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## **ORIGINAL ARTICLE**

# Age at onset of different pubertal signs in boys and girls and differential DNA methylation at age 10 and 18 years: an epigenome-wide follow-up study

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STUDY QUESTION: Is the age of onset of pubertal markers related to subsequent changes in DNA methylation (DNAm)?

**SUMMARY ANSWER:** We identified 273 cytosine-phosphate-guanine (CpG) dinucleotides in girls and 67 CpGs in boys that were related to puberty and that were replicable in two other investigations.

**WHAT IS KNOWN ALREADY:** Previously, 457 CpGs (not gender-specific) and 347 (in girls) and 50 (in boys), respectively, were found to be associated with puberty, according to investigations of studies from Denmark (20 girls and 31 boys) and North America (30 girls and 25 boys).

**STUDY DESIGN, SIZE, DURATION:** The study was based on a birth cohort of 1456 participants born in 1989/90, with follow-up at age 10 and 18 years.

**PARTICIPANTS/MATERIALS, SETTING, METHODS:** The follow-up included 470 participants with information on DNAm and age of pubertal onset (244 girls and 226 boys). Age of pubertal onset was ascertained retrospectively at age 18 years. Using the Pubertal Development Scale, both genders were asked about ages of onset of growth spurt, body hair growth and skin changes. Ages at voice deepening and growth of facial hair were inquired from boys; ages at breast development and menarche from girls. Blood samples were collected at 10 and 18 years of age. DNA was extracted using a standard salting out procedure. The methylation level for each CpG site was assessed using one of two different platforms. DNAm was measured by a ratio of intensities denoted as  $\beta$  values for each CpG site. After quality control, 349 455 CpG sites were available for analysis. M values were calculated (log2( $\beta/(1-\beta)$ ) to approximate a normal distribution, and their levels were adjusted for blood cell proportions. Linear mixed models were applied to test the association between age of pubertal markers and repeated measurement of DNAm at 10 and 18 years.

**MAIN RESULTS AND THE ROLE OF CHANCE:** In girls, a total of 63 019 CpGs statistically significantly changed after occurrence of any of the five pubertal events and 13 487 were changed subsequent to all five events: the respective number is boys were 3072 and 301. To further exclude false-positive findings, we investigated which CpGs were replicable in prior studies from Denmark or North America, resulting in 273 replicable CpG in girls and 67 CpGs in boys (236 and 68 genes, respectively). Most identified genes are known to be related to biological processes of puberty; however, genetic polymorphisms of only four of these genes were previously linked to pubertal markers in humans.

**LIMITATIONS, REASONS FOR CAUTION:** The relative age of pubertal onset to the age of DNAm measurements does not allow causal inference, since DNAm at an earlier age may have affected the pubertal age or pubertal age may have altered later DNAm. This investigation concentrates on autosomes. CpGs on X and Y chromosomes are not included in the current study.

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This is an Open Access article distributed under the terms of the Creative Commons Attribution Non-Commercial License (http://creativecommons.org/licenses/by-nc/4.0/), which permits non-commercial re-use, distribution, and reproduction in any medium, provided the original work is properly cited. For commercial re-use, please contact journals.permissions@oup.com **WIDER IMPLICATIONS OF THE FINDINGS:** Assessment of biological processes involved in pubertal transitions should include epigenetic information. Differential DNAm related to puberty needs to be investigated to determine whether it can act as an early marker for adult diseases known to be associated with puberty.

**STUDY FUNDING/COMPETING INTEREST(S):** This work was supported by NIH grants R03HD092776 (Epigenetic characterization of pubertal transitions) and R01AI121226. The 10-year follow-up of this study was funded by National Asthma Campaign, UK (Grant No 364), and the 18-year follow-up by a grant from the National Heart and Blood Institute (R01 HL082925). The authors have no conflicts to report.

Key words: puberty / epigenetics / gender / DNA methylation / birth cohort / Pubertal Development Scale

# WHAT DOES THIS MEAN FOR PATIENTS?

In girls and boys, puberty shows up with a variety of signs such as menstrual bleeding, growth spurt, facial and body hair growth, deepening of the voice and breast development. Age at puberty is also related to many medical conditions occurring later in life such as asthma, obesity, cardiovascular diseases and cancer. To identify biological indicators of puberty that could be used for early detection of such medical risks, the researchers investigated DNA-methylation.

DNA-methylation describes a change made to the DNA in genes that does not involve alterations to the DNA sequence itself (epigenetics). Methyl groups are either added to or removed from the DNA, which then alter the function of genes. The researchers explored whether genome-wide changes of DNA methylation between the age of 10 and 18 years can be linked to age of onset of different pubertal events. To this end, the researchers used a large number of children born on the Isle of Wight in the UK (244 girls and 226 boys at either age 10 or 18 years). To exclude false-positive findings, the researchers then checked which DNA-methylation changes had also been found in earlier, smaller studies from Denmark and North America.

The researchers identified 273 DNA-methylation changes in girls and 67 in boys that had also been found in earlier studies, covering 236 and 68 genes, respectively. Most of the genes identified are related to biological processes of puberty; however, in genetic studies only four of the identified genes had previously been linked to pubertal markers in humans. This novel epigenetic information will be used in future studies to investigate which of these markers are risk factors for adult medical conditions and whether they be altered in order to reduce the risk of disease.

# Introduction

Puberty is defined as a series of biological and physical transformations initiated by central activation of the hypothalamic-pituitary-gonadal axis (Dorn and Biro, 2011). Pulsatile release of GnRH signals the pituitary gland to secrete the gonadotrophins LH and FSH, and testosterone in females and males, respectively, which trigger the development of secondary sexual characteristics. Age at pubertal onset has been related to a wide variety of conditions including asthma, overweight and obesity, metabolic changes, blood pressure, cardiovascular diseases, stroke, type 2 diabetes, cancer, mortality and psychological well-being in women (Hsieh et al., 1990; de Ridder et al., 1992; van Lenthe et al., 1996; Guo et al., 1997; Remsberg et al., 2005; Varraso et al., 2005; Jacobsen et al., 2007; Mendle et al., 2007; Jacobsen et al., 2009; Lakshman et al., 2009; Baek et al., 2015; Gomes et al., 2015; Ryu et al., 2015; Chang et al., 2016; Gemelli et al., 2016; Yoo, 2016; He et al., 2017; Li et al., 2017; Platt et al., 2017; Yang et al., 2017), men (Hardy et al., 2006; Maule et al., 2012; Mendle and Ferrero, 2012; Ong et al., 2012; Platt et al., 2017) or both sexes (Hardy et al., 2006; Day et al., 2015; Mahmoud et al., 2018).

One potential factor regulating the timing of pubertal onset is DNA methylation (DNAm) (Demetriou et al., 2013; Lomniczi and Ojeda, 2016), which is considered to serve as an archive of markers of early developmental exposures, such as maternal smoking. In addition, DNAm may also change as a consequence of pubertal onset and in turn may affect adult diseases. Methylation is a dynamic process of addition of methyl groups to cytosine where cytosine–phosphate–guanine (CpG) dinucleotides occur in the DNA sequence, while demethylation occurs both actively and passively (Chen and Riggs, 2011).

