Dissection of Protein–Protein Interaction and CDK4 Inhibition in the Oncogenic versus Tumor Suppressing Functions of Gankyrin and P16

Anjali Mahajan1†, Yi Guo2†, Chunhua Yuan3, Christopher M. Weghorst4 Ming-Daw Tsai1,2,3,5 and Junan Li1,4⁎

1Departments of Chemistry and Biochemistry, The Ohio State University, Columbus OH 43210, USA
2Ohio State Biochemistry Program, The Ohio State University, Columbus OH 43210, USA
3Campus Chemical Instrument Center, The Ohio State University Columbus OH 43210, USA
4Division of Environmental Health Sciences, College of Public Health, The Ohio State University, Columbus OH 43210, USA
5Genomics Research Center and Institute of Biological Chemistry, Academia Sinica Taipei, Taiwan

Received 5 April 2007; received in revised form 22 July 2007; accepted 16 August 2007
Available online 22 August 2007

Edited by M. F. Summers

Keywords: P16; gankyrin; CDK4 binding; protein/protein interactions; ankyrin repeat

Protein–protein interactions usually involve a large number of residues; thus it is difficult to elucidate functional and structural roles of specific residues located in the interface. This problem is particularly challenging for ankyrin repeat proteins (ARs), which consist of linear arrays of small repeating units and play critical roles in almost every life process via protein–protein interactions, because the residues involved are discontinuously dispersed in both the ARs and their partners. Our previous studies showed that while both specific CDK4 inhibitor p16INK4A (P16) and gankyrin bind to cyclin-dependent kinase 4 (CDK4) in similar fashion, only P16 inhibits the kinase activity of CDK4. While this could explain why P16 is a tumor suppressor and gankyrin is oncogenic, the structural basis of these contrasting properties was unknown. Here we show that a double mutant of gankyrin, L62H/I79D, inhibits the kinase activity of CDK4, similar to P16, and such CDK4-inhibitory activity is associated with the I79D but not L62H mutation. In addition, mutations at I79 and L62 bring about a moderate decrease in the stability of gankyrin. Further structural and biophysical analyses suggest that the substitution of Ile79 with Asp leads to local conformational changes in loops I–III of gankyrin. Taken together, our results allow the dissection of the “protein–protein binding” and “CDK4 inhibition” functions of P16, show that the difference between tumor suppressing and oncogenic functions of P16 and gankyrin, respectively, mainly resides in a single residue, and provide structural insight to the contrasting biological functions of the two AR proteins.

© 2007 Elsevier Ltd. All rights reserved.

Abbreviations used: AR, ankyrin repeat; BSA, bovine serum albumin; CDK4, cyclin-dependent kinase 4; D1/2, the denaturant concentration at the midpoint of transition during chemical-induced unfolding; G bead, reduced glutathione-agarose (Sigma); GdnHCl, guanidinium hydrochloride; GST, glutathione-S-transferase; HMBC, heteronuclear multiple-bond correlation; HSQC, heteronuclear single-quantum coherence; INK4, inhibitors of CDK4; IPTG, isopropyl-β-D-thiogalactopyranoside; NOE, nuclear overhauser enhancement; NOESY, NOE spectroscopy; P16, specific CDK4 inhibitor p16INK4A, also called CDKNA2 or MTS1; Rb, retinoblastoma susceptible gene product; Tm, the temperature at the midpoint of transition during thermal denaturation; WT, wild-type.

*Corresponding author. E-mail address: li.225@osu.edu.
† A.M. and Y.G. contributed equally to this work.
Introduction

Ankyrin repeat (hereafter, AR) proteins are a class of proteins in which multiple conserved repeats, namely ankyrin repeats, stack in a linear array to form a repetitive skeletal architecture with variable molecular surfaces.\(^1\) They are involved in numerous physiological processes, including cell signaling, cytoskeleton integrity, transcription, cell cycle control, apoptosis, inflammatory response, development and differentiation, and transport, through mediating protein–protein interactions. While protein interactions such as SH2 or SH3 binding to their target proteins usually involves a localized region of SH2 or SH3 and a very short motif in the target (phosphorylated Tyr and polyproline stretches, respectively), AR proteins are not known to bind to a specific motif in the binding partner.\(^3\) It is thus a very intriguing question as to how AR proteins control their functional specificity.

Specific CDK4 inhibitor p16\(^{\text{INK4A}}\) (P16) and gankyrin are two AR proteins (consisting of four and seven ARs, respectively) with opposite biological functions. P16 specifically binds to cyclin-dependent kinase 4 (CDK4) and inhibits the latter’s retinoblastoma susceptible gene product (Rb) phosphorylating activity.\(^4\) Such inhibition precludes the release of transcription factors E2Fs from incompetent Rb/E2F complexes, thus blocking the transactivation of downstream genes required for entry into the S phase. It has been well established that P16 is a tumor suppressor whose genetic inactivation through deletion, methylation, or mutation has been found in almost every type of human cancers.\(^5\) In contrast, binding of gankyrin to CDK4 renders the kinase resistant to P16 inhibition,\(^6,7\) which results in enhanced Rb-phosphorylation and stimulation of E2F transcription activity, thus effecting cell cycle progression. Gankyrin is also able to directly bind Rb and facilitate Rb phosphorylation and degradation.\(^8\) Furthermore, gankyrin interacts with MDM2 and promotes the ubiquitination and targetting of P53 to the proteasome for degradation.\(^9\) Therefore, gankyrin functions as a negative regulator of three prominent tumor suppressors, P16, Rb, and P53,\(^10,11\) and overexpression of gankyrin leads to cell transformation.\(^8\) In addition, it has been reported that gankyrin is competent in binding to S6 ATPase of the 26 S proteasome\(^6\) and MAGE A4,\(^12\) and gankyrin is overexpressed in most of human hepatocellular carcinomas.\(^13,14\) Taken together, these findings indicate that gankyrin is an oncogenic protein involved in the development of human cancers through down-regulation of CDK4, Rb, and P53.

