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Debrisoquin hydroxylase (CYP2D6) is a cytochrome P450 enzyme that catalyzes the metabolism of > 30 commonly prescribed medications. Deficiency in CYP2D6 activity, inherited as an autosomal recessive trait, was found to be significantly less common in American blacks (1.9%) than whites (7.7%). To determine the genetic basis for this difference, inactivating CYP2D6 mutations were assessed by allele-specific PCR amplification and RFLP analyses of genomic DNA from 126 unrelated whites and 127 unrelated blacks. Blacks had a twofold lower frequency (8.5 versus 23%, P = 0.001) of the CYP2D6(B) mutation (point mutation at intron 3/exon 4 splice site), while complete deletion of the CYP2D6 gene (5.5% blacks, 2.4% whites), and the CYP2D6(A) mutation (single nucleotide deletion in exon 5; 0.24% blacks, 1.4% whites) were not different between the two groups. The prevalence of heterozygous genotypes was significantly lower in blacks (25 versus 42% of extensive metabolizers, P = 0.009), consistent with the observed prevalence of the deficient trait in blacks and whites. We conclude that the same CYP2D6 mutations lead to a loss of functional expression in blacks and whites, but American blacks have a lower prevalence of the deficient trait due to a lower frequency of the CYP2D6(B) mutation. This could explain racial differences in drug effects and disease risk.

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Genetic Basis for a Lower Prevalence of Deficient CYP2D6 Oxidative Drug Metabolism Phenotypes in Black Americans

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Abstract

Debrisoquin hydroxylase (CYP2D6) is a cytochrome P450 enzyme that catalyzes the metabolism of >30 commonly prescribed medications. Deficiency in CYP2D6 activity, inherited as an autosomal recessive trait, was found to be significantly less common in American blacks (1.9%) than whites (7.7%). To determine the genetic basis for this difference, inactivating CYP2D6 mutations were assessed by allele-specific PCR amplification and RFLP analyses of genomic DNA from 126 unrelated whites and 127 unrelated blacks. Blacks had a twofold lower frequency (8.5 versus 23%, P < 0.001) of the CYP2D6(B) mutation (point mutation at intron 3/exon 4 splice site), while complete deletion of the CYP2D6 gene (5.5% blacks, 2.4% whites), and the CYP2D6(A) mutation (single nucleotide deletion in exon 5; 0.24% blacks, 1.4% whites) were not different between the two groups. The prevalence of heterozygous genotypes was significantly lower in blacks (25 versus 42% of extensive metabolizers, P = 0.009), consistent with the observed prevalence of the deficient trait in blacks and whites. We conclude that the same CYP2D6 mutations lead to a loss of functional expression in blacks and whites, but American blacks have a lower prevalence of the deficient trait due to a lower frequency of the CYP2D6(B) mutation. This could explain racial differences in drug effects and disease risk. (J. Clin. Invest. 1993. 91:2150–2154.) Key words: CYP2D6 • pharmacogenetics • ethnic differences • oxidative metabolism

Introduction

Debrisoquin hydroxylase (CYP2D6) is a cytochrome P450 enzyme that catalyzes the metabolism of >30 commonly prescribed medications, including beta receptor antagonists (e.g., metoprolol, propranolol), antiarrhythmics (e.g., encainide, propafenone), antidepressants (imipramine, desipramine), analgesics (e.g., codeine), antitussives (e.g., dextromethorphan), and antipsychotics (e.g., haloperidol, thioridazine). Previous population studies have shown that ~7–10% of United States and European Caucasians are deficient in CYP2D6 activity, (1–3) resulting in significantly impaired metabolism of medications that are substrates solely for this enzyme (4–6).

