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Catalytic Mechanism of Non-Target DNA Cleavage in CRISPR-Cas9 Revealed by Ab-Initio Molecular Dynamics

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Abstract

CRISPR-Cas9 is a cutting-edge genome editing technology, which uses the endonuclease Cas9 to introduce mutations at desired sites of the genome. This revolutionary tool is promising to treat a myriad of human genetic diseases. Nevertheless, the molecular basis of DNA cleavage, which is a fundamental step for genome editing, has not been established. Here, quantum-classical molecular dynamics and free energy methods are used to disclose the two-metal-dependent mechanism of phosphodiester bond cleavage in CRISPR-Cas9. Ab-initio MD reveals a conformational rearrangement of the Mg²⁺-bound RuvC active site, which entails the relocation of H983 to act as a general base. Then, the DNA cleavage proceeds through a concerted associative pathway fundamentally assisted by the joint dynamics of the two Mg²⁺ ions. This clarifies previous controversial experimental evidences, which could not fully establish the catalytic role of the conserved H983 and the metal cluster conformation. Overall, the non-target DNA cleavage catalysis described here resolves a fundamental open question in the CRISPR-Cas9 biology, and provides valuable insights for improving the catalytic efficiency and the metal-dependent function of the Cas9 enzyme, which are at the basis of the development of novel genome-editing tools.



Introduction

CRISPR (clustered regularly interspaced short palindromic repeats)-Cas9 is a revolutionary genome editing tool, which is bearing innovative discoveries across the field of Life Sciences. 1 By enabling the correction of DNA mutations, this technology offers the promise to treat a priori a myriad of human genetic diseases. Recent advances have shown the first cancer patients being infused with CRISPR-Cas9-modified T cells to increase the natural antitumor immune response.² The CRISPR-Cas9 "molecular scissors" uses the RNA-guided endonuclease Cas9 to precisely cleave DNA sequences of interest.³ In spite of a ground-breaking impact and widespread utilization, the catalytic mechanism of DNA cleavage, which is a fundamental step in genome editing, is not understood, potentially limiting the use of CRISPR-Cas9 for further biomedical applications. Its knowledge is paramount for improving the catalytic efficiency of the Cas9 enzyme and, in turn, for the development of novel and more specific Cas9-based genome editing tools.

Similarly to several other nuclease enzymes, Cas9 is a metal-dependent nuclease that performs Mg²⁺-aided DNA cleavages and is inhibited by Ca²⁺. ^{3,4} In the activated form of the enzymatic complex, the DNA binds Cas9 by matching a guide RNA with one strand (the target strand), while the other non-target strand is displaced and also accommodated in the protein. Then, two nuclease domains - HNH and RuvC - cleave the target and nontarget DNA strands, respectively (Figure 1A). The RuvC domain shares the structural fold of the RNA Ribonuclease H (RNase H) and is likewise thought to perform phosphodiester bond cleavages through a two-metal-ion mechanism (Figure 1A).^{4,5} Nevertheless, the mechanism of DNA cleavage as operated by RuvC remains nebulous and appears controversial from the available structural and biophysical studies. Structures of CRISPR-



Cas9 have shown that the RuvC active site hosts three carboxylates (D10, D986 and E762), 4,6-8 which constitute the highly conserved DDE (or DEDD) motif, a fingerprint of the two-metal-dependent nucleases (Figure 1B).9 The metal-bound X-ray structure obtained in the presence of Mn²⁺ ions revealed that the DDE motif coordinates the two catalytic metals, ^{3,6} while the H983 residue also approaches the active site coordinating the A-site metal (MeA) with the nitrogen ligand. Previous computational studies employing extensive classical molecular dynamics (MD)^{10–12} conveyed the observation that one water molecule stably coordinates MgA, in close proximity to the scissile phosphate and suitable to act as a nucleophile (Figure 1B, Movie S1). In this scenario, critical open questions remain unmet. First, it is unknown how the catalytically relevant Mg²⁺ ions, rather than the crystallographic Mn²⁺, would rearrange within the active site to accomplish the catalysis. Moreover, the intricate features of the chemical mechanism, including the origin of the nucleophile initiating the chemical step, and the type of catalysis leading to phosphodiester bond cleavage, have not been described. Although biochemical experiments have suggested that H983 could have a role in this chemical step, ⁷ crystallographic data trapping H983 in the coordination sphere of MeA arise debates on how this residue could engage in the catalysis. The efficiency and specificity of the enzyme depend on the clarification of these and on other unresolved mechanistic aspects, such as the dynamical pathway leading from the reactants (**R**) to the products (**P**), the exact mechanism for transition state (TS^{\ddagger}) stabilization, and the pivotal role and dynamics of the catalytic metals.^{5,13,14} Their understanding is crucial to improve the enzyme specificity and reduce off-target cleavages, a key goal of biomedical applications of CRISPR-Cas9 genome editing. 15



