Genome-wide association meta-analysis of individuals of European ancestry identifies new loci explaining a substantial fraction of hair color variation and heritability

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Hair color is one of the most recognizable visual traits in European populations and is under strong genetic control. Here we report the results of a genome-wide association study meta-analysis of almost 300,000 participants of European descent. We identified 123 autosomal and one X-chromosome loci significantly associated with hair color; all but 13 are novel. Collectively, single-nucleotide polymorphisms associated with hair color within these loci explain 34.6% of red hair, 24.8% of blond hair, and 26.1% of black hair heritability in the study populations. These results confirm the polygenic nature of complex phenotypes and improve our understanding of melanin pigment metabolism in humans.

Human pigmentation refers to coloration of external tissues due to variations in quantity, ratio, and distribution of the two main types of the pigment melanin: eumelanin and pheomelanin. Most melanin is produced by melanosomes, large organelles specialized in melanin synthesis and transportation, located mainly in the epidermis, hair, and iris, as well as the central nervous system. Early humans had darkly pigmented skin, which protected against high ultraviolet radiation (UVR) and its consequences, such as skin cancer and folate depletion. European and Asian populations evolved lighter skin pigmentation as they migrated toward northern latitudes with lower UVR. The lighter pigmentation maximizes UVR absorption needed to maintain adequate vitamin D levels. In Europeans, pigmentation of skin, hair, and/or eyes has characteristic geographic distributions because of natural selection and perhaps genetic drift; a role for sexual selection has been debated.

Hair color is one of the most prominent traits in humans. Twin studies suggest that up to 97% of variation in hair color may be explained by heritable factors, and genome-wide association studies.
have identified several chromosomal regions associated with hair color and related pigmentation traits. Except for red hair, known variants have a relatively low predictive value, and the heritability gap remains relatively large, which suggests that many hair color genes remain undiscovered.

Here we report the results of a meta-analysis of two GWAS carried out in two large discovery cohort studies: 157,653 participants from the 23andMe, Inc. customer base and 133,238 individuals from the UK Biobank (UKBB). Participants in both studies self-reported their natural color of hair in adulthood (Supplementary Fig. 1 and Supplementary Note). For the purpose of this work, each hair color category collected (black, dark brown, light brown, red, and blond) was assigned numerical values ranging from lowest (blond) to highest (black). These codes were used as the outcome variable in linear regression-based GWAS analyses. To minimize population admixture and stratification, the analyses were restricted to individuals of European ancestry (Supplementary Figs. 2 and 3) and adjusted for the first ten principal components of the genotype matrix, as well as for age and sex.

The analyses confirmed a strong association between hair color and principal components, especially in the less ethnically homogeneous 23andMe dataset, which includes participants of more varied European origin, in line with the known north–south cline in hair color variation and other regional differences in hair color across Europe (Supplementary Table 1). The strongest associations in both groups were with sex (Table 1). Women had higher odds ratios (ORs) and were more likely to report blond (OR = 1.20 and OR = 1.29 in the 23andMe and UKBB participants, respectively) or red hair (OR = 1.72 and OR = 1.40, respectively) than any other color and three to five times less likely to report black hair (OR = 0.35 and OR = 0.20, respectively) compared to men.

Genomic inflation factors (λ_{GC}) from the 23andMe and the UKBB GWAS were 1.147 and 1.146, respectively, in line with expectations of high power to detect large polygenic effects in these large samples (Supplementary Fig. 4). Meta-analyzed GWAS results reached conventional genome-wide significance (P < 5 × 10^{-8}) in many regions, primarily clustering around 123 distinct autosomal genomic single-nucleotide polymorphisms (SNPs) and one X-chromosomal locus (Fig. 1 and Supplementary Table 2), mostly new (Table 2). In line with power expectations (Supplementary Fig. 5), 75 of these regions were significant genome-wide in at least one of the two cohorts and always at least nominally significant (P < 0.01) in the other.

