Transient $\alpha$-helices in the disordered RPEL motifs of the serum response factor coactivator MKL1

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The megakaryoblastic leukemia 1 (MKL1) protein was originally identified in a study of chromosomal translocation in infants and children with acute megakaryocytic leukemia1,2. It has been implicated in cancer cell migration and invasion3-6 as well as in the regulation of neurite outgrowth7-10 and dendritic complexity11-13. MKL1 is a member of the myocardin-related transcription factor family, which regulates essential biological processes, including the development and differentiation of cells. MKL1 functions as a transcriptional coactivator of the serum response factor (SRF) in the cell nucleus. MKL1 also functions as a G-actin-binding protein. The localization of MKL1 in cells is regulated by the monomeric G-actin level in the cytoplasm—namely, actin polymerization. Actin polymerization is induced by the activation of RhoA signaling, which decreases the monomeric G-actin level in the cytoplasm. The depletion of G-actin in the cytoplasm results in the dissociation of MKL1 from G-actin. As a result, MKL1 translocates from the cytoplasm to the nucleus through the importin $\alpha$/$\beta$1 heterodimer11. In the nucleus, MKL1 collaborates with SRF to induce the transcription of a number of genes, including actin, c-fos, and SRF itself12-17.

Rat MKL1 (GenBank accession number: BAN82605.1) is a 1038 amino acid protein that has an N-terminal actin-binding RPEL domain, basic boxes, a glutamine-rich domain, an SAP domain, a leucine zipper-like domain, and a transactivation domain17,18. The RPEL domain consists of three RPEL motifs, each of which functions as an actin-binding element19. RPEL2 and RPEL3 each have a core sequence of RExxxEL, while RPEL1 has a non-canonical RExxxxEL core sequence (Figure 1a). The crystal structures of RPEL1 and RPEL2 in complex with G-actin have been reported20. In these complexes, the RPEL motif adopts two $\alpha$-helices (helices $\alpha$1 and $\alpha$2) and binds to the hydrophobic cleft and hydrophobic ledge of G-actin (Figure 1b). The complex structure also suggests that side chains of L136, K139, I140, R143, L149, I154, and L155 of RPEL2 are essential for the interaction with G-actin (Figure 1c)20.

Although the structure of the RPEL motif in complex with G-actin has been reported20, little is known about the conformational properties of RPEL motifs in the free state. This information is necessary to fully understand the translocation of MKL1 from the cytoplasm to the nucleus, since the translocation requires the dissociation of MKL1 from G-actin. In this study, the structure of the free RPEL motif is investigated by nuclear magnetic resonance (NMR) and circular dichroism (CD) spectroscopy.
Results

CD analysis of RPELs. We analyzed the secondary structure of RPELs using CD spectroscopy. The CD spectra of RPELs exhibited a strong negative band near 200 nm and a weak negative shoulder at 220 nm, which is characteristic of unfolded polypeptides. This is supported by the prediction of structural disorder by IUPred, which indicates that RPEL1, RPEL2, and RPEL3 are disordered. The negative band near 200 nm is stronger in the order of RPEL3, RPEL2, and RPEL1. Although free RPEL motifs are largely unstructured, analysis of the helix content shows that RPEL adopts a small but significant amount of α-helix conformation. The helix content is higher in the order of RPEL1, RPEL2, and RPEL3.

Chemical shift analysis of RPEL1. The conformational properties of RPEL1 in the free state were investigated by NMR spectroscopy. Backbone resonance assignments were obtained with standard triple-resonance NMR experiments. The 1H-15N HSQC spectrum of RPEL1 shows the backbone amide resonances within 8.0–8.6 ppm in the 1H dimension. The narrow chemical shift dispersion in the 1H dimension is characteristic of intrinsically disordered proteins (IDPs). NMR chemical shifts can be used to probe the propensity of proteins to adopt an α-helical conformation and extended (β-strand) conformation at the residue level. In order to investigate the conformational properties of RPEL, we utilized the secondary structure propensity (SSP) program. The SSP program combines chemical shifts from different nuclei into a single score indicating the secondary structure propensity. A positive SSP score indicates a propensity for α-helical conformation, and a negative score indicates disorder.

