Clinically observed deletions in SARS-CoV-2 Nsp1 affect its stability and ability to inhibit translation

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(Received 17 November 2021, revised 21 March 2022, accepted 23 March 2022, available online 25 April 2022)
doi:10.1002/1873-3468.14354
Edited by Urs Greber

Nonstructural protein 1 (Nsp1) of SARS-CoV-2 inhibits host cell translation through an interaction between its C-terminal domain and the 40S ribosome. The N-terminal domain (NTD) of Nsp1 is a target of recurring deletions, some of which are associated with altered COVID-19 disease progression. Here, we characterize the efficiency of translational inhibition by clinically observed Nsp1 deletion variants. We show that a frequent deletion of residues 79–89 severely reduces the ability of Nsp1 to inhibit translation while not abrogating Nsp1 binding to the 40S. Notably, while the SARS-CoV-2 5' untranslated region enhances translation of mRNA, it does not protect from Nsp1-mediated inhibition. Finally, thermal stability measurements and structure predictions reveal a correlation between stability of the NTD and the efficiency of translation inhibition.

Keywords: COVID-19; Nsp1; pathogenicity; ribosome; SARS-CoV-2; virus

Coronaviruses (CoVs) are a family of enveloped, positive-sense single-stranded RNA viruses subdivided into four genera: Alpha-, Beta-, Gamma-, and Delta-coronaviruses [1]. Four so-called common-cold coronaviruses that cause only mild disease are prevalently circulating in the human population: HCoV-OC43, HCoV-HKU1, HCoV-229E, and HCoV-NL63 [2]. The situation has changed dramatically in the recent years with the zoonotic introduction of three highly pathogenic Betacoronaviruses into humans: SARS-CoV in 2002 [2], MERS-CoV in 2012 [2], and, most recently, SARS-CoV-2 in 2019 [3–6]. SARS-CoV-2 is the cause of the COVID-19 pandemic which to date has caused nearly five million confirmed deaths (https://coronavirus.jhu.edu/map.html, retrieved 2021-10-21).

Coronaviruses have unusually large genomes for positive-sense single-stranded RNA viruses. At 27–32 kb, they are an approximately threefold larger than other representatives of this group. This genomic expansion is thought to be made possible by a more complex, proofreading-capable polymerase [7], and it has allowed coronaviruses to acquire a larger toolbox

Abbreviations
COVID-19, coronavirus disease 2019; CTD, C-terminal domain; Nsp1, nonstructural protein 1; NTD, N-terminal domain; pLDDT, per-residue prediction quality score; SARS-CoV-2, severe acute respiratory syndrome coronavirus 2; TSA, thermal shift assay; UTR, untranslated region.
of host cell-manipulating proteins. One such key pathogenicity factor is the nonstructural protein 1 (Nsp1). This protein has no known enzymatic activity, and while betacoronaviruses deleted for the nsp1 gene can replicate in cell culture, they are strongly attenuated in vivo [8]. The 180 amino acid-long SARS-CoV Nsp1 (with which SARS-CoV-2 Nsp1 shares 84% sequence identity and displays conservation of all known key motifs) was found to suppress the antiviral host cell interferon response through a dual mechanism: it mediates the cleavage of host cell mRNAs by an unknown ribonuclease [9], and it suppresses translation through direct interaction with the small (40S) subunit of the ribosome [10]. Distinct mutations in SARS-CoV nsp1 have been described that selectively abrogate either of these two effects: substitutions of Arg124-Lys125 in the folded N-terminal domain of SARS-CoV were shown to abolish the Nsp1-induced mRNA cleavage [11], and residues Lys164-His165 in the C-terminal part of Nsp1 have been shown to be essential for binding to the 40S subunit of the ribosome (Fig. 1A) [12,13]. SARS-CoV-2 Nsp1 binds empty small ribosomal subunit, as well as the 43S preinitiation complex and the full 80S ribosome, in all cases only efficiently engaging the open conformation of the small subunit [14]. In cryo-EM structures of SARS-CoV-2 Nsp1 bound to the human 40S, the C-terminal ~30 residues (148–180 in PDB ID 6ZLW) were present in a sufficiently rigid conformation that an atomic model could be built [15–17]. This model shows that the C-terminal part of Nsp1 folds as two α-helices that form a hairpin-like arrangement inside the mRNA tunnel, incompatible with concomitant binding of mRNA. Consistent with the functionally crucial location of the C-terminal domain in the Nsp1:40S complex, both its truncation (A118–180) and K164/H165A substitution abrogate the Nsp1 interaction with the small ribosomal subunit [18]. Off the ribosome the C-terminal region (131–180) is unstructured, suggesting...
a possibility that the structure is attained upon recruit-
ment to the ribosome [19]. The N-terminal domain
could be located in the cryo-EM maps but at too low a
resolution to allow model building, presumably due to
flexibility with respect to the 40S. The cryo-EM derived
location of the Nsp1 N-terminal domain in the riboso-
mal complex is further supported by in situ cross-
linking mass spectrometry identifying multiple cross-
links between Nsp1 and ribosomal protein S3 [18,20].

