Key role of the REC lobe during CRISPR–Cas9 activation by ‘sensing’, ‘regulating’, and ‘locking’ the catalytic HNH domain

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Abstract

Understanding the conformational dynamics of CRISPR (clustered regularly interspaced short palindromic repeat)–Cas9 is of the utmost importance for improving its genome editing capability. Here, molecular dynamics simulations performed using Anton–2—a specialized supercomputer capturing micro-to-millisecond biophysical events in real time and at atomic-level resolution—reveal the activation process of the endonuclease Cas9 toward DNA cleavage. Over the unbiased simulation, we observe that the spontaneous approach of the catalytic domain HNH to the DNA cleavage site is accompanied by a remarkable structural remodeling of the recognition (REC) lobe, which exerts a key role for DNA cleavage. Specifically, the significant conformational changes and the collective conformational dynamics of the REC lobe indicate a mechanism by which the REC1–3 regions ‘sense’ nucleic acids, ‘regulate’ the HNH conformational transition, and ultimately ‘lock’ the HNH domain at the cleavage site, contributing to its catalytic competence. By integrating additional independent simulations and existing experimental data, we provide a solid validation of the activated HNH conformation, which had been so far poorly characterized, and we deliver a comprehensive understanding of the role of REC1–3 in the activation process. Considering the importance of the REC lobe in the specificity of Cas9, this study poses the basis for fully understanding how the REC components control the cleavage of off-target sequences, laying the foundation for future engineering efforts toward improved genome editing.

Introduction

CRISPR (clustered regularly interspaced short palindromic repeat)–Cas9 is a bacterial adaptive immune system, which has revolutionized life sciences through the introduction of a facile genome editing technology (Doudna and Charpentier, 2014; Jinek et al., 2012). In this system, the endonuclease Cas9 can be programmed with single-guide RNAs to site-specifically recognize and cleave any DNA sequence bearing a protospacer adjacent motif (PAM) sequence, which serves as a key recognition element across the genome. This enables genetic engineering of biological systems with unprecedented efficiency, resulting in transformative applications in the life sciences, including the fields of medicine and biotechnology.

During CRISPR–Cas9 activation, the DNA binds Cas9 by matching the guide RNA with one strand (the target strand, TS) and forming an RNA:DNA hybrid, while the non-target strand (NTS) is displaced. Two nuclease domains, HNH and RuvC, catalyze the cleavage of the TS and NTS, respectively. The Cas9 protein comprises a recognition (REC) lobe, which mediates the nucleic acid binding through three recognition domains (REC1–3), and a nuclease lobe including the RuvC and HNH catalytic cores (Fig. 1). At the protein C-terminus, a domain structurally similar to type II topoisomerase constitutes the PAM interacting (PI) region (Jiang and Doudna, 2017; Nishimasu and Nureki, 2017). Structural and biophysical studies have revealed that relevant conformational changes occur upon binding of the nucleic acids (Chen and Doudna, 2017). Specifically, RNA binding primes the protein for subsequent DNA binding (Jiang et al., 2015), while the dynamics of the HNH exert conformational control over Cas9 nuclease activity (Sternberg et al., 2015). Upon DNA binding, the HNH domain undergoes a structural transition from an inactivated state, in which the catalysis is hampered,
to an activated state prone for the cleavage of the TS. While the inactivated state has been well characterized via X-ray crystallography (Anders et al., 2014; Nishimatsu et al., 2014), high-resolution data for the activated state are missing. Indeed, the most complete X-ray structure of Cas9 from Streptococcus pyogenes in complex with the nucleic acids captured a pre-activated state of the system, with the HNH catalytic H840 located ∼19 Å from the cleavage site on the TS (Fig. 1b) (Jiang et al., 2016). Clearly, the activation of the system toward catalysis requires a further conformational change of the HNH domain (Fig. 1b, lower panel). Very recently, a novel cryo-EM structure has been reported at a 5.2 Å resolution (EMD-8236) (Huai et al., 2017), in which the HNH domain approaches REC1. The difficulty in experimentally capturing the active conformation of CRISPR–Cas9 and the associated conformational transitions reflects the ‘striking flexibility’ of the protein, as arising by the interplay with the nucleic acids, during recognition, association, and cleavage (Palermo et al., 2016; Sternberg et al., 2015). As a support to the structural characterizations, extensive experimental efforts, including Forster Resonance Energy Transfer (FRET) and structural comparisons with homologous systems, identified the conformational requirements of a catalytically competent Cas9 (Chen and Doudna, 2017). In an initial study, Sternberg et al. used bulk FRET to reveal that the conformational dynamics of the HNH domain controls DNA cleavage (Sternberg et al., 2015). Subsequently, single-molecule FRET (smFRET) experiments characterized the conformational features of the activated HNH docked at the cleavage site (Dagdas et al., 2017), and also revealed that the high flexibility of the REC lobe facilitates the activation of the HNH domain (Chen et al., 2017; Osuka et al., 2018). Moreover, computational studies have contributed in understanding the conformational dynamics of HNH from the apo form of Cas9 up to the DNA-bound state (Huai et al., 2017; Palermo et al., 2017a, 2017b; Zuo and Liu, 2017). However, the detailed conformational rearrangements leading up to the catalytically active Cas9 protein have not been clarified. Specifically, it is unclear how HNH would dock at the cleavage site and, importantly, how the REC domains would facilitate this process mechanistically. Understanding the role of the REC lobe in the activation of Cas9 is of key importance for improving the system toward controlled functionality. Indeed, mutations within the REC lobe have been shown to reduce off-target cleavage events in newly evolved CRISPR–Cas9 systems (Casini et al., 2018; Chen et al., 2017; Kleinstiver et al., 2016).

