The intrinsically disordered C-terminal domain of the measles virus nucleoprotein interacts with the C-terminal domain of the phosphoprotein via two distinct sites and remains predominantly unfolded

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Abstract

Measles virus is a negative-sense, single-stranded RNA virus within the Mononegavirales order, which includes several human pathogens, including rabies, Ebola, Nipah, and Hendra viruses. The measles virus nucleoprotein consists of a structured N-terminal domain, and of an intrinsically disordered C-terminal domain, NTAIL (aa 401–525), which undergoes induced folding in the presence of the C-terminal domain (XD, aa 459–507) of the viral phosphoprotein. Within NTAIL, an α-helical molecular recognition element (α-MoRE, aa 488–499) involved in binding to P and in induced folding was identified and then observed in the crystal structure of XD. Using small-angle X-ray scattering, we have derived a low-resolution structural model of the complex between XD and NTAIL, which shows that most of NTAIL remains disordered in the complex despite P-induced folding within the α-MoRE. The model consists of an extended shape accommodating the multiple conformations adopted by the disordered N-terminal region of NTAIL, and of a bulky globular region, corresponding to XD and to the C terminus of NTAIL (aa 486–525). Using surface plasmon resonance, circular dichroism, fluorescence spectroscopy, and heteronuclear magnetic resonance, we show that NTAIL has an additional site (aa 517–525) involved in binding to XD but not in the unstructured-to-structured transition. This work provides evidence that intrinsically disordered domains can establish complex interactions with their partners, and can contact them through multiple sites that do not all necessarily gain regular secondary structure.

Keywords: measles virus; nucleoprotein; phosphoprotein; intrinsic disorder; induced folding; NMR; CD; SAXS

Measles virus (MV) is an enveloped RNA virus within the Morbillivirus genus of the Paramyxoviridae family. Its nonsegmented, negative-sense, single-stranded RNA genome is encapsidated by the viral nucleoprotein (N) within a helical nucleocapsid. This N–RNA complex is used as a template for both transcription and replication. These latter activities are carried out by the viral polymerase complex, which consists of two components, the large protein (L) and the phosphoprotein (P) (for review, see Lamb and Kolakofsky 2001).
Nucleoproteins of *Paramyxoviridae* are divided into two regions: a structured N-terminal moiety, *N*<sub>CORE</sub> (aa 1–400 in MV), which contains all the regions necessary for self-assembly and RNA-binding (Buchholz et al. 1993; Curran et al. 1993; Bankamp et al. 1996; Liston et al. 1997; Myers et al. 1997b, 1999; Karlin et al. 2002a; Kingston et al. 2004b), and a C-terminal domain, *N*<sub>TAIL</sub> (aa 401–525 in MV). *N*<sub>TAIL</sub> is intrinsically unstructured (i.e., it lacks any stable secondary and tertiary structure in physiological conditions) (Longhi et al. 2003) and is exposed at the surface of the viral nucleocapsid (Heggeness et al. 1980, 1981). The presence of a flexible region protruding from the viral nucleocapsid allows the establishment of interactions with numerous different viral partners and with several cellular proteins (Moyer et al. 1990; De and Banerjee 1999; tenOever et al. 2002; Zhang et al. 2002; Laine et al. 2003). In *Morbilliviruses* and *Respiroviruses*, *N*<sub>TAIL</sub> is responsible for binding to *P* (Curran et al. 1993; Harty and Palese 1995; Bankamp et al. 1996; Liston et al. 1997; Longhi et al. 2003; Kingston et al. 2004b), to the polymerase complex L–P, and to the matrix protein (Coronel et al. 2001). Beyond viral partners, *N*<sub>TAIL</sub> also interacts with several cellular proteins, including the interferon regulatory factor 3 (tenOever et al. 2002) and the heat-shock protein Hsp72 (Zhang et al. 2002). Moreover, *N*<sub>TAIL</sub> within viral nucleocapsids released from infected cells also binds to a yet unidentified protein receptor expressed at the surface of human thymic epithelial cells (Laine et al. 2003, 2005).

The *P* protein of *Paramyxovirinae* plays multiple roles in both transcription and replication: It is an essential subunit of the viral polymerase complex and acts as a bridge between the nucleocapsid template (*N*<sup>NUC</sup>) and the polymerase complex L–P. *P* is a modular protein, consisting of an intrinsically unstructured N-terminal moiety (PNT) (Karlin et al. 2002b, 2003), and of a well conserved C-terminal moiety (PCT) which contains all the regions required for transcription (Curran 1996). *Paramyxovirinae* PCT have a modular organization, consisting of alternating disordered and structured regions (Karlin et al. 2003). In particular, they possess a coiled-coil domain (referred to as PMD, for P multi-merization domain) responsible for both oligomerization and binding to L (Smallwood et al. 1994; Liston et al. 1995), and a C-terminal globular region (referred to as XD), that is involved in binding to both monomeric and assembled forms of N (Curran et al. 1994, 1995a,b; Harty and Palese 1995; Tuckis et al. 2002; Johansson et al. 2003; Kingston et al. 2004b). Figure 1 shows a schematic representation of the MV *N*<sup>NUC</sup>–*P* complex, which highlights the role of *N*<sub>TAIL</sub> in the recruitment of *P* via the binding to XD.

We have previously reported the crystal structure of the C-terminal globular domain of MV *P* (XD, aa 459–507): It consists of a monomeric protein composed of three α-helices, forming an anti-parallel three-helix bundle (Johansson et al. 2003). We have also shown that *N*<sub>TAIL</sub> undergoes an induced folding in the presence of XD and that this unstructured-to-structured transition implies a gain of α-helicity (Johansson et al. 2003). Using a combination of computational and biochemical approaches, we have identified within *N*<sub>TAIL</sub> an α-helical molecular recognition element (α-MoRE, aa 488–499) involved in binding to *P* and in the induced folding of *N*<sub>TAIL</sub> (Bourhis et al. 2004). The α-MoRE has been modeled in the crystal structure of XD, in the long hydrophobic cleft delimited by helices α2 and α3 (Johansson et al. 2003). In this model, the contact of *N*<sub>TAIL</sub> with a hydrophobic patch at the surface of XD would be the driving force in the induced folding of the α-MoRE of *N*<sub>TAIL</sub> by burying apolar residues at the protein–protein interface. Very recently, Kingston et al. (2004a) have reported the crystal structure of a chimeric protein composed of MV XD and a peptide corresponding to residues 486–505 of MV N. The structure of the complex is a four-helix bundle in which the α-helix of N is bound in the reverse orientation with respect to the model proposed by Johansson et al. (2003). Using NMR and crystallographic studies, Kingston et al. (2004a) deduced that the induced folding of *N*<sub>TAIL</sub> is restricted to only 18 residues (out of 125), although the analysis was restricted to the N region encompassing residues 477–505.
In this paper we examine this localized induced folding event in the context of the entire NTAIL domain. A low-resolution structural model of the complex between XD and NTAIL shows the presence of a bulky globular region and of an extended and elongated shape. The model shows that the N-terminal region of NTAIL (residues 401–488) remains predominantly unfolded, and the 489–525 region is packed against XD, thus suggesting that beyond the α-MoRE the C terminus may play a role in the interaction with XD. Indeed, we present several lines of experimental evidence confirming that beyond the region encompassing the α-MoRE, an additional NTAIL region encompassing residues 517–525 also contributes to binding to XD, although without undergoing any gain of regular secondary structure.

Results

Small angle X-ray scattering (SAXS) studies of the NTAIL–XD complex

Small angle X-ray scattering (SAXS) is a valuable technique for the study of flexible, low compactness macromolecules in solution, which has already been successfully used to characterize NTAIL (Longhi et al. 2003). Therefore, we used SAXS to study the NTAIL–XD complex. To this endeavor, we cloned, expressed, and purified from the Escherichia coli an N-terminally histidine-tagged form of NTAIL (i.e., NTAILHN) (data not shown). The identity of the recombinant products was confirmed by immunoprecipitation (IP) studies using anti-N Cl25 and anti-hexahistidine tag monoclonal antibodies (mAbs) (data not shown).

Beforehand, we checked whether XD possesses the same structure in solution as in the crystal. SAXS experiments performed on XD showed that it has a globular shape with a radius of gyration (Rg), extrapolated at a nil concentration, of 12.1 ± 0.8 Å and a maximum diameter Dmax of 41 ± 1 Å (data not shown). Comparison with the crystal structure using the program CRYSOL (Svergun et al. 1995) indicates that the scattering profile calculated for the crystal structure (PDB code 1OKS) is identical to the experimental one, fitting the data with a χ² of 1.4 and giving a theoretical Rg of 12.3 Å (data not shown). These results suggest that the overall conformation of XD in solution is similar to that observed in the crystal.

