prior to the meta-analyses. Double genomic control⁶⁵ was applied: firstly, to adjust the statistics generated within each cohort and secondly, to adjust the overall meta-analysis statistics. Results are reported as a change in standard deviation (SD) units per effect allele as reported in **Table 1**.

Selection of SNPs for stage 2 follow up.

All loci reaching $P < 1 \times 10^{-7}$ from stage 1 GWAS of each early growth trait were selected for follow-up in stage 2. These included the two SNPs associated with Age-AR in the *FTO* locus (rs1421085) and in the intergenic region between *RANBP3L* and *SLC1A3* (rs2956578), and the SNP associated with BMI-AP in *LEPR/LEPROT* (rs9436303). Four further SNPs (one SNP associated with BMI-AP near *PCSK1* (rs10515235), one SNP associated with Age-AR in *TFAP2B* (rs2817419), and two SNPs associated with BMI-AR near *GNPDA2* (rs10938397) and in *DLG2* (rs2055816)) were selected for follow-up on the basis of showing an association with an early growth trait at $P < 1 \times 10^{-5}$ and being in/near genes with established links to adiposity and metabolic phenotypes except for *DLG2*, a possible candidate gene involved in glucose metabolism⁶⁶. In addition, one locus with a plausible association ($P = 5.91 \times 10^{-5}$) with PWV, near *TMEM18* (rs2860323), was also selected for follow-up based on previous reports showing an association with severe early onset obesity¹⁹ and its association with BMI in adulthood²⁹ and childhood⁶ (**Supplementary Table 3**). No loci for PHV or AGE-AP passed the p-value threshold or other selection criteria used for follow up. **Supplementary Table 3** shows the SNP selection criteria and proxies used in more detail.

Stage 2 follow up of lead SNPs in single SNP association analyses

For follow up of lead signals selected from stage 1 we used data from up to 16,550 children of European descent from 12 additional population based studies (up to 11 studies for each early growth trait), namely the Avon Longitudinal Study of Parents and Children (ALSPAC, United Kingdom), Cambridge Baby Growth Study (CBGS, United Kingdom), Children's Hospital of Philadelphia (CHOP, United States of America), Copenhagen Prospective Study on Children (COPSAC, Denmark), Danish National Birth Cohort (DNBC, Denmark), Étude des Déterminants pré- et postnatals du développement et de la santé de l'Enfant (EDEN, France), The Exeter Family Study of Childhood Health (EFSOCH, United Kingdom), Infancia y Medio Ambiente Project (INMA, Spain), Lifestyle-Immune System–Allergy Study (LISA (R), Germany), Northern Finland Birth Cohort Study 1986 (NFBC1986, Finland), The Physical Activity and Nutrition in Children (PANIC, Finland) and Southampton Women's Survey (SWS, United Kingdom). We used directly genotyped or imputed data for the eight SNPs (or proxies of $r^2 > 0.8$) selected from stage 1 and tested their association in a total of 5367, 16550, 12256, 12192 children of European ancestry with PWV, BMI-AP, Age-AR, BMI-AR respectively (**Figure 2**). Direct genotyping was performed in some follow-up studies by KBiosciences Ltd. (Hoddesdon, UK) using their own novel system of fluorescence-based competitive allele-specific PCR (KASPar). The call rates for all genotyped SNPs were >95%. Study characteristics, genotyping platform, imputation and association test software used, as well as sample and genotyping and imputation quality control steps in each stage 1 study are given in **Supplementary Table 2**. We used the same methods as in stage 1 for sample selection, genotyping quality control, association testing and meta-analysis.

Combined analysis of stage 1 and stage 2 samples. All stage 1 and 2 results were meta-analysed using the inverse-variance fixed effects method in either METAL⁶⁴ or R (version 3.2.0; <http://www.r-project.org/>). In these combined analyses, loci reaching $P < 5 \times 10^{-8}$ were considered as genome-wide significant and loci reaching $P < 5 \times 10^{-6}$ were considered as a suggestive association. Heterogeneity between studies was tested by Cochran's Q tests and the proportion of variance due to heterogeneity was assessed using I^2 index for each individual SNP at each stage.

Estimation of genetic variance explained

The variance explained (VarExp) by each SNP was calculated using the effect allele frequency (f) and beta (β) from the meta-analyses using the formula $\text{VarExp} = \beta^2 (1 - f)2f$.

Analysis of the association of the index SNP associated with early growth phenotypes in other GWAS data sets

We looked up the index SNP or a proxy associated with each early growth trait in publicly available published meta-analysis data sets to assess their associations with other GWAS traits using the Pheno Scanner available at <http://www.phenoscanter.medschl.cam.ac.uk/phenoscanter>⁶⁷. We used a P -value cut off of $< 5 \times 10^{-8}$ for displaying the association results and r^2 cut off of > 0.6 for proxy SNP lookups from 1000G.

Conditional analyses

To test whether the index SNP (rs9436303) in *LEPR/LEPROT* locus associated with BMI at AP have an effect on BMI at AP independent of the other trait associated SNPs in the same locus we first, tested the individual association of our index SNP and each of the five previously published genome-wide significant SNPs associated with obesity and metabolic traits in the *LEPR/LEPROT* locus with BMI at AP using a linear regression model adjusted for sex and gestational age in 3459 children from the NFBC1966 study using the same exclusion criteria described above. We next tested whether our index SNP has an independent effect on BMI at AP, by adjusting the linear regression model between the index SNP and BMI at AP for each of the five previously published genome-wide significant SNPs, sex and gestational age.

To test whether our index SNP (rs2817419) and the adult BMI and waist circumference associated index SNP (rs2207139) at the *TFAP2B* locus in GIANT have independent effects on Age at AR we tested the association between rs2817419 and Age at AR in a linear regression model adjusting for rs2207139 and sex and *vice versa* in the NFBC1986 study.

Expression quantitative locus (e-QTL) analysis.

To study the molecular mechanisms underlying the significant genetic variants associated with growth patterns, we searched for cis eQTL using results obtained on liver, skin, whole blood and subcutaneous and omental fat living tissue made available by the MuTHER⁶⁸, KORA⁶⁹, DeCode⁷⁰ and Lee Kaplan⁷¹ studies. Additional lookups were conducted in 44

post-mortem tissues from GTEx consortium³⁶. The association analysis were performed with the index SNP. The analysis of eQTL was limited to genes in cis within a +/- 1Mb window of the index SNP

