CRISPR-Cas systems: role in cellular processes beyond adaptive immunity

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Abstract
Clustered regularly interspaced short palindromic repeats and associated Cas proteins (CRISPR-Cas) are the only known adaptive immune system in prokaryotes. CRISPR-Cas system provides sequence-specific immunity against invasion by foreign genetic elements. It carries out its functions by incorporating a small part of the invading DNA sequence, termed as spacer into the CRISPR array. Although the CRISPR-Cas systems are mainly responsible for adaptive immune functions, their alternative role in the gene regulation, bacterial pathophysiology, virulence, and evolution has started to unravel. In several species, these systems are revealed to regulate the processes beyond adaptive immunity by employing various components of CRISPR-Cas machinery, independently or in combination. The molecular mechanisms entailing the regulatory processes are not clear in most of the instances. In this review, we have discussed some well-known and some recently established noncanonical functions of CRISPR-Cas system and its fast-extending applications in other biological processes.

Keywords
CRISPR-Cas · CRISPR-Cas application · CRISPR-Cas in gene regulation · Genome remodeling · CRISPR-Cas alternative roles

Introduction
CRISPR-Cas system defends the prokaryotes from invasion by mobile genetic elements (MGEs) including phages, plasmids, and transposons. This system was first observed in the various archaeal genomes in the late 1980s. Its apparent prevalence in the broad range of bacterial and archaeal lineages suggested the common function of CRISPR-Cas systems in prokaryotes (Jansen et al. 2002; Mojica et al. 2000). After initial discovery, owing to the unique sequence and structure of a repeat-spacer array, inference of CRISPR-Cas function in prokaryotes took almost two decades. Detailed sequence analysis and bioinformatics studies revealed that the spacers in the CRISPR array target the phage and plasmid sequences (Bolotin et al. 2005; Pourcel et al. 2000). After initial discovery, owing to the unique sequence and structure of a repeat-spacer array, inference of CRISPR-Cas function in prokaryotes took almost two decades. Detailed sequence analysis and bioinformatics studies revealed that the spacers in the CRISPR array target the phage and plasmid sequences (Bolotin et al. 2005; Pourcel et al. 2005). Thereafter, experimental evidence showed that CRISPR-Cas prevented phage infection in Streptococcus thermophilus as well as averted the invasion by plasmid in Staphylococcus epidermidis, in a sequence-specific manner (Barrangou et al. 2007; Marraffini and Sontheimer 2010).

The brief outline of the CRISPR-Cas mechanism of action is shown in the following Fig. 1. The defining feature of the CRISPR-Cas system is the presence of a repeat-spacer array preceded by an AT-rich promoter sequence and adjoining Cas operon lying nearby in the genome (Charpentier et al. 2015). The CRISPR-Cas mediates its function by integrating a small nucleic acid sequence known as a spacer, acquired from invading foreign genetic element into the CRISPR array and thus confers the cell a unique and heritable form of adaptive immunity. This embodies the first adaptation step of the CRISPR-Cas mechanism. This is followed by the expression step, whereby transcription of CRISPR loci takes place which generates pre-crRNA (pre-CRISPR RNA). Pre-crRNA is recognized by ribonuclease and cleaved into smaller repeat-spacer units to generate crRNAs. This crRNA in combination with Cas proteins forms monomeric/multimeric complexes, which are part of the interference complex. In the last interference step, crRNA hybridizes to the complementary target sequence from the invading genetic element and thus recruits the interference complex onto the target sequence leading to its degradation (Charpentier et al. 2015).

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Over the past 15 years, impeccable progress has been achieved in understanding the molecular function of the CRISPR-Cas system. Such developments helped in unraveling the alternate functions of CRISPR-Cas systems. The alternate functions of CRISPR-Cas systems are described in detail in various published articles (Newsome et al. 2021; Ratner et al. 2015; Sampson and Weiss 2013; Sampson and Weiss 2014; Westra et al. 2014). CRISPR-Cas systems are incredibly versatile owing to the unique spacers of diverse origin, diversity in adaptation and effector complex, and a large repertoire of cas family genes (Makarova et al. 2002). The diversity in CRISPR-Cas function is mainly attributed to the versatility found in cas gene encoded diverse proteins with different enzymatic domains (Makarova et al. 2011). CRISPR-Cas is a multifaceted system and targets not only dsDNA (double-stranded DNA) but ssDNA (single-stranded DNA) and ssRNA (single-stranded RNA) as well. CRISPR type with RNA-targeting potential has been instrumental in the realization that the CRISPR-Cas system can play
a role in endogenous gene regulation, likewise the RNAi system in eukaryotes (Westra et al. 2014). In addition, the observation that CRISPR-Cas can occasionally incorporate spacer from the chromosomal region has also given rise to the perspective that CRISPR-Cas systems might have gene regulatory function (Stem et al. 2010).