Here, multi-microsecond length molecular dynamics (MD) simulations reveal the activation process in ∼16 µs of continuous simulation. Remarkably, the active conformation reached via MD simulations matches the structural and conformational transitions indicated by smFRET, while also being in accord with the newly available cryo-EM data. We show that the transition of the HNH domain depends on the structural remodeling of the REC domains, and is driven by favorable interactions with the REC lobe that form ‘on the fly’ during MD, eventually leading to the stable docking of HNH at the cleavage site. The observed conformational changes of the REC components pinpoint on an atomic scale how the recognition domains REC1–3 ‘sense’ nucleic acids, ‘regulate’ the HNH conformational change, and ultimately ‘lock’ HNH at the cleavage site, contributing to its catalytic activation. Finally, tight coupling between the REC lobe and the HNH domain is observed upon activation, ultimately ensuring catalytic competence (Chen et al., 2017; Dagdas et al., 2017).

Results

Conformational transition of the HNH domain

Several studies have been conducted to understand the kinetics of the HNH conformational change, revealing a slow overall conformational transition (i.e. from the inactivated to activated state over milliseconds to seconds) (Raper et al., 2018; Shibata et al., 2017; Singh et al., 2016). However, the final conformational adjustment of the pre-activated crystal structure (Fig. 1) could rapidly occur (i.e. within micro-to-milliseconds), leading to an activated Cas9 (Jiang et al., 2016). Due to the high flexibility of the HNH domain (Sternberg et al., 2015), the precise determination of its conformational change and the associated kinetics has been limited. In order to test the hypothesis of a fast sub-millisecond conformational change, in our previous study, we have carried out MD simulations of the pre-activated CRISPR–Cas9 (5F9R.pdb) (Jiang et al., 2016) in an enhanced sampling regime (Palermo et al., 2017a). We
employed a Gaussian-accelerated MD (GaMD) approach, which can access conformational states of proteins and nucleic acids over milliseconds (and in some cases beyond) by running much shorter simulations (i.e. of hundreds of nanoseconds) (Miao and McCammon, 2016a, 2018; Miao et al., 2015). During ∼400 ns of GaMD, the catalytic H840 approached the scissile phosphate on the TS (i.e. phosphate −3) at a distance of ∼15.0 Å, from its original location at ∼19.4 Å (Fig. S1). Although the catalytic domain remained beyond the range required for catalysis, these simulations have highlighted the tendency for a fast conformational change of HNH in the late step of activation, in agreement with experiments (Dagdas et al., 2017; Shibata et al., 2017; Sternberg et al., 2015). Encouraged by these outcomes, we here performed continuous simulations with the aim of capturing the activation process of the HNH domain in real time and on an atomic scale. Noteworthy, recent studies have employed enhanced sampling methods to access kinetic information (Stelzl and Hummer, 2017), while the efficiency of accelerated MD methodologies in exploring the conformational space has been used in conjunction with Markov models to construct solid kinetic models (Paul et al., 2017). However, while enhanced sampling MD describes well thermodynamic properties and conformational ensembles, it does not directly provide kinetic information, the latter is however preserved via unbiased simulations (Abrams and Bussi, 2014; Miao and McCammon, 2016b).

In order to obtain a continuous MD trajectory encompassing the microsecond time scale, we carried out MD on a specialized supercomputer – Anton-2 – that enables for micro-to-millisecond length simulations (Shaw et al., 2014). CRISPR–Cas9 was simulated for ∼16 µs revealing that the HNH domain approaches the cleavage site on the NTS after ∼7 µs of MD and stably reaches a catalytically competent state after ∼10 µs of MD (Fig. 2a–b).

