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The potentially deleterious functional variant flavin-containing monooxygenase 2*1 is at high frequency throughout sub-Saharan Africa

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Abstract

Background—The drug-metabolizing enzyme flavin-containing monooxygenase 2 (*FMO2*) is the predominant *FMO* isoform present in the lung of most mammals, including non-human primates. All Europeans and Asians tested have been shown to be homozygous for a nonfunctional variant, *FMO2**2A, which contains a premature stop codon due to a single-nucleotide change in exon 9 (g.23238C > T). The ancestral allele, *FMO2**1, encodes a functionally active protein and has been found in African-Americans (26%) and Hispanics (2% to 7%). Possessing this variant increases the risk of pulmonary toxicity when exposed to thioureas, a widely used class of industrial compounds. *FMO2* may also be involved in the metabolism of drugs that are used to treat diseases that are prevalent in Africa.

Results and Conclusion—We conducted a survey of g.23238C > T variation across Africa that revealed that the distribution of this SNP is relatively homogeneous across sub-Saharan Africa, with approximately one third of individuals possessing at least one *FMO2**1 allele, though in some populations the incidence of these individuals approached 50%. Thus many sub-Saharan Africans may be at substantially increased health risk when encountering thiourea-containing substrates of *FMO2*. Analysis of HapMap data with the Long-Range Haplotype test found no evidence for positive selection of either 23238C > T allele and maximum-likelihood coalescent analysis indicated that this

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mutation occurred some 500,000 years before present. This study demonstrates the value of performing genetic surveys in Africa, a continent in which human genetic diversity is thought to be greatest, but where studies of the distribution of this diversity are few.

Keywords

drug metabolism; flavin-containing monooxygenase; flavin-containing monooxygenase 2; long-range haplotype test; pharmacogenetics; sub-Saharan Africa

Introduction

Flavin-containing monooxygenases

Flavin-containing monooxygenases (FMOs, EC1.14.13.8) catalyze the nicotinamide adenine dinucleotide phosphate-dependent oxidative metabolism of a variety of foreign chemicals that contain, as their site of oxidation, a soft nucleophilic heteroatom, such as nitrogen, phosphorous, sulphur or selenium [1–4]. Substrates include therapeutic drugs, dietary-derived compounds and environmental pollutants.

Humans possess five functional *FMO* genes, designated *FMO1–FMO5* [5–7]. All but the *FMO5* gene is present within a 220-kb cluster on chromosome 1q24.3 [7]. *FMO5* is located about 26 Mb closer to the centromere at 1q21.1 [7]. A sixth gene, *FMO6*, present within the cluster, does not produce a correctly spliced mRNA and thus seems to be a pseudogene [8]. A second *FMO* gene cluster, containing five pseudogenes, *FMO7P–FMO11P*, is located about 4 Mb centromeric of the *FMO* gene cluster [7].

Earlier studies on flavin-containing monooxygenase 2

In most mammals, including nonhuman primates, *FMO2* is the major isoform expressed in the lung [6,9–12]. A single nucleotide polymorphism (SNP) (g.23238C > T, dbSNP #rs6661174) in exon 9 that converts a glutamine codon at position 472 to a stop codon (Q472X), resulting in the production of a truncated polypeptide that is functionally inactive [10], has been identified in humans. In populations of European ($n = 79$) and Asian ($n = 118$) origin, all individuals tested have been found to be homozygous for this allele (*FMO2*2A*) [10,13]. An allele, *FMO2*1*, that has, however, been shown earlier to encode a full-length, functionally active protein [10,14] has been found in African-Americans (26%, $n = 180$) [10,13,15] and Hispanics (2–7%, $n = 280$ and 327) [16].

Substrates of human *FMO2* include thioether-containing organophosphate pesticides, such as phorate and disulfoton [17]. In this case, products of the *FMO2*-catalyzed reaction are substantially less toxic than the parent compounds [18] and as a consequence the enzyme has a protective role. *FMO2* has also, however, been shown to catalyze *S*-oxygenation of thiourea and some of its derivatives, such as phenylthiourea, α -naphthylthiourea and ethylenethiourea [19], producing sulfenic and/or sulfinic acid metabolites, which are more toxic than the parent compound [18]. Sulfenic acid derivatives of thioureas can deplete glutathione, leading to oxidative stress [20]; they can also bind to sulphhydryl groups on proteins and thus may directly perturb cell function [21]. Thus, if exposed to thiourea or its derivatives, individuals who possess an *FMO2*1* allele are predicted to be at increased risk of pulmonary toxicity. With an estimated global production of 10 000 ton [22], thioureas are present in a wide range of industrial, household and medical products and, consequently, exposure to these chemicals is widespread.

FMOs are also involved in the metabolism of therapeutic drugs, including several that are used to treat multidrug-resistant tuberculosis [23–25], which is a major health problem in Africa,

with an estimated 544 000 deaths in 2005 (<http://www.who.int/mediacentre/factsheets/fs104/en/>). Evidence that at least one of these drugs, ethionamide, is a substrate for human *FMO2* is observed [4], but it is not known whether metabolism of the drug by *FMO2* will increase or decrease its efficacy or toxicity.

