Interaction of an anticaner peptide fragment of azurin with p53 and its isolated domains studied by atomic force spectroscopy

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Abstract: p28 is a 28-amino acid peptide fragment of the cupredoxin azurin derived from Pseudomonas aeruginosa that preferentially penetrates cancerous cells and arrests their proliferation in vitro and in vivo. Its antitumor activity reportedly arises from post-translational stabilization of the tumor suppressor p53 normally downregulated by the binding of several ubiquitin ligases. This would require p28 to specifically bind to p53 to inhibit specific ligases from initiating proteosome-mediated degradation. In this study, atomic force spectroscopy, a nanotechnological approach, was used to investigate the interaction of p28 with full-length p53 and its isolated domains at the single molecule level. Analysis of the unbinding forces and the dissociation rate constant suggest that p28 forms a stable complex with the DNA-binding domain of p53, inhibiting the binding of ubiquitin ligases other than Mdm2 to reduce proteasomal degradation of p53.

Keywords: AFS, cancer physics, unbinding forces

Introduction
The p53 protein is a transcription factor that acts as a tumor suppressor by playing an essential role in preventing inappropriate cell proliferation and in maintaining genome integrity. It is stabilized by post-translational modifications in response to different stress signals, increasing its intracellular levels and activating the transcription of downstream target genes that regulate cell-cycle arrest, DNA repair, and apoptosis.1,2 p53 is a 393-residue protein with three functional domains: an N-terminal domain (NTD, aa 1–93), a core DNA-binding domain (DBD, aa 102–292), and a C-terminal domain (CTD, aa 293–393) that are responsible for its transcriptional activation, DNA-binding, and tetramerization functions, respectively.3,4 The activity of p53 is tightly regulated through post-translational modification, localization, and degradation.5 Its major negative regulator is the Mdm2 oncoprotein that inhibits its transcriptional activity,6 favors its nuclear export,7 and acts as an E3 ubiquitin ligase, targeting p53 for proteasomal degradation.8 The regulatory function of Mdm2 is exerted through formation of a complex with p53.6,9

The central role of p53 in safeguarding the genome integrity provides an attractive target for anticancer drugs that can stabilize it, interfere with its downregulatory pathway,10,11 and enhance its tumor-suppressor function in cancer cells. One such agent, azurin, a copper-containing, electron-transfer protein secreted by the opportunistic pathogen Pseudomonas aeruginosa, has significant anticancer activity in vitro12–15 and in vivo.12,14 The antiproliferative activity of azurin is based on the stabilization and subsequent increase in intracellular concentration of p53.12–15 A single-molecule
atomic force spectroscopy (AFS) experiment has shown that a stable complex is formed between full-length p53 and azurin. However, biological as well as computational studies have reported that azurin binds to either the NTD of p53 or its DBD. Site-direct mutagenesis and computational investigations have revealed that the two methionines, located at positions 44 and 64 within a hydrophobic patch of azurin are crucial for the interaction with p53.

Although azurin preferentially enters cancer relative to normal cells, a 128-amino acid protein could display some immunogenicity, a potentially significant side effect, that compromises its pharmaceutical efficacy. Since peptide fragments of azurin could provide therapeutic molecules with the same cytotoxicity, delivery, and target specificity of the whole protein, but with potentially fewer side effects, truncated versions of azurin have been investigated for anticancer activity. Amino acids 50–77 of azurin form a 2.9 kDa peptide fragment that encompasses the azurin α-helix and retains the preferential penetration of the whole protein, but also its antitumor activity in vitro and in vivo. The antiproliferative activity appears to result from aa residues 11–18 of p28 (61–69 of azurin) binding to p53 in a region that does not inhibit the binding of Mdm2 or subsequent ubiquitination. Since the molecular details and the kinetics of its interaction with p53 and, more importantly, with what domain(s) have not yet been clarified, a detailed study of the p28–p53 interaction could provide significant information on p28 action at the molecular level.

