INFLAMMATION
Familial autoinflammation with neutrophilic dermatosis reveals a regulatory mechanism of pyrin activation

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Pyrin responds to pathogen signals and loss of cellular homeostasis by forming an inflammasome complex that drives the cleavage and secretion of interleukin-1β (IL-1β). Mutations in the B30.2/SPRY domain cause pathogen-independent activation of pyrin and are responsible for the autoinflammatory disease familial Mediterranean fever (FMF). We studied a family with a dominantly inherited autoinflammatory disease, distinct from FMF, characterized by childhood-onset recurrent episodes of neutrophilic dermatosis, fever, elevated acute-phase reactants, arthralgia, and myalgia/myositis. The disease was caused by a mutation in MEFV, the gene encoding pyrin (S242R). The mutation results in the loss of a 14-3-3 binding motif at phosphorylated S242, which was not perturbed by FMF mutations in the B30.2/SPRY domain. However, loss of both S242 phosphorylation and 14-3-3 binding was observed for bacterial effectors that activate the pyrin inflammasome, such as Clostridium difficile toxin B (TcdB). The S242R mutation thus recapitulated the effect of pathogen sensing, triggering inflammasome activation and IL-1β production. Successful therapy targeting IL-1β has been initiated in one patient, resolving pyrin-associated autoinflammation with neutrophilic dermatosis. This disease provides evidence that a guard-like mechanism of pyrin regulation, originally identified for Nod-like receptors in plant innate immunity, also exists in humans.

INTRODUCTION
Autoinflammatory diseases are characterized by recurrent episodes of fever with systemic and organ-specific inflammation, as well as uncontrolled activation of the innate immune response in the apparent absence of a infectious trigger (1). Familial Mediterranean fever (FMF, OMIM ID: 249100) is the most common of the monogenic autoinflammatory diseases, characterized by short (24 to 72 hours) episodes of fever associated with serositis, progressing to amyloidosis if untreated (2). FMF is an autosomal recessive disease caused by mutations in both alleles of the MEFV (Mediterranean fever) locus, which encodes the protein pyrin (3, 4). Normally, pyrin functions as an innate immune sensor that can trigger formation of an inflammasome, allowing the production of inflammatory mediators during infection. Pyrin activation is mediated by oligomerization via the central BCC (B-box and coiled-coil) domain, which results in signaling through the N-terminal PYD (pyrin) domain to the inflammasome. Studies in mice indicate that FMF mutations are a gain of function (5); however, it is unclear how FMF mutations, typically in the C-terminal B30.2/SPRY domain, initiate the activation process. As a potent checkpoint for the initiation of inflammation, the molecular mechanisms of pyrin regulation are critical and yet still poorly understood.

RESULTS
We studied a three-generation Belgian family of 22 individuals, of whom 12 developed autoinflammatory disease (Fig. 1A). The disease was characterized by neutrophilic dermatosis, childhood-onset recurrent episodes of fever lasting several weeks, increased levels of acute-phase reactants, arthralgia, and myalgia/myositis (Fig. 1B). The neutrophilic dermatosis comprised a spectrum of clinical manifestations, including severe acne, sterile skin abscesses, pyoderma gangrenosum, and neutrophilic
small-vessel vasculitis (Fig. 1, C and D). Pathological examination of affected skin showed a dense, predominantly neutrophilic, vascular, perivascular, and interstitial infiltrate (Fig. 1D). Serum cytokine analysis revealed elevated inflammatory mediators such as interleukin-1β (IL-1β), IL-6, and tumor necrosis factor-α (TNF-α) and cytokines induced by inflammation such as IL-1 receptor antagonist (IL-1Ra) (Fig. 1E and

