The Frequency of Familial Mediterranean Fever Gene Mutations and the Correlations between Phenotype and Genotype in Turkish Children

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Authors’ contributions

This work was carried out in collaboration among all authors. Authors HE and ACS designed the study, performed the statistical analysis, wrote the protocol and wrote the first draft of the manuscript. Author OG did the genetic study. Authors AE, DDS and FU managed the analyses of the study. All authors did the literature searches and helped write. All authors read and approved the final manuscript.

ABSTRACT

Aim: Familial Mediterranean Fever is an autosomal recessive disease characterized by recurrent inflammatory attacks of serosal membranes. The aim of the current study was to determine the frequency of the Mediterranean fever (MEFV) gene pathogenic variants in 158 children (78 male, 80 female) diagnosed with Familial Mediterranean Fever (FMF) and to compare the phenotype-genotype correlation.

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Methods: In our retrospective case-control study, 158 FMF patients (78 males, 80 females) who were diagnosed with MEFV gene mutation in Bursa Yuksek Ihtisas Training and Research Hospital, Department of Pediatrics between January 2018 and June 2019 were included in the study. Mutation screening of the MEFV gene was performed for 12 mutations and the 8 most common mutations were taken into the study.

Results: Abdominal pain (77.8%), fever (74%) and arthralgia (46.2%) were the most prevalent clinical features in our patients. The most frequent mutations were M694V, E148Q, V726A, M680I and P369S. In cases with M694 mutation, it was noted that the incidence of arthritis was 2.5 times, appendectomy frequency 3.1 times higher, and early diagnosis probability 3.2 times higher. The frequency of chest pain was 2.9 times higher in the M680I mutation, and the frequency of arthralgia was 2.2 times higher in the P369S mutation.

Conclusion: Patient’s mutations in FMF patients are important for clinical expectations, and some mutations such as P369S are not as innocent as expected. However, reevaluation of phenotypes of mutations that are rare with more patients will be significant.

Keywords: Familial mediterranean fever; MEFV gene mutations; phenotype-genotype correlations.

1. INTRODUCTION

Familial Mediterranean Fever (FMF) is an autosomal recessive autoinflammatory disorder characterized by recurrent attacks of fever and serosal inflammation [1]. The disease affects patients in the Mediterranean and Middle Eastern populations such as Jews, Turks, Arabs, Armenians with genetic prevalence of 1 to 6% [2]. The estimated prevalence of FMF is about 1/400-1/1000, and the carrier frequency is 20% in the Turkish population [3-4]. FMF is caused by mutations in the Mediterranean fever (MEFV) gene, which is located on the short (p) arm of chromosome 16 at position 13.3 [5]. Phenotype-genotype correlations have not been completely determined in patients with FMF. Types of MEFV mutations and clinical picture of FMF change in different geographic regions and ethnic groups. Several researchers have reported an increased disease severity and the development of renal amyloidosis in patients having specific MEFV gene mutations such as homozygous M694V [6]. We conducted the current study to analyze the correlation of variable clinical presentations and MEFV genotypic distributions in Turkish children.

2. METHODS

2.1 Patients

In our retrospective case study, 158 FMF patients (78 males, 80 females) who were diagnosed with MEFV gene mutation in Bursa Yuksek Ihtisas Training and Research Hospital, department of Pediatrics between January 2018 and June 2019 were included in the study. Epidemiological features such as age, gender, ethnicity, FMF story in family, consanguinity between mother and father, age of diagnosis and clinical features as fever, abdominal pain, chest pain, arthralgia, arthritis, erysipelas like erythema, myalgia, oral aphthae, headache during attack, appendectomy story and response to colchicine are noted in the cases.

2.2 Mutation Analysis

2-3 ml peripheral venous blood was collected into hemogram tubes for DNA isolation from all individuals evaluated by FMF pre-diagnosis. Isolation was performed in the MagPurix DNA isolation robot using the MagPurix Blood DNA Extraction kit. The obtained DNA samples were subjected to Polymerase Chain Reaction (PCR). For each case 2 μl DNA sample and 23 μl PCR mix were used. PCR protocol was performed in GenePro (Bioer) brand gradient system thermal cycler.

