Conserved binding of GCAC motifs by MEC-8, couch potato, and the RBPMS protein family

HEDDY SOUFARI1,2 and CAMERON D. MACKERETH1,2

1University of Bordeaux, Institut Européen de Chimie et Biologie, F-33607 Pessac, France
2Inserm U1212, CNRS UMR 5320, ARNA Laboratory, F-33076 Bordeaux, France

ABSTRACT

Precise regulation of mRNA processing, translation, localization, and stability relies on specific interactions with RNA-binding proteins whose biological function and target preference are dictated by their preferred RNA motifs. The RBPMS family of RNA-binding proteins is defined by a conserved RNA recognition motif (RRM) domain found in metazoan RBPMS/Hermes and RBPMS2, Drosophila couch potato, and MEC-8 from Caenorhabditis elegans. In order to determine the parameters of RNA sequence recognition by the RBPMS family, we have first used the N-terminal domain from MEC-8 in binding assays and have demonstrated a preference for two GCAC motifs optimally separated by >6 nucleotides (nt). We have also determined the crystal structure of the dimeric N-terminal RRM domain from MEC-8 in the unbound form, and in complex with an oligonucleotide harboring two copies of the optimal GCAC motif. The atomic details reveal the molecular network that provides specificity to all four bases in the motif, including multiple hydrogen bonds to the initial guanine. Further studies with human RBPMS, as well as Drosophila couch potato, confirm a general preference for this double GCAC motif by other members of the protein family and the presence of this motif in known targets.

Keywords: ITC; NMR; X-ray crystallography; protein–RNA; structure

INTRODUCTION

The regulation of mRNA processing involves protein factors that recognize specific RNA elements in order to modulate various properties of the mature mRNA. RNA-binding proteins mediate transport of the final mRNA to the cytoplasm and can alter its stability or define a precise sub-cellular localization. Initial processing steps of the pre-mRNA are also targets of specific protein factors. During splicing, the definition of intron boundaries involves specific recognition of intronic RNA sequences by constitutive splicing factors. An additional set of protein factors can be expressed during development or in response to an external stimulus to alter the inclusion or exclusion of exons to change mRNA stability and function, as well as the function of the encoded protein isoforms. In order to understand the mechanism and target selection by these RNA-binding proteins, both the protein factors and RNA motif elements need to be characterized at the molecular level.

One set of RNA-binding proteins, the RBPMS family, is named after the vertebrate RNA-binding protein gene with multiple splicing (RBPMS) (Shimamoto et al. 1996). The first member cloned was the couch potato gene from Drosophila (Bellen et al. 1992a) followed by the mec-8 gene from C. elegans (Lundquist et al. 1996). Vertebrate RBPMS and the paralog RBPMS2 are expressed in several tissues including heart, muscle, kidney, liver, lung, eye, adipose, and ovary (Wilmore et al. 2005; Kwong et al. 2010; Farazi et al. 2014). RBPMS affects synapse density and axon formation in the retina ganglion cells (Piri et al. 2006; Hornberg et al. 2013; Rodriguez et al. 2014) and colocalizes with NonO, PSF, and G3BP1 in mouse neuron cells (Furukawa et al. 2015). Expression of RBPMS2 in the intestine may play a role in motility disorders (Notarnicola et al. 2012) and gastrointestinal tumors (Hapkova et al. 2013). The connection to cancer and cell cycle control is also suggested by an interaction with cFOS to regulate AP-1 target genes (Fu et al. 2015). A role in heart development was observed for chicken RBPMs and the Xenopus orthologs (Gerber et al. 1999, 2002) and Xenopus RBPMS/Hermes is further implicated in early oocyte maturation (Chang et al. 2004; Zearfoss et al. 2004; Song et al. 2007; Aguero et al. 2016).

The founding member of the RBPMS family is couch potato from Drosophila, originally identified in a genetic screen.
for components involved in neuron differentiation and in particular the sensory organs of the peripheral nervous system (Bellen et al. 1992b). Similar to human RBPMS, the *couch potato* gene exhibits alternative splicing and is relatively large (>100 kb). Flies with a mutant *couch potato* allele are less active (Bellen et al. 1992a,b) and display other neurological defects (Glasscock and Tanouye 2005). In addition, natural variation in the gene may underlie seasonal and geographical adaptation to environmental stress in several *Drosophila* species (Schmidt et al. 2008; Salminen et al. 2015).

Finally, the RBPMS family member in *C. elegans*, *mec-8* (mechanosensory abnormality gene 8), was first described following the characterization of touch-insensitive mutants (Chalfie and Sulston 1981). Subsequently, *MEC-8* was found to regulate the alternative splicing of the perlecanc homolog *unc-52* in the hyperdermis (Lundquist and Herman 1994; Spike et al. 2002) and an isoform of the stomatin homolog *mec-2* required for function of the six mechanosensory neurons (Calixto et al. 2010).

