Multiple Roles of SARS-CoV-2 N Protein Facilitated by Proteoform-Specific Interactions with RNA, Host Proteins, and Convalescent Antibodies

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ABSTRACT: The SARS-CoV-2 nucleocapsid (N) protein is a highly immunogenic viral protein that plays essential roles in replication and virion assembly. Here, using native mass spectrometry, we show that dimers are the functional unit of ribonucleoprotein assembly and that N protein binds RNA with a preference for GGG motifs, a common motif in coronavirus packaging signals. Unexpectedly, proteolytic processing of N protein resulted in the formation of additional proteoforms. The N-terminal proteoforms bind RNA, with the same preference for GGG motifs, and bind to cyclophilin A, an interaction which can be abolished by approved immunosuppressant cyclosporin A. Furthermore, N proteoforms showed significantly different interactions with IgM, IgG, and IgA antibodies from convalescent plasma. Notably, the C-terminal proteoform exhibited a heightened interaction with convalescent antibodies, suggesting the antigenic epitope is localized to the C-terminus. Overall, the different interactions of N proteoforms highlight potential avenues for therapeutic intervention and identify a stable and immunogenic proteoform as a possible candidate for immune-directed therapies.

KEYWORDS: mass spectrometry, protein, SARS-CoV-2, COVID-19, coronavirus, virus, nucleocapsid

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) is the etiological agent of coronavirus disease 2019 (COVID-19), which reached pandemic status in fewer than 3 months following its discovery. As of March 2021, there were >100 million infected and more than 2.5 million deaths. SARS-CoV-2 packages a large RNA genome of ~30 kb, which encodes for 25 nonstructural and 4 structural proteins (spike, nucleocapsid, membrane, and envelope proteins). The structural and genetic makeup of SARS-CoV-2 is highly homologous to SARS-CoV, the virus responsible for the 2002 SARS pandemic. Despite striking similarities between SARS-CoV and SARS-CoV-2, the unique biological and molecular features that have contributed to increased infectivity of SARS-CoV-2 are not well-understood.

Much attention has focused on the spike glycoprotein trimer, as it is the largest structural protein that plays a critical role in virus attachment to host cells. The nucleocapsid (N) protein, on the other hand, is one of the most abundant viral proteins; hundreds of copies of N make up the viral core that encapsulates the genomic RNA. N protein consists of two structured domains separated by a long flexible linker, giving rise to a high degree of conformational freedom. The separated domains allow the N protein to carry out many functions in the viral lifecycle such as RNA replication/transcription, virion assembly, and immune system interference. Interestingly, the nucleocapsid and spike proteins are the main immunogens in COVID-19 patients. Quantitative measurements of plasma or serum from SARS-CoV-2 patients found that the adaptive immune response to the N protein is more sensitive than to the spike protein, making it a better indicator of early disease. Antibodies against N protein are sensitive targets for COVID-19 diagnostics; however, the modes of action of anti-N antibodies in immunity and viral clearance are not well-understood.

Efforts to elucidate the structure of N protein aim to understand how the individual domains interact with RNA and other proteins. However, N protein is highly sensitive to proteolysis due to the intrinsically disordered linker region, and therefore, the structure of the full-length nucleocapsid protein remains elusive. Proteolysis is a key strategy in viral proliferation and often begets a change in protein function. Understanding the unique biological features of SARS-CoV-2,
and the ability to target specific viral processes, relies on an understanding of the sequence of interactions between viral and host proteins, inclusive of viral proteoforms.

Here, we present a comprehensive analysis of the SARS-CoV-2 N protein using native mass spectrometry (MS), top-down fragmentation, and bottom-up sequencing. We find that the full-length N protein undergoes proteolysis at highly specific sites to generate at least five unique proteoforms. We identify various stoichiometries of the proteoforms that are influenced by pH and evaluate the propensity for N proteoforms to bind different RNA sequences. We find that only the dimeric form of full-length N binds to RNA, suggesting it is the functional unit of ribonucleoprotein assembly. In addition, we show that an N-terminal proteoform directly interacts with cyclophilin A, a highly abundant cytosolic host protein implicated in viral replication. Moreover, cyclosporin A, an immunosuppressive drug, abolishes the interaction between N and cyclophilin A. Finally, using convalescent plasma from patients >6 months from initial COVID-19 diagnosis, we found that N proteoforms produced significantly different responses to IgM, IgG, and IgA antibodies. Notably, the C-terminal proteoform exhibited a heightened interaction with convalescent antibodies, suggesting the antigenic epitope is localized to the C-terminus. Our results contribute proteoform-specific information that may guide some of the many therapies against COVID-19 that are under investigation.