The rationale for studying flavin-containing monooxygenase 2 in Africans

It has been shown earlier that most African-Americans have a significant European contribution to their ancestry (about 4–30% [26–28]) so it is likely that functional *FMO2* will be found at an even higher incidence in sub-Saharan Africans than in African-Americans. As this may be important in regard to drug efficacy and public safety we assessed the distribution of the *FMO2**1 and *FMO2**2A alleles in multiple populations across Africa. Samples from the Middle East (Turkey and Yemen) were also characterized to determine whether the *FMO2**1 allele was present at appreciable frequencies in populations outside but close to Africa.

In addition, we used the Long-Range Haplotype test [29], which examines the level of allele-specific haplotype linkage disequilibrium, to analyze data from the International HapMap project for evidence of positive selection at the g.23238C > T SNP and used sequence data from the National Institute of Environmental Health Sciences (NIEHS) SNP program to estimate the time of origin of the *FMO2**2A allele. This will help to provide preliminary insights into the evolutionary history and future of the *FMO2* enzyme.

Materials and methods

Sample collection

DNA samples were prepared from buccal swabs from a sample of 18-year-old males unrelated at the paternal grandfather level from the following locations in and around Africa: Algeria–Mostaganem ($n = 43$), Algeria–Port Say ($n = 118$), Cameroon–Mayo Darle ($n = 119$), Cameroon–Lake Chad ($n = 76$), Ethiopia–Gambella ($n = 106$), Ethiopia–Addis Ababa ($n = 24$), Ethiopia–Borena (and surrounding area) Wollo ($n = 36$), Ethiopia–Dessie (and surrounding area) Wollo ($n = 26$), Ghana–Sandema ($n = 90$), Ghana–Navrongo ($n = 45$), Malawi–Lilongwe ($n = 144$), Malawi–Mangochi ($n = 60$), Malawi–Mzuzu ($n = 56$), Morocco–Ifrane ($n = 70$), Mozambique–Sena ($n = 84$), Nigeria–Calabar ($n = 88$), Senegal southern region ($n = 94$), Senegal–Dakar ($n = 95$), South Africa–Pretoria ($n = 41$), Sudan northern region ($n = 136$), Sudan southern region ($n = 126$), Tanzania–Kilimanjaro ($n = 50$), Turkey–East Anatolia ($n = 31$), Turkey–West Anatolia ($n = 28$), Uganda–Ssesse Islands ($n = 39$), Yemen–Sena ($n = 34$), Yemen–Hadramaut region ($n = 83$), Zimbabwe–Mposi ($n = 34$). All samples were collected anonymously with informed consent. Sociological data, including age, current residence, birthplace, self-declared cultural identity and religion of the individual and of the individual's father, mother, paternal grandfather and maternal grandmother were also collected. In addition, the African populations sampled were grouped into four geographic regions (north Africa-NA, west Africa-WA, central east Africa-CEA, south-east Africa-SEA), as delineated in Table 1. The two Anatolian–Turkish samples were considered to be from a single region (TU), as were the two Yemeni samples (YE).

g.23238C > T genotyping

A 68-base pair (bp) region containing the g.23238C > T SNP was amplified by PCR using the primers *FMO2*-1414-UM (5'-TGG CTG TGA GAC TCT ATT TCG GAC CCT GCA ACT CCG A-3') and *FMO2*-1414-LM (5'-CCA TTG CCC AGG CCC AAC CAG GCG ATA TT-3'). Each primer contained a single mismatch to its target sequence at the 3'-end penultimate nucleotide (underlined). The design of the primers was such that the amplification product would contain recognition sites for the restriction endonucleases (REs) *Mbo*I (GATC), if the

target sequence contained a C at position 23238, and *MseI* (TTAA), if the target sequence contained a T at position 23238.

DNA was amplified in 10- μ l reaction volumes containing 0.4 μ mol/l of each primer, 0.13 units *Taq* DNA polymerase (HT Biotech, Cambridge, UK), 9.3 nmol/l TaqStart monoclonal antibody (BD Biosciences Clontech, Oxford, UK), 200 μ mol/l dNTPs and reaction buffer supplied with the *Taq* polymerase. The cycling parameters were: 5 min of preincubation at 93°C, after 37 cycles of 93°C for 1 min, 55°C for 1 min and 72°C for 1 min.

The resultant PCR product was used for two independent, complementary RE digestions that each targeted one of the two introduced RE sites. RE digestions were carried out in 10 μ l volumes containing 4 μ l of PCR product, 0.7 units RE (*MboI* or *MseI*), BSA and reaction buffer according to the supplier's recommendations (New England Biolabs, Hitchin, UK). All reactions were incubated overnight at 37°C. After RE digestion DNA fragments were resolved by electrophoresis through a 3.5% agarose gel. When full-length PCR product is digested with *MboI*, *FMO2*1* alleles are cleaved, resulting in two fragments of length 35 and 33 bp, respectively. When full-length PCR product is digested with *MseI*, *FMO2*2A* alleles are cleaved, resulting in two fragments of length 38 and 30 bp, respectively.

Data analysis

Tests for departure of observed genotype frequencies from those expected under Hardy–Weinberg equilibrium [30] were performed using Arlequin software (Laurent Excoffier, Guillaume Laval and Stephan Schneider, Zoological Institute, University of Bern, Bern, Switzerland) [31]. Pairwise F_{ST} values were estimated from analysis of molecular variance F_{ST} values [32].