![Fig. 2.](https://www.cambridge.org/core/coreflo/5F9R.pdb, starting configuration for MD) and in the model structure obtained via cryo-EM fitting of the EMD-8236 map (5Y36.pdb) (Huai et al., 2017; Jiang et al., 2016). Transparent boxes are used to indicate the range assumed by the distances in single molecule Förster Resonance Energy Transfer (smFRET) experiments (Chen et al., 2017; Dagdas et al., 2017).
During the conformational transition, the distance between the Ca atom of the HNH catalytic residue (H840) and the scissile phosphate on the target DNA strand (i.e. the phosphate atom at position ~3) shows a gradual decrease starting at ~7 Å, stabilizing at a distance of ~8 Å from the cleavage site after ~10 µs of MD (Fig. 2b, Fig. S2B). Remarkably, upon ~7 µs of unbiased MD, H840 approached the scissile phosphate at ~15.0 Å, reaching the configuration previously observed via GaMD simulations (Fig. S1). This result further establishes the ability of GaMD to capture long time scale events. However, longer GaMD trajectories might be required to access the complete conformational transition. In the final configuration, the catalytic H840 Ca is located ~8 Å from the cleavage site, while the imidazole side chain (the reactive nitrogen) locates at a distance of ~5–6 Å from the cleavage site, priming the HNH domain for the hydrolysis of the TS (Fig. S3) (Jinek et al., 2014). This configuration of the HNH domain agrees well with smFRET experiments (Raper et al., 2018; Sternberg et al., 2015). Indeed, the S867–S355 distance, previously used to characterize the activated state of the HNH domain, reaches the experimental value of ~21 Å (Fig. S4). In the course of the simulation, we detect significant conformational plasticity of the REC lobe, in agreement with previous characterizations, performed using smFRET experiments (Chen et al., 2017; Dagdas et al., 2017). Figure 2c–d reports the time evolution during MD of the E60–D273 and S960–S701 distances, which have been used in smFRET to distinguish the conformational states adopted by the REC2 and REC3 domains, respectively (Chen et al., 2017; Dagdas et al., 2017). Here, we observe a conformational change of REC2 (Fig. 2a–c). After ~7 µs, simultaneously with the initiation of the conformational rearrangement of HNH, REC2 starts an outward transition that, upon ~11 µs of MD, results in an overall translation by ~8 Å relative to the starting position (i.e. the E60–D273 distance reaches ~40 Å from the initial 32.6 of the X-ray structure 5F9R.pdb). REC3 also shows significant conformational transitions (Fig. 2d). Indeed, the S960–S701 distance broadly fluctuates but overall increases by ~3–3.5 Å (from the initial value of 40.3 of the X-ray structure 5F9R.pdb), resulting in the opening of the groove hosting the RNA:DNA hybrid. Remarkably, in the final configuration the E60–D273 and S960–S701 distances reach values that well agree with the smFRET ranges (Figure 2c–d) (Chen et al., 2017; Dagdas et al., 2017). Interestingly, the transitions of HNH and REC2 appear to be concerted, as it starts for both domains upon ~7 µs of simulation (Fig. 2b–c). However, HNH reaches an active conformation by 10 µs, while REC2 fully adopts the conformational transition at ~11 µs. Together, these observations suggest that the conformational changes of the recognition lobe assist the conformational activation of the HNH domain. Specifically, we observe an opening of REC3 and the mutual conformational adaptation of REC2 and HNH, whereby HNH approaches the cleavage site on the TS and REC2 moves apart with an outward translation of ~8 Å, enabling the active site to access the cleavage site.

A recent cryo-EM structure, solved at a 5.2 Å resolution (EMD-8236), shows that the HNH domain is closer to the recognition lobe than in the pre-cleavage state crystal structure (Hui et al., 2017). The final configuration obtained from MD is in good agreement with the all-atom model fitted in the EMD-8236 EM map (5Y36.pdb, Fig. 2b–d). In detail, in the 5Y36.pdb, the E60–D273 and S960–S701 distances reach values of 38.4 and 42.6 Å, respectively. MD simulations access these values by ~11 µs (for the E60–D273 distance, Fig. 2c) and ~10 µs (for the S960–S701 distance, Fig. 2d). In the fitted structure, the H840 Ca atom remains at a distance of 10.1 Å from the scissile phosphate (Fig. 2b), which is beyond the range required for catalysis (Fig. S3). In summary, the continuous MD simulations overall access the conformation obtained via cryo-EM, and further explore the conformational space. In this respect, it is worth noting given its limited resolution (5.2 Å), the available EMD-8236 structure can in fact represent multiple conformational states that could not be resolved (Nogales, 2016). As such, the fluctuations captured by the extensive MD simulations can be considered representative of the conformational landscape surrounding the EMD-8236 structure.