The scattering profile of the NTAILHN–XD complex was obtained as described in Materials and Methods. Analysis of the curve in the low q region with the Guinier approximation gave an Rg of 32.7 ± 0.7 Å. The molecular mass (MM) calculated from the forward scattering intensity I(0) is 23.5 ± 2 kDa, in agreement with the value expected for a 1:1 stoichiometric complex (21.3 kDa), thus suggesting that the complex did form in solution. The high value of Rg indicated that the overall structure of the NTAILHN–XD complex is not compact. The distances distribution function inferred from the scattering curve of the NTAILHN–XD complex exhibits a maximum at 20 Å, with a shoulder at about 30 Å and a long tail up to 146 Å, typical of an elongated object (see Fig. 2A). The bump most probably corresponds to the intramolecular distances within the globular portion of the complex (see below), while the tail indicates that NTAIL possesses regions with an extended conformation.

The overall envelope of the complex was restored ab initio from its scattering profile using the program GASBOR (Svergun et al. 2001). Several independent runs yielded different shapes with recurrent features: they were all elongated, with a globular cluster of always the same size at one extremity and an elongated protuberance with varying bends and cross-sections. The quality of the fit to the experimental data was similar in all cases, with a χ² of 1.3 to 1.6. The globular part most probably corresponds to XD packed against the folded region of NTAIL, while the outgrowth corresponds to unfolded regions of the latter. The crystal structure of the XD–NTAIL486–505 complex (PDB code 1T6O), which encompasses the α-MoRE (residues 488–499), was inserted in the shape using the program SUPCOMB (Kozin and Svergun 2001) (Fig. 2B). Interestingly, only one protuberance was observed, corresponding to the N-terminal area of NTAIL, and no other protuberance was restored from the shape determination in the opposite region of the globular cluster, thus suggesting that the C-terminal region of NTAIL (aa 506–525), although not visible per se, is packed inside the bulky portion of the complex (Fig. 2B). In order to get further insights into the conformation that the regions of NTAIL encompassing residues 401–485 and 506–525 adopt in the complex with XD, we used the program package CREDO, which is an extension of GASBOR and calculates the position of missing loops in crystal structures. The results obtained with several independent runs using the crystal structure of XD–NTAIL486–505 as the template, revealed at one side a peptide chain in an extended conformation and at the other side a chain packed against the bulky portion of the XD–NTAIL486–505 complex. The extended conformation, which protrudes from the globular cluster of the complex and points to the solvent, corresponds to the 92-residue-long N-terminal region of the NTAILHN construct. In contrast, the more compact peptide chain, corresponding to the 20-residue-long C-terminal region of NTAIL (aa 506–525), always packs against the XD–NTAIL486–505 complex but at varying distances and positions. An example of one of the calculated conformations is shown in Figure 2C. The experimental data were
fitted by these models with similar $\chi^2$ of about 0.8–1.0 (see Fig. 2C, inset). The various possible conformations provided by the program CREDO together with the extended structural properties of the complex (as indicated by the distance distribution function, the $R_g$ and the restored shape, with respect to the number of amino acids of the complex) are typical of disordered polypeptide chains lacking a stable regular structure. Accordingly, the structure shown in Figure 2C represents only one possible conformation found by CREDO among many others. Indeed, CREDO restores only one conformation per run, and therefore cannot account for multiple conformations due to disorder when fitting the data. Therefore, although a unique conformation of the N$_{TAIL}$ region encompassing residues 506–525 cannot be derived, all the solutions provided by CREDO consistently revealed a packing of the C-terminal region of N$_{TAIL}$ against XD. The recurrence of these packed conformations suggests that the C-terminal region of N$_{TAIL}$ interacts with XD.

Moreover, we checked the influence of the Box3 conformation on the scattering curve by calculating the scattering profile of pseudomodels of the complex. In these pseudomodels, the N-terminal region of N$_{TAIL}$ is in the same conformation as in the credo model shown in Figure 2C. The only difference concerns the orientation of Box3, which points out of the complex, in different solvent-exposed conformations. Noteworthy, the theoretical scattering curves calculated from these pseudomodels using CRY SOL (Svergun et al. 1995) poorly fit with the experimental curve, with an average $\chi^2$ of $\sim$7.0 (data not shown). This confirms that the C terminus of N$_{TAIL}$ does contribute to the observed scattering profile, thus attesting the reliability of the model provided by CREDO.

Cloning, expression, and purification of N$_{TAIL}$ deletion constructs

In order to further explore the possible contribution of N$_{TAIL}$ regions other than the $\alpha$-MoRE to XD binding, we have looked at the regions of homology conserved amongst members of the Morbillivirus genus (Di a llo et al. 1994). These regions of homology are herein referred to as Box1, Box2, and Box3 and span N residues 401–420, 489–506, and 517–525, respectively (see Fig. 3A), with the $\alpha$-MoRE being located within Box2. We have then designed three deletion constructs bearing different combinations of these homology boxes (Fig. 3A).

The gene fragments encoding the different N$_{TAIL}$ deletion proteins were cloned into the pDest14 vector (Invitrogen) to yield N-terminally histidine-tagged and C-terminally flag-tagged recombinant products, the expression of which is under the control of the T7 promoter. In all cases, most recombinant protein was recovered from the soluble fraction of bacterial lysates (Fig. 3B, lanes SN). The N$_{TAIL}$ deletion proteins were purified to
The number of different conformations of the N<sub>Tail</sub> deletion proteins is limited, as indicated by the sharpness of the peaks observed in gel filtration (data not shown). As expected for intrinsically disordered protein subdomains, the Stokes radius (R<sub>S</sub>) values, as inferred by gel filtration (27 ± 3 Å, 22 ± 3 Å, and 27 ± 3 Å for N<sub>Tail3</sub>, N<sub>Tail2,3</sub>, and N<sub>Tail1</sub>, respectively), are consistent with extended conformations (see Materials and Methods). Thus, these deletion proteins share similar hydrodynamic properties with full-length N<sub>Tail</sub>- all possessing an elongated shape.

**Surface plasmon resonance studies**

In order to directly measure the contribution of the different N<sub>Tail</sub> boxes to binding, we have studied binding reactions between XD and N<sub>Tail</sub> deletion proteins. Changes in surface plasmon resonance were monitored in real time as the N<sub>Tail</sub> proteins passed over sensor chips to which XD was covalently coupled. This analytical approach is ideally suited to study reversible, low-affinity protein–protein interactions that typify interactions involving intrinsically disordered proteins (Wright and Dyson 1999; Dunker and Obradovic 2001; Dunker et al. 2001; Uversky 2002). Moreover, reaction rate and equilibrium constants were calculated for reactions with different protein substrates, allowing differences in XD binding affinities to be readily quantified.

Binding affinities between XD and N<sub>Tail</sub> constructs were established using 180–225 RU of immobilized XD and N<sub>Tail</sub> concentrations ranging from 0.1 to 10 µM (see Materials and Methods). Dosage-dependent binding was observed in this range. Reactions conforming to a 1:1 ligand-substrate (Langmuir) binding model, exhibiting an excellent fit (i.e., a χ<sup>2</sup> value < 1 and residuals within the range of ±2) following global analysis of sensorgrams. Binding reactions between XD and N<sub>TailHNC</sub> exhibit an equilibrium dissociation constant of 80 nM (see Table 1). The XD binding affinity for N<sub>Tail1</sub> (50 nM) is similar to that of N<sub>TailHNC</sub> indicating that Box1 does not participate in binding. In contrast, removal of either Box3 alone or Box2 plus Box3 results in a strong decrease (three

### Table 1. Calculated equilibrium dissociation constants (K<sub>D</sub>) between XD and N<sub>Tail</sub> proteins

| Analyte     | Residuals | χ<sup>2</sup> | K<sub>D</sub> (M) |
|-------------|-----------|--------------|-----------------|
| N<sub>TailHNC</sub> | -1.5–1.5  | 0.676        | 8.1 × 10<sup>−8</sup> |
| N<sub>Tail3</sub>  | -0.6–0.4  | 0.043        | 1.2 × 10<sup>−5</sup> |
| N<sub>Tail2,3</sub> | -2.4–2.4  | 0.919        | 4.1 × 10<sup>−5</sup> |
| N<sub>Tail1</sub>  | -1.2–1.2  | 0.239        | 4.9 × 10<sup>−8</sup> |
orders of magnitude) in the equilibrium dissociation constant, where $N_{T\text{AIL}1\Delta}$ and $N_{T\text{AIL}2\Delta}$ display similar binding affinities (see Table 1). The strong decrease in the affinity resulting from removal of Box3 clearly indicates that Box2 is not the sole region involved in binding to XD, and attests that Box3 also plays a role in the interaction with XD, as already suggested by SAXS studies.

