

Cite this: *RSC Adv.*, 2017, 7, 29713

Conformation dynamics of the intrinsically disordered protein c-Myb with the *ff99IDPs* force field†

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The intrinsically disordered protein c-Myb plays a critical role in cellular proliferation and differentiation. Loss of *c-myb* function results in embryonic lethality due to failure of fetal hepatic hematopoiesis. The conformation dynamics of the intrinsically disordered c-Myb are still unknown. Here, molecular dynamics (MD) simulations with the intrinsically disordered protein force field *ff99IDPs* were used to study the conformation dynamics. In comparison with *ff99SBildn*, *ff99IDPs* can reproduce more diverse disordered conformers of c-Myb. The predicted secondary chemical shift under *ff99IDPs* is more close to that of experiment data than that under *ff99SBildn*. Therefore, *ff99IDPs* can sample native molten globule, native pre-molten globule and native coil conformers for c-Myb. These results are consistent with those of other intrinsically disordered proteins. Kinetic analysis of MD simulations shows that c-Myb folds *via* a two-state process and indicates that c-Myb folds in the order of tertiary folding and helical folding. The folding nucleus of KEL plays an essential role in stabilizing the folding state with dynamic correlation networks. The influences of solvent models for TIP3P, TIP4P-EW and TIP5P were also investigated and it was found that TIP3P and *ff99IDPs* are the best combination to research the conformer sampling of c-Myb. These results reveal the conformation dynamics of c-Myb and confirm that the *ff99IDPs* force field can be used to research the relationship between structure and function of other intrinsically disordered proteins.

Received 12th April 2017
Accepted 30th May 2017

DOI: 10.1039/c7ra04133k

rsc.li/rsc-advances

Introduction

The MYB family of transcriptional regulators, including A, B and C Myb proteins, plays a critical role in cellular proliferation and differentiation. Furthermore, the *c-myb* protein (c-Myb) is significantly expressed in different hemopoietic lineages^{1–4} of both normal and tumoral immature cells.

*c-myb*⁵ was originally identified as an oncogene carried by the chicken leukemia viruses avian myeloblastosis virus and E26,^{6–8} and is a significant conserved transcription factor.⁹ The previous work reports that high levels of *c-myb* are expressed in immature hematopoietic cells,¹⁰ while its expression is down-

regulated during differentiation.¹¹ This indicates that the *c-myb* gene plays a significant role in cellular proliferation of immature hematopoietic cells.

Three functional domains are found for c-Myb: an N-terminal highly conserved DNA binding domain, a central transcriptional activation domain, and a C-terminal negative regulation domain.¹² c-Myb is one of intrinsically disordered proteins (IDPs),¹³ since it does not fold spontaneously into well-organized globular structures in the absence of stabilizing interactions. The conformation dynamics of intrinsically disordered c-Myb is still unknown, therefore we used molecular dynamics (MD) simulations to study the conformation dynamics. Dynamic conformation plays a key role in the function of intrinsically disordered proteins.^{14–18}

However, the accuracy of the force field remains an issue in applications of MD simulation, while most traditional widely used force fields can't well reproduce intrinsically disordered proteins, especially on backbone dihedrals.¹⁹ In order to better reproduce the disordered conformation, a set of AMBER force fields *ff99IDPs*^{20,21} and others²² were used, which were developed specially for IDPs with grid-based energy correction maps (CMAP) term^{23,24} to correct the dihedral of backbone for eight disordered-promotion residues of alanine, arginine, glycine, glutamine, serine, glutamic acid, lysine, and proline. By adding CMAP energy term, the Φ/Ψ distributions of the eight disorder-

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† Electronic supplementary information (ESI) available. See DOI: 10.1039/c7ra04133k



promoting residues have been improved with respect to the benchmark data of IDPs, while the root mean squared percentage deviation is less than 0.15% between the benchmark and the simulation.²⁰

In this study, we compared the performances of *ff99IDPs* and *ff99SBildn* to reproduce intrinsically disordered c-Myb. Based on the results, *ff99IDPs* was used to reveal the conformation dynamics of c-Myb.