CYP2D6 is under monogenic control and the poor metabolizer (PM) phenotype (i.e., CYP2D6 deficiency) is inherited as an autosomal recessive trait (2). Deficient CYP2D6 activity in PM livers is associated with the absence of immunodetectable CYP2D6 protein (7), which is caused by one of several gene defects (e.g., gene deletion or point mutations/nucleotide changes) (8, 9). In Caucasians, two prevalent inactivating mutations in the CYP2D6 gene (CYP2D6) are a point-mutation at the 3′ splice site consensus sequence of intron 3 (CYP2D6(B) mutation), and a single nucleotide (adenine) deletion in exon 5 (CYP2D6(A) mutation) (10–12). Either of these mutations lead to defective mRNA and protein (9), and a deficiency in CYP2D6 activity. A third variant of the CYP2D6 locus leading to the absence of CYP2D6 activity, is a complete deletion of the CYP2D6 gene (13), which is detectable by RFLP analysis (11.5-kb XbaI restriction fragment) (14).

In a study of 468 unrelated white and 105 unrelated black subjects, we found a significantly lower prevalence of the CYP2D6 deficient trait in American blacks (1.9%) versus whites (7.7%) (3). The fourfold lower prevalence of PM phenotypes in blacks indicates that blacks are less likely to experience exaggerated effects of drugs that are principally inactivated by CYP2D6 (e.g., metoprolol [5], propafenone [4]), are less likely to fail therapy with drugs that must be activated by CYP2D6 (e.g., codeine [15]), and would have a different risk of diseases associated with CYP2D6 phenotype. The present study was undertaken to define the genetic basis for this racial difference in the CYP2D6 poor metabolizer trait.

Methods

Human subjects. Subjects from whom blood was collected were enrolled from the population of patients, families, and employees at St. Jude Children’s Research Hospital, from local schools and from healthy volunteer blood donors. The protocol was approved by the institutional review board for clinical trials. Informed consent was obtained from all subjects over 12 yr of age; parental consent was obtained for all subjects <18 yr of age. For subjects who had CYP2D6 phenotype determined, a complete medication history for the previous 2 wk and basic demographic and family pedigree data were collected by a nurse. Excluded from the study were subjects with known renal or hepatic disease, those related to another study subject, and those taking

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1. Given the uncertainty of true ancestry, “black” and “white” are used within this manuscript to designate individuals of presumed African and Caucasian ancestry, respectively.

2. Abbreviations used in this paper: A or CYP2D6(A), nonfunctional CYP2D6 with single nucleotide deletion in exon 5; B or CYP2D6(B), nonfunctional CYP2D6 with point mutation at intron 3/exon 4 splice site; CYP2D6, debrisoquin hydroxylase cytochrome P450; del, complete deletion of CYP2D6 gene; EM, extensive metabolizer phenotype; PM, poor metabolizer phenotype; wt, wild type CYP2D6 allele without known mutations.
medications known to induce or inhibit cytochrome P450 enzymes or to be substrates for CYP2D6 (within 2 wk of the study). Subjects were classified as either black or white, based on their statement of race on a questionnaire and its agreement with skin color.

DNA extraction, RFLP, and PCR-based detection of CYP2D6 mutations. In all subjects, 15–30 ml of blood was collected in potassium EDTA tubes; DNA was isolated from leukocytes after proteinase K digestion, phenol-chloroform extraction, and precipitation with isopropanol/sodium acetate (16).