RESULTS AND DISCUSSION

N Protein Undergoes Proteolysis in the Vicinity of the Linker Region

Nucleocapsid proteins of coronaviruses share similar topological organization and show high sequence homology among related coronaviruses. The N protein is characterized by two major structural domains, the RNA binding and oligomerization domains. Similar to other coronavirus nucleocapsid proteins, the two domains are separated by a long and flexible linker region thought to be devoid of secondary structure. An N-terminal arm and a C-terminal tail flank the RNA binding and oligomerization domains, respectively. We constructed plasmids consisting of the full-length nucleocapsid protein with a cleavable N-terminal...
purification tag and expressed the constructs in both *Escherichia coli* and human embryonic kidney (HEK) 293T cells.

For the *Escherichia coli* construct, we purified the protein and verified the cleavage of the affinity tag via SDS-PAGE (Figure S1). Before removal of the purification tag, three distinct protein bands were detected at ~49, ~38, and ~28 kDa. Following removal of the tag, all three protein bands migrated by ~3 kDa less, or the mass of the tag (Figure S1B). We confirmed that each band corresponded to N protein by in-gel trypsin digestion followed by LC–MS-based bottom-up proteomics. All three bands contained peptides from the N protein, resulting in 78.1, 49.9, and 43.2% sequence coverage for bands 1, 2, and 3, respectively. To determine the representation of protein domains in each gel band, we plotted the distribution of the peptides detected across the five protein domains (Figure S1C). As anticipated, we observe an unbiased distribution of peptides across all five domains for band 1, consistent with the expectation that tryptic peptides would be reasonably distributed across the full-length protein. Over 50% of the total peptides detected in bands 2 and 3 were localized to the RNA binding domain, suggesting that the proteins are predominantly N-terminal derivatives. However, in all three bands, peptides located in the S8-residue C-terminal domain were detected, indicating the purified protein is made up of a diverse mixture of N proteoforms and not biased to N-terminal species due to the location of the purification tag.

We recorded a native mass spectrum to identify the stoichiometry of N protein from *E. coli* (Figure 1A) and observed two main charge state envelopes centered at 13+ and 20+. Deconvolution of the *m/z* signals provided experimental masses of 45 769 ± 1 and 91 537 ± 2 Da, respectively, which are in excellent agreement with the theoretical monomeric (45 769.83 Da) and dimeric (91 539.66 Da) masses of full-length N protein. A second distribution of monomers and dimers was observed and found to have deconvoluted masses of 28 735 ± 2 and 57 534 ± 1 Da, respectively.

Additional N proteoforms were observed immediately upon protein purification and continued to develop further over time (Figure S2). The mass spectrum evolved to reveal a series of peaks corresponding to five unique protein distributions (Figure 1B). The high baseline in this mass spectrum suggests the presence of additional proteolytic products; however, their low signal-to-noise precludes their characterization. Furthermore, it is likely that C-terminal proteoforms, containing the oligomerization domain, participate in the formation of higher-order oligomers that become insoluble and are therefore not detected at later time points. We instead sought to characterize the most stable and prominent proteoforms.

To determine the identity of each proteoform, we adapted a two-tiered tandem mass spectrometry approach to determine the intact mass and amino acid sequence for each series of peaks in the mixture. An individual peak in the mass spectrum was first isolated and subjected to charge reduction via electron-transfer dissociation (ETD) under conditions that do not result in the formation of fragments but instead produce a series of charge-reduced peaks (Figure 1C). The charge-reduced spectrum was necessary to confirm the assignment of the charge state series for each proteoform. The assigned charge states were then used to obtain deconvoluted masses of each proteoform present in solution (Table 1).

We tentatively assigned the proteoforms based on their intact masses and then generated sequence ions to confirm our assignment. Individual proteoforms were subjected to fragmentation by higher-energy collision induced dissociation (HCD), which accelerates the isolated ions into an inert gas to induce fragmentation along the amide backbone. The fragmentation products were then used to determine the molecular composition and to localize the exact site of cleavage. The HCD spectrum results in a series of singly, doubly, and triply charged sequence ions exemplified by the fragmentation of the 9+ charge state at *m/z* 2616.79 (Figure 1D). Fragmentation of intact proteins under native conditions is expected to yield 3–10% sequence coverage at the termini, and the propensity for fragmentation differs depending on several criteria (e.g., mass, charge, composition, structure) for natively folded proteins. Here, we achieved ~4% sequence coverage of the proteoforms by native top-down MS (Tables S1–S5). Fragmentation at the termini was complementary to our goal—to confirm the chemical identity of the distinct proteoforms of N protein.

