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The burgeoning field of innate immune-mediated disease and autoinflammation.

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The burgeoning field of innate immune-mediated disease and autoinflammation.

(short title “Innate immune-mediated disease and autoinflammation”)

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Abstract

Immune-mediated autoinflammatory diseases are occupying an increasingly prominent position among the pantheon of debilitating conditions that afflict mankind. This review focuses on some of the key developments which have occurred since the original description of autoinflammatory disease, in 1999, and focuses on underlying mechanisms that trigger autoinflammation. The monogenic
autoinflammatory disease range has expanded considerably during that time, and
now includes a broad spectrum of disorders, including relatively common conditions
such as cystic fibrosis and subsets of systemic lupus erythematosus. The innate
immune system also plays a key role in the pathogenesis of complex inflammatory
disorders. We have proposed a new nomenclature to accommodate the rapidly
increasing number of monogenic disorders, which predispose to either
autoinflammation or autoimmunity or, indeed, combinations of both. This new
terminology also encompasses a wide spectrum of genetically determined
autoinflammatory diseases, with variable clinical manifestations of immunodeficiency
and immune dysregulation/autoimmunity. We also explore some of the ramifications
of the breakthrough discovery of the physiologic role of pyrin and the search
for identifiable factors that may serve to trigger attacks of autoinflammation. The
evidence that pyrin, as part of the pyrin inflammasome, acts as a sensor of different
inactivating bacterial modification Rho GTPases, rather than directly interacting with
these microbial products, sets the stage for a better understanding of the role of
micro-organisms and infections in the autoinflammatory disorders. Finally, we
discuss some of the triggers of autoinflammation as well as potential therapeutic
interventions aimed at enhancing autophagy and proteasome degradation pathways.
Introduction

“La fixité du milieu intérieur est la condition de la vie libre et indépendante”


“The constancy of the internal environment is the condition for a free and independent life” in (Lessons on the physiological properties and pathological changes of body fluids)

Since the discovery of mutations in the pyrin protein as the cause of familial Mediterranean fever (FMF), in 1997 [1,2], a veritable treasure trove of susceptibility genes, with associated signalling pathways and potential disease mechanisms have been unearthed, which, in turn, has provided some essential guidelines on the most effective therapies for these debilitating conditions [3,4]. The term “autoinflammation” was first proposed by Dan Kastner, in 1999, [5] to differentiate between the pathogenesis of various hereditary periodic fever syndromes (HPFs), which are uncommon causes of recurrent fevers in clinical practice, and that of autoimmune diseases, characterized by the presence of autoantibodies and autoantigen-specific T and B cells. In particular, autoinflammation describes the type of inflammation mediated by the innate immune system [6], and the expression of pyrin in key cells of this system, including neutrophils, monocytes, dendritic cells, and serosal fibroblasts reflects this. Mutations in other central regulators of the innate immune system, as described below, have subsequently been found to underlie a range of other monogenic conditions as well as polygenic autoinflammatory diseases [7], such as Behcet’s and Crohn’s disease [8,9] (Figs. 1).
With relatively recent advances in massively parallel sequencing and wider use of this technology, we have witnessed the discovery of a succession of monogenic disorders, predisposing to either autoinflammation or autoimmunity or, indeed, combinations of both, further revealing the complex functioning of the human immune system [3,9,10]. These novel monogenic diseases may be of limited clinical impact, in the overall scheme of things, but they do represent true experiments of nature that continue to provide unique pathogenic insights into the hierarchy and levels of regulation of organ-specific immune defence responses. To quote directly from DJ Weatherall “if the severity of their phenotypes can be reduced by genetic or even environmental factors, it may be possible to reproduce these effects pharmacologically” [11].

Furthermore, functional studies of these disorders have generated many new and surprising biological concepts; for example, the discovery that autosomal recessive mutations of the mevalonate kinase gene (MVK), a key step in the cholesterol pathway, caused hyperimmunoglobulinemia D with periodic fever syndrome (HIDS) [12,13], has prompted closer examination of the broader interactions between inflammation and overall lipid signalling. The expanding list of novel autoinflammatory diseases and associated susceptibility genes has already been extensively covered [3,14]; in this review we propose to describe a selection of these diseases in order to illustrate some of the many unanticipated developments in this field, which have arisen as a result of the study of genetic causes of autoinflammation, often in quite rare conditions.

Interleukin 1 (IL-1)/NLRP3-mediated autoinflammatory diseases
In 2002, the late Jurg Tschopp’s laboratory reported on the identification of an intracellular complex called the NOD-like receptor family, pyrin domain containing 3 (NLRP3) inflammasome that triggered activation of inflammatory caspases, with pro-interleukin 1β (pro-IL-1β) processing and subsequent secretion of pro-inflammatory IL-1β [15] (Fig. 2). The genetic basis of familial cold autoinflammatory syndrome (FCAS) [16], Muckle-Wells syndrome (MWS) [17,18] and chronic infantile neurologic, cutaneous, articular syndrome/ neonatal-onset multisystem inflammatory disease (CINCA/NOMID) [19,20], were all found to be associated with mutations in the NLRP3/CIAS1 gene, and evidence that release of IL-1β was central to the pathogenesis of MWS came with the demonstrated efficacy of interleukin-1 receptor antagonist (IL-1Ra), anakinra, in 2 patients with MWS [21]. Collectively, the spectrum of these conditions soon became known as cryopyrin associated periodic syndrome (CAPS), reflecting a shared aetiopathogenesis (Table 1). Furthermore, as it quickly became apparent that this collection of conditions responded exquisitely to IL-1 blockade [21-23], so too it gradually emerged that IL-1 inhibition was also effective in other HPFs, like TNF receptor-associated periodic syndrome (TRAPS) [24], HIDS and FMF [25], although the response was less predictable in some cases. So it was proposed that caspase-1 activation with release of IL-1β was a pathway common to many autoinflammatory conditions; the mutated NLRP3 produces a gain of function, with lack of feedback inhibition, that results in constitutive activation of the NLRP3 inflammasome with IL-1β and IL-18 release [3,26]. The interleukin-1 receptor antagonist (IL-1Ra) provides a “biological brake” on inflammation driven by either endogenous IL-1α or IL-1β; deficiency of IL-1Ra (DIRA) [27] and deficiency of IL-36 receptor antagonist (IL-
36Ra) (DITRA) lead to unopposed IL-36 signalling and pustular psoriasis [28,29]
(Table 1).

