Zinc-binding properties of Junín virus nucleocapsid protein

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The arenavirus nucleocapsid protein (N) is a highly basic 63 kDa protein with a dual function during the virus life-cycle. First, it is involved in essential steps of genome replication, promoting the synthesis of the full-length antigenomic copy of S RNA, and second it associates with the genomic RNA to form the nucleocapsid. We have expressed the N protein of Junin virus in E. coli and shown that it binds zinc in vitro. This property is in agreement with the presence in the carboxy-terminal region of the N protein of the CX2HX23CX4C sequence, which resembles a classical zinc-finger motif. The specificity for zinc binding was demonstrated by competition with other divalent metal ions. The ability of the predicted motif to bind zinc was established by analysis of a series of N mutants, including truncated variants and amino acid substitutions. In addition, alternative zinc-binding sites were found.

Introduction

Arenaviruses are enveloped viruses with a genome composed of two single-stranded RNA species, designated L (large, ca. 7 kb) and S (small, ca. 3.5 kb). Junin virus, a South American arenavirus, is the aetiological agent of a severe endemico-epidemic disease called Argentine haemorrhagic fever. The complete nucleotide sequence of the S RNA of Junin virus has been determined for different strains (Ghiringhelli et al., 1991, 1997; Albariño et al., 1997, 1998). The S RNA encodes the major structural proteins of the virion, which are the precursor of the envelope glycoproteins (GPC) and the viral nucleocapsid protein (N). Open reading frames are arranged in opposite orientations (ambisense coding strategy; Auperin et al., 1984) and are separated by a non-coding intergenic region that folds into a stable stem-loop secondary structure (Ghiringhelli et al., 1991). Once the virus enters a cell, the genomic form of the S RNA is transcribed and yields only two antigenomic forms: the 1.8 kb N mRNA and a 3.4 kb full-length antigenomic S RNA. The N mRNA is the first RNA species to be transcribed in the infected cell, and the structure of the intergenic region seems to be the transcriptional termination signal (Franze-Fernández et al., 1993; Tortorici et al., 2000). The N protein is a highly basic protein, translated from an antigenomic sense mRNA species that is encoded in the 3′ half of the viral S RNA (Romanowski, 1993). N displays a dual function during the viral life-cycle: it associates tightly with the genomic RNA to form the nucleocapsids and, as recently demonstrated, it is essential for genome replication, since it promotes the synthesis of the genomic replicative form in the viral life-cycle (Lee et al., 2000; Tortorici et al., 2000).

In general, RNA–protein interactions involve both specific and non-specific contacts and a number of protein sequences and/or structural motifs have evolved to allow such interaction. Among these motifs are the classical zinc fingers or the general RING fingers, which are autonomously folding domains that require zinc for proper folding and RNA-binding activity (Nolte et al., 1998; McColl et al., 1999). We found one classical zinc-finger domain near the carboxy terminus of the N protein sequence (Paris et al., 1996). In this work we demonstrate that N protein binds zinc specifically. Furthermore, we analysed a series of mutant N proteins to characterize the zinc-binding region. The experimental results demonstrate that the zinc-binding activity is located at the carboxy-terminal region of the N protein and that the substitution of critical amino acid residues eliminates binding in the isolated zinc finger, but alternative binding sites located in the immediately adjacent or partially overlapping region, may bind Zn²⁺ with a slightly lower affinity.
**Methods**

**Molecular cloning and oligonucleotides.** Plasmids were propagated in *E. coli* DH5α grown in Luria–Bertani broth. Standard procedures were used for DNA manipulations (Sambrook *et al.*, 1989) according to manufacturers’ protocols. All DNA oligomers are listed in Table 1, as are the positions of the nucleotide sequences used to obtain the mutant proteins (translation initiation codon, translation stop codon and amino acid substitutions).

