Mobile Genetic Elements and Evolution of CRISPR-Cas Systems: All the Way There and Back

Eugene V. Koonin* and Kira S. Makarova

National Center for Biotechnology Information, National Library of Medicine, National Institutes of Health, Bethesda, Maryland

*Corresponding author: E-mail: koonin@ncbi.nlm.nih.gov.

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Abstract

The Clustered Regularly Interspaced Palindromic Repeats (CRISPR)-CRISPR-associated proteins (Cas) systems of bacterial and archaeal adaptive immunity show multifaceted evolutionary relationships with at least five classes of mobile genetic elements (MGE). First, the adaptation module of CRISPR-Cas that is responsible for the formation of the immune memory apparently evolved from a Casposon, a self-synthesizing transposon that employs the Cas1 protein as the integrase and might have brought additional cas genes to the emerging immunity loci. Second, a large subset of type III CRISPR-Cas systems recruited a reverse transcriptase from a Group II intron, providing for spacer acquisition from RNA. Third, effector nucleases of Class 2 CRISPR-Cas systems that are responsible for the recognition and cleavage of the target DNA were derived from transposon-encoded TnpB nucleases, most likely, on several independent occasions. Fourth, accessory nucleases in some variants of types I and III toxin and type VI effectors RNases appear to be ultimately derived from toxin nucleases of microbial toxin–antitoxin modules. Fifth, the opposite direction of evolution is manifested in the recruitment of CRISPR-Cas systems by a distinct family of Tn7-like transposons that probably exploit the capacity of CRISPR-Cas to recognize unique DNA sites to facilitate transposition as well as by bacteriophages that employ them to cope with host defense. Additionally, individual Cas proteins, such as the Cas4 nuclease, were recruited by bacteriophages and transposons. The two-sided evolutionary connection between CRISPR-Cas and MGE fits the “guns for hire” paradigm whereby homologous enzymatic machineries, in particular nucleases, are shuttled between MGE and defense systems and are used alternately as means of offense or defense.

Key words: CRISPR-Cas systems, mobile genetic elements, casposons, CRISPR adaptation, CRISPR effector modules.

Introduction

Thanks to the striking success of the Cas9 endonucleases as new generation of genome editing tools, in recent years, comparative genomics, structures, biochemical activities and biological functions of Clustered Regularly Interspaced Palindromic Repeats (CRISPR)-CRISPR-associated proteins (Cas) systems and individual Cas proteins have been explored with an unprecedented intensity (Sorek et al. 2013; Barrangou and Marraffini 2014; Doudna and Charpentier 2014; Hsu et al. 2014; Mohanraju et al. 2016; Wright et al. 2016; Barrangou and Horvath 2017; Komor et al. 2017; Koonin et al. 2017). The CRISPR-Cas are adaptive (acquired) immune systems with memory of past encounters with foreign DNA that is stored in unique spacer sequences derived from viral and plasmid genomes and inserted into CRISPR arrays. Transcripts of the spacers, along with portions of the surrounding repeats, are utilized as guide CRISPR (cr)RNAs to recognize the cognate sequences in foreign genomes and thus direct Cas nucleases to their unique cleavage sites.

Because CRISPR-Cas are programmable immune systems that can adapt to target any sequence, they are not subject to extreme diversifying selection that led to the evolution of the immense variety of restriction-modification enzymes, the most abundant form of innate immunity in prokaryotes (Pingoud et al. 2016). Nevertheless, CRISPR-Cas systems evolve in a regime that is common to all defense system, namely continuous arms race with genetic parasites, primarily viruses, resulting in rapid evolution of at least some cas gene sequences (Takeuchi et al. 2012). Furthermore, the notable diversity of the gene compositions and genomic architectures of the CRISPR-cas loci translates into diversification of the molecular mechanisms of defense (Makarova et al. 2011, 2015).

