SARS-CoV-2 Nsp13 is a viral RHIM protein promoting cell death linked to Z-RNA sensing and ZBP1-RIPK3 signaling

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Running title: Bat-associated RNA viruses employ viral RHIMs and regulate host cell death.

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**ABSTRACT**

Interferons and regulated cell death pathways counteract virus spread and mount immune responses, but their deregulation often results in inflammatory pathologies. The RIP-homotypic interaction motif (RHIM) is a conserved protein domain critical for assembling higher-order amyloid-like signaling complexes inducing cell death. A few DNA viruses employ viral RHIMs mimicking host RHIMs to alleviate cell death-mediated antiviral defenses. Whether RNA viruses operate such viral RHIMs remains unknown. Host RHIM-protein signaling promotes lung damage and cytokine storm in respiratory RNA virus infections, arguing the presence of viral RHIMs in RNA viruses. Here, we report the identification of novel viral RHIMs in Nsp13 and Nsp14 of SARS-CoV-2 and other bat RNA viruses and provide a basis for bats as the hosts for the evolution of RHIMs in RNA viruses. Nsp13 expression promoted CoV-RHIM-1-dependent cell death after SARS-CoV-2 infection, and its RNA-binding channel conformation was critical for cell death function. Nsp13 interacted and promoted the formation of large insoluble complexes of ZBP1 and RIPK3. Unlike DNA virus RHIMs, SARS-CoV-2 Nsp13 did not restrict host RHIM-dependent cell death. Instead, it promoted ZBP1-RIPK3 signaling-mediated cell death dependent on intracellular RNA ligands. Intriguingly, SARS-CoV-2 genome fragments showed high Z-RNA forming propensity which bound to Z-RNA sensing Zα domains and promoted Nsp13-dependent cell death. Our findings reveal the functional viral RHIMs in RNA viruses and the role of SARS-CoV-2 Nsp13 in cell death associated with Z-RNAs and ZBP1-RIPK3 signaling, allowing the understanding of mechanisms of cellular damage and cytokine storm in respiratory virus infections and COVID-19.
**INTRODUCTION**

Pathogenic RNA virus infections often result in uncontrolled tissue damage and inflammatory responses which prime disease pathogenesis (Balachandran and Mocarski, 2021; Diamond and Kanneganti, 2022; Flerlage et al., 2021). Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) infection is associated with mild to severe respiratory infections and cause coronavirus disease 2019 (COVID-19) (Yang et al., 2020; Zhou et al., 2020). COVID-19 patients develop pneumonia and acute respiratory distress syndrome (ARDS), which lead to the failure of respiratory function (Flerlage et al., 2021; Yang et al., 2020; Zhou et al., 2020). However, the host and viral mechanisms underlying disease pathogenesis in COVID-19 and other human CoV infections are unclear. Aberrant cell death and cytokine storm have been reported to be associated with COVID-19 pathogenesis (Diamond and Kanneganti, 2022; Flerlage et al., 2021; Nagaraja et al., 2022). RNA viruses, including CoVs, are adapted to evade host defense responses to promote their spread and disease severity in the infected hosts (Garcia-Sastre, 2017; Minkoff and tenOever, 2023). SARS-CoV and SARS-CoV-2 infections delay type I interferon (IFN) production and cause dysregulated immune and proinflammatory responses in the lung (Flerlage et al., 2021; Minkoff and tenOever, 2023).

Regulated cell death promotes innate host defense responses to virus infections (Balachandran and Mocarski, 2021; Basavaraju et al., 2022; Kaiser et al., 2013). Regulated inflammatory cell death not only removes the viral replication niche by eliminating the virus-infected cells and but mounts protective immune responses and repair processes (Flerlage et al., 2021; Thomas et al., 2020). Virus infection-induced cytokines and type I IFNs also promote inflammatory cell death as a protective host defense mechanism (Karki et al., 2021a; Thomas et al., 2020). Human receptor-interacting serine/threonine protein kinase 1 (RIPK1), RIPK3, Toll/IL-1 receptor domain-containing adapter protein-inducing IFN-β (TRIF), and Z-nucleic acid binding protein 1 (ZBP1) are inflammatory cell death signaling proteins (Kesavardhana et al., 2020b; Mompean et al., 2019). These proteins have a conserved receptor interacting protein (RIP)-homotypic interaction motif.
(RHIM), which confers protein-protein interactions of these cell death proteins. RHIM-mediated protein-protein interactions trigger the formation of higher-order necosome complexes that structurally resemble an amyloid fibrillar assembly (Li et al., 2012; Mompean et al., 2019; Sun et al., 2002). The necosome complex further drives mixed lineage kinase domain-like (MLKL) mediated inflammatory cell death, called necroptosis and apoptosis. Necroptosis is essential for eliminating virus-infected cells, blocking viral spread and promoting aggravated inflammation and tissue damage in some cases (Balachandran and Mocarski, 2021; Basavaraju et al., 2022; Kaiser et al., 2013; Thomas et al., 2020). RIPK3 and ZBP1 are the key triggers of necroptosis during viral infections (Guo et al., 2018; Kesavardhana et al., 2020a; Kuriakose et al., 2016; Nogusa et al., 2016; Thapa et al., 2016; Upton et al., 2010, 2012; Wang et al., 2014; Zhang et al., 2020). These proteins are also associated with activating inflammasome-driven pyroptosis, and a multifaceted cell death modality called PANoptosis (Basavaraju et al., 2022; Christgen et al., 2020; Karki and Kanneganti, 2023; Nagaraja et al., 2022). RHIM-driven cell death pathways have evolved to destroy viral replication platforms and activate immune responses. Thus, viruses, in turn, have evolved to modulate the host RHIM protein mediated signaling (Kaiser et al., 2013; Mompean et al., 2019). Few DNA viruses have evolved to mimic host RHIMs to mediate host evasion strategies. These viral RHIMs compete for RHIM-mediated interaction with RIPK3 to prevent its interaction with RIPK1 or ZBP1 and the formation of cell death signaling complexes (Guo et al., 2018; Guo et al., 2015; Pham et al., 2019; Steain et al., 2020; Upton et al., 2010, 2012; Wang et al., 2014). Thus, these viral RHIM decoys inhibit host cell death mechanisms by controlling functional cell death signaling complex assembly and help continue viral replication and spread. It is unknown whether similar viral RHIMs are operated by RNA viruses, particularly pathogenic RNA viruses that have caused human outbreaks and pandemics. Aberrant activation of apoptosis and necroptosis, severe inflammation, and necrotic lung damage are the clinical signs observed during pathogenic influenza and SARS-CoV-2 infections (Flerlage et al., 2021; Karki and Kanneganti, 2023; Li et al., 2020; Li et al., 2020; Li et al., 2023; Nagaraja et al., 2022). This aberrant
cellular damage and inflammation suggest uncontrolled activation of regulated cell death in the lung and a possible lack of viral-RHIM traits in pathogenic respiratory viruses.

Here, we report the identification of novel viral RHIMs of CoVs (CoV-RHIMs) in Nsp13 and Nsp14 proteins. CoV-RHIM in Nsp13 of pathogenic CoVs (SARS-CoV, MERS-CoV and SARS-CoV-2) is highly conserved and retain RHIM-sequence features compared to other human-infected CoVs, whereas CoV-RHIMs in Nsp14 were conserved in less pathogenic human CoVs. Also, our observations indicate bat RHIM-proteins in driving the evolution of viral RHIMs in RNA viruses. Unlike RHIM-proteins of DNA viruses, CoV-RHIM of SARS-CoV-2 Nsp13 promoted cell death, and the RNA binding channel of Nsp13 is critical for this function. Also, SARS-CoV-2 Nsp13 formed complexes and promoted RIPK3 and ZBP1 higher-order complex formation suggesting its ability to form RHIM-driven signaling complexes. Influenza A virus (IAV) infection and nonviral activation of ZBP1-RIPK3-dependent cell death pathway in cells lacking ZBP1 expression suggested that Nsp13 promoted cell death in the absence of ZBP1, suggesting ZBP1-like functioning of Nsp13. Further biochemical studies suggest that intracellular RNA ligands promote Nsp13 cell death activation. We show the Z-RNA hotspots of the SARS-CoV-2 RNA genome and demonstrate that SARS-CoV-2 Z-RNAs bind to Z-nucleic acid binding domains. Our observations demonstrate the evolution and operation of viral RHIMs in RNA viruses that might be linked to cellular damage and cytokine storm.

**RESULTS**

**Identification of novel CoV-RHIMs in SARS-CoV-2 Nsp13 and Nsp14 proteins**

RHIM-proteins oligomerize into stacks of β-sheets with a compact hydrophobic core and grow into a functional amyloid structure to activate signaling processes (Li et al., 2012; Mompean et al., 2019; Pham et al., 2019). RHIMs consist of core tetrad (I/V-Q-I/V/L/C-G) residues promoting the stacking of β-sheet structures (Eisenberg and Jucker, 2012; Kajava et al., 2014; Li...
et al., 2012; Mompean et al., 2019; Pham et al., 2019). Mutating these core tetrad residues inhibits the formation of functional amyloid structures by host RHIM-proteins necessary for activating cell death (Jiao et al., 2020; Kesavardhana et al., 2020a; Li et al., 2012; Newton et al., 2016; Pham et al., 2019; Steain et al., 2020; Thapa et al., 2016; Upton et al., 2010, 2012). A recent study also found that mutating core tetrad sequences, in some cases, may not completely abolish functional amyloid formation (Pham et al., 2019; Steain et al., 2020). This establishes the requirement of additional residues in RHIM other than the core tetrad that promote stacking interfaces for amyloid stability (Mompean et al., 2019; Pham et al., 2019). We sought to identify the viral RHIMs in SARS-CoV-2 and other CoVs by protein sequence and structure-based analysis. Interestingly, we identified three novel RHIM-like sequences in Nsp13 (RNA helicase) and Nsp14 (exoribonuclease) of the SARS-CoV-2 and other CoVs, which were named as CoV-RHIMs, (Figure 1A, B). These CoV-RHIMs were highly conserved in all the SARS-CoV-2 variants. The core tetrad in CoV-RHIM-1 and -3 of the SARS-CoV-2 showed high similarity with other viral and host RHIM-proteins, although the CoV-RHIM-2 was mutated in its core tetrad residues (Figure 1B, 1D-F). In addition, a Val/Ile (V/I) upstream of the core tetrad was noticeable in SARS-CoV-2 RHIM-proteins, similar to that of the host and other viral RHIM-proteins (Figure 1B). However, a conserved Asn (N) downstream of the core tetrad was not seen in CoV-RHIMs of the SARS-CoV-2 and other previously identified viral RHIM-proteins (Figure 1B). More importantly, CoV-RHIMs of SARS-CoV-2 attained β-sheet conformation in crystal structures of Nsp13 and Nsp14 proteins and are surface accessible to be able to form complexes with other RHIM proteins when they are not in the replication complex (Figure S1A) (Liu et al., 2021; Newman et al., 2021). These observations suggest that SARS-CoV-2 proteins show RHIM-like features and indicate their potential role in modulating host cell death and inflammatory responses.