A broad range of autoinflammatory diseases is currently being treated with IL-1 cytokine blockade, with marked attenuation of symptoms and disease progression. Canakinumab is a high affinity fully human monoclonal anti-human interleukin 1β antibody and rilonacept (IL-1 Trap) is a long-acting dimeric fusion protein IL-1 blocker. Clinical trials have been undertaken in CAPS, gouty arthritis, and systemic juvenile idiopathic arthritis (sJIA) [30-33]. There is a growing literature supporting the use of these agents in a wide spectrum of autoinflammatory conditions, including gout, Schnitzler syndrome, and Blau syndrome [34]. While multiple studies are ongoing, these agents have already been approved by for the treatment of CAPS and sJIA by a number of drug regulatory bodies.

Finally, somatic mosaicism has been reported in a number of autoinflammatory conditions. Since the first ever report of somatic mosaicism, in a Japanese patient with CINCA/NOMID in 2005 [35], it has subsequently been reported in several cases of CAPS, as well as FMF [36] and TRAPS [37].

**Interferon (IFN) mediated autoinflammatory diseases**

Aicardi-Goutières syndromes (AGS) constitute a collection of rare inflammatory disorders, associated with aberrant sensing of DNA/RNA, and usually affecting the brain and skin with clinical onset, most often, in early childhood. Since the initial description, of mutations in genes encoding the 3′→5′ exonuclease TREX1 in patients with AGS1 [38,39], in 2006, a total of seven AGS susceptibility genes have been identified to date, and this wide range of genetic mutations all lead to excessive interferon (IFN)-producing responses, known as type I interferonopathies.
A variety of disease mechanisms are involved: AGS 1-6 are of autosomal recessive inheritance and the AGS 7 patients have autosomal dominant gain-of-function mutations in the interferon induced with helicase C domain 1 (IFIH1) gene.

TREX1 is induced as part of the IFN-stimulatory DNA (ISD) response, an antiviral pathway that detects DNA, triggering immune activation through IRF3 [41]. Both TREX1 and SAMHD1 (AGS5) act as a negative regulators of the ISD response [42]. The genotype-phenotype spectrum of TREX1 is remarkably broad and complex [43]. Familial chilblain lupus, systemic lupus erythematosus (SLE) and retinal vasculopathy with cerebral leukodystrophy have all been associated with mutations in TREX1 [44], in addition to the AGS1 phenotype, which, in its more severe form, is characterized by intracranial calcifications, cerebral atrophy, leukodystrophy, chronic cerebrospinal fluid (CSF) lymphocytosis, increased CSF alpha-interferon (IFNα) and negative serologic investigations for prenatal infections.

Individuals with AGS7 also have severe neurologic impairment and immunological disease, particularly SLE [45]. However, clinical variability and non-penetrance are notable features of some AGS7 patients, despite the presence of IFN up-regulation (increased expression of type I IFN regulated genes, referred at as an IFN signature).

A variety of therapies have been used to treat the chronic excessive IFN production in AGS patients. Anti-inflammatory therapies, including Janus kinase (JAK) inhibitors, such as baricitinib and tofacitinib, and IFN pathway-blocking drugs, such as sifalimumab, have all been been used in AGS [46,47]. If AGS progresses to antibody-mediated disease then anti-B cell therapy, such as rituximab may be of benefit. Reverse transcriptase inhibitors (RTIs) are also being used to treat severely affected AGS patients and results are awaited with interest.
Apart from AGS there is a growing list of interferonopathies, due to gain-of-function mutations in genes such as the PSMB8, present in most patients with chronic atypical neutrophilic dermatosis with lipodystrophy and elevated temperature/proteasome-associated autoinflammatory syndrome (CANDLE/PRAAS) syndrome [48]. Liu et al. have demonstrated that mutations in the stimulator of interferon genes (STING) lead to constitutive STING–IFN-β pathway activation in patients with STING-associated vasculopathy with onset in infancy (SAVI) (STING is also known as transmembrane protein 173) [49]. A clinical trial aiming to assess the effect of JAK inhibitors, in SAVI and other related autoinflammatory syndromes, is currently ongoing (ClinicalTrials.gov number, NCT01724580).

It has been proposed that IL-1β and type I IFN are the main drivers, respectively, of autoinflammation and autoimmunity, acting as counterregulators of each other by activating specific metabolic signalling pathways to limit either innate or adaptive immune responses [50]. However, the fine details of such regulatory networks remain to be established.

**Autophagy in autoinflammation.**

Autophagy is emerging as a major pathway involved in the pathogenesis of autoinflammatory disease. The MVK mutation, and the subsequent depletion in isoprenoid synthesis, reduces functional autophagy in HIDS. However, this is not the only autoinflammatory disease where defective autophagy contributes to disease pathogenesis. Autophagy is a cellular process that maintains homeostasis by the clearance of redundant or damaged cellular components. There is a close relationship between autophagy and the inflammasomes, with evidence that
autophagy has a role in inhibiting the inflammasomes. This evidence not only suggests that autophagy clears inflammasome activators, such as ROS [51,52], mtDNA [53], HMGB1-DNA [54] and β-amyloid plaques in Alzheimer’s disease [55], but also clearance of the inflammasome itself [56]. Studies inhibiting autophagy observe increased NLRP3 inflammasome activation due to ROS accumulation [57].

