

## PAPER

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[View Journal](#) | [View Issue](#)Cite this: *RSC Mechanochem.*, 2024, 1, 78Enhanced HP1 $\alpha$  homodimer interaction *via* force-induced salt bridge formation: implications for chromatin crosslinking and phase separation†Shingo Tsukamoto,<sup>a</sup> Mohammad Khavani,<sup>a</sup> Nya Domkam<sup>a</sup> and Mohammad R. K. Mofrad<sup>\*ab</sup>

Recent studies have underscored the potential role of Heterochromatin Protein 1 $\alpha$  (HP1 $\alpha$ ) in chromatin crosslinking, phase separation, and the orchestration of nuclear mechanics. One of the cornerstones of HP1 $\alpha$  functionality lies in its homodimerization through the chromoshadow domain (CSD), which is crucial for these processes. Nevertheless, it has remained unknown how HP1 $\alpha$  can foster condensations responding to mechanical force and induce phase separation in the mechanically unfavorable heterochromatin region. To elucidate the biophysical basis of HP1 $\alpha$ , we used full atomistic molecular dynamics (MD) simulations, focusing on the CSD–CSD dimer of HP1 $\alpha$  under a pulling force. Notably, force application resulted in a stronger, more stable interaction at the  $\alpha$ -helix interface of the CSD–CSD. This enhanced interaction was attributed to a force-induced salt bridge formation on the  $\alpha$ -helix interface, emerging from an angle alteration of a lysine residue that enables closer proximity to a glutamic acid residue on the paired CSD. This study reveals an intriguing facet of HP1 $\alpha$  mechanics: its mechanical sensitivity, wherein dimerization strength is enhanced by mechanical force. The molecular dynamics of the CSD–CSD dimer under force provide novel insights into HP1 $\alpha$  mechanics, contributing to our understanding of chromatin mechanics and phase separation.

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## Introduction

Mechanical forces, originating both inside and outside the cell, such as cell substrate stretching and actin contraction, are conveyed to the nucleus *via* the cytoskeleton and the LINC complex.<sup>1–3</sup> The active and passive responses of the nucleus and chromatin to these mechanical forces impact gene

transcription by modulating the accessibility of mechano-sensitive transcription factors or by directly altering chromatin and DNA structures.<sup>4,5</sup> Unlike the relatively stable structure of DNA, chromatin exhibits high mechanochemical dynamism, readily undergoing structural changes.<sup>6,7</sup> Understanding the mechanical responses of chromatin is thus a challenging and crucial aspect of mechano-genomic regulation.

Heterochromatin, characterized by its compact, transcriptionally silent, methylated, and mechanically rigid nature,<sup>8,9</sup> plays a pivotal role in providing mechanical stability and shielding DNA from mechanical damage.<sup>10,11</sup> Histone modifications can modulate chromatin condensation states, with histone methylation promoting increased heterochromatin content and nuclear stiffness.<sup>9,12–16</sup> However, our previous study revealed that local strain distribution caused by cell substrate stretching remained unaffected by trichostatin A (TSA) treatment, a histone deacetylase inhibitor.<sup>17</sup> Moreover, there is growing evidence that chromosomal proteins may exert distinct effects on nuclear and chromatin mechanics compared to histone methylation.<sup>18</sup> These findings suggest that histone modification alone may not fully encompass the intricacies of intranuclear mechanics.

In recent years, Heterochromatin Protein 1 $\alpha$  (HP1 $\alpha$ ), a chromosomal protein, has garnered increasing attention for its indispensable roles in maintaining nuclear and chromatin mechanics.<sup>18–21</sup> HP1 $\alpha$  accumulates within heterochromatin,

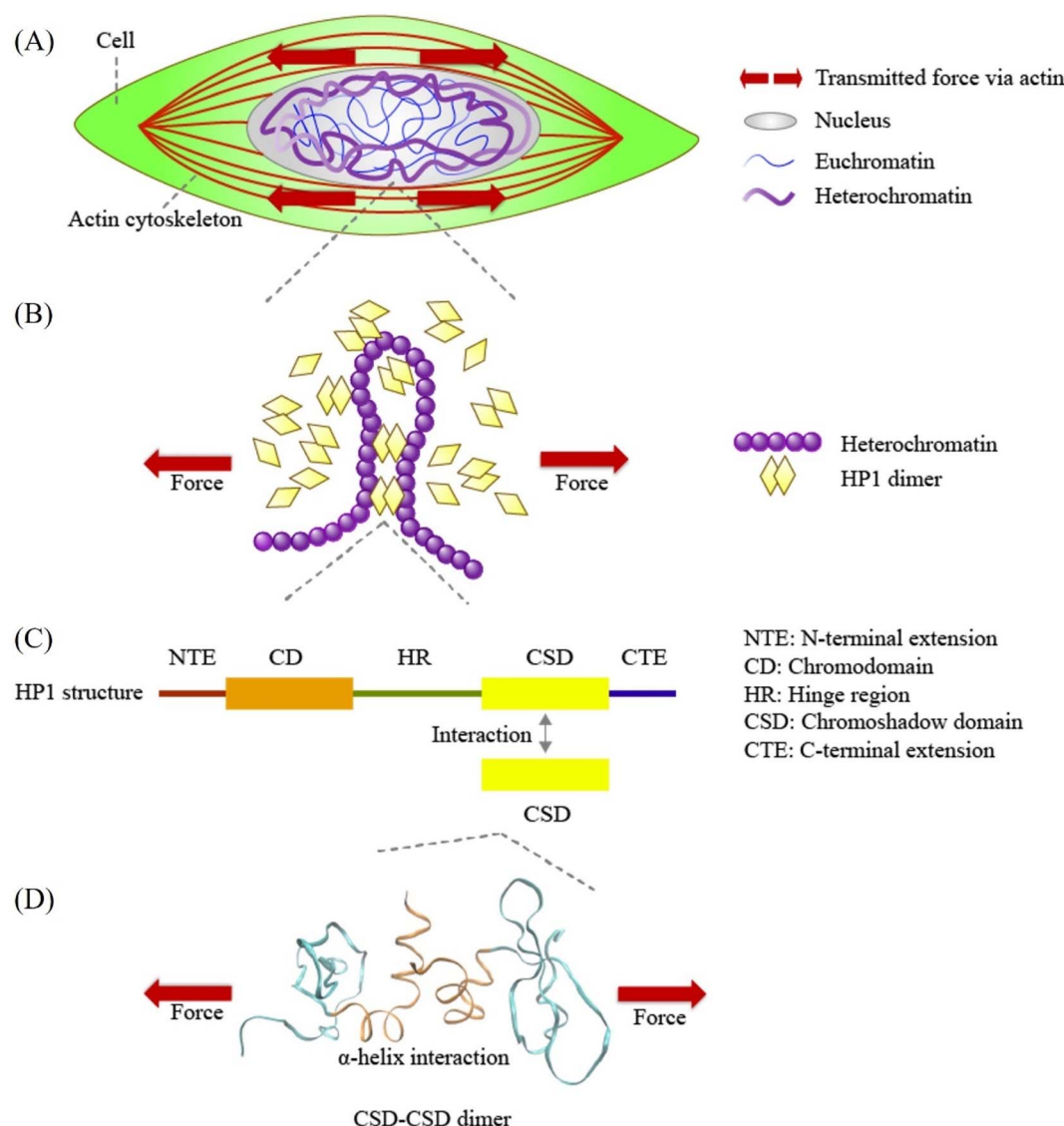
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† Electronic supplementary information (ESI) available: The following analyses and data were conducted and obtained within this study: root mean square deviation (RMSD) profiles for the entire chromoshadow domain (CSD) dimer and individual monomers, as well as each  $\alpha$ -helix within the CSD dimer and monomer, over the course of a 200 ns relaxation period. Visual representation of the CSD–CSD dimer, highlighting the salt bridge residue pairs <sup>1</sup>Lys154–<sup>11</sup>Glu169 and <sup>11</sup>Glu118–<sup>11</sup>Lys159. Frequency histograms depicting the distribution of distances between the salt bridge pair <sup>11</sup>Glu118–<sup>11</sup>Lys159. Analysis of hydrogen bond lifetime distribution for the <sup>11</sup>Glu162–<sup>11</sup>Asn157 pair. Frequency histograms illustrating the distribution of angles between the two  $\alpha$ -helices within the CSD–CSD dimer. Frequency histograms depicting the distribution of lengths for the residue pairs <sup>1</sup>Glu169–<sup>11</sup>Lys154, <sup>1</sup>Glu169– $z$ -axis, and <sup>11</sup>Lys154– $z$ -axis. Frequency histograms illustrating the distribution of distances between <sup>1</sup>Lys154–<sup>11</sup>Glu169, generated without applying any position constraints during a 20 ns duration. See DOI: <https://doi.org/10.1039/d3mr00011g>.

where it compacts chromatin by bridging chromatin fibers, thereby contributing to the mechanical rigidity of the HP1 $\alpha$  accumulates inside heterochromatin and compacts chromatin by bridging the chromatin fibers, contributing to the mechanical rigidity of the nucleus<sup>18,21–23</sup> (Fig. 1B). Studies utilizing coarse-grained polymer models of chromatin have suggested that chromatin crosslinking can alter chromatin's mechanical response and properties.<sup>13,24</sup> Additionally, HP1 has been shown to recruit proteins related to heterochromatin and facilitate liquid–liquid phase separation *in vitro*, a process that substantially influences nuclear stiffness.<sup>25–27</sup> Importantly, the mechanical context in which phase separation occurs significantly affects the process.<sup>28,29</sup> For instance, Shin *et al.*

demonstrated the mechanical sensitivity of nuclear liquid condensation in regions characterized by lower mechanical energy, softer properties, and lower density, such as euchromatin.<sup>29</sup> These findings contrast with the phase separation of HP1 in heterochromatin, which promotes chromatin compaction.<sup>26,27,29</sup> Furthermore, optical trapping experiments applying external forces in stretch-relaxation cycles to HP1 $\alpha$ -DNA condensation surprisingly revealed enhanced DNA compaction by HP1 $\alpha$  following force application.<sup>20</sup> However, several mechanochemical questions concerning how the mechanical properties and responses of HP1 $\alpha$  enable its phase separation within heterochromatin and enhance the stability of HP1 $\alpha$ -DNA condensation after force application remain unanswered. A



**Fig. 1** Schematic representation of force transmission at various scales, from cellular forces to Heterochromatin Protein 1 $\alpha$  (HP1 $\alpha$ ). (A) Mechanical forces are transmitted from the cell through the cytoskeleton to the nucleus. (B) HP1 $\alpha$  plays a pivotal role in binding and bridging chromatin fibers, contributing to phase separation and influencing chromatin structure and mechanics. (C) Human HP1 $\alpha$  protein structure, highlighting its key domains: N-terminal extension (NTE), chromodomain (CD), hinge region (HR), chromoshadow domain (CSD), and C-terminal extension (CTE). CSD interacts with another CSD in the HP1 $\alpha$  monomer, forming the HP1 $\alpha$  homodimer. (D) A visual representation of the CSD–CSD dimer (PDB ID: 3I3C) in VMD software, indicating the application of schematic force. The orange and sky-blue regions represent the  $\alpha$ -helix interface and other residue segments of the CSD–CSD dimer, respectively.



detailed molecular analysis of HP1 $\alpha$  mechanics can provide insights into these mechanochemical questions.