Fig. 1. Clinical and immunological features of PAAND. (A) Dominant inheritance of PAAND over three generations, with complete penetrance observed for the S242R mutation (MEFV c.C726G, denoted CG). Trios indicated by red boxing were subjected to exome sequencing. (B) Twenty-two family members were assessed for neutrophilic dermatosis, recurrent childhood-onset fever, acute-phase reactants, arthralgia, myalgia, cardiomyopathy, anemia, pyogenic arthritis, and serositis across three generations (G1, G2, and G3). ND, not determined; "*", recent onset. (C) Typical macroscopic features (top: pyoderma lesions on the arm; bottom: pustular acne lesions on the face) (representative from patient II.7). (D) Histologic presentation of inflammatory infiltrate in the subcutis, involving subcutaneous vessels and extending into the surrounding panniculus and deep dermis. Scale bar, 500 μm. Higher magnification shows predominantly polymorphonuclear vascular and perivascular infiltrate (representative from patient II.7). Scale bar, 50 μm. (E) Serum levels of IL-1β, IL-1Ra, and IL-6 in patients during inflammatory episodes (n = 6) and healthy controls (n = 121), compared by t test. For healthy controls, box and whiskers represent ranges of 25 to 75% and 5 to 95%. For patients, mean and SD are shown. Patients with values >95th percentile of the healthy range are annotated as "elevated."
arginine substitution at position 242 (S242R) in the pyrin protein. Because of the genetic causation by pyrin mutation (see below) and the characteristic features of systemic inflammation and dermatosis, we termed this disease pyrin-associated autoinflammation with neutrophilic dermatosis (PAAND).

PAAND disease inheritance indicated a dominant mode of transmission (Fig. 1A). We therefore performed genome-wide linkage analysis on 20 family members, revealing a genome-wide significant LOD (logarithm of the odds ratio for linkage) score of 3.6 for a single region (1 to 6,283,321 on hg19) in chromosome 16 (Fig. S2). Exome sequencing was performed on two linked trios within the pedigree (I.4, I.5, and II.6; II.8, II.9, and III.3), and we subjected the 8720 variants within the linkage region to our filtering pipeline, identifying a single synonymous variant in RPL3L, and a single missense mutation in MEFV (rs104895127) (fig. S3). Inheritance of this mutation was directly tested in the remaining members of the pedigree, displaying complete segregation with disease (Fig. 1A). The identified mutation was a C-to-G substitution at c.726 in exon 2 of MEFV, which results in a serine-to-arginine substitution at position 242 (S242R) in the pyrin protein. MEFV mutations are associated with the development of FMF; however, the S242R mutation falls in a protein region distinct from that of typical FMF-associated mutations (fig. S3). Some less common variants in MEFV have been reported near S242; however, their pathogenicity has not yet been evaluated. A few cases of dominantly inherited FMF have been reported in the literature (6), but these are also not near S242R. Although not falling within a characterized protein domain, Ser242 is highly conserved across mammals (fig. S3), unlike some of the more typical FMF variants that naturally occur in other species (9). Despite the association of MEFV mutations with FMF, the disease described here with S242R mutation is clinically distinct from FMF, with the absence of serositis and amyloidosis even after lifelong disease, the presence of severe recurrent neutrophilic dermatosis, and bouts of fever lasting several weeks rather than days (Fig. 1B). Clinical cutaneous manifestations evoked a closer resemblance to PAPA (pyrogenic arthritis, pyoderma gangrenosum, and acne), which is caused by gain-of-function mutations in the pyrin-activating partner PSTPIP1, although here too it was distinct, with a lack of pyrogenic arthritis (Fig. 1B).