**Folding propensities of $N_{T\text{AIL}}$ deletion proteins**

The far-UV circular dichroism (CD) spectra of $N_{T\text{AIL}}$ deletion proteins at neutral pH are typical of unstructured proteins, as seen by their large negative ellipticity at 198 nm and very low ellipticity at 185 nm (Fig. 4A–C). The solvent 2,2,2-trifluoroethanol (TFE) mimics the hydrophobic environment experienced by proteins in protein–protein interactions, and is therefore widely used as a probe to unveil disordered regions having a propensity to undergo an induced folding (Hua et al. 1998). Thus, we have recorded CD spectra of $N_{T\text{AIL}}$ deletion proteins in the presence of increasing concentrations of TFE (Fig. 4A–C). All proteins show an increasing gain of $\alpha$-helicity upon addition of TFE, as indicated by the characteristic maximum at 190 nm and minima at 208 nm and 222 nm (Fig. 4A–C). In the case of $N_{T\text{AIL}3\Delta}$ and $N_{T\text{AIL}1\Delta}$, most unstructured-to-structured transitions take place in the presence of 20% TFE, a concentration at which the $\alpha$-helical content is estimated to be about 13% (using the ellipticity at 222 nm) (Fig. 4D). On the other hand, in the case of $N_{T\text{AIL}2\Delta}$, TFE concentrations as high as 30% are required for most pronounced unstructured-to-structured transitions to take place (see Fig. 4B). Moreover, in the presence of 30% TFE, the $\alpha$-helical content of $N_{T\text{AIL}2\Delta}$ is not only lower (11%) than that (16%) of $N_{T\text{AIL}1\Delta}$ and $N_{T\text{AIL}3\Delta}$, but also lower than the $\alpha$-helical content observed at 20% TFE for the two other $N_{T\text{AIL}}$ deletion proteins (Fig. 4D). Therefore, $N_{T\text{AIL}2\Delta}$ displays the lowest $\alpha$-helical potential, while $N_{T\text{AIL}1\Delta}$ and $N_{T\text{AIL}3\Delta}$ exhibit a folding propensity similar to that observed for $N_{T\text{AIL}}$ (Longhi et al. 2003; Bourhis et al. 2004). These results, beyond confirming that the region spanning residues 489–516 affects the folding potential of $N_{T\text{AIL}}$ (Bourhis et al. 2004), suggest that neither Box1 nor Box3 contribute to the $\alpha$-helical propensity of $N_{T\text{AIL}}$, in agreement with the secondary structure prediction provided by PSI-PRED (McGuffin et al. 2000) and PHD (Rost 1996), which both predict an $\alpha$-helix (residues 489–504, see Fig. 3A) as the sole secondary structure element within $N_{T\text{AIL}}$.

![Figure 4](image-url)
Induced folding of $N_{TAIL}$ deletion proteins
in the presence of XD

In order to investigate whether the $N_{TAIL}$ deletion proteins retained the ability to undergo induced folding in the presence of XD, we have used far-UV CD spectroscopy. The $N_{TAILHNFC}$ protein, bearing an N-terminal hexahistidine tag and a C-terminal flag was purified from the soluble fraction of E. coli (data not shown) and used as the reference in these experiments to allow direct comparison with the $N_{TAIL}$ truncated proteins. Noticeably, the CD spectrum of $N_{TAILHNFC}$ is fully superimposable on that of $N_{TAILHN}$ (data not shown), thus ruling out the possibility that the flag sequence might affect the folding properties of the protein.

The far-UV CD spectrum of XD (Fig. 5A–D, gray line) is typical of a structured protein with a predominant α-helical content, as indicated by the positive ellipticity between 185 nm and 200 nm, and by the two minima at 208 nm and 222 nm. After mixing $N_{TAILHNFC}$ with different molar excesses of XD, the observed CD spectra differed from the corresponding theoretical average curves calculated from the individual spectra. Since the theoretical average curves correspond to the spectra that would be expected if no structural variations occur, deviations from these curves indicate structural transitions. The results obtained in the presence of a threefold molar excess of XD indicate a random coil to α-helix transition, as judged by the much more pronounced minima at 208 nm and 222 nm, and by the higher ellipticity at 190 nm of the experimentally observed spectrum compared to the corresponding theoretical average curve (see Fig. 5A). Although even more dramatic structural transitions of full-length $N_{TAIL}$ have been previously observed with a twofold molar excess of XD (Johansson et al. 2003), the results obtained with a threefold molar excess have been selected for presentation in order to allow direct comparison with the $N_{TAIL}$ deletion proteins (see below).

Figure 5, panels B–D, show the results obtained for the three $N_{TAIL}$ deletion proteins in the presence of a threefold molar excess of XD, a condition leading to the most dramatic structural transitions. Removal of Box3 significantly reduces, but does not abrogate, the folding ability of $N_{TAIL}$. Indeed, the experimental CD spectrum does not significantly deviate from the average curve in the 200–260 nm region, but it considerably deviates from the average curve in the 185–195 region (58% mean increase of ellipticity) (Fig. 5B), thus supporting partial folding ability of $N_{TAIL3}$ in the presence of XD.

Figure 5. Induced folding on $N_{TAIL}$ deletion proteins in the presence of XD. Far-UV CD spectra of $N_{TAILHNFC}$ (A), $N_{TAIL3}$ (B), $N_{TAIL2,3}$ (C), and $N_{TAIL1}$ (D) either alone (black line) or in the presence of a threefold molar excess of XD (full circles). The CD spectrum of XD alone (gray line), as well as the theoretical average curves calculated by assuming that no structural variations occur (see Materials and Methods) are also shown (open circles).
Further removal of Box2 results in a truncated NTAIL form, which has completely lost its ability to fold in the presence of XD, as indicated by the good superimposition between the experimental and the average spectra (Fig. 5C). Conversely, removal of Box1 does not affect the folding ability of NTAIL, as the deviations from the average spectrum are similar to those observed with the full-length form (Fig. 5, cf. D and A). The mean increase in ellipticity in the 185–195 nm region (77%) and the decrease in the ellipticity value at 220 nm (46%) observed with NTAILΔ1, are comparable to the corresponding values observed with NTAILHNF (70% and 46%, respectively).

As a control, we recorded CD spectra of NTAILHNF in the presence of lysozyme (data not shown). The absence of significant structural variations even with molar excesses as high as 5, confirms the specificity of the deviations observed upon addition of XD to the NTAIL proteins.

A two- and threefold molar excess of XD is required to induce the most pronounced structural transitions of full-length and truncated forms of NTAIL, respectively. In contrast, an equimolar amount of PCT is sufficient to produce the same effect (Longhi et al. 2003; Bourhis et al. 2004). This difference can be accounted for by a lower affinity of NTAIL towards XD, compared to PCT, rather than to the formation of a 1:2 or 1:3 stoichiometric complex. Indeed, formation of a 1:1 stoichiometric complex between XD and a peptide corresponding to residues 477–505 of N, has been documented using isothermal titration calorimetry (Kingston et al. 2004b). The higher affinity of NTAIL for PCT compared to XD could be ascribed either to cooperativity phenomena among the different XDs within the PCT tetramer (Rahaman et al. 2004) or to the possible contribution of other PCT regions to binding. This latter possibility can be ruled out based on recent data pointing out XD as the sole PCT region contributing to binding (Kingston et al. 2004b; S. Longhi and M.J. Oglesbee, unpubl.).

In conclusion, these results indicate that (1) Box1 is fully dispensable for binding to XD and induced folding, (2) Box2 is strictly required for induced folding to take place, and (3) Box3 contributes to binding to XD, as pointed out by the reduced ability of NTAILΔ3 to undergo induced folding. Moreover, the fact that NTAILΔ3, NTAILΔ1, and NTAIL (Longhi et al. 2003; Bourhis et al. 2004) share similar folding propensities suggests that the contribution of Box3 to the interaction can be accounted for more in terms of binding rather than of induced folding.

