Atomic-scale insights into allosteric inhibition and evolutionary rescue mechanism of \textit{Streptococcus thermophilus} Cas9 by the anti-CRISPR protein AcrIIA6

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\section*{A B S T R A C T}

CRISPR-Cas systems are prokaryotic adaptive immunity against invading phages and plasmids. Phages have evolved diverse protein inhibitors of CRISPR-Cas systems, called anti-CRISPR (Acr) proteins, to neutralize this CRISPR machinery. In response, bacteria have co-evolved Cas variants to escape phage's anti-CRISPR strategies, called anti-anti-CRISPR systems. Here we explore the anti-CRISPR allosteric inhibition and anti-anti-CRISPR rescue mechanisms between \textit{Streptococcus thermophilus} Cas9 (St1Cas9) and the anti-CRISPR protein AcrIIA6 at the atomic level, by generating mutants of key residues in St1Cas9. Extensive unbiased molecular dynamics simulations show that the functional motions of St1Cas9 in the presence of AcrIIA6 differ substantially from those of St1Cas9 alone. AcrIIA6 binding triggers a shift of St1Cas9 conformational ensemble towards a less catalytically competent state; this state significantly compromises protospacer adjacent motif (PAM) recognition and nuclease activity by altering interdependently conformational dynamics and allosteric signals among nuclease domains, PAM-interacting (PI) regions, and AcrIIA6 binding motifs. Via \textit{in vitro} DNA cleavage assays, we further elucidate the rescue mechanism of efficiently escaping AcrIIA6 inhibition harboring St1Cas9 triple mutations (G993K/K1008M/K1010E) in the PI domain and identify the evolutionary landscape of such mutational escape within species. Our results provide mechanistic insights into Acr proteins as natural brakes for the CRISPR-Cas systems and a promising potential for the design of allosteric Acr peptidomimetics.

\section*{1. Introduction}

The evolutionary arms race between bacteria and viruses (phages) has been ongoing for millions of years [1–5]. Suffering from constant predation by viruses, bacteria have evolved a number of either innate or adaptive immune mechanisms to protect themselves against frequent assault [6–10]. Among the diverse anti-viral defences, the clustered regularly interspaced short palindromic repeats (CRISPR) system, which cleaves invading genetic elements in a sequence-specific manner, is a form of bacterial adaptive immunity against bacteriophages and represents a milestone in this coevolutionary arms race [11–14]. This specificity and easily programmable nature of the single-protein (type II) CRISPR-Cas9 endonuclease led to its exploitation by researchers to achieve facile gene manipulation [11,15–22].

CRISPR-Cas9 function begins with assembly of the surveillance complex, in which the Cas9 endonuclease associates with single guide RNA (sgRNA) carrying CRISPR RNA (crRNA) transcribed from previously encountered invading genetic elements to form a ribonucleoprotein complex (RNP) [13,23,24]. Subsequent double-strand DNA (dsDNA) recognition and interference by Cas9 requires identification of the protospacer adjacent motif (PAM) located

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immediately downstream of the target sequence before further substrate DNA targeting [24–31]. Previous structural characterisation of Streptococcus thermophilus Cas9 (St1Cas9), the model system that kicked off the CRISPR craze, revealed a bilobed architecture composed of a recognition (REC) and nuclease (NUC) lobes connected through an arginine-rich bridge helix (BH) [2,32–35] (Fig. 1 A–D). The REC lobe primarily engages in nucleic acid binding, mediating sgRNA loading and RNA-DNA heteroduplex recognition [24,36–38]. The NUC lobe includes two important catalytic HNH and RuvC nuclease domains that perform synchronised site-specific cleavage of the corresponding target DNA (tdNA) and nontarget DNA (ntDNA) strands, and the two nuclease domains are interconnected by L1 and L2 linkers. It also includes an α/β wedge (WED) and PAM-interacting (PI) domains [collectively referred to as topoisomerase-homology (TOPO) and C-terminal domains (CTD)] connected to the catalytic cores via a phosphate lock loop (PLL). The PAM-containing DNA duplex is recognised and accommodated by WED and PI domains. Identification of a matching PAM sequence initiates dsDNA unwinding and heteroduplex formation, which ultimately triggers St1Cas9 conformational activation characterised by prominent RuvC and HNH relocation for efficient substrate DNA cleavage [2,39–42].

To counteract this robust anti-viral immunity, phages utilise anti-CRISPR proteins (Acrs) that bind and interfere with CRISPR-Cas function in a sequence-independent manner [5,43–48]. As natural antagonists of Cas endonucleases, Acrs have garnered intense research interest for their potential in restraining the powerful CRISPR-Cas machinery and enhancing the precision at which genetic perturbations can be made [49–51]. To date, ~45 diverse Acrs targeting different CRISPR-Cas systems have been identified [45,52–54], and bioinformatic analysis has suggested that Acrs are likely far more widespread than expected [55–58]. Acrs are found to interfere with CRISPR-Cas in a variety of manners [53,59], including inhibiting nucleic acid recruitment during RNP assembly (AcrIF3) [60], competing with sgRNA loading (AcrIIc2) [61,62], PAM binding (AcrIIa2, AcrIIa4) [63–65], DNA hybridisation (AcrIF1) [66], and blocking catalytic residues (AcrIIC1) [47]. Among the identified Acrs (AcrIIA1–AcrIIA6 and AcrIIC1–AcrIIC3) that function against the type II CRISPR-Cas system [43,46,67], only the newly reported AcrIIa6 possesses specific inhibition towards St1Cas9 [2,45,55]. A cryo-electron microscopy (cryo-EM) structure shows that AcrIIa6 binds to the back face of St1Cas9 RNP which is not primarily engaged in substrate DNA recognition and binding, nonoverlapping with the sgRNA and dsDNA binding sites (Fig. 1E). AcrIIa6 consists of 183 amino acids and predominantly exists as a symmetric homodimer (monomers referred to as heterodimers) to elucidate the conformational reorganisation of St1Cas9, a system that forms a single-stranded 400-nt RNA strand (sgRNA) (Supplementary Fig. 2). Each system underwent 3 μs × 5 independent runs with random initial velocities, leading to a cumulative simulation timescale of 120 μs. After reaching relative equilibrium after 500–ns MD simulation, a ~1.3-fold increase in root-mean-square deviation (RMSD) value for St1Cas9 in the St1Cas9-sgRNA-RNP was observed in the presence of AcrIIa6, indicating considerable conformational variation compared with that of the initial structural complex (Supplementary Results 2.1 and Supplementary Fig. 3). Moreover, comparing the average structures of SpCas9-sgRNA-AcrIIa6 and SpCas9-sgRNA throughout the simulations revealed that, owing to the absence of the WED domain in SpCas9 that was involved in both PAM recognition and AcrIIa6 interaction in St1Cas9, AcrIIa6 was incapable of binding to SpCas9-sgRNA, as well as other bacterial species lacking the WED domain (Supplementary Fig. 4). This result was consistent with the biolayer interferometry experiments that revealed the insensitivity of SpCas9 to AcrIIa6 [2].