The structure of Nsp1 bound to the ribosome raises
the question how any translation is possible if Nsp1
blocks the mRNA channel. Somewhat paradoxically,
experiments with in vitro translation systems have
shown that SARS-CoV-2 Nsp1 can efficiently inhibit
translation of both host and SARS-CoV-2 mRNAs,
with some studies showing slightly smaller and others
showing equal relative inhibition of viral mRNA
[14,17,21]. However, with SARS-CoV-2 5′ untranslated
region (UTR) acting as a translational enhancer,
mRNAs with the viral 5′ UTR are inherently trans-
lated more efficiently than host mRNAs [17]. There-
fore, even if translation of viral mRNA is inhibited to
a similar relative degree by Nsp1, viral mRNA still
has higher relative translational efficiency during infec-
tion. Furthermore, Tidu et al. [21] suggested that viral
mRNA is less sensitive to inhibition by Nsp1, while
similar degree of inhibition of viral mRNA
was reported by Schubert et al. [17] and
Lapointe et al. [14]. An interaction between Nsp1 and
the stem-loop region 1 (SL1) of the viral 5′ UTR was
implicated in partial escape of SARS-CoV mRNAs
from Nsp1 suppression [22], but the lack of affinity of
SARS-CoV-2 Nsp1 for the viral 5′ UTR remains to be
reconciled with such a mechanism [21]. The N-
terminal region of Nsp1 was shown to play a role in
translational shutdown, and its deletion (Δ1–117)
attenuating both ribosome and mRNA binding of
SARS-CoV-2 Nsp1 [18].

In the course of the ongoing COVID-19 pandemic,
extensive genetic characterization has been carried out
of new emerging mutants and virus variants. Nsp1 dis-
plays one of the highest degrees of diversity among
SARS-CoV-2 proteins [23–25], with several circulating
viruses with deletions in Nsp1 reported [25,26]. A
three-residue deletion (A141–143) was described as
appearing in clinical isolates from different geographi-
cal localities [26]. These three residues are located close
to, albeit not in, the region which forms an ordered
structure inside the 40S mRNA tunnel. It thus seems
possible that this shortening of the linker region
between the ribosome-inserting part and the stably
folded N-terminal β-barrel domain [27,28] may affect
the association of this altered protein with the
ribosome, and thus its effectiveness in translational
shutoff. Another cluster of deletions is located around
residues 79–89 of Nsp1 [25]. This results in various
length deletions around a loop in the atypical β-barrel
of the folded N-terminal domain. The longest deletion,
Δ79–89, is reported to be the fifth most common dele-
tion in SARS-CoV-2 Nsp1 in a worldwide comparison
[25]. The 79–89 deletion, as well as shorter deletions in
the same region of Nsp1, correlates with higher cycle
threshold (Ct) values in patients (i.e. lower viral load),
less severe disease outcome, and a weaker interferon
response as measured by lower serum levels of IFN-β
[25]. The biochemical basis for the altered disease
response associated with Nsp1 deletions is not estab-
lished. Here, we purified a number of Nsp1 proteins
corresponding to these circulating mutations in SARS-
CoV-2 and compared their potency in inhibiting trans-
lations using a human cell in vitro translation lysate.
We correlate the findings to thermal stability assays
and protein fold predictions of the mutants. The result
shed light on the clinical and cellular findings related
to Nsp1-mutated SARS-CoV-2 isolates.

