Migraine and Genetic Polymorphisms: An Overview

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Abstract: The relationship between genetic polymorphisms and migraine as a cause of an increased risk of thrombotic disorders development is still debated.

In this respect, factor V Leiden, factor V (H1299R), prothrombin G20210A, factor XIII (V34L), β-fibrinogen, MTHFR (C677T), MTHFR (A1298C), APO E, PAI-1, HPA-1 and ACE I/D seem to play a determinant role in vascular diseases related to migraine.

The present review analyzes both the incidence of the above genetic vascular mutations in migraineurs and the most recent developments related to genetic polymorphisms and migraine.

Keywords: Genetic Polymorphisms, Migraine, Thrombosis, Vascular Disease.

INTRODUCTION

Migraine is a common neurovascular disorder. During the last decade, many clinical studies have given evidence of an association between migraine, particularly migraine with aura, and ischaemic stroke. There are several pathophysiological mechanisms implicated in the genesis of ischaemic events in migrainous patients [1].

Recent studies assess the role of migraine as a risk factor for endothelial dysfunction, responsible not only for a reduced availability of vasodilatators and for an increase of vasoconstrictor agents, but also for a release of procoagulant, proinflammatory and proliferative factors, predisposing migraineurs to atherogenesis [2].

Endothelial dysfunction is due to an increased oxidative stress, promotor of inflammatory processes, proposed as implicated in the pathogenesis of migraine [3].

Migraine is related to an intracranial vasospasm followed by the maximum dilatation of extra and intra-cranial arteries responsible for pain [2-4]. Considering that platelet activation and plasma coagulability are increased during migraine attacks [5], it is strongly supported that genes affecting vascular endothelial function could play a significant role in cerebral blood flow changes occurring in migraineurs, so, contributing to the aetiology of the migraine [5,6].

In this respect, factor V Leiden, factor V (H1299R), prothrombin G20210A, factor XIII (V34L), β-fibrinogen, MTHFR (C677T), MTHFR (A1298C), APO E, PAI-1, HPA-1 and ACE I/D genetic polymorphisms seem to play a determinant role in vascular diseases in migraine [2-6].

Therefore, the association between genetic polymorphisms and migraine may induce an increased risk of thrombotic disorders development [2-6].

The present review analyzes both the incidence of the above genetic vascular mutations in migraineurs and the most recent developments related to genetic polymorphisms and migraine.

MIGRAINE AND GENETIC POLYMORPHISMS

Recently, several angiotensin I-converting enzyme (ACE) inhibitors and an angiotensin II receptor blocker were demonstrated to have a clinically important prophylactic effect in migraine. ACE is one of the key enzymes in the renin-angiotensin-aldosterone system, which modulates vascular tension and blood pressure [7]. In humans, serum ACE levels are strongly genetically determined. Individuals who were homozygous for the deletion (D) allele showed increased ACE activity levels. To investigate the role of ACE polymorphism in headache, the ACE insertion (I)/deletion (D) genotypes of 54 patients suffering from migraine with aura (MwA), 122 from migraine without aura, 78 from tension-type headache (TH), and 248 non-headache healthy controls was studied by Kowa et al. [7]. The ACE D allele were significantly more frequent in the MwA than controls. The incidence of the D/D genotype in MwA (25.9%) was significantly higher than that in controls (12.5%). No differences in the remaining groups were found. These findings indicate that the D allele and the D/D genotype in the ACE gene is a genetic risk factor for MwA thus suggesting a possible relationship between ACE activity and the pathogenesis of migraine [7].