The CRISPR-Cas belong to the class of nucleic acid-guided defense systems, along with eukaryotic RNAi and prokaryotic...
Argonaute-based machinery (Hutvagner and Simard 2008; Shabalina and Koonin 2008; Hur et al. 2014; Swarts et al. 2014; Koonin 2017). Unlike the Argonaute systems and most of the forms of the eukaryotic RNAi but similarly to the piRNA branch of RNAi, CRISPR-Cas mediates bona fide adaptive immunity (Makarova et al. 2006; Koonin and Makarova 2009; van der Oost et al. 2009; Marraffini and Sontheimer 2010). The CRISPR-cas genomic loci are modified to target the genome of a unique pathogen or its closest relatives with exceptional specificity and efficiency. These loci typically consist of a CRISPR array, that is, from several to several hundred direct, often partially palindromic, exact repeats (25–35 bp each), separated by unique spacers (typically, 30–40 bp each) and the adjacent cluster of multiple cas genes that are organized in one or more operons. The CRISPR-Cas immune response consists of three stages: 1) adaptation, 2) expression, and 3) interference. At the adaptation stage, a distinct complex of Cas proteins binds to a target DNA molecule, migrates along that molecule and, typically after encountering a distinct, short (2–4 bp) motif known as Protospacer-Adjacent Motif (PAM), cleaves out a portion of the target DNA, the protospacer, and inserts it into the CRISPR array between two repeats (most often, at the beginning of the array) so that it becomes a spacer (Amitai and Sorek 2016; Jackson et al. 2017). Some CRISPR-Cas systems possess an alternative mechanism of adaptation, namely spacer acquisition from RNA via reverse transcription by a reverse transcription complex or Cas2 (Li et al. 2014; Kunne et al. 2016; Fagerlund et al. 2015; Amitai and Sorek 2016). In many variants, additional Cas proteins, such as Cas4 or Cas3 also contribute to the adaptation stage, in some cases forming fusions with Cas1 or Cas2 (Li et al. 2014; Kunne et al. 2016; Fagerlund et al. 2017).

In a sharp contrast to the relatively simple and uniform organization of the adaptation module, the effector modules are highly diverse, and their variation forms the basis of the current classification of CRISPR-Cas systems (Makarova et al. 2015; Koonin et al. 2017). On the basis of the organizational principles of the effector modules, all CRISPR-Cas systems are divided into Class 1, with multisubunit effector complexes comprised of several Cas proteins, and Class 2, in which the effector is a single, large, multidomain protein. Among other distinctions, Class 1 and Class 2 CRISPR-Cas systems substantially differ in the mechanisms of pre-crRNA processing. In Class 1 systems, the crRNAs are generated by a dedicated complex of multiple Cas proteins (Brouns et al. 2008; Wiedenheft et al. 2011; Rouillon et al. 2013; Spilmann et al. 2013; Staals et al. 2013). In Class 2 systems, processing is catalyzed either by an external bacterial enzyme, RNase III, with the help of an additional RNA species, the transactivating CRISPR (tracr) RNA (Deltcheva et al. 2011; Jinek et al. 2012, 2013a; Makarova et al. 2015; Shmakov et al. 2015; Jiang and Doudna 2017; Liu et al. 2017a), or by the same effector protein that is involved in target cleavage (East-Seletsky et al. 2016; Fonfara et al. 2016; Liu et al. 2017b; Zetsche et al. 2017).

The differences between Class 1 and Class 2 CRISPR-Cas systems extend to the interference stage. In Class 1, the processing complex containing the guide crRNA recognizes the target site and recruits an additional Cas protein (Cas3 in type I and Cas10 in type III) that contains the nuclease domain directly responsible for the target cleavage (Sinkunas et al. 2011; Gong et al. 2014; Redding et al. 2015). In Class 2, cleavage is performed by the nuclease domain(s) of the large effector protein (Deltcheva et al. 2011; Gasiunas et al. 2012; Jinek et al. 2012; Zetsche et al. 2015; Abudayyeh et al. 2016; Mohanraju et al. 2016; Wright et al. 2016; Barrangou and Horvath 2017; Jackson et al. 2017; Komor et al. 2017; Koonin et al. 2017; Nishimasu and Nureki 2017).

At the molecular level, the CRISPR-Cas systems possess a readily definable modular organization (Makarova et al. 2013a; Makarova et al. 2015). The two principal parts of the CRISPR-Cas systems are the adaptation and effector modules that consist, respectively, of the suites of genes encoding proteins involved in spacer acquisition (adaptation) and genes encoding Cas proteins involved in pre-crRNA processing that is followed by the target recognition and cleavage (interference). In most of the CRISPR-Cas systems, the adaptation module consists of the Cas1 and Cas2 proteins that form a complex, in which Cas1 is the endonuclease (integrase) involved in the cleavage of both the source, protospacer-containing DNA and the CRISPR array, whereas Cas2 forms the structural scaffold (Nunez et al. 2014, 2015; Wang et al. 2015; Amitai and Sorek 2016). In many variants, additional Cas proteins, such as Cas4 or Cas3 also contribute to the adaptation stage, in some cases forming fusions with Cas1 or Cas2 (Li et al. 2014; Kunne et al. 2016; Fagerlund et al. 2017).
**Class 1**

