

Solution structure of the two RNA recognition motifs of hnRNP A1 using segmental isotope labeling: how the relative orientation between RRM influences the nucleic acid binding topology

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Abstract Human hnRNP A1 is a multi-functional protein involved in many aspects of nucleic-acid processing such as alternative splicing, micro-RNA biogenesis, nucleocytoplasmic mRNA transport and telomere biogenesis and maintenance. The N-terminal region of hnRNP A1, also named unwinding protein 1 (UP1), is composed of two closely related RNA recognition motifs (RRM), and is followed by a C-terminal glycine rich region. Although crystal structures of UP1 revealed inter-domain interactions between RRM1 and RRM2 in both the free and bound form of UP1, these interactions have never been established in solution. Moreover, the relative orientation of hnRNP A1 RRM is different in the free and bound crystal structures of UP1, raising the question of the biological significance of this domain movement. In the present study, we have used NMR spectroscopy in combination with segmental isotope labeling techniques to carefully analyze the inter-RRM contacts present in solution and subsequently determine the structure of UP1 in solution. Our data unambiguously demonstrate that hnRNP A1 RRM interact in solution, and surprisingly, the relative orientation of the two RRM observed in solution is different from the one found in the crystal structure of free UP1 and rather resembles the one observed in the nucleic-acid bound form of the protein. This strongly supports the idea that the two

RRMs of hnRNP A1 have a single defined relative orientation which is the conformation previously observed in the bound form and now observed in solution using NMR. It is likely that the conformation in the crystal structure of the free form is a less stable form induced by crystal contacts. Importantly, the relative orientation of the RRM in proteins containing multiple-RRMs strongly influences the RNA binding topologies that are practically accessible to these proteins. Indeed, RRM domains are asymmetric binding platforms contacting single-stranded nucleic acids in a single defined orientation. Therefore, the path of the nucleic acid molecule on the multiple RRM domains is strongly dependent on whether the RRM are interacting with each other. The different nucleic acid recognition modes by multiple-RRM domains are briefly reviewed and analyzed on the basis of the current structural information.

Keywords Segmental isotope labeling · Expressed protein ligation · Intein · Structural biology · HnRNP A1 · UP1 · RRM · NMR · Inter-domain interaction · ¹³C-edited half-filter NOESY · Domain orientation · RNA recognition modes

Abbreviations

GB1	B1 domain of streptococcal protein G
hnRNP	Heterogeneous nuclear ribonucleoprotein
HSQC	Heteronuclear single quantum coherence
MESNA	Sodium 2-mercaptoethanesulfonate
<i>Mxe</i> GyrA	GyrA gene from <i>Mycobacterium xenopi</i>
NOE	Nuclear Overhauser effect
NOESY	Nuclear Overhauser effect spectroscopy
PABP	Polyadenylate binding protein
r.m.s.d.	Root-mean-square deviation
RRM	RNA recognition motif
UP1	Unwinding protein 1

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Introduction

Eukaryotic mRNAs are transcribed as precursors (pre-mRNAs) containing intervening sequences (introns) that are subsequently removed such that the flanking regions (exons) are spliced together to form mature mRNAs. In addition to constitutive splicing, alternative splicing generates different mRNAs encoding distinct proteins, and hence increases protein diversity (Nilsen and Graveley 2010). For efficient splicing, most introns require a conserved 5' splice site, a branch point sequence followed by a polypyrimidine tract and a 3' splice site. In addition, other signal sequences along alternatively spliced exons, or their flanking introns are targeted by two large families of proteins that finely regulate alternative splicing: the SR-protein family (serine-arginine protein family) and the hnRNP family (heterogeneous nuclear RiboNucleoProtein family).

The hnRNP protein family consists of at least 20 proteins in humans that have been characterized as components of protein complexes bound to pre-mRNA (hnRNP complexes) (Dreyfuss et al. 1993). Most proteins of the hnRNP family contain at least one RRM domain (RNA recognition motif) (Maris et al. 2005), from one RRM in hnRNP C up to four RRMs in hnRNP I (also called Polypyrimidine Tract Binding protein, PTB). They also contain additional domains, like glycine-rich domains or aspartate-glutamate-rich domains that either contribute to RNA recognition and/or mediate protein/protein interaction. Among this family, hnRNP A1 is one of the most abundant and best-characterized components of hnRNP complexes. It is well established that the multi-functional hnRNP A1 protein plays an active role not only in alternative pre-mRNA splicing (Mayeda and Krainer 1992; Cáceres et al. 1994; Yang et al. 1994), but also in the maturation of some micro-RNA precursors (Guil and Cáceres 2007; Michlewski et al. 2008; Michlewski and Cáceres 2010), in nucleo-cytoplasmic mRNA transport (Piñol-Roma and Dreyfuss 1992), in promoting RNA strand annealing (Pontius and Berg 1990), and in telomere biogenesis and maintenance (LaBranche et al. 1998; Zhang et al. 2006; Flynn et al. 2011).

Human hnRNP A1 is a 320-amino-acid protein composed of two closely related RRM domains in its N-terminal region followed by a highly flexible glycine-rich C-terminal region (45 % of glycine). The N-terminal region, which includes RRM1 and RRM2 and spans residues 1 to 196, is also known as unwinding protein 1 (UP1). Interestingly, the two RRMs of hnRNP A1 are neither redundant nor functionally equivalent, in spite of their similar sequences and overall structure. Indeed, chimeric protein construction by duplication, deletion or swap of the RRMs differently affects the hnRNP A1 alternative splicing function (Mayeda et al. 1998). The relative position of

the two RRMs is therefore crucial for the alternative splicing activity of hnRNP A1.

To date, several high-resolution crystal structures of the two tandem RRMs of hnRNP A1 have been solved both in their free form and bound to repeats of telomeric DNA fragments. Three structures of free UP1 with resolution ranging from 1.1 to 1.9 Å have been refined in the P2₁ space group from an identical monoclinic crystal form (pdb accession codes 1UP1, 1HA1 and 1L3K) (Xu et al. 1997; Shamoo et al. 1997; Vitali et al. 2002). These three structures are almost indistinguishable with an average pairwise root-mean-square deviation (r.m.s.d.) for backbone atoms of 0.21 ± 0.04 Å. In these structures, the two RRMs are interacting with one another, mainly via two Arg-Asp salt bridges, and hence adopting a single relative orientation. The two RRMs are oriented in an anti-parallel manner, meaning that the two RNA binding surfaces are discontinuous and could bind to RNA strands having opposite 5'–3' polarity. It is important to notice that this fixed relative orientation of the RRMs strongly influences the repertoire of RNA binding topologies that can be formed with hnRNP A1. These different accessible topologies have been discussed previously (Xu et al. 1997; Shamoo et al. 1997). However, since the inter domain interaction surface is relatively modest (~ 630 Å²) and the residues involved in the Arg-Asp salt bridges are not absolutely conserved in the hnRNP A1-like proteins, the possibility that the association between RRM1 and RRM2 might be the result of crystal packing forces has been pointed out (Shamoo et al. 1997).

In addition to these free structures, eleven structures of UP1 bound to wild-type or diverse mutated repeats of telomeric DNA fragments have been solved with resolution ranging from 1.8 to 2.6 Å. All these structures have been refined in the P4₃2₁2 space group from an identical tetragonal crystal form (pdb accession codes 2UP1, 1PGZ, 1PO6 and 1U1K to 1U1R) (Ding et al. 1999; Myers et al. 2003; Myers and Shamoo 2004). These bound structures are almost indistinguishable with a calculated average pairwise r.m.s.d. for protein backbone atoms of 0.19 ± 0.04 Å. Interestingly, the overall interface between RRM1 and RRM2 as observed in this different crystal form is globally conserved as compared with the interface in the free form of the protein. For instance, the two Arg-Asp salt-bridges and other important contacts are similarly present in the structure of the telomeric DNA bound form. However, the relative orientation of the two RRMs is significantly altered and the free and bound structures do not perfectly overlay. Indeed, overall backbone r.m.s.d. between the free and bound UP1 structures is as high as 1.70 Å whereas individual RRMs display a much better agreement (0.32 and 0.45 Å for RRM1 and RRM2, respectively). These significant conformational changes

correspond to a rotation of $\sim 15^\circ$ of one RRM compare to the other (Ding et al. 1999). This reorientation of the RRMs has been attributed to the binding to the DNA substrate (Ding et al. 1999). However, one could also imagine that this relative movement of the RRMs is due to differences in the protein–protein contacts with neighbouring proteins in the two different crystal lattices (*i.e.* monoclinic and tetragonal crystals). In any case, the interactions at the inter-RRM interface in UP1 are probably quite weak, since the orientation of the two RRMs can be influenced by nucleic acid binding or by contacts with neighbouring molecules in the crystal lattice. Together with the aforementioned doubt brought up with the analysis of the free crystal structure (Shamoo et al. 1997), it directly raises the question whether the contacts between the two RRMs are present at all in solution. In addition, several biochemical studies performed in solution have been favouring a model of two independent RRMs connected by a flexible inter-RRM linker (Casas-Finet et al. 1991; Shamoo et al. 1994, 1995). In conclusion, there is to date no direct and unequivocal evidence of the existence, in solution, of an inter-RRM interaction in hnRNP A1.

Importantly, the relative orientation of different RRMs in proteins containing multiple-RRMs strongly influences the modes of RNA binding that are practically accessible to these particular proteins. Indeed, RRMs are asymmetric binding platforms contacting single-stranded nucleic acids in a single defined orientation, namely the 5' extremity towards β -strand 4 and the 3' towards β -strand 2 (Maris et al. 2005) (Supplementary Figure 1). Therefore, RRMs forming a discontinuous and anti-parallel platform may induce a looping in the nucleic acid target, as observed in the case of PTB RRM34 (Oberstrass et al. 2005; Lamichhane et al. 2010); and RRMs interacting to form a continuous binding platform can bind to longer stretches of nucleotides as seen in the structure of the polyA-binding protein (Deo et al. 1999) (See Supplementary Figure 1 for schematic illustrations of these cases). To date, structural information on the spatial organization of multiple RRMs in proteins containing at least two RRMs is still quite limited, since the structures of only a dozen of proteins with multiple RRMs have been solved in their free and/or nucleic acid bound form. These structures revealed that a limited number of distinct situations are actually occurring and exploited to achieve distinct biological functions. These features will be presented and discussed in a latter paragraph of this article, on the basis of the available structures.

In the present study, we have used segmental isotope labeling to determine whether the two RRMs of hnRNP A1 are interacting in solution. Segmental isotope labeling is a very attractive technique to reduce the complexity of NMR spectra with a large number of potential applications for the

study of large and/or multi-domain proteins (Skrisovska et al. 2010). Different methods, known as native chemical ligation (NCL), expressed protein ligation (EPL), and protein trans-splicing (PTS), are available for segmental isotope labeling of proteins (David et al. 2004; Muralidharan and Muir 2006). To date, isotope segmental labeling has not been extensively applied in NMR, although different studies have already demonstrated that segmental labeling is a very elegant and relevant approach to investigate large proteins (Yagi et al. 2004; Minato et al. 2012), to study conformational changes and ligand binding (Anderson et al. 2005), to investigate inter domain interactions within multi-domain proteins (Camarero et al. 2002; Zhang et al. 2007), and also to enable precise protein structure determination of multi-domain proteins (Vitali et al. 2006; Chen and Wang 2011; Chen et al. 2011).