Pathogenicity and virulence

Almost three years later after the initial discovery of CRISPR-Cas adaptive immune function, in Pseudomonas aeruginosa, it was observed that type I-F CRISPR system was involved in the regulation of biofilm-forming capacity and swarming motility of this pathogen (Palmer and Whiteley 2011; Zegans et al. 2009). The first well-documented, direct evidence of regulation of pathogenicity and virulence was discovered in Francisella novicida. It has developed a mechanism to subvert the host immune response by modulating the gene expression with the help of CRISPR-Cas system. Once inside the macrophage, Francisella novicida enters the phagosomes containing abundant receptors for the recognition of pathogen (Jones et al. 2012). Toll-like receptor 2 (TLR2) is one such receptor that recognizes bacterial lipoprotein (BLP) and further elicits the immune response of the host (Aliprantis et al. 1999; Brightbill et al. 1999). Francisella novicida, instead, employs Cas9, tracer RNA (trans-activating), and scaRNA (small CRISPR-associated RNA) to lower the expression levels of BLP protein on the outer envelope. This results in the reduced TLR2 activation and milder immune response, thus allowing the organism to thrive in the host (Jones et al. 2012). Neisseria meningitidis in the human lung epithelial model was also shown to utilize Cas9 for the infection of the host. Cas9 was shown to be essential for attachment to the cell wall, invasion, and replication inside the host (Heidrich et al. 2019; Sampson et al. 2013). Likewise in Streptococcus pyogenes, Cas9 was observed to impact the epithelial cell binding and growth in human blood cells. In addition, it was observed that it attenuated the virulence of the pathogen in murine necrotizing skin infection model. In addition, Cas9 was also shown to help Campylobacter jejuni to attach and invade the host in the colorectal epithelial cell model (Louwen et al. 2013). Transcriptome analysis of the wild-type and cas9 mutant strains of Campylobacter jejuni further revealed that Cas9 was involved in the upregulation of virulence factors. It was shown experimentally that the cas9 mutant strain was unable to form biofilm. Further, it lacked the ability of intracellular invasion and adhesion, survival inside the host and motility, suggesting the wide-ranging impact of cas9 on the survival and pathogenicity of the organism (Shabbir et al. 2018). The cas9 gene was found to affect the expression level of virulence determinants in several other microorganisms as well, suggesting its involvement in alternate functions other than the adaptive immunity (Gao et al. 2019).

In Legionella pneumophila, type II-B CRISPR-Cas expression was observed to be induced during intracellular growth in macrophages of amoebas, including Acanthamoeba castellanii and Hartmannella vermiformis. The cas2 gene was found to be important for the growth of organism inside the macrophages. It was reported that the Cas2 protein from this organism comprised of DNase as well as RNase activity. Further, Gunderson and colleagues argued that, as of now, it is not clear that whether the RNase or DNase or both catalytic activities are required for infection; however, the RNase activity appeared to be the dominant in their study. It was established that the nuclease activity was essential for the initiation of infection in the host. It was also established experimentally that the introduction of cas2 in Legionella pneumophila strain lacking CRISPR-Cas led to on an average 60-fold more infectivity (Gunderson and Ciancioatto 2013). In Acinetobacter baumannii, harboring type II system, the CRISPR-Cas was shown to be associated with high biofilm-producing ability. These strains in addition to containing genes exclusive for biofilm formation also lacked plasmids, which suggested that the CRISPR-Cas has dual role of adaptive immune function as well as a modulator of group behavior (Mangas et al. 2019). Moreover, Cas3 protein from the type I system, in addition to type II system, was found to modulate the pathogenicity and virulence of several microorganisms. For instance, in Salmonella enteric, cas3 deletion was shown to decrease the biofilm-forming capacity (Cui et al. 2020). In addition, loss of cas3 also affected the intracellular invasiveness inside the host. Transcriptome analysis revealed that cas3 deletion affected quorum-sensing genes, type three secretion systems (T3SS), salmonella pathogenicity island-1 (SPI-1), and genes related to flagella formation (Cui et al. 2020). Cas3 was also reported to regulate the biofilm formation in addition to fluoride resistance in Streptococcus mutans (Tang et al. 2019). In contrast to other studies, cas3 deletion in Porphyromonas gingivalis increased the virulence inside the THP-1 cells. These results also concurred with the earlier studies on Galleria mellonella infection model, whereby infection with cas3 deletion mutant led to high mortality compared to when infected with wild-type strain (Solbhat et al. 2020). The table (Table 1) given below includes some of the other examples of noncanonical functions performed by CRISPR-Cas system in the prokaryotes.

