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The solution structure and dynamics of the DH-PH module of PDZRhoGEF in isolation and in complex with nucleotide-free RhoA

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Received 23 June 2009; Revised 24 July 2009; Accepted 27 July 2009

DOI: 10.1002/pro.219

Published online 7 August 2009 proteinscience.org

Abstract: The DH-PH domain tandems of Dbl-homology guanine nucleotide exchange factors catalyze the exchange of GTP for GDP in Rho-family GTPases, and thus initiate a wide variety of cellular signaling cascades. Although several crystal structures of complexes of DH-PH tandems with cognate, nucleotide free Rho GTPases are known, they provide limited information about the dynamics of the complex and it is not clear how accurately they represent the structures in solution. We used a complementary combination of nuclear magnetic resonance (NMR), small-angle X-ray scattering (SAXS), and hydrogen-deuterium exchange mass spectrometry (DXMS) to study the solution structure and dynamics of the DH-PH tandem of RhoA-specific exchange factor PDZRhoGEF, both in isolation and in complex with nucleotide free RhoA. We show that in solution the DH-PH tandem behaves as a rigid entity and that the mutual disposition of the DH and PH domains remains identical within experimental error to that seen in the crystal structure of the complex, thus validating the latter as an accurate model of the complex *in vivo*. We also show that the nucleotide-free RhoA exhibits elevated dynamics when in complex with DH-PH, a phenomenon not observed in the crystal structure, presumably due to the restraining effects of crystal contacts. The complex is readily and rapidly dissociated in the presence of both GDP and GTP nucleotides, with no evidence of intermediate ternary complexes.

Keywords: RhoA; guanine nucleotide exchange factors; protein structure; protein dynamics; nuclear magnetic resonance; small angle X-ray scattering; deuterium-hydrogen exchange mass spectrometry

Additional Supporting Information may be found in the online version of this article.

Jakub Gruszczyk was on leave from Faculty of Biochemistry, Biophysics and Biotechnology, Jagiellonian University, Krakow, Poland. Marta Kasterka was on leave from Faculty of Biology and Environmental Protection, University of Lodz, Lodz, Poland.

Grant sponsor: NIH; Grant numbers: PO1 HL48807, RO1 GM86457; Grant sponsor: EU design study SAXIER; Grant number: RIDS 011034

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Introduction

Rho (Ras homology) GTPases are a homologous family of 22 proteins that act in pathways regulating various aspects of cell physiology, including remodeling of the cytoskeleton and gene transcription. The most ubiquitous and consequently the most studied among them are RhoA, Cdc42, and Rac1.1-3 All are poor enzymes with very low k_{cat} values, so that the GTP-bound form (i.e., enzyme-substrate complex) has a long half-life. The conformation of the protein in this complex is distinct from that in the enzyme-product complex (Rho•GDP), so that the bound GTP nucleotide confers biological activity on RhoA enabling it to bind to effectors. The stability of the substrate-bound and product-bound states of Rho GTPases allows them to serve as cellular molecular switches, controlled by accessory proteins that can load GTP at the beginning of the cycle and accelerate its hydrolysis to GDP to terminate the biological signal.

The loading of GTP is catalyzed by guanine nucleotide exchange factors, or GEFs. 4-6 These large, multidomain cytosolic proteins contain a tandem of DH (Dblhomology) and PH (plekstrin-homology) domains, which together constitute the minimal functional module responsible for catalysis. It is generally accepted that the tandem functions by binding to and stabilizing nucleotide-free Rho, which upon dissociation is more likely to bind GTP than GDP because the concentration of the former in the cell is an order of magnitude higher. Other types of protein domains found in many GEFs serve additional regulatory purposes, including targeting to specific cellular sites or inhibition via intramolecular interactions. Given the biological importance of Rho GTPases and their role in various diseases, including cancer,7 there is considerable interest in the elucidation of molecular mechanisms that govern the DH-PH catalytic activity and specificity.

The RGSL family of GEFs, comprises p115,⁸ the leukemia associated RhoA-specific guanine exchange factor (LARG),^{9,10} and the PDZ-containing RhoA exchange factor (PDZRhoGEF).¹¹ All three proteins show high specificity for RhoA and are activated by select G-protein coupled receptors.¹² Their common feature is the presence of the RGSL (regulator of G-protein signaling-like) domain upstream of the DH-PH tandem. This domain interacts with the α -subunit of trimeric G-proteins, allowing for a unique cross-talk between these receptor-coupled GTP-ases and the cytosolic Rho proteins. In addition, LARG and PDZRho-GEF contain a PDZ domain at the N-terminus.

The RGSL GEFs exert their biological function in a variety of tissues, including smooth muscle, where they regulate contractility through Ca²⁺ independent pathways, a phenomenon known as Ca²⁺-sensitization.^{13–15} The DH-PH tandems of LARG and PDZRho-GEF were characterized by X-ray crystallography: the former has been studied both in isolation and in complex with RhoA,¹⁶ while the PDZRhoGEF was reported in complex with RhoA.¹⁷ Although in both complexes

the mutual disposition of the DH and PH domains is similar, it is different from the structures of homologous tandems in other GEFs. 18-22 This structural variation, along with the limited interactions typically seen in complexes between the PH domain and the GTPases, raises the possibility of distortion of the DH-PH/RhoA complex structures by crystal packing forces. Although single domain proteins are rarely distorted in any significant way by packing interactions, multidomain systems in which supramodular architecture is dependent on flexible linkers may crystallize in a number of low energy conformational states, not necessarily representing the one in solution.23 Thus, it is not clear if the DH and PH domains have fixed relative orientations in solution, or if they assume this tertiary structure only in a complex with the GTPase. Further, the constraints of the crystal lattice critically limit insight into structure dynamics in solution.

To address these questions we investigated the structure of the DH-PH tandem of the PDZRhoGEF in solution, both in isolation and in complex with RhoA, using high field-heteronuclear NMR, small angle X-ray scattering (SAXS), and H/D exchange mass spectrometry (DXMS). We find that in solution the DH-PH tandem assumes a tertiary architecture that is observed in the crystal structure of the complex, thus enabling it to readily bind the GTPase. However, we also show that in solution the DH-PH/RhoA complex displays sophisticated dynamics that are not revealed by the crystal structure. The nucleotide-free RhoA is significantly disordered when bound to the DH-PH tandem, a phenomenon that cannot be inferred from crystallographic data. Finally, we provide evidence that the complex dissociates rapidly in the presence of either GTP or GDP, virtually ruling out the presence of stable intermediates.

