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The Changing Concepts regarding the Mediterranean Fever Gene (MEFV): Toward a Spectrum of Pyrin-Associated Autoinflammatory Diseases (PAAD) with Variable Heredity

Guilaine Boursier, PharmD, PhD, Veronique Hentgen, MD, Guillaume Sarrabay, MD, Isabelle Koné-Paut, MD, Isabelle Touitou, MD, PhD*

1: Department of Medical Genetics, Rare Diseases and Personalized Medicine, CHU Montpellier, Montpellier, France
2: Department of general pediatrics, CH Versailles, CEREMAIA, France
3: Stem cells, Cellular plasticity, Regenerative medicine and Immunotherapies, INSERM, University of Montpellier, Department of Medical Genetics, Rare Diseases and Personalized Medicine, CEREMAIA, CHU Montpellier, Montpellier, France
4: University Paris Sud-Saclay, UVSQ, Kremlin Bicetre, France; AP-HP, CHU de Bicetre, Pediatric Rheumatology, CEREMAIA, Le Kremlin Bicetre, France.
5*Corresponding author: I Touitou, Stem cells, cellular plasticity, regenerative medicine and immunotherapies, INSERM, University of Montpellier, Department of Medical Genetics, Rare Diseases and Personalized Medicine, CEREMAIA, CHU Montpellier, Montpellier, France

Isabelle.touitou@inserm.fr
Tel: +33 4 67 33 58 59

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Key words: FMF, autoinflammatory disorder, PAAD, PAAND

Abbreviations and acronyms:

IL-1: interleukin 1
FMF: Familial Mediterranean fever
MEFV: Mediterranean fever gene
PAAD: Pyrin-associated autoinflammatory disease
PAAND: Pyrin-associated autoinflammatory disease with neutrophilic dermatosis
PAPA: Pyogenic arthritis, pyoderma gangrenosum and acne
PFAPA: Periodic fever' apthous-stomatitis, cervical pharyngitis and adenitis
SAID: Systemic autoinflammatory disease
WES: Whole-exome sequencing

Familial Mediterranean fever (FMF) is a genetic disease characterized by recurrent fever and serositis attacks. It is the oldest known hereditary recurrent fever and the first systemic autoinflammatory...
disease (SAID) whose gene (MEFV for Mediterranean FeVer, NM_000243.2) has been identified.\textsuperscript{3, 4} The two decades that followed this discovery have been fruitful in clinical, genetic and functional developments, making FMF a unique and fascinating model for understanding this disease, and the innate immune defense in general. FMF usually begins at pediatric age and can be seen by a wide variety of pediatricians: generalists, rheumatologists, infectious disease specialists, gastroenterologists or dermatologists.

We will retrace the impressive scientific and medical journey that has switched a little-known rare disease gene into a field of intense interest leading from basic research to personalized medicine. We will explain how a gene initially identified from individuals meeting well-defined criteria (FMF) is now recognized as being at the origin of a wide and unanticipated range of clinical phenotypes jointly referred to as pyrin-associated autoinflammatory disease (PAAD).

The evolution of knowledge has led to a better interpretation of the impact of mutations in the various clinical pictures associated with the MEFV gene, resulting in increasingly relevant diagnoses.

From benign paroxysmal peritonitis to FMF

The various names given to FMF since its description in the early 1900s reflect the main features of the disease. Historical cases of FMF were reviewed in a previous dedicated study.\textsuperscript{1} Figure 1 summarizes the key steps of the clinical and genetic discoveries in FMF. The first indisputable report of FMF was by Janeway, in 1908 (paroxysmal syndrome, with recurrent vomiting).\textsuperscript{5} Heller, in 1955, coined the current name “familial Mediterranean fever”,\textsuperscript{2} which was recently validated by an international consortium for taxonomy and definition of autoinflammatory diseases.\textsuperscript{6}

From classical FMF to the clinical criteria for its diagnosis

FMF is a recessive disease rare in Europe or USA, but relatively common in some populations of the Mediterranean basin, with prevalence of healthy carriers (ie, individuals carrying one pathogenic mutation) reaching up to one in five in Armenians.\textsuperscript{7} The estimated worldwide prevalence of FMF is
The classical FMF phenotype is characterized by self-limited and short-duration episodes of fever lasting 2 to 3 days. The mean age at disease onset is 4 years, but symptoms can occur as early as the first year of life or late in adulthood. Symptoms of acute inflammation include sterile peritonitis, pleural inflammation, arthritis, skin rashes, and elevated levels of acute phase reactants related to inappropriate secretion of interleukin 1 (IL)-1, IL-6 and tumor necrosis factor α (TNFα) reflecting the activation of the innate immune system. Absence of autoantibodies and auto-reactive T cells denotes the absence of abnormal stimulation of adaptive immunity and provides an additional argument for recurrent autoinflammatory fever. Colchicine at an adequate dose and full compliance prevents inflammatory attacks and occurrence of renal AA amyloidosis in approximately 95% of cases. IL-1 blockers can be effective in refractory cases. FMF is diagnosed on a clinical basis and can be confirmed by genetic testing (see below). The most commonly used clinical criteria are those of Tel Hashomer, which are established in the Jewish adult population but also validated in children of other ethnicities. The Livneh criteria are also widely used, but they lack specificity in the pediatric population. A Turkish group has proposed new criteria for FMF diagnosis in children, which were validated in several ethnic groups. New, more comprehensive diagnostic criteria, are under development.