**FIGURE 1.** Characterization of *MEC-8* RRM1 dimer and ligand sequence. (A) RNA recognition motif (RRM) domains within *C. elegans* *MEC-8* (UniProt G5ECJ4), *Drosophila melanogaster* *couch potato* (CPO; UniProt Q01617), human RBPMS (UniProt Q93062), and human RBPMS (UniProt Q6ZRY4). Residue numbers are indicated. Note that all four proteins exist as multiple splice isoforms, with only the N-terminal RRM domain common to all *MEC-8*, *couch potato*, and RBPMS variants. (B) Sequence alignment of the conserved RRM domain of the RBPMS family. Regions of β-strands (black arrows) and α-helices (white cylinders) are indicated above the *MEC-8* sequence, based on the crystal structure of the unbound *MEC-8* RRM1 domain. Residues involved in dimerization are indicated with asterisks. Residue sidechains involved in ligand binding are indicated with a black circle, and interactions with backbone atoms have a white circle. Identical residues in all four sequences are highlighted in black and those that are homologous by sidechain properties are shaded in gray. (C) NMR spectroscopy characterization based on residue-specific 15N relaxation measurements of T1 (light green), T2 (purple), the combined ratio of T1/T2 and [1H]15N heteronuclear NOE. Predicted values of monomeric and dimeric RRM1 are indicated by dashed and solid lines, respectively (Daragan and Mayo 1997). Values of [1H]15N heteronuclear NOE below 0.6 correspond to backbone amides with significant conformational disorder. (D,E) Structure overview and close-up of the dimerization interface of the unbound *MEC-8* RRM1 dimer (PDB 5BJR). (F) Relative affinity of ligands with individual nucleotide substitutions binding to *MEC-8* RRM1 by using isothermal titration calorimetry (ITC). The heights of the A, C, G, and U in each position are relative to the determined values of the association constant, $K_a$. Representative raw data and derived $K_a$ values can be found in Supplemental Figure S2. (G) Representative data from ITC measurements of *MEC-8* RRM1 with the RNA ligands AGACA and AGCAGAUUUUUAGCACA.
Agüero et al. 2016; Teplova et al. 2016). Disruption of the domain interface in chicken RBPM2 abolished normal protein function related to intestinal development, and in vitro the mutant protein showed reduced binding to a segment of NOGGIN mRNA (Sagnol et al. 2014). Dimerization mutants of human RBPM5 failed to localize to arsenite-triggered stress granules, and similar to RBPM2, reduced RNA binding was observed by the mutant (Teplova et al. 2016). Although the residues involved in dimerization are further conserved in MEC-8 and couch potato, the oligomeric state of these proteins had not been investigated.

In terms of the RNA sequences recognized by RBPM5, a PAR-CLIP approach identified RNA targets with multiple CAC motifs separated by varying lengths (Farazi et al. 2014). Additional electrophoresis mobility shift assays demonstrated in vitro interaction of RBPM5 and RBPM2 with AC-rich RNA ligands, as well as oligonucleotides based on wild-type and mutated segments of mRNA from UBE2V1, SRM, ETF1, and NDUA6. A crystal structure of the RRM domain of human RBPM5 in complex with a short RNA confirmed recognition of the trinucleotide CAC (Teplova et al. 2016). In contrast to these observations, a complementary approach involving RNA-compete identified a longer 4-nucleotide (nt) GCAC motif recognized by other members of the RBPM5 family, namely couch potato from Drosophila and MEC-8 from C. elegans (Ray et al. 2013).

In this study, we use crystal structures and binding data from isothermal titration calorimetry to show that the N-terminal RRM domain from MEC-8 is a dimer and interacts with highest affinity to a double GCAC motif. Additional experiments on the RRM domains from couch potato and RBPM5 demonstrate shared binding preferences by this family of RNA-binding proteins.

RESULTS AND DISCUSSION

Based on a sequence alignment of MEC-8, Drosophila couch potato, human RBPM5 and human RBPM2 (Fig. 1B), we have cloned the first RRM domain (RRM1) from MEC-8 as a construct from residues 28–123 (hereafter MEC-8 RRM1). Characterization of the purified protein by NMR spectroscopy reveals that MEC-8 RRM1 forms a stable and soluble dimer in solution, similar to the situation with human RBPM5 and RBPM2 (Sagnol et al. 2014; Teplova et al. 2016). Specifically, NMR spectroscopy allows for the estimate of molecular size from measurement of the relaxation properties of the amide nitrogens in the protein backbone. The T1 and T2 $^{15}$N relaxation values (Fig. 1C) are consistent with the values predicted for a dimer (solid lines) as opposed to values for a monomer (dotted lines) (Daragan and Mayo 1997). The measurements also allow for an approximate calculation of molecular size by using the ratio of T1 over T2 to give an average value of 26 ± 7 kDa, again close to the calculated dimer mass of 22 kDa. As a final characterization by NMR spectroscopy, we looked at conformational disorder of the protein backbone by measuring $^{15}$N heteronuclear NOE, which reveals that residues from Val117 to the C terminus are highly flexible (values <0.6). We therefore made a second construct in which the last six residues were removed (MEC-8 RRM128–117) without otherwise perturbing the structure or dimerization. This second construct was successful in producing a crystal that diffracted to 2.6 Å resolution (complete statistics in Materials and Methods). The atomic details of the MEC-8 RRM128–117 dimer reveal an overall interface involving residues Asp41, Lys43, Arg45, Glu46, Tyr48, Leu49, Leu50, and Arg52 of helix α1, and residues Val96, Phe98, Asp99, and Pro100 from the loop between helix α2 and strand β4 (Fig. 1E). This dimer interface is consistent with that observed for both human RBPM5 (Teplova et al. 2016) and human RBPM2 (Sagnol et al. 2014) and confirms that MEC-8 forms a similar dimer arrangement to other members of the RBPM5 family. Furthermore, we did not observe any evidence of a monomer in the concentrations used for NMR spectroscopy or ITC, and in contrast to RBPM5 and RBPM2, attempts to disrupt dimerization via comparable interface mutations (Arg45Glu or Leu49Glu) instead resulted in a loss of the protein fold (Supplemental Fig. S1). Dimer formation for MEC-8 may therefore be required, at least in vitro, for stabilizing the RRM1 domains.