**Sequence alignments.** Alignment of arenavirus N protein sequences was done using the Clustal X program (Thompson *et al.*, 1997) and the substitution matrix PAM 250. The parameters used for the pairwise and multiple alignment were GAP opening = 10, GAP extension = 0.1 and GAP opening = 10, GAP extension = 0.05, respectively.

**Construction of deletion and substitution mutants of N gene.** Plasmid pBJUNS (Ghiringhelli *et al.*, 1991) containing the entire N ORF of Junin virus (strain MC2) was the progenitor of plasmids used in this work. The N ORF was amplified by PCR using primers N1 and N2 (Table 1). The PCR product was inserted into pGEM-T (Promega). The N ORF was recovered by digestion with *Nco*I and *Bam*HI, filled-in using the Klenow fragment of DNA polymerase I, and inserted by ligation into the filled-in *Nde*I site of pET-22b(+) under the control of a bacteriophage T7 promoter (Novagen). The putative zinc-finger domain was generated by PCR using primers ZfD and N2 (Table 1). The PCR product was ligated to pET-22b(+) vector following the same strategy used for the N ORF. Mutations which resulted in carboxy-terminal deletions of the N ORF were made using the Erase-a-Base System (Promega). ExoIII/S1 deleted products containing the authentic methionine codon of the N protein were modified to include stop codons immediately after the last N amino acid. To this end, oligonucleotides ST1 and ST2 (see Table 1) were designed to incorporate stop codons in each of the open reading frames. The oligonucleotide mixture (25 µM each) was heated at 100 °C for 5 min, annealed in 100 mM NaCl, 10 mM Tris–HCl pH 7.5, 0.1 mM EDTA and the ST1/ST2 heteroduplex ligated to the ExoIII products. The deleted genes were finally generated by PCR, using primers N1 and ST2, and the amplified fragments were inserted into the *Nde*I site of pET-22b(+). Mutations that resulted in amino acid substitutions in the N gene were introduced by site-directed PCR mutagenesis and PCR combinations of overlapping fragments using *Pfu* polymerase and the appropriate combinations of primers listed in Table 1 (Higuchi, 1989; Higuchi *et al.*, 1989).

### Table 1. DNA oligonucleotides used in the study

All the mutagenized codons are indicated in bold.

<table>
<thead>
<tr>
<th>Name</th>
<th>Size (nt)</th>
<th>Sequence</th>
<th>Use and description</th>
</tr>
</thead>
<tbody>
<tr>
<td>N1</td>
<td>30</td>
<td>AGATCTGATCCATGGCACACCTCCAAAGAG</td>
<td>5′ end primer used to generate N(1–564). The translational initiation codon is indicated in bold and the <em>Nco</em>I site is underlined.</td>
</tr>
<tr>
<td>N2</td>
<td>31</td>
<td>GAGATCTGATCCCTACAGTGCATAGGCTGC</td>
<td>3′ end primer used to generate N(1–564), N(478–564) and point mutant genes. The <em>Bam</em>HI site is underlined. The sequence complementary to the translational stop codon is indicated in bold.</td>
</tr>
<tr>
<td>ZfD*</td>
<td>28</td>
<td>GCATGCCATGGGAGAACAAAGCCAGGAGA</td>
<td>5′ end primer used to generate N(478–564). Translation initiation codon is indicated in bold and the <em>Nco</em>I site is underlined. S to G conservative change was introduced to generate the <em>Nco</em>I site (mutagenized codon is boxed).</td>
</tr>
<tr>
<td>ST1</td>
<td>11</td>
<td>TAATTTGATTA</td>
<td>These complementary oligos were designed to incorporate stop codons (in bold) in the three open reading frames into the deletion mutant.</td>
</tr>
<tr>
<td>ST2†</td>
<td>15</td>
<td>GATCCCTAATCAAATT</td>
<td></td>
</tr>
<tr>
<td>ZfD(C525A)V</td>
<td>26</td>
<td>CTAGCAGACAGCCGTTGGCTCAG</td>
<td>5′ end primer used to generate a mutant with the C⁵₂⁵ → A substitution.</td>
</tr>
<tr>
<td>ZfD(C530A)V</td>
<td>31</td>
<td>ACACTGACTGAAACATTTAGCGCTACAGCAG</td>
<td>5′ end primer used to introduce a C⁵₃₀ → A substitution.</td>
</tr>
<tr>
<td>ZfD(C497A; H500N)V</td>
<td>29</td>
<td>ACTCCATTGTTATGTTGGCTAGTGGCC</td>
<td>5′ end primer used to introduce the C⁴₉⁷ → A substitution and the H⁵₀₀ → N substitution.</td>
</tr>
<tr>
<td>ZfD(C525A)VC</td>
<td>26</td>
<td>CAGTGGCACCACCGCTGCTCTGCTAG</td>
<td>3′ end primer used to introduce a C⁵₂⁵ → A substitution.</td>
</tr>
<tr>
<td>ZfD(C530A)VC</td>
<td>31</td>
<td>CTGCTAGACGCCATAATGTTTCAGTCTAG</td>
<td>3′ end primer used to introduce a C⁵₃₀ → A substitution.</td>
</tr>
<tr>
<td>ZfD(C497A; H500N)VC</td>
<td>29</td>
<td>GGCACACTAGCCACAAATACAAATGAGT</td>
<td>3′ end primer used to introduce the C⁴₉⁷ → A substitution and the H⁵₀₀ → N substitution.</td>
</tr>
</tbody>
</table>

