<table>
<thead>
<tr>
<th>Title</th>
<th>Familial Mediterranean fever phenotype progression into anti-cyclic citrullinated peptide antibody-positive rheumatoid arthritis: a case report</th>
</tr>
</thead>
<tbody>
<tr>
<td>Author(s)</td>
<td>Yago, Toru; Asano, Tomoyuki; Fujita, Yuya; Migita, Kiyoshi</td>
</tr>
<tr>
<td>Citation</td>
<td>Fukushima Journal of Medical Science. 66(3): 160-166</td>
</tr>
<tr>
<td>Issue Date</td>
<td>2020</td>
</tr>
<tr>
<td>URL</td>
<td><a href="http://ir.fmu.ac.jp/dspace/handle/123456789/1360">http://ir.fmu.ac.jp/dspace/handle/123456789/1360</a></td>
</tr>
<tr>
<td>Rights</td>
<td>© 2020 The Fukushima Society of Medical Science. This article is licensed under a Creative Commons Attribution-NonCommercial-ShareAlike 4.0 International license.</td>
</tr>
<tr>
<td>DOI</td>
<td>10.5387/fms.2020-07</td>
</tr>
<tr>
<td>Text Version</td>
<td>publisher</td>
</tr>
</tbody>
</table>
Familial Mediterranean fever phenotype progression into anti-cyclic citrullinated peptide antibody-positive rheumatoid arthritis: a case report

Toru Yago, Tomoyuki Asano, Yuya Fujita and Kiyoshi Migita

Department of Rheumatology, Fukushima Medical University School of Medicine

(Received March 13, 2020, accepted October 14, 2020)

Abstract
Familial Mediterranean fever (FMF) is caused by dysfunction of the MEFV gene product, pyrin. Here we report a case of FMF phenotype which developed into rheumatoid arthritis (RA), based on a positive result for anti-cyclic citrullinated peptide (CCP) antibody (Ab). A 42-year-old woman presented to our clinic with more than 6 months of intermittent arthralgia in the wrists, feet, and fingers associated with menstruation. No fever was reported and there was no family history of FMF or other autoimmune diseases. Laboratory tests revealed elevated C-reactive protein (CRP) and rheumatoid factor (RF). Tests for autoantibodies including anti-CCP Ab, antinuclear Ab, and anti-DNA Ab were all negative. Genetic analysis identified an R304R homozygous mutation in MEFV; however, the pathological significance is unclear because this mutation does not cause amino acid substitution. We diagnosed incomplete FMF phenotype despite the lack of fever due to periodic arthritis, lack of autoantibodies, and complete resolution of arthritis following colchicine treatment within a day. Several months later, increased stiffness and arthralgia persistently occurred in finger joints on both sides. Ultrasonography revealed synovitis at the metacarpophalangeal and metatarsophalangeal joints. Laboratory analysis revealed the patient to be positive for anti-CCP Ab. Therefore, we finally diagnosed RA. Her arthritis diminished following administration of methotrexate and salazosulfapyridine. We consider the possibility that pyrin dysfunction may have affected the acquired immunity, contributing to the onset of RA as an autoimmune disease. This is an interesting case of equivalent FMF progressing into RA and will be valuable to raise awareness of a continuum from autoinflammatory to autoimmune disease.

Key words: Familial Mediterranean fever, Rheumatoid arthritis, Anti-CCP antibody, Autoinflammatory disease, Autoimmune disease

Introduction
Autoinflammatory diseases are characterized by systemic inflammation such as fever and rash, which differentiates them from autoimmune diseases such as rheumatoid arthritis (RA) or infections. In terms of pathology, autoinflammatory diseases are caused by innate immunity while autoimmune diseases are caused by adaptive immunity, meaning that the two disorders are on opposite ends of a disease spectrum even though they share many clinical characteristics. Familial Mediterranean fever (FMF) is the most common autoinflammatory disease, characterized by recurrent attacks of fever and polyserositis. Recently, pyrin — encoded by the MEFV gene — has been revealed to be directly involved in inflammasome activation and enhancement of interleukin (IL)-1β production. Genetic and clinical differences have been reported between Japanese and non-Japanese patients with FMF,
indicating that a relationship exists between genotype and clinical phenotype. It is common for FMF to co-exist with established rheumatic disease; however, FMF phenotype in the context of RA is very rare. We report the case of a patient with identical FMF who presented with joint synovitis in whom anti-cyclic citrullinated peptide ( CCP) antibody (Ab) began to be detected during the disease course, with eventual progression into RA.