For identification of the CYP2D6(A) and CYP2D6(B) mutations, mutation-specific PCR amplification was performed as previously described (10), with the following modifications. The magnesium concentration was 1.5 mM; the concentration of each oligonucleotide primer was 0.125 µM; Taq polymerase was obtained from Perkin-Elmer Cetus (Norwalk, CT); and PCR was carried out using a thermocycler (Ericalc, Inc., San Diego, CA). The sequences (5′–3′) of oligonucleotide primers to detect the CYP2D6(A) mutation were as follows (10): first round amplification to isolate appropriate segment of CYP2D6 (2098–3200, numbered according to Kimura et al. [17]) from CYP2D7 and CYP2D8 with primer A1 = GCCGAGCCAGA-GACGGAGGA and primer A2 = CCGGCCTGACACTTCCTCT; second round amplification with A2 plus A3 = GTAACTGAG- GCACA (to amplify only wild type sequences) or A2 plus A4 = GTAACGTGACCCG (to amplify only mutant sequences). The sequences (5′–3′) of oligonucleotide primers to detect the CYP2D6(B) mutation were as follows: first round amplification to isolate the appropriate segment of CYP2D6 (1385–2122) from CYP2D7 and CYP2D8 with primer B1 = ATTTCGGAGCTGGATCC and B2 = GAGACT- CTCGCTTCTCT; second round amplification with B1 plus B3 = CGAAAGGGCGTCTCC (to amplify only wild type sequence) or primer B1 plus B4 = CGAAAGGGCGTCT (to amplify only mutant sequence).

For identification of the CYP2D6 gene deletion, RFLP analysis was performed (14). DNA (5–10 µg) was digested to completion with XbaI restriction endonuclease under conditions recommended by the source (New England Biolab, Beverly, MA), and subjected to agarose gel (0.5%) electrophoresis. Southern blotting was performed by alkaline transfer to GeneScreen Plus™ membranes using conditions recommended by the source (DuPont, Boston, MA). The 1.6-kb CYP2D6 cDNA probe (kindly provided by Dr. Frank Gonzalez, National Institutes of Health) was prepared by nick translation with [32P]dCTP (6,000 Ci/mmol; Amersham Corp., Arlington Heights, IL) to a specific activity of 1–3 x 10⁹ dpm/µg DNA, hybridized to digested DNA, and the restriction fragment length patterns examined after film exposure for 4–10 d at 80°C. The presence of a band at 11.5 kb identified the CYP2D6 gene deletion (14).

Determination of CYP2D6 phenotype. After collection of a blank urine sample, dextromethorphan (30 mg) was administered orally to a subset of these subjects, as previously described (18). Urine was collected over the next 4 h, aliquots frozen, and later thawed and assayed by high performance liquid chromatography for dextromethorphan and its polymorphically formed metabolite, dextrorphan (19). The molar concentration ratio of dextromethorphan/dextrorhan was used to assign phenotype, which has been previously shown to exactly co-segregate with that assigned using debrisoquin itself (19). A urinary metabolic ratio < 0.3 defined extensive metabolizers (EMs) and values ≥ 0.3 defined subjects with the CYP2D6 deficiency (PMs), consistent with the antimode previously established by population studies of blacks and whites in our laboratory (3) and by others (20). Where dextromethorphan was undetectable because of very extensive metabolism to dextrorphan, the ratio was assigned a value equal to the ratio calculated using the lowest measurable dextromethorphan concentration for each sample; this occurs only in extensive metabolizers.

Statistical analysis. Chi-square analysis was used to assess the difference in proportions (e.g., proportions of homozygous EM genotypes in the blacks versus whites), except when the number of observations was small (e.g., count < 5 in a cell), for which Fisher’s exact test was used. For testing hypotheses about population proportions, such as whether the observed proportion of heterozygous EM genotypes in each racial population agrees with the predicted proportion (based on Hardy-Weinberg calculations), normal approximation to the binomial distribution was used.

Results

PCR and RFLP analyses were performed on 253 unrelated subjects, 126 whites and 127 blacks. The proportion of males and females was not different in blacks (43% females) versus whites (45% females). Assignment of genotype, based on mutation-specific PCR amplification, is exemplified in Fig. 1 for each of the five genotypes detected by PCR. There were 106 whites and 123 blacks with at least one wild type CYP2D6 allele (i.e., EM genotype), and 20 whites and 4 blacks with two mutant alleles for CYP2D6 (i.e., PM genotype). The higher proportion of PM genotypes in the present study, compared to the expected 7.7 and 1.9%, is reflective of efforts to obtain DNA from all consenting subjects identified as a PM phenotype.