Accordingly, we have designed an N-terminally hexahistidine-tagged NTAIL variant form bearing a tyrosine to tryptophan substitution at position 518 (see also Fig. 3A). Introduction of a tryptophan residue in Box3 allowed binding events to be followed by fluorescence spectroscopy, while maximizing the conservative nature of the substitution (note that neither NTAIL nor XD contain any tryptophan residue).

Most recombinant product was purified from the soluble fraction of the bacterial lysate (Fig. 6A). The identity of the recombinant product was confirmed by IP studies using anti-N Cl25, and anti-hexahistidine tag mAbs (data not shown). As already observed in the case of full-length NTAIL and NTAIL deletion proteins, NTAILWS518 migrates in SDS-PAGE with an apparent MM higher than expected (Fig. 6A). The mutated protein, displays the same gel filtration elution profile as the wt form (data not shown), leading to an estimated Rs of 27 ± 3 Å.

As shown in Figure 6B, the far-UV CD spectrum of NTAILWS518 is almost perfectly superimposable on that of NTAILHNF, thus indicating that the introduction of the tryptophan residue does not affect the overall secondary structure content of the protein. In order to investigate whether the variant form retained the ability to undergo induced folding in the presence of XD, we have added various molar excesses (ranging from 1 to 4) of XD to NTAILWS518, and recorded the corresponding far-UV CD spectra (data not shown). These latter studies indicate that the tyrosine to tryptophan substitution does not affect the ability of the protein to undergo induced folding in the presence of the partner, thus supporting the biochemical relevance of this variant form.

Fluorescence spectroscopy studies showed that NTAILWS518 has a maximum of emission at 356 nm, indicating that Trp 518 is fully exposed to the solvent (data not shown). Addition of gradually increasing XD concentrations triggers an increase in the fluorescence intensity in a dose-dependent manner, which indicates a modification in the pattern of interactions with neighboring groups. At the same time, addition of XD causes a progressive shift in the emission maximum from 356 nm to 352 nm (data not shown), thus indicating that Trp 518 becomes only slightly less exposed to the solvent.

No significant variations are observed in the fluorescence spectrum obtained after addition to NTAILWS518 of a 2 μM solution of an irrelevant protein (SARS virus, unclassified protein 5) of similar size and devoid of tryptophan residues (data not shown). After plotting the relative fluorescence intensity increase as a function of the XD concentration (Fig. 6C), an apparent constant equilibrium dissociation (K_Dapp) value of 133 ± 32 μM is derived. This value is in good agreement with the value obtained by surface plasmon resonance studies with wt

Fluorescence spectroscopy studies

In order to further characterize the contribution of Box3 to binding, we have used fluorescence spectroscopy.
Although the environment of Trp 518 remains mostly polar upon binding to XD, the observed increase in the fluorescence intensity indicates that the chemical environment of Trp 518 is affected, thus further suggesting that the C terminus of NTAIL interacts with XD.

Two-dimensional Heteronuclear Magnetic Resonance (NMR)

In order to further explore the nature of the interaction established between Box3 and XD, we have used NMR spectroscopy. To this endeavor, we have recorded a HSQC spectrum of 15N uniformly labeled NTAILHN either alone or in the presence of a twofold molar excess of XD, as well as of 15N uniformly labeled NTAILD3 in the same conditions.

This analysis allowed a quantitative estimation of the number of residues involved in the interaction with XD by following chemical shift changes in the backbone amide and proton resonances upon addition of unlabeled XD. Upon addition of XD to NTAILHN, 16 correlation peaks are displaced. Among them, 11 undergo an upfield shift (see Fig. 7, stars), which indicates a random coil to α-helix transition, and two correspond to the side chain of either a Gln or an Asn. Additionally, at least seven additional peaks undergo a less dramatic displacement (see Fig. 7, diamonds). Because of its small amplitude, this shift most likely does not reflect a conformational change. Nevertheless, it is indicative of a change in the chemical environment of these residues resulting either by direct interaction with the partner or by local magnetic perturbations of residues spatially close to the newly formed α-helix. Based on this experiment, however, the determination of NTAIL residues involved in these two types of shifts is not possible. The precise identification of these residues would require the full assignment of the NTAIL HSQC spectrum, which is complicated by the strong signal overlapping inherent in the predominantly unfolded nature of NTAIL.

Comparison of present results to those of the NMR spectroscopy studies of Kingston et al. (2004a) suggest that the 11 residues undergoing the random coil to α-helix transition are most likely located within the 486–503 region. In order to assess whether the additional NTAIL residues undergoing a less pronounced displacement upon binding to XD are located within Box3, we recorded a HSQC spectrum of 15N uniformly labeled NTAILD3 either alone or with a twofold molar excess of XD. As shown in Figure 7, the same peaks undergoing the large displacement in the NTAILHN–XD complex are also observed in the NTAILD3–XD complex. In particular, among these peaks, the occurrence in both complexes of the 11 peaks that undergo the random coil to α-helix transition (see Fig. 7, stars), provides further
support that the helical folding occurs within Box2. On the other hand, the seven peaks that undergo a less dramatic displacement upon complex formation with XD (see Fig. 7, diamonds) are not present in the spectrum recorded on the N\textsuperscript{TAIL}\textsuperscript{3}–XD complex. This latter observation indicates that these seven peaks correspond to residues that are located within Box3.

In conclusion, these experiments reveal that complex formation between N\textsuperscript{TAIL} and XD implies two types of interaction: one, mediated by residues belonging to Box2, involves a significant gain of α-helicity, while the other, attributable to Box3 residues, is not accompanied by a significant structural transition.

Discussion

In this paper we show that N\textsuperscript{TAIL} remains predominantly unfolded after binding to XD, with two distinct sites being involved in the interaction. Although the XD-induced gain of regular secondary is restricted to Box2 (aa 489–504), the region encompassing residues 489–504 is not the only N\textsuperscript{TAIL} region involved in binding to XD. In particular, we present several lines of evidence indicating that the extreme C terminus of N\textsuperscript{TAIL} (Box3, aa 517–525) also contributes to binding, without however gaining any regular secondary structure.

The C terminus of N\textsuperscript{TAIL} is an additional XD binding site

SAXS studies suggest that the C terminus of N\textsuperscript{TAIL} interacts with XD. In particular, the calculated overall envelope of the complex shows the presence of a globular cluster of invariant size at one extremity and of an elongated protuberance with varying shapes. The elongated protuberance corresponds to the 92-residue-long N-terminal region of N\textsuperscript{TAIL} while the more compact region accommodates the structure of the XD–N\textsuperscript{TAIL}\textsuperscript{486–505} complex as well as the 20 C-terminal residues of N\textsuperscript{TAIL}. Data from overall shape calculations thus clearly indicate that the C terminus is not protruding towards the solvent, and remains close to XD. On the other hand, attempts to more precisely model the conformation adopted by the C-terminal region of N\textsuperscript{TAIL} within the complex led to an ensemble of solutions. In all these solutions, the C-terminal region of N\textsuperscript{TAIL} always packs against the XD–N\textsuperscript{TAIL}\textsuperscript{486–505} complex, rather than being extended and exposed to the solvent. Identification of the possible gain of regular secondary structure within the C-terminal region of N\textsuperscript{TAIL} is beyond the resolution limits of SAXS. However, results provided by CD, fluorescence spectroscopy, and heteronuclear NMR all converge to suggest that the C-terminal region of N\textsuperscript{TAIL} does not gain any regular secondary structure (see below). Besides supporting a role for the C-terminal region of N\textsuperscript{TAIL} in the interaction with XD, these data also

Figure 7. 2D-HSQC NMR spectra. HSQC spectrum of a 0.5 mM solution of purified \textsuperscript{15}N-N\textsuperscript{TAIL}HN alone (black) or in the presence of a twofold molar excess of XD (red), of a 0.125-mM solution of purified \textsuperscript{15}N-N\textsuperscript{TAIL}\textsuperscript{3} alone (green) or in the presence of a twofold molar excess of XD (blue). All proteins were in 10 mM sodium phosphate at pH 7. All spectra are recorded at 283 K. ppm quotes for resonance shifts in parts per million of the spectrophotometer frequency. Some peaks are labeled as follows: stars indicate peaks present in spectra of both complexes only, while diamonds quote for peaks present in the spectrum recorded with the \textsuperscript{15}N-N\textsuperscript{TAIL}HN–XD complex, but not with the \textsuperscript{15}N-N\textsuperscript{TAIL}\textsuperscript{3}–XD complex. The inset shows the purified \textsuperscript{15}N-N\textsuperscript{TAIL}\textsuperscript{3} and \textsuperscript{15}N-N\textsuperscript{TAIL}HN proteins.
NMR experiments show that upon complex formation with XD, 11 N\textsubscript{TAIL} residues, belonging to Box2, undergo a random coil to \(\alpha\)-helix transition, while at least seven additional residues undergo a shift in their chemical environment not accompanied by the gain of regular secondary structure elements. Our data show that these latter peak displacements correspond to residues belonging to Box3, as indicated by the absence of such peaks in both the spectra of N\textsubscript{TAIL3} alone and in complex with XD. Thus, the NMR studies, beyond confirming the role of Box3 in the interaction with XD, also highlight that binding of N\textsubscript{TAIL} to XD implies two types of interaction, where gain of regular secondary structure is restricted to only one of the binding determinants, Box2.