Our further analysis of CoV subtypes (including bat-originated beta-CoVs) showed notable differences and distinct evolutionary patterns among them (Figure 1C-F) (Menachery et al., 2015; Zhou et al., 2020). The CoV-RHIM-1 in Nsp13 was conserved among SARS-CoV-1 (SARS-CoV
BJ01), MERS-CoV, SARS-CoV-2 and Bat-CoV RaTG13, which is one of the closest associates of SARS-CoV-2 in bats (Figure 1D). Moreover, the CoV-RHIM-1 appeared to be conserved in several other beta-CoVs from bats, suggesting the occurrence of CoV-RHIMs in bat-originated CoVs (Figure 1D). Human-CoV HKU1 is a beta-CoV that infected humans after the SARS-CoV-1 outbreak but is associated with mild respiratory illness (Cui et al., 2019). We observed mutations in the core tetrad and its upstream conserved V/I position of HKU1 CoV-RHIM-1, which are critical residues for forming a compact hydrophobic interface in the RHIM-amyloid structure (Figure 1D). CoV-RHIM-1 phylogenetic analysis showed that the HKU1 was distantly related to the pathogenic beta-CoVs (SARS-CoV-1, MERS and SARS-CoV-2), despite being classified initially as a member of beta-CoVs (Figure 1C and 1D). Human-CoV 229E and NL63 are alpha-CoVs causing mild respiratory symptoms like HKU1 (Cui et al., 2019). These CoVs also had a mutation at the first position of the core tetrad (V/I→F) (Figure 1D). These observations further suggested an intriguing variation in CoV-RHIM-1 of the infected human CoVs that distinguishes mild and severe human CoV infections. Gamma-CoVs are not known to infect humans so far. We observed that avian-CoV, one of the gamma-CoVs, consisted of a conserved core tetrad, like pathogenic beta-CoVs, indicating its potential to mimic pathogenic human CoVs (Figure 1D).

We identified two additional CoV-RHIMs in Nsp14 of the CoVs (Figure 1E and 1F). Gln (Q) and Gly (G) at positions 2 and 4 of the core tetrad are mutated in SARS-CoV-2, SARS-CoV-1, MERS, HKU1, and bat beta-CoVs (Figure 1E). Unlike beta-CoVs, the alpha-CoVs (229E and NL63) consisted of the conserved core tetrad and adjacent residues promoting RHIM-mediated amyloid formation (Figure 1E). These observations further established the CoV-RHIM variations of alpha and beta-CoVs, which were zoonotically transmitted to humans but diverged in their pathogenicity. Our observations also showed a notable variation in CoV-RHIM-1 (Nsp13) and CoV-RHIM-2 (Nsp14) of pathogenic versus less severe human CoVs, which implicate a potential link between CoV-RHIM-1 in Nsp13 and the severity of SARS-CoV-1, MERS and SARS-CoV-2.
infections \textit{(Figure 1D,E)}. The CoV-RHIM-3 in Nsp14 is highly conserved across CoVs, except in gamma-CoVs. In addition, CoV-RHIM-3 of alpha-CoVs showed high similarity to human-RHIMs. Even though alpha-CoVs are phylogenetically close to beta-CoVs, they were found to be distantly related when the phylogenetic analysis was done based on CoV-RHIM-2 and -3 of Nsp14. Nevertheless, the conserved tetrad in CoV-RHIM-3 of alpha and beta-CoVs indicate its critical requirement for CoV adaptation in host cells and perhaps to regulate host defense responses.

\textbf{Bats as the source of viral RHIM-evolution in RNA viruses}

Why CoVs consist of multiple RHIM-like sequences in nonstructural proteins, and what factors drove the evolution of viral RHIMs in CoVs? Bats are the primary reservoirs of human-infecting beta-CoVs and host several viruses like SARS-CoV-2 and other pathogenic RNA viruses (Irving et al., 2021; Menachery et al., 2015; Nagaraja et al., 2022). Identifying bat CoV-RaTG13 and other related viruses as the closest relative of the SARS-CoV-2 further elucidated the association of bat CoVs with severe human infections and pandemics (Irving et al., 2021; Zhou et al., 2020). Also, bats are the natural reservoirs for several pathogenic RNA viruses which trigger cellular damage and uncontrolled inflammatory response (Irving et al., 2021; Nagaraja et al., 2022). Bats are mammals and consist of immune gene networks like humans (Mandl et al., 2015; Pavlovich et al., 2018; Zhang et al., 2013). Bats also express RIPK3 and ZBP1, which are the potential targets for viral RHIM-mediated host modulation (Dondelinger et al., 2016; Irving et al., 2020). We observed that the core tetrad and surrounding residues in the RHIMs of bat RIPK3 and ZBP1 are conserved in both Yangochiroptera and Yinpterochiroptera suborders and are nearly identical to the RHIMs of human RIPK3 and ZBP1 \textit{(Figure 1G,1H)}. This raises the possibility that CoVs might have encountered the antiviral effects of bat RIPK3 and ZBP1 prior to their zoonotic transfer into humans. This suggests that zoonotic RNA viruses originating in bats might evolve to operate viral RHIMs. In line with this hypothesis, we found that lyssavirus rabies, picornavirus, and hantaviruses originated from bats encode RHIM-like sequences \textit{(Figure 1I)}. These observations
thus suggest the likely evolution of viral RHIMs in bat RNA viruses. When the bat RNA viruses infect humans, they may either counteract or promote human RHIM-mediated cell death contributing to virus propagation and severe pathology.

CoV-RHIM-1 of 1B domain and RNA binding channel promote Nsp13-mediated human cell death

To examine whether CoV-RHIMs are associated with cell death regulation in human cells, we generated lentiviruses expressing SARS-CoV-2 Nsp13-WT. We noticed robust cell death in human HT-29 cells transduced with Nsp13-expressing lentiviruses but not EGFP-expressing lentiviruses, followed by puromycin selection (Figure 2A). The detection of Nsp13 protein expression in HT-29 cells indicated that the cell death was not due to defective Nsp13 lentiviruses (Figure 2B). Human cell lines, A549 and HCT116, also showed robust cell death after Nsp13-lentivirus transduction (Figure 2A,2B). This suggests that SARS-CoV-2 Nsp13 expression promotes cell death in multiple human cell types. We also observed robust cell death in HT-29 cells transduced with SARS-CoV-1 and MERS-CoV NSP13 lentiviruses (Figure 2C).

SARS-CoV-2 Nsp13 consists of two RecA ATPase domains (Rec1A and Rec2A) which can unwind DNA and RNA (Chen et al., 2020; Yan et al., 2020). This helicase activity of Nsp13 is critical for the progressive elongation of the RNA-dependent RNA polymerase (RdRp) on a highly structured SARS-CoV-2 RNA genome (Chen et al., 2020; Chen et al., 2022a; Yan et al., 2020). In addition to RecA domains, Nsp13 consists of a Zinc-binding domain (ZBD), a stalk region (S), and an inserted domain 1B (here after called 1B) (Figure 2D). Rec1A, Rec2A, 1B, and a part of the S domain of the Nsp13 form an RNA binding channel, and the CoV-RHIM-1 spans to domain 1B (Figure 2D,2E) (Chen et al., 2020; Chen et al., 2022a; Yan et al., 2020). To understand how Nsp13 induces cell death, we generated strategic domain deletion constructs of the Nsp13 (Figure S1B). Immunoblotting analysis showed that all the domain deletion constructs of the
Nsp13 expressed at the protein level (Figure S1C). Deletion of 1B, Rec1A and Rec2A or only Rec1A and Rec1B showed significant inhibition of Nsp13-mediated cell death (Figure S1D). This suggested the critical role of 1B, Rec1A, and Rec2A in Nsp13-dependent cell death. In addition, deletion of the ZBD or 1B domain did not diminish Nsp13-mediated cell death (Figure S1D). Deletion of ZBD and S domains decreased the magnitude of Nsp13-mediated cell death (increased cell survival). Interestingly, deletion of 1B and ZBD-S domains showed diminished Nsp13-mediated cell death (Figure S1D). These observations indicate that 1B domain deletion in combination with Rec1A-Rec2A or ZBD-S domain deletions abolished Nsp13-dependent cell death. Thus, 1B domain role in Nsp13-dependent cell death in human cells requires RNA binding channel (formed by Rec1A, Rec2A, 1B and a part of S) function.

To further understand the role of CoV-RHIM-1 in Nsp13-mediated cell death, we mutated the core tetrad residues of the CoV-RHIM-1 to alanine (VQIG \rightarrow AAAA, named Nsp13-Tet-mut). We also swapped the CoV-RHIM-1 core tetrad of the SARS-CoV-2 Nsp13 with less pathogenic HKU1-CoV residues (VQIG \rightarrow TVLG, named Nsp13-Swap-mut)). We observed that Nsp13-Tet-mut lentiviruses did not promote cell death in HT-29, A549, and HCT-116 cells, but Nsp13-Swap-mut induced cell death similarly to Nsp13-WT (Figure 2A,2B). This implies that, unlike the viral RHIMs of DNA viruses, CoV-RHIM-1 is critical for Nsp13-mediated cell death in human cells. Nsp13-Swap-mut did not affect Nsp13-mediated cell death, perhaps due to conserved CoV-RHIM-1 conformation. Although domain deletion experiments suggested the requirement of 1B and RNA binding channel for Nsp13-mediated cell death, it was intriguing that mutating CoV-RHIM-1 in domain 1B was sufficient to abolish Nsp13-mediated cell death. We then asked why mutating CoV-RHIM-1 tetrad is sufficient to abolish Nsp13-mediated cell death. Recent structural studies indicate that the SARS-CoV-2 mini replication-transcription complex (RTC) consists of two Nsp13 molecules, one in apo-form (Nsp13-Apo) and the other in RNA-bound conformation (Nsp13-Rbound) (Figure 2E) (Chen et al., 2020; Chen et al., 2022a; Yan et al., 2020). The two Nsp13 molecules are in proximity due to an interacting interface at domain 1B. Nsp13-Rbound shows a
shift in its 1B domain orientation compared to NSP13-Apo, which is critical for facilitating fully open conformation of RNA binding channel for RNA entry (Figure 2D-F). Using AlphaFold, we predicted the structure of Nsp13-Tet-mut and found that it attains similar conformation as Nsp13-Apo and Nsp13-Rbound (Figure S1E-G). However, we observed striking conformational perturbations in the vicinity of the RNA binding channel of Nsp13-Tet-mut (Figure 2F-H and S1E-G). Structural superimposition indicated that Nsp13-Tet-mut differed from Nsp13-Apo in RNA binding channel conformation (Figure S1F-G). Nsp13-Tet-mut attained open RNA binding channel conformation similar to Nsp13-Rbound because of the significant shift in domain 1B orientation (Figure 2H). Furthermore, the Rec2A domain of the Nsp13-tet-mut showed conformational alterations compared to Nsp13-Rbound conformation. This implicated that tetrad mutations in CoV-RHIM-1 altered the conformation of domain 1B and Rec2A in the RNA binding channel of Nsp13, which explains the Nsp13-Tet-mut mediated resistance to cell death. In support of this, SARS-CoV-2 min-RTC structural studies imply that inter-Nsp13 contact through the 1B domain triggers a structural shift in one of the two Nsp13 (Nsp13-2) 1B domains leading to fully open conformation of RNA binding channel (Yan et al., 2020). Domain deletion experiments and structural analysis data further suggest that, in addition to domain 1B that spans CoV-RHIM-1, RNA binding channel was also critical for Nsp13-mediated cell death. Perhaps, RNA interaction with Nsp13 might facilitate RHIM-mediated cell death.