In order to track the large-scale collective motions of the REC lobe during the long time scale dynamics, we performed Principal Component Analysis (PCA) (Amadei et al., 1993). This analysis captures how the protein domains move with respect to each other, highlighting conformational changes that are difficult to observe by visual inspection of the MD trajectory. As a result, PCA confirms large amplitude motions for HNH and the REC2–3 domains and the concerted nature of the conformational changes, while also revealing a conformational change for REC1 (Movie S1, Fig. S5). The latter moves in an opposite direction with respect to REC2 and REC3, toward the HNH domain. This result agrees well with the experimental vector map of global Cas9 conformational changes, from the RNA-bound state (4ZT0.pdb) to the DNA-bound state (5F9R.pdb, Fig. S4D) (Chen et al., 2017). During the transition, the HNH domain forms a series of salt-bridge interactions with the REC lobe (Fig. 3). These ionic interactions mainly involve the REC1–2 regions, while HNH and REC3 remain separated from each other. This result well agrees with the novel cryo-EM structure (EMD-8236) (Huai et al., 2017). In this structure, HNH approaches the REC1 domain, but moves away from REC3 (Fig. S6). At ~7 µs of MD, charged residues of HNH and REC1 start engaging in ionic interactions, which increase in strength and number along the dynamics, stabilizing after ~10 µs, locking HNH at the TS (Fig. 3a). These interactions are key for the stable docking of HNH at the cleavage site. REC2 is also involved in ionic interactions, which are maintained along the simulation, indicative of the concerted conformational change of the two domains (Fig. 3b). Remarkably, interactions between HNH–REC1 are maintained in the activated state (i.e. during the last ~4 µs of MD), ensuring the positioning of HNH at the cleavage site. REC1 therefore cooperates with REC2 and REC3 in favoring HNH activation.

**Insights on the on-target specificity**

Here, we monitored the dynamic interactions of key positively charged residues, which belong to the HNH domain and intervene in the specificity of CRISPR–Cas9 (Fig. 4). Among them, K810 and K848 have been shown to reduce off-target cleavage events when mutated to alanine (Slaymaker et al., 2016). K913 has been shown to bind the NTS in shorter time scales (i.e. ~0.8 µs), suggesting a possible role in facilitating the approach of HNH toward the scissile phosphate (Palermo et al., 2016). In the pre-activated state (Jiang et al., 2016), these residues do not interact with the nucleic acids. With the approach of HNH to the cleavage site on the TS (i.e. phosphate ~3), K810 binds the phosphate ~4 (Fig. 4a). Besides, K848 establishes multiple interactions with the nucleic acids and with the protein residues (Fig. 4b), finally contacting the DNA:RNA hybrid via the RNA phosphate backbone. Remarkably, in the 5Y36.pdb (all-atom
model of the EMD-8236 map), K810 approaches the TS, while K848 binds the RNA backbone similarly to the configuration obtained upon ∼15 µs of MD (at position −8) (Huai et al., 2017). K913 engages the NTS at several positions throughout the simulations (Fig. 4c). This finding confirms previous evidences from shorter MD simulations, suggesting that the interactions between the NTS and the loop formed by 906–918 residues would favor the approach of HNH toward the TS (Palermo et al., 2016), and also clarifies smFRET experiments showing that the docking of HNH at the TS in its active configuration requires the presence of the NTS (Dagdas et al., 2017). Overall, these residues act as anchors of HNH at the DNA, favoring its docking at the TS cleavage site. As such, the disruption of these interactions with K810A/K848A substitutions may destabilize HNH at the cleavage site, thereby altering the dynamics of its conformational activation and consequently its cleavage activity (Chen et al., 2017; Slaymaker et al., 2016)