Methylation of gene promoter regions is considered to modulate gene expression (Shenker and Flanagan, 2012), but methylation occurring at other genomic regions (e.g. gene body, intergenic) may also influence transcription (Jones, 2012). DNAm also regulates alternative splicing (Oberdoerffer, 2012; Lev Maor et *al.*, 2015), which results in different messenger RNAs and different biological functions (Black, 2003). In girls, precocious puberty has been found to be associated with both increased and decreased methylation levels of different genes (Bessa et *al.*, 2018).

There is a need to better understand changes in DNAm following the onset of different pubertal markers such as menarche, thelarche, growth spurt and voice deepening. Hence, a first step is to explore which CpGs can be linked with age of onset of different pubertal events. Two recent longitudinal cohort studies using epigenome-wide analyses investigated associations between puberty timing and DNAm (Almstrup et al., 2016; Thompson et al., 2018). The first study included healthy Danish children, 20 girls and 31 boys, who were examined twice including measurements of DNAm before and after achievement of pubertal Tanner state B2 (Marshall and Tanner, 1969; Marshall and Tanner, 1970). Testicular volume ( $\geq$ 4 ml) for boys and Tanner breast stage for girls (both indicating a Tanner state of B2 and higher) were utilized as a means of measuring pubertal stage for both genders separately (Almstrup et al., 2016). When boys and girls were analyzed together while adjusting for age and the different cell counts in analyzed blood samples, the Danish study identified 457 CpGs that changed as a consequence of onset of puberty with a false discovery rate (FDR) of 0.05. Associated methylation levels were either increased or decreased before and after pubertal events. A limitation of the Danish investigation is that the first measurement of DNAm was close to pubertal onset and thus methylation levels may already have been affected by the process of pubertal transition. The second study was based on 30 girls and 25 boys from North America with DNAm measured at 8 and 14 years of age (Thompson *et al.*, 2018). Puberty was assessed at both ages by Tanner staging. With separate analyses of girls and boys, the study identified 347 CpGs in girls and 50 CpGs in boys that changed as a consequence of puberty. In addition, 48 differentially methylated CpGs were found in the combined sample of boys and girls. The Thompson *et al.* (2018) study did not adjust for age at sample collection or cell composition of the samples drawn at 8 and 14 years. Thus, some DNAm changes could be related to aging but not pubertal changes. Moreover, not adjusting for cell composition, which is likely to change between 8 and 14 years, may have led to some CpGs remaining undetected.

Since age at menarche is not part of the Tanner's Sexual Maturation Scale (Marshall and Tanner, 1969; Marshall and Tanner, 1970), information on age at menarche, a frequently used indicator of puberty in girls (James et al., 2018), also in genetic studies (Elks et al., 2010; Perry et al., 2014; Willemsen and Dunger, 2016), was not assessed in the above investigations. As genome-wide associations explain only 2.7% of the variance of age at menarche (Perry et al., 2014), it is important to explore whether DNAm could be associated with age at menarche.

We previously investigated variables collected by the Pubertal Development Scale (PDS) (Petersen *et al.*, 1988) for association with child growth (Yousefi *et al.*, 2013). In this study, based on a larger birth cohort of children born on the Isle of Wight (IoW) in the UK (244 girls and 226 boys at either age 10 or 18 years), we investigated whether ages of pubertal events are associated with DNAm at 10 and 18 years. To this end, we used the age of onset of different pubertal events (Bond *et al.*, 2006). Since puberty-related processes vary among girls and boys, we conducted separate analyses for both sexes and adjusted for age and cell composition of the DNAm collected at both ages, 10 and 18 years. We identified CpGs, and their respective genes, that are associated with age of different pubertal markers. In addition, we compared our findings with those published by Almstrup *et al.* (2016) and Thompson *et al.* (2018). Finally, we identified biological pathways represented by the genes whose CpGs gained statistical significance.

## **Materials and Methods**

Data analyzed in this study were from a birth cohort composed of children born between I January 1989 and 28 February 1990 on the IoW, UK (Arshad *et al.*, 2018). The IoW is a semi-rural island located close to the British mainland. Of the 1536 children born in this period, 1456 were recruited, and 1313 participated in both follow-ups at ages 10 and 18 years. In subsamples of these children, DNAm was measured at ages 10 and 18 years.

### Assessment of puberty

The National Institute of Child and Human Development (NICHD) questionnaire from the Study of Early Child Care and Youth Development based on the PDS was used (Petersen *et al.*, 1988). At age 18 years, children self-reported their development using five markers of pubertal growth. Both genders were asked about ages of onset of growth spurt, body hair growth and skin changes. Ages at voice deepening and growth of facial hair were inquired from boys, while ages

at breast development and menarche were reported by girls. Since we asked 18-year-old participants, we focused on ages of onset which is a subpart of the five questions of the NICHD questionnaire (Yousefi et al., 2013). Ages at which pubertal events occurred were provided as years in whole numbers (i.e. age 13 years).

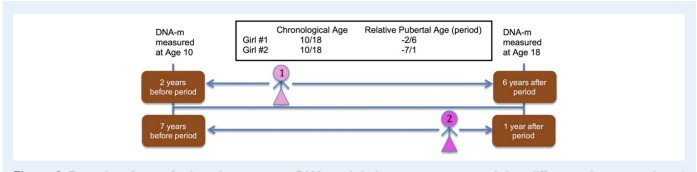
The varying ages of onset of a specific pubertal event provide unique pubertal timing information in relation to DNAm measurements. For instance, if a girl's first period occurs at age 12 years, then DNAm measurement at age 10 is 2 years before onset of menarche; DNAm measurement at 18 years is 6 years later (Fig. 1). If another girl's first period occurs at age 17 years, then DNAm measurement at age 10 and 18 is 7 years before and 1 year after, respectively (Fig. 1). Although DNAm was measured at the same chronological age, 10 and 18 years, the timing of puberty in relation to DNAm measurements of both girls is different (-2 and +6 in the first girl, -7 and +1 in the second). In our statistical analyses, we followed the approach by Almstrup et al. (2016) by using the difference in years of the age at methylation assessment and age at pubertal onset ( $\Delta$ years) for identification of differential CpG methylation. For example, in our repeated measurement analysis, -2and +6 were used as differences in time ( $\Delta$ years) between DNAm measurement and onset of specific pubertal markers (Fig. 1). Using repeated DNAm measurements and their differences to pubertal age allowed us to integrate ages at pubertal events that occurred before the first DNAm measurement at age 10 years (e.g. when breast development started at age 8 years).

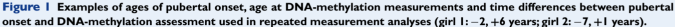
## **DNA** methylation analysis

Blood samples were collected at ages 10 and 18 years. DNA was extracted using a standard salting out procedure (Miller et al., 1988). DNA concentration was estimated by fluorometric quantitation. The methylation level for each CpG site was assessed using one of two platforms, the Illumina Infinium HumanMethylation450 BeadChip (Illumina, Inc., San Diego, CA, USA) or the Infinium MethylationEPIC BeadChip (Illumina, Inc., San Diego, CA, USA). For each sample, I µg of DNA was bisulfite-treated for cytosine to thymine conversion using the EZ-96 DNA methylation kit (Zymo Research, Irvine, CA, USA), following the manufacturer's protocol. HumanMethylation450 BeadChip interrogates >484000 CpGs and MethylationEPIC BeadChip >850000 CpGs, both associated with over 24 000 genes. Arrays were processed using a standard protocol (Bibikova and Fan, 2009) with multiple identical control samples assigned to each bisulfite conversion batch to assess array variability. DNA samples were randomly distributed on the arrays, aiming to eliminate batch effects. DNAm was measured by a ratio of intensities, denoted as  $\beta$  values for each CpG site, which is a ratio of methylated (M) over the sum of methylated and unmethylated (U) probes ( $\beta = M/[c + M + U]$ ), where c is used as a constant to prevent zero in the denominator. M values were calculated via  $\log 2(\beta/(1-\beta))$ , which better approximate a normal distribution.

