Introduction of CRISPR/Cas system, as a genetic interference pathway and essential factor in genome evolution

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ABSTRACT

The clustered regularly interspaced short palindromic repeats / (CRISPR) associated protein system is an adaptive immune system in bacteria and archaea. This system mimics eukaryotic RNA interference (RNAi) system and can play an important role in gene regulation and immune system. CRISPR interference (CRISPRi) defends the bacteria against invader agents, such as phages and plasmids. As a result, it can provide a genetic interference pathway depending on the integration of a small stretch of invader DNA of the bacterial or archaeal genome, which acts as a heritable genetic memory to defense against viruses or phages.

Keywords: CRISPR/Cas system, adaptive and heritable immunity, genome editing, CRISPRi

INTRODUCTION

Acquisition of a new gene, which confers a selective advantage, is an essential factor in genome evolution. A great proportion of bacterial and archaeal genomes contain genes derived from exchanging the genetic material among related or unrelated species [1]. This phenomenon is known as Horizontal Gene Transfer (HGT). HGT occurs with transformation, conjugation or transduction [2]. A small fraction of acquired genes confers selective advantage, thus microorganisms evolve many genetic mechanisms to prevent HGT, such as DNA restriction or surface exclusion [2].
During the past two decades, clustered regularly interspaced short palindromic repeats (CRISPR) systems have been identified as novel genetic interference pathways limiting at least two major mechanisms of HGT consist of conjugation and transduction. Mechanisms such as eukaryotic RNAi and related pathways are analogous to CRISPRi [3, 4]. Unlike other defense mechanisms, CRISPRi is an adaptive and heritable immune system and enables bacteria to control the sequence length, and to be reprogrammed to reject the invading DNA molecule, which has not been previously encountered [3]. In this review, we discuss about the CRISPR structure and functions, and mention various applications, which have advantages in basic and medical genetics research.

**Discovery of the CRISPR system**

In 1987, Ishino and his colleagues cloned and sequenced *iap* gene, which is responsible for alkaline phosphatase isozyme conversion in *E. coli* [5]. He found that there was a set of 29 nucleotide (nt) repeat downstream of *iap* gene. These repeats were separated by non-repetitive, and similar short sequences known as spacers. Similar repeats were detected when the whole genome sequencing projects were carried out in bacteria and archaea [6, 11]. Now a days, CRISPR is introduced as a family of repeats present in many bacterial, and archaeal species. This sequence was nominated as CRISPR in 2002 [38]. In fact, this term can reflect the loose structure (Fig 1). A ‘Leader’ sequence precedes a cluster of repeats which is an AT rich region with several hundreds base pairs length which has intraspecies conservation [38].

**Leader sequence:**
- AT-rich
- several hundreds nucleotides length
- conserved intraspecies

**Spacer/ Repeat sequence:**
- Lengths of Repeat sequence (2 to several hundreds with 23- 50 nt)
- Lengths of Spacer sequence (17- 84 nt)

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*Figure 1.* CRISPR array structure. CRISPR array involves an AT-rich leader sequence with several hundreds nucleotides length, which is conserved intraspecies. Second CRISPR array segment is Repeat/spacer sequence. The number of repeat sequences is from two to a few hundreds with 23- 50nt length. Spacer sequences are 17-84nt long matched with sequences from mobile genetics elements, such as bacteriophages and plasmids.
As mentioned, CRISPR/Cas system provides adaptive immune system against invading agents [3, 4]. This immunity is based on the integration of the invader DNA stretch into the CRISPR loci as proto-spacer which acts as genetic memory for protection against viruses or phages [3, 4]. The space sequence is matched with the foreign sequences, mobile genetic elements [13, 15]. This provides the possibility of a potential base pairing that can enable a sequence-based interference with phage infection, gene expression or both. These suggest that the function of CRISPR system is analogous to eukaryotic RNAi [17, 18].

Analysis of the CRISPR loci in Yersinia pestis strains demonstrated that the form of new spacer acquisition is polarized [15], meaning that new units have been added at one end of the cluster close to the next end [15, 16]. Note that Cas genes are essential for CRISPR function. In 2008, Marraffini et.al showed that limitation of Plasmid conjugation can be induced using CRISPRi in Staphylococcus epidermidis [19]. This research demonstrated an association between CRISPR/Cas gene loci and prevention of HGT in bacteria; Thus, CRISPR/Cas system is known as the immunity system.

