Meta-analysis of loci associated with age at natural menopause in African-American women

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Age at menopause marks the end of a woman’s reproductive life and its timing associates with risks for cancer, cardiovascular and bone disorders. GWAS and candidate gene studies conducted in women of European ancestry have identified 27 loci associated with age at menopause. The relevance of these loci to women of African ancestry has not been previously studied. We therefore sought to uncover additional menopause loci and investigate the relevance of European menopause loci by performing a GWAS meta-analysis in 6510 women with African ancestry derived from 11 studies across the USA. We did not identify any additional loci significantly associated with age at menopause in African Americans. We replicated the associations between six loci and age at menopause (P-value < 0.05): AMHR2, RHBDL2, PRIM1, HK3/UIMC1, BRSK1/TMEM150B and MCM8. In addition, associations of 14 loci are directionally consistent with previous reports. We provide evidence that genetic variants influencing reproductive traits identified in European populations are also important in women of African ancestry residing in USA.

INTRODUCTION

Menopause marks the end of a woman’s reproductive life. It can occur naturally or be induced. Clinically, natural menopause is defined as the irreversible cessation of menses for 12 months with no obvious cause, such as pregnancy or lactation (1–3). Surgical menopause occurs when both ovaries are removed or damaged by radiation or chemotherapeutic drugs used in cancer treatments (3). Age at natural menopause occurs at ~50–55 years of age in Caucasian women (1–3). Studies have shown that the timing of menopause is a risk factor for osteoporosis, coronary heart disease and cancers. For example, early menopause is associated with increased mortality from cardiovascular diseases (4–8). It has been estimated that for every year’s delay in menopause, the risk for cardiovascular disease is decreased by 2% (9). Women who suffer from premature menopause, defined as menopause prior to age 40, have a 50% higher mortality rate than those reporting menopause above 50 years (4,10,11). To better assist women in their reproductive decisions and their overall health, it is important to determine factors influencing the timing of menopause.

Age at natural menopause is a strongly heritable trait, with more than half of the variation attributed to genetic factors (10,12). Women were at an increased risk of early or late menopause if their mother or non-twin sister had an early or late menopause (13). The odds ratio of a woman having an early menopause if their mother had experienced one was 6.02 (1). Estimates for heritability from mother–daughter and non-twin sister studies range from 42 to 85% (13,14). Environmental factors, such as current cigarette smoking, socio-economic status and alcohol consumption, also associate with the onset of menopause but to a lesser extent (2,3,15,16).

Recent genome-wide association studies (GWAS) in 38 968 women of European ancestry identified numerous single nucleotide polymorphisms (SNPs) associated with age at menopause. These SNPs clustered ~21 loci: RHBDL2, EXO1, FNDC4, TLK1, METAP1D, HELQ, RNF44, UIMC1/HK3, SYCP2L, chr13q34, BAT2, ASH2L, FSHB/MPPED2, PRIM1, KPN13, TDRD3, POLG, GSPT1, BRSK1/TMEM150B, NLRP11 and MCM8 (17–19). SNPs in most of these clusters have also been shown to be associated with early menopause (20,21). Mutations in genes NOBOX, GDF9, BMP15 and FOXL2 have also been associated with premature ovarian insufficiency in several candidate gene studies (22–26). Most of these genes are implicated in DNA repair and immune pathways and suggest that ovarian aging is an early indicator of overall aging (19). Candidate gene studies in European women have also identified six additional loci associated with age at menopause: F5, CYP1B1, ESR1, AMHR2, HSD11B1 and APOE (27–33).

Epidemiological studies have shown that age at natural menopause varies across ethnic groups. African and African-American women were reported to experience menopause at
least 6–12 months sooner when compared with women of European descent (34–37). To date, however, the majority of the genetic studies on age at menopause have involved only women of European ancestry. Genetics studies in various ethnic groups are especially important in elucidating and distilling the genes controlling the timing of this universal reproductive event. To extend our understanding of genetic variants influencing menopause in African-American women, the Women’s Health Initiative (WHI), the Candidate-gene Association Resource (CARe) Consortium, the African American Breast Cancer Consortium (AABC), the Genetic Epidemiology Network of Arteriopathy Study (GENOA), the Health Aging and Body Composition Study (Health ABC), the Healthy Aging in Neighborhoods of Diversity across the Life Span study (HANDLS) and the Multi-Ethnic Study of Atherosclerosis (MESA) Family Study collaborated in this first genome-wide association meta-analysis of GWAS on age at menopause in African-American women and to examine the implicated loci identified in European ancestry populations in African-American women to replicate loci identified in European ancestry populations in African-American women and to examine the implicated regions for secondary signals.

RESULTS

Definition of age at natural menopause

Age at natural menopause was defined as the irreversible cessation of menses for at least 12 months in women who did not undergo hysterectomy with or without oophorectomy (1,2). Women with premature menopause, defined as having menopause prior to 40 years of age, and women who reported menopause after 60 years of age were excluded. Women with hormone use prior to menopause or women who underwent chemotherapy/radiation were also excluded. The current study involved women from 11 population-based studies. The detailed characteristics of participants in these cohorts can be found in Supplementary Material, Table S1. After applying exclusion criteria, a total of 6510 women were included in the analysis. Ancestry analysis showed that the percentage of African ancestry in these women ranged from 75.5 to 83.5%. The mean age at natural menopause ranged from 48.2 to 50.2 years old across cohorts. The majority of the women were recruited from the southern part of the USA.

Meta-analysis for age at natural menopause

We analyzed the association of ~3.25 million genotyped and imputed SNPs with age at natural menopause in two steps. First in each cohort, we performed linear regression analyses, using an additive genetic model, with each SNP coded as a count of the number of minor alleles (0, 1 or 2 for genotyped SNPs and raw dosage values for imputed SNPs) and age at menopause as the response variable. To correct for global ancestry effects, ancestry background of participants was estimated separately in each cohort using principal component analyses (PCA). Recruitment sites and up to the first 10 principal components (PCs) were used as covariates. We performed meta-analysis across all 11 studies using association results from each cohort. SNPs found in only one cohort or genotyped in fewer than 1000 participants were excluded. Manhattan and quantile–quantile plots from the meta-analysis can be found in Supplementary Material, Figure S1. None of the associations achieved genome-wide significance ($P$-value $< 5 \times 10^{-8}$). Results for the top 10 SNPs with the lowest $P$-values are shown in Supplementary Material, Table S2. rs2165706, located in the intergenic region on chromosome 17, had the lowest $P$-value at $1.65 \times 10^{-6}$.