Pathway enrichment analysis

To explore the pathways associated with early growth phenotypes, we applied a Meta-Analysis Gene-set Enrichment of variaNT Associations (MAGENTA (version 2))⁴³ to the stage 1 GWAS results. Briefly, each gene in the genome is mapped to a single SNP with the lowest *P*-value within a 110 Kb upstream or 40kb downstream window of the gene. The corresponding *P*-value, representing each gene, is corrected for confounding factors such as gene size, LD patterns, SNP density and other genetic factors. The adjusted *P*-values are ranked and the observed number of genes in a given pathway above a specified *P*-value threshold (75th and 95th percentiles used) is calculated. This number is compared with that from repeating the process based on 10000 randomly permuted pathways of identical size. In doing so, an empirical gene set enrichment association (GSEA) *P*-value for each pathway is computed. In our study, individual pathways with a FDR < 0.05 and nominal GSEA *P* < 0.05 were deemed significant, and, unless otherwise stated, results for the 95th percentile cut-off analysis are reported. To test whether SNPs in LD with the index SNP were driving the enrichment of the significant pathways the top 10 intragenic SNPs associated with the relevant early growth phenotype (*P*<0.05) within genes of the significant MAGENTA pathways were selected by the regression models (**Supplementary Note 1**). These were then examined for association with cis eQTL data in the liver, skin, subcutaneous and omental fat dataset as described above, and a proxy (*r*²>0.5) was used if the top SNP was not available in the expression study. In addition, using all top intragenic SNPs and the index SNPs associated with the significant MAGENTA pathways, we searched the regulomeDB⁵² and Haploreg⁵³ database for relevant functional data, including: coding variation, regulatory chromatin marks, DNaseI hypersensitivity, protein binding and motif alteration data.

The association of BMI genetic risk score with early growth traits

We calculated the weighted genetic risk scores in children from NFBC1966 and NFBC1986 studies separately using the 97 SNPs associated with BMI at genome-wide levels of significance in the GIANT consortium in studies of up to 339,224 individuals¹. A proxy SNP (*r*²>0.8) was used if the index SNP associated with BMI was not available in the NFBC studies. Only 95/97 were used in the NFBC1986 cohort as no suitable proxies were available for two SNPs (rs6477694 and rs2112347). We used the same exclusion criteria applied in the GWAS analyses described above and calculated the weighted score of the 95/97 SNPs using the formula 1 below to account for varying effect sizes of the SNPs. The SNPs were recoded to reflect the number /dosage of the BMI increasing alleles for that SNP. The weight (*W*) for each locus is the effect size of the BMI increasing allele of each SNP from Locke *et al*¹.

$$\text{Weighted score} = W_1 \times \text{SNP}_1 + W_2 \times \text{SNP}_2 + \dots \cdot W_n \times \text{SNP}_n \quad (1)$$

The weighted genetic risk score was rescaled to reflect the number of BMI- increasing alleles using formula 2 as described in Lin *et al*⁷².

$$\text{Genetic risk score} = \frac{\text{Weighted score} \times \text{Number of available SNPs}}{\text{Sum of weights of the available SNPs}} \quad (2)$$

We then created quintiles of weighted genetic risk score in each study containing approximately 20% children in each of the five groups, and tested the mean difference in each early growth phenotype between the lowest and the highest quintile of the weighted genetic risk score in a linear regression model. In the linear regression model we used the quintiles of weighted genetic risk score as the independent variable, and each early growth trait as the dependent variable adjusted for sex and gestational age except for Age-AR and BMI-AR where the regression model was only adjusted for sex. The genetic risk score analyses were carried out in StataMP 13 for Windows (StataCorp, Brownsville, TX). The results from NFBC1966 and NFBC1986 cohort was meta-analysed using the fixed effect inverse variance estimator implemented in the Stata command, “metan”.

Linkage-disequilibrium (LD) score regression analyses

The use of LD score regression in estimating the genetic correlation between two traits have been described previously^{73,74}. Briefly the LD score regression uses GWAS meta-analysis summary statistics of several million SNPs of the two traits under comparison (here the GWAS meta-analysis summary statistics of each early growth trait and the other trait under comparison) and calculates the cross product of the test statistics at each SNP. This cross product of the test statistics is then regressed against the LD score of each SNP (i.e.: sum of the LD r^2 between a variant and all the variants in a 1 cm region in the genome) and the slope of this regression line gives the genetic correlation between the two traits of interests. We primarily used the LD hub⁷⁵ available at <http://ldsc.broadinstitute.org> to quantify the genetic correlation between each of the five early growth traits, and health related outcomes. LD hub is a centralised database of summary-level GWAS results from 36 GWAS consortia, and provides a web interface to automate multiple LD score regression analyses in a single run. The analyses can be conducted on selected phenotypes of interest or can be carried out on all traits available on the database in hypothesis free tests. We first reformatted GWAS summary statistics for each of the five early growth traits according to the sample input format provided on the developer’s website prior to uploading on their server. The diseases/traits of interest for LD score regression analyses were selected from the pre compiled list of GWAS available on LD hub where summary statistics are available in the required format for running LD score regression. The LD hub uses pre-calculated LD scores based on European samples. We selected a total of 49 disease/traits of interest from 33 GWAS studies in the following pre-compiled categories: education, anthropometric traits, lipids, glycaemic traits, bone mineral density, neurological / psychiatric diseases and other traits (including adiponectin, CAD, T2D, menarche) and submitted the request for LD score regression for each of the five early growth traits at a time. Following each analysis a genetic correlation matrix between the early growth trait and the selected

disease/traits were returned which included the genetic correlation value (r_g) its standard error and the corresponding P -value for each trait comparison.

However, LD hub currently does not harbour GWAS summary statistic data for lung function measures or blood pressure traits. To quantify the genetic correlation between early growth phenotypes and these measures we obtained GWAS summary statistics data from the SpiroMeta consortium for the lung function measures of FEV₁, FVC, and FEV₁/FVC ratio⁷⁶. To generate genetic association statistics for SBP and DBP we carried out GWAS of systolic and diastolic blood pressure (SBP and DBP) using 125,334 subjects from the UK Biobank study⁷⁷ (**Supplementary Note 3**). For these LD score regression analyses we used the Python scripts provided on the developer's website at <https://github.com/bulik/ldsc>. Prior to running the LD score regression analyses each summary statistics file was reformatted using the `munge_sumstats.py` Python script which filtered the SNPs to HAPMAP 3 SNPs as recommended on the developer's website to minimise any bias from poor imputation quality. SNPs were also excluded if MAF<0.01, ambiguous strand, duplicate rsID and reported sample size is less than 60% of the total available. If the sample size for each SNP was available we used the `-N-col` to specify the relevant sample size column in the GWAS summary statistics file, and when no sample size column was available we used the maximum sample size reported in the GWAS meta-analysis. After the GWAS summary statistics files were reformatted we then used the `ldsc.py` Python script to run the LD score regression analyses between each of the six early growth traits, blood pressure and lung function measures. The pre-computed European LD scores calculated from 1000G data available on the developer's website was used for LD score regression.

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AUTHOR CONTRIBUTIONS

All authors contributed in reviewing the paper. **Supplementary Table 15** presents the contribution of each individual author. The names of individuals who contributed to specific statistical analysis and drafting the paper are given below.

Writing and discussion group on analyses specific to the project: N.M.G.D.S., S.S., S.D., A.D.S.C.A., U.S., M.K., I.J.M., J.B., R.C., H. R.T., A.M.L., A.R., A.B., V.W.V.J., J.F., P.O., K.O., S.G., M.I.M and M.-R.J.