Multiple repeats along the conformational change
Continuous MD simulations of huge macromolecular systems reaching the microsecond time scale are challenging to be achieved via conventional supercomputers and the data obtained using Anton-2 are at the limit of the state-of-the-art technology.
As such, the MD performed using Anton-2 in the present study, as well as in the previous studies by our and other research groups (Lindorff-Larsen et al., 2016; Mouchlis et al., 2015), enabled to capture conformational changes and biophysical processes in a single trajectory. While this approach certainly preserves the kinetic features of the system, considering the stochastic nature of the biomolecular processes, multiple simulations are required to precisely estimate the time scale of the events. However, due to the high computational cost of each single trajectory, it is challenging to produce additional and/or longer simulation runs, recovering the observed events. In order to cope with this issue and to understand the statistical relevance of the observed conformational changes, we have extracted eight equally distributed snapshots (at times $1, 3, 5, 7, 9, 11, 13,$ and $15 \mu s$ of the Anton-2 trajectory) to perform independent MD simulations of $\sim 300$ ns in two replicas (reaching additional $\sim 4.8 \mu s$ of aggregate statistics). Figures S6–S8 report the time evolution along the simulated runs of the H840–P$_{DNA}$, E60–D273, and S960–S701 distances, enabling comparison with the continuous simulation obtained with Anton-2 (Fig. 2). As a result, in the systems extracted at $1, 3,$ and $5 \mu s$, in which Cas9 assumes a pre-activated configuration, the computed distances stably oscillate around the initial values. The three distances also remain stable in the activated configurations (i.e. extracted at $11, 13,$ and $15 \mu s$). In the case of the systems extracted at $7$ and $9 \mu s$, at which the system undergoes the conformational transition, we observe higher fluctuations in the time evolution of the computed distances. Moreover, in the hundreds-of-nanoseconds, these latter follow the trend observed in the microsecond length dynamics, which shows the transition toward the activated state (Figs S6–S8). This highlights the tendency for the conformational transition toward activation, supporting the results obtained with single trajectory obtained with Anton-2 and previous enhanced sampling simulations (Fig. S1) (Palermo et al., 2017a). In order to understand the factors underlying this consistent behavior, we examined the structures extracted at times $1, 3, 5, 7, 9, 11, 13,$ and $15 \mu s$ of the Anton-2 trajectory and used as a starting point of the independent MD runs (Fig. S9). As a main difference between the pre-activated systems (i.e. extracted at times $1, 3,$ and $5 \mu s$) and the system initiating the conformational transition (i.e. extracted at time $7 \mu s$), in this latter we observe the approach of the side chains of charged residues belonging to HNH and REC1 (Fig. S9D), preluding their engagement in salt-bridge interactions, which will be fully formed at time $9 \mu s$. At this point (9 $\mu s$, Fig. S9E), the outward transition of REC2 with respect to HNH is observed. Considering that these newly formed salt-bridge interactions are stably maintained in the activated state (i.e. for snapshots extracted at 11, 13, and $15 \mu s$, as well as throughout the last $\sim 6 \mu s$ of the continuous MD simulation, Fig. 3), these results suggest that the approach of the side chains of the charged residues of HNH and REC1 at $\sim 7 \mu s$ is the first triggering event making the system prone to the larger conformational change leading to its final activation.

**Cooperative domain dynamics in the activated state**

The activated state reached via MD simulations remains stable over the last $\sim 6 \mu s$ (Fig. 2), allowing its conformational dynamics to be analyzed. To characterize the inter-dependent motions of the protein residues and understand the cooperative dynamics of the Cas9 domains in the activated state, we employed the Lange and Grubmüller method, which captures the overall correlations (i.e. both linear and non-linear) among protein residues (Lange and Grubmuller, 2006). The generalized correlation (GC) matrix is a sensitive method for detecting the interdependence in the motions of two spatially distant residues, and provides a measure of how much the motion of one residue is dependent on that of another residue. GC matrices have been computed for the activated configurations, over the last $\sim 4 \mu s$ of MD. Visual inspection of the GC matrix shows that the HNH domain strongly correlates with the REC lobe in the active state (Fig. 5a). A quantitative evaluation of the inter-dependent couplings established by HNH with the REC lobe has been obtained by computing the inter-domain GC scores (GCs), which
accumulate the GC coefficients of the single residues over each protein domain of interest (Fig. 5b, full details in the Methods section). The GCs are a measure of the overall inter-domain correlations, indicating the most relevant coupled motions occurring across the system during the simulations. The GCs have been useful in the characterization of the allosteric effects in CRISPR–Cas9 (Palermo et al., 2017b) and other protein/nucleic acid complexes (Ricci et al., 2016), as well as in describing the inter-dependent conformational dynamics between the protein and RNA components of the human spliceosome (Casalino et al., 2018). As a result, the HNH and REC3 domains display the highest overall GCs of 0.64 (Fig. 5b), which indicates a highly cooperative conformational dynamics in the activated state. Notably, both MD simulations and recent cryo-EM data (EMD-8236) (Huai et al., 2017) show that HNH and REC3 are not in direct contact with each other in the activated configuration (Figs 2d and 5c). Considering that the RNA:DNA hybrid locates within the groove formed by HNH and REC3 and directly interacts with both domains, the high inter-dependency of the domains motions suggests that the hybrid mediates the coupling between the two domains. This provides a plausible explanation for recent experimental observations, showing that REC3 allows for HNH nuclease activation upon recognition of the formation of the RNA:DNA hybrid (Chen et al., 2017). High inter-domain correlations are also detected for the HNH and REC2 (GCs of 0.48). This inter-dependent conformational dynamics corroborate the experimental evidence for tight coupling between the HNH and REC2 domains to ensure catalytic competence (Chen et al., 2017). Specifically, smFRET has shown reciprocal changes in the conformational states assumed by HNH and REC2 across multiple DNA substrates, indicating that the conformational dynamics of HNH and REC2 is tightly coupled to ensure catalysis. Notably, the HNH domain has high conformational plasticity, as shown by biochemical and biophysical experiments (Osuka et al., 2018; Sternberg et al., 2015) and previous shorter MD simulations (Palermo et al., 2016). Additionally, in the cryo-EM structure EMD-3277, the HNH and REC2 are observed at a lower resolution (8–10 Å) than the overall structure (6 Å), highlighting their mobility (Jiang et al., 2016). In light of this evidence, the data reported here explain how these flexible protein components are coupled to enable function. Finally, high correlations are also observed between the REC2 and REC3 (GCs of 0.47), indicating that in the activated state, the REC lobe engages in a tight interplay to enable function.

