



Research paper

Population and evolutionary genetics of the *PAH* locus to uncover overdominance and adaptive mechanisms in phenylketonuria: Results from a multiethnic study



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ABSTRACT

Background: Phenylketonuria (PKU) is the most common inborn error of amino acid metabolism in Europe. The reasons underlying the high prevalence of heterozygous carriers are not clearly understood. We aimed to look for pathogenic *PAH* variant enrichment according to geographical areas and patients' ethnicity using a multiethnic nationwide cohort of patients with PKU in France. We subsequently appraised the population differentiation, balancing selection and the molecular evolutionary history of the *PAH* locus.

Methods: The French nationwide PKU study included patients who have been referred at the national level to the University Hospital of Nancy, and for whom a molecular diagnosis of phenylketonuria was made by Sanger sequencing. We performed enrichment analyses by comparing alternative allele frequencies using Fisher's exact test with Bonferroni adjustment. We estimated the amount of genetic differentiation among

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populations using Wright's fixation index (*Fst*). To estimate the molecular evolutionary history of the *PAH* gene, we performed phylogenetic and evolutionary analyses using whole-genome and exome-sequencing data from healthy individuals and non-PKU patients, respectively. Finally, we used exome-wide association study to decipher potential genetic loci associated with population divergence on *PAH*.

Findings: The study included 696 patients and revealed 132 pathogenic *PAH* variants. Three geographical areas showed significant enrichment for a pathogenic *PAH* variant: North of France (p.Arg243Leu), North-West of France (p.Leu348Val), and Mediterranean coast (p.Ala403Val). One *PAH* variant (p.Glu280Gln) was significantly enriched among North-Africans (OR = 23.23; 95% CI: 9.75–55.38). *PAH* variants exhibiting a strong genetic differentiation were significantly enriched in the 'Biopterin_H' domain (OR = 6.45; 95% CI: 1.99–20.84), suggesting a balancing selection pressure on the biopterin function of *PAH*. Phylogenetic and timetree analyses were consistent with population differentiation events on European-, African-, and Asian-ancestry populations. The five *PAH* variants most strongly associated with a high selection pressure were phylogenetically close and were located within the biopterin domain coding region of *PAH* or in its vicinity. Among the non-*PAH* loci potentially associated with population divergence, two reached exome-wide significance: *SSPO* (SCO-spondin) and *DBH* (dopamine beta-hydroxylase), involved in neuroprotection and metabolic adaptation, respectively.

Interpretation: Our data provide evidence on the combination of evolutionary and adaptive events in populations with distinct ancestries, which may explain the overdominance of some genetic variants on *PAH*.

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1. Introduction

Phenylketonuria (PKU) is an inherited autosomal recessive disorder of phenylalanine metabolism caused by a deficiency in the enzyme phenylalanine hydroxylase (*PAH*). The *PAH* gene (HGNC: 8582) located in 12q23.2 codes for the *PAH* enzyme that converts phenylalanine into tyrosine in the presence of molecular oxygen and catalytic amounts of tetrahydrobiopterin (BH4) [1,2]. Clinical manifestations of PKU are related to the toxic accumulation of phenylalanine in the blood and brain and, if left untreated, could lead to irreversible severe physical, neurological, and cognitive abnormalities.

During human peopling of the world and the different processes of human migration over time, several subgroups have progressively differentiated and experienced specific stresses and environmental conditions [3–5]. Numerous factors, including geographic isolation and adaptation, led to a divergence between human populations with regards to their genetic heritage, a concept known as population stratification [3–5]. The study of population stratification helped to better evaluate the influence of genetic determinants on human phenotypes, including disease phenotypes [6]. Fine-scale population stratification has been described in Europe and France with rare genetic variants that tend to be more geographically localised than common variants [7,8]. The geographical borders that surround the French territory and the several waves of population migration (e.g., Celts, Barbarians, Roman empire) have impacted the genetic structure of the modern French population [7,8].

PKU is the most common inborn error of amino acid metabolism in Europe, with an estimated prevalence of heterozygous carriers of *PAH* pathogenic variants around 2–3% [9,10]. From an evolutionary perspective and population genetics point of view, Krawczak & Zschocke nicely demonstrated that the high prevalence burden of heterozygous carriers for pathogenic *PAH* variants could not have resulted from a high mutation rate in the *PAH* locus or a high initial frequency of *PAH* gene mutations [10]. In this context, the use of the PKU model as a study framework for population genetics seems relevant, mainly because of the large variability of mutations in the *PAH* gene, an aggregated alternative allele frequency (AAF) of pathogenic variants on the *PAH* gene ~ 1% in Caucasian and Asian Oriental populations, and the ethnic diversity of populations affected by the PKU.

Several studies have listed the *PAH* gene variants observed among patients with PKU [11–20]. These studies assessed the

genotype-phenotype correlation between the *PAH* gene variants and the PKU phenotype. None of these studies was explicitly devoted to the evaluation of population divergence on the *PAH* locus. In the present study, we combined Sanger sequencing and exome-wide association study on a multiethnic nationwide cohort of patients with PKU in France to assess the effects of population divergence and balancing selection on the *PAH* locus. We also performed phylogenetic and evolutionary analyses using whole-genome and exome-sequencing data on healthy individuals and non-PKU patients to better appraise the sequence of phylogenetic events on the *PAH* locus.

2. Patients and methods

2.1. Patients population

The French nationwide PKU study included patients who have been referred at the national level to the Division of Biochemistry and Molecular Biology of the University Hospital of Nancy, and for whom a molecular diagnosis of phenylketonuria was made by Sanger sequencing (Figure S1). Parental and patient consent was obtained after oral and written information. All eligible patients were included in the study, and no patient declined to participate.

2.2. Study aims

The aims of the study were: i) to describe the pathogenic *PAH* variants and to look for potential pathogenic variant enrichment according to main geographical areas in France; ii) to perform ancestry marker analysis and look for potential pathogenic *PAH* variant enrichment according to patients' ethnicity; and iii) to assess the potential involvement of balancing selection on the *PAH* locus using population divergence analysis.

2.3. Sanger sequencing of the *PAH* locus

Genomic DNA from EDTA-treated peripheral blood samples was isolated using the Nucleon™ BACC3 Kit (GE Healthcare, Aulnay-sous-Bois, France). The 13 exons of the *PAH* gene were sequenced by bi-directional sequencing using the BigDye® Terminator v.1.1 cycle sequencing (Applied Biosystems, Foster, Ca, USA) and capillary electrophoresis (Applied Biosystems 3130 Genetic Analyzer). The PCR primer sequences are shown in Supplemental Table S1.

Research in context

Evidence before this study

Phenylketonuria (PKU) is the most common inborn error of amino acid metabolism in Europe, with an estimated prevalence of heterozygous carriers of *PAH* pathogenic variants around 2–3%. The reasons underlying the high prevalence of heterozygous carriers are not clearly understood. Numerous factors, including geographic isolation and adaptation, led to a divergence between human populations with regards to their genetic heritage, a concept known as population stratification. In the present study, we combined Sanger sequencing and exome-wide association study on a multiethnic nationwide cohort of patients with PKU in France to assess the effects of population divergence and balancing selection on the *PAH* locus. To better appraise the sequence of phylogenetic events on the *PAH* locus, we also performed phylogenetic and evolutionary analyses using whole-genome and exome-sequencing data from healthy individuals and non-PKU patients, respectively.

Added value of this study

Our study confirmed the high mutation burden on the *PAH* gene by reporting more than 130 pathogenic variants. Interestingly, within the *PAH* locus, the variants that were subject to a high population divergence were significantly enriched in the C-terminal catalytic 'Biopterin_H' domain by comparison to the N-terminal regulatory 'ACT domain', suggesting a potential role of balancing selection pressures in the shaping of the *PAH* locus. Phylogenetic and time-tree analyses on the *PAH* locus were consistent with population differentiation events on European-, African-, and Asian-ancestry populations. The five *PAH* variants most strongly associated with a high selection pressure were phylogenetically close and were located within the biopterin domain coding region of *PAH* or in its vicinity. Among the loci potentially associated with population divergence, two reached exome-wide significance and were positioned on the *SSPO* (SCO-spondin) and *DBH* (dopamine beta-hydroxylase) genes. *SSPO* and *DBH* are involved in neuroprotection and metabolic adaptation, respectively. Interestingly, the balancing selection pressure on the *PAH* locus, especially on the biopterin domain, may reflect a mechanism of adaptation to environmental factors but also to other enzymatic reactions that may involve the common BH4 cofactor in the *PAH* and *NOS* pathways.

Implications of all the available evidence

Our findings suggest a contributing role of population divergence and balanced selection to uncover heterozygote advantage and overdominance mechanisms on the *PAH* gene. Furthermore, our data provide a new paradigm regarding the combination of evolutionary and adaptive events in various populations, which may explain the overdominance of some genetic variants on the *PAH* gene. Future studies are needed to decipher the adaptive mechanisms that underlie the influence of *PAH*, *SSPO*, and *DBH* genes, including those related to overdominance and heterozygote advantage, in response to the selection pressures exerted on the *PAH* gene.

2.4. Exome-wide genotyping

High-throughput genotyping of DNA samples from patients included in the study was performed at the INSERM Unit UMR_S 1256 (NGERE) (UMS2008 IBSLor) using the Illumina Infinium HumanExome BeadChip v1.2 (Illumina Inc., San Diego, CA, USA)

according to the manufacturer's recommendations. The Illumina HumanExome BeadChip includes 247,870 markers focused on protein-altering variants selected from >12,000 exome and genome sequences representing multiple ethnicities and complex traits [21]. Additional array content includes variants associated with complex traits in previous GWAS, HLA tags, ancestry-informative markers ($n = 4,388$), markers for identity-by-descent estimation, and random synonymous SNPs [21]. Bead intensity data were processed and normalised for each sample Illumina's GenomeStudio Genotyping Module. The Genome Reference Consortium GRCh37 (hg19; Feb. 2009) map was used (Illumina manifest file ImmunoBead-Chip_11419691_B.bpm), and normalised probe intensities were extracted for all samples that passed standard laboratory quality-control thresholds. Genotype calling was performed with the GenomeStudio GenTrain 2.0 algorithm (Illumina's GenomeStudio data analysis software). Sample quality control measures included: sample call rate (>90%), overall heterozygosity, and relatedness testing. Duplicated samples and those showing cryptic relatedness were assessed by calculating identity by descent (IBD). We estimated IBD sharing and carried out PCA on an LD-pruned set of autosomal variants obtained by removing large-scale high-LD regions [22,23], variants with a call rate <0.99, variants with a minor allele frequency (MAF) <5% or variants with Hardy-Weinberg equilibrium (HWE) P -value < 10^{-4} . By estimating the probability of sharing zero, one, or two alleles for any two individuals as $P(Z = 0)$, $P(Z = 1)$, and $P(Z = 2)$, respectively, a proportion of IBD was calculated using the following formula: $PI-HAT = PI = P(Z = 1)/2 + P(Z = 2)$. A PI-HAT threshold >0.2 was used to suggest cryptic relatedness. Genetic variants were removed from the primary analysis if they had a call rate <95%, a significant departure from Hardy-Weinberg equilibrium (exact HWE- $P < 10^{-4}$), or a MAF <0.1%.