Logistic regression analysis was carried out to evaluate the differences in the *FMO2*1* allele frequency among subgroups within regions and among regions in which the subgroups had similar allele frequencies. This was undertaken by first testing for fit of the subgroup frequencies to a model, which allowed only for regional differences in the *FMO2* allele frequencies. Pearson χ^2 tests were subsequently carried out to test for overall heterogeneity within individual regions. If significant heterogeneity was found in a region, further pairwise comparisons of the subgroups within the region were made by Fisher's exact tests. For logistic regression analysis and post-hoc region and subgroup comparisons, individuals were categorized into two groups on the basis of whether or not they possessed at least one *FMO2*1* allele. (In this way the sample size equaled the number of individuals studied, n , rather than the number of chromosomes, $2n$, thus ensuring that the observations were truly independent).

Principal coordinates analysis was performed, using GENSTAT5 software (VSN International Ltd, Hemel Hempstead, United Kingdom), on pairwise similarity matrices. Here similarity was quantified as being equal to the value of the genetic distance subtracted from 1.0 ($1-F_{ST}$). Values along the main diagonal, representing the similarity of each population sample to itself, were calculated from the estimated genetic distance between two copies of the same sample. For analysis of molecular variance-based F_{ST} distances, the resulting similarity of a sample to itself simplifies to $n/(n-1)$.

A Mantel test for the correlation between a matrix of pairwise F_{ST} values and a corresponding matrix of pairwise geographic distances was performed within the R-programming environment, using routines found in the APE package (Emmanuel Paradis, Université Montpellier II, Montpellier, France).

The Long-Range Haplotype test

The Long-Range Haplotype test was carried out using the Phase II International HapMap Project data release (<http://www.hapmap.org>) from four different populations. HapMap Phase II encompasses the following: the rel#21 YRI build, consisting of 3 241 616 SNPs genotyped in 30 parent–offspring trios from the Yoruba in Ibadan, Nigeria, the rel#21 CEU build, consisting of 1 105 072 SNPs genotyped in 30 parent–offspring trios from the Centre d'Etude du Polymorphisme Humain (Centre d'Etude du Polymorphisme Humain-Utah residents with ancestry from northern and western Europe) panel and the rel#21 CHB and JPT build, consisting of 3 305 784 SNPs genotyped in 45 unrelated Han Chinese from Beijing, China and 45 unrelated Japanese from Tokyo, Japan. On account of their high genetic similarity, it is accepted practice to pool the CHB and JPT datasets. In each dataset approximately one SNP is genotyped every 2 kb across the human genome.

The iHS method of Voight *et al.* [33] was applied to the g.23238C > T SNP in the HapMap Phase II YRI dataset and to the *FMO2* gene in the HapMap Phase II YRI, CEU and pooled JPT and CHB datasets, using the web-based tool Haplotter (<http://pritch.bsd.uchicago.edu/data.html>).

Estimating the age of the g.23238C > T mutation

Individuals of various ethnicities, including a subset of HapMap samples, have been sequenced for all exons of selected environmental response genes, including *FMO2*, as part of the NIEHS SNP program (NIEHS SNPs. NIEHS Environmental Genome Project, University of Washington, Seattle, Washington [<http://egp.gs.washington.edu>] [(June, 2007)]). We utilized *FMO2* sequencing data from the NIEHS SNP program to estimate the time when the g.23238C > T mutation occurred (see Supplementary Section 1 for a full explanation).

Results

The distribution of 23238C > T in Africa

The g.23238C > T allele frequencies and geographic locations for populations typed in this study are shown in Table 1 and Fig. 1. The overall *FMO2*1* allele frequency for all samples from Africa ($n = 1800$) was 0.153, with 28.3% of individuals having at least one *FMO2*1* allele. Across all 24 populations in Africa the observed percentage of individuals who have at least one *FMO2*1* allele ranged from 4.3 to 49.1. For samples from sub-Saharan Africa ($n = 1569$) the overall *FMO2*1* allele frequency was 0.170, with 31.4% of individuals having at least one *FMO2*1* allele and across these 21 populations the observed range of frequencies of *FMO2*1*-carrying individuals was 17.8–49.1%. The YE sample ($n = 117$) had an overall *FMO2*1* allele frequency of 0.047, with 8.5% of individuals having at least one *FMO2*1* allele. The *FMO2*1* allele was not observed in the Anatolian–Turkish sample ($n = 59$). No population deviated significantly from Hardy–Weinberg equilibrium ($P > 0.12$).

Using logistic regression on the proportion of individuals with at least one *FMO2*1* allele, significant differences were found both among regions [$P < 0.0001$, degrees of freedom (df) = 5] and among populations within regions ($P < 0.04$, df = 23). The major factor contributing to among-region differences is likely to be the noticeably lower *FMO2*1* frequencies observed in nonsub-Saharan African populations in comparison with sub-Saharan African populations. Pearson's χ^2 tests were carried out to explore within-region differences (see Table 2). The only statistically heterogeneous region was CEA ($P < 0.003$, df = 5). Exclusion of CEA from the logistic regression analysis resulted in no significant differences ($P = 0.25$, df 15) among populations within the remaining regions.