PCR samples of all cases were loaded to the NanoChip 400 using the FMF 12 Kit (Savyon Diagnostics). The results of the analysis with the NanoChip Version 7 analysis program were evaluated and the mutation status of the 12 regions (M694V, E148Q, M694I, V726A, M680I> A, M680I> C, R761H, F479L, I692del, P369S, K695R, A744S, P369S) on the MEFV (Mediterranean Fever) gene associated with FMF disease was checked. M694I, I692del, F479L mutations with allele counts less than 5 were removed from the study. M680I> A and M680I> C were considered as a single mutation.

2.3 Statistical Analysis

Descriptive analyzes of numerical data were shown as mean and standard deviation, and
3. RESULTS

Among the 158 children, 78 (49.3%) were males and 80 (50.7%) were females. The patient’s mean age was 9.37±4.41 (ranged between 1–17 years). The mean age at onset of FMF in Turkish children was 4.2±2.7 years and the age at diagnosis was 6.9±3.5. A positive family history of FMF was noted in 53 (33.6%) of the patients, whereas the parents of 15 (9.7%) patients proved to be relatives. The most frequent clinical findings were abdominal pain (77.8%), fever (74%), arthralgia (46.2%), myalgia (21.5%), arthritis (13.2%), headache (13.2%), erysipelas-like erythema (12.6%), oral aphtha (12.6%) and chest pain (8.8%). Colchicine was administered orally to all patients and a favorable therapeutic effect was seen in 122 patients (77.2%). The number of patients with appendectomy was 29 (18.3%). Amyloidosis was found only in a patient with homozygous M694V mutation (0.63%). Homozygous mutations were detected in 49 (30.8%), heterozygous mutations in 57 (36.5%), compound heterozygous mutations in 52 (32.7%) patients. The most common homozygous genotypes were M694V/M694V, heterozygous genotypes M694V/- and compound heterozygous genotypes M680I/M694V which were found in 37 (23.3%), 25 (15.7%), and 13 (8.2%) of these patients, respectively (Table 1). The most common mutation was M694V with a frequency of 50.5%, followed by E148Q (14.5%), V726A (12.9%), M680I (8.6%), P369S (4.1%), K695R (3.9%), R761H (3.1%) and A744S (1.9%) (Table 2). When patients were assessed by Logistic Regression analysis to evaluate genotype and phenotype association, it was found that arthralgia frequency in M694V mutation was 2.5 times higher than other mutations and appendectomy frequency was 3.1 times higher and it was noted that determination is 3.2 times earlier than other mutations (p<0.05). Similarly, arthralgia was found to be 2.2 times higher in the P369S mutation and chest pain was 2.9 times higher in the M680I mutation (p<0.05) (Table 3).

4. DISCUSSION

In this study, we evaluated the clinical features and the genotypes of FMF in 158 Turkish children. The hospital where the study was conducted is located in a migration-receiving region in Bursa. When the origins of the patients studied were examined, it was seen that they were from 7 regions of Turkey. Therefore, it is thought that this study reflects the general population of Turkey.

Although FMF disease is seen in the Mediterranean region, it has become a problem of the whole world due to travels and migrations. In recent studies, it is reported that the highest prevalence is among the Turks with 1 per 400 and 1 per 1000 [3-4]. One of the important reasons for this is the consanguineous marriage, which is still a serious problem for Turkey even though it has decreased over the years. Rates of consanguineous marriages vary according to regions in Turkey (The East of Turkey: 34.4%, The West of Turkey: 11-13%) [7-8]. In the present study, consanguinity rate was 9.7% (Bursa is in West of Turkey). Although some studies have reported a male predominance, the frequency of FMF was similar in both genders [9]. In our study, the frequency of FMF was found to be the same in both genders (female to male ratio was 1.02).