Replication of loci previously identified in women of European ancestry

We hypothesized that loci that have been previously identified as associated with age at menopause in women of European ancestry would also be associated with age at menopause in African-American women. We took a broad approach in selecting SNPs in these loci for replication in which we included all SNPs reported to-date in European women without removing dependent signals. As information on linkage disequilibrium (LD) was not available for all SNPs, we did not want to miss true signals by prefiltering the list of reported SNPs. Previously reported SNPs in European women included 29 discovered in GWAS and meta-analysis study with $P$-values $< 10^{-7}$ and 12 SNPs with $P$-values not reaching genome-wide significance ($<10^{-8}$) (17–19). These SNPs clustered ~30 loci: RHBDL2, EXO1, FNDC4, TLK1, METAP1D, HELQ, RNF44, UMC1/HK3, SYCP2L, BAT2, ASH2L, FSHB/MPPED2, PRIM1, KPN3, TDRD3, POLG, GSP1, BRSK1/TMEM150B, NLRP11, MCM8, CLDN11, SLC44A4, GPR124, DYNCH1H1, BANP, chr2q23.3, chr7q22.3, chr9q21.31, chr11q25 and chr13q34. We also included seven SNPs located in F3, CYP1B1, ESR1, AMHR2, HSDB1, APOE that were implicated in candidate gene studies in European women (27–33). We also sought to identify additional signals in the LD block surrounding these index SNPs.

Loci on chromosome 1 (RHBDL2), 5 (HK3/UMC1), 12 (PRIM1), 19 (BRSK1/TMEM150B), 20 (MCM8) and AMHR2 are associated with age at menopause in African-American women

HK3/UMC1 locus on chromosome 5

Five correlated SNPs (rs2278493, rs691141, rs7718874, rs365132 and rs402511) have been reported in association with age at menopause at the HK3/UMC1 locus on chromosome 5 (17). Since these SNPs were identified in different cohorts of European ancestry, we first assessed whether they represented the same signal from this region by looking at the LD relationships among these SNPs in Europeans. We used the HapMapII CEU cohort as a proxy for Europeans and found these five SNPs likely reflected the same signal, thus allowing us to perform only one test at a significance level of 0.05. Of the five SNPs, only rs402511 was significant ($P = 0.042$) while rs365132 and rs7718874 showed similar magnitude of $P$-values ($P = 0.052$ and 0.063) (Table 1). The effect sizes for rs402511, rs365132 and rs77188754 were 0.205, 0.188 and −0.196, respectively, slightly attenuated compared with those reported in European women (0.39, 0.287 and −0.39, respectively, from initial discovery stage). To fine-map the signal in this locus, we examined the LD structure in the HapMapII YRI.
<table>
<thead>
<tr>
<th>SNP</th>
<th>Chromosome (position)</th>
<th>AL</th>
<th>Freq (s.e.)</th>
<th>Effect (s.e.)</th>
<th>P-value</th>
<th>Total sample size</th>
<th>European ancestry women P-value</th>
<th>Gene</th>
<th>Genic location</th>
<th>ENCODE regulation</th>
<th>DNasel signal</th>
<th>Histone mark</th>
<th>Chromatin state</th>
</tr>
</thead>
<tbody>
<tr>
<td>rs4246511&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1 (39152972)</td>
<td>T</td>
<td>0.69 (0.013)</td>
<td>0.206 (0.095)</td>
<td>0.031&lt;sup&gt;c&lt;/sup&gt;</td>
<td>6510 0.24 9.08 × 10&lt;sup&gt;-17&lt;/sup&gt;</td>
<td>RBBDL2 Intron – – – –</td>
<td>B3K4Me1</td>
<td>Weak enhancer</td>
<td></td>
<td></td>
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<tr>
<td>rs1635501&lt;sup&gt;b&lt;/sup&gt;</td>
<td>1 (240107398)</td>
<td>T</td>
<td>0.479 (0.019)</td>
<td>0.111 (0.083)</td>
<td>0.178 6478 0.16 8.46e10&lt;sup&gt;-10&lt;/sup&gt;</td>
<td>EXO1 Intron – – – –</td>
<td>–</td>
<td>–</td>
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<tr>
<td>rs2003639&lt;sup&gt;c&lt;/sup&gt;</td>
<td>2 (27568920)</td>
<td>T</td>
<td>0.369 (0.012)</td>
<td>0.031 (0.086)</td>
<td>0.721 6510 0.17 2.25 × 10&lt;sup&gt;-14&lt;/sup&gt;</td>
<td>ENDC4</td>
<td>Intron – – – –</td>
<td>–</td>
<td>–</td>
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<tr>
<td>rs10183486&lt;sup&gt;d&lt;/sup&gt;</td>
<td>2 (17169217)</td>
<td>T</td>
<td>0.578 (0.015)</td>
<td>0.028 (0.083)</td>
<td>0.732 6502 0.19 2.21 × 10&lt;sup&gt;-14&lt;/sup&gt;</td>
<td>TLK1 Intron – – – –</td>
<td>–</td>
<td>–</td>
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<tr>
<td>rs4693089&lt;sup&gt;e&lt;/sup&gt;</td>
<td>4 (84592646)</td>
<td>A</td>
<td>0.235 (0.014)</td>
<td>0.122 (0.099)</td>
<td>0.223 6510 0.22 2.38 × 10&lt;sup&gt;-19&lt;/sup&gt;</td>
<td>HELQ</td>
<td>Intron – – – –</td>
<td>–</td>
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<tr>
<td>rs2278493&lt;sup&gt;f&lt;/sup&gt;</td>
<td>5 (167247040)</td>
<td>T</td>
<td>0.139 (0.009)</td>
<td>0.092 (0.118)</td>
<td>0.438 6489 0.3 7.20 × 10&lt;sup&gt;-8&lt;/sup&gt;</td>
<td>HK3 Intron – – – –</td>
<td>Repressed</td>
<td>–</td>
<td></td>
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<tr>
<td>rs691141&lt;sup&gt;g&lt;/sup&gt;</td>
<td>5 (16255904)</td>
<td>A</td>
<td>0.424 (0.008)</td>
<td>0.026 (0.072)</td>
<td>0.753 6495 0.36 3.90 × 10&lt;sup&gt;-12&lt;/sup&gt;</td>
<td>HK3</td>
<td>Intergenic – – – –</td>
<td>H3K4Me1</td>
<td>Weak enhancer/insulator</td>
<td></td>
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<tr>
<td>rs7362979&lt;sup&gt;h&lt;/sup&gt;</td>
<td>5 (17629594)</td>
<td>C</td>
<td>0.061 (0.008)</td>
<td>0.551 (0.171)</td>
<td>1.25 × 10&lt;sup&gt;-3&lt;/sup&gt; 6503 – – – –</td>
<td>HK3/UMC1</td>
<td>Intergenic – – – –</td>
<td>–</td>
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<tr>
<td>rs7718874&lt;sup&gt;i&lt;/sup&gt;</td>
<td>5 (176290671)</td>
<td>A</td>
<td>0.210 (0.011)</td>
<td>0.190 (0.101)</td>
<td>0.052 6509 0.39 1.30 × 10&lt;sup&gt;-13&lt;/sup&gt;</td>
<td>UMC1 Intron – – – –</td>
<td>–</td>
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<tr>
<td>rs3561321&lt;sup&gt;j&lt;/sup&gt;</td>
<td>5 (17621810)</td>
<td>T</td>
<td>0.188 (0.011)</td>
<td>0.180 (0.062)</td>
<td>0.062 6508 0.39 8.4 × 10&lt;sup&gt;-14&lt;/sup&gt;</td>
<td>UMC1 Intron – – – –</td>
<td>–</td>
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<tr>
<td>rs402511&lt;sup&gt;k&lt;/sup&gt;</td>
<td>5 (176367046)</td>
<td>A</td>
<td>0.205 (0.101)</td>
<td>0.042&lt;sup&gt;j&lt;/sup&gt;</td>
<td>6510 0.39 1.40 × 10&lt;sup&gt;-15&lt;/sup&gt;</td>
<td>UMC1 Intron – – – –</td>
<td>–</td>
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<tr>
<td>rs2135157&lt;sup&gt;l&lt;/sup&gt;</td>
<td>6 (11005474)</td>
<td>T</td>
<td>0.663 (0.091)</td>
<td>0.487</td>
<td>6492 0.29 16.0 × 10&lt;sup&gt;-8&lt;/sup&gt;</td>
<td>SYCP2 Intron – – – –</td>
<td>–</td>
<td>–</td>
<td></td>
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<tr>
<td>rs1046089&lt;sup&gt;m&lt;/sup&gt;</td>
<td>6 (31791046)</td>
<td>A</td>
<td>0.492 (0.014)</td>
<td>0.035 (0.083)</td>
<td>0.672 6469 0.21 1.63 × 10&lt;sup&gt;-16&lt;/sup&gt;</td>
<td>BAT2</td>
<td>Intergenic – – – –</td>
<td>UMC1</td>
<td>Synony-mous</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>rs2517388&lt;sup&gt;n&lt;/sup&gt;</td>
<td>8 (38968889)</td>
<td>T</td>
<td>0.099 (0.123)</td>
<td>0.942</td>
<td>6509 0.26 9.31 × 10&lt;sup&gt;-15&lt;/sup&gt;</td>
<td>ASH2L Intron – – – –</td>
<td>–</td>
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<tr>
<td>rs12294104</td>
<td>11 (3039475)</td>
<td>T</td>
<td>0.046 (0.137)</td>
<td>0.736</td>
<td>6508 0.22 1.46 × 10&lt;sup&gt;-11&lt;/sup&gt;</td>
<td>FSHB/MPPED2 Intergenic – – – –</td>
<td>–</td>
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<tr>
<td>rs2277339&lt;sup&gt;o&lt;/sup&gt;</td>
<td>12 (55432336)</td>
<td>T</td>
<td>0.255 (0.112)</td>
<td>0.022&lt;sup&gt;o&lt;/sup&gt;</td>
<td>6510 0.38 2.47 × 10&lt;sup&gt;-19&lt;/sup&gt;</td>
<td>PRIM1</td>
<td>Intergenic – – – –</td>
<td>–</td>
<td>–</td>
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</tbody>
</table>

