



REVIEW ARTICLE

Twists and turns of the genetic story of mevalonate kinase-associated diseases: A review

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Abstract Mevalonate kinase (MK)-associated diseases encompass a broad spectrum of rare auto-inflammatory conditions, all resulting from pathogenic variants in the mevalonate kinase gene (MVK). Their clinical manifestations are highly variable, ranging from more or less serious systemic disorders, such as hereditary recurrent fevers, to purely localized pathologies such as porokeratosis. The oldest condition identified as linked to this gene is a metabolic disease called mevalonic aciduria, and the most recent is disseminated superficial actinic porokeratosis, a disease limited to the skin. The modes of inheritance of MK-associated diseases also diverge among the different subtypes: recessive for the systemic subtypes and dominant with a post-zygotic somatic genetic alteration for MVK-associated porokeratosis. This review quickly retraces the historical steps that led to the description of the various MK-associated disease phenotypes and to a better understanding of their pathophysiology, then summarizes and compares the different genetic mechanisms involved in this group of disorders, and finally discusses the diverse causes that could underlie this phenotypic heterogeneity.

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Abbreviations: DSAP, Disseminated superficial actinic porokeratosis; HIDS, Hyper-IgD syndrome with periodic fever; IL-1, Interleukin 1; MA, Mevalonic aciduria; MK, Mevalonate kinase (protein); MKD, Mevalonate kinase deficiency; MVK, Mevalonate kinase (gene); PK, Porokeratosis; RP, Retinitis pigmentosa.

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Introduction

Mevalonate kinase (MK) is an enzyme involved in the isoprenoid pathway and cholesterol synthesis that converts mevalonate into 5-phosphomevalonate.¹ MK is ubiquitously expressed, notably in blood leukocytes and skin keratinocytes. The gene encoding MK, *MVK*, is located on chromosome 12q24 and contains 10 coding exons and one non-coding exon.²

Several phenotypes have been associated with pathogenic variants in *MVK*. They all belong to a heterogeneous group of conditions called autoinflammatory diseases, resulting from mutations in genes of the innate immune system.^{3,4} The first MK-associated disease described which was most obvious on a pathophysiological level was a recessive metabolic disorder named mevalonic aciduria (MA; OMIM 610377); enzyme precursors were constantly detected in patients' urine.¹ Recently, dominant *MVK* mutations were also identified in patients with porokeratosis (PK), a phenotype localized to the skin and totally different from the systemic form.

The various clinical subtypes and modes of inheritance that have emerged^{1,2} and many molecular developments that have occurred over the last 28 years have led to a better knowledge of the pathophysiology of these diseases^{5,6} and to improved patient care.^{7–9} The main clinical advances brought about by pinpointing the culprit gene was the implementation of sequencing tests for diagnosis.¹⁰ The understanding that MK-associated diseases are caused by a decrease in enzyme activity has led to the development of effective targeted therapies (e.g., the injection of interleukin 1 [IL-1] blockers for patients with the systemic subtype) and gives hope for future local therapy for those with the skin subtypes.^{8,9,11}

The literature provides abundant information on the clinical or mutational spectrum and/or therapeutic aspects of MK-associated hereditary recurrent fever and PK. Therefore, this review does not extensively cover these themes already well illustrated elsewhere^{12,13} but rather uses the *MVK* gene, instead of the phenotype, as the entry point for the study. Indeed, MK-associated diseases represent a fascinating example of the accumulation of patients and genetic knowledge overturning our initial belief that mutations in this single gene lead to a systemic disease only.¹⁴ The aim was to provide an original historical and genetic overview of MK-associated diseases, to investigate the different genetic mechanisms underlying these disorders and their respective modes of inheritance, and to address questions that have been little explored until now, such as how a single gene can be the cause of so many different clinical pictures.