In this work, we have determined the solution structure of the two RRMs of hnRNP A1 using a segmental labeling strategy in order to clearly determine whether these two RRMs are truly interacting in solution. This approach enabled us to unambiguously identify inter domain NOEs between RRM1 and RRM2 of hnRNP A1 and to calculate a precise overall structure. In our solution structure of free UP1, the two Arg-Asp salt bridges are conserved at the interface between RRM1 and RRM2, but surprisingly, the relative orientation of the two RRMs is quite different from the one found in the crystal structure of free UP1 but resembles the one observed in the nucleic-acid bound form of the protein.

Materials and methods

Cloning, expression and purification of hnRNP A1 RRM12 (UP1)

The DNA sequence encoding the two RRMs of hnRNP A1 (residues 2–196) (Uniprot entry P09651), were sub-cloned by PCR amplification from pET9d-hnRNPA1 (Mayeda and Krainer 1992) between *Bam*HI and *Xho*I cloning sites in *Escherichia coli* expression vector pET28a. The construct contains a N-terminal tag whose sequence MGSSHHHHHSSGLVPRGSHMENLYFQGG includes a 6 histidine stretch used for protein purification and a TEV-protease cleavage site used for subsequent removal of the purification tag. Proteins were overexpressed in BL21(DE3) codon-plus (RIL) cells in either LB media or M9 minimal media supplemented with $^{15}\text{NH}_4\text{Cl}$ and ^{13}C -labeled glucose. The cells were grown at 37 °C to OD600 ~ 0.4 , cooled down at 30 °C and induced at OD600 ~ 0.6 by adding isopropyl- β -D-thiogalactopyranoside to a final concentration of 0.5 mM. Cells were harvested 15 h after induction by centrifugation. Cell pellets were resuspended

in lysis buffer (Tris–HCl pH 8.0 50 mM, NaCl 1 M, EDTA 1 mM, DTT 1 mM) and lysed by sonication. Cell lysates were centrifuged 40 min at 45,000g. Supernatant was loaded on a Ni–NTA column on a ÄKTA Prime purification system (Amersham Biosciences), and the protein of interest was eluted with an imidazole gradient. The fractions containing the protein were pooled, TEV protease was added at a TEV/UP1 ratio of 1/200 (w/w), and specific cleavage of the purification tag was performed at room temperature for 16 h. His-tagged TEV protease and purification tag were further separated from the protein of interest with a Ni–NTA column. The fractions containing the protein were pooled, dialyzed against the NMR Buffer (NaPi pH 6.5 10 mM, DTT 1 mM), and concentrated to ~1.2 mM with a Vivaspin 10,000 MWCO (Sartorius Stedim Biotech).

Cloning, expression and purification of isolated RRM1 and RRM2 of hnRNP A1

The DNA sequence encoding the two individual RRMs of hnRNP A1 (*i.e.* RRM1 from residues 2–97 and RRM2 from residues 95–196), were sub-cloned by PCR amplification from pET9d-hnRNPA1 between *NheI* and *BamHI* cloning sites in *E. coli* expression vector pET28a. The constructs contain a N-terminal tag whose sequence MGSSHHH HHHSSGLVPRGSHMA includes a 6 histidine stretch used for protein purification. Proteins were overexpressed and purified as for the UP1 construct, except that the protein purification tag was not cleaved off by TEV-protease.

Cloning, expression and purification of $^{13}\text{C}/^{15}\text{N}$ -labeled RRM1 construct for protein ligation

The general approach to produce the segmentally labeled RRM12 sample is based on previously published protocols in use in our group (Vitali et al. 2006; Skrisovska and Allain 2008; Michel et al. 2013). The DNA sequence encoding RRM1 (residues 2–94) was sub-cloned from pET9d-hnRNPA1 between *NcoI* and *SapI* cloning sites of *E. coli* expression vector pEM9B (Michel et al. 2013) encoding a C-terminal fusion of the *Mxe* GyrA intein. The *SapI* restriction site naturally present in the pET9d-hnRNPA1 sequence had to be removed with a silent mutation K16K (AAG to AAA) prior to sub-cloning into pEM9B. A minimal sequence modification D94M was introduced into the inter-RRM linker to allow for efficient self-cleavage of the *Mxe* GyrA intein (Southworth et al. 1999). The protein construct was overexpressed with IPTG induction in BL21(DE3) codon-plus (RIL) cells in M9 minimal media supplemented with $^{15}\text{NH}_4\text{Cl}$ and ^{13}C -labeled glucose at 30 °C for 16 h. The intein fusion construct was purified with a Ni–NTA column on a ÄKTA Prime purification system, dialyzed against inactive

reaction buffer (Tris–HCl pH 8.0 50 mM, NaCl 200 mM, EDTA 1 mM, 2-mercaptoethanol 1 mM) and concentrated to 0.25 mM with a Vivaspin 10,000 MWCO.

Cloning, expression and purification of non-labeled RRM2 construct for protein ligation

The DNA sequence encoding RRM2 (residues 95–196) was sub-cloned from pET9d-hnRNPA1 between *NcoI* and *BamHI* cloning sites of *E. coli* expression vector pEM5B (Michel et al. 2013). The required S95C mutation and the preceding TEV-protease cleavage site were introduced with the primers during PCR amplification. The protein construct was overexpressed in LB media, purified using the same procedure as for the RRM1 intein-fusion construct, and concentrated to 0.5 mM in the inactive reaction buffer.

Intein cleavage, protein ligation and ligation product purification

Purified protein samples bearing $^{13}\text{C}/^{15}\text{N}$ -labeled RRM1 and unlabeled RRM2 were mixed with a two times excess of the unlabeled construct, and the reaction was activated by adding 100 mM sodium 2-mercaptoethanesulfonate (MESNA) and TEV-protease at a TEV/protein ratio of 1/200 (w/w). The reaction mixture was incubated at 35 °C for 48 h. The efficiency of the ligation reaction was analysed on SDS-PAGE. Purification and solubilisation tags released by TEV-cleavage and the intein protein released by self-cleavage during the ligation reaction are either retained on a Ni–NTA column or on a chitin column (see (Michel et al. 2013) for details). The reaction mixture was therefore applied on a Ni–NTA column and thereafter on a chitin column. The flow-through was then loaded on a SP-Sepharose column equilibrated with Tris–HCl pH 7.0 25 mM to separate the desired ligated product (RRM12, pI = 7.9) from the unreacted RRMs (*i.e.* RRM1 and RRM2, pI = 7.0 and 8.0, respectively). Bound proteins were eluted with a 0–400 mM NaCl gradient, fractions containing the desired ligated RRM12 were pooled and purity was evaluated by SDS-PAGE. At this stage the sample still contain about 10 % of unligated RRM2 that was further removed by applying the sample at 1 mL/min on a Superdex 26/60 HiLoad Prep Grade column (GE Healthcare) equilibrated with NMR buffer. Purity of the final sample was evaluated by SDS-PAGE to be >95 %. The sample was concentrated to 1.0 mM in 250 μL with a Vivaspin 10,000 MWCO.

NMR spectroscopy

All NMR spectra were recorded at 303 K on Bruker AVI-II-500 MHz, AVIII-600 MHz, AVIII-700 MHz, AVIII-750

MHz and Avance-900 MHz spectrometers (all equipped with a cryoprobe except for AVIII-750). The data were processed using TOPSPIN 2.1 (Bruker) and analyzed with Sparky (Goddard and Kneller 2006). Protein resonances were assigned with 2D (^1H , ^{15}N)-HSQC, 2D (^1H , ^{13}C)-HSQC, 3D HNCA, 3D HNCACB, 3D CBCA(CO)NH, 3D HNCO, 3D HN(CA)CO, 3D [^{13}C ; ^{15}N ; ^1H] HCC(CO)NH-TOCSY, 3D [^1H ; ^{15}N ; ^1H] HCC(CO)NH-TOCSY, 3D NOESY- (^1H , ^{15}N)-HSQC and two 3D NOESY- (^1H , ^{13}C)-HSQC optimized for the observation of protons attached to aliphatic carbons and to aromatic carbons, respectively. In addition, the assignment of aromatic protons was conducted using 2D (^1H , ^1H)-TOCSY and 2D (^1H , ^1H)-NOESY measured in D_2O ; histidine protonation and tautomeric form were determined from a long-range (^1H , ^{15}N)-HSQC spectrum (Pelton et al. 1993). Our assignment of RRM1 agrees for most resonances with previously published assignment (Garrett et al. 1994), yet ours is more complete. We recorded all 3D NOESY spectra with a mixing time of 100 ms and the 2D NOESY spectra with a mixing time of 60 ms. The assignment of inter domain NOEs was based on a 3D ^{13}C F1-edited, F3-filtered NOESY-HSQC spectrum ($\tau_m = 150$ ms) (Zwahlen et al. 1997) measured in D_2O on the RRM12 segmentally labeled sample with only RRM1 ^{13}C -labeled.

We measured NH RDCs from in-phase/anti-phase (^1H , ^{15}N)-HSQC experiments (Cordier et al. 1999), by comparing the peak positions of the up-field and down-field components measured in isotropic solution and in a dilute liquid crystalline phase. The alignment medium used for RDC measurements contained a mixture of *n*-dodecyl-penta(ethylene glycol) and *n*-hexanol (3 % C12E5/hexanol, $r = 0.96$) dissolved in 90 % $\text{H}_2\text{O}/10$ % D_2O (Rückert and Otting 2000). Under these conditions, a quadrupolar splitting of 25.9 Hz was observed in the ^2H spectrum.

Protein structure calculation

Automated NOE cross-peak assignments (Herrmann et al. 2002a) and structure calculations with torsion-angle dynamics (Guntert et al. 1997) were performed with the macro noeassign of the software package CYANA 3.0 (Guntert 2004). Unassigned peak lists of the four NOESY spectra were generated as input with the program ATNOS (Herrmann et al. 2002b) and manually cleaned to remove artefact peaks. In addition, a manually assigned peak list corresponding to the 3D ^{13}C -edited half-filter NOESY was also added to account for inter domain NOE measured with the segmentally labeled sample. The interdomain NOE were not manually converted into distance constraints. Instead, peak intensities were automatically calibrated and converted to distance constraints by CYANA with an optimized average-distance-parameter accounting for the presence of only long-range NOE in this particular peak-

list. The input also contained 64 hydrogen-bond restraints and 336 backbone dihedral restraints based on the chemical shift information from the program TALOS+ (Shen et al. 2009). Hydrogen bonded amides were identified as slowly exchanging protons in presence of D_2O . Their bonding partner was identified from preliminary structure calculations performed with only NOESY spectra as input. We calculated 100 independent structures that we refined in a water shell with the program CNS 1.3 (Brunger et al. 1998; Brunger 2007) including distance restraints from NOE data, hydrogen-bonds restraints, backbone dihedral restraints from TALOS+ and ^{15}N - ^1H RDC restraints as previously described (Barraud et al. 2011). The 20 best energy structures were analyzed with PROCHECK-NMR (Laskowski et al. 1996) and the iCING web server (Doreleijers et al. 2012) (<http://nmr.cmbi.ru.nl/icing/>). Overall structural statistics of the final water-refined structure are shown in Table 1. Structures were visualized and figures were prepared with program PYMOL (<http://www.pymol.org>).

