Exploring Genetic Susceptibility to Fibromyalgia

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Fibromyalgia (FM) affects 1% to 5% of the population, and approximately 90% of the affected individuals are women. FM patients experience impaired quality of life and the disorder places a considerable economic burden on the medical care system. With the recognition of FM as a major health problem, many recent studies have evaluated the pathophysiology of FM. Although the etiology of FM remains unknown, it is thought to involve some combination of genetic susceptibility and environmental exposure that triggers further alterations in gene expression. Because FM shows marked familial aggregation, most previous research has focused on genetic predisposition to FM and has revealed associations between genetic factors and the development of FM, including specific gene polymorphisms involved in the serotonergic, dopaminergic, and catecholaminergic pathways. The aim of this review was to discuss the current evidence regarding genetic factors that may play a role in the development and symptom severity of FM.

Key Words: Fibromyalgia; Genes; Polymorphism, Genetic

INTRODUCTION

Fibromyalgia (FM) is estimated to affect from 1% to 5% of the general population, and most patients with FM are women.1 FM is characterized by chronic widespread pain and is commonly accompanied by a constellation of additional symptoms such as fatigue, sleep disturbance, joint stiffness, and depression. Although the central symptom of FM, the severity of pain can vary between FM patients and pain may fluctuate in a specific patient at different times. Despite much debate regarding the classification and concept of FM in the past, FM is now recognized as a disorder.2,3

Patients with FM experience significantly impaired quality of life and their cost of care represents a significant economic burden.4 In fact, FM patients have similar health care resource requirements as patients with other common chronic diseases, such as diabetes mellitus and hypertension.5 For these reasons, FM is now recognized as a major health problem, and many recent studies have evaluated the pathophysiology of FM. Although the etiology of FM remains unknown, it is thought to involve the interaction of numerous factors, including psychological, genetic, neurobiological, and environmental factors.6,7 Owing to the occurrence of marked familial aggregation among FM sufferers, much previous research has focused on genetic predisposition to FM and has provided reasonable support for an association between genetic factors and the development of FM.8-10 In this review, we explore the current evidence for potential genetic factors playing a role in the development and symptom severity of FM (Table 1).

GENETICS OF PAIN

Pain is known to have a genetic background in that it runs strongly in families in humans, which is due to pain-related genes that affect the expression or function of protein products in a way that affects the pain response.11 Individual differences in pain and analgesia can be explained by environmental as well as heritable factors.12 Pain genes have been evaluated in large familial studies using linkage analysis and in large cohorts of individuals with and without a specific disease by using association analysis.11,13 Consequently, several genes have been shown to play major roles in pain sensitivity, including the genes encoding voltage-gated sodium-channels (Nav),
TABLE 1. Candidate genes involved in the pathogenesis of FM

| Affected systems          | Genes                              | Effect on FM                                         | References |
|---------------------------|------------------------------------|------------------------------------------------------|------------|
| Serotonergic systems      | T102C in 5-HT<sub>2A</sub> receptor gene | Development of FM                                     | 39         |
|                           | T102C in 5-HT<sub>2A</sub> receptor gene | Psychiatric symptoms                                  | 27         |
|                           | 5-HTTLPR S/L                        | Development of FM                                     | 25, 41     |
|                           | 5-HTTLPR S/L, intron2 VNTR          | Depression and psychological distress                 | 41         |
|                           | Serotonin receptor subunit genes, HTR3A and HTR3B | Anxiety-related traits                                | 25         |
| Catecholaminergic systems | Rs4680 of COMT gene                 | Development of FM                                     | 17, 53, 54, 55, 56, 57 |
|                           | ACCG haplotype of COMT gene         | Increased pain severity                               | 17, 57     |
|                           | DRD4 exon3 repeat polymorphism      | More maladaptive coping and pain                     | 28         |
|                           |                                    | No effect                                            | 58, 59     |
| Dopaminergic systems      |                                    | Development of FM and increased pain severity        | 17, 55     |
| Other Ion channels        | SCN9A gene-encoded DRG sodium channel polymorphism | Development of FM and severe symptoms               | 10         |
| NO metabolism             | GCH1 haplotypes                     | Development of FM and pain sensitivity               | 69         |
| Adrenergic receptors      | ADR polymorphism                    | Development of FM and different domains of FM symptoms | 74         |

FM: fibromyalgia, 5-HTTLPR: serotonin transporter (5-HTT) promoter region, S/L: short/long, VNTR: variable number tandem repeat, COMT: catechol-O-methyl transferase, DRD4: dopamine-D4-receptor, DRG: dorsal root ganglia, NO: nitric oxide, GCH1: GTP cyclohydrolase 1, ADR: adrenergic receptor.