At the molecular level, the formation of chromatin crosslinks and phase separation orchestrated by HP1 relies on HP1 dimerization.<sup>18,26,30–33</sup> HP1 forms a homodimer through its chromoshadow domain (CSD)<sup>34</sup> (Fig. 1C). The CSD monomer primarily interacts with another CSD through their  $\alpha$ -helices to form this homodimer<sup>34</sup> (Fig. 1D). Disruption of the CSD–CSD interactions leads to abnormal nuclear shape and inhibits DNA-driven phase separation.<sup>18,26</sup> Despite the significance of CSD–CSD interactions in chromatin mechanics and phase separation demonstrated in previous studies, numerous questions remain unanswered. One key question is how the CSD–CSD  $\alpha$ -helix-mediated interactions in the HP1 dimer maintain or alter their interactional strength in response to mechanical forces. Understanding the mechanical properties of CSD–CSD  $\alpha$ -helix-mediated interactions in the HP1 dimer is crucial for unraveling the molecular and chromatin assemblies and the mechanical properties of chromatin and the nucleus.

To investigate the detailed molecular mechanisms underlying HP1 $\alpha$ , we conducted molecular dynamics simulations to analyze the mechanical behavior and strength of CSD–CSD  $\alpha$ -helix-mediated interactions under force application. Our simulations unveiled a remarkable sensitivity of the CSD–CSD interaction of HP1 $\alpha$  to mechanical forces, resulting in the formation of force-induced salt bridge interactions. This newfound mechanical sensitivity provides fresh insights into HP1 $\alpha$ 's mechanical resilience, phase separation, and mechano-genomic regulatory mechanisms.

## Methods

### Molecular dynamics simulation

GROMACS19.6 software package with the CHARMM36 force field was employed for the all-atom molecular dynamics (MD) simulation and corresponding analyses.<sup>35–38</sup> The crystal structure of the CSD–CSD of the chromobox protein homolog 5 (CBX5), the human HP1 homolog alpha, is available on the protein database (PDB ID: 3IC3). Since there is a discrepancy in residue numbering between the PDB file and full-length sequences (UniProt ID: P45973), the residues were renumbered corresponding to the full-length sequence. The CSD–CSD structure was inserted into a long rectangular box ( $6 \times 6 \times 12$  nm<sup>3</sup>) with periodic boundary conditions in all three directions, sufficiently large space in the direction of pulling to ensure an uninterrupted pulling process, free from interactions with the periodic images of the system. The system was solvated by using the CHARMM-modified TIP3P water model.<sup>39</sup> The sodium and chloride counter ions were added to the system, representing the 150 mM concentration of NaCl where chromatin can be well aggregated.<sup>40,41</sup> The energy minimization was performed for 50 000 steps with an energy tolerance of 1000 kJ mol<sup>−1</sup> nm<sup>−1</sup> to ensure the system has no steric clashes or inappropriate geometry. To increase the temperature and equilibrate the solvent and ions around the protein, the NVT (isothermal–isochoric) and NPT (isothermal–isobaric) ensembles with V-rescale thermostat at a constant temperature of 300 K were applied to

the system for 5 ns for each with 1 fs time step. The Parrinello–Rahman barostat was employed for NPT ensembles to maintain the desired pressure at 1 atm. The damping parameters for the V-rescale thermostat and the Parrinello–Rahman barostat were set to 0.1 ps and 2 ps, respectively. The position of the heavy atoms of the protein was restrained during NVT and NPT equilibration to eliminate the additional variable of structural changes in the protein. Then, the protein relaxation was performed for 200 ns (with 2 fs time steps) as the product step without any position restrains in the equilibrated system. The particle mesh Ewald method was used for long-range electrostatic interactions with a short-range electrostatic cutoff distance of 1.0 nm. The cutoff for van der Waals interactions was 1.0 nm. The CSD–CSD dimer reached a very stable state after around 100 ns, even compared to a CSD monomer (ESI Fig. S1†). The obtained relaxed structures after 200 ns full atomistic simulations were used as the initial input for the next steered molecular dynamics (SMD) simulations.

Center-of-mass pulling force was applied to the CSD–CSD dimer to obtain satisfied information of the  $\alpha$ -helix binding interactional behaviors against force and simultaneously reduce computational costs.<sup>42</sup> The harmonic potential was used for the pulling simulation, which enables the force to adjust based on the nature of the interactions of the CSD–CSD. Each monomer was pulled apart along the  $z$ -axis at a pull rate of 0.05 nm ns<sup>−1</sup> while the edge residue of one monomer was fixed so that force was applied to the dimer in the same direction during the simulation (Fig. 2A). We defined the unconstrained, more natural state CSD monomer as <sup>1</sup>CSD and the position-constrained CSD monomer as <sup>11</sup>CSD. The deliberate application of a slow pulling rate serves to mitigate the potential disruption to the secondary structure of the protein, thereby facilitating the preservation of its conformational properties. The pulling simulation, the force-applied condition, was performed for 20 ns. The no-force condition, a control group, was defined as the further 20 ns relaxation with the edge residue constraint same as the force condition after the first 200 ns relaxation. The simulation was repeated three times ( $N = 3$ ) for each force/no-force condition. The visualization of the simulation was performed by using VMD software.<sup>43</sup>

### Trajectory file obtainment for analysis

The trajectory files saved every 10 ps were used for the analyses in force and no-force simulations. The  $\alpha$ -helix interface of CSD–CSD was chosen for the analysis to obtain the important mechanical and chemical responses of HP1 $\alpha$  dimer interaction. All analyzed results were plotted using Python.

### RMSD and RMSF

Root mean square deviation (RMSD) values of each  $\alpha$ -helix of <sup>1</sup>CSD (unconstrained) and <sup>11</sup>CSD (constrained) throughout the simulation were computed by using GROMACS software (eqn (1)).

$$\text{RMSD}(t_1, t_2) = \frac{1}{\sum_{i=1}^N m_i} \sqrt{\sum_{i=1}^N m_i \|r_i(t_1) - r_i(t_2)\|^2} \quad (1)$$



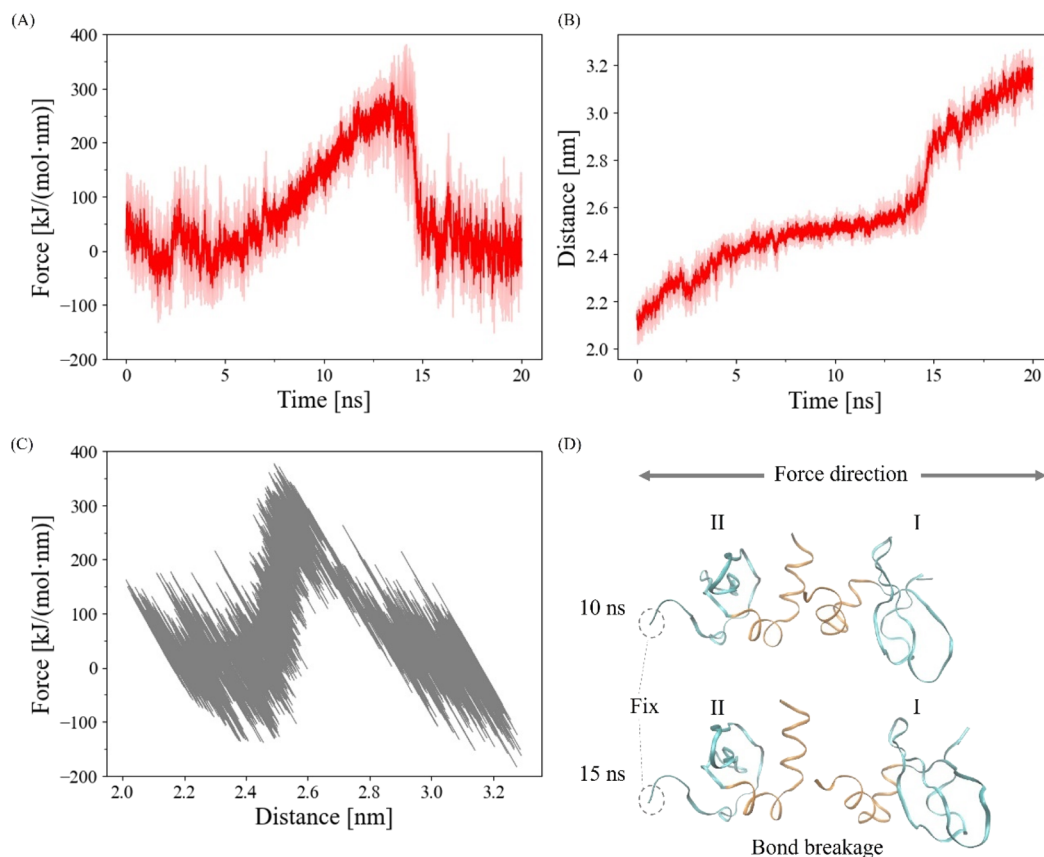


Fig. 2 (A) The averaged pulling harmonic force of three simulations ( $N = 3$ ) with shaded areas representing standard deviations. (B) The averaged distance between the centers of mass of each CSD monomer of three simulations ( $N = 3$ ) with shaded areas representing standard deviations. (C) The distance–force curve over the 20 ns. Data from all three simulations ( $N = 3$ ) was plotted on the graph in gray. (D) Visualized images of the CSD–CSD dimer at 10 ns and 15 ns. The interaction through the  $\alpha$ -helix interface remains intact at 10 ns but is disrupted by 15 ns. The terminal residue of <sup>11</sup>CSD was constrained to ensure that force was applied to the dimer in the same direction throughout the simulation.

where  $r_i(t)$  is the position of atom  $i$  at time  $t$ . The initial time frame structure ( $t_2 = 0$ ) was used as the reference. Root mean square fluctuation (RMSF) of the  $\alpha$ -helix interface was obtained by computing the standard deviation of the atomic positions from the mean positions over time steps on GROMACS (eqn (2)).

$$\text{RMSF} = \sqrt{\langle (r_i - \langle r_i \rangle)^2 \rangle} \quad (2)$$

where  $r_i$  denotes the coordinates of atom  $i$  and  $\langle r_i \rangle$  represents the ensemble average position of the atom. The least-squares fitting was conducted before calculating RMSD and RMSF to remove the effect of global rotation and translation of the protein.