Materials and methods

Molecular dynamics simulation

The atomic coordinates of the c-Myb were extracted from the ternary complex of PDB database (PDB code: 2AGH).²⁵ AMBER12 was used to perform efficient simulations with periodic boundary conditions.²⁶ Hydrogen atoms were added using the LEaP module of AMBER12. Counter-ions were used to maintain system neutrality. C-Myb was solvated in a truncated octahedron box of TIP3P, TIP4P-EW, and TIP5P solvent models with a buffer of 10 Å, respectively.^{27–29} The pairwise interactions (van der Waals and direct Coulomb) were computed with a cutoff distance of 8 Å. Particle Mesh Ewald (PME) was employed to treat long-range electrostatic interactions in AMBER12.³⁰ The *ff99SBildn* and *ff99IDPs* force fields were used for the intramolecular interactions, respectively. The Langevin thermostat was used in the preparation runs with a friction constant of 1 ps⁻¹ and the Berendsen thermostat was used in the production runs.³¹ All MD simulations were accelerated with the CUDA version of PMEMD in GPU cores of NVIDIA® Tesla K20.

To relieve any further structural clash in the solvated systems, initial minimization was performed with 2000-step conjugate gradient minimization. After minimization, a 400 ps' heating up in the NVT ensemble and a 200 ps' equilibration in the NPT ensemble at 298 K were performed before MD simulation. To compare the performances between *ff99IDPs* and *ff99SBildn* for the sampling of c-Myb, multiple independent trajectories with the same initial structure but different velocities under each force field were simulated. Total 5.8 μs trajectories were collected at 298 K. The simulation conditions are gathered in Table 1.

Data analysis

The PTRAJ module in AMBER12 and Amber Tools 13 were used to calculate root-mean-square deviations (RMSDs) and root-

mean-square fluctuations (RMSFs) for c-Myb.³² Kclust program in the MMTSB toolset were used to analysis the structural cluster on the basis of phi/psi dihedral.³³ Native contact was assigned using in-house tools.^{34–36} In order to illustrate the convergence for sampling the disordered c-Myb, cluster numbers within 5 ns period along all the trajectories were plotted. Dictionary of Secondary Structure of Proteins (DSSP) algorithm was applied to assign the secondary structures of c-Myb. Experimental Cα chemical shift data were offered by Arai *et al.*³⁷ SPARTA version 1.01 was used to calculate the secondary chemical shift data for the simulated structures.³⁸ Radius of gyration (RG) and RMSD were both separated into eight bins. All of the structural visualizations were generated using PyMOL 1.7.

Dynamics correlation network

Every amino acid was defined as one node for dynamics correlation network. The fluctuation correlation between any pair of nodes *i* and *j* was calculated with eqn (1).

$$C_{ij} = \frac{\langle \Delta \vec{r}_i(t) \cdot \Delta \vec{r}_j(t) \rangle}{\sqrt{\langle (\Delta \vec{r}_i(t))^2 \rangle \langle (\Delta \vec{r}_j(t))^2 \rangle}} \quad (1)$$

where $\Delta \vec{r}_i(t) = \vec{r}_i(t) - \langle \vec{r}_i(t) \rangle$, $\vec{r}_i(t)$ is the position of node *i* at time *t*, and $\langle \cdot \rangle$ represents a time averaging. These elements were conveniently organized as a covariance matrix for simulated system. This general method has been successfully applied in several systems.^{39–42} In the current study, the covariance matrix for each system was constructed using snapshots (every 2 ps) of all simulated trajectories. Besides nodes, “edge” that transfers allosteric information from one node to another is defined between any two nodes without covalent bond and the distance between two heavy atoms from two nodes are closer than 4.5 Å over 75% sampling time. The strength of the edge between nodes *i* and *j* is defined as the absolute value of the inter-node correlation (C_{ij}). The number of connected edges at each node is defined as the degree of the node. Correlation-weighted degree, which is the summation of strengths of all edges connected to a given node, indicates the importance of the node. After the network construction, Cytoscape3.1.1 was used to calculate the network topological parameters.⁴³ The shortest path between any two nodes in the network was identified with the Floyd-Warshall algorithm.⁴⁴

Results and discussion

Comparison of force fields

MD simulations were performed to model the conformation of c-Myb. The RMSD, Cα fluctuation (RMSF), structural clustering, secondary Cα chemical shift, helicity content, and other structural indices was calculated under *ff99IDPs* and *ff99SBildn*, respectively. For each force field, the RMSD and Cα fluctuation (RSMF) were calculated to ensure that both force field were able to sample rational conformations. The RMSDs relative to initial structure are shown in Fig. 1. This figure indicates that that 100

Table 1 Simulation condition for c-Myb

Force field	Solvent	Trajectories	Time (ns)
<i>ff99SBildn</i>	TIP3P	10	100
	TIP4P-EW	5	
	TIP5P	5	
<i>ff99IDPs</i>	TIP3P	10	190
	TIP4P-EW	5	
	TIP5P	5	



ns simulations are sufficient for the sampling conformer for c-Myb at room temperature under *ff99SBildn* and 190 ns under *ff99IDPs*. The $C\alpha$ fluctuation (RMSF)s under *ff99SBildn* and *ff99IDPs* are shown in Fig. 2. The results suggest that the fluctuation of *ff99IDPs* is higher than that of *ff99SBildn*.