Figure 1 | (a) Amino acid sequences of RPEL motifs from rat MKL1. The positions of helices α1 and α2 of RPEL1 in complex with G-actin are indicated. The sequences of RPxxxEL and RRxxxEL are underlined. Numbering is based on GenBank accession number BAN82605.1. Asterisks indicate the essential residues for actin-binding. (b) Three-dimensional structure of RPEL2 bound to G-actin (PDB entry 2V52). The positions of helices α1 and α2 are indicated. (c) The interaction region between RPEL2 and G-actin (PDB entry 2V52). G-actin is shown in surface and cartoon representations. RPEL2 is shown in stick representations and essential residues for actin-binding are shown in red. The side chain of K139 in RPEL2 adopts two conformations, both interacting with G-actin. (d) Prediction of disorder tendency of full-length MKL1 by the IUPred predictor. Scores above a threshold value of 0.5 are considered to be disordered. The positions of RPEL1, RPEL2, and RPEL3 are indicated.

Figure 2 | CD spectra of RPEL1 (solid line), RPEL2 (dotted line), and RPEL3 (dashed line) at pH 7.0 and 25°C in the far-UV region.
a propensity for extended (β-strand) conformation (Figure 4a). The SSP scores represent the expected fraction of α-helical or extended (β-strand) conformation at a given residue. For instance, an SSP score of 0.5 indicates that 50% of the conformers in the disordered state ensemble adopt an α-helical conformation at that particular position. In order to examine the positions of helices, we utilized the difference between 13Cα secondary shifts and 13Cβ secondary shifts (ΔδCα - ΔδCβ) (Figure 4b). Consecutive positive values of ΔδCα - ΔδCβ indicate a propensity to adopt an α-helical structure, and consecutive negative values indicate an extended (β-strand) structure. Since appropriate random coil chemical shifts are important to obtain reliable secondary structure propensities, we used the random coil chemical shifts suitable for IDPs.

The residues from Asn90 to Gln97 and from Arg102 to Ser107 have a propensity to adopt an α-helical conformation, indicating that helices α1 and α2 are formed in free RPEL1 (Figure 4b). This does not mean, however, that the helix α1 and α2 are stably formed in free RPEL1, since the averaged SSP score is ~19% for the helix α1 region (residues 90–97) and ~26% for the helix α2 region (residues 102–108). Therefore, helices α1 and α2 are transiently formed in free RPEL1.

Proline mutagenesis of RPEL1. We confirmed the transient α-helix formation of RPEL1 using proline mutagenesis. Proline mutation unfolds or greatly destabilizes the protein structure when inserted in the middle of secondary structures. Mutation of Leu94 to Pro reduced the α-helical propensity in the helix α1 region, indicating the helix formation. However, the mutation of Leu94 to Pro had little effect on the helical propensity in the α2 region (Figure 5a). In addition, the mutation of Leu105 to Pro reduced the α-helical propensity in the α2 region of RPEL1 but had little effect on the helical propensity in the α1 region (Figure 5b). These results indicate that the transient α-helices in free RPEL1 are independently formed without helix-helix interactions.

Chemical shift analysis of RPEL2 and RPEL3. We also investigated the conformational propensity of RPEL2 and RPEL3 using NMR spectroscopy (Figure 6). The residues from Thr132 to Lys139 and from Leu149 to Met152 of RPEL2 have a propensity to adopt an α-helical conformation (Figure 6a). On the other hand, RPEL3 exhibits no significant helical propensity (Figure 6b). Together, our results suggest that the helices α1 and α2 are transiently formed in RPEL1 and RPEL2, while the helix is not formed in RPEL3. The helix content is higher in the order of RPEL1 > RPEL2 > RPEL3 (Figures 4 and 6).

Discussion
Many IDPs undergo a disorder-to-order transition upon binding to their target molecule, a process that has been called “coupled folding and binding.” There are two models describing the mechanism of the disorder-to-order transition—namely, the conformational

| Helix content of the RPEL motif calculated from far-UV CD spectra |
|-----------------|-----------------|-----------------|
|                | the self-consistent method | the CONTIN method | the variable-selection method |
| RPEL1           | 16              | 13              | 7.1             |
| RPEL2           | 14              | 11              | 5.1             |
| RPEL3           | 6.7             | 5.0             | 0.7             |
selection model and the induced fit model. In both models, IDPs are largely unstructured before binding to their target molecules. On the other hand, a number of studies have shown that the disordered state ensemble contains a significant amount of native-like secondary structure. However, the significance of the native-like secondary structure in the disorder-to-order transition is not fully understood.