Classification and Structure of the CRISPR loci

CRISPR sequences are presented in approximately 40% of Bacteria and 90% of Archaea, which have at least one CRISPR locus [20]. This marked difference may be caused by a bias in sequences of bacteria. This bias is presented in long established laboratory strains that have had no bacteriophage exposure for many generations, so their genome lacks the CRISPR loci.

CRISPR sequence consists of the repeat-spacer units ranging from just a few to several hundred numbers, the average is 66. The highest number is observed in the genome of the thermophilic bacterium, Chloroflexus sp. Y400fl with 374 units in one of its four CRISPR loci. The Repeat units have various sequences, even among CRISPR loci in the same genome. They exhibit limited dyad symmetry. There are 12 categories of CRISPR loci based on similarity of the sequences and their potential to form stem-loop structure [20]. A 23-50 nt range is reported as repeat length with an average of 31 nt, and 17-84 nt range is reported as spacer sequence length with an average length of 36 nt. The presence of various sets of Cas genes increases the complexity of CRISPR loci. More than 40 Cas gene families have been identified [17, 24]. There are eight CRISPR subtypes based on the presence of 45 gene families associated with CRISPR loci (established by Haft et.al.) [24] (Table 1). Cas1-cas6 was identified as gene families, which are considered ‘core’ Cas genes. Cas1 and Cas2 are introduced in all CRISPR loci (in a few cases
Cas2 is encoded as a combined domain of Cas3 protein) [24, 25].

**Acquisition of new spacer sequences**

At the molecular level, CRISPR function can be divided into three phases during which the new spacers are incorporated in the CRISPR array: Acquisition, Expression and Interference (Fig 2a), the CRISPR RNAs (crRNAs) are expressed, processed (Fig 2b) and CRISPRi occurs (Fig 2c) [26]. During the first step, a new spacer, who is known as proto-spacer, integrates in the invader genome. This procedure allows the microorganism to increase the potential of rapid adaptation with the environmental invaders. Thus, this step can be referred to as ‘adaptation’ phase [3, 26]. The spacer incorporation has an unknown molecular mechanism in detail. In *Ecoli*, Cas1 and Cas2 are necessary for the pre-existing function of spacers [27]. Because of the widespread presence of these proteins in CRISPR/Cas systems, it is thought that they participate in adaptation. The short conserved flanking sequences of proto-spacer (2-5bp) in invader genome are named ‘CRISPR motifs’ or ‘proto-spacer adjacent motif’ (PAM), which are classified based on their CRISPR sequences [20, 25]. The motif presence indicates that selection of proto-spacer is not random and a recognition signal is needed for the selected target sequence that is considered as a new spacer [25]. Mutation in CRISPR motifs or PAM sequences can cause CRISPR immunity evading which indicates the flanking sequences role during the defense (or interference) phase [21, 25].

The integration stores the genetic information to use it as inhibitor of the invader agents, specifically during ‘defense’ phase of CRISPRi [4]. CrRNA transcription correlates the adaptation phase with the defense phase [28, 29]. Transcription of CRISPR initiates from the leader sequence. Studies on thermophiles revealed that the camp regulator protein can upregulate *Cas* genes and camp signal transduction can guide CRISPR response in this bacterium [28-30].

**CRISPR transcription and processing**

CRISPR transcript (CRISPR precursor RNA) is cleaved and processed into a small crRNA by Cas proteins. Cas proteins are multi-complex proteins known as CRISPR associated complexes for antiviral defense (cascade) [31]. The components of cascade may be different in each subtype. Experimental data revealed that pre-crRNA processing occurs endonucleolytically at a specific site of each repeat
Figure 2. Three phases of CRISPR/Cas function. (a) Acquisition, new spacers are incorporated in CRISPR array, (b) Expression, crRNAs are expressed and processed, (c) Interference. The expressed and processed RNA matches with the invader genome, Cas proteins can recognize this structure. After that, invader genome will be cleaved with Cas helicase and nuclease activity.

(8 nt upstream of the spacer). Mature crRNA is associated with the cascade. CrRNAs form a stem-loop structure because they have palindromic repeats. CrRNA involves a spacer-flanking unit with repeat sequences (Fig 2b). CRISPRs are separated by short spacer sequences matching the DNA of bacteriophage or plasmid. Therefore, crRNAs consist of a spacer sequence can be matched with invader genome and formed base pairs. Finally, specific Cas proteins identify stem-loop structure and cut the invader genome and inactive it.