In the present work, the kinetic properties of the interaction of p28 with the full-length p53 and its DBD and NTD at a single-molecule level were investigated by AFS, while the interaction with the CTD has been excluded on the basis of competition assays results. AFS is an innovative nanotechnology, suitable for measuring intermolecular forces down to the piconewton range, at single-molecule resolution, in near-physiological conditions without any labeling and using an extremely low amount of substances. As such, it provided detailed information on the interaction strength and kinetics of these biomolecular partners, complementing traditional biochemical and molecular approaches. The kinetic parameters determined from force spectroscopy experiments suggest a bio-recognition process occurs between p28 and the DBD of p53, while almost no interaction is registered with the NTD. This strongly suggests that p28 undergoes a specific interaction only with the DBD of p53 and provides a first look at the kinetics of the reaction that underlies the anticancer activity of p28.

### Methods

#### Protein expression and purification

The human p53 gene encoding aa 1–93 was amplified by polymerase chain reaction (PCR) with pUC18p53, kindly provided by Professor Nobuo Tsuchida, Department of Molecular Cellular Oncology and Microbiology, Tokyo, Medical and Dental University. Primer set used was 5'-AAA GGG GGA TCC ATG GAG GAG CCG CAT CAG ATC CT-3' and 5'-AAA AGG GAA TTC TCA CAG GGG CCA GGA GGG GGA TCC ATG GAG GAG CCG CAT CAG ATC CT-3' (Genencor, Firenze, Italy). 10 pmol of each primer was added to 20 ng of template DNA and 12.5 mL of 23 PCR HotStartTaq™ mix (Qiagen, Hilden, Germany) in a final volume of 25 mL. The amplified fragment was digested by Bam HI and Eco RI (Invitrogen, Carlsbad, CA) and ligated into pGEX-2T vector in frame with the glutathione S-transferase (GST) gene sequence. The fusion protein, expressed in Escherichia coli BL21 cells transformed with the pGEX-2T (1–93) plasmid, was purified as previously described.

Also, purification of full-length p53 (aa 1–393) and its DBD (aa 94–288) followed a standard GST method. Briefly, E. coli BL21 DE3 having pGEX-4T-ntp53 or pGEX-4T-DBP-p53 plasmid DNA were grown at 37°C to optical density measured at a wavelength of 600 nm of 0.4. Bacteria were incubated for an additional 3 hours with 0.5 mM isopropyl-1-thio-β-galactopyranoside under vigorous shaking. Cells were lysed in phosphate-buffered saline (PBS) containing 0.1% Triton X-100, 1 mM dithiothreitol (DTT), and protease inhibitors. After centrifugation, supernatant fractions were incubated with glutathione-Sepharose™ beads (G4510; Sigma, St Louis, MO). GST-fusion proteins were eluted with 50 mM Tris-HCl pH 8.0 containing 10 mM glutathione (G4251; Sigma) and 1 mM DTT, and incubated with either thrombin or Factor Xa protease (Amersham Biosciences, Piscataway, NJ). After incubation at room temperature for 16 hours, thrombin or Factor Xa protease was removed by P-aminobenzamidine-agarose or anti-factor Xa-agarose beads (Sigma). GST-free p53 and its derivatives were dialyzed against storage buffer (50 mM Tris-HCl pH 7.5, 1 mM MgCl₂, 0.5 mM DTT). The quality, correct folding, and purity of recombinant proteins were subsequently verified.