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LD score regression, BMI SNP look-ups and genetic risk score analyses: N.M.G.D.S

GWAS of blood pressure in the UKBiobank: J.R.P.

COMPETEING FINANCIAL INTERESTS: The authors declare no competing interest

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Table 1. Summary statistics of the eight independent SNPs associated with Peak Weight Velocity in infancy (PWV), BMI at Adiposity Peak in infancy (BMI-AP), Age at Adiposity Rebound (Age-AR) and BMI at Adiposity Rebound (BMI-AR) in discovery (stage 1), follow-up (stage 2) and in combined meta-analyses.

Index SNP	Chromosome Position ^a	In/near gene	Effect allele/Other allele	Effect allele frequency	Stage 1 (n=7215)		Stage 2 (n=16550)		Combined (n=22769)	
					Effect size (SE)	P value	Effect size (SE)	P value	Effect size (SE)	P value
PWV (kg/month)										
rs2860323	chr2:614210	<i>TMEM18</i>	G/A	0.12	0.09 (0.02)	5.9×10^{-5}	0.02 (0.02)	4.7×10^{-1}	0.06 (0.02)	3.9×10^{-4}
BMI-AP (kg/m²)										
rs9436303 ^b	chr1:65430991	<i>LEPR/LEP ROT</i>	G/A	0.22	0.13 (0.02)	4.7×10^{-8}	0.05 (0.01)	6.7×10^{-4}	0.07 (0.01)	8.3×10^{-9}
rs10515235	chr5:96323352	<i>PCSK1</i>	A/G	0.21	0.09 (0.02)	9.7×10^{-7}	0.03 (0.01)	1.5×10^{-2}	0.05 (0.01)	2.4×10^{-6}
Age-AR (years)										
rs1421085	chr16:53767042	<i>FTO</i>	C/T	0.25	-0.10 (0.02)	6.1×10^{-8}	-0.13 (0.01)	7.1×10^{-24}	-0.12 (0.01)	3.1×10^{-30}
rs2956578	chr5:36497552	<i>intergenic region^c</i>	G/A	0.31	0.11 (0.02)	6.7×10^{-8}	0.00 (0.01)	8.3×10^{-1}	0.04 (0.01)	1.1×10^{-3}
rs2817419	chr6:50845193	<i>TFAP2B</i>	A/G	0.76	-0.10 (0.02)	2.9×10^{-6}	-0.07 (0.01)	1.8×10^{-6}	-0.08 (0.01)	4.4×10^{-11}
BMI-AR (kg/m²)										
rs10938397	chr4:45180510	<i>GNPDA2</i>	G/A	0.35	0.09 (0.02)	5.4×10^{-6}	0.05 (0.01)	3.1×10^{-4}	0.06 (0.01)	2.9×10^{-8}
rs2055816	chr11:85406487	<i>DLG2</i>	C/T	0.25	-0.13 (0.02)	1.4×10^{-7}	-0.03 (0.02)	1.8×10^{-1}	-0.07 (0.02)	5.1×10^{-6}

^aSNP positions are according to dbSNP build 147. ^b Stage 1 meta-analysis results for this SNP is excluding the LISA-D study as this study showed heterogeneity. ^cIntergenic region between *RANBP3L* and *SLC1A3*. Results are from inverse-variance fixed-effects meta-analysis of European ancestry children. The effect allele for each SNP is labeled on the positive strand according to HapMap. The effect size is the change in standard deviations (SD) per effect allele from linear regression, adjusted for child's sex and principal components assuming an additive genetic model. BMI at AP was additionally adjusted for gestational age. None of the loci for Peak Height Velocity (PHV) passed the selection criteria for stage 2 follow up. *P* values for discovery and combined analysis are shown in bold if genome-wide significant ($P < 5 \times 10^{-8}$). The maximum sample size used in meta-analyses of each stage is shown in (). Standard Error (SE).

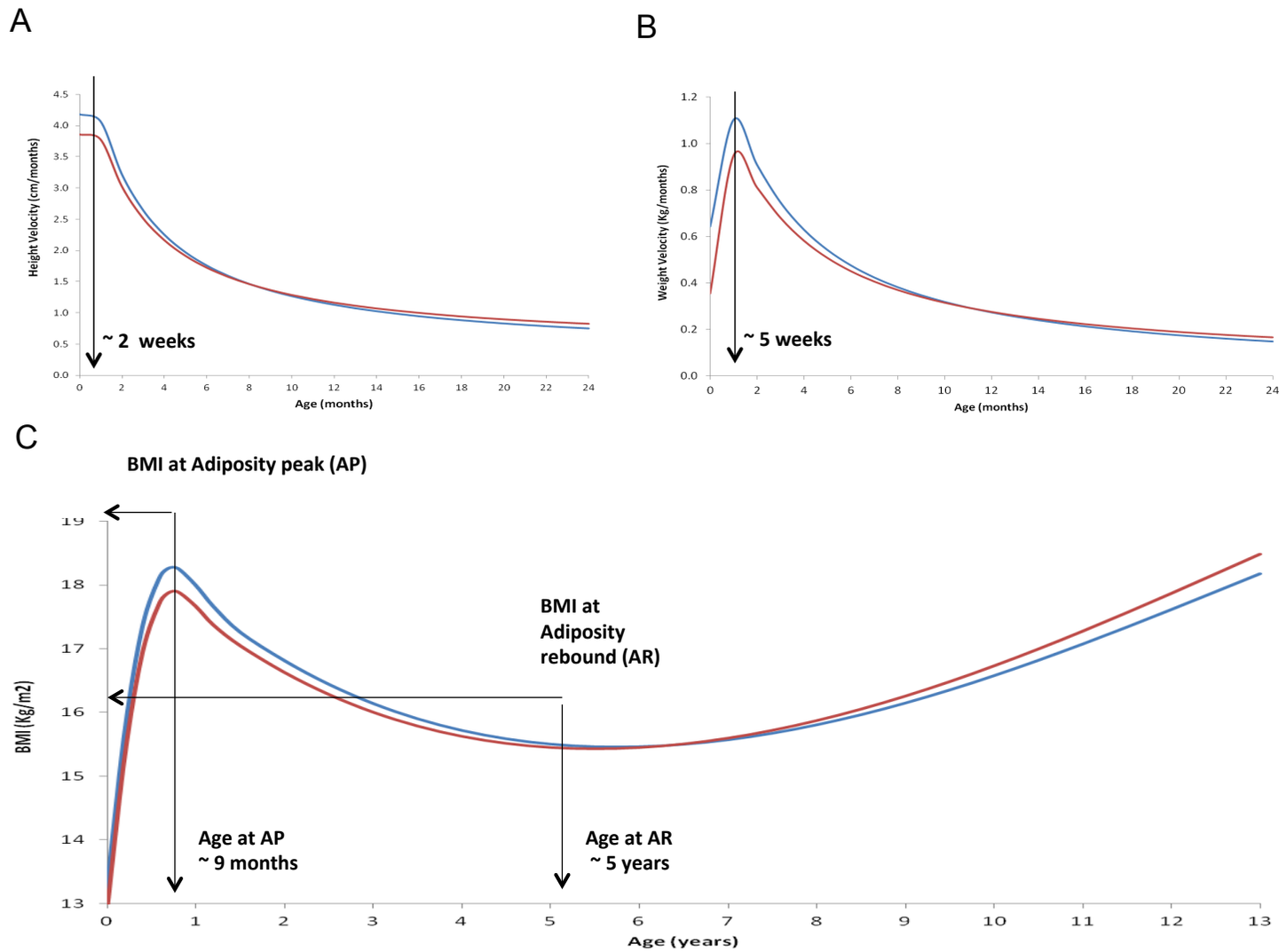


Figure 1. Graphical illustration of height and weight growth patterns and the derived measures of early growth traits used in the present study. A) Peak Height Velocity (PHV); B) Peak Weight Velocity (PWV); C) Age and BMI at Adiposity Peak (AP) and Rebound (AR). The growth curves for males are in blue and for females in red. Curves are based on the Northern Finland Birth Cohort (NFBC) 1986 data.