In order to obtain atomic information on RNA binding by MEC-8 RRM1, an initial aim was to define the RNA sequence elements necessary for a high affinity ligand. The first goal was to define the optimal RNA binding sequence for one monomer of the MEC-8 RRM1 dimer. The MEC-8 protein was previously found to recognize a GCAC motif based on the RNA compete method (Ray et al. 2013) and a shorter CAC motif was found to be enriched in targets of other RBPM5 family members by using the PAR-CLIP approach (Farazi et al. 2014). To better define the affinity of various sequences to MEC-8 RRM1, we used isothermal titration calorimetry (ITC) and a series of 6-nt ligands derived from a central GCAC motif. Individual measurements used ligands with all four bases at the 5′ and 3′ positions (Amrane et al. 2014). In addition, each of the GCAC bases was sequentially mutated to the other three bases in order to determine the importance of each nucleotide in the motif. The resulting ITC measurements therefore illustrate the relative effect on affinity for each base in the six positions (Fig. 1F; Supplemental Fig. S2). From these results, MEC-8 RRM1 does not appear to have selectivity for the first or last position in the hexamer sequences. In contrast, the CAC motif is clearly required for high affinity binding, preceded by a moderate preference for guanine in the second position of the tested ligands.

The dimeric nature of MEC-8 RRM1 adds another important aspect to RNA recognition, such that the highest affinity ligands are expected to contain two copies of this GCAC motif. We therefore used ITC to test the dimeric MEC-8 RRM1 with a series of longer RNA ligands in which two GCAC motifs are separated by 4–14 nt (Table 1; Supplemental Fig. S3). Simple extension of the AGCAC 6-mer by four uracils to
the 5′ or 3′ end resulted in only minor changes in affinity (2.3 ± 0.3 μM and 1.75 ± 0.03 μM, respectively, as compared to 3.2 ± 0.3 μM), whereas all of the double motif ligands from the 4-nt spacer to the 14-nt spacer bound with higher affinity than the KD for the single GCAC motif. As expected, the stoichiometry of the complexes shifted from a 1:1 binding to a 1:2 ratio, with one RNA ligand recognized by a dimer of MEC-8 RRM1 (Fig. 1G; a 0.5:1 ratio of RNA ligand per MEC-8 monomer, versus 1:1 for the shorter AGCACA). The highest affinity ligands contain a linker of eight or more bases (KD values of 80–150 nM) and we therefore prepared crystallization screens with a variety of these oligonucleotides.

Successful crystallization and structure determination at 1.53 Å resolution was obtained for a complex containing the shorter MEC-8 RRM1 construct (residues 28–117) and the DNA version of the 10-nt spacer ligand (AGCACA). The overall structure and domain orientation of the MEC-8 RRM128–117 dimer (Fig. 2A) is similar to the arrangement of the unbound structure (Fig. 1D), with an rmsd of 0.6 Å. Density was observed for nucleotides that correspond to each of the two AGCAC sequences within the ligand. In contrast, nine linker nucleotides are not visible, with an rmsd of 0.6 Å. Density was observed for nucleotides similar to the arrangement of the unbound structure (Fig. 1D).