The autophagy mechanism is a regulated process of ‘self-eating’ where the contents of entire organelles are recycled for other biological functions. Mutations in proteins such as NLRP3 or TNFR1, can overcome normal protein homeostatic mechanisms, resulting in autoinflammatory diseases, such as CAPS and TRAPS [58]. The inflammasomes are at the centre of the pathogenesis of autoinflammatory diseases and so the involvement of autophagy in these conditions may uncover new therapeutic targets. TRAPS is known to have inflammasome activation and individuals with TRAPS respond well to anakinra. Defective autophagy within TRAPS contributes to NF-κB signalling, ROS production and defective TNF-induced apoptosis [59,60]. Autophagy deficiency can be considered as a causal link between a pathological mutation and subsequent protein accumulation, inflammasome activation and cytokine secretion [59,60]. This is particularly relevant in inflammatory diseases with known protein misfolding and ER stress. One such example is cystic fibrosis (CF), which has been shown to have defective autophagy [61-63] and common infections of *Burkholderia cepacia complex (B. cenocepacia)*, which is able to inhibit autophagy as part of its infection machinery [64,65]. Autophagy and the inflammasomes go hand-in-hand, so in order to expose new disease mechanisms of innate immune driven diseases, both should be considered in tandem. On the other hand, genetic defects in the proteasome cause protein accumulation and proteasome dysfunction, which can trigger IFN-dependent autoinflammation. Loss-
of-function proteasome subunit mutations in CANDLE/PRAAS patients also promote type I IFN production [48,66].

**The unfolded protein response (UPR)**

Many different factors trigger activation of NLRP3; this is a 2-stage process requiring priming, usually via toll-like receptor (TLR) signalling, with a 2nd signal, typically intracellular calcium (Ca^{2+}) ion release, potassium (K^+) flux or intracellular reactive oxygen species (ROS). An ever-increasing number of molecules, in the form of whole pathogens, toxins, pathogen-associated molecular patterns (PAMPs), and DAMPs, are being found to trigger activation of the different inflammasomes, in particular the NLRP3 inflammasome (Fig. 3). It is most unlikely that these diverse agents bring about the activation by direct interactions with the intracellular NLRP3 receptor; instead, it is probable that NLRP3 is responding to generic cellular stress-signals induced by this variety of triggers. Among the cellular mechanisms that have evolved to maintain protein homeostasis include proteasome-mediated degradation of ubiquitinated proteins and the unfolded protein response (UPR). The UPR prevents protein overload in the secretory pathway and also prevents the spread of inflammation by degrading pro-inflammatory protein complexes, such as the NLRP3 inflammasome [58].

**Cystic Fibrosis (CF) as an autoinflammatory disease**

Cystic Fibrosis (CF) is a life-threatening autosomal recessive disorder of the lungs and digestive system [67,68]. The defective gene CFTR results in abnormalities in production and function of the CFTR protein, causing dysregulation of epithelial fluid transport and inflammation [69-72] and a predisposition to recurrent
pulmonary infections due to pathogens such as *Pseudomonas aeruginosa* (*P. aeruginosa*) and *B. cenocepacia*. Alterations in function and localisation of CFTR within leukocytes and epithelial tissues results in an exaggerated inflammatory response, with production of a wide spectrum of proinflammatory and chemotactic cytokines such as IL-17, IL-8, IL-6, IL-1β, IL-18, TNF, upregulation of TLRs and lipopolysaccharide (LPS) response [73]. The neutrophil is the predominant cell type infiltrating the CF lung, like a primary inflammatory response seen in acute infection, with inflammation in CF airways being driven by local environmental cells (macrophages and bronchial epithelial cells), rather than T cell derived lymphokines, as a systemic immune response. CF exhibits many hallmarks of an autoinflammatory condition [10], with infiltration by innate immune cells (neutrophils and macrophages) at target sites, and a paucity of autoantibodies or autoreactive T cells.

The physiological drive to autoinflammation in CF is due to CFTR dysfunction, which results in abnormal airway surface liquid (ASL) dehydration, reduced airway luminal pH, increased ASL glucose and hyperuricaemia [74-76]. These changes provide a milieu for activation of the NLRP3 inflammasome [77-79]. In human macrophages, IL-1β secretion and caspase-1 activation occurs following extracellular acidification, which is abolished following knockout of mRNA expression of NLRP3 receptor [79].

As well the physiological changes in epithelial ion transport, abnormal CFTR production, function and trafficking results in a state of hyperinflammation, associated with expansion of the endoplasmic reticulum (ER), located within the cytoplasm of cells, that inhibits ROS-mediated autophagy [61,80,81]. The most common mutation F508 results in a misfolded protein which is retained intracellularly and results in defective autophagy due to transglutaminase (TG2)-mediated...

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depletion of Beclin 1 and overactivation of protein kinase CK2 [81]. Normal autophagy activity suppresses activation of AIM2 and NLRP3 inflammasomes and helps regulate inflammation [82]. Reduced autophagy induces aberrant activation of the inflammasomes with accumulation of bacterial-containing phagosomes [82,83]. In CF murine airways and human macrophages, defective CFTR results in reduced levels of scaffold protein, CAV1, reduced inhibition of TLR4 signalling and hyperinflammation [84,85]. Similarly, studies in human CF broncho-epithelial cells show evidence of increased NLRP3 activation and defective NLRC4 activity, which can be inhibited by IL-1Ra (Fig. 3) [86]. Increased levels of ceramide appear to trigger the inflammasome protein complex, with upregulation of ASC protein, caspase-1 and increased production of IL-1β and IL-18 cytokines in the lungs of a CF mouse model [69].