GTP cyclohydrolase 1 (GCH1), mu-opioid receptors, and catechol-O-methyl transferase (COMT). In this context, familial aggregation and genome-wide association studies (GWAS) of chronic pain syndromes, including FM, have been performed to evaluate genetic susceptibility to disease.

FAMILIAL AGGREGATION IN FM

Several studies have observed strong familial aggregation among FM patients. Pellegrino et al. investigated 17 families of patients with FM for evidence of inherited FM. In that study, 26 (52%) of the enrolled family members (parents and siblings) showed clinical evidence of FM, and an additional 11 individuals (22%) showed abnormal muscle consistency without tender points. However, the study participants were not diagnosed as having FM according to the 1990 American College of Rheumatology (ACR) criteria for the classification of FM.

Buskila et al. evaluated the familial occurrence of FM among 58 offspring aged 5 to 46 years from 20 complete nuclear families with FM-diagnosed mothers. FM patients were diagnosed according to the ACR criteria, and 16 offspring (28%) were found to have FM. Offspring with and without FM did not differ significantly in terms of their exposure to environmental factors or other psychological or familial factors, such as anxiety, depression, global well-being, quality of life, or physical functioning. Thus, these results suggest that certain genetic factors are associated with the high familial occurrence of FM.

In another study, Buskila et al. studied female FM patients and their close relatives (parents, siblings, children, and husbands) to assess the familial aggregation of FM. The prevalence of FM among the blood relatives of patients with FM was 26%, compared to only 19% among unrelated family members (husbands). The occurrence of FM showed marked female preponderance: 41% of female relatives showed evidence of FM, compared to only 14% of male relatives. Additionally, pain sensitivity among young male and female relatives, which was measured by using mean tender point counts, was significantly higher than in controls. Moreover, adult relatives had considerably higher mean tender point counts than did controls. The authors concluded that the relatives of patients with FM demonstrated familial aggregation with respect to FM and were likely to have higher pain sensitivity than the general population.

Arnold et al. evaluated the familial aggregation of FM and pain sensitivity among 533 relatives of 78 probands with FM and compared these with 272 relatives of 40 probands with rheumatoid arthritis (RA). FM cases showed significant aggregation among the relatives of FM probands compared to those of RA probands (odds ratio [OR], 8.5; p=0.0002). The mean number of tender points was also significantly higher among the relatives of FM patients than among the relatives of patients with RA.

Familial aggregation in FM does not exclude other possible etiologic factors in the development of FM. In fact, there
is evidence that environmental factors also contribute to the development of FM. However, familial aggregation does indicate that genetic factors are important to the etiology of FM. Subsequent investigations have demonstrated that FM is associated with genetic polymorphisms affecting the serotonergic, dopaminergic, and catecholaminergic systems. Furthermore, certain genetic polymorphisms influence not only the development of FM but also symptom severity.

GENOME-WIDE ASSOCIATION AND LINKAGE STUDIES IN FM

GWAS are a recently developed research technique used to evaluate many common genetic variants in different individuals to determine whether a specific variant is associated with a given trait. As such, these studies are typically used to investigate disease heritability. GWAS usually focus on associations between single-nucleotide polymorphisms (SNPs) and traits, such as certain diseases that are associated with genetic background. A linkage study has a similar aim to GWAS and is a recombinant technology used to map genetic loci by use of observations of related individuals.

GWAS have been used to evaluate the association between FM and genetic susceptibility. Docampo et al. explored genetic susceptibility to FM through GWAS and array comparative genomic hybridization assessments of copy number variants. In that study, among 505,454 SNPs, none of the SNPs reached the genome-wide significance threshold. Nevertheless, the authors identified two associated variants, an SNP in the MYTIL gene and an intronic deletion in the NRXN3 gene, that suggested a possible role for the central nervous system in FM genetic susceptibility. Feng et al. also discovered possible candidate genes in FM by whole exome sequencing. They identified 2 nonsense mutations, W32X in C11orf40 and Q100X in ZNF77, associated with FM, one of which correlates with high plasma MCP-1 and IP-10 levels and the other with high plasma IL-12 levels. This result suggests a probable inflammatory basis of the development of the syndrome.