Recognition of the invading sequences

The base pairing potential between crRNA and the invader genome is mediated by the recognition of crRNA-Cas ribonucleoprotein (crRNP) complex. Base pairing is initiated from a seed sequence, after that pairing is promoted and finally DNAse activity is initiated which causes the cleavage of the invader genome [32-34]. It is noted that the CRISPR machinery recognizes DNA targets rather than RNA ones in numerous species; an exception is type III-B of
CRISPR/Cas system, which recognizes RNA targets [50]. In vitro analysis suggests that during RNA cleavage, 3’phosphate and 5’hydroxyl ends are generated at a 14 nt distance from the target sequence which is paired to the 3’terminal of the crRNA [3].

**CRISPR-Cas diversity**

The dynamic and selective pressures cause rapid evolution of the immune system because of the high potential of the CRISPR/Cas system to adapt with the new environmental agents rapidly. Therefore, it is acceptable that CRISPR/Cas systems are highly diverse. The comparative studies of CRISPR loci demonstrated that the sequences of the CRISPR repeats, the Cas genes and Cas operon architecture are major differences between various systems. Based on these differences, three main types of CRISPR were determined as type I, type II and type III; these three types are divided into several subtypes themselves [41]. Each type has a specific signature Cas-proteins: Cas3-helicase, Cas9-nuclease, and Cas10 (a large protein with unknown function) are defined for type I, II and III systems, respectively. Type II system is distinguished from two other types phylogenetically and structurally. Nomenclature is based on the target, cleaving the invading agents, crRNA and Cas-protein composition of crRNP complexes or cascade [41]. Cas protein composition of type I-

A to type I-F crRNP complexes are determined as cascade, but all type II crRNPs, which are divided to type II-A, II-B and II-C, are known as Cas 9 complexes. The Cas proteins involved in Type III-A and Type III-B crRNPs systems are known as Csm and Cmr complexes.

Type II system is only found in bacteria in composition with other CRISPR-Cas types, whereas Types I and III are found in different combinations among bacteria and archaea which are phylogenetically diverse [39, 42]. It is interesting to note that viral genomes and plasmids have CRISPR-Cas systems, which suggests that HGT provides the possibility of exchanging CRISPR systems [43-47]. Despite the diversity, there are four functional categories for all Cas proteins: Nucleases or recombinase, involved in acquisition of the spacer; ribonucleases, catalyzes the crRNA guides processing; proteins present in crRNA complexes and finally the proteins which have nuclease activity to degrade DNA and RNA targets.

**Type I and Type III systems**

In these two types, Cas6-like nuclease plays a role in pre-crRNA primary processing. Cas6 proteins have structural variations. Their enzymatic activity for cleaving pre-crRNA is involved in hydrolyzing a single phosphodiester bond in repeated sequences of the transcript. The
crRNA arrangement is consisted of a 5’ unusual seahorse shape (binding to DNA changes its conformation) of 8 repeated nucleotides (complete spacer sequence) derived from 3’seahorse shape with a variable size which forms a hairpin structure in some systems. The Cas6 protein delivers primary crRNAs to the respective crRNP complexes [49, 50]. In some subtypes, after transferring the crRNAs to the protein complexes, additional processing occurs to produce different mature crRNAs and the resulted crRNAs can have different nucleotide numbers [51, 52]. Cas5 protein is substituted with Cas6 in type I-C system. It shows the variation in applied proteins as Cas proteins in different types and subtypes of CRISPR systems. In type II systems, pre-cRNA processing occurs completely different; CRISPR locus includes a Cas operon, a CRISPR array and a gene encoding a trans-activating crRNA (tracrRNA) [53]. The tracrRNA is complementary to the repeat region of the pre-cRNA transcript, which is consisted of 25 nucleotides. The housekeeping ribonuclease RNase III recognizes and cleaves a double stranded region resulted from base pairing between these two RNAs. Primary crRNA processing requires Cas9 protein [53, 54]. This protein probably plays an important role in binding to the RNA molecule and positioning it for being cleaved by RNase III. The association of Cas9 with crRNA-tracrRNA hybrid follows initiation of the processing by RNase III. The 5’end of the crRNA spacer is 24-27 nt trimmed in length [53]. The mature crRNA-tracrRNA hybrid requires interfering the target. This structure is probably necessary for proper anchoring and positioning of the crRNA in Cas9. Following binding to the target nucleic acid, major rearrangements occur in Cas9 domains. These rearrangements are revealed using crystallography studies.