To make pairwise comparisons of regions using a Fisher's exact test, populations within each region were pooled, except in the case of CEA, which we had identified earlier as having statistically significant heterogeneity and therefore was excluded from this analysis. From these pairwise comparisons (Table 3) the following arrangement of regions based on frequencies of individuals with at least one *FMO2*1* allele could be discerned:

$$TU < (YE=NA) < (SEA=WA)$$

Further examination of populations in CEA, with pairwise Fisher's exact tests (Table 4), showed the populations in this region to be roughly split into two main groups, one consisting of north Sudan and the four Amharic populations and the other consisting of the Anuak and south Sudan. A principal coordinates analysis plot of pairwise F_{ST} values for all populations in Africa (Fig. 2) showed the Anuak of Gambella and south Sudan to be genetically close to each other with respect to the g.23238C > T SNP, probably because both of them possessing slightly elevated *FMO2*1* frequencies in comparison with the other African populations surveyed here. Addis Ababa seems to be somewhat separated from all populations, but this may be a stochastic effect because of its low sample size. A Pearson's χ^2 test comparing the frequencies of individuals with at least one *FMO2*1* allele in all populations in sub-Saharan Africa was significant ($P < 0.003$), but removing only the Anuak and south Sudan populations resulted in nonsignificance ($P = 0.526$), emphasizing that these two populations are outliers from the overall allele distribution observed across sub-Saharan Africa.

A significant correlation between matrices of pairwise genetic distances (F_{ST}) and geographic distances (kilometre) was found using the Mantel test when all populations typed in this study ($P < 0.001$) and only African populations ($P < 0.003$) were considered, but not when only sub-Saharan African populations were analyzed ($P = 0.741$), confirming the generally similar distribution of g.23238C > T alleles across sub-Saharan Africa.

When samples were grouped by self-declared ethnic identity [they were included as a separate group if there were 15 samples or more with the same self-declared ethnic identity (see Supplementary Table 1)], no significant differences were found between the same ethnic group living in multiple locations (Fisher's exact, $P > 0.24$) (see Supplementary Table 2), for example, the Amharic speakers who were sampled in three locations (Pearson's χ^2 , $P = 0.47$ df = 2), or among different ethnic groups collected at the same location (Fisher's exact, $P > 0.09$).

Examining flavin-containing monooxygenase 2 for evidence of natural selection

Genotyping of the g.23238C > T SNP and many neighbouring SNPs by the International HapMap project allowed us to investigate, using the Long-Range Haplotype test [29], whether a signal suggestive of positive selection of either allele at this locus could be detected. The *FMO2*1* allele frequency in the YRI dataset is 0.175, which is similar to that observed in sub-Saharan Africa. In contrast, *FMO2*1* was absent in the CEU and CHB + JPT datasets, consistent with previous studies [10,13].

The method of Voight *et al.* [33], which uses the statistic, standardized iHS, allows direct comparisons of SNPs of different frequencies and provides a measure of haplotype conservation around the target SNP in comparison with the rest of the genome. We used the web-based tool Haplotter, which applies the method of Voight *et al.* [33] on HapMap Phase II data, to look for evidence of recent positive selection at the g.23238C > T locus in the YRI dataset.

The standardized *iHS* for this locus is 1.653, a value which lies in the 95th percentile on a standard normal curve. This indicates that the increased level of haplotype homozygosity on the derived T allele (as *iHS* is positive) is not significantly different (P value > 0.05 , two-tailed test) from that we would expect from the genome as a whole [an $iHS \geq 2$ (P value ≤ 0.05), would be considered statistically significant) and therefore provides no evidence of recent positive selection for either allele.

This particular analysis was not possible using the CEU or JPT and CHB datasets because the g.23238C > T SNP is monomorphic in these populations. Examination of the whole *FMO2* gene, which, however, involves examining the proportion of SNPs in the gene that have extreme *iHS* values in comparison with other genes (see Voight *et al.* [33]), again using Haplotter, showed no evidence of selection in any population (P values: CEU = 0.084736, YRI = 0.406732, JPT and CHB = 0.608040).

Analysis of National Institute of Environmental Health Sciences flavin-containing monooxygenase 2 resequencing data

The NIEHS SNP program identified, from whole gene sequencing, 19 *FMO2* coding-region variants among the Panel 2 samples (see Table 5), 14 of which were reported earlier [15] in African-Americans. Four mutations were synonymous, nine were nonsynonymous, one was found in the 3' untranslated region, two were insertions (one of which was found in the 3' untranslated region), one was a deletion and two were premature stop codons (including 23238C > T).

After haplotype inference of the 12 NIEHS Yoruba individuals (24 chromosomes), four chromosomes were shown to possess the 23238C allele (see Table 5). Three of these chromosomes had identical haplotypes (haplotype 1), with two synonymous changes (g.13733G > A, g.22027G > A) in comparison with an ancestral reference sequence (elucidated from chimpanzee and macaque data), one of which was found only on a 23238C background (g.22027G > A). The fourth chromosome had an additional, nonsynonymous, mutation [g.18237G > A (R238Q)] that was only found on a 23238C background (haplotype 2).

