In the 1970s, searches for the underlying mechanisms of fever led to the discovery of a class of molecules referred to initially as leukocytic pyrogens. These secreted proteins gained notoriety for their potency in triggering inflammation, and were ultimately given the names IL-1α and IL-1β upon their purification and cloning in the 1980s. Since that time, these two unique inflammatory mediators, both of which share a common IL-1 receptor (IL-1R1), became recognized as members of a much larger IL-1 family of cytokines, including 11 pro-inflammatory and anti-inflammatory cytokines that share a common inactive precursor structure and 10 multi-chain cytokine receptors. The IL-1 family also includes two decoy receptors, an inhibitory binding protein and two receptor antagonists, the most well-known of which is IL-1 receptor antagonist (IL-1RA), a natural inhibitor of IL-1R1 activation. The presence of many proteins within this family, often with synergistic or opposing functions, suggests important biological roles requiring tight regulation. Although the IL-1 family includes disease-relevant cytokines IL-18, IL-33 and IL-36 (reviewed in Mantovani et al.), for the purposes of this Review on autoinflammatory diseases, we focus on IL-1β, IL-1α and IL-1RA.

The most consistent clinical feature of autoinflammation mediated by IL-1, besides fever, is episodic or chronic systemic and/or tissue inflammation. Common areas that are affected include the skin and musculoskeletal system, although certain inflammatory sites are unique to specific IL-1-related disorders, such as serous membranes (pleura and peritoneum), the central nervous system (CNS) and conjunctiva. Patients also frequently complain of fatigue or malaise, symptoms that are commonly associated with chronic inflammation. In general, laboratory evaluation of patients, either during symptoms and sometimes between episodes, reveals elevated acute phase markers and neutrophilia in blood and tissue. Chronic systemic inflammation can lead to anaemia of chronic disease and tissue damage, such as amyloid A amyloidosis.

The first autoinflammatory disorders described were monogenic hereditary fever disorders caused by mutations in single genes encoding for inflammasome-related proteins. This classification has since been expanded to include other monogenic autoinflammatory diseases that result in IL-1-mediated inflammation and putative polygenic disorders that appear to be mediated by IL-1, as indicated by clinical response to IL-1-targeted therapies. The monogenic disorders are generally rare, with a prevalence ranging from 1 in millions to 1 in tens of thousands, although some of these diseases,
Key points

- IL-1α, IL-1β and IL-1RA are highly regulated inflammatory mediators involved in damage- and pathogen-associated molecular pattern (DAMPs and PAMPs), and cell death pathways.
- Patients with evidence of systemic inflammation without persistent infection or autoantibodies should raise suspicion of an IL-1-mediated autoinflammatory disorder.
- Rare monogenic and common polygenic diseases with neutrophilia and inflammation might respond to targeting the IL-1 pathway.
- Genetic testing confirms IL-1-driven autoinflammatory disorders, yet new disease phenotype–genotype correlations continue to be identified.
- IL-1-targeted therapies are highly effective and safe; new therapeutics focus on targets independent of IL-1 receptor binding, including NLRP3, caspases, IRAK4 and MK2.

particularly specific autosomal-recessive conditions, may be observed more frequently in specific isolated populations (such as familial Mediterranean fever (FMF)), likely because of selective infectious pressures occurring centuries ago. Founder mutations have been described as underpinning several of these disorders in different parts of the world; however, de novo mutations are often identified in many of the autosomal-dominant diseases. In this Review, we describe the molecular biology underlying IL-1-mediated inflammation and its role in autoinflammatory disorders. We highlight pertinent clinical features relevant to the practising rheumatologist, diagnostic and management considerations, and ongoing challenges faced by physicians and patients.

The complex biology of IL-1

IL-1β, often referred to as IL-1, is expressed primarily in myeloid cells (such as monocytes, macrophages and neutrophils) in a highly controlled fashion involving numerous regulatory mechanisms. IL-1β is expressed as an inactive precursor (pro-IL-1β) that is cleaved to its active form by several proteases, including caspase 1 and neutrophil- and microorganism-derived proteases. Caspase 1-dependent cleavage is driven primarily by activation of large intracellular multi-protein complexes known as inflammasomes, which consist of a sensor protein (such as NLRP3 or pyrin) and an adaptor protein ASC that oligomerize to form polymeric caspase 1 cleavage platforms. In addition to cleaving pro-IL-1β, caspase 1 cleaves and thereby activates gasdermin D, which allows it to form membrane pores enabling the release of IL-1β from the cell and mediating pyroptosis, a pro-inflammasome form of cell death (Fig. 2). IL-1β can also be released by a caspase 8 and gasdermin E-mediated lytic process, by mixed lineage kinase domain-like protein (MLKL)-mediated necroptosis, and likely by other mechanisms that remain to be elucidated. After IL-1β is released from the cell, the pro-inflammatory effects are exerted through binding to IL-1R1 on the same cell or nearby cells, activating intracellular signalling pathways involving IRAK4, MK2 and NF-κB that ultimately lead to further expression of inflammatory cytokines and inflammasome proteins. In this manner, IL-1β is a potent activator of its own expression and release, leading to an amplified, autoinflammatory response. Though less frequently recognized, IL-1α is constitutively expressed in all cells as an active pro-form and is unique in that it localizes to the nucleus, cytoplasm and cell membrane, where it has site-specific functions (Fig. 2). IL-1α shuttles between the nucleus and cytoplasm depending on cellular conditions, such as homeostasis or infection. In the nucleus, it binds to chromatin and functions as a transcription factor regulating cytokine expression downstream of NF-κB and AP-1 including IL-6 and IL-8, whereas in the cytoplasm it binds to mitochondrial cardiolipin to regulate NLRP3 inflammasome function. At the cell membrane, IL-1α can bind to and activate IL-1R1 on adjacent cells, or can be released in membrane-bound apoptotic bodies, leading to further local and systemic inflammation. Although IL-1α can be cleaved by several proteases, it does not require cleavage for biologic activity or for secretion. The major mechanism of IL-1α release appears to be through lytic cell death (pyroptosis or necroptosis); it is thus considered an alarmin that triggers sterile inflammation locally, such as during ischaemia. Moreover, through binding to IL-1R1, IL-1α acts as a potent driver of neutrophil recruitment in several tissues including the skin and lungs. Interestingly, IL-1β can bind to IL-1α and serve as a shuttle for its secretion, suggesting an additional layer of complexity to the regulation of IL-1 release.

A third important member of the IL-1 family, IL-1 receptor antagonist (IL-1RA), appears to be a predominantly anti-inflammatory cytokine. IL-1RA is expressed in all cells and tissues, and is induced by several inflammatory stimuli, as evidenced by high levels of IL-1RA in serum from patients with inflammatory diseases. It is a natural inhibitor of IL-1-mediated inflammation, acting by competitively binding to IL-1R1 and thereby preventing the binding of both IL-1α and IL-1β. Although this function of IL-1RA is well established, it has more than one isoform that may have additional functions. Moreover, IL-1RA has anti-apoptotic effects via an intracellular non-receptor-mediated mechanism. Although a complete understanding of IL-1RA biology remains to be reached, its clinical importance became evident with the discovery of patients with deficiency of IL-1RA (DIRA), discussed below, and the widespread utility of recombinant IL-1RA (anakinra) as a therapeutic for IL-1-driven autoinflammatory disease.