I-A
I-B
I-C
I-D
I-E
I-F
IV
IV variant
III-A
III-D
III-C
III-B
III-B variant

**Class 2**

II-B
II-A
II-C
II-C variant
V-A
V-B
V-C
V-D
V-E
VI-U
VI-A
VI-C
VI-B1
VI-B2

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**Fig. 1.**—Current classification of CRISPR-Cas systems. The organization of the CRISPR-cas loci and domain architectures of the effector proteins as well as the (predicted) target (DNA or RNA, or both) are shown for each subtype. The trees reflect the latest classifications for the Class 1 and Class 2 CRISPR-Cas systems (Makarova et al. 2015; Koonin et al. 2017) and are not traditional phylogenetic trees. The block arrows represent cas genes (not to scale); homologous genes are shown by the same color. For each cas gene, the systematic name and the legacy name (if any) are indicated below the respective arrow. The adaptation and effector modules are shaded in blue and light brown, respectively. A shaded outline for an arrow depicting a gene indicates that the gene in question is present only in a subset of CRISPR-cas loci of the respective subtype. For subtype III-D, a locus with a reverse transcriptase fused to cas7 is included; other reverse transcriptase-containing variants, from subtypes III-A and III-D, are not shown. SS, small subunit; TM, predicted transmembrane segment.

Dong et al. 2016; Jiang et al. 2016a; Yamano et al. 2016; Jiang and Doudna 2017; Liu et al. 2017a, 2017b; Smargon et al. 2017) (see more below). The composition and organization of the genes encoding effector module components have been comprehensively compared with delineate 6 types and 24 subtypes within the two CRISPR-Cas classes (Makarova et al. 2015; Koonin et al. 2017) (fig. 1). Various proteins involved in ancillary roles, such as regulation of the CRISPR response and other, still poorly characterized functions, can be assigned to a third, accessory module (Makarova et al. 2013a, 2014, 2015; Mohanraju et al. 2016). The modules of the CRISPR-Cas systems are partially autonomous as demonstrated by their frequent recombination as well as by the existence of isolated adaptation and effector modules in many bacterial and archaeal genomes (Makarova et al. 2015; Silas et al. 2017). However, it is important to note that the functional separation between the modules is only a rough approximation because some Cas proteins, in particular, Class 2 effectors, appear to be involved in all stages of the CRISPR response (Heiler et al. 2015, 2017).

The currently characterized diversity and classification of CRISPR-Cas systems are covered in several recent reviews. Comparative genomics analyses aiming at the reconstruction of the origins and evolution of the CRISPR-Cas systems have revealed a pervasive trend, namely multiple contributions of several classes of mobile genetic elements (MGE) to the emergence and diversification of the CRISPR-Cas immunity (Makarova et al. 2013a, 2015; Chylinski et al. 2014; Koonin and Krupovic 2015a; Mohanraju et al. 2016; Shmakov et al. 2017). In this article, we summarize the evidence on the roles of MGE in CRISPR-Cas evolution and combine it with the
indications of “reverse flow,” that is, hijacking of CRISPR-Cas systems and their components by MGE. We conclude that in prokaryotes, MGE and defense systems, in particular, CRISPR-Cas, form a dynamic network of genetic elements that continuously exchange genes.

**Origin of the Adaptation Module and Adaptive Immunity in Prokaryotes: The Casposons**

Cas1, the endonuclease responsible for spacer integration into CRISPR arrays, is not always encoded within CRISPR-cas loci. Analysis of the genomic surroundings of the “solo” cas1 homologs led to an unexpected discovery: the genomic context of these genes closely resembled transposable elements, with terminal inverted repeats (TIR) flanking DNA regions of 12–18 kb (Krupovic et al. 2014). These putative transposons all shared two genes encoding Cas1 and a DNA polymerase of family B, and additionally, encompassed variable sets of genes, mostly various nucleases and predicted DNA-binding containing helix-turn-helix (HTH) domains (fig. 2). Given the close similarity between the mechanisms of the reactions catalyzed by Cas1 during spacer integration into CRISPR arrays and by integrases during transposon integration, a natural prediction was that Cas1 functions as a bona fide integrase, hence the name Casposons for this class of MGE and casposase for the enzyme itself (Krupovic et al. 2014; Krupovic and Koonin 2016). Indeed, the integrase activity of the casposase has been promptly validated experimentally (Hickman and Dyda 2015), and moreover, similar target site specificities of Casposon integration and CRISPR spacer incorporation have been demonstrated (Beguin et al. 2016). The invariable presence of a DNA polymerase gene indicates that Casposons are self-synthesizing transposons that catalyze their own replication during transposition via a copy-and-paste mechanism. Before the discovery of the Casposons, self-synthesizing transposons, those of the Polinton class (also known as Mavericks), have been identified only in eukaryotes (Kapitonov and Jurka 2006; Pritham et al. 2007; Krupovic and Koonin 2015). The Casposons are not one of the particularly abundant prokaryotic MGE classes but nevertheless, show considerable diversity of genome organization, with four families characterized by distinct gene compositions. Transposition of Casposons has not been demonstrated directly but comparative genomic analysis of numerous strains of the archaeon *Methanosarcina mazei* has revealed clear signs of recent mobility, leaving little doubt that at least some of the Casposons are active MGE (Krupovic et al. 2016).

