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Biophysical characterisation of the nucleocapsid protein from a highly pathogenic porcine reproductive and respiratory syndrome virus strain

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Abstract

The arterivirus nucleocapsid (N) protein is a multifunctional protein that binds viral RNA for encapsidation and has potential roles in host cell processes. This study characterised the N protein from a highly virulent North American strain of porcine reproductive and respiratory syndrome virus (PRRSV). The association with viral RNA was mapped to defined motifs on the N protein. The results indicated that disulphide bridge formation played a key role in RNA binding, offering an explanation why infectious virus cannot be rescued if cysteine residues are mutated, and that multiple sites may promote RNA binding.

1. Introduction

Coronaviruses and arteriviruses are families of positive strand RNA viruses that replicate in the cytoplasm of an infected cell. Due to similar genome organisation, replication and transcription strategies these viruses are grouped together into the Nidovirales. Many of their members cause important respiratory, gastrointestinal and reproductive diseases. These include severe acute respiratory syndrome coronavirus (SARS-CoV) [1], a zoonotic infection of humans and animals, and the arterivirus porcine reproductive and respiratory syndrome virus (PRRSV) that causes reproductive failure in pregnant sows, high mortality in piglets and respiratory disease in pigs of any age [2,3]. Although vaccines to a number of respiratory syndrome coronavirus (EAV) and PRRSV N proteins have been solved and show structural similarities to the CTD of the coronavirus N protein [16]. Due to the abundance of disordered regions and proteolytic instability of the full length N protein, the detailed structure of the N protein has not been solved in entirety using any one single technique and our picture of this is based on multiple complementary analyses.

The PRRSV genome shows rapid change and strains can vary in virulence [7], with various genes contributing to pathogenesis, including the nucleocapsid (N) protein [8–10]. For both the coronaviruses and arteriviruses the N protein is one of the most abundant virally encoded proteins and plays essential roles in the virus life cycle [11,12]. Its most prominent role is complexing with viral RNA, although the mechanisms for this for either type of viral N protein is not precisely characterised. For the coronavirus N protein phosphorylation has been shown to play a role in mediating binding to RNA [13,14] with the amino-terminal region facilitating long-range non-specific electrostatic interactions potentially allowing contacts to form via a ‘lure’ and ‘lock’ mechanism [15].

Sequence comparison initially suggested that coronavirus N proteins exhibited a three-domain structure. This includes a disordered unstructured region at the N-terminal end that is followed by the structured N-terminal domain (NTD). A disordered serine/arginine- (SR-) rich region is located in the central position of the coronavirus N protein. Similarly the extreme C-terminus is disordered with a proximally structured C-terminal domain (CTD). The arterivirus N proteins are smaller in size with regard to molecular weight (the coronavirus N protein is approximately 45 kDa whereas the arterivirus N protein is approximately 15 kDa) and consists of an N-terminal half that is mainly disordered and a structured C-terminal half which has a similar fold as the CTD of the coronaviruses [16]. Due to the abundance of disordered regions in combination with proteolytic instability of the full length N proteins the detailed structure of the N protein has not been solved in entirety using any one single technique and our picture of this is based on multiple complementary analyses.

The X-ray structures of the CTD of the arterivirus equine arteritis virus (EAV) and PRRSV N proteins have been solved and show structural similarities to the CTD of the coronavirus N protein [16,18,19]. There is a very high overall similarity between the CTD of EAV and PRRSV N proteins. The main differences exist in
their $\beta$-loops and their most C-terminal $\alpha$-helices that end about 6 residues before the C-terminus. Monomers of EAV and PRRSV N harbour two anti-parallel $\beta$-strands that are flanked by two $\alpha$-helices. Both proteins contain another short $\alpha$-helix towards the N-terminal of the CTD. Moreover, PRRSV as well as EAV N are predicted to contain an $\alpha$-helix within their otherwise unstructured N-terminal half. Similar to the coronavirus N proteins EAV and PRRSV N form dimers with a $\beta$-sheet platform that is surrounded on one side of the dimer by the $\alpha$-helices. With a view to potential RNA binding or other interactions taking advantage of electrostatic interactions, the electrostatic potential is more positive on the side of the dimer on which the helices reside. The proposed RNA binding region of the PRRSV N protein, based on the structure, is in the unstructured N-terminal half, located within a region harbouring positively charged residues. With a negative electrostatic potential found on the side of the $\beta$-sheet floor this could support correct positioning of the RNA in complex to N protein. PRRSV N is found in a head to head helical arrangement within the crystals, supporting the ribbon-like pattern observed for N using cryo-tomography [20].

To provide a deeper understanding of the role of PRRSV N protein in binding RNA and to delineate RNA binding regions a sequential alanine substitution approach was used in combination with various biophysical assays and selected further point mutation. This analysis highlighted that several sites on the PRRSV protein may act cooperatively to promote RNA binding.

