The Role of Folate Dependent Genetic Susceptibility in The Risk of Multiple Sclerosis

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Abstract

Background: Multiple sclerosis (MS) is a chronic inflammatory disease of the central nervous system (CNS) of unknown cause. Epidemiological studies implicate that genetic and environmental factors interplay major roles in the etiology of MS. Several investigators have reported elevated plasma homocysteine and reduced folate and B vitamin levels in MS. Aberrations in one-carbon metabolism have impact in the pathophysiology by genetic susceptibility and increased the risk of MS. Furthermore, abnormalities in a carbon metabolism pathway may result from environmental factors such as reduced levels of vitamin B and folate due to inadequate B vitamin and folate intake from foods or the problems on their synthesis, and genetic factors such as polymorphism in folate – dependent enzymes.

Results: Many studies showed conflictive results genetic polymorphisms on the risk of MS. In the present review, we have emphasized the genetic susceptibility to the risk of MS, especially genetic polymorphisms on one carbon metabolism pathway. The associations on individual differences on the levels of homocysteine, B vitamin, folic acid due to genetic make-up and the risk of MS will be discussed. There are numerous reasons causing the development of MS especially, individual susceptibility should not be neglected. We have focused on three gene polymorphisms namely Methylene tetrahydrofolate reductase (MTHFR) C677T, A1298C and Methionine synthase (MTR) G66A since they are key enzymes on one carbon metabolism pathway.

Conclusion: So far conducted studies showed that genetic factors on products of one carbon pathway have major roles and these factors contribute to the development of MS. Individual differences in the development and treatment of the disease should not be overlooked.

Keywords: Multiple sclerosis; Homocysteine; Methylene tetrahydrofolate Reductase (MTHFR); Methionine Synthase Reductase (MTRR); Methionine Synthase (MTR)

Introduction

Multiple sclerosis (MS) is characterized by inflammation, demyelination and axon damage. It is an autoimmune central nervous system (CNS) disease. Myelin sheaths, oligodendrocytes the axon and nerve cells are damaged. The disease often occurs in young adults. The prevalence is ranges from 2 to 200 per 100,000 depending on geographical features. It was first reported by Jean-Martin Charcot in 1868 that MS was a chronic disease [1]. MS generally starts during the most productive period of life. It leads the significant disabilities and labor loss, thus affects quality of life. Furthermore, high treatment costs cause this disease to be discussed at the social dimension as much as individual [2]. In MS, inflammatory plaques in the brain and spinal cord cause multiple attacks of multifocal neurological deficits, with attacks of illness and recurrent episodes of recovery. The plaques are characteristically numerous and vary with age. Each attack which becomes permanent causes the loss of axon and leads to progressive loss of neurological function. Initial complaints such as sense loss, numbness, double vision and walking difficulty are often weakness, however as the disease progresses, severe complications are observed from blindness to the loss of consciousness [3,4].


In spite of intensive research, both the etiology and the pathogenesis of MS are not completely resolved. One of the important and most commonly studies have been conducted on one carbon metabolism pathway (OCMP) since it has major impacts on the development of MS. Homocysteine (Hcy), folic acid (FA) and vitamin B (Vit B) have key components of OCMP. Hcy is a sulphurous amino acid, not involved in protein
structure. It is an essential amino acid and intakes from foods and also, synthesized from endogenous proteins. It is a methylated group of methionine and is metabolized by converting it back to methionine via remission, or by trans-sulphurisation, converting it into cysteine, methyl malonic and 2-methylcitric acid [5]. Elevated plasma Hcy levels cause hyperhomocysteinemia and consequently homocysteineuria. It is shown that increased plasma Hcy levels are a significant risk factor for diseases [6]. Increased Hcy levels can be taken at normal levels with FA intake. Therefore, investigating the cause of the increase in plasma Hcy which is a risk factor for many diseases and withdrawal of normal levels from food is important for health [7]. The etiology of hyperhomocysteinemia is considered to be multifactorial (genetic, nutritional, and lifestyle factors), not related to only FA intake, and there is an ongoing debate regarding the relative contribution of each factor. It is proposed that high Hcy levels are due to genetic polymorphism on enzymes which have important roles on OCMP. Causal mechanisms of MS development including the identification of the folate-mediated one carbon pathway that is directly involved in MS development have yet to be established.