#### Methylation data at age 10 and 18 years

Quality control was undertaken and quantile normalized beta values were processed following the CPACOR pipeline (Lehne et al., 2015) from a total of seven batches—two from 450K and five from the EPIC array platform (DNAm samples at 10 years) and six batches—two from 450K and four from the EPIC array platform (DNAm samples at age 18 years). These samples from different array platforms and





batches were combined, and only the shared probes between EPIC and 450K arrays were included. In both the 10- and 18-year methylation data, the R package *ComBat* (Johnson *et al.*, 2007) was used to remove the batch effect in the combined dataset. CpG sites with probe-single nucleotide polymorphisms within 10 base pairs and with a minor allele frequency greater than 0.007 in the Caucasian population (which represented ~10 subjects expected in the sample of this study cohort) were excluded. In total, 349 455 CpG sites from a total 470 participants (n = 322 at age 10 and n = 465 at age 18 years) with pubertal information were available for analysis.

#### Cell type proportions

The estimateCellCounts() function from the Minfi package (Aryee et al., 2014) using the adult reference panel (Reinius et al., 2012) and the method by Houseman et al. (2012) and Jaffe and Irizarry (2014) was utilized to estimate cell proportions for each sample at each age (10 and 18 years). White blood cell counts (CD4T, CD8T, natural killer cells, B cells, monocytes, granulocytes and eosinophils) were estimated using DNAm data of separate platforms (450 k and EPIC) and then combined.

#### **Analytical design**

Using epigenome-wide DNAm data to identify differentially methylated CpGs that were associated with the onset of the five pubertal events, we applied linear mixed models with repeated methylation measurements of individual CpGs (dependent variable) determined at age 10 (n = 322) and 18 years (n = 465). Following Almstrup's approach (Almstrup et al., 2016), the differences in years between the age of puberty onset and the age of DNAm measurements (Fig. 1) were used as independent variables ( $\Delta$ years). Analyses were run separately for girls and boys. The analysis was not restricted to paired data (DNAm measured at 10 and 18 years), since missing DNAm data at one age does not result in exclusion of observations in linear mixed models. To adjust for methylation changes that are simply due to the chronological age, the methylation of 349 455 CpGs was first regressed on age of DNAm measurement using M values. Then, the residual methylations not explained by age were regressed in the linear mixed models on differences in years ( $\Delta$ years). For each pubertal event (body hair growth, breast growth, menarche, skin changes and growth spurt in girls and body hair growth, facial hair growth, voice deepening, skin changes and growth spurt in boys), the residual methylations and  $\Delta$ years were modeled as follows:

$$\mathsf{y}_{ij} = \alpha_j + \beta * \mathsf{x}_{ij} + \rho * \mathsf{w}_{ij}, + \varepsilon_{ij},$$

where  $y_{ii}$  is the residual methylations of the *i*th individual at the *i*th time point (i.e. age 10 and 18 years) after regression out of chronological age of DNAm measurements,  $\alpha_i = \alpha_0 + U_i$  is the random intercept for each subject in the repeated measurement model with  $U_i \sim N(0, \sigma_0^2)$ ;  $x_{ii}$  represents  $\Delta$  years of the *i*th individual at the *j*th time point;  $w_{ii}$ 's are covariates such as cell proportions;  $\beta$  represents the  $\Delta$ years' effect on the residual of the DNAm for a pubertal event;  $\varepsilon_{ii}$  is the error term. To adjust for cell composition at each period (age 10 and age 18 years), cell proportions were added into the model as covariates. To prevent a large proportion of false positives (Benjamini and Hochberg, 1995) resulting from 349 455 tests and a P value on 0.05, we controlled for the FDR to correct for multiple tests. P values of the genome-wide association studies, stratified by gender, are illustrated using Manhattan plots. Volcano plots were used to show the extent of positive or negative associations between methylation of CpGs and  $\Delta$ years. Next, findings were compared with those of a Danish and a North American study (Almstrup et al., 2016; Thompson et al., 2018). Among the 453 CpGs provided by Almstrup et al. (personal communication), 389 were found among the 349 455 CpGs in the IoW cohort. Thompson et al. (2018) identified 347 CpGs in girls and 50 in boys, of which 288 for girls and 43 for boys were available for analysis in the IoW dataset. The analyses were performed using the R package (Vienna, Austria) and SAS version 9.4 (Cary, NC, USA). Finally, we investigated biological pathways for the genes of those CpGs that were replicated in either study, using the ToppFun server (http://toppgene.cchmc.org/ enrichment.jsp) (Chen et al., 2009) as a functional annotation tool. To this end, we first annotated the CpGs to their corresponding gene names by using the Infinium MethylationEPIC v1.0 B4 Manifest File. We found the nearest gene names using Snipper vI.2 (https://csg.sph. umich.edu/boehnke/snipper/) for the CpGs that had no gene names in the EPIC manifest file.

## Results

With respect to sex and socioeconomic status, there was no statistically significant difference between the 1313 offspring who were followed up at age 18 years and the 470 adolescents who had information

Variable	Total sample at age 18 years (n = 1313), %	Sample with puberty data and DNA methylation ( <i>n</i> = 470),%	P value chi-square test
Sex (girls)	50.3	51.9	0.37
Socioeconomic status of the parents			0.87
Low	14.4	15.0	
Medium	77.3	76.5	
High	8.4	8.5	
Participation at age 18 yrs			<0.0001
In person	65.8	97.7	
By telephone	32.1	2.3	
By mail	2.1	0	
Still in education at age 18 years	69.8	77.9	< 0.000

**Table I** Comparison of all participants in the Isle of Wight birth cohort at 18 years and those with information on age of pubertal onset and DNA methylation at 10 or 18 years.

Table II Age of onset of pubertal events in girls and boys.

