

Replication of genetic loci for ages at menarche and menopause in the multi-ethnic Population Architecture using Genomics and Epidemiology (PAGE) study

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STUDY QUESTION: Do genetic associations identified in genome-wide association studies (GWAS) of age at menarche (AM) and age at natural menopause (ANM) replicate in women of diverse race/ancestry from the Population Architecture using Genomics and Epidemiology (PAGE) Study?

SUMMARY ANSWER: We replicated GWAS reproductive trait single nucleotide polymorphisms (SNPs) in our European descent population and found that many SNPs were also associated with AM and ANM in populations of diverse ancestry.

WHAT IS KNOWN ALREADY: Menarche and menopause mark the reproductive lifespan in women and are important risk factors for chronic diseases including obesity, cardiovascular disease and cancer. Both events are believed to be influenced by environmental and genetic factors, and vary in populations differing by genetic ancestry and geography. Most genetic variants associated with these traits have been identified in GWAS of European-descent populations.

STUDY DESIGN, SIZE, DURATION: A total of 42 251 women of diverse ancestry from PAGE were included in cross-sectional analyses of AM and ANM.

MATERIALS, SETTING, METHODS: SNPs previously associated with ANM ($n = 5$ SNPs) and AM ($n = 3$ SNPs) in GWAS were genotyped in American Indians, African Americans, Asians, European Americans, Hispanics and Native Hawaiians. To test SNP associations with ANM or AM, we used linear regression models stratified by race/ethnicity and PAGE sub-study. Results were then combined in race-specific fixed effect meta-analyses for each outcome. For replication and generalization analyses, significance was defined at $P < 0.01$ for ANM analyses and $P < 0.017$ for AM analyses.

MAIN RESULTS AND THE ROLE OF CHANCE: We replicated findings for AM SNPs in the *LIN28B* locus and an intergenic region on 9q31 in European Americans. The *LIN28B* SNPs (rs314277 and rs314280) were also significantly associated with AM in Asians, but not in other race/ethnicity groups. Linkage disequilibrium (LD) patterns at this locus varied widely among the ancestral groups. With the exception of an intergenic SNP at 13q34, all ANM SNPs replicated in European Americans. Three were significantly associated with ANM in other race/ethnicity populations: rs2153157 (6p24.2/*SYCP2L*), rs365132 (5q35/*UIMC1*) and rs16991615 (20p12.3/*MCM8*). While rs1172822 (19q13/*BRSKI*) was not significant in the populations of non-European descent, effect sizes showed similar trends.

LIMITATIONS, REASONS FOR CAUTION: Lack of association for the GWAS SNPs in the non-European American groups may be due to differences in locus LD patterns between these groups and the European-descent populations included in the GWAS discovery studies; and in some cases, lower power may also contribute to non-significant findings.

WIDER IMPLICATIONS OF THE FINDINGS: The discovery of genetic variants associated with the reproductive traits provides an important opportunity to elucidate the biological mechanisms involved with normal variation and disorders of menarche and menopause. In this study we replicated most, but not all reported SNPs in European descent populations and examined the epidemiologic architecture of these early reported variants, describing their generalizability and effect size across differing ancestral populations. Such data will be increasingly important for prioritizing GWAS SNPs for follow-up in fine-mapping and resequencing studies, as well as in translational research.

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Key words: menopause / menarche / genome-wide association study / race/ethnicity / single nucleotide polymorphism

Introduction

Menarche and menopause define a woman's reproductive lifespan and are important risk factors for chronic disease. Menarche marks the onset of the first menstrual period and its early occurrence has been associated with behavioral and psychosocial disorders during adolescence (Hayward et al., 1997) and adverse outcomes during adulthood including type 2 diabetes, cardiovascular disease and breast, ovarian and endometrial cancers (Hsieh et al., 1990; Lakshman et al., 2008; Lakshman et al., 2009; He et al., 2010; Stockl et al., 2011; Cramer, 2012). These associations may in part be mediated by obesity in children, which is a predictor of both early menarche and adulthood obesity (Freedman et al., 2003). Ongoing decreases in age at menarche (AM) worldwide have motivated the search for factors triggering puberty, many of which are believed to be environmental (Parent et al., 2003). However, genetic factors are also important (Gajdos et al., 2010). AM heritability (h^2), which is the proportion of variation in AM explained by genetics, is relatively high with h^2 estimates ranging from 0.24 to 0.73 (Kaprio et al., 1995; Anderson et al., 2007; Gajdos et al., 2010).

Age at natural menopause (ANM) is defined as the age at cessation of natural ovarian function (not occurring as a result of surgery, i.e. bilateral oophorectomy, radiation treatment or chemotherapeutic agents). Early menopause, typically described as menopause occurrence before the age of 45 years, is associated with increased risk of osteoporosis, coronary heart disease, stroke and all-cause mortality (Kritz-Silverstein and Barrett-Connor, 1993; Cooper and Sandler, 1998; Lisabeth et al., 2009; Wellons et al., 2012) whereas late menopause, defined as menopause occurring at the age of 54 years or older, is associated with an increased risk of breast and endometrial cancers (Hsieh et al., 1990; Cramer, 2012). ANM heritability varies in the literature, ranging from 0.31 to 0.87 (Murabito et al., 2005).

Although environmental and lifestyle factors, such as smoking, have been reported to influence timing of menopause (Gold et al., 2001), in general they explain only a small proportion of the variation in ANM (van Noord et al., 1997; Voorhuis et al., 2010). Family history is one of the strongest predictors of ANM, (Morris et al., 2011) and thus motivates the search for genetic risk factors.

