Systolic and diastolic blood pressure (SBP and DBP) are complex phenotypes, with known environmental and genetic risk factors. Elevated SBP and DBP are associated with premature mortality and cardiovascular diseases. Clinical Perspective on p 278

Recent genome-wide association studies (GWAS) have identified genetic variants associated with adult SBP and DBP and hypertension. Several of these loci are in biologically plausible candidate genes, for example, those that influence the renin–angiotensin system. There are established patterns of age-related changes in BP in industrialized populations, which support a potential interaction of genetic variants with age-related changes to environmental exposures in these populations. For genetic variants that directly modulate childhood BP, effects might change with age, might differ between developmental periods (early life, childhood, and adolescence), and might also differ to those variants that act in adulthood.

Children and adolescents are rarely treated with antihypertensives, whereas from middle age onwards, an increasing proportion of adults are using such medications. Consequently, it is easier to examine genetic variants that are associated with untreated SBP and DBP in children. Variation in SBP, and to
a lesser extent DBP, in adolescence and early adulthood is associated with subsequent adult risk of coronary heart disease and stroke.\textsuperscript{28,29} Understanding the risk factors, including genetic variation, that are associated with SBP and DBP through childhood and into adolescence may therefore inform an improved understanding of the life course pathogenesis of adult hypertension and cardiovascular disease. GWAS of BP in children to date have been limited to investigations of cross-sectional BP in individual cohorts,\textsuperscript{30} often in non-European populations.\textsuperscript{31,32}

The principal aim of the current study was to identify age-specific genetic associations with SBP and DBP across childhood and adolescence. The secondary aim was to compare results from this GWAS to published results from adult GWAS of BP.

**Methods**

**EAGLE Consortium: BP Working Group**

Seven pregnancy/birth or childhood cohorts of European ancestry from the Early Genetics and Lifecourse Epidemiology (EAGLE) Consortium BP working group with completed genome-wide genotyping and measures of SBP and DBP on participants between the ages of 4 and 20 years contributed to this study.

Each study was conducted with appropriate institutional ethics approval, and written informed consent was obtained from participants or their main caregivers. Each study conducted their own GWAS analyses, and results were then pooled using meta-analysis. The methodology for recruitment of participants and measurement of BP for each cohort are described in the Data Supplement.

Only those participants who had at least one BP measurement at, or before, 20 years of age were included in analyses. With the exception of the Brisbane Longitudinal Twin Study (BLTS), all analyses were restricted to children from singleton pregnancies, the first born (with relevant genotype and phenotype data) from cohorts that included siblings, or a randomly selected child from those that included twins. BLTS, which included all twins and siblings, took into account both zygosity and relatedness using the statistical package Merlin\textsuperscript{33} during their GWAS.

**ICBP**

We used GWAS data from studies contributing to the International Consortium of Blood Pressure (ICBP) on adults who were recruited for classical or genetic epidemiological purposes from general population samples\textsuperscript{34} to compare with associations identified in EAGLE.

**Genotyping and Imputation**

Single nucleotide polymorphisms (SNPs) were genotyped on one of the following platforms: Affymetrix, Illumina, or Perlegen. Predefined marker filters were applied before imputation (Hardy–Weinberg equilibrium >10\textsuperscript{−8}, MAF >0.01, SNP call rate >95%). Each study imputed SNPs by combining their study’s genotyped SNPs with HapMap Phase II CEU SNPs samples, preferably using release 22 of HapMap build 36. Imputation results are summarized as an allele dosage distribution and a Q-statistic. Further filtering based on N-effective >70% was also used. QQ-plots for the meta-analysis of each BP outcome across each epoch and data set are presented in Figure II in the Data Supplement. A threshold of $P \leq 5 \times 10^{-8}$ was used to define genome-wide levels of significance. An association of an SNP cluster was defined as a least two variants each reaching a threshold of $P \leq 5 \times 10^{-8}$. This more liberal statistical significance level ($P \leq 5 \times 10^{-6}$) was used to ensure the inclusion of variants that could potentially have been identified in larger samples of children.

**Exploring Functionality**

We used the Encyclopedia of DNA Elements\textsuperscript{35} to investigate the functionality (transcriptional active region, H3K27Ac regulatory

**Cross-Sectional GWAS**

Cross-sectional associations were performed for SBP and DBP in 3 age epochs over childhood/adolescence. These epochs were chosen before any analyses and were based on previous evidence of the ages in European populations that girls go through menarche, boy’s voices break, and both sexes have a growth spurt and in part pragmatically, relating to available data in the participating studies. These epochs were the following: prepuberty (4 to 7 years), puberty (8 to \( \leq 12 \) years), and postpuberty (13 to \( \leq 20 \) years; Table I and Figure I in the Data Supplement).

Genetic analyses assumed an additive model for SNP-based main effects. All analyses were adjusted for age, height, and weight, consistent with analyses used in GWAS of BP in adults (adjusted for body mass index).\textsuperscript{36} The Z-score was calculated per individual as the difference between an individual’s BP measure and the average BP measure for that given age epoch and sex for each cohort. This was then divided by the standard deviation of the BP measure across the particular cohorts’ age, epoch, and sex. Those Z-scores were then regressed against each SNP (additive) adjusting for age, sex, and height of the individual. Participants could contribute to >1 epoch but only one measurement time point contributed to each epoch. The median measurement time was selected in the event that a cohort had repeatedly measured data within a single epoch.