To uncover the domain-specific influence of AcrIIa6, we analysed the per-residue root mean square fluctuations (RMSF) of St1Cas9 (Supplementary Results S2.1 and Supplementary Fig. 5),...
and projected the RMSF difference between simulation systems with dissimilar compositions onto the initial St1Cas9 structure for more intuitive representations (Supplementary Fig. 6). Despite significantly increased residue fluctuations of HNH and WED domains in the presence of AcrIIA6, the most prominent RMSF was observed within the L1 that bridges the catalytic RuvC and HNH domains. In addition, L2 exhibited diverse fluctuations under different situations, in which AcrIIA6 binding was found to enhance L2 flexibility. Notably, the catalytic HNH domain resides topologically distant from the direct AcrIIA6 binding interface. Thus, the increased HNH flexibility in St1Cas9/C1 sgRNA/C1 AcrIIA6 is a good demonstration of the allosteric effects of AcrIIA6, in which the flexible L1 and L2 linkers may play a pivotal role as allosteric transducers [76]. Furthermore, although it is involved in direct interaction with AcrIIA6, the WED domain had a more primary function as a key component in mediating target DNA recognition and binding together with the PI domains (containing TOPO and CTD), whose fluctuations remained largely unchanged after AcrIIA6 binding. The prominently altered conformational flexibility of proximal WED and distal HNH domains support the notion that AcrIIA6 acts through modifying DNA-associated conformational dynamics of the WED domain, which also allosterically enhanced plasticity of HNH and positioned the nuclease domain in an unfavourable conformation for subsequent DNA cleavage (Supplementary Results S2.1).

2.2. AcrIIA6 induces St1Cas9 conformational transitions from ‘approaching’ to ‘departing’

Principal component analysis (PCA) was performed to characterise the global conformational transition pattern and domain dynamics (Supplementary Methods). The conformational space sampled by St1Cas9 in each system was investigated using the two most representative collective principal components (PC1 and PC2). Distinct conformations were adopted by St1Cas9 dependent on AcrIIA6 binding, as evidenced by the dominant conformational clusters that were well distinguished along PC1, with PC1 < 0 corresponding to the AcrIIA6-inactivated St1Cas9 (Fig. 2A and Supplementary Fig. 7). Further insights into the domain dynamics along PC1 revealed a ‘departing’ conformation, characterised...
mainly by relocations of the HNH, WED, and PI domains in the St1Cas9/C1 sgRNA/AcrIIA6 complex relative to the ‘approaching’ conformation observed without AcrIIA6 (Fig. 2B and Supplementary Movie 1). In the departing conformation, the WED and PI domains synchronously moved towards AcrIIA6 at the back of the RNP; this movement constricted the DNA recognition groove (Fig. 2E, H), which may be too narrow to accommodate the DNA duplex when searching throughout the genomic landscape for matching PAM sequences—the prerequisite for initiating DNA unwinding and melting. Meanwhile, the catalytic HNH domain shifted away from the central concave region responsible for heteroduplex accommodation (Fig. 2D, G), which can result in increased spatial distance between the catalytic and cleavage sites and is characteristic of Cas9 under an inactivated state. Additionally, comparing the prin-
cipal motions of St1Cas9 in St1Cas9-sgRNA and St1Cas9-sgRNA-dsDNA, whose conformational clusters were sepa-
rated along PC2, indicated closure of the bilobed architecture with inward movement of the catalytic domains after dsDNA loading (Fig. 2A, C, F, I, and Supplementary Movie 2), which was in agree-
ment with the classic open-to-closed conformational transition [27,42,76,81,82]. Based on these observations, PC1 can be viewed as a parameter measuring the ability of St1Cas9 RNP to recognise PAM sequences and associate with complementary dsDNA (DNA-
relevant dynamics), as both WED and PI domains are primarily implicated in DNA recognition and accommodation-relevant events, while PC2 indicates the potential of the surveillance complex to transit into the ‘closed’ catalytically active state, in which prominent inward relocations of REC and NUC lobes occurred. Our simulation results indicate that AcrIIA6 showed compromised inhibition towards St1Cas9-sgRNA-dsDNA, as evidenced by the location of major conformational clusters for St1Cas9-sgRNA-dsDNA-AcrIIA6 with PC2 < 0, which represented the catalytically active closed conformation (Fig. 2A).

Collectively, the results indicate that association of St1Cas9 with the allosteric effector AcrIIA6 leads to altered St1Cas9 confor-
mational dynamics with departing characteristics. In addition to the prominent conformational rearrangements of WED and PI domains, AcrIIA6 triggered relocation of the nuclease HNH domain away from the central concave for heteroduplex accommodation, resulting in unfavourable PAM recognition and catalytic activation and ultimately decreasing DNA binding affinity and catalytic activity of St1Cas9 RNP (Supplementary Results S2.2).

2.3. Contraction of WED and PI domains of St1Cas9 upon AcrIIA6 binding hinders efficient PAM interrogation

Based on the results from PCA analysis where distinct behaviour of WED and PI domains that might be unfavourable for the binding of incoming substrate DNA in the presence of AcrIIA6, we con-
ducted further experiments to dissect the detailed dynamics and molecular basis underlying such conformational shift. Direct interactions of AcrIIA6 with St1Cas9 induced significant structural rearrangements of the WED and PI domains, whose back face is responsible for PAM recognition – the first of the sequential events leading to target cleavage. To investigate the dynamic features of WED and PI associated with both PAM recognition and AcrIIA6 binding, two order parameters were calculated based on structural information from the cumulative 15-μs MD sampling of each sim-
ulation system. One parameter was defined as the perimeter of the triangle formed by the WED residue K837 and two PI residues K1049 and N1079 (ÅK837-K1049-N1079), which reflects the relative degree of openness of regions associated with PAM recognition and subsequent dsDNA accommodation. The other parameter was defined by the volume of one of the major AcrIIA6 binding pockets for AcrIIA6-h1 β2-β3 hairpin insertion, which is sandwiched between WED and PI domains and constitutes a major interface for intramolecular recognition between AcrIIA6 and St1Cas9 RNP [2], and can monitor the binding affinity between St1Cas9 RNP and AcrIIA6 (graphic representation of the two order parameters is shown in Supplementary Fig. 8).

The potential of mean force (PMF) as a function of the selected order parameters indicated that PAM recognition and AcrIIA6 binding, both mediated by WED and PI domains, were mutually exclusive events to some extent, leading to activation and inhibition of St1Cas9 RNP, respectively (Fig. 3 and Supplementary Table S1). In St1Cas9-sgRNA (Fig. 3A), two distinct states were observed: one with a smaller ÅK837-K1049-N1079 perimeter but a larger AcrIIA6 binding pocket volume (C'), and the other was the opposite (Cc). The coexistence of C' and Cc coincided with the propensity of St1Cas9 to be bound by either AcrIIA6 or dsDNA, indicating an innate equilibrium between the two states with different WED and PI domain configurations (Supplementary Fig. 9A). Upon association with dsDNA (Fig. 3B), as a result of WED and PI conformational rearrangements, the pocket volume for AcrIIA6-b2-β3 hairpin insertion decreased significantly compared with that of Cc before DNA loading. This reduction was accompanied by a change in pocket shape, hampering St1Cas9-sgRNA-dsDNA complex recruitment of AcrIIA6. Meanwhile, the smaller ÅK837-K1049-N1079 perimeter in the St1Cas9-sgRNA-dsDNA compared with that of Cc in the St1Cas9-sgRNA supported the regional con-
traction of WED and PI domains for tight association with dsDNA (Supplementary Fig. 9B–D), in which residues from St1Cas9 WED, TOPO, and CTD domains (K867, S961, V962, S963, K1025, and N1026) established stable polar contacts with the phosphoribosyl backbone of the PAM sequence, especially with nucleotides dC2*, dA3*, dG4*, and da7* (Fig. 3G, H and Supplementary Fig. 10). Upon binding of AcrIIA6 (Fig. 3C), the pocket volume was retained at a high level; however, a drastic decrease in ÅK837-K1049-N1079 perimeter was observed. This indicated a shrinkage of the groove encompassed by WED and PI domains, which is insufficient for PAM recognition and dsDNA accommodation (Supplementary Fig. 9C, E, F). Even in the presence of dsDNA, contraction of the PAM rec-
novation groove was only slightly relieved (Fig. 3C, D). The flat groove resulting from AcrIIA6 binding markedly weakened overall interactions between PAM and St1Cas9 by increasing conformational plasticity of dsDNA, which underwent an approximate 5.2 Å inward shift towards St1Cas9 to avoid potential structural conflicts with the deformed PAM recognition groove (Fig. 3G, H, and Supplementary Table S2). Altogether, the results indicate that contraction of WED and PI domains in the St1Cas9-sgRNA induced by AcrIIA6 binding would compromise PAM recognition and DNA binding capability of St1Cas9 RNP, leading to inhibition of the surveillance complex (Supplementary Results S2.3).

