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ORIGINAL RESEARCH

The Diversity of CYP2C19 Polymorphisms in the Thai Population: Implications for Precision Medicine

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Introduction: *CYP2C19* plays a major role in the metabolism of various drugs. The most common genetic variants were the *CYP2C19*2* and **3* alleles (*rs4244285* and *rs4986893*, non-functional variants). In previous studies, we found that genetic polymorphisms in *CYP2C19* variants influenced the active metabolites of clopidogrel and caused major adverse cardiovascular and cerebrovascular effects. However, the distribution of *CYP2C19* varies among ethnic groups and according to adverse drug reactions. This study aimed to investigate the frequency of *CYP2C19* genetic polymorphisms in the Thai population and analyze the differences in the frequency of *CYP2C19* genetic polymorphisms between Thai and other populations.

Methods: This study enrolled 211 unrelated healthy Thai individuals in total. We performed a real-time polymerase chain reaction to genotype CYP2C19*2 (681G > A) and CYP2C19*3 (636G > A).

Results: In the Thai population, the *CYP2C19*1* allele was the most prevalent at 70.14%, while the *CYP2C19*2* and *3 alleles were found at frequencies of 25.36% and 4.50%, respectively. Conversely, the *CYP2C19*3* allele was not detected in Caucasian, Hispanic, African, Italian, Macedonian, Tanzanian, or North Indian populations. The phenotypic profile of this gene revealed that the frequency of intermediate metabolizers (IMs) is nearly equal to that of extensive metabolizers (EMs), at 42.65% and 48.82% respectively, with genotypes *1/*2 (36.02%) and *1/*3 (6.63%). Likewise, poor metabolizers (PMs) with genotypes *2/*2 (6.16%), *2/*3 (2.37%), and *3/*3 (<1%) are more prevalent in our population as well.

Conclusion: The distribution of *CYP2C19* genotype and phenotype influenced by non-functional alleles has potential as a pharmacogenomics biomarker for precision medicine and is dependent on an ethnic-specific genetic variation database.

Introduction

Cytochrome P450 (CYP) proteins form a superfamily of enzymes that are involved in the metabolism of drugs, fatty acids, steroids, and xenobiotics.^{1,2} Research has demonstrated that the metabolic activity of CYP enzymes is influenced by genetic polymorphisms.^{3,4} These polymorphisms in drug-metabolizing enzyme genes contribute to variations in pharmacological responses and the risk of adverse drug events among individuals and across different ethnic groups.⁵ The human *CYP2C* subfamily includes four members: *CYP2C8, CYP2C9, CYP2C18*, and *CYP2C19*.⁶

The *CYP2C19* gene is located on chromosome 10 (10q24.1-q24.3) and is responsible for approximately 10% of drug metabolism in clinical practice.^{7,8} *CYP2C19* plays a crucial role in metabolizing various therapeutic drugs, including clopidogrel, phenytoin, omeprazole, proguanil, diazepam, citalopram, imipramine, amitriptyline, and clomipramine.⁹ The most common genetic variant of *CYP2C19* is the *CYP2C19*2* allele (*rs4244285*, c. G681A), a single base pair mutation in exon 5 that results in a splicing defect, impairing enzyme function.¹⁰ Another significant variant, *CYP2C19*3* (*rs4986893*), is a G636A mutation in exon 4, producing a premature stop codon.¹¹ *CYP2C19* genotypes and phenotypes are categorized as follows: *1/*1 (extensive metabolizers, EMs, two functional alleles); *1/*2 and *1/*3 (intermediate metabolizers, IM, one null allele and one functional allele); and *2/*2, *2/*3, and *3/*3 (poor metabolizers, PM, two non-functional alleles).^{12–14} Numerous studies have shown a correlation between *CYP2C19* polymorphisms and enzyme activity in patients treated with relevant drugs.^{15–17}

Clopidogrel is commonly used to treat myocardial infarction (MI), stroke, acute coronary syndrome (ACS), and atherosclerotic vascular disease.¹⁸ Notably, CYP2C19 plays a crucial role in converting clopidogrel into its active metabolite, clopi-H4.¹⁹ This active metabolite inhibits the adenosine diphosphate P2Y12 receptor, thereby reducing platelet activation. According to the Clinical Pharmacogenetics Implementation Consortium (CPIC) guidelines for clopidogrel, alleles associated with the *CYP2C19* poor metabolizers phenotype (*2/*2, *2/*3, and *3/*3) are significantly linked to lower levels of active clopidogrel metabolites and an increased risk of major adverse cardiovascular and cerebrovascular events.²⁰ Furthermore, the *CYP2C19* intermediate metabolizers (IMs) phenotype also reduces the effectiveness of clopidogrel.

The distribution of poor metabolizers (PMs) with genotypes *2/*2, *2/*3, and *3/*3 vary among different populations. The highest prevalence is observed in East Asians (12.97%) and Asians (8.15%), followed by African Americans (4.05%) and Europeans (2.38%). A similar pattern is seen with intermediate metabolizers (IMs), with East Asians having the highest frequency (45.92%), followed by Asians (40.80%), African-Americans (31.39%), and Europeans (26.10%).²¹ These genetic variations in *CYP2C19* across different ethnicities are crucial for understanding individual differences in treatment effectiveness and safety, especially in the context of pharmacogenomics-based dosing guidelines. Consequently, this study aimed to investigate the prevalence of *CYP2C19* genetic variations in the Thai population and compare them with those in other populations.