IL-1 and autoinflammation

Our appreciation of the complexity of IL-1 biology originally stemmed from attempts to understand the mechanisms of fever. As such, it is fitting that the rekindling of interest in IL-1-related mechanisms at the turn of the century occurred with the discovery of the molecular pathways underlying the pathology of several rare hereditary fever disorders. These conditions are all characterized by recurrent or chronic systemic and tissue inflammation combined with fever, rash and musculoskeletal symptoms. The identification of gene mutations responsible for these immunodysregulation diseases led to the introduction of an entirely new disease classification known as autoinflammation, which encompasses inflammatory disorders driven by innate immunity in the absence of high-titre autoantibodies or antigen-specific T lymphocytes. Each discovered
autoinflammatory disease revealed new mechanisms of dysregulated IL-1 biology, such as intrinsic unregulated inflammasome function, extrinsic mechanisms of inflammasome activation, and ineffective or absent IL-1 regulatory pathways. All of these mechanisms have broader implications for our understanding of innate immune regulation.

The innate immune system requires highly regulated mechanisms to prevent constitutive or uncontrolled responses that could harm the host, while allowing for rapid inflammatory responses that are selective for pathogen- and damage-associated molecular patterns (PAMPs and DAMPs) (Fig. 2). Inflammasome proteins are generally inactive under normal conditions but are activated by specific triggers, such as pathogen toxins. Uniquely, the NLRP3 inflammasome is activated by numerous triggers including ATP and several crystals. Gain-of-function mutations in MEFV and NLRP3, which encode pyrin and cryopyrin, respectively, result in intrinsic constitutive activation of the inflammasome, or a reduced threshold for its activation. This mechanism is responsible for the inflammatory phenotype observed in patients with FMF or cryopyrin-associated periodic syndromes (CAPS).

Other hereditary fever disorders, such as mevalonate kinase deficiency (MKD), TNF-associated periodic syndrome (TRAPS), and pyogenic arthritis, pyoderma gangrenosum and acne (PAPA) syndrome, are caused by mutations in genes that encode proteins with extrinsic effects on inflammasome activation, such as MVK, TNFRSF1A and PSTPIP1, rather than the inflammasome-encoding genes themselves. Although

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**Fig. 1** | **Timeline of key events leading to current understanding and treatment of IL-1-mediated autoinflammatory diseases.** First described as pyrexin in 1943, and subsequently termed lymphocyte-activating factor and leukocytic pyrogen in the 1970s, the introduction of interleukin nomenclature united these secreted macrophage products as IL-1. The timeline shows scientific advances in yellow, the first identification of a specific gene as the cause of a given autoinflammatory syndrome in grey, and the initial approval for IL-1-targeted therapies (with agency) in red. AOSD, adult-onset Still disease; CAPS, cryopyrin-associated periodic syndromes; DIARA, deficiency of IL1 receptor antagonist; FCAS, familial cold autoinflammatory syndrome; FMF, familial Mediterranean fever; GSDMD, gasdermin D; HIDS, hyper IgD syndrome; MKD, mevalonate kinase deficiency; MWS, Muckle–Wells syndrome; NOMID, neonatal-onset multisystem inflammatory disease; PAPA, pyogenic arthritis, pyoderma gangrenosum and acne syndrome; RA, rheumatoid arthritis; sjIA, systemic juvenile idiopathic arthritis; Syn, syndrome; TRAPS, tumour necrosis factor receptor-associated periodic syndrome; UK, Medicines and Healthcare Products Regulatory Agency of the United Kingdom.

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the mechanisms of autoinflammation extend beyond IL-1 to other cytokines and immune pathways, the study of IL-1-mediated autoimmune diseases remains an exciting field that is highly relevant to the practicing rheumatologist.

**Aetiology of autoinflammatory diseases**

The description and characterization of autoinflammatory diseases has long been linked to the identification of the underlying genetic and molecular basis for inflammation. Although phenotypically distinct...
patients are typically characterized first in the literature, the description of additional patients with shared genotypes has resulted in a much broader disease spectrum than was initially appreciated. Currently, more than 30 genetically defined autoinflammatory diseases, many with direct and indirect links to the IL-1 pathway, have been described and listed in the registry of hereditary autoinflammatory disorder mutations26–32.

Monogenic autoinflammatory disorders are caused by mutations in single genes related to control of inflammation (Table 1). Despite the rarity of these syndromes, their shared features of fever, arthralgias and rashes suggest that many will present to the rheumatology clinic. Mechanistically, monogenic autoinflammatory disorders lead to persistent activation of the NLRP3 or pyrin inflammasomes and subsequent activation of caspase 1, resulting in IL-1 release and autoinflammation. Other monogenic disorders, including those caused by mutations in TNFRSF1A, MVK, PSTPIP1 and CDC42, lead to IL-1 activation through pathways other than direct inflammasome activation, such as by the accumulation of intracellular stress triggers, enhanced binding to known intracellular sensors, or by affecting immune signalling pathways11–39. Autoinflammatory diseases caused by other members of the IL-1 cytokine family (Supplementary Table 1) are beyond the scope of this review.

**IL-1-driven autoinflammatory diseases**

**NLRP3 spectrum disease.** CAPS, also known as cryopyrinopathies, represent a disease continuum caused by gain-of-function mutations in NLRP3. These heterozygous mutations were first described in families with familial cold autoinflammatory syndrome and Muckle–Wells syndrome40, and then subsequently identified in patients with neonatal-onset multisystem inflammatory disease (NOMID; also known as chronic infantile neurological, cutaneous and articular syndrome (CINCA))41. Across the spectrum of severity, patients with CAPS share symptoms of recurrent fever, urticaria-like rash with neutrophilic infiltration, headaches, joint pain and conjunctivitis, as well as serological evidence of systemic inflammation. However, unique clinical features exist and a fairly consistent genotype-phenotype correlation can be used to define where patients fall on the CAPS disease spectrum41. Emphasis on the full spectrum is now reinforced by a newly proposed taxonomy that classifies these disorders as mild, moderate and severe NLRP3-associated autoinflammatory disease (NLRP3-AID). For historical context, and consistency with international drug approvals, this Review uses both the old and the new taxonomy.