2.4. Counter-effects of WED and PI conformational dynamics on AcrIIA6

Given the significant weakening of intermolecular recognition between dsDNA and St1Cas9 in the presence of AcrIIA6, we subse-
quently probed the AcrIIA6-St1Cas9 RNP binding interface of dif-
ferent systems. Our MD simulation results demonstrated a mixed protein-nucleotide interface recognised by AcrIIA6, which inserted structural motifs from both monomers into the clefts formed within St1Cas9 RNP to facilitate stable anchoring (Fig. 4A, B). The backbone of stem loop 1 (ST1) from sgRNA (G71, C72, U73, C75) strongly interacted with the AcrIIA6-a1 β2-β3 hairpin (T118) and L9 loop (G167, R168, N170), constituting the nucleotide interface for AcrIIA6 docking in St1Cas9 RNP. The protruding hairpin (R120, Y122, A124, N127, Y128, A130) and L8 loop (G146, S147, Q151) of AcrIIA6 extensively interacted with St1Cas9 residues from WED (G943, D947) and PI (D987, Q989, K1010) domains. In addition, the AcrIIA6-a5-a6 helix residue N83 established strong polar interactions with the PI (F1019, D1115) domain. However, the density and intensity of the interaction network between St1Cas9 RNP and AcrIIA6 markedly changed in response to dsDNA binding (Fig. 4C, D, Supplementary Fig. 12, 13, and Supplementary Table 3, 4). Despite a few weakened contacts (e.g. AcrIIA6-R168 and U73; AcrIIA6-A124 and K1010), the overall intermolecular interaction network was strengthened owing to a few newly established interactions. However, the molecular mechanics/Poisson Boltzmann surface area (MM/PBSA) binding free energy calculations between AcrIIA6 and St1Cas9 RNP with/without dsDNA sug-
gested reduced affinity towards St1Cas9-sgRNA-dsDNA...
Such unfavourable binding may be attributed to a prominent decrease in the interfacial area between St1Cas9 RNP and AcrIIA6 upon dsDNA loading, in which both sgRNA and St1Cas9 engaged a smaller area for interaction with AcrIIA6 (Supplementary Table 6). Interestingly, AcrIIA6 as a dimer was also destabilised upon binding to DNA-loaded St1Cas9 RNP, as revealed by the reduced binding free energy between AcrIIA6 monomers (Supplementary Table 5). Thus, together with the compromised binding affinity towards St1Cas9-sgRNA dsDNA, the counter-effect on the AcrIIA6 dimer by conformational dynamics associated with dsDNA may contribute to the inefficient inhibition of St1Cas9-sgRNA dsDNA by AcrIIA6.

2.5. AcrIIA6 alters conformational plasticity of the distal HNH domain

The outward displacement of the nuclease HNH domain is a unique feature of the AcrIIA6-induced global conformational tran-
Fig. 4. AcrIIA6-St1Cas9 RNP interaction network in different systems. Structure of representative conformations and magnified view of the molecular interactions established between AcrIIA6–sgRNA (top) and AcrIIA6–St1Cas9 (bottom) in (A, B) St1Cas9/C1 sgRNA/AcrIIA6, (C, D) St1Cas9/C1 sgRNA/dsDNA/AcrIIA6, and (E, F) mut St1Cas9/C1 sgRNA/AcrIIA6 (K1010E mutation). The landscape was generated with the perimeter of triangle $\Delta_{337-534-1075}$ of the pocket volume for AcrIIA6 $\beta_2$-$\beta_3$ hairpin insertion. The fraction of each cluster is indicated in brackets. (G) Representative structure extracted from energy basin CDA in the St1Cas9/C1 sgRNA/C1 dsDNA/AcrIIA6 system. Nucleotide components (translucent ones) from C5 in St1Cas9 sgRNA/dsDNA is superimposed onto the nucleic backbone in C1A as contrast; the $\sim$5.2 Å inward shift of the dsDNA duplex in the presence of AcrIIA6 is denoted by a curved arrow. St1Cas9 domains are colour coded as in Fig. 1A. Molecular interactions between PAM nucleotides and St1Cas9 residues in (H) St1Cas9/C1 sgRNA/C1 dsDNA and (I) St1Cas9/C1 sgRNA/C1 dsDNA AcrIIA6.
sition to the departing conformation. We then investigated into St1Cas9 HNH domain conformational dynamics upon AcrIIA6 binding. As revealed by the PCA analysis, AcrIIA6 binding induced outward movement of the distal HNH domain. We suspected that such departure from the catalytic core would result in significant barrier for St1Cas9-sgRNA-AcrIIA6 to attain the catalytically active state even if dsDNA can be loaded onto the complex, thus, we employed the most prevalent conformation from St1Cas9-sgRNA-dsDNA (C\text{st1}) which well reflected the catalytically competent state to draw a comparison with representative St1Cas9 structure from St1Cas9-sgRNA-AcrIIA6 (C\text{ac}). Comparison between the two representatives showed that, compared with the catalytically-competent state, the HNH domain of St1Cas9 exhibited ~59° rotation around the L2 hinge that led to departure from the central concave upon AcrIIA6 binding (Fig. 5A, B, measured when superimposing the two structures with Pymol). This outward relocation further increased the distance by 12.57 Å between the catalytic residue HS99 in the HNH domain and the presumed location for scissile phosphate on the target strand (Fig. 5C). Compared with St1Cas9-sgRNA, dsDNA loading did not affect HNH conformational plasticity but significantly enhanced structural flexibility of L1 and rendered L2 less stable (Fig. 5D and Supplementary Fig. 14A, B). This allowed the inward movement of the HNH nuclease domain during conformational transition toward the closed catalytically active state in the presence of substrate DNA (also detailed in above PCA section). In contrast, the association of St1Cas9-sgRNA with AcrIIA6 reduced the original structural stability of both HNH and L1 but stabilised L2 (Fig. 5E and Supplementary Fig. 14C), thereby permitting rotation-based outward movement of the HNH nuclease domain, which was also opposite to the moving direction induced by dsDNA. The opposite structural stability of L1 and L2 linker of St1Cas9-sgRNA would indeed result in distinct HNH motion: when L2 loosens, the HNH domain trends to move towards L2 which resides near the main body of St1Cas9 and thus enables activation; while L1 loosens, the HNH domain falls to the rim of St1Cas9 nuclease as L1 lies in the relatively outer part and ultimately contributed to inhibition (Fig. 5D, E). In the presence of either dsDNA or AcrIIA6 binding, interactions between HNH and the remaining domains in St1Cas9 decreased significantly—as revealed by the reduced number of hydrogen bonds established between them (Supplementary Fig. 15) – with greater HNH flexibility in St1Cas9-sgRNA-AcrIIA6.

