The p53 tetramer shows an induced-fit interaction of the C-terminal domain with the DNA-binding domain

Original Article

The Trp53 gene is the most frequently mutated gene in all human cancers. P53 is a powerful transcription factor that can activate different biochemical pathways and affect the regulation of metabolism, senescence, DNA damage response, cell cycle and cell death. The understanding of its function at the molecular level could be of pivotal relevance for therapy. Investigation of long-range intra- and interdomain communications in the p53 tetramer–DNA complex was performed by means of a structural model that included the tetramerization helices in the C-terminal domain, the DNA-binding domains and a consensus DNA-binding site of 18 base pairs. Nonsymmetric dynamics are illustrated in the four DNA-binding domains, with loop L1 switching from inward to outward conformations with respect to the DNA major groove. Direct intra- and intermonomeric long-range communications between the tetramerization and DNA-binding domains are noted. These long-distance conformational changes link the C terminus with the DNA-binding domain and provide a biophysical rationale for the reported functional regulation of the p53 C-terminal region. A fine characterization of the DNA deformation caused by p53 binding is obtained, with ‘static’ deformations always present and measured by the slide parameter in the central thymine–adenine base pairs; we also detect ‘dynamic’ deformations switched on and off by particular p53 tetrameric conformations and measured by the roll and twist parameters in the same base pairs. These different conformations can indeed modulate the electrostatic potential isosurfaces of the whole p53–DNA complex. These results provide a molecular/biophysical understanding of the evident role of the C terminus in post-translational modification that regulates the transcriptional function of p53. Furthermore, the unstructured C terminus is able to facilitate contacts between the core DNA-binding domains of the tetramer.

INTRODUCTION

The Trp53 gene is the most frequently mutated gene in all human cancers. P53 is a powerful transcription factor that is able to activate different biochemical pathways that affect the regulation of cell death and the cell cycle. The quaternary structure of p53 in its tetrameric form was reconstructed using a combination of NMR, small-angle X-ray scattering, electron microscopy and FRET (fluorescence resonance energy transfer) and showed that an open cross-shaped structure was formed in the absence of DNA, with loosely coupled dimers interacting via the DNA-binding domain (DBD: residues 102–292). In contrast, the structure rigidifies upon DNA binding and becomes more compact. This multitechnique experimental approach showed that the folded core domains interacted with a DNA response element and the tetramerization domains (TETs: residues 323–356) in the C-terminal domain, resulting in conformations that are compatible with those previously described based on experimental data from the separate fragments. All of the remaining portions of the protein (i.e., the linkers connecting DBD with TET, the whole N-terminal domain and the 30 residues at the C terminus) are intrinsically disordered according to the current structural information.

There is an absence of information on the additional components of the transcription machinery; therefore, a molecular dynamics (MD) investigation could allow an incremental advance in understanding the possible intermolecular interaction at the tetramer level.

We previously described the MD of the full-length p53 monomer, which showed a very stable structure for up to 850 ns in the DBD but significant flexibility both in the N-terminal transactivation domains (TAD1 and TAD2) and the C-terminal region, including the tetramerization domain. This flexibility disclosed several hydrogen bonds between the N and C termini or between them and the DBD. Clearly, these interactions may change in the tetramer, and therefore we have undertaken a subsequent investigation herein to analyse the interactions at the level of the p53 tetramer.

As noted by Lubin et al., destabilizing mutations cause the DBD to misfold only when it is part of the tetramer, but not when it is in its monomeric state. The building of an atomic model and its conformational sampling by state-of-the-art MD simulations can furnish structural information not obtainable by any experimental technique and help unravel the biological mechanism of p53 oncogenic mutants. In silico experiments on truncated forms of p53 have already proven their usefulness.

