The third RNA recognition motif of Drosophila ELAV protein has a role in multimerization

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ABSTRACT

ELAV is a neuron-specific RNA-binding protein in Drosophila that is required for development and maintenance of neurons. ELAV regulates alternative splicing of Neuroglian and erect wing (ewg) transcripts, and has been shown to form a multimeric complex on the last ewg intron. The protein has three RNA recognition motifs (RRM1, 2 and 3) with a hinge region between RRM2 and 3. In this study, we used the yeast two-hybrid system to determine the multimerization domain of ELAV. Using deletion constructs, we mapped an interaction activity to a region containing most of RRM3. We found three conserved short sequences in RRM3 that were essential for the interaction, and also sufficient to give the interaction activity to RRM2 when introduced into it. In our in vivo functional assay, a mutation in one of the three sequences showed reduced activity in splicing regulation, underlining the functional importance of multimerization. However, RRM2 with the three RRM3 interaction sequences did not function as RRM3 in vivo, which suggested that multimerization is not the only function of RRM3. Our results are consistent with a model in which RRM3 serves as a bi-functional domain that interacts with both RNA and protein.

Such a case was initially reported for the interactions between U2 small nuclear ribonucleoprotein particle (snRNP) B′ and A′ proteins (10–13), and later structural studies, including that of other proteins, revealed that the protein–protein interactions often, but not always, occur through amino acids on the α-helices of the RRM (8,14–21). This dual interacting capability of the RRM domain might explain its prevalence.

The ELAV protein family is a conserved RNA-binding protein family found in animals from Caenorhabditis to humans. Its characteristic structure is the three RRRs (RRM1, 2 and 3) with a 'hinge' region of about 60–80 amino acids separating the second and third RRRs. Mammals have four members of the family called Hu proteins (HuR, HuB, HuC and HuD) that have been implicated in stabilization and/or translation activation of mRNAs containing AU-rich elements (AREs) (22–24). Drosophila has three ELAV-family proteins (ELAV, Rbp9 and Fne), ELAV is specifically expressed in all neurons in Drosophila, and elav null mutants show embryonic lethal phenotype with developmental defects in the nervous system (25,26). It has been shown that ELAV regulates alternative splicing of Neuroglian (Nrg) and erect wing (ewg) transcripts, by binding to their introns to produce neuron specific isoforms (27–30). Additionally, ELAV has been shown to autoregulate its own message (31–33).

Recently, ELAV has been shown to form a multimeric complex on the ewg intron RNA. It appears that ELAV in solution is mainly in a tetramer-configuration without the substrate RNA, and is assembled into a larger dodecameric complex upon binding to the target RNA (34). Mammalian ELAV family proteins have also been shown to form multimers (35–37). Therefore, it is likely that multimerization is a common feature shared by the ELAV-family proteins, and an important part of the mechanism by which they exert their effect on the bound RNA.

In this study, we show that one of the major sites for ELAV–ELAV interaction activity is located in RRM3 and a small adjacent region of the hinge. Three short sequences in RRM3 are shown to be necessary for the interaction,
MATERIALS AND METHODS

Plasmid construction

To construct plasmids for the yeast two-hybrid assay, we used pGBK7T and pGADT7 vectors (Clontech) for the GAL4 DNA-binding-, and transcription-activation-domain fusions, respectively. Both fusions were N-terminal fusions; i.e. the DNA-binding, or activation domain was fused to the N-terminus of the protein to be tested. The full-length and truncated elav coding sequences were amplified by PCR from an elav cDNA clone, digested with EcoRI and BglII, and subcloned between EcoRI and BamHI sites of pGBK7T and pGADT7 vectors.

For construction of hinge-RRM1 (eH1) and hinge-RRM2 (eH2) cDNAs, the elav RRM1 and 2 coding sequences were PCR-amplified with primers that have the 3' end sequence of the hinge region fused to the 5' region of the RRM1 or 2 coding sequence. The fused hinge sequence would anneal to the corresponding site of the hinge region during the next round of PCR. The secondary PCRs were performed in which eH1 or eH2 were amplified from mixed templates of the first PCR product and pGADT7-eH. The resulting PCR fragments were digested with EcoRI and BglII, and subcloned into EcoRI/BamHI sites of pGADT7.

