Evidence for cooperative tandem binding of hnRNP C RRMs in mRNA processing

ZUZANA CIENIKOVÁ, SANDRINE JAYNE, FRED FRANZ DAMBERGER, FRÉDÉRIC HAI-TRIEU ALLAIN, and CHRISTOPHE MARIS
Department of Biology, Institute of Molecular Biology and Biophysics, ETH Zürich, 8093 Zürich, Switzerland

ABSTRACT
The human hnRNP C is a ubiquitous cellular protein involved in mRNA maturation. Recently, we have shown that this protein specifically recognizes uridine (U) pentamers through its single RNA recognition motif (RRM). However, a large fraction of natural RNA targets of hnRNP C consists of much longer contiguous uridine stretches. To understand how these extended sites are recognized, we studied the binding of the RRM to U-tracts of 8–11 bases. In vivo investigation of internal translation activation of unr (upstream of N-ras) mRNA indicates that the conservation of the entire hnRNP C binding site, UC(U)₈, is required for hnRNP C-dependent IRES activation. The assays further suggest a synergistic interplay between hnRNP C monomers, dependent on the protein’s ability to oligomerize. In vitro spectroscopic and thermodynamic analyses show that isolated RRMs bind to (U)₁₁ oligomers as dimers. Structural modeling of a ternary double-RRM/RNA complex indicates additionally that two RRM copies can be accommodated on the canonical sequence UC(U)₈. The proposed tandem RRM binding is in very good agreement with the transcriptome-wide recognition of extended U-tracts by full-length hnRNP C, which displays a cross-linking pattern consistent with a positively cooperative RRM dimer binding model.

Keywords: hnRNP C; uridine; alternative splicing; IRES; RRM

INTRODUCTION
The heterogeneous nuclear ribonucleoprotein C1/C2 (hnRNP C) was first characterized in the 1980s as an abundant, RNA-associated nuclear protein (Choi and Dreyfuss 1984). Today, hnRNP C is established as a major player in the mRNA maturation pathway: Its function spans nascent RNA packaging (Rech et al. 1995), sorting (McCloskey et al. 2012), and splicing regulation (Venables et al. 2008; König et al. 2010). More recently, the protein was also detected in the cytoplasm, associating with internal ribosome entry sites (IRES) of several eukaryotic mRNAs (Zaidi and Malter 1995; Kim et al. 2003; Schepens et al. 2007). Located within the 5’ untranslated regions (UTR), IRES sites serve to recruit the ribosome and associated trans-acting factors, and initiate protein synthesis in a cap-independent manner.

The human hnRNP C homolog is ~300 residues long and is present in vivo in two variants, C1 and C2; the latter includes a 13 amino acid insertion (Merrill et al. 1989). While the endogenous hnRNP C protein acts as a (C1),₂(C2) oligomer (Barnett et al. 1989), it has been shown that C1 or C2 homotetramers retain the wild-type structural and functional properties (McAfee et al. 1996b). The oligomerization is dependent on a central helical domain folding into antiparallel coiled-coil tetramers (Whitson et al. 2005). Two hnRNP C subdomains are thought to be involved in RNA binding: The amino-terminal RNA recognition motif (RRM) recognizes five contiguous uridines, with the sequence stringency increasing gradually in the 5’–3’ direction (Wan et al. 2001; Cieniková et al. 2014); a basic region preceding the oligomerization domain, rich in positively charged amino acids, provides additional, sequence-independent affinity for RNA (McAfee et al. 1996a; Soltaninassab et al. 1998). One tetramer binds ~200 to 300 ribonucleotides in a strongly cooperative fashion (McAfee et al. 1996b). A large-scale cross-linking study of hnRNP C revealed that the protein’s association on RNA transcripts undergoing splicing is coordinated by regularly spaced uridine tracts (König et al. 2010). Uridine pentamers, corresponding to the sequence specifically recognized by the RRM domain, were the most represented sites among hnRNP C cellular targets. However, uridine stretches considerably longer than 5 nucleotides (nt) were also highly enriched...
among the identified binding sites. Further supporting the importance of longer uridine tracts, the activation of internal translation initiation of upstream of N-ras (unr) mRNA during mitosis was shown to be regulated by hnRNP C (Sella et al. 1999) migrating into the cytoplasm and binding to a UC(U)$_n$ tract located at position −102 to −93 upstream of the translation start site (Cornelis et al. 2005; Schepens et al. 2007). An additional, weaker poly(U) hnRNP C-binding site was identified at position −190 to −180. A thermodynamic investigation of the RRM-poly(U) RNA interaction (Cieniková et al. 2014) revealed that register sampling of the RRM on extended uridine tracts leads to a gain of binding affinity which could be the reason for the conservation of such long binding sites. Alternatively, König et al. (2010) proposed that hnRNP C tetramer binds long uridine stretches via two RRMs in tandem.

To understand the recognition of extended uridine-rich sites by hnRNP C, we investigated the binding of the RRM to a (U)$_{11}$ oligomer using NMR spectroscopy and isothermal titration calorimetry (ITC). Using the insights derived from these in vitro studies and from molecular modeling driven by the experimental NMR and thermodynamic data, we extend our thermodynamic model of hnRNP C binding to account for tandem RRM binding to long U tracts in vivo and characterized the functional requirements for this binding mode using hnRNP C-dependent unr IRES activation in cell culture assays.