**Nsp13 promotes cell death in SARS-CoV-2 infected cells and is dependent on CoV-RHIM-1 and RNA binding channel**

Mutations in Nsp13 are associated with species-specific and geographical adaptation of SARS-CoV-2, and targeting Nsp13 is known to impair SARS-CoV-2 replication (Fang et al., 2007; Garvin et al., 2020; Grimes et al., 2023; Naderi et al., 2023). Also, substitutions in Nsp13 of CoVs severely affect replication and viral propagation (Fang et al., 2007; Grimes et al., 2023; Sonnleitner et al., 2022; Yue et al., 2022; Zhang et al., 2015). Thus, generating Nsp13 mutants of SARS-CoV-2 is
challenging. To understand the role of Nsp13 in SARS-CoV-2-induced cell death, we ectopically expressed Nsp13-WT, Nsp13-Tet-mut, and Nsp13 lacking RNA binding channel spanning domains (Δ1B-Rec1A-Rec2A) or only 1B domain (Δ1B) in A549 cells expressing human ACE2 (A549-ACE2). These cells were infected with SARS-CoV-2 (Hong Kong/VM20001061/2020), and cell death was measured in infected cells using Sytox green staining. Immunoblotting for SARS-CoV-2 nucleocapsid (N) protein in infected cells suggested productive SARS-CoV-2 infection and replication (Figure S1H). We found uninfected Nsp13-WT expressing A549-ACE2 cells showed increased basal-level cell death (Figure 2I). Upon SARS-CoV-2 infection, Nsp13-WT expressing A549-ACE2 cells showed significantly increased levels of cell death than infected mock cells (Figure 2I). However, Nsp13-Tet-mut, Δ1B-Rec1A-Rec2A, and Δ1B expressing cells showed diminished cell death than Nsp13-WT, similar to mock cells after SARS-CoV-2 infection (Figure 2I). This further indicates that Nsp13 is critical in promoting SARS-CoV-2-mediated cell death. To determine whether cell death is associated with SARS-CoV-2 virus propagation, viral titers in infected cell supernatants were measured using Vero-E6 cell-based plaque assay. Nsp13-WT expressing cells showed lesser viral titers in supernatant compared to mock or Nsp13-Tet-mut, Δ1B-Rec1A-Rec2A, and Δ1B expressing cells upon SARS-CoV-2 infection (Figure 2J). Cell death rate and viral titers in SARS-CoV-2 infected Nsp13-WT expressing cells indicate that Nsp13-mediated cell death restricted SARS-CoV-2 viral propagation. Overall, these observations corroborate in vitro overexpression studies and suggest that Nsp13 promotes cell death in SARS-CoV-2 infected cells, and CoV-RHIM-1 and RNA binding channel function are critical for Nsp13-mediated cell death.

SARS-CoV-2 Nsp13 interacts with human RHIM-proteins and promotes the formation of large insoluble complexes
RHIM-RHIM interactions promote protein-protein interactions to assemble cell death signaling complexes like necosome and ripoptosome to activate apoptosis and necroptosis (Cho et al., 2009; Kesavardhana et al., 2020b; Mompean et al., 2019; Mompean et al., 2018). Viral-RHIMs of MCMV and HSV-1 inhibit necroptosis by directly interfering with host protein RHIM-RHIM interactions (Guo et al., 2018; Guo et al., 2015; Kaiser et al., 2013; Upton et al., 2010, 2012). To understand Nsp13 CoV-RHIM-1 role in host-RHIM-driven cell death, we ectopically expressed Nsp13 and its mutants in HT-29 cells and treated them with TNF and SMACmimetic (TNF+SMACmim) for TNF-induced apoptosis and TNF+SMACmim+zVAD for TNF-induced necroptosis (Figure 3A). As expected, these triggers induced robust cell death in HT-29 cells (Figure 3A). Expression of Nsp13-WT, Nsp13-Tet-mut, or Nsp13-Swap-mut did not diminish TNF-induced apoptosis and necroptosis in HT-29 cells (Figure 3A). RHIM-mediated RIPK1 and RIPK3 interactions are critical for TNF-induced apoptosis and necroptosis (Kesavardhana et al., 2020b; Newton et al., 2016; Weinlich et al., 2017). Viral RHIM-proteins, M45 and ICPs, directly interact with RIPK3 and ZBP1 through RHIM-homotypic interactions and restrict virus-induced necroptosis (Guo et al., 2018; Kaiser et al., 2013; Upton et al., 2010, 2012). ORF20 of VZV also interacts with ZBP1 to regulate virus-induced apoptosis (Steain et al., 2020). We sought to determine the ability of SARS-CoV-2 Nsp13 to interact with human RIPK1, RIPK3, and ZBP1 by co-expressing these proteins in HEK-293T cells, followed by immunoprecipitation. HEK-293T cells express RIPK1 endogenously but do not express RIPK3 and ZBP1 proteins. When Nsp13, RIPK3 and ZBP1 were co-transfected in HEK-293T cells, immunoprecipitation with anti-HA-tag (RIPK3-HA) and anti-ZBP1 antibodies showed RIPK3-RIPK1-ZBP1 complex formation without Nsp13 (Figure 3B). Nsp13 did not interact with RIPK1 and RIPK3, corroborating the dispensable role of Nsp13 in TNF-induced apoptosis and necroptosis (Figure 3B). However, Nsp13 immunoprecipitation showed its interaction with ZBP1 (Figure 3B). Immunoprecipitation with anti-ZBP1 antibody did not show the interaction of Nsp13, perhaps due to antibody binding site steric constraints. These observations suggest that Nsp13 may not interfere with host RHIM-
protein interactions but can interact with ZBP1 in an ectopic expression system. To further establish SARS-CoV-2 Nsp13 association with host-RHIM proteins, we tagged Nsp13 with EGFP (Nsp13-EGFP) for capturing Nsp13 association with RHIM proteins in imaging studies (Figure 3C and S2A). Tagging EGFP did not alter Nsp13-mediated cell death and protein expression, indicating the retention of Nsp13 native conformation upon EGFP fusion (Figure 3D and S2A). Nsp13-EGFP and ZBP1-dTomato were colocalized upon co-expression, and the colocalized regions appeared as complexes (Figure 3E). We further co-expressed Nsp13-EGFP with ZBP1 tagged with HA-Tag (ZBP1-HA) and found that Nsp13 and ZBP1 colocalize in HEK-293T cells (Figure 3F). Although we did not observe Nsp13 and RIPK3 interaction in immunoprecipitation experiments, imaging studies show that Nsp13 colocalized with RIPK3 (Figure 3F). However, Nsp13 association with RIPK1 was not detectable in HEK-293T cells (Figure S2B). These observations further indicate the association of Nsp13 with the ZBP1 and RIPK3 and higher-order complex formation.

RHIM-RHIM interactions of host and viral proteins form fibrillar insoluble oligomeric complexes (Chen et al., 2022b; Mompean et al., 2019; Mompean et al., 2018; Pham et al., 2019). Recent studies demonstrated the formation of insoluble amyloid-like complexes by RIPK1, RIPK3 and ZBP1 upon interaction with viral-RHIM proteins (Chen et al., 2022b; Mompean et al., 2019; Mompean et al., 2018; Pham et al., 2019). To further understand Nsp13 association with ZBP1 and RIPK3, we visualized Nsp13, ZBP1, and RIPK3 in HEK-293T cells using DNA-PAINT imaging that enables nanometer-scale resolution for monitoring complex formation. DNA-PAINT imaging suggested spatial localization of Nsp13 with ZBP1 and RIPK3 (Figure 3G-H and S2C). A recent study showed the formation of round and rod-shaped RHIM-protein amyloid complexes using super-resolution microscopy (Chen et al., 2022b). ZBP1 appeared as rod-shaped complexes in Nsp13-ZBP1 complexes compared to Nsp13 (Figure 3G-H and S2C). Nsp13 appeared to have resided in the pockets of ZBP1. RIPK3 appeared as both round and rod-shaped structures in Nsp13-RIPK3 complexes (Figure 3G-H and S2C). The DNA-PAINT imaging indicated the
formation of round and rod-shaped complexes by ZBP1 and RIPK3. Nsp13 is in proximity to them, although Nsp13 appeared to form smaller and distinct structures than ZBP1 and RIPK3. To monitor whether SARS-CoV-2 Nsp13 can assemble amyloid-like complexes, we expressed Nsp13 and its mutants alone or in combination with RIPK3 and ZBP1 in HEK-293T cells. To preserve oligomeric complexes, cell lysates were crosslinked using a DSP cross-linker, which has a built-in disulfide in its spacer region that allows decoupling of crosslinked oligomers through treatment with disulfide reducing agents such as beta-mercaptoethanol (BME). Nsp13-WT expression led to the formation of NP-40 soluble higher-order complexes with undetectable oligomers in the insoluble fraction (Figure S2D). We observed an increased complex formation by Nsp13-Tet-mut and Nsp13-Swap-mut than Nsp13-WT in NP40-soluble fractions (Figure S2D). Nsp13-Swap-mut also formed detectable NP-40 insoluble oligomers. Thus, SARS-CoV-2 Nsp13 appeared to oligomerize into large complexes upon overexpression and mutating or swapping core tetrad of CoV-RHIM-1 further enhanced homo-oligomer formation. This indicated the predominant monomeric nature of the Nsp13-WT protein, which might oligomerize when 1B domain conformation is perturbed.

Since Nsp13 interacted with ZBP1 in HEK-293T cells, we further monitored whether Nsp13 affects ZBP1 oligomerization. We observed detectable but low levels of higher-order oligomers of ZBP1 in NP40-insoluble fraction upon transfection (Figure S2D). Co-expression of ZBP1 with Nsp13-WT showed significantly increased levels of ZBP1 oligomers in NP-40 insoluble fraction (Figure S2D). However, Nsp13-Tet-mut and ZBP1 co-expression showed a lesser magnitude of ZBP1 oligomer formation (Figure S2D). In contrast to the predominant monomeric form of Nsp13-WT, its co-expression with ZBP1 showed detectable levels of Nsp13-WT complexes in NP40-insoluble fraction and increased levels of oligomers in NP40-soluble fraction (Figure S2D). These crosslinking experiments suggested that Nsp13 assembles into CoV-RHIM-dependent higher-order complexes, and Nsp13 promotes higher-order oligomerization of ZBP1. As Nsp13-Tet-mut
already forms non-native oligomers, its expression with ZBP1 did not facilitate ZBP1 oligomerization (Figure S2D).

We further co-expressed Nsp13-WT with RIPK3 and ZBP1 and monitored higher-order complex formation. The co-expression of ZBP1 and RIPK3 led to the formation of NP-40 soluble and insoluble RIPK3 oligomers (Figure 3I). These oligomers were dissolved into monomers upon BME treatment (Figure 3I). However, ZBP1 did not show detectable oligomerization in both fractions (Figure 3I). Similarly, the co-expression of Nsp13-WT with RIPK3 triggered RIPK3 complex formation in NP40-insoluble and soluble fractions (Figure 3I). Higher order complexes of the RIPK3 were observed even when RIPK3 was expressed with Nsp13-WT and ZBP1. The magnitude of oligomer formation was lesser than that was seen in Nsp13-WT and RIPK3 co-expressed cell lysates (Figure 3I). ZBP1 formed NP40-insoluble complexes when co-expressed with Nsp13-WT or Nsp13-WT and RIPK3 (Figure 3I). Even though BME treatment decoupled the complexes of RIPK3 and ZBP1, a fraction of complexes was not completely dissolved after BME treatment, suggesting the formation of insoluble fibrillar amyloid-like oligomers with hydrophobic interfaces as reported previously for host RHIM-proteins (Figure 3I) (Mompean et al., 2018; Pham et al., 2019). These observations suggest that Nsp13 forms soluble higher-order complexes when co-expressed with RIPK3 or ZBP1. Nsp13 co-expression promotes insoluble amyloid-like complexes of RIPK3 and ZBP1. Thus, Nsp13 may preferably form soluble oligomers which further nucleate RIPK3 and ZBP1 insoluble fibrillar amyloid-like necrosome assemblies. Together, these observations indicate that Nsp13 employs viral RHIMs to promote higher-order complex formation and RNA-binding dependent cell death.

SARS-CoV-2 Nsp13 promotes ZBP1 function and mimics ZBP1 triggered cell death in the absence of ZBP1 expression

Higher-order complex formation by ZBP1 and Nsp13 after co-expression led us to examine the role of Nsp13 in ZBP1-RIPK3 signaling-specific cell death activation. ZBP1 activation triggers
IAV-induced necroptosis, apoptosis and pyroptosis cell death programs. Most transformed human cell lines do not express RIPK3 and ZBP1. L929 cells (mouse fibroblast cell line) retain ZBP1 expression and show ZBP1-dependent cell death after IAV infection. Thus, we generated Zbp1−/− L929 cells to study the role of Nsp13 in ZBP1-dependent cell death pathways. We transiently expressed Nsp13 in WT and Zbp1−/− L929 cells and infected these cells with IAV to study the role of Nsp13 in IAV-induced necroptosis (Figure 4A).