Paterna et al. [8] evaluate if the DD genotype could also be associated with the frequency and duration of migraine...
without aura, in 302 patients suffering from migraine without aura, the genotypes of the ACE gene, plasma ACE activity, and the frequency (weekly) and duration of migraine attacks were evaluated. No drugs were given before (4 weeks) and during the study. The same evaluations were performed in 201 subjects without migraine. Patients with migraine without aura showed higher incidence of the ACE-DD gene (48.34%) than control subjects (37.32%). The frequency of migraine (average attacks per week) was higher in patients with DD (2.11 +/- 1.9) than in patients with ID (1.54 +/- 1.44). No difference in duration of migraine attacks (hours per week) was observed. Plasma ACE activity was increased in patients with the ACE-DD gene. These data suggest that ACE-DD gene polymorphism could have an important role in determining migraine attacks and the frequency of these attacks [8].

Also, angiotensin converting enzyme (ACE) gene has been implicated as a genetic factor associated with migraine [9]. A case-control study was performed to investigate the association between ACE and migraine in 240 migraine patients and 200 healthy controls [9]. There was no significant difference in allelic frequency (I and D) and genotype polymorphism (DD, DI and II) of the ACE gene in migraine patients and controls. Analysis of the difference in ACE polymorphism stratified by gender revealed that male migraine patients with the homozygote DD genotype (ACE-DD) were significantly fewer than that of male controls. There was no existence of a difference among the frequency and duration of headache in each subgroup of migraine patients stratified by ACE genotype. These findings indicate that ACE-DD may have a slight protective effect against migraine in male patients [9].

Another study was conducted to determine whether the ACE I/D gene variant is involved in migraine risk and whether this variant might act in combination with the previously implicated MTHFR C677T genetic variant in 270 migraine cases and 270 matched controls [10]. Statistical analysis of the ACE I/D variant indicated no significant difference in allele or genotype frequencies. However, grouping of genotypes showed a modest, yet significant, over-representation of the DD/ID genotype in the migraine group (88%) compared to controls (81%). Multivariate analysis, including genotype data for the MTHFR C677T, provided evidence that the MTHFR (T/T) and ACE (ID/DD) genotypes act in combination to increase migraine susceptibility [10]. This effect was greatest for the MA subtype where the genotype combination corresponded to an OR of 2.89 (95% CI:1.47-5.72, P = 0.002). In Caucasians, the ACE D allele confers a weak independent risk to migraine susceptibility and also appears to act in combination with the C677T variant in the MTHFR gene to confer a stronger influence on the disease [10].

Even the role of factor XIII (V34L), APO E gene, MTHFR C677T and A1298C polymorphisms in migraine patients with cerebrovascular risk was considered [11-16] for migraine’s vulnerability.

In fact, the common Val34Leu polymorphism of the A-chain factor XIII gene, associated with variations in factor XIII activity, has been suggested to play a significant role in the development of arterial and venous thrombotic disorders [11]. It was analysed the prevalence of this polymorphism in 17 patients with coexisting ischemic cerebrovascular disease and migraine (5 with aura, and 12 without aura), 89 patients with migraine (43 with aura, and 46 without aura), 116 patients with ischemic cerebrovascular disease, and 467 healthy Caucasian controls [11]. The factor XIII Leu 34 variant was present in 47.1; 40.5; 34.9; and 35.1% of patients with coexisting ischemic cerebrovascular disease and migraine, ischemic cerebrovascular disease, migraine, and control subjects, respectively. These data suggest that the factor XIII Leu 34 allele does not play a protective role against these disorders in our population [11].

Rainero et al. investigated the role of apolipoprotein E (APOE) polymorphisms in migraine by analyzing the APOE genotypes of 241 migraine patients and 587 controls [12]. The results indicated that APOE alleles were similarly distributed between cases and controls. However, a significant increase of the varepsilon2-varepsilon4 genotype in the group of migraine patients was also observed. Patients were divided into three subgroups: migraine with aura; migraine without aura; and mixed headaches (migraine associated with tension-type headache). Subgroup analysis showed that the varepsilon2-varepsilon4 genotype was significantly increased only in patients with mixed headaches. The stratification of patients by APOE status did not reveal significant associations with the clinical features of the disease. In conclusion, this study showed no significant association between APOE polymorphisms and migraine [12].