* ZfD, zinc-finger domain.
† ST1/ST2, three frame stop adapters.
Zinc binding by Junin virus N protein

Fig. 1. Multiple sequence alignment of the arenavirus nucleocapsid protein. The alignment was done using the Clustal X program. Gaps were included to maximize alignment of the homologous amino acids. The sequence C497H500C525C530 represents the amino acids involved in zinc-coordination in a zinc finger-like domain (shadowed). The numbers represent the amino acid positions in the N protein sequence of Junin virus. Arrows indicate positions of alternative amino acid residues that could coordinate zinc. Abbreviations: LAG, Lassa-Nigeria; LAJ, Lassa-Josiah; LCA, LCM-Armstrong; LCE, LCM-WE; MOP, Mopeia; CD1, Junin-Candid 1; MC2, Junin-MC2; MAC, Machupo; OLI, Oliveros; PIC, Pichinde; SAB, Sabia; TAC, Tacaribe.

Horton et al., 1990). In addition, carboxy-terminal deletions were introduced in the N ORF containing the point mutations using N1 primer plus ZfD(C497A; H500N)V and N1 plus ZfD(C530A)V (see Table 1). The N gene containing four amino acid substitutions in the full-length N ORF was used as a template. The ST1/ST2 heteroduplex was ligated to the amplified fragments. The mutant genes were finally generated by PCR using primers N1 and ST2 and were inserted into the NdeI site of pET-22b(+) vector. All these constructs were designed to express the heterologous protein starting from its own methionine codon and with no fused sequences at its carboxy terminus.

DNA sequencing. The identity of mutants was confirmed by sequencing via the chain-termination method (Sanger et al., 1977) using the fmol DNA Sequencing System (Promega).

Protein expression. E. coli BL21(DE3) cells were transformed with recombinant plasmids, and recombinant cells were cloned. Protein expression was routinely induced with 1 mM IPTG for 4 h. E. coli cells were collected by centrifugation, rinsed with PBS, resuspended in 50 mM Tris–HCl pH 8–0, 100 mM NaCl, 1 mM EDTA, 0.5% Nonidet P-40 plus 0.2 mM PMSF and disrupted by sonication. After centrifugation, the recombinant proteins were collected from the insoluble fraction and analysed by SDS–PAGE (Laemmli, 1970).

Immunoblotting. Samples of bacterial extracts were separated by SDS–PAGE (10–20% gradient polyacrylamide gel), blotted onto nitrocellulose membranes (Schleicher & Schuell), and probed with a 1/100 dilution of human antiserum specific for Junin virus. Filters were then incubated with a goat anti-human antibody conjugated to alkaline phosphatase (Sigma) and developed using the chromogenic substrates NBT–BCIP (Promega).