**Case report**

A 42-year-old woman presented with swelling of the dorsum of the left foot and intermittent arthralgia in both hands lasting for 6 months with no fever. Her medical history included a malignant sweat gland tumor in the head when she was 30 years old. She had been previously diagnosed with iron deficiency anemia because of myoma uteri, for which she was taking iron supplements. No urinary tract infection or diarrhea had occurred before the onset of arthralgia. The joint swelling and arthralgia only occurred during menstruation. She was referred to our clinic for further examination with intermittent polyarthritis. She had no family history suggestive of FMF or other autoimmune diseases including RA. Cardiovascular and respiratory examinations were unremarkable with no enlargement of the liver or spleen noted. However, physical examination revealed swelling and tenderness of the left foot as well as redness and tenderness in the metacarpophalangeal joint of the left thumb occurring only during menstruation. In addition, her arthritis was not a typical synovitis characterized as “fusiform swelling” that can be seen in RA, but rather, the swelling of an entire digit. No fever or oral ulcers were observed and blood tests showed normal hemoglobin levels and normal leukocyte and platelet counts. Laboratory tests revealed C-reactive protein and rheumatoid factor levels to be elevated (0.41 mg/dL and 64 IU/mL, respectively). Anti-CCP Ab, anti-nuclear Ab (ANA), anti-DNA Ab, anti-SS-A Ab, anti-aminocyl tRNA synthetase Ab, myeloperoxidase (MPO)-anti-neutrophil cytoplasmic Ab (MPO-ANCA), and proteinase-3 (PR3)-ANCA were all negative. Immunoglobulin G (IgG), IgA, IgM, and IgD levels were normal and parvovirus B19-IgM was negative. Urinalysis showed no proteinuria or occult blood. Genetic analysis revealed a R304R homozygous mutation in the MEFV gene; however, the pathological significance is unclear because no amino acid substitution was identified. The patient did not fulfill the EULAR/ACR 2010 criteria for classification of RA at that time. Following colchicine treatment (0.5 mg/day), the arthritis entirely disappeared within a day. Then, we applied international Tel-Hashmer criteria by Livneh et al. to the present case, matching two minor criteria, including arthritis and favorable response to colchicine. Therefore, despite the lack of fever, we diagnosed equivalent incomplete FMF phenotype, based on periodic arthritis, complete disappearance of arthritis within a day of initiating oral colchicine treatment, and lack of autoantibodies. Four months later, stiffness and arthralgia persistently developed in some joints despite colchicine treatment up to 1.5 mg/day, and each joint was also fusiform; therefore, we performed ultrasonographic examination which revealed synovitis including the second metacarpophalangeal joints of the right side and the first metatarsophalangeal joints of both sides (Figure 1). Because of the observation of typical synovitis, we assayed anti-CCP Ab again, which was positive at this time (titer 11.2 U/mL). Therefore, the diagnosis of RA emerging from FMF phenotype was made at that time because the symptoms fulfilled the EULAR/ACR 2010 criteria for classification of RA including anti-CCP Ab positivity. Treatment with 5 mg/day prednisolone, methotrexate, and salazosulfapyridine was initiated, and arthritis improved within two months.

**Discussion**

The pathology of FMF is caused by dysregulation of inflammation due to abnormal function of the protein pyrin. Patients with FMF develop many clinical manifestations such as recurrent attacks of fever, periodic arthritis, and polyserositis. The most characteristic symptom is recurrent fever attacks of more than 38°C lasting several days. The condition usually affects pre-adolescent individuals; however, a Japanese nationwide survey found the mean age at onset of clinical symptoms to be 19.6 ± 15.3 years in Japanese patients, with around one-third of patients (37.3%) experienced their first attack after they were 20 years old, indicating that Japanese patients with FMF develop their clinical manifestations later compared to non-Japanese cohorts. In the present case, the age of onset was 42 years, which is similar to that in Japanese patients with FMF. Migita et al. reported the incidence of fever, chest pain as a result of pleuritis, and arthritis in FMF among a mainly Japanese cohort to be 95.5%, 35%, and 31.3%, respectively, which was
comparable to reports from non-Japanese cohorts\textsuperscript{11}). It should be noted that about 5% of Japanese FMF patients are afebrile. In contrast, the prevalence of abdominal pain as a result of peritonitis and AA amyloidosis are higher in non-Japanese than Japanese patients with FMF. Arthritis is one of the common clinical features of FMF; therefore, the possibility of FMF should be considered in patients presenting with arthritis, especially in the absence of RF or anti-CCP Ab, regardless of ethnicity. However, the diagnostic criteria vary for different populations; the Japanese criteria are based upon modified Tel-Hashmer criteria, which are simplified from the diagnostic criteria proposed by Migita’s group\textsuperscript{9}, which include one major criterion (recurrent febrile episodes) and eight minor criteria (febrile attack with abdominal pain from peritonitis, chest pain from pleuritis, monoarthritis, pericarditis, scrotal pain from orchitis, headache from aseptic meningitis, or a favorable response to colchicine treatment). On the other hand, the Livneh’s criteria\textsuperscript{7}, also derived from Tel-Hashmer criteria, provide an established international diagnosis of FMF. Importantly, in contrast to the Japanese criteria, periodic fever is not required for diagnosis of FMF in Livneh’s criteria. Livneh’s criteria are based on for major criteria involving typical attacks (peritonitis, pleuritis or pericarditis, monoarthritis, and fever) and five minor criteria involving incomplete attacks (abdomen, chest, joint, exertional leg pain, and favorable response to colchicine). If the patient fulfills two minor criteria, we can diagnose incomplete FMF even without periodic fever. Moreover, a diagnosis of FMF also should be concluded after exclusion of differential diagnoses such as infection, malignancy, or other autoinflammatory disease. It is highly important that a diagnosis of FMF be based on clinical manifestations, not solely genetic analysis, according to EULAR recommendations\textsuperscript{12}). Mindful that this case was afebrile, we diagnosed equivalent incomplete FMF by Livneh’s criteria, since the patient had arthritis that rapidly resolved following colchicine treatment.