<sup>1</sup> At University of Washington on May 29, 2014 http://hmg.oxfordjournals.org/ Downloaded from
rs6139882

20 (5902202) T 0.052 (0.156) −0.850 (0.265) 1.36 × 10⁻³⁸ 6060 − − MCM8 Intron GR (112) − − −

\( P \)-value is significant at \( \alpha = 0.05 \) (for previously identified SNPs), or at \( \alpha = 0.05/N \), where \( N \) is the number of SNPs on the LD block (for SNPs with the lowest \( P \)-value on the LD block).

AL, allele; Freq, frequency; s.e., standard error.

Positions are in NCBI36 coordinates.

Average frequencies and effect sizes of minor alleles across all cohorts are listed.

Units in years.

If the SNP is in the intergenic region, the closest upstream and downstream genes are listed.

Data related to regulation of transcription from the ENCODE project (74).

CHiP-seq results with identities of transcription factors. Highest signals are shown in brackets.

Regions where the chromatin is hypersensitive. Regulatory regions tend to be DNaseI sensitive. DNaseI sensitivity is shown as the absolute density of \( in vivo \) cleavage sites across the genome mapped.

Modification of histone proteins is suggestive of enhancer (H3K4Me1 and H3K27Ac) and promoter activity (H3K4Me3, H3K4Me1 and H3K27Ac).

Computed using Hidden Markov model by Broad Institute, MIT and MGH. Fifteen states are possible (76).

SNPs identified in previous meta-analyses (19). Effect sizes and \( P \)-values were from the published combined results.

SNPs identified in previous GWAS (17,18).

SNPs identified in the LD block analysis in current study.

rs6139882 is not in LD with rs236114 (\( r^2 = 0.000 \)). rs6139882's LD with these two SNPs suggests that the functional SNP must be in LD with these two SNPs. Rs6139882 is not in LD with rs402511 or rs3762971 at \( \alpha = 0.05 \). We found that rs3762971 may be an independent signal from rs402511. We sought to test all SNPs in high LD with rs402511 in both European and African ethnic backgrounds. Due to the extent of LD in this region, we were only able to narrow the location of the signal down to 167.8 kb (chr5: 176,255,944–176,423,163, \( r^2 = 0.28 \)).
Americans, indicating that rs6139882 is likely a secondary signal from this locus.

RHBDL2 locus on chromosome 1, PRIM1 locus on chromosome 12 and AMHR2 locus

Of the 16 SNPs identified only by a meta-analysis of European cohorts, we replicated two loci (rs4246511 at RHBDL2 on chromosome 1, \( P = 0.031 \); rs2277339 at PRIM1 on chromosome 12, \( P = 0.022 \)) in African-American cohorts (Table 1). The effect sizes for rs4246511 and rs2277339 were 0.21 and 0.36 in our cohorts, similar to those reported previously (0.24 and 0.38, respectively, from the combined analysis (19)). In addition, 11 loci showed direction of effects consistent with previous reports.