Without Nsp13 expression, Zbp1−/− cells showed significantly less cell death than WT cells after IAV infection or IAV combined with zVAD treatment (IAV + zVAD, for activating necroptosis). Nsp13-WT expression in WT cells did not alter IAV or IAV+zVAD-induced cell death and promoted faster kinetics of cell death activation (Figure 4A). Interestingly, NSP13-WT expressing Zbp1−/− cells showed a higher magnitude of cell death, which is comparable to WT-cells, suggesting the role of Nsp13 in promoting IAV-induced cell death in the absence of ZBP1 (Figure 4A). Immunoblotting analysis showed that Nsp13-WT expression in Zbp1−/− cells led to increased phosphorylated MLKL levels than Zbp1−/− cells without Nsp13 expression (Figure S2E). This indicated IAV+zVAD-mediated necroptosis activation by Nsp13 despite lacking ZBP1 expression. Also, we observed increased cleaved caspase-3 in Nsp13-WT expressing Zbp1−/− cells and no apparent differences in phosphorylated MLKL levels (Figure S2E). However, Nsp13-Tet-mut expression in Zbp1−/− cells did not reduce IAV or IAV+zVAD-induced cell death, suggesting that CoV-RHIM-1 mutations do not appear to abolish Nsp13-induced cell death in IAV infected Zbp1−/− cells (Figure 4A). Nsp13-WT and Nsp13-Tet-mut expressing WT and Zbp1−/− cells showed a very low magnitude of cell death without infection than IAV and IAV+zVAD treatment (Figure 4A).

To further understand whether Nsp13 promotes ZBP1-dependent cell death in human cells, we infected HT-29 cells with IAV+zVAD. HT-29 cells express RIPK3 but show undetectable ZBP1 expression. Expression of human ZBP1 in HT-29 cells led to increased necroptosis activation after IAV+zVAD treatment (Figure 4B). Nsp13-WT co-expression with ZBP1 enhanced IAV+zVAD-induced necroptosis in HT-29 cells (Figure 4B). In addition, staining HT-29 cells with
Annexin-V+ and Sytox green indicated that the expression of Nsp13 in HT-29 cells preferably induced necrotic-like cell death in the absence of stimulation or infection, and this cell death was further enhanced in the presence of ZBP1 (Figure S2F,S2G). These observations indicate that Nsp13 does not interfere, but perhaps promotes, ZBP1-induced cell death during IAV infection. Intriguingly, Nsp13 promoted IAV-induced cell death in the absence of ZBP1 expression.

Leptomycin-B (LMB), in combination with IFNs, is known to trigger ZBP1 activation and cell death (Jiao et al., 2020). A recent study also showed that curaxin (CBL0137) promotes ZBP1 activation and necroptosis by forming Z-DNAs (Zhang et al., 2022). We examined LMB+IFN-β and curaxin-mediated cell death in WT and Zbp1−/− L929 cells to further probe the role of SARS-CoV-2 Nsp13 in ZBP1 function (Figure 4C,4D). As expected, LMB+IFN-β treatment triggered cell death and lack of ZBP1 significantly abolished this cell death (Figure 4C). However, upon Nsp13-WT expression in Zbp1−/− cells, increased cell death was observed after LMB+IFN-β treatment compared to those without Nsp13-WT expression (Figure 4C). Curaxin treatment in WT and Zbp1−/− cells showed ZBP1-dependent cell death activation and Nsp13-WT expression led to increased cell death in Zbp1−/− cells similar to that of LMB+IFN-β treatment (Figure 4D). Nsp13-WT overexpression further enhanced cell death in WT cells after LMB+IFN-β or curaxin treatment (Figure 4C, 4D). These results further suggest that SARS-CoV-2 Nsp13 promotes ZBP1-mediated cell death, and in the absence of ZBP1 expression, Nsp13 triggers cell death in response to ZBP1 activating ligands. Why might Nsp13 have evolved to regulate ZBP1-RIPK3 signaling and induce host cell death? Type I interferons (IFN) promote ZBP1 and RIPK3 signaling and activation of cell death in viral infections (Jiao et al., 2020; Kesavardhana et al., 2017; Kuriakose et al., 2016; Newton et al., 2016; Thomas et al., 2020). Delayed type I IFN production and restricted IFN signaling due to Nsp13 and other proteins of SARS-CoV-2 facilitates host modulation and efficient virus propagation (Flerlage et al., 2021; Minkoff and tenOever, 2023). We anticipate that the dampened type I IFN signaling might restrict early-on activation of ZBP1-
RIPK3 signaling and cell death, which preferentially destroy viral replication platforms. Perhaps, Nsp13 has evolved to trigger cell death, due to insufficient ZBP1-RIPK3 signaling activation, at the late stages of the infection to facilitate viral spread, tissue damage, and inflammation. Indeed, recent studies demonstrate IFN and inflammatory cytokine-driven cell death, tissue damage, and disease pathogenesis during SARS-CoV-2 infection at later stages (Flerlage et al., 2021; Karki et al., 2022; Karki et al., 2021a; Laurent et al., 2022; Wong and Perlman, 2022).

Intracellular RNA ligands and type I IFNs regulate SARS-CoV-2 Nsp13-mediated cell death

Type I IFNs upregulate ZBP1 and MLKL expression and promote necroptosis and other programmed cell death activation in viral infections (Jiao et al., 2020; Kesavardhana et al., 2017; Newton et al., 2016; Thomas et al., 2020). SARS-CoV-2 Nsp13 is associated with innate immune evasion and restricts type I IFN signaling activation (Minkoff and tenOever, 2023). We hypothesize that type I IFN treatment may regulate Nsp13-mediated cell death. To test this, we treated L929 cells ectopically expressing Nsp13 with IFN-β to monitor cell death. Ectopic expression of Nsp13 in L929 cells promoted spontaneous cell death at the basal level and IFN-β treatment reduced this Nsp13-induced cell death (Figure S3A,S3B). This suggested that Nsp13-mediated cell death might be restricted by type I IFN-IFNAR signaling. Nsp13-mediated cell death requires CoV-RHIM in 1B domain and RNA binding channel (spanning 1B, Rec1A and Rec2A domains), suggesting RNA binding as a critical step for cell death activation by Nsp13.

Intracellular delivery of Poly(I:C) (transfection) increased Nsp13-mediated cell death, but the addition of Poly(I:C) to the cell surface did not alter Nsp13-mediated cell death (Figure S3C,S3D). As expected, IAV infection in Nsp13-expressing cells promoted increased cell death (Figure 4E,4F). However, unlike IFN-β-dependent inhibition of Nsp13-mediated cell death, IFN-β treatment after IAV infection in Nsp13-expressing cells did not reduce cell death (Figure 4E,4F). Without Nsp13 expression, IAV and IFN-β treatment showed a further increase in cell death.
levels. Furthermore, LMB treatment in Nsp13-expressing cells showed a higher magnitude of cell
death activation than in Nsp13-expressing untreated cells (Figure 4E,4F). LMB+IFN-β treatment
did not alter Nsp13-mediated cell death. These observations suggested that IFN-β did not inhibit
Nsp13-mediated cell death during IAV infection or LMB treatment, which generates intracellular
RNA ligands. Thus, these results further indicate the requirement of RNA ligand binding in Nsp13-
mediated cell death and the regulation of this cell death by type I IFNs.

The SARS-CoV-2 genome shows Z-RNA signatures and binds to Zα domains of ZBP1
The role of Nsp13 in promoting cell death, interacting with host RHIM-proteins, and its ability to
promote ZBP1 signaling suggest the association of Nsp13 with Z-RNAs. We hypothesize that the
SARS-CoV-2 RNA genome consists of base repeats favoring Z-RNA conformation and are
generated during its replication process. The repeats of purine and pyrimidine (Pu:Py),
predominantly (CG)n, (UG)n/(TG)n, (CA)n, favor Z-RNA/Z-DNA conformation (DeAntoneo et al.,
2023; Herbert et al., 2021; Kesavardhana and Kanneganti, 2020; Maelfait and Rehwinkel, 2023).
In addition, recent studies show that short interspersed nuclear elements (SINEs) with inverted
repeat sequences increase the propensity of RNA to attain Z-conformation (de Reuver et al.,
2022; Jiao et al., 2020; Tang et al., 2021; Zhang et al., 2022). We have devised a comprehensive
analysis, incorporating alternate Pu:Py, inverted tandem repeats, SINEs, and the sequences
favoring double-strand RNA formation, for identifying Z-RNA segments in the SARS-CoV-2
genome. We found that Z-RNA-like sequences were distributed across the SARS-CoV-2 genome,
but most of these sequences span ORF1a and ORF1b of the genome (Figure 5A). We further
predicted the RNA secondary structures of these potential Z-RNA hotspots using RNAstructure
prediction server (Figure 5B, S4A). Among these Z-RNA forming double-stranded RNA (dsRNA)
structures, we selected dsRNAs in which alternate Pu:Py repeats stabilized the secondary
structure. We identified several SARS-CoV-2 genome hotspots with Z-RNA-favoring sequences
using this approach. Recent studies experimentally determined the secondary structure of the SARS-CoV-2 genome in infected cells (Huston et al., 2021; Lan et al., 2022; Sun et al., 2021). The predicted secondary structures of Z-RNA sequences were mapped on the experimentally solved SARS-CoV-2 genome secondary structure to check if they retained the same structures as the prediction. Using this secondary structure comparison, the candidate Z-RNAs which retain the dsRNA conformation in experimentally solved SARS-CoV-2 genome structure were identified (Figure 5B, S4A). The identified Z-RNAs forming secondary structures span the SARS-CoV-2 Nsp2, Nsp3, Nsp4, Nsp6, Nsp13, and Nsp16 coding regions of ORF1ab (Figure S4A). By examining the experimentally determined SARS-CoV-2 genome secondary structure, we further identified additional genome sequences with Z-RNA forming propensity in Nsp1 and N-protein coding regions. Together, our rational sequence and structure-based mining of the SARS-CoV-2 genome identified the sequences with high Z-RNA forming propensity.