2.5. Next-generation sequencing (Nancy Reference Centre Population)

We used the next-generation sequencing (NGS) data that was generated from the molecular diagnostics activity conducted on non-PKU patients in the Department of Molecular Medicine at the University Hospital of Nancy, Reference Centre for Inborn Errors of Metabolism (Nancy, France). NGS analysis was carried out on an Illumina MiSeq sequencer using the TruSight One sequencing panel, targeting 4813 genes, including the *PAH* gene (Illumina, San Diego, USA). Among the genetic variants with validated quality metrics (read depth >20, alt-read ratio >0.2, and genotype quality of 99), only those on the *PAH* gene were considered for the analysis in the present study. Variants were aligned to GRCh37. All the variants retrieved on the *PAH* locus were visually inspected by checking the BAM files alignment.

2.6. Extraction of whole-genome sequencing data of ancestral populations (1 kG population)

We extracted whole-genome sequencing data from the 1 kG project phase 3 which provides a database of genomic variants of 2504 individuals across five ancestral populations: Africans (AFR, $n = 661$), Europeans (EUR, $n = 503$), Admixed Americans (AMR, $n = 347$), East Asians (EAS, $n = 504$), and South Asians (SAS, $n = 489$) [24]. Genomic and phenotypic data were downloaded at <http://www.internationalgenome.org/data/>. Variants were aligned to GRCh37. The total number of genetic variants available in chromosome 12 was 3,985,172, including 2387 markers in the *PAH* gene.

2.7. Principal component analysis and ancestry inference

Principal-component analysis (PCA) was performed on the samples merged with 1092 subjects from the 1 kG phase 1 super populations as reference populations (Africans (AFR), $n = 246$; Americans (AMR), $n = 181$; Asians (ASN), $n = 286$; and Europeans (EUR), $n = 379$)

(data were retrieved at: <http://www.internationalgenome.org/data/>). PCA analysis was performed under the additive model on 116,880 markers to find up to 10 components (Eigenvalues). PCA analysis results were reported using two-dimensional plots based on the two top Eigenvalues. PKU patients were plotted against all the 1092 subjects of the 1 kG phase 1 project. We used the same approach for PCA analysis on NGS data from patients included in the Department of Molecular Medicine at the University Hospital of Nancy.

2.8. Statistical analysis

We performed the enrichment analysis by comparing AAFs using Fisher's exact test with Bonferroni adjustment. Enrichment odds ratios and their 95% CI were reported. We estimated the amount of genetic differentiation among populations using the Wright's fixation index F_{st} (co-ancestry Coefficient θ) [25–27]. We calculated the 95% CI around the F_{st} using a percentile- t bootstrapping technique [28]. We estimated the fixation index F_{st} between pairs of subpopulations and the overall F_{st} index across all the studied populations. We used the following thresholds for F_{st} interpretation according to Hartl and Clark [26]: $F_{st} < 0.05$: little genetic differentiation; $F_{st} [0.05–0.15]$: moderate genetic differentiation; $F_{st} [0.15–0.25]$: great genetic differentiation; $F_{st} > 0.25$: very great genetic differentiation. To better visualise F_{st} indexes between pairs of subpopulations, we used dendrogram with heat map representation according to the following criteria: distance metric for top dendrogram: Euclidean; linkage algorithm for top dendrogram: weighted; distance metric for side dendrogram: Euclidean; and linkage algorithm for side dendrogram: weighted. To assess the haplotype structure of the *PAH* locus, we performed haplotype block analysis using the following criteria: upper confidence bound: 0.98; lower confidence bound: 0.7; reject criteria threshold: 0.9; MAF threshold: 0.05; Maximum number of markers in a block: 30; maximum length of a block: 160; display threshold: 0.01, maximum EM iterations: 50; and EM convergence tolerance: 0.0001. Linkage disequilibrium (LD)-pairwise analysis was performed on all adjacent pairs of genetic variants in a chromosome or within a haplotype block using a matrix output for both the expectation-maximization algorithm (EM) and the composite-haplotype method (CHM) [29,30]. R^2 and D' values were used in the Linkage disequilibrium plots. We performed the exome wide-association study through haplotype trend regression analysis to assess potential genetic loci associated with population differentiation and balancing selection pressure. For the haplotype association test, we used a moving window with a dynamic width of 10 kb and calculated the association per haplotype on 37,933 haplotype blocks. We estimated haplotype frequencies using the expectation/maximization (EM) algorithm with maximum EM iterations of 50 and an EM convergence tolerance of 0.0001 [31]. We compared haplotype frequencies using the Chi-squared test with Bonferroni and false discovery rate (FDR) adjustment. In a second step, we performed per-variant analysis on top significant loci obtained from exome-wide haplotype regression analysis by using linear regression with Bonferroni and FDR adjustment for assessing the same outcome. All quality controls and statistical analyses were performed using the SNP & Variation Suite (v8.8.3; Golden Helix, Inc., Bozeman, MT, USA).

2.9. Phylogenetic analyses of the *PAH* locus

We used two datasets for phylogenetic analyses: i) the 1 kG dataset that included all the *PAH* variants that exhibit the highest population divergence ($F_{st} > 0.50$ for AFR- vs EAS-ancestry differentiation) ($n = 64$) and ii) the 'Nancy Reference Centre Population' dataset that included all the *PAH* variants that were reported through NGS analysis ($n = 35$). We extracted all the potential DNA sequences on the *PAH* locus taking into account the *PAH* variants in their reference and alternative forms to produce DNA

FASTA files using the SeqTailor tool [32]. Phylogenetic trees were developed using the maximum likelihood-based method on aligned sequences. The evolutionary history was inferred by using the maximum likelihood method and the Tamura-Nei model [33]. Initial trees for the heuristic search were obtained automatically by applying Neighbour-Join and BioNJ algorithms to a matrix of pairwise distances estimated using the maximum composite likelihood approach and then selecting the topology with superior log likelihood value. Bootstraps with 500 replicates were applied. The phylogenetic tree was drawn to scale, with branch lengths measured in the number of substitutions per site. We conducted a timetree analysis using the RelTime method [34]. Divergence times for all branching points in the topology were calculated using the maximum likelihood method and Tamura-Nei model [33]. The timetree was drawn to scale, with branch lengths measured in the relative number of substitutions per site. The final timetree, showing relative node times, was normalised such that the maximum relative node time is 1. Phylogenetic and molecular evolutionary analyses were conducted using MEGA version X [35].

3. Results

3.1. Description of the pathogenic *PAH* variants found in the French Nationwide PKU Study

Between January 2004 and December 2012, 696 patients had a molecular diagnosis of PKU at the Division of Biochemistry and Molecular Biology of the University Hospital of Nancy using Sanger sequencing. Patients were recruited from seven geographical areas: North, North-East, North-West, Paris area, South-East, South-West, and Mediterranean coast (Figure S2). The descriptive statistics regarding the geographical origin and parent's ethnicity are available in Table 1. A total of 132 pathogenic *PAH* variants were found (Supplemental Table S2). The top three pathogenic

Table 1
Characteristics of the 696 patients included in the French Nationwide PKU Study.

Gender	Number of patients	Percentage
Female	364	52.3
Male	332	47.7
Geographical area / Main recruitment centre	Number of patients	Percentage
Mediterranean coast	158	22.7
North-West of France	151	21.7
North-East of France	115	16.5
North of France	96	13.8
South-East of France	87	12.5
Paris area	55	7.9
South-West of France	34	4.9
Ethnicity of the Mother*	Number of patients	Percentage
European Caucasian	395	71.3
North African	82	14.8
South European	37	6.7
Turkish	17	3.1
East European	10	1.8
Other	6	1.1
African sub-Saharan	3	0.5
Admixed: Caucasian/South European	3	0.5
Admixed: Caucasian/ North Africans	1	0.2
Ethnicity of the Father†	Number of patients	Percentage
European Caucasian	405	73.6
North African	78	14.2
South European	25	4.5
Turkish	17	3.1
East European	14	2.5
Other	8	1.5
Admixed: Caucasian/South European	3	0.5

* data available in 554 cases.

† data available in 550 cases.

PAH variants were: rs5030855 (7.97%; intronic variant, c.1066–11G>A), rs5030858 (6.11%; Missense, c.1222C>T, p. Arg408Trp), and rs5030861 (4.60%; splice donor variant, c.1315+1G>A). All the details related to the genomic description and the AAFs of the 132 PAH variants are reported in Supplemental Tables S2 and S3, respectively.

3.2. Enrichment analysis according to the geographical area

We performed an enrichment analysis to assess whether some pathogenic PAH variants were significantly enriched in a prespecified geographical area. Three geographical areas showed significant enrichment for a pathogenic PAH variant: rs62508588 (c.728G>T, p. Arg243Leu) in the North of France (OR = 22.41; 95% CI: 4.31–116.59; $P = 1.96 \times 10^{-4}$), rs62516092 (c.1042C>G, p.Leu348Val) in the North-West of France (OR = 4.29; 95% CI: 2.08–8.85; $P = 8.16 \times 10^{-5}$), and rs5030857 (c.1208C>T, p.Ala403Val) in the Mediterranean coast (OR = 4.88; 95% CI: 2.73–8.73; $P = 1.14 \times 10^{-7}$) (Supplemental Table S4, Fig. 1 and Supplemental Figure S2). The AAFs of all the PAH variants are reported in Supplemental Table S3. Enrichment analyses for all geographical areas are reported in Supplemental Tables S5–S11.

3.3. Enrichment analysis according to ethnicity

Among the 696 patients included in the study, 576 were studied using the Illumina Infinium HumanExome BeadChip v1.2. We performed PCA analysis on the 576 samples merged with 1092 subjects from the 1 kG phase 1 super populations (AFR, AMR, EAS, EUR, and

SAS) as reference populations (Fig. 2). The PCA analysis demonstrated that the study population was represented by two main categories: a subpopulation with a European-background and a subpopulation with a genomic background that lies between European-ancestry subjects and those with African-ancestry subjects (Fig. 2). In sensitivity analyses, the geographical areas of the recruiting centre were not associated with a specific clustering pattern on the PCA plot (Figure S3). When the ethnicity of PKU patients was considered as a classification variable, the PCA plot exhibited a typical clustering pattern with Caucasian-ancestry PKU patients that clustered around the EUR-ancestry HapMap populations and the North-African PKU patients that clustered between the EUR- and the AFR-ancestry HapMap populations (Fig. 3A).