The goal of this study was to understand the structural basis of the contrasting properties of P16 and gankyrin, and to use these studies to further understand how AR proteins control their biological specificity. As revealed in the crystal structure of P16/CDK6\(^{15}\) (a close homolog of CDK4), and other biochemical studies\(^16-18\) (Figure 1), contacts between P16 (or other INK4 proteins, in general) and CDK4 occur in discontinuous patches, and a number of residues located in both loop and helical regions of P16 contribute to CDK4 binding through electrostatic, hydrogen bonding, and van der Waals interactions.\(^3\) While mutations of most of CDK4-interacting P16 residues only lead to mild decrease in its inhibitory activity, the D84H mutant of P16 (a mutant found in human cancers), or the corresponding mutant of P18, D76A, loses all inhibitory activity even though these two mutants are still able to bind to CDK4.\(^16,18\) On the other hand, it has been reported that CDK4 R24C, a mutant frequently found in human cancers, retains its kinase activity but is resistant to P16 inhibition.\(^19,20\) Further analysis of the crystal structure of the P16/CDK6 complex indicates that there is a strong electrostatic interaction between the side-chains of D84 of P16 (D76 of P18) and R24 of CDK4 (R26 of CDK6) at the kinase active site, and mutation in either D84 of P16 or R24 of CDK4 disrupts this interaction, thus abolishing P16 inhibition.

Gankyrin competes with P16 for CDK4 binding,\(^7\) but it does not influence the kinase activity of CDK4, suggesting that gankyrin acts as a P16 blocker rather than a CDK4 inhibitor or activator. The structure of gankyrin in complex with CDK4 is not available, although the structure of free gankyrin has been solved by both NMR\(^21\) and X-ray.\(^22,23\) Our structure-based sequence homology analysis showed that the residue corresponding to D84 of P16 is I79 in gankyrin (Figure 2). Using site-directed mutagenesis, we show that the I79D mutant of gankyrin, as well as a double mutant L62H/I79D, binds to CDK4 and inhibits its kinase activity similar to P16. Further structural analyses suggest that these substitutions bring about notable local conformational changes in gankyrin, most likely facilitating the positioning of a negatively charged residue into the proximity of positively charged R24 of CDK4. Analysis of conformational stability also suggests that the CDK4-inhibitory activity likely comes at the expense of stability. Overall, our studies provide insights into the molecular mechanisms underlying interactions between AR proteins and their partners.

Results

Structure-based protein engineering

Previous studies in our, as well as other, laboratories showed that gankyrin binds to CDK4 in a way similar to that of P16 in vitro and in vivo, and this binding counteracts the CDK4-inhibitory activity of P16.\(^6,7\) Further fragmentation experiments demonstrated that the first four ARs of gankyrin are required and sufficient for CDK4 binding and P16 counteraction.\(^7\) To understand the differences between the functions of gankyrin and P16, structural and sequence alignments of the two proteins were performed as shown in Figure 2 ((a) and (b), respectively). As shown in Figure 2(b), four negatively charged CDK4-binding residues, E26, E27, D74, and D92 of P16 align perfectly with E20, E21, D70, and E87, respectively, of gankyrin thus
justifying this structure-based sequence homology analysis (Figure 2(c)). These residues are very likely involved in the binding of gankyrin with CDK4.21

One interesting notion of this analysis is that D84 of P16 corresponds to I79 in gankyrin. Apparently, the hydrophobic side-chain of I79 is incapable of interacting via a salt bridge or a hydrogen bond with the positively charged side-chain of R24 at the active site of CDK4, although a much weaker non-polar contact could not be ruled out. Therefore, this residue is an interesting candidate to explore the functional role and to see if introducing a negatively charged Asp residue in place of I79 of gankyrin could rescue its CDK4-inhibiting activity. H66 is another important residue in P16, which is not conserved between P16 and gankyrin. The crystal structure of the P16/CDK6 complex showed that this residue does not directly contact CDK4; however, mutations of this residue led to the loss of 80% of the CDK4-inhibitory activity of P16,21 suggesting that H66 contributes indirectly to inhibition, likely through stabilizing the structure of P16 and facilitating the positioning of D84. The residue corresponding to H66 in P16 is L62 in gankyrin, and it can be reasoned that introducing a L62H mutation in gankyrin may influence the stability of gankyrin but not its inhibitory ability, as long as the critical Asp is absent at position 79 of gankyrin. To confirm the above notion, two single mutants, L62H and I79D, and one double mutant, L62H/I79D of gankyrin were generated by site-directed mutagenesis, and their biochemical and biophysical properties were subsequently evaluated.

