HYBRID REVIEW

Inflammasome regulation in driving COVID-19 severity in humans and immune tolerance in bats

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Summary sentence: Human SARS-CoV-2 infection-induced inflammasome activation is associated with COVID-19 severity, whereas bats, which host pathogenic CoVs, show dampened inflammasome activation and immune tolerance to viral infections.

Abstract
Coronaviruses (CoVs) are RNA viruses that cause human respiratory infections. Zoonotic transmission of the SARS-CoV-2 virus caused the recent COVID-19 pandemic, which led to over 2 million deaths worldwide. Elevated inflammatory responses and cytotoxicity in the lungs are associated with COVID-19 severity in SARS-CoV-2-infected individuals. Bats, which host pathogenic CoVs, operate dampened inflammatory responses and show tolerance to these viruses with mild clinical symptoms. Delining the mechanisms governing these host-specific inflammatory responses is essential to understand host–virus interactions determining the outcome of pathogenic CoV infections. Here, we describe the essential role of inflammasome activation in determining COVID-19 severity in humans and innate immune tolerance in bats that host several pathogenic CoVs. We further discuss mechanisms leading to inflammasome activation in human SARS-CoV-2 infection and how bats are molecularly adapted to suppress these inflammasome responses. We also report an analysis of functionally important residues of inflammasome components that provide new clues of bat strategies to suppress inflammasome signaling and innate immune responses. As spillover of bat viruses may cause the emergence of new human disease outbreaks, the inflammasome regulation in bats and humans likely provides specific strategies to combat the pathogenic CoV infections.

KEYWORDS
Bat immunity, Coronaviruses, COVID-19, Immune tolerance, Inflammasome, NLRP3, SARS-CoV-2

INTRODUCTION

Coronaviruses (CoVs) are single-stranded RNA viruses in the family of Coronaviridae. Zoonotic transmission of CoVs causes mild to severe human infections. Seven different CoVs have entered humans so far. Human infections of OC43-CoV, 229E-CoV, NL63-CoV, and HKU1-CoVs cause seasonal colds and are mostly asymptomatic, whereas infections of severe acute respiratory syndrome CoV (SARS-CoV) and Middle East respiratory syndrome CoV (MERS-CoV)
led to severe disease outbreaks.\textsuperscript{1–3} SARS-CoV-2 infection caused the ongoing pandemic and led to more than 2 million deaths worldwide. SARS-CoV-2 infects the upper respiratory tract and binds to host receptor protein, angiotensin-converting enzyme 2 (ACE2), through the spike protein (S protein).\textsuperscript{4–6} The membrane-bound host serine protease, transmembrane protease serine 2 (TMPRSS2), activates S protein to facilitate the entry of SARS-CoV-2 into the host cells (Fig. 1).\textsuperscript{4} The RNA genome of the SARS-CoV-2 will be released into the cytoplasm, where it replicates to generate new progeny of virions.\textsuperscript{1,2} New virion particles transport through the secretion pathway to be released into extracellular space. In most cases, the SARS-CoV-2 infection is asymptomatic and, in some cases, it spreads to the lower respiratory tract causing pneumonia.\textsuperscript{2,6} SARS-CoV-2 infection in some patients becomes severe and develops into acute respiratory distress syndrome leading to severe respiratory abnormalities and mortality.\textsuperscript{2,6} The SARS-CoV-2-induced moderate and severe disease in humans is collectively named coronavirus disease 2019 (COVID-19).\textsuperscript{2,6}

The SARS-CoV-2-driven pathogenesis in human patients is a complex process. It is increasingly evident that excessive cytokine responses and infiltration of inflammatory cells, particularly inflammatory monocytes and neutrophils, are associated with COVID-19 severity.\textsuperscript{7–9} Virus-induced cytopathic effects of lung epithelial cells in association with excessive inflammatory cytokines trigger acute lung injury and loss of respiratory function during COVID-19.\textsuperscript{9,10–12} Endothelial barrier damage further leads to viral dissemination and spread to other organs. Dysregulated immune responses and increased inflammatory cytokine signatures also occur in other pathogenic RNA virus infections.\textsuperscript{8,11} Inflammation-driven excessive cytokine response in response to SARS-CoV-2 infection is highly associated with severe COVID-19 and a great effort has been put to understand and target these responses for alleviating COVID-19 severity. Several studies report the association of excessive levels of proinflammatory cytokines, IL-1α, IL-1β, IL-6, IL-8, and TNF with COVID-19 severity.\textsuperscript{10,12–16} The mechanisms triggering these specific cytokine responses during SARS-CoV-2 remain elusive.