Previously known pigmentation loci were all strongly associated in the meta-analysis results: HERC2 (rs12913832), IRF4 (rs12203592), and MCIR (rs1805007), as well as others, showed some of the strongest statistical evidence for association ever published for human complex traits. Strong associations were found for genes whose mutations reportedly cause impairment of pigmentation, such as Waardenburg (EDNRB, rs1279403, P < 10^{-10}; MITF, rs9823839, P < 10^{-10}); Hermansky–Pudlak (HPS5, rs201668750, P = 4.68 × 10^{-11}), trichomégal (FGF5, rs7681907, P = 5.684 × 10^{-20}), or Aplehamon macromastia (TWIST2, rs11684254, P = 1.233 × 10^{-20}) syndromes. Many polymorphisms significantly (P < 5 × 10^{-8}) associated with hair color in our meta-analysis had existing entries in the GWAS catalog. In previous publications, they were associated with several phenotypes, including pigmentation traits (Supplementary Table 3).

Among the associated loci, some of the strongest effects were observed for two solute carrier 45A family members (SLC45A1, rs80293268, P < 10^{-100}; and SLC45A2, rs16891982, P < 10^{-100}); polymorphisms near a third solute carrier gene were also significantly associated with hair color (rs60086398 upstream of SLC7A1, P = 4.93 × 10^{-9}). In addition, forkhead box family genes (FOXO6, rs3856254, P = 4.0 × 10^{-10}; and FOXE1, rs3021523, P = 4.23 × 10^{-20}) and sex-determining region Y (SRY)-box genes (SOX5, rs9971729, P = 8.8 × 10^{-12}; and SOX6, rs1531903, P = 9.1 × 10^{-16}) were among those highlighted in our results. An additional locus, located on chromosome X in the second intron of the collagen type IV alpha 6 gene, was also significantly associated (COL46A1, rs1266744, P = 5.03 × 10^{-12}). Chromosome X information was not analyzed. Notably, given the observed strong association of hair color with sex, there was no particular difference in effect sizes observed for these loci among men and women in either cohort (Supplementary Table 4 and Supplementary Fig. 6); only one SNP significantly associated with hair color in the meta-analysis showed significant

### Table 1 | Effect of sex on the hair color phenotypes in the 23andMe and UK Biobank cohorts

|        | 23andMe | UKBB |
|--------|---------|------|
| Blond  | 1.285   | 0.108 |
| Red    | 1.395   | 0.126 |
| Light brown | 1.101   | 0.114 |
| Dark brown | 0.993   | 0.11 |
| Black  | 0.195   | 0.033 |

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Table 2 | A selection of genes newly associated with hair color