Materials and methods

Cloning and mutagenesis

For expression of wild-type Nsp1, the nsp1 gene was ampli-
fied from the construct pLVX-EF1alpha-SARS-CoV-2-
nsp1-2xStrep-IRES-Puro (Addgene) [29] and inserted into a
1B vector (Macrolab, UC Berkeley) using In-Fusion clon-
ing kit (Takara Biosciences, San Jose, CA, USA). Deletion
mutations were generated from this plasmid by standard
site-directed mutagenesis methods. The reporter plasmid
T7-5′SARS-CoV-2 UTR-Firefly luciferase was generated by
inserting the T7 promoter upstream of SARS-CoV2 5′
UTR fused to the firefly luciferase coding sequence (ampli-
fied from T7-cmvtrans-Fluc-poly A) into the destination
vector pUC19. The reporter plasmid with nonviral UTR
was also based on the plasmid T7-cmvtrans-Fluc-poly A
(Addgene). All plasmids were sequenced to confirm cloning
of the correct sequence.

Protein expression and purification

Wild-type Nsp1 and all deletion variants were expressed
and purified as follows. The plasmid was transformed into
Escherichia coli BL21 (DE3) cells for overexpression. An
overnight culture was grown at 37 °C until the OD600
reached 0.4, after which the incubator temperature was
changed to 25 °C to let the cells cool down to induction
temperature 25 °C. At OD600 of around 0.8–0.9 the protein
expression was induced by addition of 0.5 mM Isopropyl β-
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D-1-thiogalactopyranoside (IPTG) and the protein was expressed at 25 °C overnight. Cells were harvested by centrifugation at 7000 g (rotor JLA-8.1000; Beckman Coulter, Brea, CA, USA) for 60 min. After discarding the supernatant, the cell pellet was washed with lysis buffer (50 mM Hepes-NaOH, pH 7.4, 300 mM NaCl, 0.1 mM THP, 10 mM imidazole, and 5% glycerol) and stored at −80 °C.

Cell mass was thawed and resuspended in lysis buffer supplemented with DNase I and protease inhibitor cocktail (1 mM benzamidine, 0.2 mM phenylmethylsulfonyl fluoride, and 5 μM leupeptin). Homogenized suspension was then passed twice through a cell disruptor (Constant System Limited, Daventry, UK) at a pressure 27 kPsi. The lysate was clarified by centrifugation at 36 000 g (rotor JA-25.50; Beckman Coulter) for 1 h, and the supernatant passed through a 0.22-μm syringe filter. One milliliter Ni-Sepharose Fastflow resin (Cytiva, Umeå, Sweden) pre-equilibrated with lysis buffer was combined with clarified lysate and incubated for 2 h at 4 °C under gentle agitation. Next, the resin was loaded onto a gravity-flow column. The protein-bound resin was washed twice with 20 mL wash buffer (50 mM Hepes-NaOH, pH 7.4, 300 mM NaCl, 0.1 mM THP, 30 mM Imidazole, and 5% glycerol) twice. The resin was then resuspended in 4 mL lysis buffer, supplemented with TEV protease (approx. 70 μg·mL⁻¹) and incubated overnight at 4 °C on a rotator wheel. The cleaved protein was collected as flowthrough. An additional wash with 5 mL lysis buffer was performed to collect the residual cleaved protein. For production of His₆-tagged Nsp1, the protein was instead eluted from the resin using wash buffer supplemented with 250 mM imidazol. The proteins were then further purified by anion exchange chromatography. To do so, after diluting with buffer A (50 mM Hepes-NaOH, pH 7.4, 100 mM NaCl, 0.1 mM THP, and 5% glycerol), dilute sample was filtered using 0.22-μm syringe filter (VWR, Radnor, PA, USA) and loaded onto a HiTrap Q HP 1ml column (Cytiva) pre-equilibrated with lysis buffer supplemented with DNase I and protease inhibitor cocktail (20 mM Hepes-KOH, pH 7.4, 100 mM KOAc, 2 mM Mg(OAc)₂, 0.1 mM THP, and 2.5% glycerol). Protein elutions were subjected to SDS/PAGE gel electrophoresis and the fractions corresponding to the center of the Nsp1-containing peak were pooled and concentrated. Aliquots were then flash frozen in liquid N₂ and stored at −80 °C. The identity of all the proteins was confirmed by trypsin-digestion mass spectrometry. His₆-tagged protein was used for western blotting of density gradient fractions, whereas all other experiments were done with untagged protein.