\section*{2 | INFLAMMASOME ACTIVATION}

Inflammasomes are heteromeric multiprotein complexes that promote the activation of caspase-1 (CASP1) protease.\textsuperscript{17,18} The CASP1 activation in turn triggers the release of proinflammatory IL cytokines, IL-1β and IL-18, and damage-associated molecular patterns (DAMPs) like IL-1α, high mobility group box protein 1 (HMGB1), and lactate dehydrogenase (LDH)\textsuperscript{19,20} (Fig. 1). Activation of innate immune receptors, such as Nod-like receptors (NLRs) and AIM2-like receptors (ALRs), recruits an adaptor protein, apoptosis-associated speck-like protein containing a CARD (ASC), to assemble the inflammasome complex and trigger proximity-induced CASP1 activation\textsuperscript{18,21} (Fig. 1). The widely studied NOD-like receptor family pyrin domain (PYD) containing 3 (NLRP3) inflammasome requires 2 signals for its activation. The first signal provides priming of the NLRP3 inflammasome through inflammatory stimuli (like TLR or TNF receptor (TNFR)) engagement that promote NF-kB dependent expression of the NLRP3 and IL-1β.\textsuperscript{22,23} The priming signal also facilitates NLRP3 post-translational modifications favoring inflammasome assembly.\textsuperscript{24–26} The second signal, known as the activation signal, is provided by pathogen-associated molecular patterns or cellular damage signals.\textsuperscript{17,27,28} The second signal triggers NLRP3 inflammasome assembly, ASC recruitment, and activation of the CASP1. The ASC recruitment to NLRP3 inflammasome assembly, ASC recruitment, and activation of the CASP1. The ASC recruitment to NLRP3 inflammasome assembly, ASC recruitment, and activation of the CASP1. The ASC recruitment to NLRP3 inflammasome assembly, ASC recruitment, and activation of the CASP1. The ASC recruitment to NLRP3 inflammasome assembly, ASC recruitment, and activation of the CASP1. The ASC recruitment to NLRP3 inflammasome assembly, ASC recruitment, and activation of the CASP1.
homotypic interactions between them \(^{18,29}\) (Fig. 1). CASP1 cleaves pro-IL-1\(\beta\) and pro-IL-18 into mature IL-1\(\beta\) and IL-18 forms. CASP1 also cleaves a pore-forming protein Gasdermin D (GSDMD) at the linker region of its N- and C-terminal domains to liberate it from its autoinhibitory conformation \(^{30-32}\) (Fig. 1). The N-terminal domain of the GSDMD is transported to the plasma membrane and oligomerize into a pore structure whose pore size is sufficient to enable the passage of small molecules and mature IL-1\(\beta\) and IL-18. \(^{18,19}\) The formation of GSDMD pores in turn promotes osmotic lysis and a lytic form of cell death called pyroptosis \(^{30-32}\) (Fig. 1). Earlier it was considered that the plasma membrane rupture and osmotic lysis that occur after GSDMD pore formation is a passive event. A recent study challenged this assumption and demonstrates that the plasma membrane rupture during pyroptosis is a programmed phenomenon regulated by nerve injury protein 1 (NINJ1) \(^{33}\) (Fig. 1). Thus, inflammasome activation triggers GSDMD pore formation and plasma membrane rupture for releasing large molecular weight DAMPs like HMGB1 and LDH. A recent study demonstrates that a lectin protein, Galectin-1, is also a bona fide DAMP released in inflammatory cell death and promotes inflammation. \(^{34}\) Serum levels of DAMPs are considered as a measure of inflammatory cell death activation in disease settings. Inflammasome activation facilitates antimicrobial host defense during microbial infections and compromised activation leads to severe pathology. However, inflammasomes also drive chronic inflammatory responses in autoimmune-inflammatory and metabolic diseases. \(^{28,35}\) Dysregulated or excessive inflammasome activation accelerates microbial pathogenesis through elevated proinflammatory responses or cytokine storm. \(^{21,23,28-30,35}\)

### 3 THE INFLAMMASOME ACTIVATION AND COVID-19 SEVERITY

The NLRP3 inflammasome is a widely studied complex and is known to be activated by RNA viruses. \(^{17,18,23,36}\) Inflammasome responses play an essential role in clearing viruses and virus-infected cells and promote tissue repair responses. \(^{27,36-40}\) In many RNA virus infection cases, viral RNAs generated during the replication process are sufficient to activate the NLRP3 inflammasome. \(^{37,36}\) In some cases, viral proteins also regulate the activation of the NLRP3 inflammasome. The inflammasome-induced cell death and the release of proinflammatory IL-1\(\alpha\), IL-1\(\beta\), and IL-18 cytokines drive the recruitment of inflammatory immune cells to the infection site and promote viral clearance and tissue repair processes. \(^{11,27,36-40}\)