At the mild end of the CAPS spectrum, patients with familial cold autoinflammatory syndrome (mild NLRP3-AID) experience brief flares, often less than 24 h in duration, induced by exposure to cold temperatures42. More severe clinical features on the CAPS spectrum include longer duration of episodes and stronger neurological symptoms due to CNS inflammation. In the moderate NLRP3-AID phenotype, Muckle–Wells syndrome, patients experience longer episodes of 2–3 days and often develop a progressive sensorineural hearing loss beginning in the first or second decade of life.

| Table 1 | Monogenic autoinflammatory diseases driven by IL-1 |
|---------|---------------------------------------------|
| **Disease** | **Gene** | **Presentation** | **Therapeutic target** |
| FCAS | NLRP3 | Cold urticaria, chills, conjunctivitis, myalgia/arthritis, fever | IL-1 |
| MWS | NLRP3 | Sensorineural hearing loss, uveitis, myalgia/arthritis, fever | IL-1 |
| NOMID | NLRP3 | CNS inflammation, cold urticaria, arthritis, myalgia/arthralgia, fever | IL-1 |
| FMF | MEFV | Neurological, cutaneous and articular syndrome, arthralgia, fever | IL-1 |
| PAAND | MEFV | Sterile skin abscesses, myalgia, myositis, rash, fever | IL-1, TNF |
| HIDS | MVK | Triggered by vaccination, abdominal pain, vomiting, rash, myalgia/arthralgia, fever | IL-1, TNF |
| MA | MVK | Developmental delay, FTT, dysmorphic features, recurrent fever | IL-1, TNF |
| TRAPS | TNFRSF1A | Painful centrifugal rash, periorbital oedema, recurrent fever, abdominal pain, headache, myalgia/arthralgia, fever | IL-1, TNF |
| DRA | IL1RN | Pustular rash, sterile osteomyelitis, periostitis, hepatosplenomegaly, fever | IL-1 |
| PAPA | PSTPIP1 | Pancytopenia, perinatal defects, facial dysmorphism, recurrent infection, rash, MAS/HLH, fever | IL-1 |
| Hz/Hc | PSTPIP1 | Rash, FTT, neutrophilic infiltration, neutropenia | IL-1, TNF |
| CDC42/NOCARH | CDC42 | Pancytopenia, perinatal defects, facial dysmorphism, recurrent infection, rash, MAS/HLH, fever | IL-1 |

CNs, central nervous system; DRA, deficiency of IL1 receptor antagonist; FCAS, familial cold autoinflammatory syndrome; FMF, familial Mediterranean fever; FTT, failure to thrive; HIDS, hyper IgD syndrome; Hz/Hc, hyperzincemia/hypercalprotectinemia; HLH, haemophagocytic lymphohistiocytosis; MA, mevalonic aciduria; MAS, macrophage activation syndrome; Mkd, mevalonate kinase deficiency; MWS, Muckle–Wells syndrome; NOCARH, neonatal-onset cytophenia with dysshamatoipoiesis, autoinflammation, rash, and HLH; NOMID, neonatal-onset multisystem inflammatory disease; PAAND, pyrin-associated autoinflammation with neutrophilic dermatitis; PAPA, pyogenic arthritis, pyoderma gangrenosum and acne; TNF, tumour necrosis factor; TRAPS, tumour necrosis factor receptor-associated periodic syndrome.
Neutrophilic dermatosis
Inflammatory skin disorder characterized by predominant neutrophilic infiltrate.

The most severe NLRP3-AID phenotype, NOMID, is characterized by nearly persistent systemic inflammation with additional neurological symptoms including chronic aseptic meningitis and cognitive impairment44,45. In these patients, increased intracranial pressure may lead to papillary oedema and optic disc atrophy46. Moreover, skeletal abnormalities, including frontal bossing and a distinctive distal femur overgrowth, are also typically observed46,47.

The constellation of symptoms, combined with frequent genetic testing, has also resulted in the identification of NLRP3 variants in other related syndromes. For example, somatic variants in NLRP3 have been described in some patients with Schnitzler syndrome48. CAPS-like phenotypes have also been observed in patients with variants in other autoinflammatory genes including NLRP12 (REF.49), NLRC4 (REF.50) and F12 (Factor XII)51. Thus, the phenotypic spectrum and genotypic aetiology of inflammasome-mediated diseases remain active areas of clinical and translational research.

MEFV spectrum disease. Mutations in MEFV, which encodes pyrin, underlie two distinct autoinflammatory syndromes: pyrin-associated autoinflammatory diseases-FMF and pyrin-associated autoinflammation with neutrophilic dermatosis (PAAND). FMF, likely the most well-known of the autoinflammatory syndromes, is characterized by discrete episodes of fever with serositis, synovitis and rash5. The most frequently reported symptom is abdominal pain, which can resemble an acute abdomen in presentation and on physical examination5. Although the disease phenotype is classically described as inherited in an autosomal-recessive fashion, there are increasing reports of patients with only one identifiable heterozygous MEFV mutation, with clear autosomal-dominant inheritance53,54. Murine studies have confirmed that MEFV mutations observed in patients are gain-of-function, consistent with inflammatory phenotypes in patients with a single identified heterozygous variant55. The most concerning consequence of uncontrolled inflammation in FMF is the development of systemic amyloid A (SAA) amyloidosis. Owing to the prevalence of MEFV mutations in certain parts of the world, it is also evident that specific variants may carry a greater risk of SAA amyloidosis dependent on the underlying genetic background. For example, patients with homozygosity for the M694V missense mutation may experience particularly severe disease with increased frequency of attacks, associated co-morbidities and reduced response to therapy56,57. The goal of therapy for any patient with FMF is reduction of inflammation and ultimately prevention of amyloidosis.

More recently, PAAND has been described as a distinct, autosomal-dominant syndrome caused by mutations in MEFV58,59. Affected patients experience recurrent inflammatory episodes with fever, neutrophilic dermatosis, arthralgia, myalgia and myositis beginning in early childhood. Elevation in serum acute-phase reactants is observed during inflammatory episodes. To date, two families with PAAND have been identified, exhibiting decreased 14-3-3 binding to pyrin, resulting in inflammasome activation and IL-1β and IL-18 secretion, as well as pyroptotic cell death58,59.

Deficiency of the IL-1 receptor antagonist. Autosomal-recessive missense mutations and large deletions in IL1RN were more recently described in patients with deficiency of the IL-1 receptor antagonist (IL-1RA), and are primarily due to founder effects60–62. In the case of deletion mutations, the size of the deletion may affect the phenotype, as has been observed in Puerto Rican patients with DIRA, where a 175-kb genomic deletion eliminates not only IL1RN, but also five IL-1-related genes63,64. In all cases, the result is an absent or truncated IL-1RA protein that is not secreted, and ultimately unable to inhibit IL-1α and IL-1β inflammatory responses. Uniquely, patients with DIRA have systemic inflammation with neutrophilia and elevations in serum inflammatory markers, although fever may be absent. Patients present near birth with a neutrophilic pustular rash that can be triggered by mechanical stress. Skin biopsy samples show neutrophilic infiltration of the dermis and epidermis, superficial folliculitis with pustule formation along hair follicles, acanthosis and hyperkeratosis65,66. Osteopenia with sterile lytic bone lesions, epiphyseal ballooning of the long bones and widening of the anterior rib ends, periostal reaction, joint swelling and fusion of the cervical vertebrae have also been described in the majority of patients60–63. Infants frequently demonstrate hypoxaemia and dyspnoea due to interstitial pneumonia, localized ground-glass opacities and areas of atelectasis or gastrointestinal reflux. Thrombosis has also been described.