**Meta-Analyses**

Separate genome-wide meta-analyses were run for SBP and DBP for each epoch using the inverse-variance weighting method in Meta-Analysis (software for GWAS) (www.sph.umich.edu/csg/abecasis/metaetal).\textsuperscript{36} Before meta-analysis, rare variants were excluded (MAF>0.01; Table I in the Data Supplement). Double genomic control correction was applied (once during the study-specific analyses [before the meta-analysis] and repeated on the statistics resulting from the meta-analysis). Heterogeneity between results from individual studies was assessed using I\textsuperscript{2} and a Q-statistic. Further filtering based on N-effective >70% was also used. QQ-plots for the meta-analysis of each BP outcome across each epoch and data set are presented in Figure II in the Data Supplement. A threshold of $P \leq 5 \times 10^{-8}$ was used to define genome-wide levels of significance. An association of an SNP cluster was defined as at least two variants each reaching a threshold of $P \leq 5 \times 10^{-8}$. This more liberal statistical significance level ($P \leq 5 \times 10^{-6}$) was used to ensure the inclusion of variants that could potentially have been identified in larger samples of children.

**Table 1. List of Participating Cohorts With GWAS and SBP and DBP Data Available in Children**

| Cohort         | Time Frames, epochs |
|----------------|---------------------|
|                | Prepubertal, (4–7 y) | Pubertal, (8–12 y) | Postpubertal, (13–20 y) |
| Raine          | 1276                | 1251               | 1009               |
| GenerationR    | 1847                |                    |                   |
| ALSPAC         | 5967                | 5750               | 4050               |
| Lisa Plus      | 282                 |                    |                   |
| YFS            | 400                 | 842                |                   |
| INMA           | 600                 |                    |                   |
| BLTS           | 10090               | 8423               | 5176               |

Number of subjects per cohort per time frame used in analyses. ALSPAC indicates The Avon Longitudinal Study of Parents and Children Bristol, UK; BLTS, Brisbane Longitudinal Twin Study, Brisbane, Queensland, Australia; GenerationR, The Generation R Study Group, Rotterdam, The Netherlands; INMA, Spanish INMA—Infancia y Medio Ambiente, Barcelona, Catalonia, Spain; Lisa Plus, Influence of lifestyle factors on the development of the immune system and allergies in East and West Germany Plus the influence of traffic emissions and genetics, Neuenberg, Germany; Raine, The Western Australian Pregnancy (Raine) Cohort, Perth, Western Australia; YFS, The Cardiovascular Risk in Young Finns Study, Turku, Finland.
elements, DNase hypersensitivity transcription factor–binding site information from chromatin immunoprecipitation (ChIP) analysis, and 5′-C-phosphate-G-3′ (CpG) methylation levels) of identified BP-related SNPs.

Comparing Findings Across Epochs and in Adults

Any variants identified as associated with SBP or DBP at \( P \leq 5 \times 10^{-3} \) in any age epoch was examined in (1) the other 2 epochs in our study; (2) in adult participants from the ICBP, and (3) in SNPs identified as associated with adult CHD using the results from Coronary Artery Disease Genome-Wide Replication and Meta-Analysis (CARDiogramPlusC4D).36 A consortium designed to identify novel susceptibility loci for coronary artery disease and myocardial infarction.35

We also examined whether variants associated with adult BP in ICBP at \( P \leq 5 \times 10^{-3} \) for association with BP in any of the 3 epochs in our study.

Meta-analyses were conducted in Meta-Analysis (software for GWAS). The statistical software package R version 2.12.139 was used to produce all Manhattan Plots. Regional association plots were produced using LocusZoom.40

Results

Seven cohorts contributed to meta-analyses, a total of 10 090 individuals for the prepubertal epoch, 8423 individuals for the pubertal epoch, and 5176 individuals for the postpubertal epoch (Table 1). Manhattan plots for SBP and DBP by epoch are presented in Figure III in the Data Supplement. Regional association plots for the most significant SNP (by outcome and epoch, as listed in Table 2) are shown in Figure IV in the Data Supplement. Regional association plots for SNP clusters are presented in Figure V in the Data Supplement (SBP) and Figure VI in the Data Supplement (DBP). Regional association plots and linkage disequilibrium plots for significant SNP clusters for SBP and DBP are presented by epoch in Figures VII to IX in the Data Supplement. Forest plots for all significant SNP clusters and most significant association by outcome and epoch are presented in Figures X to XV in the Data Supplement. Overview figures (forest plot, regional association plot, and Manhattan plot) are presented by epoch and outcome in Figures XVI to XVIII in the Data Supplement. Plots relating to the Encyclopedia of DNA Elements functionality are displayed in Figures XXIb and XXIc in the Data Supplement. The statistical software package R version 2.12.139 was used to produce all Manhattan Plots. Regional association plots were produced using LocusZoom.40

Systolic Blood Pressure

Genome-Wide Significance

Two SNPs reached genome-wide levels of significance \((P \leq 5 \times 10^{-8})\) for childhood SBP (Table 2): in the prepubertal epoch, rs1563894 (chromosome position, 15.64422829; gene, ITGA11; per allele mean difference in SBP = −0.093SD [95% CI: −0.126, −0.060]; \( P = 2.86 \times 10^{-8} \); Tables 2 and 3 and Figures VI and XV in the Data Supplement); and during the pubertal epoch, rs872256 (chromosome position, 9.2496480; gene, unknown; \( \beta = 0.096SD [95\%\ CI: 0.063, 0.129]; P = 8.67 \times 10^{-6} \); Table 2). No SNPs reached genome-wide levels of significance for SBP in the postpubertal analyses.