A long story in a nutshell

The first major step taken to result in our current understanding of MK-associated diseases was the identification of the human *MVK* gene, in 1992 (Fig. 1).¹ The link between *MVK* mutations and serious MA disease^{15,16} as well as its recessive transmission was established in 1997.¹⁷ Two years later, *MVK* variants were also linked to a milder, seemingly different, inflammatory disease called at the time "hyper-

IgD syndrome with periodic fever" (HIDS; OMIM 260920) because high amounts of IgD were found in the serum of most patients.^{2,18} It is now suggested that the term "HIDS" be dropped because elevated IgD level is neither specific nor constant and may well be a result rather than a cause of the disease.¹⁹ Phenotypes of intermediate severity have been described as well, revealing a continuum between the milder and severe disease, which thus represent the two extremes of the same genetic disorder.^{20,21}

Decreased enzyme activity has been demonstrated in all systemic forms, which therefore are strongly suggested to be grouped under the common umbrella "MK deficiency" (MKD), mild or severe.³ Recently, and somewhat unexpectedly, whole-exome sequencing in Chinese families with disseminated superficial actinic porokeratosis (DSAP) revealed that dominant pathogenic *MVK* variants were responsible for this purely dermatological disease.²² A second somatic hit was eventually discovered in the epidermis lesions of some DSAP patients.^{23,24}

The clinical and genetic landscape of MK-associated diseases

Systemic MK-associated diseases (MKDs) are recessively inherited

Although MKD was first described in the Netherlands and France, it is now recognized worldwide.^{12,25} The first symptoms occur early in life, generally in the first 2 years and include inflammatory symptoms such as fever and elevated acute-phase reactants.²⁶ Severe MKD is characterized by a massive and constitutive urinary excretion of metabolites accumulated upstream of the defective enzyme. The activity of the defective enzyme in cultured fibroblasts or leukocytes generally decreases to about 0.5%–2% of that of controls and is sometimes undetectable below 0.5%. Complete enzyme deficiency seems incompatible with life. The inflammatory symptoms are chronic and obscured by severe neurological signs (cerebellar ataxia, seizures, and mental, motor and growth retardation).^{15,20} Eye manifestations such as cataract, conjunctivitis and retinitis pigmentosa (RP) are common.^{27–30}

In the milder form, bouts of fever last 3–7 days and are typically associated with lymphadenopathy, diarrhea, vomiting and aphthosis. Flares are often triggered by vaccination, fatigue and stress. Mild MKD also features decreased enzyme activity and abnormal urinary excretion of mevalonic acid, but to a much lesser extent, with mevalonic aciduria detectable only during febrile attacks. When patients have sufficient residual enzyme activity, about 10% of that of controls, symptoms appear as attacks and tend to improve from adolescence onward.^{26,31,32}

Patients with atypical MKD features have been documented. Ocular symptoms may occur in both MKD subtypes, although they are more frequent in MKD-MA. Non-syndromic RP caused by *MVK* pathogenic variants has been described but almost always with some systemic MKD features.²⁹ Other atypical presentations that may result in delayed diagnosis include prominent liver³³ or cardiorespiratory disease,^{34,35} inflammatory bowel disease³⁶ or amyloid A amyloidosis.^{37–39}