We performed an enrichment analysis to assess whether some variants were significantly enriched among Caucasian- and North-African ancestry PKU patients. Among the 132 PAH variants, only one (rs62508698 [c.838G>C, p.Glu280Gln]) was significantly enriched among North-African PKU patients (OR = 23.23; 95% CI: 9.75–55.38; $P = 8.47 \times 10^{-13}$). The same variant was significantly under-represented among Caucasian-ancestry PKU patients (OR = 0.11; 95% CI: 0.05–0.26; $P = 3.55 \times 10^{-8}$) (Table 2). Enrichment analyses, according to the patient's ethnicity, are reported in Supplemental Tables S12–S14.

3.4. Assessment of the population divergence on the PAH locus

Given the highly discriminant pattern of the PAH variant rs62508698 between Caucasian-ancestry PKU and North-African PKU patients, we performed fixation index analysis by calculating the F_{st}

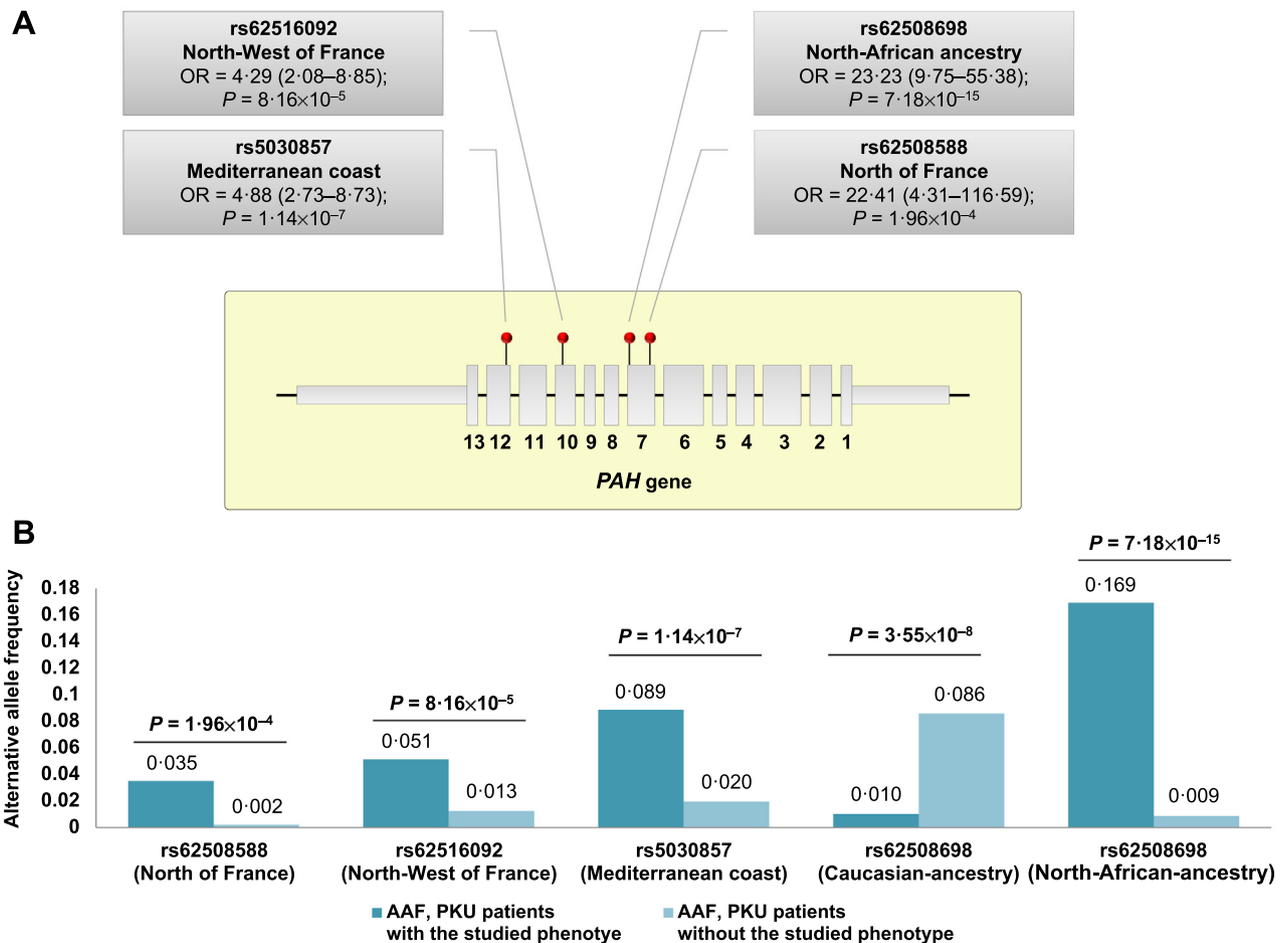


Fig. 1. (A) Genomic position and (B) alternative allele frequencies of the three pathogenic PAH gene variants enriched in the North (rs62508588), North-West (rs62516092) and Mediterranean coastal (rs5030857) areas of France and the variant enriched among patients with North-African ancestry (rs62508698).

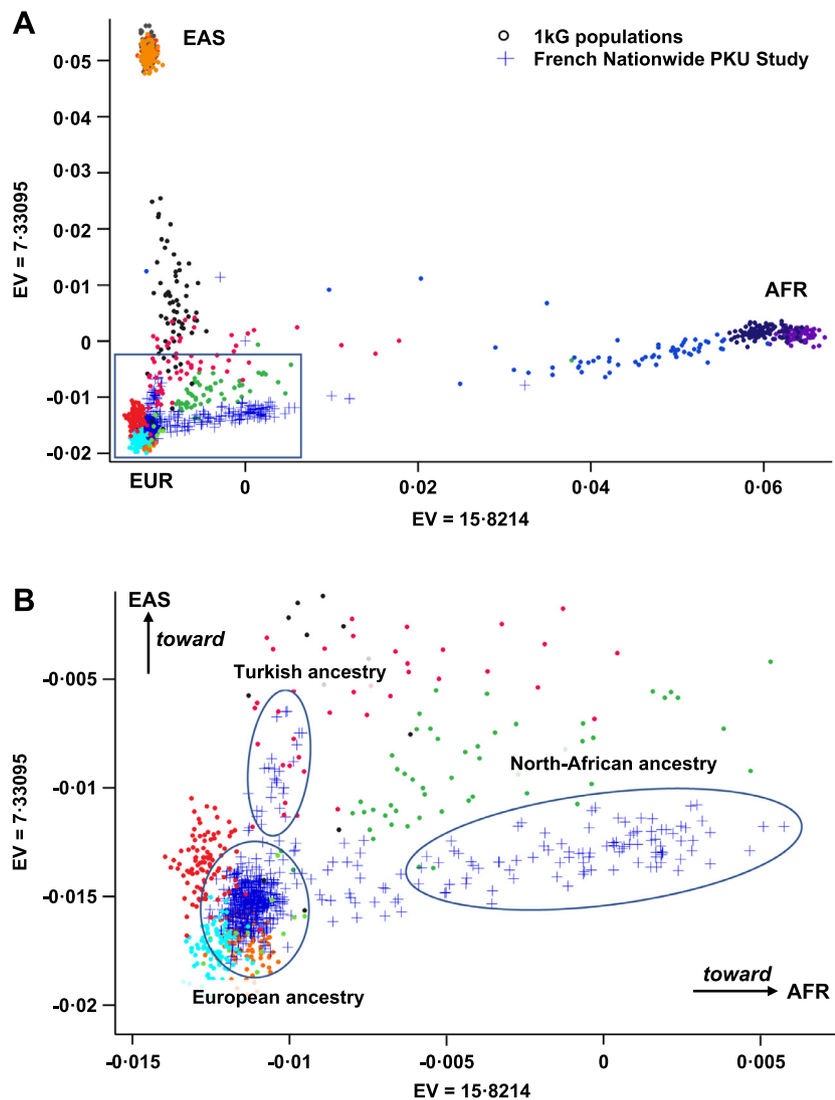


Fig. 2. (A) Principal component analysis reporting patients included in the French nationwide PKU study. (B) Zoomed view of the Fig. part defined by the inset in panel A. According to PCA-defined ancestry analysis, European-, North-African, and Turkish-ancestry patients represent the three main patients' ethnicities in the study.

index—which quantifies genetic differentiation between populations—on the 132 pathogenic *PAH* variants found in the study. The fixation index analysis confirmed that the rs62508698 variant exhibited the top *F_{st}* value (overall *F_{st}* = 0.249) across all ethnicities, which suggests a strong genetic differentiation [26] (Fig. 3B). The *F_{st}* value was even higher when the Caucasian- and North-African ancestries were considered (*F_{st}* = 0.280) (Supplemental Table S15). Considering the 132 variants in the *PAH* locus, the highest *F_{st}* values were observed in the 'Biopterin_H' domain (Figure S4A).

We used WGS data from the 2504 subjects included in the 1 kG project phase 3 and confirmed that the density of genetic variants with an *F_{st}* value ≥ 0.25 (strong genetic differentiation [26]) was significantly higher in the 'Biopterin_H' domain (1 variant per 0.936 kb) when compared to that observed in the 'ACT' domain (1 variant per 6.033 kb) with an OR of 6.45 (95% CI: 1.99–20.84; $P = 1.27 \times 10^{-4}$). The relatively high density of variants exhibiting an *F_{st}* value ≥ 0.25 in the 'Biopterin_H' domain is suggestive of a balancing selection pressure while the relatively low *F_{st}* values on the ACT domain was rather suggestive of a neutral selection across populations (Figure S4B).

We used WGS data from the 1KG project to calculate the overall *F_{st}* values using the 2387 genetic variants on the *PAH* locus by comparing the five 1 kG super populations: AFR, AMR, EAS, EUR, and SAS.

The *PAH* locus exhibited a population differentiation pattern, which was consistent with a significant balancing selection pressure with an overall *F_{st}* value of 0.16, indicating a great genetic differentiation (Supplemental Table S16). The top *F_{st}* values were observed between the AFR–EAS (0.34, 95% CI: 0.31–0.38), EAS–AMR (0.30, 95% CI: 0.27–0.33), and EUR–EAS (0.23, 95% CI: 0.20–0.25) indicating great to very great genetic differentiation (Fig. 4). We assessed the haplotype structure of the *PAH* locus in the whole 1 kG population and each super population using the same dataset of WGS. The haplotypic structure of the *PAH* locus significantly differed according to the 1 kG super population with a more complex structure in the AFR population (37 haplotype blocks) when compared to the less complex haplotype structure in the European population (21 haplotype blocks) (Fig. 5).

