EFFECTS OF CYP2C19 AND P2Y12 GENE POLYMORPHISMS ON CLINICAL RESULTS OF PATIENTS USING CLOPIDOGREL AFTER ACUTE ISCHEMIC CEREBROVASCULAR DISEASE

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ABSTRACT

The CYP2C19 and P2Y12 gene polymorphisms are responsible for resistance to clopidogrel, known as drug unresponsiveness. In this study we researched the effect of gene polymorphism on clinical results of patients who began clopidogrel therapy after acute ischemic cerebrovascular disease.

The study included 51 patients. The patient group included patients who had begun prophylactic clopidogrel due to acute ischemic cerebrovascular disease in the last 2 years. All patients were monitored by the Neurology Outpatient Clinic at Çanakkale Onsekiz Mart University Research Hospital, Çanakkale, Turkey, and only those monitored for at least 1 year were included in the study.

When the *1, *2 and *3 alleles of the CYP2C19 gene polymorphism were evaluated, two patients were homozygotes for *2/*2, 13 patients were heterozygous for *1/*2 and 36 patients were homozygotes for the wild type *1/*1. No patient had the *3 allele. Three heterozygous patients, one for *2/*2 and two for *1/*2, stopped clopidogrel therapy due to repeated strokes and began taking warfarin. When evaluating P2Y12 52 (G>T) and 34 (C>T) polymorphisms, all alleles were of the wild type.

The CYP2C19 and P2Y12 gene polymorphisms may cause recurring strokes linked to insufficient response to treatment of ischemic cerebrovascular disease. In our patient group, three patients suffered repeated strokes and these patients had the CYP2C19*2 gene polymorphism. As a result, before medication use, genetic testing is important for human life, quality of life and economic burden.

Keywords: Ischemic cerebrovascular disease (ICVD); clopidogrel; CYP2C19 and P2Y12 gene polymorphisms.

INTRODUCTION

Clopidogrel is used as a prophylactic monotherapy to prevent the development of a new attack of ischemic cerebrovascular disease (ICVD) [1]. Stroke is a major cause of disability and death, and affects nearly 16 million people globally each year [2]. As ICVD cause a high rate of morbidity and mortality, the importance of effective prophylactic treatment increases.

The clinical benefits of clopidogrel occur due to its inhibition of thrombocyte activation and aggregation [3,4]. Clopidogrel is actually a prodrug and transforms to active metabolite through CY2C19 enzyme in the cytochrome P450 (CYP450) enzyme family in the liver. As a result of CYP2C19 gene polymorphism, the transformation of clopidogrel to active metabolite reduces [5]. The CYP2C19 *1 allele
is a wild type allele linked to enzyme activity [6]. The most frequently seen variant causing reduced enzyme activity are the \textit{CYP2C19*2} allele and the less frequently seen \textit{CYP2C19*3} allele [7]. The \textit{*1/*1} wild type produces normal metabolism, heterozygous \textit{*1/*2} and \textit{*1/*3} produce mild metabolic disorders and \textit{*2/*2}, \textit{*2/*3} and \textit{*3/*3} produce severe metabolic disorders [8]. \textit{P2Y12} receptors are found in the platelet membranes and play a role in adenosine diphosphate (ADP)-induced platelet aggregation [9]. The aim of the active clopidogrel metabolite is the ADP platelet receptor \textit{P2Y12}, thus irreversibly inhibiting ADP linkage to platelets. As a result platelet activation and aggregation is inhibited [10,11]. The ADP-induced platelet aggregation varies between individuals [12]. The causes of these individual differences are genetic factors [13].

Patients with a \textit{P2Y12} gene polymorphism using clopidogrel with peripheral artery disease have been shown to develop more cerebral ischemic events than those without polymorphism [14]. Again, it has been reported that myocardial infarctus, stent thrombosis, and ischemic stroke development were higher in those with \textit{CYP2C19} gene polymorphisms compared to those without a polymorphism [15]. In acute coronary syndrome, stent patients with \textit{P2Y12} and \textit{CYP2C19} single gene polymorphisms have been shown to have negative clinical results linked to unresponsiveness to clopidogrel; however, patients with a combination of both polymorphisms had worse clinical results [16]. In our study, we planned to research the clinical effects of \textit{CYP2C19} and \textit{P2Y12} gene polymorphisms in our ICVD patients who take clopidogrel.