**Structural and functional insights into the interaction between N\textsubscript{TAIL} and XD**

The accommodation of the 486–505 region of N within XD (Kingston et al. 2004a) triggers some minor rearrangements at the surface of the latter, compared to the crystal structure of the uncomplexed form (Johansson et al. 2003). In particular, the largest movements are observed for the side chains of residues Arg15 and Glu17 of XD (corresponding to residues 472 and 474 of P). The Arg15 NH2 atom moves 9.7 Å away in the chimeric structure compared to the structure of XD alone, while the Glu17 C\& atom undergoes a shift of 2.5 Å. These two residues are both located within the loop connecting \(\alpha1\) and \(\alpha2\) helices, and occur at least 9 Å away from Box2, thus ruling out the possibility that the observed spatial rearrangements they undergo could be ascribed to Box2 embedding. The significant displacement they undergo in the complex therefore indicates that binding of Box2 induces local conformation changes in XD that could favor interaction with Box3. The surface displaced residues of XD could, in fact, be part of a Box3 binding site.

We tentatively propose that binding to XD might take place through a sequential mechanism involving successive binding of disordered domains, as it has been recently reported for p27 upon binding to the CdK2-cyclin A complex (Lacy et al. 2004).

The \(K_D\) value between N\textsubscript{TAIL} and XD, as measured by both fluorescence spectroscopy and surface plasmon resonance, is in the 100 nM range. Surprisingly, this value is considerably lower than that reported by Kingston et al. (2004b) (13 \(\mu\)M) and derived from isothermal titration calorimetry studies. A weak binding affinity, associated with a fast association rate, would ideally fulfill the requirements of a polymerase complex, which has to cartwheel on the nucleocapsid template during both transcription and replication. However, a \(K_D\) in the \(\mu\)M range would not seem to be physiologi-

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**The additional XD binding site does not gain any regular secondary structure**

Spectroscopy studies showed that Box3 contributes to binding to XD but does not undergo any gain of regular secondary structure. Although Box3 does not affect the folding potential of N\textsubscript{TAIL}, as indicated by CD studies in the presence of TFE, the removal of Box3 significantly reduces the ability of N\textsubscript{TAIL} to undergo induced folding in the presence of XD. This supports a role for Box3 in binding to XD. Further removal of Box2 results in a truncated form that has a significantly decreased folding potential and has lost the ability to undergo induced folding in the presence of XD. These results are consistent with the unique \(\alpha\)-helical forming potential of Box2 and its role as primary binding site for XD.

Contribution of Box3 to binding without dramatic structural change is supported by fluorescence spectroscopy data, which show an increase in the fluorescence intensity of N\textsubscript{TAIL} upon addition of XD. Increases in the fluorescence intensity upon binding to a partner/ligand have been already documented, for both folded (Bette et al. 2002) and intrinsically unstructured proteins (Raggett et al. 1998), and indicate that the chemical environment of the Trp, although remaining mostly polar, is changed as a result of the addition of the partner. However, the fluorescence data do not enable us to discriminate between a direct Trp 518–XD interaction and secondary effects resulting from local perturbations triggered by a direct Box2–XD interaction.

That complex formation with XD involves both Box2 and Box3, and that only Box2 undergoes a gain of \(\alpha\)-helicity, is confirmed by NMR studies. Heteronuclear
cally relevant considering the low intracellular concentrations of P in the early phases of infection. Moreover, such a weak affinity is not consistent with the ability to readily purify nucleocapsid–P complexes using rather stringent techniques such as CsCl isopycnic density centrifugation (Robbins and Bussell 1979; Stallcup et al. 1979; Robbins et al. 1980; Oglesbee et al. 1989). Our data support a higher affinity between P and N, resulting in a stable P–NTAIL complex that would be predicted to hinder processive movement of P along the nucleocapsid template. In agreement with this prediction, Box3 has been shown to have an inhibitory role upon transcription and replication, as indicated by previous minireplicon experiments, where deletion of Box3 enhanced basal reporter gene expression (Zhang et al. 2002). We can speculate that the transient nature of the NTAIL–XD interaction might be ensured by the possible intervention of cellular and/or viral cofactors. Indeed, the requirement for cellular or viral cofactors in both transcription and replication has been already documented in the case of measles (Vincent et al. 2002), respiratory syncytial (Ferns and Collins 1999), and Ebola viruses (Hartlieb et al. 2003). These cofactors may serve as processivity or transcription elongation factors and could act by modulating the strength of the interaction between the polymerase complex and the nucleocapsid template. Such a mechanism may explain the stimulatory effect of hsp72 on MV transcription and genome replication, an effect mediated by Box3–hsp72 interaction (Zhang 2002). In this capacity, hsp72 may neutralize the contribution of Box3 to a more stable complex between P and NTAIL, thereby promoting successive cycles of P binding and release that are essential to polymerase processivity.

**Conclusion**

Using different physico-chemical approaches we have shown that the interaction of NTAIL with XD involves an additional, previously unreported site located at the extreme C terminus of NTAIL. While the primary site (i.e., Box2) gains α-helical structure upon binding to XD, this additional site does not gain any regular secondary structure elements. Nevertheless, it plays a crucial role in stabilizing the complex. We have previously shown that NTAIL belongs to the premolten globule subfamily, e.g., it possesses a certain extent of residual secondary and/or tertiary structure (Longhi et al. 2003; Bourhis et al. 2004). In agreement with these findings, NMR studies showed that the NTAIL region encompassing residues 486–503 is not a statistical random coil (Kingston et al. 2004a). These results, supporting a restricted conformational freedom of this NTAIL region, are in agreement with the speculation that residual structure within NTAIL may play a role for efficient binding to P (Longhi et al. 2003; Bourhis et al. 2004). That residual structure within intrinsically disordered proteins may favor the folding process triggered by binding to a partner has already been reported (Bienkiewicz et al. 2002; Fuxreiter et al. 2004; Csizmok et al. 2005).

NTAIL provides an interesting model system for the study of the interaction of an intrinsically disordered protein and its partners. Its multiple-site mode of interaction well illustrates the complexity of the contacts established by intrinsically disordered proteins and emphasizes the need for thorough investigations of the molecular mechanisms underlying recognition of the partner. Indeed, binding of disordered regions to their targets involves different types of interactions, such as binding coupled to folding, binding with no gain of regular secondary structure, or coexistence of both interaction mechanisms (for a review, see Dyson and Wright 2005).

Binding coupled to folding has been reported for the phosphorylated kinase-inducible domain (pKID) of CREB, which undergoes a coil to α-helix folding transition upon binding to the KIX domain of the transcription coactivator CBP (CREB Binding Protein) (Radhakrishnan et al. 1997). The amphipathic helix αB of pKID interacts with a hydrophobic groove defined by helices α1 and α3 of the partner. The other pKID helix, αA, contacts a different face of the α3 helix. This mode of interaction is reminiscent of that occurring between NTAIL and XD. Similarities concern the involvement of two distinct sites within the disordered domain (where Box2 and Box3 resemble helices αB and αA, respectively) and embedding of an α-helix within a hydrophobic cleft delimited by α-helices from the structured partner. The difference concerns the mode of interaction of the second site, where Box3 of NTAIL does not gain any regular secondary structure element, contrary to αA of pKID. Nevertheless, complexes with dual interaction have already been described in the literature. Among them, we mention the case of the binding of p27 to the cyclinA–Cdk2 complex (Lacy et al. 2004), and that of the activation domain of CITED2 to the TAZ1 domain of CBP (De Guzman et al. 2004).