To further validate the ability of SARS-CoV-2 Z-RNAs (SC2-zRNA) for binding Z-RNA sensing proteins, we purified Zα domains of human ZBP1 (Zα1-Zα2) to probe its binding to SC2-zRNAs using real-time biolayer interferometry and electrophoretic mobility shift assays (EMSA) (Figure 5C). SC2-zRNAs segments were generated by in vitro transcription for binding studies with Zα1-Zα2. We mutated one of the surface-accessible serine residues in Zα2 domain to cysteine (S106C) for labeling purified Zα1-Zα2 protein. Using biolayer interferometry (BLI-Octet) we examined the real-time binding of in vitro transcribed SARS-CoV-2 RNA segments with high Z-RNA propensity (SC2-zRNA-1,2,3,6,9 and 10) to purified Zα1-Zα2 (Figure 5D,5E). In this analysis, we included poly(I:C) and a Pu:Py repeat RNA that attain Z-RNA conformation (control zRNA) when incorporated with modified guanine bases (Balasubramaniyam et al., 2018; Zhang et al., 2022). The control zRNA bound to Zα1-Zα2 protein with high association rates (despite lacking modified guanines) and poly(I:C), and buffer controls bound with very low association rates (Figure 5D,5E). Perhaps Zα1-Zα2 binding might have driven the attainment of stable Z-
conformation in control zRNA despite lacking modified bases. The SC2-zRNA bound to Zα1-Zα2 protein with higher association rates than poly(I:C) and buffer control but less than the control zRNAs (Figure 5D,5E). Notably, the dissociation of control zRNA and SC2-zRNAs from Zα1-Zα2 protein was similar (Figure 5D,5E). These observations indicate that the selected SARS-CoV-2 RNA genome segments attain z-RNA conformation and bind to Zα1-Zα2 protein without the requirement of incorporating modified bases in SC2-zRNAs. Furthermore, we performed EMSAs to monitor SC2-zRNAs interaction with Zα1-Zα2 protein. Although we attempted to label Zα1-Zα2 protein by cysteine maleimide labelling for fluorescence imaging of EMSA gels, the labeled protein was degraded rapidly. We further proceeded to use unlabeled protein for EMSA studies. We observed that control zRNA and SC2-zRNAs showed detectable shifts in zRNA sizes, indicating the binding of SC2-zRNAs with Zα1-Zα2 protein (Figure 5F). These band shifts were observed at higher Zα1-Zα2 protein concentrations, perhaps because of the faster dissociation rates observed in real-time BEI-Octet analysis (Figure 5F). The EMSA experiments further establish that SARS-CoV-2 RNA genome segments attain Z-RNA conformation and bind to Z-RNA sensing Zα domains. Since intracellular RNA ligands regulate Nsp13-mediated cell death, we tested whether SC2-zRNAs promote Nsp13-mediated cell death. Nsp13-expressing L929 cells showed enhanced cell death levels after intracellular delivery of SC2-zRNAs or Poly(I:C) compared to mock-transfected Nsp13-expressing cells (Figure S4B). In particular, SC2-zRNA-2, SC2-zRNA-6, and SC2-zRNA-10 promoted higher cell death rates than other SC2-zRNAs (Figure S4B). Control zRNAs also promoted Nsp13-mediated cell death. However, this was not as robust as SC2-zRNA induced cell death, perhaps Nsp13 preferentially binds to SC2-zRNAs. These observations suggest that SC2-zRNAs regulate SARS-CoV-2 Nsp13-mediated cell death.
DISCUSSION

Our findings in this report indicate that Nsp13 and Nsp14 of CoVs harbor putative RHIMs. Viral RHIMs reported to date were found in DNA viruses, and our study characterized viral RHIMs of RNA viruses for the first time. SARS-CoV-2 Nsp13 is a viral RHIM protein promoting human cell death programs. Our analysis further reveals bats as an essential source in evolution of viral RHIMs in pathogenic RNA viruses. It was intriguing to find that, unlike the viral RHIMs reported to date, SARS-CoV-2 Nsp13 did not inhibit cell death and instead promoted it in specific conditions. A recent study indicated a RHIM-like sequence signature in SARS-CoV-2 Nsp13 based on protein sequence analysis (Herbert and Poptsova, 2022). Our protein sequence and structure-based analysis of the CoV family revealed CoV-RHIMs in Nsp13 and Nsp14 proteins and their association with bat-originated CoVs. Moreover, the CoV-RHIMs in Nsp13 and Nsp14 showed a distinct pattern of evolution. CoV-RHIM-1 in Nsp13 showed a conserved RHIM sequence pattern in pathogenic human-CoVs and bat-associated beta-CoVs, whereas CoV-RHIMs of Nsp14 are conserved in less pathogenic human-CoVs. Our analysis might facilitate RHIM sequence-based annotation of new bat-CoVs or CoVs that have not infected humans for their possible pathogenic potential. How did Nsp13 and Nsp14 acquire RHIMs mimicking host RHIM-proteins? Bats express RIPK3 and ZBP1 proteins. The conserved CoV-RHIMs in bat-originated CoVs raise the possibility that bat RIPK3 and ZBP1 might have driven the evolution of RHIMs in CoVs to modulate their function. Also, how bats show only mild clinical symptoms despite hosting SARS-CoV-2-like viruses is unknown. RHIM-mediated cell death, the evolution of viral RHIMs, and RHIM-driven virus-host interactions might provide new clues for understanding viral tolerance in bats.

Nsp13 is a helicase and a component of the SARS-CoV-2 replication-transcription complex (Chen et al., 2022a; Newman et al., 2021; Yan et al., 2020). SARS-CoV-2 mini replication-transcription complex harbors two Nsp13 molecules in distinct conformation (apo and RNA-bound). 1B domain of the Nsp13 that constitutes CoV-RHIM-1 is essential for the interaction of apo and RNA-bound
Nsp13 molecules in the replication-transcription complex (Chen et al., 2022a; Yan et al., 2020). The RNA binding channel of Nsp13 comprises domain 1B, Rec1A, and Rec2A and is critical for holding incoming RNA genome. Of note, studies on SARS-CoV-1 and other CoV Nsp13 suggested that alterations or mutations in Nsp13 abolish its helicase activity and restrict viral propagation (Fang et al., 2007; Seybert et al., 2005; Zhang et al., 2015). Although the role of Nsp13 in SARS-CoV-2 replication and RNA genome unwinding has been described, its replication-independent functions have not been studied. Our cell death signaling, and biochemical studies indicate a critical role for the CoV-RHIM-1 and RNA-binding channel in Nsp13-mediated cell death. Also, intracellular RNA ligands and SC2-zRNAs further promoted Nsp13-mediated cell death. These observations suggest a replication-transcription complex-independent role of Nsp13 in modulating host responses, and the conformation of the RNA binding channel is critical in this case. Defective viral genomes (DVGs) generated during RNA virus replication are known to affect host innate immune responses (Genoyer and Lopez, 2019; Vignuzzi and Lopez, 2019). DVGs are the source of Z-RNAs in IAV infection, and ZBP1 senses DVGs to trigger IAV-induced cell death (Zhang et al., 2020). Also, endogenous Z-RNA sensing by ZBP1 in physiological condition triggers immunopathology (de Reuver et al., 2022; Hubbard et al., 2022; Karki et al., 2021b; Tang et al., 2021). Our in vitro functional studies revealed a possible ZBP1-like function of Nsp13 to promote programmed cell death. Recent studies show that SARS-CoV-2 infection in vitro and in infected patients generate DVGs associated with innate immune activation (Girgis et al., 2022; Xiao et al., 2021; Zhou et al., 2023). Nsp13 might act as a sensor when it is not an integral part of the replication-transcription complex and detect specific RNA ligands formed in infected cells and trigger cell death like ZBP1. Perhaps, Nsp13 detects specific RNA patterns of the SARS-CoV-2 RNA genome or DVGs that might attain Z-RNA conformation to activate cell death. An intriguing question is why Nsp13 has evolved to promote cell death, unlike other viral RHIM proteins which inhibit cell death? We anticipate that SARS-CoV-2 mediated delayed type I IFN response limit ZBP1-RIPK3 signaling and exerts suboptimal cell
death activation at an early stage of infection. Nsp13 might have evolved to trigger RHIM-dependent cell death at later stages of infection to promote virus spread. Nevertheless, future studies need to examine this question.

While this manuscript was in preparation, a new study reported that SARS-CoV-2 forms Z-RNAs in infected cells and activate ZBP1-RIPK3 signaling and cell death (Li et al., 2023). This work demonstrates that ZBP1-RIPK3 signaling promotes lung damage and inflammation during SARS-CoV-2 infection in mice. This study suggests the formation of Z-RNAs in SARS-CoV-2 infection, however, specific RNA genome regions of SARS-CoV-2 attaining Z-RNA conformation and their interaction with Z-nucleic acid binding domains (Zα domains) was unknown. Our stringent pipeline for identifying Z-RNA sequences led us to annotate high-propensity Z-RNA forming sequences in the SARS-CoV-2 genome (SC2-zRNAs). Our biochemical and real-time binding studies demonstrate the binding of SC2-zRNAs to purified Zα1Zα2 domains of human ZBP1 protein. These observations reveal SARS-CoV-2 RNA genome-derived Z-RNA ligands that might activate ZBP1 and Nsp13-mediated cell death.

METHODS

Protein sequence and structure analysis

CoV-RHIM sequences in coronaviruses, SARS-CoV-2, and other bat RNA viruses were determined based on well-characterized human and viral RHIM protein sequences and amino acid propensities that favor β-sheet amyloid structures. Coronavirus protein sequence alignments were performed using Clustal Omega multiple sequence alignment tool (from EMBL-EBI). Putative RHIM sequences were compared to annotate variation among coronaviruses. Maximum likelihood phylogenetic analysis of protein sequences was done using PhyML 3.1/3.0aLRT in Phylogene.fr, or MEGA-X and the analysis output was generated in Newick format. Phylogenetic trees were illustrated in MEGA-X (v10.1.7) using PhyML-generated Newick output formats.
Crystal structures of Nsp13 (PDB ID: 6ZSL), Apo and RNA-bound forms of Nsp13 (PDB ID: 7CXM), and Nsp14 (PDB ID: 6N0B) of coronaviruses were visualized, and ribbon and surface structure models were prepared using Chimera 1.14 or ChimeraX 1.4 protein model visualization tool.

**Plasmids and constructs**

pLVX-EF1alpha-SARS-CoV-2-Nsp13-2xStrep-IRES-Puro, pLVX-EF1alpha-SARS-CoV-Nsp13-2xStrep-IRES-Puro, and pLVX-EF1alpha-MERS-Nsp13-2xStrep-IRES-Puro were a kind gift from Prof. Nevan Krogan’s lab (University of California, San Francisco). Nsp13-Tet-mut (residues 193-196, VQIG→AAAA) and Nsp13-Swap-mut (residues 193-196, VQIG→TVLG) constructs were generated by site-directed mutagenesis using overlap extension PCR. The construct expressing Nsp13-EGFP was generated by subcloning the sequence of EGFP from pLVX-EF1alpha-EGFP-2xStrep-IRES-Puro to pLVX-EF1a-SARS-CoV-2-Nsp13-2xStrep-IRES-Puro using overlap extension PCR. Nsp13 domain deletions constructs were generated using overlap extension PCRs in the same vector background of the SARS-CoV-2 Nsp13 expression construct. The coding sequence of human ZBP1 (hZBP1) was amplified from pCMV-Human-ZBP1 cDNA clone expression plasmid (Sino Biological Inc., HG19385-UT) and subcloned into pLVX-EF1alpha-2xStrep-IRES-Puro backbone using EcoRI and BamHI restriction sites. The HA-tag sequence was fused at the C-terminus of ZBP1 through PCR primers. The construct expressing hZBP1-dTomato was generated by fusing dTomato coding sequence at the C-terminus of ZBP1 by overlap extension PCR. Other plasmids used in the study, pcDNA3-HA-RIPK3 (78804) and pcDNA3-FLAG-RIPK1 (78842) were procured from Addgene.

**Cell Culture, transfections, and stimulations**
Cell lines used in this study, A549 (lung carcinoma), HT-29 (human colorectal adenocarcinoma), HCT-116 (human colorectal adenocarcinoma), L929 (mouse connective tissue fibroblast), HEK-293T (human embryonic kidney) and MDCK (Madin-Darby canine kidney) were obtained from National Centre for Cell Science (NCCS) cell repository and were authenticated by STR profile analysis. Cell lines were tested to be negative for mycoplasma contamination. A549-ACE2 and Vero-E6 cells were obtained from Dr. Shashank Tripathi’s lab (Indian Institute of Science). All the cell lines were cultured at 37°C and 5% CO$_2$ in DMEM growth medium (Thermo Fisher Scientific, 11995040) supplemented with 10% v/v FBS (Thermo Fisher Scientific, 10270106), Antibiotic-Antimycotic (Thermo Fisher Scientific, 15240062) and non-essential amino acids (Thermo Fisher Scientific, 11140-050). For transient expression of proteins, plasmids were transfected into specific cell lines using Xfect (TakaraBio, 631318) or Lipofectamine 2000 (Thermo Fisher Scientific, 11668-019) transfection reagents in reduced serum media Opti-MEM (Thermo Fisher Scientific, 31985070).