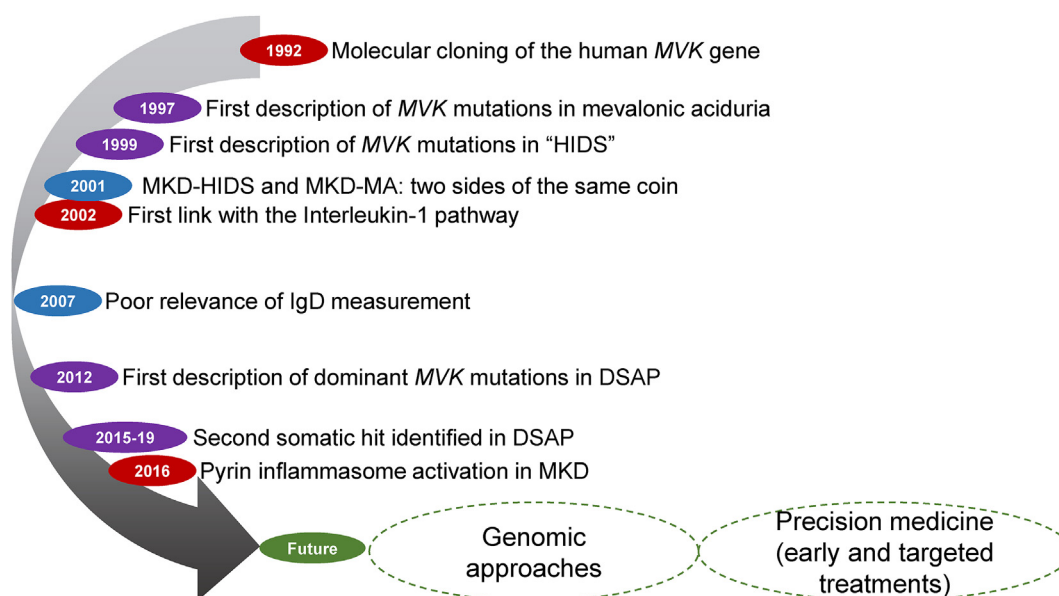


Figure 1 Key milestones of the clinical and genetic discoveries in mevalonate kinase deficiency (MK-associated diseases). The pathophysiology, clinical, and genetic steps are depicted in red, black and purple, respectively. MA: mevalonic aciduria; HIDS: hyper-IgD syndrome; DSAP: disseminated superficial actinic porokeratosis.

The pathophysiology of MKD, although better understood now, has not yet revealed why certain mutations lead to one phenotype rather than another. In brief (Fig. 2), pathogenic variants impede the folding and stability of MK and therefore its enzymatic activity, which results in

mevalonic acid accumulation and decreased production of downstream geranyl-geranyl-pyrophosphate, itself leading to defective prenylation.^{13,15,40} Defective prenylation in turn leads to inactivation of RhoA GTPase and subsequent defective activation of specific kinases, namely, the serine-