RESULTS

We built a detailed atomistic model for residues 92–360 of p53 in its tetrameric form and in interaction with a consensus
Figure 1. (a–c) The 3D structure model of p53 is shown here in its tetramer form bound to a full DNA consensus site extracted from the MD simulation. The four monomers are coloured in black, red, green and blue. The 18 bp DNA structure is coloured in purple. (d) Per-residue root-mean-square fluctuations (RMSFs) for each p53 monomer (colours as in a–c). Arrows highlight the residues belonging to the DBD, the tetramerization domain (TET) and the tetramerization helix (TE). As previously described for full-length p53, the DBD region is significantly more rigid, with relatively low RMSF values. (e) Secondary structure as a function of simulation time. The four monomers (M1–M4) show a very stable structure over time in the DBD and TE, whereas specific regions are able to switch between α-helix (blue), turn (yellow) and bend (green).
DNA-binding site that was 18 base pairs in length (Figures 1a–c). To the best of our knowledge, this is the only model of p53 in its tetramer form that includes the tetramerization helices and Zn-binding ions. Our previous study of full-length p53 showed a key role for the Zn-coordinated helix H1 at the centre of the long-range interactions with: (1) the N-terminal domain through His178; (2) the REG region (residues 363–393) in the C-terminal domain through His179; and the β-sandwich and the loop–sheet–helix regions in DBD through the Zn-coordinated cysteine residues.13 In this work, we focus our attention on these last interactions in the p53–DNA complex.

Structural stability of the p53 tetramer

After an initial equilibration where weak restraints were applied to the DNA and tetramerization helices, four independent all-atom MD simulations were performed that totalled 100 ns. The structure and dynamics of the tetramer are characterized for three different regions: DBDs, TETs and the loop connecting them (residues 293–322). As expected, the per-residue root-mean-square fluctuation shows that the loops are the more flexible regions of the protein complex (Figure 1d), whereas the DBDs and TETs remain quite rigid. Moreover, inspection of their secondary structure contents along the trajectories (Figure 1e) confirms this behaviour. Per-residue root-mean-square fluctuation and secondary structure analysis indicate that while the overall structure is well conserved in the four monomers, subtle dynamic differences are observed, particularly in monomer 4 (M4) compared with the others. For example, the region of maximum flexibility in this monomer is not located around residues 312–330 as it is for the other monomers but in residues 295–305. Both of these intervals are located at the N termini of the TEs (tetramerization helices; residues 335–355). Moreover, an α-helix propensity for residues 175–180 is shown by M4 alone during the 100 ns timeframe (Figure 1e). Note that two of the four Zn-binding residues are present in these ranges (i.e., Cys176 in loop 2 and His179 in the H1 helix) (Supplementary Figure S1).

Long-range interdomain communications

Owing to the subtle thermodynamic regulation of the p53 tetramer that has also been associated with a mechanism that is able to prevent cancer-induced misfolding in p53 oncogenic mutants,14 we investigated long-range coupling dynamics to highlight how the domains were (dynamically) coupled and describe the key regions characterizing the overall tetramer motion. To extract these biologically relevant motions, we calculated the Dynamic Cross Correlation (DCC) Map and performed Essential Dynamic (ED) analyses.19

The DCC Map for the p53 tetramer is shown in Figure 2a. This analysis furnishes an overall picture of the correlated motions that occur between protein residues during the simulation. Highly positive peaks of the elements of the map (Cij) are indicative of a strong correlation between the movement of residues i and j, whereas negative Cij values denote that the two residues move in opposite directions (anticorrelated motion). Both cases are relevant when investigating biological macromolecules, particularly in couples of residues that are located far apart in the three-dimensional (3D) structure. Long-range correlated movements are often linked to the biological functions of the protein.20–23 The analysis was performed on the 1076 Cα carbons of the tetramer (269 for each monomer) because they contain sufficient information to describe the largest protein motions. Note that the map is symmetrical, so the choice of highlighting a particular element in the upper left or lower right is only because of visual effects. The four squares along the black diagonal (each residue has a correlation of 1 with itself) contain the movement correlation among the residues of each monomer. The figure clearly shows highly correlated movements among the DBD residues in all monomers with differences in intensity (the DBD in M4 shows the highest correlation among its residues). In contrast, only M3 shows a strong anticorrelation between the DBD and TET residues. Regarding the intermonomer correlations in DBD, M2 and M4 exhibit the highest (anti)correlation, whereas M1 and M3 have a similar but lower signal. A positive correlation is observed between the DBDs in M3 and M4. No significant correlations are observed between the DBDs in the couples M1–M2, M1–M3 and M2–M3.