To date, genome-wide association studies (GWASs) have identified genetic variants associated with reproductive lifespan traits, but have been conducted almost exclusively in European-descent populations (He et al., 2009; Perry et al., 2009; Stolk et al., 2009; Sulem et al., 2009) or in limited samples of non-European women (Liu et al., 2009; Chen et al., 2012) despite the fact that AM (Freedman et al., 2002; Wu et al., 2002; Anderson et al., 2003; Gajdos et al., 2010) and ANM (Bromberger et al., 1997; Gold et al., 2001) vary widely by race/ethnicity. For example, natural menopause occurs at a median age of 51.4 years in women of European ancestry while, on average, it occurs earlier in African American and Hispanic women, and later in women of Japanese ancestry (Bromberger et al., 1997; te Velde and Pearson, 2002; Henderson et al., 2008; He and Murabito, 2012). Reasons for racial differences in reproductive traits remain uncertain, though they may be a result of genetic variation, nutritional status (Parent et al., 2003) or secular trends in anthropometric traits, such as height and weight, which also vary by race (Freedman et al., 2002). Additional research is needed to understand the genetic variation underlying reproductive traits in diverse populations. Therefore, we sought to replicate GWAS findings for AM and ANM in our large sample of European Americans and to investigate whether these associations are also consistent in women of diverse race/ethnicity, including American Indians, African Americans, Asians, Hispanics and Native Hawaiians from the Population Architecture using Genomics (PAGE) study (Matise et al., 2011).

Methods

Study participants

The PAGE study consists of a coordinating center and four study sites: (i) Causal Variants Across the Life Course (CALiCo), a consortium of epidemiologic cohorts including the Atherosclerosis Risk in Communities Study (ARIC), Coronary Artery Risk Development in Young Adults (CARDIA), Cardiovascular Health Study (CHS), the Hispanic Community Health Study/Study of Latinos (SOL) and Strong Heart Study (SHS); (ii) Epidemiologic Architecture for Genes Linked to Environment (EAGLE) accessing three National Health and Nutrition Examination Surveys (NHANES); (iii) the Multiethnic Cohort (MEC) study and (iv) the Women's Health Initiative (WHI).

The current analysis includes European American (EA), African American (AA), Hispanic, American Indian (Am Ind) and Native Hawaiian women from ARIC, CARDIA, CHS, SHS, EAGLE (NHANES III only), MEC and WHI with available genotype and AM or ANM data and who consented to use of genetic data. The PAGE cohorts have been described previously (Matise *et al.*, 2011), but briefly: ARIC is a prospective study of atherosclerotic cardiovascular disease in 15 792 men and women aged 45–64 at the baseline examination in 1987–1989 (The ARIC Investigators, 1989). ARIC recruited predominantly EA and AA individuals from four communities: Forsyth County, NC; Jackson, MI; suburban areas of Minneapolis, MN and Washington County, MD. CARDIA is a multicenter longitudinal study of the development and determinants of cardiovascular disease over time in 5115 young adults initially aged 18–30 years at the baseline examination in 1985–1986 (Hughes *et al.*, 1987; Cutter *et al.*, 1991). AA and EA adults were recruited from four US cities (Birmingham, AL; Chicago, IL; Minneapolis, MN; and Oakland, CA). The CHS is a population-based longitudinal study of risk factors for cardiovascular disease and stroke in adults 65 years of age or older, recruited at four field centers (Forsyth County, NC; Sacramento County, CA; Washington County, MD; Pittsburgh, PA) (Fried *et al.*, 1991). Overall, 5201 predominantly EA individuals were recruited in 1989–1990 from random samples of Medicare eligibility lists, followed by an additional 687 AA recruited in 1992–1993 (total $n = 5888$). The EAGLE study accessed DNA samples ($n = 7159$) from NHANES III, phase 2, collected between 1991 and 1994 (Chang *et al.*, 2009). NHANES is a US population-based, cross-sectional survey of Americans (infants to the elderly) ascertained regardless of health status, conducted by the National Center for Health Statistics at the Centers for Disease Control and Prevention. DNA samples collected for NHANES participants aged 12 years or older are linked to surveys of demographic, health and lifestyle factors as well as to data from a physical examination and laboratory measures. MEC is a population-based prospective cohort study of over 215 000 men and women in Hawaii and California aged 45–75 years at baseline (1993–1996) and primarily of five ancestries (Kolonel *et al.*, 2000). Participants eligible for the current analysis were females who provided blood samples for nested case–control studies of breast, colorectal, ovarian, endometrial or other cancers for genetic association and biomarker studies in PAGE ($n = 7216$). The SHS is a population-based study of risk factors for cardiovascular disease among Am Ind (Lee *et al.*, 1990; North *et al.*, 2003). Participating communities include seven tribes in southwestern Oklahoma (Apache, Caddo, Comanche, Delaware, Fort Sill Apache, Kiowa and Wichita), three tribes in Arizona (Gila River and Salt River Pima/Maricopa and Akchin Pima/Papago) and three tribes from South/North Dakota (Oglala Sioux, Cheyenne River Sioux and Spirit Lake Communities). The WHI is a large national health study of post-menopausal women which includes a series of clinical trials designed to assess chronic disease prevention and an observational study designed to study risk factors for chronic diseases. A total of 161 808 women

aged 50–79 years old were recruited from 40 US clinical centers between 1993 and 1998 (The Women's Health Initiative Study Group, 1998). A subset of women ($n \sim 24000$) from the WHI clinical trials and observational study were selected for genotyping in PAGE and are included in these analyses.