Results

The solution structure of the free DH-PH tandem

As in other structurally characterized GEFs, 16,17,19,20,24 the DH and PH domains of PDZRhoGEF are connected by a long α -helix, that is, helix α_6 of the DH domain. Thus, the relative disposition of the two domains depends on the stability of the connecting α -helix, which in the crystal structure is well resolved with low displacement parameters (B factors). To assess if this helix is rigid in the absence of stabilizing intermolecular packing forces in the crystal, we investigated its dynamics in solution.

We used secondary chemical shifts of backbone atoms²⁵ to probe the dynamics of isolated DH and PH domains, and of the intact DH-PH tandem. The backbone order parameters (S^2) were computed using the RCI server²⁵ based on our recently determined assignment of individual domains and DH-PH tandem²⁶ (Fig. 1). As expected, the order parameters in individual domains are very similar to those of the tandem. Lower

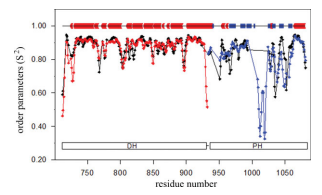


Figure 1. Order parameters (S^2) derived from backbone chemical shifts of DH (red), PH (blue), and DH-PH (black) using RCI server.²⁵ The diagram above the plot shows the location of secondary structure elements in the crystal structure of DH-PH in complex with RhoA; α-helices are shown in red and β-strands are blue.

 S^2 values in the tandem indicate enhanced backbone flexibility and correlate well with the location of surface loops. The PH domain exhibits significantly higher intrinsic dynamics than the DH domain, with several surface loops showing particularly high mobility and in particular the loop encompassing residues 1008–1021, which is completely disordered in the crystal. With respect to the α_6 -helix, which connects the DH and PH domains, the chemical shifts do not indicate any increase in dynamics, although the assignment could

not be extended beyond Ala923 except for shifts assigned to residues 935–937 within the PH-domain portion of the helix (Fig. 1). We have additionally measured $^{15}\mathrm{N}\{^1\mathrm{H}\}$ NOE spectra and found that Gly935 does not show any decrease in the heteronuclear NOE effect, consistent with a well ordered conformation (Supporting Information Figure 1). These results strongly suggest that the α_6 -helix remains intact in the DH-PH tandem in solution, with the concomitant fixed relative disposition of DH and PH domains. To obtain independent experimental confirmation, we used two complementary approaches: residual dipolar couplings (RDCs) and small angle SAXS.

Residual dipolar couplings (RDCs) provide accurate measure of the orientation of backbone N-H groups (1D_{HN}) and allow for direct comparison with X-ray coordinates.27-29 We measured 1DHN RDCs for the DH-PH tandem aligned in a positively charged polyacrylamide gel (Fig. 2). Despite considerable broadening of the signals, we obtained a set of 69 $^{1}D_{HN}$ values, that is, $\sim\!30~\%$ of all assigned backbone amides, including 35 within the DH domain and 34 within the PH domain (see Table I). First, we assessed if the ¹D_{HN} values of the individual domains are consistent with the DH-PH/ RhoA crystal structure, using the Q quality factors.³⁰ Fitting the experimental RDCs to the two crystallographically independent DH domains yielded Q-factors of 36.1% and 37.1%, a good agreement assuming an experimental error of 4 Hz. For the two PH domains the

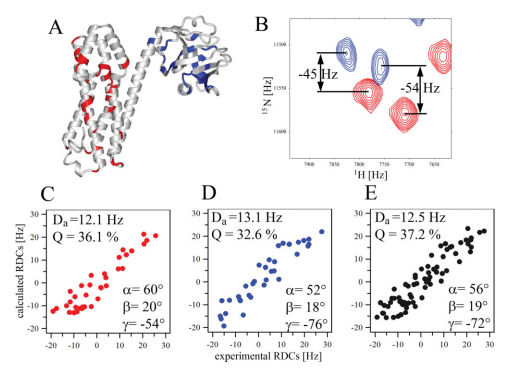


Figure 2. Domain orientation in DH-PH studied by RDCs. (A) DH-PH from the crystal structure of the complex showing the locations of $^1D_{HN}$ RDCs measured for residues in DH (red) and PH (blue); (B) an example of 1H - ^{15}N TROSY-HSQC (red); and 1H - ^{15}N HSQC (blue) for 2H , ^{13}C , ^{15}N -DH-PH aligned in positively charged polyacrylamide gel used to extract $^1D_{HN}$ RDCs. (C–E) Fitting of experimental RDCs for DH, PH, and DH-PH, respectively, to the crystal structure. Magnitude of the alignment tensor (D_a), Q-factors and Euler angles are shown for each analysis.

Table I. Analysis of Alignment Tensor Parameters Based on RDCs Measured for DH-PH and DH-PH/RhoA Complex

	Q [%]	$D_{\mathrm{a}} \left[\mathrm{Hz} \right]$	α	β	γ	N
DH ^a PH ^a DH-PH ^a DH-PH/RhoA ^b	36.1 ± 2.6 32.6 ± 2.1 37.2 ± 1.7 39.0 ± 2.3	12.1 ± 0.4 13.1 ± 0.5 12.5 ± 0.3 16.5 ± 0.6	59.6 ± 1.1 52.4 ± 1.3 56.0 ± 0.8 57.5 ± 2.3	20.5 ± 1.7 18.5 ± 1.2 19.0 ± 0.9 19.3 ± 1.2	-54.5 ± 16.5 -75.8 ± 2.6 -71.7 ± 3.9 -78.5 ± 3.2	35 34 69 37

Calculations of Q-factors (Q), magnitude of the alignment (D_a) and Euler angles (α , β , γ) defining rotations of molecular coordinates about x, y, and z axes, relative to principal axes frame of alignment tensor were carried out employing program PALES31 based on the crystal structure of DH-PH/RhoA complex (PDB code 1XCG, molecule A) and using N experimental ¹D_{HN} RDCs (see Materials and Methods for details).