Positive correlations are present between nearly all of the TET couples (highlighted by ellipses in A). Interestingly, we observe the presence of a strong anticorrelation signal between the TET in M2 and DBD in M3 and M4 and between the TET in M3 and DBD in M4 (highlighted by the three rectangles in B). These last interactions, together with the strong anticorrelation between DBD and TET in M3, demonstrate that the role of TETs in the tetramer is not over after the formation of the complex with DNA. Instead, communication between TETs and DBDs is always

Figure 2. (a) DCC Map for the p53 tetramer. Four squares delimit the intra-DBD interactions in monomers M1–M4. High correlated motions between couples of TET are highlighted by ellipses A. High anticorrelated motions between TET-DBD couples are highlighted by rectangles B. (b) 2D projection of the concatenated DBD trajectory on the essential subspace along eigenvectors 1 and 2. The structural basin visited by the different monomers is highlighted in different colours (M1–M4 colours as in Figures 1a–d).
maintained in the p53 tetramer. This is consistent with the hypothesis that the DBD can hop on/off the DNA to search for responsive elements. Moreover, structural communications between TETs and DBDs are likely to occur during the p53–DNA dissociation step. The analysis of the DCC Map clearly highlights the significant asymmetry of the monomers, with M3 at the centre of nearly all of the primary long-range correlations between the p53 tetramer elements.

Asymmetric motion of the four DBDs
We investigated the structural basin generated by the four DBDs by concatenating their trajectories and then performing an ED analysis on the obtained 100x4 ns long sampling. Root-mean-square fluctuations of the C-α atoms along the first two ED eigenvectors (Supplementary Figure S2) indicate that their principal motions are dominated by the fluctuations of the two segments connecting the β-strands as in Supplementary Figures S6 and S7 (residues 208–213; see Supplementary Figure S1 for secondary structure nomenclature of the DBD) and Supplementary Figures S6 and S8 (residues 220–229) located in regions opposite the DNA and exposed to the solvent (Supplementary Figure S2). Moreover, the different motion of loop L1 (residues 117–122) in the four monomers is well captured by the second eigenvector. This dynamic information is further discussed in the context of protein–DNA interactions.

The projection of the concatenated DBD trajectory on the essential subspace defined by the first two eigenvectors (Figure 2b) shows a similar conformational behaviour for the M1 and M3 monomers that sample the same subspace region. In contrast, the M2 and M4 monomers visit different regions, thereby reflecting differences in the conformations sampled along eigenvectors 1 and 2. This analysis also indicates that the second eigenvector is able to discriminate between the two couples of monomers (i.e., M1–M3 vs M2–M4), implying that protein regions that are significantly active in this eigenvector (e.g., loop L1) are likely to have a key role in determining the similarities (M1–M3)
and differences (M1–M3 vs M2 vs M4) in monomer conformational behaviour. In accordance with the DCC map (Figure 2a), a break in the tetramer symmetry in the DBDs is observed.

### P53–DNA interactions

The high correlations between key p53 regions demonstrate the presence of structural and dynamic long-range communications in which DBD is central. We investigated its role in the interaction with DNA by identifying the long-lasting (longer than 60 or 80% of the simulation time) direct hydrogen bonds between p53 and the 18 bp DNA (Figure 3a). In accordance with the X-ray data, all monomers strongly interact with the phosphate backbone of T5-G6 or T15-G16. Some contacts are well conserved in all of the monomers, such as the interaction between Arg273 with T15 in M1 and M3 and with T5 in M2 and M4 or the interaction between Ala276 and Ser241 with G16 in M1 and M3 and G6 in M2 and M4. Other interactions are present only in some monomers, such as that between Arg248 and T5 in M2 and M4 and Asn239 with G6 and G16 in M2 and M3, respectively.