All studies collected self-identified race/ethnicity information by questionnaire. Additional details about the Asian and Hispanic race/ethnicity groups follow. In MEC, 100% of Asian participants were of Japanese descent. Of the WHI participants self-identifying as 'Asian or Pacific Islander', 93% provided more specific ancestry information; the majority reported Japanese ancestry, though mixed and other Asian ancestry (including Chinese, Filipino, Korean and Vietnamese) were also reported. While EAGLE included only self-identified Mexican Americans, Hispanics in WHI and MEC were predominantly Mexican American, but may also include other Latino groups. Institutional review boards at each participating study site approved all procedures, and written informed consent was obtained from all participants.

Outcomes

Similar to previous GWAS, our goal was to measure genetic associations within the normal population distributions of AM and ANM. AM was defined as the age when menstrual periods started, in years. AM was treated as a continuous variable in analyses, however, in MEC, values were recorded in categories defined by 2-year intervals, for example 15–16 years. In this case, we assigned AM as the mean value of these intervals, for example 15.5 years. Data collection in WHI was continuous except for the lower and upper categories for which women were assigned a value of 0.5 years lower or higher than the category value, respectively. For example, for the maximum category of 17 years or older, women were assigned a value of 17.5 years. In studies with continuous data for AM, women reporting extreme values (AM < 8 years or > 19 years) were excluded from the analyses.

ANM was defined as the age at cessation of regular menstrual periods. Women with hysterectomy before onset of natural menopause or who reported menopause onset as a result of surgery (bilateral oophorectomy), radiation treatment or chemotherapy and women reporting ANM < 40 years or > 60 years were excluded from these analyses. In addition, in WHI and CHS, women who reported use of hormone replacement before the onset of natural menopause were also excluded. In MEC, ANM was collected as a categorical variable. As described for AM, the mean value of the category interval was assigned for ANM in MEC.

Genotyping

Using a Catalog of Published Genome-Wide Association Studies (Hindorf *et al.*, 2009) (www.genome.gov/gwastudies), we selected SNPs associated with AM and ANM in prior GWAS (as of February 2010) for genotyping in the PAGE study. Thus, SNPs reported in recent GWAS of these traits (Elks *et al.*, 2010; Stolk *et al.*, 2012) are not included. Some reported SNPs were in high linkage disequilibrium (LD) in the studied populations, in which case, we chose only one SNP. In total, we investigated three menarche SNPs (rs314277, rs314280 and rs7861820) reflecting two loci and five menopause SNPs (rs1172822, rs16991615, rs2153157, rs365132 and rs7333181) reflecting five loci. Study-specific DNA extraction, genotyping methods and quality control are described in detail elsewhere (Matise *et al.*, 2011). As reported previously (Matise *et al.*, 2011), principal components of continental ancestry (African, European and Asian) were derived from a panel of ancestry informative genetic markers (Kosoy *et al.*, 2009) or from existing genome-wide SNP data in each PAGE study separately. Genetic ancestry information was not available for the EAGLE samples.

Statistical analysis

SNPs were coded assuming an additive mode of inheritance. Race/ethnicity-specific linear regression models were used to test for SNP associations with the continuous outcomes of AM or ANM. Both AM and ANM models were adjusted for principal components of ancestry to control for potential confounding by population stratification, and also for other cohort-specific covariates, such as enrollment site, as appropriate. In addition, ANM models were adjusted for birth year. In sensitivity analyses, we investigated adjustment for birth year and BMI in AM models to consider potential SNP associations (controlling for BMI) in which adult BMI serves as a weak proxy of childhood BMI. Tests of association were performed separately in each of the participating cohorts and then combined in race-specific fixed effect meta-analyses for each outcome. Of note, data for Native Hawaiians and for Am Ind (AM outcome) were only available in one study, MEC or WHI respectively, and thus did not undergo meta-analysis. All meta-analyses were performed using inverse-variance weighted models in METAL (Willer et al., 2010), and between-study effect size consistency was assessed using the Q -test (χ^2 P -value) and the I^2 metric (Higgins and Thompson, 2002), where low I^2 suggests that there is little between-study variability that cannot be explained by chance. Higgins et al. (2003) suggest the classification of low and high I^2 at 25 and 75%, respectively. For SNPs with $I^2 \geq 60\%$ or χ^2 P -value < 0.05 , we present study-specific results. For replication and

generalization analyses, significance was defined at $P < 0.017$ and $P < 0.010$ to account for multiple tests ($n = 3$ and 5) for each of the AM and ANM outcomes, respectively.

MEC and WHI samples included large numbers of cancer cases and controls. Because female-related cancers, such as breast cancer, may be related to our outcomes, in sensitivity analyses we tested for interaction by cancer case/control status in the MEC and WHI samples. Additionally in WHI, we compared the cancer case/control sample results with the large portion of the sample not selected with respect to cancer status. No significant differences in SNP associations by cancer case/control status (data not shown) were found in either study and thus cancer samples were retained in the analyses.

Using Quanto (Gauderman and Morrison, 2006), we estimated the statistical power for each SNP in each racial/ethnic population (Supplementary data, Table S1) to detect the previously reported effect size based on the European descent discovery population.

Results

Age at menarche

The mean AM by race/ethnicity ranged from 12.5 to 13.2 years (Table 1). We replicated the AM association for the intergenic 9q31

Table 1 Baseline characteristics of women in the age at menarche analyses.