In vitro transcription

Capped and polyadenylated RNA transcripts were synthesized from linearized plasmid (T7–5’-SCoV2-UTR–firefly Luciferase) using the mMESSAGE mMACHINE T7 Ultra kit (ThermoFisher, Watham, MA, USA) following the manufacturer protocol. Briefly, all the reagents including the 5′cap analog were gently mixed with linearized plasmid and transcription was performed by T7 RNA polymerase at 37 °C for 2 h. Then, 1 μL of Turbo DNase was gently mixed with the transcription mixture and incubated further at 37 °C for 15 min. In order to add poly (A) tail to the 3′ end of the in vitro transcribed RNA, E. coli Poly(A) Polymerase I (E-PAP) was added along with other provided reagents to the transcription mixture and after mixing gently incubated at 37 °C for 45 min. According to the manufacturer, this poly(A) tailing adds at least 150 adenosines to the 3′ end of the mRNA. Finally, the RNA prep was recovered by lithium chloride precipitation and quality-checked by running a sample on a denaturing agarose gel, which resulted in a single band. The capped and poly(A)-tailed RNA was aliquoted in 10 μL volumes and flash frozen in liquid N₂ and stored at −80 °C. For simplicity, the reporter mRNA with SARS-CoV-2 5′ UTR is referred to as ‘viral mRNA’ and the one with non-SARS-CoV2 5′UTR is referred to as ‘nonviral mRNA’ throughout the text.

Preparation of HEK293F translation lysate

In vitro translation lysates were prepared from HEK293F cells using a previously described protocol [30–32]. Cells were scraped and collected by centrifugation for 5 min at 600 r.p.m. at 4 °C. Cells were washed once with cold PBS (137 mM NaCl, 2.7 mM KCl, 100 mM Na₂HPO₄, 2 mM KH₂PO₄) and resuspended in Lysolecithin lysis buffer (20 mM Hepes-KOH, pH 7.4, 100 mM KOAc, 2.2 mM Mg(OAc)₂, 2 mM DTT, and 0.1 mg·mL⁻¹ lysolecithin), using 1 mL for 8 × 10⁶ cells. Cells were incubated for 1 min on ice, then immediately centrifuged for 10 s at 10 000 g at 4 °C. The pellet was resuspended in cold hypotonic extraction buffer (20 mM Hepes-KOH, pH 7.5, 10 mM KOAc, 1 mM Mg(OAc)₂, 4 mM DTT, and Complete EDTA-free protease inhibitor cocktail (Roche, Basel, Switzerland) at an equal volume to the size of the pelleted cells. After 5 min of incubation on ice, cells were transferred into a precooled Dounce homogenizer and lysed by 20–25 strokes. The lysate was centrifuged at 10 000 g for 10 min at 4 °C, and the supernatant transferred to a fresh tube. Aliquots were flash frozen in liquid N₂ and stored at −80 °C.