Analysis of the structures

Backbone r.m.s.d. between the different UP1 structures and solvent accessible surface areas for the evaluation of the interaction surfaces between interacting RRM1s, were calculated with program superpose and areaimol of the CCP4 program suite (Winn et al. 2011). For the analysis of the ^{15}N - ^1H RDCs, proton atoms were added to the crystal structures with the program CNS (Brunger et al. 1998; Brunger 2007). RDC restraints were analysed with the program MODULE (Dosset et al. 2001), and back-calculated after best-fitting the alignment tensor to the different NMR and crystal structures with CYANA 3.0. Quality factors (Q) (Bax et al. 2001) and correlation coefficients (R) were also evaluated with CYANA 3.0 (Guntert 2004).

NMR dynamics

For the NMR dynamics study, ^{15}N T1 and T2 measurements were recorded at 303 K at a ^1H frequency of 500 MHz with established methods (Kay et al. 1989; Skelton et al. 1993). ^{15}N T1 values were derived from six (^1H , ^{15}N)-spectra with different delays: 100, 250, 500, 750, 1500 and 2,000 ms. Similarly, ^{15}N T2 values were derived from (^1H , ^{15}N)-spectra with six different delays: 12.5, 25, 50, 75, 100 and 125 ms. T1 and T2 values were extracted by a curve-fitting subroutine included in the program Sparky (Goddard and Kneller 2006). Overall correlation times (τ_c) were derived from T1/T2 ratio of dispersed and rigid amide resonances, assuming isotropic motion (Fushman et al. 1994).

Table 1 NMR experimental restraints and structural statistics

Distance restraints		
Total NOE	5,354	
Intra-residue	1,108	
Sequential	1,361	
Medium range ($ i - j < 5$ residues)	1,015	
Long range ($ i - j \geq 5$ residues)	1,806	
Interdomain NOE	64	
Hydrogen bonds	64	
Distance restraints violations (mean \pm SD)		
Number of NOE violations $> 0.2 \text{ \AA}$	3.1 \pm 1.0	
Maximum NOE violation (\AA)	0.27 \pm 0.04	
TALOS+ derived dihedral restraints		
ϕ	168	
ψ	168	
Dihedral violations (mean \pm SD)		
Number of dihedral violations $> 5^\circ$	0.55 \pm 0.51	
Maximum dihedral violation ($^\circ$)	3.5 \pm 3.3	
RDC restraints		
Number of ^{15}N - ^1H RDC restraints	101	
RRM1	56	
RRM2	45	
RDC violation (mean \pm SD)		
Number of RDC violations $> 2 \text{ Hz}$	6.0 \pm 2.3	
Maximum RDC violation (Hz)	3.1 \pm 0.3	
R.m.s.d. from average structure (\AA)		
Backbone ^a	0.71 \pm 0.16	
Heavy atoms ^a	1.22 \pm 0.22	
	RRM1 ^b	RRM2 ^c
Backbone	0.40 \pm 0.08	0.58 \pm 0.18
Heavy atoms	0.98 \pm 0.18	1.17 \pm 0.30
Deviation from ideal covalent geometry (mean \pm SD)		
Bond lengths (\AA)	0.0041 \pm 0.0001	
Bond angles ($^\circ$)	0.54 \pm 0.01	
Impropers ($^\circ$)	1.36 \pm 0.05	
Ramachandran analysis		
Most favored region	90.1 %	
Allowed region	9.8 %	
Disallowed region	0.1 %	
CING red/orange/green scores		
R/O/G (%)	12/31/57	

^a Protein r.m.s.d. was calculated using residues 11–89, 105–111, 117–139, 146–180 for the ensemble of 20 refined structures

^b RRM1 r.m.s.d. was calculated using residues 11–89

^c RRM2 r.m.s.d. was calculated using residues 105–111, 117–139, 146–180

Results

Initial chemical shift mapping between the individual domains and RRM12

In the context of investigating RNA binding specificity of each individual RRM of hnRNP A1 (manuscript in preparation), we produced ^{15}N -labeled NMR samples of isolated RRM1 and RRM2, as well as RRM12 (UP1). Surprisingly, ^1H , ^{15}N -HSQC spectrum of UP1 was virtually indistinguishable from the superposition of the two HSQC spectra coming from the isolated RRMs (Fig. 1a–d). Indeed, in the case of interacting RRMs within PTB (RRM3 and RRM4), marked differences between the ^1H , ^{15}N -HSQC footprint of the single RRMs and the one of the double-domain construct RRM34 were observed (Vitali et al. 2006). On the contrary, nearly identical ^1H , ^{15}N -HSQC spectra have been observed between isolated RRMs and their combined double-domain construct in the context of non-interacting RRMs, for example in the case of Npl3p RRM1 and RRM2 (Skrisovska and Allain 2008) and hnRNP F qRRM1 and qRRM2 (Dominguez and Allain 2006). Based solely on such considerations one might suggest that RRM1 and RRM2 of hnRNP A1 do not interact in solution contrary to what was observed in the crystal structures (Xu et al. 1997; Shamoo et al. 1997). However, after careful inspection of the overlays (Fig. 1c, d) we could identify a small number of peaks (*i.e.* one in RRM1 and three in RRM2) with small chemical shift perturbations between 0.1 and 0.15 ppm (Fig. 1e, f). Anyhow, puzzled by the very small extent of these chemical shift variations, we decided to determine the structure of UP1 in solution with NMR spectroscopy, and in order to unambiguously assess the existence of inter-domain contacts, if any, we prepared a segmentally labeled RRM12 sample with an expressed protein ligation approach.

Segmental isotope labeling of hnRNP A1 RRM12 by expressed protein ligation

Implementation of expressed protein ligation requires a reactive thioester at the C-terminus of RRM1 and a cysteine at the N-terminus of RRM2. Since the interdomain linker contains no natural cysteine, a cysteine was introduced by substituting serine 95 (S95C), and was thus taken as the N-terminus of RRM2. In addition, the preceding aspartate residue (D94) was mutated to methionine (D94M) to allow for efficient self-cleavage of the *Mxe* GyrA intein (Southworth et al. 1999). The resulting amino-acid sequence at the desired ligation site was then ${}_{91}\text{SREMCQRP}_{98}$. The reactive C-terminal thioester was obtained with RRM1 fused N-terminally to the *Mxe* GyrA

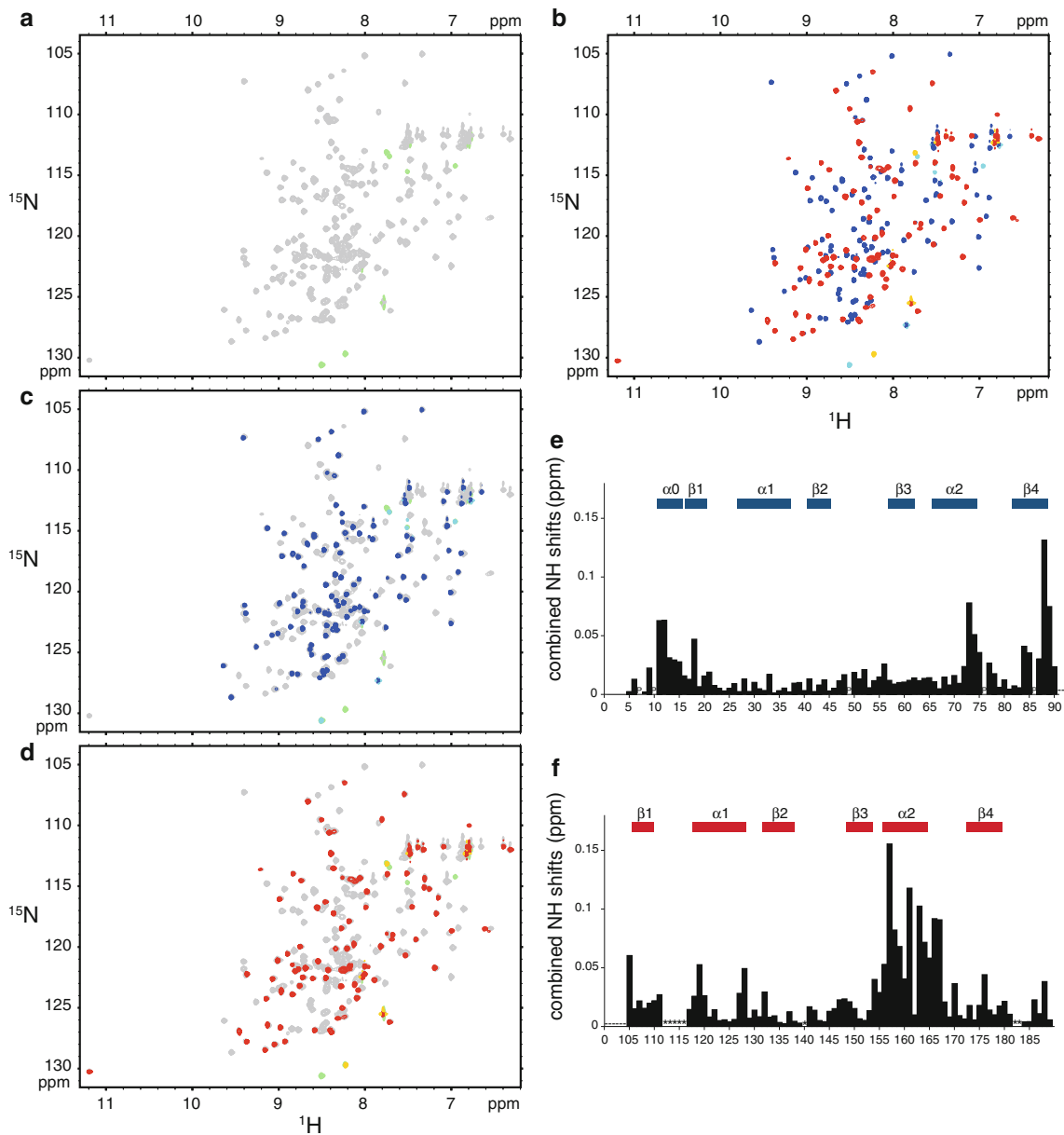


Fig. 1 NMR footprint of UP1 and single RRM1 and RRM2 from hnRNP A1. **a** ($^{15}\text{N}, ^1\text{H}$)-HSQC spectrum of UP1 (residues 2–196). Positive signals are *in grey*, and negative signals from aliased peaks are *in green*. **b** Overlay of two ($^{15}\text{N}, ^1\text{H}$)-HSQC spectra from single RRM1 (residues 2–97) *in deep blue* (positive signals) and *cyan* (negative signals) and from single RRM2 (residues 95–196) *in red* (positive signals) and *yellow* (negative signals). **c, d** Overlays of spectra to facilitate chemical shift comparison in UP1 and single RRM domains. Signals have same colours as in **a** and **b**. **c** Overlay of