As summarized in Table 1, a higher percentage of blacks (74.8%) had a homozygous EM genotype when compared with whites (58.5%). Chi-square P = 0.009. As follows, the percentage of heterozygotes among EMs was lower in blacks (25.2%) than in whites (41.5%), Chi-squared P = 0.009. The percent-

![CYP2D6 PCR](image-url)

Figure 1. Mutation-specific PCR amplification of the two major inactivating mutations in CYP2D6 gene (i.e., CYP2D6(A) and CYP2D6(B)). Shown are results using genomic DNA from three individuals who have an extensive metabolizer (EM) phenotype (two heterozygous and one homozygous EMs) and two who have the poor metabolizer (PM) trait. In the upper portion of figure: B and A, initial oligonucleotide primers to isolate the segments of CYP2D6 (from CYP2D7 and CYP2D8) which contain either the B or A mutations (i.e., either the region around the intron 3/exon 4 junction for B, or the region around exon 5 for A). After these segments of CYP2D6 have been amplified in the first round of PCR (not shown), different sets of primers are used to detect the presence or absence of mutations: W, PCR with oligonucleotide primers identical to the wild type genomic sequence; M, oligonucleotide primers identical to the respective mutant gene sequences. In the lower part of the figure indicating genotype: Wt, wildtype genomic sequence; A, amplification with primers recognizing mutation in exon five (single A nucleotide deletion); B, amplification with primers recognizing mutation at intron 3 exon 4 splice site (point mutation). Exact sequences of each primer are given in Methods. As depicted, this PCR method allows the assignment of genotype and differentiation of homozygous (wt/ wt) EM genotypes, heterozygous EM genotypes (wt/A, wt/B), and PM genotypes. RFLP analysis is required to determine the complete deletion of CYP2D6, the third major inactivating mutation at the CYP2D6 locus.
Table I. CYP2D6 Genotypes in American Blacks and Whites

<table>
<thead>
<tr>
<th>EM genotypes</th>
<th>Whites (%)</th>
<th>Blacks (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>wt/wt</td>
<td>62 (58.5)</td>
<td>92 (74.8)*</td>
</tr>
<tr>
<td>wt/A</td>
<td>1 (0.9)</td>
<td>0</td>
</tr>
<tr>
<td>wt/B</td>
<td>38 (35.9)</td>
<td>19 (15.5)</td>
</tr>
<tr>
<td>wt/del</td>
<td>5 (4.7)</td>
<td>12 (9.8)</td>
</tr>
<tr>
<td>PM genotypes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>B/B</td>
<td>14 (70)</td>
<td>1 (25)</td>
</tr>
<tr>
<td>A/A</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>A/B</td>
<td>5 (25)</td>
<td>0</td>
</tr>
<tr>
<td>B/del</td>
<td>1 (5)</td>
<td>2 (50)</td>
</tr>
<tr>
<td>A/del</td>
<td>0</td>
<td>1 (25)</td>
</tr>
<tr>
<td>Total</td>
<td>20 (100)</td>
<td>4 (100)</td>
</tr>
</tbody>
</table>

* Proportion of homozygous EMs was significantly higher in blacks than in whites. $P = 0.009$, chi-square.

Table II. CYP2D6 Allele Frequencies in American Blacks and Whites

<table>
<thead>
<tr>
<th>PM genotype</th>
<th>EM genotype</th>
<th>Total population*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Allele</td>
<td>Blacks</td>
<td>Whites</td>
</tr>
<tr>
<td></td>
<td>$n = 8$</td>
<td>$n = 40$</td>
</tr>
<tr>
<td>Wt</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>A</td>
<td>0.125</td>
<td>0.125</td>
</tr>
<tr>
<td>B</td>
<td>0.50±</td>
<td>0.85</td>
</tr>
<tr>
<td>Del</td>
<td>0.375±</td>
<td>0.025</td>
</tr>
</tbody>
</table>

* Allele frequencies in total population based on a 1.9% and 7.7% prevalence of the poor metabolizer trait in blacks and whites, respectively (i.e., sum of values for EMs multiplied by 0.923 [whites] or 0.981 [blacks], and values for PMs by 0.077 [whites] or 0.019 [blacks]). Numbers in parentheses are the number of alleles assessed by PCR and RFLP analyses. $^\dagger$ $P < 0.05$, blacks versus whites.