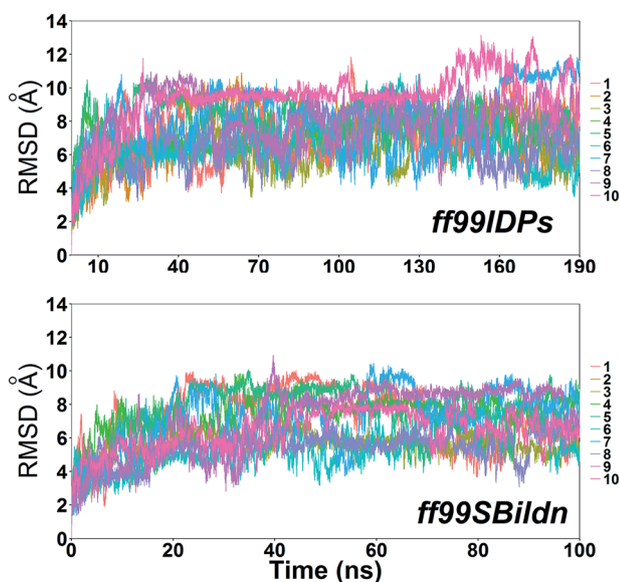


Fig. 1 RMSD under *ff99SBildn* and *ff99IDPs* force fields.

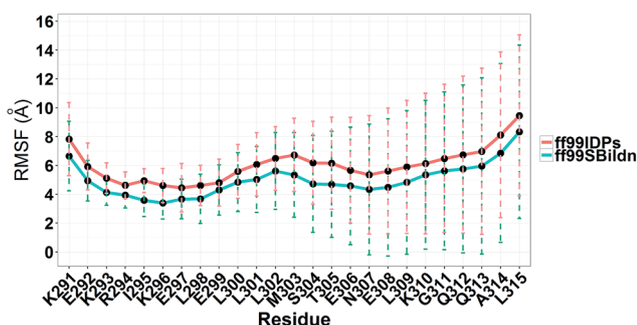


Fig. 2 RMSF under *ff99SBildn* and *ff99IDPs* force fields.

In order to ensure the convergence of the conformation sampling, the numbers of cluster that occupy 70% of the conformations over accumulated simulation time under both force fields are shown in Fig. 3. The conformer cluster suggests that the numbers of clusters do not change significantly at the end of the simulations. This indicates that the chosen simulation lengths are appropriate for sampling c-Myb.

The clustering, secondary $C\alpha$ chemical shift, helicity of c-Myb under *ff99IDPs* and *ff99SBildn* are shown in Fig. 4–6. Apparent differences were found in the structural clustering. The top 10 clusters under *ff99SBildn* occupy 58.75% of the total conformations, of which the top 14 clusters account for 70%. However, under *ff99IDPs*, the top 10 clusters occupy only 15.26% of the total conformations, while the top 195 clusters account for 70%. It was apparent that the clusters under *ff99IDPs* have a higher ratio of population for disordered structures than those under *ff99SBildn*. Most conformers show disordered state and include few secondary structural elements especially in C-terminal domain. The PMF free energy landscapes in respect to RMSD and RG show that the distribution of conformers from *ff99SBildn* is located at RMSDs from 2 to 10 Å and RGs from 8 to ~15 Å, while that from *ff99IDPs* is located at RMSDs from 2 to 12 Å and RGs from 8 to ~20 Å, indicating that *ff99IDPs* samples more flexible conformers than *ff99SBildn*, which is consistent with the clustering result. In order to compare with the experimental data, secondary $C\alpha$ chemical shifts were calculated from the clustered representative structures that occupy no less than 70% of the conformations and their occupancies. The full-length RMSD between the data under *ff99SBildn* and the experimental data was 1.123 ppm, while it is 0.573 ppm between the data under *ff99IDPs* and the experimental data (shown in Fig. 6A). This shows that the performance of *ff99IDPs* is much better than that of *ff99SBildn* to simulate c-Myb. The main difference between simulation under *ff99IDPs* and experiment is focused on the N-terminal domain by the motif of structure promoting residues of KRI. Force field parameters based on the CMAP correction of these residues indeed improve the sampling of c-Myb. *ff99IDPs* force field can sample more flexible conformers than *ff99SBildn*.