2.6. Correlated motion between functional domains mediates AcrIIA6 allosteric inhibition

Generalised correlation (GC\textsubscript{i,j}) analysis, which captures both linear and nonlinear correlations based on Shannon entropy, was conducted to identify dynamic correlations between residues [83]. The results showed that binding of the allosteric inhibitor AcrIIA6 reshaped the overall correlation structure and strengthened the correlated motions within St1Cas9 (Supplementary Fig. 16A, C, G), as evidenced by the altered GC\textsubscript{i,j} distribution in St1Cas9-sgRNA (~0.67) to a more correlated state in St1Cas9-sgRNA-AcrIIA6 (~0.72). This increase in coupled motion was consistent with the allosteric characteristic of AcrIIA6, which relied on concerted motions between distal sites for global conformational reassembly. Notably, compared with substrate DNA binding, an association with AcrIIA6 induced more significant enhancement of correlated movements between residues, especially between REC components and the remaining part of St1Cas9. Previous studies have suggested that REC motions dynamics play a role in the open to closed conformational transitions of Cas9 [38]. The altered correlation in our study suggests that AcrIIA6 binding may disrupt REC-mediated motional correlation and hinder further conformational transitions underlying endonuclease activation.

We further investigated interdependent motion between functional domains based on the obtained residue GC\textsubscript{i,j} matrix (Fig. 6). The St1Cas9-sgRNA complex possessed the weakest inter-domain correlation network, in which only the REC lobe displayed a relatively strong correlation with domains from the NUC lobe (Fig. 6A). Loading of dsDNA increased overall domain connectivity (Fig. 6B), with enhanced correlation between HNH and PI domains, confirming the ability of direct AcrIIA6-binding sites to allosterically regulate the catalytic activity of remote nuclease domains. Meanwhile, HNH and RuvC nuclease domains established stronger motion correlation upon dsDNA loading, which is the basis of synchronised DNA double-strand cleavage. AcrIIA6 binding also strengthened the correlation map within St1Cas9 and particularly enhanced the interdependent conformational dynamics between the REC and NUC lobes (Fig. 6C), indicating the role of REC in coordinating nuclease activity in AcrIIA6-induced allostery. In addition to the strong correlation between the nuclease RuvC and HNH domains, WED and PI domains also displayed stronger inter-correlation, which was supported on the structural level by their synchronous movement towards a contracted conformation with a shrunken DNA binding groove during AcrIIA6-induced conformational structural transition to the departing state.

Collectively, the results indicate that changes in domain motion correlations induced by either substrate DNA or AcrIIA6 binding underlie the structural transition to the catalytically active closed or inactive departing state as well as the coupled motion mediated by WED and PI along the direct AcrIIA6 binding interface. Furthermore, the RuvC and HNH nuclease domains are recognised as key elements in inhibitor-induced allostery.

2.7. AcrIIA6 induces community reorganisation and allosteric signalling

Dynamic network analysis was performed to uncover the correlation structure in different simulation systems, and closely related residues were clustered into ‘communities’ inter-connected with each other. The obtained community networks were visualised as spheres connected by sticks, whose widths were proportional to the connectivity strength of the edges bridging them (Fig. 7A–F). In St1Cas9 RNP, community 9 (Fig. 7A), composed predominantly of BH residues, served as a major hub for information transfer between the REC and NUC lobes, while community 10, which represented PI domains, only loosely interacted with the remaining contact network (domain composition for each community is listed in Supplementary Table 7). The recognition and loading of dsDNA reduced the community number and enhanced allosteric information transfer (Fig. 7B). Under such circumstances, the PI domain-containing community 10 established stronger correlation with both WED (community 7) and RuvC (community 16) domains, while the latter exhibited enhanced connection with community 14, which largely consisted of the HNH nuclease domain and L2 linker through community 13. This strong signal transduction was in consensus with the experimentally identified tight correlations between the catalytic activity of the two nuclease cores mediated by the linker bridging them. Binding of AcrIIA6, as shown in Fig. 7C, fragmented the community structure of St1Cas9 RNP, but strengthened global allosteric signalling. AcrIIA6 was mainly found deposited in two communities (community Acr\textsuperscript{A} and Acr\textsuperscript{B}), each of which consisted predominantly of residues from an AcrIIA6 monomer. AcrIIA6\textsuperscript{A} was found to establish extensive contacts with St1Cas9 residues/nucleotides, reflected by its coexistence with WED, TOPO, CTD, and sgRNA components in the same community (community 4, 6, and 10), whereas AcrIIA6\textsuperscript{B} exhibited
Fig. 5. AcrIIA6 alters conformational plasticity of the distal HNH domain. Molecular surface representation with L1, HNH, and L2 in representative conformations from (A) C_p in St1Cas9-sgRNA-dsDNA and (B) C_e in St1Cas9-sgRNA-AcrIIA6. (C) Probability distribution map of the distance between catalytic H599 and scissile phosphate (T3) in St1Cas9-sgRNA-dsDNA (orange) and St1Cas9-sgRNA-AcrIIA6 (yellow; presumed cleavage site inferred from structural alignment with St1Cas9-sgRNA-dsDNA). All five productive runs were calculated as DSSP plots for real-time secondary structure transitions within the L1, HNH, and L2 regions in (D) St1Cas9-sgRNA-dsDNA and (E) St1Cas9-sgRNA-AcrIIA6. Position of catalytic H599 is indicated by a blue outlined arrow along the vertical axis indicating residue number. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

Fig. 6. Inter-domain correlation of St1Cas9 in different simulation systems. Interdomain correlations calculated by accumulating per-residue CS\textsubscript{inter} within each domain in (A) St1Cas9-sgRNA, (B) St1Cas9-sgRNA-dsDNA, (C) St1Cas9-sgRNA-AcrIIA6, (D) St1Cas9-sgRNA-dsDNA-AcrIIA6, (E) mut St1Cas9-sgRNA, and (F) mut St1Cas9-sgRNA-AcrIIA6. The BH and PLL domains were left out for clarity. RuvC collectively refers to RuvC-I, RuvC-II, and RuvC-III, while PI contains both TOPO and CTD domains.
more exclusive allosteric information transfer through the CTD domain, which was consistent with the stable interaction observed between AcrIIA6 B N83 and CTD residues in our analysis. The presence of AcrIIA6 further wired community 16 as the signal transduction hub in the community network of St1Cas9 RNP, in which it received regulatory signals from community 10 and further allosterically promoted them toward the distal REC lobe (community 1, 3) and catalytic HNH domain (community 14). Under both dsDNA-loaded and AcrIIA6-bound conditions, community 13, whose major components were L1 residues, mediated informational flow between the two catalytic domains (community 14 and community 16), which is in line with previous studies showing L1 as an allosteric transducer [42,81]. The bidirectionally curved black arrow indicates the information transfer pathway; sky blue arrow indicates AcrIIA6-specific signalling pathways; signalling pathways shared between catalytic residues and PIRs and those from AcrIIA6 are highlighted with sky-blue contour. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

Similar effects on the community network of St1Cas9 RNP were observed by either AcrIIA6 binding or dsDNA loading—both exhibited enhanced signal transduction between communities composed of WED, PI, and more distal subgroups within HNH and REC via the RuvC domain (community 16). Such similarities in community reorganisation may contribute to the prominent HNH relocation induced by both dsDNA and AcrIIA6, although in opposite directions. Nonetheless, the inhibitor indeed strengthened the crosstalk between communities to a greater extent, as represented by thicker inter-community sticks, which may well underlie the more prominent outward displacement/rotation of HNH domain during AcrIIA6-induced transition from approaching to departing conformation.