Table II summarizes the CYP2D6 allele frequencies in all black and white subjects with PM genotypes, EM genotypes, and values extrapolated to the total population of blacks and whites. The frequency of mutant alleles was 14.3% for the black population versus 26.9% for the white population ($P < 0.001$). The lower frequency of mutant alleles in blacks was due almost entirely to a lower frequency of the CYP2D6(B) mutation (8.5% vs 23%, $P < 0.001$).

The mutations found in all black ($n = 4$) and white ($n = 20$) subjects with a PM genotype are also summarized in Table II. The CYP2D6(B) mutation was again the most prevalent mutation in both blacks and whites, accounting for 85% of mutant alleles in white PMs and 50% in black PMs (Fisher's exact $P = 0.047$). The CYP2D6(A) mutation accounted for 12.5% of alleles in both black and white subjects with PM genotypes, while the complete deletion of CYP2D6 represented 37.5% of alleles in black PMs and 2.5% of alleles in white PMs (Fisher's exact $P = 0.012$).

CYP2D6 phenotype was determined using dextromethorphan in 89 whites and 70 blacks with EM genotypes, and in 20 whites and 2 blacks with PM genotypes. The remainder of subjects were blood donors who did not participate in the clinical phenotyping studies ($n = 72$). In the subset of 181 subjects in whom clinical phenotype was determined, all 22 subjects with PM genotypes had a PM phenotype, while 154 of 159 subjects (96.9%) with EM genotypes had EM phenotypes. The lack of concordance between genotype and phenotype in five subjects (3%) with a wild type allele but a PM phenotype (four whites, one black), could be due to imprecision in phenotype assignment or to the presence of rare CYP2D6 inactivating mutations, which are not detected by current PCR and RFLP analyses (11).

Discussion

This study has established that blacks in the southeast United States have a lower frequency of CYP2D6 mutant alleles, yielding a lower prevalence of the autosomal recessive debrisoquin...
hydroxylase (CYP2D6) poor metabolizer trait. The present molecular studies were initiated after our clinical study revealed a 1.9% prevalence of the CYP2D6 PM trait in American blacks compared to a 7.7% prevalence in whites (3). According to Hardy-Weinberg calculations, the expected proportion of homozygous EMs in the entire population would be 74% for blacks and 52% for whites, if the PM trait has a true prevalence of 1.9 and 7.7%, respectively. Thus, the expected proportion of homozygous individuals within only the subset of individuals who are EMs would be 75% for blacks and 56% for whites, values in close agreement with those found in the present study (i.e., 74.8 and 58.5%). Therefore, the present study establishes a genetic basis for our clinical finding of a significant difference in CYP2D6 drug metabolism phenotypes in these two ethnic groups in the United States.

The present study has also established that the three major CYP2D6 inactivating mutations previously recognized in Caucasian populations (11, 12, 21), are also present in the black population. The lower overall prevalence of mutant alleles in blacks was due to a significantly lower frequency of the CYP2D6 (B) mutation in blacks (8.5 vs. 23%) while the frequencies of the CYP2D6 (A) mutation and the gene deletion were similar in both groups. That all evaluable subjects who carried two of these mutations were found to have a PM phenotype, demonstrates that these are inactivating mutations in both racial groups.