In addition to these nonspecific DNA–protein interactions, some base-specific interactions are also present. Arg280 in helix H2 forms very stable contacts with G16 in M1 or G6 in M2. Notably, M3 and M4 also participate in these latter interactions but for a percentage of time that is shorter than the chosen 60% threshold (compare >85% of the simulation time in M1 and M2 vs 36–55% in M3 and M4). Ala276 in M3 contacts N4 of G18. The very stable base-specific interactions between Lys120 and the G11–G12 bases in M4 (63 and 65%, respectively) are also present in M2 with a permanence time of 44–57% and M1 with a permanence time of 36–39%. M2 also possesses a very stable interaction of Lys120 with the A10 backbone atoms. Interestingly, the X-ray study of the p53 tetramer highlighted two distinct conformational states for the loop L1 (residues 117–122) in M2 and M4 in which L1 tucked into the major groove, in contrast with M1 and M3 where L1 adopted an outward projection away from the major groove. In our 100 ns simulation, analysis of the direct Lys120–DNA interactions shows that while L1 in M4 maintains an interaction with DNA for nearly the entire simulation time, these are weaker in M2; moreover, M3 maintains the outward conformation for the entire simulation time, and we observe a change in the orientation of L1 in M1 that allows the formation of base-specific interactions for nearly 40% of the simulation time.

Therefore, observation of the hydrogen bond network formed by Lys120 in the four DBDs can increase our understanding of the dynamic behaviour of loop L1, which is captured by the second eigenvector in the previously described DBD ED analysis. Examining the conformational space visited by the four monomers along this principal motion direction, we observe that M4 visits a smaller space (blue area in Figure 2b), which is in accordance with the observed persistence of the loop L1–DNA contact. M3 (green area in Figure 2b), which does not show a relevant interaction with DNA, visits a conformational region far from the M4 region. The conformational regions visited by M1 (black area in Figure 2b) and M2 (red area in Figure 2b) along the second eigenvector largely overlapped with those of M2 and M3, respectively. This again is in accordance with their initial loop L1 conformations captured by X-ray. However, in contrast with the other two monomers during the simulation, M1 and M2 visit conformations that are different from the starting conformations (which again is in accordance with the relatively larger and overlapping essential subspace visited by these monomers). The 3D conformations of L1 in the starting X-ray conformation and in

### Table 1. Local base-pair step parameters for the average MD and starting X-ray structure (Chen et al.25)

| Step   | Shift | Slide | Rise  | Tilt  | Roll | Twist |
|--------|-------|-------|-------|-------|------|-------|
| GC/GC  | 0.11±0.46 | 0.51±0.43 | 3.29±0.25 | −0.8±4.1 | −1.5±3.98 | 24.7±3.3 |
| CA/TG  | 0.6±0.45  | 0.52±0.45  | 3.2±0.26  | 0.99±4.4  | 6.4±4.95  | 41.99±3.3  |
| AT/AT  | −0.27±0.59 | −0.32±0.34 | 3.2±0.27  | 1.99±3.85 | 3.54±4.15 | 25.4±3.5  |
| TG/CA  | 0.47±0.39  | 0.51±0.40  | 3.31±0.28 | −2.45±4.35 | 3.65±5.16 | 41.9±3.1  |
| GC/GC  | 0.21±0.39  | −0.83±0.46 | 3.6±0.29  | 2.65±4.16 | −6.7±4.97 | 35.95±4.91 |
| CC/GG  | −0.79±0.53 | −0.40±0.66 | 3.64±0.38 | −2.76±4.24 | 1.9±5.04  | 33.47±4.3  |
| CT/GA  | 0.78±0.55  | 0.38±0.54  | 3.39±0.35 | 4.04±4  | −2.12±7.6 | 37.23±5.55 |
| TA/TA  | −0.08±0.69 | 1.26±0.48  | 3.09±0.32 | 0.15±5.9 | −0.08±6.1 | 36.87±8.03 |
| AG/CT  | 0.56±0.53  | −0.04±0.5  | 3.19±0.29 | −2.87±3.34 | −0.14±4.74 | 32.25±4.1  |
| GC/GC  | 0.74±0.38  | −1.15±0.36 | 3.67±0.29 | 0.92±3.7  | −0.24±4.58 | 27.87±4.46 |
| GC/GC  | −0.34±0.34 | −1.0±0.39  | 3.38±0.24 | 0.43±3.6  | −2.3±3.96 | 36.6±3.7  |
| CA/TG  | 0.60±0.49  | 0.47±0.37  | 3.41±0.27 | 1.12±4.2  | 5±4.8  | 44.1±2.6  |
| AT/AT  | 0.00±0.57  | −0.12±0.3  | 2.93±0.25 | −0.36±3.99 | 5.19±3.53 | 21.38±3.11 |
| TG/CA  | −0.58±0.42 | 0.69±0.4  | 3.31±0.27 | −1.54±4.39 | 6.1±4.6  | 43.78±2.67 |