Several candidate gene studies in women with European ancestry identified seven SNPs located in six loci (F5, CYP1B1, ESR1, AMHR2, HSD1, APOE) as associated with age at menopause (27–33). We investigated whether these SNPs are also associated with this trait in African-American women. Of the seven SNPs, only rs2002555 located in AMHR2 was shown to be associated with our phenotype (\( P = 0.0062 \)).

DISCUSSION

Genetic factors play an important role in determining the timing of natural menopause. These factors have been investigated most extensively in women of European ancestry but relatively little genetic research has been done in African-American women. Our study in African-American women is the largest and most comprehensive conducted to-date. In this study, we report that the mean age of women who underwent natural menopause ranged from 48.2 to 50.2 years of age across cohorts, which is earlier than women of European ancestry (1–3). Prior epidemiological studies have also shown that African-American women tend to experience menopause sooner (34,37). Differences in the timing of this universal event suggest that different modifiers might exist in different ethnic backgrounds.

Previous GWAS in women of European ancestry identified 21 loci associated with age at menopause (17–19). We investigated the relevance of these loci in 6510 African-American women and replicated rs4246511 in RHBDL2 on chromosome 1, rs2277339 in PRIM1 on chromosome 12, HK3/UMC1 locus on chromosome 5, BRSK1/TMEM150B locus on chromosome 19 and MCM8 on chromosome 20. Rs4246511 is located in the...
Table 2. Replication of loci identified from candidate gene studies in European cohorts

<table>
<thead>
<tr>
<th>Gene</th>
<th>SNP</th>
<th>Chromosome (position)</th>
<th>Alleles&lt;sup&gt;b&lt;/sup&gt;</th>
<th>Freq (s.e.)</th>
<th>Effect (s.e.)</th>
<th>P-value</th>
<th>Total sample size</th>
<th>Previous candidate studies</th>
</tr>
</thead>
<tbody>
<tr>
<td>F5</td>
<td>rs6020</td>
<td>1 (167785736)</td>
<td>T/C</td>
<td>0.275 (0.018)</td>
<td>-0.067 (0.094)</td>
<td>0.473</td>
<td>6195</td>
<td>-</td>
</tr>
<tr>
<td>CYP1B1</td>
<td>rs1056836</td>
<td>2 (38151707)</td>
<td>C/G</td>
<td>0.755 (0.016)</td>
<td>-0.025 (0.097)</td>
<td>0.794</td>
<td>6293</td>
<td>+/-</td>
</tr>
<tr>
<td>CYP1B1</td>
<td>rs1800440</td>
<td>2 (38151643)</td>
<td>T/C</td>
<td>0.954 (0.006)</td>
<td>-0.069 (0.21)</td>
<td>0.744</td>
<td>5918</td>
<td>+/+</td>
</tr>
<tr>
<td>ESR1</td>
<td>rs2234693</td>
<td>6 (152205028)</td>
<td>T/C</td>
<td>0.462 (0.014)</td>
<td>0.159 (0.081)</td>
<td>0.050</td>
<td>6502</td>
<td>-/+</td>
</tr>
<tr>
<td>AMHR2</td>
<td>rs2002555</td>
<td>12 (52103504)</td>
<td>A/G</td>
<td>0.871 (0.008)</td>
<td>0.332 (0.122)</td>
<td>0.006&lt;sup&gt;a&lt;/sup&gt;</td>
<td>6508</td>
<td>+/+</td>
</tr>
<tr>
<td>HSDB1</td>
<td>rs2830</td>
<td>17 (37958089)</td>
<td>A/G</td>
<td>0.686 (0.015)</td>
<td>-0.131 (0.091)</td>
<td>0.151</td>
<td>6510</td>
<td>-/+</td>
</tr>
<tr>
<td>APOE</td>
<td>rs7412</td>
<td>19 (50103919)</td>
<td>T/C</td>
<td>0.098 (0.003)</td>
<td>-0.334 (0.425)</td>
<td>0.431</td>
<td>720</td>
<td>-</td>
</tr>
</tbody>
</table>

<sup>a</sup>P-value is significant at α = 0.05.<n>^b^Minor/major alleles. Average frequencies and effect sizes of minor alleles across all cohorts are listed.

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from the ENCODE project suggested that this region may have weak enhancer activity (39). Rs2277339 is a missense SNP located in PRIM1, a protein involved in DNA replication (40). Results from the ENCODE project showed that it is an active regulatory site where many transcription factors, such as TAF1, HEY1, PAX5-20 and histone markers, such as H3K4Me3 and H3K27Ac, bind. Since both RHBLD2 and PRIM1 are involved in DNA repair, we hypothesize that there may be some interaction between the DNA repair pathway and the pathway leading to menopause. We uncovered two signals at HK3/UIMC1 locus on chromosome 5: rs402511 and rs3762971. Both of them are located in the intergenic region, one upstream of HK3 and one downstream of UIMC1. The histone-binding patterns indicate that they may have weak enhancer activities and rs3762971 may be a part of an insulator sequence. All of the SNPs we replicated at BRSK1/TMEM150B locus are located in the introns and have been predicted to have weak enhancer and repressor activities. Rs16991615, a missense SNP at the MCM8 locus on chromosome 20, has the lowest P-value in studies of Caucasian women; however, it did not replicate in our cohort. One reason could be due to the different MAF in two ethnic groups (7% in Caucasians and 1.3% in Africans), requiring a far larger African-American sample to detect its effect. The effect size for rs16991615 in African Americans in our studies was 0.346 (s.e. = 0.452) while it was reported to be 1.07 (from combined analysis) in Caucasians (19). We, however, did uncover another signal at MCM8 (rs6139882 located in the intronic region of MCM8). It is plausible that there are multiple signals at this locus, resulting in a similar effect on MCM8 regulation. A recent study in African Americans also could not replicate the association between rs16991615 and age at menopause while another study in Hispanics showed the association, supporting a likely hypothesis that the causal variant at the MCM8 locus is in LD with rs16991615 in Caucasians and with rs6139882 in African Americans (41,42). MCM8 is expressed within the oocytes in the follicles of the human ovary and that at least in Caucasian women, rs16991615 is associated with antral follicle count, which reflects ovarian reserve (43). These findings all support the hypothesis that MCM8 is involved in the pathway leading to menopause.