Figure 1 demonstrates genetic polymorphisms which have major roles on OCMP. This pathway is named for one carbon folic acid or one carbon metabolism pathway since a carbon unit from serine or glycine is transferred to tetrahydrofolate (THF) to form methylene tetrahydrofolate (MTHF). Folate-activated one carbon is required for the de novo synthesis of purines and thymidylate and for the remethylation of Hcy to methionine. Methionine is used for protein synthesis or transforming S-adenosine methionine (SAM) which is required for polyamine synthesis and for numerous methylation reactions.

**Figure 1** Production of homocysteine as part of the amino acid and purine biosynthesis pathway (DHF: Dihydrofolate, THF: Tetrahydrofolate, MTHFR: Methylene Tetrahydrofolate-Reductase, TS: Thymidylate Synthase, MTR: Methionine Synthase, MTRR: Methionine Synthase Reductase, SAM: S-Adenosine Methionine, SAH: S-Adenosine Homocysteine).

Methylene tetrahydrofolate-reductase (MTHFR) is an enzyme and have impact on the levels of Hcy and metabolism of methionine. It catalyzes reduction of 5, 10-methylene THF (5, 10-MTHF) to 5-methyl THF (5-THF). 5- THF acts as a methyl donor during synthesis of methionine from Hcy. In case of genetic polymorphism, MTHFR enzyme activity decreases and failure to synthesize methionine from Hcy thus, leads high levels of Hcy [8].

**The impacts of Hcy, FA and VitB12**

Increase in Hcy levels due to FA and VitB12 deficiency causes the decrease in cellular S-adenosine methionine (SAM) level and consequently the inhibition of SAM dependent methylation reactions in the CNS. Therefore, this pathway is important for the risk of neurogenerative diseases such as MS. Disruptions or impairments in folate metabolism are associated to several diseases. Impairments in the folate-dependent one-carbon network can arise from a primary folate deficiency, secondary VitB12 nutrient deficiencies and genetic variations influence cellular folate accumulation and/or utilization.

So far, it has been known very well that high levels of Hcy and FA are associated to risk of many diseases not only impact on the development of MS. In a case-control study, it was reported that increased levels of FA reduced the risk of colorectal cancer by 50% [9]. There are numerous published
studies which report higher levels of Hcy and lower levels of folate were risk factors for cardiovascular diseases [9,10]. Diabetes, renal insufficiency, different types of cancers, ovarian failure is associated with hyperhomocysteinemia [11-13]. However, the most important disease group, which is associated with FA deficiency and hyperhomocysteinemia, is neurological and psychiatric diseases. In recent years, there have been a number of publications that FA is associated with diseases such as Parkinson’s, MS, Alzheimer’s, stroke, anxiety disorders, depression, schizophrenia, bipolar disorder, congenital neural tube defects [14-16]. High levels of Hcy are toxic to neural cells therefore Hcy contributes to the pathogenesis of neurodegenerative disorders. Furthermore, psychiatric disorders via neuronal degeneration due to Hcy take place. As mentioned Figure 1 (above), inhibition of methylation causes to the decreased methylation of phospholipids in neuronal membranes, thus leads the impaired stability of neuronal cell membranes. It has been discussed and proposed hypothesis that impaired methylation of Myelin Basic Protein (MBP), the major component of the myelin sheath, triggers the risk of MS. It was reported that these effects contribute to the pathogenesis of various psychiatric and neurological disorders [14,15,17] and also many reports showed that high levels of Hcy were observed in MS [18-24].