Addition of the 15 phased NIEHS African-American samples (30 chromosomes) showed that a further seven chromosomes possessed the 23238C SNP. Six of the seven had the two synonymous mutations whereas the other lacked the g.13733G > A variant (haplotype 3). The R238Q variant was also found in two 23238C African-American individuals whereas an additional nonsynonymous mutation [g.19910G > C (R391T)] was found in a further two 23238C chromosomes (haplotype 4).

The 23238T-possessing chromosomes found in the Yoruba, African-American, European, Hispanic and Asian NIEHS samples possessed a number of variants including nonsynonymous and synonymous mutations as well as insertions and deletions, often in combination. For example, the g.7702_7703insGAC insertion is found on the same background as a deletion (g.10951delG), a stop codon (g.23238T) and two nonsynonymous mutations [g.7731T > C (F81S) and g.13732C > T (S195L)] ($n = 15$, haplotypes 21, 22 and 23).

Utilizing phased *FMO2* genomic data for the Yoruba NIEHS samples produced an estimate of the time of occurrence of the 23238C > T mutation of 502 404 years before (lower boundary: $2 \times 4889 \times 0.816 \times 19.4 = 154790$ years before, upper boundary: $2 \times 8751 \times 1.648 \times 36.1 = 1\ 041\ 243$ years before), using the coalescent-based method described by Griffiths and Majoram [34] (see Supplementary Section 1 for a full explanation).

Discussion

Functional flavin-containing monooxygenase 2 is found at high frequency throughout sub-Saharan Africa

The g.23238C > T SNP allele distribution reported in this study is consistent with our expectation based on the proportion of *FMO2*1* in African–American and Hispanic individuals. The ancestral allele of g.23238C > T is present at even higher frequencies in most sub-Saharan populations than in the admixed populations of the Americas, with approximately one-third of individuals possessing this variant.

Our results suggest that frequencies of g.23238C > T alleles are fairly similar throughout most of sub-Saharan Africa. Two groupings, the Anuak and south Sudan, which, however, display significantly higher frequencies of the ancestral allele (including geographically neighbouring populations) are present. The Anuak in Ethiopia are thought to be an immigrant population associated with a larger group of Anuak, who reside in southeastern Sudan (personal correspondence Ambaye–Ogato). This may go some way to explaining the similar high 23238T allele frequencies observed in these two populations.

The substantial difference in *FMO2*1* allele frequencies between northern African and sub-Saharan African populations is consistent with other genetic studies [35,36] which show the Saharan desert acting as a major barrier to gene flow. The presence of the *FMO2*1* allele at a low frequency in the Maghreb as well as in the Yemen could be a consequence of the Arab slave trade during the 8–19th centuries [37,38]. The absence of *FMO2*1* from the Turkish datasets is in agreement with earlier work, which has shown that the *FMO2*1* allele is not present in populations that are not of recent African descent [10,13].

The possible consequences of flavin-containing monooxygenase 2 functionality in Africans

Given the observed similarity in the distribution of the g.23238C > T polymorphism across sub-Saharan Africa, [population approximately 726 million World Bank estimate (<http://www.worldbank.org>)] and assuming the *FMO2*1* allele in Africans results in a fully functional *FMO2* enzyme [14]) we estimate that some 220 million individuals may express a functional enzyme. A considerable number are therefore potentially at risk of thiourea toxicity. This need to urgently establish the scale of potential risk is reinforced by the current widespread use of ethionamide in the treatment of tuberculosis. Drugs that are primarily metabolized by FMOs may, in general, have certain advantages over those metabolized by cytochrome P450 enzymes, because FMOs are not as readily inhibited or induced, thus reducing the risk of drug–drug interactions [39]. If, however, *FMO2* is involved in the metabolic pathway of drugs used to treat common diseases in Africa and if products of enzymatic activity have a toxic effect then great caution should be applied in the distribution and use of such drugs.

The evolution of flavin-containing monooxygenase 2 in humans

Sequence data from the NIEHS SNP programme for Yoruba and African–American samples support the suggestion that the *FMO2*1* allele results in functionally active *FMO2*. Although, we offer no statistical support, because of uncertainty in regard to certain aspects of the NIEHS data (i.e. there is possible error in haplotype inference because of the presence of very rare variants and there are large regions where successful sequencing coverage in all samples has not been achieved), it would seem that a large majority of variants that may affect the functional activity of the enzyme lie on an *FMO2*2A* background. This suggests that chromosomes possessing this allele are in mutational free fall because of the loss of function caused by the g.23238C > T mutation, whereas chromosomes with *FMO2*1* may have been evolutionarily conserved as they still retain enzymatic activity. Given the small number of g.23238C-

possessing individuals ($n = 4/12$) in the NIEHS Yoruba dataset it is, however, necessary to be cautious in drawing conclusions about *FMO2* activity in Africa from these data alone.

