DNA condensation by the nucleocapsid protein of HIV-1: a mechanism ensuring DNA protection

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ABSTRACT
The nucleocapsid (NC) protein NCp7 of the immunodeficiency virus type 1 is a small basic protein with two zinc finger motifs. NCp7 has key roles in virus replication and structure, which rely on its interactions with nucleic acids. Although most interactions involve RNAs, binding to the viral DNA is thought to be of importance to achieve protection of the DNA against cellular nucleases and its integration into the host genome. We investigated the interaction of NCp7 with plasmid DNA as a model system. The fluorescence probe YOYO-1 was used as the reporter. Binding of NCp7 to DNA caused DNA condensation, as inferred from the dramatic decrease in YOYO-1 fluorescence. Efficient condensation of DNA required the full length NCp7 with the zinc fingers. The fingerless peptide was less efficient in condensing DNA. Binding of both these NC peptides led to freezing of the segmental dynamics of DNA as revealed by anisotropy decay kinetics of YOYO-1. The truncated peptide NC(12–55) which retains the zinc fingers did not lead to DNA condensation despite its ability to bind and partially freeze the segmental motion of DNA. We propose that the histone-like property of NCp7 leading to DNA condensation contributes to viral DNA stability, \textit{in vivo}.

INTRODUCTION
The nucleocapsid (NC) protein, NCp7, of the immunodeficiency virus type 1 (HIV-1) is a small basic protein characterized by two copies of a highly conserved retroviral-type zinc finger CCHC motif, which binds Zn\textsuperscript{2+} with high affinity (1). About 2000 copies of NCp7 coat the genomic RNA in the infectious virus (2,3) and protect the genomic RNA against degradation by RNAases (4). Moreover, NCp7, as the mature protein or as the Pr55 polyprotein precursor, plays critical roles in several steps of the viral life cycle. Notably, NCp7 specifically interacts with the \(\Psi\) sequence of the genomic RNA, enabling its selective encapsidation into assembling particles (5–7). NCp7 is also critical for the efficient and complete proviral DNA synthesis by promoting both the initiation and the two strand transfer steps during reverse transcription (8–17). This is related to the NCp7 chaperone properties that enable rearrangement of a nucleic acid molecule into the most stable conformation by catalyzing the breakage and re-formation of base pairs (18–21).

The major functions of NCp7 rely on its specific and non-specific interactions with nucleic acids. Initially, based on the strong affinity of murine leukemia virus nucleocapsid protein NCp10 for single-stranded DNA-cellulose and its weak affinity for double-stranded DNA-cellulose, retroviral NCs have been considered as single-stranded nucleic acid-binding proteins (22,23). Later, this idea was challenged by the observation of significant affinity of NCp7 for various double-stranded DNA sequences (24–26). Moreover, addition of NCp7 to plasmid DNA at a ratio of one molecule per 4 bp caused a complete protection from restriction nuclease digestion, suggesting that DNA is coated by the NC protein (24). Both the binding extent and the level of protection were found to depend upon the presence of both the zinc fingers and the flanking clusters of basic amino acids (24,27–31). From these observations and recent \textit{in vivo} studies, it has been inferred that NCp7 binds to the newly synthesized viral DNA and protects it from cellular nucleases during reverse transcription and its transport to the nucleus up to the integration step (16,17,28,29).

With the aim of exploring the relationships between HIV-1 NCp7 and the proviral DNA we chose to characterize the structure and dynamics of complexes resulting from the interaction of NCp7 with a double-stranded plasmid DNA, taken as a model system. By using the bis-intercalating fluorescent dye YOYO-1 (32), we analyzed DNA condensation and dynamics, as well as the exposure of base pairs to solvent. We find that binding of NCp7 to DNA resulted in DNA condensation in a manner similar to that brought out by...
gene delivery vectors, freezing of the nanosecond segmental dynamics of DNA and protection of base pairs from solvent. Moreover, using three NCp7 derivatives (Fig. 1A), we explored the comparative influence of the zinc fingers and the flanking basic domains on the structural features of the complex.