Clusters of variants in several genes were associated with SBP at \( P \leq 5 \times 10^{-3} \) (Table 3, Figure 1, and Figure V in the Data Supplement): ITGA11 (puberty), ANLN and OR51V1 (puberty), and C1GALT1 and UGP2 (postpuberty). A cluster of SNPs was also identified surrounding SNP rs1563894 (ITGA11; puberty), highlighted above as showing genome-wide levels of significance with prepubertal SBP. We illustrate this significant SNP cluster in more depth (Figure 2).

Two SNPs were associated with the largest increases in SBP across these epochs: SNP rs3735398 (ANLN) during the pubertal epoch \((\beta = 0.116SD, 95\%\ CI: 0.073, 0.159; P = 1.28 \times 10^{-7}; \) Table 3 and Figure VIII in the Data Supplement) and a cluster surrounding SNP rs3901287 (LOXL2; \( \beta = 0.108SD, 95\%\ CI: 0.059, 0.157; P = 1.24 \times 10^{-7}; \) Table 3 and Figure IX in the Data Supplement) was associated with postpubertal SBP.

Look-Up of Functionality of Childhood SBP Results

Several SNPs were found to be located either directly in or in close proximity to functionally active regions. For SNPs associated with SBP, rs1563894 was located on an active H3K27Ac, DNase hypersensitivity, transcription factor ChIP, and CpG methylation site (Figure 3 and Figure XIXa in the Data Supplement). Rs1010366 was located downstream and in close proximity to densely active H3K27Ac, DNase hypersensitivity, transcription factor ChIP, and CpG methylation site (Figure XIXc in the Data Supplement). Rs3735398 was located in a region of active transcription and DNase hypersensitivity. Rs3787159 was located directly upstream from areas with active transcription DNase hypersensitivity and transcription factor ChIP (Figure XXIb in the Data Supplement). Rs4538187 was located in a region of transcription factor ChIP and methylation and upstream from an active transcription site and downstream from an active H3K27Ac mark with dense methylation and transcription factor active H3K27Ac mark, DNase hypersensitivity, transcription factor ChIP, and CpG methylation site, whereas rs3901287 was located in close proximity to areas of active transcription, H3K27Ac mark, DNase hypersensitivity, transcription factor ChIP, and methylation (Figure XXIc in the Data Supplement).

Comparisons Across Epochs and With Adult Outcomes

No SNPs reached associations of \( P \leq 5 \times 10^{-3} \) in all 3 epochs. Variants in chromosome 2 (rs13025174, rs13032473 [TMEM247], and rs10186089 [FSHR]) were associated with elevated SBP during prepubertal and pubertal epochs (Table 4). Several SNPs

Comparisons Across Epochs and With Adult Outcomes

No SNPs reached associations of \( P \leq 5 \times 10^{-3} \) in all 3 epochs. Variants in chromosome 2 (rs13025174, rs13032473 [TMEM247], and rs10186089 [FSHR]) were associated with elevated SBP during prepubertal and pubertal epochs (Table 4). Several SNPs
in HORMAD2, HORMAD2-AS1, and MTMR3 were associated with SBP during the pubertal and postpubertal epochs (Table 4 and Table XIII in the Data Supplement).

None of the SNPs found to be associated (P ≤ 5 × 10^{-3}) with childhood SBP were associated with adult SBP in the ICBP (Table 3). However, several loci previously found to be associated with adult SBP in ICBP were also associated with SBP in at least one epoch of childhood in EAGLE children (Table 5): (1) B3GNTL1 and KLHL1 (prepuberty); (2) DOK6, NKAIN2, ARHGGEF10, and MECOM (puberty) and (2) FGD5 and CSMD2 (postpuberty). One SNP that was associated with SBP in the pubertal epoch showed some possible association with CHD in adults (Table VII in the Data Supplement; rs3735398; P = 0.039).

Diastolic Blood Pressure

**Genome-Wide Significance**

No SNPs reached genome-wide levels of significance (P ≤ 5 × 10^{-8}) for childhood DBP (Table 2).

Clusters of variants in 7 loci were associated with childhood DBP (P ≤ 5 × 10^{-3}; Table 3) and were investigated further using LocusZoom. Regional associations for all of these associations are presented in Figure VI in the Data Supplement. SNPs associated with the largest positive increases in DBP per epoch included a prepubertal epoch cluster led by rs1714524 (LOC100996447; β = 0.154SD, 95% CI, [0.081, 0.227]; P = 8.32 × 10^{-5}; Figure 2 and Figure VIa in the Data Supplement). For the pubertal epoch, a cluster led by rs1387977 (TRHDE; β = 0.127SD, 95% CI, 0.064,
For the postpubertal epoch, a cluster led by rs6949619 (gene unknown, near NPY; $\beta = -0.092$SD, 95% CI, $-0.133$, $-0.051$; $P = 1.13 \times 10^{-5}$; Figure VIc in the Data Supplement).