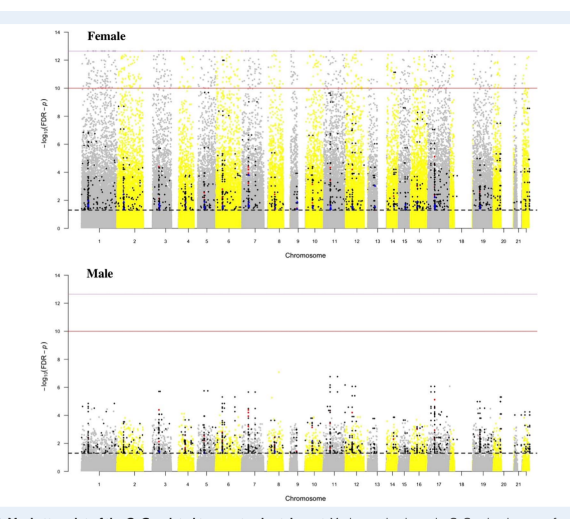
		Observations with methylation at age 10 or 18 years (n)	Mean (years)	Median (years)	SD	Minimum (years)	Maximum (years)
Girls	Growth spurt	194	12.4	12	1.82	5	16
	Body hair growth	230	12.2	12	1.45	7	16
	Skin changes	176	13.1	13	1.59	9	17
	Breast growth	230	12.3	12	1.59	8	16
	Age at menarche	237	12.3	13	1.36	9	17
Boys	Growth spurt	204	13.5	14	l.65	7	16
	Body hair growth	214	13.3	13	1.4	8	17
	Skin changes	179	13.9	14	1.3	10	18
	Deepening of the voice	218	14.1	14	1.2	10	17
	Facial hair growth	223	15.3	15	1.12	11	18

on both ages at pubertal events and DNAm at age 10 or 18 years. (Table I).

At age 18 years, puberty information was collected from subjects through in-person visits (843 of 864, 97.6% provided puberty information), telephone interviews (391/421, 92.9% with puberty information) or mailed questionnaires (27/28, 96.4% with puberty information). Of the 470 participants with pubertal and DNAm information at ages 10 or 18 years, 226 were boys and 244 were girls. In total, 322 participants had pubertal information and DNAm at age 10 years, and 465 at age 18 years. In girls, ages at menarche, breast growth and body hair growth were recalled by most participants (Table II). In boys, ages at facial hair growth and deepening of the voice were questions with a larger number of recalls. Regarding growth spurt, body hair growth and skin change, girls started about I year earlier than their male counterparts. The ages of onset of different puberty events were significantly correlated among boys and girls. All puberty variables had high internal consistency; the overall raw Cronbach's alpha coefficient was 0.80 in boys and 0.86 in girls (Yousefi et al., 2013).

We identified more CpGs that were significantly changed related to age at puberty in girls than in boys (Figs 2 and 3). Almstrup *et al.* (2016) did not provide gender-specific CpGs associated with pubertal events; hence, we included the CpGs identified by them in the Manhattan plots of both sexes (Fig. 2). For girls, Fig. 2 shows that we determined more significantly associated CpGs than the two prior studies, in particular for *P* values of less than  $e^{-10}$ . In boys, the Manhattan plot using the loW data demonstrates more significant gender-specific CpGs than Thompson (blue triangles). Since Almstrup did not separate gender-specific associations, the red dots in the Manhattan plot for boys may present CpGs related to puberty in both genders.

Separately for girl and boys, the Venn diagram (Fig. 3) characterizes CpG detected with a FDR *P* value <0.05 for each event and their overlap. In girls, a total of 63 019 CpGs changed related to any of the five pubertal ages and 13 487 were associated with all five ages. The number of CpGs that changed related to age at menarche is largest. Age at menarche also shows the largest overlap of detected CpGs with age at breast development, body hair growth and skin changes. In boys, the total and overlapping numbers of CpGs among the five pubertal

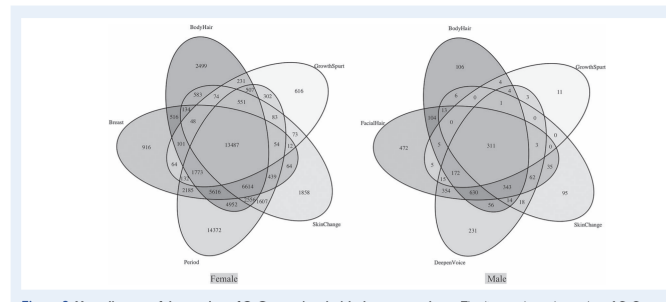


**Figure 2** Manhattan plot of the CpGs related to age at pubertal onset. Manhattan plot shows the CpGs related to ages of pubertal onset in the lsle of Wight (loW) birth cohort (gray and yellow dots) including a plot of the statistical significant CpGs determined in the loW study replicated by Almstrup *et al.* (2016) or Thompson *et al.* (2018). The black diamonds in the Manhattan plot are the CpGs identified in the loW study and replicated in either Almstrup *et al.* (2016) or Thompson *et al.* (2018). The red dots are CpGs identified as statistically significant by Almstrup *et al.* but not our study; the blue triangles were detected by Thompson *et al.* but not our study. The dashed black line represents a false discovery rate (FDR) *P* value of 0.05; the solid red line presents a *P* value of 1 e<sup>-10</sup>, and the solid violet line gives the lower limit of 2.44 e<sup>-13</sup>. The gray and yellow dots above the dashed black line characterize CpGs identified in the IoW study but not replicated by either Almstrup *et al.* or Thompson *et al.* (2016, Thompson *et al.*, 2018). Note: The negative log of adjusted *P* values on the Y-axis stretches the scale and makes it easier to visualize the significant CpGs.

events are 3072 and 311 CpGs, respectively, and are much smaller. The largest overlap in boys is seen for facial hair growth and deepening of the voice. In total, we identified more CpGs significantly related to any puberty event in girls than in boys (gray and yellow dots in Figs 2 and 3). To investigate overestimations of statistical significance due to inflation of test statistics in this epigenome-wide study, we used Q-Q plots and the genomic inflation factor (Hinrichs *et al.*, 2009). Except for age at menarche (inflation factor = 1.13), all other inflation factors for the ages at pubertal events were below 1, indicating that all our reported findings were not inflated and no adjustment was necessary. Regarding age at menarche, an inflation factor of 1.13 indicates only a minor inflation of results.

Inspecting the directions of effects and strength of the regression coefficients using volcano plots shows that there are equal numbers of positive and negative effects of puberty on methylation in boys and girls (Supplementary Fig. S1). However, there are many more CpGs in girls strongly linked to  $\Delta$ years than in boys. A positive coefficient between  $\Delta$ years and DNAm at ages 10 or 18 years means that the puberty event is related to an increase in DNAm over time. A negative coefficient describes methylation levels that are higher prior to the experience of puberty and decrease thereafter. Supplementary Figs S2 and S3 each provide four examples, two CpGs for each gender, showing an increase and decrease in DNAm over  $\Delta$ years, with and without adjustment for blood cell composition.