For apoptosis and necroptosis activation, HT-29 cells were treated with 30µM Z-VAD(OMe)-FMK (Cayman Chemical, 14463) and 500nM SMAC mimetic SM-164 (MedChemExpress, HY-15989). 2 h after treatment, 100ng ml$^{-1}$ recombinant human TNF-alpha (Abclonal, RP00001) was added. Influenza A virus (IAV)-infected L929 cells were treated with 30 µM Z-VAD(OMe)-FMK for inducing necroptosis. To activate ZBP1-specific cell death, L929 cells were treated either with a combination of 5ng ml$^{-1}$ Leptomycin-B (Sigma-Aldrich, L2913) and 100ng ml$^{-1}$ IFN-b (Abclonal, RP01076) and or with CBL0137 (Cayman Chemicals, 19110).

**IAV infection**

Influenza A virus (A/WSN/1933) was generated using 8 plasmid reverse genetics system and propagated in the MDCK cell line to obtain progeny 1 (P1) virus stocks. For IAV infection experiments, cells were seeded in DMEM (Thermo Fisher Scientific, 119905040) supplemented with 10% FBS. 24 h after seeding cells, the media was replaced with DMEM lacking sodium...
pyruvate (Sigma Aldrich, D6171), and cells were infected with IAV. 2 h post-infection, 10% FBS was added to the cells and real-time cell death analysis was performed using IncuCyte S3 Live-Cell Analysis instrument (Sartorius).

**Lentivirus transduction for generating Zbp1-knockout L929 cells and stable transgenic protein expression**

All the cell lines were maintained in DMEM containing 10% FBS and 1% Antibiotic-Antimycotic solution. Lentiviral particles expressing Cas9 were generated by transfecting HEK-293T cells with LentiCas9-Blast, psPAX2, and pMD2.G plasmids using Xfect transfection reagent in Opti-MEM media. Lentiviral supernatants were harvested 48 hours post-transfection. L929 cells were then infected with Cas9-lentiviruses in the presence of polybrene (Sigma-Aldrich, TR-1003-G) to obtain cells stably expressing Cas9 protein. The cells were selected with Blasticidin (Thermo fisher Scientific) and maintained in culture. Lentivirus stocks were prepared using transfer plasmid encoding guide-RNA (gRNA) targeting Zbp1 (Sequence: GTCCTTTACCGCCTGAAGA). L929 cells stably expressing Cas9 were infected with Zbp1 gRNA-lentiviruses in the presence of polybrene and selected with Puromycin (Sigma-Aldrich). Loss of ZBP1 protein expression after the selection was confirmed by immunoblotting analysis with anti-ZBP1 antibody.

To stably express Nsp13 and other domain deletion constructs of Nsp13 in A549, HCT-116, and HT-29 cells, lentivirus particles for transducing the cells were generated in HEK293T cells by transfecting Nsp13 expression constructs along with pCMV-VSVG and psPAX2 plasmids. Lentivirus supernatants were harvested 48 h after transfection, supplemented with an additional 10% FBS and stored at -80°C. The cells were transduced with lentivirus in the presence of 12mg ml⁻¹ of polybrene and subjected to selection with Puromycin (Sigma-Aldrich, P8833).

**SARS-CoV-2 Infection and virus titer assays**
Constructs expressing SARS-CoV-2 Nsp13-WT, Nsp13-Tet-Mut, Δ1B-Rec1A-Rec2a, Δ1B were transfected in A549 cells stably expressing ACE2 (A549-ACE2). 24 h after transfection, cells were infected with the SARS-CoV-2 virus (Isolate Hong Kong/VM20001061/2020, NR-52282, BEI Resources, NIAID, NIH) at an MOI of 3. Virus inoculum was prepared in DMEM (Thermo Fisher Scientific, 119905040) supplemented with 2% FBS and adsorption onto cells was allowed for 1 hour at 37°C. After 1 h, DMEM containing 2% FBS was added to cells. 48 h after infection, cells were fixed using 4% formaldehyde and subjected to SYTOX green staining for cell death analysis. Whole cell lysates were collected for immunoblotting analysis of SARS-CoV-2 N-protein.

Viral titers were estimated by plaque assay using Vero E6 cells. At 90% confluency after seeding, cells were incubated at 37°C, with 150 μl of increasing dilutions of virus supernatant collected from SARS-CoV-2 infected A549-ACE2 cells transiently expressing Nsp13-WT, Nsp13-Tet-Mut, Δ1B-Rec1A-Rec2A, Δ1B proteins. The virus inoculum was removed after 1 h and the cells were overlaid with 0.6% Avicel (Dupont, RC-591) prepared in DMEM containing 2% HI-FBS. After 48 h of incubation at 37°C, the cells were fixed with 4% paraformaldehyde. Following fixation, the cells were stained with Crystal Violet (Sigma Aldrich, C0775) for 20 minutes, and plaques were visualized.

Ethics statement

This study was performed according to the guidelines and in compliance with institutional biosafety guidelines (IBSC/IISc/11/2020; IBSC/IISc/32/2021). SARS-CoV-2 experiments were conducted at viral Biosafety level-3 facility and IAV-WSN experiments were conducted at Biosafety level-2 facility at Indian Institute of Science (IISc).

Real time cell death analysis

Real-time cell-death assays were performed using a two-color IncuCyte S3 Live-Cell Analysis instrument (Sartorius). Cell lines were seeded in 12 well plates and treated with cell death
stimulating agents or infected with viruses. Dead cells were stained with 20nM Sytox Green (Thermo Fisher Scientific, S7020), a cell-impermeable DNA-binding fluorescent dye rapidly entering dying cells after membrane permeabilization and fluoresces green. The resulting images and the fluorescence signals were analysed using IncuCyte S3 software, which provides a count for the number of Sytox Green-positive cells. This data was plotted using GraphPad Prism 9.0 software.

**Immunoprecipitation studies for probing Nsp13 association with host RHIM-proteins**

HEK-293T cells were seeded in 100mm dishes and transfected with the constructs expressing Nsp13, RIPK3 and ZBP1 for co-immunoprecipitation studies. 48 h after transfection, the cells were lysed in NP-40 lysis buffer (1.0 % NP40, 50mM Tris, pH 8.0, 150mM NaCl supplemented with protease inhibitor and phosphatase inhibitor cocktails), and lysates were cleared by centrifugation at 12000 rpm for 20 min. The whole cell lysates were incubated with 10 μg of anti-Streptag-Antibody and 2 μg of the indicated primary antibodies for 4–5 h at 4°C. Protein A/G PLUS-Agarose (Santa Cruz Biotechnology, 38221990) beads were washed with 1XPBS, incubated for 30 min with 3% BSA for blocking, and added to the samples. After 2 h of incubation at 4°C, the agarose beads were then collected by centrifugation at 1000 rpm and washed three times with 1:1 PBS and NP-40 lysis buffer. Immunoprecipitates were eluted from the beads in the sample buffer and subjected to immunoblotting analysis.

**Capturing oligomeric complexes in soluble and insoluble fractions of cellular lysates**

HEK-293T cells transiently transfected with indicated clones were subjected to lysis with NP-40 lysis buffer (1.0% NP40, 50mM HEPES, pH 8.0, 150mM NaCl with protease inhibitor and phosphatase inhibitor cocktails) containing 2mM Pierce Premium Grade DSP (Thermo Fisher Scientific, PG82081). Whole cell lysates were collected and incubated at room temperature for...
45 min. The reaction was stopped by adding 1M Tris-HCl and incubating for 15 minutes. Whole cell lysates were centrifuged at 10,000 rpm for 15–20 min to separate insoluble fractions (pellet) and soluble fractions. Insoluble fractions (pellets) were washed using lysis buffer and dissolved in NP-40 lysis buffer. Soluble and insoluble lysates were mixed with 4x sample buffer with or without β-mercaptoethanol (BME) and subjected to SDS-PAGE and immunoblotting analysis to monitor the oligomeric status of complexes.

**Immunoblotting analysis**

The cells were lysed using NP40-lysis buffer and added sample-loading buffer containing SDS and BME. The lysates were resolved in 8-12% SDS-PAGE gels, and the gels were subjected to transfer onto the PVDF membrane. The membranes were blocked in 5% skimmed milk at room temperature for 1 h and further incubated with primary antibodies overnight at 4°C. The membranes were washed using Tris-buffered saline with tween 20 (TBST), subjected to horseradish peroxidase (HRP)-conjugated secondary antibodies for 1h at room temperature, and then washed with TBST. The primary antibodies used in the study were anti-ZBP1 (Adipogen, AG-20B-0010-C100), anti-Strep Tag (Qiagen 34850), anti-p-MLKL (Cell Signaling Technology, 37333), anti-IAV N-Protein (Invitrogen, PA5-32242), anti-CASP3 (Cell Signaling Technology, 9662), anti-HA Tag (Invitrogen,26183), anti-Flag (Invitrogen, MA1-91878),anti-RIPK1 (Cell Signaling Technology, 3493), anti-GAPDH (Invitrogen,15738), anti-SARS-CoV-2 N protein (Cell Signaling Technology, 33717). HRP-conjugated secondary antibodies used in this study were Jackson Immuno Research Laboratories anti-rabbit (111-035-047) or anti-mouse (315-035-047) antibodies. Blots were developed using Immobilon Forte Western HRP substrate (Millipore, WBLUF0500) and visualized using Image Quant LAS500 or Image Quant 800 (Cytiva, Amersham).
Immunofluorescence and DNA-PAINT imaging

HEK-293T cells were transfected with the respective plasmids (Zbp1-HA, ZBP1-dTomato, RIPK1-FLAG, and RIPK3-HA) along with Nsp13-EGFP. 20-22 h after transfection, the cells were fixed with 4% paraformaldehyde in 1X PBS (ChemCruz, sc-281692) for 20 minutes at room temperature. After fixing, cells were permeabilized using 0.1% Triton X-100 for 10 min at room temperature. Nonspecific binding was blocked using 3% BSA (GBiosciences, RC1021). Cells were incubated with anti-HA Tag (Invitrogen, 26183) or anti-FLAG Tag antibody for 1 h at room temperature. The cells were then subjected to staining with Alexa Fluor 568-conjugated anti-mouse IgG (Invitrogen, A-11004; 1:500) for 30 mins at room temperature. After each step, the cells were washed twice with 1X PBS. The cells were counterstained with DAPI. All images were acquired using Olympus FV 300 Confocal microscope system.

For DNA points accumulation for imaging in nanoscale topography (DNA-PAINT) imaging, HEK-293T cells were seeded and co-transfected with constructs expressing Nsp13-EGFP along with ZBP1-HA and RIPK3-HA separately. 20-22 h after transfection, the cells were fixed for 15 min using 4% paraformaldehyde (Electron Microscopy Sciences, 15710), preheated to 37°C and subjected to 5 washes with PBS, pH 7.4. 1 mg ml⁻¹ sodium borohydride solution in PBS was used to quench the free aldehyde groups, followed by 5 washes with PBS. The cells were then permeabilized with 0.5% Triton X-100 in PBS for 15 min, followed by 2 washes with PBS. The cells were then incubated with 3% Bovine Serum Albumin (Sigma, A4503) in PBS for 45 min for blocking, followed by 2 washes with PBS. 1:200 dilution of anti-GFP (Invitrogen, A-11122) and anti-HA tag (Invitrogen, 26183) primary antibodies were prepared in 3% Bovine Serum Albumin, added to the wells, and incubated overnight in the dark at 4°C followed by 2 washes with PBS. 1:200 dilution of the docking-strand conjugated anti-mouse and anti-rabbit secondary antibody in 3% Bovine Serum Albumin was added to the wells and incubated for 45 minutes at room
temperature, followed by 2 washes with PBS. Gold nanoparticles diluted at a 1:5 ratio in PBS was added to the chamber and incubated for 5 min followed by 5 washes with PBS.