**I79D and L62H/I79D of gankyrin bind and inhibit CDK4**

The binding of these gankyrin mutants to CDK4 was first investigated using pull-down assays with
glutathione-S-transferase (GST)-tagged gankyrin proteins and the CDK4/cyclin D2 holoenzyme. Instead of CDK4, the CDK4/cyclin D2 holoenzyme was used in this assay mainly due to the fact that the holoenzyme is the biological active form and there is no interaction between gankyrin and cyclin D2. As shown in Figure 3(a), CDK4 was detected in the reaction mixtures containing GST–gankyrin wild-

![Image](image_url)

**Figure 2** (legend on next page)
type (lane 3) and mutant proteins (lanes 4–6) as well as in the reaction mixture containing GST–P16 (lane 2), which was used as positive control in this assay, suggesting that all three gankyrin mutants bind to CDK4. Even though this assay should not be used to quantitatively evaluate protein/protein interaction, the amounts of CDK4 in the pull-down products were comparable to each other, indicating that there are no significant differences among the wild-type and the mutant gankyrin proteins in their CDK4-binding affinity.

The CDK4-modulating activities of gankyrin and the three mutants were then assessed using an in vitro kinase assay as described. As shown in Figure 3(b), neither gankyrin nor the L62H mutant brought about any significant change in CDK4-mediated phosphorylation of Rb, suggesting that wild-type gankyrin and L62H mutant do not influence the activity of CDK4. However, in the reaction mixtures containing mutant I79D or L62H/I79D, as the concentration of I79D or L62H/I79D increased, CDK4-mediated phosphorylation of Rb decreased, indicating that unlike wild-type gankyrin, both I79D and L62H/I79D mutants inhibit the kinase activity of CDK4. As a positive control, increasing amounts of P16 in the reaction mixtures also led to

Figure 2. Structure-based sequence homology between P16 and gankyrin. (a) Surface charge distribution of P16 (PDB code 1DC2; left) and gankyrin (PDB code 1TR4; right) indicating the location of the two residues of interest. (b) Structure-based sequence alignment of human P16 and the first four ARs of gankyrin. Four negatively charged CDK4-binding residues, E28, E29, D74, and D92 of P16 (highlighted in green) are conserved in gankyrin as E20, E21, D70, and E87, respectively (green). Two residues of interest, H66 and D84 of P16 (gold) as well as their corresponding residues, L62 and I79 (green) in gankyrin are highlighted.
a decrease in the kinase activity of CDK4. Further quantitative analyses yielded IC_{50} values of 170(±32) nM and 120(±27) nM for I79D and L62H/I79D mutants, respectively (Figure 4), only slightly higher than that for P16 (72(±15) nM), suggesting that the CDK4-inhibitory activities of I79D and L62H/I79D are comparable to that of P16. In contrast, the IC_{50} values for wild-type gankyrin and L62H mutant were higher than 3.0 μM, indicating that gankyrin wild-type and L62H mutant do not inhibit CDK4 under the experimental conditions used. Interestingly, our previous studies showed that while mutations at D84 of P16 completely abolished its CDK4-inhibitory activity, mutations at H66 of P16 also led to an eightfold decrease in the CDK4-inhibitory activity, suggesting that H66 of P16 is able to affect the CDK4-inhibitory activity through certain ways. In contrast, the gankyrin L62H mutation does not exhibit any detectable CDK4-inhibitory activity, and the IC_{50} values for gankyrin I79D and L62H/I79D are almost identical. Taken together, an Ile → Asp substitution at position 79 of gankyrin does not significantly affect the binding of gankyrin to CDK4 but enables gankyrin to inhibit the kinase activity of CDK4. In comparison, a Leu → His mutation at position 62 apparently affects neither the binding nor the inhibition.

I79D and L62H/I79D mutations bring about substantial perturbation to the local conformation of gankyrin

2D 1H,15N heteronuclear single-quantum coherence (HSQC) NMR spectroscopy was subsequently used to investigate the possible structural perturbation caused by the gankyrin mutations described above. As shown in Figure 5(a), the spectrum of L62H is almost identical to that of wild-type gankyrin. The few notable chemical shift changes are mapped to residues Lys30, Lys35, Thr34, Thr42, Phe58, and Val64, all of which are located in the first two ARs, and thus in close vicinity of L62H mutation (Figure 6(a)). Hence, L62H mutation brings about only minor changes in local environments rather than any significant change in the global structure of gankyrin. In contrast, there are considerable and extensive changes in the spectra of I79D and L62H/I79D in comparison to the spectrum of wild-type (WT) gankyrin (Figure 5(b) and (c)). In addition, the spectra of I79D and L62H/I79D are virtually superimposable, further supporting our previous observation that the L62H mutation does not cause any significant change in the global structure of gankyrin.

On the basis of the results of 2D 1H homonuclear NOESY recorded on wild-type (WT) gankyrin, 21 a total of 168 backbone amides have been assigned, among which the residues in the AR2 and AR3 are least assigned due to the largest chemical shift perturbations. It can be concluded that while the residues in AR5-AR7 retain virtually the same chemical shifts, the residues perturbed as a result of the Ile → Asp substitution are located in the flexible loop regions within the first four ARs, including Val10, Arg35, Arg37, Ser40, Thr42, Ala43 (the first loop), Gly73, Trp74, Ser75, Ser82 (the second loop), Ala101, Val102, Gly106, and Thr108 (the third loop) (Figure 6(b)). Interestingly, all these residues are part of the TPLH interaction network of central ankyrin repeats of gankyrin. These results suggest that the conformations of loops I to III are perturbed in the I79D mutant, and raise the possibility that the observed functional change upon this mutation may not be a simple effect of a side-chain substitution. Instead, the I79D substitution possibly leads to local structural adjustments that help to position the side chain for the inhibitory function.