Here, we characterize the mechanism of phosphodiester bond cleavage within the RuvC active site through quantum-classical QM(Car-Parrinello)/MM MD simulations (ab-initio MD)¹⁶ and free energy methods. Ab-initio MD reveals a conformational change of the RuvC active site upon substitution of the crystallographic Mn²⁺ ions with the catalytically relevant Mg²⁺, which results in the loss of the H983 ligand by the A-site ion (MgA) and in the formation of a stable catalytic site, prone for the in-line nucleophilic attack. Then, the two-metal-dependent catalytic mechanism proceeds through a concerted (S_N2-like) associative pathway, activated by the conserved H983 and fundamentally assisted by the joint dynamics of the catalytic Mg²⁺ ions. Interestingly, a second set of abinitio simulations retaining the crystallographic configuration reveals an alternative phosphate-mediated cleavage, which proceeds through a concerted dissociative pathway. Although the two proposed mechanisms are found to be energetically comparable, mutagenesis experiments revealing that the H983A mutation abolishes the RuvC catalysis ⁷ support the active role of H983 and advocate for the H983-activated mechanism. Notably, the dynamics of the catalytic metals remarkably differs in the two chemical pathways and can be considered a feature distinguishing the nature of phosphodiester bond cleavage in the two-metal-ion enzymes. Comparison with the two-metal-dependent enzymes further supports the identified mechanism and also suggests a common catalytic strategy for genome editing and recombination. Overall, the findings and chemical details of DNA cleavage proposed in this work provide foundational knowledge on how the endonuclease Cas9 cleaves nucleic acids, and pose the basis for future investigations aimed at completing the understanding of the CRISPR-Cas9 mechanism of action. This work also



represents a valuable contribution for future mechanism-based design of improved CRISPR-Cas9 genome-editing tools.

Results

Conformation of the Reactant (R). QM/MM MD simulations have been based on the most complete X-ray structure of CRISPR-Cas9 (PDB ID: 5F9R). ⁶ The active site has been treated at the DFT-BLYP^{17,18} level of theory, while the surrounding molecular environment has been described by MMforce fields, resulting into a box of ~340,000 atoms. The QM part (105 atoms) includes the scissile and adjacent DNA phosphates groups, the two metal ions and their coordination sphere, composed by the DDE motif and H983 (protonated in epsilon, as arising from previous investigations). 4,11,12 Six solvation waters have also been treated at the QM-BLYP level of theory (details are reported in the Supporting Information (SI)). The choice of simulating an enlarged QM part – including 6 solvation waters and protein residues, as well as the adjacent DNA phosphate groups – enabled to assess whether these chemical groups engage in the catalysis and the related proton transfer events. This is a key point because the activation of the nucleophile could occur in different ways, such as through a phosphate group 19-22 or an amino acid 23,24 serving as a general base, or even through the bulk water. 25,26 Notably, these ab-initio simulations have been performed substituting the crystallized Mn²⁺ ions with the catalytically active Mg²⁺ ions.⁴

The investigation of the reactant (R) state has been initially performed through unconstrained ab-initio MD (two replicas of ~40 ps) in order to assess its stability and ensure that the chemical requirements for a two-metal-ion catalysis were satisfied.

As a result, we observe a conformational rearrangement of the catalytic site, whereby H983 detaches from MgA, while the nucleophilic water molecule simultaneously moves in between H983 and the scissile phosphate (Figure 1B). This change of the interaction network established by H983 and by the water nucleophile (Figure 2C) is observed in both simulation replicas (Figure S3). However, during these events the water nucleophile preserves the contact with MgA, whose coordination sphere is saturated by the side-chain oxygen of S15 (Figure S3). In the final configuration, the DDE motif steadily coordinates the Mg²⁺ ions, while the scissile phosphate locates in between the two metal ions, which are jointly coordinated by the pro-Sp oxygen. This is a key condition for the catalysis, as it allows the scissile phosphate to assume a configuration prone for an in-line nucleophilic attack.^{5,27} Noteworthy, a similar structural organization of the catalytic site had been also observed in a previous study performed by us as a result of ~2.4 µs of force field-based accelerated MD followed by ~40 ps of ab-initio MD (Figure S4). 11 Moreover, the detachment of the H983 ligand from MgA has also been reported in some classical MD simulations, including ours, ^{10,12} which however do not allow conclusive remarks on the catalysis and on the metal ions geometry, given the well-known limitations of the employed classical force-field models.²⁸ Interestingly, Mg²⁺ poorly binds to nitrogen, whereas Mn²⁺ shows high affinity. ^{29–31} In fact, it has been suggested that Mn²⁺ can contact nitrogen atoms in crystallization buffers.³¹ Although one cannot generalize,^{31,32} this supports the loss of the Mg²⁺-N coordination patterns observed during classical and ab-initio MD. To cross-validate this observation within the RuvC site, we performed unconstrained ab-initio MD also in the presence of the crystallographic Mn²⁺ ions. As a result, MnA preserves the nitrogen ligand and the crystallographic coordination



sphere over ~40 ps (Figure S5). Taken together, these findings suggest that the coordination of MgA by means of H983 is unlikely in the RuvC site. Yet, H983 interacts with the attacking water, which in turn is stably positioned in proximity to the scissile phosphate and coordinates MgA (Figure 1B). The resulting stable configuration of the metal cluster and of the surrounding protein residues in the reactant state (R) laid the foundation for the investigation of the catalytic mechanism.