Microscopy was performed on a Nikon Ti-2 eclipse microscope equipped with a motorized HTIRF, perfect focus system (PFS), and a Teledyne Photometrics PRIME BSI sCMOS camera. Illumination using 561 nm wavelength lasers was done using the L6cc Laser combiner from Oxxius Inc., France. Imaging was done under Total Internal Reflection conditions. Imager sequences used for DNA-PAINT were 5’-AGGAGGA-Cy3B-3’ and 5’-AGAGAGA-Cy3B-3’. For 2-color imaging of Nsp13-EGFP and RIPK3-HA or ZBP1-HA, the imager solution from the first round of imaging was washed 5 times with buffer C (1× PBS and 500 mM NaCl) with 2 min of incubation. The second imager solution was prepared in buffer C, added to the wells, and imaged at the same plane. For all DNA-PAINT imaging rounds, the laser power was set at 313W/cm² on the imaging plane, and the image acquisition rate was set to 10 Hz in HiLo mode. The obtained raw fluorescence data was reconstructed using ‘Localize’ tool embedded in the Picasso Software suite to obtain a super-resolved image (Schnitzbauer et al., 2017). Drift correction was performed by redundant cross-correlation (RCC) followed by correction with fiducial markers. RCC was also used to align the super-resolved structures from the two imaging channels. Reconstructed data was rendered in Picasso:Render. For 2-color data, reconstructed files for both rounds of imaging were individually drift-corrected and loaded into ‘Render’ together using RCC or gold nanoparticles to align the two images.

Protein Purification

The bacterial codon-optimized coding sequence of ZBP1- Zα1Zα2 domains was synthesized (Thermo Fisher Scientific) and subcloned into N terminally His-Tagged pNIC-ZB vector. The plasmid expressing pNIC-ZB-ZBP1-Zα1Zα2 was transformed into Rosetta DE3 (kind gift from Prof. B Gopal from MBU, IISc) and grown overnight on LB agar plates with kanamycin and chloramphenicol. Single colonies from the plate were inoculated 20ml LB media and incubated
overnight at 37°C at 180 rpm. 20ml overnight cultures were inoculated into 1000 ml of LB containing kanamycin and chloramphenicol. Cultures were grown at 37°C, induced at 0.6 OD (600nm) by adding 0.3 mM IPTG, then incubated for 4 h at 37°C. The pellet lyed in the lysis buffer 50 mM HEPES (pH 7.5), 500 mM NaCl, 5 % glycerol, 1 mM PMSF, 1 mM DTT and 5 mM imidazole). Cells were disrupted by subsequent sonication until a clear solution was obtained. The cells were further centrifuged at 18300g for 40 min at 4°C. Ni-NTA beads (Gbiosciences, 786-940) were loaded on a gravity flow column and washed with 50 ml of lysis buffer containing 10 mM imidazole and a further wash with 25 ml of buffer containing 1M NaCl. Proteins were then eluted with gradient elution (10 mM Tris HCl, 500 mM NaCl, 5% glycerol, 1 mM PMSF, and no imidazole to 500 mM imidazole). The eluted protein was run on a 15% SDS-PAGE gel to check the purity. The eluted protein fraction was further treated with TEV protease to exclude the Z-basic part from the purified protein. Dialysis was performed for the eluted product with 20 mM Tris 7.5, 50 mM NaCl, 2 mM DTT overnight at 4°C. The eluted fraction was passed through the Heparin column. The column was washed with 50 ml of 50 mM HEPES (pH 7.5), 50 mM NaCl, 5% glycerol, 1 mM PMSF. The eluted fractions were pooled and diluted in no salt buffer (50 mM HEPES (pH 7.5), 5 % glycerol, 1 mM PMSF). These fractions were further loaded onto the Resource-S cation exchange column (Cytiva, 17117801), which was equilibrated with start buffer (20 mM HEPES pH 7.5 and 50 mM NaCl). The bound protein was eluted using gradient elution buffer (50 mM to 1 M NaCl). The eluted fractions were run on SDS-PAGE to check the purity of the protein (MW ~22.8 kDa), and the fractions that have pure proteins were pooled, snap frozen, and stored at -80°C.

**SARS-CoV-2 Z-RNA predictions.**

The SARS-CoV-2 Wuhan-Hu-1 RNA genome (NCBI, NC_045512.2) was used for predicting the possible Z-RNA forming sequences. Alternative purine-pyrimidine (Pu:Py) repeats and short interspersed nuclear elements (SINEs) with inverted repeats favor RNAs to attain Z-conformation
(Balasubramaniyam et al., 2018; Hall et al., 1984; Herbert et al., 2021; Jiao et al., 2020; Thamann et al., 1981; Zhang et al., 2020; Zhang et al., 2022). SARS-CoV-2 genome was assessed for the presence of Pu:Py repeats and SINE-like sequences using Microsatellite repeat finder, the non-B DNA Motif Search Tool (nBMST) and RNAfold developed by Mathew’s lab for predicting the secondary structures of RNAs (Cer et al., 2013; Reuter and Mathews, 2010). In the Microsatellite repeat finder, the parameters were extended to inverted repeats, tandem repeats, and Z-DNA-like sequences for identifying Z-RNA-like features. The repeat sequence length was set to a maximum 6 bases, as previous studies established that the Z-nucleic acid sequences are repeats of Pu:Py with a maximum length of 6 (Hall et al., 1984; Placido et al., 2007; Popenda et al., 2004; Thamann et al., 1981). Further, the alternative Pu:Py repeats spanning inverted repeats in the SARS-CoV-2 genome were selected, and the secondary structure was predicted using RNAfold. The nBMST was executed to delineate the sequences forming alternative DNA conformations that differ from the canonical right-handed Watson-Crick double-helix. By utilizing the tool, we predicted sequences with inverted short tandem repeats that form slipped/hairpin-like structures and Z-DNA-like motif sequences with repeats of purines and pyrimidines. The secondary structure of all the possible inverted repeats in the SARS-CoV-2 genome identified as mentioned above was predicted using RNAfold server to illustrate the canonical base pairs that RNA can attain considering free energy minima. The secondary structure of the SARS-CoV-2 genome in infected cells has been solved recently (Huston et al., 2021; Lan et al., 2022; Sun et al., 2021). Using this experimentally solved complete SARS-CoV-2 genome as a reference, we compared the secondary structures of predicted Z-RNA favoring inverted repeats with the experimentally solved secondary structure and precise base pairing events of the SARS-CoV-2 genome. Following the annotations, the SARS-CoV-2 RNA genome segments with Z-RNA forming potential were selected that retain secondary structures and base pairing like experimentally solved structures.

EMSA for probing ZBP1- Zα1Zα2 association with SARS-CoV-2 z-RNA fragments.
To perform EMSAs, SARS-CoV-2 Z-RNA (SC2-zRNA) fragments were generated using in vitro transcription method. The HiScribe T7 High Yield RNA Synthesis Kit (NEB, E2040S) was used to transcribe the template DNA per the manufacturer's instructions. The SARS-CoV-2 RNA fragments with Z-RNA forming potential were synthesized as DNA oligos with an upstream T7 promoter sequence (GGATCCTAATACGACTCACTATA). To synthesize uncapped RNA, 1 μg of DNA containing the T7 promoter upstream was mixed with the provided reaction buffer and the DNA oligos complementary to the T7 promoter. The mixture was subjected to a thermal cycling step at 95°C for 5 min. Following the thermal cycling, NTPs (provided in the kit) for RNA synthesis were added to the reaction mixture. The specific NTPs and their concentrations are typically included in the HiScribe T7 High Yield RNA Synthesis Kit. The reaction mixture containing the DNA template, reaction buffer, NTPs, and T7 RNA polymerase was then incubated at 37°C for 14-16 h. This extended incubation period allows the T7 RNA polymerase from the kit to catalyze the transcription process, resulting in the synthesis of uncapped RNA. The purified ZBP1-Zα1Zα2 domain was incubated at different concentrations with different SC2-zRNAs for 30 min at 23°C in the RNA binding buffer (10 mM HEPES pH 7.5, 50 mM NaCl). The RNA-protein mixture was then loaded onto the 6% Native PAGE gel and was run in Tris-boric acid-EDTA (TBE) buffer at 120 V for 40 min at 4°C. Once electrophoresis was completed, the gels were post-stained with ethidium bromide and imaged in the Bio-Print VILBER imaging system.

**SC2-zRNA transfections**

24 h after seeding, L929 cells were transfected with the construct expressing Nsp13-WT. 20 h after transfection, the cells were transfected with specified RNAs. 5 μg of RNA (poly I:C, Positive control Z-RNA, indicated SARS-CoV-2 Z-RNAs) was incubated with Lipofectamine 2000 transfection reagent (Invitrogen, 11668019), for 15 mins at room temperature. After incubation, the transfection mix was added to the cells in reduced serum media Opti-MEM, and cells were subjected to real-time cell death analysis using Sytox Green staining in Incucyte.
Biolayer interferometry (BLI)-Octet

The binding kinetics of ZBP1- Zα1Zα2 with SARS-CoV-2 RNAs were evaluated using the Octet RED96 instrument (Sartorius). First, amine-reactive second-generation (AR2G) biosensors were equilibrated in a buffer PBS pH 7.4. Then, the AR2G biosensor channel was activated using EDC and sulfo-NHS reagents (Sigma). Following this, ZBP1- Zα1Zα2 was diluted to a concentration of 10 µg ml⁻¹ and immobilized on activated AR2G biosensors for 300 sec in a 10 mM sodium acetate buffer, pH 4.0. The reaction was quenched by utilizing an excess of sulfo-NHS esters in 1 M ethanolamine. For monitoring SC2-zRNAs interaction with ZBP1- Zα1Zα2, black 96-well plates (Nunc F96 Microwell, Thermo Fisher Scientific) were loaded with 200 µl of the protein or the buffer, maintained at 25°C, and agitated at 1000 rpm. The baseline for each read was recorded using the buffer alone. The association of ZBP1- Zα1Zα2 with SC2-zRNAs at 100 nM and 200 nM concentration was captured for 300 sec, followed by the dissociation for 200 sec by dipping the sensor into the buffer without protein. After each kinetic assay, the biosensor chip was regenerated using 0.1 M Glycine-HCl. The acquired kinetic data was analyzed using the manufacturer’s software (Data Analysis HT v11.1). A global fitting approach was employed to analyze the data, to fit specific sensograms with a 1:1 binding model.

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DISCLOSURES

The authors declare no conflicts of interests.

AUTHOR CONTRIBUTIONS

S.K. conceptualized the study; S.M., D.J., A.A.D., S.N., and S.K. designed experiments, performed cell culture and biochemical experiments; S.K., K.B., and A.A.D. performed sequence and phylogenetic analysis; D.J. and S.K. performed Z-RNA mining and sequence analysis; D.J. and M.S. performed in vitro transcriptions, purification of Za protein, EMSA, and Octet binding studies; S.M, A.A.D., and M.A. performed imaging and DNA-PAINT studies; S.M., O.K., and A.A.D. performed SARS-CoV-2 infections and viral titer assays; S.T. assisted with SARS-CoV-2 infection experiments; M.G. helped with protein purification and DNA-PAINT experiments; S.M., D.J., A.A.D, and S.K. wrote the manuscript; M.G. and S.K. supervised the study; S.K. provided the guidance, and brought the funding; All the authors contributed in manuscript editing.