Materials and Methods

Thai Subjects

This cross-sectional study recruited 211 unrelated, healthy individuals of Thai ethnicity, with clinical data obtained from the Excellence Pharmacogenomics and Precision Medicine Centre, College of Pharmacy, Rangsit University, between November 2021 and February 2023 (<u>Supplement 1</u>). All participants were native Thais who were lived in the location of Thailand and had no history of adverse drug reactions (ADRs). We used self-identified race/ethnicity (SIRE) method to confirm the ethnicity, specifically three generations were confirmed as Thais. Thailand is centrally located in Mainland Southeast Asia, bordered by Myanmar to the west, Laos to the northeast, Cambodia to the east, and Malaysia to the south. The study received approval from the Ethics Review Board of Rangsit University (COA. No. RSUERB2021-104), and according to the Declaration of Helsinki with a written informed consent was obtained from all participants.

CYP2C19 Genotyping

Genomic DNA was extracted from EDTA whole blood using a Geneaid DNA Isolation Kit (Geneaid Biotech Ltd., New Taipei City, Taiwan). The quality and quantity of the genomic DNA were measured using an ultraviolet-visible

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spectrophotometer (Zhengzhou, Henan, China). Genotyping of CYP2C19*2 (681G > A, rs4244285, non-functional variant) and CYP2C19*3 (636G > A, rs4986893, non-functional variant) was performed using TaqMan assays (C_30634128_10 and C_27861809_10 respectively; ABI, Foster City, CA, USA). The PCR cycling conditions included 50 cycles of denaturation at 92°C for 15 seconds, and annealing and extension at 60°C for 1.30 minutes. *CYP2C19* variants were identified using a QIAGEN QIAquant 96 real-time PCR system (Qiagen, Hilden, Germany).

CYP2C19 Phenotype

According to the Clinical Pharmacogenetics Implementation Consortium (CPIC) Guideline, *CYP2C19*1/*1* is categorized as an extensive metabolizers (EMs) with two normal-function alleles. Intermediate metabolizers (IMs) have one normal and one non-functional allele (*CYP2C19*1/*2* and *CYP2C19*1/*3*). Poor metabolizers (PMs) carry two nonfunctional alleles (*CYP2C19*2/*2*, *CYP2C19*2/*3*, and *CYP2C19*3/*3*).

Statistical Analysis

The frequencies of the two common *CYP2C19* variants were analyzed using the Arlequin program version 3.1 for Hardy–Weinberg equilibrium testing. Allele and phenotype frequencies of *CYP2C19* in the Thai population were compared with those in other populations using R Statistical Software (v4.1.2; R Core Team 2021). Odds ratios for all comparisons were illustrated in a forest plot using the Meta for R package (v4.2.0; Viechtbauer 2010). A significant difference was identified for each population compared with the Thai population if the p-value was less than 0.05.

Results

The Frequency of CYP2C19 Alleles in Thai Population

A total of 211 unrelated healthy Thai participants were included in this study, comprising fifty-five males (26.07%) and one hundred fifty-six females (73.93%). The mean age of the participants was 38.3 years, ranging from 19 to 75 years. We determined the allele frequency distribution of *CYP2C19* in the Thai population (Table 1). The *CYP2C19*1* allele was the most common, with a frequency of 70.14%. The frequencies of the *CYP2C19*2* and *CYP2C19*3* alleles were 25.36% (n = 107) and 4.50% (n = 19), respectively. Among the healthy Thai participants, the *CYP2C19* genotypes were distributed as follows: 103 (48.82%) were extensive metabolizers (EMs, *CYP2C19*1/*1*), 76 (36.02%) were intermediate metabolizers (IMs, *1/*2 and *1/*3), and 14 (6.63%), 13 (6.16%), and 5 (2.37%) were poor metabolizers (PMs, *2/*2 and *2/*3). The frequency of the *CYP2C19*3/*3* (PMs) genotype was less than 1.00% of the total Thai population. The allele frequencies of *CYP2C19*2* and *3 were tested for Hardy–Weinberg equilibrium (p-value < 0.05), showing no significant deviation in the Thai population.

CYP2C19 Alleles (n = 422)			Allele Frequencies, %
CYP2C19*1			70.14
CYP2C19*2, c.G681A			25.36
CYP2C19*3 c.C806T			4.50
CYP2C19 genotyping (n = 211)	Predicted phenotype	n	Genotype frequencies, %
CYP2C19*1/*1	EMs	103	48.82
CYP2C19*1/*2	IMs	76	36.02
CYP2C19*1/*3	IMs	14	6.63
CYP2C19*2/*2	PMs	13	6.16
CYP2C19*2/*3	PMs	5	2.37
CYP2C19*3/*3	PMs	0	0.00

Table I	Alleles and Genotyping	Frequencies for (CYP2C19 Genes in the	Thai Population (n = 211)
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Abbreviations: EMs, Extensive metabolizers; IMs, intermediate metabolizers; PMs, poor metabolizers.

Comparative of CYP2C19 Genotype and Phenotype Frequency Between the Thai and Other Populations

The frequencies of the *CYP2C19*2* and *3 alleles were evaluated in Thai and various other populations, as shown in Tables 2 and 3 and Figure 1A and B. This study found that the *CYP2C19*2* allele was particularly frequent. The highest frequencies of the *CYP2C19*2* allele were observed in Indians (37.90%),³⁰ Native Japanese (34.50%),³³ and Bhutanese (30.14%),³⁴ followed by North Indians (29.75%),²⁹ Koreans (28.40%),³³ Thais (25.36%), Han Chinese (24.67%),³¹