We next aimed to elucidate the detailed information transfer pathway underlying AcrIIA6-induced WED and PI contraction and HNH outward reorientation as well as how the catalytic domains coordinate with each other in this process. To this end, we computed the optimal residue-by-residue signalling pathway
connecting AcrI6 (T118Δ, A124Δ, S147Δ, N83Δ), PAM-interacting residues (PIRs, including WED: K867, S961, V962, S963; TOPO: K1025; CTD: N1026), HNH (H599), and RuvC (D9) catalytic sites. The obtained signalling pathways are summarised on both the residue (Supplementary Fig. 17, 18) and simplified domain level (Fig. 7G–L). In St1Cas9 RNP alone (Fig. 7G), signals from PIRs were transmitted to HNH via RuvC through an unchanged pathway bypassing the PLL domain that structurally connects WED and nuclease domains. Meanwhile, catalytic D9 communicated with H599 only through RuvC and HNH residues. Generally, dsDNA loading and AcrI6 binding both wired the communication pathway between the above components, mainly by enabling direct signalling from PIRs to RuvC without transmitting through PLL, and introducing L2 residues along the signalling path between D9 and H599 catalytic sites to achieve a more intense and efficient transduction (Fig. 7H, I and Supplementary Fig. 17). Specifically, PAM nucleotides were employed to enable efficient signal transmission between WED and RuvC domains in St1Cas9-sgRNA-dsDNA. However, stronger signalling was achieved by AcrI6 binding, as evidenced by the shorter path length and smaller path degeneracy, indicating that it is a more potent allosteric effector than the PAM duplex (Supplementary Fig. 17 and Supplementary Table 8).

Common signalling routes were adopted by inhibitory signals from AcrI6 when propagating within St1Cas9 RNP to the catalytic sites and PIRs; however, the presence of dsDNA hindered such efficient signalling (Fig. 7J, Supplementary Fig. 18, and Supplementary Table S8). In St1Cas9-sgRNA-dsDNA-AcrI6, inhibitory signals from AcrI6 were transmitted via an elongated path consisting of more residues before reaching the AcrI6-St1Cas9 RNP interface. Such compromised inhibitory signalling may underlie the preservation of St1Cas9-sgRNA-dsDNA catalytic activity when bound by AcrI6. In general, PI residues K1010 and F1019 served as direct recipients of the AcrI6-derived inhibitory signal, whereas L601 (HHN), N683 (L2), and S961 (RuvC) maintained the mediation of allosteric information transfer within St1Cas9 RNP, suggesting that mutagenesis of these residues is a potential modulatory strategy for fine-tuning the activity and selectivity of St1Cas9.

2.8. Bacteria Cas9 employ residue variations to protect against AcrI6 assault

The presence of St1Cas9 strains carrying point mutations (G993K/K1008M/K1010E) that’s capable of evading the phage invading system depicted the ongoing coevolution between species. Based on the previous findings [2], we further conducted multiple sequence alignment of PI domains of all 53 *S. thermophilus* genomes during the evolution of its Cas9 endonuclease to bypass AcrI6 inhibition. Using large-scale MD simulations, we further clarified the molecular basis underlying such triple mutational escape (G993K/K1008M/K1010E, referred to as mut St1Cas9). Based on the dynamics exhibited by mut St1Cas9-sgRNA throughout 15 μs sampling (Supplementary Figs. 3, 5, and 6) and its similarity with St1Cas9 RNP in terms of conformational distribution in the principal subspace (Fig. 2A and Supplementary Fig. 7), the triple mutations were unable to significantly disturb the overall conformation; this was predictable because residue variations are naturally present in *S. thermophilus* with normal endonuclease activity. In addition, the original approaching (along PC1) and open

(along PC2) essential dynamics were well preserved along each order (Supplementary Fig. 20), with relaxed PAM recognition grooves encompassed by WED and PI domains as well as HNH nucleosome domain docking near the cleavage site. Nonetheless, triple mutations weakened the innate correlation within mut St1Cas9 (Fig. 6E and Supplementary Fig. 16E, G) and altered its community network by enhancing direct informational flow from mutation-harbouring community 10 to community 16 (representing the RuvC nucleosome domain), ultimately resulting in weakened signalling between HNH and RuvC catalytic sites and regions relevant to PAM recognition in WED and PI. The original WED and PI architecture associated with PAM recognition was largely retained in the mutated systems, as evidenced by the similar ΔK867–K1029–N1027 perimeter in the major conformational clusters from mut St1Cas9 RNP under both AcrI6 unbound/bound states (Fig. 3E, F; C for mut St1Cas9-sgRNA, Cmut and Cmut for mut St1Cas9-sgRNA-AcrI6) as that in the unmutated systems (Fig. 3A, B and Supplementary Table 1). However, introduction of the triple residue substitutions reduced the AcrI6Δb (j2–j3) binding pocket volume by inducing greater counter-movement between WED and PI domains resembling the closure of a nutcracker, which resulted in disrupted mut St1Cas9 RNP shape complementarity as well as decreased interface area for AcrI6 recognition (Supplementary Fig. 21 and Supplementary Table 6). In contrast, the MM/PBSA binding free energy demonstrated that the affinity between St1Cas9 and AcrI6 was significantly compromised in the presence of triple mutations, further destabilising the AcrI6 dimer (Supplementary Table 5). Indeed, the original interaction network between St1Cas9 RNP and AcrI6 was markedly weakened upon triple residue substitutions, with complete loss of the strong polar interaction between K1010 and AcrI6Δb (Fig. 4E, F, Supplementary Fig. 12C, 13C, and Supplementary Tables 3, 4). K1010 was also directly implicated in AcrI6 allosteric signalling toward St1Cas9 as directly recipient of signals from AcrI6Δb A124 (Fig. 7 and Supplementary Fig. 18); its charge-reversal substitution from lysine to glutamic acid resulted in a switch to a more roundabout and less effective lane (through TOPO residue D987 or K984) for inhibitory signals to be transduced from AcrI6 to the mut St1Cas9.

2.9. Determination of key mutations in St1Cas9 that affect AcrI6 inhibition

To explore the molecular determinants of coevolution of phage-host interactions, we monitored the role of naturally occurring mutations of St1Cas9 in conferring AcrI6-mediated inhibition. For this, we generated three single mutants, G993K, K1008M, and K1010E, three double mutants, G993K/K1008M, G993K/K1010E, and K1008M/1010E, and one triple mutant, G993K/K1008M/K1010E, and subsequently stably expressed them in *E. coli* cells. To exclude the effect of mutations on the thermodynamic stability of St1Cas9, thermal shift assays were performed by the quantitative reverse transcription PCR (RT-qPCR). The results showed that the melting temperature (*T*~m~) of all variants was comparable to that of wild-type St1Cas9 (Fig. 8A and Supplementary Fig. 22), suggesting that the mutations had a minor effect on the thermodynamic stability of St1Cas9.