In the present study, we investigated the conformational properties of RPEL1-3 in the free state using NMR and CD spectroscopy. The isolated RPEL motifs are largely disordered in solution, while RPELs acquire the tertiary structure upon binding to the monomeric G-actin. According to the crystal structures of RPEL1 and RPEL2 in complex with G-actin, the RPEL motif adopts two α-helices and binds to the hydrophobic cleft and the hydrophobic ledge of G-actin. The essential residues for actin binding are completely conserved between RPEL2 and RPEL3, while the essential residues are partially different between RPEL1 and RPEL2: Ile140 and Leu155 of RPEL2 are substituted with Leu and Met in RPEL1, although Leu136, Lys139, Arg143, Leu149, and Ile154 of RPEL2 are conserved in RPEL1.

Our experimental data indicate that free RPELs contain α-helical structures. The α-helical structure is partially and transiently formed in the regions where the helices α1 and α2 are induced upon binding to G-actin. Generally, the structural disorder of IDP in solution raises the entropic penalty when IDP undergoes the disorder-to-order transition upon binding to its target molecule. The preformed α-helices of IDP may decrease the entropic penalty of the disorder-to-order transition.

Mouilleron et al. have shown that RPEL1 and RPEL2 bind to G-actin tightly, with dissociation constants (Kd) of 1.0 and 1.9 μM, respectively. They also showed that RPEL3 binds weakly to G-actin with a Kd value of 28.9 μM, although the essential residues for the interaction with G-actin are conserved in RPEL3. From these Kd values, the Gibbs free energy changes (ΔG°) of binding are 8.2 kcal/mol, 7.8 kcal/mol, and 6.6 kcal/mol for RPEL1, RPEL2, and RPEL3, respectively. The helical propensity is higher in the order of RPEL1, RPEL2, and RPEL3. These results suggest that the amount of preformed structure may correlate with the ΔG° value of binding. It is conceivable that the binding strength between IDP and the target molecule is modulated by the preformed structural elements in free IDP. A previous study suggests that the preformed secondary structure in IDP is an important determinant for the interaction between IDP and its target molecule.

Proline mutagenesis shows that the transient α-helices in RPEL1 are locally formed without helix-helix interactions. In contrast to our observation, long-range helix-helix interactions are formed in the

Figure 5 | ΔΔCα - ΔΔCβ secondary chemical shifts of the (a) L94P and (b) L105P mutant of RPEL1. The positions of helices α1 and α2 are indicated in each panel. The positions of the mutations are indicated by arrows.

Figure 6 | ΔΔCα - ΔΔCβ secondary chemical shifts of (a) RPEL2 and (b) RPEL3. The positions of helices α1 and α2 are indicated in each panel.
unfolded state of other proteins such as ACTR\(^a\), acyl coenzyme A binding protein\(^b\), and hepatitis C virus protein NSSA\(^c\). A key difference seems to be that these other proteins have amphipathic helices with a clear hydrophobic side that is likely to form helix-helix interactions. The difference may also result from the length of \(\alpha\)-helices. The \(\alpha\)-helices of RPEL are shorter than those of the other unfolded proteins.

The RPEL motif of MKL1 is an ensemble of conformations ranging from random coils to native-like \(\alpha\)-helices. Our results suggest that the amount of preformed structure may correlate with the binding strength between IDP and the target molecule. In order to fully understand the mechanism of the disorder-to-order transition, it will be useful to investigate the binding kinetics of RPEL with varying helical propensity to G-actin\(^d\).