Early growth phenotypic traits – Study flow-chart

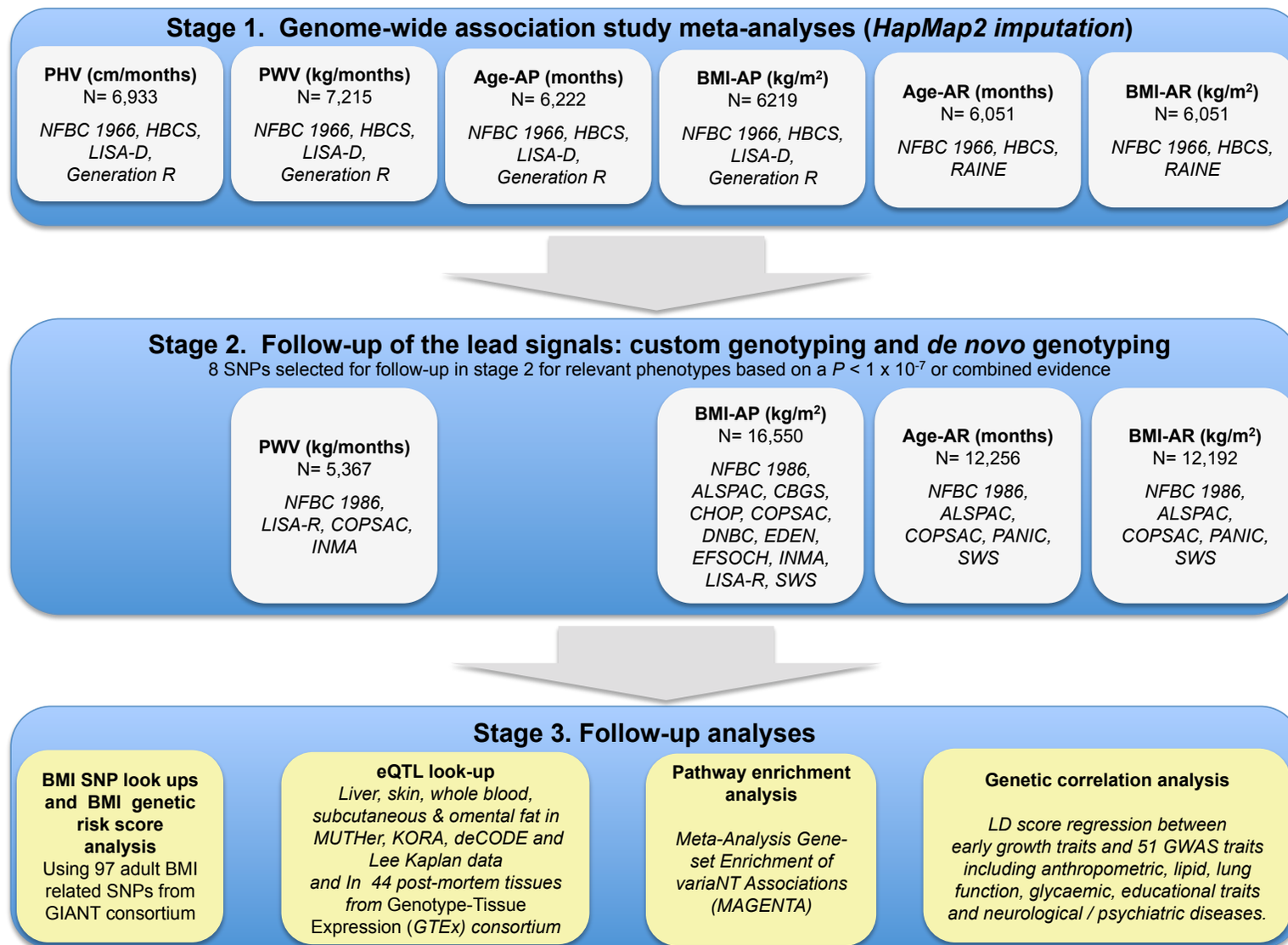
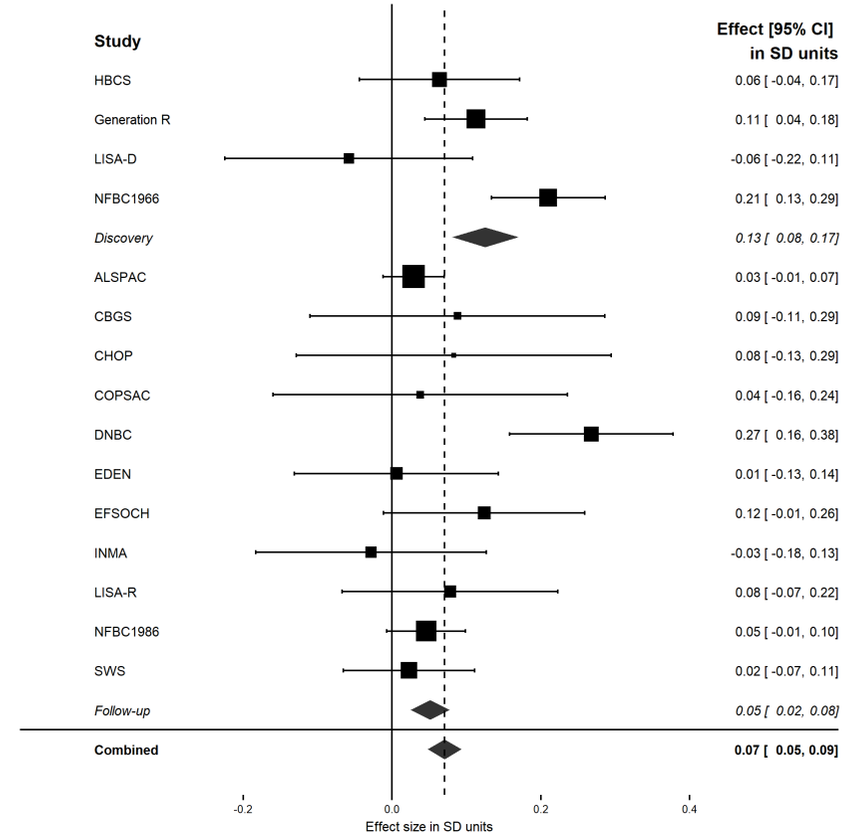
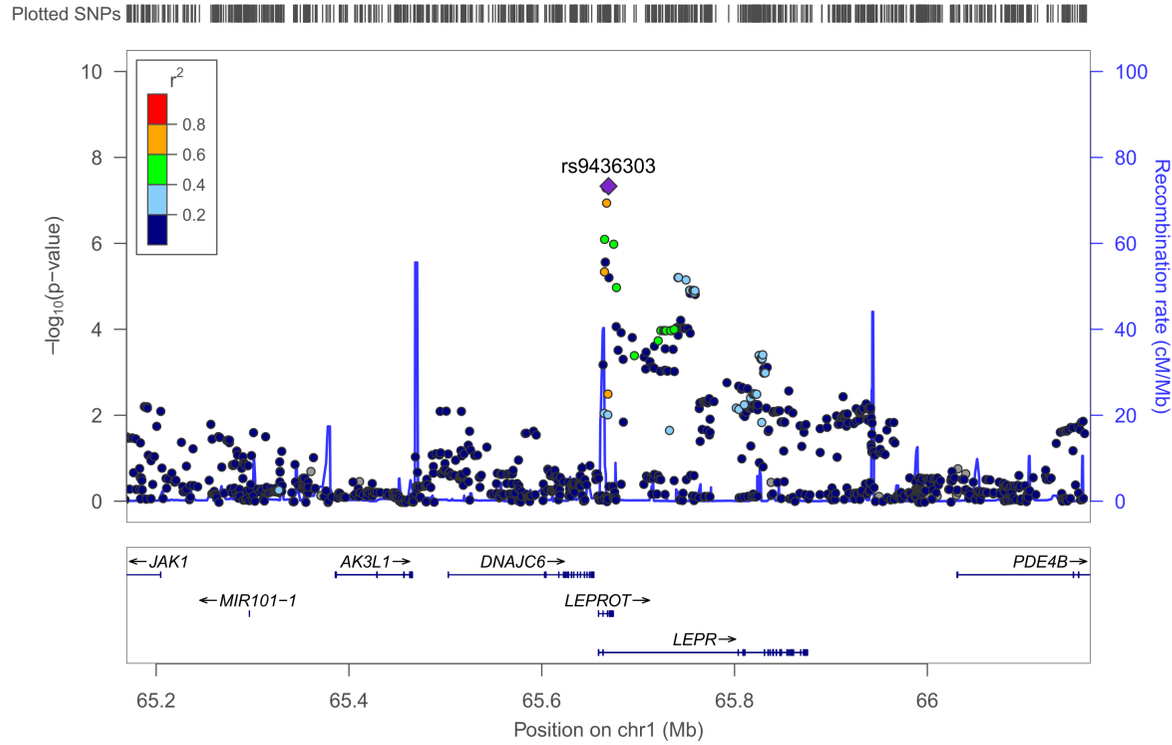