two ($^{15}\text{N}, ^1\text{H}$)-HSQC spectra from single RRM1 (residues 2–97) and from UP1 (residues 2–196). **d** Overlay of two ($^{15}\text{N}, ^1\text{H}$)-HSQC spectra from single RRM2 (residues 95–196) and from UP1 (residues 2–196). **e, f** Backbone amide chemical shift difference between UP1 (spectrum of panel **a**) and isolated RRM1 **e** and RRM2 **f** (spectra of panel **c** and **d**). (P) corresponds to proline residues (-) to residues of the interdomain linker, and (*) to missing amide signals. Secondary structure elements are drawn above the histograms

intein and the N-terminal cysteine with a properly engineered TEV-protease cleavage site in front of RRM2 (Fig. 2a and “Materials and methods” for details). Each construct was expressed separately in *E. coli* allowing for different labeling scheme for each domain (namely $^{13}\text{C}/^{15}\text{N}$ -labeled for RRM1 and unlabeled for RRM2). Ligation was conducted at 35 °C for 48 h with a twofold

excess of the unlabeled RRM2 in order to increase the ligation efficiency with respect to the $^{13}\text{C}/^{15}\text{N}$ -labeled RRM1 domain. The RRM12 ligated construct was then purified from remaining unligated domains with ion exchange and size-exclusion chromatography (see “Materials and methods” and Supplementary Figure 2). We finally obtained ~6 mg of ligated RRM12 (from 2L of

culture in M9 medium) and could prepare a concentrated NMR sample (~ 1.0 mM in 250 μ L). This sample was well-folded and we could confirm with 1D ^1H NMR that the ligated UP1 adopts the same overall structure as the conventional recombinant UP1 protein (data not shown). This is further supported by the comparison of ^1H , ^{15}N -HSQC spectra of ligated UP1 and conventional recombinant UP1, where signals of RRM1 perfectly overlay in each spectra (compare Figs. 2b, 1c). This shows that mutations introduced for the ligation reaction in the linker region ($_{94}\text{DS}_{95}$ to $_{94}\text{MC}_{95}$) do not affect the structure of UP1.

We could therefore measure 3D ^{13}C F1-edited, F3-filtered NOESY-HSQC in order to unambiguously detect interdomain NOE crosspeaks, if true contacts exist between the two RRMs in solution. In this 3D NOESY spectrum, one could clearly see several interdomain NOEs in several cross-sections (Fig. 3). For example, clear contacts are

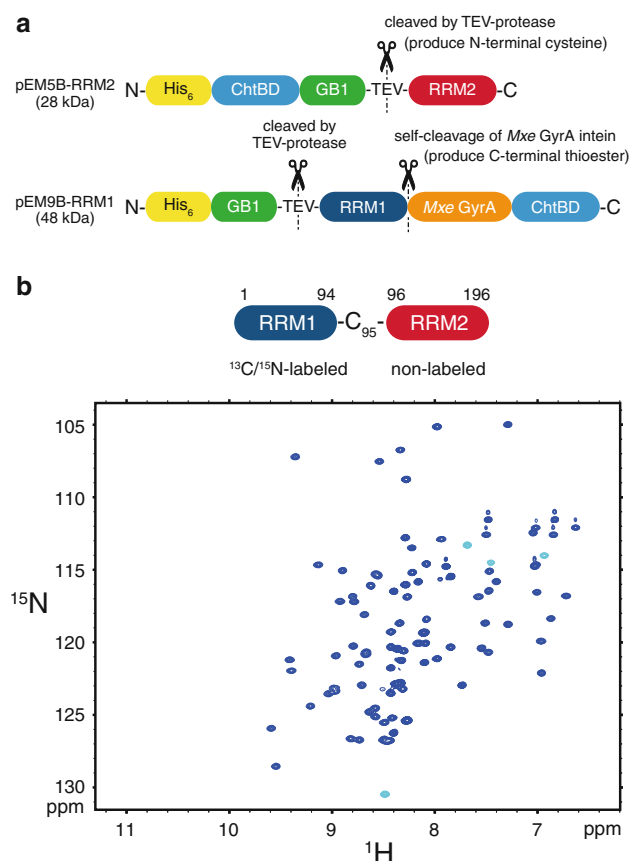


Fig. 2 Segmentally labeled hnRNP A1 RRM12 for NMR structural investigations. **a** Constructs used to generate the C-terminal fragment (RRM2) with an N-terminal cysteine and the N-terminal fragment (RRM1) with a C-terminal reactive thioester. See also the “[Material and method](#)” section and Supplementary Figure 2 for details on ligation reaction and product purification. **b** (^{15}N , ^1H)-HSQC spectrum of ligated RRM12 with $^{13}\text{C}/^{15}\text{N}$ -labeled RRM1 and unlabeled RRM2. Positive signals are in *deep blue*, and negative signals from aliased peaks are in *cyan*. See also Fig. 1 for a comparison of the (^{15}N , ^1H)-HSQC spectra

seen between the side-chains of Leu13 in RRM1 and Ile164 in RRM2 as well as between those of Met72 in RRM1 and His156 and Asp157 in RRM2. Also, multiple contacts are seen between Lys87 in RRM1 and His156 in RRM2 as well as multiple residues of the interdomain linker (*i.e.* His101, Leu102 and Thr103). In addition, multiple NOEs are seen between the side-chains of Val90 in RRM1 and of several residues in RRM2 (namely, Lys179, Asp160, Val163 and Ile164). We want here to briefly mention that some methyl–methyl contacts would be expected between aliphatic residues for which contacts are clearly seen between a methyl group on one hand and other types of aliphatic protons on the other hand (see for instance Leu13- $\text{H}\delta 1$ s NOEs to Ile164 $\text{H}\alpha$, $\text{H}\beta$ and $\text{H}\gamma 1$ s on Fig. 3a). However, it is difficult to unambiguously observe and assign these methyl–methyl contacts since they are often overlapping with the strong doublets of the diagonal peaks. For this reason, only few methyl–methyl inter domain NOE were confidently identified in the 3D ^{13}C -edited half-filter NOESY. Overall, we could unambiguously assign 64 interdomain NOEs that were thereafter converted into long-range inter-proton distances and used to precisely determine the structure of UP1 in solution. Importantly, this clearly demonstrates that RRM1 and RRM2 are truly interacting in solution.

Structure determination of hnRNP A1 RRM12

In order to precisely analyse the atomic details of the interdomain interface present in solution, and especially to compare this interface with the previously determined crystal structures, we solved the solution structure of hnRNP A1 RRM12 using NMR. A total of 5,354 distance constraints were derived from NOESY spectra. This includes 64 interdomain NOE that have been unambiguously assigned with the use of the segmentally labeled UP1 sample (Fig. 3). This large number of constraints allowed us to obtain a precise structure with a backbone r.m.s.d. over the entire domain of 0.71 ± 0.16 \AA for the ensemble of 20 conformers (Fig. 4a and Table 1). Constraints also include 64 hydrogen-bond restraints, backbone dihedral restraints derived from TALOS+ predictions for 168 residues and 101 ^{15}N – ^1H amide RDC restraints obtained from measurements in a partially oriented sample. Hydrogen bonded amides were identified as slowly exchanging protons in presence of D_2O . Their hydrogen-bond acceptors were identified from preliminary structures. Further details on the assignment and structure calculation procedures are reported in the “[Materials and methods](#)” section. NMR experimental constraints, refinement and structural statistics are presented in Table 1. In addition, as a further analysis of the quality of the structure, we determined the structure of the protein with the same calculation and

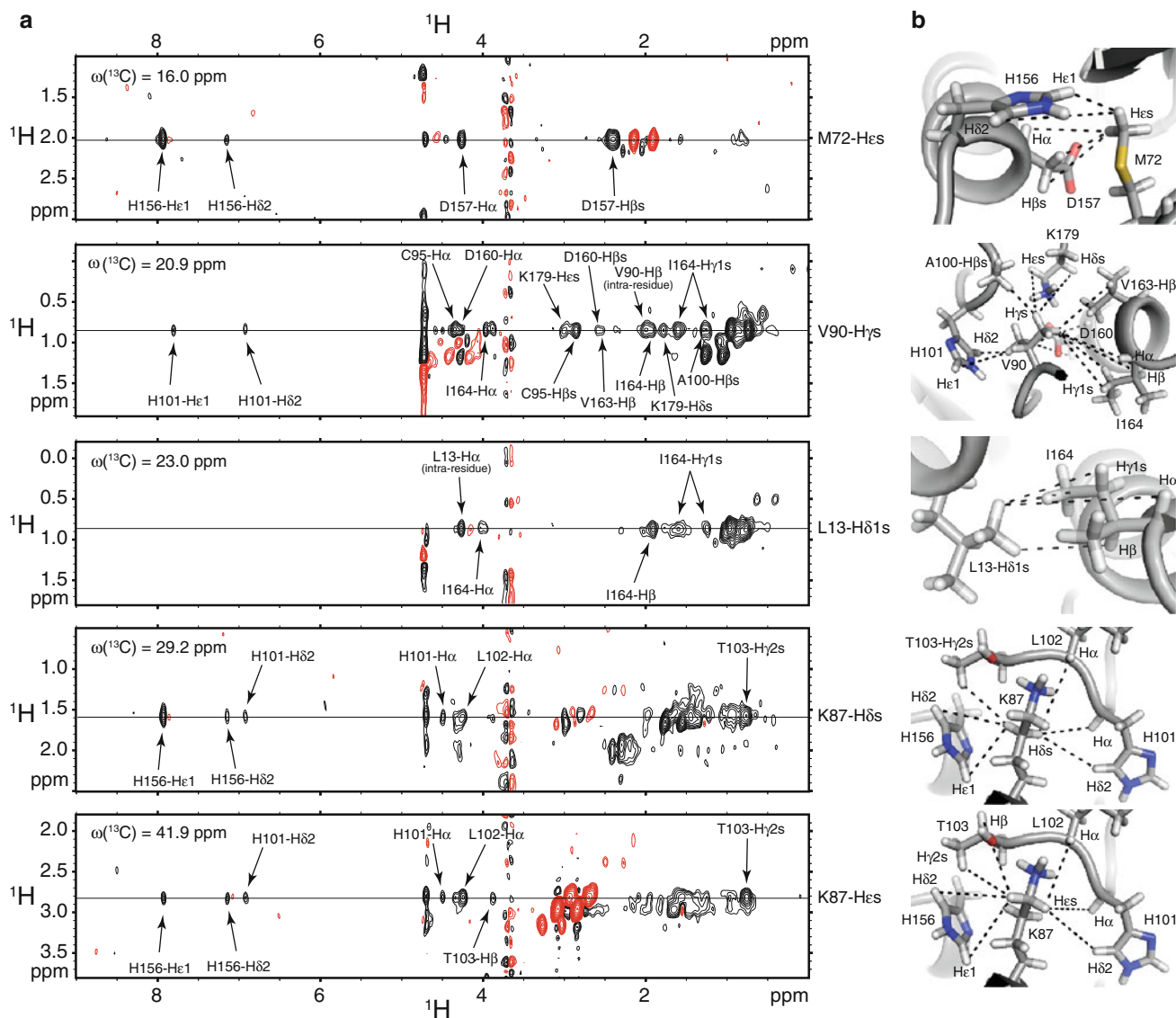


Fig. 3 Interdomain contacts as determined with segmentally labeled hnrNP A1 RRM12. **a** Series of cross-sections from the 3D ^{13}C -edited half-filtered NOESY measured on the segmentally labeled hnrNP A1 RRM12 with RRM1 $^{13}\text{C}/^{15}\text{N}$ labeled and RRM2 unlabeled. Assignment of interdomain NOEs are reported on the spectra and illustrated as dashed lines on the solution NMR structure (see panel **b**). ^{13}C