It remains unclear why the present case was afebrile during the disease course; however, FMF without fever has been described in several case reports\textsuperscript{13–15}). It may be that dysfunction of cofactors such as microsomal prostaglandin E synthase-1 (mPGES-1), a terminal enzyme required for PGE\textsubscript{2} synthesis, may suppress fever as pro-inflammatory cytokines are necessary for febrile response. An in vivo study demonstrated that lipopolysaccharide (LPS)-stimulated febrile response is attenuated in chimeric mice when mPGES-1 expression is restricted to hematopoietic cells\textsuperscript{16}). Thus, the synergistic actions of IL-1β, IL-6, and mPGES1 in non-hematopoietic cells regulate the LPS-induced febrile response. In our case,
insufficient expression of mPGES-1 in non-hematopoietic cells may have resulted in the afebrile presentation; however, further investigation is required to elucidate the exact mechanisms underlying the lack of fever in FMF.

The present case experienced recurrent arthritis but joint swelling and pain was completely absent when she was not menstruating. This phenomenon is an important indicator of FMF, because menstruation is known to be a trigger for clinical attacks of FMF. Therefore, in cases such as ours, the patient’s arthralgia can be characterized as FMF-related arthropathy.

This case had a homozygous mutation of MEFV (R304R), which does not cause amino acid substitution. The MEFV gene is responsible for FMF and more than 300 sequence variants have been identified; of the 14 reported in 2012, nine are pathogenic and five have unknown significance. It has been reported that 86.5% of Japanese patients with FMF have one or more MEFV mutations or polymorphisms, the most frequent being M694I/E148Q (19.8%)20. However, about 15% of Japanese FMF patients have no MEFV mutations. On the other hand, the frequency of the MEFV mutations in RA has been reported by Migita’s group21. They demonstrated that allele frequencies of R408Q, P369S, E148Q, L110P mutations account respectively for 5.6%, 6.7%, 24.2%, 9.5% in RA patients and that the overall mutation rate was comparable between RA and healthy controls, indicating that MEFV mutations are not rare in RA. Their findings also suggest that MEFV mutations could not be a genetic factor affecting the susceptibility to RA. In addition, most importantly, recommendations allow a diagnosis of FMF even without MEFV mutations22. Based on these factors, it is possible to diagnose our case as FMF phenotype.

Migita et al. reported that, among 192 Japanese patients with FMF, 26 (13.5%) experienced accompanying rheumatic diseases. While the coexistence of autoinflammatory and autoimmune diseases is not rare23, reports of coexisting FMF and RA are relatively rare23, with anti-CCP Ab-positive RA being particularly rare. Monoarthritis associated with FMF usually occurs in large joints such as the knee or hip and protracted arthritis is rare24; in contrast with early RA where polyarthritus frequently develops in the hand or ankle. On the other hand, Endo et al. reported that late-onset FMF patients with onset ≥40 years present significantly more frequently with arthritis and myalgia compared to those with onset <40 years25. The present case is also late-onset FMF phenotype, therefore occurring predominantly with arthritis, consistent with Endo’s report.

