Characterization of the Binding Interaction between the Oncoprotein Gankyrin and a Grafted S6 ATPase

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Supporting Information

ABSTRACT: A complex with the C-terminal portion of the proteosomal subunit S6 ATPase is the only available structure of a protein−protein interaction involving the oncoprotein gankyrin. However, difficulties associated with recombinant expression of S6 ATPase alone, or truncations thereof, have limited our understanding of this assembly. We replaced the C-terminal portion of FtsH from Escherichia coli with the structurally homologous C-terminal portion of S6 ATPase and used this grafted protein to characterize the gankyrin−S6 ATPase binding interaction by isothermal titration calorimetry.

Overexpression of gankyrin (Figure 1a, orange) is directly linked to the onset, proliferation, and/or metastasis of breast, liver, oral, pancreatic, and colorectal cancers. In addition, gankyrin plays an essential role in Ras-initiated tumorigenesis, which is operative in ∼30% of all cancers. Protein−protein interactions (PPIs) involving gankyrin are of great interest in basic research and as therapeutic targets.

Gankyrin is reported to bind both cyclin-dependent kinase 4 (CDK4) and MDM2, resulting in increased efficiency of pRb phosphorylation and p53 polyubiquitination and degradation, respectively. However, the structural basis for these interactions has not yet been reported, and neither the targets nor their putative gankyrin binding domains are expressed as soluble recombinant proteins in Escherichia coli. A more promising venue for studying gankyrin−protein interactions is the co-crystal structure with a C-terminal portion of the S6 ATPase from the 26S proteasome, reported by Yokoyama and co-workers (Figure 1a). Preliminary characterization of the recognition interface by Yokoyama and co-workers (Figure 1a) suggested that the C-terminal portion of the S6 ATPase, the two proteins have similar tertiary structures [root-mean-square deviation of ∼1.4 Å over 74 main chain residues (Figure 1c)]. Expanding on this finding, we set out to determine if a grafted protein, in inaccessible protein.

Our initial efforts to identify such a scaffold relied on the recognition by Yokoyama and co-workers that, while the C-terminal portion of FtsH from E. coli has a low level of sequence homology (∼25%) with S6 ATPase, the two proteins have similar tertiary structures [root-mean-square deviation of ∼1.4 Å over 74 main chain residues (Figure 1c)]. Expanding on this finding, we set out to determine if a grafted protein, in

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which the C-terminal ATPase domain of FtsH is replaced with S6 ATPase, is expressed as a soluble protein in *E. coli* that mimics the native S6ATPase–gankyrin interaction.

Grafted FtsH-S6 ATPase and wild-type FtsH (wt-FtsH) were expressed as His₆-tagged proteins in *E. coli* as soluble proteins (Supporting Information, Figure S1). Circular dichroism spectra of the two proteins are virtually identical (Figure 2a), suggesting no appreciable structural change to the proteins (Supporting Information, Figure S1). Circular dichroism spectra of wild-type FtsH (wt-FtsH, top) and FtsH-S6 ATPase (bottom). (b) Co-purification of wild-type gankyrin-His₆ and FtsH-S6 ATPase mutants: wt-S6 FtsH-S6 ATPase (lane 1), R342A (lane 2), R338A/R342A (lane 3), R338A/R339A/R342A (lane 4), E356A/E357A (lane 5), D359A/D362A (lane 6), and K397E (lane 7). (c) Co-purification of wild-type S6 ATPase and gankyrin-His₆ mutants: wt-gankyrin (lane 1), R41A (lane 2), K116A (lane 3), R41A/K116A (lane 4), D39A/D71A (lane 5), and E182A (lane 6).

FtsH scaffold or grafted S6 ATPase domain. The affinity of this grafted protein for gankyrin was first assessed using a pull-down assay in *E. coli*. Binding face residues on gankyrin or FtsH-S6 ATPase were mutated to alanine, on the basis of the rationale is a lower energy of desolvation for the alanine residue compared to that of the native protein. While the molecular effects that do not dramatically lower, or completely abolish, complex stability cannot be probed using this assay. Only through the described grafting strategy are we able to create a soluble and stable mimic of S6 ATPase, which permits the use of more sensitive biophysical methods for probing this important binding interaction.

We used isothermal titration calorimetry (ITC) to obtain the full thermodynamic signature (ΔH, ΔS, and ΔG) and stoichiometry (N value) of this interaction, as well as characterize mutational effects on complex stability. Gankyrin binds the grafted FtsH-S6 ATPase with a dissociation constant (Kᵦ) of ∼67 nM (Table 1, entry 1). The observed changes in enthalpy (ΔH) and entropy (−ΔS) for this binding interaction were −28.7 and 19.0 kcal/mol, respectively. Gankyrin does not bind wild-type FtsH with any appreciable affinity (Supporting Information, Figure S2), which is unsurprising, given that the S6 and FtsH ATPase subdomains share only ∼24% sequence homology.