On the other hand, binding without any concomitant gain of regular secondary structure, has been also reported. This is the case of the unfolded proteins 4E–BP1 (4E binding protein 1), an inhibitor of translation. Its binding to eIF4E does not involve any transition to stable regular secondary structure (Fletcher et al. 1998). Interestingly, when eIF4E binds to another partner, namely eIF4G, it induces a coil-to-helix transition in the latter (Marcotrigiano et al. 1999), highlighting the
diversity of the structural transitions that can be triggered by a structured partner.

Finally, there are a few examples of proteins that undergo different structural transitions as a function of the partner they bind. Notably, HIFα (hypoxia-inducible factor α) possesses an intrinsically disordered domain, which can interact with two different partners. This unstructured domain adopts an α-helical structure when bound to the TAZ1 domain of CBP. However, binding of the same region of HIFα to an asparagine hydroxylase results in a highly extended conformation, which is required for the enzymatic activity of the latter. These findings provide an elegant example of the plasticity that intrinsically disordered proteins display for different partners involved in different functions. We can speculate on a similar behavior in the case of NTAIL. In particular, Box2 and Box3 of NTAIL interact both with at least two distinct partners, P and Hsp72, and competition between XD and Hsp72 for binding to NTAIL has been recently shown (Zhang et al. 2005). It is conceivable that binding of NTAIL to P or to Hsp72 may involve a different structural transition, as in the case of HIFα.

Last, but not least, our findings, beyond contributing to elucidate the dynamics of the interactions established by intrinsically disordered proteins, provide an interesting target site for mutational studies aimed at exploring further the mode of interaction between NTAIL and XD.

Materials and methods

Bacterial strains and media

The E. coli strains DH5α (Stratagene) was used for selection and amplification of DNA constructs. The E. coli strains Rosetta [DE3] pLysS (Novagen) and C41 [DE3] (Avidis) were used for expression of recombinant proteins. E. coli was grown either in Luria-Bertani (LB) medium, or in minimal M9 medium supplemented with 15NH4Cl.

Chemicals and antibodies

Pfu polymerase was from Promega. Primers were purchased from Invitrogen. The anti-hexahistidine tag mAb was purchased from Qiagen. The anti-flag mAb was purchased from Sigma. The anti-N CL25 (Giraudon et al. 1988; Buckland et al. 1989) mAb was kindly provided by D. Gerlier.

Construction of protein expression plasmids

The XD gene construct, encoding residues 459–507 of the MV P protein (strain Edmonston B) with an hexahistidine tag fused to its N terminus, has already been described (Johansson et al. 2003).

All NTAIL constructs were obtained by PCR using the MV N gene, strain Edmonston B, as template. The NTAILHNC construct, encoding residues 401–525 of the MV N protein with a hexahistidine tag fused to its N terminus, was obtained using the plasmid pET21a/N (encoding the MV N protein; Kurlin et al. 2002a) as template. Forward primer (5'-gtgctacaaggatgagatcactatcatcatcatcatcatctacaagctgtggagac-3') was designed to introduce a hexahistidine tag encoding sequence (upper case) at the N terminus of NTAIL, while reverse primer (5'-ggggaccactttgctgcggatcttggctggtcactactattcttcggtggtcactac-3') was designed to introduce an AttB2 site (bold). The PCR amplification product was further used as template in a second PCR step, using forward primer (5'-ggggaccactttgctgcggatcttggctggtcactactattcttcggtggtcactac-3') and XD primer (5'-ggggaccactttgctgcggatcttggctggtcactactattcttcggtggtcactac-3') designed to introduce an AttB1 site (bold), and reverse primer as above. After purification (PCR Purification Kit; Qiagen), the PCR product was cloned into the pDest14 vector (Invitrogen) using the Gateway recombination system (Invitrogen). The final construct is referred to as pDest14/NTAILHNC.

The NTAILHNFCTAIL2 gene construct, encoding residues 401–525 of the MV N protein with an N-terminal hexahistidine tag and a C-terminal Flag sequence (dykddddk) (Brizzard et al. 1994), was obtained using the plasmid pDST14/NTAILHNC as template. Forward primer (5'-ggggaccactttgctgcggatcttggctggtcactactattcttcggtggtcactac-3') was designed to introduce an AttB1 site (bold), and reverse primer (5'-gtgctacaaggatgagatcactatcatcatcatcatcatctacaagctgtggagac-3') was designed to introduce a hexahistidine tag encoding sequence (upper case). The PCR amplification product was further used as template in a second PCR step, using the same forward primer as above, and reverse primer (5'-ggggaccactttgctgcggatcttggctggtcactactattcttcggtggtcactac-3') designed to introduce an AttB2 site (bold). After purification, the PCR product was cloned into the pDest14 vector to yield pDest14/NTAILHNFCTAIL2.

The NTAILHNFCTAIL3 gene construct, encoding residues 401–516 of the MV N protein with an N-terminal hexahistidine tag plus a C-terminal Flag sequence, was obtained using the plasmid pet21a/N as template. Forward primer (5'-gtgctacaaggatgagatcactatcatcatcatcatcatctacaagctgtggagac-3') was designed to introduce a hexahistidine tag encoding sequence (upper case) at the N terminus of NTAIL, while reverse primer (5'-gtgctacaaggatgagatcactatcatcatcatcatcatctacaagctgtggagac-3') was designed to introduce a hexahistidine tag encoding sequence (upper case). The PCR amplification product was further used as template in a second PCR step, using forward primer (5'-gtgctacaaggatgagatcactatcatcatcatcatcatctacaagctgtggagac-3') and XD primer (5'-ggggaccactttgctgcggatcttggctggtcactactattcttcggtggtcactac-3') designed to introduce an AttB1 site (bold), and reverse primer (5'-ggggaccactttgctgcggatcttggctggtcactactattcttcggtggtcactac-3') designed to introduce an AttB2 site (bold). After purification, the PCR product was cloned into the pDest14 vector to yield pDest14/NTAILHNFCTAIL3.
product was cloned into the pDest14 vector to yield pDest14/$N_{TAIL\text{L}}$.  

The $N_{TAIL\text{L}}$ gene construct, encoding residues 421–525 of the MV N protein with an N-terminal hexahistidine tag plus a C-terminal Flag sequence, was obtained using the plasmid pET21a/N as template. Forward primer (5’-gatagaaacctgaTCTCATCATCATCACGGTTCCTCCTT-3’) was designed to introduce a hexahistidine tag encoding sequence (upper case) at the N terminus of $N_{TAIL\text{L}}$, and reverse primer (5’-gttcatTTTGTGTCACTCGTCCTTATAACTGTCGTTGTTTG-3’) was designed to introduce a Flag-encoding sequence (upper case). The PCR amplification product was further used as template in a second PCR step, using the same pair of primers as in the second PCR step, which yielded both $N_{TAIL\text{L}}$ and $N_{TAIL\text{L,23}}$ amplification products. After purification, the PCR product was cloned into the pDest14 vector to yield pDest14/$N_{TAIL\text{L}}$.

The $N_{TAIL\text{L,518}}$ gene construct, encoding residues 401–525 of the MV N protein with a Trp → Tyr substitution at position 518, and with a hexahistidine tag fused to its N terminus, was obtained by PCR, using the plasmid pET21a/N as template. Two separate PCR steps were carried out in parallel, yielding amplification products A and B. Product A was obtained using forward primer (5’-gatagaaacctgaTCTCATCATCATCACGGTTCCTCCTT-3’) designed to introduce a hexahistidine tag-encoding sequence (upper case) at the N terminus of $N_{TAIL\text{L}}$, while reverse primer (5’-aagaTCTCATCACCATCCTGCTAAGAACATG-3’) was designed to introduce a Trp at position 518 (bold and upper case) and a silent mutation at nucleotide position 1545 (upper case) thus resulting in the introduction of a BstXI site (underlined). Product B was obtained using forward primer (5’-gacacccgAAGATCT-3’) designed to introduce a Trp at position 518 (bold and upper case) and a silent mutation at nucleotide position 1545 (upper case) thus resulting in the introduction of a BstXI site (underlined), and reverse primer (5’-ccgatgagATCTGCTAAGAACATG-3’) designed to introduce a hexahistidine tag-encoding sequence (upper case) at the N terminus of $N_{TAIL\text{L}}$, and reverse primer (5’-gacacccgAAGATCTGCTAAGAACATG-3’) was designed to introduce an AtrB1 site (bold), and reverse primer (5’-gggaccaTCTCATCACGGTTCCTCCTT-3’) was designed to introduce an AtrB2 site (bold) and a Trp at position 518 (upper case). After purification, the PCR product was cloned into the pDest14 vector using the Gateway recombination system. The final construct is referred to as pDest14/ $N_{TAIL\text{L,518}}$. Candidate clones bearing the desired mutation were selected on the basis of the ability of their recombinant plasmids to be restricted by BstXI, a unique restriction site introduced by PCR together with the Tyr to Trp substitution.