MATERIALS AND METHODS

Materials

The plasmid DNA pCMV-luc (5.2 kb) was propagated and purified as described (35). Branched chain polyethyleneimine (PEI) (25 kDa) was a gift from Professor J.-P. Behr, Illkirch, France. YOYO-1 (491/509) was obtained from Molecular Probes Inc. NCp7 and the related peptides were synthesized by solid-phase chemistry as described (36). Purity of the peptides analyzed by mass spectrometry was >98%. Peptide concentration was determined on a Cary 400 spectrophotometer, using an extinction coefficient at 280 nm of 12 700, 7000 and 5700 M⁻¹ cm⁻¹ for NC(1–72), NC(1–72)dd and NC(12–55), respectively. To protect the highly oxidizable Cys residues of NC(1–72) and NC(12–55), the peptides were stored lyophilized in their zinc-bound forms.

Samples

Plasmid DNA–YOYO-1 complexes were made by mixing equal volumes of solutions of DNA and YOYO-1 in 100 mM
NaCl, 20 mM HEPES, pH 7.4. The mixed solution was incubated at room temperature for at least 5 h before use. This procedure ensured uniform distribution of YOYO-1, which tightly binds to DNA. Complexes of plasmid DNA–YOYO-1 with NCP7 and its derivatives were formed by adding the peptide to a solution of DNA–YOYO-1 complex. In the case of equilibrium measurements the samples were kept for at least 90 min for completion of condensation.

**Spectroscopic measurements**

An SLM 48000 spectrofluorimeter was used for steady-state fluorescence measurements.

Time-resolved fluorescence intensity and anisotropy measurements on DNA–YOYO-1–NCP7 complexes were performed by using the frequency doubled output of a Ti-Sapphire laser (Spectra Physics). The excitation wavelength was 480 nm and the emission was collected at 515 nm by using a single photon counting microchannel plate photomultiplier (Hamamatsu R3809U). Other details are given elsewhere (37).

The resolution of the time-correlated single photon counting set-up was 25.5 ps and the width of the instrument response function was ~40 ps.

Fluorescence intensity decays obtained at the magic angle were deconvoluted with the instrument response function and analyzed as a sum of exponentials:

\[
I(t) = \Sigma \alpha_i \exp(-t/\tau_i)
\]

where \( I(t) \) is the fluorescence intensity collected at the magic angle at time \( t \) and \( \alpha_i \) is the amplitude of the \( i \)th lifetime \( \tau_i \) such that \( \Sigma \alpha_i = 1 \).

Time-resolved fluorescence anisotropy decays were analyzed by the following equations

\[
I_{//}(t) = I(t) \left[ 1 + 2r(t) \right]/3
\]

\[
I_{\perp}(t) = I(t) \left[ 1 - r(t) \right]/3
\]

\[
r(t) = \left[ I_{//}(t) - I_{\perp}(t) \right]/\left[ I_{//}(t) + 2I_{\perp}(t) \right] = r_0 \Sigma \beta_i \exp(-t/\phi_i)
\]

where \( r_0 \) is the initial anisotropy and \( \beta_i \) is the amplitude of the \( i \)th rotational correlation time \( \phi_i \) such that \( \Sigma \beta_i = 1 \). \( I_{//} \) and \( I_{\perp} \) are the intensities collected at emission polarizations parallel and perpendicular, respectively, to the polarization axis of the excitation beam. While analyzing the kinetics of anisotropy decay using equations 1 and 2, values of \( \alpha_i \) and \( \tau_i \) [obtained from the analysis of \( I(t) \)] were kept fixed in order to reduce the number of floating parameters. This procedure results in better estimates of anisotropy decay parameters.

**RESULTS**

**Condensation of DNA by NCP7**

Condensation of plasmid DNA by NCP7 was monitored by following the fluorescence of the bis-intercalating fluorophore YOYO-1. Indeed, we have recently shown that YOYO-1 is a simple and very sensitive probe for monitoring the condensation of DNA by a variety of condensing agents (32). The quantum yield of YOYO-1 increases by several orders of magnitude during the transition from aqueous solution to its intercalating site in double-stranded DNA (32). Subsequently, when the double-stranded DNA undergoes the transition from extended (either supercoiled or relaxed) conformation to a condensed form catalyzed by a condensing agent, the quantum yield of bound YOYO-1 undergoes a drastic reduction (32). Furthermore, we had shown that this reduction is a result of a combination of formation H-dimers of YOYO-1 and fluorescence resonance energy transfer between these dimers and YOYO-1 monomer (32).