Mechanism of phosphodiester bond cleavage. To probe the catalytic mechanism of phosphodiester bond cleavage, we performed QM/MM MD simulations in combination with free energy methods. Specifically, we employed the "blue moon ensemble" method. along with thermodynamic integration. ¹⁶ In this approach, the reaction mechanism is investigated along a selected and appropriate Reaction Coordinate (RC). This is progressively evolved from the **R** to the **P** through a series of sequential steps, wherein its value is kept fixed. By integrating the average (converged) constraint forces obtained along the pathway we derived the associated free energy profile (details are reported in the SI). In the present study, we used as a RC the difference in distance between the breaking (O3'DNA-PDNA) and forming (OWAT-PDNA) bonds (Figure 2A). The appropriateness of this RC for the study of phosphodiester bond cleavage has been shown in several studies of RNA/DNA processing enzymes 19,20,25,33 and is discussed in the SI. Overall, this method has been widely employed to investigate biochemical reactions, ¹⁶ including the two-metalion mechanism in several enzymes,13 such as the prototypical RNase H,19,20 and noncoding RNAs like group II intron ribozyme²⁵ and the spliceosome.³⁴ This enables direct comparison with the current investigations of the catalytic mechanism in CRISPR-Cas9.

The simulations show that the system evolves from the reactants (R) to the products (P), separated by a transition state (TS[‡]) maximum (Figure 2 and S6). The reaction is activated by H983, which acts as a general base, and proceeds through a concerted ($S_N 2$ -like) associative mechanism (Movie S2). 35,36 In detail, at a RC = 0 Å, a proton transfer (PT) event is observed, where H983 activates the nucleophilic water into a hydroxide ion by abstracting a proton right before the TS^{\ddagger} . Indeed, from RC = 0 Å (corresponding to the PT) to RC = 0.2 Å (highest peak of the free energy profile, corresponding to the TS[‡]), the free energy profile displays a plateau, indicating that the barrier for the proton abstraction by H983 is negligible (i.e., free energy barrier for PT of less than 1 kcal mol⁻¹) with respect to the **TS**[‡] (details are reported in the SI). In support of this observation, a similar histidine-activated mechanism and similar energetics have been recently observed in the catalysis of the HIV-1 RNase H.²⁴ Upon activation, OH⁻ becomes fully aligned with the scissile P-O3' bond, resulting in the formation of TS[‡]. At RC = 0.4 Å, the reaction evolves downhill toward $\bf P$ formation. A full description and cross-validation of this mechanism, including the calculation of the PT mechanism, is given in the SI (Figure S7). The associative phosphodiester bond cleavage proceeds with an overall Helmholtz free energy ($\Delta F\#$) of $\sim 16.55 \pm 1.22$ kcal mol⁻¹. The backward free energy profile (i.e., obtained by sampling along the RC in the opposite direction) was also computed, revealing no large hysteresis between the forward and backward process (Figure S6A). The calculated activation barrier of $\sim 16.55 \pm 1.22$ kcal mol⁻¹ is in agreement with the experimental catalytic rate of 3.5 s⁻¹ (corresponding to $\Delta G\# \sim 16/17$ kcal mol⁻¹).³⁷ This rate constant has been measured distinguishing the catalysis from nucleic acid binding and providing a measure for the RuvC domain, enabling proper comparison with our

computations. Importantly, biochemical experiments performed in a previous study showed that the H983A mutation abolishes the RuvC activity, therefore supporting the active role of H983 in the catalysis.⁷

The variation of the significant interaction distances along the investigated reaction step (Figure 3A) provides insights into the role of the catalytic metals. During multiple picoseconds dynamics of the **R** state and following windows, the Mg-Mg distance remains stable around ~ 4.9 Å. At the TS^{\ddagger} the two Mg^{2+} ions move closer to each other, displaying a ~1 Å shorter separation distance. At this point, the pro-Sp (O_{Sp}) oxygen approaches MgB (i.e., d5 in Figure 3A decreases). This is consistent with the two-metal-ion catalysis, 5,13 where the two metal ions get closer and cooperatively coordinate the reactant groups (i.e., leaving group, nucleophile and electrophile) to stabilize the TS^{\ddagger} . During the evolution from the \mathbf{R} to the \mathbf{TS}^{\ddagger} , MgA establishes a close contact with the water's oxygen (as shown by constant values of d3). In this way, MgA acts as a Lewis acid, properly orienting and activating the nucleophile. Subsequently (from the TS[‡] to the P), MgA detaches from the OH group (i.e., d3 increases). Simultaneously, the pro-Sp (O_{Sp}) oxygen detaches from MgB (i.e., d5 increases), while MgB stably interacts with O3' (i.e., d6 slightly decreases), facilitating its exit as leaving group. Finally, starting from a RC = 0.4 Å, when the reaction evolves toward **P** formation, the Mg-Mg distance increases, as a sign that the Mg^{2+} ions move apart to facilitate the **P** release.