Ethnicity	Allele Frequency, %	OR (95% CI)	p-value	References
Thai (n = 211)	25.36	Present study		
Italian (n = 360)	11.10	0.36 (0.23-0.58)	1.21x10 ⁻⁵	[22]
Macedonia (n = 184)	14.40	0.47 (0.27-0.82)	0.0056	[23]
Caucasian (n = 216)	16.20	0.56 (0.34-0.93)	0.0176	[24]
Hispanic (n =500)	12.80	0.43 (0.28-0.66)	6.04x10 ⁻⁵	[25]
African (n = 99)	20.20	0.74 (0.39–1.35)	0.3208	[26]
African-American (n = 500)	19.40	0.70 (0.47-1.05)	0.0711	[25]
Venezuelan (n = 281)	19.70	0.71 (0.45–1.11)	0.1249	[27]
Tanzanian (n = 212)	17.90	0.64 (0.39–1.04)	0.0600	[28]
North Indians (n = 121)	29.75	1.23 (0.72-2.08)	0.4426	[29]
Indian (n = 112)	37.90	1.74 (1.03–2.93)	0.0298	[30]
Han Chinese (n = 96)	24.67	0.97 (0.53-1.740)	1.0000	[31]
Native Japanese (n = 200)	34.50	1.53 (0.98–2.39)	0.0529	[32]
Korean (n = 271)	28.40	1.15 (0.75–1.77)	0.5362	[33]
Bhutanese (n = 443)	30.14	1.26 (0.86-1.86)	0.2307	[34]
Vietnamese (n = 275)	20.50	0.74 (0.47–1.17)	0.1899	[35]
Malaysian (n = 62)	5.70	0.20 (0.05-0.58)	0.0007	[36]

Table 2 The Allele Frequencies of CYP2C19*2 in Many Populations

Notes: Bold font indicates significantly different (p-value < 0.05), allele frequency compared between different ethnicities and Thais. Abbreviations: OR, odds ratio; 95% CI, 95% Confidence Interval.

Ethnicity	Allele Frequency, %	OR (95% CI)	p-value	References
Thai (n = 211)	4.50	Present study		
Italian (n = 360)	0	0 (0.00-0.29)	0.0001	[22]
Macedonia (n = 184)	0	0 (0.00–0.57)	0.0042	[23]
Caucasian (n = 216)	0	0 (0.00-0.48)	0.0016	[24]
Hispanic (n =500)	0	0 (0.00-0.21)	1.58×10^{-5}	[25]
African (n = 99)	0	0 (0.00-1.06)	0.0620	[26]
African-American (n = 500)	0.40	0.09 (0.01-0.44)	0.0005	[25]
Venezuelan (n = 281)	1.26	0.32 (0.07-1.18)	0.0847	[27]
Tanzanian (n = 212)	0	0 (0.00-0.49)	0.0018	[28]
North Indians (n = 121)	0	0 (0.00–0.87)	0.0291	[29]
Indian (n = 112)	2.20	0.41 (0.04–2.03)	0.3412	[30]
Han Chinese (n = 96)	3.27	0.72 (0.12-2.99)	0.7595	[31]
Native Japanese (n = 200)	9.00	2.22 (0.92-5.75)	0.0718	[32]
Korean (n = 271)	10.10	2.48 (1.10-6.13)	0.0224	[33]
Bhutanese (n = 443)	15.69	4.20 (2.04–9.78)	8.94 x 10 ⁻⁶	[34]
Vietnamese (n = 275)	2.50	0.59 (0.18–1.80)	0.3151	[35]
Malaysian (n = 62)	6.50	1.55 (0.34–5.79)	0.5002	[36]

Table 3 The Allele Frequencies of CYP2C19*3 in Many Populations

Notes: Bold font indicates significantly different (p-value < 0.05), allele frequency compared between different groups ethnicities and Thais.

Abbreviations: OR, odds ratio; 95% Cl, 95% Confidence Interval.

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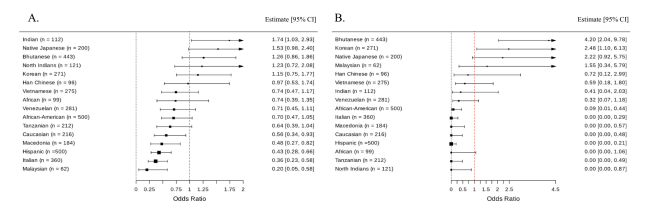


Figure I Forest plot displaying the odds ratios for the frequencies of CYP2C19*2 (A) and CYP2C19*3 (B) alleles across multiple populations in comparison to the Thai population allele frequency.

Vietnamese (20.50%),³⁵ and Africans (20.20%)²⁶ (Table 2 and Figure 1A). Conversely, the Bhutanese population had the highest observed frequency of the *CYP2C19*3* allele (15.69%)³⁴ (Table 3 and Figure 1B). High distributions of the *CYP2C19*3* allele were also observed in Koreans (10.10%),³³ Native Japanese (9.00%),³² Malaysians (6.50%),³⁶ and Thais (4.50%), while the *CYP2C19*3* allele was not found in Caucasian, Hispanic, African, Italian, Macedonian, Tanzanian, and North Indian populations.^{22–26,28,29}

A comparison of *CYP2C19*2* and *3 frequencies between the Thai population and other populations is presented in Tables 2 and 3 and Figure 1A and B. The distribution of *CYP2C19* variations was similar among the Thai, Han Chinese, Vietnamese, and Venezuelan populations (p-value > 0.05). However, the *CYP2C19*2* allele showed significant differences between the Thai population and others, such as Italian (p-value = 1.21×10^{-5}), Macedonian (p-value = 0.0056), Caucasian (p-value = 0.0176), Hispanic (p = 6.04×10^{-5}), Indian (p-value = 0.0298), and Malaysian populations (p-value = 0.0007). A strong significant difference was also observed for *CYP2C19*3* frequencies when comparing Thai with Bhutanese and Hispanic populations (p-value = 8.94×10^{-6} and 1.58×10^{-5} , respectively).