Q-factors were 32.6% and 29.3%, respectively. Next, to establish the mutual orientation of the DH and PH domains we calculated their respective alignment tensors (D_a) from the fit of experimental RDCs to the crystal structure. Their magnitudes are very similar and the Euler angles representing the average orientation of the molecules relative to the alignment tensor frame are also similar (Fig. 2, Table I). Simultaneous fitting of the full set of ¹D_{HN} RDCs to the DH-PH tandem in the crystal structure yields a Q-factor of 37.2%, comparable to the fit for the individual domains. Thus, the NMR data confirm that the two domains in the tandem are mutually oriented in the same way as in the crystal structure. However, these data do not exclude a change in relative translation.

We used small angle SAXS to determine the overall shape of the molecule in solution. The composite experimental scattering pattern from DH-PH is presented in Figure 3, and the overall structural parameters computed from the SAXS data are given in Table II. Several independent runs of the ab initio program DAMMIN³² yielded reproducible molecular shapes. The average model generated by DAMAVER33 is consistent with the crystal structure of the DH-PH tandem in complex with RhoA (Fig. 3). The value of normalized spatial discrepancy (NSD) between the two is 1.0, that is, the two molecular envelopes are virtually identical at low resolution. Furthermore, a simulated scattering curve computed by CRYSOL³⁴ using the atomic coordinates of DH-PH fits very well to the experimental data (Table II).

The structure of the DH-PH/RhoA complex in solution: Overview of the NMR strategy

To gain further insight into the structure and dynamics of the DH-PH/RhoA complex in solution, we analyzed two samples: 2H,13C,15N-labeled DH-PH bound to unlabeled RhoA; and 2H,13C,15N-labeled RhoA bound to an unlabeled DH-PH tandem. This approach allowed for independent analyses of the DH-PH tandem and RhoA in the complex. Despite the size of the complex (64 kDa), the 1H-15N TROSY-HSQC spectra were of high quality [Fig. 4(A,C)]. When either of the two labeled proteins was in excess, the corresponding spectra showed two sets of signals (not shown) consistent with slow exchange kinetics, typical of strong

complexes with dissociation constants in the nM range. As expected, the formation of the complex results in strong chemical shift perturbations for both RhoA and DH-PH (Fig. 4). Importantly, changes in the spectra of RhoA upon binding to the DH-PH tandem are more pronounced compared to those occurring in the tandem, consistent with significant structural perturbations of RhoA

To obtain chemical shift assignments for quantitative analysis of the two proteins in the complex, we conducted TROSY-based HNCO, and HNCA experiments and 15N-edited NOESY at 900 MHz, for samples in which either the DH-PH tandem or RhoA were triple-labeled. The assignment was based on sequential information derived from HNCA and NOESY experiments and comparison of backbone chemical shifts to those of isolated proteins. Using this strategy, we assigned 184 of the 258 residues previously assigned for free DH-PH. This was primarily due to peak

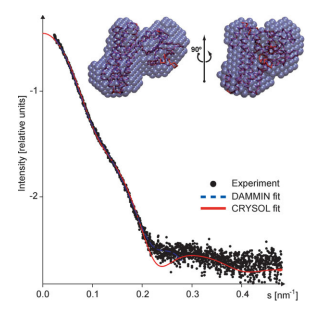


Figure 3. SAXS profiles of DH-PH (see text for details). Dots denote the experimental data, the fits obtained by DAMMIN³² and CRYSOL are shown as blue dashed and red solid lines, respectively. Inset: The averaged ab initio shape obtained from SAXS data is presented by blue beads, the crystallographic model is shown as red Cα-trace.

The RDCs measured for DH-PH were fit to coordinates of separate DH, PH domains, and DH-PH tandem.

^b The analysis of RDCs measured for DH-PH in the complex with RhoA.

Table II. Overall Structural Parameters of DH-PH Obtained by SAXS

R _g (nm)	D _{max} (nm)	$V_{\rm p}~({\rm nm}^3)$	MM (kDa)	χ_{s}	χcryst
2.9 ± 0.1	9.5 ± 0.5	60 ± 15	40 ± 10	1.6	1.6

 $R_{\rm g},\,D_{
m max},\,V_{
m p},$ and MM are radius of gyration, maximum size, excluded volume, and molecular mass, respectively.

 χ_s and χ_{cryst} are discrepancy between the experimental data and computed scattering curves from ab initio shape and atomic model, respectively.

overlap although a number of resonances were strongly perturbed and difficult to assign unambiguously. With respect to RhoA, we were only able to assign 89 residues in the complex, although in this case the analysis was complicated by strong broadening of numerous resonances (Figure 4). In particular, we could not assign any of the residues from Switch I and Switch II that are in direct contact with DH domain in the crystal structure.

The structure of the DH-PH tandem in complex with RhoA

To assess how the two proteins are affected by complex formation, we mapped the chemical shift perturbations onto the crystal structure of the DH-PH/RhoA

complex [Fig. 5(A)]. In the DH-PH tandem, the most pronounced changes affect, as expected, those residues within the DH domain which are in the direct contact with RhoA in the crystal structure. A total of 43 amides within the DH domain showed resonances that were shifted to such degree that they could not be assigned. In the PH domain, the chemical shift perturbations were much smaller than those in the DH domain and were limited to two surface patches: one involving Met966, and Glu969 and some neighboring residues, in proximity of both RhoA and the DH domain, and the second, involving Ser1065 and its neighbors, in contact with Glu97 of RhoA. The small magnitude of the chemical shift changes observed for PH amides in the latter patch indicate that the contact is weak.

We also conducted cross-saturation experiments,³⁵ with saturation of the protonated RhoA and monitoring of deuterated DH-PH (not shown). We found that the only residues exhibiting significant cross-saturation effect were within the DH domain. No cross saturation was detected between RhoA and PH, confirming that these contacts are weak.