**Methods**

**Protein expression and purification.** MKL1(85–116), MKL1(129–160), and MKL1(173–204) were expressed as GST-fusion proteins\(^e\). The DNA encoding MKL1(85–116), MKL1(129–160), and MKL1(173–204) were inserted into a pOP3C plasmid. Sequencing of the inserted DNA was performed on an ABI PRISM 3130 Genetic Analyzer (Applied Biosystems). The GST-fusion proteins were expressed in C41(DE3)RIPL cells harboring the pOP3C plasmid. The cells were grown in M9 minimal medium supplemented with \(15\) mM NaCl and \(15\) mM-glucose. Protein expression was induced by the addition of isopropyl-\(\beta\)-D-thiogalactopyranoside at a final concentration of \(0.3\) mM. After 5 h cultivation, the cells were collected by centrifugation at \(3765\) \(g\) for 15 min at \(4\) °C. The C41(DE3)RIPL cells were resuspended in buffer A (20 mM Tris-HCl, 100 mM NaCl, and 1 mM dithiothreitol at pH 7.5) and lyzed by sonication on ice. The proteins were detected in a soluble fraction after centrifugation at \(9400\) \(g\) for 60 min at \(4\) °C. The GST-fusion proteins were purified with a glutathione Sepharose 4B column (GE Healthcare Bio-Sciences) equilibrated with buffer A. The purified GST-fusion protein was digested on a column with 3C protease for 15 h at \(4\) °C to remove the GST region. The flow-through containing RPELs was collected by adding buffer A to the column, and RPELs were further purified by high-performance liquid chromatography with a COSMOSIL C-18 AR-II column (Nacalai Tesque). The solutions containing RPELs were collected and dialyzed against buffer B (10 mM sodium phosphate and 50 mM NaCl at pH 7.0). The purified RPELs have an N-terminal extension (GPHM) derived from the pOP3C plasmid. Matrix-assisted laser desorption ionization time-of-flight mass analysis (MALDI-TOF MS) was used to confirm the molecular weights of the RPELs. MALDI-TOF MS was performed on a Bruker Daltonics Autoflex-T1 mass spectrometer.

**NMR spectroscopy.** For NMR experiments, the proteins in buffer B were concentrated to \(0.12–0.50\) mM and D$_2$O was added to a final concentration of \(7\%\). The NMR samples also contained 1 mM NaN$_3$ and 20 \(\mu\)M 2,2-dimethyl-2-silapentane-5-sulfonate sodium salt. All NMR experiments were performed on a Bruker Avance 800 MHz spectrometer equipped with a cryoprobe. \(^1H\), \(^13C\), \(^15N\), and \(^1H\)C\(^15N\) resonances were assigned using standard three-dimensional triple resonance experiments including C(CA)(CN), CBCA(CO)(NH), HN(CA)CO, HN(CO)CA, and HN(CA)CO\(^h\). The assignments have been deposited in the BioMagResBank under accession numbers 11564, 11565, 11566, 11567 and 11568. The \(^1H\) chemical shifts were directly referenced to the resonance of 2,2-dimethyl-2-silapentane-5-sulfonic acid salt, while the \(^15N\) and \(^13C\) chemical shifts were indirectly referenced with the absolute frequency ratios. NMR data were processed with NMRPipe\(^i\) and analyzed with NMRView\(^i\).

The random coil chemical shifts of Kjaergaard et al.\(^j\) were used to calculate the \(^1H\)C secondary shifts and the \(^1H\)C secondary shifts. The SSP scores were calculated with the random coil chemical shifts\(^k\) and the average secondary shifts for the fully formed secondary structure\(^l\) as described previously\(^m\). An averaging window of five residues was applied in the SSP analysis as described previously\(^n\). The SSP scores were calculated using calibrated \(^1H\)C and \(^15N\) chemical shifts as inputs.

**CD spectroscopy.** CD spectra were measured using a J-805 spectropolarimeter (JASCO) at 25 °C. Sample solutions contained 0.25–0.39 mM RPEL, 10 mM sodium phosphate, and 50 mM NaCl at pH 7.0. A quartz cell with a 0.2 mm path length was used for all measurements. The content of the secondary structure was estimated using the CONTIN method, the self-consistent method, and the variable-selection method\(^o\)–\(^s\). The Dichroplot program\(^t\) and the Dichropert server\(^u\) were used for the calculation.
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Acknowledgments
This study was supported by Grants-in-Aid for Scientific Research in Innovative Areas (project number: 21113003) from the Ministry of Education, Culture, Sports, Science, and Technology of Japan.

Author contributions
M.M. wrote the paper. M.M. and T.F. performed the NMR experiments and analysis. T.F., T.O. and M.I. prepared the plasmids. T.F. and T.O. purified the proteins. M.T. and A.T. designed the project and reviewed the manuscript.

Additional information
Funding This study was supported by Grants-in-Aid for Scientific Research in Innovative Areas (project number: 21113003) from the Ministry of Education, Culture, Sports, Science, and Technology of Japan.

Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Mizuguchi, M. et al. Transient α-helices in the disordered RPEL motifs of the serum response factor coactivator MKL1. Sci. Rep. 4, 5224; DOI:10.1038/srep05224 (2014).

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