Alternative Reaction Pathways. To examine possible alternative reaction pathways that could take place within the RuvC active site, we performed a second set of QM/MM MD simulations always starting from the same crystal structure (PDB ID: 5F9R).⁶ This time, with the aim of preserving the original crystallographic active site conformation,

the initial ~40 ps-long equilibration was performed keeping the distance between H983 and MgA constrained to the crystallographic value (details in the SI). As a result of the ab-initio MD equilibration, the water molecule does not directly interact with H983, but it forms a hydrogen bond with the adjacent phosphate (Figures 4 and S6). The obtained reactant was then subjected to QM/MM free energy simulations following the same protocol adopted in the H983-activated chemical mechanism and using the same RC (details are reported in the SI). Interestingly, the catalysis proceeds though a concerted (S_N2-like), phosphate-mediated pathway (Figure 4).^{35,36} Peculiarly, this mechanism shows a dissociative TS^{\ddagger} , at RC = 0.5 Å, where the breaking of P_{DNA} -O3' bond is more advanced than the formation of $O_{WAT}\!\!-\!\!P_{DNA}$ bond (Figure 3B). Immediately after the \mathbf{TS}^{\ddagger} , the water nucleophile spontaneously releases its proton to the adjacent phosphate (Figure 3A, PT'), leading the chemical step to completion. Full description and cross-validation, including the PT mechanism, are reported in the SI (Figure S8). The overall computed Helmholtz free energy reaches a ΔF # value of ~18.55 ± 1.26 kcal/mol, including no large hysteresis (Figure S6B). This pathway appears to be energetically slightly unfavorable when compared to the H983-activated pathway (Figure 4). However, the difference in energy (~2 kcal mol⁻¹) lies within the statistical error of the simulations, thus suggesting that both mechanisms are theoretically possible.

In light of the in-depth description of phosphoryl transfer reactions previously reported^{35,36} our H983-mediated mechanism displays a concerted (S_N2-like) associative pathway, with a similar extent of partial bond formation to the nucleophile and partial bond cleavage to the leaving group at the transition state (TS^{\ddagger} , Figures 2 and 3A). The phosphate-mediated pathway displays instead a shifted TS^{\ddagger} , with slightly more bond

cleavage to the leaving group than bond formation to the nucleophile (Figures 3B and 4). This was defined as a concerted (S_N2-like) dissociative pathway.³⁵ By inspecting the variation of the significant interaction distances along the phosphate-mediated pathway (Figure 3B), we note that, differently to the H983-mediated path (Figure 3A), MgA loosely binds the attacking water, as a sign that it does not act as a Lewis acid. Intriguingly, the distance between MgA and the water nucleophile (d3' in Figure 3B) shows a similar trend to the distance between the pro-Sp oxygen and MgB (d5'). This indicates that when the pro-Sp oxygen of the scissile phosphate loses coordination to MgB and fully moves onto MgA, the water nucleophile concertedly moves away from MgA. These evidences suggest that in the dissociative mechanism, the Mg²⁺ cluster activates the electrophile (i.e., the scissile phosphate), rather than the nucleophile. A similar synchronized dynamics of the MgA-O_{WAT} and MgB-O_{pro-Sp} coordination distances has also been observed in a group II intron ribozyme, ²⁵ which displayed an analogous concerted dissociative pathway. Therefore, this suggests that this dynamical pattern of the metal cluster, and of MgA in particular, might be a hallmark of the concerted (S_N2-like) dissociative cleavage of phosphodiester bonds. On the other hand, analogously to the associative catalysis, MgB stabilizes the oxyanion leaving group. Our simulations show that throughout the entire chemical step (i.e., from R' to P') MgB remains in close contact with O3', with the d6' distance displaying a slight decrease toward products formation. Overall, in agreement with previous findings, 5,13,25 the observed differences in the dynamics of the catalytic metals can be seen as a signature of phosphoryl transfer events in the two-metal-ion enzymes, facilitating the interpretation of their chemical nature.