**Assembly of crRNP complexes**

Stable crRNP complexes are consisted of mature crRNAs and Cas proteins. The multi-subunit surveillance complexes of type I systems are called Cascade, which are the first identified crRNP complexes in *E. Coli* type I-E system [34, 84]. The structural studies have shown that there are architectural similarities between type I and type III crRNPs [54-57], but Cas9 is the only protein component that presents in type II crRNP complexes [54-57].

**Type I crRNP complexes**

Cas5, Cas6, Cas7 and a single 61-nucleotide crRNA are known as the core complex of type I-E crRNP of *E. Coli*. These complex proteins are less tightly associated with the Cse1 and Cse2 subunits. One Cse1, two Cse2s, one Cas5, six Cas7s and Cas6 (in I-E and I-F subtypes) are the complete complexes that have uneven subunit
stoichiometries. Features of type I and III crRNP complexes are provided at the time of alternative translation of encoding polycistronic mRNAs [58]. The helical backbone is composed of Cas7 (Csy3 in type I-F system), Cas8 (known as Csy1 in type I-F system), Cas5 (known as Csy2 in type I-F system) and the crRNA is consisted of Cascade complexes of type I-C [59] and type I-F systems [60]. Cas3’ and Cas3’’ as truncated domains of Cas3, act as a helix, and a nuclear domain respectively, and form different parts of the complex.

**Type II crRNP complexes**

Analysis of Cas9 proteins from the type II-A system of *Streptococcus pyogenes* shows that Cas9 has a conserved architecture consisting of two distinct lobes, the recognition lobe (the α-helix structure) and the nuclease lobe. The first lobe is involved in coordinating with the guild RNA (gRNA), and the second lobe is responsible for PAM recognition and the target DNA cleavage. Reorientation of two structural lobes following binding to the single guide RNA (sgRNA) causes a central channel formation and then DNA substrates will become able to attach [56]. The groove located at the interference surface of two lobes has a positive charge and filled with an RNA-DNA hetero-duplex. As a result, Cas9-sgRNA-ssDNA target structure is performed. HNH and Ruv C, as two nuclear domains, are not sufficient and cleavage does not happen. R-Loop configuration is required to reach the ultimate cleavage-competent state of Cas9. Because of CRISPR/Cas9 importance in genetic regulating, editing and engineering, performance of type II CRISPR/Cas9 is illustrated as a schematic figure (Fig 3).

**Type III crRNP complexes and similarities with type I systems**

Csm and Cmr complexes were determined from type III-A crRNA of *S. solfataricus* and two type III-B crRNP complexes from *Pyrococcus furiosus* and *Thermus thermophilus*, respectively (61). Csm3 in type IIIA and Cmr4 in type III-B systems are consisted of a multi-copy backbone that belongs to Type III complexes. This complex is a structural homolog of Cas7. The Cmr complexes from *P. furiosus* are the same as crRNA in cascade because of their positioning along the backbone. The Subunits of Type I and III systems can be introduced as small and large subunits.
Figure 3. Three phases of Type II CRISPR/Cas9 function. (A) Invader DNA is recognized by Cas1 and Cas2, and processed (red) and integrated at the leader end of the CRISPR array (red) (adaptation or acquisition phase, blue). (B) For protection against invader DNA in the next attack, CRISPR is transcribed as a single transcript (pre-crRNA), then it is matured into a small targeting crRNA using RNase III and tracrRNA. CrRNA and tracrRNA complex associate with Cas9, and then hybridization of the complementary DNA sequence is mediated by spacer sequence. Cas9 cleaves the targeted DNA at downstream of PAM (red circle) (Effector or expression and interference phases, pink).

The small subunit of Type III-A complexes consists of three copies of Csm2 and Cmr5 in Type III-A and Type III-B complexes, respectively [61, 62]. Type I and Type III crRNA complexes have conserved position of the large substitution, which reveals that their roles are analogue. The small subunits of type I and Type III systems are conserved in some parts of the structure [61-63].