The mutant hinge-RRM3 constructs (eH3mu-7) were generated by site-directed PCR mutagenesis. For each mutation, a complimentary pair of primers was designed at the target site to replace a certain RRM3 sequence with the corresponding sequence from RRM2. In the primary PCRs, the 5' and 3' parts were amplified separately using one of the complimentary mutagenic primers at one end. The PCRs produced the 5' and 3' parts of the constructs overlapping with each other for the length of the mutagenic primers. The 5' and 3' parts were joined together by the secondary PCRs in which a mixture of the two primary PCR products was used as a template. The secondary PCR fragments were digested with EcoRI and BglII, and subcloned into EcoRI/BamHI sites of pGADT7.

The eH3mu8 was constructed in a similar way as the other mutant hinge-RRM3 constructs were made, but with three separate primary PCRs; from the 5' end of the hinge to the first two mutations (ETEE and W), from the first two mutations to the third mutation site (MTNY), and from the third mutation to the 3' end of RRM2. The three primary PCR fragments were joined together by the secondary PCR with the primers at both ends of the eH3mu8. The secondary PCR fragment was digested with EcoRI and BglII, and subcloned into EcoRI/BamHI sites of pGADT7.

For Rbp9 and fne constructs, the coding sequences of Rbp9 and fne were amplified by PCR from cDNA prepared from adult Drosophila RNA. The amplified Rbp9 coding sequence was digested with BglII and XhoI, and subcloned into BamHI/SalI and BamHI/XhoI sites of pGBK7T and pGADT7, respectively. The fne coding sequence was digested with EcoRI and BglII, and subcloned into EcoRI/BamHI sites of pGBK7T and pGADT7.

For the upstream activating sequence (UAS) constructs, PCRs were performed to put a mutant RRM in place of RRM3 in the full length elav cDNA. The pGADT7 with a mutant hinge-RRM insert (eH3mu4, eH3mu8, or eH2) was digested with HpaI, and used as a template together with pGBK7T-eQ12H. With the primers at the 5' end of elav coding sequence and downstream to the cloning site of pGADT7, the full length elav coding sequences with a mutant RRM3 were amplified. The PCR fragments were digested with EcoRI and XhoI, and subcloned first into EcoRI/XhoI sites of pGADT7, then cut out by BglII and XhoI digestion, and subcloned into BglII/XhoI sites of the pUAST vector (38). Likewise, the control elav coding sequence was excised from the pGADT7-eQ12H construct with BglII and XhoI digestion, and subcloned into BglII/XhoI sites of pUAST. All the UAS-elav constructs have hemagglutinin (HA) epitope tag sequence at the N-terminus that is derived from pGADT7.

All the subcloned PCR fragments were fully sequenced to confirm the absence of inadvertent mutations introduced during PCR. See Supplementary Data for all the primer sets used for the PCRs (Table S1) and their sequences (Table S2).

Yeast two-hybrid assay

The Matchmaker Two-Hybrid System 3 (Clontech) was used for the two-hybrid interaction assays. The Saccharomyces cerevisiae Y187 and AH109 strains (Clontech) were transformed with the pGBK7T and pGADT7 constructs, respectively. To test two-hybrid interactions, Y187 with a pGBK7T construct and AH109 with a pGADT7 construct were mated, and the resulting diploids were tested for two-hybrid reporter gene activation. Each mating culture was plated on a SD/-Ade/-His/-Leu/-Trp testing plate to examine the activation of both ADE2 and HIS3 reporter genes, as well as on a SD/-Leu/-Trp plate to confirm mating success. Colony formation on the SD/-Ade/-His/-Leu/-Trp plate was scored after 6 days of incubation at 30°C. In all figures, ‘+’ means yeast colony formation indicating interaction between the two proteins, and ‘−’ means no or very few colony formations indicating no or very weak interaction (see Figure 3B for examples of ‘+’ and ‘−’ results). All the constructs were tested to confirm that they did not activate the reporter gene expressions by themselves.