To determine experimentally the orientation the two domains in DH-PH module in the complex, we

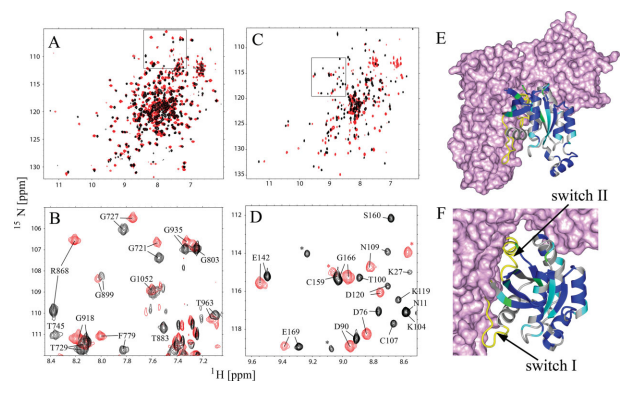


Figure 4. Solution NMR of the DH-PH/RhoA complex. (A) Comparison of the $^1\text{H}-^{15}\text{N}$ TROSY-HSQC spectra of DH-PH (black) and DH-PH/RhoA (red); (C) comparison of the $^1\text{H}-^{15}\text{N}$ TROSY-HSQC spectra of RhoA (black) and RhoA-DHPH (red); (B,D) assigned fragments of the squared regions in (A) and (C), respectively; unassigned residues are labeled with asterisk; (E,F) mapping of the assigned resonances onto the model of RhoA (ribbon) in the complex with DH-PH (surface representation); residues exhibiting small chemical shift perturbations ($\Delta\sigma < 0.06$ ppm) are blue, medium (0.06 ppm $< \Delta\sigma < 0.15$ ppm) are cyan and strongest ($\Delta\sigma > 0.15$ ppm) are green; residues that were not assigned are gray; chemical shift perturbations were calculated using the following formula $\text{sqrt}(\Delta\sigma_{\text{HN}}^2 + 0.1 \ \Delta\sigma_{\text{N}}^2)$. Switch I and II regions of RhoA which were not assigned are shown in yellow and labeled.

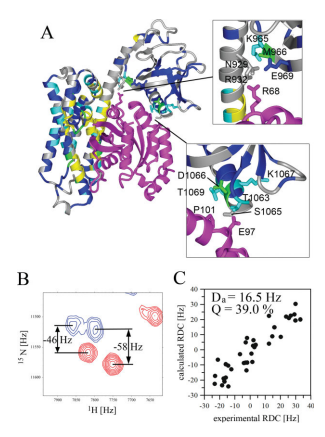


Figure 5. Characterization of the structure of DH-PH/RhoA complex in solution; (A) mapping of chemical shift perturbations within DH-PH upon binding to RhoA (magenta); two regions showing contacts involving RhoA and PH domain are shown in insets; residues exhibiting small chemical shift perturbations ($\Delta \sigma < 0.05$) are blue, medium (0.05 $< \Delta \sigma <$ 0.10) are cyan and strongest ($\Delta \sigma >$ 0.10) are green; residues that could be assigned only for free DH-PH and experience very strong perturbations in the complex are yellow; residues without assignment for free DH-PH are gray; chemical shift perturbations were calculated using the following formula sqrt($\Delta \sigma_{HN}^2 + 0.1 \Delta \sigma_N^2$). (B) An example of $^1H^{-15}N$ TROSY-HSQC (red) and ¹H-¹⁵N HSQC (blue) for ²H, ¹³C, ¹⁵N-DH-PH/ RhoA aligned in positively charged polyacrylamide gel, used to extract ¹D_{HN} RDCs; (C) fitting of RDCs measured for the DH-PH in the complex with RhoA to the crystal structure.

used residual dipolar couplings (RDCs), measured for the complex aligned under conditions similar to those used for the free DH-PH tandem [Fig. 5(B,C)]. Because of the strong broadening and overlap of the signals, we were able to measure accurately only 37 ¹D_{HN} RDCs including 16 and 21 RDCs for DH and PH, respectively. Fitting of these RDCs to the crystal structure yielded the Q-factor of 39%. Although the number of RDCs included in this calculation is relatively small, this result strongly suggests that the relative orientation of the DH and PH domains in the complex in solution is accurately represented by the crystal structure.

Structure and dynamics of RhoA in complex with DH-PH

Close examination of assigned chemical shifts for RhoA in complex with DH-PH reveals significant structural perturbations throughout the protein, including sites distant from DH-PH binding surface [Fig. 4(C)]. We find that in solution the subset of RhoA resonances in the DH-PH bound state exhibit significant degree of broadening, consistent with a much higher level of dynamic disorder, with ~20% of amides, including all of those in Switch I and Switch II, broadened beyond detection in the ¹H-¹⁵N TROSY-HSQC spectrum. Such resonance broadening is not caused by overall increase in molecular weight of RhoA in the complex but most likely it results from backbone motions occurring on the microsecond to millisecond timescale. In contrast, the crystal structure of RhoA in complex with DH-PH of PDZRhoGEF³⁶ shows relatively little disorder, with temperature (B) factors elevated slightly for residues in the RhoA's Switch I region.

To further probe the dynamics in the DH-PH/ RhoA complex, we used DXMS. This technique is useful for monitoring the dynamics of interacting surfaces in protein-protein complexes on a significantly longer timescale than that probed by NMR.37,38 When the quenched complex is proteolysed, the peptides with the comparatively lower deuterium content map the interacting surfaces or regions with increased stability. In our case, proteolytic digestion yielded 44 and 24 reporter peptides, covering 94% and 96% of the DH-PH tandem and RhoA sequences, respectively (Supporting Information Figure 2). The H/D experiments were carried out for the isolated DH-PH tandem, isolated RhoA•GDP, and for the DH-PH/RhoA complex. The results were mapped on the crystal structure of the complex [Fig. 6(A)]. For both proteins there were several overlapping reporter peptides that provided a consistency test.

We identified three sets of peptides with H/D exchange rates that are lower in the complex than in the free proteins (Fig. 6). The first set maps to the contact between helices 3101a and α1a of the DH-PH tandem, and the Switch I region of RhoA; these peptides show a modest, but measurable ~5 % reduction in deuterium labeling compared to free proteins. A second set of reporter peptides corresponds to the contact between fragment comprising helices α4a, 3104b, and α5a of DH-PH, and Switch II region of RhoA; in this case we observe a much more dramatic, \sim 20 % decrease in the H/D exchange. These results are in apparent contrast to the NMR spectra which show conspicuous absence of both Switch I and Switch II amide resonances in the 1H-15N TROSY-HSQC spec-(see above). However, broadening of NMR resonances results from dynamics on microsecond to millisecond timescale, which is significantly faster than the time required to obtain DXMS data. The