RESULTS

Affinity enhancement due to multiregister binding of the RRM is insufficient to account for the in vivo hnRNP C occupancy on long uridine tracts

We recently developed a structurally driven thermodynamic model to predict the binding of hnRNP C RRM to poly (U) sites (Cieniková et al. 2014). This model explained very well the in vivo ligand saturation obtained by iCLIP for uridine tracts shorter than 7 nt (König et al. 2010), indicating that one RRM binds such sites in multiregister mode (Fig. 1A, green curve). In the present study, we extended this analysis to longer uridine stretches. However, we observed that the saturation deviates sharply from this model when uridine tracts exceed 8 nt in length. Since one RRM binds four to five consecutive uridines (Cieniková et al. 2014), this change of binding behavior could indicate that two hnRNP C RRMs bind side-by-side on such long tracts.

In vitro, a pair of isolated hnRNP C RRMs can bind an 11-nt uridine tract

To determine whether two RRMs of hnRNP C can simultaneously bind to a long pyrimidine tract, we characterized spectroscopically and thermodynamically the interaction of the hnRNP C RRM with an extended uridine stretch. We recorded NMR relaxation spectra of the RRM in the presence of a synthetic (U)$_{11}$ oligomer at 1:1 and 2:1 stoichiometric ratios. The chemical shifts of the corresponding protein amides in the 1:1 and 2:1 RRM:(U)$_{11}$ complexes are expected to mostly overlap due to the degenerate nature of the RNA sequence and therefore the relaxation behavior of these signals is a superposition of the 1:1 and 2:1 complexes. The average rotational correlation time $\tau_C$ of the RRM, determined from $^{15}$N relaxation data, is 10.6 ± 0.4 nsec (mean ± SD) in the 1:1 sample and it increases to 12.6 ± 1.0 nsec in the 2:1 sample (Fig. 1B). This difference is statistically significant ($P$-value <10$^{-13}$ using Student’s $t$-test), suggesting that in the latter sample a significant fraction of the RRM is involved in the more slowly tumbling 2:1 protein:RNA complex. However, NMR signals belonging to the free RRM form remain observable in the $^1$H-$^15$N HSQC spectrum obtained at the 2:1 ratio, consistent with the interpretation that the 2:1 species is only partially saturated (Supplemental Fig. S1A).

Additional insights are provided by a calorimetric titration of the RRM to the (U)$_{11}$ RNA (Fig. 1C). An initial fit of the ITC data to the simplest binding model (one set of...
$N$ identical sites) suggests a binding stoichiometry $N \approx 1.5$. A visual inspection of the binding isotherm reveals biphasic behavior, indicating two association events of different macroscopic affinity. A model presuming two identical, sequentially bound ligands increases $10^6$-fold the fit quality, as indicated by the value of the corrected Akaike information criterion (Table 1). The fitted macroscopic affinity constant of the first binding step displays an $\sim 11$-fold enhancement compared with the expected microscopic single site affinity ($K_s \approx 5 \times 10^5$ M$^{-1}$ [Cieniková et al. 2014]), confirming a strong multiregister binding of the RRM ligand. The second binding step manifests a weak negative cooperativity (interaction factor $a \approx 0.4$), when accounting for the existence of $\sim 4$ binding registers on the (U)$_{11}$ RNA at 2:1 stoichiometry. This 2.5-fold reduction in affinity is likely to stem from adjustments required to avoid steric clashes between the adjacent RRM molecules.

**Structural modeling suggests that a 10-nt site containing eight consecutive uridines can accommodate a dimer of hnRNP C RRMs**

To obtain insight into how two tandem RRMs could be accommodated on a longer U-tract, we performed structural modeling guided by constraints derived from the solution NMR structure of a single RRM binding the five uridines of the AUUUUUC RNA (Cieniková et al. 2014). Since the cross-linking data show that hnRNP C binding is enhanced for uridine tracts of eight or more bases, and the NMR structures previously obtained indicate that one hnRNP C RRM contains 5-nt binding pockets, we calculated using torsion angle dynamics a model of two RRMs binding in tandem on a sequence of 10 nt containing a (U)$_8$ tract. The sequence UC(U)$_8$ was chosen because it corresponds to the known hnRNP C regulatory site in mtr $5^\prime$ UTR. The proposed ternary complex model (Fig. 2A) satisfies the minimal specificity requirements of the protein (three strongly uridine-selective pockets at the 3′ end of each RRM). A cytidine occupies the second position of the upstream RRM in the model, instead of a uridine as in the solved structure. To confirm that this substitution can be accommodated by the RRM in vitro, we compared the AUCUUUC- and AUUUUUC-RRM complexes by NMR (Fig. 2B,C; Supplemental Fig. S1B). The similarity of the protein and RNA chemical shifts indicates that the AUCUUUC-RRM complex adopts a similar overall conformation and binds in the same register as the solved AUUUUUC-RRM complex, with the first cytidine positioned in the aforementioned pocket 2. The torsion angle dynamics simulation of the double-RRM-UC(U)$_8$ complex structure (Fig. 3) shows additionally that the proposed arrangement of two RRMs on the UC(U)$_8$ oligomer is indeed possible without major steric clashes, since we observe $<40\%$ increase in the average CYANA target function measuring the severity of violations of the NMR-derived input constraints compared with the sum of equivalent simulations with two isolated RRMs (Table 2). Our analysis therefore indicates that a sequence of 10 nt containing a tract of eight uridines constitutes the minimal specific binding site for a hnRNP C RRM dimer.