Discussion

The mechanism of phosphodiester bond cleavage has been studied in several twometal-ion enzymes, ^{13,19–21,23,24,26} enabling comparison with the current investigations. These studies conveyed on the associative nature of the chemical step, which requires the activation of the nucleophile, before the leaving group is dissociated. Differences in the activation of the nucleophile – i.e., whether mediated by a phosphate group, ^{19–22} an amino acid ^{23,24} or the bulk water²⁶ as a general base – have been shown to be crucial for accomplishing the specific enzymatic function, and for the mechanism-based design of artificial enzymes and/or selective inhibitors.³⁸ These previous studies support our finding of a concerted (S_N2 like) associative mechanism of DNA cleavage in CRISPR-Cas9, proceeding upon activation by H983 general base and fundamentally assisted by the joint dynamics of the two catalytic Mg²⁺ ions (Figure 2). A recent study of the HIV-1 RNase H,²⁴ employing a finite-temperature string method and the B3LYP functional,^{18,39} disclosed a reaction mechanism similar to our H983-activated catalysis, with comparable energetics. In that study, a thermally accessible (i.e., with a PT free energy barrier less than ~1 kcal mol⁻¹, and a plateau in the free energy profile of the hydrolysis) proton abstraction by an active site histidine occurs right before a TS[‡] exhibiting the same extent of bond cleavage to the leaving group and bond formation to the nucleophile, analogously to the H983-mediated mechanism described here (Figure 2). A similar associative two-metal-ion mechanism involving an active site histidine has also been found in the DNA polymerase III with exonuclease activity.²³ Taken together, these studies support an associative catalysis of DNA cleavage activated by H983 (a full discussion is reported in the SI).



Our ab-initio investigations also revealed a phosphate-mediated cleavage, which proceeds through a concerted dissociative pathway energetically comparable to the H983activated catalysis (Figure 4). However, mutagenesis and DNA cleavage experiments have probed that the H983A substitution abolishes the RuvC catalysis in CRISPR-Cas9,⁷ supporting the active role of H983 evinced in our H983-activated catalytic mechanism (Figure 2). Moreover, a dissociative pathway has been rarely observed in the two-metaldependent enzymes as either aided by Zn2+,40 promoted by a non-Mg2+-coordinated water^{41,42} or assisted by a single Mg²⁺ ion as recently suggested for GTP hydrolysis.⁴³ A similar pathway has also been reported in a group II intron ribozyme, 25 where a solventmediated mechanism has been proposed. However, nucleolytic ribozymes commonly react through an associative catalysis, activated by guanine nucleobases. 27,44-46 These previous studies indicate that the dissociative catalysis is rare in the two-Mg²⁺-dependent enzymes. suggesting that the alternative phosphate-mediated dissociative mechanism proposed here could be less likely for the RuvC catalysis in CRISPR-Cas9. Notably, RuvC active site resembles that of RuvC resolvase, 47 which processes Holliday junctions during genetic recombination and post-replication repair. In this enzyme, Mg²⁺-coordinating ligands strongly match the DDE motif in Cas9 (i.e., D7, E70, and D146 correspond to the Cas9 residues D10, E762, and D986), while the catalytic H143 locates in the position of H983 in Cas9 (Figure S10). This hints to a similar histidine-activated mechanism, advocating for a common catalytic strategy for genome editing and recombination. Remarkably, a conserved histidine flanking the DDE motif composes the active site of other Cas9 species, 8 most likely implying a similar two-metal-dependent mechanism. It is also important to mention that we cannot exclude that a third metal could engage in the RuvC



catalysis, as suggested for RNA hydrolysis. 48 However, the lack of biochemical and structural data currently hampers the investigation of this hypothesis in CRISPR-Cas9. Finally, it is interesting to note that both chemical pathways identified in this study (Figure 2 and 4) display an endothermic nature, in which, according to the Hammond's postulate, the TS^{\ddagger} more closely resembles the products. Hence, the protonation of the O3' could have an important contribution to the overall drive of the reaction. Considering that the pKa of phospho-monoester groups is ~7, while that of an alkoxide is ~16, it is reasonable to expect that the proton transfer to the O3' would lead to a gain in free energy of ~12-13 kcal mol⁻¹ (accordingly to the general relation $\Delta G = -RT \ln K$). Moreover, in CRISPR-Cas9, upon initial phosphodiester bond cleavage, RuvC can further process the DNA strand, due to not yet clearly characterized conformational changes within the active site. 49 This suggests that the post-cleavage events could further contribute to the stabilization of the product. Notably, conformational changes occurring upon initial phosphodiester bond cleavage have been shown to prime the catalytic function for completion in other two-Mg²⁺ dependent enzymes, as observed in a group II intron ribozyme.⁵⁰ This grants future investigations employing both classical and quantum mechanical approaches to better portray the post-cleavage events.

Overall, our *ab-initio* approach characterizes the two-metal-ion mechanism of nonarget DNA cleavage in the RuvC active site of CRISPR-Cas9, as mediated by the catalytically relevant Mg²⁺, in agreement with experimental data.