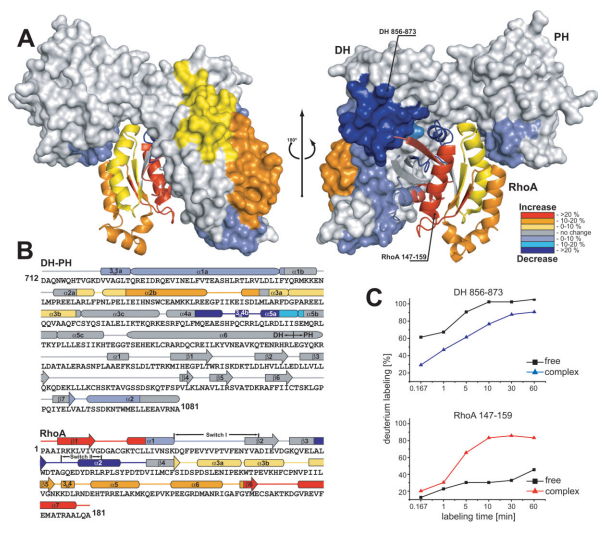


Figure 6. The deuterium exchange map of the DH-PH/RhoA complex. (A) Deuteration level of the DH-PH/RhoA complex shown after 60 min of ion-exchange reaction. Regions of DH-PH (surface) and RhoA (ribbon) affected by the formation of complex are color-coded, based on the change in the deuteration level, suggested by H/D exchange. Areas of the protein with significant decrease in exchange upon complex formation are colored dark blue (deuteration reduction of >20%), blue (reduction of 10–20%), and slate (reduction of 0–10 %). Regions of increased solvent accessibility are shown in red (deuteration increase of >20%), orange (10–20%), and yellow (0–10%). Parts of the protein with no significant changes in deuteration level are shown in grey (PDB ID: 1XCG). (B) The secondary structure elements of DH-PH and RhoA are shown above the protein sequences, and the color-coding represents changes in the levels of deuteration. (C) Two examples of the differences in deuteration levels as a function of time in free DH-PH and RhoA, and after formation of the binary complex. Reduction in deuteration was observed for peptide 856–873 from DH-PH tandem, whereas H/D exchange level increased for peptide 147–159 from RhoA.

interaction involving Switch I of RhoA appears to be generic for a variety of RhoGEFs, ^{19,20,24,36} while the Switch II interaction was suggested to be largely responsible for RhoA selectivity in the p115 family of GEFs. ¹⁷

Finally, the third area of contact between DH-PH and RhoA involves Ser1065 and Asn1068 from the helix $\alpha 2$ of the PH domain (residues 1060–1073), which form hydrogen bonds with Glu97 of RhoA.¹⁷ The DXMS results show modest ($\sim 5\%$) reduction in H/D exchange in the $\alpha 2$ helix. Unexpectedly, large portions of both DH-PH and RhoA exhibited higher H/D exchange in the complex than in isolation. In the DH domain, helices $\alpha 2a$, $\alpha 2b$, $\alpha 3a$, and $\alpha 3b$, compris-

ing residues 776–832 and distal to the RhoA-binding surfaces, all show $\sim\!\!10$ % increase in the H/D exchange, suggesting that they are destabilized as a result of complex formation with RhoA. The GTPase, on the other hand, exhibits marked overall 5–35 % increase in H/D exchange compared to its GDP-bound form, and in agreement with the enhanced dynamics suggested by the NMR data (above). The only exceptions are the two Switch regions, both of which interact with the DH domain.

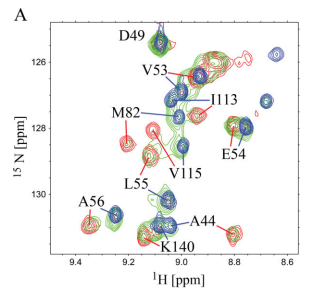
A comparison of the DXMS data for the DH-PH/RhoA complex with temperature (B) factors from the crystal structure reveals poor correlation (Supporting Information Figure 3). This is particularly evident in

the DH domain helices α2a, α2b, α3a, and α3b (residues 776-832). The ~10 % increase in H/D exchange in this region is inconsistent with relatively low crystallographic B-factor values. However, the most dramatic contrast between the DXMS and X-ray crystallographic data, relates to Switch I and II regions of RhoA. DXMS indicates that both of these regions become significantly shielded from H/D exchange upon formation of a complex with DH-PH. On the other hand, temperature (B) factors show increase in the Switch I region and only mild decrease for Switch II compared to the RhoA•GDP structure, while the NMR data is consistent with very strong conformational fluctuations of both Switches on a short timescale.

The interpretation of the data was not straightforward because we were unable to obtain NMR data for the nucleotide-free RhoA. The differences between RhoA in complex with DH-PH and RhoA•GDP are not really representative of the simple transition from nucleotide-free RhoA to GEF-bound form, but rather of a two-step transition from GDP-bound RhoA to nucleotide-free RhoA and then to complex with DH-PH. The overall destabilization of RhoA observed by DXMS in complex with DH-PH is most probably a direct consequence of the loss of nucleotide. Both Switch regions remain shielded in the complex from solvent, and the complex is quite stable on a long timescale, but within the complex the Switches undergo rapid conformational exchange. The loss of conformational entropy in RhoA as a result of complex formation may be in part compensated by the partial destabilization of the DH domain (helices $\alpha 2$ and $\alpha 3$), resulting in an "entropy transfer" effect. Overall, the temperature (B) factors of the complex in the crystal structure are not representative of true complex dynamics, because they are constrained by extensive crystal contacts, particularly between the DH domain and RhoA from a symmetry related complex.

Dissociation of the DH-PH/RhoA complex by guanine nucleotides

The dynamic behavior of RhoA in the complex may be an important functional feature which facilitates nucleotide exchange. To directly observe whether guanine nucleotides are able to bind to and dissociate from the complex, we titrated the DH-PH/2H,15N-RhoA complex with GDP and GTP (Fig. 7). The addition of even substoichiometric amounts of nucleotides, in the presence of 1 mM Mg2+ gave rise to a second set of signals characteristic of free RhoA. Thus, in the presence of the guanine nucleotide, either GDP or GTP, the complex undergoes rapid dissociation and regeneration of the nucleotide-bound RhoA. We see no evidence of any transition complex involving DH-PH and nucleotide-bound RhoA, as was suggested by others,39 indicating that even if such a transition state exists



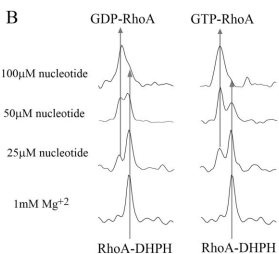


Figure 7. Nucleotide induced dissociation of the DH-PH/ RhoA complex. (A) fragment of the superimposed ¹H-¹⁵N TROSY-HSQC spectra showing the comparison of the of 80 μ M nucleotide free DH-PH/²H, ¹⁵N-RhoA (red), the 80 μ M DH-PH/2H, 15N-RhoA complex in the presence of 50 μM GDP (green) and 100 μM ²H, ¹⁵N RhoA•GDP (blue). (B) extraction of 1D traces from ¹H-¹⁵N TROSY-HSQC spectra used to follow the dissociation of the complex by guanine nucleotides; the arrows show chemical shifts of the Cterminal Ala181 amide proton of RhoA bound to DH-PH (upfield signal) and nucleotide (downfield signal); left and right panels shows the titration of DH-PH/RhoA complex with GDP and GTP, respectively.

transiently, it is unstable and undergoes very rapid dissociation.