**In vivo, tandem hnRNP C RRM binding is positively cooperative**

The in vitro experiments support the hypothesis of double-RRM binding on sites carrying at least eight contiguous uridines. To address whether this binding behavior reflects how hnRNP C interacts with cellular transcripts, we implemented a structure-based thermodynamic model of uridine lattice saturation by tandem RRMs and then compared the transcriptome-wide distribution of hnRNP C cross-links on uridine tracts up to 12-nt long found by iCLIP (König et al. 2010) with the predictions of this model. The saturation model is an extension of the uridine lattice binding model that we used previously to describe the single RRM’s interaction with short tracts (Cieniková et al. 2014). It was parameterized with the affinity weight values obtained for equimolar binding but extended to account for 2:1 RRM:RNA stoichiometry. The introduction of the 2:1 stoichiometry species manifests itself by an inflexion point on the saturation curve for tracts long enough to allow binding of two RRM ligands ($n \approx 8$ or 9) (Fig. 1A, red curve). The gain in saturation due to the bipartite binding improves the overall fit to the in vivo data but does not fully account for the observed saturation. A good fit to the in vivo cross-link data for uridine lattice lengths between 1 and 12 nt is obtained only after the introduction of a positive interaction $\alpha$ between the two RRMs (Fig. 1A, blue curve). The experimental data are therefore consistent only with a model assuming that the cross-links found in tracts of 8–12 uridines originate from cooperative binding of two RRMs, each occupying five contiguous nucleotides. The value of the affinity enhancement at 2:1 RRM:RNA stoichiometry obtained from the fit to the iCLIP data set is $\approx 14$. The binding at uridine tract lengths above 13 nt cannot be quantitatively investigated, since the experimental data for very long uridine tracts are sparse and therefore the cross-link frequencies unreliable. A more recent iCLIP of hnRNP C, performed with increased sequencing depth, shows however that the uridine site occupancy saturates at tract

| TABLE 1. | ITC titrations of hnRNP C RRM to (U)$_{11}$ |
|-------------------|-------------------|
| Binding model | $N$ | $K_1$ (M$^{-1}$) | $K_2$ (M$^{-1}$) | AICc (d.o.f.) |
| One set of sites | $1.53 \pm 0.04^a$ | $(2.13 \pm 0.25) \times 10^5$ | – | 145.7 (43) |
| Sequential sites | $2^b$ | $(5.71 \pm 1.00) \times 10^5$ | $(7.54 \pm 0.53) \times 10^4$ | 75.5 (42) |

$^a$Uncertainties in the fitted parameters were estimated from the fit residual.  
$^b$Fixed parameter of the model.
lengths of 11 or more uridines (Supplemental Fig. S1B of Zarnack et al. 2013), suggesting that there is no positive cooperativity driving binding of additional RRMs beyond the first dimer. However, fitting to this data set is hindered by the fact that it also includes cross-links with only one detected cDNA, which appear to predominantly correspond to nonspecific interactions.

Further insights into the arrangement of the two side-by-side binding RRMs can be gained by analyzing the position-specific distribution of hnRNP C cross-links within the uridine tracts (Fig. 4; Supplemental Fig. S2). Our thermodynamic model for the binding of two RRMs in tandem predicts correctly the existence of two binding maxima and their absolute and relative position. However, while the modeled maxima are of similar height, the experimental iCLIP data show a decrease or broadening of cross-link deposition at the 5′ end of long uridine tracts, indicating that a feature in the binding of the upstream RRM is not captured by the model. Introducing an additional dynamic component into this upstream binding which permits Gaussian sampling of registers centered on the predicted position broadens the maximum and improves considerably the fit to the observed cross-link distribution.

Disruptions within the extended poly(U) site in unr IRES site abolish the translation activation by hnRNP C

We have shown that two hnRNP C RRM domains are able to bind in tandem on a uridine stretch of ~8 nt. We observed a synergistic binding behavior in the context of the full-length, oligomerizing protein interacting with cellular transcripts in vivo, in contrast to isolated RRM domains binding to (U)11 in vitro which did not manifest positive cooperativity at 2:1 stoichiometry. To understand how the requirement
TABLE 2. CYANA calculation statistics for hnRNP C RRM structural models

| Domain           | TF| Upper distance | Van der Waals | Torsion angle |
|------------------|---|----------------|---------------|--------------|
|                  | Number | Maximal violation (Å) | Number | Maximal violation (Å) | Number | Maximal violation (Å) |
| RRM-UCU_{i}      | 4.0 ± 0.8 | 7 ± 4 | 0.41 ± 0.11 | 6 ± 2 | 0.37 ± 0.03 | 0 ± 0 | 0.49 ± 1.47 |
| RRM-UB_{i}       | 3.8 ± 0.8 | 6 ± 3 | 0.38 ± 0.14 | 5 ± 1 | 0.38 ± 0.04 | 0 ± 0 | 0.76 ± 2.24 |
| Double RRM-UCU_{i} | 10.8 ± 1.6 | 22 ± 7 | 0.65 ± 0.19 | 15 ± 4 | 0.41 ± 0.05 | 0 ± 1 | 4.08 ± 4.74 |

*Mean and SD of CYANA target function values over 50 best structures.

AVERAGE number of restraints violated by more than the following amounts: distance and Van der Waals restraints 0.2 Å, torsion angle restraints 5°.

for side-by-side binding of hnRNP C RRM s affects IRES activity in vivo, we studied the nucleotide requirements of hnRNP C-dependent internal translation activation of unr mRNA in HEK293T cells (Schepens et al. 2007). We designed mRNAs with mutations within the unr IRES UC(U)_{i} site (Fig. 5A). The wild-type variant (WT) carried the central (U)_{8} tract; in the mutants (A)_{2}(U)_{6} and (A)_{4}(U)_{4}, the first two, respectively four uridines of the (U)_{8} site were replaced by adenosines; in (U)_{3}A(U)_{4}, the fourth uridine was replaced with an adenosine; finally, the (A)_{8} mRNA, with adenosines instead of the eight uridines, was used as our negative control. We show that shortening or interrupting the (U)_{8} tract leads to a 20%-30% decrease in translation activation (Fig. 5B). Moreover, these altered IRES variants are unresponsive to overexpressed ectopic hnRNP C1, in contrast to the WT unr IRES where the translation initiation is strongly promoted. All these mRNA variants (except the negative control) were designed to contain between four and six contiguous uridines, which is sufficient for binding one hnRNP C RRM, as indicated by the results of electrophoretic mobility shift assays (Supplemental Fig. S3). Nevertheless, the introduced modifications interfere with the activator role of hnRNP C, indicating that an interaction spanning the full uridine-rich binding site is required to promote UNR translation initiation.