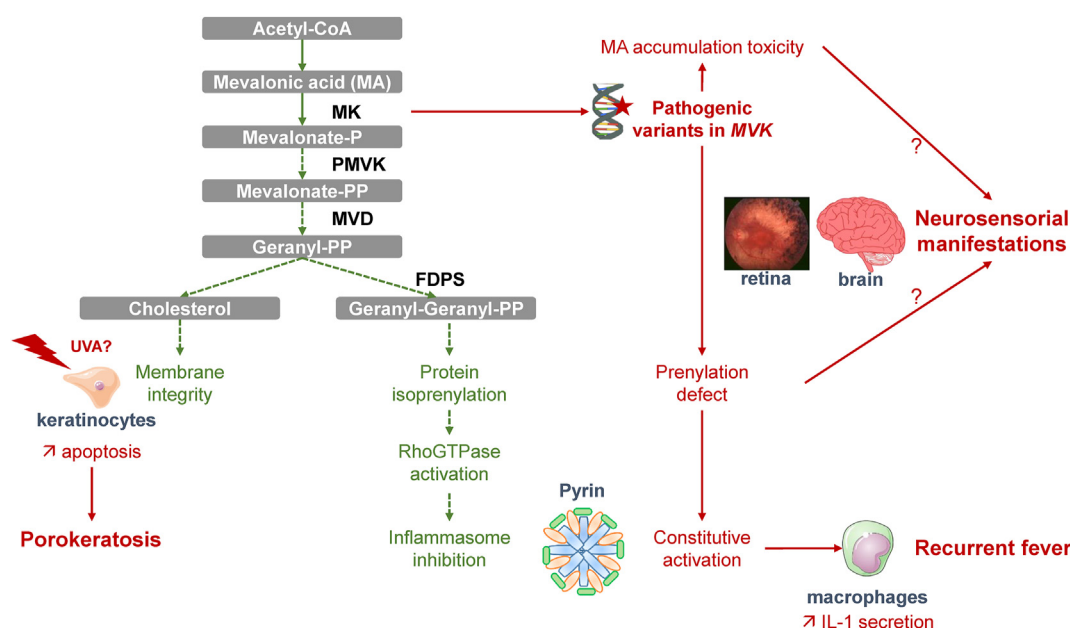


Figure 2 Schematic representation of the MK pathway and various consequences of mevalonate kinase gene (*MVK*) pathogenic variants for the different MK-associated disease subtypes. Pathogenic variants impede the folding and stability of MK and therefore its enzymatic activity, which results in mevalonic acid accumulation, decreased production of geranyl-geranyl-pyrophosphate, and ultimately activation of the pyrin inflammasome. Dashed arrows represent multiple steps in the pathway; green and red arrows and words depict normal MK (protein) pathway and pathogenic consequences of *MVK* (gene) mutations, respectively. PP: pyrophosphate; MVD: mevalonate diphosphodecarboxylase; PMVK: phosphomevalonate kinase; FDPs: farnesyl diphosphate synthase; IL-1: interleukin 1.

threonine kinases protein kinase N1 (PKN1) and PKN2.^{5,6} These latter bind and phosphorylate pyrin, blocking activation of the pyrin inflammasome.⁴¹ Inflammasomes are protein platforms that aggregate upon detection of a pro-inflammatory signal to activate secretion of potent pro-inflammatory cytokines such as IL-1.⁴² Constitutive activation of pyrin and NOD-like receptor family, pyrin domain containing 3 (NLRP3) inflammasomes has been established in two other hereditary recurrent fevers, familial Mediterranean fever (FMF) and NLRP3-associated autoinflammatory diseases, but remained unsuspected in MKD until recently. The inflammatory features of MKD are caused by excessive production of IL-1,⁴³ but the neurological or ocular symptoms seen in MKD-MA are less understood.^{29,44} Elevated mevalonic acid level could be toxic, or isoprenoids could be differentially regulated in the brain and retinal tissues (Fig. 2).

Patients with systemic MKD have two recessively inherited mutations, in the homozygous or the compound heterozygous state. More than 140 pathogenic MKD-associated variants are known and are registered in an online database dedicated to autoinflammatory mutations (<https://infevers.umai-montpellier.fr/web>).⁴⁵ Most variants (115/147; 72%) are substitutions, and others are small intragenic rearrangements, including three deletions of exons 2, 3 or 5. The first reported, p.(Ala334Thr), was repeatedly associated with MA and is therefore considered a severe variant.¹⁷ The most frequent variant, p.(Val377Ile), is derived from a founder effect.⁴⁶ Although this latter variant was mainly reported in the milder phenotype,⁴⁶ p.(Val377Ile) homozygosity was also reported in one asymptomatic individual⁴⁷ or was found combined with another mutation in some patients with severe manifestations such as amyloidosis.³⁹ Few studies of phenotype–genotype correlation are available,^{20,25,32,48,49} and larger series, taking into account not only the two *MVK* alleles but also possible modifiers and epigenetic factors, are needed.

Localized MK-associated diseases (PK) are dominantly inherited but recessively expressed following a second post-zygotic somatic event

In PK, the autoinflammatory manifestations are restricted to the skin. PK represents a group of dominantly inherited disorders with clinical and genetic heterogeneity,²³ mainly reported in people of Asian origin.⁵⁰ These are diseases of epidermal keratinization classified according to lesion type and location, with the disseminated DSAP form being the most common.⁵⁰ Keratotic lesions usually appear in adolescents and young adults and are mainly localized on sun-exposed skin areas, but new lesions may arise later in life, which explains the age-dependent penetrance and late onset of the disease.⁵²