Bacterial physiology

In Myxococcus xanthus, three cas genes including cas8, cas7, and cas5 known as devT, devR, and devS, respectively, were found to be essential for sporulation and fruiting body
| Function | CRISPR-Cas type | Mechanism | Self-targeting | Organism | Ref |
|----------|----------------|-----------|----------------|----------|-----|
| Regulation of pathogenicity and virulence | Type I-F | Spacer targeting of prophage gene | Yes | *P. aeruginosa* | Zegans et al. (2009) |
| | Type II-C | Cas9 binding to noncanonical targets | No | *N. meningitis* | Heidrich et al. (2019) |
| | Type II | *cas1*, *cas2*, and *csn1* affect the virulence by unknown mechanism | No | *C. jejuni* | Louwen et al. (2013) |
| | Type II | *cas9*-dependent regulation of virulence factors | No | *S. pyogenes* | Gao et al. (2019) |
| | Type II-B | *cas2* nuclease activity | No | *L. pneumophila* | Gunderson and Cianciotto (2013) |
| | Type II | *cas9*-mediated regulation of virulence associated genes | No | *C. jejuni* | Shabbir et al. (2018) |
| | Type II | Cas9-linked presence of biofilm-forming genes | No | *A. baumannii* | Mangas et al. (2019) |
| | Type I-E | *Cas3*-mediated regulation of quorum-sensing gene, biofilm-forming genes, and secretion system (T3SS) | No | *S. enterica* | Cui et al. (2020) |
| | Types I-C & II-A | Cas3-mediated regulation of virulence genes | No | *S. mutans* | Tang et al. (2019) |
| | Type I | Cas3-mediated regulation of virulence genes | No | *P. gingivalis* | Solbiati et al. (2020) |
| Regulation of physiology | Type I-B | Cas8 affects *fraA* gene regulation | No | *M. xanthus* | Viswanathan et al. (2007) |
| | Type I-E | Cas5 and Cas2 affect OmpC protein levels, and Cas3, Cas1, Cas6a, and Cse2 affect OmpF protein levels | No | *S. typhi* | Medina-Aparicio et al. (2021) |
| | Types I-C & I-F | Spacer target the glycogen phosphorylase (glp) gene | Yes | *A. actinomycetemcomitans* | Jorth and Whiteley (2012) |
| | Type II | Spacer targeting two-component systems, a transcriptional regulator, and iron transport gene | Yes | *L. monocytogenes* | Mandin et al. 2007; Sesto et al. (2014) |
| Stress response | Type I–E | CasABCDE component of cascade complex-mediated gene silencing of excretory pathway | No | *E. coli* | Perez-Rodriguez et al. (2011) |
| | Type II | *Cas9* overexpression | No | *S. thermophilus* | Young et al. (2012) |
| | Type I-A, types I-D & III-B | Overexpression of type I-A and type I-D associated Cas proteins | No | *S. islandicus* | Quax et al. (2013) |
| | Types II-A & I-C | Regulation of *cas* genes expression by VicR/K system differentially | No | *S. mutans* | Serbanescu et al. 2015 |
| | Types II-A & I-C | c-di-AMP-mediated differential regulation of CRISPR-Cas expression | No | *S. mutans* | Cheng et al. (2016) |
| | Types I-C & I-B | Unknown | NO | *B. cereus* | Zheng et al. (2020) |
development (Thony-Meyer and Kaiser 1993). Further, it was observed that devT mutation affected the aggregation, sporulation, and chemotaxis. It was found that the levels of devT affected the transcript levels of the fruA gene, responsible for fruiting body formation (Boysen et al. 2002). It was suggested that the FruA protein has an affinity for the regulatory site in the cas locus, which induces the devT expression and subsequently leads to the positive feedback loop. The role of the CRISPR array in the regulation of fruiting body formation is not clear, though two self-targeting spacers were observed in the CRISPR array in Myxococcus xanthus. Among the two, one was targeting prophage gene encoding integrase, and the other was targeting the cas gene. From the observation, it was inferred that the both cas gene and CRISPR array might be functioning together to alter the physiology of the organism