In vitro translation assays

In vitro translation reactions were performed as previously described [30–32], with modifications relating to the Nsp1 addition. HEK293F-cell lysate was pre-incubated with...
increasing concentrations (from 0 to 3 μM final concentration) of recombinant Nsp1 (wild-type and variants) for 15 min on ice. Translation buffer (20 mM HEPES-KOH, pH 7.6, 1 mM DTT, 0.5 mM spermidine-HCl, 1 mM Mg(OAc)₂, 8 mM creatine phosphate, 1 mM ATP, 0.2 mM GTP, 150 mM KOAc, 25 μM of each amino acid, and 2 units of human placental ribonuclease inhibitor (Fermentas, Burlington, Canada) was then added followed by 1 μL of the reporter mRNA (0.5 pmol) to give a total reaction volume of 13 μL. Final RNA concentration in the reaction mixture was kept at 38 nm. Translation reactions were incubated at 30 °C for 3 h, samples were flash frozen in liquid N₂ and kept at −80 °C. Luciferase assays were performed using the Steady-Glo Luciferase Assay kit (Promega, Madison, WI, USA) following the manufacturer’s protocol. Luminescence was measured using the M200 finite series microplate reader (TECAN, Manching, Switzerland). Samples for all concentrations of each protein were prepared in triplicates and measured.

Sucrose gradient fractionation and western blotting

20 A₂₆₀ units of HEK293F translation lysate were supplemented with either wild-type or Δ79–89 variant N-terminally His-tagged Nsp1 (final concentration of 1 μM) in hypotonic extraction buffer or just hypotonic hypotonic extraction buffer and kept on ice for 15 min. One hundred microlitres or the resultant mixture was loaded onto 10–50% sucrose gradient in HEPES:Polymix buffer pH 7.5 (5 mM Mg²⁺, 95 mM KOAc, 2 mM Mg(OAc)₂, 0.1 mM THP, and 2.5% glycerol. Each sample was prepared in triplicates. Samples were dispensed into FrameStar 96-well PCR plate (4titudE) sealed afterward with PCR optical Seal (4titudE). Thermal scanning (10–95 °C at 1.5 °C-min⁻¹) was performed using a real-time PCR instrument C1000 Touch Thermal Cycler (CFX96 from Bio-Rad) and fluorescence intensity was measured after every 10 s. According to the described protocol, raw data were truncated in Microsoft Excel to remove postpeak quenching [34]. A nonlinear fitting of the truncated dataset to a Boltzmann Sigmoidal equation was performed to obtain the melting temperature (Tₘ) using PRISM 9 (GraphPad Software, San Diego, CA, USA).

Structure predictions

All structure predictions were performed using ALPHAFOLD2 [35] as implemented in COLABFOLD (https://colab.research.google.com/github/sokrypton/ColabFold/blob/main/AlphaFold2.ipynb) with default settings and allowing the structures to relax using the built-in amber [36] functionality.

Molecular graphics and visualization

Cartoon representations of Nsp1 structures were generated using CHIMERAX [37]. Schematic representations of structure topology were drawn in TOPDRAW [38].

Results

Circulating deletions in SARS-CoV-2 Nsp1 differ in their inhibitory effect on translation

Several amino acid deletions are reported in SARS-CoV-2 Nsp1: Δ85, Δ82–83, 85 and Δ79–89 in the N-terminal domain (NTD) [25] and Δ141–143 in the C-
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terminal domain (CTD) [26] (Fig. 1A). These deletions have been correlated to clinical characteristics, but the mechanistic biochemical basis of why they alter the course of COVID-19 has not been determined. We wished to assess the effect of these deletions on Nsp1’s ability to shut down host translation. As a first step toward this analysis, we purified wild-type Nsp1 and a series of deletion mutants to homogeneity and monodispersity (Fig. S1). We then established a translation assay based on lysates of the human cell line HEK293F. Since Nsp1 has been reported to reduce translation of viral mRNAs to a lesser degree than cellular mRNAs [14,17,21], we reasoned that a reporter mRNA resembling a viral mRNA would provide a more sensitive assay. Thus, we measured translation efficiency of a reporter mRNA encoding firefly luciferase, equipped with the SARS-CoV-2 5’ UTR, a 5’ cap, and 3’ poly(A) sequence, in the presence of increasing concentrations of both wild-type recombinant Nsp1 and its deletion variants (Fig. 1B).