From the clinic to genetics

The observation of familial FMF cases fueled gene hunting in the early 1990s. The MEFV gene contains 10 exons and encodes the pyrin protein. Four founding mutations in exon 10 (p.M680I, p.M694I, p.M694V, and p.V726A) account for almost 80% of clinically and ethnically typical cases. To date, nearly 100 times more sequence variants have been recorded in the dedicated online database in fevers (https://infevers.umai-montpellier.fr/). Homozygosity for p.M694V confers the most severe phenotype (more joint and renal damage, earlier onset), whereas other low-penetrant variants such as p.E148Q in exon 2, are considered susceptibility factors for inflammation.
From classical recessive inheritance to possible dominance and gene-dosage effect

Initial familial segregation analyses demonstrated autosomal recessive inheritance, exacerbated by consanguinity, a cultural habit in Mediterranean countries. However, in about one-third of typical clinical cases, only one mutation is detected. The recognition of a large series of heterozygous individuals with a clinical picture of FMF supports that some single mutations (mainly in exon 10) may induce clinical signs. Many alternative explanations have been proposed, such as the existence of a second hit in another region of the gene or in another gene implicated in the regulatory pathways of the immune system, epigenetics or environmental factors, but none of these hypotheses have been definitely proven to date. The non-detection of a second mutation does not necessarily confirm a dominant transmission. True dominant transmission is rare in FMF and requires proof of segregation over several consecutive generations, i.e. that patients with overt disease do carry the pathogenic mutation, whereas asymptomatic individuals do not. Anecdotal cases of proven vertical transmission of p.M694I and p.M694del in clinically ascertained FMF have been described (Table; available at www.jpeds.com), thus challenging the paradigm of pure recessive inheritance.

A large study from an international database of patients with SAID (Eurofever) has provided clues to begin to understand this apparent contradiction. The authors highlighted a qualitative (severe mutations vs polymorphisms) and quantitative (2, 1, 0 mutations) molecular gradient paralleling the clinical picture (i.e., ranging from typical FMF to periodic fever, aphthous-stomatitis, cervical pharyngitis and adenitis (PFAPA), a disease for which no causative gene has been pinpointed). The authors suggested a dose effect (genetic dosage) involved in the phenotype and hypothesized that, under certain circumstances (e.g. presence of variants of still unidentified modifier genes), high penetrance MEFV mutations could give rise to symptom onset even in the presence of one wild-type allele.
From gene identification to the understanding of the Pyrin function

The apparently recessive nature of FMF has led to the postulation that the underlying molecular mechanism was a loss-of-function of the mutated pyrin. In asymptomatic carriers, the defective function of a mutated allele is compensated by the second wild-type allele and pathogenic mutations in each parental gene are classically necessary for the disease to be expressed. The demonstration that MEVF mutations are in fact gain-of-function reconverted this prevailing concept. The introduction of various exon 10 mutations into mouse pyrin (knock-in) induced a phenotype similar to human FMF, whereas deletion of mouse pyrin (knock-out) did not result in enhanced inflammation.

Pyrin is a key component of a specific inflammasome. Inflammasomes are multiprotein platforms leading to activation of the pro-inflammatory IL-1 and IL-18 following a pro-inflammatory event. RhoA GTPases regulate pyrin by activating serine-threonine kinases PKN1 and PKN2, which in turn, phosphorylate pyrin at two specific serines (S208 and S242 in humans). GTPase-mediated pyrin phosphorylation results in inhibitory binding by cellular 14-3-3 proteins (Figure 2). Upon toxin stimulation or bacterial infection, RhoA GTPases are inactivated, which leads to serine dephosphorylation, 14-3-3 dissociation, and subsequent pyrin inflammasome activation.