| Function          | CRISPR-Cas type | Mechanism                                                                 | Self-targeting | Organism            | Ref             |
|-------------------|-----------------|---------------------------------------------------------------------------|----------------|---------------------|-----------------|
| Gene regulation   | Type II         | Cas9-mediated repression of BLP-producing gene                            | No             | *F. novicida*       | Jones et al. (2012) |
|                   | Type I          | *his* gene repression by self-targeting spacer                           | Yes            | *P. carbinolicus*   | Aklujkar and Lovley (2010) |
|                   | Type III        | Cmr complex-mediated targeting of RNA                                    | No             | *P. furiosus*       | Hale et al. (2012) |
|                   | Type III        | Cmr complex-mediated targeting of RNA                                    | No             | *P. gingivalis*     | Endo et al. (2015) |
| Genome remodeling | Type I-F        | Removal of 40–75 kb to entire 100 kb HAI2 pathogenicity island by self-targeting spacer | Yes            | *P. atrosepticum*   | Vercoe et al. (2013) |
|                   | Type II         | Removal of either targeted *emrB* gene or entire 66 kb integrated plasmid by self-targeting spacer | Yes            | *E. faecalis*       | Hullahalli et al. (2018) |
|                   | Type III        | Deletion of 16 kb fragments in the targeted region                       | Yes            | *S. aureus*         | Guan et al. (2017) |
|                   | Type II-A       | Removal of spacer targeted mobile genetic element (MGE)                  | Yes            | *S. thermophilus*   | Canez et al. (2019) |
|                   | Types I-E & II-A | Deletion of 34 kb genomic locus mediated by self-targeting spacer        | Yes            | *S. thermophilus*   | Selle et al. (2015) |
|                   | Type I-E        | DNA repair protein RecB and RecG are involved in naïve and primed adaptation processes | No             | *E. coli*           | Killelea and Bolt (2017) |
|                   | Type I-E        | Cas1 interacts with DNA repair machinery                                 | No             | *E. coli*           | Ivancic-Bace et al. (2015) |
|                   | Type I-E        | Adaptation process dependent on the activity of RecBCD dsDNA break repair complex | No             | *E. coli*           | Babu et al. (2011); Levy et al. (2015) |
|                   | Type I-E        | RecA loading onto RecBCD repair complex inhibits the adaptation          | No             | *E. coli*           | Radovcic et al. (2018) |
|                   | Types I-E & II-A | Positive association between RecBCD DNA repair system and type I-E CRISPR-Cas system and negative association between type II-A CRISPR-Cas system and NHEJ | No             | *Proteobacteria and Firmicutes* | Bernheim et al. (2019) |
Response to stress

In many bacterial species, it has been observed that in response to envelop stress, the expression of the CRISPR-Cas system is induced. This phenomenon was first observed in *Escherichia coli*, where the overexpression of membrane targeting protein activated the expression levels of the CRISPR-Cas system (Perez-Rodriguez et al. 2011). It was also observed that the levels of reporter green fluorescent protein (GFP), fused to the excretory molecule (ssTorA), were reduced in cells lacking chaperone DnaK. Deletion of type I-E cas operon, as well as the BaeSR system, restored the expression of ssTor-GFP. BaeSR, a two-component system, is well-known to upregulate the cas genes expression in response to membrane stress (Baranova and Nikaido 2002; MacRitchie et al. 2008). It was suggested that in *E. coli*, the combined effect of induced Cas levels and occurrence of ssTorA-targeting spacer (partially matching) in CRISPR array led to reduction in the levels of membrane protein, thereby impacting the transport channel across the cell (Perez-Rodriguez et al. 2011). In addition, in *Streptococcus thermophilus* and Archaean *Sulfolobus islandicus*, the CRISPR-Cas system was found to be induced in the aftermath of phage induced envelope stress (Quax et al. 2013; Young et al. 2012). In *Myxococcus xanthus*, dev operon corresponding to type I-B CRISPR cassette is also found to be activated during stress (Viswanathan et al. 2007). In *Streptococcus mutans*, it was observed that type II-A and type I-C CRISPR systems are involved in temperature stress tolerance. At higher temperature, the double mutants lacking both CRISPR types showed reduced capabilities to survive in comparison to single CRISPR mutant. Moreover, type II-A CRISPR deletion mutant showed lower growth rate during membrane stress and oxidative stress as compared to type I-C mutants. Intriguingly, it was observed that two-component stress response regulator VicK/R regulated these two CRISPR systems differently. This suggested that various stress factors modulate the expression levels of CRISPR systems in the organism through the channel of signaling molecules with one such being the VicK/R tw-component system. VicK/R was observed to negatively regulate the type II-A system and positively regulate the type I-C system (Serbanescu et al. 2015). In another study, in *Streptococcus mutans*, deletion of cdaA gene, coding for diadenylate cyclase required for the synthesis of c-di-AMP molecule, led to upregulation of CRISPR 1 locus and downregulation of CRISPR 2 locus. Moreover, deletion of cdaA affected the sensitivity to hydrogen peroxide and production of extracellular polymeric substances. The authors suggested that the CRISPR-Cas systems are indirectly linked to the stress response mechanisms in this organism (Cheng et al. 2016). In *Bacillus cereus*, the introduction of the CRISPR-Cas in CRISPR-Cas negative strains resulted in decreased tolerance to various stresses as well as decreased pathogenicity (Zheng et al. 2020). In conclusion, it can be remarked that indeed these CRISPR-Cas systems are involved in functions other than the adaptive immunity, though most of the molecular mechanisms pertaining to these processes are still obscure and under investigation. The graphic description of the various environmental stress factors that activate the CRISPR-Cas system is depicted in the following Fig. 2.