Ethnic differences in drug metabolism phenotypes were recognized >30 yr ago, and have now been documented for N-acetyltransferase in Asians and Caucasians (22), mephenytoin hydroxylation in Asians and Caucasians (23), and debrisoquin hydroxylase (CYP2D6) phenotypes in African blacks and whites (3) and in Asians and Caucasians (23). The genetic basis for these ethnic differences in cytochrome P450 enzymes has not been precisely defined. The present study has established that the difference in CYP2D6 drug metabolism phenotypes in American blacks and whites is due to genetic differences, and not environmental or other causes. Specifically, this ethnic difference in CYP2D6 phenotypes is due to a twofold lower frequency of the CYP2D6 (B) mutation in blacks. This mutation of CYP2D6 has been reported to comprise 22 and 27% of alleles in two studies of European Caucasians (12, 20), an allele frequency similar to that found in our white subjects (23%), but greater than the 8.5% of alleles in our black population (Table II). The frequencies of CYP2D6 (A) mutations and of complete deletion of the CYP2D6 gene were low in the European Caucasian populations, as was the case in our white and black populations. The frequency of wild type CYP2D6 alleles was 74% in both studies of European Caucasians, similar to the 73% found in our white population, but lower than the 86% frequency of wildtype alleles in our black population. Thus, CYP2D6 mutations in our white population are comparable to reports of two European Caucasian populations, but the frequency of CYP2D6 (B) mutations in our black population differs from all white populations reported to date.

The anthropology of these genetic differences is not known, but could be related to the introduction of Caucasian genes into the genome of American blacks. It has been reported that 20–25% of genes in some American black populations are from Caucasian ancestry, based on either ABO or Fy(a) allele frequencies (24). Moreover, several clinical studies have reported a very low prevalence of the CYP2D6 PM trait in African black populations. For example, the prevalence of CYP2D6 PM phenotype was found to be 0 in a study of 138 Nigerians (95% confidence interval = 0–1.4%, using 0.5 instead of 0 to estimate confidence interval), using both debrisoquin and metoprolol to determine phenotype (25). Likewise, Eichelbaum and Woolhouse (26) found 0 of 154 Nigerians to be CYP2D6 poor metabolizers when sparteine was used to assign phenotype. If the true prevalence of the CYP2D6 PM trait in African Americans of American blacks, then a 25% penetrance of Caucasian-derived genes in the American black population would be consistent with results in the present study, wherein the same inactivating mutations were present in both populations, but the overall frequency of mutant CYP2D6 alleles was significantly lower in blacks versus whites.

It should be noted that the proportion of Caucasian genes in American black populations may not be the same in all geographical locations in the United States (24). For example, using Fy(a) gene frequencies, the proportion in 1969 of Caucasian genes in blacks was higher in nonsouthern cities (e.g., New York, Detroit, Oakland; 19–26%) compared to southern locations (e.g., Georgia; 11%) (24). It is not known whether these differences persist in 1992, given the migratory nature of the United States population. Thus, it may be necessary to assess CYP2D6 genotypes and phenotypes in additional black populations, before the present findings can be uniformly extrapolated to other black populations in North America.

Since individuals with the PM trait are more likely to have exaggerated effects with some medications (e.g., metoprolol [5], propafenone [4]) and are more likely to fail therapy with drugs requiring activation by this enzyme (e.g., codeine [15]), these racial differences can be of great therapeutic importance. Moreover, the risk of lung cancer in smokers has been linked to high CYP2D6 activity (27, 28) and Parkinson’s disease has been associated with a higher frequency of CYP2D6 mutant alleles (29, 30) suggesting that ethnic differences at the CYP2D6 locus may also be important for understanding racial differences in the risk of certain diseases. For example, it is an intriguing possibility that the higher prevalence of CYP2D6 wt/wt genotypes contributes to the higher risk of lung cancer among blacks (31). The present study has established that black Americans have a lower frequency of mutant CYP2D6 genes, leading to ethnic diversity in the expression of an enzyme that metabolizes >30 commonly prescribed medications, a finding that has both therapeutic and toxicologic implications.

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