#### Tip functionalization

The 28-amino acid (Leu⁵⁰-Asp⁷⁷, 2.9 kDa) fragment of azurin was anchored to the AFS silicon nitride cantilever (Veeco Instruments, Santa Barbara, CA) by means of a cysteine residue conjugated to the NH₂-terminus to create...
p28 (Cys-p28, 3.0 kDa). The tips were cleaned in acetone for 10 minutes, dried with a stream of nitrogen, and ultraviolet (UV) irradiated for 30 minutes to expose hydroxyl groups. Tips were then immersed in a solution of 2% (v/v) 3-aminopropyl-triethoxysilane (APTES) (Acros Organics, Geel, Belgium) in chloroform, incubated for 2 hours at room temperature, rinsed in three changes of chloroform, and dried with nitrogen. Silanized tips were immersed in 1 mM N-hydroxysuccinimide-polyethylene glycol-maleimide (NHS-PEG-MAL), molecular weight 1395 Da, 9.5 nm in length (Thermo Scientific Inc, Waltham, MA) and dissolved in dimethylsulfoxide (DMSO) for 3 hours at room temperature. The PEG spacer contains an NHS-ester group at one end, which reacts with amino-silane molecule to form an amide bond and a maleimide group at the other end, which reacts with the sulphydryl group of cysteine residue linked to the NH2-terminus of p28. Tips were then washed in three changes of DMSO to remove the unbond linker, rinsed with Milli-Q® (Millipore, Bellerica, MA) water, dried with nitrogen, then incubated with 50 µL of a 10 µM solution of p28 in 50 mM PBS pH 7.5 overnight at 4°C. The tips were then gently rinsed and stored in buffer at 4°C. A schematic representation of tip functionalization is shown in Figure 1A.

Preparation of the protein substrates
The full-length p53, its DBD, and its NTD, were individually immobilized on glass slides previously cleaned for 5 minutes in acetone, dried under a stream of nitrogen and then UV irradiated for 30 minutes. After immersion in 0.3 M APTES in chloroform and incubation for 3 minutes at room temperature, they were rinsed in three changes of chloroform and dried with nitrogen. The glass slides were subsequently incubated with a solution of 1% glutaraldehyde (Sigma-Aldrich, St Louis, MI) in Milli-Q water for 10 minutes at room temperature, rinsed carefully with Milli-Q water, and dried with nitrogen. Fifty µL of a 0.8 µM p53, p53 DBD or p53 NTD in 50 mM PBS pH 7.5 were poured onto this amine-reactive surface, incubated overnight at 4°C, gently washed with PBS, and stored in buffer at 4°C. A schematic representation of the substrate functionalization is shown in Figure 1B.

Atomic force microscopy (AFM) imaging and force spectroscopy
A Nanoscope IIIa/Multimode atomic force microscope (Digital Instruments, Santa Barbara, CA) was used to perform force spectroscopy. Imaging of full-length p53, p53 DBD, and p53 NTD substrates were imaged by tapping mode AFM, with an amplitude set point corresponding to the 95% of the free amplitude value. The cantilever nominal spring constant, $k_{nom}$, was 0.5 N/m. The substrate was scratched to get a qualitative indication about the protein monolayer height as described by Funari et al.

Force measurements were carried out in PBS buffer (50 mM K$_3$PO$_4$, 150 mM NaCl, pH 7.5) using force calibration mode AFM. The cantilevers used to perform force spectroscopy studies had a nominal spring constant, $k_{nom}$, of 0.01 N/m. The effective spring constant, $k_{eff}$, was determined by the procedure reported by Hutter and Bechhoefer. A relative trigger of 50 nm was applied to limit the maximum contact force applied by the tip on the protein.

**Figure 1** Immobilization strategies of peptide and proteins to the tip or substrate. (A) p28 is bound to the amino-silanized tip via a PEG crosslinker. (B) p53, or alternatively its DBD or NTD, is immobilized on glass slides via a chemical platform involving sequentially linked amino-silane and glutaraldehyde, thus targeting aminic groups of lysine residues exposed on the protein surfaces (see Materials and methods section for details).