**Intact hnRNP C RRM and oligomerization domains are required for the translation activation**

To understand the role of the hnRNP C subdomains in IRES-dependent UNR translation initiation, we overexpressed hnRNP C1 deletion variants or full-length constructs containing point mutations within the RRM domain (Fig. 5C, D). When endogenous hnRNP C is silenced, the IRES activity decreases to levels comparable to those observed with an abolished unr uridine site in the absence of silencing (Fig. 5B). Wild-type ectopic hnRNP C1 rescues translation initiation, while this activation is not observed for the ectopically expressed amino-terminal hnRNP C1 deletion variant ΔN1-9 as well as for RRM domain mutants F19A and F54A lacking aromatic side-chains at the binding interface. This demonstrates that the RRM and the N-terminal residues are essential for internal translation initiation of UNR. Previously we showed (Cieniková et al. 2014) that the mutated or deleted residues are involved in uridine binding; the loss of function seems therefore to be connected to the inability of the mutants to recognize their uridine target sites in the unr IRES. Additionally, we observe that a shortened hnRNP C1 variant retaining only the RRM domain and the basic region (“R + B”, residues 1 to 179) is unable to promote translation initiation, despite the fact that it comprises all subdomains thought to drive the binding to RNA. Since this variant lacks the central helical domain and is therefore unable to multimerize, these data suggest that oligomerization is required for IRES activation.

To investigate the role of cooperative interactions between hnRNP C monomers in IRES activation, we designed an experiment where the full-length exogenous WT hnRNP C1 or one of the following mutants, F19A, F54A, or ΔN1-9, were coexpressed alongside the endogenous hnRNP C. As expected, overexpression of the WT variant promotes IRES activation, as demonstrated by the 20%-30% increase in translation initiation observed for the WT unr IRES in comparison to the mutant variants (Fig. 5B). These results highlight the importance of cooperative interactions between hnRNP C monomers for efficient translation initiation.

**FIGURE 4.** Comparison between the cross-link frequencies observed within a uridine tract of 10 nt and the predicted register occupancy of such a tract by the RRM dimer. (A) The two thermodynamic models are superimposed on the iCLIP data (figure adapted with permission from Macmillan Publishers Ltd. © 2010 [König et al. 2010]). The iCLIP positions are color-coded to match the uridines with nucleotide pockets in the structural model (B) of the RRM dimer binding the (U)_{10} tract in the favored register. The two cross-linking maxima, highlighted by orange arrows, match the RRM pockets where the cross-link formation was predicted (pocket 3, Cieniková et al. 2014).

www.rnajournal.org 1935
activation (Fig. 5E). On the other hand, because the mutants do not activate translation at all when the endogenous hnRNP C is depleted (Fig. 5D), we anticipated little or no response when expressing them in the presence of the endogenous hnRNP C. Contrary to expectations however, the expression of these hnRNP C mutants leads to partial or full activation of the unr IRES relative to the activation level observed upon overexpression of WT hnRNP C1. Clearly, the effects of these overexpressed mutants and of the intact endogenous protein are nonadditive, meaning that the wild-type and mutant full-length hnRNP C monomers act synergistically. It is very likely that this effect is achieved through their interaction within the tetramer.

In conclusion, our results show that to promote the internal translation initiation of UNR, hnRNP C requires an intact U8 stretch in the UC(U)8 site on human unr mRNA. Combining structural modeling, calorimetry and NMR dynamics measurements, we deduce that the wild-type, 10-nt-long uridine-rich site can accommodate two hnRNP C RMs binding side-by-side. The RRM domains in isolation are however unable to drive their own multimerization on the poly(U) RNA. Therefore, the positive cooperativity observed in vivo between two hnRNP C monomers must arise from the tetramerization of the full-length protein.

**DISCUSSION**

To date, about a dozen structures of RRMs bound to poly(U) sequences have been solved (Table 3). It is noteworthy that, with the exception of Daz-l, Nab3, and hnRNP C, all characterized proteins contain tandem RRM domains that act in concert and bind side-by-side on uridine-rich tracts of ~7 nt. Moreover, although the Daz-l and Nab3 proteins carry a single copy of the RNA-binding domain, they are reported to form dimers with a second RRM; Daz-l homodimerizes (Ruggiu and Cooke 2000) and Nab3 heterodimerizes with the single RRM domain of Nrd1 (Carroll et al. 2007). Both the Daz-l homodimer and the Nab3-Nrd1 heterodimer bind 9 nt uridine-rich sequences (Jenkins et al. 2011; Hobor et al. 2011). Similar to Daz-l and Nab3, hnRNP C contains a single RRM but forms oligomers in vivo. While the isolated RNA-binding domain of hnRNP C has been shown to recognize uridine pentamers (Cieniková et al. 2014), studies of the hnRNP C binding targets in vitro (Görlich et al. 1994) and in vivo (König et al. 2010) revealed an enrichment of tracts with eight or more contiguous uridines. This observation led to the hypothesis that the RRM domains assembled within the tetramer bind their RNA target sites in pairs. Our structural model of the ternary complex formed by two RRMs bound to poly(U) sequences (supported by the 10-fold weaker affinity of the RRM for AUCUUUC RNA compared to AUUUUUC [Cieniková et al. 2014]) and possibly steric hindrance between the domains indicated by the moderate increase in NMR constraint violations in our calculation hint that efficient tandem binding likely requires slightly longer uridine tracts. This is experimentally supported by the results of the transcriptome-wide UV cross-linking study (König et al. 2010): Within the pool of enriched RNA sequences, only targets carrying 10 or
TABLE 3. Poly(U)-binding RRM domains with known atomic structures