Zinc-binding. The zinc-binding assay was carried out basically as described by Keck et al. (1993) with some modifications. Briefly, after electrophoresis equivalent gels were either stained with Coomassie brilliant blue or transferred to nitrocellulose membranes. The transferred proteins were renatured on the membranes by incubation at room temperature for 1 h in buffer A (100 mM Tris–HCl pH 7–5, 50 mM NaCl, 1 mM dithiothreitol). The membranes were briefly washed in the metal binding buffer (100 mM Tris–HCl pH 7–5, 50 mM NaCl) and probed for 1 h with 15 µM "&ZnCl₂ (3–12 mCi/mg; NEN Dupont) in the same buffer. The membranes were then washed twice over a 30 min period in buffer A. In competition experiments, divalent metal ions (10 mM) were included in the metal binding buffer during the incubation with "&ZnCl₂. Finally, nitrocellulose membranes were wrapped in Saran Wrap and autoradiographed using Kodak X-Omat film.

Results

Zinc-binding activity of the N protein in vitro

As shown in Fig. 1, a region with the potential to form a classical zinc-finger motif was found in the N protein amino acid sequence: C₄ₓHₓ₂₃CₓₓC (residues 497–530) (Parisi et al., 1996). In order to test the zinc-binding activity of the N protein, the N gene was expressed in E. coli. The proteins from a gel similar to the one in Fig. 2(A), were transferred to a nitrocellulose membrane, allowed to renature and probed with classical zinc-finger.
The results showed that the 63-kDa protein encoded by the N ORF had zinc-binding activity (Fig. 2B).

To further characterize the metal-binding properties of the N protein, competition experiments were done to determine the specificity of the zinc-binding activity. Identical samples obtained from lysates of E. coli expressing the 63 kDa protein were loaded onto equivalent lanes of a polyacrylamide gel, electrophoresed and blotted onto a membrane. After transfer, the membrane was cut into equivalent strips, each representing one lane, and probed with 15 µM $^{65}\text{Zn}^{2+}$ in the presence or absence of unlabelled competitor metal ions. The results showed in Fig. 3 indicate that the competition efficiency was $\text{Zn}^{2+} \approx \text{Co}^{2+} > \text{Cu}^{2+} > \text{Mn}^{2+} > \text{Mg}^{2+}$.

The carboxy-terminal region of the N protein binds zinc in vitro

The predicted zinc-finger domain of the N protein CX$_2$HX$_{29}$CX$_4$C is located in the carboxy-terminal region, comprising amino acid residues 497 to 530. To determine whether this region of the 63 kDa protein binds zinc, an artificial gene encoding the zinc-finger domain was made by PCR. Carboxy-terminal deletion variants of N lacking the zinc-finger domain were also generated. The amplified products were inserted into pET-22b($\varnothing$). The heterologous proteins were expressed in E. coli, recovered from the insoluble fraction and separated on a triplicate set of 10–20% gradient gels.
Zinc binding by Junın virus N protein

Fig. 4. Zinc binding by truncated N proteins. pET-22b(+) constructs containing the full-length N(1–564) and N(478–564) genes or truncated forms of N ORFs were expressed in E. coli and analysed by electrophoresis on an SDS–polyacrylamide gradient gel (10–20%). Gels were stained with Coomassie brilliant blue (A), blotted and probed with a Junın virus-specific antibody (B), or blotted and probed with $^{65}$Zn$_{2}$$^{+}$ (C). Truncated variants of N are identified by the positions of the amino- and carboxy-terminal amino acid residues indicated in parentheses. The full-length protein is designated N(1–564); the truncated forms of the N protein are designated N(1–393) and N(1–93) and the protein with the putative zinc-finger domain is designated N(478–564) (lanes 1, 2, 3 and 4, respectively). The horizontal grey bars represent the N-derived polypeptides. Cysteine and histidine residues in the predicted zinc-finger motif are indicated by C and H. The relative zinc-binding capacities of the proteins are summarized on the right (D).