It appears likely that the entire adaptation module and perhaps even additional Cas proteins were contributed to the emerging CRISPR-Cas system by the ancestral Casposon (Koonin and Krupovic 2015a). The phylogenetic tree of the Cas1 family splits into two major branches, one of which consists of the casposases and the other one of CRISPR-associated Cas1 proteins, a topology that is compatible with a founding role of the Casposase in the evolution of CRISPR-Cas. Although the currently identified Casposons do not encode Cas2, some encode PD-DExK family nucleases homologous to Cas4, which is a component of the adaptation module in several CRISPR-Cas variants, and additional nucleases (Krupovic et al. 2014; Krupovic and Koonin 2016). Conceivably, the ancestral Casposon architecture including a gene for a Cas2 homolog remains to be discovered. Furthermore, the prototype CRISPR repeats and the leader sequence could have originated from either the TIRs or a duplicated target site of the ancestral Casposon (Krupovic et al. 2017).

The key role of Casposons in the origin of CRISPR-Cas is strongly supported by the close mechanistic similarity between the reactions catalyzed by the casposase during integration and by Cas1 during spacer incorporation into CRISPR arrays (Beguin et al. 2016). The founding event of CRISPR-Cas evolution could have been a random insertion of a Casposon in a vicinity of a putative ancestral innate immunity locus followed by immobilization of the inserted transposon and loss of some of its genes including the DNA polymerase (but see an alternative scenario below). The nature of the putative ancestral innate immunity system is perhaps the hardest puzzle of CRISPR-Cas evolution (Makarova et al. 2013a; Shmakov et al. 2015). Potentially, this locus could have encoded an ancestral Class 1 effector module that functioned without an adaptation module and CRISPR arrays, by directly employing guide RNA derived from transcripts of foreign genomes, analogously to Argonaute-based defense systems (Hur et al. 2014; Swarts et al. 2014). The organization of such potential ancestral innate immunity loci could resemble type IV CRISPR-Cas (Makarova et al. 2015) and the “minimal” variants of type I encoded by a distinct family of Tn7-like transposons (fig. 2; and see below).

**CRISPR-Cas Adaptation Modules Containing RTs**

A distinct variety of the CRISPR-Cas adaptation module that is widely represented in type III includes a RT that, in many cases, forms a fusion protein with Cas1 or alternatively, is encoded by a separate gene adjacent to cas1 (fig. 2). It has been shown that the RT-containing CRISPR-Cas systems are capable of acquiring spacers from RNA via reverse transcription, in addition to the spacer acquisition from DNA that is universal in CRISPR-Cas (Silas et al. 2016). Phylogenetic analysis of the RT superfamily indicates that most of the CRISPR-associated RTs form a monophyletic group that is affiliated with the RTs of Group II introns (Toro et al. 2014; Silas et al. 2017). The RT-Cas1 fusion represented in diverse type III loci appears to have emerged at a single point in evolution, conceivably, subsequent to a random insertion of a Group II intron into a type III CRISPR-cas locus. There is little if any correlation between
the RT phylogeny and the CRISPR-Cas subtypes that are defined primarily from the organization of the effector modules (fig. 1). Furthermore, many RT-cas fusions are not associated with an effector module subunits but often are located adjacent to a CRISPR array. Taken together, these observations indicate that the RT-containing adaptation modules can combine or function in trans with most Type III systems. Thus, the Cas1-RT combination itself can be viewed as a MGE with an organization resembling that of Group II introns that combine RT with a homing endonuclease and eukaryotic retroelements encoding RT and integrases (Eickbush and Jamburuthugoda 2008; Zimmerly and Wu 2015; McNeil et al. 2016) although this element lacks the typical RNA structures of Group II introns and is not a Group intron per se (Silas et al. 2017).

In addition, two cases of apparent independent acquisition of RT by CRISPR-cas loci have been detected (Silas et al. 2017).
One of these includes a small group of type III-B loci that are represented primarily in mesophilic archaea of the genus Methanosarcina. Phylogenetic analysis of the RT indicates that these loci have independently acquired the RT gene from a Group II intron. The second case includes type I-E CRISPR-Cas systems in bacteria of the genus Streptomyces to which the RT gene was apparently transferred from a type III CRISPR-cas locus. In these minor groups of the RT-containing CRISPR-Cas systems, RT is not fused to Cas1 or any other Cas protein, and the type III-B loci of Methanosarcina lack the cas1 gene altogether suggesting that RT interacts with Cas1 in trans. The independent fixation of the captured RT in these groups of archaea and bacteria seems to be indicative of the utility of the RT-mediated acquisition of spacers from RNA for increasing efficiency of the CRISPR response.