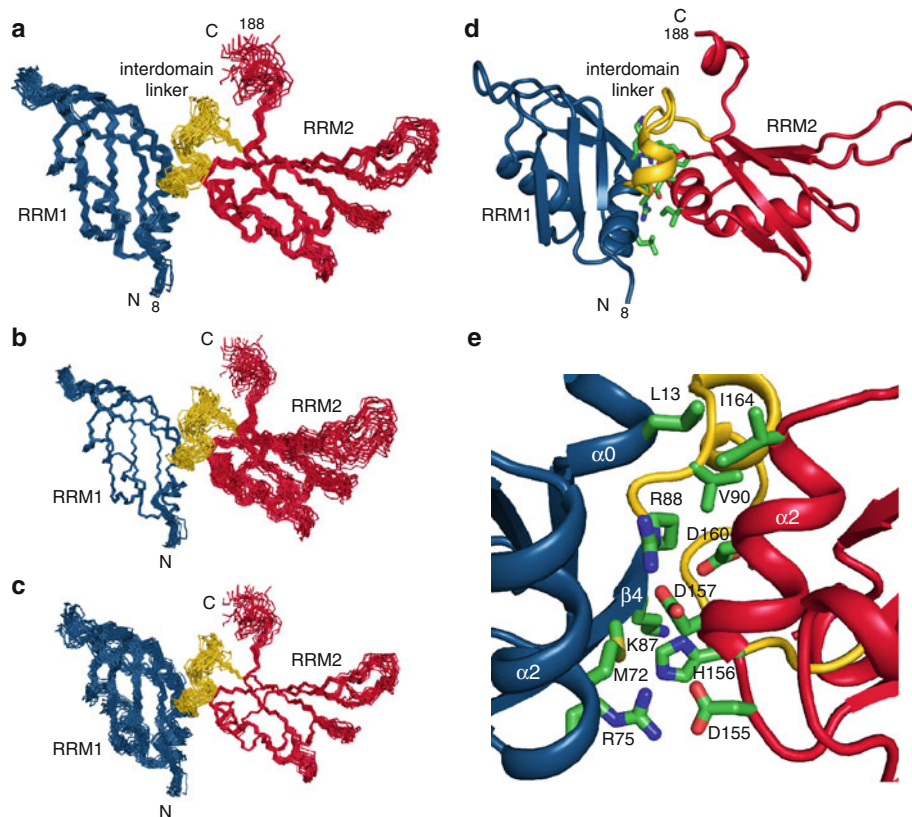
refinement protocols, but excluding the orientational information obtained from the ^{15}N - ^1H amide RDCs. We then evaluated the agreement between this NMR ensemble calculated without RDC restraints and the measured RDCs. We calculated an average Q factor (Bax et al. 2001) of 49.3 %, which is in the range of typical Q factors calculated for structures determined only from NOE information (see also Supplementary Table 1 and Supplementary Figure 3).

The structure ensemble together with dynamic information obtained from $^{15}\text{N}\{^1\text{H}\}$ -NOE values revealed that UP1 folds as two typical RRM domains separated by a

chemical shifts are indicated for each cross-section. Positive signals are in *black* and negative ones in *red*. Strips of signals at 4.72 and 3.68 ppm correspond to residual water and buffer signals, respectively. **b** Illustration of interdomain NOEs of each cross-section from panel **a** on the solution NMR structure. NOEs are represented as *dashed lines* between the corresponding protons or groups of protons

rather flexible interdomain linker. The N- and C-termini are also flexible as well as the long β 2- β 3 loops in each RRM, this character being more pronounced in RRM2 (see Fig. 4a and Supplementary Figure 4). Each individual RRM is very well defined, with average backbone r.m.s.d. of 0.40 and 0.58 Å for RRM1 and RRM2, respectively (Table 1). This value is slightly increased for the entire UP1 domain (0.71 ± 0.16 Å), as reflected on the three different overlays of Fig. 4a-c. Overall, the structure of each individual RRM is very similar to the previously determined crystallographic structures (Shamoo et al. 1997; Xu et al. 1997) but small local variations exist

Fig. 4 NMR solution structure of hnRNP A1 RRM12 **a–c** NMR ensemble. Overlay of the 20 final structures with RRM1 in *blue*, RRM2 in *red*, and the interdomain linker in *yellow*. Structures were overlaid over the entire protein (residues 8–188) (**a**), or over each individual RRM, namely residues 8–88 for RRM1 (**b**) and residues 106–188 for RRM2 (**c**). **d** Cartoon drawing of a representative structure of the NMR ensemble. Side chains involved in the interdomain interface are shown as sticks in *green*. **e** Close-up view of the interdomain interface. The interdomain interface involve residues, shown as sticks in *green*, from $\alpha 0$, $\alpha 2$ and $\beta 4$ in RRM1 and from $\alpha 2$ in RRM2. In addition, residues from the interdomain linker are also participating in the interface. See also Fig. 3b



between our solution NMR structure and crystal structures and will be described in details in the following sections.

Description of the interface between RRM1 and RRM2

The nature of the interface in our solution structure is at first sight similar to the interface observed in previous crystal structures (Shamoo et al. 1997; Xu et al. 1997; Ding et al. 1999). It involves residues from $\alpha 0$, $\alpha 2$ and $\beta 4$ in RRM1 and residues from $\alpha 2$ in RRM2 (Fig. 4e). Note that the regions of interaction perfectly match with the very small chemical shifts differences observed between UP1 and the isolated RRM1 and RRM2 (Fig. 1e, f). These interactions may be divided in three elements: (1) a small hydrophobic cluster involving Leu13, Ile164, Val90 and the aliphatic part of Arg88 side chain on one side of the interface (*top* of Fig. 4e); (2) two Arg-Asp salt-bridges, namely Arg88-Asp157 and Arg75-Asp155, in the middle and on the other side of the interface, respectively (*bottom* of Fig. 4e); and (3) a central residue, H156, which is sandwiched between the hydrophobic part of Lys87 on one side and the Arg75-Asp155 salt-bridge on the other side and also interacts with Met72 side chain (Fig. 4e). Most of these contacts led to a direct spectroscopic evidence in the 3D ^{13}C -edited half-filter NOESY (Fig. 3a, b). However, no direct spectroscopic information could be obtained on the two salt-bridges, as the closest observable protons across a

salt-bridge of this type, namely Arg-H δ s and Asp-H β s, are about 7–8 Å apart. The position of these side chains was therefore defined from electrostatic properties and NOEs to surrounding side-chains, and no additional restraints were included to force the formation of these salt-bridges. Nevertheless, each Arg-Asp salt-bridge is present in about 2/3 of the 20 structures of the final NMR ensemble, which strongly support their existence in solution, similarly to what was seen in the crystal structures.

Dynamical study of hnRNP A1 RRM1 and RRM2

In order to bring additional evidence that hnRNP A1 RRM1 and RRM2 are indeed interacting in solution, we wanted to evaluate this aspect using an independent method that would corroborate the experiments obtained with the segmental labeling approach. We therefore performed with NMR a dynamic study for both the single RRM1 and RRM2 constructs (*i.e.* RRM1 and RRM2) and the RRM12 double-domain construct (UP1). We measured for this three constructs, ^{15}N T1 and T2 relaxation times, as described in the “Materials and methods” section. Overall correlation times for each constructs (τ_c) were derived from T1/T2 ratio of dispersed and rigid amide resonances, assuming isotropic motion (Fushman et al. 1994). Overall rotational correlation times of 8.2 ± 0.4 and 10.9 ± 1.1 ns were obtained for RRM1 and RRM2 in isolation,

respectively, whereas τ_c increases significantly up to 15.8 ± 0.6 ns for the double-domain construct UP1 (Table 2). These values are in good agreement with reported values for domains of these sizes (*i.e.* 10.9, 11.2 and 22.1 kDa for RRM1, RRM2 and UP1, respectively) (Dayie et al. 1996) and definitely support that RRM1 and RRM2 are interacting in solution.

Comparison of the interdomain interface and of the relative orientation of the individual RRMs between the NMR solution structure and crystal structures of UP1

As reported in a previous paragraph, the different crystal structures of free UP1 are almost indistinguishable and this is also the case for the different crystal structures of UP1 bound to DNA. To simplify our analysis of the differences between our NMR structure and the different crystal structures, we decided to retain only one crystal structure for each form of the protein, free and bound, and to keep the most representative of each class, namely the ones that displayed the lowest backbone r.m.s.d to the other structures of their group. Interestingly, they correspond in both cases to structures with the highest resolution, *i.e.* 1.1 Å for free UP1 (pdb code 1L3K (Vitali et al. 2002)), and 1.8 Å for bound UP1 (pdb code 1U1R (Myers and Shamoo 2004)). Similarly, a representative structure of our NMR ensemble was chosen using the similar criterion, *i.e.* the lowest backbone r.m.s.d to the other structures of the ensemble, and remarkably it was also the lowest energy structure of our ensemble. These structures will be thereafter called UP1_{free}, UP1_{bound} and UP1_{NMR}.

The three structures have a very similar overall organization and can be superimposed on the entire UP1 domains with a relatively good agreement (Fig. 5a). However, it has already been mentioned that there is a significant conformational change between UP1_{free} and UP1_{bound} that corresponds to a rotation of $\sim 15^\circ$ of one RRM in respect to the other (Ding et al. 1999). In order to emphasize this conformational change between UP1_{free} and UP1_{bound}, we also superimposed the three structures on RRM1 only and could then visualize better the differences in the orientation of RRM2 (Fig. 5b, c). Surprisingly, it appeared very clearly that the relative orientation of the two RRMs in UP1_{NMR}, which is a free structure of UP1,

more closely resembles the crystal structure of UP1_{bound}, and less the one of UP1_{free} (Fig. 5b, c). To confirm numerically our visual observation, we calculated pairwise backbone r.m.s.d. between all the different RRMs and UP1s structures. These data are presented in Table 3. It confirmed that UP1_{NMR} is much closer to UP1_{bound} than to UP1_{free} (backbone r.m.s.d of 1.18 and 2.18 Å, respectively). Importantly, the comparison of the individual RRMs confirmed that these large differences are not due to local differences in the RRM themselves, but really to the different orientations adopted by the RRMs in the different structures (Table 3).