PAGE cohort	ARIC	CARDIA	EAGLE	MEC	WHI	Total
EA (n)	4368	898	1546	2121	16 091	25 024
AM (in years)	12.9 ± 1.6	12.7 ± 1.4	12.8 ± 1.7	13.0 ± 1.6	12.6 ± 1.5	
Enrollment (age in years)	53.9 ± 5.7	25.5 ± 3.4	50.2 ± 22.1	59.7 ± 8.4	64.3 ± 7.0	
BMI (in kg/m ²)	26.5 ± 5.4	23.1 ± 4.4	26.3 ± 6.1	25.7 ± 5.4	28.0 ± 5.8	
AA (n)	2333	556	1179	2123	2350	8541
AM (in years)	12.9 ± 1.7	12.5 ± 1.6	12.6 ± 1.9	13.2 ± 1.7	12.6 ± 1.7	
Enrollment (age in years)	53.3 ± 5.7	24.5 ± 3.9	35.6 ± 17.7	60.5 ± 8.8	61.4 ± 7.0	
BMI (in kg/m ²)	30.8 ± 6.5	25.9 ± 6.5	28.3 ± 7.4	29.0 ± 5.9	31.6 ± 6.8	
Asians (n)	n/a	n/a	n/a	2650	1297	3947
AM (in years)				13.1 ± 1.7	12.7 ± 1.6	
Enrollment (age in years)				59.7 ± 8.2	62.1 ± 7.3	
BMI (in kg/m ²)				24.0 ± 4.0	25.5 ± 5.0	
Hispanics (n)	n/a	n/a	1026	1693	1053	3772
AM (in years)			12.6 ± 1.7	13.1 ± 1.7	12.7 ± 1.6	
Enrollment (age in years)			36.5 ± 18.3	59.3 ± 7.4	60.4 ± 6.8	
BMI (in kg/m ²)			27.5 ± 6.2	28.4 ± 5.7	29.2 ± 6.2	
Am Indians (n)	n/a	n/a	n/a	n/a	233	233
AM (in years)					12.8 ± 1.8	
Enrollment (age in years)					62.0 ± 7.2	
BMI (in kg/m ²)					30.3 ± 6.1	
N Hawaiians (n)	n/a	n/a	n/a	734	n/a	734
AM (in years)				12.8 ± 1.7		
Enrollment age (in years)				56.2 ± 7.5		
BMI (in kg/m ²)				29.3 ± 6.9		

PAGE, population architecture using genomics and epidemiology study; AM, age at menarche; EA, European Americans; AA, African Americans; Am Indians, American Indians; N Hawaiians, Native Hawaiians; ARIC: atherosclerosis risk in communities study; CARDIA, coronary artery risk development in young adults; EAGLE, epidemiologic architecture for genes linked to environment; MEC, multiethnic cohort study; WHI, women's health initiative. Characteristics are presented as the mean ± SD unless otherwise indicated.

SNP, rs7861820, in our European descent sample (Table II; $P = 1.5E-6$). Betas were similar (same direction) in Asians, Hispanics and Native Hawaiians, but were not significant. We also tested two SNPs located in the 6q21 region, rs314277 and rs314280, for their associations with AM. Rs314280, located upstream of *LIN28B*, was significantly associated with AM in EA ($P = 2E-08$) and Asians ($P = 0.001$). In AA, the effect size was comparable, but not significant ($P = 0.16$). Effect sizes for this SNP were larger, but also not significant in Native Hawaiians ($P = 0.02$) and Am Ind ($P = 0.15$); in Hispanics, the association was also null ($P = 0.98$). Findings for rs314277, located in intron 2 of the *LIN28B* gene, also varied by race/ethnicity. The SNP was significantly associated with AM in EA ($P = 0.001$) and Native Hawaiians ($P = 0.016$). In EA, Hispanics and Asians, effect sizes were of similar magnitude to those reported by He *et al.* (2009) in that each additional A allele was associated with an $\sim 1-2$ month(s) increase in AM. These *LIN28B* SNPs are not strongly correlated in the PAGE populations, with r^2 ranging from 0.09 to 0.26.

In conditional analyses including both *LIN28B* SNPs, we found that adjustment of intronic rs314277 for rs314280 substantially attenuates the association in EA and Asians, both reducing the effect estimate and also rendering rs314277 insignificant in the model (Table III). However, adjustment of rs314280 with rs314277 only slightly attenuates the association of rs314280 with AM in these groups. Similar, but weaker findings are seen in the smaller sample of Native Hawaiians. In Hispanics, associations are stronger for rs314277 than rs314280 in both models. Allele frequencies for both SNPs varied widely by ancestral group in PAGE. For example, the T allele frequency for rs314280 was 0.75 in AA and 0.30 in Asians. We also observed population variation in the LD (r^2) between these SNPs and in the region (not shown).

In sensitivity analyses, adjustment of AM models for year of birth and BMI at study enrollment had only small effects on results (Supplementary data, Table SII).

Age at natural menopause

Mean ANM ranged from 48.0 to 51.1 years in the PAGE populations (Table IV). All five ANM loci, except for rs7333181 (13q34), replicated in EA and of those replicating, three SNPs were significantly associated with ANM in the other race/ethnicity groups: rs2153157 (6p24.2/*SYCP2L*), rs365132 (5q35/*UIMC1*) and rs16991615 (20p12.3/*MCM8*) (Table V). We observed significant heterogeneity ($P < 0.05$) for the synonymous *UIMC1* SNP rs365132 in EA, which was associated with ~ 4.8 months later ANM (on average) in MEC and WHI, but was not significant in CHS or ARIC. Between-study heterogeneity ($I^2 = 74$) was also seen for the smaller Am Ind sample, though the study-specific results were not significant. While rs1172822 (19q13/*BRSK1*) was not significant in the non-EA groups, effect sizes in all groups (except AA) were comparable to the EA estimates. In AA, we observed between-study heterogeneity, though results for all studies were insignificant. The coded allele frequency (CAF) for this SNP was notably different in the Asian population (average CAF = 0.05) compared with 0.25–0.36 in other populations. The most robust associations were found for rs16991615 (20p12.3/*MCM8*) in which we replicated the association in EA ($P = 8E-15$), as well as observed significant associations in Hispanics ($P = 0.0004$) and Am Ind ($P = 0.001$). Notably, in the Am Ind population, each additional copy of the A allele was associated with a 2.6 ± 0.8 year increase in menopause age.