3.5. Phylogenetic analyses on the *PAH* locus

The phylogenetic analysis of the 1 kG dataset highlighted five clades (Fig. 6). Among them, one exhibited clustering of five intronic *PAH* variants that were highly differentiated between African- and East Asian-ancestry populations with an *F_{st}* > 0.75. Three variants located in the 'Biopterin_H' domain coding region (12:103265899-G-

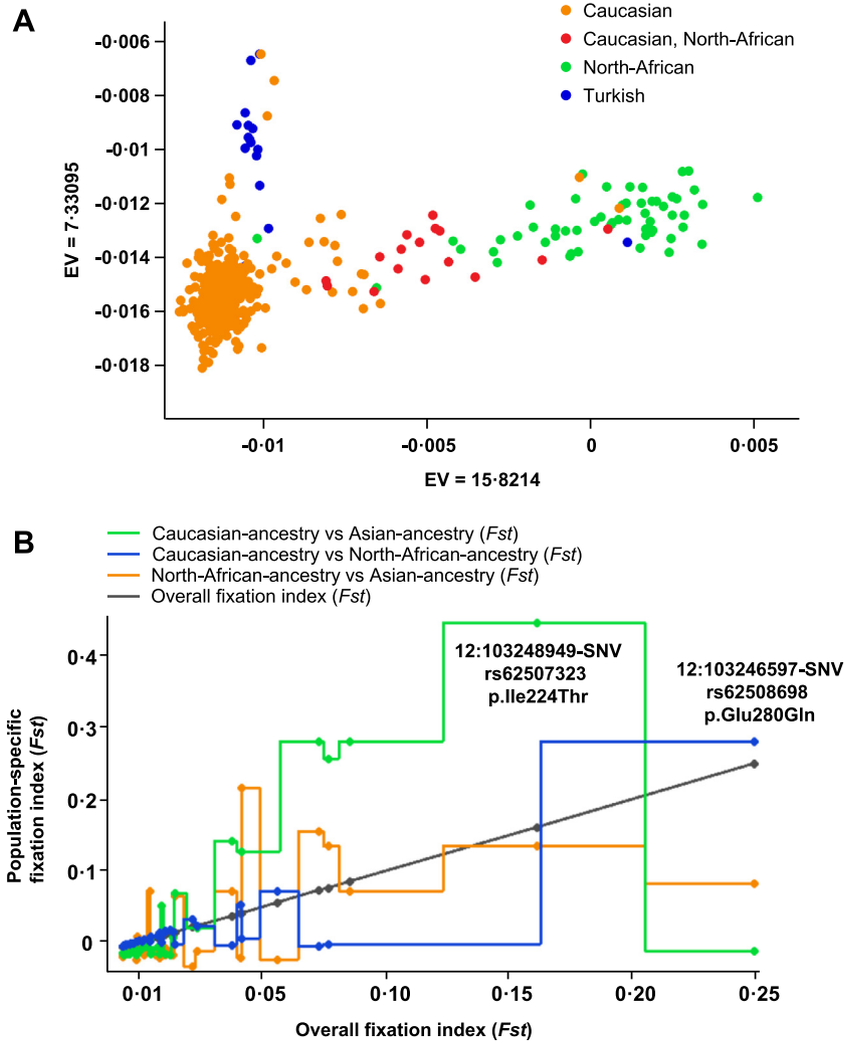


Fig. 3. (A) Principal component analysis according to patients' ethnicity. (B) The PAH gene variant rs62508698 was significantly enriched among North-African ancestry patients and significantly under-represented among patients with European-ancestry.

A, 12:103238949-T-C, and 12:103260273-T-G) are underrepresented among Africans (AAF \approx 0.05, gnomAD Genomes database) and over-enriched among East Asians (AAF \approx 0.75) (Fig. 7 and Supplemental Table S17). The two remaining variants are located outside the 'Bioplerin_H' domain coding region (12:103275872-T-C, 12:103275901-C-T) and are in high linkage disequilibrium ($D' = 1$) (Fig. 7). To better estimate the relative ordering of population differentiation events, we conducted a timetree analysis using the two top differentiated PAH variants between African- and East Asian-ancestry populations as outgroup sequences (12:103275872-T-C, 12:103275901-C-T). The timetree analysis confirmed the anteriority of divergence events on the phylogeny of the two top differentiated PAH variants with a branch length of 0.64 (relative time) (Supplemental Figure 5).

The 'Nancy Reference Centre Population' dataset included a total of 409 patients (European-ancestry, $n = 319$; North-African-ancestry, $n = 49$; and Asian-ancestry, $n = 41$). Two markers were associated with population differentiation between European- and North-African-ancestry populations and clustered under the same clade in phylogenetic analysis (Fig. 8A). The first variant corresponds to a splice acceptor variant (12:103234294-C-A, c.1200-1G>T), and the second is a synonymous variant (12:103234215-A-G, p.Asn426=). The splice acceptor variant (12:103234294-C-A) has an AAF of 1.59×10^{-5} (2/125,568) in the TOPMed (Trans-Omics for Precision Medicine) database [36] and is not reported in the gnomAD Genomes database [37]. The AAF of this variant was 7.18×10^{-4} (1/1392; enrichment ratio \approx 45) among all PKU patients in the French Nationwide PKU Study,

Table 2
Enrichment analysis for pathogenic PAH gene variants according to patients' ethnicity.

Marker	Fisher's P-value	Fisher's Bonf. P-value	OR (95% CI), Alternative allele	AAF, PKU Nationwide	AAF, Studied ethnicity	AAF, Other ethnicities	gnomAD Exomes AAF
Caucasian-ancestry*							
rs62508698	3.55×10^{-8}	4.19×10^{-6}	0.11 (0.05–0.26)	3.08×10^{-2}	1.04×10^{-2}	8.58×10^{-2}	5.57×10^{-5}
North-African-ancestry*							
rs62508698	7.18×10^{-15}	8.47×10^{-13}	23.23 (9.75–55.38)	3.08×10^{-2}	1.69×10^{-1}	8.70×10^{-3}	5.57×10^{-5}

AAF: alternative allele frequency; Bonf: Bonferroni; OR: odds ratio.

* No significant variant enrichment was found in patients' subgroups of other ethnicities.

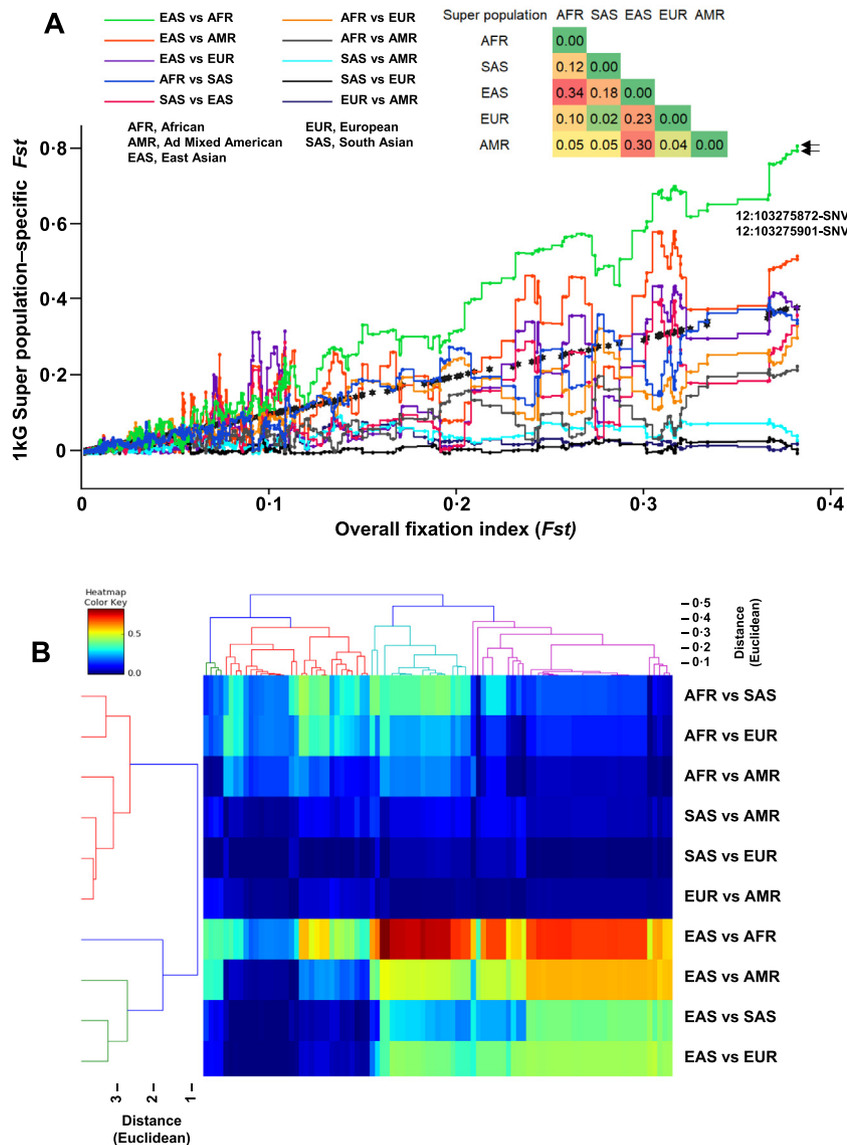


Fig. 4. (A) Two-dimension scatter plot reporting the pairwise population differentiation analysis using the F_{st} index against the overall F_{st} index (black stars) for *PAH* gene variants using whole-genome sequencing data from the 2504 subjects of the 1000 Genomes Project phase 3. The matrix at the top of the panel reports the pairwise population Fixation index. (B) Dendrogram with heat map reporting the pairwise population differentiation analysis for *PAH* gene variants using whole-genome sequencing data from the 1000 genomes project phase 3.

1.27×10^{-3} (1/395; enrichment ratio ≈ 159) among Caucasian-ancestry PKU patients in the French Nationwide PKU Study, 2.44×10^{-3} (2/818; enrichment ratio ≈ 154) in the ‘Nancy Reference Centre Population’, and 2.04×10^{-2} (2/98; enrichment ratio ≈ 1281) among North-African-ancestry subjects of the ‘Nancy Reference Centre Population’.