In order to strengthen this observation, we sought at finding unambiguous spectroscopic evidences that would demonstrate that these differences between UP1_{free} and UP1_{NMR} are not due to indirect effects of the structure calculation methodology, but really reflect true differences between solution and crystal structure. We first used the RDC information to evaluate the agreement between the measured NH RDCs and the two different crystal structures. We back-calculated the NH RDCs after best-fitting the alignment tensor to both crystal structures (UP1_{free} and UP1_{bound}), and compared them with the set of measured RDCs. Overall, UP1_{bound} agrees better with the measured RDCs than UP1_{free} (Q factors of 47.4 and 55.4 % for UP1_{bound} and UP1_{free}, respectively—Supplementary Table 1). This shows that the measured RDCs could discriminate, independently of any NMR structure determination, the subtle differences associated with the two different relative orientations of the RRMs in the two different crystal structures. This further supports our conclusion regarding the origin of the domain re-orientation between the two crystal structures. In addition, similarly to what we observed for the UP1_{NMR} structure refined with RDCs, the NMR structure calculated without RDC restraints more closely resembles UP1_{bound} than UP1_{free} and gives comparable RDC Q factor (Supplementary Table 1 and Supplementary Figure 5), indicating that the inter-domain NOE data obtained from the segmentally labeled sample would have been sufficient to notice the differences regarding the structural agreement of our NMR structure with UP1_{free} and UP1_{bound}. This means that the inter-domain NOE data should contain enough information to discriminate the two conformations. Therefore, we next analysed the inter-domain NOE cross peaks. Due to the geometrical property of this domain rotation, the N-terminal end of helix $\alpha 2$ in RRM2 is barely affected whereas its C-terminal end shows larger amplitude deviations (Fig. 5c). Interestingly, the distances between Val90 and Ile164 in UP1_{free} are large (between 8 and 11 Å, Fig. 5d) and therefore should not result in any NOE cross-peaks between these side-chains, whereas these two residues are much closer in the structures UP1_{bound} and UP1_{NMR} (Fig. 5d) potentially enabling NOE cross peaks to be measured. Such NOE cross peaks were indeed observed between Val90 and Ile164 in the 3D ¹³C-edited half-filter

Table 2 ¹⁵N T1, T2 and overall correlation time of hnRNP A1 RRM1, RRM2 and RRM12

Protein construct	T1 (ms)	T2 (ms)	T1/T2	τ_c (ns)
RRM1	500 ± 20	89 ± 8	5.6 ± 0.5	8.2 ± 0.4
RRM2	495 ± 30	55 ± 7	9.1 ± 1.8	10.9 ± 1.1
UP1	640 ± 30	36 ± 4	17.5 ± 1.2	15.8 ± 0.6

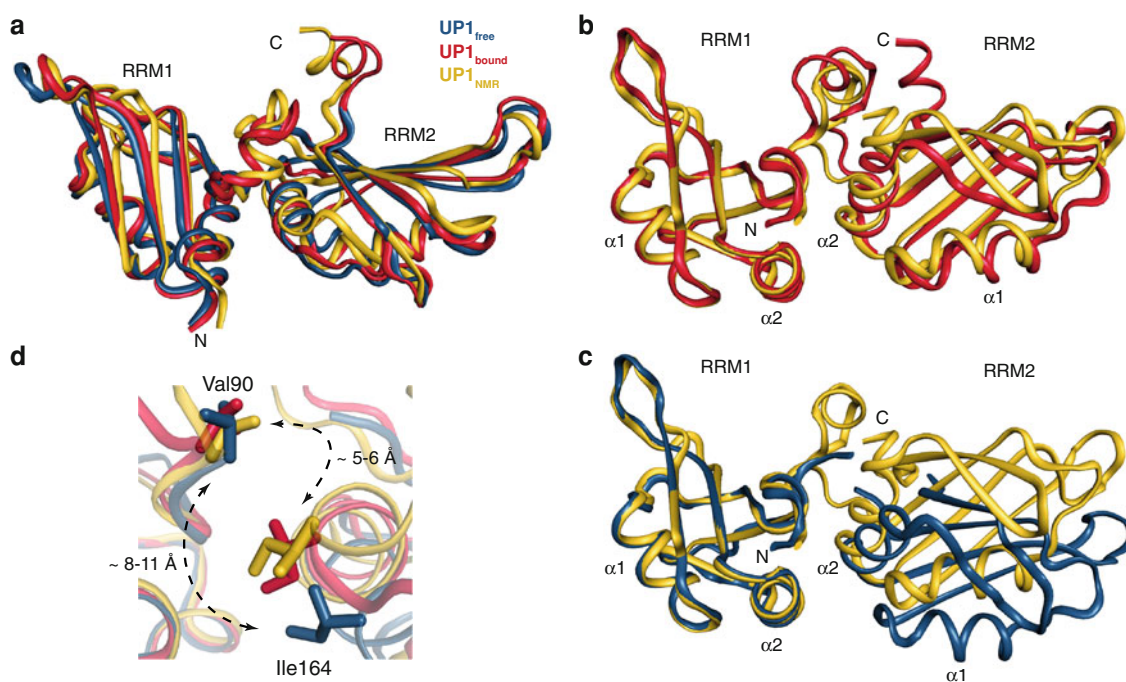


Fig. 5 Structural comparison of UP1 solution NMR structure with UP1 crystal structures. **a** Overlay of UP1_{free} in blue (*i.e.* free UP1 crystal structure, pdb code 1L3K), UP1_{bound} in red (*i.e.* DNA bound UP1 crystal structure, pdb code 1U1R), and UP1_{NMR} in yellow over the entire domain (residues 11–89 and 105–180). **b, c** Two different overlays of UP1_{NMR} in yellow with UP1_{bound} in red (**b**), and with UP1_{free} in blue (**c**). The structures are superimposed onto RRM1 (residues 11–89) to emphasize the differences in the relative orientation of the two RRMs in the different structures. The NMR structure overlays better with the UP1_{bound} structure, see for example

the large differences in the position of helices $\alpha 1$ and $\alpha 2$ in RRM2. See also Table 3 for a numerical comparison of the structures. **d** Close-up view of the large differences in the relative positions of residues Val90 and Ile164 between the different structures. The distances in UP1_{bound} and UP1_{NMR} are compatible with the observation of NOE transfer between these residues, whereas they are too far apart in UP1_{free} for efficient NOE transfer. See also Fig. 3 and Supplementary Figure 6 for an illustration of NOE transfers observed between these two residues

Table 3 Comparison of UP1_{free}, UP1_{bound} and UP1_{NMR} structures

Structures	Backbone r.m.s.d. (Å) ^a
UP1	
UP1 _{free} /UP1 _{bound}	1.70
UP1 _{free} /UP1 _{NMR}	2.18
UP1 _{bound} /UP1 _{NMR}	1.18
RRM1	
RRM1 _{free} /RRM1 _{bound}	0.32
RRM1 _{free} /RRM1 _{NMR}	0.55
RRM1 _{bound} /RRM1 _{NMR}	0.57
RRM2	
RRM2 _{free} /RRM2 _{bound}	0.45
RRM2 _{free} /RRM2 _{NMR}	0.79
RRM2 _{bound} /RRM2 _{NMR}	0.89

^a UP1 r.m.s.d. was calculated using residues 11–89 and 105–111, 117–139, 146–180; RRM1 r.m.s.d. was calculated using residues 11–89; RRM2 r.m.s.d. was calculated using residues 105–111, 117–139, 146–180

NOESY (Fig. 3 and Supplementary Figure 6), confirming that our NMR measurements are incompatible with the UP1_{free} crystal structure. This further emphasizes the

efficiency of the segmental labeling approach for the precise determination of multi-domain protein structures.

Altogether, our NMR structure of UP1 free revealed that RRM1 and RRM2 are truly interacting in solution, and that the relative orientation of these RRMs most closely resembles the one observed in the DNA bound form of UP1. Therefore, it strongly suggests that the two RRMs of hnRNP A1 have in solution a single defined relative orientation that is unchanged upon DNA binding, and that the relative domain movement observed between the two crystal forms of the protein is not a consequence of nucleic acid binding, but rather of differences in the protein–protein contacts between neighbouring proteins in the two different crystal lattices.

Discussion

In this study, we have used segmental isotope labeling to determine the solution structure of the two RRMs of hnRNP A1. This labeling strategy was crucial to unambiguously prove that hnRNP A1 RRMs interact in solution,

and this important point could not have been derived solely on the basis of chemical shift difference between the tandem construct and the isolated domains. In addition, the solution structure revealed that the difference in relative orientation of the two RRM, as observed in the two different crystal forms of the protein (free and bound to DNA), is very likely due to the different crystal lattices rather than due to the binding of nucleic acid. Although the two crystal structures and the NMR solution structures are very similar overall, some local differences exist and their importance will be discussed below. Additionally, this study established unequivocally that in solution, the two RRM of hnRNP A1 interact with each other in the free form of the protein. This does not represent the most common situation in proteins containing multiple RRM, and we propose to review hereafter how different arrangement between several RRM can influence nucleic acid recognition on the basis of the available multiple RRM structures, free and bound to nucleic acid.

Interacting domains and NMR spectroscopy

When we recorded the first NMR spectra of hnRNP A1 RRM12 construct (UP1) and of the isolated RRM, we were surprised to observe an almost perfect overlay for these two RRM although several crystal structures showed that the RRM could interact with each other (Shamoo et al. 1997; Xu et al. 1997). By experience, when ^1H , ^{15}N -HSQCs overlay almost perfectly, like observed for hnRNP A1 RRM, one often conclude that the two domains are independent domains (Skrisovska and Allain 2008; Dominguez and Allain 2006; Oberstrass et al. 2005). However, one can validate or not such conclusions using NMR relaxation measurements (^{15}N T1 and T2 relaxation times), and the case of hnRNP A1 prove the need of performing such additional analyses. Indeed, although marked differences between the NMR footprints of isolated and combined domains can be taken as a strong evidence of an interdomain interaction, one should not deduce from a virtually perfect overlay that the domains are independent. In such case, there is a real need for unambiguous methods like NMR relaxation measurements or segmental isotope labeling in combination with interdomain NOE measurements, in order to conclude about possible interdomain interactions. Furthermore, in this particular case of hnRNP A1 RRM12, automated structure calculations performed on the only basis of regular NOESY spectra measured on fully labeled samples did not converge towards a compact globular domain with interacting RRM, but showed independent folded RRM separated by a flexible linker (data not shown). Similarly, structure calculations performed with the RDC information, but without the interdomain NOE data, did not converge towards a single

relative position of the two RRM but led to independent RRM separated by a flexible linker. In other words, structure calculations performed without the information of the interdomain contacts obtained with the segmentally labeled sample failed to reliably detect and assign interdomain NOEs, demonstrating the need for unambiguous methods and the importance of our segmental labeling strategy. In the case of hnRNP A1 RRM, we believe that the almost perfect overlays of the NMR footprints (Fig. 1) is due to the particular nature of the interface between the RRM, as compared with PTB RRM3 and RRM4 interface for instance (Oberstrass et al. 2005; Vitali et al. 2006). In PTB RRM34, the interdomain interface involves many hydrophobic side chains forming an important hydrophobic core (Vitali et al. 2006). In hnRNP A1, there is not such an extended hydrophobic core, with an interface composed primarily of a small hydrophobic patch and two Arg-Asp salt-bridges (Fig. 4e). In such interface, backbone amide resonances do not seem to be a very sensitive NMR probe. Chemical shifts of the side chains directly involved in the interdomain interface would probably experience larger changes, but it is rather difficult to monitor such changes in initial sample evaluations that are often performed with only ^{15}N -labeled samples. The proper decision on whether domains are interacting or not is crucial as it can latter strongly influence and restrict our understanding of the different mode of nucleic acid binding accessible to a particular multi-domain protein (see Supplementary Figure 1 and paragraph below), and one should therefore pay particular attention to this aspect. This point is of very broad relevance, since among eukaryotic proteins, the presence of multiple RRM is very common and is estimated to occur in about 44 % of the proteins containing at least one RRM (Maris et al. 2005). In addition, the RRM is one of the most abundant protein domains, and proteins with multiple RRM are estimated to be present in about 1 % of human gene products.