**Type II and Type V Effectors: Parallel Capture of Transposon-Encoded Nucleases**

The common feature of all type II and type V effectors (Cas9 and Cas12 proteins, respectively) is the presence of a RuvC-like endonuclease domain (fig. 3). However, the sequence similarity between the RuvC-like domains of Cas9 and Cas12, and even between different subtypes within each type is very low such that these proteins can be recognized as homologs only by highly sensitive profile searches or structural comparisons. Moreover, other portions of Cas9 and Cas12 show no similarity to each other and appear not to be homologous (Shmakov et al. 2015, 2017). The structures of several Cas9 variants (Jinek et al. 2014; Nishimasu et al. 2014, 2015; Hirano et al. 2016), Cas12a (Cpf1) (Dong et al. 2016; Yamano et al. 2016), and Cas12b (C2c1) (Liu et al. 2017a; Yang et al. 2016) complexed with the guide RNA, target DNA and, in the cases of Cas9 and Cas12b, tracrRNA have been reported. All these effector proteins are similar in size and general shapes, that is, a bilobed, “jaw-like” structure that accommodates the target DNA and the guide RNA between the lobes. However, the structures cannot be superimposed apart from the RuvC-like domains (Lewis and Ke 2017). Furthermore, the RuvC-like domains of Cas9 and Cas12a contain inserts, in similar but not identical positions, that represent nonhomologous domains, the HNH family nuclease and a novel NUC domain, respectively. In type II systems, the HNH domain of Cas9 cleaves the target DNA strand (i.e., the strand complementary to the crRNA) whereas the RuvC-like nuclease cleaves the nontarget strand (Gasiusnas et al. 2012; Jinek et al. 2012). It has been initially concluded that in Cas12a, the NUC domain is the nuclease responsible for the cleavage of the target strand (Dong et al. 2016; Yamano et al. 2016). However, a subsequent, detailed biochemical study has suggested that the RuvC-like nuclease of Cas12a was responsible for the cleavage of both strands whereas the NUC domain facilitated the cleavage of the target strand in a nonenzymatic fashion (Swarts et al. 2017). Similarly, although the RuvC-like nuclease of Cas12b also contains a unique inserted domain, both DNA strands apparently are cleaved by the RuvC-like nuclease, which undergoes a major conformational change triggered by the initial, nontarget strand cleavage (Liu et al. 2017a; Yang et al. 2016). Thus, comparative analysis of the sequences and structures of type II and type V effectors reveals a remarkable interplay between homology, diversification and convergence which reflects their complex evolutionary history.

The key insight into the evolution of the type II and type V effectors has been the observation that, among other members of the RuvC-like nuclease family, these effectors, particularly Cas12, show the highest sequence similarity to TnpB proteins of the IS605 and other related families of transposons (Chylinski et al. 2014; Shmakov et al. 2015). The tnpB genes are extremely abundant in bacterial and archaeal genomes and belong to autonomous transposons, which additionally encode a transposase (TnpA) but more frequently nonautonomous transposons, in which TnpB is the only protein product including the eukaryotic Fanzor transposons (Bao and Jurka 2013). The role of TnpB in the transposon life cycle remains unclear given that this protein is not required for transposition (Pasternak et al. 2013), but the conservation of the RuvC-like endonuclease catalytic sites in most TnpB sequences indicates that these proteins are active nucleases.

Remarkably, the effectors of type II and different subtypes of type V (Cas9 and Cas12a, 12b, 12c, respectively) showed the highest similarity to different groups of TnpB proteins, suggesting independent origins for the effectors of different types and subtypes of Class 2. In these cases, informative phylogenetic trees could not be constructed due to the low sequence conservation. Nevertheless, the specific ancestry of Cas9 could be readily traced to a distinct family of transposons (denoted ISC, after Insertion Sequences Cas9-related) thanks to the shared domain architectures of the IscB and Cas9 proteins (fig. 3), in which an HNH endonuclease domain is inserted into the RuvC-like domain (Chylinski et al. 2014; Kapitonov et al. 2015).

The likely scenario for the origin of type V effectors from TnpB has become clearer with the identification of a distinct variety of putative CRISPR-Cas systems, denoted subtype V-U (after Uncharacterized), that lack adaptation modules and consist of TnpB homologs encoded next to CRISPR arrays (Shmakov et al. 2017). The putative V-U effector proteins are much smaller than Cas9 or Cas12 and similar in size or only slightly larger than typical, transposon-encoded TnpB. Moreover, unlike Cas9 and Cas12, the TnpB homologs from the V-U loci show highly significant similarity to the transposon proteins, allowing construction of robust phylogenetic trees, which strongly support independent origin of Class 2 effectors from different TnpB subfamilies (Shmakov et al. 2017). The functionality of the type V-U systems so far has not been demonstrated in direct experiments.
Nevertheless, multiple lines of evidence, namely evolutionary conservation in diverse groups of bacteria, the difference between the spacer complements of even closely related bacteria, and the presence of spacers homologous to phage genome sequences, indicate that at least some V-U variants are active immune systems. The close relationship between the putative V-U effectors and bona fide TnpB implies that V-U are recently evolved CRISPR-Cas variants. The current scenario for type II and type V evolution includes multiple, random insertions of nonautonomous TnpB-encoding transposons next to CRISPR arrays, with subsequent convergent “maturation” of the effectors that involved acquisition of additional domains (Shmakov et al. 2017). The newly gained domains are unrelated between different subtypes, but in