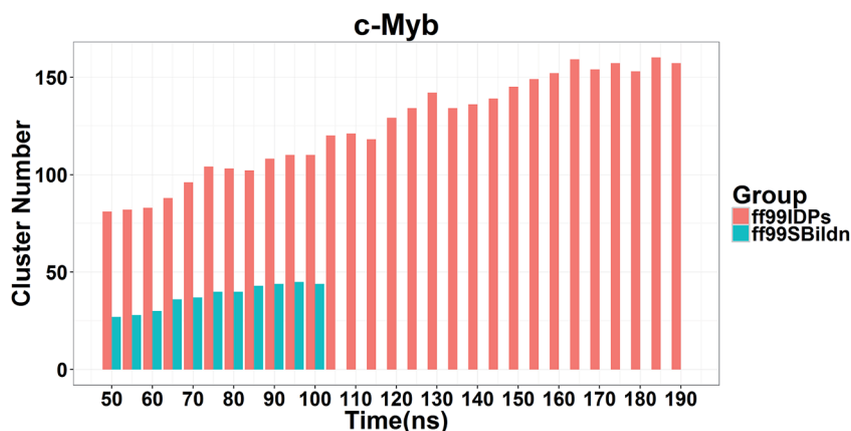


Fig. 3 Numbers of clusters that can occupy 70% of the conformations within accumulated 5 ns period under both force fields.



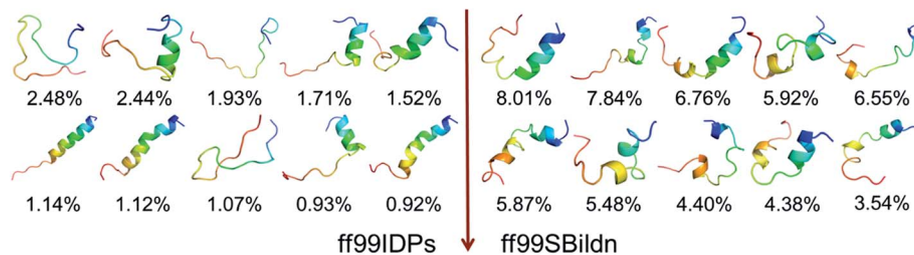


Fig. 4 Representative conformers of the top 10 clusters and their occupations.

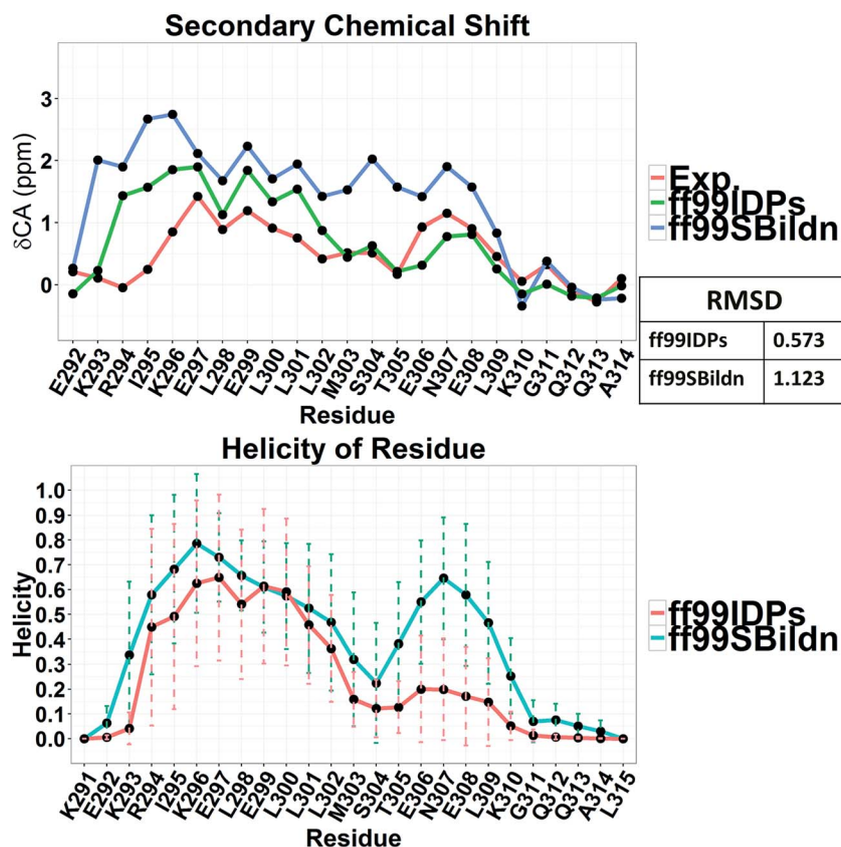


Fig. 5 RMSD between experimental and prediction data for secondary chemical shift and helicity under force fields of *ff99SBildn* and *ff99IDPs*.