Table II Age at menarche SNP association results in women of diverse ancestry.

SNP, locus, coded/other allele	GWAS reference study ^a		PAGE study results for AM					
	CAF	Beta \pm SE (in years)	Race/ethnicity	n	Ave CAF	^b Beta \pm SE (years)	P-value	I^2
rs314277 <i>LIN28B</i> (intron2) 6q21 A/C	0.14	0.16 \pm 0.02	EA	24 932	0.15	0.07 \pm 0.02	0.0001	20
			AA	7994	0.38	0.002 \pm 0.03	0.9589	52
			Asians	3929	0.05	0.18 \pm 0.08	0.0304	0
			Hispanics	3743	0.11	0.09 \pm 0.06	0.1482	0
			Am Indians	233	0.17	-0.07 \pm 0.24	0.7757	n/a
			N Hawaiians	729	0.06	0.47 \pm 0.20	0.0160	n/a
rs314280 <i>LIN28B</i> (upstream) 6q21 T/C	0.45	0.09 \pm 0.02	EA	24 819	0.45	0.08 \pm 0.01	2.0E-08	0
			AA	7999	0.75	0.05 \pm 0.03	0.1551	30
			Asians	3934	0.30	0.13 \pm 0.04	0.0012	60
			Hispanics	3751	0.37	0.001 \pm 0.04	0.9843	0
			Am Indians	231	0.42	0.25 \pm 0.17	0.1446	n/a
			N Hawaiians	728	0.39	0.22 \pm 0.10	0.0218	n/a
rs7861820 intergenic 9q31.2 C/T	0.48	-0.09 \pm 0.02	EA	23 371	0.49	-0.07 \pm 0.01	1.5E-06	0
			AA	6830	0.89	0.02 \pm 0.05	0.6461	0
			Asians	3920	0.78	-0.04 \pm 0.05	0.4370	42
			Hispanics	2734	0.49	-0.09 \pm 0.05	0.0643	0
			Am Indians	232	0.50	0.12 \pm 0.17	0.4854	n/a
			N Hawaiians	726	0.55	-0.09 \pm 0.09	0.3123	n/a

Ref, reference; CAF, coded allele frequency; Ave, average.

P-values significant after multiple testing correction for three tests are given in bold.

^aReference genome-wide association study (GWAS) of AM described for all SNPs is He *et al.* (2009).

^bBeta reflects change in AM in years for each additional copy of the coded allele.

Table III *LIN28B* SNPs conditional model results in the PAGE Study.^a

Race/ethnicity group	n	LD between SNPs (r^2) ^b	SNP	Single SNP model meta-analysis results		Conditional model ^c meta-analysis results	
				Beta \pm SE	P-value	Beta \pm SE	P-value
EA	24 720	0.21	rs314277	0.073 \pm 0.019	1.20e-4	0.030 \pm 0.021	0.1587
			rs314280	0.077 \pm 0.014	1.46e-8	0.067 \pm 0.015	1.10e-5
AA	7974	0.21	rs314277	-0.001 \pm 0.030	0.9871	-0.021 \pm 0.031	0.4968
			rs314280	0.044 \pm 0.032	0.1610	0.055 \pm 0.035	0.1177
Asians	3942	0.13	rs314277	0.185 \pm 0.083	0.0256	0.095 \pm 0.089	0.2851
			rs314280	0.133 \pm 0.041	0.0012	0.114 \pm 0.044	0.0106
Hispanics	3722	0.23	rs314277	0.110 \pm 0.062	0.0781	0.137 \pm 0.070	0.0501
			rs314280	0.003 \pm 0.040	0.9416	-0.038 \pm 0.045	0.4067
Am Indians	231	0.26	rs314277	-0.049 \pm 0.240	0.8372	-0.363 \pm 0.283	0.1993
			rs314280	0.253 \pm 0.174	0.1446	0.328 \pm 0.202	0.1034
N Hawaiians	735	0.09	rs314277	0.470 \pm 0.197	0.0168	0.379 \pm 0.205	0.0650
			rs314280	0.206 \pm 0.095	0.0301	0.154 \pm 0.099	0.1214

LD, linkage disequilibrium.

^aOnly women with data for both SNPs are included in conditional analyses and so numbers may differ from Table I.

^bLD estimates based on the study with the largest sample size within a given race/ethnicity group.

^cConditional model includes both rs314277 and rs314280 using coded alleles as described previously.

Discussion

Insights into the genetic basis of the reproductive lifespan may provide a better understanding of mechanisms related to puberty and menopause and their variation in diverse populations. In this study, we evaluated associations of known genetic variants with normal variation in AM and ANM in diverse race/ethnicity populations. We successfully replicated several GWAS findings for menarche (*LIN28B* and 9q31.2) and menopause (*BRSK1*, *MCM8*, *SYCP2L* and *UIMC1*) in European descent participants and found that several, but not all, loci also generalized to diverse race/ethnicity groups. In general, SNPs that replicated in our EA population had effect sizes similar to the original reports; SNPs generalizing to other race/ethnicity groups had comparable effect sizes although were less precise, due to smaller sample sizes.