Fig. 3.—Contributions of mobile genetic elements to the origin and evolution of CRISPR-Cas systems: Class 2 effector proteins. The genes for Class 2 effector proteins and the homologous proteins encoded by transposons or toxin–antitoxin modules are shown by block arrows (roughly to scale). Different colors denote distinct domains or uncharacterized regions in the effector proteins. The curved arrows show putative ancestor–descendent relationships. RuvC I, II, III are distinct amino acid motifs that jointly comprise the catalytic site of the RuvC-like nuclease.
each case, the outcome is the emergence of a full-fledged effector protein capable of accommodating the complex of the crRNA and the target DNA (fig. 3).

**Contributions of Toxin–Antitoxin Modules to CRISPR-Cas Evolution**

In addition to two distinct classes of transposons and group II introns, multiple contributions to the evolution of CRISPR-Cas systems apparently come from a very different type of MGE, the toxin–antitoxin (TA) modules. These modules consist of a toxin which, in most common variants, is an RNase (known as interferase) that specifically cleaves ribosome-associated mRNAs and an antitoxin that forms a complex with the toxin and reversibly inhibits its activity (Gerdes et al. 2005; Makarova et al. 2009; Van Melderen and Saavedra De Bast 2009; Van Melder 2010; Gerdes 2012). The TA modules lack mechanisms of active mobility but are typically transferred on plasmids and are “addictive” to the host cells, which die if they do not receive the TA-carrying plasmid upon segregation. This postsegregational cell killing occurs because antitoxins are unstable proteins, compared with toxins, with a much shorter half-life, and therefore, unless the toxin and antitoxin are continuously produced in stoichiometric amounts, the unleashed toxin exerts its deleterious effect on the cell lacking the TA genetic locus. Apart from plasmids, numerous TA loci are carried by bacterial chromosomes, and in addition to their selfish properties, appear to perform defense functions, namely inducing dormancy or programmed cell death (PCD) as an “altruistic” defense strategy.

Notably, at least two unrelated classes of TA modules contributed to the evolution of CRISPR-Cas (fig. 3). The structural subunit of the adaptation complex, Cas2, is derived from the VapD family of interferases, which adopt a distinct version of the RNA Recognition Motif fold (Makarova et al. 2006). The interferase catalytic site is conserved in Cas2 proteins from some CRISPR-Cas systems but disrupted in others, and the function of the demonstrated nuclease activity of Cas2 (Beloglazova et al. 2008; Ka et al. 2014; Dixit et al. 2016) in the CRISPR-mediated defense remains obscure given that this activity is not required for adaptation. One possibility appears to be that Cas2 actually functions as a toxin that could induce dormancy or PCD in cases when the immune function of CRISPR-Cas fails (Makarova et al. 2012; Koonin and Zhang 2017). A different, unrelated toxin RNase, the HEPN domain, is present in several Cas proteins. The HEPN domain-containing Cas proteins include the effectors of the RNA-cleaving type VI CRISPR-Cas systems effectors (Cas13), which contain two diverged HEPN domain copies (Shmakov et al. 2015, 2017; Abudayeh et al. 2016; Liu et al. 2017b; Smargon et al. 2017), and the Csm6 and Csx1 proteins that are responsible for the ancillary RNA cleavage in type III-A systems and type III-B systems, respectively (Elmore et al. 2016; Jiang et al. 2016b; Niewoechner and Jinek 2016; Sheppard et al. 2016). A highly diverged HEPN domain is contained also in a small protein that has been shown to regulate the activity of the effector protein Cas13b in one of the variants of the type VI-B CRISPR-Cas systems (Smargon et al. 2017). In the Csm6 and Csx1 proteins, the HEPN RNase is fused to a CRISPR-Associated Rossmann Fold (CARF) domain, which implies regulation by a nucleotide ligand (Makarova et al. 2014). In a striking recent development, the ligands of the CARF domain of Csm6 have been identified as cyclic oligoadenylates that are synthesized by the Cas10 proteins, the polymerase-cyclase large subunit of type III effectors complexes, in response to the target RNA recognition (Kazlauskiene et al. 2017; Niewoechner et al. 2017).