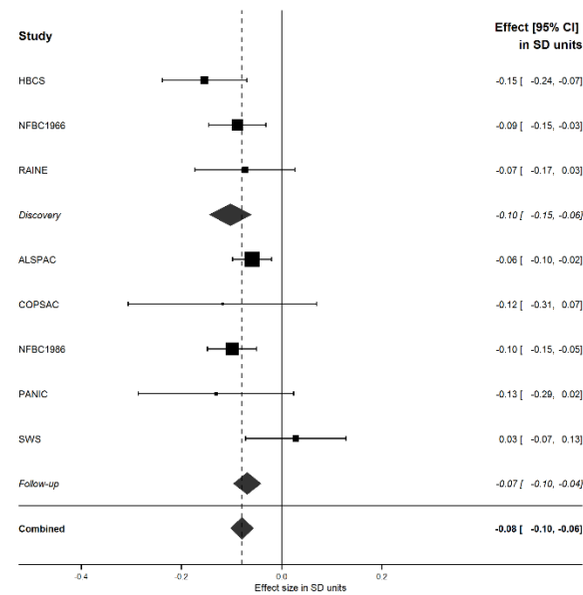
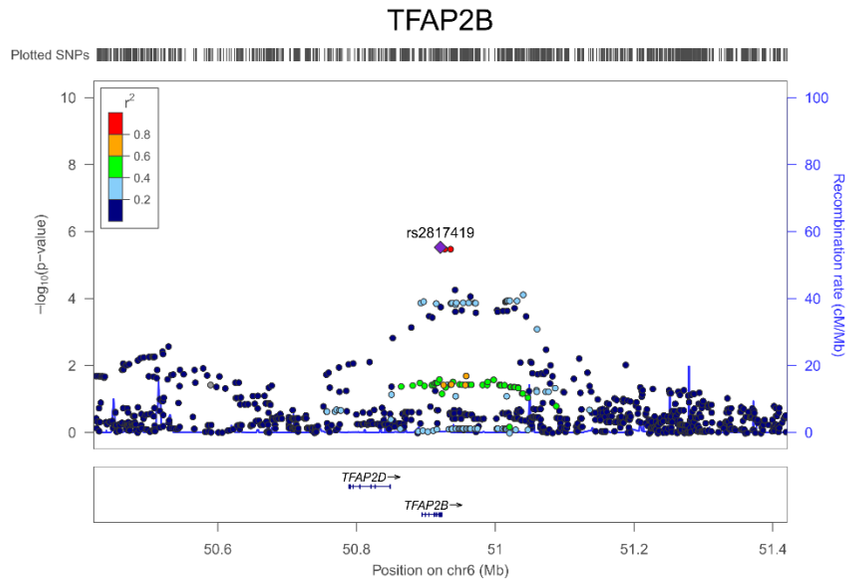
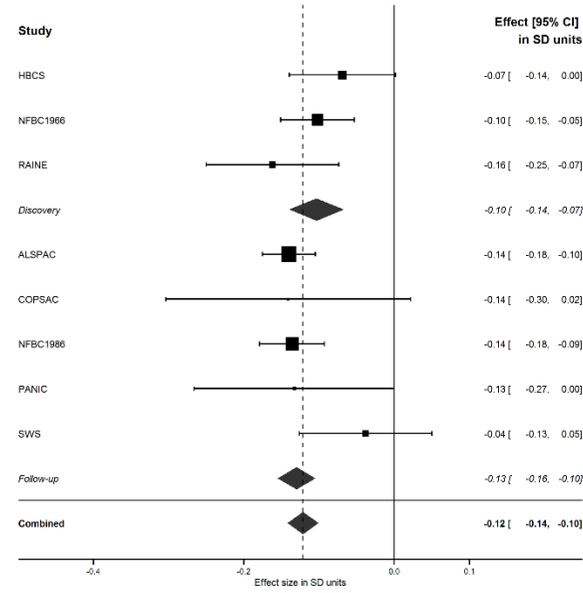
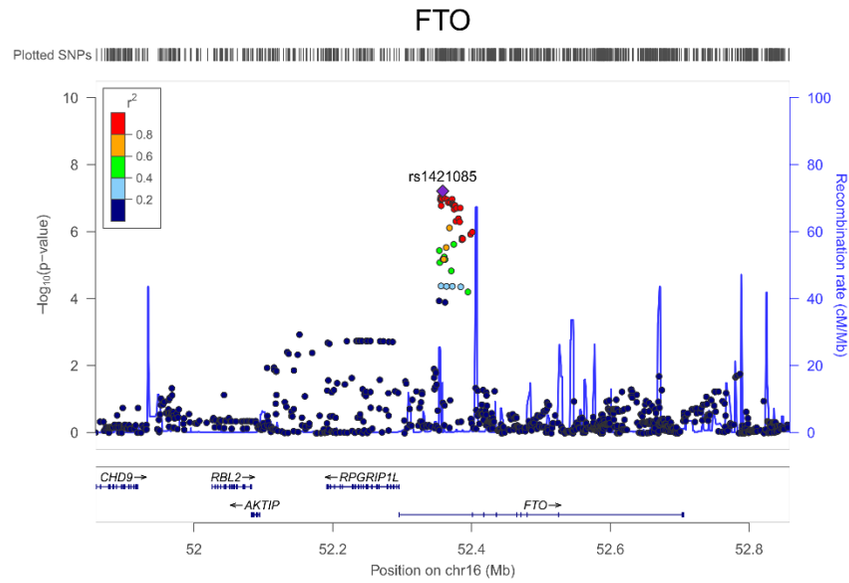
Figure 2. Summary of study design. The Flowchart shows the studies and total sample size used in Stage 1 genome-wide association meta-analyses and in Stage 2 follow up for each early growth trait leading to Stage 3 follow-up analyses. Stage 1 included up to 7215 European ancestry children from up to four population-based studies and stage 2 follow up included up to 16,550 European ancestry children from up to 11 studies.