To further investigate the physiological result of pyrin S242R mutation, we searched for other occurrences outside the investigated family. The variant was only seen in 1 of 122,836 chromosomes in 65000 Exome Aggregation Consortium (ExAC 65000) (allele frequency = 8 × 10⁻⁶), in a Colombian male for whom clinical information was not available (1000 Genomes phase 1 database), and was absent in 1087 unrelated Belgian exomes (available on NGS-Logistics). Two other patients with the same mutation in MEFV had, however, been reported in the registry of mutations responsible for hereditary autoinflammatory diseases (InFevers, http://fmf.igh.cnrs.fr/ISSAID/infivers/) (11), and a third independent family was identified. The first was an individual from France with a heterozygous S242R mutation in the absence of any other coding MEFV mutations. Disease presentation was similar to the Belgian family, with recurrent inflammatory skin nodules associated with neutrophilic infiltrate and small-vessel vasculitis, fever, arthralgia/myalgia, and the absence of characteristic FMF symptoms, namely, serositis, and of amyloidosis (fig. S4A). The second individual was from Lebanon and carried the S242R mutation in conjunction with a known FMF variant M694V. New clinical analysis was not possible; however, retrospective assessment of clinical manifestations included periodic fevers of 3- to 5-day duration and joint pain, partially controlled by colchicine treatment, as can be seen in FMF. Additional symptoms included recurrent aphthous ulcers, prompting an initial diagnosis of Behçet disease, chronically elevated C-reactive protein (CRP), and transient skin rashes/nodules (fig. S4B). A third independent family was identified in the UK and also demonstrated dominant inheritance of the PAAND phenotype (fig. S4C). Overall, the combination of inheritance in a pedigree exceptional by size and confirmation of the mutation in three additional pedigrees with autoinflammatory features strongly support the S242R mutation in pyrin as a causative driver of PAAND.

The unique clinical presentation of the S242R mutations suggested a distinct regulatory mechanism of pyrin-mediated activation of the inflammasome. Therefore, we analyzed ASC (apoptosis-associated speck-like protein containing a CARD) speck formation, which assembles as an intense cytoplasmic aggregate in response to pyrin activation (12). Green fluorescent protein (GFP)–ASC was overexpressed in human embryonic kidney (HEK) 293T cells at a level causing minimal spontaneous speck formation. With this system, the coexpression of pyrin colocalized with ASC and led to increased triggered speck formation, which was further enhanced by the S242R mutation (Fig. 2A and fig. S5, A and B). Quantification of ASC specks was performed by flow cytometry, using mCherry-tagged pyrin and further controlling for cells with similar expression levels of wild-type versus S242R pyrin (Fig. 2B and fig. S5C). The elevated speck formation of the S242R mutation was not observed from nearby heterozygous variants (R202Q and G304R) recently found in two Japanese patients who developed Sweet syndrome (Fig. 2C) (13). Typical FMF variants such as M680I and M694V did not lead to a significant increase in ASC specks in this assay; however, when double-mutant S242R/M694V pyrin was expressed, a significant increase was observed over the S242R single-mutant protein (Fig. 2C).

Downstream of ASC, the S242R mutation resulted in caspase-1 being activated, with increased production of the p20 subunit (Fig. 2D and fig. S6A) and incorporation into the inflammasome “speck” (fig. S6, B and C). To measure IL-1β production, we used the human THP-1 monocytic cell line, with retroviral overexpression of wild-type or S242R pyrin. Lipopolysaccharide (LPS) was added for 24 hours to induce the synthesis of pro–IL-1β, and a significant increase in mature, secreted IL-1β was observed from cells expressing S242R pyrin compared to wild type (Fig. 2E and fig. S6D). This inflammasome activation was independent of NLRP3, as demonstrated by treatment with the NLRP3–specific inhibitor MCC950 (fig. S6E). CRISPR (clustered regularly interspaced short palindromic repeats)–mediated deletion of endogenous human pyrin (MEFV−/−) confirms that loss-of-function mutation of pyrin does not trigger inflammasome activation and that endogenous pyrin was not required for pyrin S242R activation. By contrast, deletion of endogenous caspase-1 (CASP1−/−) prevented IL-1β production due to pyrin S242R (Fig. 2E). Another known consequence of increased caspase-1 activation is pyroptotic cell death, which was also increased by S242R expression (Fig. 2F).