| Chr | Position (Build 37) | SNP ID | Ref. allele | Freq | Nearest gene | n | Beta | Standard error | P | Beta | Standard error | P | Beta | Standard error | P |
|-----|---------------------|--------|------------|-------|--------------|---|------|---------------|---|------|---------------|---|------|---------------|---|
| 1   | 8207579             | rs80293268 | G          | 0.047 | SLCA5A1     | 122,221 | 0.194 | 0.009 | 1.54×10⁻⁴ | 157,651 | 0.157 | 0.009 | 1.29×10⁻⁷ | 0.175 | 0.007 | 1.9×10⁻⁶ |
| 2   | 205201062            | rs2369633 | T          | 0.089 | DSYK       | 132,887 | -0.071 | 0.007 | 9.20×10⁻⁴ | 157,651 | -0.077 | 0.006 | 3.15×10⁻⁴ | 0.075 | 0.005 | 3.44×10⁻⁴ |
| 2   | 28613302             | rs74443018 | G          | 0.039 | PDGF1       | 126,428 | 0.133 | 0.010 | 2.34×10⁻³ | 157,651 | 0.148 | 0.012 | 4.18×10⁻⁵ | 0.139 | 0.008 | 1.36×10⁻⁵ |
| 9   | 126808006            | rs58979950 | T          | 0.109 | UHCG       | 132,883 | 0.089 | 0.006 | 1.03×10⁻⁴ | 157,651 | 0.083 | 0.005 | 9.93×10⁻⁴ | 0.086 | 0.004 | 1.40×10⁻⁴ |
| 13  | 78393757             | rs1279403 | T          | 0.406 | ESRB       | 133,238 | -0.086 | 0.004 | <10⁻⁷ | 157,651 | -0.074 | 0.004 | 4.57×10⁻⁷ | 0.080 | 0.003 | <10⁻⁷ |
| 15  | 448246648            | rs1446664 | G          | 0.022 | SHC4       | 133,238 | 0.388 | 0.069 | 0.006 | 157,651 | 0.289 | 0.030 | 1.2×10⁻³ | 0.273 | 0.028 | 1.24×10⁻³ |
| 17  | 39951099             | rs17612447 | T          | 0.029 | XKR7       | 133,238 | 0.063 | 0.011 | 2.95×10⁻⁸ | 157,651 | 0.064 | 0.011 | 2.09×10⁻⁸ | 0.063 | 0.008 | 3.29×10⁻⁸ |
| 20  | 62660168             | rs73152911 | T          | 0.046 | JCAST      | 132,836 | 0.089 | 0.009 | 6.78×10⁻⁹ | 157,651 | 0.046 | 0.008 | 2.54×10⁻⁷ | 0.064 | 0.006 | 5.85×10⁻⁷ |

The selection was based on the strength of their effect, which is defined as the standardized linear regression coefficient. Results are given for the UK Biobank, 23andMe, and their meta-analysis, as well as for the meta-analysis results from the VisiGen Consortium. Linear models were generated from these results, and effect sizes (beta) are given in s.d. units. A, C, T, and G under the ‘Reference allele’ (ref. allele) field denote the nucleotide of the allele for which the effect size and allele frequencies (freq.) are reported. Frequencies are given for the reference allele and are the average of observed frequencies in the 23andMe and UK Biobank. Associations with P values of less than 10⁻⁸ are reported as P < 10⁻⁸.
for all hair colors, not just for the extremes of the light–dark hair color spectrum. Third, hair color is a trait that follows special geographic distribution patterns and therefore is prone to issues of population structure bias, which may be controlled in several ways. A comparison of different methodologies (Supplementary Fig. 10) shows that our approach is roughly equivalent to others. Fourth, annotation of the associated genetic regions based on physical distances most likely underestimates the number of regions involved in hair pigmentation. For example, the involvement of OCA2 and HERC2 genes in human pigmentation is not simply due to linkage disequilibrium6, yet because of their proximity, both loci in our study were assigned to the same association region. This would, however, not affect the conditional analysis at a marker level, which discriminates separate effects arising from within the same region.

In conclusion, this large GWAS meta-analysis has improved our knowledge of the genetic control of human hair pigmentation by bringing the number of known loci into the hundreds. The newly identified genetic loci explain substantial portions of the phenotypic variance of hair color spectrum and therefore is prone to issues of population structure bias, which may be controlled in several ways. A comparison of different methodologies (Supplementary Fig. 10) shows that our approach is roughly equivalent to others. Fourth, annotation of the associated genetic regions based on physical distances most likely underestimates the number of regions involved in hair pigmentation. For example, the involvement of OCA2 and HERC2 genes in human pigmentation is not simply due to linkage disequilibrium6, yet because of their proximity, both loci in our study were assigned to the same association region. This would, however, not affect the conditional analysis at a marker level, which discriminates separate effects arising from within the same region.