The increased levels of specific cytokines during severe COVID-19 suggest dysregulated inflammatory responses in pathogenesis progression. \(^{7,10}\) Immune response profiling of moderate and severe COVID-19 patients indicates a correlation of inflammasome-associated proinflammatory cytokines IL-1\(\alpha\) and IL-1\(\beta\) with the severity of the disease. \(^{8,10,12,13,41}\) Although several studies indicate the inflammasome role in severe COVID-19 disease progression, the activation of inflammasome in response to SARS-CoV-2 infection was obscure. A recent study demonstrated that the SARS-CoV-2 infection in primary human monocytes triggers CASP1 activation and IL-1\(\beta\) production, providing direct evidence for the activation of the inflammasome and proinflammatory cytokine release. \(^{41}\) Similar to other RNA virus infections, replication of the SARS-CoV-2 is critical for the activation of the inflammasome. \(^{39-42}\) Experiments with MCC950, an inhibitor specific to NLRP3, suggest the specific role of NLRP3 inflammasome in SARS-CoV-2 induced CASP1 activation and IL-1\(\beta\) release. \(^{41}\) Importantly, increased levels of LDH were seen in severe COVID-19 disease individuals indicating the lytic cell death programs (such as necroptosis and pyroptosis) after SARS-CoV-2 infection. \(^{41}\) SARS-CoV-2 infection in primary human monocytes triggers LDH release; however, MCC950 fails to suppress its release. This suggests the possibility of the NLRP3 inflammasome-independent mechanisms triggering the lytic form of cell death and the release of DAMPs, like LDH, in SARS-CoV-2 infection (Fig. 2). \(^{41,43}\) The activation of NLRP3 inflammasome and LDH release indicates potential activation of the pyroptosis along with other lytic cell death programs during SARS-CoV-2 infection and in COVID-19. Additional studies monitoring the activation of GSDMD and testing SARS-CoV2 infection in Gsdmd \(^{-/-}\) cells deliver the specific contribution of pyroptosis and other cell death mechanisms in mediating the release of LDH and other DAMPs. Mouse hepatitis virus (MHV) is a mouse-adapted lab strain, extensively used to understand host responses during CoV infections. \(^{3}\) A recent study demonstrates that the MHV infection in primary macrophages triggers pyroptosis, apoptosis, and necroptosis (PANoptosis), suggesting the ability of CoVs to engage multiple inflammatory lytic cell death programs. \(^{44}\) It remains to be established whether pathogenic human CoVs, like SARS-CoV-2, engage PANoptosis. The prominent activation of the NLRP3 inflammasome is observed in response to the SARS-CoV-2 infection and is active in COVID-19 patients who require hospitalization. \(^{41,45}\) Higher levels of CASP1, IL-1\(\beta\), and IL-18 are seen in COVID-19 patients compared with healthy individuals. \(^{12,15,41,45}\) Analyzing peripheral blood mononuclear cells isolated from COVID-19 patients indicates the active inflammasome and CASP1 in lethal cases of COVID-19. \(^{41}\) Also, NLRP3 inflammasome and ASC speck formation were observed in the area of lung injury and postmortem tissues from COVID-19 patients. \(^{31}\) Thus, SARS-CoV-2 infection in human lungs engages the inflammasome activation, which may be a likely therapeutic target for alleviating inflammation and severity of COVID-19. \(^{30,41,45,46}\) The excessive release of inflammasome cytokines may trigger uncontrolled immune cell infiltration and severe pathology in the lung. Furthermore, these findings corroborate other studies supporting the use of inflammasome activation as a marker of COVID-19 severity.

Older males with comorbidities show severe COVID-19 pathology after SARS-CoV-2 infection. \(^{37}\) The comorbidities, like diabetes, obesity, cardiovascular diseases, cancers, digestive and respiratory system disease, and neurologic disorders, make older males more susceptible to COVID-19. \(^{47-49}\) Physiologic aging is associated with chronic low-grade inflammation, which increases the risk of age-related diseases. \(^{49,50}\) The ligands that appear in aging promote increased NLRP3 inflammasome activation and chronic low-grade inflammation in older individuals. \(^{50}\) Thus, NLRP3 inflammasome and chronic low-grade inflammation in older individuals predispose them to COVID-19 severity. \(^{51}\) The higher risk of males compared with females indicates the role of...
steroid hormones in determining COVID-19 severity. Female reproductive hormones, estrogen and progesterone, suppress inflammatory immune responses and NLRP3 inflammasome function. The steroid hormone-induced anti-inflammatory responses in females may have a protective role in COVID-19.