IL-1-associated autoinflammatory diseases
TNF-associated periodic syndrome. TRAPS is an autosomal-dominant disease caused by mutations in the TNFRSF1A gene. Symptoms of TRAPS include long episodes of fever (>7 days), a tender centripetal migratory skin rash and abdominal pain. Periorbital oedema and musculoskeletal symptoms, especially myalgia of the lower extremities, have also been documented. The high severity of symptoms has resulted in the misdiagnosis of acute abdomen and unnecessary surgical intervention in some patients67. As in the cryopyrinopathies, the TRAPS phenotype severity spectrum is determined to some degree by the type of TNFRSF1A mutations68.

The mechanisms underlying the extended febrile episodes in TRAPS have long been subject to investigation. Initial studies suggested that mutations affecting amino acid residues critical for protein folding lead to a failure of TNF receptor shedding69,70, a cellular process that desensitizes cells to TNF action. However, more recent data support a role for NLRP3 inflammasome activation through a number of potential mechanisms, including intracellular accumulation of misfolded mutant TNF receptor protein, leading to elevated generation of reactive oxygen species, induced cell death or impaired autophagy71,72. This proposed mechanism is supported by the successful treatment of patients with TRAPS with IL-1-targeted therapy71,72.

MVK spectrum disease. MKD results from autosomal-recessive, loss-of-function mutations in the MVK gene encoding mevalonate kinase73,74. It is a disease spectrum, comprising hyperimmunoglobulinaemia D and periodic
fever syndrome (HIDS) at the mild end, and mevalonic aciduria at the severe end. Patients with the milder phenotype are characterized by early onset episodes of fever, rash (macular–papular, urticarial, nodular or petechial), abdominal pain, oral ulcers and adenopathy, lasting nearly a week on average. Those with the more severe mevalonic aciduria phenotype also have recurrent febrile episodes, but additionally present with severe developmental disabilities. Inflammatory flares in MKD may be triggered by routine immunization, infection or physical stress.

MVK mutations result in reduced or absent enzymatic activity of mevalonate kinase, the first enzyme in the 3-hydroxy-3-methylglutaryl (HMG)-CoA reductase pathway required for cholesterol and isoprenoid synthesis. Lack of mevalonate kinase function results in a shortage of isoprenoid lipid precursors, which are required to diminish the inflammatory response, even in response to mild stimuli. Consistently, in vitro studies of cells from patients with HIDS demonstrate a reduced ability to clear antioxidant stress, as well as mitochondrial dysfunction and deficiency in autophagy, all of which are implicated in inflammasome activation. More recent evidence in human and murine models suggests that the post-translational modification and subsequent attachment of isoprenoids to GTPases, including the Rho, Rac and Rap families of GTPases, are significantly affected by reduced mevalonate kinase activity, ultimately leading to increased IL-1β secretion from monocytes, likely in a pyrin inflammasome-dependent fashion. The successful use of IL-1-targeted therapy in HIDS is evidence that supports these pathogenic mechanisms.

PSTPIP1 spectrum disease. PAPA syndrome is an autosomal-dominant disorder resulting from gain-of-function mutations in PSTPIP1, which encodes proline-serine-threonine phosphatase-interacting protein 1 (PSTPIP1). Mechanistically, disease-associated variants result in hyperphosphorylation of PSTPIP1 and enhanced assembly of the pyrin inflammasome, with subsequent uncontrollable IL-1β release. Symptom onset is prior to 10 years of age, typically with the appearance of sterile pyogenic arthritis first, followed by dermatological features including pyoderma gangrenosum and acne.

Similar to the MEFV spectrum, mutations in PSTPIP1 result in a spectrum of disease beyond PAPA syndrome. Hyperzincaemia and hypercalprotectinaemia (also known as PSTPIP1-associated myeloid-related proteinemia inflammatory syndrome) is characterized by severe systemic and cutaneous inflammation, hepatosplenomegaly, arthritis, pancytopenia and failure to thrive owing to the accumulation of zinc. Dermatological symptoms share clinical and histopathological similarities to pyoderma gangrenosum and include vasculitis, furuncle-like ulcers, and eczematous and necrotic lesions that typically affect the lower limbs symmetrically, although eyelids may also be involved. In contrast to PAPA, the hyperzincaemia and hypercalprotectinaemia/PSTPIP1-associated myeloid-related proteinemia inflammatory phenotype has only been associated with a single amino acid change in PSTPIP1, altering the electrostatic potential and enhancing binding to pyrin.

CDC42-associated autoinflammatory disease. Recently, patients with NOMID-like disease have been reported in small cohorts, underpinned by mutations in CDC42, a small Rho-family GTPase that regulates intercellular adhesion, cytoskeleton formation, cell cycle and cell proliferation. Patients present near birth with growth restriction, recurrent febrile episodes, urticaria-like rashes, multiple cutaneous lesions, and hepatosplenomegaly. The presence of aseptic meningitis, papilloedema and facial dysmorphism, including mild frontal bossing, is reminiscent of NOMID presentation. Consistently, skin biopsy samples demonstrate perivascular lymphocytic and neutrophilic infiltration without vasculitis.

Although the first patients described with CDC42-associated autoinflammatory disease had a remarkable response to IL-1 blockade, more recent reports have noted that treatment with high dose IL-1 inhibition may not be sufficient to completely resolve symptoms, especially in the face of infection-driven macrophage activation syndrome. In contrast to other autoinflammatory diseases, patients described to date have elevations in IL-18, suggesting a role for either the pyrin or NLRC4 inflammasome in CDC42-mediated disease. Even though fewer than 10 cases have been described to date, the severity of disease phenotypes highlights the importance of an accurate diagnosis, and the name neonatal-onset cytophenia with dyshaematopoiesis, autoinflammation, rash and haemophagocytic lymphohistiocytosis syndrome has been proposed for CDC42-associated autoinflammatory diseases.

NLRP1 spectrum disease. Although NLRP1 was the first NOD-like receptor (NLR) reported to form an inflammasome, its genetic variation and relationship to disease remains unclear. Recently, patients carrying gain-of-function mutations in NLRP1 have been described, with some variability in phenotype depending on the mutated domain. Mutations in the PYRIN domain have been described in two families with autosomal-dominant corneal dyskeratosis, with distal mutations have been associated with systemic skin and mucosal symptoms (including respiratory or laryngeal papillomatosis, warts, exfoliation and plaque development), as well as inflammatory arthritis. Studies of patient serum, peripheral blood mononuclear cells and primary keratinocytes have demonstrated increased levels of IL-1β and IL-18, consistent with autoinflammatory inflammasome activation. At least two patients with NLRP1 spectrum disease have demonstrated some benefit to IL-1 blockade, although the response was not as uniform as for other disorders, perhaps suggesting a role for IL-18 in disease pathogenesis.