**Discussion**

**Activation mechanism**

In the activation process of CRISPR–Cas9, a conformational change of the catalytic HNH domain is required to trigger catalysis (Fig. 1) (Jiang et al., 2016). Here, MD simulations show that the docking of the HNH domain at the TS is facilitated by favorable interactions and by the conformational changes of the REC lobe. We show an opening of REC3 and an outward translation of REC2, which moves apart from its original position, resulting in the exposure of TS to the HNH active site for cleavage (Fig. 2). This result agrees well with the available experimental data (Chen and Doudna, 2017; Dagdas et al., 2017; Huai et al., 2017) and reveals that the activation of HNH is dependent on the conformational changes of the REC domains. Specifically, REC3 shows significant conformational transitions that result in the opening of the groove accommodating the RNA:DNA hybrid. This finding well agrees with recent cryo-EM data (Huai et al., 2017) and with the available smFRET experiments (Chen et al., 2017; Dagdas et al., 2017), showing that the activation of the HNH domain for catalysis requires a change in the conformational state of REC3. In the activated configuration, the conformational dynamics of the HNH and REC3 domains is highly coupled (Fig. 5). Considering that both HNH and REC3 directly interact with the RNA:DNA hybrid, the high inter-dependency of the domains motions suggests that the hybrid mediates the coupling between the two domains. This finding provides further clarification on recent experiments, suggesting that REC3 would allow for HNH activation upon ‘sensing’ the formation of a complete RNA:DNA hybrid (Chen et al., 2017). Accordingly, the simulations show that, while ‘sensing’ (i.e. interacting with) the RNA:DNA hybrid, REC3 affects the conformational dynamics

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Fig. 5. (a) Matrix of the generalized correlation (GC), calculated over the last ∼4 µs of MD of CRISPR–Cas9. The strength of the computed correlations is color-coded from green (highly correlated motions) to gray (poorly correlated motions). A box is used to highlight the high correlation between the HNH domain and the REC lobe. (b) Inter-domain GC scores (GCs) plotted for each Cas9 domain (REC1-3, HNH, RuvC, and PI domain) in a two-by-two matrix, measuring the strength of the overall correlation that each domain establishes with the others (details in the Method section). (c) The highest GCs are plotted on the three-dimensional structure of CRISPR–Cas9, revealing a tight coupling between HNH and the REC2-3 domains. Cas9 is shown in a molecular surface, highlighting the HNH and REC2-3 domains with different colors. The RNA (orange) and the target DNA (TS, magenta) forming a RNA:DNA hybrid, as well as the non-target DNA (NTS, cyan) are shown as ribbons.