To confirm that N proteoforms exist in a human-derived cell line, we expressed N protein in HEK 293T cells. We assessed N protein expression by Western blot using a monoclonal antibody raised against the highly homologous SARS-CoV N protein with cross-reactivity to SARS-CoV-2 N protein (Figure S3). Analysis of the whole cell lysate of cells stably expressing N protein produced mainly full-length N protein with evidence of lower-molecular-weight proteoforms (Figure S3A). Notably, when N is coexpressed with the membrane and envelope viral structural proteins, N proteoforms are generated in high abundance (Figure S3B). The enhanced cleavage of the full-length N protein in the presence of known interactors suggests that N proteoforms are primed for viral protein–protein interactions.

Having demonstrated that a number of proteoforms of N are generated in *E. coli* and HEK 293 cells, we continued our studies using protein derived from *E. coli* to characterize the proteoforms generated as a consequence of N protein primary structure. We reasoned that N from human cells is subject to additional post-translational modifications depending on the presence of co-interactors and the biological state of the cell during an infection. Considering the proteoforms identified in *E. coli*, we note that three result from cleavage after alanine (residue pairs All, All, and AllA), one results from cleavage following an arginine (RIM), and one results from cleavage in the C-terminal tail following a valine (VIT), (Figure 1E). Proteoforms N1−209 and N1−220 contain primarily the RNA binding domain, as they cleave within the flexible linker region. The major component of N1−273 is also the RNA binding domain.

### Table 1. Deconvoluted and Sequence Masses of N$_{FL}$ and N Proteoforms

| protein | deconvoluted mass$^d$ ± s.d. (Da) | sequence mass (Da) |
|---------|----------------------------------|--------------------|
| N$_{FL}$ | 45 769 ± 2 | 45 769.83 |
| N$_{FL}$ dimer | 91 537 ± 2 | 91 539.66 |
| N$_{156−419}$ | 28 735 ± 2 | 28 696.12 |
| N$_{156−419}$ dimer | 57 534 ± 1 | 57 392.24 |
| N$_{1−209}$ | 22 611 ± 1 | 22 612.71 |
| N$_{1−220}$ | 23 540 ± 0.3 | 23 541.73 |
| N$_{1−273}$ | 29 402 ± 0.6 | 29 382.43 |
| N$_{1−392}$ | 42 922 ± 1 | 42 918.74 |

$^d$ Determined using at least three adjacent charge states.
domain but, in this case, is followed by the linker region and a small portion of the oligomerization domain. Conversely, N156-419 comprises mainly the oligomerization domain while still retaining a small portion of the RNA binding domain.

Although we find no sequence similarity in the residues that flank each cleavage site, a commonality is that cleavage occurs immediately adjacent to a hydrophobic residue. The specificity of cleavage at the residues identified here, despite no common motif, suggests that there may be a structural or conformational component directing proteolysis; however, the exact mechanism is unclear. While the exact chemical identity of N proteoforms implicated in SARS-CoV-2 infection may vary, we suggest that proteolytic processing of N protein, which liberates the two major domains, produces functionally relevant proteoforms that are important antiviral targets.

**Oligomeric States of N Proteoforms Are Influenced by pH**

To better understand the role of the individual proteoforms, we expressed and purified four of the proteoforms identified by top-down MS. As almost all biological processes are influenced by pH, we recorded native mass spectra of N1−220, N1−273, and N156-419 from solutions at different pH (Figures S4–S8) to mimic the pH of different intracellular environments that N might encounter during viral infection. Mass spectra for N1−220 and N1−273 were recorded at pH 5.0, 7.4, and 8.0 reveal highly abundant charge state series centered at 9+ and a low-abundance distribution of signals centered at the 13+ charge state corresponding to monomers and dimers, respectively. The mass spectra for N1−273 show that it is predominantly monomeric with no significant change in charge state distribution or oligomeric state across the range of pH values tested. However, we observe two low-abundance charge state series corresponding to N1−209 and N1−220 suggesting that N1−273 continues to undergo cleavage at the previously mapped residues (Figure S6).

In contrast to the N-terminal proteoforms, mass spectra for N156-419 reveal multiple charge state distributions at pH 5.0, 7.4, and 8.0 (Figure S7). At pH 8, the mass spectrum reveals three charge state distributions centered at 10+, 18+, and 27+ with average masses corresponding to monomers, trimers, and a low population of hexamers (Table S6). At pH 7.4, monomers, dimers, and trimers persist. At the lowest pH (pH 5.0) N156-419 is exclusively trimeric. Finally, the mass spectra for full-length N at pH 5.0 and 8.0 reveal broadened and featureless peaks suggesting that NFL is likely aggregated (Figure S8). A low-abundance series of highly charged peaks centered at 18+ at pH 8.0 indicates some protein unfolding. Overall, we conclude that N1−209, N1−219, and N1−273 do not undergo significant pH-dependent changes in oligomeric state, while the C-terminal proteoform is highly sensitive with trimers predominating under both high- and low-pH conditions.