The Long-Range Haplotype test revealed no evidence for positive selection on either allele at the g.23238C > T SNP in any of three HapMap populations (YRI, CEU, CHB and JPT). Therefore, on the basis of current knowledge it is not possible to conclude that the high frequency of the derived *FMO2**2A allele is a consequence of selective advantage. Sabeti *et al.* [29] have, however, indicated that the extended haplotype homozygosity statistic is unable to detect positive selection that has occurred more than 30 000 years ago, so we cannot dismiss the possibility that a strong selective pressure existed before this date which resulted in the increase in *FMO2**2A allele frequency and the complete loss of the *FMO2**1 allele outside of Africa. Though not statistically significant the relatively high iHS value for the g.23238C > T SNP in the Yoruba, the almost significant P value for the *FMO2* gene in the Europeans ($P = 0.085$) and the marked difference in European/Asian and Africa 23238C frequency suggests that this scenario may be a possibility. Additional analysis and resequencing is required (using an approach similar to that of Xue *et al.* [40]) to establish whether or not there is evidence of selection acting on either allele, either within sub-Saharan Africa or outside this region.

If selection is not the cause of the elevated frequency of the derived allele then, given the presence of *FMO2**1 throughout sub-Saharan Africa at roughly similar frequencies, the most likely explanation for why the *FMO2**1 allele is not present outside Africa is because it was lost in a bottleneck when anatomically modern humans migrated out of Africa sometime after 65 000 years ago [41,42] and that therefore the g.23238C > T SNP must have a sub-Saharan African origin before this event. Dating when the g.23238C > T SNP arose, through the use of NIEHS sequencing data, seem to be, notwithstanding the need to apply somewhat crude assumptions, to support the ancient origin of this SNP with a time of 502 404 years before, well before any estimates of the first exodus of modern humans from Africa.

Conclusion

Sub-Saharan Africa is thought to possess more human genetic diversity than the rest of the world combined. It is, however, not yet clear how this diversity is distributed and indeed what part of that diversity is not present outside the continent. Paucity of such knowledge can lead to inappropriate therapeutic, prophylactic and diagnostic intervention and increase the risk of an adverse drug reaction. Surveys such as the one performed here are not only of benefit to the indigenous populations of Africa, but are also of increasing importance in the planning of healthcare in the developed world, where the number of individuals of recent African descent is growing and, in some areas, such as the Americas and Europe, is already substantial. A need for more studies on human genetic diversity in Africa is observed; research from which all people of recent African descent, wherever living, should benefit.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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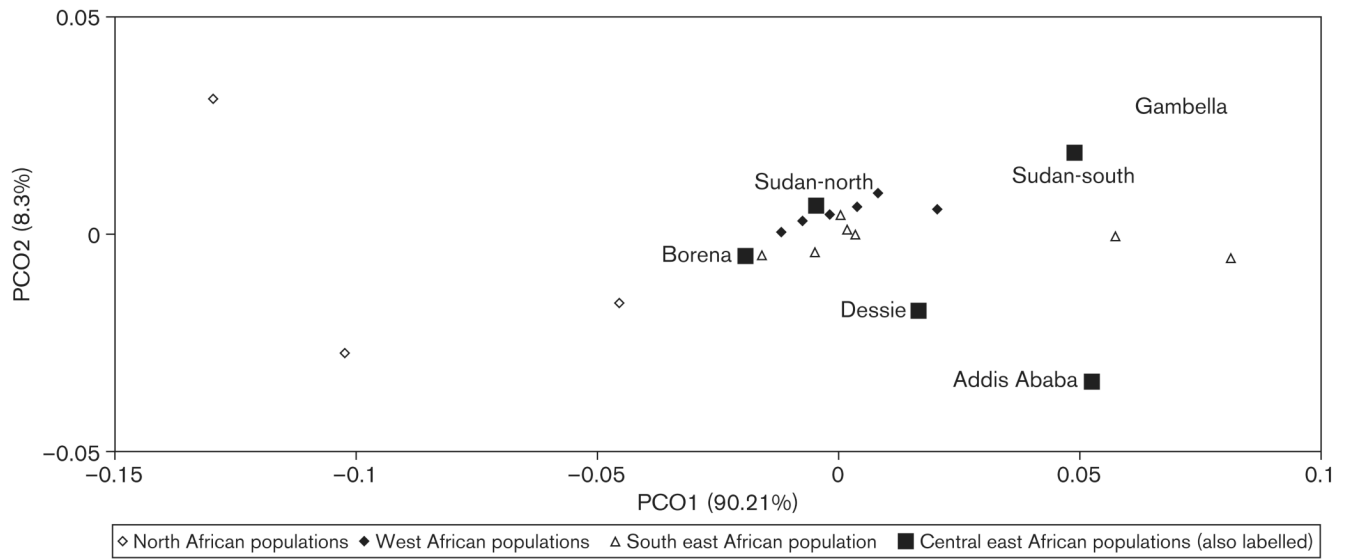


Fig. 2. Principal coordinates analysis (PCO) plot of 23238 C > T-based population F_{ST} values.