From FMF to PAAD

Patients with FMF criteria now benefit from the invaluable progress brought about by the discovery of the MEVF gene. However, sequencing of specific exons has sometimes revealed unusual phenotypes, variants and inheritance. For example, p.P369S and p.R408Q substitutions in exon 3, which most often come from the same parental chromosome (cis variant or complex allele), were associated with a highly variable phenotype, and were infrequently associated with typical FMF symptoms. The associated phenotypes included atypical FMF, PFAPA-like and entero-Behçet
disease. In general, the patients are colchicine-responders. In a three-generation Spanish kindred, five affected members fulfilling the Tel-Hashomer criteria presented with a severe periodic inflammatory disorder segregating with the rare p.H478Y MEFV variant located in exon 5. Thus, the authors questioned whether this familial phenotype represented a case of unusual FMF phenotype or another MEFV-associated periodic inflammatory disorder because of its dominant inheritance, the prolonged febrile episodes, and the resistance to colchicine. Recently, next generation sequencing of a panel of known SAID genes or whole-exome sequencing (WES) facilitated the identification of new mutations and SAID genes but also yielded unexpected results. Two such examples are worth noting (Table and Figure 2). In a family of British descent with a colchicine-responsive autosomal-dominant periodic fever syndrome, WES sequencing was conducted and revealed a c.1730C>A missense mutation in exon 8 resulting in the p.T577N substitution. Mutations at the same codon were found in three other individuals of various ethnic backgrounds. Their disorder was similar to FMF (fever, systemic inflammation, response to colchicine or IL-1 blockade) but somewhat distinct to it (inheritance pattern, duration of attacks lasting one week, evanescent urticaria-like skin rash). The authors hypothesized that mutations at codon 577 lead to a structural change significantly affecting pyrin function or oligomerisation. Dominant inheritance of genetic variants leading to p.T577N was also observed in a Japanese family with PAAD. In individuals with severe neutrophilic dermatosis resembling pyogenic arthritis, pyoderma gangrenosum and acne (PAPA) syndrome WES investigations disclosed a dominant mutation within and in the immediate vicinity of the 14-3-3 protein pyrin binding domain (Table 1). This new pyrin-associated autoinflammatory disease with neutrophilic dermatosis was named PAAND. Only three PAAND families, all of European origin, have been reported so far: two harbored the p.S242R substitution and the third had the p.E244K substitution. Both mutations abolished the ability of 14-3-3 to bind pyrin, which resulted in a constitutive inflammasome formation.
Conclusion

Pan-genomic approaches have revolutionized the identification of the pathophysiological pathways involved in SAID, an understanding of their regulation but also their better nosological classification. PAAD is an “umbrella” name chosen by the taxonomy committee to include all diseases caused by pyrin defects or MEFV mutations.\(^6\) This general term of PAAD was developed to link different subtypes of disorders of the same genetic origin.\(^6\) Butbul Aviel et al found that PFAPA (with and without FMF features) may be more common in patients from Mediterranean ancestry and suggested that the incidence and features of PFAPA may be influenced by the presence of the common MEFV variants.\(^{40}\) Association of MEFV variants with multifactorial disorders has often been reported (see review).\(^{41}\) However, these variants are thought to act as susceptibility factors with a weak individual effect, and cannot be considered as directly causative of the disease. Therefore, PFAPA does not belong to the PAAD family. In contrast, the molecular mechanism involving the MEFV gene as the cause of the disease is clearly established in PAAND which therefore represents a PAAD subtype.\(^38\)

Comprehensive analysis of mutations in the MEFV gene revealed that symptom severity is associated with sequence variants in specific exons, at least in typical Armenian individuals.\(^42\) The exon in which the genetic variant is located also seems to affect the hereditary mode of transmission of the PAAD concerned, although the exact molecular mechanism underlying these observations remains elusive. Variable modes of transmission need to be considered for genetic counseling.

We have depicted the steps that led to the elucidation of the role of defective pyrin and have pinpointed the importance of the interpretation of mutations in less commonly affected exons, specifically in non-Mediterranean individuals, or in those in whom only one MEFV mutation has been detected. As a result, retro-phenotyping, which involves the molecular geneticist verifying the clinical presentation of the patient after the mutations have been identified to check for consistency, is becoming increasingly challenging. The reason is that quite often, genetic variants predicted as
pathogenic based on familial segregation, mutation type, frequency in the general population, or in silico tools are found in genes that do not seem relevant to the phenotype of the patient. However, effective and targeted treatment depends on the correct genetic diagnosis.

Acknowledgments

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References

Legends to figures

**Figure 1: Key milestones of the clinical and genetic discoveries in familial Mediterranean fever (FMF)**

Three main periods can be delineated. The first steps were clinical (in red), basically devoted to the disease description, with definitive naming and clinical criteria as key events. The second important period required both clinical and genetic investigations (in purple). Identification of the FMF gene and its protein pyrin was the “diamond” of 1997. The last decade (in blue) saw the advent of high throughput sequencing approaches pinpointing pyrin as being involved in many more phenotypes described under the “umbrella” pyrin-associated autoinflammatory disease (PAAD). The unraveling of the normal pyrin function as well as the demonstration that mutations are gain of function with gene dosage helped in understanding the clinical spectrum of PAAD.