**RNA Sequence Influences Binding Stoichiometry**

RNA binding and ribonucleoprotein complex formation is the primary function of coronavirus N proteins. The N protein binds nucleic acid nonspecifically; however, the production of infectious virions relies on N protein forming specific interactions with viral RNA among an abundance of different cellular RNA species. With knowledge of the stoichiometries of N proteoforms, we sought to determine the propensity for N to bind specific RNA sequences. We created single-stranded RNA oligonucleotides consisting of 20 nucleotides of repeating sequences (4 × −GAUGG, 4 × −GAGAA). Considering the promiscuity of N protein, we chose sequences shown to interact with the human immunodeficiency virus (HIV) polyprotein (which includes a nucleocapsid domain) at the different stages of virus assembly and hypothesized that N proteins of different oligomeric states would exhibit similar bias toward artificial RNA motifs.

We incubated N proteoforms and RNA oligonucleotides at a molar ratio of 4:1 protein/RNA and recorded native mass spectra for all N protein–RNA complexes. The mass spectrum for N1−209 bound to 4 × −GAUGG (Figure 2A) reveals three charge state distributions with masses that correspond to the apo N1−209 monomer and N1−209 bound to one and two 4 × −GAUGG RNA oligonucleotides bound to NFL dimer. The scheme in the inset of (C) depicts NFL dimer bound to RNA as the functional unit of the ribonucleoprotein assembly.

![Figure 2](https://doi.org/10.1021/jacsau.1c00139)

Figure 2. RNA sequence influences the binding stoichiometry to N protein. (A) Mass spectrum of N1−220 after incubation with 4 × −GAUGG RNA oligonucleotides in a molar ratio of 1:4 RNA:protein. Two additional charge state distributions are observed that correspond to one and two RNA oligonucleotides bound to N1−220. The mass spectrum at m/z > 2700 was magnified 3× and offset for clarity (red trace). (B) Mass spectrum of N1−220 after incubation with 4 × −GAGAA RNA oligonucleotides in a molar ratio of 1:4. One additional charge state distribution is observed that corresponds to one RNA oligonucleotide bound to N1−220. The mass spectrum at m/z > 2700 was magnified 3× and offset for clarity (red trace). (C) Mass spectrum of NFL after incubation with 4 × −GAUGG RNA oligonucleotides in a molar ratio of 1:4. Monomers and dimers of NFL (red circles) and NFL−GAUGG (blue circles) are observed. An additional peak series between 4300 and 5600 m/z corresponds to two 4 × −GAUGG RNA oligonucleotides bound to NFL dimer. The scheme in the inset of (C) depicts NFL dimer bound to RNA as the functional unit of the ribonucleoprotein assembly.
−GAUGG RNA oligonucleotides. The charge state distribution corresponding to N$_{1−209}$ bound to two 4×−GAUGG RNA predominates over the single RNA-bound protein. Conducting the same experiment with a different oligonucleotide (4×−GAGAA) reveals only one additional charge state distribution for N$_{1−209}$ bound to one oligonucleotide (Figure 2B). Similar RNA binding stoichiometries are observed for N$_{1−220}$ and N$_{1−273}$; the mass spectra for N$_{1−220}$ and N$_{1−273}$ reveal distributions corresponding to the binding of one and two 4×−GAUGG oligonucleotides. Only one additional distribution is observed for N$_{1−220}$ or N$_{1−273}$ incubated with a 4×−GAGAA oligonucleotide, which corresponds to one oligonucleotide bound (Figures S9−10, Table S7), confirming the preference for the 4×−GAUGG sequence for all N-terminal constructs.

To examine if this preference was also observed for full-length N, we incubated the protein with a 4×−GAUGG oligonucleotide. A series of peaks was identified corresponding to the N protein dimer bound to two 4×−GAUGG oligonucleotides (Figure 2C). Similarly, in the presence of the 4×−GAGAA RNA oligonucleotide, an additional RNA-bound distribution is observed; however, the deconvoluted mass indicates that only one 4×−GAGAA oligonucleotide is bound to the N$_{FL}$ dimer (Figure S11, Table S7). No RNA binding to the monomeric form of N$_{FL}$ is observed, regardless of oligonucleotide sequence. Considering the different properties of the two RNA oligonucleotides, the 4×−GAUGG oligonucleotide contains three GGG motifs, which form short stem loops and are known to contribute additively to the efficiency of genome packaging in related viruses. Furthermore, selective RNA packaging has been described as a feature of innate immune response evasion. Our results emphasize that RNA sequence, and likely the secondary structure, is important for interactions with the N-protein. Notably, only the N$_{FL}$ dimer binds RNA, suggesting that the dimer is the functional unit of the SARS-CoV-2 ribonucleoprotein assembly. The preference for the RNA sequence known to form stem loops also has implications in efficient genome packaging and likely contributes to an optimized packing density in the intact virion.