In terms of atomic details, Ade1 and Ade15 stack against the following guanine but do not otherwise directly contact the protein surface. In contrast, specific binding of Gua2 or Gua16 is due to four hydrogen bonds each to MEC-8. A common set of hydrogen bonds connects the guanine N2 atom to both the backbone carbonyl of Ser73 and the sidechain hydroxyl of Thr65 (Fig. 2B). Due to a slight change in conformation between the bound guanines, the N1 atoms in Gua2 and Gua16 interact with either the backbone carbonyl or sidechain hydroxyl of Ser73, respectively. Another difference is that Gua2 has a hydrogen bond between N7 and the sidechain of Arg107, whereas in Gua16 this fourth hydrogen bond is between O2 and the backbone amide of Gly37. For the remaining CAC trinucleotide, the contacts are mainly similar on each MEC-8 RRM1 monomer. Cyt5/Cyt17 stack on Phe34 and are further stabilized by hydrogen bonds to the sidechain of Glu109 and backbone atoms of Leu110 and Lys112. Ade4/Ade18 stack on Phe77 with a hydrogen bond to the sidechain of Thr65 (Fig. 2B). Due to a slight change in conformation between the bound guanines, the N1 atoms in Gua2 and Gua16 interact with either the backbone carbonyl or sidechain hydroxyl of Ser73, respectively. Another difference is that Gua2 has a hydrogen bond between N7 and the sidechain of Arg107, whereas in Gua16 this fourth hydrogen bond is between O2 and the backbone amide of Gly37. For the remaining CAC trinucleotide, the contacts are mainly similar on each MEC-8 RRM1 monomer. Cyt5/Cyt17 stack on Phe34 and are further stabilized by hydrogen bonds to the sidechain of Glu109 and backbone atoms of Leu110 and Lys112. Ade4/Ade18 stack on Phe77 with a hydrogen bond to the backbone carbonyl of Ser73.

A comparison with the previously determined structure of RBPMS bound to a UCAC sequence (Teplova et al. 2016) reveals a high similarity in the recognition mode of the common CAC trinucleotide (Fig. 2C). In contrast, there is a significant difference in the position and degree of hydrogen bonding toward the initial guanine in the MEC-8 complex versus the corresponding uracil in the structure of bound RBPMS (Fig. 2D). In RBPMS, the uracil in each model is stabilized by only one direct hydrogen bond (<3.4 Å) as compared to the four hydrogen bonds between the guanine

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**Table 1. ITC data for MEC-8 RRM1**

| RNA ligand | Sequence | KD (µM) | N | ΔH (kJ/mol) | ΔS (J/mol) |
|------------|----------|---------|---|-------------|-------------|
| 6-mer      | AGCACA   | 3.2 (0.3) | 0.96 (0.06) | −18.8 (0.7) | −11.3 (0.8) |
| 4U + 6-mer | UUUUAGCACA | 2.3 (0.3) | 1.03 (0.05) | −17 (1) | −10 (1) |
| 6-mer + 4U | AGCACAUUU | 1.75 (0.03) | 1.13 (0.08) | −18 (1) | −10 (1) |
| 4-nt spacer | AGCACAUUUAGCACA | 0.38 (0.09) | 0.52 (0.02) | −17.3 (0.6) | −8.5 (0.8) |
| 6-nt spacer | AGCACAUUUAGCACA | 0.33 (0.01) | 0.47 (0.03) | −16.8 (0.0) | −7.9 (0.0) |
| 8-nt spacer | AGCACAUUUUGAGCACA | 0.18 (0.04) | 0.48 (0.01) | −14 (0.1) | −5 (1) |
| 10-nt spacer | AGCACAUUUUGAGCACA | 0.15 (0.04) | 0.53 (0.01) | −15.3 (0.9) | −6.0 (0.9) |
| 12-nt spacer | AGCACAUUUUGAGCACA | 0.08 (0.02) | 0.51 (0.00) | −17.5 (0.8) | −7.8 (0.9) |
| 14-nt spacer | AGCACAUUUUGAGCACA | 0.11 (0.07) | 0.52 (0.06) | −18.9 (0.6) | −9 (1) |
| 10-nt spacer (G2U/G16U) | AUCACAUUUUGAAGCACA | 1.29 (0.04) | 0.77 (0.03) | −23.5 (0.6) | −15.4 (0.6) |
| 10-nt spacer (G2A/G16A) | AAGCACAUUUUGAAGCACA | 0.9 (0.1) | 0.60 (0.01) | −17.4 (0.5) | −9.1 (0.5) |
| 10-nt spacer (G2C/G16C) | AACCACAUUUUGAAGCACA | 1.56 (0.01) | 0.60 (0.04) | −18.3 (0.2) | −10.4 (0.2) |
| 10-nt spacer (WT/G16U) | AGCACAUUUUGAAGCACA | 0.26 (0.08) | 0.60 (0.07) | −23.7 (0.7) | −18.3 (0.6) |
| 10-nt spacer (WT/C19U) | AGCACAUUUUGAAGCACA | 0.45 (0.03) | 0.55 (0.08) | −8.4 (0.1) | −3.8 (0.1) |
| 6-mer (G2U) | AGCACA | 20.4 (0.8) | 1.00 | −13 (2) | −7 (2) |
| 6-mer (C5U) | AGCACA | nd | nd | nd | nd |
| U1A stem-loop | AUAUGACUCGCCGAUUU | 0.24 (0.02) | 0.81 (0.17) | −2.7 (0.2) | 6.4 (0.2) |

*Values represent the mean of two or more independent measurements, with standard error in parentheses. Representative raw data can be found in Supplemental Figure S3.

For these longer ligands, a low affinity binding site also exists from single-site binding.