Phenotyping of *CYP2C19*1/*1* (extensive metabolizers, EMs) was highly prevalent in several populations (Table 4 and Figures 2 and 3). The highest frequencies of *CYP2C19*1/*1* were observed in the Italian population (79.44%) compared to the Thai population (p-value = 8.22×10^{-14}).²² Intermediate metabolizers (*CYP2C19 *1/*2* and **1/*3*) showed the following phenotype distribution: Italian (18.89%), Macedonian (19.02%), Caucasian (18.52%), Hispanic (18.00%), African (19.19%), African-American (23.60%), and Malaysian (14.52%) populations.^{22–26,36} Significant differences in IMs prevalence were observed when compared with Thais (p-value < 0.05), with the highest frequency detected in the Bhutanese population (79.01%; OR = 5.05, 95% CI = 3.49–7.34, p-value = 7.83×10^{-20}).³⁴

The distribution of poor metabolizers (PMs) of *CYP2C19* (*2/*2, *2/*3, and *3/*3) was 19.00% in Native Japanese,³² 14.02% in Koreans,³³ 10.42% in Han Chinese,³¹ 9.82% in Indians,³⁰ 8.53% in Thais, 7.44% in North Indians,²⁹ 6.32% in Bhutanese,³⁴ 6.06% in Africans,²⁶ 4.84% in Malaysians,³⁶ 4.63% in Venezuelans,²⁷ 4.40% in African-Americans,²⁵ 4.17% in Caucasians,²⁴ 4.00% in Vietnamese,³⁵ 3.77% in Tanzanians,²⁸ 2.72% in Macedonians,²³ 1.67% in Italians,²² and 1.60% in Hispanics.²⁵ The PMs prevalence was significantly higher in Thai patients compared to Italian (p-value = 0.0001), Macedonian (p-value = 0.0168), and Hispanic (p-value = 0.0006) patients. Conversely, the PMs prevalence in the Thai population was lower than in the Native Japanese population (OR = 2.51, 95% CI = 1.34–4.86, p-value = 0.0024). The odds ratios for these comparisons are illustrated in Figure 2A and B using forest plots.

Discussion

According to the Clinical Pharmacogenetics Implementation Consortium (CPIC) guidelines, the *CYP2C19* gene is a significant pharmacogenomic polymorphism in various populations.^{20,37} In clinical settings, *CYP2C19* plays a pivotal role in metabolizing approximately 8–10% of commonly used medications, including proton pump inhibitors, anticonvulsants, antiplatelet agents, and antidepressants.^{14,24} Notably, patients carrying homozygous alleles (PMs) and

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Ethnicity	Phenotyping Frequencies, n (%)	OR (95% CI)	p-value	References
EMs (*1/*1)				
Thai (n = 211)	103 (48.82)	Present study		
Italian (n = 360)	286 (79.44)	4.04 (2.75-5.98)	8.22 x 10 ⁻¹⁴	[22]
Macedonia (n = 184)	77 (41.85)	0.76 (0.49–1.15)	0.1882	[23]
Caucasian (n = 216)	89 (41.20)	0.74 (0.49–1.09)	0.1204	[24]
Hispanic (n = 250)	124 (49.60)	1.03 (0.70–1.51)	0.9255	[25]
African (n = 99)	33 (33.33)	0.53 (0.31–0.89)	0.0139	[26]
African-American (n = 250)	88 (35.20)	0.57 (0.38–0.84)	0.0033	[25]
Venezuelan (n = 281)	177 (62.99)	1.78 (1.22–2.61)	0.0018	[27]
Tanzanian (n = 106)	72 (67.92)	2.21 (1.3 3–3.75)	0.0013	[28]
North Indians (n = 121)	58 (47.93)	0.97 (0.60–1.55)	0.9094	[29]
Indian (n = 112)	33 (29.46)	0.44 (0.26–0.73)	0.0009	[30]
Han Chinese (n = 96)	50 (52.08)	1.14 (0.68–1.90)	0.6238	[31]
Native Japanese (n = 200)	62 (31.00)	0.47 (0.31–0.72)	0.0003	[32]
Korean (n = 271)	97 (35.79)	0.59 (0.39-0.86)	0.0051	[33]
Bhutanese (n = 443)	254 (57.34)	1.41 (1.00–1.99)	0.0439	[34]
Vietnamese (n = 275)	160 (58.18)	1.46 (1.00-2.13)	0.0436	[35]
Malaysian (n = 62)	41 (66.13)	2.04 (1.09–3.89)	0.0203	[36]
IMs (*1/*2 and *1/*3)				1
Thai (n = 211)	90 (42.65)	Present study		
Italian (n = 360)	68 (18.89)	0.31 (0.21-0.47)	2.17 x 10 ⁻⁹	[22]
Macedonia (n = 184)	35 (19.02)	0.32 (0.19-0.51)	4.96 x 10 ⁻⁷	[23]
Caucasian (n = 216)	40 (18.52)	0.31 (0.19-0.48)	6.37 x 10 ⁻⁸	[24]
Hispanic (n =250)	45 (18.00)	0.29 (0.19-0.46)	7.04 x 10 ⁻⁹	[25]
African (n = 99)	19 (19.19)	0.32 (0.17-0.58)	4.25 x 10 ⁻⁵	[26]
African-American (n = 250)	59 (23.60)	0.42 (0.27-0.63)	1.61 x 10 ⁻⁵	[25]
Venezuelan (n = 281)	90 (32.03)	0.63 (0.43-0.93)	0.0180	[27]
Tanzanian (n = 106)	30 (28.30)	0.53 (0.31-0.90)	0.0142	[28]
North Indians (n = 121)	54 (44.63)	1.08 (0.67–1.74)	0.7315	[29]
Indian (n = 112)	68 (60.71)	2.07 (1.27–3.41)	0.0023	[30]
Han Chinese (n = 96)	32 (33.33)	0.67 (0.39–1.14)	0.1325	[31]
Native Japanese (n = 200)	98 (49.00)	1.29 (0.86–1.94)	0.1998	[32]
Korean (n = 271)	128 (47.23)	1.20 (0.82–1.76)	0.3564	[33]
Bhutanese (n = 443)	350 (79.01)	5.05 (3.49-7.34)	7.83 x 10 ⁻²⁰	[34]
Vietnamese (n = 275)	99 (36.00)	0.76 (0.52–1.11)	0.1591	[35]
Malaysian (n = 62)	9 (14.52)	0.23 (0.09–0.50)	3.81 x 10 ⁻⁵	[36]
PMs (*2/*2, *2/*3 and *3/*3	3)			
Thai (n = 211)	18 (8.53)	Present study		
Italian (n = 360)	6 (1.67)	0.18 (0.06-0.49)	0.0001	[22]
Macedonia (n = 184)	5 (2.72)	0.30 (0.08–0.86)	0.0168	[23]
Caucasian (n = 216)	9 (4.17)	0.47 (0.18–1.13)	0.0745	[24]
Hispanic (n =250)	4 (1.60)	0.17 (0.04–0.54)	0.0006	[25]
• • •		0.69 (0.22–1.89)	0.5038	[26]
African (n = 99)	0 (0.00)	0.07 (0.22-1.07)	0.5050	20
African (n = 99) African-American (n = 250)	6 (6.06) I I (4.40)	0.49 (0.21–1.14)	0.0833	[25]