3.6. Exome-wide association study: unveiling other potential loci associated with population differentiation

For the exome-wide high-throughput genotyping, among the 696 patients included in the study, DNA samples were available for 574 (83%). We performed an exome-wide association study to assess potential genetic loci associated with the population differentiation and balancing selection pressure between Caucasian-ancestry PKU and North-African PKU patients. We performed linear haplotype trend regression using the rs62508698 as the response variable and a moving window with a dynamic width of 10 kb. We adjusted the

regression model for the three top principal components (EV = 15.8214, EV = 7.33095, EV = 2.12178) to account for potential confounding bias due to population stratification. Two loci reached statistical significance and were located in the *SSPO* (SCO-spondin, HGNC:21998) and *DBH* (dopamine beta-hydroxylase, HGNC:2689) genes (Fig. 9 and Supplemental Table S18). The genotype association study on the 37 variants located in the *SSPO* and *DBH* loci confirmed a significant association with four coding variants on the *SSPO* gene (rs73727633, p.Phe3209Leu; rs73727632, p.Cys3204Arg; rs56921891, p.Pro2393=; rs73727650, p.Arg5042Gly) and one coding variant on the *DBH* gene (rs4531, p.Ala318Thr) (Supplemental Table S19).

4. Discussion

In 2007, Charles Scriver stated that: “The PKU story contains many messages, including what the human *PAH* gene tells us about human population genetics and evolution of modern humans” [1]. Our study suggests a contributing role of population divergence and balanced

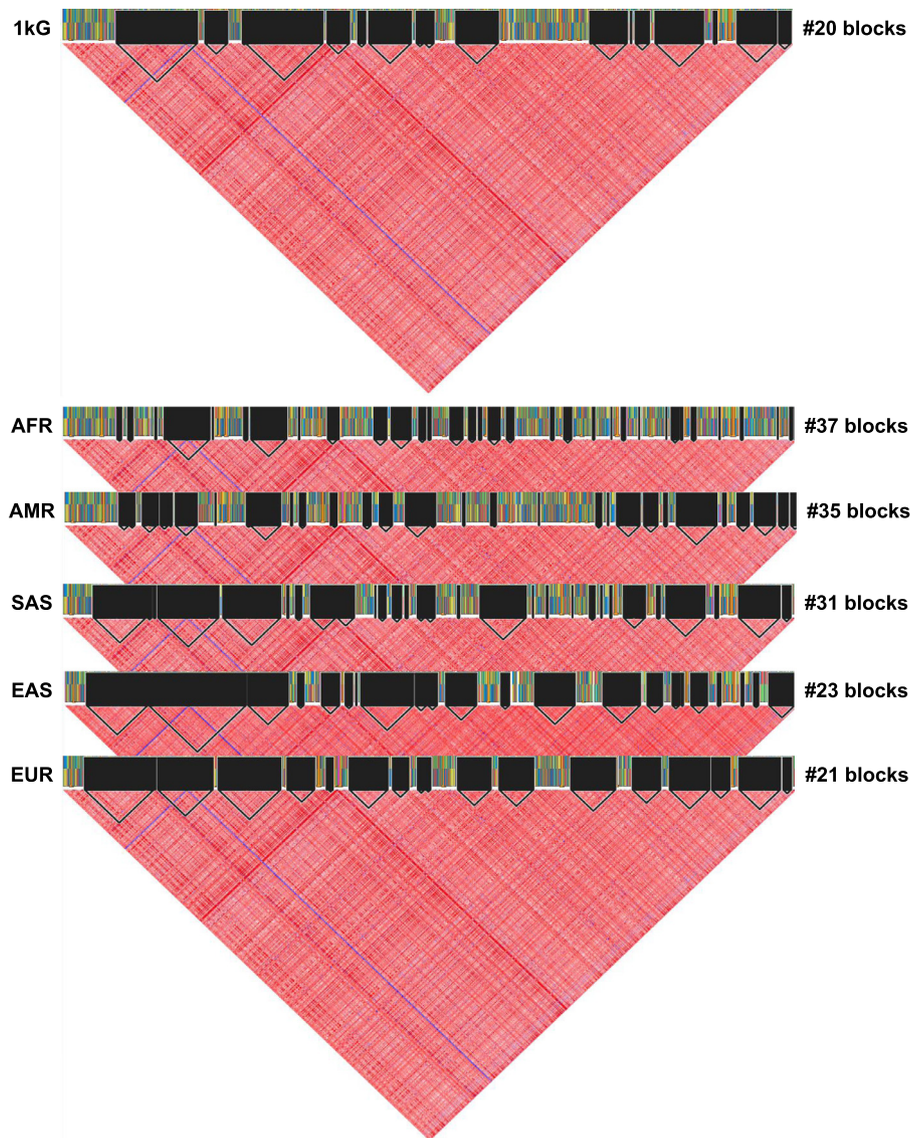


Fig. 5. Haplotype block analysis of the *PAH* gene using whole-genome sequencing data from the 2504 subjects of the 1000 Genomes Project phase 3. The number of haplotype blocks is reported for each super population.

selection to uncover heterozygote advantage and overdominance mechanisms on the *PAH* gene, in a multiethnic nationwide sample population of patients with PKU in France.

Our study confirmed the high mutation burden on the *PAH* gene by reporting more than 130 pathogenic variants. These mutations are not distributed randomly across the studied geographical areas in France. Interestingly, within the *PAH* locus, the variants that were subject to a high population divergence were significantly enriched in the C-terminal catalytic Biopterin_H domain by comparison to the N-terminal regulatory ACT domain, suggesting a potential role of balancing selection pressures in the shaping of the *PAH* locus. One variant (rs62508698) was associated with a substantial population divergence with a significant enrichment among North-Africans and an under-representation among Europeans. The exome-wide association study that addressed potential loci related to population divergence emphasised two top significant loci, namely *SSOP* and *DBH*. These data could lead to a better understanding of the adaptive mechanisms in response to the selection pressures exerted on the *PAH* gene and pave the way for new hypotheses regarding the

adaptive advantages conferred by *PAH* heterozygous mutations in the face of specific environmental factors, such as infectious diseases [38,39].

The reasons underlying the high prevalence of *PAH* heterozygous carriers are not clearly understood. One of the hypotheses is that the increased prevalence in heterozygotes is linked to an overdominant selection process [10,40–45]. Overdominance could be described as a heterozygous advantage, wherein heterozygous individuals have higher fitness than either homozygous individuals [46]. Heterozygous advantage represents a balanced selection mechanism by which multiple alleles at a specific locus are actively maintained in the gene pool of a population longer than expected from genetic drift alone [39,47]. Woolf et al. reported that women who were heterozygous for *PAH* mutations were at a lower risk of miscarriage when compared to wild-type, homozygous women [40]. It is well known that host-pathogen interactions have shaped genetic variation in modern populations [39]. The development of high-throughput approaches, as well as analytical methods and expanding public data resources, has led to new insights on how human infectious disease influenced

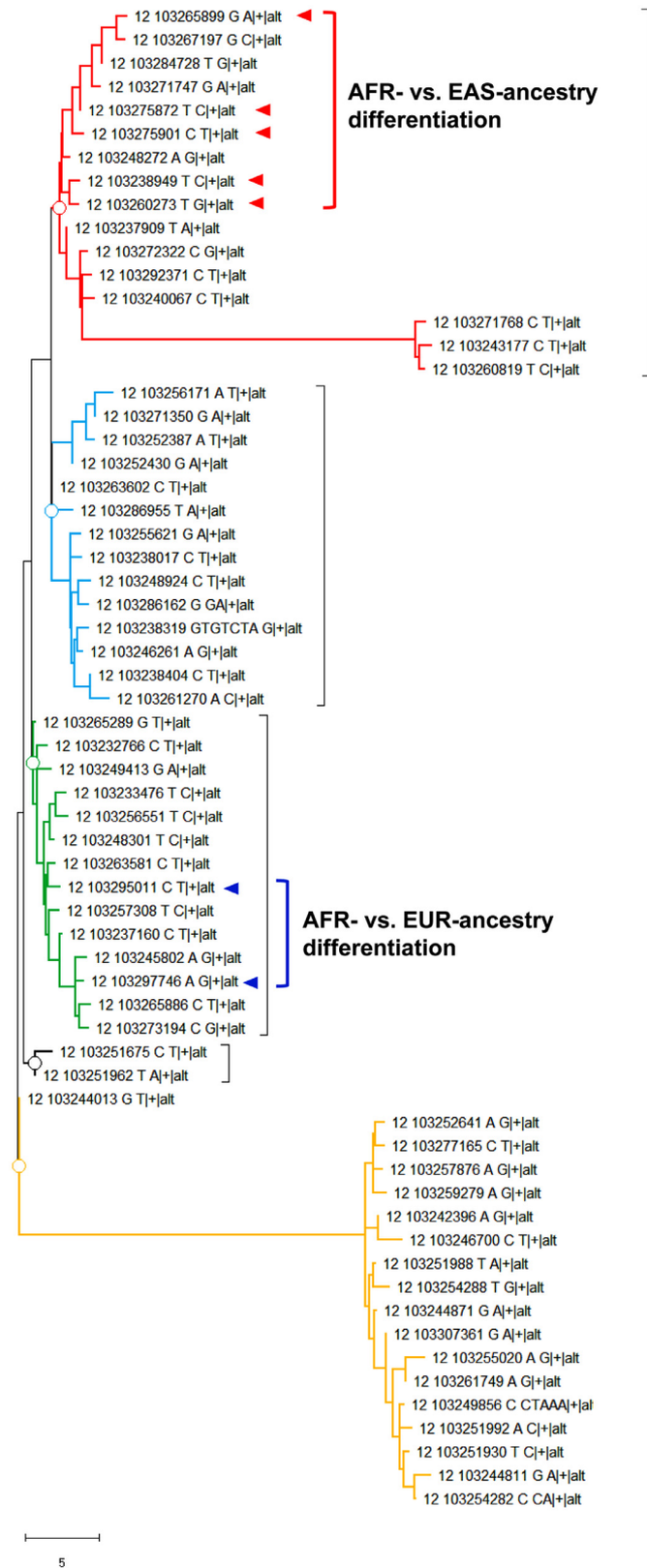


Fig. 6. Phylogenetic analysis of the *PAH* locus using 1 kG whole-genome sequencing data.

the natural selection and several adaptive mechanisms, including autoimmune and metabolic disorders [39]. Several examples of adaptive metabolic mechanisms providing benefit to defence against pathogens have been reported. An example regarding the

overdominant selection in PKU could be the consequence of higher phenylalanine levels in heterozygous than in wild-type homozygous individuals on host-pathogen interactions [48]. The potential mechanism was that phenylalanine acts as a competitive inhibitor of ochratoxin A, which is a teratogenic mycotoxin produced by *Aspergillus* spp. in mouldy grains and lentils [10,38]. Ochratoxin A is an N-acyl derivative of phenylalanine that can cross the placenta and induce spontaneous miscarriage through the inhibition of phenylalanyl-tRNA synthetase-catalysed reaction and protein synthesis [38]. Other examples of inherited disorders suggesting that overdominance provides resistance to a specific infectious disease include sickle trait (*HBB*) and resistance to malaria, cystic fibrosis (*CFTR*) and resistance to cholera, Tay-Sachs disease (*HEXA*) and resistance to tuberculosis, congenital disorder of glycosylation type IIb (*MOGS*) and resistance to viral infections, and Niemann-Pick disease type C1 (*NPC1*) and resistance to filoviruses (e.g., Ebola, Marburg) [38].