Our case exhibited symmetrical arthritis of the finger and ankle that resolved with colchicine treatment in early phase, with ultrasonographic evidence of synovitis in the joint, which is an important indicator of early RA in late phase. However, the present case did not fulfill the EULAR/ACR 2010 classification criteria for early RA without anti-CCP Ab positivity in early phase. When the present case developed arthritis just intermittently, each joint exhibited swelling of an entire digit, which made it hard precisely localize synovitis. EULAR/ACR 2010 criteria refer to target patients who have at least 1 joint with “definitive clinical synovitis”. Therefore, the present case was not classifiable as having RA with a score <6/10 (namely, presence of high titer of RF, and elevated levels of CRP). There are very few reports investigating the association FMF with RF positivity. Guler et al. showed that the proportion of RF-positive FMF was 4% although the number of patients was small. They also demonstrated that RF positivity was not different between FMF and healthy controls.

The anti-CCP Ab is a highly specific biomarker for RA (>95% specificity) that indicates a high risk of progressive joint destruction. A case complicated by anti-CCP Ab-positive RA and FMF has been reported by Migita et al.25. In their case, arthritis refractory to conventional synthetic disease-modifying antirheumatic drugs (csDMARDs) was completely improved by using colchicine. The relationship between FMF and anti-CCP Ab is controversial. One study reported a relationship between FMF and anti-CCP Ab, while another study reported no such relationship. Importantly, Karatay’s report demonstrated that anti-CCP Ab was negative in all FMF patients. However, arthritis caused by FMF may disappear with colchicine treatment. In our case, arthritis in early phase was temporarily resolved by colchicine. In contrast, arthritis was refractory to colchicine after anti-CCP Ab was detected, and csDMARDs such as MTX were effective. Moreover, each joint swelling became “fusiform,” which is typical for RA synovitis. Therefore, we diagnosed RA when anti-CCP Ab positivity was observed. This change in antibody status presents an interesting case where RA developed and overlapped with equivalent FMF; recently, “autoinflammatory-autoimmune continuum” has been reported as a new disease spectrum. Autoinflammatory diseases are systemic.
disorders caused by dysregulation of innate immunity, whereas autoimmune disorders are caused by dysregulation of adaptive immunity. However, symptoms of systemic juvenile idiopathic arthritis often include fever and polyarthritis, and genome-wide analysis has demonstrated that autoimmune arthritis can develop from a predominantly autoinflammatory disease phenotype. Furthermore, patients with haploinsufficiency of A20 (HA20), caused by heterozygous germline mutations in the TNFAIP3 gene, present with clinical manifestations of autoinflammatory diseases such as Behçet disease, often coexisting with various autoimmune disorders. Innate immunity is accelerated in RA; for example, activation of the NLRP3 inflammasome, a major component of autoinflammatory pathogenesis, reportedly contributes to autoinflammatory diseases as well as autoimmune diseases including RA.

The mechanism underlying the development of RA in the present case is unclear; one possible explanation is involvement of the "pyrin inflammasome," which is a recently proposed concept suggesting that pyrin directly enhances inflammation following inactivation of Rho GTPases by bacterial toxins and other effectors, leading to dephosphorylation of pyrin. As a result, the 14-3-3 protein disassociates from pre-active pyrin resulting in pyrin activation. The inflammasome is then activated through polymerization of activated pyrin and apoptosis-associated speck-like protein containing a CARD (ASC). Finally, the pyrin inflammasome induces IL-1β production and inflammation. Thus, pyrin may function as a sensor of inactivation of Rho GTPases induced by pathogens like intestinal flora.

Patients with FMF could be more sensitive to dysbiosis (disturbance of intestinal bacterial flora and loss of protection from various pathogens), provoking systemic innate or adaptive immunity involving intestinal epithelial cells and other immune cells. Indeed, dysbiosis has been detected in patients with RA, as dysbiosis can lead to post-translational modification of autoantibodies and subsequent development of RA. Thus, dysbiosis of gut bacteria could drive adaptive immunity and autoimmunity leading to RA and innate immunity leading to FMF.

We encountered a rare case of equivalent FMF which developed to RA, providing an interesting "hybrid" phenotype which cannot be classified as a purely autoinflammatory or autoimmune disease. Furthermore, our findings suggest that therapy for RA should be decided according to the conditions which induce the progression to autoinflammatory or autoimmune disease, especially when biological agents are considered. For example, the IL-6 inhibitor tocilizumab has been reported to be beneficial for colchicine-resistant FMF as well as RA among Japanese patients. The present case adds valuable information to current knowledge relating to pathophysiological categorization of autoinflammatory and autoimmune disorders.

References