PAAND: pyrin-associated autoinflammatory disease with neutrophilic dermatosis

**Figure 2: Schematic representation of the pyrin protein and mutations resulting in pyrin-associated autoinflammatory disease (PAAD)**

The B30.2 domain contains the most common pathogenic familial Mediterranean (FMF) mutations, which classically result in autosomal recessive FMF. Dominant mutations in exons 5, 8 and 10 are responsible for FMF-like and non-FMF PAAD. The precise molecular mechanism involved in these PAADs is still mainly unsolved. In contrast, dominant mutations in exon 2 in or close to the 14-3-3 binding domain are now known to impair binding of 14-3-3 to phosphorylated pyrin and trigger spontaneous pyrin-inflammasome formation, resulting in pyrin-associated autoinflammatory disease with neutrophilic dermatosis (PAAND). Ex: Exon.
### Table 1. Families with proven dominant inheritance associated with pyrin mutation compared to classical recessive FMF

<table>
<thead>
<tr>
<th>Origin (N families; N relatives)</th>
<th>Mediterranean</th>
<th>British (1:2)</th>
<th>Japanese (1:3)</th>
<th>Northern-Europe** (4:10)</th>
<th>Spanish (1:5)</th>
<th>British (1:7)</th>
<th>Turkish (1:8)</th>
<th>Dutch (1:3)</th>
<th>Japanese (1:3)</th>
<th>Belgium (1:12)</th>
<th>British (1:2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exon Mutation(s) ***</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>p.M694V (c.2080A&gt;G) most frequent</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>5</td>
<td>8</td>
<td>8</td>
<td>8</td>
<td>8</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Age at onset</td>
<td>childhood</td>
<td>Typical</td>
<td>7-30</td>
<td>6-35</td>
<td>9-13</td>
<td>6-10</td>
<td>ND</td>
<td>ND</td>
<td>3-16</td>
<td>childhood</td>
<td>ND</td>
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<tr>
<td>Attack duration</td>
<td>2-4 days</td>
<td>Typical</td>
<td>ND</td>
<td>2.5-6 days</td>
<td>6-12 days</td>
<td>Days to weeks</td>
<td>1-1.5 weeks</td>
<td>weeks</td>
<td>1day-6 months</td>
<td>Several weeks</td>
<td>ND</td>
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<tr>
<td>Fever</td>
<td>Yes</td>
<td>Typical</td>
<td>Yes</td>
<td>8/10 alive patients</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>No or low grade</td>
<td>Yes</td>
<td>Yes</td>
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<tr>
<td>Serositis</td>
<td>Yes</td>
<td>Typical</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Arthritis</td>
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<tr>
<td>Skin lesions</td>
<td>Erysypelas-like erythema</td>
<td>Typical</td>
<td>Typical</td>
<td>Yes</td>
<td>Yes</td>
<td>Erysipelas</td>
<td>Yes</td>
<td>Evanescent urticaria-like skin rash</td>
<td>Yes</td>
<td>Evanescent urticaria-like skin rash</td>
<td>No</td>
</tr>
<tr>
<td>Response to colchicine</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>Yes or ND</td>
<td>ND</td>
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<tr>
<td>Biological inflammation</td>
<td>Yes</td>
<td>Typical</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>Yes or ND</td>
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<td>Yes</td>
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<td>22</td>
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<td>35</td>
<td>35</td>
<td>36</td>
<td>37</td>
<td>37</td>
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</tbody>
</table>

FMF: familial Mediterranean fever; ND: not determined

* A patient harboring a c.1729A>T (p.T577S) mutation was also detected in a Dutch patient with an autoinflammatory phenotype but no familial nor clinical data was available.

** Sporadic patients described in this paper are not shown as dominant transmission is not ascertained.

*** E148Q was detected in some patients but is not mentioned as it is now recognized as a variant of unknown significance.
Pyrin-associated autoinflammatory disease (PAAD)

<table>
<thead>
<tr>
<th>PAAND</th>
<th>Severe periodic fever</th>
<th>FMF-like</th>
<th>FMF</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ex2</td>
<td>Ex5</td>
<td>Ex8</td>
<td>Ex10</td>
</tr>
</tbody>
</table>

### Mutations
- **PAAND**: S242R, E244K
- **Severe periodic fever**: H478Y
- **FMF-like**: T577A, T577N, T577S
- **FMF**: M694del, M694I

**Protein Domains**
- PYD
- Inflammasome
- Coiled Coil
- B30.2