To more closely examine the molecular mechanism by which S242R regulates pyrin activity, we immunoprecipitated glutathione S-transferase (GST)–tagged wild-type or S242R mutant protein from HEK293T cells. Eluates were run out on polyacrylamide gels and developed with a sensitive fluorescent protein stain, revealing a band that was specifically bound to the wild-type and not to the mutant protein (fig. S7A). Mass
spectrometry (MS) analysis identified this band as containing the protein 14-3-3 (six of seven human isoforms; fig. S7B). This interaction was then confirmed directly by Western blot, using a pan−14-3-3 antibody (Fig. 3A). Previously, two isoforms of 14-3-3 [epsilon (ε) and tau (τ)] were observed to interact with pyrin, dependent on phosphorylation of S242 and S208 (14). In agreement with this, we could observe the phosphorylation of S242 in wild-type pyrin by MS (fig. S7C). Recently, bacterial effectors that target Rho guanosine triphosphatases (GTPases), such as Clostridium difficile toxin B (TcdB), were reported to trigger pyrin activation (15). We therefore tested whether their mechanism of action was similar to the S242R mutation, by releasing the inhibitory 14-3-3 protein, and this was indeed the case (Fig. 3B). To confirm the phosphorylation status of S242, we used an antibody that recognizes the pSer 14-3-3 consensus motif (R/K x x pS x P, where x is any amino acid). Although the motif surrounding S242 lacks the +2 proline, the antibody did detect a serine-phosphorylated form of pyrin, and this was destroyed by the S242R mutation (Fig. 3C and fig. S7D). TcdB treatment also has the same effect, because the pSer 14-3-3 motif in pyrin can no longer be observed after TcdB exposure (Fig. 3C). Together, these data reveal that nonphosphorylated S242, and removal of 14-3-3, is the mechanism by which inhibition of Rho GTPases and the S242R mutation both activate the innate immune receptor pyrin.

In contrast to the PAAND mutation, we found that FMF mutations M680I, M694V, and V726A had no appreciable effect on 14-3-3 binding or the pSer 14-3-3 motif in pyrin (fig. S8A). Moreover, we made a double-mutant pyrin, S208A/S242R, which cannot be phosphorylated at either proposed 14-3-3 binding site, and the FMF mutation M694V was still able to further enhance inflammasome activation (fig. S8B). Therefore, FMF mutations are likely to affect pyrin activity via a mechanism separate from 14-3-3 binding, such as a structural rearrangement in the protein that facilitates inflammasome formation after pyrin activation, allowing FMF mutations to enhance disease presentation (fig. S4B).
and ASC speck formation (Fig. 2C). Notably, the effect of S208 mutation on enhanced inflammasome activation was minor compared to S242R mutation, indicating a functional hierarchy of importance between these two phosphorylation sites (fig. S8B).

These functional studies suggest that the S242R mutation drives disease in patients by removing the 14-3-3 guard from pyrin, allowing excessive IL-1β production. To determine whether this process occurred in PAAND patients, we assessed mature IL-1β production by patient monocytes. Monocytes from healthy individuals did not produce significantly increased levels of mature IL-1β in response to a short duration of LPS treatment (Fig. 4A) because of the requirement of a second signal for inflammasome activation to allow cleavage of pro–IL-1β into the mature secreted form. The treatment of monocytes from healthy individuals with both LPS, to induce pro–IL-1β production, and TcdB, to activate pyrin and thus the inflammasome, resulted in high levels of mature IL-1β secretion (Fig. 4A). By contrast, patient monocytes demonstrated increased spontaneous inflammasome activation, because LPS alone was sufficient to induce a moderate level of mature IL-1β (Fig. 4A). No significant difference in IL-1β secretion was observed between healthy and patient monocytes after TcdB addition alone, indicating similar maximal activities between the dephosphorylated wild-type allele and the S242R allele (Fig. 4A).