Autoinflammation and Infection

The UPR, is activated in airways of patients by recurrent bacterial infections [30]. The ER stress responses involve atypical UPR induction, with lack of PERK-eIF2α response to *P. aeruginosa* [87]. This atypical UPR fails to resolve ER stress in CF and sensitises innate immunity to respond vigorously to microbial challenge. This persistent autoinflammatory response is associated with CF arthropathy in 9% of adults, which in some cases is associated with a fever and rash [88]. The complex relationship between inflammation, CFTR, innate immunity and infection is poorly understood and may be related to macrophage dysfunction, abnormal phagocytic killing of *P aeruginosa* [89] and impaired degranulation of antimicrobial proteins through defective activation of GTP-binding protein, Rab27a [90]. In addition, CFTR dysfunction results in an increase sensitivity to LPS (a major constituent of the outer
membrane of Gram-negative bacteria) stimulation, altered inflammatory signaling
due to abnormal neutrophil extracellular trap formation [89,91,92] and activation of
micro-RNAs (miRNAs) [93] and NF-κB. These bacteriae trigger the NLRP3
inflammasome through cytosolic receptors resulting in increased caspase 1 protease
(CASP-1) and IL1B and IL18 production (Fig. 3). Triggers of NLRP3 inflammasome
include the common CF lung pathogens *Staphylococcus aureus, Haemophilus influenza*, *P. aeruginosa, B cepacia complex, rhinovirus, influenza* and *Aspergillus fumigatus* [94-99].

Viruses activate inflammasome-mediated innate immunity through recognition
of viral RNA [100] by TLR7 and other triggers including altered ion flux with activation
of NLRP3 and NLRC5. *P. aeruginosa* and *Burkholderia cenocepacia* (*B. cenocepacia*) are two major pathogens which when isolated in sputum of patients
with CF are associated with clinical deterioration. *B. cenocepacia* is particularly
pathogenic and can result in acute clinical deterioration with uncontrolled
inflammation, necrotizing pneumonia and bacteraemia. *B. cenocepacia* accentuates
inflammation via upregulation of mononuclear cell IL-1β processing and inhibition of
autophagy [101,102]. Stimulation of autophagy with rapamycin in the CF lungs
mouse model reduces both inflammation and infection induced by *B. cepacia* [102].
LPS, L-Ala-γ-D-Glu-m-diaminopimelic acid (m-DAP), muramyl dipeptide (MDP)
present in gram-negative and some gram-positive bacteria are also involved
inactivation of the innate immune systems, though TLR and Nod-like receptor (NLR)
proteins. Furthermore, a number of chemicals can induce structural changes in LPS,
and subsequently modify the inflammatory response [103]. CF-associated ER stress
responses involve atypical UPR induction, with lack of PERK-eIF2α response to the
*P. aeruginosa* organism [87]. This shows that the atypical UPR fails to resolve ER
stress in CF and sensitises innate immunity to respond vigorously to microbial
challenge.

A key component of the UPR is the IRE1 enzyme, activated by ER stress. IRE1 induces conversion of the transcription factor XBP1u mRNA (unspliced) to spliced XBP1 (XBP1s), the active form. Martinon et al. proposed a pro-inflammatory role for IRE1, with TLR2 and TLR4 activating IRE1 to induce sXBP1 [104]. In macrophages, IRE1 activation exacerbates secretion of proinflammatory cytokines such as IL-6, TNF and IFNβ [105]. Furthermore, the effects of defective XBP1 functioning in autoinflammatory diseases may be augmented by concomitant defects that heighten cellular stress, including mitochondrial ROS or dysregulated microRNA regulation of XBP1 mediated inflammatory processes in TRAPS [106,107]. Thus, via both direct and indirect mechanisms, XBP1 dysregulation may be an important step in the cascade of intracellular events contributing to the pathogenesis of a number of autoinflammatory diseases. Indeed, there is also evidence of a UPR mediated by the XBP1s isoform in the airway epithelium of CF patients [108]. On the other hand, an in-vitro study from Italy shows that the degree of P. aeruginosa-dependent mitochondrial dysfunction is strictly dependent on defective expression of the CFTR channel and on a flagellin-activated TLR5-dependent pathway [109].

The NLRP3 inflammasome complex also senses mitochondrial dysfunction [110] and intracellular ROS is a crucial element for inflammasome activation. Anakinra reduced endotoxin-induced airway inflammation in healthy volunteers [111], so we postulate that spontaneous NLRP3 inflammasome activation occurs in in CF patients [112]. Recent studies have linked IRE1 to NLRP3 activation [113] and have also shown that XBP1 modulates innate immune responses of alveolar macrophages in CF patients [114]. IL-1 and the NLRP3 inflammasome activation
cause arthropathy and the IRE1/XBP1 axis has been implicated in synovial macrophages and fibroblasts of RA patients [107,115].

One of the unique features of CF as an autoinflammatory disease is that it is the only such condition to have a “laboratory proven” association with bacterial infections, including *P. aeruginosa* and *B. cenocepacia*. The NLRP3 and NLRC4 inflammasomes serve different functions in regulating inflammatory responses in mice and humans with CF (Fig. 4). While both NLRP3 and NLRC4 inflammasomes contribute to pathogen clearance, NLRP3 contributes to a greater extent than NLRC4 to deleterious inflammatory responses in CF and correlates with defective NLRC4-dependent IL-1Ra production. Also IL-1 blockade markedly reduces inflammasome-dependent inflammation in murine and human CF [116].