In agreement with earlier reports [17,21], wild-type Nsp1 efficiently abrogates production of firefly luciferase in concentration-dependent manner. Already at 0.1 μM, wild-type (wt) protein reduces the translational efficiency by more than 95% (Fig. 1C). An effect similar to the wt is observed for the NTD deletion variant Δ85 and the CTD deletion variant Δ141–143. The NTD deletion variant Δ82–83, 85 shows slightly reduced suppression of translation compared to the wild-type. In contrast, the longest deletion in the NTD, Δ79–89, is substantially weakened in its ability to inhibit translation as compared to wt Nsp1 (Fig. 1C). We observed that even at 3 μM protein concentration, the translation efficiency reduces only to 40–50%, establishing that the Δ79–89 protein is less effective in translation shutdown than the wild-type and other deletion variants. Notably, the Δ79–89 protein variant is still able to bind the human 40S ribosomal subunit (Fig. 2).

Having investigated how Nsp1 variants inhibit the translation of viral mRNA, we next turned to the question of whether they are differentially efficient in inhibiting viral and nonviral (host) mRNA. We thus repeated the translation assay with mRNAs having either SARS-CoV-2 or a nonviral 5’ UTR (Fig. 1B). Consistent with previous report by Schubert et al. [17], the viral mRNA was more strongly translated in the absence of Nsp1 (Fig. 1D). We then repeated this assay in the presence of wt and Δ79–89 Nsp1, both used at 0.5 μM final concentration. Both Nsp1 variants had the same relative effect on translation of viral and nonviral mRNA, thus leading to a higher absolute level of translation of the viral mRNA in presence of either Nsp1 (Fig. 1D). Note that the experiments presented in Fig. 1C,D were conducted with different batches of translation lysate. They can thus be qualitatively, but not quantitatively compared to each other. Taken together, these data establish a biochemical basis for correlating COVID-19 disease progression to the translational shutdown efficiency of circulating deletion mutants in Nsp1.

Deletions in Nsp1 lead to altered protein stability that correlates with translation inhibition

None of the investigated nsp1 mutations was in the region previously reported to interact with the ribosome, and yet the longer deletions in the NTD were clearly altered in their translation shutdown capacity. We wanted to investigate whether this effect could be due to the destabilizing effects of the deletions on the NTD. We first noted that all proteins were soluble and monodisperse when purified from E. coli, but the Δ79–89 construct eluted at a lower volume in size exclusion chromatography, indicating a larger hydrodynamic radius (Fig. S1). To probe the effects of the deletions on protein stability, we assessed the thermal stability of wt Nsp1 and deletion variants by thermal shift assay (TSA) [39] (Fig. 3). In this assay, SYPRO orange dye is used as a probe to
estimate the extent of unfolding of the protein with increasing temperature, and the melting temperature (T_m) from each curve is determined as described in the experimental procedure section. Raw TSA data (Fig. 3A,B) demonstrate that all mutants of SARS-CoV-2 Nsp1 showed similar, wild-type-like unfolding profile, apart from the longest deletion variant (Δ79–89). Wt Nsp1 and all altered proteins, with a notable exception of Δ79–89, allowed for fitting of a simple melting curve (Fig. 3C) indicating a single transition temperature that is comparable to that of the wt protein (Fig. 3E). The thermal denaturation curve of Δ79–89 (Fig. 3A,B) was shallow and could not be perfectly recapitulated by fit to a single melting temperature (Fig. 3D). However, using the first part of the curve gave a fit that is consistent with the qualitative appearance of the curve (Fig. 3D) and a lowered thermal stability. This suggests that already at room temperature the structural integrity of Δ79–89 variant is compromised. In summary, our results establish a correlation between structural stability of Nsp1’s NTD and its ability to inhibit translation.