Figure 1B shows the effect of NCP7 on the fluorescence emission spectra of YOYO-1 intercalated into the DNA. In these samples the ratio of YOYO-1 to DNA phosphate was kept at 1:50 in order to visualize the condensation effect (32). In analogy to DNA-condensing agents such as PEI or cationic detergents (32), NCP7 induced a dramatic reduction in the quantum yield of YOYO-1. Figure 2 shows the titration of the fluorescence intensity with NCP7 and its derivatives at two different DNA concentrations. A progressive decrease of YOYO-1 fluorescence was observed with increasing NC(1–72) concentrations. This decrease levelled off at a plateau value that slightly depends on the oligonucleotide (nt)
Table 1. Parameters associated with decay of fluorescence intensity and anisotropy

<table>
<thead>
<tr>
<th>Sample a</th>
<th>Fluorescence lifetimes, ns (amplitude)b</th>
<th>Rotational correlation times, ns (amplitude)c</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(\tau_1 (\alpha_1))</td>
<td>(\tau_2 (\alpha_2))</td>
</tr>
<tr>
<td>1. DNA–YOYO-1 uncondensed</td>
<td>5.1 (0.50)</td>
<td>2.35 (0.50)</td>
</tr>
<tr>
<td>2. DNA–YOYO-1 + NC(1–72)</td>
<td>4.65 (0.43)</td>
<td>2.05 (0.57)</td>
</tr>
<tr>
<td>3. DNA–YOYO-1 + NC(1–72)dd</td>
<td>5.25 (0.69)</td>
<td>1.90 (0.31)</td>
</tr>
<tr>
<td>4. DNA–YOYO-1 + NC(12–55)</td>
<td>5.20 (0.42)</td>
<td>2.46 (0.58)</td>
</tr>
</tbody>
</table>

aThe concentration of plasmid DNA was 10 \(\mu\)M of phosphate and that of YOYO-1 was 5 nM for all the samples. The concentration of NC peptides was 4 \(\mu\)M. The medium was 20 mM HEPES, 100 mM NaCl at pH 7.4.
bThe errors associated with various parameters were as follows: \(\alpha_1 \pm 0.02\), \(\alpha_2 \pm 0.02\), \(\phi_1 \pm 0.30\), \(\phi_2 \pm 4.0\), \(\phi_3 \pm 0.03\), \(\beta_1 \pm 0.03\), \(\beta_2 \pm 0.03\) and \(\beta_3 \pm 0.03\). The value of reduced \(\chi^2\) was in the range of 1.0–1.3 in all the cases. Also, the pattern of residuals was found to be random for all the fits.cInitial anisotropy \(r_0\) was fixed at 0.31 \(\pm 0.01\), the value obtained for YOYO-1 in glycerol, during the analysis.dMean lifetime \(\tau_{\text{mean}} = \Sigma \alpha_i \tau_i\).eSince the uncertainty in the value of \(\phi_1\) was too large only the upper limit is given.

Figure 3. Titration of fluorescence intensity of DNA-bound YOYO-1 with NC(1–72) at various DNA concentrations. DNA phosphate concentrations were 0.1 \(\mu\)M (squares); 0.2 \(\mu\)M (circles); 0.4 \(\mu\)M (triangles) and 0.8\(\mu\)M (inverted triangles). Dye:phosphate ratio was 1:50.

By further investigating the dependence of the extent of DNA condensation on the concentrations of NC(1–72) and DNA-nt (Fig. 3), we found that the profile of condensation largely depended on the DNA to NC molar ratio. Nearly 80% of the total decrease in fluorescence intensity occurred at [NC(1–72)]/[DNA-nt] approximately 0.2 at all the concentrations of DNA-nt (0.1–0.8 \(\mu\)M) used. This would suggest an apparent binding site of \(2–3\) bp per NCp7 molecule, which is significantly different from the binding site size of \(9\) bp estimated earlier (26). However, the slight deviation of the titration curves from each other (Fig. 3) could be interpreted as an indication of a deviation from stoichiometric binding. Such non-stoichiometric binding is expected to lead to the presence of unbound NC indicating that the apparent dissociation constant of NC(1–72) for plasmid DNA is in the micromolar range. However, it should be borne in mind that the observed decrease in fluorescence intensity (Figs 1–3) arose from condensation of DNA and not from a direct effect of protein binding as shown by the following observation: quenching was not observed when the YOYO-1 to DNA phosphate ratio was reduced from 1:50 to 1:2000 (see for example, the mean lifetime in Table 1). These observations are similar to those seen earlier with either PEI or CTAB (32). Furthermore, direct support for condensation of DNA by the binding of NC(1–72) comes from the following observations also: (i) binding of NC to DNA leads to the formation of high molecular mass nucleoprotein complexes, which can be pelleted by centrifugation (33); and (ii) dark field electron micrographs obtained in samples having NC and DNA shows the presence of nucleoprotein complexes as dense aggregates having diameter in the range of a few hundred nanometers (34). Thus, these observations when combined with our present work indicate that YOYO-1 fluorescence is a convenient spectroscopic marker for DNA condensation.