4 | MECHANISMS OF PATHOGENIC CORONAVIRUS MEDIATED NLRP3 INFLAMMASOME ACTIVATION

How SARS-CoV-2 activate inflammasome, pyroptosis, and other lytic forms of cell death? Currently, the precise mechanisms of SARS-CoV-2 induced inflammasome activation are unclear. Potassium (K\(^+\)) efflux from cytosol or endoplasmic reticulum (ER) stress-triggered calcium (Ca\(^{2+}\)) leakage into the cytosol promotes NLRP3 inflammasome activation. RNA virus proteins that act like viroporins or ion channels are linked to the activation of NLRP3 inflammasome and release of IL-1\(\beta\). Also, SARS-CoV viroporins are associated with the activation of the NLRP3 inflammasome. SARS-CoV envelope protein (E protein) is a viroporin that is localized to the ER and the Golgi complex and functions as an ion channel to promote Ca\(^{2+}\) leakage and NLRP3 activation. SARS-CoV open reading frame 3a protein (ORF3a) is also a viroporin that promotes K\(^+\) efflux. This further promotes mitochondrial reactive oxygen species and activation of the NLRP3 inflammasome and release of IL-1\(\beta\). Both E protein and ORF3a of SARS-CoV provide a priming signal to activate NF-kB signaling, which up-regulates the transcription of NLRP3, IL-1\(\beta\), and other proinflammatory cytokines (Fig. 2). In addition to K\(^+\) efflux and NF-kB activation, ORF3a also facilitates the TNFR-associated factor 3 (TRAF3)-mediated ubiquitination of the ASC to engage the NLRP3 inflammasome. The amino acid residues of E protein and ORF3a, critical for ion channel activity, NF-kB activation, and NLRP3 inflammasome assembly, are conserved in SARS-CoV and SARS-CoV-
2 (Supplemental Fig. S1). E protein and ORF3a may likely trigger NLRP3 inflammasome activation in response to SARS-CoV-2 infection and promote COVID-19 severity. A new study demonstrates that the ORF3a of SARS-CoV-2 activates the NLRP3 inflammasome by regulating its expression and ASC-mediated inflammasome assembly.61 The role of SARS-CoV-2 ORF3a in inflammasome-induced pathogenicity remains to be studied. Nevertheless, viroporins of pathogenic CoVs, which form ion channels, retain the ability to activate the NLRP3 inflammasome like other RNA viruses (Fig. 2). In addition to promoting the NLRP3 inflammasome, the SARS-CoV ORF-3a can also trigger necrotic cell death.62 The dual function of ORF-3a in promoting inflammatory cell death pathways (pyroptosis and necroptosis) may explain the inflammasome-independent release of LDH during SARS-CoV-2 infection.41,62 In addition to E protein and ORF3a, the spike protein (S protein) and ORF7a of the SARS-CoV promote NF-kB activation and inflammasome priming (Fig. 2).57,63 The SARS-CoV-2 S protein also triggers inflammasome priming in COVID-19 patient-derived macrophages.64 It appears that SARS-CoV-2-induced inflammasome activation is a complex process that involves multiple viral components. A recent study demonstrates that nonstructural protein (NSP) 1 and NSP13 of SARS-CoV-2 abolish NLRP3 inflammasome-mediated CASP1 activation and IL-1β release (Fig. 2).65 Thus, the SARS-CoV-2 also operates viral mechanisms counteracting inflammasome activation.

ORF8b protein of SARS-CoV acts like a virulence factor and is reported to interfere with IFN responses.66,67 SARS-CoV ORF8 exists in 3 different forms, ORF8ab, ORF8a, and ORF8b. A single ORF encodes ORF8ab; however, a 29-nucleotide deletion in ORF8ab results in the generation of overlapping ORF8a and ORF8b. Deletion mutants of ORF8 in SARS-CoV associate with decreased replicative fitness.1,68,69 SARS-CoV ORF8b protein forms insoluble aggregates enabling altered cellular functions.70 SARS-CoV ORF8b promotes NLRP3 inflammasome activation in macrophages and triggers cell death in epithelial cells and macrophages.70 The ORF8b directly interacts with leucine-rich repeat (LRR) domain of the NLRP3 and localizes to NLRP3-ASC inflammasome compartments.70 Whether SARS-CoV-2 ORF8 activates the inflammasome and IL-1β release is not investigated so far. A circulating variant of SARS-CoV-2 with a 382-nucleotide deletion in its ORF8 encoding genome (ORF8-Δ382) is reported recently, which affects its expression.71,72 Patients infected with ORF8-Δ382 variant of SARS-CoV-2 show milder symptoms compared with wild-type (WT)-virus infected individuals.71 Also, lower systemic levels of proinflammatory cytokines are observed in individuals infected with the ORF8-Δ382 variant.71 In vitro studies in nasal epithelial cells show similar transcription patterns after WT SARS-CoV-2 and ORF8-Δ382 mutant infection.73 Thus, the ORF8 function appeared to be associated with the regulation of proinflammatory responses originated from immune cells during SARS-CoV-2 infection. Of note, SARS-CoV and SARS-CoV-2 display striking variation in ORF8 protein sequences (Supplemental Fig. S1). Bat CoVs, which are phylogenetically related to SARS-CoV, also show variation in the ORF8 region.1 It appears that the SARS-CoV-2 variations associate with host proinflammatory responses and may determine the outcome of COVID-19 severity. It remains to be established the precise role of ORF8 in SARS-CoV-2 inflammasome activation and its impact on host proinflammatory responses. Except ORF8, SARS-CoV and SARS-CoV-2 proteins that promote inflammasome activation are conserved. As several proteins of the SARS-CoV-2 are associated with inflammasome regulation, the precise contribution of each viral protein or combination of proteins that determine SARS-CoV-2-induced inflammasome activation or suppression needs to be investigated. Defining specific signatures in SARS-CoV and SARS-CoV-2 proteins, which are critical for triggering inflammasome activation, may facilitate the prediction of the pathogenesis of CoVs in bats that have not infected humans so far. In vivo functional studies with SARS-CoV-2 mutant viruses, such as ORF8 mutant restricting its binding to TRAF3, in animal models (mice, hamsters or monkeys) will help to annotate the key elements of SARS-CoV-2 orchestrating inflammasome activation and COVID-19 severity.