Recent GWAS have identified the 6q21/*LIN28B* locus association with AM in European descent populations (He et al., 2009; Ong et al., 2009; Perry et al., 2009; Sulem et al., 2009; Elks et al., 2010). We tested two *LIN28B* SNPs: rs314277 located in intron 2 and rs314280 in the regulatory region rich with transcription factor-binding sites (Ernst et al., 2011) upstream of *LIN28B*. The lin-28 homolog B (*LIN28B*) gene is an important member of the Lin28/let-7 axis, which plays a key role in growth, proliferation and metabolism at the cellular level, as well as in development, insulin sensitivity and oncogenesis in whole organisms (Thornton and Gregory, 2012). In humans, *LIN28B* has also been associated with body height (Gudbjartsson et al., 2008; Lettre et al., 2008), which is consistent with the observation that rapid height and weight increases precede puberty (Parent et al., 2003). Interestingly, Ong et al. (2009) identified a *LIN28B* SNP rs314276 (not tested in PAGE) associated with earlier menarche, faster pre-pubertal growth spurt in girls and shorter adult height in women; however, in their analysis of European women, rs314277 was not significantly associated with AM. The *LIN28B* SNPs replicated (at $P < 0.017$) in our EA, Asian and Native Hawaiian

populations. Our effect sizes for these SNPs (betas = 0.07–0.13) were comparable to previous reports (He et al., 2009; Sulem et al., 2009). However, findings for these SNPs in AA, Hispanics and Am Ind were null, in spite of reasonable power to detect the reported association in AA (>90%) and Hispanics (>71%). Rs314280 is in moderately high LD ($r^2 = 0.63$ –0.96) in HapMap CEU and Asian (CHB + JPT) populations with other intronic and 5' region *LIN28B* SNPs, including rs314276 which was also associated with AM (Ong et al., 2009) as described above, rs167359 and rs314274, but is in lower LD with these SNPs in HapMap3 Mexican Americans ($r^2 = 0.44$ –0.48) and AA ($r^2 = 0.18$ –0.30). Our conditional analyses of the *LIN28B* SNPs suggest that the AM association for intronic SNP rs314277 does not appear to be independent from rs314280. Also, adjustment of rs314280 for rs314277 attenuates its association with AM, though it is still significant, suggesting that the regulatory region SNP rs314280 is more likely to be in high LD with the causal variant(s). However, this observation does not appear to be consistent for PAGE Hispanics. Overall, neither of the *LIN28B* SNPs appear to have strong and consistent effects across all populations; fine mapping of this region may help to identify common causal variant(s).

The lack of association for the *LIN28B* SNPs that we observed in AA (in spite of high power) is consistent with data suggesting that LD extends over significantly shorter distances in African populations than in non-African populations (Reich et al., 2001). Additionally many GWAS SNPs, such as the ones we studied, tend to be proxies of untyped functional variants, due to typical chip design (Bhangale et al., 2008). Thus, the lack of generalization to AA may be due to differential LD with the actual causal SNP or, potentially, differential genetic background effects in AA as described in Carlson et al. (in revision 2012).

In European descent women, we replicated all ANM loci reported in two recent GWAS (He et al., 2009; Stolk et al., 2009) with the exception of one intergenic SNP at 13q34. This SNP was also not replicated in a more recent GWAS published in 2012 (Stolk et al., 2012),

Table IV Baseline characteristics of women in the age at natural menopause analyses.

PAGE Cohort	ARIC	CHS	MEC	SHS	WHI	Total
EA (n)	1628	959	1037	n/a	7563	11 187
ANM (in years)	48.4 ± 3.7	49.3 ± 4.3	50.7 ± 3.1		50.5 ± 4.0	
Enrollment age (in years)	57.7 ± 4.3	72.4 ± 5.5	61.4 ± 7.4		65.0 ± 6.8	
BMI (in kg/m ²)	26.5 ± 5.2	26.0 ± 4.8	25.3 ± 5.2		28.0 ± 5.9	
Current smoker (n (%))	385 (23.7)	110 (11.5)	126 (12.2)		659 (8.7)	
AA (n)	626	196	735	n/a	876	2433
ANM (in years)	48.2 ± 3.9	48.9 ± 4.7	50.6 ± 3.2		50.2 ± 4.4	
Enrollment age (in years)	57.3 ± 4.5	73.5 ± 5.9	62.8 ± 7.3		61.9 ± 6.8	
BMI (in kg/m ²)	31.0 ± 6.7	29.2 ± 6.1	28.8 ± 5.9		31.2 ± 6.8	
Current smoker [n (%)]	146 (23.3)	22 (11.2)	121 (16.5)		115 (13.1)	
Asians (n)	n/a	n/a	1317	n/a	707	2024
ANM (in years)			51.1 ± 3.0		50.3 ± 4.0	
Enrollment age (in years)			62.1 ± 6.7		62.9 ± 7.2	
BMI (in kg/m ²)			23.7 ± 3.8		25.2 ± 5.0	
Current smoker [n (%)]			88 (6.7)		38 (5.4)	
Hispanics (n)	n/a	n/a	737	n/a	420	1157
ANM (in years)			50.5 ± 3.0		49.7 ± 3.9	
Enrollment age (in years)			60.3 ± 6.5		60.5 ± 6.9	
BMI (in kg/m ²)			28.2 ± 5.5		29.2 ± 6.2	
Current smoker [n (%)]			79 (10.7)		33 (7.9)	
Am Indians (n)	n/a	n/a	n/a	861	76	937
ANM (in years)				48.0 ± 4.2	49.8 ± 4.2	
Enrollment age (in years)				57.6 ± 7.6	61.6 ± 7.2	
BMI (in kg/m ²)				31.6 ± 6.9	29.6 ± 6.5	
Current smoker [n (%)]				225 (27.0)	7 (9.2)	
N Hawaiians (n)	n/a	n/a	300	n/a	n/a	300
ANM (in years)			50.8 ± 3.1			
Enrollment age (in years)			58.7 ± 6.7			
BMI (in kg/m ²)			28.2 ± 6.4			
Current smoker [n (%)]			55 (18.3)			