The carboxy-terminal region of the N protein has alternative residues that bind zinc in vitro

To establish whether the zinc-binding activity was due to the classical zinc-finger motif, point mutations were made by PCR in the full-length N(1–564) and in the N(478–564) ORFs. Each cysteine of the motif (CX$_{2}$HX$_{23}$CX$_{4}$C) was changed to alanine (C → A) and the histidine residue was changed to asparagine (H → N). The PCR products containing these point mutations were inserted into pET-22b(+) and expressed in E. coli. The approximate masses of the wild-type or mutant N(1–564) and N(478–564) proteins were determined by SDS–PAGE and Coomassie brilliant blue staining (Fig. 5A). Then, similar molar amounts of these proteins were applied onto an SDS–polyacrylamide gel, transferred to a nitrocellulose membrane and probed with $^{65}$Zn$_{2}$$^{+}$. As shown in Fig. 4(C), no zinc-binding activity was detected in the truncated mutants N(1–393) and N(1–93) (Fig. 5B, lanes 5 and 6, respectively). When all four residues were changed in the N(478–564) protein, no zinc-binding activity was detected (Fig. 5B, compare lanes 7 and 8). Thus, the zinc-finger motif, located between amino acids 497 and 530, appears to be both necessary and sufficient for binding zinc.

Nevertheless, the zinc-binding activity was not abolished when these same four amino acid residues were changed in the full-length N protein N(1–564) quadruple mutant protein (Fig. 5B, lane 2). This unexpected result required an additional analysis, since the region immediately adjacent to the zinc-finger domain was not included in the series of truncated mutants used in Fig. 4. To verify if this region was also involved in zinc binding, we expressed two additional truncated proteins, designated N(1–536) quadruple mutant and N(1–502) double mutant, which also included the point mutations that abolished the zinc-binding activity of the N(478–564) quadruple mutant. Interestingly, both of these mutants bound zinc in vitro (Fig. 5B, lanes 3 and 4, respectively). These data indicate that amino acid residues between positions...
Fig. 5. Zinc blot analysis of N mutant proteins. Mutant proteins were analysed by electrophoresis on an SDS–polyacrylamide gradient gel (10–20%) and stained with Coomassie brilliant blue (A). A duplicate gel was transferred to a nitrocellulose membrane and probed with $^{65}$Zn$^{2+}$ (B). The relative zinc-binding abilities are summarized on the right. Cysteine-to-alanine mutations and histidine-to-asparagine mutations are indicated above the bars with dots (C). (D) N protein sequence of the region responsible for the zinc-binding activity: the amino acids that could be involved in zinc-coordination are indicated in bold and the amino acids involved in the zinc-finger domain are underlined.

394 and 502 can also be involved in zinc binding and explain the results observed with the N(1–564) quadruple mutant. In accordance with these results, this region has several potential coordination sites which could be involved in zinc coordination (Fig. 5D).