Furthermore, the helical content under *ff99IDPs* is also lower than that under *ff99SBildn* and consistent with the results of chemical shift (shown in Fig. 6B). The result is consistent with the property of IDPs, therefore, the trajectories under *ff99IDPs* were chosen to study the conformation dynamics for intrinsically disordered c-Myb.

The previous work reports that IDPs were grouped into three structurally different subclasses, native molten globules, native pre-molten globules and native coils.⁴⁵ According to the content of secondary structure for the conformers of c-Myb, these structure subclasses are shown in Fig. 7. The native molten globule states include high properties of helical structure, pre-molten globule with partly helical conformer and native coil corresponding to extended disordered c-Myb.

Conformation dynamics of c-Myb

Native tertiary contacts (Q_f) and native helical content (Q_h) were used to monitor the tertiary and secondary folding kinetics. Time evolutions of Q_f and Q_h for c-Myb are shown in Fig. 8. Apparently, the tertiary and secondary folding kinetics can be represented well by single exponential function (Table 2). This indicates that the tertiary and secondary folding process obeys first order kinetics at room temperature. Our kinetics analysis shows that the folding half-time of tertiary structure is 8.694 ± 0.783 ns, and the unfolding half-time of helical structure is 21.975 ± 1.546 ns. This indicates that the tertiary unfolding is much faster than the helical structure, that is, the secondary structure unfolding of c-Myb depends on the tertiary unfolding.



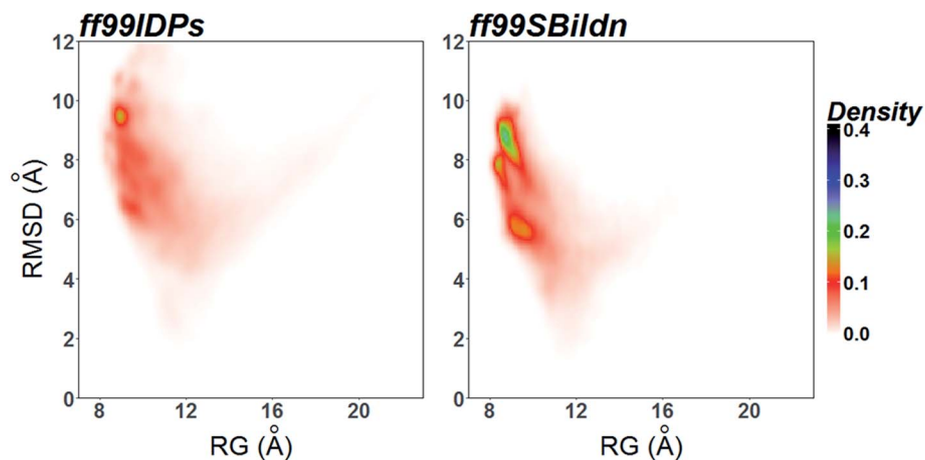


Fig. 6 PMF free energy landscapes on the 2D space of radius of gyration (RG) and root-mean-square deviation (RMSD).

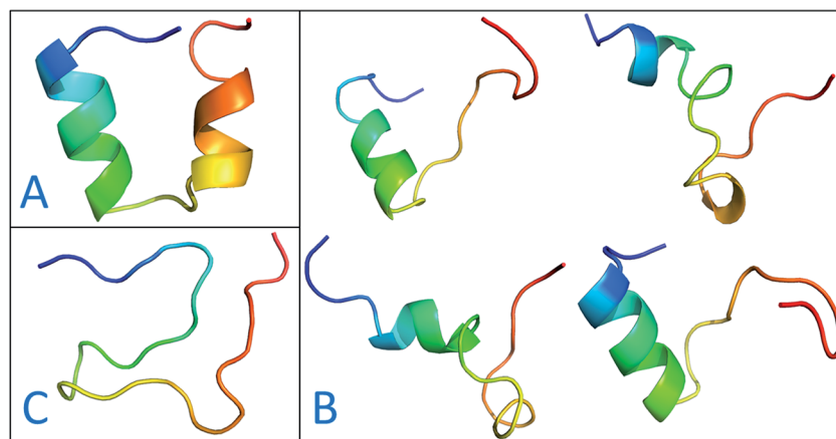


Fig. 7 Representative structures of intrinsically disordered proteins from *ff99IDPs*. (A) Native molten globules. (B) Native pre-molten globules. (C) Native coils.