Look-Up of Functionality of Childhood DBP Results

For SNPs associated with DBP, rs13040824 was located downstream from an active H3K27Ac mark and located between 2 regions of DNase hypersensitivity and transcription factor ChIP (Figure XXa in the Data Supplement). Rs7897969 was located between 2 regions that had high levels of DNase hypersensitivity, transcription factor ChIP, and methylation (Figure XXb in the Data Supplement). Rs1236530 was located in a region of DNase hypersensitivity and transcription factor ChIP (Figure XXc in the Data Supplement). Rs241264 was located between 2 areas of high methylation, whereas rs1714524 was located in an area of active transcription (Figure 3 and Figure XXIIa in the Data Supplement). Rs1687522 was located in an area of active transcription and methylation (Figure XXIIb in the Data Supplement). Rs229038 was located within a cluster of DNase hypersensitivity and upstream for a highly active region of transcription and regulation (Figure XXIIc in the Data Supplement).

Look-Up of DBP Results Across Epochs in Childhood/Adolescence

No SNPs were significantly associated ($P \leq 5 \times 10^{-3}$) with DBP across all epochs. Opposing effects were observed for rs13004438 ($CCDC141$) for increasing DBP ($\beta = 0.082$SD, 95% CI, $0.034$, $0.13$; $P = 8.31 \times 10^{-4}$) during the prepubertal epoch but reducing it during the pubertal epoch ($\beta = -0.053$SD, 95% CI, $-0.115$, $-0.030$; $P = 8.10 \times 10^{-4}$; Table 4). Variants in $POT1$ and $CNTNAP4$ were associated with consistently reducing DBP during the pubertal and postpubertal epochs (Table 4 and Table XIV in the Data Supplement).

None of the SNPs found to be at least associated ($P \leq 5 \times 10^{-3}$) with childhood DBP were associated with adult BP in the ICBP (Table 4). A number of loci previously found to be associated with adult DBP in ICBP were also associated with DBP in at least one epoch of childhood in EAGLE children (Table 5): $USP4$ and $GRK5$ (all pubertal).

We also investigated SNPs from Tables 2 and 3 in CARDIoGRAMplusC4D36 and found 2 significant hits for rs12365302 ($P = 0.008$) and rs16875222 ($P = 0.037$; Table VII in the Data Supplement).

### Additional Analyses

As a sensitivity analyses, we repeated the GWAS analyses separately in females and males and found similar directions and magnitudes of associations, though given the smaller sample sizes within these 2 subgroups were not as well powered to report on the sex-stratified associations (Tables VIII–XII in the Data Supplement).

| Time Frame | Marker Name | Allele | CHR | POS | MAF | Gene | Beta (95% CI) | P Value | Direction | ICBP p | N Effective |
|------------|-------------|--------|-----|-----|-----|------|--------------|---------|-----------|--------|-------------|
| Systolic blood pressure | Prepubertal | rs1563894 | A/G | 15 | 66422829 | 0.19 | ITGA11 | -0.093 (-0.126, -0.060) | 2.86E-08* | -------- | 0.46 | 10 090 |
| Pubertal | rs3753938 | A/G | 7 | 36412646 | 0.12 | ANLN | 0.116 (0.073,0.159) | 1.28E-07 | ++++ | 0.75 | 8423 |
| | rs3787159 | T/C | 20 | 56252573 | 0.46 | PPP4R1L | -0.064 (-0.091, -0.037) | 8.76E-06 | -------- | 0.46 | 8423 |
| | rs9667878 | T/C | 11 | 5180326 | 0.22 | OR51V1 | 0.135 (0.0740,0.196) | 9.74E-06 | ++++ | 1.00 | 8423 |
| Postpubertal | rs1010366 | T/C | 7 | 7196351 | 0.39 | C1GALT1 | 0.098 (0.057,0.139) | 3.31E-06 | +++ | 0.90 | 5176 |
| | rs4538187 | A/G | 2 | 63957245 | 0.16 | UGP2 | 0.102 (0.057,0.147) | 5.96E-06 | +++ | 0.70 | 5176 |
| | rs3901287 | A/T | 8 | 23240509 | 0.28 | LOXL2 | 0.108 (0.059,0.157) | 1.24E-05 | +++ | 0.68 | 5176 |

Diastolic blood pressure

| Prepubertal | rs241264 | T/C | 1 | 4518898 | 0.31 | Between LOC284661 and AJAP1 | -0.082 (-0.0120, -0.043) | 2.95E-05 | -------- | 0.48 | 10 090 |
| rs1714524 | T/C | 3 | 159755790 | 0.44 | LOC100996447 | 0.154 (0.081,0.227) | 8.32E-05 | -------- | 0.99 | 10 090 |
| rs16875222 | A/T | 8 | 107955966 | 0.11 | Near ABRA, OXR1 | -0.119 (-0.179, -0.060) | 8.21E-05 | -------- | 0.72 | 8423 |
| rs12237240 | T/G | 9 | 28329306 | 0.19 | LING02 | 0.068 (0.034,0.102) | 8.71E-05 | ++++ | 0.89 | 8423 |
| rs1387977 | T/G | 12 | 71307607 | 0.14 | TRHDE | 0.127 (0.064,0.190) | 6.92E-05 | ++++ | 0.90 | 8423 |
| Postpubertal | rs6949619 | T/C | 7 | 24396900 | 0.19 | Near NPY | -0.092 (-0.133, -0.051) | 1.13E-05 | -------- | 0.45 | 5176 |
| | rs229038 | C/G | 21 | 27127300 | 0.22 | ADAMTS1 | 0.213 (0.115,0.311) | 2.26E-05 | +++ | 0.77 | 5176 |

Bold text highlights SNPs ($P \leq 5 \times 10^{-3}$) represented in regional association plots shown in Figures VII–IX in the Data Supplement. Beta values are in terms of Z-scores, the number of standard deviations away from the mean. CHR indicates chromosome; MAF, minor allele frequency; and POS, position.