**Metabolic/mitochondrial mechanisms of autoinflammation.**

The relationship between inflammation and metabolism constitutes a delicate balance, with pathways from both systems converging to preserve the “milieu interieur” of the cell. This balance is maintained by short-term adaptive measures to keep these systems in check, but there may be a detrimental outcome when one arm becomes overactive and suppresses the other in the longer term. HIDS is a classic example of a monogenic autoinflammatory disease, with a metabolic defect at its core. This disease is caused by two mutations in the mevalonate kinase (MVK) gene [12,13,117] and presents with increased excretion of urinary mevalonic acid and raised immunoglobulin (Ig)-D and IgA levels in the serum [118]. Symptoms are often neurological in nature with increased mental retardation, ataxia, seizures and ocular problems. Fevers usually last around 5 days and are often triggered by traumas, illnesses or vaccine reactions. Although not consistently successful IL-1
antagonists are the most effective treatment for HIDS, with steroids having limited
efficacy [118,119]. The mutated MVK gene translates into reduced levels of the
enzyme mevalonate kinase, which normally converts mevalonic acid into mevalonate
-5-phosphate, an intermediate in isoprenoid and sterol synthesis. The exact
pathogenic molecular mechanism in HIDS is not clear but recent publications,
describing the pyrin inflammasome and its detection of bacterial modifications of Rho
GTPases, are promising avenues of exploration, as the causal biochemical
deficiency of isoprenoid synthesis in HIDS reduces RhoA prenylation [120]. As IL-1
antagonists, such as anakinra and canakinumab, are able to reduce fever frequency
and severity, the NLRP3 inflammasome is a key pathway of interest although it is not
the only possible source of IL-1β [121]. Research advances into how the
inflammasomes are controlled by ROS and autophagy, and their links to Rho
GTPase prenylation, also offer significant insights into the precise metabolic and
mitochondrial mechanisms of autoinflammatory disease.

Recently, Celsi et al. described an increase in NLRP3 activity in a HIDS
mouse cell model, using siRNA mvk silencing, when cells are treated with LPS and
lovastatin, a statin drug used to lower cholesterol [122]. However, complete
knockdown of mvk did not induce an increase in NLRP3 activity. This lead to the
conclusion that increased mutated mvk protein levels may trigger NLRP3 activity by
initiating the UPR due to protein accumulation. This hypothesis is supported by a
HIDS THP-1 macrophage cell line model [123]. This cell model produced increased
IL-1β and IL-18 levels, as well as an altered redox state. An important role for this
altered redox state was revealed as it was associated with increased mitochondrial
membrane potential, increased mitochondrial damage and increased mtDNA in the
cytosol, all linked to a defective autophagy pathway. Autophagy would ordinarily be
activated in the situation of an altered redox state to clear defective mitochondria and reduce ROS-dependent damage; however, in this cell model, autophagy was found to be defective. The mutations in MVK, with subsequent reduction in isoprenoid synthesis, causes reduced prenylation of small GTPases, which are key upstream proteins involved in autophagosome formation [123]. The authors suggest a model whereby defective autophagy, due to reduced prenylation of small GTPases, occurs upstream of increased mitochondrial damage and the increased ROS, in turn, activates the NLRP3 inflammasome [124]. Interestingly, when these small GTPases, specifically the Rho family, become modified they trigger the pyrin inflammasome [125]. The link between HIDS and reduced prenylation of Rho GTPases activating the pyrin inflammasome has been suggested to offer an effective therapeutic target [120]. RhoA activates PKN1 and PKN2 serine threonine kinases, which in turn phosphorylate pyrin. Phosphorylated pyrin is bound to 14-3-3 proteins that restrict pyrin from forming its inflammasome. Arachidonic acid is a known activator of PKN kinases and is a potential future therapeutic option for innate immune-mediated inflammation. Therefore, changes in post-translational modifications of Rho GTPases, in diseases such as HIDS or FMF, produce a reduced pyrin inhibitory capacity as well as defects in autophagy. In addition, autophagy has been shown to not only degrade ROS and mitochondrial debris in the cytosol, but also targets the NLRP3 inflammasome and pro-IL-1β for autophagosomal degradation [126,127]. Further evidence for disruption in metabolic pathways triggering the inflammasomes exists with hexokinase. Hexokinase is a glycolytic enzyme located on mitochondrial membranes. When inhibited, hexokinase dissociates from the membrane and allows release of mitochondrial DNA, activating the NLRP3 inflammasome [128]. Metabolic conditions in which hexokinase function is impaired cause NLRP3 activation.
Bacterial peptidoglycan-derived N-acetylglucosamine is detected by mitochondrial membrane-bound hexokinase, causing membrane dissociation and NLRP3 activation [129].

The UPR, metabolic pathways and associated therapies

The interplay between various metabolic pathways and the UPR has raised the possibility that key points in specific metabolic pathways could be targeted in autoinflammatory diseases. XBP1s acts a transcriptional activator of the hexosamine biosynthetic (HBP) pathway [130]; the UPR-HBP axis is triggered in a variety of stress conditions, including ischemia-reperfusion (I/R) injury, where stimulation of Xbp1s induces cardio-protection by induction of HBP. Ischemic accumulation of succinate has been shown to control reperfusion injury through mtROS [131]. Therefore the prevention of succinate accumulation could be a therapeutic goal in a range of autoinflammatory diseases that are resistant to standard therapies.

The rapid advances in the pathogenesis of autoinflammatory diseases and recognition that altered protein homeostasis contribute an innate immune component to many common diseases, underlines the unmet need for novel therapies for these conditions. For such therapies to be effective they would need to prevent protein accumulation, suppress ROS generation, and enhance of clearance mechanisms thereby preventing the development of (auto)inflammation. Therapies that succeed in augmenting the UPR could prove to be highly beneficial, as protein misfolding within the ER leads to activation of the UPR, with associated inflammation and increased disease severity. Anti-oxidants could be prescribed as adjunct therapies for diseases with aberrant ROS production and oxidative stress, like TRAPS [131].
Since both autophagy and proteasome degradation have anti-inflammatory properties, possible therapeutic interventions will be directed towards enhancing these pathways to effectively reduce NLRP3 activation [132]. Small molecules that block the NLRP3 inflammasome and related signalling pathways have recently shown promise in pre-clinical studies [133-135]. Clinical trials of agents that modulate proteotoxic stress and deactivate the inflammasome(s), combined with traditional therapies, such as IL-1 antagonists, will provide new insights into the connections between protein homeostasis and autoinflammation.