We next used NMR spectroscopy to compare the relative abilities of GDP and GTP to dissociate RhoA from the complex with DH-PH. The advantage of this approach was that we monitored the behavior of the proteins in the presence of unmodified nucleotides. For this purpose, we quantified the amount of dissociated RhoA based on the intensities of the amide proton resonance of C-terminal Ala181. As shown in

Figure 7(B), in the presence of GTP the dissociation of RhoA is somewhat more efficient than in the presence of GDP. We estimated that GTP has 1.5 times higher affinity towards the complex than GDP. We have also obtained similar results in the independent fluorescence-based assay titrating DH-PH/RhoA complex with N-methylanthraniloyl (mant) analogues of GTP and GDP (data not shown). The slightly higher affinity of GTP to the complex is not sufficient to drive the formation of the active RhoA•GTP complex in the cell. Therefore, the major factor is the ten times higher cellular level of GTP compared to GDP.

Discussion

Using heteronuclear NMR and SAXS we showed that the two domains within the isolated DH-PH tandem retain fixed relative position in solution, so that the protein tumbles as a monomeric and rigid body. Although we were not able to fully assign all the amide resonances within the connecting α_6 -helix of the free tandem, we see no indication of disorder or noticeably increased dynamics within this helix that would be consistent with independent motions of the two domains. It is not immediately obvious if the rigid architecture is a product of cooperative effects, or if the α_6 -helix provides an intrinsically stable scaffold, but the data in Figure 1 suggest that in the absence of the PH domain the C-terminus of the helix unfolds.

The rigid architecture of the DH-PH tandem sets it apart from a number of other multidomain proteins, such as calmodulin in which the two globular domains are connected through a linker which adopts a well defined α -helical conformation in the crystal structure but is clearly very flexible in solution.⁴⁰ Further, we show that the mutual disposition of the two modules in the isolated tandem is identical, within the experimental error, to that observed in the crystal structure of the DH-PH/RhoA complex.¹⁷

The stable mutual disposition of the DH and PH domains raises the question of its biological significance. Other structural studies of DH-PH tandems in various GEFs, both in isolation and in complexes with cognate GTPases, seem to support the notion that this moiety functions as a rigid unit. In some cases, this architecture primes the tandem for binding of GTPase in a cooperative fashion, so that both the DH and PH domains engage the GTPase. This appears to be the case for Dbs and Trio. 18,41 Recently, two independent structural investigations have shown that in Vav1, a cysteine-rich domain (CRD) located downstream of PH, packs together with the latter against the C-terminal helix of the DH domain to stabilize its conformation.42,43 However, the impact of this architecture on the nucleotide exchange is indirect, because neither the PH domain nor the CRD domain engages the GTPase.

It has been shown previously that in the absence of the PH domain, the catalytic efficiency of the isolated DH domain of PDZRhoGEF is impaired, and the rate of nucleotide exchange reaction is lowered 3fold.36 However, the mutations of the PH residues Ser1065 and Asn1068 to alanines, both of which are in the contact with RhoA, have no impact on the nucleotide exchange kinetics.36 The modest PH-RhoA interface and small perturbations of PH chemical shifts upon RhoA binding suggest that the interaction is weak and transient. The second site within the PH domain which exhibits chemical shift changes upon RhoA binding is found in the region bridging DH, PH, and RhoA. In the crystal structure, Asn929 (DH domain) forms hydrogen bonds with Arg68 (RhoA) and with the backbone carbonyl of Met966 (PH domain). Mutation of Asn929 to Ala results in significant reduction of nucleotide exchange rate.³⁶ Thus, the PH domain seems to have no direct role in the nucleotide exchange process, but indirectly affects the DH interaction with RhoA. Also, neither the PH domain of PDZRhoGEF nor the homologous PH domains of the related LARG and p115 exchange factors, bind phospholipids, ruling out their involvement in membrane targeting of the GEFs. However, we note that the electrostatic potential surface of the rigid DH-PH tandem may be of importance in orienting the DH domain in a favorable position to interact with membrane anchored RhoA, assuming that the translocation of the GTPase to the membrane from the RhoGDI/RhoA complex precedes the nucleotide exchange event.

It is noteworthy, that the DH-PH tandem of LARG shows a significant difference in the position of its PH domain between the crystal structures of the isolated tandem and of the complex with RhoA: the PH domain is laterally translated by more than 10 $Å^{16}$. Assuming that the results of our solution studies extend to LARG, this can only be rationalized by a distortion of the LARG tandem due to crystal contacts. Indeed, we find a unique set of crystal contacts in the structure of the isolated DH-PH tandem, so that each DH domain contacts six symmetry related DH domains, while each PH domain contacts only one other PH domain. While crystal packing forces rarely have a significant impact on the conformation of single domain globular proteins, their energy is sufficient to distort the mutual disposition of domain connected by relatively flexible secondary structure elements.²³

Finally, using NMR and DXMS, we show that RhoA shows significantly increased dynamics in solution in complex with DH-PH, specifically within the Switch I and Switch II regions, a phenomenon not evident from the crystal structure which is effectively stabilized by both the crystal contacts and the low temperature of the experiment. Molecular dynamics play an important role in protein-protein interactions in solution, but are not well represented by the crystallographic models. Destabilization of one of the partners within a complex may be a relatively common occurrence. In the case of the nucleotide exchange reaction,

this may be particularly important, as exemplified by the study of Rab8, which undergoes partial unfolding during such exchange catalyzed by its exchange factor MSS4.44 Similar effect of conformational dynamics was observed for the AN 1-4 ankyrin repeats in iκB, which become destabilized upon interaction with NFκB.⁴⁵ In the case of the DH-PH/RhoA complex, the GTPase shows increased dynamics on the microseconds to milliseconds scale, particularly within the Switch I and Switch II regions. However, both these regions are protected from H/D exchange in the complex on a longer time scale, indicating that the integrity of the complex is not compromised by the dynamics of the GTP-ase. It is also interesting to note that portions of the DH domain distal to the RhoA-binding site seem to be destabilized in the complex, compared to isolated state, as judged by the DXMS analysis. One possible explanation is that unfavorable loss of entropy by the GTPase on binding the DH-PH tandem is compensated by increase in the entropy in the DH domain. This suggests that specific interactions of other domains within intact GEFs, with the distal face of the DH domain, may provide additional regulatory mechanism, effectively lowering the affinity of the DH domain for RhoA, and inhibiting the nucleotide exchange rate.