In total we tested 349455 CpGs measured at 10 and 18 years of age (Table III). In the analysis, we used the distance of pubertal onsets relative to 10 and 18 years ( $\Delta$ years) and residual DNAm (adjusted for age of DNA methylation measurements) and controlled for cell composition. Table III shows that the majority of statistically significant CpGs were associated with age of menarche (k = 55230) in girls and

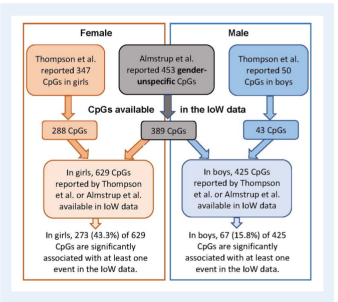


**Figure 3 Venn diagram of the number of CpGs associated with changes at puberty.** The diagram shows the number of CpGs associated with ages of changes of body hair, and skin, growth spurt, breast growth (girls), menarche (girls), facial hair (boy) and deepening of the voice (boy) at a FDR cut-off *P* value of 0.05 in boys and girls.

facial hair growth in boys (k = 2524). In girls, 18% of all CpGs were statistically significantly associated with age of at least one pubertal event and 8.8% in boys (FDR P value <0.05). In girls, 3.9% of the CpGs showed a differential methylation for all five events, and 0.089% in boys. Almstrup et al. (2016) provided 453 statistically significant CpGs (FDR < 0.05) from the combined samples of 51 boys and girls. After data cleaning, we found 389 of these CpGs in our list of 349455 CpGs. In the sample of 55 North American adolescents, Thompson et al. (2018) reported 347 statistically significant CpGs in girls and 50 in boys, of which 288 CpGs in girls and 43 CpGs in boys, respectively, are available in our IoW data as shown in the flowchart (Fig. 4). Hence, for girls, the two previous studies provided 745 statistically significant CpGs, of which 629 CpGs are available in our IoW data (288 CpGs from Thompson et al. 2018 combined with 389 CpGs genderunspecific CpGs from Almstrup et al., excluding those CpGs identified by both investigations). In boys, a total of 496 individual CpGs were detected by the two studies (Fig. 4), of which 425 CpGs are available in the IoW data (43 from Thompson et al. and 389 gender-unspecific CpGs from Almstrup et al., after excluding those CpGs identified by both investigations and those not available in the IoW data). Of these 629 and 425 in girls and boys, respectively, we detected 43.4 and 15.8% as statistically significant related to at least one pubertal event (Fig. 4).

Tables IV and V show the top 50 most significant and replicated CpGs associated with  $\Delta$ years among the five pubertal events. The respective FDR-adjusted *P* value ranges from e<sup>-13</sup> to e<sup>-5</sup> in girls and from e<sup>-4</sup> to 0.03 in boys. Supplementary Tables SI and SII provide all significant and replicated CpGs (273 CpGs in girls and 67 in boys) with details such as pubertal event and regression coefficients. In total, we identified 236 genes in girls and 68 genes in boys (at least one CpG was related to two gene names). The CpGs are almost evenly distributed across all chromosomes.

In girls, the identified 273 CpGs represent 236 genes (Supplementary Table SI). Inspecting biological processes (European Bioinformatics



**Figure 4** Flowchart of the replication approach addressing the number of available and statistically significant **CpGs in three different studies, for girls and boys separately.** The flow chart is for the present study, Thompson et al. (2018) and Almstrup et al. (2016).

Institute—https://www.ebi.ac.uk/) related to the revealed genes using as a functional annotation tool the ToppFun server (Chen *et al.*, 2009), showed I I biological processes (Supplementary Table SIII). The I I biological processes are related to growth and pubertal changes and can be grouped into five developments: bones (bone development GO:0060348, endochondral bone morphogenesis (GO:0060350), bone morphogenesis (GO:0060349)); related to adhesion (cell

Sex	Age at:	Number of significant CpGs associated with each event	Number of CpGs significantly associated with at least one pubertal event (% of 349 455 CpGs)	Number of CpGs significantly associated with all five pubertal events (% of 349 455 CpGs)	Number of significant CpGs (at least one event) that overlapped with those detected by Almstrup et al. (2016) or Thompson et al. (2018)	Number of significant CpGs (all five events) that overlapped with those detected by Almstrup et al. (2016) or Thompson et al. (2018)
Girls	Growth spurt Body hair growth Skin changes	18 108 40 242 28 237	63019 (18.0)	13 487 (3.86)	273 (43.4% of 629 CpGs detected by Almstrup or Thompson)	112 (17.8% of 629 CpGs detected by Almstrup or Thompson)
	Breast growth Menarche	32 I 55 55 230				
Boys	Growth spurt Body hair growth Skin change Deepening of the voice Facial hair growth	534 1769 901 2217 2524	3072 (8.8)	311 (0.089)	67 (15.8% of 425 CpGs detected by Almstrup or Thompson)	10 (2.3% of 425 CpGs detected by Almstrup or Thompson)

Table III Number of CpGs significantly associated with pubertal events based on analysis of 349 455 CpGs.

adhesion (GO:0007155), and biological adhesion (GO:0022610)); related to neural changes (neurogenesis (GO:0022008), neuron differentiation (GO:0030182), retina layer formation (GO:0010842), and retina development in camera-type eye (GO:0060041)); related to anatomical structure formation involved in morphogenesis (GO:0048646, which describes the initial formation of an anatomical structure from unspecified parts); and responses to progesterone (GO:0032570, which characterizes a change in activity as a result of a progesterone stimulus). The identification of these biological processes in girls provides credibility for the 273 puberty-related CpGs that were also replicated in at least one of the two other studies.

Of the genes in boys replicated by either of the previous studies (Almstrup et al., 2016; Thompson et al., 2018), functional annotation analysis showed no significant biological processes for boys using 68 genes of 67 CpGs (Supplementary Table SII). However, to inspect individual genes, we then focused on genes whose CpGs were related to at least two pubertal events (Supplementary Table SIV) [number of events] (ABI3BP [5], CBX4 [3], CNTN1 [3], DDO [3], ELOVL2 [5], GRM2 [5], HOXB7 [5], KCNAB3 [3], and LIMCH1 [2]).

# Discussion

In agreement with recent reports that epigenetic changes are crucial in the development of puberty (Lomniczi and Ojeda, 2016), we identified a large number of DNA methylation changes related to puberty. In 244 girls and 226 boys, we identified more CpGs in girls significantly linked with ages at all five pubertal events (k = 13487) than in boys (k = 311). Of these, 273 CpGs in girls and 67 CpGs in boys were also identified previously (Almstrup et al., 2016; Thompson et al., 2018). In other words, of the 629 and 425 found in girls and boys, respectively, in prior studies (Fig. 4), we detected 43.4% in girls and 15.8% in boys, as being statistically significantly related to at least one

pubertal event. Given that the sample sizes of the two prior studies were smaller (51 participants in Denmark and 55 participants in North America) compared to our cohort of 470 adolescents, a low replication proportion is not unusual. Nevertheless, the much lower proportion of replication in boys may be attributed to less obvious markers of puberty in boys than in girls (age at menarche, etc.). However, an argument against this conclusion is that of the work by Thompson et al. (2018) (30 girls and 25 boys), which is based on clinical measures independent of recall, also identified more significant CpGs in girls (347 CpGs) than in boys (50 CpGs).

Ages at pubertal events are associated with both an increase and decrease in DNAm. In addition, the strength of the identified associations is stronger in girls. The majority of CpGs were associated with age of menarche (k = 55230) in girls and facial hair growth in boys (k = 2523), and these events are not included in the Tanner stages (Marshall and Tanner, 1969; Marshall and Tanner, 1970). The large difference in the number of CpGs in boys and girls may be related to more pronounced pubertal transition in girls compared to boys (Gunnar et al., 2009) or to differential reporting of ages at puberty. Almstrup et al. (2016) analyzed data from girls (n = 22) and boys (n = 32) separately, but only found significant CpGs in boys, which may be due to the larger sample size in boys. The results of our study agree with the work by Thompson et al. (2018) (30 girls and 25 boys) that identified more significant CpGs in girls (347 CpGs) than in boys (50 CpGs).