Another study [13] was performed to determine the prevalence of the MTHFR mutation in patients with migraine and tension-type headache (TH). Seventy-four patients with migraine headaches (22 with aura and 52 without aura), 47 with THs, and 261 normal controls were recruited. This study showed that the incidence of the homozygous transition (T/T) in migraine sufferers (20.3%) was significantly higher than that in controls (9.6%). Moreover, the frequency of the T/T genotype in individuals with migraine headaches with aura was remarkably high (40.9%). The MTHFR T allele was more frequent in the migraine group than in the control group. Therefore, these results support the conclusion that the MTHFR gene, causing mild hyperhomocysteinemia, may be a genetic risk factor for migraine [13].

Finally, in a case-control study, was studied the prevalence of two common MTHFR polymorphisms, C677T and A1298C, in 102 migraine patients (23 migraine with aura, 70 migraine without aura and nine with tension-type headache) and compared it to that of 136 healthy controls [14]. The frequencies of the T allele of MTHFR677 and the C allele of MTHFR1298 were significantly higher in the total migraine population (33.82%, 33.82%) than in controls (25.38% and 24.26%), respectively [14]. The genotypes T677T and C1298C were the only genotypes significantly associated with migraine. Patients with migraine with aura with C1298C and C677/C1298C genotypes were even more profoundly associated with migraine risk than others [14]. However patients with migraine without aura with T677T and C1298C genotypes showed the same susceptibility. Patients with C1298C and C677/C1298C genotypes may also predispose to tension-type headache.
Although the above studies were conflicting, they suggested a higher frequency of some prothrombotic genetic abnormalities in migrainous patients [11-14].

While the exact etiology of migraine headaches is unknown, several theories have been proposed. The vascular theory, as proposed in 1938 by Graham and Wolff [15], attributes migraines to an initial intra-cranial arterial vasoconstriction, resulting in reduced blood flow to the visual cortex, followed by a period of extra-cranial vasodilation [16]. Modern imaging techniques have shown that during a common migraine attack there are in fact only minor changes in cerebral blood flow, and the proposed initial vasoconstrictive phase may actually last much longer than the aura [17]. It has also been hypothesised that migraine sufferers have an inherent vasomotor instability and are more susceptible to the vasodilatory effects of certain physical and chemical agents. This point of view has been reinforced by the observation that organic nitrates, which are capable of delivering nitric oxide, trigger migraine attacks in migraineurs, at low doses, ineffective in normal subjects [18].

Moskowitz's theory involves the trigeminovascular complex, which links the aura and the headache of migraine [19-20]. In this theory the trigeminovascular neurons release substance P and other neurotransmitters in response to various triggers (Fig. 1).

Taken together, these events configure the so-called phenomenon of “neurogenic inflammation” considering that both excessive trigeminal discharge and neurovascular inflammation of meninges occur in migraine headache.

The relationship between hypercoagulability state and migraine is another interesting pathogenetic aspect to be considered. In this regard homocysteinemia is believed to be a clear risk factor for thrombosis and it can be associated with an increased stroke and cardiovascular risk [16-20].

Vascular disease seems to be mediated by a growth of oxidative stress that, in its turn, could show the way to vascular disease by direct and indirect mechanisms [21].

A direct association has been shown between oxidative plasmatic capacity and homocysteine plasmatic levels. Among these being the production of endothelial nitric oxide, one of the most important risk factors for vascular disease and, at the same time, a mediator involved in migraine genesis [22].