N$_{1−209}$ Interacts Directly with Cyclophilin A

Cyclophilin A (CypA), a highly abundant immunophilin found in host cells, has been implicated in the replication cycle of coronaviruses and is known to aid in assembly of other viruses. We sought to determine if CypA could play a role in SARS-CoV-2 infection through monitoring direct interactions of CypA with N proteoforms. We incubated N$_{1−209}$ and CypA in a 1:1 molar ratio and used native mass spectrometry to measure possible interactions. The mass spectrum reveals charge state distributions that correspond to monomeric N$_{1−209}$ and CypA and three distinct charge state distributions at m/z > 3000 (Figure 3A). The three higher m/z distributions correspond to (i) heterodimers of N$_{1−209}$ and CypA, (ii) homodimers of CypA, and (iii) a low population of homodimers of N$_{1−209}$ (Table 2). Notably, we do not observe evidence of CypA interacting with N$_{1−220}$, N$_{1−273}$, nor N$_{FL}$ under the same conditions (Figure S12−15).

To determine if the interaction between N$_{1−209}$ and CypA could be inhibited by an approved immunosuppressant cyclosporin A (CsA), we incubated the N$_{1−209}$-CypA complex with a 2-fold molar excess of the drug (Figure 3B). The mass spectrum reveals three abundant charge state distributions: (i) a distribution centered at 9+ corresponding to N$_{1−209}$ monomers, (ii) a highly abundant distribution centered at a charge state of 8+ that corresponds to CypA bound to CsA, and (iii) a low-abundance distribution that corresponds to monomeric CypA (Table 2). Very low-abundance distributions of N$_{1−209}$ homodimers, CypA homodimers, and N$_{1−209}$-CypA heterodimers are barely detected following magnification >3000 m/z. Therefore, we can conclude that the 1:1 interaction between CypA and N$_{1−209}$ can be inhibited by CsA binding to CypA.

Antigenic Regions of N Are Located at the C-Terminus

Since N protein is detected by antibodies with higher sensitivity than any other structural proteins of SARS-CoV-2, we sought to determine if all N proteoforms were similarly recognized by immunoglobulins. We first incubated NFL with a monoclonal antibody (mAb) raised against the full-length protein in molar ratios of 1:1 (Figure 4A). The mass spectrum reveals three distributions >6000 m/z with deconvoluted masses that correspond to the apo antibody as well as one and two NFL proteins bound (Table 2, Figure S16). When the same mAb was incubated with three N-terminal proteoforms (N$_{1−209}$, N$_{1−220}$, and N$_{1−273}$) we did not observe any mAb binding in the mass spectrum (Figure 4B). To validate these
Table 2. Deconvoluted and Sequence Masses for N Proteoform Complexes

| protein/complex                  | deconvoluted mass ± s.d. (Da) | expected mass ± (Da) |   |
|----------------------------------|-------------------------------|----------------------|---|
| 4 × −GAUGG RNA                   | --                            | 6622.00              |   |
| 4 × −GAGAA RNA                   | --                            | 6650.20              |   |
| N1−209 + one 4 × −GAUGG RNA      | 29 233 ± 0.4                  | 29 234.71            |   |
| N1−209 + two 4 × −GAUGG RNA      | 35 917 ± 10                  | 35 856.71            |   |
| N1−209 + one 4 × −GAGAA RNA      | 29 262 ± 1                   | 29 262.91            |   |
| N1FL dimer + two 4 × −GAUGG RNA  | 105 131 ± 60                 | 104 783.66           |   |
| cyclophilin A (CypA)             | 20 084 ± 0.4                 | 20 175.82            |   |
| CypA dimers                      | 40 373 ± 62                  | 40 351.64            |   |
| N1−209 + CypA                    | 42 867 ± 15                  | 42 696.71            |   |
| N1−209 dimers                    | 45 356 ± 15                  | 45 225.42            |   |
| cyclosporin A                    | 120 285 ± 0.0                | 120 263.63           |   |
| CypA + CsA                       | 21 287 ± 0.5                 | 21 286.63            |   |
| mAb (monomer)                    | 137 990 ± 125                | --                   |   |
| N1FL + mAb                       | 183 931 ± 102                | 183 759              |   |
| 2N1FL + mAb                      | 229 984 ± 36                 | 229 528              |   |
|                                  |                               | 236 731              |   |