### H-bond calculations

The number of inter-strand hydrogen bonds formed between the two  $\alpha$ -helices of the CSD–CSD dimer and the H-bond lifetime were computed on GROMACS. H-bonds were counted when the distance between the possible acceptor (OH and NH groups) and donor (O and N atoms) pair was within 0.35 nm, and the angle between the hydrogen, oxygen, and nitrogen was below 30°. The histogram for the frequency of H-bond numbers between the  $\alpha$ -helix interface before bond breakage was

computed using Python. The histogram was also fitted by kernel density estimation (KDE) with a Gaussian kernel to visualize the distribution trend. The H-bond lifetime distribution was obtained by calculating the frequency of H-bond breakage within each lifetime interval divided by the total number of H-bonds that have broken over time. Lifetime intervals are every 10 ps, starting at 5 ps.

### Electrostatic energy, salt bridge pair distance, and salt bridge ratio calculations

The short-range inter-strand coulombic energy between the two  $\alpha$ -helices of CSD–CSD with a 1.0 nm cutoff distance was calculated over simulation time on GROMACS and reported as the electrostatic interactions. Distances are measured between possible salt bridge pairs of Lys-NZ and Glu-CD (connected to both OE1 and OE2) in the  $\alpha$ -helix interface (ESI Fig. S2†). The frequencies of the coulombic energy and salt bridge pair distance with kernel density estimation before the interaction breakage were plotted using Python. The salt bridge ratio was defined as the frequencies of salt bridge pairs within 0.4 nm divided by the total number of data over time before the interaction breakage.





## Force distribution analysis

The TRFDA GROMACS tool<sup>44</sup> was used for the force distribution analysis for MD simulation. The coulombic energy was chosen for the force analysis to see the effect of salt bridge interactions. Assuming atom  $i$  is an atom in residue  $r_i$  and  $j$  is in residue  $r_j$ , the residue pairwise force acting on their center of mass was computed in eqn (3).

$$\vec{F}_{r_i, r_j} = \sum_{i \in r_i, j \in r_j} \vec{F}_{ij} \quad (3)$$

The punctual stress with units of force was defined as the sum of the absolute values of the magnitude of the vector pairwise force on a single atom or residue, enabling the identification of where pairwise forces accumulate and the detection of atomic-level hot-spots. We used the term “punctual stress” followed by the original paper of the TRFDA GROMACS tool.<sup>44</sup> In the calculation of punctual stress, the force acting on a dimensionless point instead of an area was utilized due to the difficulty of defining geometrical properties. Punctual stress heatmaps and averages on the  $\alpha$ -helix interface for each residue were made by using Python.

## Structural analysis of residue pairs

The Lys-NZ or Glu-CD (ESI Fig. S2†) and  $\alpha$ -carbon of a certain residue were chosen to define a vector. Angles before the interaction breakage cutoff were calculated between the two vectors of interest of the residues or between one vector and the z-axis in the pulling force direction on GROMACS. The angle frequencies with kernel density estimation before bond breakage were plotted using Python.

## Results

Previous experimental studies employing force application have underscored the significance of Heterochromatin Protein 1 $\alpha$  (HP1 $\alpha$ ) in the mechanics of the nucleus and chromatin.<sup>18,20</sup> Central to the mechanical roles of HP1 is its homodimerization through the chromoshadow domain (CSD), a process that underpins HP1's ability to establish chromatin crosslinks and drive phase separation.<sup>18,25,26,30–33</sup> To delve deeper into the molecular intricacies governing the response of HP1 $\alpha$  dimerization to applied force, our investigation was directed toward the CSD–CSD dimer. We utilized Root Mean Square Deviation (RMSD) and Root Mean Square Fluctuation (RMSF) analyses to gain insights into the specific residues and interactions that underpin the CSD–CSD dimer interaction. Subsequently, we investigated the detailed dynamics of the interplay between the CSD–CSD dimer interaction under conditions with and without applied force. We employed Force Distribution Analysis (FDA) to scrutinize how forces are distributed throughout the  $\alpha$ -helix interfaces of the CSD–CSD dimer. This analysis aimed to provide a comprehensive understanding of the mechanical effects of pulling forces on individual residues within the dimer. Ultimately, our structural analysis elucidated the alterations in residue angles induced by the applied force, thus revealing the

emergence of force-induced interactions within the CSD–CSD dimer.

## The disruption of CSD–CSD dimer interaction under pulling force

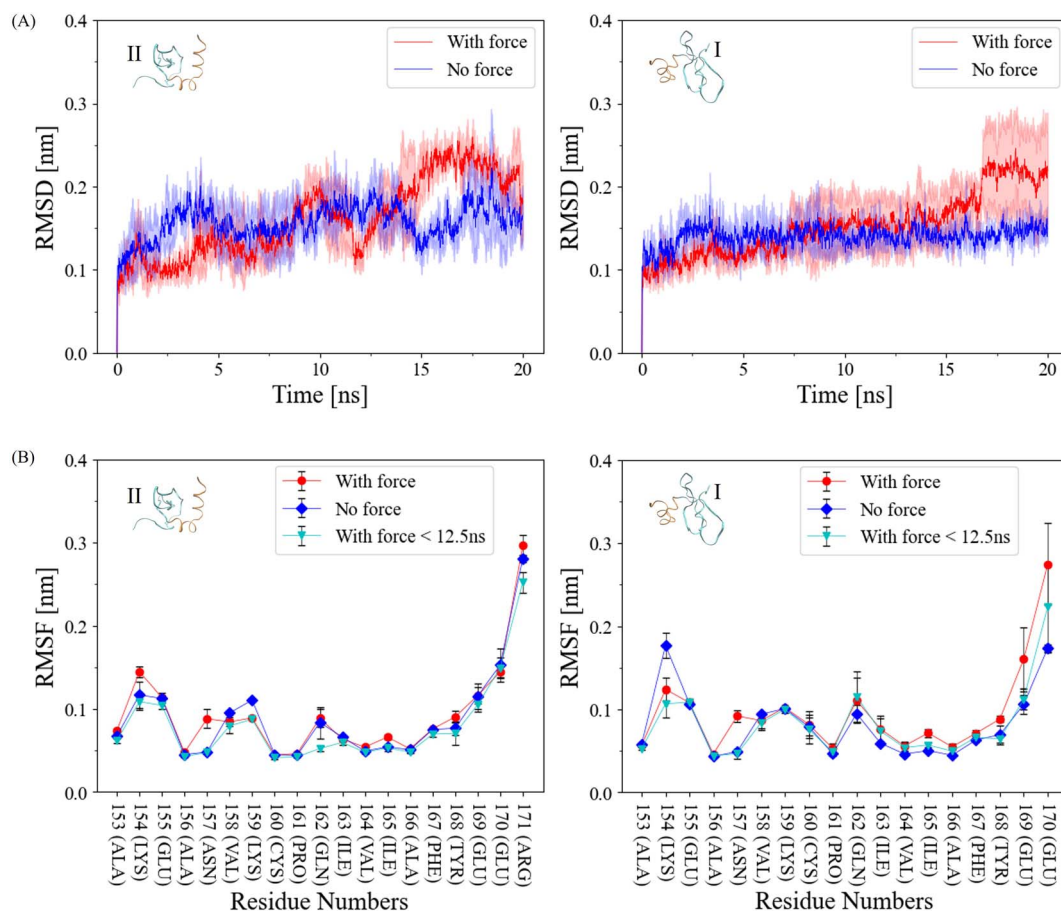
To validate the proper execution of our simulation, we applied a harmonic pulling force using GROMACS, as depicted in Fig. 2. In Fig. 2A and B, we present the averaged harmonic forces and distances between the centers of mass of each CSD monomer from three simulations along with the standard deviations indicated by the shaded areas. Notably, the CSD–CSD dimer interaction *via* the  $\alpha$ -helix interface experienced a rupture within the 20 ns simulation period. The distance–force curve showed an increase in force between distances of 2.4 and 2.6 nm, followed by a sharp decrease upon reaching approximately 300 kJ mol<sup>−1</sup> nm<sup>−1</sup> at 2.6 nm (Fig. 2C). The maximum forces leading to interaction breakages were consistently observed between 13 and 15 ns, displaying notable fluctuations across all simulations. Consequently, to eliminate the influence of abnormal fluctuations resulting from the interaction breakage process and to effectively observe the impact of the pulling simulation on the  $\alpha$ -helix interaction, we established a 12.5 ns cutoff for bond breakage.

## The role of amide groups and charged residues in interaction stability under force

In our pursuit to comprehend the fluctuation dynamics at the  $\alpha$ -helix interface of CSD–CSD induced by mechanical force, we calculated the Root Mean Square Deviation (RMSD) of this interface and compared the results between force-applied and no-force conditions (Fig. 3A). In the no-force condition, the RMSD remained relatively stable at around 0.15 nm throughout the simulation. In contrast, the RMSD under force exhibited a substantial increase after the interaction breakage, which occurred at the 12.5 ns cutoff. Curiously, RMSD values under force prior to this cutoff were relatively lower than those in the no-force condition. To explain this intriguing observation, we can postulate that certain regions within the proteins exhibit reduced flexibility when subjected to external forces. Consequently, these regions demonstrate reduced fluctuations during the simulation before reaching the breaking point, in contrast to systems devoid of applied forces.

The Root Mean Square Fluctuation (RMSF) of each interacting  $\alpha$ -helix within CSD–CSD was assessed based on individual residues (Fig. 3B). While most residues exhibited similar RMSF values between the force and no-force conditions, notable distinctions were observed in specific amide groups, such as residue Asn157, and certain charged residue groups, such as Lys154. Asn157 exhibited almost the same RMSF values in the no-force and the force-applied condition before the 12.5 ns bond breakage cutoff, but in the over 20 ns simulation, RMSF was higher in the force-applied condition. Interestingly, Lys154 presented a contrasting behavior, demonstrating a lower RMSF value under force-applied conditions than in the no-force condition, even before the bond breakage cutoff. The increased RMSF values under force throughout the simulation





**Fig. 3** (A) Root mean square deviation (RMSD) of each  $\alpha$ -helix of the <sup>1</sup>CSD (right, unconstrained) and <sup>11</sup>CSD (left, constrained) throughout the 20 ns simulation, with and without force, and standard deviations indicated by shaded areas.  $N = 3$  for each force and no-force condition. (B) Root mean square fluctuation (RMSF) of each  $\alpha$ -helix of the <sup>1</sup>CSD (right, unconstrained) and <sup>11</sup>CSD (left, constrained) over the 20 ns for both force and no-force conditions, and prior to the 12 ns bond breakage cutoff with force. The results are presented as mean  $\pm$  standard deviation.  $N = 3$  for each force and no-force condition.

can be attributed to the significant bond breakages involving the corresponding residues, resulting in heightened fluctuations. Conversely, the lower RMSF values under force before the 12.5 ns cutoff may suggest the development or stabilization of bonds involving these specific residues.