The HEPN superfamily, which includes primarily RNases involved in various defense-related functions in both prokaryotes and eukaryotes, in general, and the CRISPR-associated HEPN domains in particular, show extreme sequence divergence such that their identification often requires careful manual examination of protein alignments (Anantharaman et al. 2013; Shmakov et al. 2015; Smargon et al. 2017). Due to this low sequence conservation, reliable phylogenetic analysis and confident identification of the specific ancestral relationships of the HEPN domains are impractical. Nevertheless, with the exception of the CRISPR-associated HEPN RNases, all other prokaryotic HEPN domains appear to belong to TA or Abortive Infection modules, which are extremely abundant in many bacteria and archaea, especially thermophiles (Makarova et al. 2009, 2013b). These TA modules are, in general, poorly characterized, but the toxin activity of one of the HEPN domains from such a module has been demonstrated experimentally (Yao et al. 2015). Thus, there is little doubt that the CRISPR-Cas systems have originally coopted a toxin RNase on at least one but possibly multiple occasions (fig. 3).

There is an intriguing possibility that the toxicity of the HEPN domain RNases remains relevant in the context of the CRISPR-mediated defense as indicated by the recent characterization of the type VI interference activity. Once activated by the recognition of the cognate target RNA, the Cas13a and Cas13b proteins become promiscuous RNases that cleave RNA nonspecifically (Abudayeh et al. 2016; Smargon et al. 2017). Expression of the Cas13 proteins together with the guide and target RNAs is toxic for bacteria, suggesting induction of dormancy or PCD in bacteria (Abudayeh et al. 2016). This function of the type VI effectors remains to be studied in detail but appears compatible with the hypothesis on coupling between immunity and dormancy induction and/or PCD in prokaryotes (Makarova et al. 2012; Koonin and Zhang 2017).

**Capture of CRISPR-Cas Systems and Individual Cas Genes by MGEs**

In the preceding section, we described the recruitment of proteins, primarily nucleases, from widely different MGE for...
functions in the CRISPR-mediated adaptive immunity. Here we address the reverse gene flow, from CRISPR-Cas systems to MGE. The most wide spread case of such recruitment includes a distinct group of Tn7-like transposons, which contain “minimal” type I-F CRISPR-Cas systems that consist of a reduced effector module and a short CRISPR array (fig. 4) (Peters et al. 2017). Several smaller groups of Tn7-like transposons encompass similarly truncated type I-B CRISPR-Cas systems. Notably, these type I variants lack the Cas3 helicase and the associated HD domain that are required for target cleavage (Beloglazova et al. 2011; Mulepati and Bailey 2011; Huo et al. 2014). Accordingly, they are predicted to be active in pre-crRNA processing yielding mature crRNAs as well as target binding but not interference that requires target cleavage. Phylogenetic analysis indicates a single, ancestral capture of a type I-F CRISPR-cas locus by the founder of a specific subfamily of Tn7-like transposons and two independent instances of type I-B loci capture (Peters et al. 2017). The transposon-associated CRISPR arrays contain multiple spacers homologous to plasmid and bacteriophage sequences, and in several cases, also bacterial chromosomal sequences adjacent to the transposon. Taken together, these observations prompt the hypothesis that the transposon-encoded CRISPR-Cas systems generate displacement (R-loops) in the cognate DNA sites, targeting the transposon to these sites and thus facilitating their spread, particularly, via plasmids and phages (Peters et al. 2017). Experimental study of the transposon-encoded CRISPR-Cas systems and their involvement in the transposon life cycle can be expected to shed new light on the functional and evolutionary interplay between CRISPR-Cas and MGE. More specifically, we do not currently understand what features of the Tn7 biology make this particular

![Diagram of CRISPR-Cas systems and mobile genetic elements.](image-url)
transposon family prone to capture and maintain CRISPR-Cas systems. Elucidation of the role of CRISPR-Cas in the life cycle of Tn7 is likely to uncover such features.

Similarly to the above cases, many type IV CRISPR-Cas systems are encoded on plasmids and might contribute to the plasmid–host interaction, in particular inhibiting host defense including resident CRISPR-Cas. The complete type IV systems encompass genes for the effector complex, Cas6 and CRISPR arrays but many lack an array, suggesting that they rely on arrays in the host genome (fig. 1) (Makarova et al. 2015).