Comparison between hnRNP A1 solution and crystal structures

We showed in this study that the relative orientation of the two RRM of hnRNP A1 in solution closely resembles the one observed in the crystal structure of UP1 bound to DNA (Fig. 5), indicating that the two RRM of hnRNP A1 have a single defined relative orientation in solution, and that the small domain movement observed between the two crystal structures of the protein (free and bound) is not resulting from nucleic acid binding as previously proposed. The only differences in the side chains interaction that we could find to rationalize the fact that UP1_{bound} would be more stable than UP1_{free} are located in the small hydrophobic cluster formed by Leu13, Ile164 and Arg88. In UP1_{bound}, Ile164 is

more tightly packed in between the side chains of Leu13 and Arg88, as seen also in $UP1_{NMR}$, whereas in $UP1_{free}$, these side chains are further apart (Supplementary Figure 7). This could explain that $UP1_{bound}$ would be the native conformation of UP1, as observed in solution by NMR, and $UP1_{free}$ a destabilized structure induced by crystallization.

In addition to this difference in domain orientation observed between the free solution structure ($UP1_{NMR}$) and the free crystal structure ($UP1_{free}$), some small local variations exist between these two structures (Supplementary Figure 8). $RRM1_{NMR}$ and $RRM1_{free}$ are almost indistinguishable (backbone r.m.s.d. of 0.55 Å for the entire domain, *i.e.* residues 11–89, Table 3). Even the long $\beta 2$ – $\beta 3$ loop of $UP1_{NMR}$ agrees quite well with $UP1_{free}$ (Supplementary Figure 8). According to the heteronuclear $^{15}N\{^1H\}$ -NOE values (Supplementary Figure 4), residues directly following $\beta 2$ or directly preceding $\beta 3$ are relatively rigid and only 4 residues have NOE values <0.72 . Differences between the two structures are more apparent in RRM2 where the $\beta 2$ – $\beta 3$ and the $\beta 1$ – $\alpha 1$ loops differ significantly (Supplementary Figure 8). These two loops are also regions where the precision of the structure is lower (Fig. 4c). This conformational heterogeneity observed in the $\beta 2$ – $\beta 3$ loop correlates with the heteronuclear $^{15}N\{^1H\}$ -NOE values, which are overall lower than the corresponding NOE values in RRM1 $\beta 2$ – $\beta 3$ loop. The heterogeneity observed in the $\beta 1$ – $\alpha 1$ loop only reflects the lack of observable NMR signals and therefore the lack of chemical shift assignment for this region. This loop is most probably also dynamic, but we do not have clear evidence to support this point. Outside these two loop regions, $RRM2_{NMR}$ and $RRM2_{free}$ overlay quite well (backbone r.m.s.d. of 0.79 Å for residues 105–111, 117–139 and 146–180, Table 3). The only structural difference between rigid residues in $UP1_{NMR}$ and $UP1_{free}$ is located in the loop following helix $\alpha 2$ and involve residues Lys166 and Tyr167 (Supplementary Figure 8). In $UP1_{free}$, Lys166 makes intermolecular contacts with Glu66 of a symmetry related molecule (Shamoo et al. 1997; Xu et al. 1997), leading to a large distortion of the protein backbone for these two residues. Similarly, in the $UP1_{bound}$ structure, Lys166 and Tyr167 makes intermolecular contacts with Asp94 and the carbonyl group of Ile164, respectively, of a symmetry related molecule, leading to comparable distortions of this region (Ding et al. 1999). In some structures of the $UP1_{NMR}$ ensemble, Lys166 makes intramolecular contacts with Glu93 of the interdomain linker. As a consequence, the long side chains of Lys166 and Tyr167 come in close proximity to the $\beta 4$ strand which could have some implication for nucleic acid recognition, since this strand forms the main region of sequence specific contacts involved in nucleic acid recognition (Ding et al. 1999). Importantly, this large distortion might be related to the relative domain movement observed between $UP1_{free}$ and

$UP1_{bound}$. Indeed, this region directly follows helix $\alpha 2$, which makes most of the interdomain contacts from RRM2, and Lys166 is very close in sequence to Ile164, which experience the largest displacement in this domain rotation of $\sim 15^\circ$ (Fig. 5c). Furthermore, Lys166 and Tyr167 are engaged in different protein–protein contacts in the two different crystal lattices of $UP1_{free}$ and $UP1_{bound}$ structures. However, also the differences in this region might be coupled with the interdomain movement, we cannot exclude that the re-orientation would come from additional crystal contacts or from totally different contacts in other regions of the protein.

Finally, two regions of the protein are not seen in the electron density maps of $UP1_{free}$, and become structured upon DNA binding as a consequence of being directly involved in nucleic acid recognition. These two regions correspond to the interdomain linker (residues Arg92 to Leu102), and the C-terminal segment after RRM2 (from residue Ser182). This perfectly correlates with the dynamic information obtained in solution (Supplementary Figure 4), where these residues have $^{15}N\{^1H\}$ -NOE values reflecting flexible residues. But interestingly, even if these regions are structurally heterogeneous (Fig. 4a–c), the secondary structured elements that appear upon DNA binding in these regions—namely a short α helical turn in the interdomain linker, involving residues 93–96, and a C-terminal α -helix after RRM2 involving residues 183–188—seem to some extent to be already present in the free form. The helical propensity of these two regions is supported by backbone chemical shift and few NOE cross-peaks (data not shown).

Repertoire of nucleic acid recognition modes in proteins containing multiple RRM domains

Since RRM domains are asymmetric binding platforms, the relative orientation of different RRMs in proteins containing multiple-RRMs strongly influences the modes of nucleic acid binding by these particular proteins. In other words, the path of the nucleic acid molecule bound to multiple RRMs is strongly dependent on whether these RRMs are interacting with each other and adopt a single defined relative orientation. A limited number of simplified situations can be used to describe the interplay between two RRMs and a nucleic acid molecule, depending on whether the RRMs are interacting or not in their free state, and whether they interact or not in the nucleic acid bound state. To date there is no example of RRMs interacting in their free state and not in their bound state, but all the other situations are supported by structural work (Fig. 6a–e). Additionally, the dynamic aspects related to nucleic acid recognition are definitely essential for multi-domain proteins and will be briefly mentioned below. However, readers interested in more details on these aspects may also refer to the excellent review by Mackereth and Sattler (2012).

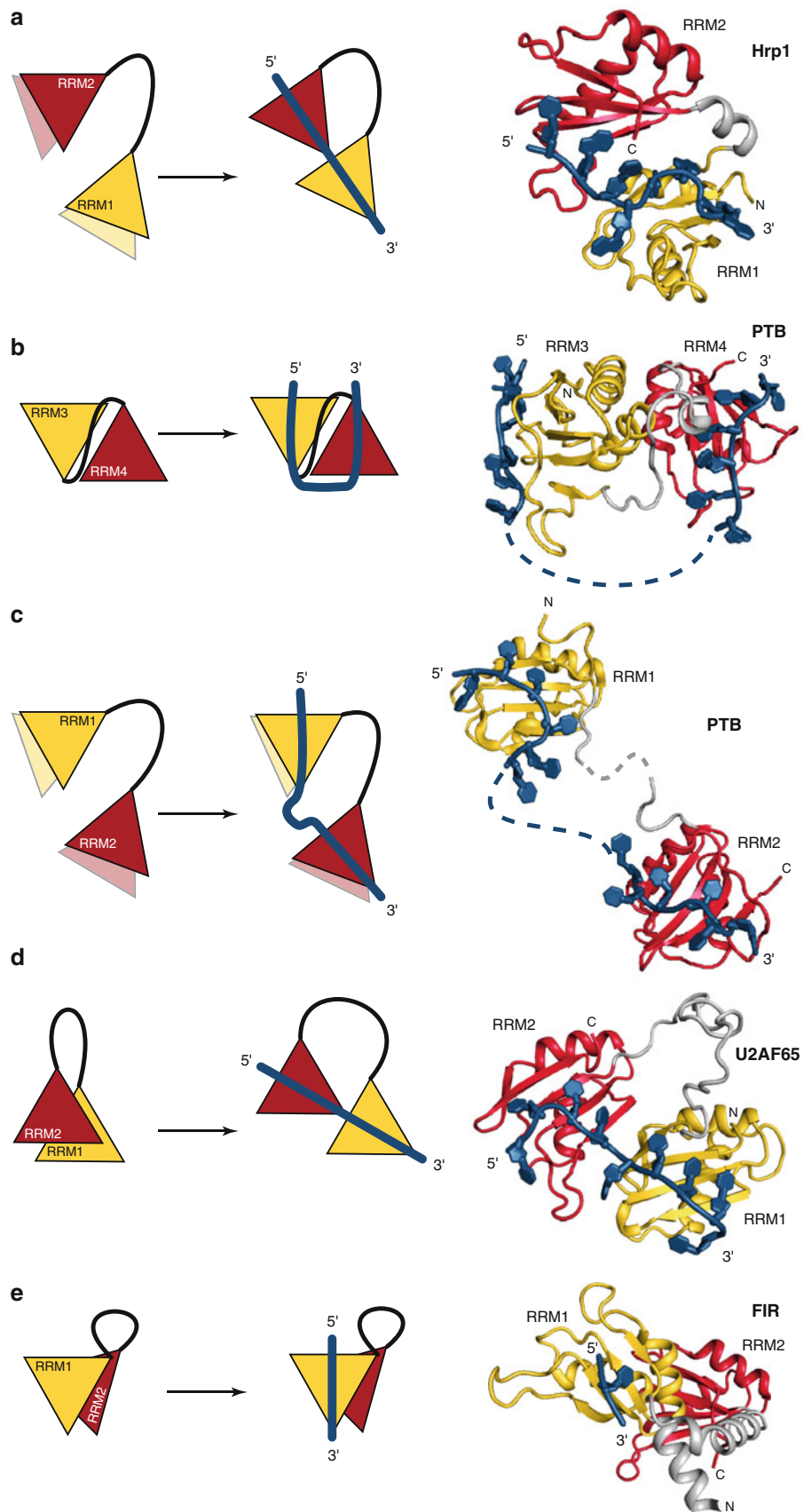
In the situation #1, the RRM s are not interacting in the free form and interact upon nucleic acid binding to form a continuous binding platform (Fig. 6a). Many different proteins can be classified in this group, and structural data are very abundant for this particular case, with for instance the different structures of nucleolin (Allain et al. 2000b, 2000a; Johansson et al. 2004; Arumugam et al. 2010), PABP (Deo et al. 1999; Safaee et al. 2012), Hrp1 (Perez-Canadillas 2006; Leeper et al. 2010), Sex-lethal (Handa et al. 1999; Crowder et al. 1999) and HuD (Wang and Tanaka Hall 2001). In these cases, the association of the two RRM platforms allows the continuous recognition of longer nucleic acid stretches (6–10 nucleotides), which often strongly increases the binding affinity compare to isolated domains. Interestingly, in all these different cases, RRM2 binds the 5' end of the RNA and RRM1 the 3' end. Remarkably, the relative orientation of the RRM s and the path of the RNA on the RRM platforms are very similar for different unrelated proteins of this class, namely Sex-lethal, Hrp1 and HuD. In these structures, the interdomain interface is relatively small ($\sim 350 \text{ \AA}^2$) and the interaction is mediated by a limited set of contacts, mainly one single salt-bridge and few hydrogen bonds, and might therefore be quite weak. This might explain why these contacts are induced by nucleic-acid binding and are not present in the free forms of the proteins. In addition, in cases where tandem RRM s could not be crystalized bound to the same RNA molecule (CUGBP1 for example), models have been proposed that have orientation similar to HuD/Sex-lethal/Hrp1 or to PABP (Teplova et al. 2010). However, there is to date no evidence to support one model or the other and it is therefore possible that none of these two models adequately describe CUGBP1 RRM12 binding to RNA. Overall, according to the currently available structural informations, this situation of independent RRM s interacting upon RNA binding seems to be the most common in tandem RRM s although the number of example is still scarce. It is very possible that this particular case might be over-represented in the solved structures since such stable and compact complexes would be more susceptible to crystalize than independent and dynamic RRM s.