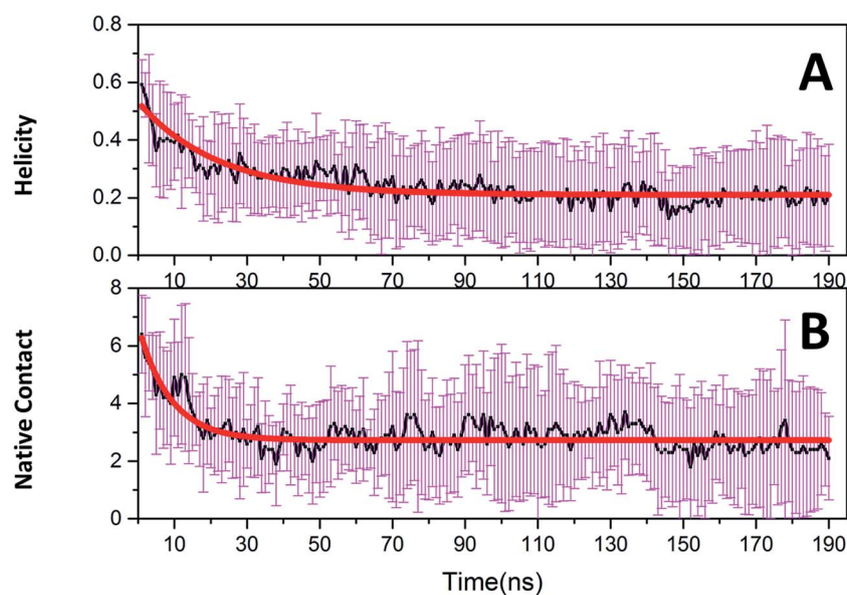


Fig. 8 Folding kinetics of *c-Myb* for native contact and helicity content under *ff99IDPs*. (A) Native contact; (B) helicity content.



Table 2 Kinetics parameters for c-Myb

$y = A \times \exp(-x/\tau) + B$		
	Native contact	Helicity
A	3.994 ± 0.209	0.323 ± 0.012
B	2.725 ± 0.032	0.210 ± 0.004
τ	8.694 ± 0.783	21.975 ± 1.546
R^2	0.760	0.831

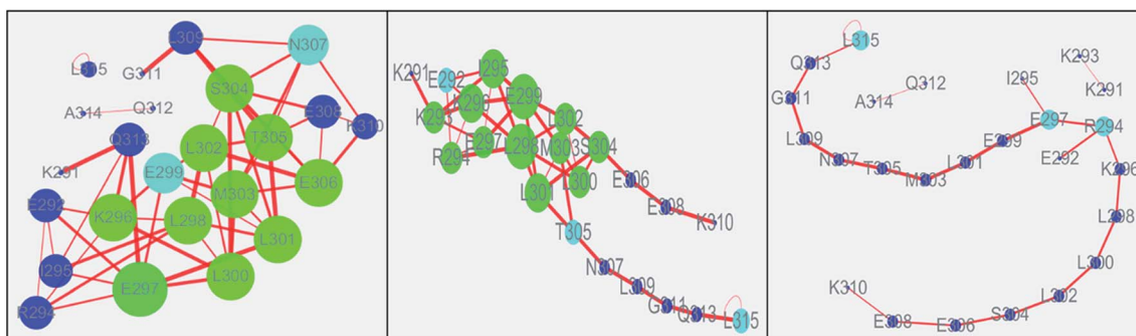
That is, the unfolding of the tertiary structure happened before the issue of the secondary structure.

In order to further reveal the folding kinetics, the dynamic correlation network analysis was used to illustrate the residue fluctuation correlation. To construct the correlation network, the covariance matrices were first calculated. Then fluctuation correlation networks were built. The topology parameters of network for three stages according to the time scale of folding kinetics are listed in Table 3.