Type I IFNs promote efficient activation of the inflammasome and pyroptosis.74,75 Virus-induced type I IFNs promote the activation of inflammasome responses in various RNA virus infections.27,74 Type I IFNs at the acute stage of the infection or before peak SARS-CoV or MERS-CoV viral titers robustly inhibit the dysregulated inflammatory response and lung damage and marginally affect viral titers.76,77 MERS-CoV, SARS-CoV, and SARS-CoV-2 viruses employ highly evolved viral mechanisms to restrict type I IFN secretion in infected host cells.78–80 This early on type I IFN antagonism during CoV infection helps increase in viral titers and spread. In lethal SARS-CoV infection, a robust increase in viral titers and delayed type I IFN signaling drive pathologic inflammatory response and lung damage.76,77 In particular, delayed type I IFNs promote the recruitment of inflammatory monocyte and macrophages, resulting in an uncontrolled release of inflammatory cytokines and chemokines in the lung.76,77 The subsequent loss of epithelial integrity and endothelial damage results in lung pathology and viral spread to other organs. Delayed type I IFNs during lethal SARS-CoV-2 infection may trigger elevated inflammasome activation, pyroptosis, and release of IL-1β and IL-18. Both pyroptosis and proinflammatory cytokines may drive the loss of epithelial and endothelial integrity and lung damage.

Dysregulated cytokine responses or cytokine storms occur in systemic bacterial infections that lead to organ failure and mortality.81,82 RNA virus infections like influenza, Zika, and Ebola are associated with dysregulated inflammatory responses and severe pathology.83–93 Excessive cytokine responses during highly pathogenic influenza infections, H1N1 and H5N1, are associated with severe pulmonary pathology. Do SARS-CoV-2 inflammatory cytokine signature mimics other pathogenic RNA virus infections? Recent studies show that the levels of IL-1α and IL-1β are significantly lower in COVID-19-infected patients when compared with influenza-infected individuals.14,41,80,94 In support of these studies, the analysis of gene expression profiles in the upper respiratory tract of COVID-19 patients with other respiratory virus infections further suggests a striking decrease in the expression of inflammasome genes.15 However, IL-1α and IL-1β levels were higher in COVID-19 patients compared with healthy controls suggesting a correlation of inflammasome activation with SARS-CoV-2-mediated disease severity.10,12,15 Even though the cytokine storm is considered
a critical feature of COVID-19, these new studies reveal a specific set of inflammatory cytokine signatures responsible for the pathogenesis and inflammasome cytokines are among them. Proinflammatory IL-1 cytokine (IL-1α and IL-1β) signaling often promotes host defense responses during RNA virus infections. IL-1 receptor antagonist (IL-1Ra) is a natural inhibitor of IL-1 cytokine signaling. IL-1Ra-mediated regulation of IL-1 cytokines enables balanced control of inflammation and retain homeostasis. Like other pathogenic RNA virus infections, increased levels of IL-1Ra were reported in individuals with severe COVID-19 disease. It is not clear yet why inflammation dampening IL-1Ra is elevated in SARS-CoV-2 infection. In the case of Ebola and Zika virus infections, where increased IL-1Ra levels are reported, IL-1Ra is required for protective immune responses and its deletion leads to severe proinflammatory responses.

5 | BAT CORONAVIRUSES, INFLAMMASOME REGULATION, AND IMMUNE TOLERANCE

Bats are the only mammals that fly and host several pathogenic RNA viruses like CoVs, Ebola, and Nipah viruses. Bats exhibit mild clinical symptoms or often asymptomatic despite high viral titers. Pathogenic human CoVs, SARS-CoV, SARS-CoV-2, and MERS-CoV, likely originated from bats and zoonotic transmission of these CoVs caused severe human outbreaks and pandemics. Bats host diverse CoVs that are phylogenetically related to the pathogenic human CoVs. This indicates a potential threat of a spillover of bat-originated CoVs that may cause future outbreaks. Bats display high metabolic rates, however, their life span is long. Bats appear to show efficient antiviral mechanisms mainly because of constitutive expression of type I IFNs and dampened inflammatory responses. As bats are mammals, they express innate immune receptors and inflammasome-associated proteins like humans. How bats, unlike humans, alleviate inflammatory responses while hosting several RNA viruses? Do CoVs engage inflammasome activation in bats? Despite the difficulty in performing experiments in the bat system, recent studies attempted to understand the inflammasome activation in bats and provide a basis for regulating inflammation in bats.