By further investigating the dependence of the extent of DNA condensation on the concentrations of NC(1–72) and DNA-nt (Fig. 3), we found that the profile of condensation largely depended on the DNA to NC molar ratio. Nearly 80% of the total decrease in fluorescence intensity occurred at [NC(1–72)]/[DNA-nt] approximately 0.2 at all the concentrations of DNA-nt (0.1–0.8 \(\mu\)M) used. This would suggest an apparent binding site of \(2–3\) bp per NCp7 molecule, which is significantly different from the binding site size of \(9\) bp estimated earlier (26). However, the slight deviation of the titration curves from each other (Fig. 3) could be interpreted as an indication of a deviation from stoichiometric binding. Such non-stoichiometric binding is expected to lead to the presence of unbound NC indicating that the apparent dissociation constant of NC(1–72) for plasmid DNA is in the micromolar range. However, it should be borne in mind that the observed decrease in fluorescence intensity (Figs 1–3) arose from condensation of DNA (32) subsequent to the binding of NC and not due to the binding of NC to DNA per se (see ‘Discussion’). Hence, data presented in Figures 1–3 cannot be rigorously used for the estimation of binding site size and binding constant.

Figure 4 shows a typical time-dependence of the condensation process induced by either NC(1–72) or the fingerless peptide NC(1–72)dd. The time dependence shows that condensation is a two-step process: one that occurred within the mixing time of a few seconds and another with a half time of \(~5–20\) min. The rate of the slow phase and the extent of contribution of the fast phase were higher in the case of NC(1–72) when compared with NC(1–72)dd. Furthermore, the rate of the slow phase showed only a small dependence on the concentration of DNA (compare Fig. 4A and B), suggesting that the slow phase may not represent a bimolecular process of binding of NC to the DNA. A bimolecular process would have resulted in a 5-fold decrease in the rate of the slow process.
when the concentration of DNA was decreased by a factor of 5. In contrast, the decrease in the apparent rate was significantly <5-fold. However, we cannot rule out, during the slow phase, a more complex process, which involves coming together of several complexes of DNA±NC in forming a large nucleoprotein structure.

Base solvent-accessibility and dynamics of NCp7±DNA complexes

In order to check whether the bases get exposed to solvent in the condensed form of DNA, quenching of YOYO-1 fluorescence, intercalated into DNA bases by acrylamide, was monitored (Fig. 5). Solvent exposure of DNA bases could be estimated from the level of quenching by acrylamide (32). Although the accessibility of intercalated YOYO-1 to acrylamide was very low in both the extended and condensed forms of DNA (as shown by the very low levels of quenching), a small but reproducible increase of base exposure to solvent (as observed from the increase in slope) was observed in the presence of NCp7 (Fig. 5). In contrast to NCp7, condensation by PEI, a cross-linked polymer used in gene delivery systems, led to a much larger exposure of bases (Fig. 5) as seen earlier (32). In these experiments the ratio of YOYO-1 to oligonucleotides was kept very low (1:2000) to ensure that YOYO-1 fluorescence mainly reflected the changes in the property of individual YOYO-1 molecule rather than reflecting condensation-induced interaction among bound YOYO-1 molecules (32) seen at a high ratio such as 1:50 (Figs 1–4). Under the former condition, fluorescence lifetime ($\tau$) did not change significantly on condensation (see later) and hence any change in slope ($= K_{SV} = \tau K_Q$) of plots in Figure 5 reflects mainly change in accessibility (represented by the quenching constant, $K_Q$) of YOYO-1 to acrylamide (32).