**Abbreviations:** APTES, 3-aminopropyl-triethoxysilane; DBD, DNA-binding domain; MAL, maleimide; NTD, N-terminal domain; PEG, polyethylene glycol.
monolayer to 0.5 nN. A ramp size of 150 nm was set, and an encounter time of 100 ms established. The approach velocity was set at a value of 50 nm/s, with the retraction velocity increased from 50 to 4190 nm/s. Loading rate, defined as \( r = \frac{dF}{dt} \), was determined according to the relationship \( r = \frac{dF}{dt} = kv \), where \( k \) is the cantilever spring constant. Correspondingly, the nominal loading rate, obtained by using the nominal spring constant, \( k_{nom} \), was found in the range 0.5–42 nN/s. The effective loading rate was determined by replacing the nominal cantilever constant with the effective one, \( k_{sys} \), to take into account that molecules (ie, proteins and/or linkers) tied to an AFM tip make the cantilever spring constant change. The \( k_{sys} \) values were obtained from the slope of each retraction curve immediately prior to the unbinding event.29–31 All blocking experiments were conducted at the nominal loading rate of 7 nN/s.

Results
Interaction of p28 with full-length p53
All experiments were performed using AFS, a single-molecule technique that complements traditional proteomic and molecular biology approaches for the functional analysis of biorecognition events.25,32 In a typical AFS experiment, a ligand is anchored to the AFS tip while the receptor is immobilized onto a substrate. Force–distance curves are thus performed cyclically on the tethered system, as sketched in Figure 2 and described in the legend. Briefly, the tip is approached to the substrate until the ligand and the receptor are brought into close proximity and can interact to form a complex. The cantilever then retracts from the surface, and when the force that it exerts overcomes the stability of the complex, a sudden jump in the deflection occurs as a consequence of the complex dissociation.

Hence, one key requisite to investigate ligand–receptor interaction by AFS is a robust attachment of ligand and receptor on their respective AFM supports, preferably through covalent bonds, which are generally stronger than those characterizing protein–protein interaction.25,33 Strong immobilization of the two partners ensures the stability of the tethered system over time allowing repeated approach/retraction cycles. In this respect, the NH₂-terminal of p28 is conjugated to a cysteine residue whose SH group reacts with the MAL group of a flexible 9.5 nm-long PEG polymer covalently linking the p28 peptide to the AFS tip (Figure 1A). The linker increases the flexibility and the re-orientation freedom of the peptide when the AFM tip approaches the protein monolayer. This favors the biorecognition process.34 Moreover, during the tip-retraction, the linker undergoes a stretching process whose unique features assist in discriminating specific and nonspecific unbinding events.25,33

Full-length p53 molecules are immobilized on a glass slide by targeting the amine groups of the lysine residues exposed on the protein surface (Figure 1B). The presence of several lysine residues available for the reaction is susceptible to generate statistically random orientations of the proteins on the substrate. This immobilization strategy could generate some configurations that might result unfavorably for the bio-recognition event (see below).

Before proceeding with AFS measurements, the morphology of p53 proteins immobilized on the glass slides were analyzed by imaging AFM using a bare tip. The presence of a homogenous layer of p53 with discernible single proteins was observed. To verify the presence of a monolayer and define its height, a bare tip was thus used to scratch the substrate, working in contact mode as described in Funari et al27 (results not shown).

With a functionalized tip thousands of force curves were thus recorded on such a substrate with an intermediate degree of coverage (condition also suggested by Gilbert et al),35 at several distinct points in which a corresponding percentage of specific events were observed. The approach speed was kept constant while the retraction one, \( v \), was varied from 50 to 4190 nm/s.
As collected force curves can significantly differ in AFS unbinding experiments in the single molecule regime,25,32 force curves were accepted whose retraction portion, before the jump-off, exhibited a nonlinear trend starting and ending at the zero-deflection line where the nonlinear trend is attributed to the viscoelastic properties of the PEG linker under stretching.26,37 When multiple jumps were observed, which could be due to subsequent rupture of the complex bonds, the curves were accepted if the last jump started and ended at zero deflection, with the last jump taken as representative of the unbinding process.