Amide and charged residue groups play critical roles in the formation of hydrogen bonds and salt bridge interactions. To further elucidate the determinants behind the distinct fluctuation behaviors under force, we delved into the dynamics of hydrogen bonds (H-bonds) and salt bridge interactions in the subsequent sections.

#### Pulling force does not alter overall H-bond properties on the $\alpha$ -helix interface of the CSD–CSD dimer

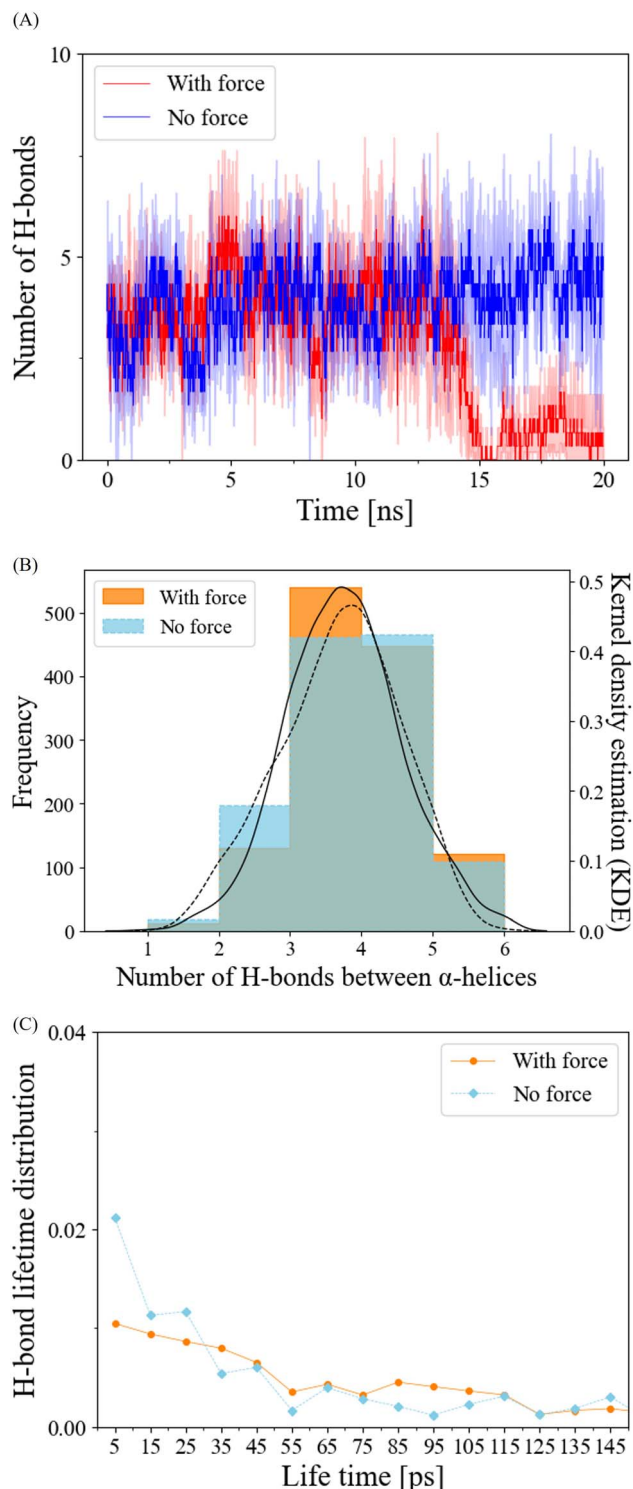
In our exploration of H-bond dynamics at the  $\alpha$ -helix interface, we conducted an analysis of both the number and lifetime of H-bonds (Fig. 4). For the no-force condition and prior to the interaction breakage cutoff at 12.5 ns in the force condition, the number of H-bonds remained relatively constant, typically ranging from 3 to 5 (Fig. 4A and B). Following the cutoff, the force-applied condition exhibited a pronounced decrease in the

number of H-bonds (Fig. 4A). When we examined the frequency distribution of the number of H-bonds using kernel density estimation (KDE) before the breakage cutoff, both the force and no-force conditions displayed similar trends, with peak frequencies observed at 3 to 5 H-bonds (Fig. 4B). The lifetime distribution of H-bonds before the breakage cutoff demonstrated that H-bond breakages occurred in a manner that was largely independent of the application of pulling force (Fig. 4C). These findings collectively suggest that the mechanical responses of H-bonds are not the primary factors contributing to the reduced RMSD and the stabilization of  $\alpha$ -helix interface interactions in the dimerization of HP1.

#### Discovery of a newly formed salt bridge under mechanical force enhancing the $\alpha$ -helix interface interaction in the CSD–CSD dimer

To gain insights into the behavior of salt bridge interactions between the  $\alpha$ -helix interfaces under force, we initiated our investigation by calculating short-range coulombic energy changes within these interfaces over the course of the simulation (Fig. 5A). As expected, a significant decline in coulombic





**Fig. 4** Hydrogen bonds (H-bonds) properties at the  $\alpha$ -helix interface of the CSD–CSD dimer. (A) Number of H-bonds at the  $\alpha$ -helix interface with/without force during the 20 ns simulation, with standard deviations shown as shaded areas.  $N = 3$  for each force and no-force condition. (B) Histogram showing the frequency of H-bond numbers at the  $\alpha$ -helix interface breakage cutoff of 12.5 ns under force and no-force conditions comprising data from all six simulations ( $N = 3$  for each force/no-force condition). The histogram was also fitted by kernel density estimation (KDE). The black solid line in the histogram represents the KDE for the force-applied condition, while the black dashed line corresponds to the KDE for the no-force condition. (C) Distribution of H-bond lifetime with/

energy was observed after the 12.5 ns breakage cutoff. Interestingly, prior to this cutoff, attractive coulombic energy between the  $\alpha$ -helix interfaces was more prominent when force was applied compared to the no-force condition. For a more quantitative analysis, we computed the distribution of coulombic energy before the breakage cutoff (Fig. 5B). In the absence of force, the coulombic energy displayed the highest frequency at approximately  $-200 \text{ kJ mol}^{-1}$ . In contrast, the peak of the Kernel Density Estimation (KDE) of the coulombic energy frequency with force was situated at about  $-240 \text{ kJ mol}^{-1}$ . The average coulombic energy values were  $-196 \text{ kJ mol}^{-1}$  and  $-236 \text{ kJ mol}^{-1}$  in the no-force and force conditions, respectively. These results clearly indicate that the application of a pulling force strengthens coulombic energy interactions at the  $\alpha$ -helix interfaces of the CSD–CSD dimer.

Salt bridge interactions are facilitated by coulombic attraction with water molecules.<sup>45</sup> To validate the sequence data of the  $\alpha$ -helix and visualize the residues, we identified potential residue pairs that could form salt bridge interactions (Fig. 5C). The residue  $^1\text{Lys154}$  can interact with residue  $^{11}\text{Glu169}$ , and reciprocally,  $^1\text{Glu169}$  and  $^{11}\text{Lys154}$  are capable of making salt bridges as well.

To quantitatively assess the dynamics of these salt bridge interactions, we calculated the distances between each residue pair. The positively charged atom involved is Lys NZ1, while the negatively charged atoms are Glu OE1 and OE2, contributing to the salt bridge. Since both Glu OE1 and OE2 atoms can participate in interactions with Lys NZ1, we selected Glu CD, which connects to both Glu OE1 and OE2, for distance calculations (ESI Fig. S2†).

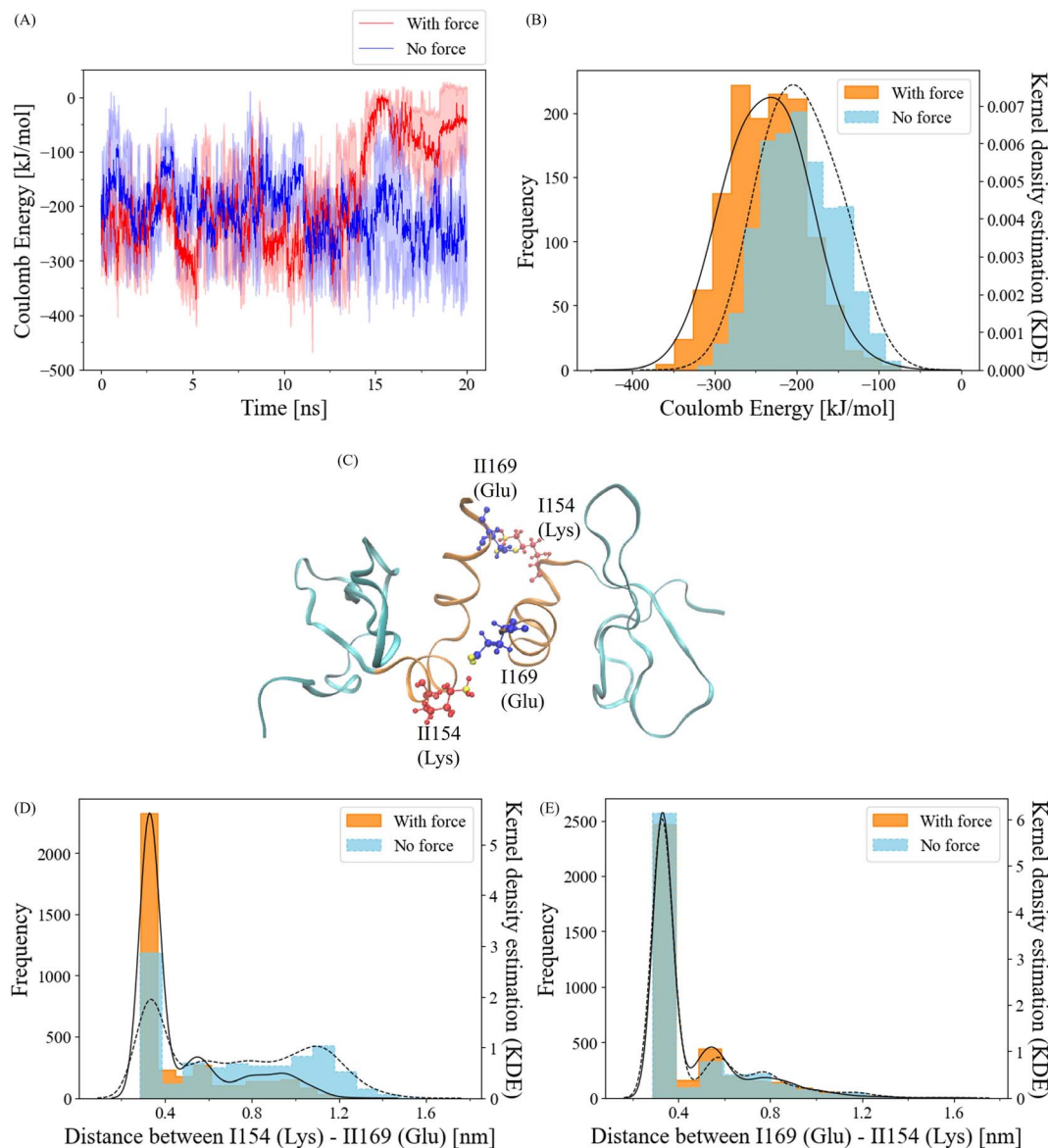
Frequency distributions of the distance between Glu CD and Lys NZ1 for  $^1\text{Lys154}$ – $^{11}\text{Glu169}$  and  $^1\text{Glu169}$ – $^{11}\text{Lys154}$  were determined (Fig. 5D and E). Prior to the 12.5 ns cutoff, the  $^1\text{Glu169}$ – $^{11}\text{Lys154}$  residue pair exhibited similar frequencies within the salt bridge distance cutoff of  $0.4 \text{ nm}^{46}$  under both force and no-force conditions (Fig. 5E). The salt bridge ratios with and without force were approximately 66% and 69%. In contrast, the  $^1\text{Lys154}$ – $^{11}\text{Glu169}$  residue pair exhibited a notably higher frequency within  $0.4 \text{ nm}$  with force compared to the no-force condition (Fig. 5D). While the salt bridge ratio without force was only 32%, it increased to 66% when force was applied. These findings strongly suggest that the  $^1\text{Lys154}$  and  $^{11}\text{Glu169}$  residue pair is responsible for the salt bridge interaction, which responds to the application of pulling force.