In addition, Arnold et al. performed a linkage scan, evaluating 341 markers in members of 116 families from the Fibromyalgia Family Study. They detected a signal on chromosome 17p11.2-q11.2, indicating a susceptibility locus for FM. However, subsequent research on this linkage study has not yet been reported.

The results of association and linkage studies have expanded our understanding of the genetic basis of FM and have suggested several genetic markers as susceptibility genes for FM. However, studies are not sufficient and thus these findings remain inconclusive. Therefore, further well-designed studies consisting of large study populations are needed.

CANDIDATE GENE ANALYSIS IN FM

1. Genes involved in serotonergic metabolism

Earlier research on genetic susceptibility in FM focused on the serotonergic pathway. Accumulating evidence supports the concept of disturbances in serotonin (5-HT) metabolism and neurotransmission in FM. This is based on several recent findings. Russell et al. found that the level of serotonin metabolites was lower in serum and cerebrospinal fluid in FM patients. Similarly, Wolfe et al. showed that serum levels of 5-HT in FM patients were significantly lower than in controls. The serotonin transporter (5-HTT) is responsible for transport of 5-HT from synaptic spaces into presynaptic neurons, thus playing a key role in serotonergic neurotransmission. For these reasons, genetic variation in the serotonergic system has received attention as a possible genetic factor in the etiology of FM.

Bondy et al. investigated a silent polymorphism (T102C) in the 5-HTT receptor gene in 168 FM patients and 115 healthy controls and found significant changes in genotype distribution among FM patients compared with the controls. Specifically, FM patients showed a decrease in the T/T genotype and an increase in both the T/C and C/C genotypes. In addition, pain severity was significantly higher in patients with the T/T genotype. Gürsoy et al. also investigated the role of the T102C polymorphism in the development of FM. In contrast to the findings of Bondy et al., the T102C polymorphism of the 5-HTT receptor gene did not differ significantly between patients and controls. However, the genotypes influenced symptom severity in FM patients. The T/T genotype was associated with psychiatric symptoms of FM, as measured by using the Symptom Checklist-90-Revised (SCL-90-R) test. In addition, patients with the T/T genotype demonstrated the lowest pain thresholds. Furthermore, a meta-analysis demonstrated that the 5-HTT receptor T102C polymorphism conferred susceptibility to FM.

Offenbaecher et al. analyzed the genotypes of the promoter region of the 5-HTT gene in patients with FM. The frequency of the S/S genotype of 5-HTT was higher among FM patients (31%) than among healthy controls (16%). Furthermore, FM patients with the SS genotype exhibited severe symptoms of depression and psychological distress, as evidenced by higher scores on the Beck Depression Inventory (BDI) and the SCL-90-R compared to patients in the L/L and L/S groups. These results support the notion that altered serotonin metabolism is related to the development of FM and its symptoms.

Cohen et al. further evaluated the role of the 5-HTT promoter region (5-HTTLPR) polymorphism. Ninety-nine female FM patients from two Israeli ethnic groups (Jewish and Arab) were genotyped. In that study, each patient was assessed by using the Tridimensional Personality Questionnaire (TPQ), a self-reported instrument consisting of 100 yes/no questions. The results showed an association between FM and the 5-HTTLPR polymorphism in both
Israeli ethnic groups. Furthermore, the 5-HTTLPR polymorphism was associated with the TPQ scores of the FM patients. There was a significant association between 5-HTTLPR genotype and the TPQ harm avoidance trait, which is consistent with the results of an earlier study by Lesch et al. that revealed that the short allele of this polymorphism is related to anxiety-related traits.

Gürsöy et al. reported that the 5-HT gene polymorphism was not associated with FM patients who had normal psychiatric status. In that study, 53 mentally healthy FM patients and 60 healthy controls were included. However, the 5-HTTLPR polymorphism and the VNTR variant of the 5-HT gene were not significantly different between the patients and controls. Additionally, Frank et al. showed no association between FM and either of the serotonin receptor subunit genes, HTR3A and HTR3B, in 48 patients with FM. Moreover, the association between the 5-HTTLPR polymorphism and FM was not observed in a meta-analysis.

Thus, although the roles of specific genes involved in the serotonergic pathway in the pathophysiology of FM have not yet been confirmed, much evidence does support an association between genes involved in serotonin function and FM. In fact, studies have also shown that 5-HT3 receptor antagonists, such as ondansetron, granisetron, and tropisetron, are potential treatment options in the therapy of FM patients. Because the serotonergic pathway influences psychiatric symptoms, such as depression, anxiety, and fatigue, which occur commonly in patients with FM, it is possible that serotonergic gene polymorphisms indirectly affect variable clinical features of FM.