Several small candidate gene studies suggested F5, CYP1B1, ESR1, AMHR2, HSDB1 and APOE may affect the timing of menopause. A recent study in a large number of women with...
European ancestry did not replicate any of these genes (33). Of these genes, we only found rs2002555 at AMHR2 to be significantly associated with age at menopause. AMHR2 is a gene coded for AMH (anti-Mullerian hormone) receptor, type II. A missense SNP in this locus is associated with sex determination in the tiger pufferfish, suggesting that AMHR2 played a role in the reproductive pathway across species (44). In humans, AMH binds to AMHR2. AMH is expressed by granulosa cells of the ovary during the reproductive years, setting the pace at which follicles reenter meiosis and growth (45). Hence, AMH level is associated with the rate of depletion of the primordial follicle pool and has been shown to be a good predictor for age at menopause (46).

Results from our meta-analysis of African-American women confirm some, but not all, of the findings in women of European ancestry. One reason is our modest sample size. With our sample
size, however, we have at least 80% power to replicate SNPs with magnitude of effect as low as 0.2 in Caucasian women (Fig. 4; Supplementary Material, Table S4). These SNPs were located in five of the 21 loci identified from previous GWAS. Another three loci could be replicated with at least 60% power. Another reason is that even though menopause is a universal event determined mostly by genetic factors, there exist many modifiers in the associated loci. The complex genomic structures in African Americans make it more difficult to distinguish modifiers with smaller effects from background mutation due to the existence of more variants. Due to the different LD patterns in African Americans, the identified GWAS hits could be proxies for the causal variant, which may not be well genotyped or imputed in our data. Our meta-analysis supports the notion that genetic variants influencing menopause timing identified in European populations are also important in women of African ancestry residing in the USA. More functional experiments are needed as a direct way to test the effects these SNPs have on ovarian aging.

**MATERIALS AND METHODS**

**Cohort descriptions**

*Women’s Health Initiative (WHI)* is a long-term national health study that focuses on strategies for preventing common diseases such as heart disease, cancer and fracture in postmenopausal women. A total of 161,838 women aged 50–79 years old were recruited from 40 clinical centers in the USA between 1993 and 1998. WHI consists of an observational study, two clinical trials of postmenopausal hormone therapy (estrogen alone or estrogen plus progestin), a calcium and vitamin D supplement trial, and a dietary modification trial (47). Study recruitment and exclusion criteria have been described previously (48). Study protocols and consent forms were approved by the institutional review boards at all participating institutions. Medical history was updated annually (for women in the observational study) or semiannually (for women in the clinical trials) by mail and/or telephone questionnaires. The WHI SNP Health Association Resource (WHI-SHARE) includes 8515 self-identified African-American women from WHI who provided consent for DNA analysis. Approximately 1% of samples could not be genotyped. Samples in the following categories were excluded: with call rates <95%, in duplicates, or with genotype calls on the Y chromosome. The relatedness among participants was investigated and there were 97 sets of monozygotic twins, parent–offspring, full-siblings and half-siblings. After retaining only one individual from each set, our final sample size was 8421 African-American women. Genotyping and quality control: all WHI samples, plus 2% blinded duplicates, were genotyped at Affymetrix Inc. (www.affymetrix.com) on the Genome-wide Human SNP Array 6.0 (909,622 SNPs). SNPs that were located on the Y chromosome or were Affymetrix quality control probes (not intended for analysis) were excluded (n = 3280). We also excluded SNPs that had call rates <95% and concordance rates <98%, leaving us 871,309 SNPs available for use in this study. The average concordance for blinded duplicate samples was 99.8%, and the average sample call rate after SNP exclusions was 99.8%. Imputation: imputation for WHI samples was performed in MACH 1.0.16 (http://www.sph.umich.edu/csg/abecasis/mach/) using a reference panel composed of 60 unrelated CEU and 60 unrelated YRI samples from HapMap release 22 (49). Phenotype definition: the questions related to menopausal status on WHI enrollment questionnaires were: ‘How old were you when you last had any menstrual bleeding? (If you are still having menstrual bleeding or periods, enter your current age.)’, ‘When was the last time you had any menstrual bleeding or spotting?’, ‘Did you ever have a hysterecomy?’ and ‘Did you ever have an operation to have one or both of your ovaries taken out?’ Eligible WHI participants were defined as those who had at least one ovary intact before the cease of menses and who reported having last menstrual bleeding or spotting over 12 months ago at the time of the study. WHI Participants with endometrial and cervical cancer history were removed. Population Stratification Correction: EIGENSTRAT was used to perform separate PCA to compute the first 10 PC for each individual (50). The reference panel was composed of 475 ancestors from HapMap populations and the Human Genome Diversity Panel. One hundred and nine CEU and 108 YRI individuals were taken from the HapMap. PCA analyses were performed on these two groups separately using EIGENSTRAT. Each population was subjected to five rounds of outlier removal based on the resulting PC information. In total, two outliers were identified in each group, bringing the final population sizes to 107 CEU and 106 YRI. Sixty-three individuals in HGDP sampled from the Americans (8 Surui, 22 Mayans, 13 Karitiana, 14 Pima, 6 Colombian) and 225 East Asians were subjected to the same EIGENSTRAT PCA-based outlier removal procedure as for the HapMap individuals, leaving 200 East Asian individuals and 62 Native Americans. 178 101 SNPs overlapping the HGDP, HapMap and WHI subjects were used in EIGENSTRAT. To complement the PCA analysis, *frappe* was used to estimate the relative contributions of all four ancestral groups to every admixed individual (51). A total of 656,852 autosomal SNPs were used in the *frappe* analysis.

The *GENOA* is a community-based study of hypertensive sibships that aims to identify genes influencing blood pressure.

![Figure 4.](http://hmg.oxfordjournals.org/) Comparisons of effect magnitude of loci associated with age at menopause between women with European and African ancestry. Each point represents a previously identified locus associated with age at menopause in women with European ancestry. Loci replicated in the current African-American cohorts are colored in black. The dash line indicates the theoretical $x = y$ line.
In the initial phase of the GENOA study (9/1995 to 6/2001), sibships containing ≥ 2 individuals with essential hypertension diagnosed before age 60 years were selected for participation. At the Jackson, MS field center, 1854 African Americans were enrolled. Participants returned in a second phase of GENOA (December 2000 to June 2004) for physical examination and measurement of non-conventional and novel risk factors. Genotyping and Quality Control: all samples were genotyped on Affymetrix Genome-wide human SNP array 6.0. Imputation: a single step approach was performed in MACH 1.0.16 (http://www.sph.umich.edu/csg/abecasis/mach/) using a reference panel composed of 60 unrelated CEU and 60 unrelated YRI samples from HapMap release 22 (49). Phenotype definition: subjects were asked whether or not they had reached menopause, defined as the cessation of natural menses for at least 12 months. Subjects that reported reaching menopause were asked ‘In what year or how old were you when you reached menopause?’. Subjects with menopause induced by surgery, radiation or chemotherapy were excluded. Population Stratification Correction: PCs for individuals were calculated based on all SNPs in the dataset without a reference panel using SNP and Variation Suite 7 (SVS7) from Golden Helix (http://www.goldenhelix.com/SNP_Variation/). The relative contribution of CEU and YRI in GENOA individuals were assessed on a panel of 385 ancestry informative markers using the STRUCTURE program (54).