*SNPs which have reached genome-wide levels of significance ($P \leq 5 \times 10^{-8}$).
Discussion

To our knowledge, this is the first GWAS to examine genetic associations with BP cross-sectionally across childhood and adolescence. This builds on our previous work in which we found that an allelic score summing the established adult GWAS hits for SBP was associated with SBP at mean age 6 years but not with age-related change in SBP between age 6 and 17.41,42 In the current study, we have identified 2 novel SNPs: one associated with SBP measured prepubertally and one with SBP measured during puberty. We did not find any SNPs reaching genome-wide significance with DBP or with either SBP- or DBP-measured postpuberty, but when we reduced our statistical significance threshold to $P \leq 5 \times 10^{-3}$, there were several additional variants in gene clusters relating to SBP or DBP across the 3 epochs. At this more liberal $P$ value threshold, we found some evidence of overlap in novel SNPs across childhood/adolescent age epochs within our study and some overlap with those that have been found by previous GWAS to be associated with adult BP.

*ITGA11* was genome-wide associated with SBP during the prepubertal epoch and has been shown to be associated with hypertrophic cardiomyopathy43 and coronary artery disease.44 An SNP in close proximity to the *SMARCA* and *VLDLR* genes (rs872256) was genome-wide associated with SBP during the pubertal epoch. *SMARCA* is a member of the *SWI/SNF* family of proteins and is highly similar to the brahma protein, where it has been hypothesized that cardiac hypertrophy and the fetal gene expression program are associated with distinguishable binding of brahma and *SMARCA4* on genes.45 From animal studies, brahma gene expression is found to be restricted to mesodermal tissues involved in early vasculogenesis and heart morphogenesis.46 *VLDLR* has been shown to be associated with obesity from both animal studies47–49 and human GWAS.50 Recently, a pathway analyses based on results from a GWAS has identified plausible biological links between *VLDLR* with vascular endothelial growth factor, which is known to affect angiogenesis and atherosclerosis.51 Even with a reduced $P$ value threshold of $P \leq 5 \times 10^{-3}$, these 2 genome-wide novel variants did not overlap with variants at this threshold in any other age group.

Some of the variants that we identified as associated with SBP or DBP in any of the different age epochs were in gene clusters that also had some evidence of potentially relevant functionality. *FSHR* which was associated with SBP in prepubertal and puberty has been shown to influence SBP vascular responses in hypertensive rats with hyperhomocysteinemia52 and is known to be involved in the regulation of systemic arterial BP (GeneCards). *POT1* which was associated with lower DBP in pubertal and postpubertal children is an important molecular marker for biological aging.53 *MTMR3* which was associated with lower SBP in puberty and postpuberty has been hypothesized to be a mediator for miR-4513, which is significantly associated with BP and related metabolic outcomes, such as cholesterol.54

The fact that we have found more variants associated with SBP than with DBP may reflect differences in changes in SBP and DBP across childhood and adolescence. SBP increases across childhood to a peak at around age 15/16 years and thereafter levels off, whereas DBP seems to increase monotonically across childhood into late adolescence/early twenties.26,27 This is the first GWAS of BP in children at different ages, and we have maximized the sample size by collaborating across several studies with relevant data. We limited analyses...
Figure 2. Most significant SNP clusters resulting from the sex-combined meta-analyses genome-wide association study (GWAS). From top to bottom: forest plot, regional association plot (−log10 \( P \) values), linkage disequilibrium plots (LOD scores), and Manhattan plots (−log10 \( P \) values)). Green boxed areas on Manhattan plots highlight chromosome regions illustrated in regional association plots. Letters signify the following data subsets: SBP (A) and DBP (B) for the prepubertal epoch. ALSPAC indicates The Avon Longitudinal Study of Parents and Children Bristol, UK; EAGLE: Early Genetics and Lifecourse Epidemiology; GenR, The Generation R Study Group, Rotterdam, The Netherlands; ICBP, International Consortium of Blood Pressure; INMA, Spanish INMA—Infancia y Medio Ambiente, Barcelona, Catalonia, Spain; Raine, The Western Australian Pregnancy (Raine) Cohort, Perth, Western Australia; YFS; The Cardiovascular Risk in Young Finns Study, Turku, Finland.
to Europeans only to minimize population stratification and followed-up hits by looking up for functionality and for overlap with SNPs from GWAS in adults.

We used age groups to define epochs during which BP was measured as prepubertal, pubertal, and postpubertal. We acknowledge that some participants will have been incorrectly categorized by this method. However, it was not possible to use Tanner scores for all participants, and to have used those data would have compromised our sample size. Furthermore, assessment of pubertal stages using self- (or parental) assessment of Tanner scales, which were the methods used in most cohorts in EAGLE, is also prone to misclassification. Misclassification because of using age is likely to be random, whereas Tanner scores could be systematic in relation to characteristics, such as body mass, that are related to BP.