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of HNH and, in turn, its activation for cleavage. MD simulations also pinpoint a key role for the REC2 domain. During the conformational activation of HNH, REC2 rotates outward to enable the HNH to approach the cleavage site on the TS (Fig. 2). As well, the continuous simulation shows that during the activation process, the opening of REC2 and the approach of HNH to the TS start simultaneously (at ∼7 μs, Fig. 2), indicating that the two domains cooperatively relocate to facilitate catalysis, likely in response to the ‘sensing’ of REC3 (Sung et al., 2018). In the activated state, HNH and REC2 show highly cooperative conformational dynamics (Fig. 5), corroborating the experimental evidence of their tight coupling to ensure catalytic competence (Chen et al., 2017). Taken together, these findings suggest that REC2 functions as a ‘regulator’ for HNH function (Chen and Doudna, 2017; Chen et al., 2017). MD simulations further reveal an unexpected role for REC1, which approaches HNH by establishing a series of ionic interactions, thereby enabling HNH to dock at the cleavage site on the TS. These observations suggest that REC1 acts as a ‘lock’ for HNH (Fig. 3). Overall, the tight interplay observed among the REC1–3 domains and the HNH domain reveals how the ‘sensor’ (REC3), the ‘regulator’ (REC2) of HNH activation (Chen and Doudna, 2017; Chen et al., 2017), together with the ‘lock’ (REC1), contribute to the formation of an activated CRISPR–Cas9 complex (Movie S2).

Conclusions

Here, we disclose the mechanism leading to the formation of a catalytically active CRISPR–Cas9 nucleosome following substrate DNA binding. Unbiased all-atoms MD simulations reveal the approach of the HNH domain to the cleavage site on the TS in ∼16 μs of continuous simulation. We characterize on an atomic scale the molecular determinants leading to the HNH conformational transition and the key role of the REC lobe in enabling the docking of HNH at the cleavage site. The remarkable conformational changes and the collective conformational dynamics of the REC1–3 domains enable ‘sensing’ of the nucleic acid binding, ‘regulation’ of the HNH conformational transition and ‘locking’ of the HNH at the cleavage site, eventually leading to the formation of a catalytically active CRISPR–Cas9 (Movie S2). Good agreement with biochemical experiments and structural data highlights the consistency of the activated conformation. As such, these simulations add detailed mechanistic information on the CRISPR–Cas9 activation process, clarifying the mechanistic role of the REC lobe components. Considering the key role of the REC lobe in the specificity of Cas9 (Casini et al., 2018; Chen et al., 2017; Kleinstiver et al., 2016), this study provides the foundation for understanding how the REC lobe domains control the cleavage of off-target sequences. Toward this aim, extensive characterization is currently ongoing in our laboratories. Overall, the knowledge on the HNH activation process and the role of the recognition lobe components, deciphered here, establishes a framework for future studies and novel structure-based engineering efforts for improved genome editing.

Material and method section

Structural model

MD simulations were based on the X-ray structure of the S. pyogenes Cas9 in complex with RNA and DNA (5F9R.pdb), solved at 3.40 Å resolution (Jiang et al., 2016). The model system was embedded in explicit waters, while Na+ ions were added to neutralize the total charge, leading to an orthorhombic periodic simulation cell of ∼180 · 120 · 140 Å³, for a total of ∼300 000 atoms.

MD simulations

A simulation protocol tailored for RNA/DNA endonucleases was adopted (Palermo et al., 2015; Sponer et al., 2018), embracing the use of the Amber ff12SB force field, which includes the ff99bsc0 corrections for DNA (Perez et al., 2007) and the ff99bsc0 + χOL3 corrections for RNA (Banas et al., 2010; Zgarbova et al., 2011). The TIP3P model was employed for explicit water molecules (Jorgensen et al., 1983). The Åqvist (Åqvist, 1990) force field was employed for Mg ions, as in previous studies on similar Mg-aided RNA/DNA nucleases (Casalino et al., 2016; 2017; Palermo et al., 2013). An integration time step of 2 fs was used. All bond lengths involving hydrogen atoms were constrained using the SHAKE algorithm (Ryckaert et al., 1977). Temperature control (300 K) was performed via Langevin dynamics (Turq et al., 1977), with a collision frequency γ = 1. Pressure control was accomplished by coupling the system to a Berendsen barostat (Berendsen et al., 1984), at a reference pressure of 1 atm and with a relaxation time of 2 ps. The system was subjected to energy minimization to relax water molecules and counter ions, keeping the protein, the RNA, DNA, and Mg ions fixed with harmonic position restraints of 300 kcal/mol · Å². Then, the system was heated up from 0 to 100 K in the canonical ensemble (NVT), by running two simulations of 5 ps each, imposing position restraints of 100 kcal/mol · Å² on the above-mentioned elements of the system. The temperature was further increased up to 200 K in ∼100 ps of MD in the isothermal–isobaric ensemble (NPT), reducing the restraint to 25 kcal/mol · Å². Subsequently, all restraints were released and the temperature of the system was raised up to 300 K in a single NPT simulation of 500 ps. After ∼1.1 ns of equilibration, ∼10 ns of NPT runs were carried out allowing the density of the system to stabilize around 1.01 g/cm³. Finally, the production runs were carried out in the NVT ensemble. Simulations were performed using the GPU version of AMBER 16 (Case et al., 2016; Salomon-Ferrer et al., 2013) and the SPFP precision model (Le Grand et al., 2013). MD simulations were performed to equilibrate the system for ∼120 ns prior to long time scale continuous MD. Indeed, the well-equilibrated system was used as a starting point for simulations on Anto-2 (described below). As well, eight equally distributed snapshots were extracted at times 1, 3, 5, 7, 9, 11, 13, and 15 μs of the Anto-2 trajectory and subjected to independent MD simulations in two replicates (i.e., ∼300 ns for each snapshot in two replicates), reaching a total of additional 4.8 μs.