Genetic susceptibility and the risk of MS

Investigations on genetic polymorphisms provide data for the role of individual susceptibility to the development of MS. It is essential to solve its underlying mechanism. There are numerous reasons causing the development of MS especially, individual susceptibility should not be neglected. We have focused on three gene polymorphisms namely Methylene tetrahydrofolate reductase (MTHFR) C677T, A1298C and Methionine synthase (MTR) G66A since they are key enzymes on OCMP. There are two reasons to show the association with the MTHFR C677T polymorphism to disease. First, the disease might influence the levels of Hcy and there might be effect modification by the MTHFR gene polymorphism. Second, the genotype might be associated with disease risk, possibly mediated by altered metabolism of folates and Hcy [6,25]. The MTHFR gene consists of 11 exons and has identified 14 mutations. C677T and A1298C point mutations are the most common among them and result in loss of enzyme activity. In C677T polymorphism, cytosine (C) at the position 4th exon 677 transits to thymine (T). As a result, valine amino acid is substituted for the 226th amino acid domain of the enzyme structure. This polymorphism of the enzyme leads to loss of activity. The loss of activity of enzymes are related to the development of MS. The prevalence of C677T gene polymorphism differs from among ethnic groups such as 1% or less is in sub-Saharan African and US blacks and 25% or more is for Italians, US Hispanics [26]. Table 1 demonstrates the distribution of these gen polymorphisms on different populations. Other commonly studied gene polymorphism is A1298C polymorphism. The nucleotide at 1298th nucleotide in the 7th exon changes from A (Adenine) to C (Cytosine). This point mutation causes the change of glutamine to alanine in MTHFR protein. In this mutation, MTHFR activity is reduced as it is in other mutation types [27]. MTR is the enzyme that synthesizes methionine from Hcy by transferring methyl and consists of 1265 amino acids. MTR gene locates at chromosome 1q43 with 33 exons. One of the most studied polymorphisms in the MTR gene is A2756G and A>G transition at 27. exon 2756 leads the amino acid substitution of glycine to aspartic acid. This polymorphism decreases enzyme activity as the VitB12 cofactor regulates its reactivation and role in methylation [28,29]. Methionine synthase reductase (MTRR) plays a critical role in maintaining VitB12 in an active form. It has pivotal role on total plasma Hcy level. Elevated Hcy contributes to the CNS dysfunction, neurodegenerative, and cerebrovascular disease [30,31]. In the A66G gene polymorphism, isoleucine residues have been reported to be the remnants of methionine and has been reported to increase plasma homocysteine levels [28,32].

There are conflicting results in the literature about the association between C677T, A1298C MTHFR, MTR A2756G and A66G MTRR gene polymorphisms and the risk of the development of MS. Çevik et al. study MTHFR C677T gene polymorphism in Turkish population included 130 MS patients and 150 group-matched controls. Results show that the genotype and allele frequencies of the gene polymorphism are statistically significant different between MS patients and controls. Furthermore, T allele of C677T polymorphism was associated with MS susceptibility in Turkish population [33]. Fekih Mrissa et al. conducted a case–control (case n=39 and controls n=104) study in Tunisia population. They found a statistically significant difference in the frequency of MTHFR A1298C between controls and MS patients. Subjects with CC genotype was about 4 times higher the risk of MS. On the other hand, no significant difference was found in the MTHFR C677T gene polymorphism between controls and MS patients [34]. In UK Caucasian population, similar study was performed with larger sample size (n=104 patients) and it was found that the frequencies of MTHFR A1298C mutant genotypes were higher in MS patients compared to controls however it could not reach the significance. They suggested that it needed to be performed further studies with larger sample size to confirm their study results [35]. The association between MTHFR C677T and A1298C missense variants and the risk of MS was investigated in a Southern Iranian population (controls n=231 and patients n=180). Compared with controls, a strong association between two gene variants and the risk of MS was found. Subjects carrying 677T allele (CT and TT genotypes) had higher MS risk than subjects carrying CC genotype (2.9 times and 6.23, respectively). The variant 1298AC genotype also increased the risk for MS and 2.14 times higher MS risk was observed subjects carrying mutant genotypes [36]. Two polymorphisms MTRR A66G and MTHFR A1298C were investigated in an Australian case-control population (140 controls and 140 patients) study. No significant allelic frequency difference was observed between cases and controls [37]. The genotype frequencies of the missense variant MTHFR 1298A > C investigated study in Germany were significantly different between patients (1298AA: 1298AC and 1298 CC and 0.34: 0.55: 0.11) and controls (0.52: 0.36: 0.12).
These results suggest that homozygosity for the A allele of MTHFR 1298A > C may be protective against the incidence of MS [38]. As result in MTHFR C677T polymorphism C allele is protective, CT and TT genotypes have increased MS risk. For MTHFR A1298C polymorphism A allele is protective, AC and CC genotypes have increased the risk of MS. However, results do not support a major role for either functional gene mutation in MS susceptibility. But for MTR A2756G and MTRR A66G polymorphisms, there are not enough studies. Hence, further studies are need to be conducted (Table 1) [39-58].

Table 1 The prevalence of MTHFR C 677T, A1298C, MTRR A66G and MTR A2756G in different populations.

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Conclusion

In conclusion, genetic susceptibility should not be neglected to the development and treatment of MS. Although there are many conflictive results, when compile the data on the relation between genetic polymorphisms and the development of MS, genetic factors have important impact. Meta-analyses and studies with larger sample size needs to be conducted and enzyme activities should be detected and correlations with genetic polymorphisms needs to be made to show the underlying mechanism of MS.

Conflict of Interest

There is no conflict of interest.

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