Our experiments underscore the power of solution structural studies as a complement to crystallographic investigations of regulatory complexes.

Materials and Methods

Protein expression and purification

The DH-PH tandem of human PDZ-RhoGEF (residues 712-1081) was expressed as fusion protein with GST in E. coli BL21(DE3)-RIL (Stratagene) and purified as described previously.46 The purified protein was concentrated in 50 mM Tris-HCl, pH 7.5, 150 mM NaCl, and 1 mM DTT. Human truncated RhoA (residues 1-181), was expressed in pETUni vector⁴⁷ in fusion with a His6-tag. RhoA was purified by Ni-NTA-agarose affinity chromatography in 50 mM Tris-HCl, 400 mM NaCl, 1 mM MgCl₂, 5 mM imidazole, pH 8.0, and eluted with 150 mM imidazole buffer, before final purification using gel filtration. Proteins labeled with stable isotopes (2H, 13C, 15N) were expressed in minimal media with (15NH₄)₂SO₄ as sole source of nitrogen and ¹³C- or ²H, ¹³C-glucose as a source of carbon. The media were enhanced by the addition of labeled Bio-Express (Cambridge Isotope Labs). Four types of labeling scheme were used: 15N; 13C, 15N; 2H, 15N; and 2H, ¹³C, ¹⁵N.

Preparation of protein samples for NMR experiments

For assignment experiments a sample of 0.9 mM¹³C, ¹⁵N RhoA was prepared in 25 mM TRIS buffer, pH 7.5 with 50 mM NaCl, 5 mM MgCl₂, 1 mM DTT

and saturated with GDP. Samples of DH-PH for NMR experiments contained between 0.2 and 0.35 mM protein exchanged to 200 mM MOPS/TRIS, pH 7.5 buffer.²⁶ The DH-PH/RhoA complex was prepared by mixing approximately equimolar amounts of the DH-PH tandem and RhoA, with 10% excess of the unlabelled component and subsequent dialysis of the sample to remove GDP and Mg⁺² ions. We found that long term solubility of DH-PH/RhoA complex in 200 mM MOPS/TRIS, pH 7.5 is slightly lower than that of DH-PH tandem, but the samples of the complex were nevertheless stable at 0.25 mM for the period of a week at 25°C.

NMR experiments

Measurements for ¹³C, ¹⁵N-RhoA were carried out using Varian Inova 500 MHz at 25°C. To assign backbone chemical shifts we recorded standard triple resonance experiments [HNCO, HNCA, HN(CO)CA, HNCACB, CBCA(CO)NH and 3D 15N-edited NOESY with 200 ms mixing time] and we have been able to assign 148 of 168 (88 %) backbone amides. To assign the DH-PH and RhoA in the complex we prepared two 0.35 mM samples of 2H,13C,15N-labeled RhoA complexed to unlabeled DH-PH and ²H, ¹³C, ¹⁵N-labeled DH-PH in complex with unlabeled RhoA. The spectra (TROSY-HNCA, TROSY-HNCO, 3D ¹⁵N-edited TROSY-NOESY with 250 ms mixing time) were measured using Varian Inova 900 MHz spectrometer at 28°C. Assignment of the binary DH-PH/RhoA complex was based on comparison of HN, N, C', Ca backbone chemical shifts to the chemical shifts of free proteins.26 The sequential connectivities were additionally verified using correlations from TROSY-HNCA and TROSY-NOESY experiments. Overall, we assigned the total of 50% and 53% of backbone amides for the complexed DH-PH and RhoA, respectively.

The TROSY-based ¹⁵N{¹H} heteronuclear NOE experiment with 3 s saturation was measured for free DH-PH at 28°C using a Bruker Avance III 800 MHz spectrometer.

To measure cross-saturation effects we modified the 1H-15N TROSY-HSQC experiments to introduce saturation of aliphatic resonances.35 The cross-saturation spectra for 2H,13C,15N-RhoA bound to unlabeled DH-PH were measured at 28°C with 0.6 and 1.2 s saturation times using Varian Inova 600 MHz. Experiments for the 2H,13C,15N-DH-PH complexed with unlabeled RhoA were carried out at 28°C with saturation times of 0.3, 0.6, and 1.0 s using Varian Inova 900 MHz.

Measurement and analysis of RDCs

The alignment of the DH-PH and DH-PH/RhoA complex was successfully obtained using a positively charged gel: 50% (3-acrylamidopropyl)-trimethylammonium chloride/50% acrylamide (referred to hereafter as 50+M).48 The RDCs were measured for 0.5 and 0.35 mM samples of 2H,13C,15N-labeled DH-PH and 0.35 mM2H,13C,15N-labeled DH-PH/RhoA, respectively. Introduction of weak alignment resulted in signal broadening and decreased the quality of the spectra, making it impossible to measure J-couplings by HNCO-based triple resonance experiments. Instead, we measured ¹H-¹⁵N HSQC and ¹H-¹⁵N TROSY-HSQC in interleaved manner and extracted the anisotropic ¹J_{HN} coupling from the peak positions in ¹⁵N dimension. All measurements were carried out using Varian Inova 900 MHz spectrometer at 30°C. To calculate the ¹D_{HN} RDCs we assumed the isotropic ¹J_{HN} value of -94 Hz.49 The calculated 1DHN ranged from -19 to +28 Hz and -23 to +33 Hz for DH-PH and DH-PH/ RhoA, respectively. On the basis of the repeated experiments, we estimated 4 Hz error in ¹D_{HN} determination.