Immunoprecipitation

For co-immunoprecipitation (co-IP), adult flies of the following genotypes were prepared: elav52, elavDvORF, CyO, elav52, elavDvORF/+; elav−13/+; elav52, elav−13, TM3 Ser, UAS-eQ12H3/+; Hsp70-GAL4/+ and
Hsp70-GAL4/+ . The elav\textsuperscript{5} (26) is a null allele of elav, elav\textsuperscript{DvORF} (39) and elav\textsuperscript{--} (40) are elav transgenes, UAS-eQ12H3 is HA-tagged elav under the control of UAS, and Hsp70-GAL4 (P[GAL4-Hsp70.PB]; Bloomington Drosophila Stock Center) is GAL4 driven by the Heat shock protein 70 (Hsp70) promoter. A 37°C heat-shock was applied for 30 min to UAS-eQ12H3/+; Hsp70-GAL4/+ and Hsp70-GAL4/+ flies 3 h and 30 min before head-collection. Eighty heads were collected for each genotype and homogenized in 250 μl of immunoprecipitation buffer [IPB; 50 mM Tris–HCl pH 7.5, 50 mM NaCl, 0.1% Triton X-100, Complete Mini Protease Inhibitor Cocktail Tablet (Roche)]. The homogenate was centrifuged, the supernatant was divided into two equal volumes of 110 μl, and the RNase T1 was added to one (RNase +). Both RNase + and − homogenates were incubated at room temperature (RT) for 30 min. After the incubation, the homogenate was centrifuged again and 100 μl of the supernatant was recovered. The immobilized Protein A (Pierce Biotechnology) was pre-washed with IPB, then incubated with either the mouse anti-ELAV monoclonal antibody (mAb) 7D (for ELAV-DvORF) or the mouse anti-HA mAb with either the mouse anti-ELAV monoclonal antibody (Jackson ImmunoResearch Laboratories) at the dilution of 1:200 each in PBT for 2 h and 30 min at RT. The wing discs were mounted with 70% glycerol in PBS after washing with PBT.

**Confocal microscopy and image analysis**

Immunofluorescent images were acquired by Leica TCS SP2 mounted on Leica DM IRE2 inverted microscope. Intensities of fluorescence were measured by ImageJ software (National Institutes of Health) from average Z-series projection images of the wing discs. The ELAV (Cy5) and GFP (fluorescein) signals in the dpp expression pattern as well as in the background were measured for eight wing discs for each genotype. The background signal was subtracted from the signal in the dpp-pattern to obtain the compensated signal intensity for each disc. The compensated GFP signal was normalized by the compensated ELAV signal for each disc, and the average and confidence interval of the normalized GFP signal was calculated for eight wing discs of each genotype.

**RESULTS**

All Drosophila ELAV-family proteins interact with each other

It has been shown that ELAV forms multimeric complexes in the presence or absence of substrate RNA (34), and mammalian Hu proteins form homo- and hetero-multimers (35-37). In our yeast two-hybrid screening of a Drosophila embryonic cDNA library for ELAV-interacting proteins, we identified ELAV itself and Fne, another ELAV-family protein in Drosophila, as the strongest ELAV-interacting proteins (Toba and White, our unpublished data). This suggests that Drosophila ELAV-family proteins also interact with each other as Hu proteins do. To examine interactions among all three Drosophila ELAV-family proteins, we cloned the cDNA of elav, Rbp9 and fne into yeast two-hybrid vectors. The interactions between the proteins in yeast were assessed by looking at colony formation on the testing plate that requires activation of both of the two reporter genes (see Materials and Methods section for details). Interactions were positive in all the combinations (Figure 1A), which suggests that Drosophila ELAV-family proteins may form homo- and hetero-multimers under these conditions.

**ELAV–ELAV interaction requires RNA**

To examine ELAV multimerization in the in vivo environment, we conducted co-IP from Drosophila extracts in two different ways: (i) A protein extract was prepared from flies expressing both the elav\textsuperscript{DvORF} (39) and elav\textsuperscript{--} (40) transgenes over the elav\textsuperscript{5} (null) background. The elav\textsuperscript{DvORF} expresses ELAV of *D. viliris* that has a larger molecular weight than *D. melanogaster* ELAV. The ELAV\textsuperscript{--} protein has a 13 amino acid deletion in the first RRM that includes the epitope recognized by the monoclonal anti-ELAV antibody. The ELAV\textsuperscript{DvORF} was IP-ed with the monoclonal antibody to see if the
Figure 1. (A) Yeast two-hybrid interactions between Drosophila ELAV-family proteins. In the matrix, ‘+’ and ‘−’ indicate positive and negative interactions in the yeast two-hybrid assay, respectively. The results show that all Drosophila ELAV-family proteins, ELAV, Rbp9 and Fne, interact with themselves as well as each other. ‘Vector only’ means empty pGBK7 and pGADT7 vectors for the DNA binding and activation domain fusions, respectively. ND, not determined. (B and C) ELAV–ELAV interaction is sensitive to RNase treatment. (B) Both ELAVDvORF and ELAV−13 were expressed in the elav null mutant flies. Immunoprecipitation (IP) was performed with the anti-ELAV mAb 7D which does not recognize ELAV−13, and western blot was probed with the anti-ELAV polyclonal antiserum. ELAV−13 was co-IPed with ELAYDvORF (lane 6), and the co-IPed ELAV−13 became undetectable with RNase treatment (lane 7). The elav5; elavDvORF/CyO (lanes 1, 4 and 5) and elav5; elav−13/TM3 Ser (lanes 3, 8 and 9) are negative controls in which ELAVDvORF and ELAV−13 were solely expressed, respectively. (C) HA-tagged ELAV was expressed in the flies with Hsp70-GAL4 driver, and an anti-HA mAb was used for IP. Western blot was probed with the anti-ELAV polyclonal antiserum. The endogenous ELAV was co-IPed with HA-ELAV (lane 3), and the interaction was disrupted by RNase treatment (lane 4). The Hsp70-GAL4 (lanes 2, 5 and 6) is a negative control that does not have the HA-tagged ELAV (UAS-eQ12H3) transgene.