The physiologic role of pyrin

The discovery of the physiologic role of pyrin by Feng Shao’s group represents a major advance in the field of autoinflammation [125,136]. The raison d’etre of the innate immune system is to protect the population from infection (Fig. 2); however, mutations in these protective genes can also lead to autoinflammatory disease. Shao and colleagues presented evidence that pyrin, as part of the pyrin inflammasome, acts as a sensor of different inactivating bacterial modification RHO GTPases, rather than directly interacting with these microbial products. This guard mechanism of pathogen detection has previously reported for pathogen recognition receptor (PRRs) in plants. Several Rho-inactivating bacterial toxins have been reported, including the TcdB toxin from Clostridium difficile the C3 toxin from Clostridium botulinum and the pertussis toxin from Bordetella pertussis, and, in the context of this review B. cenocepacia deamidates RhoA at Asn41 [125,136].

More recent developments in this field include the discovery that RhoA activates the serine-threonine kinases PKN1 and PKN2 that bind and phosphorylate pyrin
[120]. This activation of PKN1 and PKN2 was found to decrease IL-1\(\beta\) release from peripheral blood mononuclear cells (PBMCs) of patients with FMF or HIDS. Defective prenylation, as seen in HIDS, was associated with RhoA inactivation and pyrin inflammasome activation (Fig. 4). Thus, the authors propose a novel molecular connection between FMF and HIDS.

Masters et al. have described an autoinflammatory disease, labelled pyrin-associated autoinflammation with neutrophilic dermatosis (PAAND), caused by a mutation in pyrin, which disrupts pyrin regulation and mimics the effect(s) of pathogen sensing by pyrin, leading to proinflammatory IL-1\(\beta\) production [137]; the disease resolved in one patient by targeting IL-1\(\beta\). These data reveal a regulatory mechanism of pyrin activation and suggest that it is regulated through a guard-like mechanism, which prevents the development and progression of autoinflammation.

A number of fundamental questions arise from these fascinating discoveries, including the precise molecular mechanisms of pyrin inflammasome activation and whether specific environmental factors may trigger attacks in patients with autoinflammation.

**New diseases and mechanisms**

Gain-of-function mutations in the NLRC4 gene a novel inflammasome disorder associated with predisposition to macrophage-activation syndrome (MAS) and highly elevated IL-18 levels [14] (Table 1). Aksentijevich and colleagues [138] found that TNFAIP3 mutations cause haploinsufficiency of A20 (HA20), with reduction of NF-\(\kappa\)B [139] and IL-1 signalling leading to A20 haploinsufficiency, in an early-onset autoinflammatory disease, where the phenotype resembles Behcet’s disease [140]. A paper in press by the same group describes another NF-\(\kappa\)B mediated disease,

http://mc.manuscriptcentral.com/jpath
caused by loss-of-function mutations in OTULIN/FAM105B gene, encoding a
deubiquitinase with linear linkage specificity. These patients have a very severe
phenotype, surprisingly resembling CANDLE, but clinically responsive to TNF
inhibitors [141]. Together with HA20 these two diseases described a new category of
autoinflammatory diseases, due to dysregulated ubiquitination. Thus the
ubiquitination pathway has assumed greater important in the investigation of
systemic autoinflammatory disorders of undefined etiology (SAIDs).

Mutations in the TNFRSF11A gene have been reported in patients with a
disease that has clinical similarities to TRAPS [142]. A report of a novel digenic
pattern of inheritance in CANDLE/PRAAS patients, has provided insights into
proteasome dysfunction and associated IFN production [66].

Triggers of autoinflammation

Autoinflammatory diseases are mainly driven by proinflammatory cytokines,
usually generated as a result of cellular stress, and especially oxidative stress with
associated mitochondrial DNA (mtDNA) damage. The resulting release of metabolic
mediators such as mitochondrial ROS, which acts as a DAMP for the NLRP3
inflammasome activation [143]. The search for identifiable (exogenous) factors that
might serve to trigger attacks of autoinflammation involves careful the patient’s
environment, diet, or lifestyle [110]. Some known triggers known to influence the
effects of individual mutations include

1. Generalised exposure to cold may precipitate attacks of fever in familial cold
autoinflammatory syndrome (FCAS).
2. Attacks of HIDS may be triggered by trauma, illnesses or vaccine reactions [117,144]. A severe inflammation reaction following vaccination against Streptococcus pneumoniae has been described in patients with CAPS [142].

3. Urate and CPP crystals cause NLRP3 inflammasome activation in gout and calcium pyrophosphate deposition disease (CPPD) [77].

4. The pyrin inflammasome is activated upon bacterial toxin-induced modification of host Rho GTPases [125].

5. Dying cells have the capacity to activate the innate immune system and induce a sterile inflammatory response [145,146]; necrotic cells are sensed by the Nlrp3 inflammasome with subsequent release of IL-1β [147]. In a mouse model mitochondria were critical to activation of the Nlrp3 inflammasome by direct binding of Nlrp3 to the inner mitochondrial lipid cardiolipin.

6. The relationship between IFN-α and brain pathology in AGS is poorly understood [148]. Viral infection and replication introduces single-stranded RNA (ssRNA), double-stranded RNA (dsRNA) and DNA:RNA hybrids, with induction of type I IFN genes. The AGS phenotype may resemble congenital viral and individual subsets of SLE [43,44].