Hyperhomocysteinemia is an independent risk factor for coronary and artery disease [23,24] and is also associated with cerebrovascular disease [25,26]. Multiple genetic and environmental factors can lead to hyperhomocysteinemia. In particular a polymorphism in the 5,10-methylenetetrahydrofolate reductase (MTHFR) gene has been described and individuals homozygous for the mutation C677T, who exhibit reduced enzymatic activity and elevated homocysteine concentration [27]. The decreased activity of MTHFR, in case of mutated enzyme, determines a reduced cellular availability of methyl THF, the substrate for methionine synthase. This enzyme catalyses the transfer of a methyl group from methylenetetrahydrofolate to homocysteine, producing THF and

![Fig. (1). Integrated pathogenetic model of migraine.](image-url)
methionine. By this way C677T mutation in MTHFR gene can lead to elevated homocysteine concentration. Two different studies have investigated the prevalence of this mutation in migraineurs patients [14,28]. The MTHFR T allele seems to be more frequent in the migraine group than in the control group in both studies. Furthermore Kara et al. [14] determined the prevalence of another MTHFR polymorphism (A1298C) in migraine patients. The C allele of MTHFR1298 was significantly higher in migraine and tension-type headache patients. Interestingly migraine with aura patients with C1298C and C677T/C1298 genotypes were even more profoundly associated with migraine risk than others. The authors have not analysed blood homocysteine levels, but elevated serum levels of homocysteine have been reported in patients suffering of migraine with aura not in tension-type headache patients. Hering-Hanit et al [29] examined blood homocysteine in a significant number of patients with migraine but they did not find its levels augmented. To the best of our knowledge no study has been conducted on the possible correlation between blood homocysteine levels and migraine severity and on the possible role of hyperhomocysteinemia as a causative factor in the predisposition to migraine. In fact it is well known that hyperhomocysteinemia may lead to an excessive production of homocysteic acid [30]. In this view elevated homocysteic acid levels, secondary to hyperhomocysteinemia, could sensitize dura mater and cerebral arteries and/or favour the activation of trigeminovascular system, thus predisposing a patient to migraine attacks or increasing disease’s severity [30].

However, an impairment of mitochondrial oxidative metabolism might also play a role in the pathophysiology of migraine, by influencing neuronal information processing. Biochemical assays of platelets and muscle biopsies performed in migraine sufferers have shown a decreased activity of the respiratory chain enzymes [31]. Studies with phosphorus magnetic resonance spectroscopy ((31)P-MRS) have demonstrated an impairment of the brain oxidative energy metabolism both during and between migraine attacks [31]. However, molecular genetic studies have not detected specific mitochondrial DNA (mtDNA) mutations in patients with migraine, although other studies suggest that particular genetic markers (i.e. neutral polymorphisms or secondary mtDNA mutations) might be present in some migraine sufferers [31].

Also, mitochondrial DNA (mtDNA) haplogroup U, defined by the polymorphism 12308A>G, may constitute a risk factor for a stroke in migraine [31]. Finnila et al. [32] identified 14 patients with an occipital stroke and with 12308A>G by determining complete mtDNA coding region sequence for the patients and for population controls by conformation sensitive gel electrophoresis (CSGE) and direct sequencing [32]. Sequence information was used to construct a phylogenetic network of mtDNA haplogroups U and K, which was found to be composed of subclusters U2, U4, U5 and a new subcluster U7, as well as cluster K. Five patients with a migraine stroke belonged to subcluster U5 (P=0.006; Fisher's exact test). Many unique mutations were found among the patients with an occipital stroke including two tRNA mutations that have previously been suggested to be pathogenic. Analysis of mtDNA sequences by CSGE and comparison of the sequences through phylogenetic analysis greatly enhances the identification of mtDNA clusters in population and detection of mtDNA mutations in patients [32].

In this respect, it is very interesting the link between migraine, mitochondrial pathology and drugs used in relation to particular genetic mutations. The association between migraine and abnormal mitochondrial function has been suggested by clinical similarities between this form of headache and some mitochondrial diseases such as MELAS [33] in which 10 different mutations have been identified, but 80% of cases the mutation is 3243A>G in the leucine tRNA gene (tRNA Leu) and it is typically associated with migraine attacks with aura, vomiting and acute migraine. The mutation is transmitted by maternal inheritance. Other examples are MERRF and NARP [34]. MERRF syndrome (myoclonic epilepsy with ragged red fibers) is a mitochondrial encephalomyopathy, characterized by myoclonic seizures in combination, sometimes, sensorineural deafness, optic atrophy, short stature, or peripheral neuropathy. In some cases were observed lipomatosis, cardiomyopathy, retinitis pigmentosus, ophthalmoparesis and or pyramidal signs. The disease is progressive, with worsening of epilepsy and other symptoms, such as ataxia, deafness, muscle weakness and dementia. Over 80% of patients present with the mutation 8344A>G in the gene coding for the RNA transport of lysine (MTTK). Have also been identified other mutations in other genes coding for RNA transport and in the gene MTND5. These mutations can cause overlapping clinical syndrome (MERRF / MELAS) with infantile-like episodes.