Collectively, insufficient research is available to fully understand the role of serotonergic gene polymorphisms in FM. Further studies are needed to better understand the role of genes involved in serotonergic metabolism in the development of FM.

2. The COMT gene in FM

Major advances in our understanding of the pathophysiology of FM have resulted from the recognition of central sensitization. Central sensitization represents an enhancement in the function of neurons and circuits in central nociceptive pathways and thus results in pain hypersensitivity. The development of central sensitization is related to the monoamine neurotransmitters, and COMT is a major enzyme that inactivates catecholamine neurotransmitters such as dopamine, epinephrine, and norepinephrine. Genetic mutations in the COMT gene can induce functional impairment in the COMT enzyme and alterations in COMT activity. Thus, polymorphism in the COMT gene has been suggested as a genetic factor associated with FM susceptibility and symptom severity.

The most widely investigated COMT gene polymorphism is the SNP rs4680. This SNP occurs in codon 158, resulting in a valine (Val) to methionine (Met) transition. Rs4680 can cause three possible genotypes and, as a result, alter the activity of the COMT enzyme. The H/H (GG; Val-158-Val) genotype gives rise to an effective enzyme, whereas the H/L (AG; Met-158-Val) genotype produces an intermediate-activity enzyme. The L/L (AA; Met-158-Met) genotype causes a defective enzyme that is incapable of effectively clearing catecholamines.

To date, data regarding the association between rs4680 and susceptibility to FM are inconclusive. However, much evidence supports an association between this SNP and the development of FM. Matsuda et al. revealed an association between rs4680 and FM susceptibility in a Brazilian population. Similar associations in a variety of ethnic backgrounds have been described by several investigators: Gürsöy et al. in a Turkish population, Valgas-Alarcón et al. in a Spanish population, and Cohen et al. in an Israeli population. Furthermore, the role of rs4680 was not limited to increased FM susceptibility. Barbosa et al. showed that rs4680 was not only associated with the development of FM, but also with increased pain severity in Brazilian FM patients. In that study, the individual carriers of the homoygous mutant genotype for rs4680 presented a higher Fibromyalgia Impact Questionnaire (FIQ) score. In another study performed in a Spanish population, it was found that rs4680 played a key role in pain severity in FM patients. FM patients with the Met158Met genotype showed higher pain hypersensitivity to thermal and pressure stimuli than patients with the Val158Met genotype. Finan et al. also showed that rs4680 may affect FM pain symptoms in a 30-day electronic diary assessment in which patients recorded daily ratings of pain and two forms of maladaptive coping: pain catastrophizing and pain attention. The authors found that FM patients with the Met158Met genotype reported more maladaptive coping and pain when pain catastrophizing or pain attention was elevated, relative to those with the Val158Val genotype. This result suggests that genetic variation in the Val158Met polymorphism may contribute to FM pain symptoms through pathways of pain-related cognition.

However, there is also evidence of no association between rs4680 and the development of FM. In a recent meta-analysis, stratification by ethnicity failed to reveal an association between rs4680 and FM in European or Turkish populations. Using an unpublished dataset, we also investigated the association between rs4680 and FM susceptibility in a Korean population and found that rs4680 was not associated with FM susceptibility.

Although other COMT gene polymorphisms have been identified, including the SNPs rs6269, rs4633, and rs4818, the roles of these variants in FM are not yet fully understood and have been contradictory in some cases. Thus, investigators have paid attention to the haplotype of the COMT gene rather than individual SNPs. The haplotype refers to the set of alleles of a group of closely linked individual SNPs that are usually inherited as a unit and is critical for evaluating the genetic basis of diseases. Certain haplotypes have been associated with FM susceptibility and also pain sensitivity. Diatchenko et al. revealed that specific haplotypes of the COMT gene were as-
associated with low, average, and high pain sensitivity, and the ACCG haplotype of the COMT gene was defined as a high pain sensitivity haplotype. Likewise, the ACCG haplotype of the COMT gene was more frequent among FM patients and was also associated with higher pain responsiveness to stimuli in a haplotype analysis.17 A previous report also showed associations between FM patients with the ACCG haplotype and higher FIQ scores.55

In conclusion, the COMT gene is one of most investigated genetic polymorphisms regarding susceptibility to FM. Although there are some negative results, COMT remains a highly likely candidate gene associated with FM susceptibility. Moreover, research is still in progress to evaluate the role of COMT gene polymorphisms in the disease course and the outcomes of FM. Further prospective large-scale studies are needed to connect clinical practice and the development of drugs for FM.