This concept of full integration of structure and function is further supported by some preliminary NOE analysis on the aforementioned 3D 15N-edited NOESY as well as parallel 2D 1H homonuclear NOESY recorded on WT, I79D, and L62H. Firstly, in

![Figure 4.](image-url) Quantitative measurement of the CDK4-inhibitory activities of gankyrin proteins. The in vitro kinase assays were performed as described for Figure 3(b), and the incorporation of 32P into the Rb substrate was quantitated using Image-Quant (Molecular Dynamics). IC_{50} was defined as the inhibitor concentration at which half of the maximum inhibition can be obtained. The experimental error for this assay was estimated to be ±30%. All assays were performed in triplicate, and P16 was used as a positive control.
the 3D NOESY data set relative NOE intensity changes have been observed for the limited number of residues that have been assigned in the second and third loop. For example, the NOE between H$\gamma_2$/T37 and HN/S40 is becoming much stronger whereas the sequential NOE between H$\alpha$/N67 and HN/D68 is comparatively weaker in I79D mutant. Secondly, in the downfield region of 2D NOESY recorded in H$_2$O (Figure 7), the I79D mutant shows a contrasting result compared with the L62H mutant concerning the H$\varepsilon_2$ of the histidine residue in a TPLH motif. The significant change in I79D is highlighted by the large perturbation on H$\varepsilon_2$/H78 resonance and more importantly, the disappearance of the H$\varepsilon_2$/H45 signal despite extensive search aided by 2D $^1$H-$^1$5N HSQC and 2D $^1$H-$^1$5N heteronuclear multiple-bond correlation (HMBC) experiments on a $^{15}$N-labeled I79D. Since H$\varepsilon_2$/H45 is the bridging proton of the inter-AR hydrogen bond between the H45 imidazole ring and the backbone oxygen of W74, its disappearance could be indicative of the disruption of the inter-AR interaction and the local conformational changes in both AR2 and AR3.

I79D and L62H/I79D destabilize the global structure of gankyrin

The effect of the above gankyrin mutations described above on the conformational stability of gankyrin was evaluated using far-UV circular dichroism (CD) spectroscopy. In guanidinium hydrochloride (GdnHCl)-induced unfolding, changes in the ellipticity at 222 nm, indicative of changes in the $\alpha$-helical content, were monitored. The unfolded fraction was derived from the raw data and plotted against the concentration of denaturant (Figure 8(a)). The unfolding curves of all three mutants as well as the wild type gankyrin can be fitted well to a model with a two-state transition between

Figure 5. Comparison of $^{15}$N-$^1$H HSQC spectra of wild-type gankyrin and its mutants. The HSQC spectra were obtained in $^2$H$_2$O solution using a 600 MHz spectrometer. (a) L62H (red) and wild-type (black); (b) I79D (blue) and wild-type (black); (c) L62H/I79D (green) and wild-type (black); (d) superimposition of spectra in (a), (b) and (c). L62 and I79 peaks in the spectra of wild-type gankyrin are in overlapping regions, hence, cannot be precisely pointed at in the respective mutant data. The peaks that show larger chemical shifts are labeled with arrows.
native and unfolded proteins; the resultant values of $\Delta G^\text{water}$ (the denaturation free energy in water), $D_{1/2}$ (the denaturant concentration at the midpoint of transition), and the slope $m$ are listed in Table 1. Since the $m$ values of the mutants and the wild-type gankyrin are almost identical, differences in the values of $\Delta G^\text{water}$ can be directly interpreted as their differences in conformation stability. Compared to wild-type gankyrin, both L62H and I79D mutations have moderately destabilized the conformation by 0.51 and 0.85 kcal mol$^{-1}$, respectively. The double mutant, L62H/I79D, which possesses the highest CDK4-inhibitory activity, has additional 0.34 kcal mol$^{-1}$ loss of conformational stability from I79D. The relative stability has been further confirmed in a heat-induced unfolding experiment (Figure 8(b)), in which the temperatures at the midpoint of two-state unfolding transition, $T_m$ values, were extracted (Table 1). Compared to gankyrin WT, all three mutants have decreased $T_m$ values, and the double mutant has the lowest $T_m$. In conclusion, it appears that the relative conformation stability of the three mutants coincides with the relative perturbations in HSQC, and the CDK4-inhibitory activity via mutation likely comes at the expense of stability.

Discussion

Possible conformational adjustments of local loops upon I79D mutation of gankyrin