Table 4 Phenotyping Frequencies of CYP2C19*2 and *3 Polymorphism Among Different EthnicitiesCompared with Thai Population

(Continued)

Ethnicity	Phenotyping Frequencies, n (%)	OR (95% CI)	p-value	References
Tanzanian (n = 106)	4 (3.77)	0.42 (0.10–1.33)	0.1595	[28]
North Indians (n = 121)	9 (7.44)	0.86 (0.33-2.10)	0.8361	[29]
Indian (n = 112)	(9.82)	1.17 (0.48–2.73)	0.6876	[30]
Han Chinese (n = 96)	10 (10.42)	1.25 (0.49–2.99)	0.6695	[31]
Native Japanese (n = 200)	38 (19.00)	2.51 (1.34–4.86)	0.0024	[32]
Korean (n = 271)	38 (14.02)	1.75 (0.94–3.36)	0.0642	[33]
Bhutanese (n = 443)	28 (6.32)	0.72 (0.38-1.43)	0.3274	[34]
Vietnamese (n = 275)	(4.00)	0.45 (0.19–1.03)	0.0518	[35]
Malaysian (n = 62)	3 (4.84)	0.55 (0.09–1.97)	0.4260	[36]

Table 4 (Continued).

Notes: Bold font indicates significantly different (p-value < 0.05), phenotyping frequencies compared between different ethnicities and Thais.

Abbreviations: OR, odds ratio; 95% Cl, 95% Confidence Interval.

heterozygous alleles (IMs) of *CYP2C19*2* and **3* variants are reported to have approximately 1.8-fold and 1.5-fold increased risk, respectively, of developing severe adverse cardiovascular events (e.g.death, myocardial infarction, stroke) following clopidogrel treatment.^{38,39} Thus, non-functional *CYP2C19* variants significantly impact the dosage requirements for clopidogrel therapy in individual patients.

In our study, the most prevalent non-functional variant was the *CYP2C19*2* allele, found in approximately 25.36% of our sample, consistent with previous reports in various populations, including Asian populations (Han Chinese, Native Japanese, Korean, Bhutanese, Vietnamese, and North Indians), Africans, African-Americans, Venezuelans, and Tanzanians.^{25–28,31–36} In contrast, Caucasian, Hispanic, Macedonian, and Italian populations exhibited lower frequencies of the *CYP2C19*2* allele compared to the Thai population.^{22–25} However, these variations were attributed to the distribution of non-functional *CYP2C19* alleles in each population and their impact on clopidogrel efficacy. Further investigations are necessary to confirm the influence of *CYP2C19*2* distribution across ethnicities and the effects on clinically relevant medications.

The allele frequency of *CYP2C19*3* (c C806T) in our study was approximately 4.50%, similar to earlier reports in the Thai population.⁴⁰ Additionally, the frequency of *CYP2C19*3* in Han Chinese was 3.27%, consistent with other Asian populations, as shown in Table 3 and Figure 1B. Interestingly, the distribution of *CYP2C19*3* in these Asian populations, including Thai, closely resembled each other, except for Bhutanese, Korean, and North Indian populations. Specifically, we observed a higher distribution of *CYP2C19*3* in the Bhutanese population compared to the Thai population (p-value = 8.94×10^{-6}). However, the

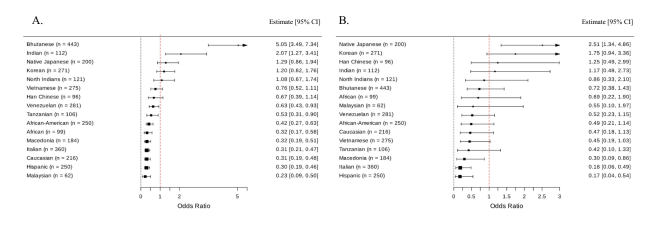


Figure 2 Forest plot displaying the odds ratios for the frequencies of CYP2C19 intermediate metabolizers (IMs; *1/2 + *1/3) diplotypes (**A**) and CYP2C19 poor metabolizers (PMs; *2/*2 + *2/*3 + *3/*3) diplotypes (**B**) across multiple populations in comparison to the Thai population diplotype frequency.