In 2012, Zhang et al.²² identified for the first time *MVK* pathogenic variants in a series of both familial and sporadic cases of DSAP. Since then, several other DSAP patients with different germline *MVK* mutations have been reported.^{53–61} In 2015, and confirmed later, heterozygous germline mutations in genes encoding three other enzymes downstream

of MK in the mevalonate pathway (i.e., mevalonate diphosphodecarboxylase [*MVD*], phosphomevalonate kinase, and farnesyl diphosphate synthase) were also found to cause DSAP and other PK subtypes (Fig. 2).^{11,23,24,62,63} One patient had both *MVD* and *MVK* mutations.²³ As compared with patients with mutations in other genes of the MK pathway, those with *MVK* mutations generally showed the widest range of phenotypes in terms of both the number and size of lesions.²³

The molecular mechanisms underlying *MVK*-associated PK are diverse and appear to involve sequential genetic alterations (Fig. 3). The first hit corresponds to a germline pathogenic variant occurring in *MVK*. Both *de novo* and inherited point mutations and large deletions have been identified by using massively parallel sequencing and copy-number variation analysis. In the initial publication, no somatic variant was found in 5/82 patients with available keratinocytes isolated from skin lesions.²² However, 3 years later, Zhang et al. identified a somatic alteration in Chinese patients with PK. Using allelic expression imbalance and cDNA mutation analyses in pairwise lesions and neighboring normal-appearing tissues, these authors demonstrated significantly reduced expression of the wild-type allele in lesions. This decreased expression of the normal allele was attributed to gene conversion of the wild type to a mutated allele, resulting in an enhanced proportion of the initial mutation in lesional tissues.²³ The same group identified a second acquired *MVK* mutation in lesions of one patient with DSAP (F38) (i.e., a post-zygotic c.1003G>A transversion resulting from RNA-editing of the wild-type allele in addition to the c.1093T>A germline mutation).

More recently, Kubo et al.²⁴ detected a second hit in genes of the mevalonate pathway that occurred solely in cutaneous lesions of seven Japanese patients with DSAP. One of these patients acquired several different *MVK* pathogenic variants in the wild-type allele as demonstrated by PCR-based cloning and sequencing of genomic DNA extracted from different biopsies of the lesional epidermis. The somatic events included various transitions (mainly C>T or G>A) as well as loss of heterozygosity due to mitotic homologous recombination. The mutations differed from the germline variant found in blood cells, differed among the cutaneous lesions in a same patient, and were absent in healthy skin. The authors concluded that each skin lesion originated from a postnatal keratinocyte clone. The proportion of cells with genetic alteration among normal cells determined the mosaicism rate in the lesional tissues.

Altogether, these results support that a second, post-natal mutation of the wild type in the mevalonate pathway genes is required for PK to develop. The loss of the normal allele was the result of a gene conversion, RNA modification, or large deletion; a homologous chromosomal recombination; or an as yet unknown epigenetic mechanism (Fig. 3). The acquired genetic alteration transforms the native heterozygous genotype into a biallelic mutated genotype in the sick cells. Thus, the apparent dominant heredity with incomplete time-dependent penetrance observed in some patients in fact reflects a somatic recessive expression due to the late occurrence of a second post-zygotic loss-of-function mutation in keratinocytes. This concept mirrors the “two-hit” hypothesis of Knudson