Of note, polymorphisms in NLRP1 have been linked to increased risk of several autoimmune disorders, including rheumatoid arthritis, psoriasis, vitiligo and type 1 diabetes (reviewed in Yu et al.). Further study is needed to determine the functional consequences of these genetic variants and whether IL-1-targeted therapy is appropriate.
**Polygenic autoinflammatory disorders**

The prominent features of fever, neutrophilia and elevated inflammatory markers observed in the monogenic autoinflammatory disorders, have fuelled a search for the genetic aetiologies behind related inflammatory disorders seen frequently by rheumatologists. Unlike monogenic diseases, many have multiple genetic aetiologies identified by case-control studies and genome-wide association studies of IL-1-related pathways, although it is often other shared clinical signs and symptoms and a lack of autoantibodies that suggest a mechanism related to IL-1-mediated disease pathways. This apparent IL-1 lack of autoantibodies that suggest a mechanism related to IL-1 has fuelled a search for activated inflammatory markers observed in the monogenic diseases, many with multiple genetic aetiologies. Nonetheless, targeted therapy may be effective in some patients with polygenic autoinflammatory disorders (Table 2), as shown in small cohorts and case reports. Equally evident is the recognition of partial responses to IL-1 blockade, consistent with the complex aetiology underlying these disorders. Nonetheless, the potential for pathway-specific therapy has been observed in some polygenic rheumatological disorders, further discussed below. Using a case-by-case approach, IL-1 blockade may be considered in these conditions.

**Chronic recurrent multifocal osteomyelitis.** Chronic recurrent multifocal osteomyelitis (CRMO) is characterized by chronic, relapsing sterile bone inflammation. The most commonly affected areas are the knee, ankle or wrist, although the vertebrae, pelvis and clavicle may also be affected. Several genes have been linked to CRMO, including FGR and FBLIM1. The recent identification of ILIRN variants in patients with CRMO-like presentations could have implications for treatment strategies, such as the use of anakinra, although this needs further exploration. The association of CRMO with congenital dyserythropoietic anaemia, known as Majeed syndrome, is caused by autosomal-recessive mutations in LPIN2. This rare syndrome has been reported in only three families to date.

### Table 2 | Summary of polygenic autoinflammatory diseases with a role for IL-1

| Disease          | Genetic associations linked with phenotype | Presentation                                      | Therapeutic targets |
|------------------|--------------------------------------------|--------------------------------------------------|---------------------|
| sJIA/AOSD        | Class II HLA locus, HDAC9, LACCC1/FAMIN     | Fever, arthritis, arthralgia, evanescent rash     | IL-1, IL-6, TNF, IL-18, IFN-γ |
| Schnitzler syndrome | NLRP3 (REF. 49)                                | Chronic urticaria, fever, arthralgia, lymphadenopathy, bone pain, skeletal hyperostosis | IL-1, TNF |
| Gout             | ADBR3, MTHFR, SLC2A11, SLC2A12, SLC2A9, SLC2A12, ABCG2, SLC17A1, SLC17A3, ALC16A9, GCGR, LRC16A, PDZK1, R3HDMI2, RREB1, ALDH16A1 (REF. 174), MAF (REF. 189), LRAF1 (REF. 174) | Recurrent flares of inflammatory arthritis, chronic arthropathy, tophaceous deposits, uric acid nephrolithiasis | IL-1 |
| Recurrent pericarditis | MEFV (REF. 190), HLA B14, DRB1*01, DQB1*0202 (REF. 190) | Pleuritic chest pain, pericardial rub, ECG changes, pericardial effusion | IL-1 |
| CRMO             | ILIRN (REF. 49)                               | Recurrent fever, arthritis, multifocal bone inflammation | IL-1, TNF |
| Hidradenitis suppurativa | PSENEN, NCSTN, PSEN1 (REF. 176)              | Inflammatory nodules, sinus tracts and open comedones in intertriginous areas | TNF, IL-1 |
| PASH             | MEFV, NOD2, NLRP3, ILIRN, PSTIP1, PSMB8 (REF. 185), NCSTN | HS lesions, pyoderma gangrenosum, acne           | IL-1, IL-18 |
| PAPASH           | PSTIP1 (REF. 185)                             | HS lesions, pyogenic sterile arthritis, pyoderma gangrenosum, acne | IL-1, TNF, IL-17A, IL-18 |
| PASS             | Unknown                                      | HS lesions, pyoderma gangrenosum, acne vulgaris, ankylosing spondylitis | IL-1, IL-18 |
| AHLE             | CFI (REF. 190)                               | Fever, neurological dysfunction, seizures, CSF pleocytosis | IL-1 |
| PFAPA            | IL12A, STAT4, IL10, CCR1–CCR3 (REF. 190), CARD9 (REF. 190), ALPK1 (REF. 190) | Recurrent fever with regular periodicity, aphthous stomatitis, exudative pharyngitis, cervical adenitis | IL-1 |
| Kawasaki disease | FCGR2A, BLK, CD40 (REFS 190,191), ITK, CASEF3 (REF. 191) | Fever, conjunctivitis, mucositis, rash, cervical lymphadenopathy, coronary artery dilatation | TNF, IL-1 |
| Behçet disease   | IL12A, IL10, STAT4, CCR1–CCR3, IL13R–IL12RB2, FUT2 (REFS 190,191), HLA-B15 (REF. 190) | Recurrent genital aphthae, uveitis or retinal vasculitis, nodular or papulopustular skin lesions, positive pathergy | IL-1, TNF |

**AHLR:** acute haemorrhagic leukocencephalitis; AOSD: adult onset Still disease; CNS: central nervous system; CRMO: chronic recurrent multifocal osteomyelitis; CSF: cerebrospinal fluid; ECG: electrocardiogram; HS: hidradenitis suppurativa; PAPASH: pyogenic arthritis, acne, pyoderma gangrenosum, and suppurative hidradenitis; PASH: pyoderma gangrenosum, acne, suppurative hidradenitis; PASS: pyoderma gangrenosum, acne and suppurative hidradenitis; PFAPA: periodic fever, aphthous stomatitis, pharyngitis, adenitis; sJIA: systemic juvenile idiopathic arthritis; TNF: tumour necrosis factor.
Hidradenitis suppurativa spectrum. Hidradenitis suppurativa is a chronic inflammatory disease affecting the epithelium of the hair follicles, whereby recurrent suppurative lesions lead to tissue destruction and fibrosis, specifically in the intertriginous areas. The pathogenesis of hidradenitis suppurativa is thought to be due to a combination of genetic and environmental factors, although a positive family history is present in nearly 30% of patients, suggesting an autosomal-dominant inheritance. More recently, hidradenitis suppurativa has been linked to a spectrum of autoinflammatory syndromes including pyoderma gangrenosum, acne, suppurative hidradenitis and ankylosing spondylitis (PASS); pyoderma gangrenosum, acne, and suppurative hidradenitis (PASH); pyogenic arthritis, pyoderma gangrenosum, acne and suppurative hidradenitis (PAPASH); and psoriatic arthritis, pyoderma gangrenosum, acne, suppurative hidradenitis (PsPASH). Although all of these syndromes share the sterile neutrophilic inflammatory lesions of hidradenitis suppurativa, the varied joint manifestations are consistent with a complex aetiology.