2. Materials and methods

2.1. Bacterial strains and construction of plasmids

The N gene from PRRSV strain NVSL #97-7895 was cloned into the pTrEx1.1Neo (Novagen) vector. Alanine block substitutions were constructed by a two-step PCR approach creating products allowing for a triple ligation with the pTrEx1.1Neo vector. Plasmid pTrExNC23S coding for an N protein carrying a cysteine (cys) to serine (mutant NC23S), and resulted in co-purification of no or very small amounts of RNA (Fig. 3B).

3. Results and discussion

3.1. The N protein interacts with RNA

To investigate RNA binding and establish an in vitro RNA binding assay for the N protein, the N gene from an infectious cDNA of the highly lethal North American PRRSV strain NVSL #97-7895 [22] was cloned into the expression vector pTrEx1.1Neo (Novagen) and the recombinant protein purified from E. coli using metal affinity chromatography as described [13]. RNA co-purification was used to indicate the RNA binding capability of N protein. The amount of RNA that could be extracted from an equal amount of purified protein decreased when the protein was incubated with increased RNase A concentration (Fig. 1A). To control for protein stability, protein samples were taken at the end of the incubation and analysed by SDS–PAGE (Fig. 1A).

The size of the co-purified RNA was estimated by electrophoretic analysis and a major species found to be approximately 150 nucleotides. To investigate whether the 150 nucleotides were bound by one N protein molecule, analytical gel filtration chromatography (GFC) was performed (Fig. 1B). These molecular weights indicated that the two peaks contained multimers and possibly monomers, respectively. From the mass deduced from the elution time it is tempting to speculate that peak 1 represents a dimer, suggesting that each monomer protected an average of 15 nt from nucleolytic cleavage.

3.2. Global structure of the PRRSV N protein

No complete structure for the PRRSV N protein is available, therefore a combination of modelling using the sequence encompassing the CTD of the N protein from the VR-2332 strain was used to map the equivalent region on the NVSL #97-7895 strain N protein using the HHpred program (pdb file 1p65; http://toolkit.tuebingen.mpg.de/hhpred) and the secondary structure prediction program Jpred (http://www.compbio.dundee.ac.uk/www-jpred/) was employed to model potential folds within the N-terminal half (NTH) (Fig 2A). Overall the structure was confirmed using FRODO. For example, these suggested 15% alpha helix, 29% beta sheet and 56% random coil by the K2d neural network programme.

3.3. Potential role of disulphide bridge formation in RNA binding

The North American PRRSV N protein contains three cys residues at positions 23, 75 and 90 and has been shown to be present as disulphide-linked dimers in virions [23,24]. However, only equivalents of Cys23 and Cys75 are present within N proteins from European PRRSV strains. Cys23 and Cys90 may play an important role for in the virus life cycle as substitution of these residues with serine in recombinant infectious virus led to abrogation of progeny virus production [25]. To determine whether the cys residues had possible roles in mediating RNA binding, N protein was purified under reducing conditions and resulted in less RNA being co-purified (Fig 3A). This indicates that the formation of disulphide linkages may play a role in RNA binding. Examination of the predicted N protein structure (Fig. 2A) indicated that Cys75 and Cys90 are too far away from their partners in the partial crystal structure to form disulphide bridges [19]. The possible role of Cys23 in RNA-binding was investigated by substituting this residue with serine (mutant Ncys23s), and resulted in co-purification of no or very small amounts of RNA (Fig. 3B).
The influence of disulphide bridge formation on RNA binding may be due to a higher valency that is achieved upon N protein.

**Fig. 1.** RNA binding by the N protein. (A) Ethidium bromide stained acrylamide gel loaded with RNA extracted from equal amounts of purified recombinant N protein. Lanes 1, 2, 3, 4, 5, 6 and 7 contain RNA extracted from protein samples treated with 0, 5, 10, 20, 40, 80 and 160 ng/ml RNase A for 10 min at 30 °C, respectively. Sizes of the ssRNA marker are indicated on the left. Below the RNA gel, protein samples taken just prior to RNA extraction and analysed by SDS–PAGE are shown as control for protein stability during RNase A treatment. (B) Chromatogram obtained during analytical gel filtration chromatography of the N protein (blue trace). The inset shows the standard curve used to estimate the molecular weight of the species within the peak. The position of peaks potentially containing decamer (Peak 1) and monomers (Peak 2) of N protein are indicated by arrows. Both graphs have the retention time (R_t) plotted on the x-axis. The y-axis contains information about the absorbance units and logarithmic molecular weights for the N protein graph and the standard curve, respectively. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

**Fig. 2.** (A) Predicted structure of PRRSV N protein which harbours one structured domain that is located in the C-terminal half. The N-terminal half of the protein is thought to be mainly unstructured besides a predicted α-helix. Shading in alternating red and green shows the positioning of the alanine block substitutions (corresponding to those in Table 1) used in this study on the predicted structure. (B) Representative CD spectra of the PRRSV protein in different concentrations of NaF. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

**Fig. 3.** RNA binding is influenced by disulphide bond formation. Shown are denaturing acrylamide gels onto which RNA was loaded which was extracted from equal amounts of protein. (A) RNA extracted from wild type N protein purified under reducing (+DTT) or non-reducing (−DTT) conditions. (B) RNA extracted from wild type N protein and the cys to serine mutant N_C23S that were purified under non-reducing conditions. Sizes of marker bands are indicated on the left of the panels.