**Autoinflammation in the more common chronic systemic conditions**

It is now accepted that innate immune-mediated inflammation plays a key role in the pathogenesis of some of the more common chronic systemic conditions, such as Crohn’s disease [4], type 2 diabetes (T2D) and a myeloid subset of rheumatoid arthritis (RA) [149], as well as in diseases not formerly considered inflammatory, such as neurodegenerative conditions [150]. There is increasing evidence that cell intrinsic or environmental alterations in protein homeostasis may contribute to the
pathogenesis in these conditions; thioredoxin-interacting P (TXNIP) serves as a functional link between ER stress, NLRP3 inflammasome activation and inflammation related to T2DM [151].

Therapies

As the field of autoinflammatory disorders has developed so rapidly clinicians and researchers have produced guidelines to optimise and disseminate recommendations for universal management of children and young adults with these disorders. An international panel of 22 experts was established to develop evidence-based recommendations for the management and treatment of CAPS, TRAPS and MKD using the European League Against Rheumatism (EULAR) standard operating procedures for developing best practice [152,153].

Proposed new Nomenclature – an expanded classification of autoinflammatory diseases

The continuously expanding number of monogenic diseases, for which susceptibility genes have been found, and which present with a range of overlapping clinical features, both autoinflammatory and autoimmune in nature, has raised the question as to how to (sub)classify those conditions, as the terms autoinflammation and/or autoimmunity are insufficient to adequately describe them. In addition to the challenge posed by these conditions with overlapping features, a range of other diseases, with variable clinical manifestations of immunodeficiency and immune dysregulation/autoimmunity have been genetically delineated. These include PLCG2-associated antibody deficiency and immune dysregulation (PLAID) [154], haploinsufficiency of CTLA-4, caused by heterozygous germline mutations [155] and
XLPDR disorder, due to deficiency of POLA1, which encodes the catalytic subunit of DNA polymerase-α [156]. This latter condition also has an associated IFN signature.

Despite these observations, a combination of both pathogenic innate and adaptive immune responses underlie the immunopathology of most inflammatory conditions. As reviewed in [46] some clinical features, like B-cell immunodeficiency, may arise in conditions which are mainly innate-immune driven, and autoinflammatory in phenotype, such as deficiency of adenosine deaminase 2 (DADA2) [157,158] but B-cell immunodeficiency may also be found in monogenic autoimmune conditions, like haploinsufficiency of CTLA-4 and PLAID. Furthermore, AGS7 has the potential to progress from being primarily innate-immune driven to becoming an antibody-mediated disease.

In light of the expanding number of overlapping syndromes of both autoinflammation and autoimmunity we propose to broaden the classification of diseases by assigning the term autoimmuno-inflammatory disease. Conditions like PLAID, where the clinical picture combines features of immunodeficiency as well as autoimmunity, and, arguably, the cold urticaria element of PLAID is innate immune related, might also be considered; following the template proposed above complex conditions of that nature could be referred to as an autoimmuno-inflammatory-immunodeficiency. However the primary immunodeficiency diseases (PI) constitute an extensively classified group of conditions, and it may not be possible to find a satisfactory all-purpose blanket term for novel complex conditions with features of immunodeficiency as well as autoimmunity and autoinflammation.
Summary of developments and outlook

The identification of a genetic aetiology for an increasing number of autoinflammatory diseases has led to a growing recognition that dysregulation of this normal defence mechanism may be more prevalent than previously realised in other diseases. Autoinflammation is likely to play a variable role in a wide spectrum of human disease, acting within a milieu of complex processes, involving innate and adaptive immunity. Understanding the role of autoinflammation in various diseases processes is essential if new targets are to be identified for future therapies.

A major part of the human immune system’s basic function is to control the host’s relationship with his/her microbiota, referring to the the totality of microorganisms that inhabit the human body in health and disease. Recent major technological advances, including single cell sampling and shotgun sequencing enables detailed study of individual microbiota and inflammatory disease can related to components of the microbiome [159] (the combined genetic material of the microorganisms), and to the intracellular pathways that pathogens within the microbiome may dysregulate survive [160]. It is most likely that the widespread influence of intracellular microbes on innate immune defences and autoinflammatory diseases will be elaborated in significant detail in the next decade.

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TS, DP, SS and MMcD wrote the manuscript.

Conflict of Interest Statement:

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Legends for Figures and Table

Figure 1: Diseases classified according myeloid (autoinflammation) or lymphoid lineage (autoimmune).

Diseases of the immune system are classified according to whether the lymphocyte responsible for the disease is of myeloid (autoinflammation) or lymphoid lineage (autoimmune). Clinical heterogeneity within immunological diseases may reflect the variable expression of autoinflammatory and autoimmune factors in disease causation.

A disease spectrum that includes rare monogenic diseases at the polar ends of the spectrum, and polygenic diseases, involving both myeloid and lymphoid cells in pathogenesis, occupying the centre [10]. This diagram adds a third variable, environmental triggers, to further define the pathogenesis of these diseases. The figure does not include all immunologically recognised diseases because of their large number.

HIDS- hyper IgD syndrome, CAPS- cryopyrin-associated autoinflammatory syndrome, FMF- familial Mediterranean fever, TRAPS- tumour necrosis factor receptor associated periodic syndrome, sJIA- systemic juvenile idiopathic arthritis, AOSD- adult onset Still’s disease, RA- rheumatoid arthritis, CF- cystic fibrosis, SLE- systemic lupus erythematosus, T1D- type 1 diabetes, APS-1- autoimmune polyglandular syndrome type 1, PLAID- PLCG2 associated antibody deficiency and immune dysregulation, ALPS- autoimmune lymphoproliferative syndrome, IPEX- immune dysregulation polyendocrinopathy enteropathy X-linked syndrome.
Figure 2: Priming, assembly and degradation of the NLRP3 inflammasome.