The bat immune cells show a striking variation in NLRP3 inflammasome activation compared with human or mouse cells. Stimulations with various NLRP3 ligands show diminished activation of the NLRP3 inflammasome in bat immune cells. Reduced priming/transcriptional up-regulation of NLRP3 is a characteristic feature of bat immune cells (Fig. 2). Also, altered LRR domain function in bats contributes to the dampened NLRP3 inflammasome function in bats. However, expression of ASC, CASP1, and IL-1β appeared to be not affected in bat immune cells compared with human cells. Because of NLRP3 priming and altered LRR domain defects, bat immune cells show lesser formation ASC speck assemblies and reduced release of IL-1β cytokine. Besides, bat cells show impaired pyroptosis and LDH release compared with human cells. Infection of bat immune cells with MERS-CoV fails to induce NLRP3 inflammasome activation and IL-1β release with unaltered viral loads. Thus, the bat’s immune system is adapted to repress NLRP3 inflammasome activation and proinflammatory responses (Fig. 2).

Flight in bats show high metabolic demand and mitochondrial oxidative phosphorylation activity. This results in DNA damage and the release of mitochondrial or nuclear self-DNA into the cytoplasm. Absent in melanoma 2 (AIM2) is a cytosolic DNA sensor that assembles inflammasome and triggers pyroptosis. Bats show complete loss of AIM2 and related Pyrin and HIN domain (PYHIN) gene family that play critical roles in nucleic acid sensing at multiple intracellular compartments. Cyclic GMP-AMP synthase (cGAS) is also a cytosolic DNA sensor that produces cyclic GMP-AMP (cGAMP) upon DNA sensing. cGAS binds to stimulator of IFN genes (STING) protein that activates type I IFN responses. Bats show dampened STING mediated IFN activation due to a mutation at the S358 phosphorylation site of the STING, which is essential for type I IFN response. Thus, bat DNA sensing system is attenuated at multiple signaling pathways to curtail inflammasome activation and type I IFN responses. In addition, a recent study unravels variation in amino acid residues of bat CASP1 (D365N and R371Q) that reduces its enzymatic activity essential for IL-1β processing and pyroptosis. A recent study by Goh et al. identifies that some of the bat species, which show intact CASP1 function, show abolished IL-1β cleavage as a complementary strategy to dampen inflammation. Identification of these mutations in bat CASP1 indicates a likely adaptation of inflammasome assembling proteins in bats to dampen inflammatory responses to RNA viruses.

Polyinosinic-polycytidylic acid (poly(I:C)) is a double-stranded RNA ligand that mimics viral RNAs. Poly(I:C) stimulation experiments indicate that bat and human cells trigger antiviral responses by up-regulating IFN-β; however, only human cells express robust TNF cytokine levels. The bat cells utilize a unique repression system to reduce TNF expression in response to viral RNA mimics. TNF signaling facilitates activation of the inflammasome, inflammatory responses, and lytic forms of cell death. Thus, TNF regulating mechanisms of bats may further limit excessive inflammasome activation and release of proinflammatory cytokines and DAMPs compared with humans.

6 | BAT VARIATIONS IN KEY INFLAMMASOME MEDIATORS—ASC, CASP1, GSDMD, AND IL-1β

Though the bats show reduced NLRP3 expression, they appear to show expression of the ASC, CASP1, and GSDMD proteins critical for the inflammasome activation by various innate immune receptors. To fully delineate the bat adaptations in the inflammasome pathway, we mapped functionally critical residues of ASC, CASP1, and GSDMD in bats to comprehensively examine variations (Fig. 3). The ASC consists of PYD and CARD domains that facilitate homotypic interactions and filamentous/speck-like structures. The assembly of ASC and its polymerization is essential to recruit and activate the CASP1. Our functional residue mapping analysis shows that the key residues essential for ASC oligomerization (PYD) and CASP1 recruitment (CARD)
FIGURE 3  Bats show variation in functionally essential residues of inflammasome components.

(A) Domain architectures of the ASC and CASP1 proteins. Asterisks indicate critical residue positions promoting homotypic interactions and oligomerization of ASC and CASP1; cleavage site residues of CASP1 are indicated in blue; the catalytically important residue of the CASP1 are indicated in red. (B) Common and scientific names of organisms that are used in panels C and D for comparing the conservation of inflammasome components. Bat species are categorized into Yangochiroptera and Yinpterochiroptera suborders. (C and D) Multiple sequence alignments of specific regions of ASC (C) and CASP1 (D) from human, mouse, and bat species. The human ASC and CASP1 protein sequences are used as references to define the numbers of amino acid residues. The critical residue positions are colored as represented in panel A. The conserved residue positions are highlighted in red, the conserved cleavage site residues are in blue, and the mutated residue positions in green. The D122 in panel D represents the cleavage site in mice that correspond to the D119 cleavage site of the human CASP1.