Our sample size was too small to definitely test for sex differences, but consistent with findings from adult GWAS\textsuperscript{17,20,55-57}; there did not seem to be notable differences between females and males. In many of our analyses, we used a $P$ value threshold that was larger than conventional genome-wide thresholds. This decision was made a priori and was intended to ensure that we did not miss potentially important associations and overlaps (between epochs and with adult GWAS findings) given the relatively modest sample size. However, we acknowledge that these findings should be treated with caution until they are replicated.

**Conclusions**

To conclude, we have identified 2 novel loci related to SBP in childhood (one at prepuberty and one during puberty), but none related to SBP postpuberty or in any age epoch for DBP in childhood, at genome-wide significance. The 2 novel SBP SNPs were specific to those epochs and did not relate to SBP in other epochs even with a higher $P$ value threshold of $P\leq5\times10^{-3}$. With this more liberal $P$ value, we identified more variants related to SBP and 2 variants related to DBP measured during puberty. Most of these were specific to the particular epoch in which they were found, though for a small number, we did find overlap with adjacent epochs and also some overlap with published adult variants. Thus, our results provide some support for age-specific associations, as well as for associations that might be present across most ages. The observed genetic associations with no previous history of association with adult BP may be true novel associations, but require further investigation and replication.

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Table 4. Comparing SBP and DBP Meta-Analyses Across Epochs in EAGLE

| Marker Name | CHR | POS   | Gene          | MAF | Allele | Prepubertal Beta (95% CI) | P Value | Direction | Pubertal Beta (95% CI) | P Value | Direction |
|-------------|-----|-------|---------------|-----|--------|--------------------------|---------|-----------|------------------------|---------|-----------|
| Systolic blood pressure | | | | | | | | | | | |
| rs13025174  | 2   | 46557999 | 0.56         | A/G | 0.075      | (0.031, 0.12)          | 9.37E-04 | ++++      | 0.054                  | (0.026, 0.082) | 1.38E-04   | ++++      |
| rs13032473  | 2   | 46559169 | 0.56         | A/G | 0.076      | (0.031, 0.121)        | 8.60E-04 | ++++      | 0.054                  | (0.026, 0.082) | 1.38E-04    | ++++      |
| rs10186089  | 2   | 49072628 | 0.83         | A/G | 0.123      | (0.056, 0.190)        | 3.07E-04 | ++++      | 0.073                  | (0.031, 0.114) | 5.45E-04    | ++++      |
| | | | | | | | | | | | |
| | | | | | | | | | | |
| Pubertal Postpubertal | | | | | | | | | | |
| rs1383450   | 8   | 56167423 | 0.87         | T/C | 0.073      | (0.033, 0.112)        | 2.85E-04 | ++++      | 0.115                  | (0.063, 0.167) | 1.47E-05   | +       |
| rs276978    | 16  | 84783954 | 0.30         | T/C | 0.073      | (0.030, 0.117)        | 9.88E-04 | ++++      | 0.098                  | (0.043, 0.153) | 5.26E-04    | +       |
| rs2293382   | 20  | 9449526  | 0.25         | C/G | 0.069      | (−0.108, −0.030)      | 5.62E-04 | −−−−−     | −0.083                  | (−0.131, −0.035) | 7.33E-04    | −        |
| rs5752974   | 22  | 28601630 | 0.96         | A/G | −0.088     | (−0.137, −0.040)      | 3.74E-04 | −−−−−     | −0.115                  | (−0.175, −0.055) | 1.79E-04    | −        |
| rs16988143  | 22  | 28704677 | 0.04         | T/C | 0.090      | (0.041, 0.138)        | 3.05E-04 | ++++      | 0.118                  | (0.057, 0.179) | 1.61E-04    | +++      |
| rs11552852  | 22  | 28754377 | 0.98         | T/C | −0.093     | (−0.142, −0.043)      | 2.68E-04 | −−−−−     | −0.123                  | (−0.185, −0.062) | 7.87E-05    | −        |
| rs16988244  | 22  | 28763373 | 0.04         | T/G | 0.091      | (0.042, 0.140)        | 2.46E-04 | −−−−−     | 0.115                  | (0.055, 0.175) | 1.71E-04    | +++      |
| rs16988333  | 22  | 28882813 | 0.04         | A/G | 0.090      | (0.041, 0.139)        | 2.84E-04 | −−−−−     | 0.110                  | (0.050, 0.170) | 3.12E-04    | +++      |
| rs5753042   | 22  | 28917972 | 0.06         | A/G | 0.088      | (0.039, 0.137)        | 3.92E-04 | −−−−−     | 0.105                  | (0.044, 0.167) | 7.64E-04    | +++      |
| Diastolic blood pressure | | | | | | | | | | | |
| rs13004438  | 2   | 179421913| 0.17         | T/C  | 0.082      | (0.034, 0.13)         | 8.31E-04 | +++       | −0.053                  | (−0.084, −0.022) | 8.10E-04    | −−−−−−−− |
| rs12542146  | 8   | 10485173 | 0.23         | A/G  | 0.100      | (0.041, 0.159)        | 9.29E-04 | ++?       | −0.073                  | (−0.115, −0.030) | 8.42E-04    | −       |
| rs9373002   | 6   | 132279972| 0.28         | A/G  | 0.101      | (0.042, 0.159)        | 7.36E-04 | ++?       | 0.072                  | (0.029, 0.114) | 8.95E-04    | +++      |
| rs294782    | 2   | 47750478 | 0.21         | A/G  | 0.075      | (0.033, 0.13)         | 4.35E-04 | +++       | 0.090                  | (0.041, 0.140) | 3.31E-04    | +++      |
| rs12532038  | 7   | 124281701| 0.39         | T/C  | −0.059     | (−0.090, −0.028)      | 2.06E-04 | −−−+      | −0.067                  | (−0.106, −0.028) | 8.25E-04    | −−−−     |
| rs734335    | 14  | 100648711| 0.40         | A/G  | −0.092     | (−0.136, −0.047)      | 5.23E-05 | −−−−−     | −0.110                  | (−0.165, −0.055) | 9.20E-05    | −−−−     |
| rs804460    | 16  | 74923996 | 0.84         | T/C  | −0.116     | (−0.164, −0.068)      | 2.10E-06 | −−−−−     | −0.099                  | (−0.157, −0.040) | 9.17E-04    | −−−−     |