The dermal manifestations and relatively modest systemic up-regulation of IL-1β (Fig. 1) suggested that the process of excessive IL-1β production may be elevated predominantly in the tissues. In support of this, immunofluorescence analysis demonstrated elevated levels of cleaved (active) caspase-1 and mature IL-1β in skin biopsies from patients with PAAND, whereas levels of pro–IL-1β remained normal (Fig. 4B). Finally, the central role of this process in the pathogenesis of PAAND was tested through a clinical trial with anakinra (recombinant IL-1Ra). One of the patients from the UK (G1) was treated with anakinra, after she failed a course of corticosteroids and methotrexate. Shortly after starting anakinra (100 mg daily), her clinical symptoms resolved and, for the first time after several years during which her inflammatory markers were regularly measured, her CRP levels dropped to within normal limits (<5 mg/ml) (Fig. 4C). These observations confirm that the consequences of constitutive pyrin inflammasome activation in PAAND are mediated by IL-1β and may be effectively controlled using IL-1β antagonism.

**DISCUSSION**

The identification of a dominantly inherited autoinflammatory disease with severe neutrophilic dermatosis driven by pyrin S242R mutation greatly expands the spectrum of pyrin inflammasome-related disorders. We propose PAAND as a distinct entity to FMF, due to differences both in clinical phenotype (with distinct characteristic features) and in the underlying molecular mechanism. The clinical presentation of distinct autoinflammatory diseases (FMF and PAAND) from different mutations in the same gene is reminiscent of the effect of NLRP3 mutation. NLRP3 mutations are associated with a spectrum of autoinflammatory diseases, including familial cold autoinflammatory syndrome, Muckle-Wells syndrome, and neonatal-onset multisystem inflammatory disease. Although distinct mutations are reportedly associated with each distinct clinical manifestation, a significant overlap is observed.

Our data suggest that clinical presentation of MEFV mutation segregates into clinical phenotype based on the molecular impact of mutation, with defective 14-3-3 binding resulting in PAAND, whereas...
Severe cutaneous abscesses (again, this patient had aspects of both FMF and PAAND, including the presence of both S242R and M694V was also previously observed in one individual with no apparent disease. However, the pedigree from Lebanon included one individual with both S242R and M694V, with a phenotype that did not fully replicate all PAAND features, and one individual with no apparent symptoms. Additional pedigrees were included from the UK, France, and Lebanon. Written informed consent was obtained from each participant, and the study was approved by the ethical committee of UZ Leuven (SS2653), the Espace Ethique/Université Saint Joseph–Liban, and the Comité de Protection des Personnes Ile de France 5 (Paris, France). Patients from the UK were consented under ethical approval for molecular genetics research studies obtained from South Yorkshire Research Ethics Committee (REC ref. no. 11/H1310/1).

Written informed consent to publish images of patients’ symptoms was obtained.

**MATERIALS AND METHODS**

**Human subjects**

We documented 22 family members from three generations, all of Flemish (Belgian) ancestry. The family included 12 patients (7 males and 5 females) with a clinical combination of childhood-onset episodes of myalgia/myositis and neutrophilic dermatosis (pyoderma gangrenosum, acne, and cutaneous small-vessel vasculitis), increasing in severity with age. The remaining 10 family members were healthy with no apparent symptoms. Additional pedigrees were included from the UK, France, and Lebanon. Written informed consent was obtained from each participant, and the study was approved by the ethical committee of UZ Leuven (SS2653), the Espace Ethique/Université Saint Joseph–Liban, and the Comité de Protection des Personnes Ile de France 5 (Paris, France). Patients from the UK were consented under ethical approval for molecular genetics research studies obtained from South Yorkshire Research Ethics Committee (REC ref. no. 11/H1310/1). Written informed consent to publish images of patients’ symptoms was obtained.
Genetic linkage and whole-exome sequence analysis

Whole-genome genotyping for 20 family members (affected and healthy) was performed using the Infinium Human Linkage-12 Genotyping BeadChip (Illumina). The BeadChip includes 6090 common single-nucleotide polymorphism markers with an average spacing of 441 kb and 0.58 cm across the genome. Linkage analysis was performed with the Merlin 1.1.2 software assuming autosomal dominant inheritance, full penetrance, no consanguinity, and a mutation allele frequency of 0.0001. Candidate genes were ranked according to expression in human peripheral blood leukocytes (23).