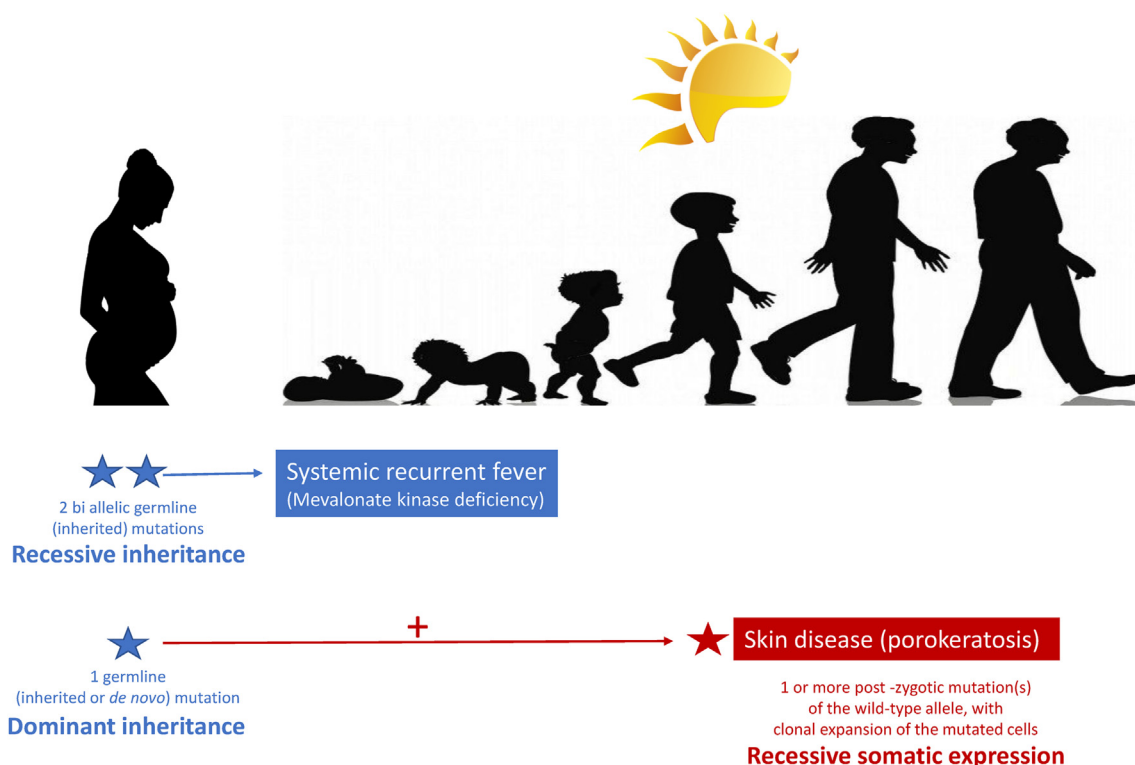


Figure 3 Genetic mechanisms involved in the different MK-associated disease subtypes. In the systemic form (mevalonate kinase deficiency [MKD]), two mutations are inherited from each parent (biallelic). In the localized skin porokeratosis form, the two (or more) mutations occur sequentially: the first is germline and dominantly inherited from one parent and the second is acquired later in life on the wild-type allele, presumably after UVA-mediated DNA alteration of epidermis cells in sun-exposed skin areas for some of the pathogenic variants.

for cancers and is consistent with the late onset of PK⁶⁴ and with the fact that certain PK patients are at greatest risk for malignant transformation.⁵¹

How are MK and PK linked? Functional studies are scarce, and levels of enzymatic activity or MA have not been measured (or at least reported) to directly prove the enzyme deficiency in PK patients. However, Liu et al.⁶¹ reported that three mutant MK proteins were less stable than the wild-type protein and that one mutation p.(Gly335Asp) resulted in the misfolding of the ATP binding domain. Zhu et al.⁵⁵ recently demonstrated decreased kinase activity, reduced cholesterol production, and increased apoptosis in cells transfected with two *MVK* pathogenic variants identified in DSAP patients. Shortage of isoprenoids could predispose patients to idiopathic inflammation of the skin because the mevalonate pathway is crucial in regulating calcium-induced keratinocyte differentiation and proliferation.^{22,23} Moreover, cholesterol, a product of the mevalonate pathway, is a key component of the extracellular lipid matrix that is important for the barrier function of the skin.⁶⁵ Overexpression of wild-type *MVK* has been associated with increased differentiation and decreased type A ultraviolet radiation (UVA)-induced apoptosis of keratinocytes.²² The authors of this study hypothesized that environmental (i.e., UVA) factors precipitate cell death in patients with impaired MK function, which may account for the clinical distribution of the skin lesions in DSAP patients.