### Salt bridge residues exhibit elevated stress distribution

In our quest to elucidate the impact of pulling force on the mechanical characteristics of the  $\alpha$ -helix interfaces within CSD–CSD, we conducted a comprehensive Force Distribution Analysis (FDA) across these interfaces. The color map representing the force-applied condition displayed notable punctual stress concentrations within the  $^1\text{Lys154}$ – $^{11}\text{Glu169}$  and  $^1\text{Glu169}$ – $^{11}\text{Lys154}$  residue pairs prior to bond breakage. These punctual stresses on  $^1\text{Lys154}$ ,  $^{11}\text{Glu169}$ ,  $^1\text{Glu169}$ , and  $^{11}\text{Lys154}$  dramatically

without force before the breakage cutoff, derived from all six simulation runs ( $N = 3$  for each force/no-force condition).





**Fig. 5** (A) Coulombic energy changes in the  $\alpha$ -helix interface during the 20 ns simulation, with standard deviations represented by shaded areas.  $N = 3$  for each force and no-force condition. (B) Histogram depicting the frequency of the Coulomb energy between the  $\alpha$ -helix interface before the interaction breakage cutoff of 12.5 ns under force and no-force conditions, based on data from all six simulation runs ( $N = 3$  for each force/no-force condition). (C) Visual representation of CSD–CSD dimer with  $^{\text{I}}\text{Lys154}$ – $^{\text{II}}\text{Glu169}$  and  $^{\text{II}}\text{Glu169}$ – $^{\text{I}}\text{Lys154}$  salt bridge residue pairs. Lysine, a positively charged amino acid, is colored red, with the NZ atom in yellow. Glutamic acid, a negatively charged amino acid, is colored blue, with OE1 and OE2 atoms in yellow. (D and E) Histograms illustrating the frequency of salt bridge pair distance between (D)  $^{\text{I}}\text{Lys154}$ – $^{\text{II}}\text{Glu169}$  and (E)  $^{\text{II}}\text{Glu169}$ – $^{\text{I}}\text{Lys154}$  before the interaction breakage cutoff of 12.5 ns under force and no-force conditions, comprising data from all six simulation runs ( $N = 3$  for each force/no-force condition). The black solid lines in the histograms represent the Kernel Density Estimation (KDE) for the force-applied condition, while the black dashed lines correspond to the KDE for the no-force condition.

decreased due to the bond breakage between the  $\alpha$ -helix (Fig. 6). In contrast, the color map for the no-force condition exhibited consistent high stress levels within the  $^{\text{I}}\text{Glu169}$ – $^{\text{II}}\text{Lys154}$  residue pair throughout the entire simulation.

To quantitatively evaluate the differences in punctual stress between the force and no-force conditions, we calculated the average punctual stress for each residue, accompanied by standard deviations, focusing on data prior to the bond breakage cutoff at 12.5 ns (Fig. 6). Notably, both the  $^{\text{I}}\text{Glu169}$  and  $^{\text{II}}\text{Lys154}$  residues exhibited high punctual stresses in both force and no-

force conditions. However, the stresses on  $^{\text{I}}\text{Lys154}$  and  $^{\text{II}}\text{Glu169}$  residues under force were approximately  $500 \text{ kJ mol}^{-1} \text{ nm}^{-1}$  higher than those observed in the absence of force. These findings provide compelling support for the development of a salt bridge interaction between  $^{\text{I}}\text{Lys154}$  and  $^{\text{II}}\text{Glu169}$  under force conditions.

Furthermore, residue  $^{\text{II}}\text{Lys159}$  displayed stress levels approximately  $500 \text{ kJ mol}^{-1} \text{ nm}^{-1}$  higher when compared to the no-force condition (Fig. 6). Notably,  $^{\text{II}}\text{Lys159}$  engaged in an internal salt bridge interaction with  $^{\text{II}}\text{Glu118}$ , a residue located





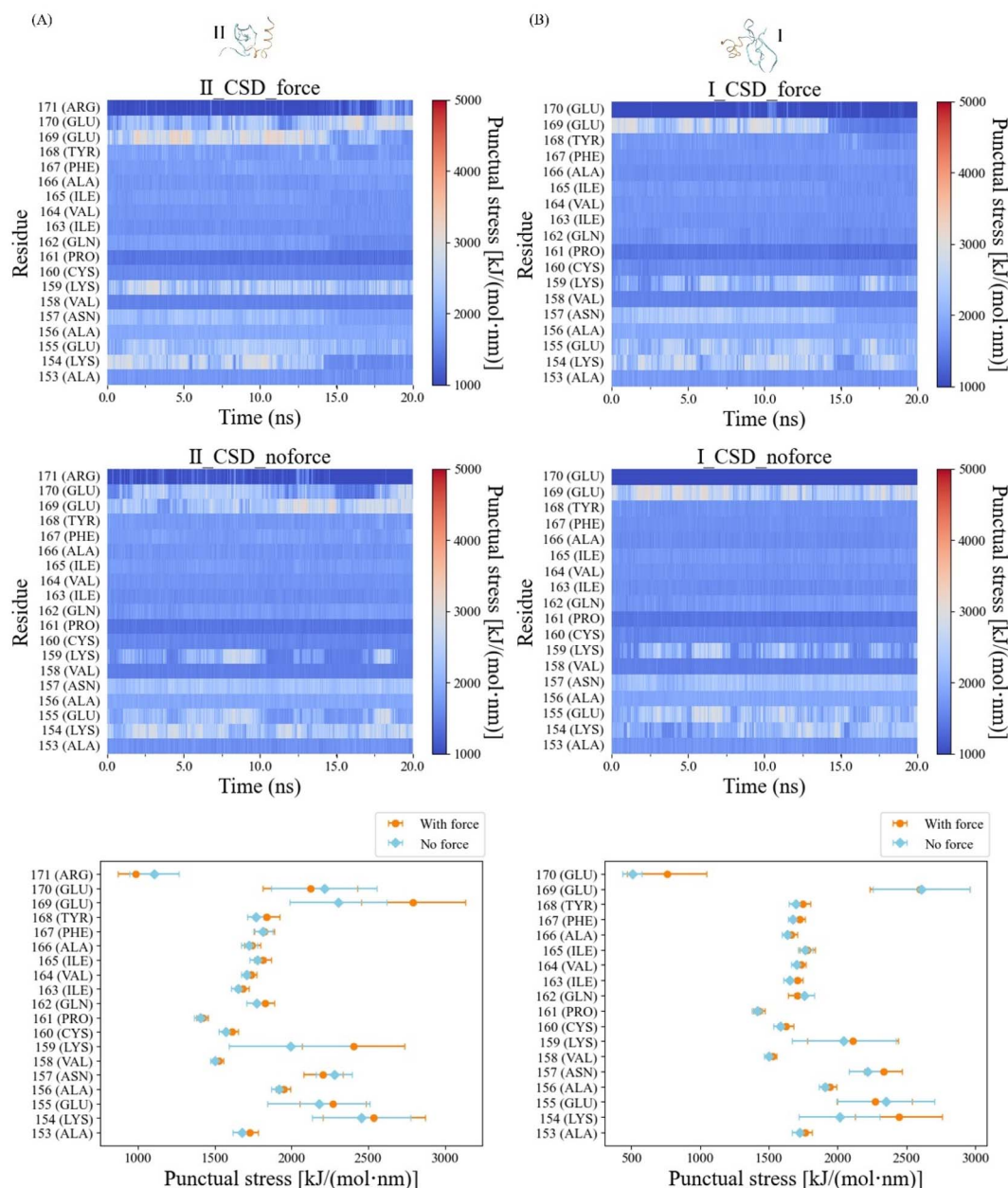


Fig. 6 Force distribution analysis for the (A) constrained  $^{11}\text{CSD}$  monomer (left) and (B) unconstrained  $^1\text{CSD}$  monomer (right) along the  $\alpha$ -helix interface with/without force over the 20 ns simulation. Punctual stress heatmaps were generated by averaging data from three simulations at each time step for both force and no-force conditions. The bar plots display the average punctual stress distributed across residues with standard deviations, for the data recorded before the bond breakage cutoff of 12.5 ns.

near the opposite edge of the same CSD monomer (ESI Fig. S3A<sup>†</sup>). The frequencies of salt bridge interactions within a 0.4 nm range in the force condition were significantly higher compared to those in the no-force condition. The salt bridge ratios, considering the presence or absence of force, were 67% and 9%, respectively (ESI Fig. S3B<sup>†</sup>).