Endogenous gene regulation

Various researchers speculated that spacers targeting the chromosome, which are being tolerated in the CRISPR array by the organism, might be involved in the gene regulation process (Sorek et al. 2008). Computational analysis of *Escherichia coli* type I-E CRISPR-Cas system, to investigate the spacer targets, revealed that CRISPR spacers have a high propensity
to target chromosomes as compared to phage genomes. In addition, it was revealed that spacers targeted transcriptionally active regions at a higher rate. The study suggested that the type I-E CRISPR-Cas system in *Escherichia coli* plays a role in endogenous gene regulation (Bozic et al. 2019). In addition to computational analysis, several experimental evidences showed that the CRISPR arrays are involved in gene regulation. In *Francisella novicida*, the type II CRISPR-Cas system was shown to modulate the expression of a protein-coding gene associated with bacterial virulence. The tracrRNA along with Cas9 and small CRISPR-associated RNA repressed the BLP expression inside the host. It was demonstrated that the bacterium avoided cell death by targeting the BLP mRNA instead of encoding gene (Jones et al. 2012; Sampson et al. 2013). The BLP protein triggers the host T cell-mediated immune response; thus lowering the expression levels of this protein. It enables the bacterium to escape the immune system and thrive inside the host (Sampson et al. 2013). Similarly, in *Pelobacter carbinolicus*, a self-targeting spacer was identified to target the histidyl-tRNA synthetase (*hisS*) and lower its expression. The crRNA having the spacer identical to *hisS* gene was found to compete with the RNA polymerase, thus hindering the transcription of *hisS* gene. The *hisS* gene unlike most of the self-targeting spacers was found at the trailer side of the 111 spacer-long CRISPR loci, indicating that the spacer is being tolerated for ages by the bacterium. This self-targeting of *hisS* gene also resulted in lower frequency of histidine amino acid in the protein molecules of *Pelobacter carbinolicus* in comparison to the other closely related strains (Aklujkar and Lovley 2010).

The type III system which targets mRNA instead of DNA is also observed to be involved in gene regulatory functions. For instance, in *Pyrococcus furiosus*, Cmr (Cas module RAMP termed as RNA associated mysterious proteins) complex was revealed to recognize and cleave the endogenous complementary RNAs, providing direct evidence that CRISPR-Cas can regulate the transcription and gene expression (Hale et al. 2012). Likewise, *Porphyromonas gingivalis* which contains the type III system also targets the mRNA and might lead to gene regulation (Endo et al. 2015). Nonetheless, Cas2 protein, part of the adaptation machinery in most CRISPR-Cas systems, can
CRISPR-Cas mediated physiological processes; CRISPR-Cas is activated due to extracellular signals such as phage infection, DNA invasion, and environmental stress. Upon activation, the CRISPR array and Cas proteins either function in combination or alone to carry out the various biological processes. CRISPR array usually mediates the pathophysiological changes or gene regulation processes by directly targeting the DNA or mRNA by a self-targeting spacer. Cas proteins modulate the cell physiology, virulence, and bacterial behavior either by activating the downstream pathways or by direct interaction with the other proteins or molecules. These processes are multifaceted, and the mechanism entailing various pathways awaits experimental elucidation.

**Bacterial genome remodeling**

In order to avoid cell death due to self-targeting spacers, organisms are shown to remodel the targeted region either by removing the protoscaler or deleting the entire targeted region. Several studies have observed that self-targeting forces the organism to evolve. In most of the circumstances, self-targeting of gene located in pathogenicity island (PI) resulted in deletion of entire PI regions. In *Pectobacterium atrosepticum*, spacer targeting of horizontally acquired island 2 (HAI2) led to deletion of the entire 100 kb HAI2 region to avoid the self-targeting of the genome (Vercoe et al. 2013). Spacer targeting erythromycin (*ermB*) gene, present in the plasmid and integrated between the flanking IS elements on both sides, also resulted in generation of deletion mutants either lacking the *ermB* gene or 75% of 66 kb plasmid (Hullahalli et al. 2018). Targeting of lacZ region both in gram-negative *Escherichia coli* carrying type I-E system and gram-positive *Streptococcus thermophilus* harboring type II-A systems also led to large genome deletions of up to 35 kb and 37.4 kb size, respectively (Canez et al. 2019; Cui and Bikard 2016). In line with these studies, in other gram positives including *Staphylococcus aureus* and *Streptococcus thermophilus*, self-targeting led to loss of large fragments of targeted regions (Guan et al. 2017; Selle et al. 2015). Thus, it can be concluded that the consequence of self-targeting
event, as suggested by various studies in diverse range of organisms comprising different CRISPR-Cas types, is the remodeling of the bacterial genome.