Other rheumatological diseases. More common rheumatological disorders with autoinflammatory features and IL-1 signatures include systemic juvenile idiopathic arthritis, adult-onset Still disease, recurrent pericarditis and microcystic arthropathies, such as gout and calcium pyrophosphate deposition disease. Despite complex genetics, the cardinal features of systemic juvenile idiopathic arthritis and adult-onset Still disease of fever, arthritis, rash and systemic inflammation are commonly also observed in the monogenic autoinflammatory diseases, and it is therefore not surprising that IL-1-targeted therapies are effective in these diseases. Recurrent pericarditis is a clinical feature observed in some patients with monogenic autoinflammatory disorders and may be the initial presentation (FMF with pericarditis). IL-1 pathway-targeted therapies, including colchicine and some IL-1 biologics, are currently approved for use in patients with pericarditis, with or without mutations in MEFV. Similarly, gout is characterized by recurrent attacks of fever and joint involvement that self-resolve. Given the role of the NLRP3 inflammasome in the recognition and perpetuation of monosodium urate- and calcium pyrophosphate-mediated inflammation, it is unsurprising that IL-1-targeted therapies have been used successfully in gout and in calcium pyrophosphate deposition disease. For other rheumatological diseases, including the seronegative spondyloarthropathies such as ankylosing spondylitis and psoriatic arthritis, IL-1 blockade has modest benefit in some patients, but may be considered in patients with refractory disease. Together, these disorders indicate that much remains to be learned regarding the genetic and environmental drivers of IL-1-mediated inflammation. Nonetheless, the availability of targeted therapies provides symptomatic relief for appropriate patients.

In paediatric patients, clinical presentations of fever and varied mucosal signs of inflammation have a broad differential diagnosis and can be seen in Kawasaki disease, periodic fever, aphthous stomatitis, pharyngitis, adenitis (PFAPA) syndrome and Behçet disease, and the latter two disorders are now postulated to exist on a genetic susceptibility spectrum (TABLE 2). PFAPA syndrome is characterized by recurrent episodes of fever, with set periodicity associated with non-infectious exudative pharyngitis, aphthous ulcers and cervical lymphadenopathy. Given the similarities with recurrent febrile episodes and mucosal inflammation, together with some shared genetic association, PFAPA has been proposed to be in a disease continuum with Behçet disease that is uniquely characterized by genitourinary ulcers and uveitis. Colchicine and IL-1 inhibitors have been used successfully in some patients on this syndromic spectrum, but incomplete responses suggest that mechanisms beyond IL-1 might play a role in disease pathogenesis. The self-limited Kawasaki disease shares several clinical features with other autoinflammatory diseases, including fever, inflammation of the skin, conjunctiva and joints, as well as pronounced neutrophilic inflammation. Interestingly, Kawasaki presentations might be the initial sign of autoinflammatory disorders. Although the aetiology of Kawasaki disease remains a source of active investigation, gene expression data and recent therapeutic success with IL-1 blockers support a major role for the IL-1 pathway in disease pathophysiology. Recent studies support a complex genetic aetiology underlying these disorders, but ongoing research offers opportunities for pharmaceutical intervention.

Management of autoinflammatory diseases

Diagnostic approach

Given the rarity of autoinflammatory diseases, many patients experience delays in diagnosis that could result in the development of complications due to uncontrolled inflammation. Failure to achieve a timely diagnosis may take a toll physically and socially on patients, further reducing quality of life. Lack of timely diagnosis is often due to poor awareness by physicians of autoinflammatory diseases; this is therefore an area in which rheumatologists can provide a patient-centred, comprehensive, team-based clinical experience and fill an unmet need.

Recognition of systemic signs, symptoms and flare patterns remains the cornerstone of autoinflammatory disease diagnosis. For many disorders, the differential diagnosis includes other autoinflammatory diseases, as well as systemic inflammatory disorders in the rheumatology realm. In general, non-specific inflammatory markers are elevated in autoinflammatory diseases, but there are few specific tests to define the diagnosis other than molecular genetic evaluations. Consequently, advances in sequencing techniques over the past few decades, including the availability of next generation approaches and commercialization of gene panels for disease classes, have improved the rate of diagnosis and dramatically increased the identification of variants in autoinflammatory disorder genes. However, although heralded as the gold standard, genetic testing is not without challenges. Increasing recognition of low penetrance variants, oligogenic or digenic presentations, somatic mosaicism and ‘mutation-negative’ patients, continues to be clinically challenging with regard to both disease diagnosis and long-term therapeutic management.
RhEumatology

for any given patient. This approach is aimed at minimizing or preventing the development of organ damage in patients, including hearing loss and vision loss, CNS inflammation and amyloidosis. It further underscores the need for ongoing evaluations, in order to make dose adjustments for weight gain and growth, changes in metabolism and the need for a multidisciplinary team to improve patient overall wellbeing.

Vaccination

Immunizations are an important component of ongoing health care. For most patients with autoinflammatory disorders, normal vaccination schedules are recommended; however, some may have exaggerated inflammatory responses to vaccines. This fact is clearly the case in patients with MKD, who characteristically (and often diagnostically) have disease flares triggered by vaccinations. In addition, patients with CAPS have been reported to have large local or systemic reactions to pneumococcal vaccinations, particularly the polysaccharide-conjugated form. This is customarily recommended for patients receiving most forms of immunosuppressive therapy, including the IL-1 inhibitors, to avoid live viral vaccines, although there is little evidence supporting this practice. There is also a theoretical risk of decreased vaccine effectiveness when patients are receiving some anti-inflammatory therapies, as it could affect the normal host immune response to vaccines. Optimization of vaccinations is therefore recommended prior to starting treatment when possible. Specifically, pneumococcal vaccines are particularly important for patients who will be on IL-1 inhibitors, as there is an increased risk of streptococcal disease.

With the exception of MKD, in which immunization is a known trigger for flares, most patients with IL-1-mediated autoinflammatory diseases tolerate routine vaccination, with no or mild symptoms. With specific regard to SARS-CoV-2 immunization, patients with autoinflammatory disorders studied to date have tolerated the SARS-CoV-2 vaccines well, with symptoms similar to those observed in large cohorts of healthy individuals, such as local arm pain, myalgia and headache. It is noteworthy, however, that although no flaring of their disease requiring hospitalization has been observed, two patients with NOMID receiving SARS-CoV-2 vaccines reported worsening of headache, leading to a temporary increase in the dose of their anti-IL-1 therapy. Clearly, longer follow-up and additional studies are needed to determine how to best balance SARS-CoV-2 vaccination and therapeutic dosing in this population.

Treatment of autoinflammatory diseases

Current therapies

The earliest and most widely utilized therapy for IL-1-mediated autoinflammatory disorders is colchicine, a plant-based medicine that was discovered...
serendipitously in the 1970s as a successful prophylactic treatment for patients with FMF. Colchicine seems to work by disrupting microtubules, which mediate intracellular organelle and vesicle movement, cytokine secretion, cell division and migration, and regulation of gene expression.

The identification of the pyrin inflammasome more than 20 years later, and the demonstration that colchicine blocks pyrin inflammasome activation, might explain why it is not as effective in the non-pyrin-mediated monogenic autoinflammatory disorders. However, there are some patients with FMF that do not respond to colchicine and others who cannot tolerate the gastrointestinal side effects or adhere adequately to a daily maintenance therapy. These patients usually respond to IL-1-targeted biologics, although the prevention of amyloidosis has not been as well established as it has with colchicine. It was the pharmacological development of the IL-1R antagonist, anakinra, initially for the treatment of sepsis, that enabled a direct means of evaluating the role of IL-1 in human diseases. This was a major advance, given that IL-1β levels are low, and IL-1α is typically below the level of detection, in the serum of patients with even the most severe autoinflammatory disease.