Table 1

Genotype and allele frequencies

| Country | Population | Cultural identity | <i>FM02*1/FM02*1</i> | <i>FM02*1/FM02*2A</i> | <i>FM02*2A/FM02*2A</i> | <i>n</i> | <i>FM02*1</i> frequency | <i>FM02*2A</i> frequency | At least one <i>FM02*1</i> allele | Latitude | Longitude |
|----------|------------|-------------------|----------------------|-----------------------|------------------------|----------|-------------------------|--------------------------|-----------------------------------|----------|-----------|
| Ethiopia | Various | Various | 2 | 39 | 78 | 119 | 0.181 | 0.819 | 34.45% | 6.542 | 11.454 |
| | Various | Various | 2 | 25 | 49 | 76 | 0.191 | 0.809 | 35.53% | 12.28 | 14.752 |
| Kenya | Bulsa | | 3 | 21 | 66 | 90 | 0.15 | 0.85 | 26.67% | 10.726 | -1.279 |
| | Kasena | | 1 | 7 | 37 | 45 | 0.1 | 0.9 | 17.78% | 10.884 | -1.085 |
| Nigeria | Igbo | | 0 | 22 | 66 | 88 | 0.125 | 0.875 | 25.00% | 4.957 | 8.314 |
| | Manji | | 3 | 25 | 66 | 94 | 0.165 | 0.835 | 29.79% | 12.986 | -15.88 |
| Senegal | Wolof | | 1 | 24 | 70 | 95 | 0.137 | 0.863 | 26.32% | 14.687 | -17.452 |
| | Various | | 5 | 47 | 54 | 106 | 0.269 | 0.731 | 49.06% | 8.25 | 34.583 |
| Tanzania | Amhara | | 1 | 7 | 16 | 24 | 0.188 | 0.813 | 33.33% | 9.006 | 38.852 |
| | Various | | 0 | 7 | 29 | 36 | 0.097 | 0.903 | 19.44% | 10.75 | 38.767 |
| Togo | Amhara | | 1 | 6 | 19 | 26 | 0.154 | 0.846 | 26.92% | 11.231 | 39.526 |
| | Various | | 2 | 37 | 97 | 136 | 0.151 | 0.849 | 28.68% | 15.213 | 33.036 |
| Zambia | Various | | 10 | 43 | 73 | 126 | 0.25 | 0.75 | 42.06% | 10.854 | 29.772 |
| | Various | | 4 | 35 | 105 | 144 | 0.149 | 0.851 | 27.08% | -13.983 | 33.774 |
| Zimbabwe | Various | | 1 | 16 | 43 | 60 | 0.15 | 0.85 | 28.33% | -14.467 | 35.267 |
| | Various | | 0 | 17 | 39 | 56 | 0.152 | 0.848 | 30.36% | -11.465 | 34.023 |
| Zimbabwe | Sena | | 2 | 22 | 60 | 84 | 0.155 | 0.845 | 28.57% | -17.442 | 35.027 |

| Country | Cultural identity | <i>FM02*1/FM02*1</i> | <i>FM02*1/FM02*2A</i> | <i>FM02*2A/FM02*2A</i> | <i>n</i> | <i>FM02*1</i> frequency | <i>FM02*2A</i> frequency | At least one <i>FM02*1</i> allele | Latitude | Longitude |
|---------|-------------------|----------------------|-----------------------|------------------------|----------|-------------------------|--------------------------|-----------------------------------|----------|-----------|
| | Bantu | 2 | 15 | 24 | 41 | 0.232 | 0.768 | 41.46% | -25.753 | 28.297 |
| | Chagga | 2 | 18 | 30 | 50 | 0.22 | 0.78 | 40.00% | -5.383 | 38.05 |
| | Bantu | 0 | 10 | 29 | 39 | 0.128 | 0.872 | 25.64% | -0.452 | 32.564 |
| | Shona | 0 | 7 | 27 | 34 | 0.103 | 0.309 | 20.59% | -17.309 | 31.328 |
| | Unspecified | 0 | 5 | 38 | 43 | 0.058 | 0.942 | 11.63% | 35.94 | 0.09 |
| | Unspecified | 0 | 10 | 108 | 118 | 0.042 | 0.958 | 8.47% | 35.083 | -2.183 |
| | Berber | 0 | 3 | 67 | 70 | 0.021 | 0.979 | 4.29% | 33.588 | -5.165 |
| | Anatolian Turks | 0 | 0 | 31 | 31 | 0 | 1 | 0.00% | 40.277 | 33.254 |
| | Anatolian Turks | 0 | 0 | 28 | 28 | 0 | 1 | 0.00% | 39.684 | 31.212 |
| | Unspecified | 0 | 4 | 30 | 34 | 0.059 | 0.941 | 11.76% | 15.409 | 44.242 |
| | Unspecified | 1 | 5 | 77 | 83 | 0.042 | 0.958 | 7.23% | 16.811 | 49.942 |
| | Total | 43 | 477 | 1456 | 1976 | 0.142 | 0.858 | 26.32% | | |

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Table 2**Pearson's χ^2 test on individual regions**

| Region | df | χ^2 | P value |
|--------|----|----------|---------|
| CEA | 5 | 18.12 | 0.003* |
| NA | 2 | 2.15 | 0.34 |
| SEA | 7 | 7.51 | 0.37 |
| WA | 6 | 7.34 | 0.29 |
| Yemen | 1 | 0.64 | 0.43 |

df, degrees of freedom.

* P value is less than 0.05.

CEA, central east Africa; NA, north Africa; SEA, south-east Africa; WA, west Africa.