Structure predictions suggest that NTD β-barrel destabilization causes the decreased stability of Δ79–89 nsp1

We next wanted to explore the possible structural basis for the decreased translation inhibition and thermal stability of the altered Nsp1 proteins. To investigate how the Nsp1 deletions impact its structure we predicted the structures of wt, Δ141–143, Δ85, Δ82–83, 85, and Δ79–89 Nsp1 using alphafold 2 [35]. Since there are available experimental structure of the free SARS-CoV-2 Nsp1 NTD (residues 10–126) [27], as well as the ribosome-inserted CTD [15–17], we could compare them to the prediction of the wt Nsp1 structure as a baseline. The predicted Nsp1 structure aligns very well with the experimentally determined N-terminal β-barrel domain, with average positional shifts (root-mean-square deviation, RMSD) of the respective Cα atoms of only 0.64 Å for residues 10–126 (Fig. 4A, Fig. S2A). In the CTD, two α-helices are correctly predicted where they are observed in the experimental structure, albeit at a different angle to each other than in the 40S-bound structures.
Overall, the per-residue prediction quality score (pLDDT) correlated with the rigidity of the fold across Nsp1, with the NTD β-barrel having higher scores than the CTD which was reported to be flexible in solution (Fig. S2B) [19]. Supported by the accurate prediction of the wt Nsp1 structure, we thus reasoned that structure predictions for the altered Nsp1 proteins may shed light on the differential effects of the deletions on protein stability. Unsurprisingly, the deletion Δ141–143, predicted to be located in a disordered region of the CTD, had no effect on the predicted fold of other parts of Nsp1 (Fig. S2). The other three deletions (Δ85, Δ82–83, 85, and Δ79–89) are all located at the beginning of the fourth β-strand of the β-barrel (β4) (Fig. 4B,D). Interestingly, despite being located in a secondary structure element, the two shorter deletions are not predicted to affect the β-barrel integrity (Fig. S2C), likely given that the hydrogen bonding between β-strands is mediated by the protein backbone and can be rescued by the residues ‘next-in-line’ to the deleted residues. The predicted integrity of the fold of these deletions correlates well with their unaltered thermal stability (Fig. 3E). The only striking effect on the integrity of the β-barrel domain is predicted to stem from the longest deletion. The Δ79–89 structure prediction suggests a near-complete dissolution of the β3–β5 strands leading to a break in the β-barrel domain, severely affecting its integrity (Fig. 4C,E). This prediction is in line with the drastic reduction in thermal stability for the Δ79–89 Nsp1 protein from 46 °C to ~27 °C (Fig. 3E). Taken together, structure predictions of the Nsp1 deletions suggest destabilization of the N-terminal β-barrel domain as the mechanism behind the altered properties of the Δ79–89 Nsp1 protein.

**Discussion**

The high transmissibility of SARS-CoV-2 and the lack of pre-existing immunity contributed to an explosive development of the COVID-19 pandemic. This happening in the age of readily available RNA sequencing has given a fine-grained view of the mutations taking place in the viral genome in the course of the pandemic. The nonstructural protein Nsp1 is a major pathogenicity factor for coronaviruses [8]. Although Nsp1 has remained unaltered in the major SARS-CoV-2 variants, certain internal deletions in Nsp1 (Fig. 1A) are regularly detected in clinical isolates and
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PK was supported by a Kempe Foundations postdoctoral fellowship, and ES was supported by a Umeå University Excellence by Choice (EC) postdoctoral fellowship. L-AC was supported by the Human Frontier Science Program (Career Development Award CDA00047/2017-C) and the Knut and Alice Wallenberg Foundation (through the Wallenberg Centre for Molecular Medicine Umeå). VH was supported by grants from Cancerfonden (20 0872 P), the Knut and Alice Wallenberg Foundation (grant 2020.0037), the Estonian Research Council (grant PRG335), and the Swedish Research council (grants 2017-03783 and 2021-01146). The project was supported through an exploratory grant for COVID-19-related research, Medical faculty at Umeå University, 2020, to L-AC and VH.