The sequence of the coding region of all expression plasmids was verified by sequencing (MilleGen).

Expression of $N_{TAIL}$ constructs

E. coli strain Rosetta [DE3] (Novagen) was used for the expression of $N_{TAIL}$ constructs. Since the MV N gene contains several rare codons with a very low frequency in E. coli, coexpression of $N_{TAIL}$ constructs with the plasmid pLYS S (Novagen) was carried out. This plasmid, which supplies six rare tRNAs, carries also the lysozyme gene, thus allowing a tight regulation of the expression of the recombinant gene, as well as a facilitated lysis. Cultures were grown overnight to saturation in LB medium containing 100 µg/mL ampicillin and 17 µg/mL chloramphenicol. An aliquot of the culture was diluted 1/25 in LB medium and grown at 37°C. At OD$_{600}$ of 0.7, isopropyl β-D-thiogalactopyranoside (IPTG) was added to a final concentration of 0.2 mM, and the cells were grown at 37°C for 3 h. The induced cells were harvested, washed, and collected by centrifugation. The resulting pellets were frozen at −20°C.

Isotopically substituted (15N) $N_{TAIL\text{L}}$ and $N_{TAIL\text{L,518}}$ were prepared by growing transformed bacteria in minimal M9 medium supplemented with 15NH$_4$Cl (0.8 g/L). A 50-mL preculture grown overnight to saturation in LB medium containing 100 µg/mL ampicillin and 17 µg/mL chloramphenicol, was harvested, washed in minimal M9 medium, and inoculated into 1 L of minimal M9 medium supplemented with ampicillin and chloramphenicol. The culture was grown at 37°C. At OD$_{600}$ of 0.5, IPTG was added to a final concentration of 0.2 mM and the cells were grown first at 37°C for 3 h, and then over night at 28°C. The induced cells were harvested, washed, and collected by centrifugation. The resulting pellets were frozen at −20°C.

Expression of tagged XD was carried out as described in Johansson et al. (2003).

Purification of $N_{TAIL}$ proteins

Cellular pellets from bacteria transformed with the different $N_{TAIL}$ expression plasmids were resuspended in 5 volumes (v/w) buffer A (50 mM sodium phosphate at pH 8, 300 mM NaCl, 10 mM Imidazole, 1 mM phenyl-methyl-sulphonyl-fluoride (PMSF) supplemented with lysozyme 0.1 mg/mL, DNase I 10 µg/mL, protease inhibitor cocktail (Sigma) (50 µL/g cells). After a 20-min incubation with gentle agitation, the cells were disrupted by sonication (using a 750 W sonicator and four cycles of 30 sec each at 60% power output). The lysate was clarified by centrifugation at 30,000 g for 30 min. Starting from a 1-L culture, the clarified supernatant was incubated for 1 h with gentle shaking with 4 mL Chelating Sepharose Fast Flow Resin preloaded with Ni$^{2+}$ ions (Amersham Pharmacia Biotech), previously equilibrated in buffer A. The resin was washed with buffer A, and the $N_{TAIL}$ proteins were eluted in buffer A containing 250 mM imidazole. Eluates were analyzed by SDS-PAGE for the presence of the desired product. The fractions containing the recombinant product were combined, and concentrated using Centricron Plus-20 (molecular cutoff, 5000 Da) (Millipore). The proteins were then loaded onto a Superdex 75 HR 10/30 column (Amersham Pharmacia Biotech) and eluted in either 10 mM sodium phosphate at pH 7 or 10 mM Tris/HCl at pH 8. The proteins were stored at −20°C.

Purification of histidine-tagged XD was carried out as described in Johansson et al. (2003). All purification steps, except for gel filtrations, were carried out at 4°C.

Apparent molecular mass of proteins eluted from gel filtration columns was deduced from a calibration carried out with LMW and HMW calibration kits (Amersham Pharmacia Biotech). The theoretical Stokes radii ($R_{S}\Lambda$) of native (Ra$\Lambda$) and fully unfolded (R$U$) protein with a MM (in Daltons) were calculated according to (Uversky 1993): $\log(R_{S}\Lambda) = 0.369 \cdot \log(MM) – 0.254$ and $\log(R_{U}\Lambda) = 0.533 \cdot \log(MM) – 0.682$.  

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**Determination of protein concentration**

Protein concentrations were calculated either using the theoretical absorption coefficients ε (mg/mL·cm) at 280 nm as obtained using the program ProtParam at the EXPASY server (http://www.expasy.ch/tools), or the Biorad protein assay reagent (Bio-Rad).

**Immunoprecipitation studies of NTAIL proteins**

IP experiments were carried out using the anti-N Cl 25, the anti-flag, and the anti-hexahistidine tag mAbs, and bacterial lysates expressing NTAIL proteins as described in Longhi et al. (2003).

**Small angle X-ray scattering**

All protein samples were prepared by dilution of the purified NTAILH and XD solutions in buffer 10 mM Tris/HCl at pH 8, 40 mM NaCl, with 1 mM DTT as radiation scavenger. The complex NTAILH–XD was prepared by mixing XD and NTAILH with a molar ratio of 2:1 in the same buffer and at a final protein concentration of 10 mg/mL. The samples were filtered prior to each measurement (Millex syringe filters 0.22 μm, Millipore) to eliminate possibly existing large aggregates.

SAXS experiments were carried out on beamline ID02 (Narayanan et al. 2001) at the European Synchrotron Radiation Facility (ESRF), Grenoble, France. The wavelength was 1.0 Å and the sample-to-detector distance was 3.0 m and 1.0 m, leading to scattering vectors q ranging from 0.02 to 0.20 Å⁻¹ and 0.05 to 0.40 Å⁻¹, respectively. The scattering vector is defined as q = 4 π/λ sinθ, where θ is the scattering angle. The detector was an X-ray image intensified optically coupled to an ESRF developed FReLoN CCD camera. Forty successive frames of 1.0 sec with a 5-sec pause between each frame were recorded for each sample. The protein solution was circulated through an evacuated quartz capillary between each frame. Thus, no protein solution was irradiated longer than 1.0 sec. Each frame was then carefully inspected to check for possible bubble formation or radiation-induced aggregation. No such effect was observed, and individual frames could then be averaged. Absolute calibration was made with a Lupolen sample. A series of measurements at different protein concentrations ranging from 1.8 to 10 mg/mL were performed for every protein (XD, NTAILH, and the mixture NTAILH–XD) to check for interparticle interaction. Background scattering was measured before or after each protein sample using the buffer solution and then subtracted from the protein scattering patterns after proper normalization and correction from detector response. All the experiments were carried out at 20°C. The data acquired at both sample-to-detector distances of 3 m and 1 m were merged and extrapolated to zero concentration for the calculations using the entire scattering spectrum.

The scattering pattern of the NTAILH–XD complex was obtained following this process: to avoid any possible bias on the absolute intensities due to the concentration, the experimental merged scattering curves obtained as described above were normalized by their theoretical I(0). The scattering pattern of XD was then subtracted twice from the scattering pattern of the mixture NTAILH–XD (molar ratio 1:2). Given the rather low Kd between NTAIL and XD (~100 nM; see Results section), we assumed that the concentration of uncomplexed NTAILH in solution is negligible at the concentration used in the SAXS experiments (2 to 10 mg/mL). The scattering pattern of the complex thus obtained was then used for further calculations.

The value of the Rg was derived from the Guinier approximation (Guinier and Fournet 1955): I(q) = I(0) exp[−q²Rg²/3], where I(q) is the scattered intensity and I(0) is the forward scattered intensity. The Rg and I(0) are inferred, respectively, from the slope and the intercept of the linear fit of Ln[I(q)] versus q² at low q values (q Rg < 1.0). The distance distribution function P(r) is the histogram of all the interatomic distances within a molecule. This function also provides the maximum dimension Dmax of the molecule, which is defined as the point where P(r) becomes zero. The P(r) function was calculated by the Fourier inversion of the scattering intensity I(q) using GNOM (Svergun 1992) and GIFT (Bergmann et al. 2000) on the entire scattering spectra.