| Protein          | RNA          | PDB ID     |
|------------------|--------------|------------|
| Human CPEB1 RRM12 | (U)<sub>2</sub> | 2MKK       |
| Human CPEB4 RRM12 | (U)<sub>2</sub> | 2MKI       |
| Human CLGBP1 RRM1 | U(U)<sub>2</sub>GUU | 3NNH       |
| Human CLGBP1 RRM2 | UGUU         | 3NMR       |
| Mouse Daz-1 RRM  | UGUUU         | 2K57       |
| Human hnRNP C RRM| A(U)<sub>2</sub>C | 2MXY       |
| Mouse HuC RRM12  | (A(U)<sub>2</sub>)<sub>2</sub> | 1FNX       |
| Human HuD RRM12  | (U)<sub>2</sub>(U)<sub>2</sub> | 1FXL       |
| Human HuR RRM12  | (U)<sub>2</sub>(U)<sub>2</sub> | 4ED5       |
| Yeast Nab3 RRM   | UCUU         | 2L41       |
| Fly Sxl RRM12    | GUUG(U)<sub>2</sub> | 1B7F       |
| Human U2AF65 RRM | (U)<sub>2</sub> | 2G4B       |

more contiguous uridines manifest two clear cross-linking maxima. However, the constant 5–6 nt spacing between these maxima supports the compact arrangement of the RRM dimer proposed in our structural model. Further insights into tandem RRM binding on extended uridine tracts are provided by the thermodynamic interaction model. The model reproduces well in vivo cross-link saturation (Fig. 1A) and position probabilities (Fig. 4; Supplemental Fig. S2). Additionally, broadening in the cross-link position distribution suggests that the upstream-binding RRM is more dynamic than the downstream RRM, occupying a larger number of registers. Similar behavior was observed previously in the case of Sx and U2AF65 tandem RRMs binding to degenerate polypyrimidine tracts, where the 5′-end RRM2 was shown to sample a larger nucleotide range than the 3′-end RRM1 (Banerjee et al. 2003). While the origin of this difference in interaction behavior is unclear, it is possible that the downstream RRM is tethered to the 3′ end of the tract by protein or RNA contacts unaccounted for in our model. In the case of U2AF65 binding to the polypyrimidine tract preceding the 3′ splice site, U2AF35 could act as such an anchoring protein (Jenkins et al. 2013). For hnRNP C, hnRNP A1 could have an analogous function. Both proteins are constitutive components of the 40S hnRNP core particle (Lothstein et al. 1985), and there is evidence that they act in similar molecular contexts, in particular in 3′ splice site definition (Fig. 6A; Domsic et al. 2003; Tavanaz et al. 2012; Zarnack et al. 2013).

Our translation assays on 5′UTR IRES as well as the modeling of the iCLIP data suggest a synergistic interaction between two hnRNP C RRMs binding side-by-side in vivo. Such positive cooperativity between tandem binding RRMs is frequently observed (Kanaar et al. 1995; Park et al. 2000; Sickmier et al. 2006; Kuo et al. 2009; Teplova et al. 2010; Kuwasako et al. 2014). Indeed, structural data indicate that two RRMs binding in tandem to RNA make more extensive contacts to their RNA targets than single RRMs (57.7% ± 3.6% of RNA surface area contacting the protein interface in the former case, compared to 44.6% ± 9.1% in the latter).

$P$-value < 10<sup>-4</sup> based on 13 tandem RRMs and 26 single RRMs poly(U) RNA complexes deposited in PDB [see Materials and Methods]]. In most of the examples studied structurally to date, the interacting RRMs are in cis within the same protein molecule. However, a recent study of the ASD-1/SUP-12/RNA ternary complex (Kuwasko et al. 2014) showed that a synergistic RNA recognition can be achieved via RRMs provided by two separate protein monomers, although the measured affinity enhancement is rather moderate (one order of magnitude). In this respect, it is similar to the value of the intermolecular cooperativity factor ($\alpha \approx 14$) estimated for the hnRNP C RRMs on the basis of the thermodynamic modeling of the iCLIP data. Investigation of the 5′UTR IRES activation indicates that the ability to oligomerize is likely to be responsible for the cooperative behavior of tandem hnRNP C RRMs. We therefore propose that the oligomerization of hnRNP C serves a similar purpose as the RRM multiplicity in other proteins. It facilitates the cooperative recognition of extended low-complexity binding sites. It remains to be determined whether the arrangement within the tetrmeric serves simply to favorably orient the RRMs so as to prime them for the dual site recognition, or whether other domains of hnRNP C, such as the basic region, also directly contribute to sequence-specific RNA binding.