**PATIENTS AND METHODS**

**Patients.** The patient group comprised of patients who began prophylactic clopidogrel 75 mg/day as a result of acute ICVD in the previous 2 years. All patients were monitored by the Neurology Outpatient Clinic at Canakkale Onsekiz Mart University Research Hospital, Canakkale, Turkey. Those who had been monitored for at least 1 year were included in the study. Patients who stopped attending our clinic, or who did not take their medication regularly were not included in the study. The study was approved by the Institutional Review Board.

**Detection of the \textit{CYP2C19} Genotype.** Venous blood samples (2 mL) were collected from each patient in EDTA tubes. Genomic DNA was extracted from the whole blood using a high-purification preparation kit (Roche Diagnostics GmbH, Mannheim, Baden-Württemberg, Germany). \textit{CYP2C19} alleles were detected by specific probes in Lightmix for the detection of human \textit{CYP2C19*1} (wild type allele), \textit{CYP2C19*2} (rs4244285) and \textit{CYP2C19*3} (rs4986893). Detection reagent (TIB-MOLBIOL GmbH, Berlin, Germany) by real-time polymerase chain reaction (RT-PCR) (LightCycler 2.0; Roche Diagnostics GmbH), according to the manufacturer’s recommendations. The G681A point mutation in exon 5 of \textit{CYP2C19*2} and G636A transition in exon 4 of \textit{CYP2C19*3} were detected. The genotypes were identified by running a melting curve analysis with specific melting points (Tm). The wild type \textit{CYP2C19*1} exhibits a Tm 54.4 °C at channel 530 and Tm 53.4 °C at channel 640. The allele variant \textit{CYP2C19*2} exhibits a Tm of 48.6 °C at channel 530 and the allele variant \textit{CYP2C19*3} exhibits a Tm of 60.8 °C at channel 640.

**Detection of the \textit{P2Y12} Genotype.** The 52 (G>T) (rs6809699) and 34 (C>T) (rs17602729) polymorphisms of the \textit{P2Y12} gene was analyzed with the PCR-RFLP (restriction fragment length polymorphism) method. The primer sets used were: 5’-AAT AAT TCA CCT CTG CGC CCG G-3’ (wild type allele), \textit{CYP2C19*2} 5’-TTT AGA GGA GGC TGT GTC CAA-3’/5’-AAT AAT GTT 3’ for the \textit{52 (G>T)} polymorphism, and 5’-TTT AGA GGA GCC TGG GTC CAA-3’/5’-AAT ATT GTT ACC AGG CGC AGG GGT GAA-3’ for the 34 (C>T) polymorphism. The PCR was performed with 25 µL DreamTaq Green PCR master mix (Thermo Scientific, Pittsburgh, PA, USA) with 5 µL (75 ng) DNA, 1 µm of forward and reverse primers and PCR grade water in a total reaction volume of 50 µL. An ABI PRISM™ 9700 thermal cycler (Applied Biosystems, Grand Island, NY, USA) was used for the PCR reactions. The thermal cycling conditions were; an initial denaturation step at 94 °C for 3 min. and 35 cycles at 94 °C for 20 seconds, 57 °C for 20 seconds, and 72 °C for 25 seconds; a final extension was performed at 72 °C for 3 min. An \textit{Smal} enzyme (Thermo Scientific) was used for digestion of the amplification product for the detection of the 52 (G>T) polymorphism. The PCR product used to detect the 34 (C>T) polymorphism was digested with Tsp509 I (synonime TasI) (Thermo Scientific). The products were size-fractionated on a 2.0% agarose gel.
RESULTS

The study included patients monitored for acute ICVD in the previous 2 years and monitored by our clinic for at least 1 year. All patients included in the study began clopidogrel after acute ICVD. A total of 51 patients [21 males (41.17%) and 30 females (58.83%)] were included. Their average age was 66.4 ± 9.6 years.

When the *1, *2, and *3 alleles of CYP2C19 were evaluated, two patients were homozygous for *2/*2, 13 patients were heterozygous for *1/*2 and 36 patients were homozygous for the wild type *1/*1 alleles. No patient carried the *3 allele. Three heterozygous patients, one for *2/*2 and two for *1/*2, stopped clopidogrel due to repeated strokes and began to take warfarin. These patients had no previous history of warfarin use.