ELAV−13 would be co-IP-ed; (ii) The HA-tagged ELAV was expressed in adult Drosophila using the GAL4-UAS system (38), and the HA-ELAV was IP-ed with an anti-HA antibody to examine if the endogenous ELAV would be co-IP-ed with it. In both cases, ELAV protein that was not recognized by the antibody was co-IP-ed with the other form of the protein (Figure 1B lane 6 and Figure 1C lane 3). However, the interactions seemed RNase sensitive since RNase treatment of the protein extract made co-IP-ed proteins undetectable in both cases (Figure 1B lane 7 and Figure 1C lane 4). This result suggests that either the interaction is mediated by RNA, or binding of the RNA allows a conformational change in ELAV that facilitates the protein–protein interaction.

Mapping of ELAV multimerization domains

ELAV consists of three RRMs, an N-terminal alanine/glutamine (AQ)-rich region, and a hinge region between the RRM2 and 3 (Figure 2A). To map the region that is responsible for the ELAV–ELAV interaction, we made a series of deletion constructs and tested interactions using the yeast two-hybrid system. Interaction activity required the presence of both the hinge region and RRM3 in all cases but of eQ12 and eQ12H constructs in Figure 2B. The eQ3 fragment interacted with itself (Figure 2B). In addition to the hinge-RRM3 region, the N-terminal AQ-rich region seemed to contribute to the interaction since the constructs that lack RRM3 or both hinge and RRM3 (eQ12 and eQ12H in Figure 2B) still retained some interaction activity as long as they had the AQ-rich region (compare eQ12 with e12, eQ12H with e12H in Figure 2B). Our results suggest the involvement of both the AQ-rich and hinge-RRM3 regions in the interaction. We resolved to pursue the activity of the hinge-RRM3 region since it is likely to be a general property of ELAV-family proteins, as the AQ-rich region is not present in any other ELAV-family protein. Additionally, the AQ-rich region is not essential to the vital function of ELAV (44).

We narrowed down the activity within the hinge-RRM3 region with deletion constructs of the hinge-RRM3 fragment. The result suggests that N-terminal two thirds of the hinge region and the C-terminal 18 amino acids of RRM3 are dispensable for the interaction (Figure 2C). The break point of the eSH3 in Figure 2C is the C-terminal end of the sequence required for nuclear localization of the protein (45). The construct eSHabba in Figure 2C that consists of most of the RRM3 and a C-terminal part of the hinge was the smallest fragment that showed the same interaction activity as the original eH3 fragment.

Three conserved short sequences in RRM3 play a critical role in ELAV–ELAV interaction

To identify amino acids responsible for the interaction in the hinge-RRM3 region, we looked for evolutionary conservation within the region where we mapped the interaction activity. While we found conserved sequences in the RRM3, the sequence in the hinge region was very divergent and we hardly found any conservation within the region where interaction activity was mapped (data not shown). Although it is still possible that the region has important structures for the interaction without apparent sequence conservation, we focused our investigation on the RRM3.
Before proceeding to the analysis of the sequence, we wanted to make sure that it is the amino acid sequence of RRM3 that determines the interaction activity, not just the presence of any RRM after the hinge. The latter could be the case since the general structure of the RRM is quite well conserved. To test the possibility, ELAV RRM1 or 2 were fused to the C-terminus of the hinge, and checked for the interaction by the two-hybrid assay. Our results showed that RRM1 or 2 do not substitute for RRM3, indicating that amino acids unique to the RRM3 are responsible for the interaction (Figure 3B column 10, data not shown).