In spite of the striking similarity between the skeletal structures of P16 and the first four ARs of gankyrin, available structures indicate that the
microenvironments around the “effector” residues, i.e. D84 and I79 for P16 and gankyrin, respectively, are different from each other. In P16, D84 is located at the beginning of the first helix of AR3 and structurally, it has more freedom than the residues in the middle of the helix. Hence, even though D84 is only slightly exposed on the surface of the free protein (Figure 2(a)), the conformationally flexible loop (loop 2) preceding D84 could influence the microenvironment around D84 and make this negatively charged residue accessible to the solvent or the positively charged side-chain of R24 at the active site of CDK4. In the crystal structure of the P16/CDK6 complex, negatively charged side-chain of D84 is in the vicinity of a positively charged environment (Figure 9). In gankyrin, I79 structurally and sequentially corresponds to D84 of P16. While I79 is located at the beginning of the first helix of AR3, its bulky, aliphatic side-chain presumably makes it less accessible to the aqueous surrounding. As shown in the docking model of gankyrin and CDK6 (Figure 9), non-polar I79 is positioned in an unfavorable electrostatic surrounding at the active site of CDK6. Additionally, the preceding H78 or other nearby residues are unable to provide any assistance in stabilizing the aliphatic side-chain of I79. Therefore, after introduction of Asp residue at position 79, the microenvironments surrounding this residue need to be adjusted in order to position the negatively charged side-chain of Asp in the proximity of the active site of CDK4 and facilitate the kinase inhibition. Such structural adjustment in gankyrin I79D mutant is supported by the substantial chemical shift changes in the HSQC spectra of gankyrin I79D and L62H/I79D mutants and the discernable changes in NOE pattern. While large chemical shift changes were observed mostly for residues located in the flexible loops connecting AR2, AR3, and AR4, significant changes were also observed for the residues constituting the TPLH network. The residues involved in the TPLH network include, for example, T42 and A43 of 42TALH45 (AR2), S75 of 75SPLH78 (AR3) and T108 of 108TPLH111 (AR4), as well as the residues preceding the TPLH sequence, such as S40, G74, W75, and G106. Most of these residues are located on the concave surface of gankyrin facing CDK4 (Figure 6). The 2D NOE SY have provided more insights about

Figure 7. 2D NOESY spectra on (a) WT, (b) I79D, and (c) L62H showing the proton downfield region. The cross-peaks designated A–C are NOEs from H^ε2/His in a TPLH motif to H^δ1 and H^ε2 of the same histidine residue, and H^α/ (His+30) residue (H^α/T207 for H177), respectively. The latter is a supporting evidence of inter-ankyrin repeat hydrogen bond between N^ε2/His and O/(His+29) (e.g. N^ε2/H177 and O/K206). It is clear from the comparisons that L62H mutant retains the TPLH features from AR2 to AR6, while I79D has large perturbation in TPLH of AR3 (H78) and ANK4 (H111), and likely disruption in 42TALH45 evidenced by the disappearance of the H^δ2/H45 signal.
the structural perturbation in TPLH motifs, 42TALH45 in particular, induced by the I79 mutation. The observation is not surprising, considering that the side-chain of I79 is pointing to AR2 and directly interacts with H45 evidenced by a NOE assigned previously between Hδ1/I79 and Hδ2/H45. It is expected that such interaction between these two side-chains help to orient the histidine imidazole ring and restrict its flexibility, enabling H45 to adopt an appropriate Nε2-H tautomeric form and engage

Figure 8. Chemical and heat-induced unfolding of gankyrin mutants monitored by far-UV CD. (a) GdnHCl-induced unfolding. Samples containing 7.5–10.0 μM proteins were incubated with different amounts of GdnHCl on ice overnight and the ellipticity at 222 nm was monitored by far-UV CD (190 nm–260 nm) at 25 °C. The fraction unfolded, defined as (the ellipticity at 222 nm at a denaturant concentration–the ellipticity at 222 nm at the native state) / (the ellipticity at 222 nm at the fully unfolded state–the ellipticity at 222 nm at the native state), was plotted against the GdnHCl concentration. The unfolding curve was fitted to a model of two-state approximation. (b) Thermal melting spectra were recorded at 222 nm by heating from 3 °C to 65 °C with a rate of 1 deg.C per minute and a 1 °C interval. Tm, the temperature at the midpoint of transition, was obtained through fitting the melting curve to a two-state transition model.

Table 1. Biochemical and biophysical parameters of gankyrin and its mutant proteins

<table>
<thead>
<tr>
<th>Protein</th>
<th>In vitro CDK4 inhibitiona IC50 (nM)</th>
<th>ΔGdwater (kcal mol⁻¹)</th>
<th>D1/2 (M)</th>
<th>m (kcal mol⁻¹ M⁻¹)</th>
<th>Heat-induced unfoldingb Tm (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gankyrin</td>
<td>&gt;3000</td>
<td>2.77</td>
<td>1.54</td>
<td>1.80</td>
<td>51.5</td>
</tr>
<tr>
<td>Gankyrin L62H</td>
<td>&gt;3000</td>
<td>2.26</td>
<td>1.38</td>
<td>1.64</td>
<td>44.5</td>
</tr>
<tr>
<td>Gankyrin I79D</td>
<td>170±32</td>
<td>1.92</td>
<td>1.14</td>
<td>1.68</td>
<td>42.0</td>
</tr>
<tr>
<td>Gankyrin L62H/I79D</td>
<td>120±27</td>
<td>1.58</td>
<td>1.01</td>
<td>1.57</td>
<td>38.5</td>
</tr>
</tbody>
</table>

a IC50 is defined as the concentration of gankyrin proteins required for 50% of the maximum inhibition. The highest concentration of gankyrin proteins tested in such an in vitro kinase assay was 3.0 μM, and an IC50 value higher than 3.0 μM indicated that at the tested concentration range of gankyrin proteins, the CDK4 kinase activity was higher than 50% of that in the absence of any gankyrin protein in the reaction mixture. The estimated error in the determination of IC50 is ±20%, and the experiments were performed in triplicate.
b ΔGdwater, D1/2, and m values were calculated according to a two-state model, and the error limit in ΔGdwater is estimated to be ±0.5 kcal/mol. The temperature at the midpoint of transition, and the error is estimated to be ±0.5 °C.
in several hydrogen bonds. A modeling structure of I79D (Figure 10) shows that the Asp side-chain is incapable of such an interaction without significant structural adjustment. Since P16 does not have the TPLH motif or its close variant, and specifically, 46RPIQ49 of P16 is aligned with 42TALH45 of gankyrin, the destabilization in TPLH motifs induced by the I79 mutation likely makes the local conformation and dynamics at the binding site mimic the counterpart in P16.