3. Dopamine receptor genes in FM

Dopamine is a key neurotransmitter involved in modulating pain perception and natural analgesia.63 Furthermore, dopamine has an important role in the descending inhibition of pain.64 In fact, altered dopamine receptor function has been demonstrated in FM patients.55,66 Malt et al.65 reported that FM patients showed an increased prolactin response (dopamine sensitivity) to the buspirone challenge test. But at the same time, FM patients presented no difference in hypothermic response or growth hormone release (5HT sensitivity), which suggests altered dopaminergic neurotransmission among FM patients. Moreover, pramipexole, a dopamine-D2/D3 agonist, demonstrated higher efficacy in FM patients compared with placebo. Treatment with pramipexole showed improvement on scores used to evaluate pain, function, fatigue, and global status in FM.67 It is in this context that researchers have investigated dopaminergic receptor genes as potential contributors to genetic susceptibility to FM. Buskila et al.68 reported that the D4 dopamine receptor exon III repeat polymorphism was associated with the personality profile of FM patients. FM patients with those polymorphisms scored low on the TPQ novelty-seeking personality trait. Again, there remains insufficient research to fully understand the role of the dopaminergic receptor genes in FM. However, on the basis of the effects of pramipexole in FM, it is worth further evaluating dopaminergic-related gene polymorphisms in FM.

4. Other genes potentially involved in FM

The genetic factors discussed so far do not fully explain the etiology of FM. Thus, researchers are also interested in other possible genetic polymorphisms. Because several ion channels contribute to the detection and processing of thermal, mechanical, and chemical pain stimuli, dysfunctional ion channels have been proposed as a possible risk factor associated with FM susceptibility. For example, one study suggested that the SCN9A gene-encoded dorsal root ganglia sodium channel polymorphism was associated with severe symptoms of FM.10

Recently, Kim et al.69 showed that certain GCH1 haplotypes had a protective role against FM susceptibility and pain sensitivity, suggesting that nitric oxide (NO) may be involved in pain sensitivity in the pathogenesis of FM. Oxidative stress and NO metabolism participate in various processes, such as vascular dilatation, neurotransmission, and immune regulation, and may also be involved in the regulation of pain in the pathogenesis of FM.70,71

FM patients frequently have symptoms related to relentless sympathetic hyperactivity, which may also explain FM symptoms unrelated to pain sensitivity.72 Adrenergic receptors (ARs) act as binding sites for catecholamines and serve important roles in the responses of the sympathetic nervous system.73 Given this background, Vargas-Alarcón et al.74 investigated the association between AR gene polymorphism and FM and found that AR gene polymorphisms were associated with the risk of developing FM and were also related to different domains of the FIQ score.

The pathophysiology and symptoms of FM are not fully described by the genetic factors highlighted in this review, even if all of the potential contributions discussed here are confirmed through further research. Thus, to deepen our understanding of the genetic basis of FM, researchers have also examined other pain-related genes. GWAS or genetic studies performed in other FM-related disorders would also be of interest.

CONCLUSIONS

The pathogenesis of FM is not yet fully understood. However, FM is currently thought to result from the interaction of numerous factors including physical, psychological, genetic, neurobiological, and environmental factors. Because familial aggregation strongly supports the likelihood of a genetic component to the etiology of FM, recent research has focused on genetic susceptibility and environmental exposures that trigger further alterations in gene expression.

In this context, certain polymorphisms involving genes from the serotonergic, dopaminergic, and catecholaminergic systems have been suggested as possible risk factors for the development of FM. Additionally, these polymorphisms may influence symptom severity and could be targets for pharmacological treatments. However, these genetic factors do not fully explain the etiology of FM. Moreover, researchers have as yet paid little attention to the effects of genetic factors on disease progression and treatment outcomes of FM. Further studies are required to evaluate the effects of genetic polymorphism on clinical outcomes in patients with FM. Indeed, many more prospective studies, involving larger numbers of FM patients, are needed to expand our understanding of the role of genetics in FM.
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CONFLICT OF INTEREST STATEMENT

None declared.

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