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FIGURE LEGENDS

Figure 1: Identification of CoV-RHIMs in Nsp13 and Nsp14 of SARS-CoV-2 and bats as hosts in viral RHIM evolution in RNA viruses. A, Schematic representation showing the genomic organization of SARS-CoV-2 and CoV-RHIMs in Nsp13 and Nsp14 proteins encoded by ORF1b. B, Amino acid residues and their relative conservation in RHIMs of human proteins (RIPK1, RIPK3, and ZBP1), M45 of murine cytomegalovirus (MCMV), ICP6 and ICP10 of herpes simplex virus (HSV), ORF20 of varicella-zoster virus (VZV) and Nsp13 and Nsp14 of CoVs (SARS-CoV-2, MERS-CoV and CoV-NL63). Human ZBP1 consists of two RHIM sequences (ZBP1-R1 and ZBP1-R2). C, Phylogenetic tree representing the relatedness of different CoV genera based on protein sequences encoded by ORF1ab. D-F, The phylogenetic tree and the relative conservation of CoV-RHIM-1 in Nsp13 (D), CoV-RHIM-2 (E), and CoV-RHIM-3 (F) in
Nsp14 across CoVs. G, Bat species from Yangochiroptera and Yinpterochiroptera suborders considered for analyzing bat RIPK3 and ZBP1 protein sequences. H, Protein sequence alignment and phylogenetic trees showing relative conservation of human and bat RHIM-sequences within RIPK3 and ZBP1 proteins. I, The potential viral RHIM signatures in lyssavirus rabies, picornavirus, and hantaviruses that originate from bats. The core tetrad residues and the proximal conserved residues of RHIMs are highlighted in bold; conserved or identical residue positions in RHIM sequences are highlighted in red; less conserved but chemically similar residue positions in RHIM sequences are highlighted in blue. Asterisk (*) symbols indicate Wuhan SARS-CoV-2 isolates.

Figure 2: Nsp13 of pathogenic beta-CoVs promotes cell death in human cells and is dependent on CoV-RHIM-1 and RNA-binding channel. A, Microscopic analysis of HT-29, HCT-116, and A549 cells infected with lentiviruses expressing SARS-CoV-2 Nsp13-WT, Nsp13-Tet-mut, Nsp13-Swap-mut and EGFP followed by puromycin treatment. Scale bar, 50μm. B, Immunoblot analysis of lysates from HT-29, HCT-116, and A549 cells showing expression of Nsp13-WT, Nsp13-Tet-mut, Nsp13-Swap-mut (Strep-Tag) and GAPDH after lentivirus transduction. C, Microscopic analysis of HT-29 cells transduced with lentiviruses expressing SARS-CoV-1 and MERS-CoV Nsp13-WT. Scale bar, 50μm. D, The annotation of domains in the Nsp13 structure (PDB ID: 6ZSL). E, Nsp13 protomers in apo or RNA bound conformation in SARS-CoV-2 mini replication-transcription complex structure (PDB ID: 7CXM). F-H, Alanine mutations in core tetrad residues of Nsp13 alter RNA binding channel conformation and abolish Nsp13-mediated cell death. F, The structure of RNA bound Nsp13 showing open RNA binding channel conformation (grey colored ellipse and yellow surface structure). G, The structure of Apo-Nsp13 showing closed RNA binding channel conformation. Apo and RNA-bound Nsp13 structures were aligned to show the displacement of 1B, Rec1a and Rec2a domains at the RNA binding channel. H, The predicted structure of Nsp13-Tet-mut shows altered RNA binding channel conformation. Alignment of Nsp13-Tet-mut and Nsp13-RNA bound structures showing the open RNA binding channel conformation of Nsp13-Tet-mut. I, Cell death levels in mock or SARS-CoV-2 infected A549-ACE2 cells ectopically expressing Nsp13 constructs as indicated. Cell death was monitored by Sytox green staining at 48 h of infection. ***P=0.0002 (comparison of SARS-CoV-2 infected mock and Nsp13-WT cells), ***P=0.0003 (comparison of SARS-CoV-2 infected Nsp13-WT and Δ1B expressing cells), **P=0.0076 (comparison of SARS-CoV-2 infected Nsp13-WT and Nsp13-Tet-mut expressing cells), **P=0.0043 (comparison of SARS-CoV-2 infected Nsp13-WT and Δ1B-Rec1A-Rec2A expressing cells), *P=0.0178, ns – not significant (one way ANOVA).
SARS-CoV-2 infectivity titers in cell supernatants measured by plaque assay. *P=0.0318 (one way ANOVA). Data shown are mean ± SEM.

**Figure 3: SARS-CoV-2 Nsp13 associate with host RHIM proteins, promoting large signaling complex formation.** A, Real-time cell death measurement by Sytox green staining of HT-29 cells transiently transfected with 500ng or 1μg of Nsp13-WT, Nsp13-Tet-mut and Nsp13-Swap-mut constructs. TNF and SMACmimetic (SMACmim) was used to induce apoptosis, and TNF, zVAD and SMACmim was used to induce necroptosis. B, Immunoblot analysis of anti-Strep-Tag (Nsp13), anti-RIPK1, anti-HA-tag (RIPK3) and anti-ZBP1 immunoprecipitates and whole cell lysates (input) from HEK-293T cell lysates expressing Nsp13 alone or co-expressing Nsp13, RIPK3 and ZBP1. C, Schematic of Nsp13 tagged with EGFP at its C-terminus. D, Microscopic analysis of A549 and HCT-116 cells infected with lentiviruses expressing SARS-CoV-2 Nsp13-WT, Nsp13-EGFP or EGFP followed by puromycin treatment. Scale bar, 50μm. E, Confocal microscopy imaging of HEK-293T cells expressing Nsp13-EGFP or ZBP1-dTomoto alone or co-expressing both the constructs. Scale bar, 10μm. F, Confocal microscopy imaging of HEK-293T cells expressing Nsp13-EGFP and in combination with ZBP1-HA or RIPK3-HA. Scale bar, 10μm. G-H, Visualization of SARS-CoV-2 Nsp13 and ZBP1 or RIPK3 in HEK-293T cells using DNA-PAINT imaging. Scale bars, 5 μm (complete cell image); 200nm (magnified images). I, Immunoblot analysis of crosslinked lysates of HEK-293T cells expressing Nsp13-WT, RIPK3 and ZBP1 individually or co-expressing Nsp13-WT+ZBP1, Nsp13-WT+RIPK3, Nsp13-WT+ZBP1+RIPK3 or ZBP1+RIPK3, in non-reduced (without BME) and reduced (with BME) conditions. O- oligomer complexes; M-Monomer.

**Figure 4: SARS-CoV-2 Nsp13 promotes ZBP1-RIPK3 signaling-dependent cell death and is regulated by intracellular RNA ligands.** A, Real-time cell death measurement by Sytox green staining of WT and Zbp1−/− L929 cells ectopically expressing Nsp13-WT and Nsp13-Tet-mut after IAV, IAV+zVAD and mock infection. ***P=0.0005, *P=0.0307 (two-way ANOVA). B, Real-time cell death measurement by Sytox green staining of ZBP1 expressing HT-29 cells after Nsp13 transfection and IAV+zVAD infection. C & D, Cell death measurement by Sytox green staining of WT and Zbp1−/− L929 cells expressing Nsp13-WT after LMB+IFN-β (C) and Curaxin (D) treatment. Data shown are mean ± SEM. E, Microscopic images of cell death in L929 treated as indicated and acquired by Incucyte imaging analysis for the cells. Scale bar, 200μm. F, Real-time analysis of cell death of mock or Nsp13 transfected L929 cells treated /infected with IFN-β alone, IAV alone, IAV + IFN-β, LMB alone or LMB + IFN-β. ****P<0.0001, **P=0.0022, *P=0.0310 (two way ANOVA, n=3). Data shown are mean ± SEM.
Figure 5: SARS-CoV-2 genome constitutes Z-RNA forming hotspots that interact with Z-RNA sensing Zα domains. A, Schematic representation of SARS-CoV-2 genome organization and hot spots of the genome with Z-RNA conformation forming potential. B, Specific SARS-CoV-2 genome segments with high Z-RNA forming dsRNA conformations. The orange box outlines represent alternate purine-pyrimidine repeats with Z-RNA forming and Zα-domain binding potential. C, SDS-PAGE gel picture representing purified human ZBP1-Zα1α2 domains. D-E, Octet binding of SARS-CoV-2 Z-RNAs (SC2-zRNA), control Z-RNA (alternate purine-pyrimidine repeats favoring Z-RNA conformation) and poly(I:C) at 100nM (D) and 200nM (E) concentration to human ZBP1-Zα1α2 domains. F, Electrophoretic mobility shift assay of control Z-RNA and SARS-CoV-2 Z-RNAs with human ZBP1-Zα1α2 domains to monitor their interaction.
Figure 1

CoV-RHIMs in different CoVs:
- Nsp13: CoV-RHIM-1
- Nsp14: CoV-RHIM-2 and CoV-RHIM-3

Viral RHIMs:
- RIPK1
- ZBP1
- MCMV M45
- HS1V ICP6
- H52 V2V ORF20
- SARS-CoV-2 Nsp13
- MERS-CoV Nsp13
- CoV-NL63 Nsp14
- SARS-CoV-2 Nsp14
- CoV-NL63 Nsp14

Bulbiviruses:
- HDaV1
- HDaV2
- HDaV3
- HDaV4
- HDaV5
- HDaV6
- HDaV7
- HDaV8

Figure 1A:
- ORF1a
- ORF1b
- Nsp13
- Nsp14
- S
- E
- M
- N

Figure 1B:
- RIPK1
- ZBP1
- MCMV M45
- HS1V ICP6
- H52 V2V ORF20
- SARS-CoV-2 Nsp13
- MERS-CoV Nsp13
- CoV-NL63 Nsp14
- SARS-CoV-2 Nsp14
- CoV-NL63 Nsp14

Figure 1C:
- beta-CoV
- alpha-CoV
- gamma-CoV
- delta-CoV

Figure 1D:
- Nsp13: CoV-RHIM-1
- Nsp13: CoV-RHIM-2
- Nsp13: CoV-RHIM-3

Figure 1E:
- Nsp14: CoV-RHIM-2
- Nsp14: CoV-RHIM-3

Figure 1F:
- Nsp14: CoV-RHIM-3

Figure 1G:
- Pteropus alecto
- Hyposideros armiger
- Myotis davidii
- Eptesicus fuscus
- Miniopterus natalensis

Figure 1H:
- ZBP1
- RIPK3
- Laiivirus glycoprotein
- Hantavirus glycoprotein
- Lyssavirus rabies glycoprotein
- Bat picornavirus VP1

Figure 1I:
- Picornavirus
- CoVs
- Lyssavirus
- Viral RHIMs

Figure 1J:
- Bat species
- Sub order
- Pteropus alecto
- Hyposideros armiger
- Myotis davidii
- Eptesicus fuscus
- Miniopterus natalensis

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Figure 4

(A) Sytox Green+ cells for IAV, IAV + zVAD, and Mock transfected with WT or Zbp1<sup>−/−</sup> cells.

(B) Time (h) showing Sytox Green+ cells for Mock, Nsp13-WT, and Nsp13-Tet-mut transfected with WT or Zbp1<sup>−/−</sup> cells.

(C) LMB + IFN-β Sytox Green+ cells over time.

(D) Curaxin showing Sytox Green+ cells over time.

(E) Images of Mock, IAV, IAV + IFN-β, Nsp13 + IAV, Nsp13 + IFN-β, Nsp13 + LMB, and Nsp13 + LMB + IFN-β.

(F) Sytox Green+ cells showing the effect of IAV, Nsp13, IAV + IFN-β, Nsp13 + IAV + IFN-β, LMB, and Nsp13 + LMB + IFN-β over time.

* P < 0.05
** P < 0.01
*** P < 0.001
**** P < 0.0001
ns Not significant
Figure 5

A

5' 5000 10000 15000 20000 25000 3'

ORF1a ORF1b S E M N

Table 1

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<tr>
<th>SC2-zRNA-1</th>
<th>SC2-zRNA-2</th>
<th>SC2-zRNA-3</th>
<th>SC2-zRNA-6</th>
<th>SC2-zRNA-9</th>
<th>SC2-zRNA-10</th>
<th>Control zRNA</th>
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<tr>
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Graph 1

Graph 2

Graph 3

Graph 4

Graph 5

Graph 6