In conclusion, our results suggest a structural adjustment in I79D, especially in the loop regions, and that such adjustment may facilitate the positioning of negatively charged D79 for an electro-

Figure 9. Docking model showing the interactions of I79 of gankyrin with the active site of CDK6. (a) Docking model of the putative gankyrin/CDK6 complex. This model was constructed by superimposing the first four ARs of gankyrin (ribbon diagram, green) onto P16 (ribbon diagram, golden) in the crystal structure of the P16/CDK6 complex (PDB code 1BI7; CDK6 is shown in solid-filled diagram). Since no structure of CDK4 is available, the crystal structure of CDK6, a close homologue of CDK4, is used in this modeling. H66 and D84 of P16, as well as L62 and I79 of gankyrin are highlighted. Surface residues of CDK6 with positive and negative charges are shown in red and blue, respectively, in the solid-filled diagram. The active site of CDK6 is framed and analyzed in (b). (b) Interactions between D84 of P16 (corresponding to I79) and the active site of CDK6. P16 is in gold, and gankyrin is in green. Apparently, interactions with the charged surface of R26 in the active site of CDK6 (R24 in CDK4) depend on the negatively charged Asp of P16.
static interaction with R24 of CDK4. Moreover, this conformational adjustment was observed for the free gankyrin I79D mutant under physiological condition. Since structural analyses show that there is no significant structural change on P16 or P18 upon binding to CDK4 (data not shown), i.e. no induced-fit event happens on the AR protein part after binding to CDK4, like P16 or P18, free gankyrin I79D mutant is present in a functional conformation (in regard to CDK4 binding and inhibition).

Structural basis for the decreased stability of the gankyrin mutants

As demonstrated by our GdnHCl and heat-induced unfolding experiments, the conformation of gankyrin mutants was destabilized in the order L62H < I79D < L62H/I79D. One structural difference between P16 and gankyrin is that there is a TPLH stretch, or its variant, present in six out of the seven ankyrin repeats of gankyrin but none as highly matched TPLH variant is present in any of the four P16 ankyrin repeats. These TPLH stretches are located within neighboring loops and form an intra and inter-AR interaction network that stabilizes the global structure of gankyrin. Besides the potential electrostatic and steric disturbance caused by the above mutations, impairment to the TPLH interaction network could be the major cause for structural destabilization of gankyrin mutants. As discussed earlier, an Ile → Asp substitution at position 79 brings about considerable conformational changes in residues within/around these TPLH stretches, which perturb the TPLH interaction network and subsequently destabilize the global structure of gankyrin. As for L62H, such mutation only caused “minor” conformational changes in residues within the first two ARs (including the first TPLH stretch), and the “impairment” to the TPLH network is minor and local (Figures 6(a) and 7(c)). This can explain why L62H mutation destabilizes the conformation to a lesser extent and does not affect the function in CDK4 binding and modulation. In contrast, due to the absence of the TPLH interaction network, the corresponding H66Y mutation in P16 with marginal stability could strongly destabilize the structure, which consequently can impair its CDK4-inhibitory activity as evidenced by a loss of 80% of CDK4-inhibitory activity of this mutant.16

Structural basis for the functional diversity of the AR proteins

The residues in P16 could play three major roles: (1) maintaining the skeletal structure of P16, e.g. L63 and L64, which are conserved in almost all AR motifs;1,28 (2) CDK4 binding (but not inhibition), e.g. E28 and E29;16 and (3) inhibition (may also contribute to binding), e.g. D84. Our studies suggest that these three functions are inter-related. For example, residues in groups 1 and 2 that are responsible for generating the skeletal structure, may facilitate the positioning of D84 to R24 at the active site of CDK4, thus inhibiting its kinase activity (Figure 9). Therefore, mutations in group 1 and 2 residues could affect the inhibition by destabilizing the global structure and/or impairing CDK4 binding, whereas mutation of D84 directly abolishes the CDK4-inhibitory activity. On the other hand binding of gankyrin to CDK4 brings the extremely hydrophobic I79 residue in close proximity of R24 of CDK4 (Figure 9), which

![Figure 10](image_url)
precludes any electrostatic interaction between the side-chains of these two residues and as a result gankyrin just acts as a P16 blocker rather than a CDK4 inhibitor. However, when a negatively charged Asp residue is introduced at position 79 of gankyrin, following certain adjustments in the local environment, its electrostatic interaction with R24 of CDK4 resumes, and so does the inhibition.