**Target surveillance and interference**

The invader genome targeting proceeds in a stepwise manner [26]. This process contains several steps: (1) Scanning the invader genome for Proto-spacer sequence to find out which one complements crRNA. (2) Discriminating itself from outsider. (3) Base pairing between the seed region and the complementary proto-spacer (7-8nt length) following extension of pairing causes strand displacement [64-66]. Hybridization of crRNA, and target strand forms R-Loop structure which cause changes in crRNP complex
conformation [61, 67-69]. This transition in structure provides a signal to recruit a trans-acting nuclease (for type I and type III-A systems) or leads to activate of intrinsic nuclease activity (for type II and type III-B systems). Nonspecific interaction between crRNP complex and the invader genome results in scanning the invader genome. PAM is recognized by one of the subunits of cascade, this association between crRNP, and PAM motifs may destabilize the duplex of DNA. Thus, crRNA is enabled to hybridize with target DNA. Pairing occurs between the seed region of crRNA spacer (close to the PAM motif), and the proto-spacer complementary to the target. Hybridization between crRNA, and target DNA generates R-Loop, which causes a major change in the conformation of cascade and local bending of target DNA [70, 71]. Recruitment of Cas3 nuclease-helicase is caused by these structural changes. Helicase activity of Cas3, which depends on ATP which is combined with metal-dependent nuclease activity, causes completely degradation of target DNA [71, 72]. The displaced strand of R-Loop is cleaved by endonucleotide cleavage followed by a 3’ to 5’ exonucleotide degradation [71-73], the other DNA strands undergo endonucleolytic, and exonucleolytic degradations [71, 72-75].

Type II systems
Type II system interference is completely different from type I and type III systems. Cas9-RNP complex is consisted of Cas9, crRNA and tracrRNA which perform the interference in this type [53-55, 76]. Cas9 targets a DNA fragment in a step-wise manner [77]. SgRNA works as a leader and causes structural rearrangements that lead to a central channel formation binding to the target DNA [56]. The DNA is scanned for a PAM motif by Cas9-RNA complex. Type I, type II systems scan, and find DNA using similar pathways. PAM motif is located on displaced strand of target DNA and positioned to the 3’ seahorse shape of crRNA guide. Cas9 nuclease lobe determines PAM recognition using the flexible loops contained two tryptophans in its C-terminal domain [77]. DNA binding, DNA strands displacement and initiating of R-Loop formation in PAM require interaction with the PAM motif. Base pairing initiates in seed sequence for over 12 nt and proceeds to the distal end of the target sequence [77]. At the final stage, Cas9 cleaves the DNA using the nucleolytic activity of two active sites in nuclease lobe. The DNA strand paired with the crRNA is cleaved by a HNH-like nuclease domain and also the displaced DNA strand is cleaved by a Ruv-like nuclease domain [54, 56]. So, a blunt double-stranded end at a specific site with 3 nt distance
of 3’end of the proto-spacer is resulted [76]. Cas9 is known as a single-turnover enzyme [77].

**Type III systems**

Type III-A system complex is named Csm and contains Csm1-Csm5 and crRNA [78, 79]. Details of how the type III crRNP complexes target the proto-spacer is unknown. Type III-A systems discriminate self-DNA from non-self DNA in a PAM-independent manner [80]. This process is performed with Cas10-like protein Csm1 (large subunit) mediation. It is proposed that crRNA contains a seed sequence at the 5’end of the spacer region [38]. The Csm6 (a helicase and nuclease associated with Csm complex) is involved in DNA interference. Csm6 is known as Csx1, and its analogue with Cas3 in type I systems.

The complex of type III-B systems is Cmr which contains Cmr1-Cmr6 [78, 79]. The special feature of these complexes is that they target RNA rather than DNA [81]. RNA cleavage proceeds from 3’end to 5’end [62]. The spacer contains a seed-like sequence located at the 5’end. Results have shown that these complexes have multiple catalytic sites. Cmr-α, and Cmr-β are two Cmr complexes in *Sulfolobus islandicus*. Cmr-α is composed of Cmr1-Cmr6, that target plasmid DNA using a Csx1-dependent mechanism that requires targets transcription [82]. This seems to be a functional analogue of the aforementioned Csm complex of III-A systems. But Cmr-β complex is composed of Cmr1-Cmr7 in *Sulfolobus islandicus* and targets RNA in vitro (51), but its catalytic mechanism is different from the mechanism of III-B system in *P. furiosus* and *T. Thermophiles* [61, 62]. These results reveal that the mechanisms and functions of heteroduplex formation in type III systems are diverse [82]. Molecular details of crRNA binding by crRNP complexes are diverse between different CRISPR/Cas systems. Interestingly, rice protein mediates base-pairing pattern. Strings of RecA molecules, in complex with an ssDNA template, invade a dsDNA helix. After the displacement of the non-complementary strand, the new hybrid formed of complementary DNA strands provide a space that is locally allowed to adopt a class B conformation. This process resembles the crRNA–target hybrid configuration of the Cascade R loop.