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Author contributions

P.G.H., A. Valdes, and F.L. jointly wrote the manuscript, coordinated meta-analyses, and performed prediction modeling. N.A.F., D.M.E., V.B., A. Visconti, G.H., G.M., S.M.R., D.I.B., N.G.M., and M.F. contributed populations samples and data used for analyses. M.K. and T.D.S. jointly coordinated the work and participated in manuscript preparation. D.A.M., D.I.B., N.G.M., and M.F. each conducted part of the analyses described in this work. G.D.S., P.G., C.M.v.D., M.A.I., D.T., M.C., A.R., S.Y., A.W.H., Y.C., C.Z., A.G.U., M.A.H., T.N., M.F., and D.A.H. each contributed to the analyses. N.A.F., D.M.E., V.B., A. Visconti, G.H., G.M., S.M.R., D.I.B., N.G.M., and M.F. contributed populations samples and data used for analyses. M.K. and T.D.S. jointly coordinated the work and participated in manuscript preparation.

Competing interests

N.A.F. and D.A.H. are employees of the 23andMe Inc., a consumer genetics company.

Additional information

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Methods

The 23andMe cohort. All research participants were drawn from the customer base of 23andMe, Inc., a consumer genetics company. This cohort has been described in detail previously11. All participants included in the analyses provided informed consent as approved surveys online according to our human subject protocol, which was reviewed and approved by Ethical & Independent Review Services, a private institutional review board (http://www.eandireview.com). Hair color phenotypes were used to create an ordinal trait with values: 0, light blond; 1, dark blond; 2, red; 3, light brown; 4, dark brown; 5, black. DNA extraction and genotyping were performed on saliva samples by CLIA-certified and CAP-accredited clinical laboratories of Laboratory Corporation of America. Samples were genotyped on one of four genotyping platforms. The V1 and V2 platforms were variants of the Illumina HumanHap550 + BeadChip, including about 25,000 custom SNPs selected by 23andMe for a total of about 500,000 SNPs. The V3 platform was based on the Illumina OmniExpress + BeadChip, with custom content targeted to target the overlap with our array, with a total of 950,000 SNPs. The V4 platform in current use is a fully custom array, including a lower-redundancy subset of V2 and V3 SNPs, with additional coverage of lower-frequency coding variations, and about 570,000 SNPs. Samples that failed to reach a 98.5% call rate were re-analyzed. For the GWAS only, participants with >97% European ancestry, as determined through an analysis of local ancestry, were included. For the purposes of ethnic categorization, an algorithm first partitioned phased genomic data into short windows of about 100 SNPs and used a support vector machine (SVM) to classify individual haplotypes into one of 31 reference populations. The SVM classifications then fed into a hidden Markov model (HMM) that accounts for switching errors and inserted gaps, and gives probabilities for each reference population in each window. The reference population data are derived from public datasets (the Human Genome Diversity Project, HapMap, and 1,000 Genomes), as well as 23andMe customers who have reported having four grandparents from the same country. A maximal set of unrelated individuals was chosen for each analysis using a segmental identity-by-descent (IBD) estimation algorithm12. Individuals were defined as related if they shared more than 700 cM IBD, including regions where the two individuals share either one or both genomic segments identical-by-descent. This level of relatedness corresponds approximately to the minimal expected sharing between first cousins in an outbred population.

Participant genotype data were imputed against the September 2013 release of 1,000 Genomes Phase 1 reference haplotypes, phased with ShapeIt26. We phased and imputed data for each genotyping platform separately. We phased using an internally developed phasing tool that implements the Beagle haplotype graph-based phasing algorithm26, modified to separate the haplotype graph construction and phasing steps.

SNPs with Hardy–Weinberg equilibrium $P < 10^{-8}$, call rate < 95%, or with large allele frequency discrepancies compared to European 1,000 Genomes reference data were excluded from imputation. Imputation was done against all-ethnicity 1,000 Genomes haplotypes (excluding monomorphic and singleton sites) using Minimac27. For the X chromosome, separate haplotype graphs were built for the non-pseudautosomal region and each pseudautosomal region, and these regions were phased separately. Male and females were imputed together using Minimac27, as with the autosomes, targeting males as homozygous pseudo-diploids for the non-pseudautosomal region.