Unanswered questions and perspectives

This review quickly summarizes nearly three decades of intensive studies on MK-associated diseases, focusing on the most recent breakthroughs that have emerged from modern genetics. The understanding, at least in part, of the genetic mechanisms involved in this group of disorders has had many twists and turns, which will have a major impact on both our fundamental knowledge and genetic counseling for patients. Indeed, for the first identified phenotype (i.e., systemic MKD), the recessive mode of transmission, a feature common to most enzyme disorders, has been clear and never discussed, but the latest data on the purely cutaneous PK form have contradicted this paradigm and have revealed an unexpected genetic mechanism, which involves, at least in some patients, an initial germline mutation followed by a second somatic event. This second post-zygotic hit explains why, despite the dominant heredity of PK, its clinical expression is somatically recessive in nature. Nevertheless, several issues remain that have been little explored until now.

One issue concerns a possible phenotype–genotype correlation between the *MVK* variants carried by patients and the MK-associated subtypes they ultimately display. The fact that a same mutation (e.g., p.Gly202Arg) can be found in both systemic and skin forms is puzzling.⁴⁵ Is there a link between the gene location or type of the mutation(s) and the clinical expression of the disease? Some general

trends have already been suggested, with p.(Val377Ile) and p.(Ala334Thr) associated with mild and severe or complicated systemic phenotypes, respectively, but exceptions exist.^{20,25,32} One clue has recently been provided by a multi-omic approach revealing the role of signal transducer and activator of transcription 1 (STAT1) as a modifier gene.⁶⁶ STAT1 is central in the regulatory pathway of inflammation mediated by IFN- γ . This study revealed a variant, p.(Arg241Gln) that was differentially expressed in two sisters homozygous for the p.(Val377Ile) genotype who had radically opposite (asymptomatic vs. strongly symptomatic) phenotypes. The authors suggested that the presence of both the homozygous MVK p.Val377Ile mutation and the heterozygous STAT1 p.R241Q mutation is required to elicit clinical symptoms.

With respect to PK, UVA, known to be an important trigger in the development of DSAP, seems to be an environmental factor that interacts significantly with genetic factors to influence the clinical expression of skin disease. Still, differential exposure to the sun cannot explain why patients with the systemic form do not have PK skin lesions. An extensive review of all published patients could help resolve the issue of MVK-related phenotype–genotype associations.

A second issue concerns the fact that, although the two loss-of-function variants are present in all cells, including keratinocytes, because they are inherited in systemic MKD, no porokeratotic lesions have ever been described in MKD patients. Regardless, such patients can display other types of rashes such as maculo-papules or urticaria in 39% and 15% of cases, respectively.²⁵ Conversely, the fact that PK patients or simple carriers, parents of MKD patients for example, do not have a fever is likely explained by the presence of half a dose of functional enzyme in heterozygous individuals, which is therefore compatible with normal inflammasome regulation.

A third issue is whether the two sequential molecular alterations (Knudson concept) represent a general pathophysiological mechanism of MVK-associated PK. Indeed, no second hit was detected in several patients with PK.^{22–24} Both technical limitations (low-rate mosaicism undetected) and yet-unknown epigenetic factors have been suggested.^{22–24,67}

Finally, does the two-hit hypothesis account for other cases of recessive autoinflammatory diseases such as FMF, with heterozygous patients sometimes being symptomatic?⁶⁸ This hypothesis has not been formally investigated, but two cases of acquired FMF have been reported.^{69,70}

In conclusion, the example of the genetic story of MVK-associated diseases retraces how amazing and various are the mechanisms underlying phenotypic heterogeneity. Large-scale integrated genomic technologies currently under development⁶⁶ will undoubtedly provide more surprises in the near future and bring new diagnostic and therapeutic solutions to patients in the context of precision medicine.

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