Stoichiometry set at 1:1 for fitting.

(nl) Not determined due to insufficient signal.
and residues within MEC-8. This raises the question as to the general importance of the guanine in the RNA motif recognized by MEC-8 RRM1 and for the rest of the RBPMs family. To address this aspect, we first determined the effect of mutating both guanine bases in the 10-nt-spacer RNA ligand. Mutation of both guanine bases to uracil (UCAC), adenine (ACAC), or cytosine (CCAC) reduced the affinity by a factor of six to ten (K_D values of 1.29 ± 0.04 µM, 0.9 ± 1.56 µM and 1.56 ± 0.01 µM, respectively, as compared to a K_D of 0.15 ± 0.04 µM for the ligand with guanine; Table 1). In contrast, maintaining at least one GCAC motif ensures relatively high affinity even when the second motif was mutated to the moderately or severely reduced-binding UCAC or GCA sequences, respectively (Table 1).

To determine whether the RNA binding preferences exhibited by MEC-8 RRM1 were shared with other members of the RBPMs family, we repeated the ITC experiments with the RRM domains from D. melanogaster couch potato (residues 447–543) and human RBPMs (residues 11–111). Starting with the tandem GCAC ligands, both couch potato and RBPMs RRM domains displayed highest affinity when the two motifs were separated by 10–12 nt (Table 2; Supplemental Figs. S4, S5). The common mode of binding between these three members of the RBPMs family is clear when the individual K_D values are normalized for each protein based on the affinity to the 6-mer AGCACA ligand (Fig. 2E). This binding similarity also extends to thermodynamic trends for the spacer-length series of RNA ligands. For all three proteins, the enthalpy of binding (ΔH) becomes less exothermic from the 4-nt to the 8-nt spacer ligand (Tables 1, 2), at which point it reverses and increasing linker length binds with more negative ΔH. This pattern is
TABLE 2. ITC data for couch potato RRM and RBPMs RRM

| RNA ligand       | Sequence                  | $K_D$ (µM) | $N$   | $\Delta H$ (kJ/mol) | $\Delta S$ (J/mol) |
|------------------|---------------------------|------------|-------|---------------------|-------------------|
| Couch potato     | AGCAC                     | 1.09 (0.09)$^a$ | 0.95 (0.05) | $-22.1 (0.9)$ | $-14.0 (0.8)$ |
| 6-mer            | AGCACUUULU                | 0.6 (0.1)  | 1.05 (0.12) | $-20.9 (0.0)$ | $-12.3 (0.1)$ |
| 4U + 6-mer       | UUUUAGCACA               | 0.50 (0.03) | 1.00 (0.02) | $-19.1 (1.1)$ | $-10.1$ |
| 4-nt spacer      | AGCACAUUAGCACA           | 0.12 (0.04) | 0.49 (0.10) | $-24 (3) $    | $-14 (3)$ |
| 6-nt spacer      | AGCACAUUUUAGCACA         | 0.16 (0.01) | 0.49 (0.06) | $-22.4 (0.3)$ | $-13.1 (0.4)$ |
| 8-nt spacer      | AGCACAUUUUUUAGCACA       | 0.11 (0.00) | 0.52 (0.01) | $-22 (2)$     | $-12 (2)$ |
| 10-nt spacer$^a$| AGCACAUUUUUUUAGCACA       | 0.05 (0.01) | 0.51 (0.09) | $-24.1 (0.5)$ | $-14.2 (0.4)$ |
| 12-nt spacer$^a$| AGCACAUUUUUUUUUAGCACA    | 0.08 (0.01) | 0.50 (0.06) | $-27 (3)$     | $-17 (3)$ |
| 14-nt spacer$^a$| AGCACAUUUUUUUUUUUUUAGCACA | 0.14 (0.09) | 0.52 (0.06) | $-30 (2)$     | $-21 (2)$ |
| 10-nt spacer (G2U/G16U) | UGCACAUUUUUUUUUUACUCA   | 0.37 (0.06) | 0.63 (0.00) | $-24.9 (0.1)$ | $-16.1 (0.2)$ |
| 10-nt spacer (G2A/G16A) | AACACAUUUUUUUUACCACA   | 0.34 (0.08) | 0.54 (0.03) | $-19.9 (0.4)$ | $-11.0 (0.2)$ |
| 10-nt spacer (G2C/G16C) | ACCACAUUUUUUACCA       | 0.68 (0.01) | 0.52 (0.00) | $-21 (1)$     | $-12 (1)$ |
| 10-nt spacer (WT/G16U) | AGCACAUUUUUUUUACUCA    | 0.09 (0.01) | 0.55 (0.03) | $-32.0 (0.7)$ | $-22.5 (0.6)$ |
| 10-nt spacer (WT/C19U) | AGCACAUUUUUUUUAGCAUA   | 0.21 (0.05) | 0.54 (0.06) | $-14.7 (0.9)$ | $-6 (1)$ |
| U1A stem-loop   | AUUCCAUUGCACUCCGGAUUU   | 0.87 (0.02) | 0.92 (0.20) | $-7.3 (0.8)$  | $0.9 (0.9)$ |
| RBPMs           | AGCAC                     | 10 (1)     | 0.97 (0.01) | $-12 (2)$     | $-5 (2)$ |
| 6-mer            | AGCACUUULU               | 7 (2)      | 0.9 (0.2)   | $-9 (3) $     | $-2 (4)$ |
| 4U + 6-mer       | UUUUAGCACA              | 10 (1)     | 1.00$^b$    | $-10 (1)$     | $-4 (1)$ |
| 4-nt spacer      | AGCACAUUAGCACA           | 4.7 (0.7)  | 0.54 (0.04) | $-6.3 (0.5)$  | $0.9 (0.5)$ |
| 6-nt spacer      | AGCACAUUUUAGCACA         | 1.0 (0.2)  | 0.71 (0.06) | $-3.5 (0.7)$  | $4.7 (0.5)$ |
| 8-nt spacer      | AGCACAUUUUUUAGCACA       | 0.9 (0.2)  | 0.48 (0.01) | $-2.9 (0.2)$  | $5.4 (0.3)$ |
| 10-nt spacer     | AGCACAUUUUUUUUUAGCACA    | 0.6 (0.1)  | 0.55 (0.06) | $-4.7 (0.6)$  | $3.8 (0.6)$ |
| 12-nt spacer     | AGCACAUUUUUUUUUUUAGCACA  | 0.5 (0.1)  | 0.47 (0.01) | $-7.2 (0.7)$  | $1.4 (0.7)$ |
| 14-nt spacer     | AGCACAUUUUUUUUUUUUUAGCACA | 1.5 (0.1)  | 0.5 (0.1)   | $-9 (2)$     | $-1 (2)$ |
| 10-nt spacer (G2U/G16U) | UGCACAUUUUUUUUUUACUCA   | 1.9 (0.3)  | 0.73 (0.03) | $-17.5 (0.8)$ | $-9.7 (0.9)$ |
| 10-nt spacer (G2A/G16A) | AACACAUUUUUUUUACCACA   | 9 (4)      | 0.65 (0.03) | $-10.4 (0.5)$ | $-3.3 (0.8)$ |
| 10-nt spacer (G2C/G16C) | ACCACAUUUUUUACCA       | 4 (1)      | 0.70 (0.00) | $-10 (2)$     | $-3 (2)$ |
| 10-nt spacer (WT/G16U) | AGCACAUUUUUUUUUUACUCA  | 0.7 (0.1)  | 0.69 (0.08) | $-20 (2)$     | $-12 (2)$ |
| 10-nt spacer (WT/C19U) | AGCACAUUUUUUUUUAGCAUA | nd$^d$     | nd         | nd           | nd |
| U1A stem-loop   | AUUCCAUUGCACUCCGGAUUU   | nd         | nd         | nd           | nd |