When the patients’ alleles were evaluated in the group without recurring ischemic stroke, the *2 allele frequency was 13.54%. In the recurring ischemic stroke group, the *2 allele frequency was 66.7%. In the relative risk calculation of the recurring ischemic stroke group, the odds ratio (OR) was identified as 13.23 [95% confidence interval (95% CI) 6.45-27.11], which was significant at $p <0.0001$ (Table 1). When the P2Y12 52 (G>T) and 34 (C>T) polymorphisms were evaluated, all alleles were of the wild type.

DISCUSSION

Clopidogrel is used to prevent the development of a new ischemic attack [1]. Clopidogrel prevents ICVD; it has also been shown to be effective in patients with peripheral artery disease who use clopidogrel [14,17]. However, after using the drug, individual differences in response were observed and this situation has brought insufficiency of clopidogrel treatment to the agenda [18,19].

In our study, in spite of using clopidogrel, three of our 51 patients developed ICVD during at least 1 year of follow-up and these patients began taking warfarin instead. When these patients began clopidogrel treatment, they had no history of warfarin use. When the CYP2C19 gene was evaluated in our patients with recurring ICVD, one patient was homozygous for *2/*2, and two were heterozygous for *1/*2.

In the patients with recurring ICVD, the *2 allele frequency was significantly higher than in the group without recurring ischemia ($p <0.0001$). In our three patients, recurring ICVD and subsequent necessity to change medication, was linked to the CYP2C19 gene polymorphism. This is because clopidogrel is a prodrug and transforms into its effective metabolite through the action of CYP2C19 enzyme in the liver CYP450 enzyme family. The transformation of clopidogrel into active metabolite reduces when linked to CYP2C19 genetic variations [5].

The reasons for clopidogrel resistance are multifactorial and other reasons include medication interactions and insufficient use of the drug [20]. In medication interactions, proton pump inhibitors (PPI) attract attention because PPI, especially, are frequently prescribed with clopidogrel. Proton pump inhibitors use reduces the effects of clopidogrel on platelets. This situation is worrisome as it may cause recurring strokes [21]. In our patient group, there was no use of PPI by the patients who had recurring strokes. Similarly, patients who did not use the medication sufficiently, another cause of unresponsiveness to medication, were not included in the study.

Recently, there were increasing worries about the failure in effectiveness of clopidogrel, due to genetic variations in CYP2C19 in the CYP450 enzyme family that facilitates metabolism of the medication [20]. As a result, the American Food and Drug Administration recommended that though not required, testing should be carried out before use, due to possible low effectiveness [22]. In our patient group, none of the patients had any tests before beginning the drug as the current social security system in our country does not cover the cost of the tests. Together with covering the cost of the tests, it is estimated that about

| Parameter                  | *1 (%) | *2 (%) | *3 (%) | OR, $p$ Value |
|----------------------------|--------|--------|--------|--------------|
| Recurring ICVD [+]         | 2 (33.33%) | 4 (66.66%) | –      | 13.23, <0.0001 |
| Recurring ICVD [−]         | 83 (86.46%) | 13 (13.54%) | –      | 13.23, <0.0001 |
3.0% (2.0-14.0%) of the population do not metabolize clopidogrel well [23]. Those with P2Y12 polymorphism using clopidogrel for peripheral artery disease are reported to have four times more ICVD than those with wild type genotypes. It has been shown that medications like aspirin are not affected by variations in P2Y12. These results may be linked to genetic variations in the target receptors for clopidogrel [14]. However, in our study, no patient was identified with the P2Y12 polymorphism.

In our patient group, no single person had both P2Y12 and CYP2C19 gene polymorphisms, so haplotype analysis was not possible. As a result, the clinical effects of such a combination could not be identified in our study. However, in a previous study of stent patients with acute coronary syndrome, the clinical results of clopidogrel treatment were monitored. It was shown that a combination of P2Y12 and CYP2C19 gene polymorphisms produced worse negative clinical results than each polymorphism alone [16]. The increase in the number of polymorphisms increasing the unresponsiveness to the drug was a sign that genetic variations are very important for clopidogrel effectiveness.

The CYP2C19 and P2Y12 gene polymorphisms may cause recurring stroke attacks linked to an insufficient response to ICVD treatment. Stroke is among the top causes of mortality and morbidity. Repeated ischemic stroke events increase this risk further. As a result, genetic testing before medication use is important for human life, standard of living and economic burden.

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