Abbreviation: MD, molecular dynamics. Only the central TA bases show a deviation from B-DNA. The DNA parameters that are significantly different from their corresponding values found in standard B-DNA are shown in bold.
In our MD simulation, L1 in M1 is observed to move from the outward (Figure 3d) to an inner conformation (Figure 3e) with respect to the major groove. In contrast, L1 in M2 maintains the inner conformation of the starting X-ray for nearly 60% of the simulation time (Figure 3e), although the outward conformation is sampled as well (Figure 3d).

It is worth noting that loop L1 has been proposed to regulate p53-DNA-binding kinetics and to confer sequence specificity. Moreover, a computational simulation of the DBD tetramer performed by Pan and Nussinov before the availability of the X-ray structure used in our model (and therefore with different p53 starting positions and a different DNA sequence), nevertheless, suggested a key role for Lys120 in loop L1 as an allosteric switch for the p53 tetramer ED (Figure 3b).

DNA structure and protein-coupled motions

From the analysis of the DNA parameters obtained by our MD simulations, we found that DNA remained largely in its B-form, although we observed a slight distortion of the slide parameter of the central thymine–adenine base pairs that was in accordance with the crystallographic data. Table 1 shows the average DNA parameters from our MD in comparison with those observed in the initial X-ray structure.

The X-ray starting structure has a significant deviation in the central TA bases from the standard B-DNA in three parameters: slide, roll and twist (bold in Table 1). The slide parameter for the same couple of bases shows an analogous deviation to that observed in the X-ray structure during the entire simulation time. In contrast, a deviation from the standard B-DNA of the roll and twist parameters is only observed in a region of the conformational space sampled during the MD simulation. The projection of the trajectory along the first and second eigenvectors obtained from the essential dynamic analysis of the whole p53 core in its tetrameric form is shown in Figure 3b. We coloured the tetramer conformations in which the DNA roll and twist parameters deviate from the B-DNA in purple. Interestingly, this DNA distortion coincides with a well-defined area of the tetramer essential subspace, thereby demonstrating a clear coupling between the DNA distortion and p53 core tetramer conformations.

Therefore, our MD simulation was able to differentiate between slide distortion that is most likely due to the binding of the p53 tetramer itself and the other DNA distortion measured by the roll and twist parameters that is dynamically switched on and off as a function of the specific p53 tetramer conformation and is instantaneously visited. The ED analysis shows that this last DNA distortion is observed only when a specific region (well described by the first eigenvector) of the tetramer conformational space is visited.

Induced-fit effect in p53: how the C terminus facilitates contact with the core DBDs of p53

Recently, the unstructured lysine-rich region in the p53 C-terminal domain was shown to control site-specific DNA binding and to promote structural changes in the DBD. The findings reported in the present study provide a biophysical explanation for the biological evidence demonstrated by Laptenko et al. Collectively, both studies represent an important step towards discrimination between two non-mutually-exclusive alternatives that contribute to the regulation of specific p53–DNA complexes via the C-terminal domain. In the first scenario, the C-terminal domain provides multiple nonspecific p53–DNA interactions that stabilize the binary complex (i.e., the lysine residues function simply as an anchor set on each side of the corresponding binding site to provide extra stabilization). The second possibility is based on an induced-fit mechanism, where the C-terminal domain actually induces specific local conformational changes within the p53 tetramer upon binding to the p53 site.

In our model, the unstructured fragment containing Lys370, 372, 373, 381, 382 and 386 is not present (see the Materials and methods section); hence, this structure corresponds to the experimental Δ30 p53 form. Our results clearly demonstrate long-range effects of TET on the conformations visited by DBD, further producing a deformation in the bound DNA structure. Therefore, our data demonstrate the existence of an induced-fit mechanism that is in keeping with the experimental results obtained with Δ30 p53, which demonstrated that in the absence of the lysine fragment the effect is eventually modulated.