Association test results were computed by linear regression, assuming additive allele effects. For the imputed dosages rather than best-guess genotypes were used. Covariates for age, gender, and the top five principal components to account for residual population structure were also included into the model. Results for the X chromosome were computed similarly, with male genotypes coded as if they were homozygous diploid for the observed allele.

HLA allele dosages were imputed from SNP genotype data using HIBAG28. We imputed allele dosages for HLA-A, B, C, DPB1, DQA1, DQB1, and DRB1 loci at four-digit resolution. To test associations between HLA allele dosages and phenotypes, we performed logistic or linear regression using the same set of covariates that as used in the SNP-based GWAS for that phenotype. We performed separate association tests for each imputed allele.

The UK Biobank. The UK Biobank database includes 502,682 participants who were aged from 49–69 years when recruited between 2006 and 2010 from across the UK to take part in the project. The study was approved by the National Research Ethics Committee (REC reference 11/NW/0382). The participants filled out several questionnaires about their lifestyle, environmental risk factors, and medical history and gave their informed consent29. The participants were invited, through a computerized questionnaire, to answer the question “What best describes your natural hair color?” (If your hair color is grey, the color before you went grey?). The participants’ answers were used to create a hair color variable with values: 1, blond; 2, red; 3, light brown; 4, dark brown; 5, black; and other codes for “other,” “don’t know,” or “prefer not to answer.” The later three values were removed from analyses.

Extracted DNA was then processed in the approximate order received to produce genotype data using the Affymetrix Axiom platform, as described elsewhere (see URLs). Details on genotyping procedure and quality control can be found elsewhere (see URLs). Phasing on the autosomes was carried out using a version of the SHAPEIT2 program modified to allow for very large sample sizes. The new algorithm uses a divisive clustering algorithm to identify clusters of haplotypes and calculates HMM distances only between pairs of haplotypes within each cluster. Only haplotypes within each cluster are used as candidates for the surrogate family copying states in the HMM model. Imputation was carried out using the same algorithm as is implemented in the IMPUTE2 program. More detailed information on the imputation procedure followed can be found elsewhere (see URLs). Linear models were built for the main GWAS analysis: haircolor genotype $+ \text{age} + \text{sex} + \text{PC1} + \text{genotyping platform}$. 

Replication cohorts (the VisiGen Consortium). Subjects were individuals of European descent participating in any of the 10 studies from the International Visible Trait Genetics Consortium (VisiGen). The VisiGen participants were phenotyped through self-report, and each phenotypic category was assigned a unique numerical value within each cohort. The self-reported hair color categories were, however, highly heterogeneous across the 10 studies. Therefore, all participants ($n = 27,865$) were genotyped, imputed, and analyzed separately by each participating center, using standard techniques described in the Supplementary Note.

Meta-analyses. Results from each participating cohort (23andMe and UK Biobank) were standardized to allow for the different scales (six categories of hair color in the former but only five in the latter) and minimize differences arising from the slightly different categorizations of the same phenotype. Both weighted z-scores and inverse variance regression analyses (the latter use weighted regression coefficients and standard errors) were calculated using Meta28. The results obtained from both methods were similar, and inverse-variance results are reported throughout the manuscript for the discovery cohort. Association effect sizes, standard errors, and probabilities were taken from the association analyses software. When the significance of the association exceeded the range of float numbers determined by the system and software, the probabilities were calculated using Mathematica 11.1 computational algebra (Wolfram Research Inc.). Meta-analyses of the replication cohorts were calculated using the weighted z-score method to reflect the fact that the phenotypic definitions were not harmonized across the different participating cohorts (please refer to the Supplementary Note for more detailed population description and phenotypic definitions).