**Association with DNA repair system**

There exists a close association between the CRISPR-Cas systems and DNA repair machinery at different levels, and this has a diverse impact on the microorganisms. These two separate systems usually coincide directly at the adaptation step. In general, during the incorporation process, a nick generated in the first direct repeat by nucleases in order to incorporate a new spacer is again sealed by a DNA repair system (Killelea and Bolt 2017). It has also been shown in the type I-E system in *Escherichia coli* which utilizes the DNA polymerase from the DNA repair pathway during the spacer acquisition process (Ivancic-Bace et al. 2015). Besides, the common transcription regulator, Cas3a was seen to activate the process of CRISPR-spacer incorporation as well as DNA repair process, suggesting co-operation between the two at several stages (Liu et al. 2015). In addition, it has been shown that proteins involved in the DNA repair pathway and many of the Cas proteins share homologous regions such as helicase domain from Cas3 protein (type I system), nuclease from Cas4 protein (type I and type II), and RuvC like nuclease (type II and type V) (Hudaiberdiev et al. 2017; Shmakov et al. 2017; Sinkunas et al. 2013; Westra et al. 2012; Xiao et al. 2018; Zhang et al. 2012). Furthermore, experimental evidence in *Escherichia coli* revealed that casl deletion resulted in mutants defective in DNA repair pathway (Smith 2012).

Cas1 protein was also found to interact with several cellular repair proteins like RecB, RecC, and RuvB (Babu et al. 2011). The RecBCD system recognizes breaks in dsDNA during the replication process. This repair machinery moves along the replication fork and degrades damaged DNA till it detects the Chi site. An earlier study speculated that degraded DNA products of repair machinery are incorporated as a template for new spacer adaptation processes. Thus, RecBCD can be regarded as part of the anti-phage defense system as it degrades the linear phage DNA and facilitates the incorporation of the spacer. This system avoids the incorporation of the host genome into the CRISPR array by recognizing the Chi site which is present in the chromosomal DNA frequently at a very high rate (Levy et al. 2015). It was observed that RecB deletion hampered naïve spacer adaptation by type I-E systems in *E. coli* (Ivancic-Bace et al. 2015). According to the hypothesis, in the naïve acquisition process, RecBCD is recruited to repair the DNA damage induced by Cas1 during spacer incorporation (Ivancic-Bace et al. 2015; Levy et al. 2015). Recently, in a study, it was observed that RecG primes the adaptation of spacer from the MGEs by using the pre-existing spacer in type I systems (Heussler et al. 2016; Ivancic-Bace et al. 2015).

Despite sharing similarities at many levels and enabling each other’s successful function, there exists some antagonizing relationship between the two. The CRISPR-Cas adaptive immune function relies on the inability of the DNA repair pathway to reverse the DNA damage induced by Cas nucleases. In a comparative study, CRISPR-Cas machinery and DNA repair systems were analyzed in more than 5000 sequenced bacterial genomes. The study revealed that the type I CRISPR-Cas system has an affinity towards the RecBCD DNA repair system (Levy et al. 2015). This was found to be in concurrence with the observations made in *E. coli*, where both the systems were having synergistic interaction (Radovic et al. 2018). However, it was observed that the type II system negatively correlated with the non-homologous end joining (NHEJ) recombination pathway (Faure et al. 2019). To explain the complex relationship between the two, Bernheim and colleagues proposed a hypothesis. Accordingly, CRISPR-Cas systems are frequently subjected to transfer between the species by horizontal gene transfer (HGT). Upon introduction into some new species, CRISPR-Cas continuation in the organism is influenced by several factors, including their production cost, benefit incurred to the host, prevalent phage predation menaces in the population, and inheritance of other defense systems. In addition, genetic components especially the DNA repair system was also proposed to impact the sustenance of the CRISPR-Cas system in the organism. Proteins involved in the DNA repair system are reported to be under intense selection pressure, and Bernheim et al. proposed that in case of incompatibility, CRISPR-Cas system will be weeded out from the genome in order to maintain the repair systems. Owing to the strong selection process, organisms will wind up with different combinations of DNA repair systems and CRISPR-Cas systems. The results from this study might help to explain the checkered distribution and diversity of the CRISPR-Cas system in bacteria (Bernheim et al. 2019).