Our statistical approach used repeated measurements of DNAm with differences between age of puberty and the respective DNAm measurement at 10 and 18 years ( $\Delta$ years) as predictors (Fig. 1). The effect of chronological age was removed, and the residuals of the methylation levels (*M* values) were used. Hence, a positive slope between the time of pubertal events relative to DNAm at ages 10 or 18 years means that the puberty event is related to an increase of DNAm over time. Vice versa, a negative slope described methylation

5.30

5.25

5.25

5.21

cg15626881

cg05584950

cg04586023

cg01750375

1.2E-07

1.5E-07

I.6E-07

1.9E-07

9.3E-06

9.5E-06

1.2E-05

1.5E-05

7

2

22

20

BMPER

ATOH8

HDAC10

WISP2

33 923 464

86012847

50 690 380

43 343 365

Body

TSS1500

TSS1500

Highest Gene name UCSC ref gene CpG Smallest P P value after Chr MapInfo **Relation to UCSC** CpG island absolute t value FDR group value adjustment 8.09 6.7E-16 5.9E-13 17 cg18369972 GAS7 9940121 Body; TSS200 cg09037630 7.99 1.3E-15 1.0E-12 6 NFKBILI 31 528 239 cg08129129 -7.71 1.3E-14 7.3E-12 14 BCLIIB 99714031 Body S Shore 2.0E-10 5 S\_Shelf cg02387618 -7.19 6.4E-13 TCF7 133 453 480 Body П 5'UTR cg05895618 7.18 7.1E-13 2.3E-10 CSRP3 19222395 cg26207239 -7.131.0E-12 3.1E-10 П BIRC2 102231936 Body 7 ST7 cg07168526 7.00 2.5E-12 9.4E-10 116797907 Body 6.94 4.0E-12 9.8E-10 П 59 95 1 663 TSS1500 cg06881914 MS4A6A 2 RNF144A 7 | 7 | 869 N\_Shore cg05087455 6.83 8.7E-12 2.0E-09 Body 15 cg10403394 6.84 8.2E-12 2.6E-09 TPMI 63 349 192 Body 22 cg03834031 6.82 8.9E-12 2.8E-09 MIRLET7BHG 46 465 717 N Shore 6.70 RIPOR2 cg09248660 2.1E-11 4.3E-09 6 24 969 848 cg08059678 2.7E-11 5.4E-09 16 VAC14 70728425 -6.66Body cg23368715 6.62 3.7E-11 7.0E-09 12 CIR 7245510 TSS1500 6 cg08486432 -6.574.9E-11 9.0E-09 ITPR3 33 598 003 Body N\_Shelf 16 VAC14 cg00259097 -6.479.6E-11 1.6E-08 70770604 Body 5'UTR: 20 cg24403644 -6.362.1E-10 3.2E-08 TOX2 42 574 624 Body; IstExon cg06740227 6.55 5.6E-11 3.3E-08 12 RASSF9 86 229 804 Body cg09437994 -6.38 1.7E-10 3.4E-08 16 CBFA2T3 89 035 1 47 Body S\_Shore 2 cg15302376 -6.19 6.1E-10 8.3E-08 DNMT3A 25 560 263 5'UTR N Shelf cg27364319 6.19 6.1E-10 8.3E-08 12 RARG 53 626 837 TSS1500 S\_Shore cg01538166 6.21 5.3E-10 8.9E-08 17 SREBFI 17743987 Island cg22460123 6.15 8.0E-10 1.0E-07 12 KRT7 52 638 294 Body N\_Shore I cg03042633 -6.09 1.1E-09 1.4E-07 SKI 2 1 8 3 3 6 5 Body L cg24681499 -6.051.4E-09 1.7E-07 FOX<sub>3</sub> 42 706 997 Body cg00476955 6.05 1.5E-09 2.2E-07 П TRPC2 3648013 Body 7 cg03193328 6.01 1.9E-09 TPSTI 65 746 852 2.2E-07 Body 10 PIP4K2A 22894054 cg14215711 6.00 2.0E-09 2.3E-07 Body 7 cg27205928 -5.864.7E-09 4.9E-07 C7orf50 1062652 Body Island cg09258479 5.84 5.2E-09 6.5E-07 I PDZKIIPI 47 655 861 TSS200 cg26580413 5.80 6.6E-09 6.9E-07 16 FTO 54 025 348 Body cg10104487 5.78 7.4E-09 7.1E-07 I TRNPI 27 329 335 N\_Shelf cg25936358 5.77 7.8E-09 9.3E-07 13 ATP4B 114301996 N\_Shore cg06437747 5.72 1.1E-08 1.2E-06 I ACTRT2 2890836 S\_Shelf cg14980020 -5.651.6E-08 1.4E-06 T DCST2 154992262 Body S\_Shelf 15 cg11442608 5.64 I.7E-08 1.8E-06 CFAP161 81 456 832 12 cg16335893 -5.45 5.1E-08 3.8E-06 MPHOSPH9 123 634 159 N\_Shore 8 cg07396746 5.43 5.6E-08 LOXL2 23 | 62 295 4. I E-06 Body 20 35 220 41 5 3'UTR cg10566581 -5.406.8E-08 4.9E-06 TGIF2 cg20770512 5.34 9.4E-08 6.3E-06 6 GFODI 13396094 Body

**Table IV** Fifty CpGs most significantly associated with  $\triangle$  years of body hair growth, breast growth, menarche, skin changes and/or growth spurt in girls.

(Continued)

S\_Shore

#### Т

Table IV Continued										
CpG	Highest absolute t value	Smallest P value	P value after FDR adjustment	Chr	Gene name	MapInfo	UCSC ref gene group	Relation to UCSC CpG island		
cg00374672	5.12	3.0E-07	I.7E-05	7	SLC12A9	100 463 416	Body	N_Shore		
cg17220749	-5.09	3.5E-07	I.9E-05	I	GALNT2	230 249 292	Body			
cg01282174	5.07	3.9E-07	2.1E-05	11	PVRLI	119630144				
cg09492074	5.23	I.7E-07	2.2E-05	15	MEGFII	66 263 624	Body			
cg22740489	5.00	5.8E-07	3.0E-05	2	LIMS3- LOC440895	110706159	Body	S_Shore		
cg18451156	-4.99	6.2E-07	3.1E-05	12	CHFR	133 424 373	Body			

\*Since these CpGs represent the overlap of association with five events, the lowest P values identified for several outcomes is shown. UCSC: University of California Santa Cruz, Chr: chromosome, FDR: false discovery rate.

levels that are related to puberty but are higher due to prior experience of puberty and which decrease thereafter. The results show that both processes are related to pubertal markers, a decrease and an increase of methylation over time (Supplementary Figs S1 and S2). The advantage is that a change in methylation was compared to a baseline prior to pubertal events. Our work demonstrates associations between changes in DNAm and various pubertal events; however, we do not claim causal associations since the effects may go in either direction.