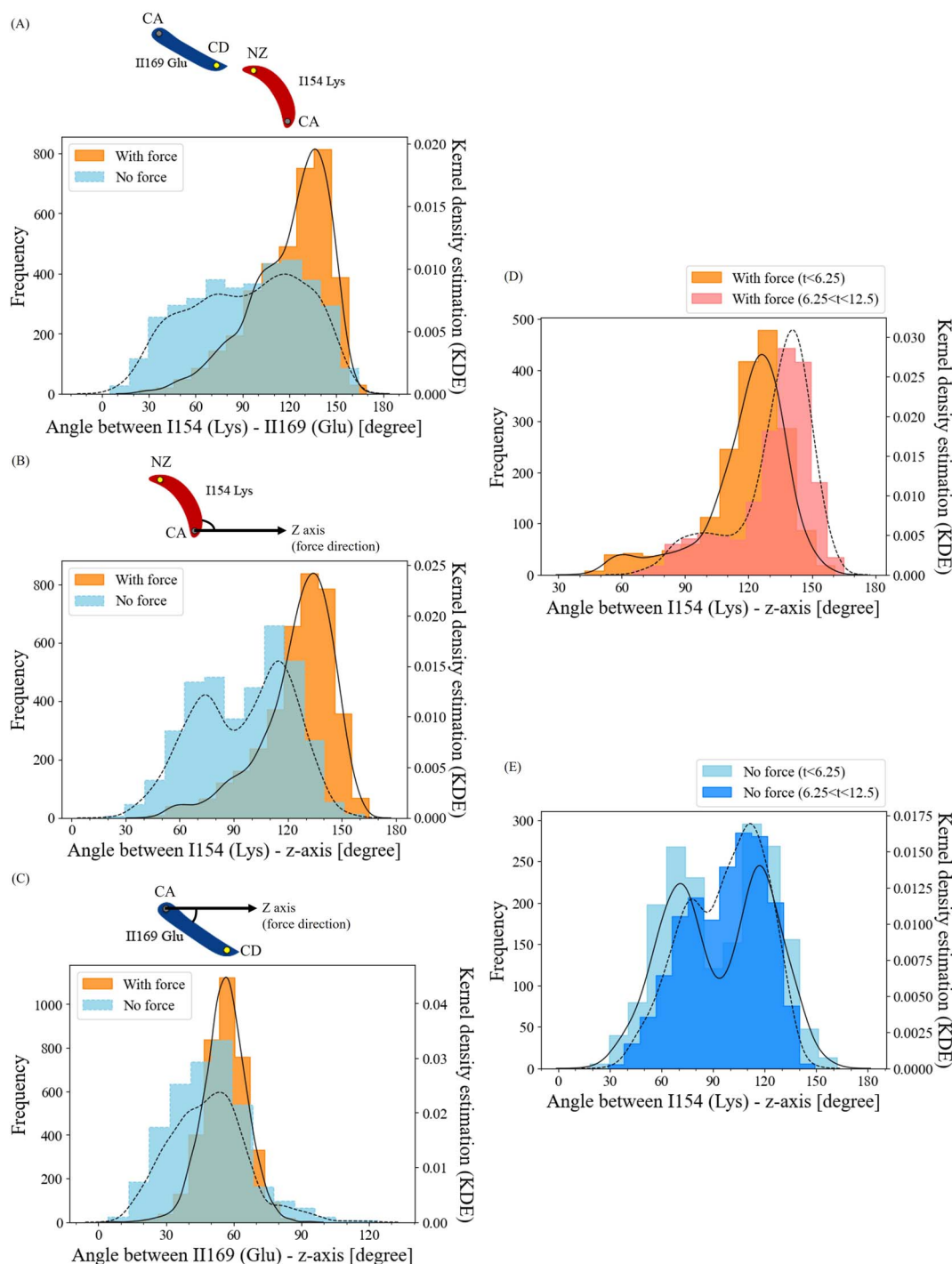
#### Force-induced angle change in residue $^1\text{Lys154}$ is the key to force-induced salt bridge interaction

To unravel how the application of force enables the  $^1\text{Lys154}$  and  $^{11}\text{Glu169}$  residue pair to form a salt bridge interaction, we

conducted an in-depth analysis of the structural alterations in this residue pair. We compared the angle frequencies between  $^1\text{Lys154}$  and  $^{11}\text{Glu169}$  with and without force (Fig. 7A). In the absence of force, the angles between these residues appeared to be randomly distributed. However, when force was applied, the angles became concentrated within the range of  $120^\circ$  to  $150^\circ$ .

To separately investigate the angle change behaviors of residues  $^1\text{Lys154}$  and  $^{11}\text{Glu169}$ , we established the z-axis of the pulling direction as a reference and calculated the angles of  $^1\text{Lys154}$  with respect to the z-axis and  $^{11}\text{Glu169}$  with respect to the z-axis (Fig. 7B and C). The angles between residue  $^{11}\text{Glu169}$  and the z-axis displayed relatively similar trends in both force





**Fig. 7** (A–C) Histograms depicting the frequency of angle measurements between (A) <sup>1</sup>Lys154–<sup>11</sup>Glu169, (B) <sup>1</sup>Lys154–z-axis, and (C) <sup>11</sup>Glu169–z-axis prior to the interaction breakage cutoff at 12.5 ns, recorded across all six simulations ( $N = 3$  for each force/no-force condition). On the schematic image above the histograms, lysine, a positively charged amino acid, is highlighted in red with atom NZ in yellow. Glutamic acid, a negatively charged amino acid, is depicted in blue with the carbon atom CD in yellow. NZ represents the nitrogen atom of the final heavy atom in lysine's sidechain, while CD corresponds to the delta carbon. The gray dot on the schematic image indicates  $\alpha$ -carbon. (D and E) Histograms illustrating the frequency of angle measurements for <sup>1</sup>Lys154–z-axis, divided into two time intervals (0–6.25 ns vs. 6.25–12.5 ns) under (D) force and (E) no-force conditions, based on data collected from all six simulations ( $N = 3$  for each force/no-force condition). Black solid and dash lines represent the Kernel Density Estimation of the force-applied and no-force conditions, respectively. The black solid lines in the histograms represent the Kernel Density Estimation (KDE) for the force-applied condition, while the black dashed lines correspond to the KDE for the no-force condition.

and no-force conditions (Fig. 7C). However, the angle frequencies of residue <sup>1</sup>Lys154 without force appeared to be relatively random, with two peaks around 70° and 110°. In contrast, when force was applied, the angles were highly concentrated in the range of 120° to 140° (Fig. 7B). Furthermore, this observation indicated that this particular residue exhibited greater flexibility over the course of the simulation, which aligns with the results obtained from the RMSF analysis. Collectively, these findings suggest that the angle change in residue <sup>1</sup>Lys154 is the key factor in facilitating the formation of the force-induced salt bridge.

To understand the dissociation process of the developed salt bridge interaction, we divided the angle frequencies between residue <sup>1</sup>Lys154 and the z-axis, considering the presence or absence of force. This division was carried out over two time intervals (0–6.25 ns vs. 6.25–12.5 ns) (Fig. 7D and E). The angles with high frequencies in the no-force condition exhibited similar patterns across both time intervals, with average angles of 93.8° for 0–6.25 ns and 96.8° for 6.25–12.5 ns (Fig. 7E). In contrast, for the force-applied condition, the peak of the angle distribution notably shifted towards higher angles over time (Fig. 7D), with the average angles being 118.7° during the 0–6.25 ns interval and increasing to 131.4° in the 6.25–12.5 ns interval. This shift towards higher angles over time indicates the disruption of the salt bridge interaction corresponded to an increase in the angle of residue <sup>1</sup>Lys154.

## Discussion

The role of Heterochromatin Protein 1 $\alpha$  (HP1 $\alpha$ ) in nuclear and chromatin mechanics has been extensively studied in prior experiments involving force application.<sup>18,20</sup> HP1 has been implicated in the formation of chromatin crosslinks and phase separation, mechanisms that are thought to provide structural rigidity to chromatin and the cell nucleus.<sup>18,25</sup> A fundamental aspect of HP1's mechanical functionality is its homodimerization, primarily facilitated through the chromoshadow domain (CSD), which is crucial for establishing crosslinks between HP1 and chromatin, as well as for promoting phase separation.<sup>18,25,26,30–33</sup> To gain a deeper understanding of the molecular intricacies of how HP1 $\alpha$  dimerization responds to applied mechanical forces, this study focused on the CSD–CSD dimer of HP1 $\alpha$ . Using full atomic molecular dynamics simulations, we applied a pulling force to this dimeric structure. While the overall state of hydrogen bonding across the  $\alpha$ -helix interface of the CSD–CSD dimer remained relatively stable (Fig. 4), a novel salt bridge interaction emerged as a response to the applied force, leading to the reinforcement of the  $\alpha$ -helix interface interactions (Fig. 5 and 6). This force-induced salt bridge formation was attributed to changes in the orientation of residue <sup>1</sup>Lys154 on the  $\alpha$ -helix interface (see Fig. 7). These findings emphasize the sensitivity of HP1 $\alpha$  homodimerization to mechanical forces and offer insights into potential mechanisms underlying the strengthening of HP1 $\alpha$ –DNA interactions, as previously observed in experimental studies.<sup>20</sup>

The RMSD results reveal that the  $\alpha$ -helix interface of CSD–CSD remains notably more stable under the influence of an applied force compared to the condition without force,

particularly before the bond breakage occurs (Fig. 3A). This heightened stability can be attributed to the development of salt bridge interactions, specifically the <sup>1</sup>Lys154–<sup>11</sup>Glu169 and <sup>11</sup>Lys159–<sup>11</sup>Glu118 pairs, which fortify the  $\alpha$ -helix interface when force is applied. In particular, the RMSF values for <sup>1</sup>Lys154 in the force-applied scenario are significantly lower before the bond breakage, in comparison to the scenario without force (Fig. 3B). Furthermore, the residues <sup>1</sup>Asn157 and <sup>11</sup>Gln162 also exhibit reduced RMSF values under force when contrasted with the no-force condition. Although the overall properties of hydrogen bonds remain relatively unaffected by the applied force, it is noteworthy that the lifetime of the <sup>11</sup>Gln162–<sup>1</sup>Asn157 H-bond pair in the force scenario shows a prolonged trend as compared to the condition without force (ESI Fig. S4†). This trend may be attributed to the positioning of the <sup>11</sup>Gln162–<sup>1</sup>Asn157 residue pair in the force scenario, where it resides between the Lys154–Glu169 salt bridge interactions, thus contributing to the local stabilization of hydrogen bonds and bolstering the structural integrity of the  $\alpha$ -helix interface within CSD–CSD in the presence of force.