In situation #2, the RRM s are interacting in the free form and adopt a single defined orientation resulting in a discontinuous antiparallel platform that is maintained in the bound form (Fig. 6b). This particular topology can induce RNA loops in the bound RNA and have a role for regulating alternative splicing. This looping capacity of interacting tandem RRM s has been demonstrated in the case of PTB RRM34 (Oberstrass et al. 2005; Lamichhane et al. 2010). Such topology with a discontinuous antiparallel platform is also found in the two tandem RRM s of hnRNP A1 suggesting that it might also be able to loop out RNA. However, there is to date no evidence to support such

mechanism of action. In these two examples, the surface of interaction between the RRM s is larger than the ones found in the situation #1 (namely ~ 630 and $\sim 850 \text{ \AA}^2$ for hnRNP A1 and PTB RRM34, respectively). These interfaces involve a combination of electrostatic and hydrophobic interactions, with even a large interdomain hydrophobic core in the case of PTB RRM34. It could have been questionable to classify hnRNP A1 in this group if our analysis was only based on the existing crystal structures, since the relative orientation could seem to be altered upon nucleic acid binding. However, our solution structure supports the fact that hnRNP A1 RRM s exist in a single defined orientation, as observed for PTB RRM34.

In situation #3, the RRM s are neither interacting in the free form nor in the bound form (Fig. 6c). This case is likely to be quite common considering the high number of cases where tandem RRM s are separated by long and disordered interdomain linkers, but structural data supporting this mode of binding are not very abundant. In most of the studies, the independence of the RRM s is well established in the free form, and is then assumed for the bound form as well, even if this point is not always clearly demonstrated. This mode of binding was proposed for instance for PTB RRM12 (Oberstrass et al. 2005), hnRNP F RRM12 (Dominguez and Allain 2006) and Npl3p RRM12 (Skrisovska and Allain 2008). In these cases, the binding of multiple RRM s to the same nucleic acid molecule increases the overall binding affinity (Shamoo et al. 1995). Since the binding of an individual RRM to RNA can be rather weak ($K_D \sim 1 \mu\text{M}$), this cooperation is an essential aspect enabling these multi-domain proteins to achieve their function in alternative splicing at low cellular concentration.

In situation #4, the RRM s are interacting in the free form such that one binding platform is occluded by the other RRM. The RRM s are also interacting in their bound form, but in a different relative orientation that forms a continuous binding platform (Fig. 6d). This case occurs in the splicing factor U2AF65, where the recognition of a poly-pyrimidine tract RNA is associated with an equilibrium between a closed state and an open state competent for RNA binding (Mackereth et al. 2011). The presence of this closed state results in an autoinhibition of binding that is used to finely tune U2AF binding to various 3' splice sites harbouring different pyrimidine tracts. In both states, the surfaces of interaction between the two RRM s are quite small (~ 460 and $\sim 320 \text{ \AA}^2$ for the closed and open states, respectively) and are stabilized by a limited set of electrostatic interactions, namely one potential salt-bridge and few contacts between polar side chains in the closed state of the protein, and a small number of contacts between polar side chains in the open state. This case illustrates the importance of the dynamics and the role of weak



◀ **Fig. 6** Repertoire of nucleic acid binding modes by proteins containing two RRM. Schematic representations of the different mode of binding of tandem RRMs to strands of nucleic acid are shown in the *left panels*, and archetypal structures corresponding to each mode of binding are shown in their bound form in the *right panels*. N-terminal RRMs are in *yellow*, C-terminal RRMs in *red* and nucleic acid molecules in *blue*. **a** (*Situation #1*) The RRMs are not interacting in the free form and interact upon nucleic acid binding to form a continuous binding platform. The RRMs do not have a defined relative orientation in the free form, but have a defined and unique relative orientation in the bound form. This case occurs in diverse proteins, namely PABP, nucleolin, Hrp1, Sex-lethal and HuD. The structure in the right panel represents Hrp1 bound to RNA (pdb code 2CJK) (Perez-Canadillas 2006). **b** (*Situation #2*) The RRMs are interacting in the free form to create a discontinuous antiparallel platform and interact in the bound form keeping the same relative orientation. This case occurs in PTB RRM34, and hnRNP A1 RRM12. The structure in the right panel represents PTB RRM3 and RRM4 bound to RNA (pdb code 2ADC) (Oberstrass et al. 2005). The path of the RNA from RRM3 to RRM4 is based on biochemical and biophysical data obtained in solution (Lamichane et al. 2010). **c** (*Situation #3*) The RRMs are neither interacting in the free form nor in the bound form. This case is expected to be quite common, but structural data are not abundant to support this mode of binding. It was proposed for instance for PTB RRM12, hnRNP F RRM12 and Npl3p RRM12. The structure in the right panel corresponds to PTB RRM1 and RRM2 bound to RNA (pdb codes 2AD9 and 2ADB) (Oberstrass et al. 2005). **d** (*Situation #4*) The RRMs are interacting in the free form such that one binding platform is occluded by the other RRM. The RRMs are also interacting in their bound form, but in a different relative orientation that forms a continuous binding platform. This case occurs in U2AF65 and its bound form is represented on the right panel (pdb code 2YH1) (Mackereth et al. 2011). **e** (*Situation #5*) The RRMs are interacting in the free form such that one binding platform is occluded by the other RRM. The RRMs are also interacting in their bound form and only one RRM can bind to nucleic acid. This case occurs in Prp24 and FIR and the bound form of FIR is represented on the right panel (pdb code 2QFJ) (Crichlow et al. 2008)

interdomain contacts for the binding of multi-RRM domains to RNA, in relation to a subtle regulation of a complex biological mechanism. This equilibrium could not have been deduced from the static crystal structure of U2AF65 (Sickmier et al. 2006), confirming the importance of solution techniques for the analysis of the interaction between multi-domain proteins and nucleic acids.

In situation #5, the RRMs are interacting in the free form such that one binding platform is occluded by the other RRM. This interaction is preserved in the bound form and thus only one RRM can bind to nucleic acid (Fig. 6e). This situation has been observed in the transcriptional repressor FIR (Crichlow et al. 2008; Cukier et al. 2010), and could possibly describe as well the situation observed in the splicing factor Prp24 (Bae et al. 2007; Martin-Tumasz et al. 2010). In FIR, the binding platform of RRM2 is occluded by RRM1 through a very large interaction surface involving RRM1 helices ($\sim 900 \text{ \AA}^2$), leaving only RRM1 binding platform available for DNA binding (Fig. 6e). A recent NMR study showed that the occluded RRM2, which

cannot bind to DNA, is involved in protein–protein interaction with the transcriptional activator FBP (Cukier et al. 2010), whereas the initial crystal structure favoured a model in which RRM2 would drive the dimerization of FIR (Crichlow et al. 2008). There is a similar arrangement in the three tandem RRMs of Prp24, where RRM2 forms extensive interdomain contacts with RRM1 and RRM3 in the crystal structure of RRM123 (Bae et al. 2007). However, whereas the RRM1–RRM2 interaction is preserved in solution (Bae et al. 2007), RRM2 and RRM3 do not interact in solution, leaving the RRM2 binding platform available for RNA binding of a segment of the U6 snRNA (Martin-Tumasz et al. 2010). The role of the occluded RRM is not as clear as in the case of FIR, but it has been proposed that RRM1 would also interact with U6 snRNA, also with a different surface than the canonical β -sheet platform (Bae et al. 2007). Further data would be needed to generalize these observations, but these two examples suggest that when an RRM binding platform is occluded by a preceding or a following RRM, the occluded RRM could function in protein–protein interaction, or could bind to nucleic acid in a non-canonical manner.

Conclusion

This overview of the structural characterization of tandem RRM proteins and their interaction with nucleic acid illustrates the large repertoire of nucleic acid recognition modes in proteins containing multiple RRM domains. Noticeably, large surfaces of interaction between tandem RRMs ($\sim 600\text{--}900 \text{ \AA}^2$) are associated with a static behaviour and a defined relative orientation of the domains (situations #2 and #5), whereas smaller surfaces ($\sim 300\text{--}500 \text{ \AA}^2$) correlate with a dynamic behaviour and a re-orientation of the RRMs upon nucleic acid binding (situations #1 and #4). In addition, several examples directly indicate that solution techniques are essential to investigate the interaction of tandem RRMs and more generally multi-domain proteins with nucleic acids. Crystallization can sometimes confine protein domains in non-functional conformations or induce artificial nucleic-acid topology of binding. To obtain reliable information, in solution, on the relative orientation and on the surface of interaction of different domains of multi-domain proteins, segmental isotope labeling is definitely a method of choice. And considering the progress that have been realized in the last years in developing new methods and/or improving existing protocols for segmentally labeled sample production, we believe that segmental labeling will be a central approach for the future investigations of multi-domain nucleic-acid-binding proteins.

Accession numbers

The chemical shifts of hnRNP A1 RRM12 have been deposited in the BioMagResBank under accession number 18728. The coordinates of the structure have been deposited in the Protein Data Bank under accession code 2LYV.

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