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H. sapiens  MGRARAILDAILAILKLTAELEKKKKLKLVEPLSDGGYRIFRAGALLSLDAHQLTDLV

M. musculus  MGRARAILDAILAILKLTAELEKKKKLKLVEPLSDGGYRIFRAGALLSLDAHQLTDLV

P. alecto  MGRARAILDAILAILKLTAELEKKKKLKLVEPLSDGGYRIFRAGALLSLDAHQLTDLV

F. vamppyrus  MGRARAILDAILAILKLTAELEKKKKLKLVEPLSDGGYRIFRAGALLSLDAHQLTDLV

H. armiger  MGRARAILDAILAILKLTAELEKKKKLKLVEPLSDGGYRIFRAGALLSLDAHQLTDLV

R. ferrumequinum  MGRARAILDAILAILKLTAELEKKKKLKLVEPLSDGGYRIFRAGALLSLDAHQLTDLV

M. davidii  MAGRARAILDAILAILKLTAELEKKKKLKLVEPLSDGGYRIFRAGALLSLDAHQLTDLV

E. fuscus  MAGRARAILDAILAILKLTAELEKKKKLKLVEPLSDGGYRIFRAGALLSLDAHQLTDLV

N. natalensis  MAGRARAILDAILAILKLTAELEKKKKLKLVEPLSDGGYRIFRAGALLSLDAHQLTDLV

H. sapiens  HRAAL1AVRTYNVEKLVDELGYCLVQYVRAETPENFSDMKLSFSFTAPKNWTNKGDLFQLRQESTYLV

M. musculus  HRAAL1AVRTYNVEKLVDELGYCLVQYVRAETPENFSDMKLSFSFTAPKNWTNKGDLFQLRQESTYLV

P. alecto  HRAAL1AVRTYNVEKLVDELGYCLVQYVRAETPENFSDMKLSFSFTAPKNWTNKGDLFQLRQESTYLV

F. vamppyrus  HRAAL1AVRTYNVEKLVDELGYCLVQYVRAETPENFSDMKLSFSFTAPKNWTNKGDLFQLRQESTYLV

H. armiger  HRAAL1AVRTYNVEKLVDELGYCLVQYVRAETPENFSDMKLSFSFTAPKNWTNKGDLFQLRQESTYLV

R. ferrumequinum  HRAAL1AVRTYNVEKLVDELGYCLVQYVRAETPENFSDMKLSFSFTAPKNWTNKGDLFQLRQESTYLV

M. davidii  HRAAL1AVRTYNVEKLVDELGYCLVQYVRAETPENFSDMKLSFSFTAPKNWTNKGDLFQLRQESTYLV

E. fuscus  HRAAL1AVRTYNVEKLVDELGYCLVQYVRAETPENFSDMKLSFSFTAPKNWTNKGDLFQLRQESTYLV

N. natalensis  HRAAL1AVRTYNVEKLVDELGYCLVQYVRAETPENFSDMKLSFSFTAPKNWTNKGDLFQLRQESTYLV
appeared to be conserved with a notable exception at F59 and E130 residues (Fig. 3). Bats that belong to Yangochiroptera suborder (M. natalensis, M. davidii, and E. fuscus) show variation at one of the critical CARD (E130N) and PYD (F59S) residues required for ASC filament formation and CASP1 interaction (Fig. 3C). We also observed a critical variation at one of the CARD residues (Y82H) of the CASP1 of multiple bat species (Fig. 3D). INCA (inhibitory CARD) is a natural human CARD only protein that shares high similarity with the CARD domain of the CASP1 and interferes with its function. Interestingly, the INCA protein shows Y82H mutation (like bat CASP1), making it unsuitable for CARD-mediated polymerization, but interfering with the CASP1 function. These variations in ASC and CASP1 may interfere in their interaction and assembly of inflammasome complexes.

CASP1 contains a CARD domain, a large (p20) and small (p10) subunits. Interdomain linker regions separate these domains.120–122 Once activated, CASP1 is self-cleaved at aspartic acid residues of the linker regions to generate enzymatically active p20 and p10 subunits.120–123 Mouse CASP1 shows variation at D119; however, D122 of the mouse CASP1, which is in close vicinity, compensates this variation and retains self-cleavage in linker regions123 (Fig. 3D). Our analysis indicates a notable variation at several cleavage site residues located in linker regions of bat CASP1 of both Yangochiroptera and Yinpterochiroptera suborders compared with human CASP1 (Fig. 3D). We also did not observe any compensatory aspartic acid residues in adjacent amino acid sequences suggesting a possible exclusion of self-cleavage at these CASP1 regions. However, the residue crucial for CASP1 catalytic activity (C285) is highly conserved in bats and humans. We observed D365N and R371Q mutations in P. vampyrus and P. alecto, which were recently reported by Goh et al.,109 to affect the catalytic activity of the CASP1 (Fig. 3D). Thus, it appears that a wide range of mutations may interfere with the bat CASP1 function following inflammasome activation.