* Determined using at least three adjacent charge states.

results with conventional methods, we turned to protein detection by Coomassie stain and immunoblotting (Figure 4B, inset). The Coomassie stain detected N1FL and all proteoforms in high abundance. In contrast, the immunoblot revealed detection of only NFL and N156 in high abundance. In contrast, the immunoblot revealed detection of only NFL and N156 in high abundance. Finally, serum IgA antibodies were detected for all N proteoforms and showed a heightened response to N156−419. These results allow us to conclude that while all three antibodies are present, IgM antibodies are the most prevalent for all N proteoforms and IgG antibodies exhibit significant preference for the C-terminal proteoform.

**CONCLUSION**

We present a comprehensive characterization of SARS-CoV-2 N protein and highlight molecular features that may influence the unique biology of SARS-CoV-2 infection (Figure 5). Specifically, we find that N protein undergoes proteolysis in the vicinity of the linker region, separating the two major domains (the RNA binding and oligomerization domains). We identify various stoichiometries of N proteoforms that are influenced by pH, explicitly N156−419, which forms oligomers under both high- and low-pH solution conditions. We also show that N proteoforms bind RNA with a preference for GGG motifs and present evidence for NFL dimers being the functional unit of assembly in ribonucleoprotein complexes. Furthermore, we determined that immunophilin CypA binds directly to N1−209, an interaction that can be inhibited through addition of the immunosuppressant cyclosporin A. To test the immunogenicity of N proteoforms, we used a recombinant antibody and immunoblot techniques to demonstrate that the antigenic site of SARS-CoV-2 N protein resides toward the C-terminus. To test this in an *in vivo* scenario, we obtained convalescent plasma from eight patients >6 months after initial COVID-19 diagnosis. We discovered a heightened IgG and IgA response for the N1FL and N156−419 relative to N-terminal proteoforms N1−209, N1−220, and N1−273.

The flexibility of the intrinsically disordered regions in proteins serves to provide conformational freedom for making favorable protein−protein interactions while making proteins highly susceptible to proteolytic cleavage. The nucleocapsid protein of SARS-CoV, responsible for the SARS pandemic of 2002, was shown to undergo similar proteolytic cleavage; however, the precise role of proteolysis in SARS-CoV N protein remains elusive.32−35 New approaches in proteomics have revealed proteolytic cleavage of multiple viral proteins during SARS-CoV-2 infection, including extensive proteolysis of the N protein.31 Furthermore, our results share striking similarities with the cleavage sites observed for both SARS-CoV and N protein from SARS-CoV-2-infected cells.31

One of the most critical steps of viral proliferation is the packaging of genomic material and its assembly into new virions. Genome packaging of RNA viruses is highly selective and depends on specific nucleotide sequences and complex structural elements called packaging signals (Pcs).32 It has recently been demonstrated that HIV polyproteins have a stronger tendency to oligomerize in complex with Psi-RNA relative to non-Psi RNA.35 Our results indicate that N1FL and N proteoforms exhibit preference for RNA sequences that mimic key structural features of the genomic RNA.36 We anticipate that the more efficient binding of N protein dimers to RNA with GGG motifs underlies the selective packaging of genomic RNA in SARS-CoV-2. The disruption of structural features of viral RNA has been proven to inhibit replication in some viruses.35 In the case of SARS-CoV-2, disruption of preferred
RNA structures, and therefore disruption of N protein—RNA interactions, presents a promising strategy to intervene in replication and the potential to develop live attenuated vaccines.
Our results demonstrate highly specific interactions between cyclophilin A and N1−209. Cyclophilin A has been found in mature virions of HIV and is known to play a key role in the replication of HIV, hepatitis B, and other coronaviruses. Interestingly, the interaction between cyclophilin A and the HIV capsid protein is known to be conformation-dependent. We anticipate such parallels with SARS-CoV-2 and speculate that other N proteoforms do not interact with CypA because of a conformational change or restricted access to the interacting domains. Intervention of the specific interactions between N proteoforms and cyclophilin A could therefore prove beneficial as a component of treatment strategies.

Considering antibody responses, proteoforms N1−209, N1−229, and N1−279 show a diminished interaction with IgG antibodies in convalescent plasma when compared to the full-length protein. These proteoforms could thereby contribute to virtually unchecked virus proliferation, depending on their functional roles and the protective properties of anti-N IgG. By contrast, N156−419 resulted in a heightened IgM response and an IgG response that could not be differentiated from the full-length protein, suggesting that the antigenic site is localized to the C-terminus. This finding is in accord with epitope mapping of the homologous SARS-CoV N protein. The significant differences in IgM, IgG, and IgA detection highlight new avenues for therapies to direct immune responses and represent potential targets for next-generation vaccine development.