Phylogenetic and timetree analyses on the *PAH* locus were consistent with population differentiation events on European-, African-, and Asian-ancestry populations. Few studies have assessed the potential relationship between population divergence, overdominance, and molecular phylogenetics in the setting of inherited metabolic disorders. Using three different populations, we have highlighted a selection pressure on the *PAH* gene with a substantial population divergence in association with specific variants. The assessment of the allelic frequencies of the most differentiated variants between Africans and East-Asians also revealed an enrichment gradient between Africans and Europeans but to a lesser extent. Phylogenetic analyses showed that the variants most strongly associated with a high selection pressure were close phylogenetically and were located within the 'Biopterin_H' domain coding region or in its vicinity. The 'Biopterin_H' domain of *PAH* contains a binding domain for the *PAH* cofactor BH4, which regulates the enzyme activity as well as being essential in catalysis [49]. Several experiments of coupled transcription-translation in vitro assay have shown that BH4 has a chaperone-like effect on *PAH* synthesis and/or is a protecting cofactor against enzyme auto-inactivation and degradation [49]. BH4 is also an essential oxidant-sensitive cofactor for the nitric oxide synthase that synthesises nitric oxide (NO) from arginine and oxygen [50]. Patients with severe malaria have BH4 depletion, which in turn compromises both NO synthesis and phenylalanine metabolism [50,51]. The decreased BH4 availability in severe malaria uncouples NO synthase and leads to impaired NO bioavailability and potentially increased oxidative stress [51]. These data may suggest the potential role of malaria as an environmental factor that may drive selection pressure. However, malaria could not explain the genetic differentiation between Latinos and East Asian populations since the distribution of the infectious agent is similar in both geographical regions [52]. On the other hand, several studies have shown a clear genetic differentiation between East-Asian and Latino populations in terms of hypoxic adaptation to altitude [53–57]. Indeed, balancing selection on *NOS1* and *NOS2* has been shown to participate in the adaptive mechanisms associated with higher oxygen delivery in Andean and Tibetan populations [56,57]. The balancing selection pressure on the *PAH* locus, especially on the biopterin domain, may reflect a mechanism of adaptation to environmental factors but also to other enzymatic reactions that may involve the common BH4 cofactor in the *PAH* and *NOS* pathways. Thus, a combination of evolutionary and adaptive events in various populations may potentially explain the overdominance of some genetic variants on the *PAH* gene.

In humans, few phylogenetic studies have been reported in the setting of inherited metabolic disorders. One example is an in silico study that identified significant evolutionary events on the *NPC1* gene, which is associated with the Niemann-Pick disease type C

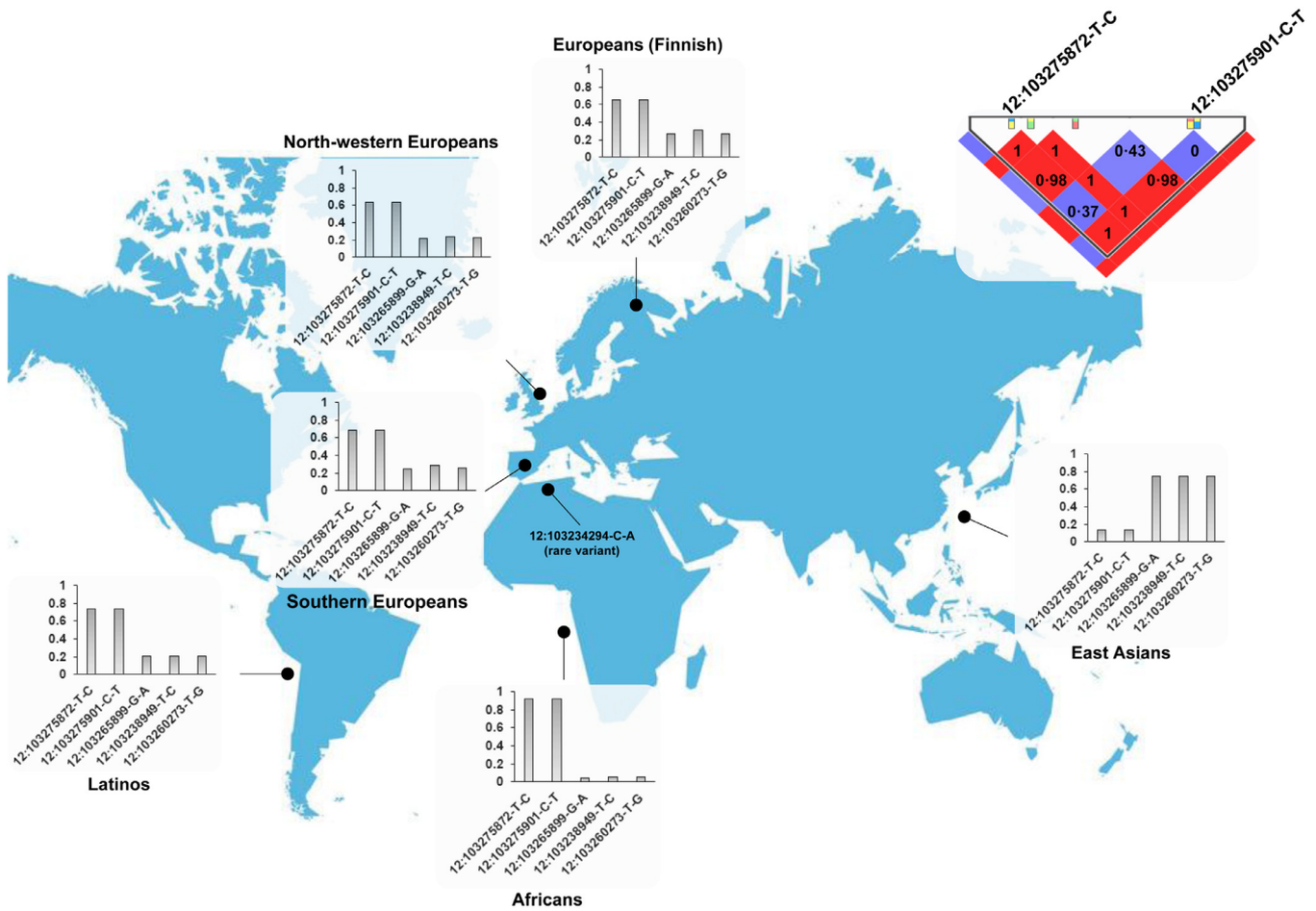


Fig. 7. World map representation of the alternative allele frequencies of the top differentiated (African- vs East Asian-ancestry populations) and phylogenetically clustered *PAH* variants.

disease [58]. This study did not evaluate the involvement of *NPC1* genetic variants in population differentiation or their potential role in overdominance. The precise model that links the evolutionary aspects of a gene from a phylogenetic point of view with population differentiation and overdominance remains to be clarified. Future studies on genes in which polymorphisms are known to be maintained by selection will help better understanding the role of balancing selection in the setting of population differentiation and phylogenetic evolution [59].

Our data also suggest that other loci, potentially associated with population differentiation and balancing selection, may contribute to the adaptive mechanisms of overdominance and heterozygote advantage. The exome-wide association study that addressed potential loci in association with population divergence on the *PAH* locus unravelled two top significant loci on *SSPO* and *DBH* genes. The *SSPO* gene codes for the subcommissural organ spondin (SCO-spondin) protein which is involved in the modulation of neuronal aggregation and the formation of the central nervous system. In vitro, SCO-spondin has been shown to stimulate neuronal differentiation and neurite growth [60]. The thrombospondin type-1 (TSP1) repeat (TSR) domain supports the main neurogenic properties of the SCO-spondin protein [60]. In vitro and in vivo studies have shown that the NX210 oligopeptide –which was designed from SCO-spondin's

specific TSR domain– was able to prevent neural cell death and to promote neurite outgrowth [60]. NX210 has been granted Orphan Drug designation by the European Medicine Agency for the treatment of spinal cord injury with an ongoing phase 1 study planned in patients with spinal cord injury. The association of a *PAH* gene variant serving as a marker of high balancing selection pressure with a gene locus involved in neuroprotection provides new insight into the potential adaptive mechanisms to *PAH* variants overdominance among populations. This hypothesis deserves further studies. The second locus associated with population divergence on *PAH* was on the *DBH* gene. The *DBH* gene codes for the dopamine beta-hydroxylase, which catalyses the conversion of dopamine to norepinephrine. The dopamine molecule is a down product of phenylalanine metabolism. It has been shown that the synthesis of dopamine is impaired in the human brain of patients with PKU [61]. The most relevant biochemical abnormalities among PKU patients are a decrease in the activity of tyrosine and tryptophan hydroxylases, which lead to deficiencies in dopamine and serotonin, which result in neurological abnormalities such as abnormal muscle tone, irritability and lethargy, seizures, and progressive developmental delay [62]. The treatment of PKU patients with a restricted phenylalanine diet induces a reduction of phenylalanine levels and an increase of dopamine and serotonin levels [61]. Consistently,