$^a$Values represent the mean of two or more independent measurements, with standard error in parentheses. Representative raw data can be found in Supplemental Figures S4, S5 for couch potato and RBPMs, respectively.

$^b$For these longer ligands, a low affinity binding site also exists from single site binding.

$^c$Stoichiometry set at 1:1 for fitting.

$^d$(nd) Not determined due to insufficient signal.

countered by the entropic term ($T\Delta S$) which is most unfavorable for the shortest and longest ligands and highest for the 8-nt linker.

RNA binding preference for the guanine is also conserved among the three proteins. Similar to the situation with MEC-8, mutation of the two guanine bases to uracil (UCAC), adenosine (ACAC), or cytosine (CCAC) in the 10-nt linker RNA caused a reduction in binding by a factor of 7–14 for couch potato RRM, and by a factor of three to 15 for RBPMs RRM (Table 2). With respect to RBPMs/Hermes, evidence for in vivo importance of the guanine in a GCAC motif was previously suggested by the study of the 240-nt Xenopus oocyte mitochondrial cloud localization signal (MCLS). The MCLS contains six repeats of a UCAC sequence and a mutation of all six to UUCAC prevents proper RNA localization of Xcat2 RNA to the germinial granules (Chang et al. 2004). The localization of nanos RNA to germinal granules is also dependent on the MCLS, and colocalization studies demonstrate that both nanos RNA and RBPMs/Hermes accumulate in similar clusters (Aguero et al. 2016). In addition, it was shown that intact UGCAC repeats of the MCLS are required for interaction with RBPMs/Hermes.