Figure 6 shows the rotational dynamics of bound YOYO-1 as seen from the kinetics of decay of fluorescence anisotropy. These decay kinetics were fitted satisfactorily to a sum (equation 3) of three exponentials resulting in three correlation times (Table 1) in all the cases. The very short rotational correlation time ($\tau_1 < 0.1$ ns) arises due to the necessity of fixing the value of initial anisotropy ($r_0$) at 0.31 ± 0.01, which was obtained for YOYO-1 in a viscous solvent. It is likely that $\tau_1$ represents the internal fast rotation of YOYO-1 and $\tau_2$ and $\tau_3$ originate from segmental dynamics of DNA backbone (32). It can be seen that the observable part of anisotropy decay kinetics is largely dominated by $\tau_3$. Hence the value of $\tau_3$ could be used in characterizing the changes in motional dynamics of DNA backbone brought about by the binding of NCp7 and its derivatives. In the uncondensed form of DNA, its backbone is highly dynamic as shown by the value of $\tau_3 \approx 11$ ns similar to our earlier observations (32). Binding of NCp7 and the related peptides resulted in dramatic changes in the anisotropy decay profile reflecting changes in the motional dynamics of the DNA backbone (Fig. 6). Both NC(1–72) and the fingerless NC(1–72)$\text{dd}$ caused significant decrease in the rate of anisotropy decay as shown by significant increase in the value of $\tau_3 (> 100$ ns, this limit being the result of the limited time window offered by the fluorescence lifetime of YOYO-1) (Fig. 6 and Table 1). This could be interpreted as significant rigidification of DNA upon binding of either NC(1–72) or NC(1–72)$\text{dd}$. However, binding of the truncated peptide NC(12–55) retarded the decay of anisotropy (and hence DNA flexibility) to a lesser extent as shown by $\tau_3 \approx 33$ ns.
DNA can be effectively condensed upon binding of HIV-1 condensation probe YOYO-1 (32) that double-stranded DISCUSSION peptides at these low levels of YOYO-1 (1:2000). The ratio of YOYO-1 to DNA-nt was quite low in these samples for reasons mentioned above. Fluorescence intensity decay curves were satisfactorily fitted to a sum of two exponentials in all the bound forms (Fig. 2). The ratio of YOYO-1 to DNA-nt was quite low (1:2000) in these samples for reasons mentioned above. Fluorescence intensity decay curves were satisfactorily fitted to a sum of two exponentials in all the bound forms (Table 1) similar to the behavior in uncondensed DNA (32). Furthermore, intensity decay kinetics were largely insensitive to the binding of NC peptides although some minor changes were seen (Table 1). Thus, the effect of binding NC peptides was seen mainly in anisotropy decay kinetics rather than in their fluorescence lifetimes. Concomitantly, fluorescence intensity also did not show any change on binding of NC peptides at these low levels of YOYO-1 (1:2000).

**DISCUSSION**

In this work we have shown by using the new DNA condensation probe YOYO-1 (32) that double-stranded DNA can be effectively condensed upon binding of HIV-1 NCp7 and its fingerless (without Zn^{2+}) mutant. Our results (Fig. 2) point out that efficient condensation of DNA by NC requires the complete NC with the zinc fingers. The requirement for a complete NC is probably due to the fact that basic amino acids, expected to be a major driving force for condensation (through electrostatic interactions with the negatively charged DNA phosphate groups) are present fairly uniformly over the entire NC molecule (Fig. 1A). The requirement for the zinc fingers may be considered as counterintuitive especially when one notes that basic polypeptides and polycations are efficient condensing agents (32,38,39). The zinc finger-induced enhancement of condensation probably originates from specific interaction(s) between zinc finger side chain(s) and DNA bases. Stacking interaction of Trp37 side chain with bases (notably G) in DNA or RNA sequences as evidenced by NMR studies (40–42) was found to strongly contribute to the binding energy (26,43–45). Furthermore, the observation that zinc finger binding caused DNA bending (46) suggests that unique interactions are taking place between the zinc fingers and the DNA.

Moreover, the observation that NC(12–55) was unable to elicit the condensation signal (Fig. 2) shows that the basic domain of NCp7 is required for condensation. Since previous studies have shown only limited differences between the binding constants of NC(12–55) and NCp7 at 0.1 M NaCl concentration (43,44), the inability of NC(12–55) to condense DNA is probably unrelated to its binding energy. In addition, clear evidence of NC(12–55) binding to DNA came from the significant changes in the anisotropy decay parameters that followed addition of NC(12–55) to DNA (Fig. 2D and Table 1). The ability of the various NC derivatives to condense DNA correlates well with their ability to protect DNA from exonuclease digestion. While NC(1–72) was found to achieve a good protection toward nucleases (24), only a partial protection was observed with NC(1–72)_{del} (24) or NC mutants with mutated finger motifs (16,17,28,29) and almost no protection was afforded by NC(12–55) (Lener and Darlix, unpublished data). Moreover, in line with the hypothesis of Carteau et al. (47), NC-induced condensation of DNA may play a critical role in promoting proviral DNA integration, and notably the coupled joining of the two ends of the viral DNA at precise positions in the host cell DNA. In fact, condensation of duplex DNA by NC might facilitate association of the two proviral DNA ends and facilitate coupled integration. Tight packing of DNA has been proposed to assist end ligation by repair enzymes in radiation-resistant bacteria (48) supporting our model of NC involvement in proviral DNA integration.