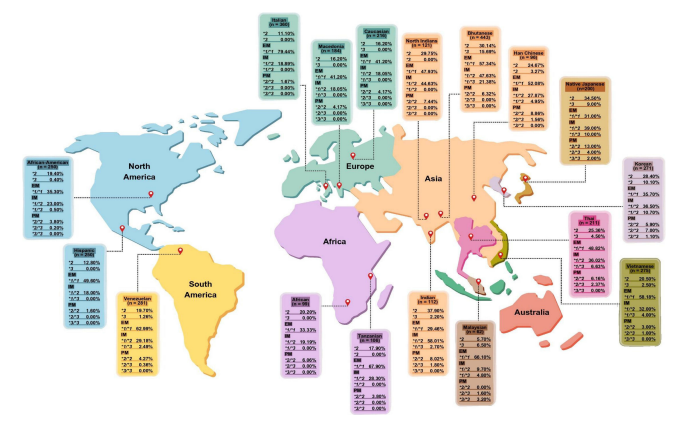


Figure 3 Distribution of CYP2C19 polymorphisms in Thais and ethnic groups.

prevalence of the *CYP2C19*3* allele was absent in Italian, Macedonian, Caucasian, Hispanic, African, Tanzanian, and North Indian populations.^{22–26,28,29} This genetic variability across racial and ethnic groups underscores the importance of *CYP2C19*3* allele frequency as a pharmacogenomics marker for screening different ethnicities to prevent therapeutic failures in individual patients, particularly in the Asian population.

None of the Thai participants in our study carried the homozygous CYP2C19*3/*3 allele. However, PMs phenotypes based on CYP2C19*2/*2 and *2/*3 genotypes were present in approximately 6.16% and 2.37% of the sample, respectively. Our findings align with previous research indicating a PMs distribution of approximately 19% in Asian populations compared to around 2% in Caucasians.⁴¹ Notably, substantial variation in the distribution of these CYP2C19 genotypes was observed across populations, with less than 1% occurrence of these variants (*1/*3, *2/*3, and *3/*3) in European and African American populations, contrasting with frequencies of *1/*3 (7.34%), *2/*3 (3.33%), and *3/*3 (0.44%) in East Asian populations.⁴² Given the genetic diversity of CYP2C19 variants across ethnicities, determining the CYP2C19*3 genotype and phenotype before treatment initiation is particularly advantageous for Asian populations. Moreover, the understanding of CYP2C19 polymorphisms can significantly impact patient prognosis and outcomes, particularly in the context of drugs like clopidogrel, where genetic variations affect drug metabolism and efficacy. However, the relationship between CYP2C19 phenotype frequency and drug efficacy or adverse drug reactions (ADRs) warrants further investigation in other populations.

Conversely, the *CYP2C19*17* variant (c. -806C>T) demonstrates ultra-rapid *CYP2C19* activity and holds a significant role in the metabolic pathways of therapeutic drugs, which could heighten the risk of treatment failure with this substrate. From a clinical standpoint, the *CYP2C19*17* variant strongly correlates with an increased risk of clopidogrel-induced bleeding in patients with cardiovascular and cerebrovascular diseases (OR = 1.89, 95% CI: 1.09– 3.25, p-value = 0.02). This association is attributed to the heightened transcriptional activity of the CYP2C19 enzyme and its exaggerated response to clopidogrel's prodrug.⁴³ Conversely, this variant exhibits a notable association with a reduced risk of major adverse cardiovascular and cerebrovascular events in patients with coronary artery disease (OR = 0.76, 95%

CI: 0.60–0.98, p = 0.03).⁴³ Furthermore, the *CYP2C19*17* genotype has been observed at higher frequencies among European and African populations, approximately 15.0–25.0%, whereas the Asian population exhibits a much lower frequency, approximately 4%.⁴⁴ Given these trends, further exploration of the frequency of *CYP2C19*17* variants and their association with adverse drug reactions is warranted across various populations.

One limitation of our study stems from the genotyping method employed for *CYP2C19*. We utilized real-time PCR to detect two single nucleotide polymorphisms (SNPs), *rs4244285* (*2) and *rs4986893* (*3). In cases where individuals did not carry these two variants, we presumed the genotype to be *1. However, this assumption may not accurately represent the true *1 status, as other untested variants could be present. This methodological approach has the potential to affect the precision of our genotyping Results and subsequent interpretations of *CYP2C19* metabolic phenotypes. While real-time PCR is a highly reliable technique, the importance of validation cannot be overstated. To mitigate this concern, future studies should consider validating genotyping results using DNA sequencing methods, which offer a more comprehensive analysis of *CYP2C19* variants and minimize potential errors. By integrating such validation steps, we can enhance the accuracy of genotyping and ensure a more dependable identification of metabolic phenotypes, thereby enhancing the relevance of our findings in clinical practice. In future research, we plan to include clinical data to investigate how these genetic variations impact drug response and adverse reactions in real-world scenarios.

In conclusion, the distribution of the *CYP2C19* gene via non-functional alleles holds promise for implementation in clinical practice and relies on an ethnic-specific genetic variation database. Specifically, the genotype and phenotype of *CYP2C19*3* could serve as a pharmacogenomics biomarker in Thai and Asian populations for screening, mitigating genetic associations with adverse drug reactions induced by certain medications.

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Disclosure

The authors have no relevant affiliations or financial involvement with any organization or entity with a financial interest or financial conflict with the subject matter or materials discussed in this manuscript.