Long time scale MD simulations of the CRISPR–Cas9 complex were performed using Anto-2 (Shaw et al., 2014), a special-purpose supercomputer for micro-to-millisecond length MD, starting from a well-equilibrated configuration, obtained after ∼120 ns of conventional MD (see above). Simulations on Anto-2 were performed using the same force-field parameters used for conventional MD simulations. A reversible multiple time step algorithm (Tuckerman et al., 1992) was employed to integrate the equations of motion with a time step of 2 fs for short-range non-bonded and bonded forces and 6 fs for the long-range non-bonded forces, for a total of ∼16 μs of simulations. Simulations were performed at constant temperature (300 K) and pressure (1 atm) using the multigrator integrator as implemented in Anto-2 (Lippert et al., 2013). The k-Gaussian split
Ewald method (Shan et al., 2005) was used for long-range electrostatic interactions. Hydrogen atoms were added assuming standard bond lengths and were constrained to their equilibrium position with the SHAKE algorithm (Ryckaert et al., 1977). Overall, molecular simulations have been carried out for an overall sampling time of >20 μs.

**Principal component analysis**

In PCA, the covariance matrix of the protein Ca atoms is calculated and diagonalized to obtain a new set of generalized coordinates (eigenvectors) to describe the system motions. Each eigenvector, called Principal Component (PC), is associated to an eigenvalue corresponding to the mean square fluctuation contained in the system’s trajectory projected along that eigenvector. The first PC1 corresponds to the system’s largest amplitude motion, and the dynamics of the system along PC1 is usually referred to as ‘essential dynamics’ (Amadei et al., 1993). In this work, each conformation sampled during MD was projected into the collective coordinate space defined by the first two eigenvectors (PC1 and PC2), characterizing the essential conformational sub-space sampled by Cas9. Full details are in the SI.

**Correlation analysis**

Cross-correlations of residues in the Cas9 protein were computed based on mutual information between all Ca atoms using the generalized correlation analysis approach developed by Lange and Grubmüller (Lange and Grubmüller, 2006), which is explained in detail in the SI. The g_correlation module in the Gromacs 3.3 (Lindahl et al., 2001) package has been employed. Based on the computed correlations, the Generalized Correlation score (GCs) has been employed to define quantitatively the inter-dependent inter-residue couplings established by HNH with the REC lobe. The GCs has been originally introduced by Ricci et al. in the study of the conformational dynamics of the PPARY–RXRa nuclear receptor complex (Ricci et al., 2016), and used to describe the allosteric effects in CRISPR-Cas9 (Palermo et al., 2017b) and the inter-dependent dynamics of the human spliceosome (Casalino et al., 2018). The GCs are computed by processing the GC matrix as follows. For each amino acid residue a GCs can be defined:

$$GC_{ij} = \sum_{j \neq i} GC_{ij},$$

representing a measure of both the number and the intensity of the GC coefficients displayed by each residue. To filter non-trivial correlations and eliminate the noise due to uncorrelated motions, per-residue GCs were computed considering only highly positive correlations (GC ≥ 0.60). GCs were used to detail the overall inter-domain correlations as follows: GCs were calculated for each residue i belonging to a specific protein domain (e.g. HNH, Rec1–3), with the residues j belonging to another protein domain of interest. Then, GCs were accumulated over all residues j of each specific Cas9 domain and normalized by the number of coupling residues i and j, which display GC ≥ 0.60. This resulted in a set of per-domain GCs, ranging from 0 (not-correlated) to 1 (correlated), measuring the strength of the overall correlation that each domain establishes with the others. The overall GCs are a measure of the most important correlated motions among protein domains taking place in the simulations, which help in identifying how specific protein regions mechanically intervene in the overall correlation network (Palermo et al., 2017b; Ricci et al., 2016).

**Supplementary material**

The supplementary material for this article can be found at https://doi.org/10.1017/S0033583518000070

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**References**