In studies conducted in countries where FMF is frequent and in our country, abdominal pain frequency is expressed as 91-96%, fever frequency is 92-100%, chest pain frequency is 31-84%, arthritis frequency is 33-70% and erysipelas like erythema frequency is 3-40% [10]. In our study, abdominal pain (77.8%), fever (74%) and joint findings (59.4%) were most common findings. Joint findings are present in the most common form of asymmetric, non-destructive arthritis in about 75% and the initial symptom in about 10% of the FMF patients. The incidence of arthritis in Turks, Armenians and Arabs is significantly lower than that reported in Jews [6,11]. In our patients, it was found that the rate of patients with arthralgia clinic was 46.2% and the incidence of arthritis was 13.2%. Amyloidosis was detected in only one patient with homozygous M694V genotype.
Table 1. Genotype distribution of patients

| Mutations               | Genotypes     | n  | %    |
|-------------------------|---------------|----|------|
| Homozygous              | M694V/M694V   | 37 | 23.41|
|                         | E148Q/E148Q   | 7  | 4.43 |
|                         | V726A/V726A   | 2  | 1.26 |
|                         | M680I/M680I   | 2  | 1.26 |
|                         | R761H/R761H   | 1  | 0.63 |
|                         | Subtotal      | 49 | 30.99|
| Heterozygous            | M694V/-       | 25 | 15.82|
|                         | E148Q/-       | 11 | 6.96 |
|                         | V726A/-       | 7  | 4.43 |
|                         | P369S/-       | 6  | 3.79 |
|                         | K695R/-       | 4  | 2.53 |
|                         | A744S/-       | 3  | 1.89 |
|                         | M680I/-       | 1  | 0.63 |
|                         | Subtotal      | 57 | 36.05|
| Compound Heterozygous   | M680I/M694V   | 13 | 8.22 |
|                         | M694V/V726A   | 10 | 6.32 |
|                         | E148Q/M694V   | 5  | 3.16 |
|                         | M680I/V726A   | 4  | 2.53 |
|                         | E148Q/P369S   | 3  | 1.89 |
|                         | E148Q/V726A   | 3  | 1.89 |
|                         | K695R/M680I   | 3  | 1.89 |
|                         | R761H/V726A   | 3  | 1.89 |
|                         | A744S/R761H   | 1  | 0.63 |
|                         | A744S/V726A   | 1  | 0.63 |
|                         | E148Q/R761H   | 1  | 0.63 |
|                         | K695R/M694V   | 1  | 0.63 |
|                         | K695R/P369S   | 1  | 0.63 |
|                         | K695R/V726A   | 1  | 0.63 |
|                         | M680I/R761H   | 1  | 0.63 |
|                         | M694V/P369S   | 1  | 0.63 |
|                         | Subtotal      | 52 | 32.83|
| TOTAL                   |               | 158| 100  |

Table 2. Allele frequencies of the MEFV gene mutations

| Allele  | Number of alleles | Frequency (%) |
|---------|-------------------|--------------|
| M694V   | 129               | 50.58        |
| E148Q   | 37                | 14.50        |
| V726A   | 33                | 12.94        |
| M680I   | 22                | 8.62         |
| P369S   | 11                | 4.31         |
| K695R   | 10                | 3.92         |
| R761H   | 8                 | 3.13         |
| A744S   | 5                 | 1.96         |
| 255     |                    | 100          |

Mutation analysis in FMF disease, which depends on criteria such as typical clinical findings for diagnosis, response to family history and colchicine, is of great importance in terms of confirmation of diagnosis. In the current study, 12 MEFV mutations were screened among 158 Turkish children. The most common mutation was M694V (allele frequency was 50.5%). The most frequent mutation reported in Turkey in multicenter studies is reported as M694V and the allele frequency is reported at various rates of 14.6- 51.4% [8,12,13]. Although the ranking according to ethnicity is different in studies conducted both in Turkey and in countries where FMF is frequent, the 5 most common mutations are reported as M694V, E148Q, V726A, M680I, M694I [4,6,14]. In our study, the frequency of M694I was found to be low, and instead of it the P369S mutation was detected among the first 5 mutations (allele frequency was 4.1%). In 2017, Cekin and et al. conducted a study with 514 patients in Istanbul, indicating that the P369S allele frequency was 3.3%, the fifth most frequent mutation [15]. When genotypes of our patients were evaluated, M694V/M694V (23.3%), M694V/- (15.7%) and M694V/M680I (8.2%) were the most common genotypes. E148Q was the second most common mutation in our study (14.5%).
Table 3. Logistic regression models between MEFV gene mutations and clinical parameters

