
Genetic polymorphisms of the Cytochrome P450 isof orm 2C19 (CYP2C19) gene—known to cause poor metabolism of important prescription drugs (e.g., anticonvulsants such as phenytoin, antidepressant, cancer chemotherapy, anti-malaria drugs, anti-ulcer proton-pump inhibitors, and methadone)—were examined in a Puerto Rican newborn population. At least seven single nucleotide polymorphisms (SNPs) in the CYP2C19 gene have been shown to influence enzyme activity. The CYP2C19*2 and CYP2C19*3 variants are most common and best characterized clinically. They both are considered non-functional alleles associated with lack of enzymatic activity. Patients with these variants will require significantly lower doses in order to avoid overexposure and increased risk of adverse effects to standard doses of CYP2C19 substrates. The frequencies of variant alleles for the CYP2C19 marker have been reported to differ among ethnic groups. 2C19*2 is 30% in general population; 85% in Asians and about 15% in Caucasians. 2C19*3 is less common, about 6-10% Asians; but rare (<1%) in Caucasians and African Americans (1-3). The allele frequencies of these two variants in the Puerto Rican population remain to be determined.

The objectives of this study were to determine the prevalence of clinically relevant allele variants (CYP2C19*2, splicing defect G681A SNP and CYP2C19*3, stop codon G636A SNP) in a Puerto Rican study population and finally to determine whether the allele distributions for gene mutations meet Hardy–Weinberg equilibrium (HWE).

Genomic DNA were extracted from leukocytes in 122 dried blood filter samples from different regions of Puerto Rico, which were kindly provided by the Puerto Rico Newborn Screening Program. Samples were analyzed by the PCR-RFLP technique at the Molecular Genetic Core facilities of the School of Medicine, UPR as well as by the Physiogenomic-array with the Illumina BeadArray™ platform, at Laboratory of Personalized Health (LPH), Hartford, CT (4-6).

We were able to obtain preliminary results for the CYP2C19*2 allele from 122 individuals (figures 1 and 2). The results for the CYP2C19*2 allele have showed that 90 of these DNA samples (73.7%) were found to be homozygous for the wild-type allele; and 30 DNA samples (24.6%) were single carrier for the allelic variant *2 (heterozygous); whereas, two DNA sample (1.64%) were double-carrier for this clinically relevant mutation. Genotype analysis also revealed that the allele frequency for CYP2C19*2 was 0.139. Notably, these findings were confirmed by Illumina BeadArray™ technology genotyping platform using highly multiplexed ligation.
assays (i.e., minor allele frequency of 0.1406 was detected using a Genomas proprietary PG-array consisting of 384 SNPs from 222 genes). It is known that the prevalence of the poor metabolizer (PM) phenotype in Caucasians ranges from 2 to 5% and that the most common gene defect is the CYP2C19*2 variant, which accounts for 75-83% of the defective alleles in PMs. Single CYP2C19*2 carrier frequency in the Puerto Rican newborn tested was 24.6%. Although the HW goodness-of-fit χ²-test was the only test applied, no major deviations from HWE were claimed (χ² = 0.28; p = 0.87). Statistical analysis showed that no significant differences in genotypic frequencies were found between our sample and those reported for Caucasians where 2% are PMs and 0.75% harboring the *2 allele (χ² = 0.723; p = 0.70). We also examined 72 of these samples for the CYP2C19*3 mutant allele (null metabolizer), but we could not detect any carriers or individual homozygous for this mutation, which suggests that the frequency for the CYP2C19*3 allele is lower than that for the CYP2C19*2 allele in the Puerto Rican population. Be advised that this mutation is usually considered rare in non-Asian descendant population (7-8). A larger number of samples need to be genotyped for these two allelic variants in order to validate our findings in Puerto Ricans.

Little pharmacogenetic-guided recommendations of clinical relevance are available due in part to limited information in target populations. Accordingly, there is a need to start translating pharmacogenetic data into specific clinical recommendations for labelling and optimal dosage calculation in each individual. Considering the observed prevalence of clinically relevant CYP2C19*2 polymorphisms in Puerto Ricans, it is necessary to educate pharmacists and healthcare providers in CYP2C19-related pharmacogenetics in order to achieve better outcomes by developing DNA-guided individual (personalized) CYP2C19 substrates dosing regimens.

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References