To find the amino acids responsible for the interaction, we looked for RRM3-specifically conserved residues. Figure 3A shows an amino acid sequence alignment of RRRMs from three Drosophila, and four human ELAV-family proteins. RRM3-specific conservations are indicated by magenta, which are defined as the amino acids conserved among RRM3s, but not in RRM1 or 2. Conserved amino acids among the RRM3s that are not specific to RRM3 are indicated by blue. We chose the underlined sequences (numbers 1–7) as the targets for the mutational analysis. The sequence number 5 is not an RRM3-specific conservation, and was chosen as a negative control.

The eH3 construct was mutated by replacing one of the underlined sequences in Figure 3A with the corresponding sequence of ELAV RRM2. The interaction with eH3 was tested by the two-hybrid assay. While mutation in sequences 1, 2, 5 and 7 did not affect the interaction (Figure 3B columns 2, 3, 6 and 8), mutations in sequences 3, 4 and 6 caused loss of interaction (Figure 3B columns 4, 5 and 7), which indicated that the mutated amino acids were essential for the interaction. The three essential sequences 3, 4 and 6 are the ‘ETEE’ sequence adjacent to the putative α-helix 1, the single tryptophan on the α-helix 1, and ‘MTNY’ close to the putative α-helix 2, respectively. Next, we asked if the three sequences are sufficient to provide interaction activity to RRM2 when introduced into the eH2 construct at the corresponding sites. The hinge-RRM2 construct with the RRM3 sequences 3, 4 and 6 (eH3mu8) interacted with eH3...
Given that our results indicate RRM3 has a critical role in ELAV–ELAV interaction, it is possible that RRM3 serves mainly as a protein–protein interaction domain, not as an RNA-binding domain. To test the possibility, we examined the in vivo function of ELAV with mutant RRM3 by expressing them using the GAL4-UAS system. We made four UAS-elav constructs: (i) the wild-type control (eQ12H3); (ii) the W419E mutant (eQ12H3mu4) that is presumably interaction defective; (iii) a mutant in which RRM2 replaces RRM3 except for the sequences 3, 4 and 6 (eQ12H3mu8) and (iv) a mutant in which RRM2 replaces RRM3 completely (eQ12H2) (see Figure 5C for schematics). To evaluate the in vivo function of the mutant proteins, we employed a GFP reporter gene (UnGA) whose expression is dependent upon ELAV-regulated whose expression is dependent upon ELAV-regulated
have shown that mutant proteins lacking RRM3 still bind ARE efficiently (37,47,48). Yet, evolutionary conservation of the RRM3 sequence is the strongest among the three RRMs, which indicates an important function for RRM3. In this study, we showed that ELAV RRM3 has a central role in multimerization of the protein, and we believe that the multifunctional aspect of RRM3 makes it the most conserved RRM of the protein. It is noteworthy that RRM3 of HuC and HuB has been shown to have a dominant negative effect on neural-phenotype-inducing activity of the proteins (49). An explanation for this effect is competition between the full-length protein and the RRM3 fragment for binding to the RNA substrates, as shown by Gao and Keene (36). Considering our results, however, one can conceive another possible mechanism that the over-expression of the RRM3 fragment interferes with multimerization of the full-length proteins.

Kasashima et al. (35) studied multimerization of Hu proteins. They found that RRM3 of HuC showed the strongest interaction activity in the yeast two-hybrid system when full length HuB was used as bait, basically agreeing with our result. However, when HuC RRM3 was used as bait, a fragment containing both RRM1 and 2 also showed strong interaction activity (35). In our yeast two-hybrid experiment, similar constructs (e12 construct in Figure 2B) did not show interaction with either the full length ELAV or hinge-RRM3 fragment. This discrepancy may come from the presence of an arbitrary threshold in our system; i.e. we evaluate interaction by scoring colony formation of the yeast, and weak interactions may fail to be scored in principle. Therefore, it is possible that RRM1 and 2 also contribute to multimerization. In fact, we found that the RRM1 + 2 fragment showed limited interaction activity when the N-terminal AQ-rich region was added to it (eQ12 construct in Figure 2B). Nonetheless, our results suggest that the interaction activity of RRM3 is at least stronger than that of RRM1 or 2. The three sequence elements in the RRM3 we identified in this study are likely to be important for multimerization of all ELAV-family proteins, since they were identified as conserved sequences among the family.