Of the 273 replicable CpGs in girls representing 235 genes, three genes (TCF7—transcription factor 7, FTO—alpha-ketoglutarate dependent dioxygenase, UBA7—ubiquitin like modifier-activating enzyme 7) were identified in prior genetic studies and one of the 68 genes representing 67 CpGs was also found in boys (EVI5L—ecotropic viral integration site 5 like) (Young et al., 2012; Cousminer et al., 2014; Hagen et al., 2014; Ojeda and Lomniczi, 2014; Perry et al., 2014; Perry et al., 2015; Livadas and Chrousos, 2016; Avendano et al., 2017). The detection of a large number of differently methylated CpGs on various genes in girls and identification of 11 biological processes in girls but none in boys suggests that pubertal changes in girls may be related to physiologically linked developments. It is possible that pubertal processes in boys represent more isolated developmental changes (without important biological pathways).

In boys, concentrating on genes that were associated with ages of at least two pubertal events may increase their biological significance. Of the nine genes included in Supplementary Table SII, four were also identified in girls related to multiple pubertal events (ABI3BP-ABI family member 3 binding protein [5], DDO—D-aspartate oxidase [5], HOXB7—homeobox B7 [2] and KCNAB3—potassium voltage-gated channel subfamily A regulatory beta subunit 3 [5], Supplementary Table SI). Gene Ontology annotations related to ABI3BP include heparin and glycosaminoglycan binding (https://www.genecards.org/). Heparin-binding proteins (HBPs) originate from secretions of the male accessory glands; seminal plasma contains multiple proteins that bind heparin (Yadav et al., 2013). HBPs content decreased in castrated rats and was restored by testosterone replacement (Nass et al., 1990). Epigenetic regulatory actions of CBX4 (Chromobox 4) seems to be related to feminization of the zebrafish (Ribas et al., 2017). CNTN1 (Contactin 1) encodes the neuronal membrane protein Contactin I that functions as a cell adhesion molecule. It was reported that a complex including Contactin is involved in pubertal processes in female rats (Srivastava et al., 2011). DDO (D-aspartate oxidase) encodes a flavoprotein that catalyzes the oxidative deamination of D-aspartate and N-methyl D-aspartate (Setoyama and Miura, 1997). D-aspartate seems to operate differently in female and male gonads of frogs (Raucci and Di Fiore, 2011), and the N-methyl D-aspartate receptor has been suggested to play a role during sexual development of male rats (Hsu et al., 2005). ELOVL2 codes a fatty acid elongase 2; this enzyme plays a role in male fertility of Drosophila (Ng et al., 2015). Glutamate metabotropic receptor 2 encoded by GRM2 plays a role in male sexual performance (Li et al., 2013). HOXB7 encodes a nuclear protein that is involved in cell proliferation and differentiation. In mice, mutations of HOXB7 are related to a delay in urinary tract maturation (Murawski et al., 2007). We did not identify any association with puberty for the KCNAB3 (potassium voltage-gated channel subfamily a regulatory beta subunit 3) and LIMCH1 (LIM and calponin homology domains 1) genes. However, sex hormones have played a role of KCNE (potassium voltage-gated channel subfamily E) potassium channel subunits, which were discovered as hormonally regulated potassium channel components (Duffey and Bett, 2011). Hence, in boys there may be no concerted biological processes. However, a pubertal role for seven of the nine individual genes with replicated CpGs associated with more than one pubertal event cannot be excluded.

A strength of our study is the use of the PDS method to characterize age of pubertal onset. The PDS was developed as a self-report of pubertal growth status. In comparing the self-reported Sexual Maturation Scale (SMS) (developed by Tanner et al.) and the PDS, in other studies more missing data resulted from the SMS (13%) than from the PDS (4%) method (Bond et al., 2006). Given a moderate to high concordance between the PDS and physician ratings (Brooks-Gunn et al., 1987; Carskadon and Acebo, 1993) and difficulties with the acceptance of anatomical drawings in other questionnaires, the PDS was recommended for epidemiological studies (Bond et al., 2006). The PDS approach is also preferable to Tanner SMS methods (Marshall and Tanner, 1969; Marshall and Tanner, 1970) for obtaining retrospective comprehensive information on the entire pubertal period, since in SMS, the stage of puberty is assigned to the child's current age at the time of data collection. Therefore, if the assessment is not repeated during

CpG	Highest t value among five events	Smallest P value	P value after FDR adjustment	Chr	Gene name	MapInfo	UCSC ref gene group	Relation to UCSC CpG island
cg16163419	-5.70	I.2E-08	2.1E-04	3	ABI3BP	100712326	5′UTR; I stExon	
cg21899500	5.13	2.9E-07	7.9E-04	3	GRM2	51740850	TSS1500	Island
cg21213853	5.02	5.1E-07	1.0E-03	3	GRM2	51741473	5'UTR	S_Shore
cg07547765	5.19	2.1E-07	I.2E-03	17	HOXB7	46 685 448	Body	Island
cg06740227	-4.95	7.3E-07	1.2E-03	12	RASSF9	86 229 804	Body	
cg25460807	5.02	5.1E-07	1.3E-03	8	FGF17	21 908 022		S_Shelf
cg08220120	4.94	7.9E-07	I.7E-03	5	LOC389333	138 727 711	l stExon;3′UTR	Island
cg13274149	4.82	1.4E-06	1.8E-03	9	C9orf167	140 175 393	3'UTR	Island
cg16867657	5.09	3.6E-07	1.9E-03	6	ELOVL2	11044877	TSS1500	Island
cg21878650	-4.82	1.5E-06	2.2E-03	5	ADAMTS6	64 558 623	Body	
cg07164639	-4.78	1.8E-06	2.4E-03	6	DDO	110736958	TSS1500	
cg11743675	4.72	2.3E-06	2.9E-03	12	CNTNI	41 221 855	5'UTR	
cg06942979	4.71	2.5E-06	3.0E-03	22	SELM	31501382	Body	N_Shore
cg11041817	4.67	3.1E-06	3.4E-03	17	HOXB7	46 685 327	Body	Island
cg25711003	-4.65	3.3E-06	3.5E-03	3	PHLDB2; PLCXD2	111 522 050	5′UTR; Body	
cg16980380	4.71	2.5E-06	3.9E-03	17	ENDOV	78417883		Island
cg18773260	4.92	8.4E-07	4.1E-03	17	HOXB7	46 685 292	Body	Island
cg00303541	4.49	7.1E-06	4.5E-03	3	GRM2	51741280	5'UTR	Island
cg09809672	-4.50	6.8E-06	4.9E-03	Ι	EDARADD	236 557 682	TSS1500;5′UTR; IstExon	N_Shore
cg21101631	4.90	9.4E-07	5.8E-03	5	PCDHGA4; PCDHGC3;	140874572	Body	S_Shelf
cg25693132	4.40	I.IE-05	7.3E-03	3	GRM2	51740741	TSS1500	Island
cg02489552	4.36	I.3E-05	7.4E-03	19	CCDC105	15 121 531	TSS200	N_Shore
cg06783429	4.35	I.4E-05	7.5E-03	7	SNDI	127 673 038	Body	S_Shore
cg25150953	-4.34	I.4E-05	7.6E-03	4	LIMCHI	41 540 229	Body	
cg01331772	4.33	1.5E-05	7.9E-03	2		131 094 827		
cg21572722	4.38	I.2E-05	8.3E-03	6	ELOVL2	11044894	TSS1500	Island
cg15028548	-4.3 I	I.7E-05	8.4E-03	3	ABI3BP	100712322	5′UTR; I stExon	
cg09143195	-4.30	I.7E-05	8.5E-03	12	CNTNI	41 085 901	TSS1500	N_Shore
cg03131366	-4.27	2.0E-05	9.4E-03	12		2 041 707		
cg01883408	4.25	2.1E-05	1.0E-02	5	CCNI2	132 083 691	Body	Island
cg00700412	4.22	2.4E-05	1.1E-02	12		58011837		N_Shore
cg19270739	-4.28	1.9E-05	I.IE-02	Ι		I 368 846		N_Shore
cg03994651	4.15	3.3E-05	1.2E-02	19	LMTK3	49 004 677	Body	Island
cg18055623	4.16	3.2E-05	1.3E-02	19	CACNG8; MIR935	54 485 327	Body; TSS1500	Island
cg05694021	-4.15	3.4E-05	1.3E-02	12		19 699 504		
cg05697274	4.14	3.4E-05	1.3E-02	I	GALNT2	230415377	3′UTR	Island
cg27033805	-4.11	3.9E-05	I.4E-02	8	BOPI	145 499 436	Body	S_Shore
cg16387850	4.09	4.3E-05	1.5E-02	I		46 982 889		
cg13138089	4.05	5.2E-05	I.7E-02	2	ECEL1P2	233 25   770	TSS200	Island