**CRISPR-Cas applications**

**Inhibition of horizontal gene transfer**

The rise in multidrug resistance in bacteria has forced the researcher to look for newer tools to tackle the problem of antibiotic resistance. Recently, CRISPR-Cas has also been used as a tool to inhibit the transfer of antibiotic resistance genes from one organism to another. The presence of the CRISPR-Cas system in a genome negatively impacts the concentration of antibiotic resistance determinants in multidrug-resistant (MDR) strains. In one of the study, it has been reported that the CRISPR system is certainly responsible for mitigating the dissemination of mobile genetic elements (MGEs) including...
prophages and plasmids in Enterococcus faecalis (Palmer and Gilmore 2010). Similarly, a higher number of spacers in the CRISPR array in Streptococcus pyogenes resulted in lower frequency of integrated prophages in the genome (Nozawa et al. 2011). Thus, there is a potential to harness CRISPR systems to curb the dissemination of resistant determinants and virulence in MDR/extreme drug-resistant (XDR) bacteria (Marraffini and Sontheimer 2008). Progress and development in genetic engineering tools made it feasible to introduce the resistance gene targeting spacer into any CRISPR array in a wide range of hosts (Sapranaukas et al. 2011). CRISPR arrays are being designed to target the pathogenicity islands containing antibiotic or virulence genes. This technique allows targeting of only those strains which carry such pathogenicity islands and spares the non-pathogenic populations (Bikard et al. 2014; Citorik et al. 2014; Gomaa et al. 2014). CRISPR-Cas as an antimicrobial resistance tool has been explored only recently. In a study, antibiotic resistance genes were targeted by incorporating the CRISPR-Cas system in lysogenic phage in Escherichia coli. In addition, a spacer targeting the lytic phage was also incorporated in the CRISPR array, and it was postulated that it will further protect the antibiotic-sensitized cells from lytic infection by phage (Yosef et al. 2015).

**Typing tool**

Way before, their role could be explored in bacterial physiology; CRISPR-Cas systems were utilized for typing the diversity in bacterial species (Grissa et al. 2008). For instance, in Mycobacterium tuberculosis, comprising type III CRISPR system, the variability in the spacer content had routinely been used for typing purposes and study Mycobacterium tuberculosis epidemiology (Abadia et al. 2010; Comas et al. 2009; Gomgnimbou et al. 2012; Kamerbeck et al. 1997). CRISPR typing was also explored in Campylobacter jejuni to reveal the phylogenetic relationship between strains (Schouls et al. 2003). However, unlike Mycobacterium tuberculosis, in Campylobacter jejuni (type II CRISPR system), CRISPR typing was proved to be insufficient to accomplish the understanding of evolutionary relationship. Therefore, CRISPR typing was used in combination with amplified fragment length polymorphism (AFLP) and multilocus sequence typing tool (MLST) (Schouls et al. 2003). A study revealed that polymorphism in the cas gene was linked to a gene found in Campylobacter jejuni strains isolated from patients with Guillain–Barre syndrome (Louwen et al. 2013; vanBelkum et al. 2001). Later on, it was established that single-nucleotide polymorphism along with CRISPR array variations is useful for typing process in Campylobacter jejuni (Louwen et al. 2013). This process of using sequence variation among the spacers in the CRISPR array for studying the diversity is known as spoligotyping. This technique was also used for typing in Corynebacterium diphtheriae and found to be more useful than random amplification of polymorphic DNA (RAPD) and AFLP (Mokrousov et al. 2009). In Legionella pneumophila as well, this technique enabled typing among the environmental strains (Gineva et al. 2012). Likewise the CRISPR type III and type II, type I CRISPR system was used for typing of Yersinia pestis and Salmonella enterica (Fabre et al. 2012; Vergnaud et al. 2007). CRISPR-Cas-based typing in Yersinia pestis, helped to identify the origin of the ancestral strain which caused the black plague (Vergnaud et al. 2007). In Salmonella, as well, CRISPR typing in combination with other techniques has significantly enhanced the means to distinguish strains causing outbreaks (Cao et al. 2013; Liu et al. 2011). Furthermore, in several other organisms, CRISPR typing has facilitated the delineation of subgroups. For instance, in Erwinia amylovora (type I-E), CRISPR typing helped to distinguish three major groups based on geographic origin (Rezzonico et al. 2011). CRISPR typing in Propionibacterium acnes confirmed three lineages of this organism that have also been identified using other typing techniques (Bruggemann et al. 2012). In Porphyromonas gingivalis, typing based on the CRISPR system, was observed that with this approach it was possible to regroup the strains beyond the fimbrial gene clusters (Watanabe et al. 2013).