The analysis of the saturation pattern of the endogenous hnRNP C in the human transcriptome using the RRM binding model reveals that binding of two RRMs is accompanied by a strong enhancement of overall affinity, whereas no further enhancement is detected from the binding of additional RRM domains. This observation is consistent with the proposed orientation of the RRM subdomains within the hnRNP C tetramer, where the four RRMs are arranged in two interacting pairs (König et al. 2010). In agreement with this, the 5′UTR carries two extended uridine-rich sites (Fig. 6B): In addition to the described UC(U)<sub>8</sub> site, the UC(U)<sub>4</sub>G(U)<sub>4</sub> motif located 87 nt upstream of it also fits the double-RRM sequence consensus and was shown to bind hnRNP C and contribute to translation activation (Schepens et al. 2007). While the spacing between these sites is only half of the 165 nt suggested previously as the preferred span between two RRM pairs in the tetramer model, it is still sufficient considering the known dimensions of the tetramer (hydrated diameter of 13.6 nm [Huang et al. 1994]) and assuming an average distance between consecutive phosphorus atoms in a single-stranded nucleic acid of 6.3 Å (Murphy et al. 2004). The proposed positioning of the hnRNP C tetramer within the 5′UTR would leave the start codon free for ribosome binding, and is therefore in agreement with the translation activator role of the hnRNP C protein.
MATERIALS AND METHODS

Cloning and hnRNP C RRM purification for in vitro studies

The vector containing the sequence coding for the human hnRNP C RRM subdomain (AA 2-106), and the protocol of protein purification were described previously (Cieniková et al. 2014).

RNA production and purification

RNA oligomers of ~26 nt were in vitro transcribed with T7 polymerase from purchased DNA templates annealed to 18 nt-long T7 primers. The target was purified by anion-exchange chromatography under denaturing conditions, then desalted by two rounds of butanol precipitation, lyophilized, and resuspended in the appropriate buffer.

The short single-stranded RNAs (U)11, AUUUUUC and AUCU-UUC purchased from Dharmacon were deprotected according to manufacturer’s instructions and then purified as described above.

NMR spectroscopy and dynamics

NMR samples consisted of 1 mM 15N-labeled protein and unlabeled RNA present in 1:1 or 2:1 protein:RNA molar ratio, and were prepared in 250 μL of NMR buffer (20 mM Na2HPO4, 10 mM NaCl, pH 6.5, 10% D2O). Spectra were measured at 293 K on Bruker AVIII-600 MHz, AVIII-700 MHz, and AVIII-750 MHz spectrometers. All instruments except the AVIII-750 were equipped with cryoprobes. NMR data were processed with Topspin 2.1 (Bruker); the analysis of the spectra was performed in Sparky (Goddard and Kneller 1997). Chemical shift assignments were derived from 2D 1H-1H TOCSY and 1H-15N HSQC spectra recorded on the AUUUUUC and (U)11 RNA–RRM complexes and were based on the published assignments of the AUUUUUC–hnRNP C RRM complex (Cieniková et al. 2014). Longitudinal amide relaxation rates $R_1 = 1/T_1$ were extracted from a series of 2D 1H-15N HSQC spectra recorded on the AVIII-700 MHz spectrometer with nine mixing times ranging from 20 to 1500 msec. The rates for each residue were obtained by fitting the amide signal intensities to an exponential decay using the least-squares minimization method; the corresponding standard deviations (SD) were estimated with the jackknife approach (Efron and Stein 1981). Transverse relaxation rates $R_2 = 1/T_2$ were extracted from a series of 2D 1H-15N HSQC-based $T_2$ relaxation experiments (Kay et al. 1992) recorded on the AVIII-700 MHz spectrometer with nine mixing times ranging from 20 to 1500 msec. The rates for each residue were obtained by fitting the amide signal intensities to an exponential decay using the least-squares minimization method; the corresponding standard deviations (SD) were estimated with the jackknife approach (Efron and Stein 1981). Transverse relaxation rates $R_2$ were obtained from measurements of the $^{15}$N CSA/β-1H-15N dipolar cross-correlated relaxation rates $T_{15}$ using the expression $T_{2} = \kappa T_{15}$, where $\kappa = 1.33$ is a spectrometer field-dependent constant relating $T_{15}$ to the transverse relaxation rate $R_2$ devoid of the contribution from chemical exchange (Tjandra et al. 1996; Wang and Palmer 2003). We recorded a pair of 2D spectra
detecting the N\textsubscript{H\textsubscript{2}} magnetization due to cross-relaxation ($I_{CC}$) and magnetization remaining in the initial state $N\textsubscript{i}$ ($I_{BEF}$), respectively, with mixing times $t\textsubscript{m}$ of 16, 32, and 50 msec, and fitted these data with the relation $I_{CC}/I_{BEF} = \tanh\left(\frac{t\textsubscript{m}}{t_{\text{m}}}\right)$. Duplicate spectra for one of the mixing times were measured to estimate the standard deviation of the fitted cross-relaxation rates. The mean rotational correlation time $\tau_C$ of a molecule was then calculated according to Fushman et al. (1994), using the data from resolved backbone amides belonging to the RRM globular domain (residues 14–87) having $R_s/R_b$ ratio falling within one standard deviation of the mean. To test whether the determined $\tau_C$ obtained for the RRM at 1:1 and 2:1 protein:RNA ratios were the same, we used a paired one-sided $t$-test with the relation $\frac{r_i}{R_i}$ assumed to be~8 (Cieniková et al. 2014) and 4 (see following section), respectively.