**Table V** Fifty CpGs most significantly associated with  $\triangle$  years of body hair and facial hair growth, deepening of the voice, skin changes and/or growth spurt in boys.

(Continued)

Table V Continued

CpG	Highest t value among five events	Smallest P value	P value after FDR adjustment	Chr	Gene name	MapInfo	UCSC ref gene group	Relation to UCSC CpG island
cg06443034	4.38	I.2E-05	I.8E-02	17		78417973		Island
cg01426818	4.31	1.6E-05	2.0E-02	17		77 806 444		Island
cg13392258	3.97	7.1E-05	2.0E-02	17		77 806 190		Island
cg19022697	3.92	8.9E-05	2.0E-02	I	TTC22	55 247 140	Body	Island
cg02231404	4.15	3.3E-05	2.1E-02	20	SOX18	62 679 635	Body	Island
cg02046143	-3.95	7.9E-05	2.2E-02	11	IGSF9B	133 797 91 1	Body	N_Shelf
cg15840151	3.89	1.0E-04	2.5E-02	10	CI0orfI20	124 459 681	TSS1500	
cg21968765	-3.88	1.1E-04	2.5E-02	3	ABI3BP	100712345	TSS200	
cg11766468	4.16	3.2E-05	2.7E-02	19	EVI5L	7 923 041	Body	N_Shelf
cg22398226	3.81	1.4E-04	3.0E-02	4		155 662 217		N_Shore
cg19344626	-3.80	I.4E-04	3.1E-02	19	NWDI	16830749	TSS200	

\*Since these CpGs represent the overlap of association with five events, the lowest p-values identified for several outcomes is shown.

the whole course of puberty, it only provides a snapshot of pubertal stage at one age. SMS is particularly problematic when assessing age of onset of the growth spurt, because unless serial measurements are taken for a growth curve, it is not possible to ascertain the timing of the growth spurt. As shown in a prior study (Yousefi et al., 2013), all puberty variables had high internal consistency; the overall raw Cronbach's alpha coefficient was 0.80 in boys and 0.86 in girls. In addition, one advantage of this questionnaire-based study is the replication of epigenetic associations found with clinically assessed data (Almstrup et al., 2016; Thompson et al., 2018), since agreements found between these three approaches indicate that the detected associations do not depend on how the puberty event was measured. This is true in girls, for whom 43.4% of the CpGs identified in our analyses based on guestionnaire data were replicated in two studies that used a mixture of laboratory and phenotypic markers of puberty (Almstrup et al., 2016; Thompson et al., 2018). Regarding boys, further research needs to be done to improve the replicability of 15.8%.

A limitation of the repeated measurement model relative to age of pubertal onset is that there is no 'causal' timeline, since earlier DNAm may have affected the pubertal age or the pubertal age may have altered later DNAm. Future studies are recommended that use a baseline at an earlier age that is not close to the changes occurring at puberty. A second limitation is the focus on autosomes. We did not inspect 418 CpGs on the X and Y chromosomes. It is possible that DNAm of the sex chromosome is involved in pubertal development, which, requires additional analyses and other replication data. Hence, we are willing to support investigations into the importance of methylation on the sex chromosomes by sharing data. Our aim was to identify CpGs of genes whose methylation changed from age 10 to 18 years as a consequence of the age at which pubertal markers occurred. We did not attempt to identify DNAm changes subsequent to early or late onset of puberty. Using age of onset of different pubertal markers as continuous measures, we identified DNAm changes that continue from early to late puberty and thus represent the integrated overall effect of puberty. Future studies may also investigate whether the affected genes in women play a role in different disease risks later in life. Also, we did not attempt to replicate the DNAm measurements based on the Illumina 450 K array with other methods, such as pyrosequencing, since this array has been shown to be technically reliable in multiple other studies (Dedeurwaerder *et al.*, 2011; Roessler *et al.*, 2012; Glossop *et al.*, 2013; Ronn *et al.*, 2013; Hall *et al.*, 2014; Acevedo *et al.*, 2015; Gabriel *et al.*, 2015; Yang *et al.*, 2015). Finally, the use of nucleated blood cells and not specific tissues is a limitation. However, since there is no ethical way to collect brain or reproductive tissue samples in human, DNA methylation changes in blood cells may provide a window related to pubertal changes in other organs.

## Conclusion

In a large birth cohort study with DNA methylation measurement at 10 and 18 years of age, we identified 13 487 CpGs in girls significantly linked with ages of occurrence of all five pubertal events and 311 CpGs in boys. Of these, 273 CpGs in girls and 67 CpGs in boys had been replicated in two other previous investigations. Hence, in girls there are more genes with methylation changes as a consequence of puberty than boys. The affected genes seem to be related to biological processes in bones, adhesion, the neural system, anatomical structure formation and responses to progesterone. Of 303 genes with differential DNA methylation (replicated in other studies) related to puberty in girls and boys, only four genes were known to have genetic polymorphisms associated with puberty. Differential DNAm as a consequence of puberty needs to be investigated in terms of whether these act as mediating factors for adult diseases known to be associated with puberty.

# Supplementary data

Supplementary data are available at Human Reproduction Open online.

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# **Authors' roles**

S.C. conceived the biostatistical design of the study, analyzed and interpreted the data and drafted the manuscript. H.R., N.M., F.S. and Y.J. helped with the data analyses, interpretation and biological pathways. S.C. and Y.J. contributed to funding acquisition. H.A., H.Z. and J.W.H. supervised the data and sample collection and helped drafting the manuscript. S.E. generated the DNAm data, contributed to the interpretation of the data and revising the manuscript. W.K. manage the funding acquisition, contributed to data analyses, interpretation and editing. All authors critically revised the manuscript for important intellectual content. The manuscript has been read and approved by all authors.

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# **Conflict of interest**

The authors declare that they have no conflict of interests.

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