Whole-exome sequencing was performed for two trios, that is, the proband and both parents. This corresponds to three affected (family members III-3, I-4, II-6, and II-8) and two healthy family members (I-5 and II-9). Genomic DNA samples for whole-exome sequencing were prepared from heparinized peripheral blood using the FlexiGene kit (Qiagen). Exome sequence libraries were prepared using a SeqCap EZ Human Exome Library version 3.0 kit (Roche NimbleGen). Paired-end sequencing was performed on the Illumina HiSeq 2000 (Genomics Core Facility, University of Leuven, Belgium). Alignment of the sequence reads to the Human Reference Genome Build hg19, variant calling, and annotation were done with a pipeline on the basis of BWA (Burrows-Wheeler alignment) (24), GATK (Genome Analysis Toolkit) HaplotypeCaller (25, 26), and Annovar (27). The samples sequenced had an average of 98.2 (range, 57.7 to 127.8) million reads with 93% (range, 87.2 to 94.3) to 81% (range, 57.1 to 88.8) of targeted bases being covered at ≥10- or 30-fold, respectively. To select candidate genes, the following variants were considered: (i) compatible with the presumed mode of inheritance, that is, heterozygous in affected individuals and not present in healthy ones; (ii) overlapping a coding exon or within 2 base pairs of a splicing junction; (iii) not synonymous; (iv) not present in genomic duplications; and (v) allele frequency ≤0.01 in ExAC 65000; and (vi) absent in 1087 unrelated Belgian individuals without the disease. The pyrin variant identified, rs104895178, has merged into rs104895127. The variation was validated by Sanger sequencing of exon 2 on ABI 3730XL Genetic Analyzer (Applied Biosystems). Sequencing data were analyzed using the Staden package software.

Cytokine analysis

Plasma samples collected from patients and controls were stored at –80°C. Cytokine levels in plasma were quantified by an electrochemiluminescence immunoassay format using Meso Scale Discovery plates according to the manufacturer’s instructions. Data were analyzed with Discovery Workbench 4.0. Plasma IL-1Ra levels were measured using a human IL-1Ra ELISA (Thermo Fisher Scientific) according to the manufacturer’s instructions.

Monocytes were isolated from peripheral blood mononuclear cells of patients and controls using CD14 MicroBeads (Miltenyi Biotec) and cultured overnight. The next day, the cells were primed with UltraPure LPS (200 ng/ml) (InvivoGen) for 30 min before addition of TcdB. Cells were incubated for 20 hours before measuring IL-1β release by ELISA (R&D Systems) and cell death by propidium iodide (Sigma-Aldrich) staining at 1 μg/ml and flow cytometry. All cell lines were originally from the American Type Culture Collection and tested for mycoplasma contamination.

Western blotting and immunoprecipitation

For Western blotting, THP-1 monocytes were lysed in radioimmuno- precipitation assay buffer or supermatants were concentrated by methanol/ chloroform; for immunoprecipitation, HEK293T cells were lysed in NP-40 lysis buffer supplemented with a protease inhibitor (cOmplete protease inhibitor cocktail, Roche). Immunoprecipitation was performed using Gluthathione Sepharose 4B (GE Healthcare), with samples eluted in SDS-reducing sample buffer. Eluates were run on SDS polyacrylamide gels and treated as below for MS, or routine Western blotting was performed using the following antibodies: pyrin (Adipogen, rabbit α-pyrin, AL196), caspase-1 (Adipogen, mouse α-caspase-1 p20, Casper-1), IL-1β (R&D Systems, goat anti-IL-1β, AB-401-NA), GST (mouse α-GST, in-house), HA (Covance, mouse α-HA1.1), 14-3-3 (Santa Cruz Biotechnology, goat α-14-3-3, sc-629-G), α-actinin (Santa Cruz Biotechnology, goat α-actin, sc-1616), and pSer (Cell Signaling, mouse α-pSer 14-3-3 binding motif, no. 9601). Full Western blots are presented as fig. S9.