| Symptom           | MEFV gene mutations | OR    | % 95 confidence interval | p    |
|-------------------|---------------------|-------|--------------------------|------|
| Arthritis         | M694V               | 2.58  | 1.745 – 7.852            | 0.040|
| Appendectomy      | M694V               | 3.15  | 0.892 - 5.127            | 0.028|
| Early Diagnosis   | M694V               | 3.22  | 1.275 - 7.825            | 0.029|
| Arthralgia        | P369S               | 2.21  | 1.296 – 5.177            | 0.044|
| Chest Pain        | M680I               | 2.91  | 1.545 – 5.660            | 0.045|

The second most common mutation also in the majority of studies conducted in Turkey was E148Q [15,16,17]. In the multicenter study conducted by the FMF Working Group with 2838 patients in 2005, the second most frequent mutation was found M680I (14.4%) [8].

When our patients were evaluated by multiple regression analysis in terms of phenotype / genotype, in patients with M694V mutation, arthritis (OR: 2.58, CI: 1.74-7.85) and appendectomy (OR: 3.15, CI: 0.89 -5.12) were found to be more frequent than other mutations, and the diagnosis was earlier (OR:3.22, CI:1.27-7.82) (p<0.05). In 158 children with FMF, only 1 patient developed amyloidosis, in which M694V homozygous mutation was detected. Currently, the most numerous phenotype-genotype data available relate to M694V. The penetrance of M694V is very high. This genotype has been demonstrated to be correlated with a severe disease course in Jews, Arabs, Armenians and Turks (early disease onset, frequency of attacks, arthritis, high frequency of amyloidosis, high dose colchicine requirement) [18,19]. The results of our study also support this. Another mutation with severe clinical signs is reported as M680I [20,21]. M680I was the third most common mutation in our study with a rate of 8.6%. Chest pain frequency was 2.9 times higher in M680I mutations than in other mutations (OR:2.91, CI:1.54-5.66) (p<0.05). In previous studies, P369S has been described mild phenotype or incomplete penetrance in patients with FMF [6].

In our study, when joint findings were evaluated, arthritis was common in the M694V mutation and arthralgia was more frequent in the P369S mutation (OR: 2.21, CI: 1.29-5.17). Cekin and et al. also found that the incidence of abdominal pain and chest pain in patients with P369S mutation was higher than expected [15]. In both studies, although the P369S allele frequency was low (3.3% and 4.1%), the phenotype was not considered to be very slight in the P369S mutation. E148Q and V726A are the least penetrant FMF mutations and are recognized to have a mild effect on FMF patients in all reports. Especially in homozygous mutations of E148Q, 55% of the patients are asymptomatic, and amyloidosis is not reported in these patients [22,23,24]. In our study, when allele frequencies and clinical findings were evaluated by logistic regression analysis, no statistically significant relationship was found between E148Q and A726A mutations and clinical findings (p<0.05).

5. CONCLUSION

Phenotype-genotype association in FMF patients is important in terms of clinical expectations. In our study, compatible with the literature, M694V mutation was the most common mutation and it was found to be associated with severe clinical findings. The result was that the P369S mutation did not progress with very slight clinical findings as expected. However, studies with more patients are needed to further evaluate the phenotype-genotype relationship.

CONSENT AND ETHICAL APPROVAL

The study has been approved by the ethics committee of Bursa Yuksek Ihtisas Training and Research Hospital and informed consent was obtained from all patients who participated in the study.

COMPETING INTERESTS

Authors have declared that no competing interests exist.

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