Alanine mutation of S6 ATPase R342, which engages gankyrin through a salt bridge with gankyrin E182, modestly lowers complex stability [Kᵦ = 216 ± 25.8 nM (Table 1, entry 2)]. Double (R338A/R342A) and triple (R338A/R339A/R342A) mutation of a positively charged patch on the S6 ATPase face, which disrupts a salt bridge between S6ATPase R342 and gankyrin E182, dramatically lowers complex stability [Kᵦ values of 2.5 ± 0.4 and 7.5 ± 0.2 μM (Table 1, entries 3 and 4, respectively)]. Interestingly, both of these mutations result in favorable binding entropies (−ΔS values of −1.5 ± 0.8 and −4.7 ± 0.1 kcal/mol, respectively, compared to a −ΔS of 19.0 ± 0.6 for the native interaction). While the molecular mechanism for this dramatic change is unclear, one possible rationale is a lower energy of desolvation for the alanine mutants, compared to that of the native protein. While Yokoyama’s original pull-down data suggest a significant role for S6 ATPase E356/E357 in complex stability, double alanine mutants

**Table 1. Analysis of Binding Interactions between Gankyrin and FtsH-S6 ATPase Proteins by ITC**

| entry | gankyrin | FtsH-S6 ATPase | Kᵦ (nM) | ΔG (kcal/mol) | ΔH (kcal/mol) | −ΔS (kcal/mol) |
|-------|----------|----------------|--------|---------------|---------------|----------------|
| 1     | wild-type| wild-type      | 67.3 ± 5.7 | −9.8 ± 0.1 | −28.7 ± 0.5 | 19.0 ± 0.6 |
| 2     | wild-type| R342A          | 216.6 ± 25.8 | −9.1 ± 0.1 | −22.0 ± 0.8 | 12.9 ± 0.8 |
| 3     | wild-type| R338A/R342A    | 2549 ± 353 | −7.6 ± 0.1 | −61 ± 0.7 | −1.5 ± 0.8 |
| 4     | wild-type| R338A/R339A/R342A | 7471 ± 301 | −7.0 ± 0.1 | −2.2 ± 0.1 | −4.7 ± 0.1 |
| 5     | wild-type| E356A/E357A    | 71.8 ± 5.9 | −9.8 ± 0.1 | −27.3 ± 0.9 | 17.5 ± 0.8 |
| 6     | wild-type| D359A/D362A    | no binding | − | − | − |
| 7     | wild-type| K397E          | 95.2 ± 12.3 | −9.7 ± 0.2 | −25.6 ± 2.5 | 15.9 ± 2.7 |
| 8     | R41A     | wild-type      | 313.3 ± 17.6 | −8.1 ± 1.2 | −17.1 ± 1.7 | 9.0 ± 2.8 |
| 9     | K116A    | wild-type      | 71.3 ± 15.5 | −9.7 ± 0.2 | −24.0 ± 0.9 | 14.3 ± 1.1 |
| 10    | D39A/D71A| wild-type      | 93.0 ± 5.6 | −9.7 ± 0.2 | −25.0 ± 0.7 | 15.4 ± 0.8 |
| 11    | R41A/K116A| wild-type   | 3633 ± 404 | −7.4 ± 0.1 | −4.9 ± 0.6 | −2.5 ± 0.7 |
| 12    | E182A    | wild-type      | 140.6 ± 9.7 | −9.4 ± 0.1 | −28.2 ± 2.1 | 18.8 ± 2.0 |

*All errors represent the standard deviation of three separate experiments. ITC conditions were as follows: 20 mM sodium phosphate, 150 mM NaCl, and 2.5 mM 2-mercaptoethanol (pH 7.4) at 25 °C.*
mutation did not appreciably lower binding affinity (Table 1, entry 5). Conversely, removal of a negatively charged patch on the S6 ATPase binding face [D359A/D362A (Table 1, entry 6)] abolished binding. An E182A mutation in gankyrin, which further probes a salt bridge with S6 ATPase residue 342, was found to modestly lower binding affinity \( K_D = 140.6 \pm 9.7 \text{ nM} \) (Table 1, entry 12), suggesting a relatively minor role of this particular salt bridge in complex stability. Residue K397 in S6 (Table 1, entry 12), further suggesting a relatively minor role of this residue (K397E) binds gankyrin with an affinity similar to that of the native protein \( K_D = 95.2 \pm 12.2 \text{ nM} \) (Table 1, entry 7), suggesting a relatively minor role of this particular salt bridge in complex stability. While gankyrin mutation K116A and double mutation D39A/D71A had minimal effects on binding affinity (Table 1, entries 9 and 10, respectively), an R41A mutation significantly decreased affinity \( K_D = 313.3 \pm 17.6 \text{ nM} \) (Table 1, entry 8). While the single K116A mutation had a minimal effect on binding, an R41A/K116A double mutation, which is designed to test the role of a larger hydrogen bond/salt bridge network, dramatically lowers affinity \( K_D = 3.6 \pm 0.4 \mu \text{M} \) (Table 1, entry 11). The R41A/K116A mutant, however, binds gankyrin with a favorable binding entropy \( \Delta S = -2.5 \pm 0.7 \text{ kcal/mol} \), possibly due to the lower energy of desolvation for the alanine mutants, compared to that of the native protein. The binding stoichiometry \( n \) for each interaction was found to be \( n \approx 1 \) \( n = 0.91 \pm 1.02 \) (Supporting Information). Reversing the titration did not appreciably alter the binding data (data not shown).

Collectively, our findings represent the first quantitative assessment of the binding interaction, and binding thermodynamics, of a physiologically relevant complex involving the oncoprotein gankyrin. These data also potentially establish a target affinity for therapeutic reagents designed to inhibit gankyrin-dependent protein–protein interactions.\(^{11,12}\)

■ ASSOCIATED CONTENT

3 Supporting Information

Experimental details, polyacrylamide gel electrophoresis gel analysis, sequences of all proteins used in this work, and representative isothermal titration calorimetry binding isotherms for each entry in Table 1. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes

The authors declare no competing financial interest.

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