Although the sepsis trials failed, FDA approval for anakinra in rheumatoid arthritis in 2001 enabled investigators to test this antagonist in patients. Since then, two additional IL-1-targeted therapies have been developed and approved, including rilonacept, a recombinant IL-1R that binds to and inhibits IL-1α, IL-1β and IL-1RA, and canakinumab, a human monoclonal antibody that binds specifically to IL-1β. The different mechanisms of action of these three compounds should enable the unique functions of specific IL-1 family members in human disease to be understood, although the actual clinical experience has not necessarily been that clear, likely because of complex intra-IL-1 family member regulatory mechanisms. The clinical success of IL-1 blockade (targeting IL-1α or IL-1β, or both) provides the most convincing evidence of a role for IL-1 in autoinflammatory disease. To that end, empiric trials of anti-IL-1 therapies might be useful in treating patients with either genetically undefined autoinflammatory symptoms or with prominent non-infectious neutrophilia and elevated C-reactive protein or SAA.

All three approved IL-1-targeting therapies (Supplementary Table 2) have similar safety profiles, with the primary adverse effect being increased risk of non-opportunistic infections, which tend to be mild and can often be treated without withdrawing therapy. There are limited head-to-head trials of the different direct IL-1 inhibitors, and the general consensus among treating experts is that these biologic therapies have equivalent efficacy if used at appropriate doses with sufficient adherence. Nonetheless, there are clear pharmacodynamic differences between direct IL-1 inhibitors, as evidenced by the different dosing frequencies required. There is likely an advantage of anakinra and rilonacept over canakinumab in treating DIRA given that they block IL-1α and IL-1β, rather than IL-1β alone. This fact might also be the case with recurrent pericarditis and hidradenitis suppurativa.

Diseases with CNS inflammation might respond better to anakinra because of its smaller molecular size and resulting potential to penetrate the blood–brain barrier more efficiently than rilonacept and canakinumab. There are some patient groups that require higher or more frequent dosing, including patients with more severe autoinflammatory symptoms (such as NOMID and MKD), younger patients, and patients with atypical presentations or low penetrance mutations. Therapy during pregnancy remains an understudied cohort. Even for DIRA, which may be the most obvious example of links between genetics, disease and pathogenesis, patient-to-patient variability in therapeutic response still exists. Similarly, there is increasing recognition of a role for inflammasome-driven IL-18 in some diseases, such as PAPA syndrome, which may explain a lack of uniform responses to IL-1 blockade. Despite the general success in treating autoinflammation, the targeting of such a key mediator of inflammation, as well as the injectable nature of these therapeutics, has resulted in a search for additional therapies.

### New therapies targeting the IL-1 pathway

The rapid approval of rilonacept and canakinumab for CAPS caught the attention of the pharmaceutical industry. However, it was the CANTOS trial that confirmed that targeting the IL-1 pathway can have broader implications in more common diseases not traditionally considered as autoinflammatory, such as cardiovascular disease (Box 1). Although the initial focus has been on targeting single proteins in the IL-1 pathway (IL-1β, IL-1R and IL-1α) with predicted pharmacokinetic advantages, the current trend is to either combine protein targets (such as IL-1β and IL-18) or to develop small molecule inhibitors aimed at various steps in the IL-1 pathway (Table 3).

The most prevalent small molecule target for treating autoinflammatory diseases has been NLRP3, beginning with the early development of CRID3 (now known as MCC950); furthermore, multiple companies are currently developing, acquiring or studying their own NLRP3 inhibitors for various diseases, including some...
monogenic autoinflammatory disorders. Directly inhibiting a protein that is altered in a monogenic disease is theoretically a more targeted approach than use of the currently available biologics against downstream cytokines. Small molecules could also be designed to have better blood–brain barrier penetration, which would have implications for diseases with marked CNS inflammation. One study found that blocking NLRP3 decreased the risk of specific infections associated with broader IL-1 inhibitors. However, it is also conceivable that small molecules designed to bind to wild-type protein might have less avidity for mutant protein, thereby reducing clinical efficacy in patients.

Small molecule inhibitors with targets upstream and downstream of the inflammasome pathway are similarly under investigation. Two of the earliest pathway component inhibitors to be developed were compounds targeting P2X7, the receptor for ATP responsible for the potassium flux that activates NLRP3, and caspase 1, the common effector enzyme for inflammasomes. In addition to non-IL-1 pathway functions, P2X7 inhibitors might not affect intrinsically dysregulated inflammasomes that could be activated independently of P2X7. Specific caspase 1 inhibitors were developed and studied in several diseases, including autoinflammatory diseases, but pharmacodynamic issues and unexpected side effects resulted in their discontinuation. Finally, there are other drugs in development that block downstream IL-1R signalling, including IRAK4 and MK2. Inhibiting these targets could be very effective at blocking the effects of IL-1 in addition to other cytokine receptors, which could have advantages and disadvantages in terms of applicability to other diseases, and as yet unknown adverse effects.

**Ongoing challenges and future perspectives**

The identification of variants of unknown significance, which have neither been previously identified in disease cohorts nor rigorously studied, has introduced further challenges to our understanding of the complex genetic underpinnings of autoinflammatory diseases. The same is true for low penetrance variants, defined as likely benign variants that might be present at low frequency in the general population, but which may nonetheless contribute to IL-1-driven inflammation. Importantly, patients carrying these variants might present with clinical manifestations different from typical disease, and consequently might respond differently to therapeutics.

### Table 3 | Drugs in development over the past decade targeting the IL-1 pathway

| Drug | Target | Mechanism | Clinical Trial |
|------|--------|-----------|----------------|
| CE-224535 | P2X7 | Selective P2X(7) receptor antagonist | NCT00628095 |
| AZD9056 | P2X7 | Selective P2X(7) receptor antagonist | NCT00520572 |
| BMS-986299 | NLRP3 | Agonist | NCT03444753 |
| Dapansutril | NLRP3 | Small molecular inhibitor | NCT03595371 |
| IZD334 | NLRP3 | Small molecule inhibitor | NCT04086602 |
| ZYI1 | NLRP3 | Small molecule inhibitor | NCT04731324 |
| IZD174 | NLRP3 | Small molecule inhibitor, CNS penetrant | NCT04338997 |
| AC-201 | NLRP3 | Small molecule inhibitor | NCT02287818 |
| VX-765 | Caspase 1 | Small molecule inhibitor | NCT00205465 |
| Emricasan | Caspase 1 | Pan caspase inhibitor | NCT04803227 |
| Disulfiram | GSDMD | Gasdermin D inhibitor | NCT04485130 |
| Bermekimab | IL-1α | Anti-IL-1α monoclonal antibody | NCT03512275 |
| Gevokizumab | IL-1β | Anti-IL-1β monoclonal antibody | NCT01211977 |
| LY2189102 | IL-1β | Anti-IL-1β humanized monoclonal immunoglobulin G4 | NCT00380744 |
| CYT013-IL1bQb | IL-1β | Vaccine to IL-1β | NCT00924105 |
| Lutikizumab | IL-1α/IL-1β | Dual affinity monoclonal antibody to IL-1α/IL-1β | NCT01668511 |
| MAS825 | IL-1β/IL-18 | Bispecific IL-1β and IL-18 monoclonal antibody | NCT04641442 |
| sc-rAAV2.5IL-1Ra | IL-1R1 | Self-complementing, recombinant AAV carrying IL-1RA cDNA | NCT02790723 |
| EBI-005 | IL-1R1 | IL-1β and IL-1 receptor antagonist fusion protein | NCT04121442 |
| HL2351 | IL-1R1 | Human IL-1Ra-hyFc | NCT02853084 |
| MEDI8968 | IL-1R1 | Anti-IL-1R1 human monoclonal antibody | NCT01838499 |
| AMG108 | IL-1R1 | Anti-IL-1R1 monoclonal antibody | NCT00110942 |
| EBI-005 | IL-1R1 | Chimeric IL-1RA- IL-1β | NCT02082899 |
| KT-474 | IRAK4 | Oral heterobifunctional small molecule IRAK4 degrader | NCT04772885 |
| ATI-450 | MK2 | Oral small molecule MAPKAPK2 (MK2) inhibitor | NCT04524858 |