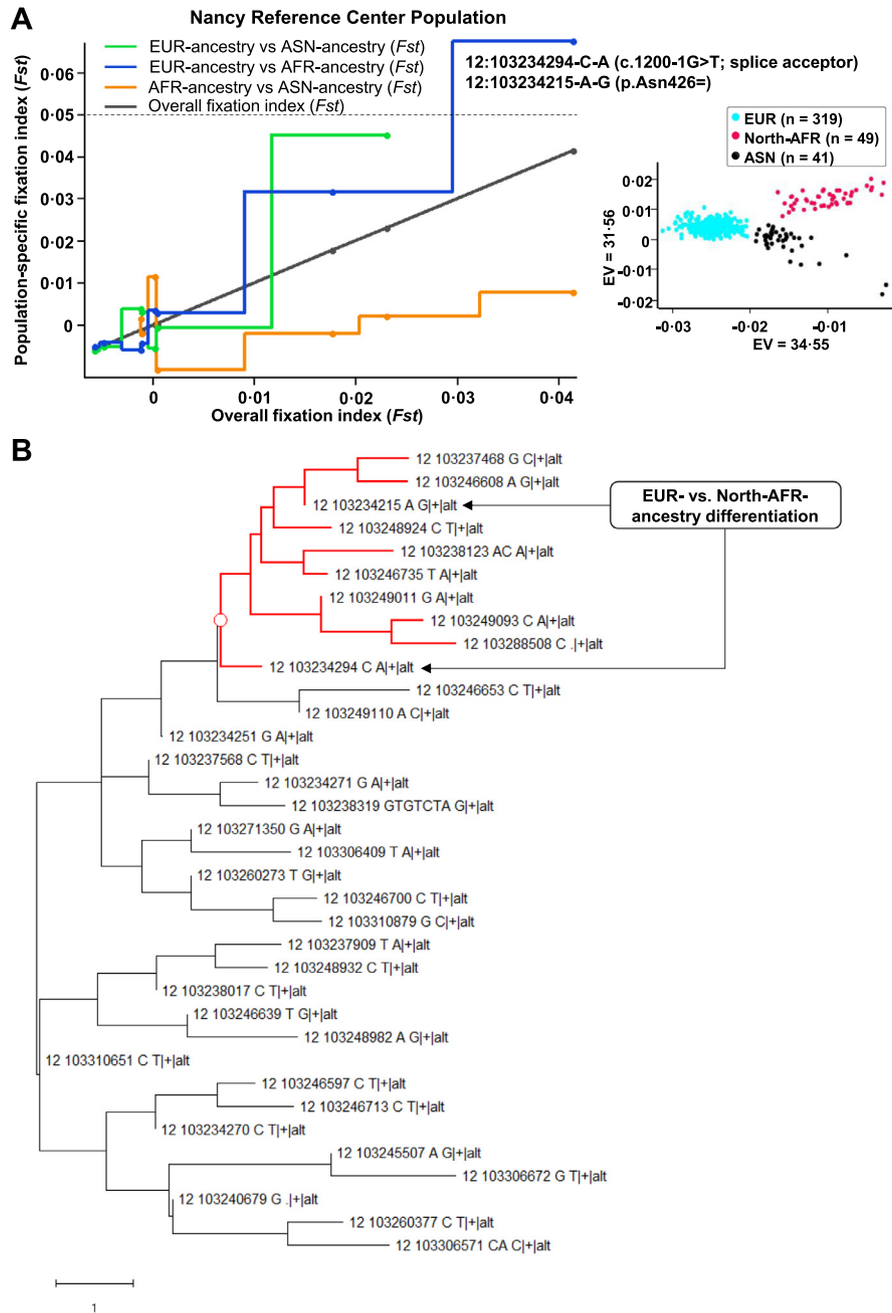


Fig. 8. (A) Two-dimension scatter plot reporting the pairwise population differentiation analysis using the F_{st} index against the overall F_{st} index for *PAH* gene variants evidenced in the 'Nancy Reference center Population'. (B) Phylogenetic and timetree analyses of the *PAH* locus using the 'Nancy Reference Centre Population' exome-sequencing data on non-PKU patients.

the mice model of PKU showed that phenylalanine diet supplemented with tyrosine induce a normalisation of dopamine levels in the brain [62,63]. The association between a selection pressure on the *PAH* locus and a genomic signature on *DBH* allows considering the hypothesis of genome plasticity with regards to environmental factors, but also adaptive metabolic factors to hyperphenylalaninemia in the setting of *PAH* overdominance. The influence of genetic variants of *DBH* on the phenotypic variability and natural history of PKU patients deserves to be studied in large cohorts.

5. Conclusion

In conclusion, our study suggests a potential role of population divergence and balanced selection in the overdominance mechanisms of the *PAH* gene. Phylogenetic and timetree analyses confirm population differentiation as a potential driver of the *PAH* gene evolution among European-, African-, and Asian-ancestry populations, notably on the biopterin domain at the crossroad between *PAH* and *NOS* pathways. Future studies are needed to decipher the adaptive mechanisms that underlie the influence of *PAH*, *SSPO*, and *DBH* genes,

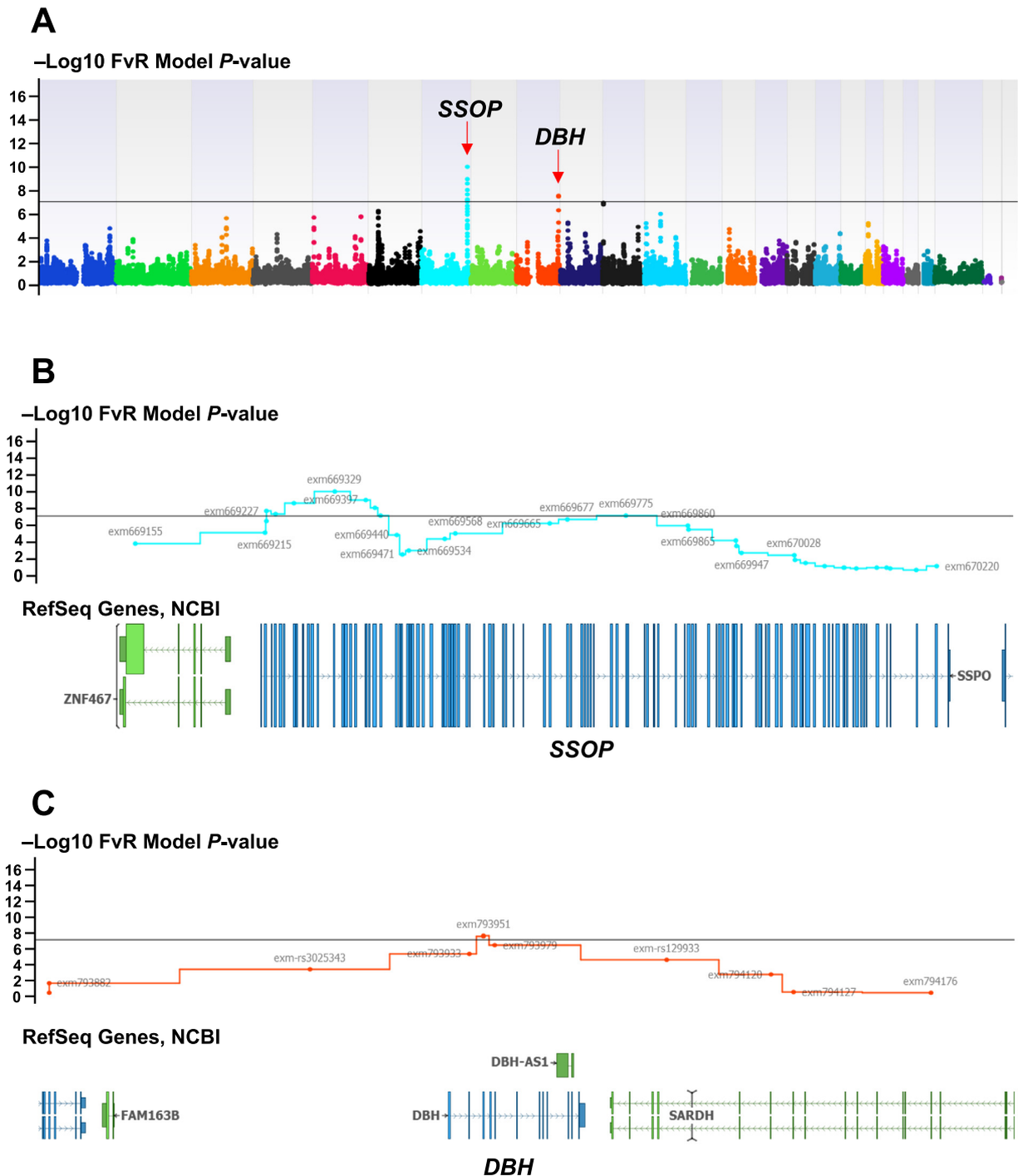


Fig. 9. (A) Manhattan plot reporting the exome-wide association study for the association with population differentiation and balancing selection pressure (*PAH* gene variant rs62508698) in haplotype trend regression analysis. The two arrows indicate the top significant loci: *SSPO* (SCO-spondin, HGNC:21998) and *DBH* (dopamine beta-hydroxylase, HGNC:2689) genes. Panels (B) and (C) report the genomic context of the *SSOP* and *SBH* genes, respectively. The *P*-values were reported after a symmetric smoothing transformation method using a window radius value of 2, as previously described [64].

including those related to overdominance and heterozygote advantage, in response to the selection pressures exerted on the *PAH* gene.

Declaration of Competing Interest

The authors who have taken part in this study declare that they do not have anything to disclose regarding conflicts of interest concerning this manuscript.

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Author contributions

Study concept and design: AO, JLG, and FN; study supervision and guarantor of the article: AO, JLG, and FN; data collection: AO, EJT, CC, PP, PR, TJ, AC, MB, AF, KM, FL, JBA, FM, CL, CD, KW, DT, PB, LDP, CG, AK, AB, GB, DL, SO, AM, RMRG, FF, JLG, FN; Sanger sequencing: CC, PP, RMRG, FN; exome-wide genotyping: CC, PR, AO, JLG; next-generation clinical exome sequencing: CC, TJ, AO, JLG, FN; statistical analysis: AO; data analysis and interpretation: AO, JLG, and FN; report drafting, review, and approval: AO, JLG, and FN; critical revision of the manuscript for relevant intellectual content: AO, JLG, and FN; approval of the submitted final draft: AO, EJT, CC, PP, PR, TJ, AC, MB, AF, KM, FL, JBA, FM, CL, CD, KW, DT, PB, LDP, CG, AK, AB, GB, DL, SO, AM, RMRG, FF, JLG, FN.

Supplementary materials

Supplementary material associated with this article can be found in the online version at doi:[10.1016/j.ebiom.2019.102623](https://doi.org/10.1016/j.ebiom.2019.102623).