The unbinding frequency, defined as the ratio between the number of accepted unbinding events and total number of the collected force curves, was found to be about 20% at a loading rate of 7 nN/s. This frequency could potentially be affected by the presence of unfavorable binding geometries and by steric hindrance.16,27,38 However, studying the frequency of unbinding as a function of the surface coverage, no significant change was found. The relatively low unbinding frequency, detected even when specific interaction occurs, was thus to be ascribed to the random orientation resulting from the immobilization strategy. Interestingly, the unbinding frequency, observed to be consistent with values previously reported for other biological interactions, was dependent on the loading rate, initially increasing, reaching a maximum, and decreasing as previously described.27

As the unbinding force for each curve is the product of cantilever deflection (d, Figure 2) and its effective spring constant ($k_{eff}$), force histograms correspond to the different loading rates at which force spectroscopy measurements were conducted. Working on the chosen intermediate substrate coverages, the histograms of the unbinding forces, ie, at a loading rate of 7 nN/s (Figure 3A), exhibit essentially a single mode distribution, slightly skewed toward high force values. The asymmetric shape, similar to that observed in other systems, could be due to some factors such as the heterogeneity in chemical bonds or spacer length, or a residual occurrence of a few multiple binding events.39,41,42 On this basis, in this present study, the most probable unbinding force was determined from the maximum of the main peak of each histogram. At a loading rate of 7 nN/s, an unbinding force value of 82 pN was found (Figure 3A), which falls within the range reported for other specific biological interactions at similar loading rates.43 Also found was that the unbinding force values and the widths of their corresponding distributions increased with an increase in loading rate, as reported previously.16,27,44

Blocking experiments on the p28/p53 complex demonstrated that the observed unbinding events arise from a specific recognition process. Indeed, after incubation of the p53-functionalized substrate with 30 µM free p28, the unbinding frequency was reduced about 55%, indicating formation of a p28/p53 complex was specific. Upon increasing the concentration of the blocking agent, a higher attenuation was not registered. The persistence of residual unbinding activity after blocking has also been reported for other force spectroscopy experiments and could be related to the forced interaction between the two partners, induced by the experimental design.45 Importantly, the similarity in force histograms before and after blocking (see Figure 3A) indicates the corresponding interactions may be similar. The specificity of the p28/p53 interaction was further verified by distribution analysis of the linker extension from zero distance to the position of the complex rupture.
in a force-distance curve. The unbinding length distribution exhibited a single-mode distribution centered at 12.0 nm with a standard deviation of 5.6 nm (Figure 3B), which is in good agreement with the extension expected from the stretching of the 9.5 nm long PEG during the unbinding process.\(^{45}\) The unbinding process which the p28/p53 complex undergoes under the influence of an external loading force can be treated within the theoretical context of the Bell-Evans model.\(^{46,47}\) According to this model, the application of an external force \(F\) modifies the energy profile of the unbinding process, lowering the activation energy barrier, resulting in an exponential increase of the dissociation rate constant \(k_{off}(F)\) with the applied force

\[
  k_{off}(F) = k_{off} \cdot \exp\left(F x_\beta / k_B T \right) \quad (1)
\]

where \(k_{off}(F)\) and \(k_{off}\) are the dissociation rate constant in the presence of and without any applied force, respectively, \(x_\beta\) is the width of the potential barrier along the direction of the applied force, \(k_B\) is Boltzmann’s constant, and \(T\) is the absolute temperature.

Assuming that the applied force \(F\) increases linearly with a constant loading rate \(r\), the most probable unbinding force, \(F^*\), at a fixed value of loading rate is given by:

\[
  F^* = k_B T x_\beta \cdot \ln\left(r x_\beta / (k_{off} k_B T)\right) \quad (2)
\]

This expression predicts a linear relationship between the most probable unbinding force, \(F^*\), and the natural logarithm of the effective loading rate \(r\). \(F^*\) has been determined from the maximum of the main peak of each unbinding force histogram, and \(r\) is given by the product between the retraction velocity and the spring constant of the entire system, \(k_{syst}\), determined from the slope of the force versus distance for each retraction curve, as described in the Methods section.