The Health ABC is a NIA-sponsored ongoing cohort study of the factors that contribute to incident disability and the decline in function of healthier older persons, with a particular emphasis on changes in body composition in old age. Health ABC enrolled well-functioning, community-dwelling black (n = 1281) and white (n = 1794) men and women aged 70–79 years between April 1997 and June 1998. Participants were recruited from a random sample of white and all black Medicare eligible residents in the Pittsburgh, PA and Memphis, TN, metropolitan areas. The key components of Health ABC include a baseline exam, annual follow-up clinical exams and phone contacts every 6 months to identify major health events and document functional status between clinic visits. Genotyping and Quality Control: genotyping was done in 1663 Caucasians and 1159 African Americans using the Illumina Human 1M-Duo BeadChip System at the Center for Inherited Disease Research (CIDR). SNPs were excluded if their call rate was < 97%, Hardy–Weinberg equilibrium \( P \)-value was \( < 10^{-6} \), or major allele frequency was < 1%. Imputation: MACH 1.0.16 was used for imputation, using HapMap CEU release 22, build 36, as the reference panel. Phenotype Definition: participants who reported hysterectomy or ovariectomy (bilateral or otherwise) were excluded. Population Stratification Correction: EIGENSTRAT was used to calculate PCs for individuals using all SNPs that passed quality control.

The HANDLS is an interdisciplinarian, community-based, prospective longitudinal epidemiologic study examining the influences of race and socioeconomic status (SES) on the development of age-related health disparities among socioeconomically diverse African Americans and whites in Baltimore. This study investigates whether health disparities develop or persist due to differences in SES, differences in race or their interaction. The HANDLS design is an area probability sample of Baltimore based on the 2000 Census. The study protocol facilitated our ability to recruit 3722 participants from Baltimore, MD with mean age 47.7 (range 30–64) years and 2200 African Americans (59%) and 1522 whites (41%). One thousand and twenty-four participants, a subset of participants self-reporting as African Americans, were genotyped at the Laboratory of Neurogenetics located in the National Institute on Aging in National Institutes of Health. Genotyping and Quality Control: samples were genotyped at 907 763 SNPs at the equivalent of Illumina 1M SNP coverage (709 samples using Illumina 1M and 1Mduo arrays, the remainder using a combination of 550K, 370K, 510S and 240S to equate the million SNP level of coverage), passing inclusion criteria into the genetic component of the study. Initial inclusion criteria for genetic data include concordance between self-reported sex and sex estimated from X chromosome heterogeneity, >95% call rate per participant (across all equivalent arrays), concordance between self-reported African ancestry and ancestry confirmed by analyses of genotyped SNPs, and no cryptic relatedness to any other samples at a level of proportional sharing of genotypes >15% (effectively excluding first cousins and closer relatives from the set of probands used in analyses). In addition, SNPs were filtered for HWE \( P \)-value of >10\(^{-7}\), missing by haplotype \( P \)-values of >10\(^{-7}\), MAF >0.01 and call rate of >95%. Basic genotype quality control and data management were conducted using PLINKv1.06 (55). Cryptic relatedness was estimated via pairwise identity by descent analyses in PLINK and confirmed using RELPAIR (56). Imputation: imputation was performed in MACH 1.0.16 (http://www.sph.umich.edu/csg/abecasis/mach/) using a reference panel composed of 60 unrelated CEU and 60 unrelated YRI samples from HapMap release 22 (49). A two-stage process was used: first estimating recombination and crossover events in a random sample of 200 participants, then based on this data and the reference haplotypes, 200 iterations of the maximum likelihood model were used to estimate genotype dosages for imputed SNPs. Phenotype Definition: age at natural menopause was defined as the age at the last menstrual period which occurred naturally. Only women with natural menopause between the ages of 40 and 60 years were included. Subjects were asked ‘what was your age at menopause in years?’ Population Stratification Correction: ancestry estimates were assessed using both STRUCTUREv2.3 (54,57,58) and the multidimensional scaling (MDS) function in PLINKv1.06 (55). In the MDS analysis, HANDLS participants were clustered with data made available from HapMap Phase 3 for the YRI, ASW, CEU, TSI, JPT and CHB populations, using a set of 36 892 linkage-disequilibrium-pruned SNPs common to each population. This set of SNPs were chosen as they are not in \( r^2 > 0.20 \) with another SNP in overlapping sliding windows of 100 SNPs in the ASW samples. HANDLS participants with component vector estimates consistent with the HapMap ASW samples for the first 4 component vectors were included. In addition, the 1024 quality controlled HANDLS samples were later clustered among themselves using MDS to generate 10 component vectors estimating internal population structure within the HANDLS study.

MESA family study is a multicenter, family based study whose goal is to locate and identify genes contributing to the genetic risk for cardiovascular disease, by looking at the early changes of atherosclerosis within families (mainly siblings). The
MESA family cohort was recruited from the six MESA Field Centers. MESA family participants underwent the same examination as MESA participants during May 2004–May 2007. In a small proportion of subjects, parents of MESA index subjects participating in MESA family were studied but only to have blood drawn for genotyping. DNA was extracted and lymphocytes immortalized for 1633 non-classic MESA family members (950 African Americans and 683 Hispanic Americans) from 594 families, yielding 3026 sibpairs. Genotyping and Quality Control: samples were genotyped on the Affymetrix Genome-wide human SNP array 6.0. Genotype calls were performed using Birdseed v1.33. We used PLINK, EIGENSTRAT and PREST-Plus software for quality control (59). Imputation: IMPUTE version 2.1.0 was used to perform imputation with HapMap Phases I and II (CEU, YRI, CHB, JPT populations) as the reference panel (release 22—NCBI Build 36 (dbSNP b126)) (60,61). Phenotype Definition: age at natural menopause was assessed retrospectively by questionnaire and included age at last menstrual period, and age at hysterectomy or ovary removal, if either. Use of hormone replacement therapy was ascertained at clinic visits from medicine bottles, as previously described (66). Women with menopause due to hysterectomy and/or bilateral oophorectomy and women using hormone replacement therapy before menopause were excluded. Population Stratification Correction: PCs were computed based on 97 404 genotyped SNPs from the Illumina Omni 1M chip (approximately every 10th SNP on the array) using EIGENSTRAT (50).