Various types of interactions, including hydrogen bonds (H-bonds), van der Waals (vdW) forces, and salt bridge interactions, contribute to the intricate network of molecular interactions within proteins. Notably, salt bridge interactions are widely recognized as one of the most robust forms of residue–residue interactions.<sup>47</sup> It is worth noting that the strength of salt bridge interactions can be influenced by a range of environmental factors, such as solvation, ion concentrations, and the spatial arrangement of charged residues on the protein's surface.<sup>47–49</sup> Moreover, it has been suggested that certain MD simulation force fields, including CHARMM, may overestimate the strength of salt bridges compared to experimental data.<sup>50,51</sup> The choice of force field can quantitatively influence the perceived strengths of these interactions.<sup>50</sup> In our study, we scrutinized the coulombic energies of the  $\alpha$ -helix interface, revealing an average value of approximately  $-236 \text{ kJ mol}^{-1}$  when force was applied, in contrast to  $-196 \text{ kJ mol}^{-1}$  in the absence of force. Quantum chemistry analyses have previously estimated that salt bridge interactions between a Glu and a Lys residue pair in an aqueous environment typically have an interaction energy of around  $-40 \text{ kJ mol}^{-1}$ .<sup>47</sup> This analysis provides compelling evidence that the observed difference in energy between the force-applied and no-force conditions primarily arises from the development of Glu–Lys salt bridge interactions. These findings align with previous observations of force-induced Glu–Lys salt bridges in actin catch–slip bonds, shedding light on a potential mechanosensing mechanism governing force-dependent actin dynamics.<sup>52</sup> Similarly, the identification of a force-induced Glu–Lys salt bridge in HP1 $\alpha$ , as demonstrated in this study, suggests a plausible mechanosensing mechanism for governing force-dependent chromatin dynamics. These insights into the mechanical behaviors of salt bridges offer valuable clues as to how external forces may influence the structural and functional aspects of proteins, with potential implications for chromatin organization in response to mechanical cues.



The formation of the force-induced Glu–Lys salt bridge hinged on the dynamic shifts in the orientation of residue <sup>1</sup>Lys154 (Fig. 7). Under the influence of the pulling force, residue <sup>1</sup>Lys154 underwent a tilting motion towards its neighboring residue <sup>12</sup>Glu169. This alteration in position brought the positively charged atom NZ1 in residue <sup>1</sup>Lys154 within close proximity to the negatively charged atoms OE1 and OE2 in residue <sup>12</sup>Glu169. As the force was consistently applied, the angle of residue <sup>1</sup>Lys154 continued to increase in a direction opposing the applied force, ultimately leading to the disruption of the  $\alpha$ -helix interface within the CSD–CSD dimer. It is worth noting that alterations in the angles and positions of lysine residues can have a significant impact on chemical interactions, a subject that has been explored in previous research.<sup>49,53</sup> A more comprehensive understanding of the structural and chemical changes in lysine under external mechanical forces can offer a more nuanced interpretation of the development of lysine salt bridges from a chemical perspective. In addition to the detailed analysis of residue <sup>1</sup>Lys154 and <sup>12</sup>Glu169, we conducted an extensive structural assessment (ESI Fig. S5†). This evaluation encompassed the measurement of angles between the two  $\alpha$ -helices, as well as the lengths of the inter-residue regions <sup>1</sup>Lys154–<sup>12</sup>Glu169 and <sup>12</sup>Lys154–<sup>12</sup>Glu169 (ESI Fig. S5†). The outcomes of this structural analysis revealed that the trends in angles and lengths were similar between the force-applied and no-force conditions, underscoring the critical role of residue <sup>1</sup>Lys154 in the mechanical response of the  $\alpha$ -helix interface within the CSD–CSD dimer. Furthermore, we extended our analysis to encompass the angles of <sup>12</sup>Glu169–<sup>12</sup>Lys154, the angles with respect to the z-axes of <sup>12</sup>Glu169 and <sup>12</sup>Lys154, which were scrutinized in the same manner as residue <sup>1</sup>Lys154 and <sup>12</sup>Glu169 (ESI Fig. S6A†). The resulting data indicated that, while there were subtle variations in the height and location of the peaks in the angle distributions between the force-applied and no-force conditions, these distinctions were not substantial (ESI Fig. S6B and C†). Collectively, these results highlight the pivotal contribution of residue <sup>1</sup>Lys154 to the mechanical response of the  $\alpha$ -helix interface within the CSD–CSD dimer, shedding light on the underlying molecular mechanisms at play.

In our FDA analysis, punctual stress derived from coulombic interactions was obtained throughout the  $\alpha$ -helix interface of the CSD–CSD to assess the mechanistic impacts of the <sup>1</sup>Lys154–<sup>12</sup>Glu169 salt bridge. We observed that while the punctual stress resulting from Lennard–Jones interactions was lower compared to that based on coulombic interactions, it exhibited similar trends (ESI Fig. S7†). Our FDA analysis of force distribution patterns revealed elevated stress levels not only in the residue <sup>1</sup>Lys154–<sup>12</sup>Glu169 under the force-applied condition but also in residue <sup>12</sup>Lys159 when compared to their respective counterparts in the absence of force, as illustrated in Fig. 6. The heightened stress experienced by residue <sup>12</sup>Lys159 was attributed to the development of a salt bridge with residue <sup>12</sup>Glu118 within the same CSD monomer, as demonstrated in ESI Fig. S3.† This internal salt bridge may indeed serve as a significant contributing factor to the overall mechanical resilience of the CSD–CSD dimer when subjected to external pulling forces. It is important to note that this study primarily focused on the

$\alpha$ -helix interface within the CSD–CSD dimer and, therefore, employed a center-of-mass pulling force approach. To gain deeper insights and to assess the realistic occurrence and mechanistic roles of the internal salt bridge, an edge–edge pulling force simulation is deemed necessary. Such an investigation would offer a comprehensive perspective on the entirety of structural changes within the CSD–CSD dimer and illuminate the functional significance of this internal salt bridge under varying mechanical conditions.

To provide a comprehensive analysis, positional constraints were employed in a manner similar to the conditions applied under force for a 20 ns relaxation simulation within the no-force scenario. The examination of angles involving <sup>1</sup>Lys154–<sup>12</sup>Glu169, <sup>1</sup>Lys154–z-axis, and the assessment of punctual stress in these residues occasionally indicated that the salt bridge interaction could potentially occur even without the presence of an external force (see Fig. 5–7). This observation raises the possibility that these findings could be influenced by constraints imposed during the simulation. In molecular dynamics simulations, atoms inherently exhibit stochastic motion in response to the surrounding solvent. In the context of our constrained conditions, one CSD monomer was allowed to move freely, while the movement of the other CSD monomer was restricted. This inherent stochastic nature may occasionally result in the free CSD monomer drifting significantly apart from the constrained CSD monomer, thereby generating conditions akin to pseudo-force effects. To explore this further, we conducted a no-force condition simulation without any position constraints and measured the distance between <sup>1</sup>Lys154 and <sup>12</sup>Glu169 (ESI Fig. S8†). The frequency distribution of distances between <sup>1</sup>Lys154 and <sup>12</sup>Glu169 within the 0.4 nm range was notably lower compared to scenarios involving force application or constraints. This no-constraint data lends support to the notion that the natural occurrence of the <sup>1</sup>Lys154–<sup>12</sup>Glu169 salt bridge interaction is unlikely and underscores the sensitivity of this interaction to external forces.

Chromatin fibers within the nucleus exhibit dynamic and multidirectional movements, subjecting HP1 to forces originating from various orientations. However, when we explored the effects of alternative directional forces, we observed that the CSD–CSD dimer underwent rotation and separation, as presented in this study, or experienced significantly longer simulation times, although this specific data is not shown. This suggests that bond breakage induced by forces acting along different directions may be less likely to occur.

Additionally, it is worth noting that the time rate of the applied force can vary depending on the method of force application to the cellular system. The effects of different pulling rates of the applied force on the mechanosensitive salt bridge formation were investigated (ESI Fig. S9†). In addition to 0.05 nm ns<sup>−1</sup> used in our simulation, 0.03 nm ns<sup>−1</sup>, 0.1 nm ns<sup>−1</sup>, and 0.5 nm ns<sup>−1</sup> pulling rates were chosen. The slower pulling rate of 0.03 nm ns<sup>−1</sup> showed a mechano-sensitivity comparable to that observed at the 0.05 nm ns<sup>−1</sup>. The salt bridge ratio of the 0.03 nm ns<sup>−1</sup> simulation was 65%, almost the same as 66% of the salt bridge ratio in the 0.05 nm ns<sup>−1</sup> simulation. Conversely, the salt bridge ratio decreased by 46% in the faster 0.1 nm ns<sup>−1</sup> pulling rate





simulation. The fastest rate of  $0.5 \text{ nm ns}^{-1}$  showed further reduction in the mechano-sensitivity for the  $^1\text{Lys154}$ – $^{11}\text{Glu169}$  salt bridge formation (ESI Fig. S9C†). The salt bridge ratio of the  $0.5 \text{ nm ns}^{-1}$  simulation was only 30%, closely mirroring the 32% observed in the no-force condition. However, such faster pulling rates would less likely occur in natural cellular system. For instance, in a prior experimental study, we endeavored to replicate conditions akin to vasoconstriction.<sup>17</sup> Considering our results of around 0.02 intranuclear strain with 0.5 Hz and assuming 15  $\mu\text{m}$  of the nucleus size,<sup>17,54</sup> this calculation translates to a velocity of approximately  $0.015 \times 10^{-5} \text{ nm ns}^{-1}$ . Notably, this velocity is considerably slower than the rate employed in our simulation, which stands at  $0.05 \text{ nm ns}^{-1}$ . Nonetheless, conducting simulations with the experimentally observed velocity remains a challenging endeavor due to computational constraints and limitations in calculation speed.

The dimerization of HP1 $\alpha$ , which links chromatin fibers together, is a fundamental process crucial for bolstering the mechanical integrity of both chromatin and the cell nucleus.<sup>18</sup> This mechanical fortification is further facilitated through the enhancement of the  $\alpha$ -helix interface of the CSD–CSD dimer, achieved by the formation of salt bridges under force conditions. This salt bridge development represents the likely mechanism by which the HP1 $\alpha$  dimer is reinforced, subsequently enhancing the resilience of chromatin against mechanical forces.