### Structural modeling

Distance and angle restraints used in the molecular dynamics calculations were identical to those published for the AUUUUC–hnRNP C RRM complex (Cieniková et al. 2014). Torsion angle dynamics was performed with CYANA 3.0 (Herrmann et al. 2002) using calculations initialized with 200 random structures of the UC(U)\textsubscript{A}–RRM, the (U)\textsubscript{A}–RRM, or the UC(U)\textsubscript{A}–double-RRM complex, referred to as A, B, and C, respectively. Constraints used for the five uridine nucleotides of A(U)\textsubscript{B}C in the structure calculation described previously were assigned to the 5 nt of UC(U)\textsubscript{A}, (U)\textsubscript{A}, and the nucleotides 1–5 and 6–10 of UC(U)\textsubscript{A}. Intra-molecular constraints were the same as in the previous work. Statistical evaluations were performed on the 50 best structures ranked by the $I_{CC}/I_{BEF}$

### Estimation of RNA interface area

The relative buried surface area of RNA oligomers was calculated using the PDBBePisa service at [http://www.ebi.ac.uk/pdbe/prot_int/pistart.html](http://www.ebi.ac.uk/pdbe/prot_int/pistart.html), version 1.51 (Krissinel and Henrick 2007). Only the nucleotides contacting the protein were taken into account in the calculation. The mean and SD reported for single RRM–RNA complexes was determined from 26 structures (PDB IDs 1AUP, 1URN, 2AD9, 2ADB, 2ERR, 212Y, 2KTY, 2KG0, 2KG1, 2KHF, 2LA1, 2LEB, 2LEC, 2MBD, 2MB0, 2RQC, 2RRA, 2RS2, 2RU3, 2XS2, 2X5S, 2XS7, 3IWN, 3MOJ, 4CLO, 467); for tandem RRM–RNA complexes the coordinates from 13 structures were considered (1B7F, 1CVJ, 1FNX, 1FXL, 1G2E, 2CJY, 2MGZ, 2MKI, 2MKK, 4BS2, 4ED5, 4FO2).

### ITC measurement and data analysis

Isothermal titration calorimetry measurements were conducted in ITC buffer (50 mM Na\textsubscript{2}HPO\textsubscript{4}, 100 mM NaCl, pH 6.5) at 303 K on a VP-ITC calorimeter (MicroCal). We followed the published protocol concerning the sample handling (Cieniková et al. 2014). Data analysis was performed with the MicroCal module of Origin 7.0 (OriginLab) using the interaction models assuming either one set of independent binding sites, or two sequential sites. Binding model selection was based on minimizing the corrected Akaike information criterion (AICc) (Burnham and Anderson 2002). In the case of the sequential, 2:1 binding model, the macroscopic constants $K_i$ are expressed as

$$
\begin{cases}
K_1 = B_1 K_3 \\
K_1 K_2 = a B_2 K_3^2
\end{cases}
$$

where $i = 1$ and 2 refer to ligand:RNA stoichiometries 1:1 and 2:1, respectively. $K_3$ is the cooperativity factor at 2:1 stoichiometry, and $B_i$ the expected number of distinct molecular species binding a uridine tract of 11 nt at stoichiometry $i$, assuming that a single RRM ligand recognizes five consecutive uridines with a microscopic affinity constant $K_C$. In the context of multiregister binding, $B_1$ and $B_2$ are assumed to be~8 (Cieniková et al. 2014) and 4 (see following section), respectively.

### Model of multistoichiometric RRM binding to a finite uniform tract

The analysis of the multivalent binding of hnRNP C RRM on uridines lattices is an extension of the previously published model (Cieniková et al. 2014). The RRM binding consensus was defined as 5’-uUUUUU-3’, with U and u representing pockets with strong and weak preferences for uridines and w and v corresponding to the affinity penalties for U and u pocket mismatch, respectively. Analogously to the ITC analysis, the binding of two interacting protein ligands was allowed; any putative higher stoichiometries were ignored since we focused on tracts shorter than 13 uridines. Under these conditions, the saturation $r$ at tract length $m$ can be expressed as

$$
r(m, a) = \frac{B_1(m, w, v) K_3[F] + 2 a B_2(m, w, v)(K_3[F])^2}{1 + B_1(m, w, v) K_3[F] + a B_2(m, w, v)(K_3[F])^2},
$$

where $K_3$ and $a$ are the affinity and the cooperativity factors defined in the previous paragraph, $[F]$ the concentration of the free protein ligand and $B_i$ the affinity-weighted number of distinct molecular species at stoichiometry $i = [1, 2]$. The expression for $B_1$ was described previously (Cieniková et al. 2014); the expression for $B_2$ is

$$
B_2(1) = 0 \\
B_2(2) = v^{-1} w^{-2} \\
B_2(3) = v^{-1} w^{-2} + 2 v^{-1} w^{-6} \\
B_2(4) = v^{-1} w^{-2} + 2 v^{-1} w^{-6} + 3 v^{-1} w^{-5} \\
B_2(5) = v^{-1} w^{-2} + 2 v^{-1} w^{-6} + 3 v^{-1} w^{-5} + 4 v^{-1} w^{-4} \\
B_2(6) = B_2(5) + w^{-4} + 4 v^{-1} w^{-3} \\
B_2(7) = B_2(5) + 2 w^{-4} + w^{-3} + 5 v^{-1} w^{-3} + 3 v^{-1} w^{-2} \\
B_2(8) = B_2(5) + 3 w^{-4} + 2 w^{-3} + w^{-2} + \sum_{j=0}^{3} 2 j v^{-1} w^{j} \\
B_2(m) = B_2(5) + \sum_{j=1}^{m-9} j + \sum_{j=1}^{4} (j + m - 9) w^{j} \\
+ \sum_{j=0}^{3} (2j + m - 8) v^{-1} w^{j} \text{ for } m \geq 9
$$

The scaled saturation function $s \times r(m)$ was fitted to the experimental hnRNP C cross-link saturation $r_{exp}(m)$ for $m = [1, 11]$. 

www.rajournal.org 1939
\( r_{exp}(m) \) was calculated as the ratio of cross-link distribution on uridine \( m \)-mers, over the transcriptome-wide uridine \( m \)-mer frequency (Fig. 3B in König et al. 2010). The values of the \( w \), \( v \) and \( K_i[F] \) parameters were previously determined and set to 19.6, 5.1, and 0.33, respectively (Cieniková et al. 2014); only the cooperativity \( \alpha \) and the scaling factor \( s \) were fitted.