Beta values are in terms of Z-scores, the number of standard deviations away from the mean. CI indicates confidence interval; DBP, diastolic blood pressure; and SBP, systolic blood pressure.

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Table 5. Comparing Systolic and Diastolic Blood Pressure Meta-Analyses in Childhood (EAGLE) and Adulthood (ICBP) for Sex-Combined Data Sets

| Time Frame   | Marker Name | CHR | POS   | Gene      | MAF        | ICBP Beta (95%CI) | PValue | EAGLE Beta (95%CI) | PValue |
|--------------|-------------|-----|-------|-----------|------------|------------------|--------|-------------------|--------|
| Systolic blood pressure | rs496146    | 17  | 78534809 | B3GNT1 | 0.3 | 0.422 (0.157,0.686) | 1.77E-03 | -0.055 (−0.086,−0.024) | 5.98E-04 |
|              | rs17810777  | 13  | 69385636 | KLHL1   | 0.11 | 0.377 (0.127,0.628) | 3.17E-03 | 0.066 (0.031,0.101) | 2.05E-04 |
| Pubertal     | rs8096788   | 18  | 65451861 | DOK6    | 0.31 | 0.65 (0.299,1.001) | 2.80E-04 | -0.095 (−0.146,−0.044) | 2.61E-04 |
|              | rs38627     | 7   | 76275543 |         | 0.27 | 0.495 (0.209,0.782) | 6.89E-04 | 0.077 (0.034,0.120) | 3.72E-04 |
|              | rs4943826   | 11  | 80730697 |         | 0.23 | 0.459 (0.163,0.755) | 2.36E-03 | 0.075 (0.032,0.118) | 5.65E-04 |
|              | rs332607    | 6   | 124765053| NKAIN2  | 0.25 | 0.317 (0.106,0.528) | 3.19E-03 | 0.054 (0.025,0.083) | 3.55E-04 |
|              | rs3824137   | 8   | 1808899 | ARHGEF10 | 0.23 | 0.358 (0.115,0.601) | 3.87E-03 | 0.058 (0.023,0.093) | 9.13E-04 |
|              | rs1771274   | 3   | 170597773| MECOM   | 0.01 | 1.404 (0.437,2.371) | 4.42E-03 | -0.400 (−0.606,−0.194) | 1.40E-04 |
|              | rs17033041  | 4   | 156610757|         | 0.14 | 0.375 (0.116,0.633) | 4.47E-03 | -0.066 (−0.105,−0.027) | 9.66E-04 |
| Postpubertal | rs293927    | 3   | 14907160 | FGD5    | 0.13 | 0.422 (0.183,0.661) | 5.42E-04 | -0.082 (−0.125,−0.039) | 2.06E-04 |
|              | rs1687304   | 3   | 14929257 | FGD5    | 0.13 | 0.404 (0.164,0.645) | 9.75E-04 | 0.089 (0.046,0.132) | 6.21E-05 |
|              | rs625757    | 1   | 33922472 | CSMD2   | 0.07 | 0.601 (0.261,0.942) | 5.43E-04 | -0.104 (−0.163,−0.045) | 4.93E-04 |
| Diastolic blood pressure | rs6760458   | 2   | 42938973 |         | 0.23 | 0.195 (0.060,0.329) | 4.63E-03 | 0.055 (0.022,0.088) | 8.67E-04 |
| Pubertal     | rs7578149   | 2   | 20175919 |         | 0.46 | 0.213 (0.087,0.339) | 9.08E-04 | -0.050 (−0.079,−0.021) | 9.83E-04 |
|              | rs11713251  | 3   | 49315011 | USP4    | 0.01 | 0.81 (0.292,1.327) | 2.19E-03 | 0.213 (0.093,0.333) | 5.03E-04 |
|              | rs7914808   | 10  | 120991173| GRK5    | 0.31 | 0.344 (0.108,0.579) | 4.20E-03 | 0.091 (0.038,0.144) | 9.07E-04 |
|              | rs1951930   | 6   | 33800633 |         | 0.15 | 0.218 (0.059,0.376) | 7.15E-03 | 0.066 (0.027,0.105) | 7.49E-04 |
| Postpubertal | rs4650457   | 1   | 80256759 |         | 0.41 | 0.205 (0.081,0.328) | 1.18E-03 | -0.063 (−0.100,−0.026) | 9.63E-04 |
|              | rs17064088  | 5   | 174293917|         | 0.05 | 0.415 (0.142,0.689) | 2.90E-03 | 0.122 (0.051,0.193) | 6.71E-04 |