To further understand this complex biological interaction, we investigated the mechanism by which p53 may transduce local DBD conformations. Specifically, we investigated the long-range effects of Lys120 that are capable of influencing or being influenced by other molecular partners. To this end, we calculated the electrostatic potential of two p53–DNA tetramer conformations that were representative of different structural basins. These conformations were chosen at the extremes of the first eigenvector of the DBD tetramer essential dynamics (purple point at the left and black point at the right in Figure 3b). The distribution of the positive and negative isosurfaces at a contour level of ± 1 KT/e is depicted in Figure 4 and Supplementary Figures S3–S9 for different orientations of the two structures. The tetramer has a dominant-negative surface in the TET region (Figures 4a and b, red colour) and a dominant-positive surface in the central DBD

Figure 4. Positive and negative electrostatic potential isosurfaces at a contour level of ± 1 KT/e are depicted in red and blue colours, respectively. (a1–c1) Different orientations of the conformation corresponding to the minimum of the first eigenvector of the DBD tetramer ED (Figure 3b). (a2–c2) Same orientations as panels a1–c1 for the conformation corresponding to the maximum of the first eigenvector of the DBD tetramer ED (Figure 3b).
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region that almost completely covers the negatively charged DNA. However, a negative surface (Figure 4c) is shown in correspondence of β-strands S1 and S3 and loop 2, connecting β-strands S3–S4 and S7–S8 (see Supplementary Figure 2S). Therefore, the complex is fully able to modulate the electrostatic surface according to the two different conformations (compare Figure 4, panels 1 vs 2). For example, when the tetramer visits the structural basins at a minimum value along the first essential eigenvector (panels 1), the negative isosurface in contact with TET completely changes its orientation and volume (compare A1 vs A2 and B1 vs B2). This is the biophysical rational for the induced-fit interaction between the C terminus and the DBD.

DISCUSSION

P53 was originally named the guardian of the genome and was thought to protect the organism from cancer, infertility and ageing[24-45] (reviewed in refs 46–49). The most significant evidence involving p53 in cancer is its high mutation rate in cancers and its ability to drive cancer progression when mutated.[50-56] At the metabolic level, p53 affects mitochondrial function, resulting in activities that directly regulate the survival of the cell.[57–63] The understanding of the highly complex signalling regulation of p53[64–70] has been further complicated by the presence of different protein isoforms.[71–74] P53 and its family members exert their actions by regulating the cellular microRNA in an intricate network.[75–84] Additionally, there is also a rather complex interaction between p53 and autophagy that acts in both directions.[85–91] Several novel drug design or high content screening investigations are attempting to use the p53 pathway for therapeutic applications in cancer,[92–95] mostly by exploiting its protein stability[96–99] rather than its transcriptional activity.[100–104]

Recently, p53 aggregation studies indicated an important regulatory effect of the conformation for the p53 protein.[105–107] Consequently, we propose that an atomistic model of the p53 tetramer including the C-terminal tetramerization helices can improve our understanding of this system.

The MD simulation of the p53 tetramer in complex with DNA highlights the presence of long-range communication between different domains that allows p53’s biological function. The inclusion of the TET at the p53 C terminus for the first time in an atomistic model permits the observation of long-range communications both at the intramonomer level between TET and the whole DBD in M3 (Figure 2a) and at the intermonomer level between M2-TET and M3-DBD and between M3-TET and M4-DBD (Figure 2a). The protein displays a double-face interaction with DNA through several nonspecific non-covalent bonds with the DNA backbone (mostly accomplished by β-strands S9 in the β-sandwich and the C-terminal end of β-strand S10 in the loop–sheet–helix motif), whereas specific sequence recognition is performed by helix H2 and loop L1, both located in the loop–sheet–helix motif (Figure 3a). Loop L1 shows nonsymmetric dynamics in the four monomers, as evidenced by the essential dynamics of the concatenated DBD trajectory along the second eigenvector (Figure 3b and Supplementary Figure S2). The simulation shows the switch from a conformation in which L1 tucks into the major groove to an outward projection away from the major groove, and vice versa (Figures 3c–e). These results provide an atomistic explanation for the regulation of p53 functions through post-translational modifications of the C terminus.[108–109]