**3D modeling**

The low-resolution shape of the NTAILH–XD complex was determined ab initio from the scattering curve using the program GASBOR (Svergun et al. 2001). This program restores low-resolution shapes of protein and calculates a volume filled with densely packed spheres (dummy residues) fitting the experimental scattering curve by a simulated annealing minimization procedure and considering the protein as an assembly of dummy residues centered on the Cn positions (spheres of 3.8 Å diameter) with a nearest-neighbour distribution constraint. Several independent fits were run with no symmetry restriction with an input of 188 dummy residues, corresponding to the 56 residues of XD and 132 residues of NTAILH. The program package CREDO (CHADD and GLOOPY) was used to restore the low-resolution model of NTAILH in complex with XD, with the crystal structure of the chimeric protein between XD and the region of NTAIL encompassing residues 486–505 (XD–NTAIL486–505) (Kingston et al. 2004a) as template. CREDO is an extension of the original program GASBOR. It calculates the structure of missing domains or loops of crystal structures from the experimental scattering curve of the entire particle and represents them by an ensemble of dummy residues forming a chain-compatible model.

**Surface plasmon resonance studies**

Binding between purified XD and purified NTAIL proteins was analyzed by using BIAcore 3000 (Amersham Pharmacia Biotech), a system for real-time biomolecular interaction analysis that is based upon surface plasmon resonance technology. Purified XD (1.4 μg/mL in acetate buffer at pH 5.5) was covalently bound to carboxy-methyl groups of CM5 sensor chips using amine-coupling chemistry (Biosensor AB, Amersham Pharmacia Biotech). The levels of immobilized XD were comprised between 180 and 225 RU (1000 RU equal a change in mass of 1 ng/mm² on the sensor surface) (Zhang et al. 2002). Kinetic and equilibrium constants were calculated by global analysis of reactions with multiple analyte concentrations (0.1, 0.2, 0.5, 1, 2, 5, and 10 μM). Reactions were performed at 25°C. Remaining flow channels on the sensor chip included a control for a nonspecific interaction with the sensor chip (i.e., activated/block flow channel) and a control for nonspecific interactions with irrelevant protein targets (i.e., lactoferrin conjugated flow channel). Protein analytes were passed over the sensor chip in HBS-P buffer (0.01 M HEPES at pH 7.4, 0.15 M NaCl, 0.005% surfactant P-20) containing 2.5 mM magnesium acetate and 2.5 mM ATP. Sensorgrams plotted changes in surface plasmon
resonance (measured in RU) as a function of time. Multiple sensorgrams representing various analyte concentrations were analyzed by using BIAlEvaluation 3.1 software. Background interaction of N_{TAIL} proteins (i.e., analytes) with the sensor surfaces were measured on flow channels that were activated and subsequently blocked under buffer conditions used to immobilize XD. This background was subtracted from all binding curves (i.e., sensorgrams) prior to global analyses. Immobilized lactoferrin was used as a specificity control, and the resultant sensorgrams ruled out high affinity interactions between N_{TAIL} constructs or peptides and this irrelevant protein ligand (data not shown). Global fitting of experimental data to well-characterized binding reactions was used to define reaction rate and equilibrium constants. Signal changes on the activated/blocked control channel were subtracted from the peptide-XD and N_{TAIL} protein-XD interactions using in-line reference and the subtracted sensorgrams were analyzed. Curves generated with serial analyte concentrations were applied globally to the 1:1 binding model, without correction for baseline drifting depending on baseline status. $\chi^2$ and residual values were used to evaluate the quality of fit between experimental data and individual binding models. Plots of residuals indicate the difference between the experimental and reference data for each point in the fit. The $\chi^2$ value represents the sum of squared differences between the experimental data and reference data at each point. A good fit between experimental and reference data has small residuals in the $\chi^2$ values and $\chi^2$ values are less than 10.

Circular dichroism

The CD spectra were recorded on a Jasco 810 dichrograph using 1-mm thick quartz cells in 10 mM sodium phosphate (pH 7) at 20°C. Structural variations of N_{TAIL} proteins were measured as a function of changes in the initial CD spectrum upon addition of either increasing concentrations of TFE (Fluka), or different amounts of XD or lysozyme (Sigma). CD spectra were measured between 185 and 260 nm, at 0.2 nm/min and were averaged from three independent acquisitions. Mean ellipticity values per residue ($[\theta]$) were calculated as $[\theta] = \frac{3300 \Delta A}{l \cdot c \cdot n \cdot m}$ where $l$ (path length) = 0.1 cm, $n$ = number of residues, $m$ = molecular mass in daltons, and $c$ = protein concentration expressed in mg/mL. Number of residues ($n$) are 140 for N_{TAILHNC}, 313 for N_{TAIL3}, 103 for N_{TAIL24}, 120 for N_{TAIL23}, 132 for N_{TAIL18}, 56 for XD, and 129 for lysozyme, while $m$ values are 15,626 Da for N_{TAILHNC}, 14,523 Da for N_{TAIL3}, 11,539 Da for N_{TAIL24}, 13,440 Da for N_{TAIL23}, 6690 Da for XD, and 14,300 Da for lysozyme. Protein concentrations of 0.1 mg/mL were used when recording spectra of both individual and protein mixtures. In the case of protein mixtures, mean ellipticity values per residue were calculated as $[\theta] = \frac{3300 \Delta A}{l \cdot c \cdot n_1 \cdot m_1 + l \cdot c_2 \cdot n_2 \cdot m_2}$, where $l$ (path length) = 0.1 cm, $n_1$ or $n_2$ = number of residues, $m_1$ or $m_2$ = molecular mass in daltons, and $c_1$ or $c_2$ = protein concentration expressed in mg/mL for each of the two proteins in the mixture. The theoretical average ellipticity values per residue ($[\theta]_{AVE}$) assuming that neither unstructured-to-structured transitions nor major structure rearrangements occur, were calculated as follows: $[\theta]_{AVE} = \frac{([\theta]_1 \cdot n_1) + ([\theta]_2 \cdot n_2 \cdot R)}{(n_1 + n_2 \cdot R)}$, where $[\theta]_1$ and $[\theta]_2$ correspond to the measured mean ellipticity values per residue, $n_1$ and $n_2$ to the number of residues for each of the two proteins, and $R$ to the excess molar ratio of protein 2. The $\alpha$-helical content was derived from the ellipticity at 220 nm as described in Morris et al. (1999).

Fluorescence spectroscopy

Fluorescence intensity variations of the single tryptophan in N_{TAILHNC} was measured by using a CarlEclipse (Varian) equipped with a front-face fluorescence accessory at 20°C, by using 2.5 nm excitation and 10 nm emission bandwidths. The excitation wavelength was 290 nm and the emission spectra were recorded between 300 nm and 540 nm. Titrations were performed in a 1-mL quartz fluorescence cuvette containing 1 µM N_{TAILHNC} in 10 mM sodium phosphate buffer at pH 7, and by gradually increasing the XD concentration from 0.05 µM to 3 µM. Experimental fluorescence intensities were corrected by subtracting the spectrum obtained with XD alone (note that XD is devoid of tryptophan residues). Data were analyzed by plotting the relative fluorescence intensities at the maximum of emission at increasing XD concentrations. The dissociation equilibrium constant ($K_{Dapp}$) value was determined from data fitted to a single exponential equation, by using the PRISM 3.02 nonlinear regression tool (GraphPad).

Two-dimensional heteronuclear magnetic resonance

2D-HSQC spectra (Mori et al. 1995) were recorded on a 500-MHz DRX Bruker spectrometer either on 0.5 mM uniformly $^{15}$N-labeled N_{TAILHNC} or on 0.125 mM uniformly $^{13}$C-labeled N_{TAIL3} in 10 mM sodium phosphate buffer at pH 7.0 containing 10% D_{2}O (v/v) alone or after addition of a twofold molar excess of XD. The temperature was set to 283 K and the spectra were recorded with 2048 complex points in the directly acquired dimension and 256 points in the indirectly detected dimension, for 6 h each. Solvent suppression was achieved by the WATERGATE 3-9-19 pulse (Piotto et al. 1992). The data were processed using the UXNMR software; they were multiplied by a sine-squared bell and zero-filled to 1 K in first dimension prior to Fourier transformation. For the spectra recorded after addition of XD, the number of scans was multiplied by 4.

Sequence analysis and secondary structure predictions

The accession number of MV N is P04851. Secondary structure predictions were carried out using both PSI-PRED (McGuffin et al. 2000) and PHD (Rost 1996).

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