An *in vitro* DNA cleavage assay was further performed to assess the inhibitory effect of AcrI6 on enzymatic activity of wild-type St1Cas9 RNP and variants. We concomitantly incubated St1Cas9 RNP with 150 nM AcrI6 and 5 nM 1664-bp long dsDNA substrates, harboring target DNA and PAM sequences. The dsDNA cleavage products were resolved using electrophoresis at different time points (Fig. 8B). When we normalized the *in vitro* DNA cleavage activities of the St1Cas9 variants to that of wild-type enzyme, we found that all the variants decreased the cleavage of target DNA.
in the presence of AcrIIA6 with varying extents. As shown in Fig. 8C and Supplementary Table S9, a moderate (~15%) reduction in AcrIIA6-dependent inhibition of St1Cas9 RNP nuclease activity was observed in both G993K and K1008M single mutants, while the K1010E mutant exhibited an appreciable (~30%) reduction in its inhibition rate in the presence of AcrlA6. This result indicated that the K1010E mutation affects a key residue, consistent with the MD results that K1010E mutation resulted in loss of strong AcrlA6–St1Cas9 interaction and hindered effective allosteric signal transduction. The two double mutants (G993K/1010E and K1008M/1010E), to a lesser extent, the G993K/K1008M double mutant, showed a similar reduction of nuclease activities upon treatment with AcrlA6 to the K1010E single mutant. Strikingly, the triple mutant (G993K/K1008M/K1010E) exhibited a marked (~40%) reduction in its nuclease activity upon treatment with AcrlA6. This implied that the St1Cas9 triple mutant can become an efficient molecular evolutionary pattern of bacteria to bypass phages AcrlA6 inhibition.

3. Discussion

The exploitation of CRISPR-Cas systems in various molecular, biotechnological, and medical applications is growing; however, fine-tuning the efficacy and safety of this toolbox is of vital importance in addressing potential problems such as off-target effects [20,84–86]. Previous attempts to prevent such unwanted activity include mutating the Cas endonuclease [87,88], fusing with regulatory domains [89,90], or employing pre-formed RNP with controllable fidelity and nuclease activity [91,92]. However, such strategies have been faced with fatal flaws in delivery modality and ortholog universality [45]. The natural bona fide CRISPR-Cas antagonist Acrs—evolved during the evolutionary arms race between bacteria and bacteriophages—represent a genetically encodable, post-translational regulation of CRISPR-derived technologies to keep this powerful machinery in check [45]. Indeed, a number of studies integrating sequence analysis, machine learning, etc. have attempted to exploit this natural CRISPR-Cas regulator repertoire [18,56–58]; moreover, computational-guided engineering of existing Acr-based structural information has been recognised as a promising approach for expanding Acr inhibition [93]. Currently, the broad spectrum of Cas endonucleases regulated by Acrs compensates for the current scarcity of regulation strategies for Cas endonucleases with different sizes, fidelity, and PAM promiscuity from that of the established CRISPR-SpCas9-based toolkit [4,7,45,46,94], which would enable controllable and customised CRISPR-based techniques for different purposes (i.e. manipulation of differently sized genes).

We explored the detailed atomic-level AcrlA6 inhibitory mechanism through extensive MD simulations, which demonstrated the AcrlA6-induced conformational transition of St1Cas9 from the approaching conformation—in which the nuclease HNH domain is positioned near the cleavage site while the WED and PI domains
adopt a stretched posture ready for PAM interrogation—to a departing ensemble—in which outward movement of the catalytic HNH core was accompanied by shrinkage of the regions responsible for PAM recognition. Furthermore, through the intensive interaction network established based on shape complementarity between AcrIIA6 and St1Cas9 RNP, inhibitory signals were propagated to the St1Cas9 endonuclease while the correlated motion between functional domains mediated further transduction of inhibitory signals; these signals allosterically resulted in increased conformational plasticity of the catalytic HNH domain residing far from the direct AcrIIA6 recognition interface, allowing for its significant relocation in the presence of the inhibitor. Moreover, by computing the residue-level communication pathway coordinating the functional synchronicity between HNH, RuvC catalytic sites, and PAM interaction regions as well as how inhibitory signals are involved, we identified the key role of K1010 in the PI domain as a direct recipient of allosteric signals from AcrIIA6, further propagating them towards the St1Cas9 RNP. Variations in K1010, along with residue substitutions in K993 and K1008, have been shown to bypass the anti-CRISPR strategy by reshaping the intermolecular recognition interface between AcrIIA6 and mut St1Cas9, rewiring the original signalling pathway. The widespread distribution of residue polymorphisms of the implicated residues, especially K1010, suggests that such variations represent a strategy of AcrIIA6 inhibition escape, reflecting the coevolution of bacteria and phages.

Notably, we observed the inhibition failure of dsDNA-loaded St1Cas9 RNP by AcrIIA6, in which the inhibitor induced a spatial shift of the substrate DNA duplex, resulting in weakened PAM-St1Cas9 interactions but retaining the nuclease HNH configuration. This observation indicates that although AcrIIA6 inhibits St1Cas9 RNP functionality by interfering with both DNA binding and nuclease activation, its more primary mode of action may be modifying the conformational dynamics of WED and PI domains associated with PAM recognition; the altered dynamics of WED and PI further resulted in HNH outward relocation, which would otherwise be located closer to the presumed cleavage site. While successfully establishing the mechanistic framework underlying AcrIIA6 allosteric inhibition, we also observed the conformational transition tendency of St1Cas9 sgRNA-dsDNA towards a catalytically active state along the 3 μs simulation; however, due to the limited simulation time scale, the fully activated St1Cas9 sgRNA-dsDNA structure for catalysis was not completely captured, posing a challenge for future studies to obtain continuous large-scale MD simulations of large macromolecular systems, such as CRISPR-Cas9.

Taken together, our findings help elucidate the driving force behind the emergence of AcrIIA6 during the coevolution of bacteria and phages as well as the detailed mechanism of allosteric inhibition of St1Cas9 RNP by the anti-CRISPR protein AcrIIA6 [95–98]. The results provide a dynamic picture of this mechanism. The identified AcrIIA6-mediated allosteric inhibitory mode can provide further guidance for designing regulatory molecules/peptidomimetics to fine-tune the bioactivity of the CRISPR-Cas machinery when performing in vivo gene manipulation, improving the safety of gene therapy. Meanwhile, the distinct allosteric mechanism displayed by AcrIIA6 further emphasises the amazing diversity of inhibitors that phages have evolved to counteract the existing challenges posed by CRISPR-Cas systems. Moreover, we provided insights into anti-CRISPR strategies exemplified by Acrs to the Acr-driven mutational escape of prokaryotes, uncovering the basis of mutational escape from AcrIIA6 inhibition through a few residue variations. The robustness of different S. thermophilus CRISPR-Cas9 systems with residue variations will be further explored to identify economically suitable strains that can resist the selective pressure imposed by rapidly reproducing phages. By providing an atomic-level understanding of AcrIIA6 allosteric inhibition of St1Cas9 and St1Cas9 anti-anti-CRISPR rescue mechanism, we can offer a guidance for devising ‘off-switches’ for the CRISPR-Cas9 machinery with improved regulatability and efficacy, enabling temporal, spatial, and conditional genome manipulation for both basic and clinical applications.