References

- 1. Reichhart DW, Feyereisen R. Cytochromes P450: a success story. Genome Biol. 2000;1(6):1-7.
- 2. Sim SC, Kacevska M, Ingelman-Sundberg M. Pharmacogenomics of drug-metabolizing enzymes: a recent update on clinical implications and endogenous effects. *Pharmacogenomics J.* 2013;13(1):1–11. doi:10.1038/tpj.2012.45
- 3. Zhou SF, Liu JP, Chowbay B. Polymorphism of human cytochrome P450 enzymes and its clinical impact. Drug Metab Rev. 2009;41(2):89–295. doi:10.1080/03602530902843483
- Trenaman SC, Bowles SK, Andrew MK, Goralski K. The role of sex, age and genetic polymorphisms of CYP enzymes on the pharmacokinetics of anticholinergic drugs. *Pharmacol Res Perspect*. 2021;9(3):1–33. doi:10.1002/prp2.775
- Zhou L, Sharma P, Yeo KR, et al. Assessing pharmacokinetic differences in Caucasian and East Asian (Japanese, Chinese and Korean) populations driven by CYP2C19 polymorphism using physiologically-based pharmacokinetic modelling. *Eur J Pharm Sci.* 2019;139:1–10.105061. doi:10.1016/j.ejps.2019.105061
- 6. Goldstein JA. Clinical relevance of genetic polymorphisms in the human CYP2C subfamily. Br J Clin Pharmacol. 2001;52(4):349–355. doi:10.1046/j.0306-5251.2001.01499.x
- 7. Wan-Po AL, Girard T, Farndon P, Cooley C, Lithgow J. Pharmacogenetics of CYP2C19: functional and clinical implications of a new variant CYP2C19*17. *Br J Clin Pharmacol*. 2010;69(3):222–230. doi:10.1111/j.1365-2125.2009.03578.x
- Ellithi M, Baye J, Wilke RA. CYP2C19 genotype-guided antiplatelet therapy: promises and pitfalls. *Pharmacogenomics*. 2020;21(12):889–897. doi:10.2217/pgs-2020-0046
- 9. Lo C, Nguyen S, Yang C, et al. Pharmacogenomics in Asian subpopulations and impacts on commonly prescribed medications. *Clin Transl Sci.* 2020;13(5):861–870. doi:10.1111/cts.12771
- 10. Santos PC, Soares RA, Santos DB, et al. CYP2C19 and ABCB1 gene polymorphisms are differently distributed according to ethnicity in the Brazilian general population. *BMC Med Genet*. 2011;12:13.1–7. doi:10.1186/1471-2350-12-13