Conclusions

Here, quantum-classical MD and free energy methods have been used to elucidate the two-metal-dependent mechanism of DNA cleavage in CRISPR-Cas9. Ab-initio MD simulations, suggest that the cleavage of the DNA non-target strand occurs through a concerted associative mechanism, activated by the conserved H983 and fundamentally assisted by the joint dynamics of the two catalytic Mg²⁺ ions. The catalysis is achieved upon a critical conformational rearrangement within the RuvC active site, which occurs in the presence of the catalytically active Mg²⁺ ions and leads H983 to properly position to act as a general base. This finding resolves prior structural and biophysical controversial evidences that could not ultimately clarify the catalytic role of the conserved H983 and the metal cluster configuration.^{3,6,7} The comparison with other two-metal-dependent enzymes further supports the identified mechanism and suggests a common catalytic strategy for genome editing and recombination. A second set of ab-initio simulations retaining the crystallographic configuration reveal an alternative phosphate-mediated cleavage. Whereas this chemical mechanism also appears energetically viable, experimental evidences depicting H983 as an essential active element for retention of catalysis in CRISPR-Cas9⁷ advocate for the H983-activated mechanism observed here. Intriguingly, the joint dynamics and the role of the metal ions is shown to remarkably differ in the in the two chemical pathways, revealing a hallmark of the enzymatic two-metal-ion mechanism. Overall, this work address a fundamental open question in the CRISPR-Cas9 biology, which has remained unclear from previous experimental studies. The chemical aspects and the mechanistic details of the DNA cleavage provided here are critical for understanding the catalytic function of CRISPR-Cas9, which is at the basis of its use as a genome editing



tool. This knowledge lays the foundation for improving the catalytic efficiency and the specificity of CRISPR-Cas9, paving the way for future therapeutic applications of this gene editing technology.

Materials and Methods

Structural Models. Molecular simulations have been carried out on the activated form of the Streptococcus Pyogenes CRISPR-Cas9 enzymatic complex, which has been based on the most complete X-ray structure of Cas9 in complex with RNA and DNA (5F9R.pdb). This model system has been embedded in explicit waters, and counterions have been added to neutralize the total charge, leading to a periodic simulations cell of \sim 180*120*140 Å³ and a total of \sim 340,000 atoms. Full details are reported in the SI.

Classical Molecular Dynamics (MD). Classical MD has been performed to equilibrate the system prior ab-initio MD, using the Amber ff12SB force field, which includes the ff99bsc0 corrections for DNA⁵¹ and the ff99bsc0+ γ OL3 corrections for RNA. 52,53 The TIP3P model has been employed for water molecules, 54 while the Åqvist force field has been used for Mg²⁺ ions,⁵⁵ in consistency with previous classical studies of two-metal aided enzymes, 13,19,25,56 and of CRISPR-Cas9. 10,11,57 An integration time step of 2 fs has been used. Hydrogen atoms were added assuming standard bond lengths and were constrained to their equilibrium position with the SHAKE algorithm. Temperature control (300 K) has been performed via Langevin dynamics, ⁵⁸ with a collision frequency $\gamma = 1$. Pressure control was accomplished by coupling the system to a Berendsen barostat, ⁵⁹ at a reference pressure of 1 atm and with a relaxation time of 2 ps. The simulations were carried out with the following protocol. The system was subjected to energy minimization



to relax the water molecules and the counter ions, keeping the protein, as well as the RNA, DNA and Mg²⁺ ions fixed with harmonic position restraints of 300 kcal/mol · Å². Then, the system was heated from 0 to 100 K in the canonical ensemble (NVT), by running two NVT 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), in which the restraint was reduced to 25 kcal/mol x Å². Subsequently, all restraints were released and the temperature was raised up to 300 K in a single NPT simulation of 500 ps. After ~1.1 ns of equilibration, ~10 ns of NPT production was carried out allowing the density of the system to stabilize around 1.01 g/cm⁻³. The obtained system has been used as a starting point for QM/MM simulations. Classical MD has been performed with the GPU version of AMBER $18.^{60}$

Ab-initio Quantum Mechanics/Molecular Mechanics (QM/MM) simulations. In the QM/MM simulations, the RuvC active site has been treated at a QM level of theory, while the rest of the system (including explicit waters) has been described using the classical force field. The OM part included the catalytic Mg²⁺ ions and their coordinating protein residues E762, D986, D10, S15, I11 and H983 (protonated in epsilon, as shown by a number of previous studies), 4,11,12 part of the DNA nucleobases C-3 and A-4 and 6 water molecules (Figure S1). This resulted in a total of 105 QM atoms, and additional 13 capping hydrogen atoms used to saturate the valence of the terminal QM atoms. The QM atoms have been described at the QM DFT/BLYP^{17,18} level, while the remaining MM part was treated using the force field described above. OM/MM simulations have been performed using the CPMD code (www.cpmd.org). The wave functions were expanded in a plane