4. Materials and methods

All simulation systems in our study were constructed based on the solved cryo-EM crystal structure of St1Cas9-sgRNA::dDNA20-AcrIIA6 (PDB ID: 6RJA), St1Cas9-sgRNA::dDNA59-ntPAM (PDB ID: 6RDJ), and St1Cas9sgRNA-AcrIIA6::dDNA59-ntPAM (PDB ID: 6RGJ) complexes at 3.0, 3.3, and 3.2 Å resolution [2]. Structural superposition of St1Cas9 from the three complexes was first performed to identify the missing residues in common. For the unsolved large segments around the HNH domain (MET510–TYR689) and RuvC-III domain (Leu751–Lys802), structural information was modelled based on sequence homology with Staphylococcus aureus Cas9 (SaCas9, PDB ID: 5AXX [99]) using SWISS-MODEL, and a structural model of reliable quality was obtained with QMEAN of −1.29 for HNH and −2.15 for RuvC domain, respectively. In addition, other sporadic missing residues in the original crystal profile were remodelled using Discovery Studio. After completion of the above-mentioned operations, the obtained systems were first subjected to 10000-step minimisation using the steepest descent algorithm typed with CHARMM forcefield.

The apo St1Cas9, St1Cas9-sgRNA, and St1Cas9-sgRNA-AcrIIA6 systems were built based on the remodelled St1Cas9-sgRNA::dDNA20-AcrIIA6 (PDB ID: 6RJA, without PAM sequence) by deleting constituent sgRNA::dDNA20-AcrIIA6, dDNA20-AcrIIA6, and dDNA20, respectively; meanwhile, the remodelled St1Cas9-sgRNA::dDNA59-ntPAM (PDB ID: 6RDJ, with PAM sequence) and St1Cas9-sgRNA-AcrIIA6::dDNA59-ntPAM (PDB ID: 6RGJ, with PAM sequence) well represented the conformation of dsDNA-loaded Cas9 RNP under AcrIIA6 unbound-(St1Cas9-sgRNA:dDNA) and bound- (St1Cas9-sgRNA:dDNA-AcrIIA6) states. Triple mutations along the AcrIIA6 binding interface (G993K, K1008M, K1010E) were introduced using Discovery Studio to construct mut St1Cas9-sgRNA and mut St1Cas9-sgRNA-AcrIIA6 systems. To further investigate the species-specific inhibition of St1Cas9 by AcrIIA6, an additional Streptococcus pyogenes Cas9 RNP model was constructed (PDB ID: 600Z [39]), whose missing residues were remodelled based on coordinate information of spCas9 under other activation states (PDB ID: 4O08); the obtained complex was also subjected to 10000-step steepest descent minimisation (Supplementary Fig. 2).

4.2. MD simulation details

The above-mentioned systems were prepared with the AMBER18 program using the ff14SB force field [100] to describe the ribonucleoprotein complex. All systems were first solvated in an orthorhombic transferable intermolecular potential three point (TIP3P) water box [101], followed by the addition of Na⁺ and Cl⁻ counterions to neutralise the system while mimicking the in vivo physiological cleavage condition (Supplementary Table S10). The generated topology and coordinate files served as inputs for subsequent MD simulations. Two rounds of energy minimisation were first carried out as previously reported [102–106], first with the whole protein scaffold fixed, followed by removing all constraints for 5000- and 10000-step maximum minimisation cycles, respectively. Subsequently, all systems were equilibrated in a canonical
ensemble (NVT) for 700 ps after heating from 0 K to 300 K within 300 ps. Finally, five independent 3-µs MD simulation replicas with random initial velocities were performed for all systems embedded in an isothermal and isobaric ensemble (NPT) with periodic boundaries, generating 40 independent trajectories and accumulating ~120-µs conformational sampling in total. Langevin dynamics using 1 ps⁻¹ collision frequency was used to control the temperature during the simulation. Long-range electrostatic interactions were analysed using the particle mesh Ewald (PME) method, and a 10 Å non-bonded cut-off was introduced for short-range electrostatics and van der Waals interactions. Covalent bond interactions involving hydrogen atoms were constrained by the SHAKE algorithm. Subsequent analysis of simulation trajectories was based on all five replicas (consisting of ~15 µs of sampling) for each simulation system to strengthen our statistical analysis. Based on the time evolution trend of the root mean square deviation of St1Cas9 (Supplementary Fig. S3), all productive runs reached relative equilibrium after the first 500-ns MD simulations.

4.3. Generalised correlation analysis

Assessment of the correlated motions within our model systems under investigation was achieved with the generalised correlation (GCij) analysis approach proposed by Oliver and Helmut [83], which enabled further insights into the allosteric signal transduction process. The GCij approach surpasses the established Pearson correlation analysis in that it enables the description of nonlinear correlations while being independent of atomic fluctuation orientations. Rest on the fundamental definition of independence of random variables, in GCij analysis the variables xi, xj are considered correlated when the product of their marginal distribution p(xi) · p(xj) is larger than their joint distribution p(xi, xj). To measure the degree of correlation between selected variables, mutual information (MI) between xi and xj was introduced and defined as:

\[ MI[xi, xj] = \int p(x_i) \ln \frac{p(x_i|x_j)}{p(x_i)} dx_i dx_j \]  

(1)

where the right side of equation (1) defines MI as closely related to the well-known Shannon entropy H(x), which is calculated as:

\[ H[x] = \int p(x) \ln p(x) dx \]  

(2)

thus, the correlation between pairs of atoms xi and xj is described by MI and calculated using the marginal Shannon entropy H(xi), H(xj), and the joint entropy term H(xi, xj) as follows:

\[ MI[xi, xj] = H[xi] + H[xj] - H[xi, xj] \]  

(3)

The g_correlation tool in Gromacs 3.3 [107] calculates the entropy terms H(xi), H(xj), and H(xi, xj) with the k-nearest neighbour distance algorithm using atomic fluctuation information. The MI[xi, xj] values were further normalised to obtain the normalised generalised correlation coefficients (GCij):

\[ GC_{ij} = \left\{ 1 - e^{-\frac{2MI[\xi, \chi_j]}{d^2}} \right\} ^{\frac{1}{2}} \]  

(4)

where d represents the dimensionality of xi and xj, which equals 3 in our analysis.

Inspired by Palermo et al. [27,76] to further represent the extent to which a generalised correlation is represented by each residue with the remaining part, we introduced the residue correlation score (CS) in our analysis. For residue i, its corresponding CSi is defined as:

\[ CS_i = \sum_{j \neq i}^N GC_{ij} \]  

(5)

where N represents the total number of residues within the protein (in our case N is 1121, total number of residues in St1Cas9). Depending on the domain to which residue j belongs, CSi,dom (residue i and j belong to the same domain) and CSi,inter (residue j belongs to domains excluding that of residue i) were calculated to describe intra- and inter-domain correlations mediated by each residue. Moreover, to gain insights into the more macroscopic inter-domain correlation, CSi,over over residues j from a specific domain were accumulated, and the obtained matrix was plotted to denote inter-domain correlations. Notably, all GCj analyses were based on the filtered GCj matrix with a minimum threshold value of 0.65.