Please note that the International Consortium of Blood Pressure (ICBP) released their GWAS results here: http://www.ncbi.nlm.nih.gov/projects/gap/cgi-bin/study.cgi?study_id=phs000585.v1.p1; however, the effects reported (beta) are absolute values of the regression coefficient. Beta values are in terms of Z-scores, the number of standard deviations away from the mean. EAGLE, Early Genetics and Lifecourse Epidemiology; and ICBP, International Consortium of Blood Pressure.
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Appendix

From the Department of Biostatistics & Epidemiology, Auckland University of Technology, Auckland, New Zealand (P.G.P.); Departments of Epidemiology (H.R.T., G.V., A.G.U., F.R., A.H., V.W.V.J., C.M.v.D.), Pediatrics (H.R.T., A.G.U., V.W.V.J.), and Internal Medicine (A.G.U., F.R.), Erasmus Medical Center, Rotterdam, The Netherlands; The Medical Research Council Integrative Epidemiology Unit (N.J.T., L.D.H., D.M.E., B.S.P., D.A.L.), School of Social & Community Medicine (N.J.T., L.D.H., D.A.L.), School of Oral & Dental Sciences (B.S.P.), and School of Experimental Psychology (B.S.P.), University of Bristol, Bristol, UK; Helmholtz Zentrum Muenchen, German Research Centre for Environmental Health, Institute of Epidemiology, Neuherberg, Germany (E.T., J.H.); Department of Clinical Chemistry, Fimlab Laboratories (T.L., L.-P.L.), and Department of Clinical Physiology (M.K.), University of Tampere, Tampere, Finland; Municipal Institute of Medical Research (IMIM), Barcelona, Catalonia, Spain (M.M.); Quantitative Genetics (P.A.L., M.J.W.), Genetic Epidemiology (J.B.W., N.G.M.), and Molecular Epidemiology (G.W.M.), QIMR Berghofer Medical Research Institute, Brisbane, Queensland, Australia; Research Centre of Applied & Preventive Cardiovascular Medicine (V.A.), and Department of Medicine (J.V.), University of Turku, Turku, Finland; Lunenfeld-Tanenbaum Research Institute, University of Toronto, Ontario, Canada (L.B.); University of Queensland Diamantina Institute, Translational Research Institute, Brisbane, Queensland (D.M.E.); School of Women’s & Infants’ Health (J.P.N., C.E.P.), and The Western Australian Pregnancy Cohort (Raine) Genetic Epidemiology Team, School of Medicine & Pharmacology Royal Perth Hospital Unit (L.J.B.), The University of Western Australia, Perth, Western Australia, Australia; Department of Biological Psychology, Vrije Universiteit Amsterdam, Amsterdam, the Netherlands (D.I.B., J.-J.H.); Department of Population Medicine, Harvard Medical School & Harvard Pilgrim Health Care Institute, Boston, MA (M.W.G.); Departments of Health Sciences & Genetics, University of Leicester, Leicestershire, UK (M.D.T.); Department of Clinical Physiology & Nuclear Medicine, Turku University Hospital, University of Turku, Turku, Finland (O.R.); Department de Salud Publica, Universidad Miguel Hernández, San Juan de Alicante, Spain (J.V.); Department of Epidemiology & Biostatistics, School of Public Health, Imperial College London, London, UK (M.-R.J.); Institute of Health Sciences (M.-R.J.), Biocenter (M.-R.J.), University of Oulu, Oulu; Unit of Primary Care, Oulu University Hospital, Kajaaninlaitur (M.-R.J.); Department of Children & Young People & Families, National Institute for Health & Welfare, Oulu, Finland (M.-R.J.); and Joanna Briggs Institute & School of Translational Health Science, University of Adelaide, Adelaide, Australia (L.J.P.).

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CLINICAL PERSPECTIVE

This study was designed to identify genetic variants associated with blood pressure in childhood and adolescence to improve our understanding of the lifecourse pathogenesis of hypertension and cardiovascular disease. Variation in systolic and diastolic blood pressure across the lifecourse is associated with subsequent adult risk of coronary heart disease and stroke. The age-related changes in blood pressure in industrialized countries suggest a varying gene environment interaction with age, but to date, relatively little is known about the genetic determinants of blood pressure in childhood and adolescence. Two novel loci were identified as having genome-wide associations with systolic blood pressure in specific age epochs. These loci were rs1563894 [ITGA11], associated with systolic blood pressure assessed prepuberty, and rs872256, which was associated with blood pressure during puberty. ITGA11 has been shown to be associated with hypertrophic cardiomyopathy and coronary artery disease. Rs872256 is in close proximity to the SMARCA and VLDLR genes. SMARCA is thought to influence cardiac hypertrophy and fetal gene expression. VLDLR has been shown to be associated with obesity and has plausible biological links with angiogenesis and atherosclerosis. We also found some evidence of gene clusters associated with blood pressure in childhood. Most of the effects we observed were specific to the particular epoch in which they were found, though a small number overlapped with adjacent epochs and with published adult variants. Our results suggest that genetic determinants of blood pressure act from childhood, develop over the life course, and show some evidence of age-specific effects.