wave basis set up to a cutoff of 75 Ry in a OM cell of dimensions ~26*22*26 Å³. The interactions between the valence electrons and ionic cores were described with normconserving Martins-Troullier pseudopotentials.⁶¹ The QM part was treated as an isolated system, and electrostatic interactions between periodic images were decoupled by the scheme of Tuckerman. 62 A rigorous Hamiltonian treatment of the electrostatic interaction between the QM and MM regions was used. 63 The QM/MM protocol consisted of an initial optimization of the wavefunction, followed by ~6 ps of careful equilibration carried out with Born-Oppenheimer (BO) MD in the canonical (NVT) ensemble using an integration time step of 20 au (~0.48 fs). The temperatures of the QM and MM subsystems were kept constant at 300 K using a Nosé-Hoover thermostat. 64,65 After this initial phase, Car-Parrinello (CP) QM/MM simulations⁶⁶ were carried out with a time step of 5 au (~0.12 fs) and a fictitious electron mass of 600 au. These unconstrained ab-initio simulations have been carried out reaching ~40 ps of sampling in two replicas, initialized from different coordinates and velocities. The final configuration has been simulated for other ~40 ps of unconstrained ab-initio MD. Additional ~40 ps of ab-initio MD have also been carried out in the presence of the crystallized Mn²⁺ ions⁴. Full details are reported in the SI.

Free energy simulations. The free energy profiles associated to phosphodiester bond cleavage have been obtained through QM/MM simulations and free energy methods. The "blue moon ensemble" method, 67,68 along with thermodynamic integration have been employed to investigate phosphodiester bond cleavage, using the difference in distance between the breaking (O3'DNA-PDNA) and forming (OWAT-PDNA) bonds as a Reaction Coordinate (RC). The appropriateness of this approach and of the employed RC have been shown in several studies of DNA/RNA processing enzymes. 19,20,24,25 and are further discussed in the SI. Starting from a value of the RC = -1.5 Å (i.e., the reactant \mathbf{R} state) we sampled in 20 sequential windows along the RC, with a resolution of 0.2 Å (0.1 Å in the region in the vicinity of the TS^{\ddagger}). A growth rate of 0.1 Å/ps has been used to move from a window to the following one. Then, each window has been simulated for ≥6 ps, reaching convergence of the constraint force, and collecting a total of ab-initio QM/MM MD of ~120 ps. By using this approach, we computed two free energy profiles, starting from two different configurations shown in Figure 1B (right and left panels, respectively). This resulted in a total of ~240 ps of sampling. In order to estimate the error associated to hysteresis, we also computed the backward reaction pathways. For each free energy profile, we selected a point after the TS^{\ddagger} was reached and we followed the backward pathway by sampling over 8 consecutive windows for ≥6 ps each (a complete description is given in the SI). This reached a total sampling of additional ~90 ps. Notably, these calculations revealed no large hysteresis between the forward and backward pathways (Figure S7). Finally, for both free energy profiles, at a RC = 1.2 Å, the constraint has been released for ~5 ps, enabling the system to spontaneously reach the product state. The derivation of all free energy profiles (both forward and backward pathways) has been performed by considering the average constraint force within each window from the points in which the force reached the convergence along the equilibrated trajectory. The statistical error at each point of the free energy profiles (both forward and backward pathways) has been computed by error propagation analysis. Finally, the overall error on the free energy barrier has been estimated as the sum of the statistical error and the error due to hysteresis between the forward and backward pathway (Figure S7). This resulted in a total

Helmholtz's free energy of $\sim 16.55 \pm 1.22$ kcal mol⁻¹ for the associative free energy profile (Figure 2) and $\sim 18.55 \pm 1.26$ kcal mol⁻¹ for the dissociative free energy profile (Figure 4). As in previous studies based on ab-initio MD, 16,19,34 further exploration of the identified chemical mechanism included the explicit calculation of the proton transfer in the H983-activated pathway. These simulations required additional ~33 ps of sampling and are described in the SI. The SI comprehensively describes the application of ab-initio MD and free energy methods.

Supporting Information

Supplementary Materials and Methods, Results, Figures, Movies and References. The Supporting Information is available free of charge on the ACS Publications website.

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Competing interests

The authors declare no competing financial interest.