References

- [1] Scriver CR. The PAH gene, phenylketonuria, and a paradigm shift. *Hum Mutat* 2007;28:831–45.
- [2] van Wegberg AMJ, MacDonald A, Ahring K, et al. The complete European guidelines on phenylketonuria: diagnosis and treatment. *Orphanet J Rare Dis* 2017;12:162.
- [3] Hellwege JN, Keaton JM, Giri A, Gao X, Velez Edwards DR, Edwards TL. Population stratification in genetic association studies. *Curr Protoc Hum Genet* 2017;95:1.22.1-1-3.
- [4] Nielsen R, Akey JM, Jakobsson M, Pritchard JK, Tishkoff S, Willerslev E. Tracing the peopling of the world through genomics. *Nature* 2017;541:302–10.
- [5] Hellenthal G, Busby GBJ, Band G, et al. A genetic atlas of human admixture history. *Science* 2014;343:747–51.
- [6] Eilbeck K, Quinlan A, Yandell M. Settling the score: variant prioritization and Mendelian disease. *Nat Rev Genet* 2017;18:599–612.
- [7] Karakachoff M, Duforet-Frebourg N, Simonet F, et al. Fine-scale human genetic structure in Western France. *Eur J Hum Genet* 2015;23:831–6.
- [8] Persyn E, Redon R, Bellanger L, Dina C. The impact of a fine-scale population stratification on rare variant association test results. *PLoS ONE* 2018;13:e0207677.
- [9] Scriver CR. Hyperphenylalaninemia: phenylalanine hydroxylase deficiency. *Metabol Mol Biol Inherit Dis* 2002;1667–724.
- [10] Krawczak M, Zschocke J. A role for overdominant selection in phenylketonuria? Evidence from molecular data. *Hum Mutat* 2003;21:394–7.
- [11] Rozen R, Mascisch A, Lambert M, Laframboise R, Scriver CR. Mutation profiles of phenylketonuria in Quebec populations: evidence of stratification and novel mutations. *Am J Hum Genet* 1994;55:321–6.
- [12] Guldberg P, Levy HL, Hanley WB, et al. Phenylalanine hydroxylase gene mutations in the United States: report from the maternal PKU collaborative study. *Am J Hum Genet* 1996;59:84–94.
- [13] Guldberg P, Rey F, Zschocke J, et al. A European multicenter study of phenylalanine hydroxylase deficiency: classification of 105 mutations and a general system for genotype-based prediction of metabolic phenotype. *Am J Hum Genet* 1998;63:71–9.
- [14] Leuders S, Wolfigart E, Ott T, et al. Influence of PAH genotype on sapropterin response in PKU: results of a single-center cohort study. *JIMD Rep* 2014;13:101–9.
- [15] Jeannesson-Thivisol E, Feillet F, Chery C, et al. Genotype-phenotype associations in French patients with phenylketonuria and importance of genotype for full assessment of tetrahydrobiopterin responsiveness. *Orphanet J Rare Dis* 2015;10:158.
- [16] Liu N, Huang Q, Li Q, et al. Spectrum of PAH gene variants among a population of Han Chinese patients with phenylketonuria from northern China. *BMC Med Genet* 2017;18:108.
- [17] Ohlsson A, Bruhn H, Nordenstrom A, Zetterstrom RH, Wedell A, von Döbeln U. The spectrum of PAH mutations and increase of milder forms of phenylketonuria in Sweden during 1965–2014. *JIMD Rep* 2017;34:19–26.
- [18] Hamilton V, Santa Maria L, Fuenzalida K, et al. Characterization of phenylalanine hydroxylase gene mutations in Chilean PKU patients. *JIMD Rep* 2018;42:71–7.
- [19] Shirzadeh T, Saeidian AH, Bagherian H, et al. Molecular genetics of a cohort of 635 cases of phenylketonuria in a consanguineous population. *J Inher Metab Dis* 2018;41:1159–67.
- [20] Li N, He C, Li J, et al. Analysis of the genotype-phenotype correlation in patients with phenylketonuria in mainland China. *Sci Rep* 2018;8:11251.
- [21] Huyghe JR, Jackson AU, Fogarty MP, et al. Exome array analysis identifies new loci and low-frequency variants influencing insulin processing and secretion. *Nat Genet* 2013;45:197–201.
- [22] Price AL, Weale ME, Patterson N, et al. Long-range LD can confound genome scans in admixed populations. *Am J Hum Genet* 2008;83:132–5.
- [23] Weale ME. Quality control for genome-wide association studies. *Methods Mol Biol* 2010;628:341–72.
- [24] Genomes Project C, Auton A, Brooks LD, et al. A global reference for human genetic variation. *Nature* 2015;526:68–74.
- [25] Wright S. The genetical structure of populations. *Ann Eugen* 1951;15:323–54.
- [26] Hartl DL, Clark AG, Clark AG. Principles of population genetics: Sinauer associates Sunderland, MA; 1997.
- [27] Weir BS, Cockerham CC. Estimating F-Statistics for the analysis of population structure. *Evolution (N Y)* 1984;38:1358–70.
- [28] Leiyang S, Hamilton MB. Properties of Weir and Cockerham's Fst estimators and associated bootstrap confidence intervals. *Theor Popul Biol* 2011;79:39–52.
- [29] Remington DL, Thornsberry JM, Matsuoka Y, et al. Structure of linkage disequilibrium and phenotypic associations in the maize genome. *Proc Natl Acad Sci U S A* 2001;98:11479–84.
- [30] Lewontin RC. The interaction of selection and linkage. I. General considerations; heterotic models. *Genetics* 1964;49:49–67.
- [31] Gabriel SB, Schaffner SF, Nguyen H, et al. The structure of haplotype blocks in the human genome. *Science* 2002;296:2225–9.
- [32] Zhang P, Boisson B, Stenson PD, et al. SeqTailor: a user-friendly webserver for the extraction of DNA or protein sequences from next-generation sequencing data. *Nucleic Acids Res* 2019;47:W623–W31.
- [33] Tamura K, Nei M. Estimation of the number of nucleotide substitutions in the control region of mitochondrial DNA in humans and chimpanzees. *Mol Biol Evol* 1993;10:512–26.
- [34] Tamura K, Battistuzzi FU, Billing-Ross P, Murillo O, Filipowski A, Kumar S. Estimating divergence times in large molecular phylogenies. *Proc Natl Acad Sci U S A* 2012;109:19333–8.
- [35] Kumar S, Stecher G, Li M, Knyaz C, Tamura K. MEGA X: molecular evolutionary genetics analysis across computing platforms. *Mol Biol Evol* 2018;35:1547–9.
- [36] Taliun D, Harris DN, Kessler MD, et al. Sequencing of 53,831 diverse genomes from the NHLBI TOPMed program. *BioRxiv* 2019:563866.
- [37] Karczewski KJ, Francioli LC, Tiao G, et al. Variation across 141,456 human exomes and genomes reveals the spectrum of loss-of-function intolerance across human protein-coding genes. *BioRxiv* 2019:531210.
- [38] Withrook IC, Anderson SJ, Jefferson MA, et al. Genetic diseases conferring resistance to infectious diseases. *Genes Dis* 2015;2:247–54.
- [39] Karlsson EK, Kwiatkowski DP, Sabeti PC. Natural selection and infectious disease in human populations. *Nat Rev Genet* 2014;15:379–93.
- [40] Woolf LI, McBean MS, Woolf FM, Cahalane SF. Phenylketonuria as a balanced polymorphism: the nature of the heterozygote advantage. *Ann Hum Genet* 1975;38:461–9.
- [41] Saugstad LF. Anthropological significance of phenylketonuria and the importance of heterozygote advantage. *Ir Med J* 1976;69:405–10.
- [42] Saugstad LF. Heterozygote advantage for the phenylketonuria allele. *J Med Genet* 1977;14:20–4.
- [43] Smith I, Carter CO, Wolff OH. Heterozygote advantage for the phenylketonuria allele. *J Med Genet* 1978;15:246–8.
- [44] Saugstad LF. Heterozygote advantage for the phenylketonuria allele. *J Med Genet* 1978;15:317–9.
- [45] Woolf LI. The heterozygote advantage in phenylketonuria. *Am J Hum Genet* 1986;38:773–5.
- [46] DeGiorgio M, Lohmueller KE, Nielsen R. A model-based approach for identifying signatures of ancient balancing selection in genetic data. *PLoS Genet* 2014;10:e1004561.
- [47] Leffler EM, Bullaughey K, Matute DR, et al. Revisiting an old riddle: what determines genetic diversity levels within species? *PLoS Biol* 2012;10:e1001388.
- [48] Woolf LI, Cranston WI, Goodwin BL. Genetics of phenylketonuria. Heterozygosity for phenylketonuria. *Nature* 1967;213:882–3.
- [49] Thony B, Ding Z, Martinez A. Tetrahydrobiopterin protects phenylalanine hydroxylase activity in vivo: implications for tetrahydrobiopterin-responsive hyperphenylalaninemia. *FEBS Lett* 2004;577:507–11.
- [50] Alkaitis MS, Ackerman HC. Tetrahydrobiopterin supplementation improves phenylalanine metabolism in a murine model of severe malaria. *ACS Infect Dis* 2016;2:827–38.
- [51] Yeo TW, Lampah DA, Kenangalem E, et al. Impaired systemic tetrahydrobiopterin bioavailability and increased dihydrobiopterin in adult falciparum malaria: association with disease severity, impaired microvascular function and increased endothelial activation. *PLoS Pathog* 2015;11:e1004667.
- [52] Dalrymple U, Mappin B, Gething PW. Malaria mapping: understanding the global endemicity of falciparum and vivax malaria. *BMC Med* 2015;13:140.

- [53] Beall CM. Two routes to functional adaptation: Tibetan and Andean high-altitude natives. *Proc Natl Acad Sci U S A* 2007;104(Suppl 1):8655–60.
- [54] Beall CM, Laskowski D, Erzurum SC. Nitric oxide in adaptation to altitude. *Free Radic Biol Med* 2012;52:1123–34.
- [55] Jeong C, Ozga AT, Witonsky DB, et al. Long-term genetic stability and a high-altitude East Asian origin for the peoples of the high valleys of the Himalayan arc. *Proc Natl Acad Sci U S A* 2016;113:7485–90.
- [56] Crawford JE, Amaru R, Song J, et al. Natural selection on genes related to cardiovascular health in high-altitude adapted andeans. *Am J Hum Genet* 2017;101:752–67.
- [57] Bhandari S, Cavalleri GL. Population history and altitude-related adaptation in the Sherpa. *Front Physiol* 2019;10:1116.
- [58] Adebali O, Reznik AO, Ory DS, Zhulin IB. Establishing the precise evolutionary history of a gene improves prediction of disease-causing missense mutations. *Genet Med* 2016;18:1029–36.
- [59] Charlesworth D. Balancing selection and its effects on sequences in nearby genome regions. *PLoS Genet* 2006;2:e64.
- [60] Sakka L, Deletage N, Lalloue F, et al. SCO-spondin derived peptide NX210 induces neuroprotection in vitro and promotes fiber regrowth and functional recovery after spinal cord injury. *PLoS ONE* 2014;9:e93179.
- [61] Guttler F, Lou H. Dietary problems of phenylketonuria: effect on CNS transmitters and their possible role in behaviour and neuropsychological function. *J Inherit Metab Dis* 1986;9(Suppl 2):169–77.
- [62] Martynyuk AE, van Spronsen FJ, Van der Zee EA. Animal models of brain dysfunction in phenylketonuria. *Mol Genet Metab* 2010;99(Suppl 1):S100–5.
- [63] Joseph B, Dyer CA. Relationship between myelin production and dopamine synthesis in the PKU mouse brain. *J Neurochem* 2003;86:615–26.
- [64] Gueant JL, Chery C, Oussalah A, et al. APRDX1 mutant allele causes a MMACHC secondary epimutation in cblC patients. *Nat Commun* 2018;9:67.