Conditional and explained heritability analyses. The program OCTA30 was used for conditional analyses30 to identify independent effects within associated loci as well as to calculate the phenotypic variance explained33 by all polymorphisms genotyped or imputed, associated with the trait after the conditional analyses. The threshold of significance was set at $P < 5 \times 10^{-8}$ and the co-linearity threshold was $r^2 = 0.8$. These estimates were derived from the UKBB cohort.

Natural selection. Results of three statistical tests for natural selection were obtained from the 1,000 Genomes selection browser31. Results from three selected tests are reported: $HS^+$ and two cross-population comparison (XP–EHH tests, CEU vs. YRI, and CEU vs. CHB) based on the extended haplotype homozygosity test32. The absolute test scores and rank scores ($-\log_{10}$ of the centile of the absolute test score across the genome) for each SNP of interest are reported.

Prediction analyses. Using the independently associated SNPs identified in the discovery GWAS meta-analysis (Supplementary Table 9) together with previously reported hair color predicting SNPs from the HirisPlex System22,28, we performed association analyses software. When the significance of the association exceeded the range of float numbers determined by the system and software, the probabilities were calculated using Mathematica 11.1 computational algebra (Wolfram Research Inc.). Meta-analyses of the replication cohorts were calculated using the weighted z-score method to reflect the fact that the phenotypic definitions were not harmonized across the different participating cohorts (please refer to the Supplementary Note for more detailed population description and phenotypic definitions).

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$\text{Prediction analyses}$. $\text{Using the independently associated SNPs identified in the discovery GWAS meta-analysis (Supplementary Table 9) together with previously reported hair color predicting SNPs from the HirisPlex System}^{22,28} \text{, we performed hair color prediction modeling in the QIMR, RS, and combined QIMR + RS datasets. For this, we randomly split the QIMR, RS, and combined QIMR + RS data into 80% training sets and 20% validation sets, respectively.} \text{This approach assures the use of totally independent datasets for predictive marker discovery, model building, and model validation. Of the 258 independently associated SNPs in the discovery GWAS meta-analysis (see Supplementary Table 9), five SNPs failed independent RS quality control in the QIMR and one in the QIMR + RS and were therefore not included in the models. In the combined QIMR + RS analysis, we used the overlapping set of SNPs. The performance of the prediction models was evaluated in the respective validation datasets using the area under the receiver operating characteristic (ROC) curve (AUC). AUC is the integral of ROC curves, which ranges from 0.5, representing total lack of prediction, to 1.0, representing perfect prediction. Prediction analyses were conducted in R version 3.2.3 using relevant packages, including lars, mnet, and pROC.}$

$\text{Effects of variants on gene expression}$. $\text{The potential eQTL effects of the variants of interest were evaluated in all 37 tissues available at the GTEx}^{8} \text{ portal (see URLs). Associations with the levels of expression of adjacent genes were assessed for all variants identified through the conditional analysis.}$

$\text{Gene set enrichment analyses}$. $\text{Gene set enrichment analyses were carried out on summary results obtained from the meta-analysis of the UK Biobank and 23andMe subjects using Magenta software}^{33}.$
Data availability. This work used data from two primary sources. The original datasets can be accessed as follows.

For UK Biobank, data can be accessed through the UK Biobank Access management (https://www.ukbiobank.ac.uk/register-apply/). The hair color data accession codes are 1747.0.0, 1747.1.0, and 1747.2.0. The participants’ age UK Biobank accession code is 21022, for sex 31.0.0, and the precomputed principal components used here are 22009.0.1 through 22009.0.10.

For the 23andMe participants, requests for summary statistics can be accessed at https://researchers.23andme.org/collaborations. There are no accession codes available.

For the TwinsUK datasets, access can be requested through http://www.twinsuk.ac.uk/data-access/, and access to the secondary source of data through the corresponding authors.