Finally, it had been previously noted that molecular similarities exist between RNA binding of CAC sequences by RBPMs and the first RRM of the U1A protein (Teplova et al. 2016). The additional contacts to the initial guanine increase this similarity. Structural details of U1A bound to the loop of hairpin II of U1 snRNA (Oubridge et al. 1994) or the U1A 3’UTR polyadenylation inhibition element (Allain et al. 1996, 1997; Varani et al. 2000) illustrate a similar RNA bound conformation for all four bases in the GCAC motif as compared to MEC-8. We therefore selected one of the U1 hairpin II RNA ligands (Fig. 2E; Oubridge et al. 1994) for assessment of binding to MEC-8 RRM1. Under the same conditions as the previous ITC measurements, this stem–loop RNA indeed bound with high affinity to MEC-8 RRM1, with a $K_D$ of 0.24
± 0.02 μM, comparable to that of the tandem GCAC ligand (Table 1) although not as tight as the interaction between U1A and the stem–loop (Lutz-Freyermuth et al. 1990; Hall and Stump 1992). Binding to couch potato RRM was relatively weaker, but still with higher affinity than to the short AGCACA RNA (Table 2; Fig. 2E). The affinity to RBPMs could not be determined due to insufficient signal to noise. Nevertheless, it is evident that at least MEC-8 and couch potato from the RBPMs family are able to recognize the GCAC motif even within the context of a stem–loop structure.

In summary, we have used a combination of structure determination and binding assays to demonstrate the binding preference of MEC-8 RRM1 for GCAC motifs, and further show that optimal targets have two motifs separated by more than six bases. These results confirm the preferred GCAC motif identified for MEC-8 and couch potato by the RNA compete method (Ray et al. 2013), but differ slightly from the smaller CAC motif obtained from the in vivo assay for RBPMs targets by the PAR-CLIP approach (Farazi et al. 2014). For the in vivo targets, it is possible that additional protein partners in the cell could enhance binding affinity to less optimal sites, such as the interaction between Xenopus RBPMs/Hermes and hnRNp1 (Aguro et al. 2016). Nevertheless, some of the identified targets such as NDUFa6 and ETF1, which were further investigated by in vitro binding assays (Farazi et al. 2014; Teplova et al. 2016), do in fact fulfill the general observed requirement of at least one GCAC motif and a separation of >6 nt to a second suboptimal motif. The region of NOGGIN mRNA that interacts with RBPMs2 (nucleotides 570–730) (Sagnol et al. 2014) also contains two GCAC sequences separated by 18 bases that could serve as the RBPMs2 binding site. Extensive separation such as this could complicate analysis, as could the allowed presentation of the GCAC sequence within RNA secondary structure elements. Precise identification of in vivo sites bound by MEC-8 and couch potato will greatly add to the description of binding site determinants, and also eventually to functional understanding of the RBPMs family of RNA-binding proteins. Finally, it should be noted that we have focused on the only RRM domain that is absolutely conserved within the entire RBPMs family and additionally in all of the generated splice isoforms of RBPMs, MEC-8 and couch potato. Nevertheless, some family member isoforms (such as the MEC-8 variant shown in Fig. 1A) also possess a second C-terminal RRM domain with the ability to bind RNA. The resulting interplay regarding RNA-binding by the first and second RRM domains, and whether or not the different isoforms are specific to a particular biological process, represents a further complexity to be investigated.

MATERIALS AND METHODS

Protein expression and purification

The N-terminal RRM domain from C. elegans MEC-8 (residues 28–123 or 28–117) was amplified from a codon-optimized synthetic gene (Bio-Basic Canada) by using PCR oligos containing NcoI or Acc65I restriction enzyme sites, and inserted into a modified PET-9d plasmid encoding a His6 purification tag followed by a tobacco etch virus (TEV) protease cleavage site. To aid in sample stability, Cys54 and Cys102 were mutated to alanine. The orthologous RRM domains from human RBPMs (residues 11–111) and Drosophila couch potato (CPO; residues 447–543) were amplified from L.M.A.G.E. clone IRAUp9691214D (BioVector) and EST clone BDGP122L092Q (BioVector), respectively, and inserted into the same modified PET-9d plasmid. MEC-8 mutant proteins were generated by using PCR amplification with a set of oligos overlapping the mutation site. All proteins were expressed in Escherichia coli BL21 LysY (New England Biolabs) using LB medium or M9 minimum medium supplemented with 15NH4Cl (1 g L⁻¹) and [13C6]glucose (2 g L⁻¹). Induction with 0.5 mM IPTG (final concentration) was followed by overnight protein expression at 20°C or 30°C for LB or M9 cultures, respectively. Bacteria were collected by centrifugation and resuspended in 50 mM Tris (pH 7.5), 500 mM NaCl, 5% (v/v) glycerol, and 5 mM imidazole. Cell lysate preparation used a combination of lysozyme and sonication followed by centrifugation at 20,000g for 30 min. Protein purification from cell lysate was achieved by using Ni²⁺-affinity chromatography with buffers composed of 50 mM Tris (pH 7.5), 500 mM NaCl, 5% (v/v) glycerol, and 5, 25, and 250 mM imidazole for the loading, wash and elution, respectively. The His6 tag was removed by addition of TEV protease following a second Ni²⁺-affinity chromatography step. The purified samples were concentrated by filter centrifugation, dialyzed overnight, and quantified by measuring the absorbance at 280 nm. The RNA was synthesized by using an Expedite 8909 (PerSeptive Biosystems). The DNA oligonucleotides were commercially produced (Eurogentec and Eurofins MWG Operon).