4.4. Dynamic network analysis

To describe and investigate residue-residue interactions within biomolecular systems, we employed concepts from network theories [108–111]. The whole ribonucleotide protein complex was defined as a set of nodes, which were assigned to the Cz atom within each amino acid residue or P atom in the nucleotide backbone for each nucleotide. Edges were drawn between nodes that stayed within a cut-off distance of 4.5 Å for at least 75% of the conformational sampling consisting of all equilibrated simulations (the last 2.5 µs in 5 replicas, 12.5 µs in total) for each system. The edges between node i and node j were further weighed by the corresponding GCij coefficients as follows:

\[ d_{ij} = -\log GC_{ij} \]  

(6)

To indicate the relative distance between nodes i and j during information transfer in the connection network. The obtained weighed dynamical network was used as a basis for further ‘community’ [112] and ‘optimal path’ analysis [113–116]. To obtain the community substructure, the Girvan-Newman divisive algorithm was applied, in which ‘edge betweenness’ that defines the number of shortest pathways across a given edge serves as an important partition criterion. By iteratively removing edges with the highest betweenness from the network and recalculating the betweenness of remaining edges until each node represents an isolated community, the optimal substructure of the network—in which connections are only dense within a community but sparse in between—can be obtained. A ‘modularity’ parameter Q indicating the quality of a given community partition was defined as:

\[ Q = \sum_m e_{mm} - \frac{(\sum m e_{mm})^2}{\sum m^2 e_{mm}} \]  

(7)

where emm indicates the fraction of edges that connect nodes in community m to nodes in community n, while \( \sum m e_{mm} \) represents the fraction of edges that links to nodes in community n. Falling within the range of 0 to 1, a larger modularity value indicates a higher quality of the obtained community structure. The package gcommunities was employed to gain insights into the community structure of the St1Cas9-sgRNA (Supplementary Table 7–1), St1Cas9-sgRNA-dsDNA (Supplementary Table 7–2), St1Cas9-sgRNA-AcrIa6 (Supplementary Table 7–3), St1Cas9-sgRNA-dsDNA-AcrIa6 (Supplementary Table 7–4), muSt1Cas9-sgRNA (Supplementary Table 7–5), and muSt1Cas9-sgRNA-AcrIa6 (Supplementary Table 7–6) systems, with their corresponding modularity values reported.

To track the detailed information transfer pathway between pairs of residues (defined as source and sink), we used the Floyd-Warshall algorithm to identify the ‘shortest pathway’ by comparing the sum of dij (path length, PL) of all edges involving in the path. The source and sink were laid on catalytic D9 in RuvC, H599 in HNH, PAM-interacting residues (PIR, including WED: K867, S961, V962, S963; TOPO: K1025; CTD: N1026), and AcrIa6 residues (T118, A124, S147, V83) to disclose their residue-by-residue inter-signalling path. The number of suboptimal pathways...
with lengths that are not larger than the PL of the shortest pathway by 2% is defined as pathway degeneracy (PD) to indicate specificity.

4.5. Plasmids preparation for protein expression in E. coli

The full-length St1Cas9 gene from Streptococcus thermophilus (strain ATCC BAA-491/LMD-9) was synthesized by Tsingke Biotechnological Technology (Beijing, China) and cloned into pET28a vector using the NdeI and BamH I restriction sites adding an N-terminal 6x histidine tag to ease protein purification. The full-length AcrIIA6 of phage D1811 was also synthesized and cloned into pET28a vector with N-terminal 6x histidine.

4.6. Site-directed mutagenesis

Point mutations were introduced into the plasmid encoding the wild-type St1Cas9 by Mut Express II Fast Mutagenesis Kit V2 (Vazyme Biotech co.,ltd, Nanjing, Jiangsu, China). The successful introduction of the mutations was verified by DNA sequencing (Personalbio, Shanghai, China). The detailed nucleotide sequences for constructing all the macromolecule models (including both wildtype and mutated complexes) were supplemented in Supporting Information Table S11.

4.7. Protein expression and purification

Plasmid was transformed into E. coli BL21-Str(DE3) pLysS cells (Weidi, Shanghai, China) for protein purification. The cells were induced with 0.5 mM isopropyl β-D-1-thiogalactopyranoside (IPTG) at 18 °C. The St1Cas9 protein was firstly purified with nickel column (GE healthcare, Little Chalfont, Buckinghamshire, UK) in the lysis buffer (50 mM HEPES pH 7.4, 500 mM NaCl, and 10 mM imidazole), 0.1 mg/mL Dnase I, 0.1 mg/mL lysozyme and 1x protein inhibitor cocktail were added before French Press. Next, the St1Cas9-containing fractions were buffer exchanged in low salt buffer (20 mM HEPES pH 7, 100 mM NaCl) and applied onto a 5 mL HiTrap SP HP column (GE Healthcare, Little Chalfont, Buckinghamshire, UK) for cation exchange chromatography. St1Cas9 containing fractions were buffer exchanged into storage buffer (20 mM Hepes, pH 7.5, 150 mM NaCl), flash-frozen in liquid nitrogen, and stored at −80 °C. Purification of mutation forms of St1Cas9 protein was carried out analogously. For AcrIIA6 protein, gel filtration chromatography was used instead of cation exchange chromatography and stored in the same storage buffer.

4.8. Preparation of sgRNA and dsDNA

The transcription templates of sgRNAs were synthesized by GENEWIZ (Suzhou, Jiangsu, China) and amplified by a PCR. sgRNAs were prepared by in vitro transcription using the T7 High Yield RNA Transcription Kit (Vazyme Biotech co.,ltd, Nanjing, Jiangsu, China) and further purified using phenol/chloroform extraction followed by ethanol precipitation. dsDNA was used as cleavage substrate was synthesized by GENEWIZ (Suzhou, Jiangsu, China) and amplified by a PCR.

4.9. In vitro DNA cleavage assay

1664-bp long dsDNA substrates containing the target (underlined) and PAM (bold) sequences were generated by PCR (target and PAM sequences inserted into the pUC19 plasmid: TTTCTGCAA-TACTTTATACGCAAG) as previously described. In vitro cleavage reactions were performed in 20 μL reaction buffer (20 mM HEPES pH 7.5, 100 mM KCl, 1 mM DTT, 10 mM MgCl2) containing 5 nM linearized dsDNA substrates, 50 ng Loading control, 150 nM St1Cas9 and 200 nM sgRNA, with or without 150 nM AcrIIA6. The reaction was incubated in 37 °C for 15 min and was quenched by adding 50 mM EDTA, 20 μg Proteinase K and 1× Loading buffer for 30 min in room temperature. Products were analysed by electrophoresis in a 2% agarose, 45 red plus dye (Sangon, Shanghai, China). Gels were imaged using a Tanon-3500 gel imaging system (Tanon, Shanghai, China) and quantification were performed using Image J. loading control dsDNA wasn’t cleaved by St1Cas9, as a correction for quantification. The experiment was independently repeated four times. Data are presented as the mean ± s.d. of three independent experiments.

4.10. Protein thermal shift assays

A total of 2 μM St1Cas9 wild-type or mutant proteins was mixed with 5x SYPRO orange dye (Sigma-Aldrich, St. Louis, MO, USA) in 1x PBS. Samples were analysed in LightCycler 480 Real-time PCR instrument System II (Roche, Basel, Switzerland). The temperature was increased at a rate of 0.05 °C/s over a range of 25–95 °C, and the fluorescence was monitored with the SYPRO orange channel. Tm values were obtained from the minimum of the first derivative (−dT/dT) plots of the unfolding protein curves through analysis using LightCycler 480 software provided by the manufacture (Roche, Basel, Switzerland). GraphPad Prism Version 7.00 (GraphPad Software, La Jolla, CA, USA) was used to normalize the dT/dT curves. At least three trials were independently performed for each experiment.

5. Ethics approval and consent to participate

Not applicable.

6. Consent for publication

All the authors have approved and agreed to publish this manuscript.

7. Availability of data and material

All data generated and analysed to support the conclusion of this study is included in this published manuscript and its additional supporting files.

8. Authors’ contributions

S.L. conceived and supervised the project, designed the experiments, and revised the manuscript; X.L. and T.C. performed MD and project oversight. All authors discussed the results and reviewed the manuscript.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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