An activating signal is required for the NLRP3 inflammasome to be assembled - examples include ATP-dependent K efflux, particulate substances, such as urate crystals entering the cell through lysosomal degradation pathways, mitochondrial damage and release of mtDNA or mtROS and intracellular pathogen recognition. The ligand for the NLRP3 inflammasome in humans is pro-caspase-1. Once activated, caspase-1 cleaves and activates inactive cytokines pro-IL-1β and pro-IL-18. A second priming signal is required to induce pro-IL-1β and pro-IL-18 expression. This is typically through NF-κB signalling, downstream of TLRs, or through XBP-1 downstream of the UPR. Once the inflammatory stimulus has subsided the NLRP3 inflammasome is cleared by autophagolysosomal degradation.

Figure 3: Cystic fibrosis as an autoinflammatory disease.

CF shares many common features of autoinflammatory diseases. Due to the mutated CFTR, there is increased ROS signalling and reduced antioxidant secretion. CF also manifests with hyperuricaemia, low airway surface pH, ASL dehydration and high glucose levels, all thought to be triggers of the NLRP3 inflammasome. CFTR mutations may cause extreme ionic imbalances, many of which have been linked with NLRP3 inflammasome activation. As the CFTR is misfolded in many genotypes of CF, this results in ER stress, UPR activation, and XBP1 signalling. Finally, increased lung infections provide frequent activation of the TLR-NF-κB inflammatory signalling pathway, priming the NLRP3 inflammasome.
When mutations in the NLRP3 inflammasome pathway or excessive/continuous stimuli interfere with its activation or priming, this inflammasome becomes the hub of life-limiting innate immune-driven diseases. Gout (yellow arrow), TRAPS (green arrow), MWS (red arrow), FMF (blue arrow) and HIDS (orange arrow) are examples of autoinflammatory conditions where the NLRP3 inflammasome is at the centre of disease pathology.

Table 1: Autoinflammatory diseases.

An update on the mechanisms involved in the autoinflammatory diseases mentioned in this review. A more comprehensive list of these diseases exists in de Jesus et al.’s review [3].
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193x275mm (300 x 300 DPI)
195x229mm (300 x 300 DPI)
278x193mm (300 x 300 DPI)
<table>
<thead>
<tr>
<th>Category</th>
<th>Name</th>
<th>Gene</th>
<th>Mechanism</th>
<th>Therapy</th>
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<tbody>
<tr>
<td>IL-1</td>
<td>HIDS</td>
<td>MVK</td>
<td>Mutated mevalonate kinase causes reduced isoprenoid synthesis, leading to reduced prenylation of RhoA, activating the pyrin inflammasome, reduced prenylation disrupts autophagy and ROS clearance, activating NLAF3.</td>
<td>IL-1 inhibition</td>
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<td>GAPPS (FCAS, MWS, CINCA, NOMID)</td>
<td>NLAF3</td>
<td>Constitutive NLAF3 inflammasome activation</td>
<td>IL-1 inhibition</td>
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<td>DIRA</td>
<td>IL1RIV</td>
<td>IL-1RA deficiency</td>
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<td>DITRA</td>
<td>IL1RIV</td>
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<td>IL-1 inhibition</td>
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<td>MAS</td>
<td>NLRC4</td>
<td>Uncontrolled macrophage activation, with increased secretion of IL-18, IFN-gamma and GM-CSF. Mechanism unknown.</td>
<td>Not defined - tocilizumab</td>
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<td>PARDP</td>
<td>MEFV</td>
<td>Pyrin inflammasome activation</td>
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<tr>
<td></td>
<td>FMF</td>
<td>MEFV</td>
<td>Pyrin inflammasome activation</td>
<td>NSAIDs, colchicine, IL-1 inhibition</td>
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<td>UPR</td>
<td>Cystic fibrosis</td>
<td>CFTR</td>
<td>Mutated CFTR, causing multisystem disease due to ionic imbalance and ER stress. NLAF3, NLRC4, UPR.</td>
<td>Antibiotics and NSAIDs</td>
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<td>TRAP3</td>
<td>TNFR1</td>
<td>TNF receptor activation, UPR, NLAF3, activation</td>
<td>IL-1 inhibition</td>
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<td>IFN</td>
<td>Aicardi-Goutières syndromes (AGS)</td>
<td>TRAF3, RNASEH2B, RNASEH2C, RNASEH2A, SAMHD1, ADAR</td>
<td>Aberrant sensing of DNA/RNA, with excessive IFN-producing responses</td>
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<td>CANDIDE/PRAAS syndrome</td>
<td>PSMB8</td>
<td>Gain of function mutation, UPR, IFN signature</td>
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<td>Bechet’s disease</td>
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<td>PDLA1</td>
<td>Dysfunctional DNA polymerase-a catalytic subunit</td>
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<td>STING</td>
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<td>CTLA4</td>
<td>Dysregulated FoxP3 Treg cells</td>
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<td>DADA2</td>
<td>CCR1</td>
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<td>AGS5</td>
<td>IFIH1</td>
<td>Dysfunctional sensing of nucleic acids</td>
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<td>PLAID</td>
<td>PLEC2, CTRA-4</td>
<td>Adaptive immunodeficiency</td>
<td>Antihistamines</td>
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<td>Granulomatous disease</td>
<td>Blau Syndrome</td>
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<td>Hyperactive NF-κB signalling</td>
<td>Steroids, anti-TNF inhibitors, IL-1 inhibition</td>
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<td>Dysregulated Ubiquitination</td>
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<td>IL-1 inhibition</td>
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<td>OTULIN</td>
<td>Dysfunctional ubiquitination and hyperactive NF-κB signalling</td>
<td>Steroids, anti-TNF inhibitors, IL-1 inhibition</td>
</tr>
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Autoinflammatory diseases

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