The determination of the alignment tensor parameters (D_a and R) and of the Euler angles (α , β , γ) defining rotations of molecular coordinates about x, y, and z axes, relative to principal axes frame of alignment tensor, was carried out using the PALES program³¹ and the crystal structure of DH-PH/RhoA complex (PDB code 1XCG). Compatibility between experimental and calculated RDC was evaluated based on quality factors Q calculated from the formula: $Q = \text{rms } (D^{\text{calc}} - D^{\text{obs}})/\text{rms}(D^{\text{obs}})$. Evaluation of the error in determination of alignment tensor and domain orientation has been done using jack-knife procedure by performing 100 cycles of calculation with random elimination of 10% of the data.²³

Titration of DH-PH/RhoA with nucleotides

For the titration of the DH-PH/RhoA complex with nucleotides, we prepared a sample of unlabeled DH-PH in complex with 2 H, 15 N-labeled RhoA, at 100 μ M concentration, in 200 mM MOPS/TRIS buffer, pH 7.5 with 1 mM MgCl₂. NMR samples were prepared by mixing the complex with stock solutions of GDP and GTP in the same buffer. Final concentration of the protein complex in NMR samples was 80 μ M and nucleotides were at 25, 50, and 100 μ M. To minimize the hydrolysis of GTP all samples were prepared and stored on ice. The 1 H- 15 N TROSY-HSQC spectra were measured at 28°C for 30 min.

Small angle X-ray scattering

SAXS measurements were performed at the X33 beamline (DESY, EMBL Outstation Hamburg, Germany)⁵⁰ using MAR345 area detector. The sampledetector distance was 2.7 m. Data were collected at protein concentrations of 1–10 mg/mL with exposure times of 120 s, at 5°C. SAXS intensity I is represented as a function of the momentum transfer modulus q [$q = 4\pi/\lambda \sin(\theta)$, where λ is the radiation wavelength(1.5 Å), and 2 θ is the scattering angle. Scattering from the buffer was collected before and after that of the sample, then averaged and subtracted from the sample

scattering curve to generate the scattering profile of the protein [I(q)]. Data at different concentrations were scaled and merged using the program PRIMUS.51 A Guinier approximation $[I(q) = I(o)\exp(-q^2R_q^2/3)]$ describing scattering intensity at a low angle region of the scattering profile was used to calculate the radius of gyration (R_g) and zero-angle scattering [I(o)] from a linear fit to the Guinier plot using the program AutoRg.52 The molecular mass (MM) of the solute was evaluated by comparison with the reference solutions of bovine serum albumin using the proportion MM $\sim I(o)/c$. In addition, the excluded (Porod) volumes $V_{\rm p}^{53}$ of the solute was also analyzed taking advantage of the fact that this parameter does not depend on concentration. The maximum size D_{\max} and real-space distance distribution function [P(r)] was determined with the program GNOM,54 which implements the indirect Fourier transform method applied to the experimental scattering plot I(q). Multiple independent reconstructions of three-dimensional shapes with the best fit to the experimental scattering curves were modeled ab initio using the DAMMIN program,32 and subsequently aligned and averaged in the program DAMAVER.33 The averaged low-resolution SAXS envelope was aligned with the high-resolution atomic structure of the DH-PH tandem (PDB ID: 1XCG) using the program SUPCOMB55 which minimizes normalized spatial deviation (NSD) between the models. The scattering from the atomic model was calculated using the program CRYSOL,34 which either predicts theoretical scattering patterns or fits the experimental data by adjusting the excluded volume and the contrast of the hydration layer.

Hydrogen-deuterium exchange mass spectrometry

Samples of the DH-PH and RhoA were prepared as described above. The DH-PH/RhoA complex was prepared by mixing DH-PH tandem and RhoA in 1:1.5 molar ratio and subsequent dialysis of the sample to 50 mM Tris-HCl, pH 7.5, 100 mM NaCl, 1 mM EDTA, and 1 mM DTT. To remove the excess unbound RhoA, the complex was subsequently purified using gel filtration chromatography.

Hydrogen/deuterium (H/D) isotopic exchange in the DH-PH fragment, RhoA, or their complex was initiated by diluting 10 μ L of the stock solution (\sim 0.8 mg/mL) into 90 μ L D₂O solvent, pH 7.4 at room temperature. The H/D exchange reaction was performed for varying times (10, 60, 300, 600, 1800, and 3600 s) before quenching. Quenching was accomplished using an equal volume of cold 0.1M NaH₂PO₃ buffer with 4M GuanHCl, pH 2.5. The deuterium-labeled proteins were then cleaved into reporter peptides on an immobilized pepsin column (\sim 100 μ L in bed volume). The resulting peptide mixture was desalted for 6 min in a C₈ peptide trapping column (Michrom Bioresources, Auburn, CA), then eluted by acetonitrile gradient from 15 to 40% over

15 min at flow rate of 5 μ L/min (after splitting). The eluted peptide mixture was then separated by a Vydac C_{18} column (0.3 × 50 mm, Grace Vydac, Hesperia, CA), and detected by an ion trap mass spectrometer. To minimize artifactual exchange during the analysis time, all the columns, loops, and lines were immersed in an ice bath during all the experiments.

The deuteration level for each reporter peptide was calculated by the following equation:

$$D\% = \frac{m - m_0}{(m_{100} - m_0) \times 0.9} \times 100\%$$

where m is the measured centroid mass of the deuterated peptide after a particular labeling time; m_0 and m_{100} are the two centroid mass limits of a reporter peptide from zero-deuteration and full-deuteration control experiments, respectively. The factor of 0.9 in the denominator accounts for the 90% deuterium in the labeling solution. Mass spectra were acquired by a linear ion trap mass spectrometer with a standard electrospray ionization source (LTQ, Thermo Electron, San Jose, CA). The capillary temperature was 250°C, spray voltage-4.3 kV, sheath gas flow-40 unit, capillary voltage-15 V, and tube lens voltage-105 V.

Acknowledgment

The authors thank Natalya Olekhnovich for help in the preparation of protein samples and outstanding technical assistance. This research benefited from access to the Southeast Collaboratory for High-Field Biomolecular NMR, a resource at the University of Georgia, funded by the National Institute of General Medical Sciences (GM66340) and the Georgia Research Alliance. We thank Dr. John Glushka and Dr. Yizhou Liu for assistance in setting up NMR experiments.

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