The influence of disulphide bridge formation on RNA binding may be due to a higher valency that is achieved upon N protein.
oligomerization and stabilization by intermolecular disulphide bridges: A single N protein may only have one RNA binding site of low affinity, whereas oligomerization may lead to the availability of several RNA binding sites that could interact with the same RNA molecule. Multi-valency is a known means to increase the affinity of low affinity binding partners [26–28]. The suggested arrangement of N protein in the ribonucleocapsid leads to the close proximity of the positively charged regions surrounding the α-helix within the NTH of the protein [20,29]. Cys23 is contained within this α-helix and therefore may stabilize this arrangement upon formation of disulphide bridges. Alternatively, mutation of Cys23 may have disrupted other functions of the protein in terms of overall structure and underlying RNA binding signals.

3.4. The CTD is important for dimer formation whereas Cys23 stabilizes higher oligomers

To investigate whether other regions of the N protein interacted to promote RNA binding, an alanine block substitution approach was adopted. Each protein variant had sequentially 10 residues mutated into alanines. Each substituted block is indicated in the predicted structure of N protein (Fig. 2A), and the predicted structure and electrostatic potential of each variant protein is shown in Supplementary (Fig. S1). As the oligomeric state and valency of molecules can influence binding affinity the mutant N proteins were assessed with regard to their oligomeric state. Wild type N protein as well as the mutants were purified under non-reducing conditions and high RNase A concentrations to prevent co-purification of RNA. RNA has previously been shown to influence the oligomeric state of N protein [24]. To assess size and purity of the recombinant proteins, samples were analysed by SDS–PAGE (Fig. 4A). Due to the highly charged character of the N protein some of the alanine substitutions that cause a change in the charge of the protein resulted in different electrophoretic mobility.

To assess the oligomeric state under native conditions, the purified proteins were separated on native blue gels (Fig. 4B), allowing electrophoretic analysis under native conditions of proteins with high isoelectric points (pIs). N protein showed monomers, dimers, trimers and tetraders. Mutant N21–30 lacking Cys23 had a different migration pattern. This mutant is mainly present as monomer and dimer suggesting that Cys23 was involved in the formation of higher order oligomers. Alanine blocks within the well-structured CTH had a more dramatic effect on oligomer formation of N protein: Mutants N61–70, N71–80, N81–90, N91–100 and N101–110 showed significantly less discrete staining. Mutants N71–80, N81–90, N91–100 and N101–110 were mainly present as monomers. Together with the finding that N111–123 could form oligomers indicated that amino acids 30–37 constituted not the only polypeptide sequence needed for dimer formation but that the CTH of N protein plays the main role in this process. As has been suggested by the crystal structure of the CTH.

3.5. Role of amino acid motifs in RNA binding

To investigate the role of different amino acids in RNA binding, the alanine block mutants were purified under non-reducing conditions that allowed co-purification of RNA with wild type N protein. RNA was extracted from purified proteins and compared to the amount extracted from equal amounts of wild type N protein. Table 1 summarises the results of this experiment. No RNA was co-purified with mutants N21–30, N41–50 and N111–123 indicating that residues within these regions may play a role in RNA binding. Yoo et al. [31] identified the region between residues 37 and 57 as containing RNA binding activity. The alanine block of mutant N41–50 is located within this region and specifically covers a region with a number of charged residues. It is possible that these positively charged residues bind to the negatively charged phosphate backbone of RNA. The alanine block of mutant N21–30 is not located within the previously identified RNA binding region. However, it is located within the predicted α-helix within the NTH of the protein (Fig. 2A). Moreover, mutant N21–30 harbours Cys23 whose mutation affects RNA binding. Also mutant N111–123 did not co-purify any RNA. As deletions of the last 11 residues of PRRSV N protein influence its conformation [30], indicating that conformational changes may inhibit RNA binding.

In summary, PRRSV N protein contains a number of different functional sites involved in both recognition of viral RNA, sub-cellular localisation and interactions with other viral and cellular proteins [12,31]. Rather than individual motifs acting in isolation, both oligomerisation of the N protein and direct RNA binding motifs are likely to act in concert to promote RNA binding.

| RNA co-purified | WT | N11–20 | N21–30 | N31–40 | N41–50 | N51–60 | N61–70 | N71–80 | N81–90 | N91–100 | N101–110 | N111–123 |
|----------------|----|--------|--------|--------|--------|--------|--------|--------|--------|--------|--------|---------|
|                | ***| *      | –      | **     | –      | *      | **     | +      | **     | *      | ND      | ±/–     |

**Table 1**

RNA binding of N protein variants. The amount of RNA binding is indicated with + and – signs where a triple *** indicates high binding affinity. ND, not determined.
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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.bbrc.2011.11.126.

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