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1. Sample size
   Describe how sample size was determined.
   This work used existing samples, the most important of which were the UK Biobank and 23andMe customers. The UK Biobank was designed specifically for research in epidemiology, is the most powered cohort to date (up to 500,000 currently, of which ~130,000 were used for the purpose of the manuscript) and together with the 23andMe cohort they have exceptional power to detect genetic variants at even effect sizes or minor allele frequencies far beyond common publication benchmarks.

2. Data exclusions
   Describe any data exclusions.
   To avoid issues related to population structure, only individuals of European origin were included in this study.

3. Replication
   Describe whether the experimental findings were reliably reproduced.
   This study is primarily focused on two major cohorts, each exceeding 100,000 subjects. These cohorts reliably replicate each-other’s discoveries, but as an additional precaution, these results were compared to results pooled from a meta-analysis of several smaller cohorts, members of the Visigen academic consortium.

4. Randomization
   Describe how samples/organisms/participants were allocated into experimental groups.
   This study did not involve any intervention.

5. Blinding
   Describe whether the investigators were blinded to group allocation during data collection and/or analysis.
   The investigators did not participate in data collection and only analyzed data made available to them. The investigators were blind to any individual genotypic or phenotypic status.

Note: all studies involving animals and/or human research participants must disclose whether blinding and randomization were used.
6. Statistical parameters

For all figures and tables that use statistical methods, confirm that the following items are present in relevant figure legends (or the Methods section if additional space is needed).

- [ ] Confirmed
- [ ] The exact sample size \((n)\) for each experimental group/condition, given as a discrete number and unit of measurement (animals, litters, cultures, etc.)
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- [ ] The test results (e.g. \(p\) values) given as exact values whenever possible and with confidence intervals noted
- [ ] A summary of the descriptive statistics, including central tendency (e.g. median, mean) and variation (e.g. standard deviation, interquartile range)
- [ ] Clearly defined error bars

See the web collection on statistics for biologists for further resources and guidance.

7. Software

Describe the software used to analyze the data in this study. Different software was used at different stages of the analyses. For example, regression models were built and assessed using PLINK, meta-analysis was run using METAL and GWAMA (as results were identical only the former were reported). Conditional analyses and estimates of the proportions of heritability explained were run using the GCTA software. Natural selection signals were assessed using data generated by others using the iHS and XP-EHH methods, downloadable from the 1000 Genomes Browser. R base packages as well as the glmnet, lars, nnet and pROC were used for the prediction models. The gene set enrichment analysis was run on the software Magenta.

For all studies, we encourage code deposition in a community repository (e.g. GitHub). Authors must make computer code available to editors and reviewers upon request. The Nature Methods guidance for providing algorithms and software for publication may be useful for any submission.

8. Materials availability

Indicate whether there are restrictions on availability of unique materials or if these materials are only available for distribution by a for-profit company.

This manuscript reports no experimental results, only statistical analyses

9. Antibodies

Describe the antibodies used and how they were validated for use in the system under study (i.e. assay and species).

No antibodies were used in this study.

10. Eukaryotic cell lines

a. State the source of each eukaryotic cell line used.

b. Describe the method of cell line authentication used.

c. Report whether the cell lines were tested for mycoplasma contamination.

d. If any of the cell lines used in the paper are listed in the database of commonly misidentified cell lines maintained by ICLAC, provide a scientific rationale for their use.

No eukaryotic cell lines were used in this study.

No cell lines were used in this study.

No cell lines and no mycoplasma was used in this study.

No cell lines were used in this study.
Animals and human research participants

Policy information about studies involving animals; when reporting animal research, follow the ARRIVE guidelines

11. Description of research animals
   Provide details on animals and/or animal-derived materials used in the study.
   This manuscript describes only an observational genetic epidemiological work.

Policy information about studies involving human research participants

12. Description of human research participants
   Describe the covariate-relevant population characteristics of the human research participants.
   This was an observational study, involving no experiment. The observations were from questionnaires answered by volunteers of European origin, who reported their natural hair color, mostly in adult, or very late childhood age.