In addition to transposons and plasmids, CRISPR-Cas systems are also scattered among bacteriophage genomes. In particular, a type I-F system, complete with an adaptation module and two CRISPR arrays, is inserted into the genomes of a group of ICP1-related phages that infect Vibrio cholerae (Seed et al. 2013). Strikingly, the majority of the spacers in the phage arrays are homologous to sequences from a host antiphage defense island, and targeting of this island by the phage-encoded CRISPR-Cas is essential for productive infection (Seed et al. 2013). Additional evidence of the presence of type I CRISPR-Cas or stand-alone CRISPR arrays in genomes of bacteriophages and archaeal viruses comes from metagenomics sequencing, and in at least some cases, the viral arrays include spacers homologous to other viral genomes (Minot et al. 2011; Garcia-Heredia et al. 2012; Bellas et al. 2015). Taken together, these findings clearly indicate that, at least sporadically, prokaryotic viruses acquire CRISPR-cas loci or CRISPR arrays alone from the hosts and deploy them as counter-defense in the virus-host arms race and/or as protection device from other viruses. This counter-defense strategy complements the dedicated anti-CRISPR proteins that are encoded by at least some and perhaps many prokaryotic viruses (Bondy-Denomy et al. 2015; Pawluk et al. 2016a, 2016b).

Finally, on multiple occasions, MGE recruit individual cas genes that either interact with the host CRISPR-Cas or are exapted for unrelated functions. The most prominent case in point is the Cas4 endonuclease that, in addition to Casposons, is encoded by numerous bacterial and archaeal viruses. In two families of Campylobacter jejuni phages, the phage-encoded Cas4-like protein affects the spacer acquisition by the host II-C CRISPR-Cas system, apparently promoting spacer capture from the host DNA (Hooton and Connerton 2015). The biological role of this effect of the phage Cas4 on adaptation is not entirely clear but the result is a stable persistence of the phage in the C. jejuni culture, so the host spacers might act as “decoys” to prevent acquisition of spacers from the phage genomes (Hooton et al. 2016). In an example of Cas4 exaptation, a virus of the archaeon Thermoproteus tenax has been shown to employ an inactivated derivative of Cas4 as the capsid protein (Krupovic et al. 2015).

An “All from the Casposon” Scenario for the Origin of CRISPR-Cas

As described in the preceding section, a large subfamily of Tn7-like transposons encode a “minimal” CRISPR-Cas system...
that is predicted to mediate crRNA processing and target recognition but not interference, and thus might facilitate the transposon targeting for integration into a new site. The incorporation of type I-F CRISPR-Cas into a Tn7-like transposon definitely was a secondary, relatively late event in evolution because it involves a specific transposon subfamily incorporating a distinct variant of CRISPR-Cas, which could occur only after the diversification of both CRISPR-Cas systems and the transposons (Peters et al. 2017). Nevertheless, the predicted involvement of CRISPR-Cas in transposition potentially might recapitulate some of the earliest stages in CRISPR-Cas history, suggesting an alternative scenario. In this version, the entire CRISPR-Cas system evolved within a Casposon such that the predecessor of the effector module initially functioned as an ancillary, RNA-guided integration mechanism or, alternatively or additionally, as a distinct defense mechanisms preventing replication of other MGE in the same host (fig. 5). Further study of the Casposon diversity can shed light on the key aspects of CRISPR-Cas origin and evolution.

Conclusions

The findings outlined in this article reveal multiple contributions of widely different classes of MGE to the origin and evolution of the CRISPR-Cas adaptive immune systems. These contributions come from two unrelated families of DNA transposons (Casposons and TnpB-encoding transposons, IS605 and ISC), retrotransposons (Group II introns) and at least two classes of TA systems. There is also substantial reverse flow of genetic information, that is, recruitment of CRISPR-Cas systems and individual cas genes by various MGE, followed by their repurposing for counter-defense or, in some cases, other functions. This multiplicity of exchanges between the immune system and MGE clearly indicates that the connection is not random but rather reflects a deep evolutionary unity that is not limited to CRISPR-Cas but involves the entirety of defense mechanisms. Indeed, simple defense systems in prokaryotes, such as TA and restriction-modification modules themselves possess properties of MGE (Kobayashi 2001; Van Melderen and Saavedra De Bast 2009; Van Melderen 2010; Furuta and Kobayashi 2011). A more complex interplay between parasitism and defense can be captured in the “guns for hire” paradigm whereby homologous proteins, such as endonucleases, are utilized as offensive and defensive “weapons,” by MGE and defense systems, respectively (Koonin and Krupovic 2015b). Recruitment of transposons or their components apparently was central not only to the evolution of CRISPR-Cas but also to the origin of adaptive immunity in vertebrates (Kapitonov and Jurka 2005; Kapitonov and Koonin 2015; Koonin and Krupovic 2015a), the system of DNA elimination and rearrangement in ciliates (Nowacki et al. 2011; Dubois et al. 2012; Bettermier and Duharcourt 2014; Allen and Nowacki 2017), and the piRNA machinery of germ line defense in animals (Aravin et al. 2007). Perhaps, there is ground for a sweeping generalization: all defense systems that are involved in some form of genome manipulation are evolutionarily linked to MGE. Elucidation of the diversity and the intricacies of the interactions between MGE and defense machineries, and development of a general theory of their coevolution are research directions for decades to come.

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