Identified based on search terms IL-1, inflammasome, NLRP3, at clinicaltrials.gov and Dinarello et al.
spectrum of disease ranging from asymptomatic, classic CAPS-associated features, or atypical presentation including gastrointestinal symptoms \(^{114}\) or more severe neurological symptoms \(^{151}\). In vitro studies suggest an intermediate phenotype with increased caspase 1 activity and IL-1β secretion compared with wild-type versions of these proteins, but with markedly less secretion than classic disease-causing mutations \(^{115}\). Patients respond at least partially to IL-1 blockade, suggesting that other mechanisms beyond IL-1 may contribute to disease pathogenesis. Nonetheless, in the small cohorts described to date, nearly half of patients experienced complete disease remission, suggesting that IL-1 inhibitors should be trialled in this unique population \(^{114}\).

In TRAPS, the most highly contested variant is traditionally known as R92Q (also known as R121Q), with a population frequency greater than 1%. Similar to the low penetrance variants in NLRP3, many R92Q carriers present with mild TRAPS symptoms with febrile episodes lasting approximately 1 week and in vitro studies demonstrating intermediate function, with reduced cytokine release from transfected cells, compared with cysteine mutations that have a more severe clinical phenotype \(^{132}\). Therapeutic responses to corticosteroids and colchicine have been described, as well as IL-1 blockade, with improvement in inflammatory episodes \(^{132,133}\).

The high prevalence of MEFV variants in certain populations adds to the challenges of variants of unknown significance and low penetrance mutations and their relationship to FMF. Variants such as E148Q have been described in up to 50% of different Jewish ethnic groups \(^{144}\). Although population frequencies of some pathogenic MEFV variants (such as V726A and M694V) are likely due to positive selection in Eastern Mediterranean populations, by conferring resistance to Yersinia pestis infection \(^{144}\), there might be an unknown heterozygote advantage to the high frequency of carriers \(^{154,155}\). However, the presence of low-penetrance heterozygous variants has led to complex clinical phenotypes including PFAPA syndrome presentations \(^{158}\), Behçet disease \(^{159}\) and risk of amyloidosis \(^{160}\). These varied clinical presentations further demonstrate the need to address patients on an individual basis.

Treating patients with autoinflammatory phenotypes with no identifiable disease-causing mutation is also challenging. Moreover, a subset of patients might have somatic mutations in an autoinflammatory gene that are restricted to a small (4%) proportion of a specific cell population. Such patients might have mild, delayed \(^{158,159}\) or atypical presentations \(^{158}\) or complete disease \(^{161}\). However, a growing number of patients have no clear genetic aetiology. Frequently referred to as syndrome of undifferentiated recurrent fever \(^{158,162}\) or undifferentiated systemic autoinflammatory disease \(^{163}\), choosing therapy for these patients is challenging. Fortunately, for patients with features of IL-1 presentations, treatment with IL-1 blockade and colchicine are successful in a significant proportion of patients \(^{158,163}\).

For patients with a poor response to IL-1 blockade due to complex genetics or lack of access to biologic therapeutics, haematopoietic stem cell transplantations have been attempted \(^{164}\). Although theoretically curative by replacing the haematopoietic cells that drive disease, the identification of patients with gain-of-function somatic mutations is instructive that only a small population of cells can drive disease phenotypes. The optimal treatment for these patients remains an active area of discussion.

Finally, the COVID-19 pandemic has brought systemic inflammatory disease into mainstream attention, with a variety of attempts to quell the cytokine storm and reduce morbidity and mortality. In the paediatric population, a multi-organ hyperinflammatory response to SARS-CoV-2 infection was described to occur weeks after infection, and is now known as multisystem inflammatory syndrome associated with coronavirus disease 2019 or paediatric multi-inflammatory syndrome temporally associated with COVID-19 (REFS \(^{165–167}\). Initially thought to be an atypical form of Kawasaki disease, it was quickly recognized that affected children were more likely to have signs of shock, gastrointestinal symptoms and coagulopathy \(^{165–167}\). Immunomodulatory therapy with intravenous immunoglobulin and glucocorticoids appears to be the most effective therapy to date, with a role for IL-1 blockade in refractory cases \(^{164–170}\).

The investigation of autoinflammatory disease and the inflammasome has involved the confluence of phenotyped patients and scientific studies at the bench. Over the past 30 years, such clinic–laboratory cooperation has been instrumental to the development of novel, targeted therapies for such patients \(^{171}\). The elucidation of inflammasome-mediated molecular pathways drove our understanding of basic innate immune mechanisms, solved several questions regarding the mechanisms of IL-1β release and led to successful therapy of many of the monogenic autoinflammatory diseases with biologics directed at the IL-1 pathway. A better understanding of the genetic and molecular mechanisms of autoinflammation and novel targeted therapies, such as biologics blocking more than IL-1 cytokine or small molecule inhibitors targeting upstream or downstream pathway elements, could lead to more effective personalized management of patients with this fascinating and sometimes frustrating family of immune dysregulation disorders.

Published online 21 June 2022
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Acknowledgements

We acknowledge the American Academy of Allergy, Asthma and Immunology Foundation (L.B.), and the NIH National Institute of Diabetes and Digestive and Kidney Diseases grant R01 DK133592-01 (L.B., I.M.H.) and R01 HL140898, R01 AI15586901 (H.M.H.) for invaluable funding.

Author contributions

The authors contributed equally to all aspects of the article.

Competing interests

L.B. is a site PI for Novartis, Inc H.M.H. is a consultant for Novartis and Kaniksa, H.M.H. has research collaborations with Regeneron, Inc., Jecure, Inc., Zomagen, Inc., Takeda.

Peer review information

Nature Reviews Rheumatology thanks E. Demirkaya, S. Masters, who co-reviewed with T. Reygaerts, and the other, anonymous, reviewer(s) for their contribution to the peer review of this work.

Publisher’s note

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Supplementary information

The online version contains supplementary material available at https://doi.org/10.1038/s41586-022-00797-1.