- 11. Ghodke Y, Joshi K, Arya Y, et al. Genetic polymorphism of CYP2C19 in Maharashtrian population. Eur J Epidemiol. 2007;22(12):907-915. doi:10.1007/s10654-007-9196-0
- 12. Sun H, Qu Q, Chen ZF, et al. Impact of CYP2C19 variants on clinical efficacy of clopidogrel and 1-year clinical outcomes in coronary heart patients undergoing percutaneous coronary intervention. *Front Pharmacol.* 2016;7:1–9. eCollection 2016. doi:10.3389/fphar.2016.00453
- Yadav AK, Chakkumkollath AK, Helna A, et al. Substantiation of a clopidogrel metabolism-associated gene (CYP2C19) variation among healthy individuals. *Indian Heart J.* 2023;S0019–4832(23). doi:10.1016/j.ihj.2023.05.005
- 14. Desta Z, Zhao X, Shin JG, Flockhart DA. Clinical significance of the cytochrome P450 2C19 genetic polymorphism. *Clin Pharmacokinet*. 2002;41 (12):913–958. doi:10.2165/00003088-200241120-00002
- 15. Bahar MA, Setiawan D, Hak E, Wilffert B. Pharmacogenetics of drug-drug interaction and drug-drug-gene interaction: a systematic review on CYP2C9, CYP2C19 and CYP2D6. *Pharmacogenomics*. 2017;18(7):701-739. doi:10.2217/pgs-2017-0194
- Chaudhry SR, Muhammad S, Eidens M, et al. Pharmacogenetic prediction of individual variability in drug response based on CYP2D6, CYP2C9 and CYP2C19 genetic polymorphisms. *Curr Drug Metab.* 2014;15(7):711–718. doi:10.2174/1389200215666141125121952
- 17. Lee SJ. Clinical Application of CYP2C19 pharmacogenetics toward more personalized medicine. Front Genet. 2012;3:1-7. doi:10.3389/ fgene.2012.00318
- Patti G, Micieli G, Cimminiello C, Bolognese L. The Role of Clopidogrel in 2020: a Reappraisal. Cardiovasc Ther. 2020;2020:1–12. doi:10.1155/ 2020/8703627
- Brown SA, Pereira N. Pharmacogenomic Impact of CYP2C19 Variation on Clopidogrel Therapy in Precision Cardiovascular Medicine. J Pers Med. 2018;8(1):1–31. doi:10.3390/jpm8010008
- 20. Lee CR, Luzum JA, Sangkuhl K, et al. Clinical Pharmacogenetics implementation consortium guideline for CYP2C19 genotype and clopidogrel therapy: 2022 update. *Clin Pharmacol Ther.* 2022;112(5):959–967. doi:10.1002/cpt.2526
- 21. CYP2C19 frequency table; Cited 29 Sept 2022. Available from: https://files.cpicpgx.org/data/report/current/frequency/CYP2C19_frequency_table. xlsx. Accessed June 27, 2024.
- 22. Serpe L, Canaparo R, Scordo MG, Spina E. Pharmacogenetics of drug-metabolizing enzymes in Italian populations. *Drug Metab Pers Ther.* 2015;30(2):107–120. doi:10.1515/dmdi-2014-0028
- Jakovski K, Nestorovska AK, Labacevski N, Dimovski AJ. Characterization of the most common CYP2C9 and CYP2C19 allelic variants in the population from the Republic of Macedonia. *Pharmazie*. 2013;68(11):893–898.
- 24. Vidović S, Škrbić R, Stojiljković MP, et al. Prevalence of five pharmacologically most important CYP2C9 and CYP2C19 allelic variants in the population from the Republic of Srpska in Bosnia and Herzegovina. *Arh Hig Rada Toksikol*. 2021;72(3):129–134. doi:10.2478/aiht-2021-72-3499
- 25. Martis S, Peter I, Hulot JS, Kornreich R, Desnick RJ, Scott SA. Multi-ethnic distribution of clinically relevant CYP2C genotypes and haplotypes. *Pharmacogenomics J.* 2013;13(4):369–377. doi:10.1038/tpj.2012.10
- 26. Santos PCJL, Soares RAG, Santos DBG, et al. CYP2C19 and ABCB1 gene polymorphisms are differently distributed according to ethnicity in the Brazilian general population. BMC Med Genet. 2011;12:1–7. doi:10.1186/1471-2350-12-13
- de Guerra DC, Flores S, Izaguirre MH. Distribution of CYP2C19*2 and CYP2C19*3 polymorphisms in Venezuelan populations with different admixture. Ann Hum Biol. 2013;40(2):197–200. doi:10.3109/03014460.2012.749946
- Dandara C, Masimirembwa CM, Magimba A, et al. Genetic polymorphism of CYP2D6 and CYP2C19 in east- and Southern African populations including psychiatric patients. Eur J Clin Pharmacol. 2001;57(1):11–17. doi:10.1007/s002280100282
- Lamba JK, Dhiman RK, Kohli KK. CYP2C19 genetic mutations in North Indians. Clin Pharmacol Ther. 2000;68(3):328–335. doi:10.1067/ mcp.2000.109365
- Adithan C, Gerard N, Vasu S, Rosemary J, Shashindran CH, Krishnamoorthy R. Allele and genotype frequency of CYP2C19 in a Tamilian population. Br J Clin Pharmacol. 2003;56(3):331–333. doi:10.1046/j.1365-2125.2003.01883.x
- Chen L, Qin S, Xie J, et al. Genetic polymorphism analysis of CYP2C19 in Chinese Han populations from different geographic areas of mainland China. *Pharmacogenomics*. 2008;9(6):691–702. doi:10.2217/14622416.9.6.691
- 32. Myrand SP, Sekiguchi K, Man MZ, et al. Pharmacokinetics/genotype associations for major cytochrome P450 enzymes in native and first- and third-generation Japanese populations: comparison with Korean, Chinese, and Caucasian populations. *Clin Pharmacol Ther.* 2008;84(3):347–361. doi:10.1038/sj.clpt.6100482
- 33. Kim KA, Song WK, Kim KR, Park JY. Assessment of CYP2C19 genetic polymorphisms in a Korean population using a simultaneous multiplex pyrosequencing method to simultaneously detect the CYP2C19*2, CYP2C19*3, and CYP2C19*17 alleles. J Clin Pharm Ther. 2010;35 (6):697–703. doi:10.1111/j.1365-2710.2009.01069.x
- 34. Dorji PW, Wangchuk S, Boonprasert K, Tarasuk M, Bangchang KN. Pharmacogenetic relevant polymorphisms of CYP2C9, CYP2C19, CYP2D6, and CYP3A5 in Bhutanese population. *Drug Metab Pers Ther.* 2019;34(4):1–14. doi:10.1515/dmpt-2019-0020
- 35. Vu NP, Thanh Nguyen HT, Bich Tran NT, et al. CYP2C19 genetic polymorphism in the Vietnamese population. Ann Hum Biol. 2019;46 (6):491–497. doi:10.1080/03014460.2019.1687750
- 36. Ang GY, Yu CY, Subramaniam V, et al. Detection of CYP2C19 genetic variants in Malaysian orang asli from massively parallel sequencing data. *PLoS One.* 2016;11(10):1–12. doi:10.1371/journal.pone.0164169
- 37. Galindo IF, Garro CC, Soares FR, et al. Interethnic variation of CYP2C19 alleles, 'predicted' phenotypes and 'measured' metabolic phenotypes across world populations. *Pharmacogenomics J.* 2016;16(2):113–123. doi:10.1038/tpj.2015.70
- 38. Mega JL, Simon T, Collet JP, et al. Reduced-function CYP2C19 genotype and risk of adverse clinical outcomes among patients treated with clopidogrel predominantly for PCI: a meta-analysis. *JAMA*. 2010;304(16):1821–1830. doi:10.1001/jama.2010.1543
- 39. Xi Z, Fang F, Wang J, AlHelal J, Zhou Y, Liu W. CYP2C19 genotype and adverse cardiovascular outcomes after stent implantation in clopidogrel-treated Asian populations: a systematic review and meta-analysis. *Platelets*. 2019;30(2):229–240. doi:10.1080/09537104.2017.1413178
- 40. Sukasem C, Tunthong R, Chamnanphon M, et al. CYP2C19 polymorphisms in the Thai population and the clinical response to clopidogrel in patients with atherothrombotic-risk factors. *Pharmgenomics Pers Med.* 2013;6:85–91. doi:10.2147/PGPM.S42332
- 41. Mizutani T. PM frequencies of major CYPs in Asians and Caucasians. Drug Metab Rev. 2003;35(2-3):99–106. doi:10.1081/dmr-120023681
- 42. Ionova Y, Ashenhurst J, Zhan J, et al. CYP2C19 allele frequencies in over 2.2 million direct-to-consumer genetics research participants and the potential implication for prescriptions in a large health system. *Clin Transl Sci.* 2020;13(6):1298–1306. doi:10.1111/cts.12830

 Huang B, Cui DJ, Ren Y, Han B, Yang DP, Zhao X. Effect of cytochrome P450 2C19*17 allelic variant on cardiovascular and cerebrovascular outcomes in clopidogrel-treated patients: a systematic review and meta-analysis. J Res Med Sci. 2017;22:1–10. doi:10.4103/jrms.JRMS_590_16

44. Nestorovska AK, Jakovski K, Naumovska Z, et al. Distribution of the most common genetic variants associated with a variable drug response in the population of the Republic of Macedonia. *Balkan J Med Genet*. 2014;17(2):5–14. doi:10.2478/bjmg-2014-0069

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