Structural studies have shown that the RRMs have a well conserved general structure; i.e. four anti-parallel β-strands form a β-sheet, and two α-helices are packed against the β-sheet (6; Figure 4). Studies on RNA-bound forms of RRMs have demonstrated that RRMs interacts with RNA through the amino acids located on the β-sheet, including those in the conserved short consensus sequences ribonucleoprotein (RNP) 1 and 2 by which RRM was initially defined (4,5,7–9). It has been shown that RRMs are also involved in protein–protein interactions. In some cases, the RRM interacts with another RRM (17,19,21), while in other cases, the RRM interacts with non-RRM proteins (8,10–16,18,20). Amino acids on the α-helices are involved in most interactions although detailed mode of interaction varies from one case to another. We identified three sequence elements that are essential for ELAV–ELAV interaction, and also sufficient to give the interaction activity to RRM2 when all of them are introduced into it. All the three elements are estimated to be located on or adjacent to the α-helices when we look
at the positions of the corresponding amino acids in other RRMs for which the tertiary structures have been determined (Figure 4). This suggests a model in which the ELAV RRM3 interacts with another RRM3 through the surface opposite to the RNA-binding surface of the domain.

In our co-IP experiment, ELAV–ELAV interaction required the presence of RNA. This observation raises the question as to whether the interaction is based on protein–protein interaction or is mediated by RNA. Although the RNA-mediated interaction model could not be excluded completely, the protein–protein interaction model is preferred for the following reasons: (i) the three identified important sequence elements for the interaction are all located away from the putative RNA binding surface; (ii) the mutations on the putative RNA binding surface (mutants eH3mu2 and 5 in Figure 3B) that potentially alter the binding specificity did not interfere with the interaction. It has also been shown that HuD–HuD interaction is greatly reduced with RNase treatment although not completely abolished (35). We hypothesize that a conformational change of the RRM upon RNA binding ensures efficient protein-protein interaction in both ELAV and HuD cases. Interestingly, a recent study on ARE-binding of HuR has revealed that RRM3 is required for cooperative assembly of HuR oligomers on RNA (37). The observation fits the idea that the initial binding of an ELAV-family protein on a target RNA facilitates the protein-protein interaction between RRM3s.

We find that RRM2 with the three critical sequences allows ELAV interaction in the yeast two-hybrid system, but still fails to support the in vivo function. Thus, additional sequences in RRM3 are also necessary for ELAV function. This is consistent with the previous study that suggests that the RNA binding ability of each RRM,
including RRM3, is essential, as transgenes that carry mutations that individually disrupt the RNA binding ability of individual RRMs were unable to provide ELAV function (40). Therefore, it is suggested that ELAV RRM3 is a bi-functional domain that interacts with both RNA and protein.

One of the three critical sequence elements for ELAV–ELAV interaction was the single tryptophan at the position 419 (W419). Coincidentally, two independent temperature-sensitive alleles of elav, elav<sup>elavts1</sup> and elav<sup>elavFliJ2</sup>, have TAG and TGA stop codons at the site of W419, respectively (50). In these two mutants, ELAV protein with an apparent normal size is produced, along with a truncated form. It is hypothesized that some form of nonsense suppression occurs in these mutants, and probably a different amino acid substitutes the original W419 (50). Further, it is suggested that the temperature sensitivity of the mutants is due to impaired function of the protein, rather than the thermolability of the protein, because the amounts of either forms of ELAV are unaffected by temperature (50). Since W419E substitution strongly reduced multimerization activity in our yeast two-hybrid experiment, we hypothesize that the amino acid substitution resulted from the nonsense suppression compromise multimerization of the protein, which leads to the temperature sensitive phenotype of the mutants.

ELAV-family proteins have been shown to form multimeric complexes on their target RNA (34–37). It is likely that multimerization is an integral part of the mechanism by which ELAV-family proteins carry out their functions. However, the structural basis of the multimer formation remains unknown. In this study, we showed the importance of RRM3 for ELAV multimerization, and identified three sequence elements that are essential for the interaction. Our results suggest a rough model for an ELAV multimer in which ELAV molecules interact with each other by their RRM3s. Further structural and biochemical studies are required to draw the complete picture of the multimeric complex, and understand the physical mechanisms of RNA processing/stabilization by ELAV-family proteins.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

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