GSDMD is a pore-forming protein that consists of a CASP1 cleavage site (D275) at the linker region of its N and C-terminal domains21,32 (Figs. 4A and 4B). CASP1-mediated cleavage of the GSDMD liberates the N-terminal domain that assembles into a pore structure on the cell membrane and triggers pyroptosis.19,30–32 Caspase-3 (CASP3) also cleaves the GSDMD N-terminal domain at D87, which abolishes pore-forming activity.30,124 (Figs. 4A and 4B). The positively charged residues in α1-helix of the GSDMD N-terminal domain (residues R7, R10, and R11) are critical for its interaction with the cell membrane and subsequent oligomerization into a pore structure125–127 (Figs. 4A and 4B). A recent study by Goh et al.,109 demonstrated the cleavage of bat GSDMD into N- and C-termini and pro-IL-1β into a mature form of IL-1β by caspases. Consistent with this study, we observed the cleavage sites of CASP1 and CASP3 in GSDMD are highly conserved in bats and humans (Fig. 4B). However, R7 and R11 residues in α1-helix of the bat GSDMD are mutated to uncharged residues suggesting a potential restriction of pore formation and pyroptosis (Fig. 4B). The caspase cleavage sites of IL-1β are conserved in bats and humans (Figs. 4A and 4C). Goh et al.,109 identified that the amino acid residues (residue positions 110, 111, and 118) surrounding the cleavage site D116 determine the IL-1β cleavage ability in M. davidii and Palecto by CASP1. The cleavage ability of the IL-1β in these bat species shows an inverse complementary relationship with their CASP1 activity.109 Previous studies indicate critical residues of IL-1β, which are necessary for binding to IL-1 receptor.128–130 The IL-1 receptor binding residues of bat IL-1β are largely conserved; however, few variations were noted at I172, K209, and E221 residues in some of the bat species (Fig. 4C). Overall, the analysis of functionally essential residues in ASC, CASP1, GSDMD, and IL-1β of bats reveals molecular adaptation of bats to restrict inflammasome mediated proinflammatory responses. Unlike the NLRP3, the expression of ASC, CASP1, and GSDMD is seen in bats, which might have forced them to acquire mutations at crucial residues to restrict inflammasome functions. What evolutionary pressures drive bat variations that dampen inflammasome activation? The causative triggers for the evolution of bat variations remain to be studied to suppress innate immune responses like inflammasome and STING activation. The dampened inflammasome in bat immune tolerance and inflammasome activation in driving COVID-19 severity suggests inflammasome as a target for alleviating disease severity in humans. Also, monitoring circulating inflammasome cytokines in humans may assist in identifying individuals who are likely predisposed for COVID-19 severity.

7 | CONCLUSIONS

Human cells operate diverse innate immune receptors, ALRs and NLRs, to engage inflammasome assembly and pyroptosis. Thus, human respiratory infections, like SARS-CoV-2, readily trigger inflammasome-mediated proinflammatory responses. The virus-induced elevated inflammatory responses ultimately drive human immunopathology despite type I IFN responses. Consistent findings showing elevated IL-1α, IL-1β, and LDH in COVID-19 patients substantiate the role of the inflammasome in promoting disease severity. This makes the inflammasome pathway a suitable target to alleviate COVID-19 severity and prevent immune overreaction. Unlike humans, bats are highly adapted to operate type I IFN responses and suppress the activation of the inflammasome and proinflammatory responses. The robust suppression of the NLRP3 inflammasome in bat immune cells is consistent with the ability to host several RNA viruses that primarily engage this NLRP3 inflammasome. Identification of bat ASC, CASP1, and GSDMD mutations suggest the possible evolution of bat inflammasome proteins that favor dampened inflammasome activation by innate immune receptors. It is likely that bat adaptation to restrict inflammasome activation allows them to tolerate RNA viruses like SARS-CoV, SARS-CoV2, and MERS-CoV (beta-CoVs) without inducing adverse inflammatory responses and tissue damage. However, the SARS-CoV-2-induced inflammasome activation drives disease pathogenesis in human infections. We still do not understand why only some of the CoVs, particularly bat-beta CoVs, that infected humans are highly pathogenic but not others. Do these infected human CoVs differ in their ability to activate the inflammasome in human cells? Do inflammasome activation signatures of the CoVs help predict the pathogenic potential
Figure 4 Variations in functionally important residues of bat CASP1 substrates, GSDMD and IL-1β proteins.

(A) Cartoon representing GSDMD, pro-IL-1β and mature form of IL-1β. Asterisks indicate critical residue positions required for IL-1β function; cleavage site residues are indicated in blue; critical lipid membrane binding residues of the GSDMD are indicated in red. (B and C) Multiple sequence alignments showing critical residues of GSDMD (B) and IL-1β protein (C). The human (H.sapiens) GSDMD and IL-1β protein sequences are used as a reference sequence to represent the amino acid residue numbers. The conserved amino acid residues are highlighted in red. The conserved cleavage site residue positions are highlighted in blue. The mutated amino acid residue positions are highlighted in green.

Overall, inflammasome activation in bats and humans provides the basis for species-specific innate immune activation, viral tolerance, and immunopathology in pathogenic CoV infections. Further exploration of bat and human immune activation mechanisms unravel fundamental mechanisms determining protective and pathologic outcomes in CoV infections.

Authorship
S.K. conceptualized and performed protein sequence analysis. S.N., D.J., and S.K wrote and edited the manuscript.

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Disclosures
The authors declare no conflicts of interest.

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