In regard to the modular and repetitive nature of AR proteins, one might assume that the AR proteins are structurally very rigid. However, the results presented in this and previous reports suggest that AR proteins are highly tolerant of structural variations, which could be the basis for its functional diversity since structural and functional properties are intimately related. Some of these properties, relating to the structural pliancy of AR proteins, are summarized here. First, there are significant structural variations in the loops linking neighboring ARs among AR proteins, even though the global topology, including the small curvature, is highly conserved. Second, some AR proteins can tolerate addition or deletion of AR repeats in the middle or at the end of AR proteins. For example, our previous biochemical studies demonstrated that the removal of up to three ARs at the C terminus of gankyrin did not affect its structural topology and CDK4-binding ability. It has also been reported that deletion of individual ARs from the Notch AR domain, or insertion of various copies of ARs from the Notch AR domain or from consensus-designed ARs, did not affect chemical and heat-induced unfolding of the Notch AR domain. Last but not least, our results from the gankyrin mutants showed that when negatively charged Asp was introduced to the site originally occupied by Ile, the local conformation of the loops adjusted accordingly (accompanied by partial loss of stability), likely to help position the Asp residue for the CDK4-inhibiting function. These properties suggest that the AR proteins are structurally pliant and are capable of conformational adjustments as a result of relatively minor changes in the sequence. This, in turn, can introduce the functional diversity seen in many of the AR proteins.

**Potential biological significance**

In its natural form, gankyrin is bi-functional in regulating the CDK4-Rb pathway: it counteracts P16 inhibition of CDK4, and facilitates the ubiquitin-mediated degradation of Rb, both of which result in forcing the cells into cell cycle progression. Through structure-based protein engineering, we have transformed gankyrin from a simple competitor of P16 in CDK4 binding into a potent CDK4 inhibitor. In addition, it is plausible that for the gankyrin I79D mutant its Rb-binding and promoting function may remain intact, since the conformational adjustments upon this mutation are limited to loops I–III, and results from our previous studies showed that the three C-terminal ARs are responsible for Rb binding, and this binding is independent of CDK4 binding. Therefore, the gankyrin I79D mutant likely remains bi-functional, but the outcomes of these two functions are now in conflict: while its Rb-binding and promoting role assists cell cycle progression, its CDK4-inhibitory role acts against cell cycle progression. It will be very interesting to see which of these two opposing functions is dominant when gankyrin I79D is overexpressed in cells. Results from such studies could further our understanding of how nature balances the two functions of gankyrin, or the contrasting functions between gankyrin and P16 in relation to cell cycle and cancer.

While this paper was in preparation, the crystal structure of the complex between mouse gankyrin and the C-terminal domain of S6 ATPase of the 26 S proteasome was reported. In this complex, most of S6 ATPase-interacting residues are located in the concave surface of mouse gankyrin, especially in loops II and III, which also contribute to binding to CDK4 as demonstrated in previous studies as well as in our current study. Furthermore, mouse gankyrin binds to S6 ATPase in the presence and absence of Rb, indicating that like CDK4 binding, S6 ATPase binding is independent of Rb binding. These findings suggest that CDK4 and S6 ATPase may compete with each other for gankyrin binding in cells. From this perspective, the similarities between CDK4 and S6 ATPase in binding to gankyrin make our question about the molecular mechanisms in which gankyrin balances and coordinates its versatile functions in cells more interesting and challenging. The significance of our current study, however, goes beyond binding, it dissects the binding and inhibition functions of gankyrin toward CDK4.

**Materials and Methods**

**Cloning, expression, and purification of human gankyrin and its mutants**

As described, human gankyrin cDNA was cloned into pGEX-6p-2 vector (Amersham) and expressed in *Escherichia coli* BL21 (DE3) Codon plus cells (Novagen) as glutathione-S-transferase-fusion proteins (GST) upon IPTG induction. GST-fusion gankyrin protein was purified from the cell lysate using a reduced glutathione-agarose column (about 15 ml of G beads, Sigma). After washing with TES buffer (10 mM Tris-β-mercaptoethanol, 0.15 M NaCl (pH 7.5)), 2.0 ml of protein-bound G beads were washed with 10 ml of TES, 50 mg/ml reduced glutathione (pH 7.5) and the elute was further purified using a Q Fastflow column (Pharmacia) to get pure GST-gankyrin. To obtain gankyrin without the GST tag, the remaining protein-bound G beads (about 13 ml) were suspended in 30 ml of TES in a 50 ml tube, and 100 units of PreScission protease (2 units /μl; Amersham) were added into the tube. After incubation at 4 °C for 24 h, the G beads were re-packed on a column and the flowthrough was further purified by an S100 column (Pharmacia) equilibrated with 5 mM HEPES, 1 μM EDTA, and 1 mM DTT (pH 7.5). After SDS-PAGE analysis, fractions containing free gankyrin were pooled, concentrated, and lyophilized for further analyses.

*Binding and Modulating of CDK4 by Gankyrin*
All gankyrin mutants were generated through PCR-based Quickchange site-directed mutagenesis (Strategene), and were expressed and purified as the wild-type.

**Pull-down assay**

To investigate the binding of gankyrin proteins to CDK4, 25 μg of GST-gankyrin proteins were incubated with 10 μg of recombinant CDK4-cyclin D2 complex (see below) and were incubated at 4 °C in 250 μl of TES (pH 7.5) for 2 h. The concentrations of the holoenzyme and GST-gankyrin proteins were 0.4 μM and 2.0 μM, respectively. Subsequently, 250 μl of fresh G beads were added into the reaction mixture, and after incubation at 4 °C for 1 h, the reaction mixture was loaded onto a spin column (Fisher Scientific) and centrifuged at 4 °C, 1500 rpm for 3 min. After washing the G beads with TES five times, 1.0 ml each time, the beads were eluted with 200 μl of TES, 50 mg/ml reduced glutathione, and the elute was further analyzed with Western blot using rabbit polyclonal anti-human CDK4 antibody (Santa Cruz, C-22). In this assay, GST–P16 was used as positive control, and GST was used as negative control.