MS analysis

Samples were run on SDS polyacrylamide gels, stained using Sypro Ruby (Bio-Rad) and subsequently Coomassie G-250. Bands of interest were excised and subjected to reduction, alkylation, and LysargiNase digestion before MS analysis essentially as previously described (32). Raw files consisting of high-resolution MS/MS spectra were processed with MaxQuant (version 1.5.2.8). Extracted peak lists were searched against the human protein sequences obtained from the UniProt, SwissProt, and Ensembl knowledge bases. Cleavage specificity was set using [X/K/R] as a simple cleavage site rule for LysargiNase, allowing up to three missed cleavages.

Fluorescence microscopy and cytometry

GFP-ASC specks and caspase-1 were imaged in HEK293T cells using an Olympus IX70 inverted microscope with 20× objective. ASC specks were quantified by flow cytometry (33). In brief, cells gated based on viability and moderate mCherry (pyrin) expression were then selected for calculation of the percentage of cells containing an ASC speck defined by GFP-width versus GFP-area profiles. For colocalization, HEK293T cells were plated on chamber µ-slides (ibidi) and transfected with mCherry-pyrin constructs and ASC-GFP. Images were acquired with a Zeiss LSM 780 confocal microscope, using a 40× oil objective. Image channels were merged and converted to TIFF using Fiji software. Immunofluorescence was performed on skin sections fixed in 10%
neutral formalin, embedded in paraffin, and antigen-retrieved with pH 9.0
tris. Sections were blocked and stained with rabbit anti-IL-1 (ab2105,
Abcam), IL-1 propeptide (615417, R&D Systems), and rabbit anti-cleaved
caspase-1 (4199, Cell Signaling) before developing with donkey anti-rabbit
488 (A-21206, Life Technologies), goat anti-mouse immunoglobulin G1
546, and DAPI (D1306, Life Technologies). Sections were mounted and
cover-slipped using Fluoromount-G (SouthernBiotech) before images
were acquired with an LSM 510 Meta confocal microscope (Zeiss).

Statistical analysis
Sample sizes were limited by patient availability and are listed in figure
legends for each assay. Statistical tests used are detailed in the figure
legends. CRP was selected as the outcome of anakinra trial, as the only
quantitative measure of inflammation with historical data available from the
patient.

SUPPLEMENTARY MATERIALS
www.sciencetranslationalmedicine.org/cgi/content/full/8/332/332ra45/DC1
Fig. S1. Inflammatory manifestations of PAAND.
Fig. S2. Results from linkage analysis.
Fig. S3. Dominant inheritance of PAAND caused by mutation in pyrin, S242R.
Fig. S4. Inheritance of S242R mutations in additional pedigrees.
Fig. S5. Fluorescent microscopy of pyrin/ASC specks.
Fig. S6. Activation of caspase-1 and IL-1β by pyrin S242R.
Fig. S7. Proteomic analysis of pyrin interactions and phosphorylation.
Fig. S8. Functional effect of FMF mutations in context of pyrin S242R.
Fig. S9. Full Western blots.
Source data (Excel files)

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Familial autoinflammation with neutrophilic dermatosis reveals a regulatory mechanism of pyrin activation


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Guarding inflammation
The innate immune system is hard-wired to protect people from infection. However, mutations in these protective genes can lead to uncontrolled inflammation, resulting in autoinflammatory disease. Now, Masters et al. describe a family with an autoinflammatory disease caused by a previously unreported mutation in pyrin. This mutation disrupts pyrin regulation and mimics the effect of pathogen sensing by pyrin, leading to proinflammatory interleukin-1β (IL-1β) production. Indeed, targeting IL-1β resolved disease in one patient. These data suggest that pyrin is regulated through a guard-like mechanism, which guards against autoinflammation in humans.