The values of topology parameters for network of the first stage are the highest among these four stages. This suggests that the characters of network for the last stages are significant different from those of other stages. The dynamics correlation networks for three stages are shown in Fig. 9. It shows that the number of nodes with weighted degree higher than 10 (more than 10 edges) is 17 in the first stage network and more than that of the other stages. Q313, E292, I295, K294, E297, L300, L298, K296, E299, M303, L302, S304, T305, E306, N307, E308, K310, with higher degree were marked in the first stage and most of them were located on the N-terminal and middle regions. At the second stage, there are just 14 nodes with high

Table 3 Topology parameters for correlation network

	0–8.7 ns	8.7–21.9 ns	21.9–190 ns
Clustering coefficient	0.213	0.157	0
Network centralization	0.132	0.134	0.056
Average number of neighbors	4.348	3.304	1.76
Network density	0.198	0.150	0.073
Network heterogeneity	0.453	0.159	0.332

Fig. 9 Dynamics correlation networks of c-Myb under *ff99IDPs* for three different stages.

weighted degree. The changed nodes are mostly focused on the middle region. For the last stage, the network is separated into three parts. This suggests that the folding of c-Myb almost finished and C-terminal and most middle regions changed into disordered. However, there are two nodes with high weighted degree remaining. These high weighted nodes might be as folding nucleus to stabilize the structure of c-Myb.

According to the results of correlation network analysis, the folding pathway corresponding to the representative structure of c-Myb is shown in Fig. 10. From the folding pathway, we can find a significant folding site, which located between K296 and L298 residues for KEL. These structures are corresponding to the initial state which includes enough native contacts, tertiary folding state which discards part of native contacts, secondary folding state with part helical structure, and folding state with folding nucleus.

Effects of solvent models in simulations of c-Myb

To illustrate the influence of solvent models with generic protein force fields in simulation of c-Myb, we tested TIP3P, TIP4P-EW, and TIP5P models with *ff99SBildn*. The results are shown in Fig. 11. These data show that the differences due to the use of different solvent models do exist, though the effect is small in the final agreement between simulation and experiment. Therefore, force field correction is necessary for improved structural sampling of c-Myb.

Nevertheless, the small effects of solvent models may be found by the biases in *ff99SBildn* force fields. We thus hypothesized that the solvent model might still play a larger role in sampling c-Myb with *ff99IDPs*. To verify this hypothesis, MD simulations with TIP3P, TIP4P-EW, and TIP5P solvent models were conducted in *ff99IDPs*, respectively. The comparison with NMR experiment is shown in Fig. 12. The analysis suggests that the *ff99IDPs* with the modern solvent models of TIP3P, TIP4P-EW and TIP5P do improve the accuracy of c-Myb conformer and better than *ff99SBildn*. In general, the result of predicted secondary chemical shift with *ff99IDPs/TIP3P* was similar to that with *ff99IDPs/TIP4P-EW*. The hydrogen bond between water and c-Myb for different solvent models is shown in ESI Fig. S1.† In general, the number of hydrogen bond for TIP5P was significant different from that for TIP3P or TIP4P-EW. This is consistent with the difference of chemical shift. In summary,



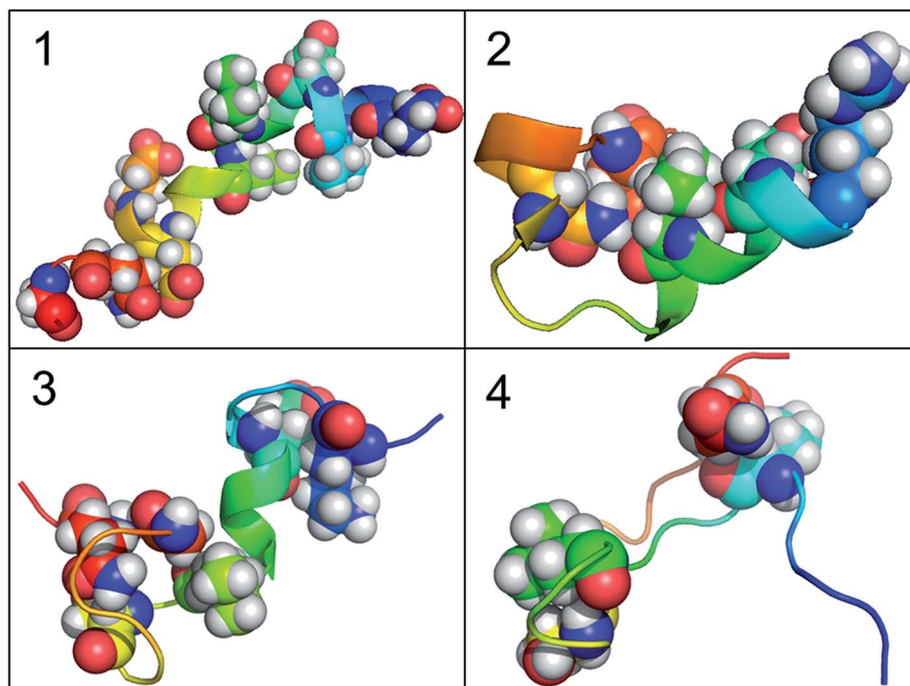


Fig. 10 The folding pathway of c-Myb. (1) Folded state. (2) Tertiary folding. (3) Helical folding (4).