ANM, age at natural menopause; CHS, cardiovascular health study; SHS, strong heart study. Characteristics are presented as the mean ± SD unless otherwise indicated.

which included ~2500 PAGE samples of European descent from ARIC and CHS. In a sensitivity analysis, exclusion of these 2500 samples from our total 11, 150 ANM samples had little effect on ANM results, with the exception of the rs365132, which became more significant ($P = 8.0E-12$) and no longer showed evidence of heterogeneity in the meta-analysis (Supplementary data, Table SIII). In spite of low allele frequencies, we observed robust associations in diverse ancestral groups for a non-synonymous SNP, rs16991615, in the mini-chromosome maintenance 8 (*MCM8*) gene. The protein encoded by *MCM8* is believed to be important in the cell division cycle; however, rs16991615 is not predicted by the sorting tolerant from intolerant algorithm (SIFT) (Kumar *et al.*, 2009) or PolyPhen2 (Adzhubei *et al.*, 2010) to be damaging to the protein. This locus also was recently replicated in a study of Hispanic women ($n = 3468$), which included about 370 women from PAGE (Chen *et al.*, 2012). Three of the five ANM SNPs generalized to diverse ancestral

groups, and findings for the intronic SNP rs1172822 in *BRSK1* showed similar trends in all ancestral groups, but were not statistically significant in the non-European descent populations. Samples sizes were smaller in the ANM than AM analyses, mainly because of exclusions for non-natural menopause.

Some limitations of our study deserve mention. We relied on self-report and recall of ages at menarche and menopause, requiring women to report events that in some cases occurred many years ago. However, recall of AM or age at menopause (natural or induced) has been shown to be fairly reproducible in mainly European descent populations with high overall validity (± 1 year) compared with recorded age at the time of occurrence (Bean *et al.*, 1979; Colditz *et al.*, 1987; den Tonkelaar, 1997). Importantly, we would not expect any potential recall bias to be differential with respect to the studied genotypes. Our study is unique in describing genetic associations in large numbers of women differing by race/ethnicity. Yet,

Table V Age at natural menopause SNP association results in women of diverse ancestry.

SNP, locus, coded/ other allele	GWAS reference study			PAGE study results for ANM						
	Ref	CAF	aBeta ± SE (in years)	Race/ ethnicity	n	Ave CAF	^a Beta ± SE (in years)	P-value	I ²	
rs365132 <i>UIMCI</i> (syn) 5q35.2 T/G	b	0.49	0.39 ± 0.052	^d EA	11 147					59
				ARIC	1625	0.50	0.01 ± 0.13	0.9602		
				CHS	954	0.51	0.09 ± 0.19	0.6261		
				MEC	1007	0.51	0.38 ± 0.14	0.0069		
				WHI	7561	0.50	0.40 ± 0.09	8.3E-5		
				AA	2237	0.79	0.46 ± 0.14	0.0011	0	
				Asians	2008	0.48	0.22 ± 0.10	0.0281	0	
				Hispanics	1145	0.42	0.25 ± 0.14	0.0904	37	
				^d Am Indians	929				74	
				SHS	853	0.27	0.31 ± 0.23	0.1867		
				WHI	76	0.46	-1.51 ± 0.87	0.0920		
N Hawaiians	299	0.57	-0.03 ± 0.25	0.9070	n/a					
rs2153157 <i>SYCP2L</i> (intron) 6p24.2 T/C	b	0.49	0.29 ± 0.052	EA	11 161	0.49	0.17 ± 0.05	0.0007	0	
				AA	2102	0.72	0.02 ± 0.13	0.9006	0	
				Asians	2018	0.69	0.25 ± 0.11	0.0262	0	
				Hispanics	1152	0.42	0.31 ± 0.14	0.0241	0	
				Am Indians	933	0.24	0.64 ± 0.24	0.0072	0	
				N Hawaiians	297	0.50	0.04 ± 0.26	0.8744	n/a	
rs7333181 intergenic 13q34 A/G	c	0.12	0.52 ± 0.093	EA	10 110	0.13	0.06 ± 0.08	0.4677	0	
				AA	1566	0.10	0.41 ± 0.27	0.1277	0	
				Asians	707	0.04	0.17 ± 0.56	0.7680	0	
				Hispanics	419	0.09	-0.06 ± 0.49	0.9042	0	
				Am Indians	936	0.05	0.08 ± 0.57	0.8839	0	
				N Hawaiians	n/a					
rs1172822 <i>BRSK1</i> (intron) 19q13.42 T/C	b,c	0.39	-0.39 ± 0.060	EA	11 146	0.36	-0.36 ± 0.05	2.3E-11	0	
				^d AA	2230				60	
				ARIC	539	0.27	0.32 ± 0.26	0.2190		
				CHS	144	0.29	-0.31 ± 0.63	0.6238		
				MEC	673	0.27	0.01 ± 0.20	0.9574		
				WHI	874	0.28	-0.69 ± 0.33	0.0990		
				Asians	2021	0.05	-0.29 ± 0.23	0.2064	0	
				Hispanics	1152	0.33	-0.27 ± 0.14	0.0560	0	
				Am Indians	924	0.33	-0.27 ± 0.21	0.1951	n/a	
				N Hawaiians	300	0.25	-0.41 ± 0.29	0.1556	n/a	
				rs16991615 <i>MCM8</i> (non-syn) 20p12.3 A/G	b	0.06	1.07 ± 0.110	EA	10 955	0.07
AA	2129	0.02	0.57 ± 0.47					0.2276	33	
Asians	1978	0.001	1.25 ± 1.72					0.4690	0	
Hispanics	1143	0.06	1.05 ± 0.30					0.0004	0	
Am Indians	934	0.02	2.63 ± 0.80					0.0010	0	
N Hawaiians	294	0.03	-0.41 ± 0.78					0.6006	n/a	