A)

LEPR/LEPROT



B)



C)

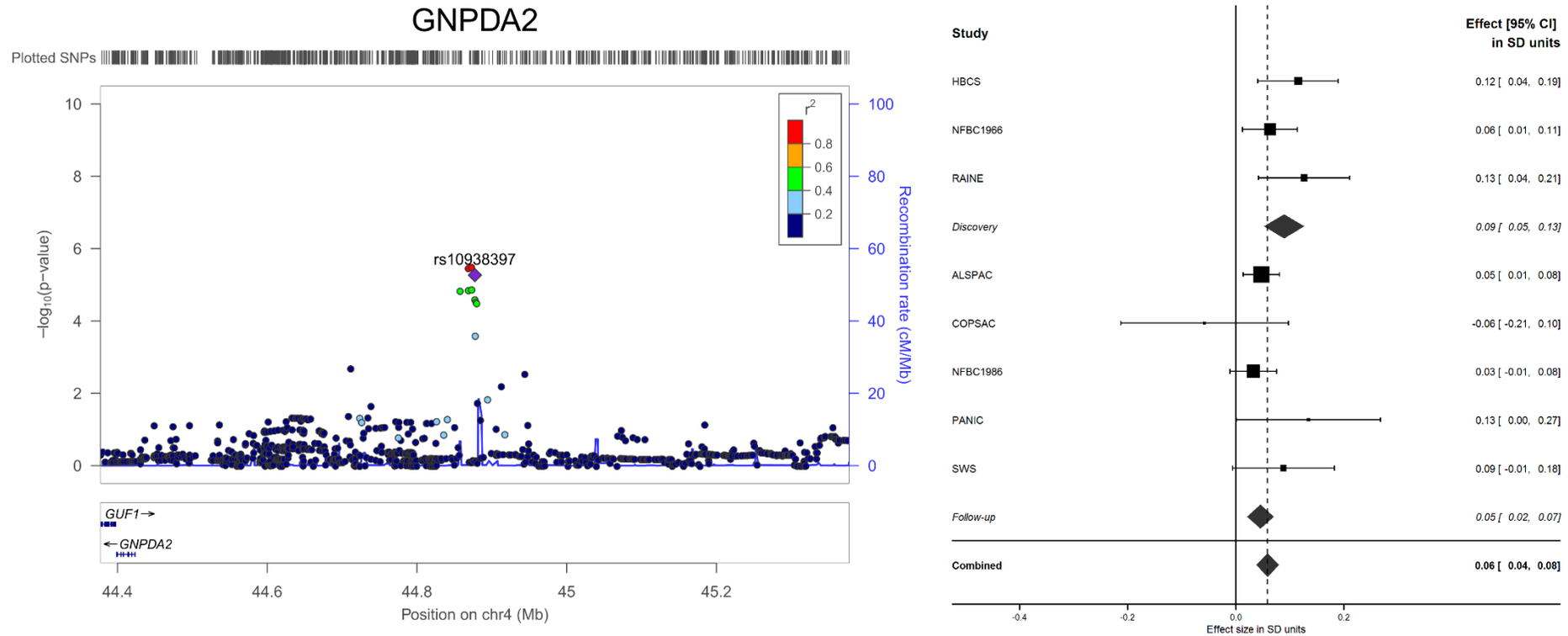
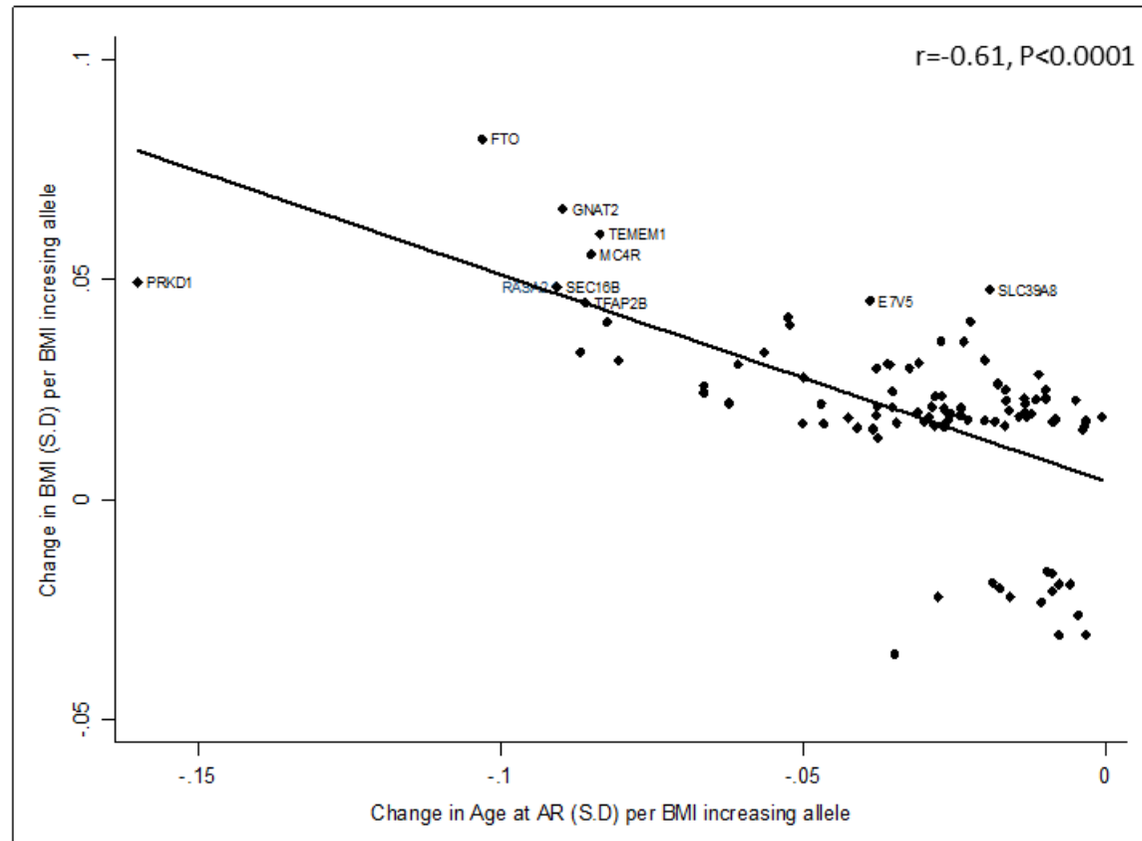