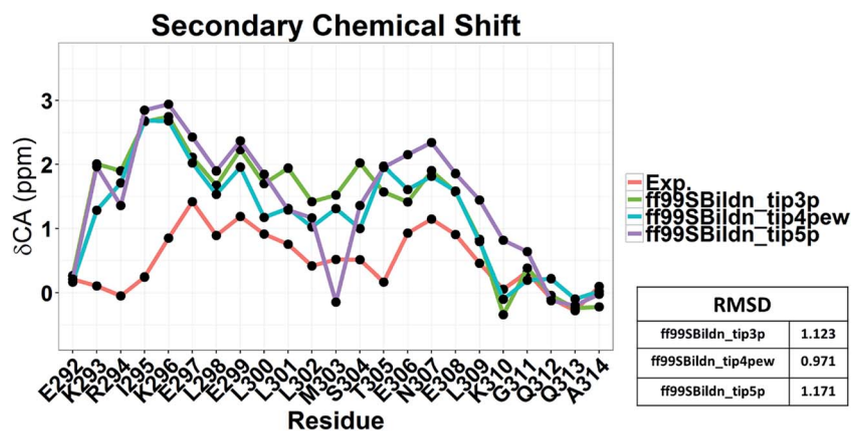


Fig. 11 Secondary chemical shift predictions from different solvent models with *ff99SBildn*.

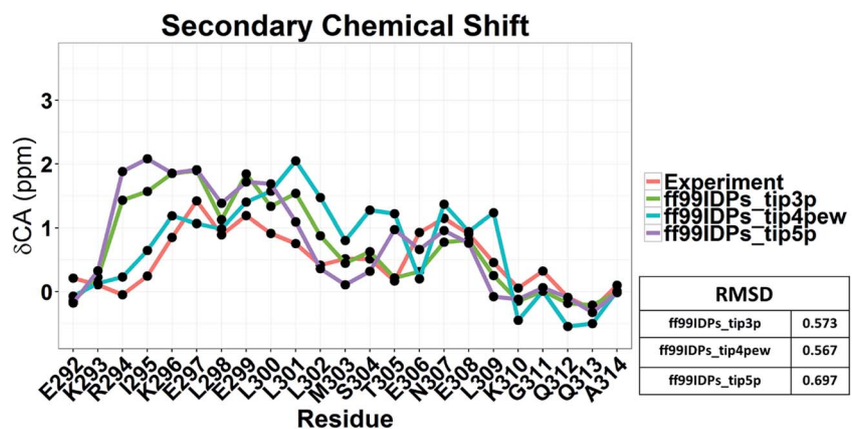


Fig. 12 Secondary chemical shift prediction from different solvent models with *ff99IDPs*.



these detailed comparison between simulation and experiment for c-Myb further supports the accuracy of *ff99IDPs* with the TIP3P or TIP4P-EW solvent in modeling IDPs structures.

Conclusion

Multiple trajectories molecular dynamics (MD) simulations with *ff99IDPs* and *ff99SBildn* force fields was used to study the conformation dynamics of c-Myb. Comparison with *ff99SBildn*, *ff99IDPs* can reproduce more diverse disordered conformers of c-Myb. The predicted secondary chemical shift under *ff99IDPs* is more approach to that of experiment than under *ff99SBildn*. Therefore, *ff99IDPs* can sample native molten globule, native pre-molten globule and native coil conformers. These results are consistent with other IDPs. Kinetic analysis of MD simulations shows that c-Myb folds *via* a two-state process and indicates that c-Myb folds in the order of tertiary folding and folding of helix. Folding nucleus of KEL plays essential role in stabilizing the folding state with dynamics correlation networks. The influences of solvent models for TIP3P, TIP4P-EW and TIP5P were also investigated and found that *ff99IDPs* and TIP3P or TIP4P-EW is the best combination to research the conformer sampling of c-Myb. These results reveal the conformation dynamics of c-Myb and confirm that *ff99IDPs* force field can be used to research the relationship between structure and function for IDPs.

Conflict of interest

The authors declare that there is no conflict of interest.

Acknowledgements

This work was supported by Center for HPC at Shanghai Jiao Tong University, the National High-tech R&D Program of China (863 Program) (2014AA021502), the National Natural Science Foundation of China (31620103901), Medical Engineering Cross Fund of Shanghai Jiao Tong University (YG2014MS47 and YG2015MS56), and National Institutes of Health/NIGMS (GM093040 & GM079383).

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