Ref, reference. P-values significant after multiple testing correction for five tests are given in bold.

^aBeta reflects change in the ANM in years for each additional copy of the coded allele.

^bReference GWAS of ANM described is He et al. (2009).

^cReference GWAS of ANM described is Stolk et al. (2009).

^dDue to high between-study heterogeneity ($I^2 > 60\%$ and/or χ^2 P-value < 0.05), study-specific results are reported separately.

we acknowledge that some heterogeneity within our specified race/ethnicity groups may exist. We also observed moderate-to-high between-study heterogeneity for a few SNPs; in particular, the ANM analyses of rs365132 showed significant between-study heterogeneity in EA and Am Ind, indicating that the meta-analysis effect size is not a good summary of the study-specific estimates. In EA, the SNP was significantly associated with ANM in MEC and WHI, but not in ARIC or CHS. Reasons for this heterogeneity are not clear; the SNP remained significant regardless of cancer status in WHI and MEC (in sensitivity

analyses). In Am Ind, coded allele frequencies notably differed between WHI and SHS, as did effect sizes. However, the WHI sample size is small and although the effect size in SHS was comparable to the other groups, the SNP did not replicate in either study and thus the interpretation (null result) remains the same.

A final limitation relates to study power to replicate previously reported findings in the non-European populations. While power for EA was excellent ($> 99\%$ for all SNPs), we had low power to replicate findings in Am Ind or Native Hawaiians (power for all SNPs was

<60%) (Supplementary data, Table S1). Power for AA ranged from very high for the AM *LIN28B* SNPs (>90%) to modest for other SNPs, ranging from 56 to 83%. Hispanics had modest power for AM SNPs (60–82%) and more variable power for ANM SNPs (19–86%). Similarly, power for Asians varied highly by SNP (8–87%) due to varying allele frequencies and sample sizes. Overall, low power for some of the SNP-race/ethnicity group analyses may have limited our ability to replicate previously published GWAS findings, which also may be inflated due to ascertainment bias, termed the winner's curse (Zollner and Pritchard, 2007).

In spite of these limitations, our study is unique in describing genetic associations with important markers of the reproductive lifespan in diverse US racial/ethnic groups. To our knowledge, such data on Am Ind and Native Hawaiians have not been presented previously and also are very limited in US Hispanics and AA (Chen *et al.*, 2012; Dvornyk and Waqar-ul-Haq, 2012; He and Murabito, 2012; Spencer *et al.*, 2013). Generalization does not appear to depend on whether SNPs are reported to be functional, though one SNP, rs16991615, which did generalize to diverse groups, is a non-synonymous variant in *MCM8*. Indeed, robust replication and generalization were seen for several intronic SNPs as well as an SNP in the regulatory region of *LIN28B*. Generalization also does not appear to be consistent by race—we observed varying generalization by SNP, outcome and race/ethnicity group and for several SNPs, observed large differences in allele frequencies between the ancestral groups. In spite of reasonable power for the AM and ANM analyses, we did not replicate most associations in the large PAGE AA population, perhaps due to different patterns in LD between African and non-African populations, as described above.

The discovery of genetic variants associated with the reproductive lifespan provides an important opportunity to elucidate the biological mechanisms involved with both normal variation and disorders of menarche and menopause. As a next step in this process, we have examined the epidemiologic architecture of these early reported variants, describing their generalizability and effect size across differing ancestral populations in the USA. Our findings suggest that many but not all reproductive trait loci transfer across women of diverse ancestry. In particular, many SNP associations do not appear to extend to AA women. Inclusion of diverse ethnic groups, such as AA, in genetic discovery studies and fine-mapping studies of GWAS-identified loci is needed to identify population-specific variants and facilitate identification of causal variants. Furthermore, given that race/ethnicity may reflect a combination of different social and environmental factors in addition to genetic factors, investigation of how environmental factors interact with genetic variants to alter timing of AM and ANM in diverse populations may be warranted.

Supplementary data

Supplementary data are available at <http://humrep.oxfordjournals.org/>.

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Authors' roles

A.M.A., C.L.C., K.L.S. contributed to study design; I.C., L.L.M., M.F., N.F., U.P., V.M.S. contributed to data collection; B.H., C.A.H., C.S.C., D.C.C., J.W.W., K.L.S. contributed to study execution; A.Y., C.L.C., K.B.G., M.F., N.W.J., R.G., S.B., V.M.S. contributed to analysis; C.L.C. contributed to manuscript drafting; A.Y., C.L.C., L.A.H., L.F.R., N.F., V.W.S. contributed to data interpretation and A.M.A., A.P., C.A.H., D.C.C., J.M., J.W.W., L.A.H., L.F.R., N.F., N.P., T.C.M., U.P. contributed to critical review of the manuscript.

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Conflict of interest

None declared.

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