**Genome editing**

Although all the CRISPR systems have the common feature of recognizing and targeting DNA, most of them are not suited for genetic engineering technologies. For instance, the DNA degradation process of the I st and IIId type CRISPR is not well characterized, and crRNA generation also requires additional Cas6 protein. In contrast, type II CRISPR systems require only a few CRISPR-Cas components for carrying out all the functions. Various studies concluded that the CRISPR-Cas9 complex is ideal for genome editing purposes. Type II targeting of phage and plasmid genetic elements was observed to introduce double-stranded breaks. It was considered to be an important feature from the genome editing point of view in mammalian cells (Garneau et al. 2010). Experimental evidence also revealed that the three elements of type II CRISPR-Cas machinery are more than sufficient to induce double-stranded breaks in mammalian DNA and facilitate the synthesis of single-guide crRNA (sgRNA). Cas9-mediated type II CRISPR systems have been used to target several DNA sequences using small targeting RNA. Small targeting RNA is also known as single-guide RNA (sgRNA) and is synthesized by fusing naturally existing tracrRNA and crRNA having sequence complementarity to the target sequence (Jiang et al. 2015). This genome editing tool has outrivaled all the other popular tools including transcription activator-like effector nucleases (TALENs) and zinc finger nucleases (ZFNs) (Wang et al. 2013). Genetic engineering involving CRISPR-Cas9 technology involves introduction of the Cas9- sgRNA construct into the cell using transformation. The co-transformation of synthesized recombination DNA template
along with the construct increases the chances of incorporating mutations into the target site (Cong et al. 2013). However, to circumvent continual fragmentation of mended homologous recombinant, the recombination DNA template must have mutations to hamper the Cas9 nuclease activity (Wang et al. 2013). Nevertheless, genome editing using Cas9 is a simple and efficient method for editing the genome of several eukaryotic as well as prokaryotic model organisms (Sander and Joung 2014).

Impeccable finding that CRISPR-Cas system can be used for genome editing technologies has opened up the prospects for the use of this system for therapeutic purposes especially for the treatment of cardiovascular diseases, genetic disorders, autoimmune, inflammatory, and immunological disorders, bacterial and viral diseases, metabolic disorders, neurodegenerative diseases, oncological disorders, and others (Bao et al. 2015; Ebina et al. 2013; Hai et al. 2014; Hu et al. 2014; Wu et al. 2016; Yan et al. 2018; Zhan et al. 2018). Furthermore, recently CRISPR-Cas technology has been used to develop SARS-CoV-2 vaccine candidates (Zhu et al. 2021). Post 2014, tremendous amount of interest had arisen among the scientist and pharmaceutical companies alike to explore and utilize this technology for gene therapy. Over the period, considerable volume of evidence has been gathered in the subject matter, substantiating the therapeutic applications, thus leading to the establishment of numerous strategic partnerships among leading pharmaceutical companies, centered around therapy development and clinical trials. Consequently, several CRISPR-Cas candidates designed for treatment of neurodegenerative and infectious diseases are currently under investigation. Numerous other clinical trials focusing oncological and hematological trials are also underway. Since 2015, close to 2000 patent applications concerning CRISPR-Cas technology has been filed or granted. Currently, based on the recent trends, the CRISPR-Cas abilities appear to be limitless, and the CRISPR-Cas revolution does not seem to end in the near future. However, despite the extraordinary potential that the CRISPR-Cas technology holds, further investigation, probing its safety and therapeutic efficacy in large diverse populations, is required (https://www.researchandmarkets.com/r/05e94f/ 09/17/2021).

**Conclusion**

The multiple studies discussed above reasonably support the perspective that CRISPR-Cas systems have a role beyond immunity. Furthermore, it is believed that CRISPR-Cas system, though initially evolved to perform the adaptive immune function in microorganisms, picked up the non-canonical functions later as an adaptation mechanism to various cues from surrounding. However, whether the system evolved as whole, i.e., CRISPR and Cas evolved simultaneously or underwent the adaptation process separately without the influence of the other component, is still uncertain. Owing to the limited amount of information, that too from select few species, it will be too early to reach at any conclusion. It can be said that CRISPR-Cas system is still evolving and its components are picking up noncanonical functions while adjusting in the newer species under novel circumstances. The flexibility and uniqueness of this system to repurpose its function as adaptive defense system of prokaryotes to the modulator of bacterial pathophysiology and several other functions suggest that these systems can be exploited to develop powerful diagnostic, therapeutic, and experimental tools.

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VD, conceptualization and writing—original draft; SC, conceptualization and reviewing; KH, reviewing.

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**Availability of data and material**  
As no datasets and material were generated in this study, therefore, this section is not applicable.

**Code availability**  
Not applicable.

**Declarations**

**Ethics approval**  
Not applicable.

**Consent to participate**  
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**Conflict of interest**  
The authors declare no competing interests.

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