The different structure of crRNA-target heteroduplex is provided by a single stretch resulted from crRNAs binding to the targets and increases the flexibility of crRNAs. In type II systems, the sgRNA base pairing with the 20 nt of target DNA is mediated by Cas9 protein [57]. The RNA targets in type III-B systems are similar to eukaryotic RNAi because of using a 21 nt guide RNA [83].
CRISPR system as a genetic editing and genetic engineering tool

In 2012, for the first time, Jennifer Doudna and her colleagues published their findings about genome editing capabilities of bacterial immunity system introduced by Danisco group [54, 76]. They showed that Cas9 protein involved in type II system can cleave the target dsDNA via detection of a base-paired structure formed between the activating tracrRNA, and the targeting crRNA. In addition, they demonstrated that Cas9 endonuclease family can be programmed by a single RNA molecule to cleave specific DNA sites; these observations raised the exciting possibility of developing a simple and versatile RNA-directed targeting, and editing. Cas9, the hallmark of type II systems, has been hypothesized to be involved in both crRNA maturation and crRNA-guided DNA interference [84]. Type II of CRISPR/Cas system has a powerful potential to be applied in genetic editing, and engineering (Fig. 4) [84]. This system has a good function in basic and applied molecular and medical genetic parallel to TALEN and ZFN (Zinc-Finger Nuclease) techniques [87-89]. Although, working with CRISPR is simple and more sufficient than TALEN and ZFN [89]. Type II of CRISPR/Cas system has been applied to discover a drug for muscle disease in zebrafish models [90].

Emergency of new era for genome sequencing reopens annotation of new genomic variations which are valuable to investigate their effects on phenotypes [85]. CsRISPR/Cas system can be applied to induce Indel variations in desired sequences using its matching and nuclease ability (Fig. 4a). In addition, CRISPR/Cas9 can be attached to activator or silencer motifs and be used as a tool to up-regulate or down-regulate the expression of interested gene (Fig. 4b). GFP-CRISPR/Cas construct is used for imaging studies to locate a genomic locus (Fig. 4c). These applications are used to study the mechanism of upstream or downstream sequences to control gene expression [85, 86].

CONCLUSION
During the past two decades, impressive progress has been made in understanding many aspects of the molecular features of CRISPR/Cas systems. CRISPR/Cas system is defined as a heritable adaptive immunity system in bacteria, and archaea. The discovery of this system reveals the genome evolution mediated by horizontal gene transfer. The molecular mechanism of this system has shown the coincide regulation to provide defense against environmental agents. This protection is prepared by a varied set of proteins [71-75]. The variations in environmental agents are driving forces of genome evolution. Understanding the details of the molecular
mechanism of CRISPR/Cas systems proceeds to point that the CRISPR/Cas applications are emerged and applied in genome editing, and engineering [85]. The CRISPR/Cas system has significant applications in basic researches, which is a good advantage for studying the molecular mechanisms. The emergence of the whole genome sequencing causes detection of the novel variations in the genome. The ability of CRISPR/Cas system in Indel variation introduction has made it as a powerful tool to perform a functional assay for new annotated variations.

Figure 4. CRISPR/Cas application in (a) Genome Editing, (b) genetic/Epigenetic Modulation and (c) Dynamic imaging of Genomic loci. Cas9 (dCas9) is a form of Cas9 mutated in nuclease domain. dCas9 is attached to the VP64 and P65 subunits of NF-κB, and KRAB domain as activator and suppressor proteins, respectively. It can be used to control the target gene expression. (C) dCas9 attached to EGFP, useful for molecular genetic studies like telomere movements studies.

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CRISPR/Cas in Genomic Evolution


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