The ARIC is a multicenter prospective investigation of atherosclerotic disease in a predominantly biracial population (67). Men and women aged 45–64 years at baseline were recruited from four communities: Forsyth County, North Carolina; Jackson, Mississippi; suburban areas of Minneapolis, Minnesota and Washington County, Maryland. A total of 15 792 individuals participated in the baseline examination in 1987–89 when blood was drawn for DNA extraction and participants consented to genetic testing. Genotype data were available on 2989 self-reported African-American individuals. Genotyping and Quality Control: samples were genotyped at the Broad Institute using the Affymetrix Genome-Wide Human SNP Array 6.0. Two methods of DNA quality control metrics were assessed on the samples (59). Genotypes were called using Birdseed v1.33. Quality control steps were performed using the software PLINK, EIGENSTRAT and PREST-Plus (59). Imputation: imputation was performed in MACH 1.0.16 (http://www.sph.umich.edu/csg/abecasis/mach/) using a reference panel composed of 60 unrelated CEU and 60 unrelated YRI samples from HapMap release 22 (49). Imputation based on Affymetrix data resulted in an allelic concordance rate of ~95.6%, calculated as 1 – 1/2^|imputed_dosage – chip_dosage|. This rate is comparable with rates calculated for individuals of African descent imputed with the HapMap 2 YRI individuals. For each imputed sample, imputation was performed in two steps. For the first step, individuals with pedigree relatedness or cryptic relatedness (pi_hat > 0.05) were filtered. A subset of individuals was randomly extracted from each panel and used to generate recombination and error rate estimates for the corresponding sample. In the second step, these rates were used to impute all sample individuals across the entire reference panel. Imputation results were filtered at an RSQ_HAT threshold of 0.3 and an MAF threshold of 0.01. Phenotype Definition: a reproductive history questionnaire was completed by all women in the ARIC study at each study visit. The questionnaire included questions on whether they were currently menstruating, their age at last menstrual period (in years) for those who were no longer menstruating, as well as questions on whether and when they had undergone surgery to remove their womb and/or ovaries, or had undergone chemotherapy or radiation treatment. Age at menopause was determined from each woman’s first report of their age at menopause (age at last period), and restricted to those that denied hysterectomy, denied removal of both ovaries prior to cessation of menses, and denied any chemotherapy or radiation treatment which stopped their menses. Only such cases between the ages of 40 and 60 years were included. Population Stratification Correction: EIGENSTRAT was used to compute the first 10 PCs for each individual (50). The reference populations were composed of 1178 European Americans and 756 Nigerians from the Yoruba region (59).
The JHS is a community-based observational study whose primary objective is to investigate causes of cardiovascular disease in an African-American population. The cohort \( N = 5301 \) was sampled from non-institutionalized African Americans 35–84 years old in urban and rural areas of three counties in Mississippi (Hinds, Madison and Rankin) and involved four components: (i) African Americans from Jackson, MS who were also participants in the ARIC cohort (31%); (ii) a family cohort of relatives of index participants (22%; all ages \( \geq 21 \) included); (iii) individuals from randomly sampled households from a commercial listing (17%) and (iv) a constrained volunteer sample designed to reflect the demographics of the overall population (30%) (68). This study was approved by the institutional review board, and all subjects provided written informed consent. Genotyping and Quality Control: samples were genotyped at the Broad Institute using the Affymetrix Genome-Wide Human SNP Array 6.0. Two methods of DNA quality control metrics were assessed on the samples (59). Genotypes were called using Birdseed v1.33. Quality control steps were performed using the software PLINK, EIGENSTRAT and PREST-Plus (59). Imputation: imputation was performed in MACH 1.0.16 (http://www.sph.umich.edu/csg/abecasis/mach/) using a reference panel composed of 60 unrelated CEU and 60 unrelated YRI samples from HapMap release 22 (49). Imputation based on Affymetrix data resulted in an allelic concordance rate of \( \sim 95.6\% \), calculated as \( 1 – 1/2^{|\text{imputed dosage} – \text{chip dosage}|} \). This rate is comparable with rates calculated for individuals of African descent imputed with the HapMap 2 YRI individuals. For each imputed sample, imputation was performed in two steps. For the first step, individuals with pedigree relatedness or cryptic relatedness (\( \text{pi}_\text{hat} > 0.05 \)) were filtered. A subset of individuals was randomly extracted from each panel and used to generate recombination and error rate estimates for the corresponding sample. In the second step, these rates were used to impute all sample individuals across the entire reference panel. Imputation results were filtered at an RSQ_HAT threshold of 0.3 and an MAF threshold of 0.01. Phenotype Definition: age at natural menopause was defined as the age at the last menstrual period which occurred naturally. Only women with natural menopause between the ages of 40 and 60 years were included. Subjects were asked ‘what was your age at menopause in years’. Population Stratification Correction: EIGENSTRAT was used to compute the first 10 PCs for each individual (50). The reference populations were composed of 1178 European Americans and 756 Nigerians from the Yoruba region (59).

Two cohorts from the AABC were included in the current study: the Carolina Breast Cancer Study (CBCS) and The Multi-ethnic Cohort Study (MEC). The other cohorts were excluded due to the small size of subjects with available data (<100). The control and case samples were treated as distinct cohorts.

The CBCS is a population-based case–control study conducted between 1993 and 2001 in 24 counties of central and eastern North Carolina (70). Cases were identified by rapid case ascertainment system in cooperation with the North Carolina Central Cancer Registry and controls were selected from the North Carolina Division of Motor Vehicle and United States Health Care Financing Administration beneficiary lists. Participants’ ages ranged from 20 to 74 years. DNA samples were provided from 656 African American cases with invasive breast cancer and 608 African-American controls. Genotyping and Quality control: genotyping in Stage 1 was conducted using the Illumina Human1M-Duo BeadChip. We removed samples with DNA concentrations <20 ng/ml. Following genotyping, we removed samples based on the following exclusion criteria: (i) unknown replicates (\( \geq 98.9\% \) genetically identical) that we were able to confirm; (ii) unknown replicates that we were not able to confirm through discussions with study investigators; (iii) samples with call rates <95% after a second attempt; (iv) samples with ≤5% African ancestry and (v) samples with <15% mean heterozygosity of SNPs on the X chromosome and/or similar mean allele intensities of SNPs on the X and Y chromosomes (these are likely to be males). In the analysis, we removed SNPs with <95% call rates or MAFs <1%. To assess genotyping reproducibility, we included replicate
samples, eliminating SNPs with genotyping concordance rates <98% based on the replicates. The final analysis dataset included 1,043,036 SNPs genotyped with an average SNP call rate of 99.7% and average sample call rate of 99.8%. Imputation: in order to generate a dataset suitable for fine-mapping, we carried out genome-wide imputation using the software MACH. Phased haplotype data from the founders of the CEU and YRI HapMap Phase 2 samples were used to infer LD patterns in order to impute un-genotyped markers. The r2 metric, defined as the observed variance divided by the expected variance, provides a measure of the quality of the imputation at any SNP, and was used as a threshold in determining which SNPs to filter from analysis ($r^2 < 0.3$). Phenotype Definition: the definition of menopause was defined in the same manner as in the Multiethnic Cohort Study (see the following section). Population Stratification Correction: we used PCs analysis to estimate global ancestry among the 5761 individuals, using 2546 ancestry informative markers. To address the potential for confounding by genetic ancestry, we adjusted for global ancestry in all analyses.