The C terminus of p53 has a strong biological effect on the transcriptional function of p53 itself because it is able to regulate DNA binding, p53 stability, p53 subcellular localization and the recruitment of cofactors (see Laptenko et al.[30]). Although this effect has been related to changes in the affinity of electrostatic interactions between the lysines at the C terminus during specific binding with other p53 regions, the in-depth mechanism has not been clarified. Although our manuscript was in evaluation, Laptenko et al.[30] demonstrated the biological value of the lysine interaction on the stability of the p53–DNA complex through the facilitation of cooperative contacts with the DBD.[30] This result can be explained by the pivotal concept of the ‘induced-fit theory’. Indeed, Petty et al.[106] showed a conformational change within the DBD that involved the L1 loop, which adopted either an extended or a recessive conformation. Our data show the dynamic flexible movement of the four monomers and demonstrate how the C terminus can modulate the electrostatic surface regardless of the presence of the lysines (which are absent in our model) (Figure 4) according to the two different conformations at the extreme of the first essential eigenvector. This is the biophysical rational for the understanding of the C-terminus interaction with the DBD.

The implications of our results are evident on the induced-fit theory (see Koshland[107] and Johnson[111]) and fully explain the results by Laptenko et al.[30] The presence of an induced-fit conformational transition will change the kinetic equilibrium, where multiple structural switches allow a better fit between p53 and DNA, thereby stabilizing the bound conformation. Halazonetis identified a role for L1 in this mechanism,[112] although we cannot exclude conformational induced fit by other association partners taking place in the transcription machinery complex.[113] As indicated above, our results in keeping with Laptenko et al.[30] demonstrate that the C terminus can create an induced-fit mechanism per se on the DBD, even in the absence of other partners. Note that long-range interdomain communications have already highlighted in the p53 full-length model[30] and preliminary results on cancer-related mutations located in DBD confirm the ability of p53 to deeply alter the dynamics of the TAD regions in the N-terminal domain and TET in the C-terminal domain (Figure 5).

Concerning the DNA conformation, a general standard B-form is maintained in the entire double strand; the only exception is the central thymine–adenine base pairs (Table 1). This result was in agreement with the description of the 3D structure obtained by X-ray diffraction. However, the simulation implies that the slide parameter is outside the range of the standard B-form for the entire simulation time, whereas the roll and twist parameters (also distorted in the starting X-ray structure) are distorted in the MD simulation only for tetrameric conformations corresponding to a specific region of the essential subspace (purple points in Figure 3b). Therefore, the MD simulation separates the ‘static’ DNA deformation that most likely occurs because of p53–DNA binding (measured by the slide parameter) from the ‘dynamic’ DNA deformation (measured by the roll and twist parameters) that is switched on and off as a function of the specific p53 tetramer conformation.

MATERIALS AND METHODS

The model of the p53 tetramer in interaction with a specific DNA response element was built using the Modeller software package[114] with the X-ray crystal structure 3KMD as a reference for the p53 core domains and DNA.[25] The NMR structures from 3SAK were used as references for the tetramerization helices.[115] The first 90 residues at the N terminus and residues 360–391 at the C terminus are intrinsically disordered.[116] We have excluded them from the model for two main reasons: (i) the inclusion of these residues in the model makes the system computationally very demanding and (ii) they would not affect the conformational behaviour of the rest of the protein complex. Forcefield parameters for the Zn ion four-ligand coordination interface were obtained from Lu et al.,[117] who performed an MD study of the p53 core domain.

The model of the p53 tetramer was solvated and the system was then minimized after a short simulation. Owing to the size of the system, the system was equilibrated by applying restraints of 1000 kJ/mol/nm² to the DNA and the tetramerization helices. Then, four independent unrestrained MD simulations were performed from the final structures of the restrained runs for a total length of approximately 100 ns. All of the MD simulations
an interface to the DSSP program.\textsuperscript{119} The DNA parameters were obtained with the Curve+ program.\textsuperscript{28} The electrostatic potential was calculated with the APBS software using 2.0 and 80 as the solute and solvent dielectric at zero salt concentration, respectively.\textsuperscript{120} The figures in the 3D structures were generated with vmd.\textsuperscript{121}

Average properties such as DCC Map, per-residue RMSF and hydrogen bonds were checked against random time windows, each containing 25 000 frames.

CONFLICT OF INTEREST

The authors declare no conflicts of interest.

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