Therefore, by plotting \(F^*\) versus the natural logarithm of the effective loading rate \(r\), the kinetic parameters \(k_{off}\) and \(x_\beta\) can be obtained from the slope and intercept of a linear fit. Figure 4 shows the dynamic force spectrum obtained by fitting the plot of \(F^*\) versus log \(r\) with Equation 2. The spectrum shows, in the range of loading rates under consideration, a single regime indicative of a single energy barrier and unique transition state of the reaction and provided values of 0.47 ± 0.02 nm for \(x_\beta\) and 0.13 ± 0.03 s\(^{-1}\) for \(k_{off}\).

The association rate constant \((k_{on}\) for the p53/p28 complex has been estimated from the expression \(k_{on} = N_A V_{eff}/t_{0.5}\), where \(N_A\) is the Avogadro’s number, \(V_{eff}\) is the effective volume of a half-sphere with radius \(r_{eff}\) around the tip, and \(t_{0.5}\) is the time for the half-maximal binding probability, given by \(t_{0.5} = 2 r_{eff} / \nu\), where \(\nu\) is the approach speed of the cantilever.\(^{16,45}\) To estimate \(t_{0.5}\), the interaction time between the proteins was varied during the force distance cycles, and an exponential increase was observed in the unbinding frequency with contact time, until a plateau was reached. A \(t_{0.5}\) of 0.05 seconds was found, and then, by assuming a \(r_{eff}\) of 4 nm, a \(k_{on}\) of 1.8 × 10\(^6\) M\(^{-1}\) s\(^{-1}\) was obtained. The dissociation constant \((K_D = k_{off}/k_{on})\) for the complex between p53 and p28 was about 7 × 10\(^{-6}\) M. Interestingly, \(K_D\) is located in an “affinity region” between the transient azurin/cytochrome c\(_{551}\) complex \((K_D\) in the range of 10\(^{-5}\)–10\(^{-6}\) M) and that reported for antigen/antibody pairs \((K_D\) in the order of 10\(^{-7}\)–10\(^{-11}\) M).\(^{38}\)

**Interaction of p28 with the p53 DBD and NTD**

The single-molecule interaction of the p28 peptide fragment of azurin with the DBD and NTD of p53 was studied. The experiments (substrate coverage, immobilization strategies, and sampling procedures) were conceived in order that the AFS results could be comparable for the different proteins immobilized on the substrate on a statistical basis. As already mentioned, the interaction with the p53 CTD has not been investigated, since competition assays with p53 site-specific antibodies demonstrate that p28 has no contact with p53 CTD aa residues.\(^{23}\)

Concerning the p28/DBD interaction, at a loading rate of 7 nN/s, an unbinding frequency of about 25% was obtained. The most probable unbinding force, extracted from the maximum of the main peak of the unbinding forces histogram (Figure 5A), was 95 pN. The unbinding
frequency and unbinding force values are slightly higher than those corresponding to the p28/p53 interaction. Blocking by incubation of the DBD-functionalized substrate with 30 µM free p28 produced a 60% reduction in the unbinding frequency (Figure 5A), indicative of the specificity of the p28/DBD complex. The force histograms, before and after blocking, also show a similar distribution. The specificity of the p28/DBD interaction is also observed in the unbinding lengths, which show a single-mode distribution centered at 12.1 nm and with a standard deviation of 3.7 nm (Figure 5B).