Figure 3. Regional association and forest plots of the four genome-wide significant loci associated with A) BMI at AP*, B) AGE at AR and C) BMI at AR. Purple diamond indicates the most significantly associated SNP in stage 1 meta-analysis, and circles represent the other SNPs in the region with colouring from blue to red corresponding to r^2 values from 0 to 1 with the index SNP. The SNP position refers to the NCBI build 36. Estimated recombination rates are from HapMap build 36⁷⁸. Forest plots from the meta-analysis for each of the identified loci are plotted abreast. Effect size [95% CI] in each individual study, discovery, follow-up and combined meta-analysis stages are presented from fixed effects models (heterogeneity of the SNP rs9436303 in *LEPR/LEPROT*, see **Supplementary Figure 4**). *At this locus there was heterogeneity between the studies in discovery ($I^2=72.1\%$, $P=0.01$) and combined stage ($I^2=59.3\%$, $P=0.002$) fixed effect meta-analyses that was mainly due to LISA-D, EDEN and the larger well-defined NFBC1966 study (**Supplementary Figure 4 a & d**). Removing the studies that showed inflated results from meta-analyses did not change the point estimate (**Supplementary Figure 4 c, f, g**). Both fixed and random effect models gave very similar results (**Supplementary Figure 4b & 4e**).

A)



B)

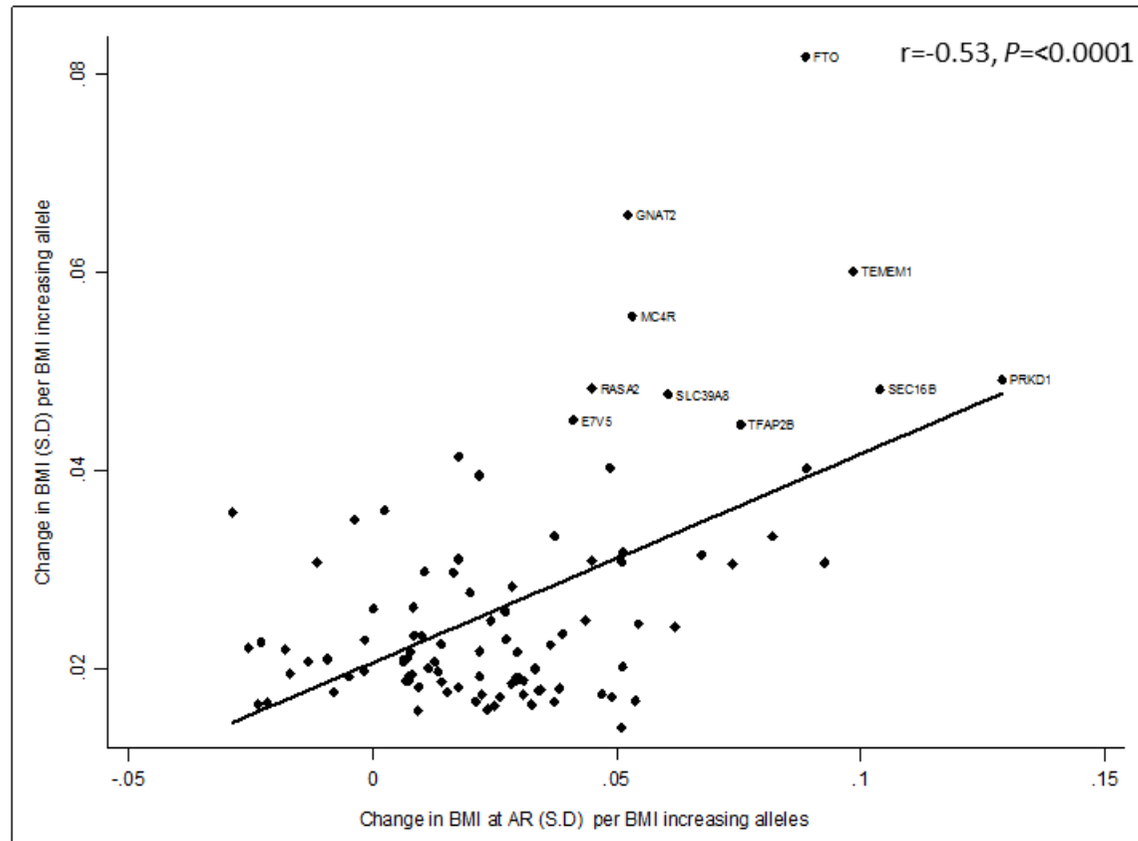


Figure 4. Scatter plots of the effect size estimates (SD units) of the 97 adult BMI associated loci on adult BMI in GIANT consortium¹ and Age at AR (A) and BMI at AR (B) in the current discovery stage GWAS meta-analyses. The effect size of the adult BMI increasing allele is plotted. The r represents the correlation coefficient between the effect size of the early growth phenotype and adult BMI and P is P-value for this correlation coefficient. The scatter plots of the other early growth phenotypes are given in **Supplementary Figure 5**.