Discussion

The evidence presented here documents the zinc-binding capacity of the N protein of Junín virus and describes the metal ion-binding specificity. In experiments designed to characterize
the zinc-binding properties of the Z protein, Salvato & Shimomaye (1989) noted that the N protein of lymphocytic choriomeningitis virus (LCMV) binds zinc in vitro. In accordance with their results, we found a zinc-finger motif conserved at the carboxy-terminal region of all arenavirus N proteins (Parisi et al., 1996). The classical zinc-finger motif is characterized by two conserved cysteines and histidines which bind to a zinc ion in a tetrahedral coordination, thereby stabilizing a structure that comprises an antiparallel two-stranded β-sheet and an α-helix. The amino-terminal end of the α-helix is responsible for establishing DNA or RNA sequence-specific interactions (Nolte et al., 1998; McColl et al., 1999). Other classes of zinc fingers have been described which differ from the classical zinc finger in the metal ion-binding motif, the number of coordinated zinc ions, or the structure of the sequence-specific nucleic acid-binding region. The most frequent of these motifs is the so-called ‘RING finger’, characterized by the presence of two zinc-binding domains. In the general RING motif, each zinc atom is ligated tetrahedrally by either four cysteines, or three cysteines and a histidine. One of these RING fingers was described in the Z protein of LCMV (Lovering et al., 1993). As shown in Fig. 1, Junin virus nucleocapsid protein has the sequence CX2H29CX3C (positions 497, 500, 525 and 530), which resembles a classical zinc-binding motif and is well conserved among arenaviruses (Parisi et al., 1996). At this point, it is important to note that the N protein of Pichinde virus does not contain the cysteine-525; instead, there is a threonine residue which could potentially form a hydrogen bond with a water molecule that may also coordinate zinc (Coleman, 1992). In line with these considerations, the complete N protein of Junin virus expressed in bacteria displays zinc-binding activity in vitro (Fig. 2B). Assessment of the metal-binding specificity of the N protein was done by competition experiments. In these experiments, N protein was probed with 65Zn2+ in the presence or absence of unlabelled competitor metal ions. The competitor efficiency was Zn2+ ≡ Co2+ > Cu2+ > Mn2+ > Mg2+ (Fig. 3). These results are consistent with a specific zinc-binding activity and are similar to data obtained with the protein encoded by the vaccinia virus A2L gene (Keck et al., 1993). A priori, many amino acid residues in the N sequence could coordinate Zn2+ in addition to the CX2H29CX3C motif. To determine if the N protein region containing the putative zinc-finger was responsible for zinc binding we made a series of deletion mutants. The results indicate that the location of the zinc-binding activity is coincident with the putative zinc-finger region (Fig. 4). To establish if the zinc-binding activity was due to the putative zinc-finger motif, we made a series of point mutants. When the four amino acids predicted as critical in the zinc finger were mutated in the N(478–564) protein, the zinc-binding capacity was abolished (Fig. 5B, lane 8). However, when the same four mutations were introduced into the full-length N protein, only a slight decrease in zinc-binding capacity was apparent (Fig. 5B, lane 2). These results suggest an alternative zinc-coordination site (or sites) in the region immediately adjacent to or partially overlapping with the classical zinc-finger domain. This amino acid sequence comprises eight histidines and a cysteine, in addition to those that participate in the classical zinc finger, that could be involved in alternative zinc binding. Some of these amino acids are conserved in the Arenaviridae (Fig. 1). According to this, the truncated N(1–502) mutant containing the indicated point mutations should interact with zinc, as shown in Fig. 5, lane 4. This combined mutant comprises the N(1–393) protein region plus 109 amino acids (not represented in any of the previously analysed deletion mutants) and only half of the predicted zinc-finger domain in which both amino acids involved in the characterized zinc-finger activity were substituted. The smaller amount of zinc bound to mutant proteins N(1–536) and N(1–502), relative to full-length N, might be the reflection of a change in the structure and/or the stability of the alternative zinc-coordination site(s) due to the lack of the carboxy-terminal region (Fig. 5B, lanes 3 and 4 compared to lane 2). In support of these speculations several reports indicate that different degrees of change in the folding of a series of proteins are due to very limited alterations at their carboxy termini (Orellano et al., 1993; Clérico et al., 2000; Derman & Agard, 2000). The complete zinc-binding domain of N might adopt one of a number of different structures. It is also possible that the N protein has two independent binding sites: the classical zinc finger and the region immediately adjacent to it. However, our experiments do not rule out the possibility that these two binding sites exclude one another or bind zinc simultaneously, acting as a unique RING-finger-like structure. Further experiments should address the relevance of these residues in zinc binding and the significance of the zinc-binding activity in the N protein interactions that lead to the formation of nucleocapsids and/or to the transcriptional antitermination activity (Tortorici et al., 2000).

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References


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