**In vitro CDK4 kinase assay**

The in vitro CDK4 activity assay was performed as described. Briefly, each reaction mixture contained 0.3 μg of recombinant CDK4/cyclin D2 holoenzyme and varying concentrations of AR proteins in 15 μl of the kinase buffer (50 mM HEPES, 10 mM MgCl2, 2.5 mM EGTA, 0.1 mM Na3VO4, 1 mM NaF, 10 mM β-glycerophosphate, 1 mM DTT, 0.2 mM AEBSF, 2.5 mg/ml of leupeptin, and 2.5 mg/ml of aprotonin). After incubation at 30 °C for 30 min, 50 ng of GST–Rh791-928 and 5 μCi [γ-32P]ATP were added in the reaction mixture which was then incubated at 30 °C for another 15 min. Proteins in the reaction mixture were separated by SDS-PAGE, and the incorporation of 32P into GST–Rh791-928 was quantitatively evaluated using a PhosphorImager (Molecular Dynamics). The concentrations of AR proteins were determined using absorbance at 280 nm in the kinase buffer and the molar extinction coefficients from ProtParam at ExPASY. The IC50 value was defined as the concentration of kinase inhibitor required for 50% of the maximal inhibition of CDK4, and measurements were repeated in triplicate.

Recombinant CDK4/cyclin D2 holoenzyme was expressed and purified as described. Briefly, human CDK4 and cyclin D2 cDNAs were cloned into pBacBak8 and pBacBak6 vectors, respectively. Of note, a His6 tag was fused to the C terminus of CDK4 to facilitate the following purification of the CDK4/cyclin D2 holoenzyme. Each construct was co-transfected into Spodoptera frugiperda SF-9 cells with Autographa california nuclear polyhedrosis virus BacPak6/BsuI-361 DNA (BD Clontech) to generate baculovirus particles. Both baculovirus particles were co-transfected into HighFive insect cells (Invitrogen), and the CDK4/cyclin D2 holoenzyme was purified through affinity chromatography using Talon resin (BD Clontech). The final product was concentrated to approximately 0.3 mg/ml in kinase buffer, and aliquots were stored at −80 °C.

**Circular dichroism (CD) analyses of gankyrin proteins**

In GdnHCl-induced unfolding, recombinant gankyrin proteins were dissolved in 20 mM sodium borate buffer (pH 7.4) containing 40 μM DTT and dialyzed against this borate buffer at 4 °C overnight. Samples containing 7.5–10.0 μM proteins were incubated with different amounts of GdnHCl (in a stock solution of 8.5 M) on ice overnight and then equilibrated at 25 °C just prior to CD analysis. The rotation at 222 nm was measured on an AVIV far-UV spectropolarimeter using a quartz microcell (Helma) of 0.1 cm light pass length, and the exact concentrations of GdnHCl were determined using its refractive index, and three scans were averaged. Here, the ellipticity at 222 nm, an indicator of the existence of α-helical secondary structure, was taken as the measure of the degree of structure present in the protein at each GdnHCl concentration, and the free energy of protein denaturation in aqueous condition was obtained on the basis of two-state approximation. To determine the protein concentration, absorbance at 280 nm was measured in the borate buffer, and the molar extinction coefficient for each protein was determined using ProtParam at ExPASY.

Heat-induced unfolding experiments were performed using 10 μM proteins in the borate buffer (pH 7.4) with 1 mM DTT and a 10 s response time. Thermal melting spectra were recorded at 222 nm by heating from 3 °C to 65 °C at the rate of 1 °C per minute and a 1 °C interval and then cooling down to 3 °C at the same rate. Tm was defined as the temperature at the midpoint of transition. The heat-induced unfolding of these gankyrin proteins was highly reversible, and for each protein, more than 90% of the CD signal was recovered after returning to the initial temperature.

**NMR analyses**

Unlabeled WT, I79D, and L62H, and uniformly 15N-labeled I79D proteins were prepared as described. The NMR samples contained 4 mM Heps, 1 mM DTT, and 5 μM EDTA in 95% H2O/5% 2H2O (pH 7.5), and about 0.4 mM protein. All NMR experiments, including 2D 1H,15N HSQC, 2D 1HHMBC, 2D 1H homonuclear NOESY, and 3D 15N-edited NOESY, were performed at 27 °C, on a Bruker DRX-600 or Bruker DRX-800 spectrometer equipped with cryo-probe. The NOE mixing time was 150 ms. Data were processed with NMRPipe and analyzed with NMRView.

**Bioinformatics analysis**

All structural modeling and comparisons were performed using DS VISUALIZER (ACCELRYS) and MOL-MOL. The molar extinction coefficient of gankyrin was calculated using ProtParam from ExPASY.

**Acknowledgements**

We thank Sandeep Kumar for critical reading of the manuscript. This work was supported by research grants from NIH (RO1 CA69472 (to M.-D. T.), R01 DE01943 and R21 DE016361 (to C. M. W.).

**References**


computer program for the visualization and analysis of NMR data. J. Biomol. NMR, 4, 603–614.