Figures

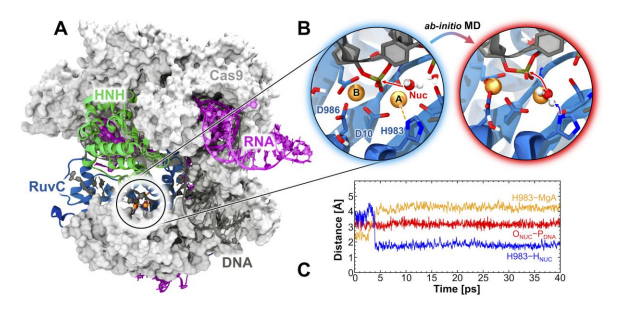


Figure 1. Overview of the CRISPR-Cas9 complex bound to a guide RNA and to a target DNA. (A) The Cas9 protein is shown in molecular surface, highlighting the HNH (green) and RuvC (blue) domains. The RNA (violet) and the DNA (black) strands are shown as ribbons. (B) Inset of the RuvC active site, displaying the catalytic metals (A and B, orange), the surrounding protein residues and water molecules (left panel). The configuration of the active site resulting from ab-initio molecular dynamics (MD) is shown on the right panel. The water nucleophile locates in between H983 and the scissile phosphate, positioning for the chemical reaction. (C) Time evolution along ~40 ps of abinitio MD of the interaction network established by H983 and by the water nucleophile. The complete set of interactions established by the metal cluster is reported in Figures S2-3.



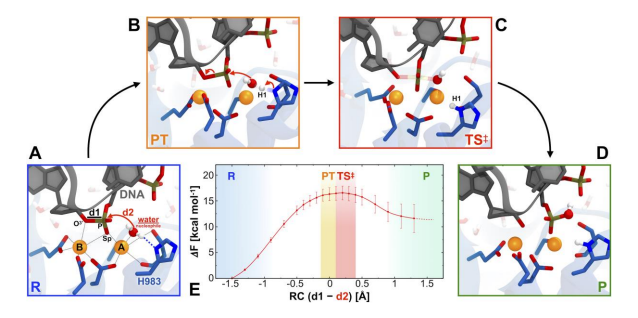


Figure 2. Structural and energetic properties of the two-Mg²⁺ aided catalysis in CRISPR-Cas9. (A) Representative snapshots of the Reactant (R), (B) Proton Transfer (PT), (C) Transition State (TS[‡]) and (D) Product (P) states along phosphodiester bond cleavage. (E) The free energy profile (ΔF #, in kcal mol⁻¹) has been computed by using the difference in distance between the breaking (O3'_{DNA}-P_{DNA}, d1) and forming (O_{WAT}-P_{DNA}, d2) P-O bonds as a Reaction Coordinate (RC = d1-d2, highlighted in panel A). The RC windows corresponding to the R, PT, TS[‡] and P states are highlighted using colored bars.



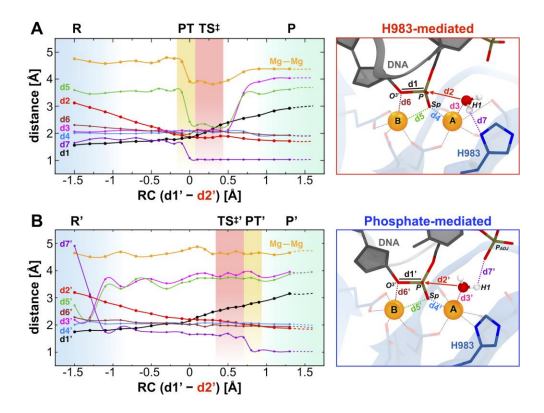


Figure 3. Interaction distances along phosphodiester bond cleavage.

(A) Variation of the significant interaction distances (i.e., the Mg-Mg and the d1-d7 distances, shown on the right panel), computed at each step of the Reaction Coordinate (RC) in the associative reaction pathway activated by H983 (described in Figure 2). (B) Variation of the critical interaction distances along the alternative dissociative reaction pathway (described in Figure 4). The RC windows corresponding to the Reactant (R), Proton Transfer (PT), Transition State (TS ‡) and Product (P) states are highlighted using colored bars. The RC (i.e., difference in distance between the breaking and forming P–O bonds) and the interaction distances are described on the R in the right panels.



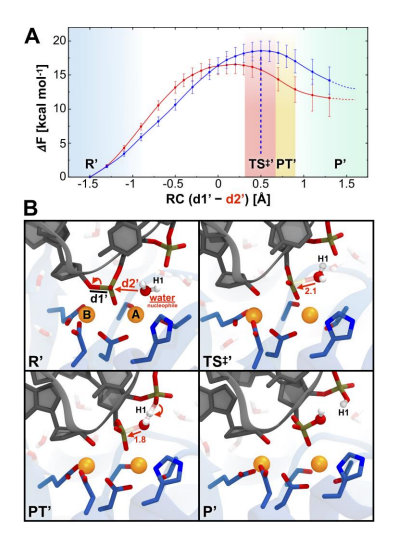


Figure. 4. Alternative phosphate-mediated reaction pathway. (A) Free energy profile ($\Delta F\#$, in kcal mol⁻¹) for the phosphate-mediated dissociative mechanism (blue line), highlighting regions corresponding to the reactant (R'), transition state (TS[‡]'), proton transfer (PT') and products (P'). The free energy profile for the associative pathway activated by H983 is also shown (red line). (B) Snapshots of the R', TS[‡]', PT' and P' states, as from the dissociative pathway.

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