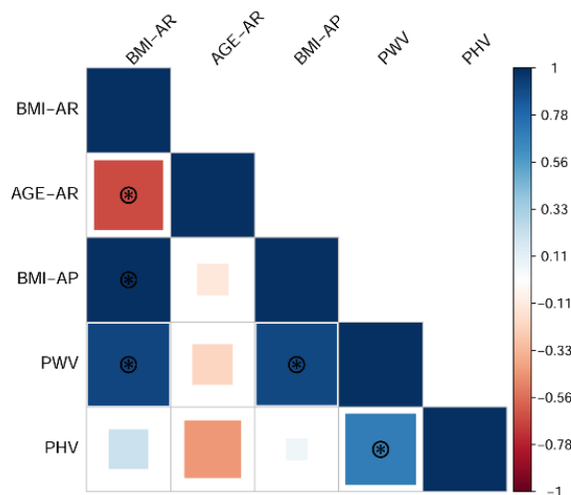
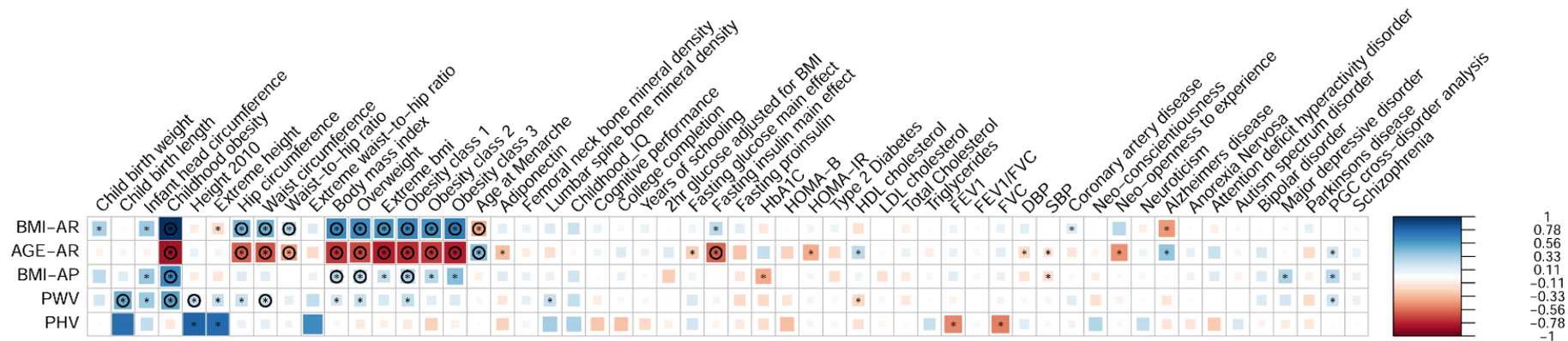


Figure 5: Genetic correlations between five early growth traits and 54 other GWAS traits analyzed by genome-wide association statistics. Blue, positive genetic correlation; red, negative genetic correlation. The correlation matrix underneath represents the genetic correlation among the five early growth traits themselves. The size of the coloured squares is proportional to the P -value where larger squares represent a smaller P -value. Genetic correlations that are different from 0 at $P < 0.05$ is marked with an asterisk. The genetic correlations that are different from 0 at a false discovery rate (FDR) of 1% FDR are marked with a circle. Genetic correlations represented here are presented in tabular format in **Supplementary Table 14**.

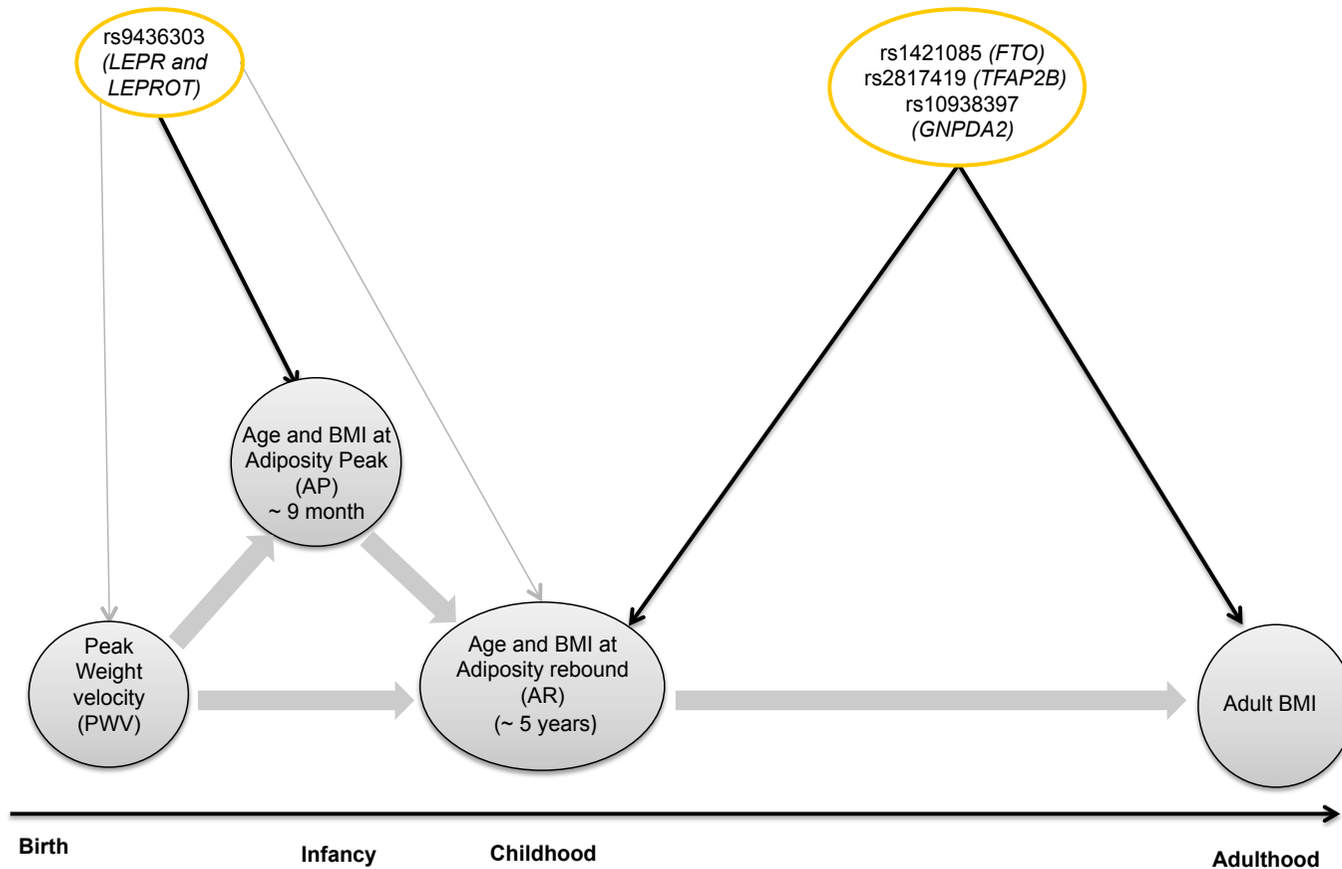


Figure 6. Schematic diagram showing the four genome-wide significant loci associated with early childhood growth phenotypes in the current study and adult BMI in the GIANT consortium. The thick grey arrows show main observational relationships between the early growth phenotypes at $P < 0.0001$ in NFBC1966 and NFBC1986 studies, and how early growth phenotypes may be linked with adult BMI. The thin arrows represent the association between the index SNPs and the early growth phenotypes in combined meta-analyses or in look-ups in GIANT consortium data¹; $P < 0.05$ (grey); $P < 0.0001$ (black).