NMR spectroscopy

Spectra were recorded at 298 K using a Bruker Avance III 700 MHz spectrometer equipped with a triple resonance-gradient standard probe. Topspin version 2.1 (Bruker BioSpin) was used for data collection. Spectra processing used NMRPipe (Delaglio et al. 1995) followed by analysis with Sparky 3 (T.D. Goddard and D.G. Kneller, University of California) or NMRViewJ 8.0 (One Moon Scientific). For backbone resonance assignment of 300 μM 13C,15N MEC-8 RRM1 the following spectra were used: 2D 1H-15NH S Q C , 3D HNCO, 3D HNCA, 3D HNHA, 3D (H)CCONH TOCSY, and 3D H(C)CONH TOCSY. All spectra were recorded in buffer containing 90% H2O/10% D2O. Amide 15N chemical shift assignments for MEC-8 RRM1 were deposited in the Biological Magnetic Resonance Data Bank (BMRB) as entry 26922.

Crystallization and structure calculation

Purified recombinant MEC-8 RRM1 was concentrated to ~10 mg/mL in 50 mM Tris (pH 7.5), 150 mM NaCl. Crystals of MEC-8 RRM1 (residues 23–117) were obtained by using 1 μL of the protein.
solution mixed with an equal volume of reservoir solution containing 0.1 M sodium acetate (pH 5.6), 0.2 M ammonium sulfate and 30% (w/v) polyethylene glycol (PEG) 4000 and using sitting drop vapor diffusion against 500 µL of reservoir solution. Crystals appeared as long plates. The ligand-bound complex was assembled by mixing the protein (10 mg/mL) with AGCACATTTTTTTT AGGACA DNA at a 2:1 molar ratio. X-shaped crystals were generated from a 2 µL 1:1 mixture of sample and reservoir buffer containing 10% (v/v) isopropanol and 10% (w/v) PEG4000 in 500 µL of reservoir. Crystal cryo-protective solution was used for 5 min in a solution composed of 6.7% (v/v) isopropanol, 6.7% (v/v) PEG4000, and 25% (v/v) glycerol. Diffraction data were collected at the European Synchrotron Radiation Facility beam line ID23-2 and processed using the program XDS (Kabsch 2010). The unbound structure was determined by molecular replacement with Phaser (McCoy et al. 2007), based on the crystal structure of the second RRM domain of yeast poly(U)-binding protein PDB 3MD1 chain A. Molecular replacement for the DNA-bound complex used the unbound MEC-8 RRM1 dimer. Phenix was used for refinement (Adams et al. 2010). A summary of data collection and structure refinement statistics is provided in Supplemental Table S1. Structure images were prepared using the program Chimera (Pettersen et al. 2004). Structural coordinates and data have been deposited in the Protein Data Bank as PDB ID codes 3BJR and 3TKZ for the unbound and ligand bound MEC-8 RRM1, respectively.

**Isothermal titration calorimetry**

ITC measurements were performed by using an ITC200 Microcalorimeter (Malvern Instruments) at 25°C with a stir rate of 500 rpm and set at high sensitivity. All proteins were dialyzed extensively against 50 mM Tris, pH 7.5, 150 mM NaCl. Buffer from the dialysis was used to solubilize the DNA oligonucleotides, or used in conjunction with NAP-5 columns (GE Healthcare Life Sciences) to change the buffer of RNA samples. The initial determination of binding preference in Figure 1F represents an average of two measurements using 160–320 µM protein in the syringe and 20–40 µM ligands in the cell, with a total of 13 injections of 3 µL following an initial injection of 1 µL. For the sequence logo, minimal binding µM ligands in the cell, with a total of 13 injections of 3 µL following an initial injection of 0.5 µL. The data were first processed following an initial injection of 1 µL. The data were first processed and the integrated data points subsequently fit by using NITPIC (Keller et al. 2012; Scheuermann and Brautigam 2015) and the integrated data points subsequently fit by using NITPIC (Keller et al. 2012; Scheuermann and Brautigam 2015).

**SUPPLEMENTAL MATERIAL**

Supplemental material is available for this article.

**ACKNOWLEDGMENTS**

We thank Gavin Collie for final preparation of crystals and synchrotron data collection. Additional assistance and advice were provided by Lionel Beaurepaire, Brice Kauffmann, Axel Innis, and Sébastien Fribourg. Brune Violet provided synthesis of the RNA oligonucleotides. Access to NMR spectrometers, equipment, and technical assistance were possible thanks to Axelle Gréard, Estelle Morvan, and the structural biology platform at the Institut Européen de Chimie et Biologie (UMS 3033). We thank the European Synchrotron Radiation Facility for access to synchrotron radiation facilities (beam line ID23-2). Financial support from the Centre Nationale de la Recherche Scientifique (IR-RMN-THC Fr3050) is gratefully acknowledged. H.S. was supported by a PhD fellowship from the Institut National de la Santé et de la Recherche (INSERM) and the Aquitaine regional government.

Received October 25, 2016; accepted December 19, 2016.

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