Table 3

Fisher's exact tests between regions

| | WA | SEA | NA | TU |
|-----|---------|---------|---------|--------|
| SEA | 0.7915 | \ | \ | } |
| NA | 0.0001* | 0.0001* | \ | } |
| TU | 0.0001* | 0.0001* | 0.0294* | } |
| YE | 0.0001* | 0.0001* | 0.8361 | 0.0321 |

* *P* value is less than 0.05.

NA, north Africa; SEA, south-east Africa; TU, Turkey; WA, west Africa; YE, Yemen.

Table 4

Fisher's exact tests between CEA populations

| | Gambella | Addis Ababa | Borena, Wollo | Dessie, Wollo | Sudan north |
|---------------|----------|-------------|---------------|---------------|-------------|
| Addis Ababa | 0.1812 | | | | |
| Borena, Wollo | 0.0018* | 0.2418 | | | |
| Dessie, Wollo | 0.0492* | 0.7598 | 0.5475 | | |
| Sudan north | 0.0013* | 0.6340 | 0.2982 | 1.0000 | |
| Sudan south | 0.2933 | 0.5008 | 0.0180* | 0.1883 | 0.0277* |

* *P* value is less than 0.05.

CEA, central east Africa.

| g. 19679A > G (E314G) | g. 19839A > G (Synon) | g. 19910G > C (R391T) | g. 22027G > A (Synon) | g. 22060T > G (N413K) | g. 23087A > G (Synon) | g. 23238C > T (Q472X) | g. 23300A > G (3' UTR)* | Ethnic identity of NIEHS samples | | | | | Total | |
|--------------------------------|--------------------------------|--------------------------------|--------------------------------|--------------------------------|--------------------------------|--------------------------------|----------------------------------|----------------------------------|----|----|----|----|-------|---|
| | | | | | | | | AA | YR | AS | EU | HI | | |
| A | A | G | G | T | A | C | A | 2 | 3 | 0 | 0 | 0 | 0 | 5 |
| | | | A | | | | | 2 | 1 | 0 | 0 | 0 | 0 | 3 |
| | | | A | | | | | 1 | 0 | 0 | 0 | 0 | 0 | 1 |
| | | C | A | | | | | 2 | 0 | 0 | 0 | 0 | 0 | 2 |
| | G | | | G | | T | G | 1 | 0 | 0 | 0 | 0 | 0 | 1 |
| | G | | | | | T | G | 1 | 4 | 1 | 2 | 3 | 11 | |
| G | G | | | | | T | | 0 | 0 | 0 | 1 | 0 | 1 | |
| | G | | | G | | T | | 0 | 1 | 0 | 0 | 0 | 1 | |
| | G | | | | | T | | 9 | 0 | 22 | 23 | 13 | 67 | |
| | G | | | | | T | G | 0 | 0 | 0 | 2 | 2 | 4 | |
| | G | | | | G | T | | 0 | 0 | 0 | 0 | 1 | 1 | |
| | G | | | | | T | | 0 | 0 | 0 | 1 | 0 | 1 | |
| | G | | | G | | T | | 0 | 1 | 0 | 0 | 0 | 1 | |
| | G | | | | | T | | 1 | 0 | 0 | 0 | 0 | 1 | |
| | G | | | | | T | | 0 | 0 | 0 | 1 | 0 | 1 | |
| | G | | | | | T | | 0 | 0 | 0 | 1 | 0 | 1 | |
| | G | | | G | | T | | 2 | 3 | 0 | 7 | 4 | 16 | |
| | G | | | | | T | | 0 | 4 | 0 | 0 | 0 | 4 | |
| | G | | | | | T | G | 0 | 0 | 5 | 3 | 6 | 14 | |
| | G | | | | | T | | 1 | 0 | 12 | 0 | 9 | 22 | |
| | G | | | G | | T | | 6 | 5 | 0 | 0 | 1 | 12 | |
| | G | | | | | T | | 2 | 0 | 0 | 0 | 0 | 2 | |
| | G | | | | | T | G | 0 | 0 | 0 | 1 | 0 | 1 | |
| | G | | | | | T | | 0 | 1 | 0 | 0 | 2 | 3 | |
| | G | | | | | T | | 0 | 1 | 8 | 1 | 1 | 11 | |
| | G | | | | | T | | 0 | 0 | 0 | 1 | 2 | 3 | |

| | | | | | | | | | Ethnic identity of NIEHS samples | | | | | Total |
|---------------------------------|---|--|--|--|--|--|--|--|----------------------------------|----|----|----|----|-------|
| | | | | | | | | | AA | YR | AS | EU | HI | |
| g. 19679A > G (E314G) | A | | | | | | | | 30 | 24 | 48 | 44 | 44 | 190 |
| g. 19839A > G (Synon) | A | | | | | | | | | | | | | |
| g. 19910G > C (R391T) | G | | | | | | | | | | | | | |
| g. 22027G > A (Synon) | G | | | | | | | | | | | | | |
| g. 22060T > G (N413K) | T | | | | | | | | | | | | | |
| g. 23087A > G (Synon) | A | | | | | | | | | | | | | |
| g. 23238C > T (Q472X) | C | | | | | | | | | | | | | |
| g. 23300A > G (3' UTR) | A | | | | | | | | | | | | | |
| g.23412_23413insT (3'UTR)* | - | | | | | | | | | | | | | |