Beyond the identification of potential vaccine targets, several additional therapeutic avenues are highlighted by this study. First, inhibition of the proteolysis reaction would prevent the formation of viral proteoforms that may be fine-tuned for various functions within the viral lifecycle. While the mechanism behind the proteolysis of N-protein is not yet understood, systematic mutation of cleavage sites defined here could lead to mechanistic and structural insights to enable small molecule screening of potential protease inhibitors. Knowledge of N-protein-structured RNA interactions could also aid the design of new therapeutics that would inhibit successful replication. However, since N protein oligomerizes in the absence of RNA, de novo drug design of assembly inhibitors is complex, as it is necessary to consider the oligomerization propensity of the various proteoforms at the pH regimes encountered in the cellular environment. CsA and cyclosporine derivatives, such as Alisporivir, however are more straightforward to track and have become attractive candidates to treat COVID-19. Disruption of the cyclophilin A−N-proteoform interactions shown here provides a convenient means of screening potential inhibitors for hit-to-lead optimization. Given the likelihood that no one intervention is likely to ameliorate the complex symptoms of COVID-19 infection, our findings contribute proteoform-specific information that may guide some of the many therapies under investigation.

### MATERIALS AND METHODS

#### Ethics

Patients were recruited from the John Radcliffe Hospital in Oxford, United Kingdom, between March and May 2020 with written and informed consent. Participants were identified from hospitalization during the SARS-CoV-2 pandemic and recruited into the Sepsis Immunomics and International Severe Acute Respiratory and Emerging Infection Consortium World Health Organization Clinical Characterization Protocol UK (IRAS260007 and IRAS126600).

Patient samples were collected at least 28 days from the start of their symptoms. Ethical approval was given by the South Central–Oxford C Research Ethics Committee in England (reference: 13/SC/0149), Scotland A Research Ethics Committee (reference: 20/SS/0028), and World Health Organization Ethics Review Committee (RPC571 and RPC572l, 25 April 2013).

#### Plasmid Construction and Cell Growth

A codon-optimized synthetic gene corresponding to the full-length nucleocapsid protein (Thermo GeneArt, Regensburg, Germany) was cloned into a modified pET28a vector using the In-Fusion cloning kit (Takara Bio Saint-Germain-en-Laye, France). The resulting plasmid encoded for an N-terminal His6 tag followed by thrombin and tobacco etch virus cleavage sequences upstream of the full-length nucleocapsid protein sequence. To generate the nucleocapsid proteoforms, the desired sequences pertaining to the truncated forms of the N protein were subcloned from the synthetic gene using polymerase chain reaction (Phusion polymerase, New England Biolabs, Hertfordshire, UK). All genes were cloned into the modified pET28 vector, and gene sequences were confirmed by Sanger sequencing.

Plasmids were transformed into BL21 (DE3) and streaked onto LB agar plates supplemented with 50 mg mL−1 of kanamycin. Several colonies were used to inoculate 100 mL of LB broth supplemented with kanamycin and grown at 37 °C overnight. Aliquots of 10 mL of the overnight precultures were used to inoculate 1 L of LB broth supplemented with kanamycin. Cell cultures were grown to OD600 ≈ 0.6 before protein expression was induced with 0.5 μg mL−1 of IPTG. Cells were grown for an additional 4 h at 37 °C before being harvested via centrifugation (5000g, 10 min, 4 °C). Cell pellets were flash frozen in liquid nitrogen and stored at −80 °C until use.

#### Protein Purification

Cell pellets were resuspended in lysis buffer (25 mM Tris-HCl pH 8.0, 500 mM NaCl, 5 mM MgCl2, 5 mM β-mercaptoethanol, 5 mM imidazole, 10% v/v glycerol) containing EDTA-free protease inhibitor tablets (Roche). Cells were lysed by five passes through a microfluidizer (preheated to 4 °C) at 20,000 psi. Cell debris was pelleted by centrifugation (20,000g, 20 min, 4 °C). The supernatant containing soluble nucleocapsid protein was passed through a 0.45 μm filter.