The MEC is a prospective cohort study of 215,000 men and women in Hawaii and Los Angeles between the ages of 45 and 75 years at baseline (1993–1996) (71). Through December, 31 2007, a nested breast cancer case–control study in the MEC included 556 African-American cases (544 invasive and 12 in situ) and 1003 African-American controls. An additional 178 African-American breast cancer cases (ages: 50–84 years) diagnosed between June 1, 2006 and December 31, 2007 in Los Angeles County (but outside of the MEC) were combined with the MEC samples in the analysis. Genomic DNA was extracted for all samples using the QIAamp Blood Mini Kit (Qiagen, Valencia, CA, USA). Genotyping and Quality control: the genotyping and quality control procedures were identical to those used for CBCS. Imputation: the imputation procedures were identical to those used for CBCS. Phenotype Definition: age at last menstrual period cannot be used to uniformly estimate age at menopause because women who use sequential EPRT often continue to have monthly menstrual periods, irrespective of their ovarian function, and women on estrogen replacement therapy and continuous combined replacement therapy can rarely distinguish breakthrough bleeding from ovarian function-determined menses. Hence, for a woman taking HRT before her reported age at last menstrual period, we set her age of menopause as the year in which she began HRT use (excluding use of progestin alone), with the rationale that HRT use was started because of menopausal symptoms. Population Stratification Correction: the method for population stratification correction was identical to those used for CBCS.

Meta-analysis

We performed additional quality control on all genotyped and imputed SNPs in each cohort by removing SNPs in each cohort if they had <1% MAF, they had fewer than five copies of minor alleles, they were genotyped in fewer than 100 participants, or they had a quality score of <0.3 if they were imputed. In total, we have 3,252,746 SNPs that were found in at least one cohort available for analyses. In all cohorts, linear regression models were used to assess the association between each SNP and age at menopause. An additive genetic model was assumed using either genotyped SNPs or imputed SNP dosages. Global ancestry was adjusted by incorporating the first 10 PCs for each individual as covariates in the regression models. In GENOA and MESA family, additional covariates were included in the models to account for family structure in the cohort. In WHI and CHS, clinical sites were included as a covariate to correct for any geographical or socio-economic differences. Meta-analyses on the linear regression results were conducted using the Meta Analysis Helper (72). The genomic control correction was turned on and the analysis was based on effect sizes and their standard errors (SCHEME STDERR). Results for SNPs found in only one cohort or <1000 total participants were discarded.

We first performed a genome-wide discovery but no statistically significant associations were detected. To replicate previous findings, we compiled a list of SNPs previously associated with age at menopause in women with European ancestry. These SNPs included 15 SNPs discovered in GWAS with $P$-values $< 10^{-7}$ (distributed across five loci), nine SNPs with $P$-values between $10^{-4}$ and $10^{-7}$ (distributed across nine loci), 17 SNPs discovered by a GWAS meta-analysis (distributed across 17 loci, of which 16 were novel) and seven SNPs implicated in candidate gene studies (distributed across six loci) (17–19,27–33). The LD relationships among the SNPs in HapMap CEU in each region were carefully examined to determine the effective number of tests. Each region was tested at an $\alpha$-level of 0.05. We examined each region further by defining a LD block for each region using the HapMap CEU population. Each LD block was defined as the largest block within 500 kb of the index SNPs on which the LD between the SNPs at both ends of the block and the index SNPs have $r^2 \geq 0.5$. The LD relationships among the SNPs on the same LD block were investigated and only one SNP from high LD pairs ($r^2 \geq 0.8$) was kept.

For index SNPs that replicated successfully, we sought to fine-map the signal by comparing the difference in LD structures between HapMap CEU and YRI samples. Graphs depicting the relationships between SNPs on the same LD block were created using LocusZoom (73). We also searched for possible secondary signals in the LD block.

In the event that the index SNP did not replicate, we identified a list of SNPs with at least $r^2 \geq 0.8$ with the index SNPs in HapMap CEU population using the Genome Variation Server (http://gversusgs.washington.edu/GVS). We performed meta-analyses for all of them to confirm that they had similar $P$-values. We then assessed whether there was any signal on the LD block by examining the SNPs on the block using a conservative Bonferroni correction based on the number of SNPs tested. When a secondary signal was found, we performed conditional analyses in each cohort with both the index SNP and the newly identified SNP in the model. Using the results from these conditional analyses, we performed a conditional meta-analysis to confirm the existence of a secondary signal.

We determined the likely function of the SNPs we investigated by utilizing the results from the Encyclopedia of DNA Elements Project (ENCODE) (74). In particular, we used the information from the ENCODE Integrated Regulation and ENCODE Chromatin State Segmentation by HMM from Broad Institute, MIT and MGH tracks from the UCSC Genome Browser (39). The ENCODE Integrated Regulation track consists of information from the following source: transcription levels assayed by RNA-seq on six-cell lines, enhancer-
and promoter-associated histone mark (H3K4Me1) on eight-cell lines, enhancer- and promoter-associated histone mark (H3K27Ac) on eight-cell lines, promoter-associated histone mark (H3K4Me3) on nine-cell lines, digital DNsase hypersensitivity clusters and transcription factor Chip-seq data.

Power calculations were performed using Quanto (version 1.2.4) using the gene-only model with a continuous trait phenotype and unrelated individuals (75). For the age at menopause, we assumed a population mean of 50 years with a standard deviation of 5.7.

The summary-level P-values for the current study can be found in Supplementary Material, Table S5.

SUPPLEMENTARY MATERIAL
Supplementary Material is available at HMG online.

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