Author contributions

IT, VH, LAC conceived the study; PK, ES, KR, TK performed experiments; PK, KR, TK, IT, VH, LAC analysed data; PK, KR, VH, LAC wrote the initial manuscript; all authors read and edited the manuscript.

Data accessibility

The data that support the findings of this study are available from the corresponding author (lars-anders.carlson@umu.se) upon reasonable request.

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have been correlated to different clinical outcomes [25,26]. The repeated appearance of certain deletions indicates that they are either better tolerated without losing infectivity, or that they are more likely to appear due to peculiarities of the RNA replication process. In this study, we used a human translation lysate assay to compare the potency of these Nsp1 deletion mutants in inhibiting translation. Notably, none of the deletions encompassed the two motifs previously identified as being necessary for Nsp1’s ability to inhibit translation [12], and mediate mRNA degradation [11], respectively (Fig. 1A). Thus, it was not clear a priori what their effect on translation would be, if any. The main finding of this study is that several of these circulating Nsp1 deletion mutants are still highly potent in inhibiting translation in vitro, except for the longest deletion, Δ79–89. Whereas still soluble and monodisperse, the Δ79–89 protein was no longer able to shut down translation completely, even when present at thirtyfold higher concentrations than the wild-type protein. We show that Δ79–89 Nsp1 also stands out from the other deletion mutants by having severely reduced thermal stability. Structure predictions of all mutants were consistent with this and suggested that the mechanism for the lost stability of the Δ79–89 protein is a rearrangement leading to a disruption of its N-terminal β-barrel domain fold.

The deletions around residues 79–89 in Nsp1 were previously characterized in terms of their effect on transcriptome and type I interferon response [25]. A side-by-side comparison of those results with our translation shutdown data reveals some interesting differences. Strikingly, the shorter deletions in this region already alter the transcriptome and lead to a much weakened type I IFN response as compared to the 180 residue version of Nsp1. These shorter deletions are not impaired in translation shutdown at the concentrations tested, indicating that this region of Nsp1 may have other functions. Notably, the deletions flank the only long loop connecting two strands of the β-barrel. This loop was reported to be flexible in the NMR structure of SARS-CoV Nsp1 [28], but it is to date unknown whether is involved in interactions with host or viral components.

In summary, we determined the translation shutdown capacity of several clinically detected deletions in SARS-CoV-2 Nsp1. The data show that a compromised stability of the N-terminal β-barrel abrogates translational shutdown, and further indicates that the shorter deletions around residue 79–89 may affect interferon response and transcriptome through a mechanism independent of translation shutdown.
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**Supporting information**

Additional supporting information may be found online in the Supporting Information section at the end of the article.

**Fig. S1.** Purification of wild type and deletion variants of SARS-CoV-2 Nsp1. (A-E) Chromatograms (280 nm absorption) of Nsp1 and its deletion variants eluting from a Superdex 75 increase 10/300 size-exclusion column. Each chromatogram has the volume corresponding to the center of the main peak noted. The A260/A280 ratio was similar (~0.6) in the main peak of all constructs. (F) 1.5 μg of each protein sample was analyzed on a 15% SDS-PAGE and stained with Coomassie blue.

**Fig. S2.** AlphaFold2-predicted Nsp1 structures. (A) (left) Structures of the experimentally determined N- (residues 10-126, PDBID: 7K7P[27]) and C-terminal (residues 148-180 PDBID: 6ZOJ[17]) domains of Nsp1. Structures are oriented as the predicted full-length structure of wt Nsp1 in (B). (right) Superposition of the predicted wt Nsp1 structure with the experimentally determined structures as in (A). (B) Predicted structure of full length wt Nsp1 shown in cartoon representation (grey) and the corresponding per-residue confidence score (pLDDT) trace for all five prediction runs. (C) Cartoon representations of the predicted structures of Δ141-143 (light blue), Δ85 (green), Δ82-83,85 (purple) and Δ79-89 (orange) Nsp1 and their respective per-residue confidence score (pLDDT) trace for each of their predictions.