Supernatant was loaded onto a Ni-NTA column pre-equilibrated in loading buffer (25 mM Tris-HCl pH 8.0, 500 mM NaCl, 5 mM MgCl2, 5 mM β-mercaptoethanol, 20 mM imidazole, 10% v/v glycerol) and allowed to pass via gravity flow. To remove common contaminating proteins, a heat-treated BL21 (DE3) E. coli lysate was loaded into a 10 mM MgATP was passed over the immobilized nucleocapsid protein using a protocol by Rial and Ceccherelli. The resin was washed with 10 column volumes of wash buffer (25 mM Tris-HCl pH 8.0, 500 mM NaCl, 5 mM MgCl2, 5 mM β-mercaptoethanol, 80 mM imidazole, 10% v/v glycerol) and then eluted twice with 10 mL of elution buffer (25 mM Tris-HCl pH 8.0, 500 mM NaCl, 5 mM MgCl2, 5 mM β-mercaptoethanol, 400 mM imidazole, 10% v/v glycerol).

The eluted protein was mixed with TEV protease in a 100:1 (w/w) ratio and loaded into a 3 kDa MWCO dialysis cassette (Thermo Fisher Scientific, United Kingdom). Cleavage of the His6-thrombin–TEV tag was carried out overnight at 4 °C in lysis buffer. The cleaved tag and TEV protease were separated from the untagged protein using reverse immobilized metal affinity chromatography on a Ni-NTA column prepared in loading buffer. The flow-through containing the untagged protein was collected and concentrated in a 10k MWCO centrifugal filter before MS analysis. Protein concentration was determined using UV−vis spectroscopy by monitoring the absorbance at 280 nm with a theoretical extinction coefficient (ε ≈ 43 890 M−1 cm−1) determined using the Exasy Protparam tool.

#### Native Mass Spectrometry

RNA oligonucleotides of repeating sequences (4 × −GAUGG, 4 × −GAGAA) were purchased from Integrated DNA Technologies. Recombinant human cyclophilin A (product ab86219) was purchased...
from Abcam (Cambridge, United Kingdom). Cyclosporine A purchased from Merck Life Science (Dorset, United Kingdom). N protein and all binding partners were buffer exchanged or diluted into 500 mM NH4OAc pH 5.0, 7.4, or 8.0. Buffer exchange was carried out using Zeba Spin Desalting Columns, 7K MWCO (Thermo Fisher Scientific, United Kingdom).

Measurements were carried out on a Q. Exactive UHMR or Orbitrap Eclipse. The Q. Exactive instrument was operated in the positive ion mode using the manufacturer’s recommended parameters for native MS. The instrument was operated at a resolving power of 12 500 (at m/z 200). An electrospray was generated by applying a slight (∼0.5 mbar) backing pressure to an in-house-prepared gold-coated electrospray capillary held at ∼1.2 kV relative to the instrument orifice (heated to ∼100 °C). The mass spectra were deconvoluted using in-house software (the Mass and Charge State Determination tool, available for download at http://www.benesch.chem.ox.ac.uk/resources.html).

An Eclipse Tribrid instrument was also used for native MS and top-down sequencing. The instrument was set to intact protein mode at a standard ion routing multipole pressure of 10 mTorr. Ion voltages were set to transmit and detect positive ions at a resolving power of 12 500 (at m/z 200). An electrospray voltage of ∼1.2 kV and backing pressure of ∼0.5 mbar were used for ion formation; desolvation was assisted using an instrument capillary temperature of ∼100 °C. To identify the accurate mass and sequence of each proteoform, we used a similar approach to that described by Huguet et al. Briefly, a desired signal was isolated using the ion trap (10 m/z isolation window, charge state set to 10) and subjected to (i) electron-transfer dissociation (ETD, 3 ms activation time, 1.0 × 105 ETD reagent target) to generate a charge-reduced series for accurate intact mass determination and (ii) higher-energy collisional dissociation (HCD) using ∼30–50 V HCD collision energy to generate fragment ions.

Monoisotopic masses of fragment ions generated by HCD having a normalized intensity of 10% or higher were fed into Prosite Lite software. A series of candidate sequences that best matched the measured intact masses for each proteoform were generated. The monoisotopic fragment masses were matched to expected ions generated in silico based on the provided candidate sequence. Internal monoisotopic fragment masses were matched to expected ions measured intact masses for each proteoform were generated. The

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacsau.1c00139.

Supporting Information includes detailed experimental methods; Figures S1–S16; Tables S1–S7 (PDF)

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Notes
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ABBREVIATIONS

SARS-CoV-2, severe acute respiratory syndrome coronavirus 2; COVID-19, coronavirus disease 2019; MS, mass spectrometry; N, nucleocapsid; HEK, human embryonic kidney; HCD, higher-energy collision induced dissociation; ETD, electron-transfer dissociation; CypA, cyclophilin A; CsA, cyclosporin A; mAb, monoclonal antibody; ELISA, enzyme-linked immunosorbent assay; Psi, packaging signals

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