The probability of binding in a given register within a uridine tract was calculated as

\[
p(\text{register } q) = \frac{[P,R]^i}{\sum_{j=1}^{2}[P,R]^j},
\]

with \( i \leq 2 \) the ligand/RNA stoichiometry. For the 1:1 species,

\[
p(q_1) \sum_{j=1}^{2}[P,R] = \frac{K_i[F]}{W_q}.
\]

where \( W_q = f(w,v) \) is the affinity penalty of the register \( q \). At \( i = 2 \), for species defined by registers \( \{q,a\} \), the probability is

\[
p(q_2) \sum_{j=1}^{2}[P,R] = \frac{\alpha(K_i[F])^2}{W_q W_u}.
\]

Dynamic binding behavior of the 5′ ligand was implemented as a Gaussian distribution of binding probability \( p(q_2) \), centered on the position \( q_2 \), with SD \( \sigma \) minimizing the residual sum of squares between the model and the data.

**In vivo assays**

The following plasmids were ordered from BCCM/LMBP (http://bccm.belspo.be/bccm/lmbp.htm): pCAGGS-Flag-hnRNPC1 expression vector (acc. no. 5271), pSV-Sport-di-RF (5253) negative control vector carrying the Firefly and Renilla luciferases, and pSV-Sport-di-RF-UNR (5247) carrying the unr IRES inserted between the luciferases (Cornelis et al. 2005; Schepens et al. 2007). The Flag-hnRNPC1 was recloned into pcDNA3.1 + vector using HindIII/Xhol restriction sites. The silencing duplex RNAs were purchased from Dharmacon: The siRNA targeting hnRNP C was designed as previously reported (Cornelis et al. 2005; Schepens et al. 2007).

**SUPPLEMENTAL MATERIAL**

Supplemental material is available for this article.

**ACKNOWLEDGMENTS**

Research of FH-TA is supported by the Swiss National Science Foundation (SNF), grant numbers 3100A0-118118, 31003ab-133134, and 31003A-149921. We thank K. Zarnack and J. Ule for providing and discussing the iCLIP data.

**REFERENCES**

Afroz T, Skrisiowska L, Belloc E, Guillen-Boixet I, Méndez R, Allain FH-T. 2014. A fly trap mechanism provides sequence-specific RNA recognition by CPEB proteins. *Genes Dev* 28: 1498–1514.

Banerjee H, Rahn A, Davis W, Singh R. 2003. Sex lethal and U2 small nuclear ribonucleoprotein auxiliary factor (U2AF65) recognize polypyrimidine tracts using multiple modes of binding. *RNA* 9: 88–99.

Barnett SF, Friedman DL, LeStourgeon WM. 1989. The C proteins of Hela 40S nuclear ribonucleoprotein particles exist as anisotropic tetramers of (C13) C2. *Mol Cell Biol* 9: 492–498.

Burnham KP, Anderson DR. 2002. *Model selection and multimodel inference: a practical information-theoretic approach*. Springer, New York.

Carroll KL, Ghirlando R, Ames JM, Corden JL. 2007. Interaction of yeast RNA-binding protein Nrd1 and Nab3 with RNA polymerase II terminator elements. *RNA* 13: 361–373.

Choi YD, Dreyfuss G. 1984. Isolation of the heterogeneous nuclear RNA-ribonucleoprotein complex (hnRNP): a unique supramolecular assembly. *Proc Natl Acad Sci USA* 81: 7471–7475.

Cieniková Z, Damberger FF, Hall J, Allain FH-T, Maris C. 2014. Structural and mechanistic insights into poly(uridine) tract recognition by the hnRNP C RNA recognition motif. 1999 *J Am Chem Soc* 136: 14536–14544.

Cornelis S, Tinton SA, Schepens B, Bruynooghe Y, Beyaert R. 2005. UNR translation can be driven by an IRES element that is negatively regulated by polypyrimidine tract binding protein. *Nucleic Acids Res* 33: 3095–3108.
Wan L, Kim JK, Pollard VW, Dreyfuss G. 2001. Mutational definition of RNA-binding and protein-protein interaction domains of heterogeneous nuclear RNP C1. J Biol Chem 276: 7681–7688.

Wang C, Palmer AG III. 2003. Solution NMR methods for quantitative identification of chemical exchange in 15N-labeled proteins. Magn Reson Chem 41: 866–876.

Wang X, Tanaka Hall TM. 2001. Structural basis for recognition of AU-rich element RNA by the HuD protein. Nat Struct Biol 8: 141–145.

Wang H, Zeng F, Liu Q, Liu H, Liu Z, Niu L, Teng M, Li X. 2013. The structure of the ARE-binding domains of Hu antigen R (HuR) undergoes conformational changes during RNA binding. Acta Crystallogr D Biol Crystallogr 69: 373–380.

Whitson SR, LeStourgeon WM, Krezel AM. 2005. Solution structure of the symmetric coiled coil tetramer formed by the oligomerization domain of hnRNP C: implications for biological function. J Mol Biol 350: 319–337.

Zaidi S, Malter J. 1995. Nucleolin and heterogeneous nuclear ribonucleoprotein C proteins specifically interact with the 3′-untranslated region of amyloid protein precursor mRNA. J Biol Chem 270: 17292–17298.

Zarnack K, König J, Tajnik M, Martincorena I, Eustermann S, Stévant I, Reyes A, Anders S, Luscombe NM, Ule J. 2013. Direct competition between hnRNP C and U2AF65 protects the transcriptome from the exonization of Alu elements. Cell 152: 453–466.