Measurement of highly probable unbinding forces over the range of five loading rates investigated (from 0.5 to 42.0 nN/s) and application of Equation 2 led to the linear plot shown in Figure 6 from which an $x_p$ of $0.46 \pm 0.05$ nm and a $k_{off}$ of $0.012 \pm 0.006$ s$^{-1}$ were extracted. While the $x_p$ value is essentially comparable to that calculated for the p28/p53 complex, the $k_{off}$ value is, remarkably, one order of magnitude lower and displays a much higher spread. On the other hand, the association rate constant, $k_{on}$, for the p28/DBD complex, measured as previously described, is $1.9 \times 10^4$ M$^{-1}$ s$^{-1}$, close to that of the p28/p53 interaction. The $k_{on}$ and $k_{off}$ values yielded for the p28/DBD complex a $K_D$ of $6.3 \times 10^{-7}$ M.

The same experimental approach used to study the p28/p53 and p28/DBD interaction, was used to investigate the possible interaction of p28 with the NTD of p53. p28 was anchored to the AFS silicon nitride tip, while the NTD was immobilized on a glass slide (Figure 1). The corresponding force curves were recorded maintaining a constant forward velocity and varying the retraction speed from 50 to 4190 nm/s. At a loading rate of 7 nN/s, there was minimal number of unbinding events which yielded a negligible unbinding frequency. Since the observed events were not statistically significant, it is reasonable to assert that there is no specific interaction between p28 and the NTD.

**Discussion**

The specific interaction of the azurin-derived peptide, p28, with full-length p53, its DBD, and its NTD was investigated for the first time at a single-molecule level by means of AFS. The technique enabled the strength and kinetics of complexes to be probed under the application of an external force, under nearly native conditions, and without any labeling. The AFS approach revealed that a specific interaction occurs between p28 and p53, leading to the formation of a stable complex. Interestingly, the unbinding force, dissociation constant, and kinetic parameters of the complex are comparable to those
measured by AFS for the p53 interaction with azurin, to which p28 belongs.16

The interaction of p28 with full-length p53 appears confined to the DBD core domain through formation of a stable complex. Interestingly, it was found that the lifetime \( \tau (= 1/k_{\text{on}}) \) of the DBD/p28 complex is ten times longer than that of the p28–p53 association, probably as a consequence of the DBD tendency to form aggregates\(^\text{48} \) that could wrap p28 and exert a steric hindrance to its exit from the complex. The possible DBD aggregation, however, seems not to affect the association phase of the interaction as the similar \( k_{\text{on}} \) calculated for the DBD-p28 and p53-p28 complexes suggest.

The finding of a specific and stable p28–DBD interaction points out more precisely which region of full-length p53 is involved in the interaction with p28, with respect to what has been previously reported in in-vitro studies.\(^23\) In addition, the results are in line with those of a computational docking study predicting that p28 and the DBD of p53 undergo a molecular association characterized by low, negative binding free energy,\(^\text{49} \) high shape complementarity, and several hydrogen bonds at the interface. Indeed, in that study the best predicted complex between DBD and p28 has been shown to involve the DBD aa residues 96–103, just close to its S1 strand and, at least in part, its L2 loop (aa 164–170).

The occurrence of a stable complex between the DBD and p28 suggests that the p28 anticancer activity may be related to its ability to inhibit the binding of E3 ligases, other than Mdm2, ie COP1, Pirh2, and perhaps TOPORS and ARF-BP1\(^\text{50–52} \) to the DBD, and reduce the proteasomal degradation of p53. In this respect, competitive assays of the DBD–p28 interaction by using these ligases, would provide significant insight into the p28 mechanism of action.

The DBD is not only involved in the control of the p53 downregulation, but it is above all, the domain necessary for the binding of the tumor suppressor to DNA. Since mutations within the DBD are often connected with p53 loss of function and subsequent tumor proliferation, it could be interesting to investigate the possibility that p28 could interact with mutated forms of DBD and enhance apoptosis. Here, p28 could potentially produce a conformational reversion of mutant p53, reactivating its wild-type function.

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Disclosure

The authors report no conflicts of interest in this work.

References