Neuropathy, ataxia, and retinitis pigmentosus (NARP) is a condition that causes a variety of signs and symptoms affecting the nervous system. NARP, is a condition related to changes in mitochondrial DNA with a mutations in the MT-ATP6 gene.

Of more recent interest is the identification of haplogroups of the mitochondrial genome related to migraine. Zaki et al [35] studied the entire mitochondrial genome and sequenced in 20 haplogroup-H patients with migraine without aura (MoA), syndrome vomiting cicling (CVS) and controls. Polymorphisms of interest were tested in 10 additional CVS subjects and in 112 haplogroup-H adults with MoA. The 16519C-->T polymorphism was found to be highly disease associated: 21/30 CVS subjects [70%, odds ratio (OR) 6.2] and 58/112 migraineurs (52%, OR 3.6) vs. 63/231 controls (27%). A second polymorphism, 3010G-->A, was found to be highly disease associated in those subjects with 16519T: 6/21 CVS subjects (29%, OR 17) and 15/58 migraineurs (26%, OR 15) vs. 1/63 controls (1.6%). The authors concluded that these polymorphisms constitute a substantial proportion of the genetic factor in migraine pathogenesis, and strengthen the hypothesis that there is a component of mitochondrial dysfunction in migraine.

Also, it was attempted to correlate a particular haplogroup to the therapeutic response, Di Lorenzo et al [36] tested in sixty-four migraineurs if the therapeutic response to riboflavin is associated with specific mitochondrial DNA (mtDNA) haplogroups, particularly, they focused their attention on haplogroup H, which is known to differ from others in terms of energy metabolism. Forty patients responded to riboflavin treatment and 24 were nonresponders. The mtDNA haplogroup H was found in 29 subjects (20 migraine
without aura, 9 migraine with aura). Riboflavin responders were more numerous in the non-H group (67.5%). Conversely, nonresponders were mostly H (66.7%). The difference between the two groups was significant ($\chi^2 = 7.07; p = 0.01$). The presence of aura had no influence on riboflavin's effectiveness ($\chi^2 = 0.113; p = 0.74$) and was not associated with a particular haplogroup ($\chi^2 = 0.55; p = 0.46$). In this pharmacogenetic study, riboflavin appears to be more effective in patients with migraine with non-H mitochondrial DNA haplotypes. The underlying mechanisms are unknown, but could be related to the association of haplogroup H with increased activity in complex I, which is a major target for riboflavin. These results may have ethnic implications, since haplogroup H is chiefly found in the European population.

CONCLUSIONS
The present review suggest that genetic polymorphism could have an important role in determining migraine attacks, and could also affect the frequency of these attacks. Considering the studies presented, it is possible to supplement the pathogenetical pattern of migraine with genetic polymorphism resulting also in an alteration of endothelial regulation tone as well as mitochondrial function. Therefore, we may conclude that genetic mutations play a crucial role in the development of migraine and in the increased risk to develop an ischaemic disease in migraineurs, but further studies are necessary. Certainly further data is needed, through further studies, especially on the biomolecular level and on the quantification of the relative risk for each polymorphism and for their combination. In the future, the genotype therapy could be considered as a diagnostic marker for therapy through the use of anti-migraine drugs. Thus, it could be possible to perform a primary prevention of this not very well known illness.

CONFLICT OF INTEREST
The authors confirm that this article content has no conflicts of interest.

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