Integration of the proviral DNA into the host genome may further require NC chaperone activity and notably, its nucleic acid destabilizing property. A limited destabilization of plasmid DNA by NC was evidenced by the slight increased level of accessibility of intercalated YOYO to acrylamide in NC(1–72)–DNA complexes (as shown by the increase in slope and K_{eq}, Fig. 5) when compared with bare DNA. This destabilizing property of NCp7 on DNA is in line with previous observations with either short (25) or long duplexes (49) or hairpin loops (50–52). Both the nucleic acid condensing and destabilizing properties of NC and their consequences on the protection of the viral DNA during its nuclear import and subsequent integration may well explain why HIV-1 with mutations in the zinc fingers or in the N-terminal basic domain of NC are replication defective (15–17,28,30).

How compact is the DNA condensed by NCp7? The observation that the extent of reduction in the fluorescence intensity of YOYO-1 (Figs 2 and 3) on condensation by NCp7 is only ~70%, which is less than the 90% reduction observed with PEI (Fig. 2) (32), suggests that the compactness of the DNA condensed by NCp7 is comparatively less. Furthermore, condensation by either NCp7 or NC(1–72)_{del} results in freezing of the nanosecond segmental dynamics of DNA.
(Fig. 6 and Table 1). The level of rigidification of the DNA backbone was higher in the cases of condensation-competent peptides NC(1–72) and NC(1–72), when compared with the incompetent analog NC(12–55). Further, the level of rigidification was similar to that observed with PEI (32) indicating this as one of the common features of DNA condensed by cationic polymers.

Interestingly, the condensation activity of NCp7 and the related peptides correlates also with the ability to form high molecular mass complexes with nucleic acids (53,54). For example the inability of NC(12–55) to give the condensation signal is in line with its inability to form high molecular mass complexes with polyA (53) and tRNA<sup>35</sup> (55) and various genomic RNA fragments (data not shown). Similarly, the reduced rate of DNA condensation by NC(1–72) [when compared with NC(1–72)] (Fig. 4) agrees with the reduced rate of formation of high molecular mass complexes (53). This suggests that the processes of condensation and formation of high molecular mass complexes are correlated with each other and probably condensation precedes formation of high molecular mass complexes. This conclusion is supported by the observation that the time course of the condensation process (half time ~20 min, Fig. 4) is significantly shorter than the time course of high molecular mass complex formation (53), which takes several hundred minutes, although the two processes were monitored with different nucleic acids. The former process could be an intramolecular one wherein the plasmid DNA, on binding to NCp7, collapses into a globule. The inability of the truncated form NC(12–55) to either condense the DNA or form high molecular mass complexes (in spite of its ability to bind) is probably due to the absence of positively charged N- and C-terminal domains. Thus, we infer that efficient formation of compact nucleoprotein globules requires both the basic residues of the N- and C-terminal regions of NCp7 and the DNA bending ability offered by the zinc finger domain. The latter suggestion requires further investigation.

In conclusion, the DNA condensing properties and the resulting DNA rigidification evidenced in this study may confer to NCp7, a histone-like activity that is in line with its critical role of NCp7 in protecting the neosynthesized DNA resulting DNA rigidiﬁcation evidenced in this study may confer to NCp7, a histone-like activity that is in line with its critical role of NCp7 in protecting the neosynthesized DNA. The inability of the truncated form NC(12–55) to either condense the DNA or form high molecular mass complexes (in spite of its ability to bind) is probably due to the absence of positively charged N- and C-terminal domains. Thus, we infer that efficient formation of compact nucleoprotein globules requires both the basic residues of the N- and C-terminal regions of NCp7 and the DNA bending ability offered by the zinc finger domain. The latter suggestion requires further investigation.

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