The development of force-induced salt bridges emerges as a determining factor in achieving stable HP1 $\alpha$ –DNA condensations. This observation aligns with the findings of a prior experimental study that applied force to HP1 $\alpha$ .<sup>20</sup> Furthermore, in the context of HP1 and chromatin phase separation, a recent coarse-grained computational investigation examined the impact of HP1 dimer–dimer interactions on phase separation.<sup>55</sup> By modulating the energy associated with these interactions, particularly in the hinge and NTE regions, with values ranging from 2kBT to 4kBT, 6kBT, and 10kBT (equivalent to approximately 5, 10, 15, and 25  $\text{kJ mol}^{-1}$  at a standard temperature of 300 K), the coarse-grained model delineated four distinct phases: a mixed state with no phase separation, liquid-like droplets, droplets with incorporated polymer, and polymer-coated regimes.<sup>55</sup> In this context, the energy derived from Lys–Glu salt bridge interactions, as indicated by quantum analysis, and the electrostatic energy difference between force and no-force conditions, which amounted to approximately 40  $\text{kJ mol}^{-1}$ ,<sup>47</sup> implies that the force-induced salt bridge formation may yield ample energy to influence the state of phase separation.

The enhanced interaction within CSD–CSD under force conditions offers valuable insights, shedding light on the counterexample of HP1 phase separation preference. Typically, phase separation occurs in regions characterized by lower mechanical energy, softness, and lower density, such as euchromatin.<sup>29</sup> However, HP1-induced phase separation predominantly takes place in heterochromatin regions. The mechanical sensitivity exhibited by the CSD–CSD interaction, driven by force-induced salt bridge development, may enable HP1 to initiate phase separation in regions with higher mechanical energy, stiffness, and greater density, such as

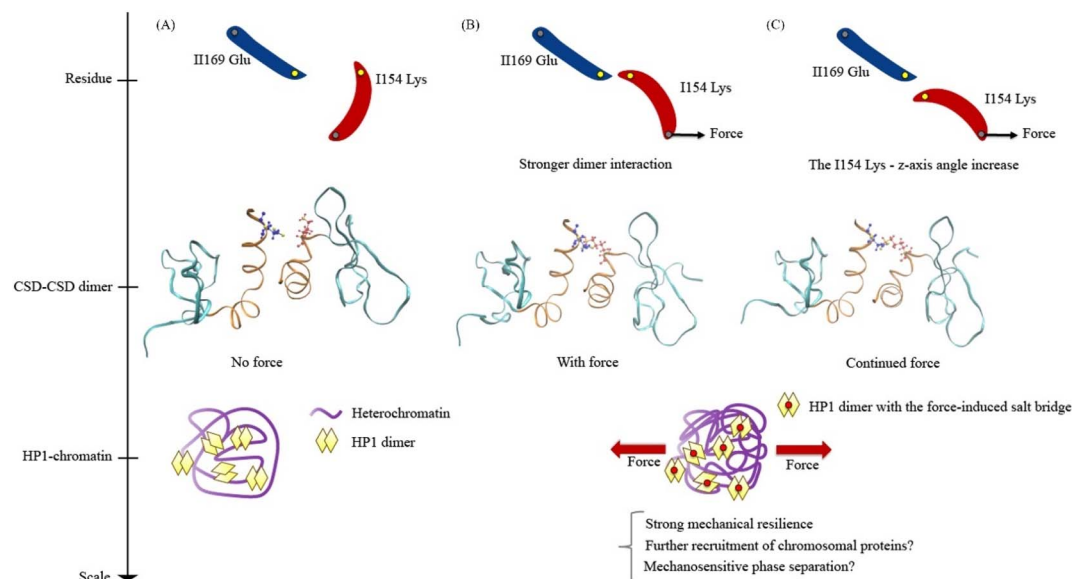
heterochromatin. The mechanical response of the CSD–CSD dimer, as elucidated in this study, implies that HP1 $\alpha$ –chromatin phase separation is mechanosensitive, making the salt bridge development a likely mechanical switch governing chromatin condensation state. This underscores the significance of conducting molecular-scale analyses of HP1 $\alpha$  mechanics.

Our simulation study has certain limitations, including the influence of positional constraints, the pulling rate, as discussed earlier, and an incomplete understanding of the significance of the force-induced salt bridge behavior in the overall HP1 structure. The mechanical force exerted by chromatin is expected to be transmitted from the chromodomain (CD) to the chromoshadow domain (CSD) through the disordered hinge region. The flexibility of the hinge region may, to some extent, dampen the transmission of force from CD to CSD. The application of homology modeling could enable the replication of CD–hinge–CSD protein regions and facilitate more precise simulations of force transmission from chromatin to CSD. Additionally, our study did not consider other protein interactions. Notably, proteins like Shugoshin (Sgo1) and the lamin B receptor (LBR) interact with the CSD dimer, exerting an influence on phase separation states.<sup>26</sup> To gain deeper insights, simulating more complex and higher-order HP1 structures, related proteins, and DNA systems is necessary, providing a comprehensive understanding of HP1-driven chromatin mechanics.

Lastly, we propose models for understanding the mechanical response of HP1 $\alpha$ , encompassing the residue scale, the CSD–CSD dimer scale, and the HP1–chromatin interaction scale (Fig. 8). In a force-absent condition, the  $^1\text{Lys154}$  angle is less likely to facilitate the formation of the  $^1\text{Lys154}$ – $^{11}\text{Glu169}$  salt bridge interaction (Fig. 8A). Consequently, the mechanical resilience of the HP1 $\alpha$  is relatively low without force. However, in the presence of applied force, the pulling force induces an alteration in the  $^1\text{Lys154}$  angle, leading to the development of the  $^1\text{Lys154}$ – $^{11}\text{Glu169}$  salt bridge interaction (Fig. 8B). As a result, the CSD–CSD dimer exhibits a stronger interaction and enhanced mechanical resilience against force, potentially allowing for the recruitment of more heterochromatin proteins and the promotion of phase separation. It is conceivable that the heterochromatin region, characterized by condensed chromatin, serves as a site for transmitting force to HP1 $\alpha$ , which may prefer a mechanically robust environment found in heterochromatin. During the dissociation process under force, the interaction breaks as the  $^1\text{Lys154}$  angle increases against the applied force direction (Fig. 8C). Continuous external pulling force disrupts the  $^1\text{Lys154}$ – $^{11}\text{Glu169}$  salt bridge interaction, ultimately leading to a high level of force-induced mechanical resilience.

Looking ahead, our future investigations will delve into the intricate dynamics of HP1–chromatin interactions, exploring their pivotal role in shaping HP1 and chromatin mechanics, as well as the state of phase separation.<sup>22,55</sup> Moreover, it is worth noting that HP1 phase separation itself is suggested to hold key mechanical functions, which extend beyond its chromatin interaction or dimerization aspects.<sup>25</sup> To comprehensively unravel the complex web of HP1 and chromatin mechanics,





**Fig. 8** Models of HP1 $\alpha$  mechanical response at residue, the CSD–CSD dimer, and HP1 $\alpha$ –chromatin scales, presented from top to bottom: (A) no-force condition: the <sup>1</sup>Lys154–<sup>11</sup>Glu169 salt bridge interaction is less likely to occur, resulting in low mechanical resilience for HP1. (B) Force condition: under applied force, the <sup>1</sup>Lys154 angle changes, leading to the development of the <sup>1</sup>Lys154–<sup>11</sup>Glu169 salt bridge interaction. This strengthens the interaction within the CSD–CSD dimer, enhancing the mechanical resilience of the HP1 $\alpha$ . (C) Dissociation processes: continuous external force disrupts the <sup>1</sup>Lys154–<sup>11</sup>Glu169 salt bridge interaction by increasing the <sup>1</sup>Lys154 angle against the applied force direction. Consequently, high mechanical resilience is compromised.

including the molecular-level mechanical behavior of the CSD–CSD dimer, we plan to conduct simulations involving the CD region interacting with H3K9me, the entire HP1 structure, and a comprehensive HP1–chromatin coarse-grained model. These efforts will provide a deeper understanding of how HP1 and chromatin's mechanical interplay is regulated and how it impacts intranuclear structure and mechanics.

## Conclusions

In our investigation, we harnessed full atomistic molecular dynamics simulations to unravel the intricate molecular mechanics governing Heterochromatin Protein 1 $\alpha$  (HP1 $\alpha$ ) when subjected to mechanical forces. Our primary focus was on the chromoshadow domain (CSD) within the HP1 $\alpha$  dimer, given its pivotal role in facilitating chromatin crosslinks and phase separation. Employing a constant velocity force, we subjected the CSD–CSD dimer to a comparative analysis between force-applied and force-free conditions. Notably, we observed a heightened attractive coulombic energy between the  $\alpha$ -helix interfaces under force, particularly before bond breakage, in comparison to the no-force scenario (Fig. 5A and B). This finding was further supported by the increased frequencies of distances between <sup>1</sup>Lys154 and <sup>11</sup>Glu169, potential salt bridge pairs, when force was applied (Fig. 5D). Additionally, force distribution analysis across the  $\alpha$ -helix interfaces indicated greater stress on <sup>1</sup>Lys154 and <sup>11</sup>Glu169 in the presence of force (Fig. 6). Through an in-depth structural analysis of these  $\alpha$ -helix interfaces, we unraveled the mechanism by which applied force induced a salt bridge formation between <sup>1</sup>Lys154 and <sup>11</sup>Glu169, driven by changes in the angle of <sup>1</sup>Lys154 (Fig. 7A–C).

Our molecular mechanistic analysis of the HP1 $\alpha$  CSD–CSD dimer unveiled a remarkable response, with <sup>1</sup>Lys154 altering its angle in reaction to mechanical force and fostering a salt bridge interaction with <sup>11</sup>Glu169. This novel insight into the strengthening of HP1 $\alpha$  CSD–CSD dimerization promises to enhance our understanding of HP1 $\alpha$  phase separation within heterochromatin regions. Furthermore, it sheds light on the heightened resilience and condensation of HP1 $\alpha$ –DNA in the presence of mechanical forces. In essence, this study underscores the potentially critical role of enhanced HP1 $\alpha$  CSD–CSD dimer interactions when force is applied, offering valuable insights into the molecular foundation of HP1 $\alpha$  dimerization, mechanosensitive phase separation, chromatin condensation and interactions, and ultimately, mechano-genomic regulation.

## Author contributions

S. T., M. K., N. D., and M. R. K. M. designed research. S. T. performed all simulations and analyses and wrote the manuscript. M. K., N. D., and M. R. K. M. reviewed and modified the manuscript.

## Conflicts of interest

Shingo Tsukamoto, Mohammad Khavani, Nya Domkam, and Mohammad R. K. Mofrad declare that they have no conflicts of interest.

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