Diverse virus-encoded CRISPR-Cas systems include streamlined genome editors

Graphical abstract

Highlights

- CRISPR pathways encoded in diverse bacteriophage are hypercompact anti-viral systems
- Phage-encoded CRISPR systems encompass all known CRISPR-Cas types
- The Cas\lambda enzyme’s compact structure is capable of plant and human cell genome editing
- These findings reveal a new source of CRISPR-Cas enzymes with value as genome editors

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In brief
A tour-de-force metagenomics analysis of phage-encoded CRISPR-Cas enzymes provides a rich resource for new CRISPR-based tools.
Diverse virus-encoded CRISPR-Cas systems include streamlined genome editors

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SUMMARY

CRISPR-Cas systems are host-encoded pathways that protect microbes from viral infection using an adaptive RNA-guided mechanism. Using genome-resolved metagenomics, we find that CRISPR systems are also encoded in diverse bacteriophages, where they occur as divergent and hypercompact anti-viral systems. Bacteriophage-encoded CRISPR systems belong to all six known CRISPR-Cas types, though some lack crucial components, suggesting alternate functional roles or host complementation. We describe multiple new Cas9-like proteins and 44 families related to type V CRISPR-Cas systems, including the Casλ RNA-guided nuclease family. Among the most divergent of the new enzymes identified, Casλ recognizes double-stranded DNA using a uniquely structured CRISPR RNA (crRNA). The Casλ-RNA-DNA structure determined by cryoelectron microscopy reveals a compact bilobed architecture capable of inducing genome editing in mammalian, Arabidopsis, and hexaploid wheat cells. These findings reveal a new source of CRISPR-Cas enzymes in phages and highlight their value as genome editors in plant and human cells.

INTRODUCTION

CRISPR-Cas systems confer resistance in prokaryotes against invading extrachromosomal elements, including viruses and plasmids (Figure 1A). To generate immunological memory, microbes capture fragments of foreign genetic elements and incorporate them into their genomic CRISPR array using the Cas1-Cas2 integrase. Subsequent transcription of the array creates CRISPR RNAs (crRNAs) that bind to and direct CRISPR-associated (Cas) nucleases to target complementary nucleic acids. These systems comprise two classes, each with three different types, defined by the architectures of their nuclease effector modules involved in crRNA processing and DNA or RNA interference.

Reports of type I-F CRISPR-Cas loci encoded in bacteriophages (phage) that infect Vibrio cholerae,3,4 or type V CRISPR-Cas loci in huge phage genomes reconstructed from microbial community DNA sequences,5 hinted at a wider distribution of phage-encoded CRISPR systems that might function in novel ways and reveal important insights into prokaryotic biology. However, these were the only reported examples of phage-encoded CRISPR systems, prompting us to perform a comprehensive study of the abundance, distribution, and diversity of CRISPR-Cas systems encoded throughout the virosphere and to begin to explore the biochemical activity of novel systems.

Here, using metagenomic analysis of microbial samples isolated from soil, aquatic, human, and animal microbiomes, we report the widespread occurrence of diverse, compact
Figure 1. Diversity of CRISPR-encoding phages and the hosts they predate

(A) An illustration of the mechanism of CRISPR interference as an anti-viral system used by bacteria and bacteriophages.

(B) Protein-clustering network analysis based on the number of shared protein clusters between the CRISPR-encoding phages in this study and RefSeq phages. The plot is composed of viral clusters where each node represents a phage genome, and each edge is the hypergeometric similarity between genomes based on shared protein clusters.

(C) Genome size distribution of circularized CRISPR-containing phages from this study (n = 152).

(D) A heatmap showing the number of CRISPR phage genomes containing each CRISPR type with respect to major bacterial phyla which they infect. "Unknown" indicates CRISPR phages that could not be assigned to any of the known types. Phyla are organized in the plot from left to right based on CRISPR array abundance and are concordant with the number of bacterial genomes available for each phylum.

See also Figure S2.
CRISPR-Cas systems encoded in phage genomes, demonstrating an unexpected biological reservoir of anti-viral machinery within infectious agents. Phage-encoded CRISPR-Cas systems include members of all six CRISPR types (types I–VI) as defined by bacterially encoded examples. We found evidence for new or alternative modes of nucleic acid interference involving phage-encoded type I, III, IV, and VI systems. In addition, the phage and phage-like sequences result in a several-fold expansion of CRISPR-Cas9 and -Cas12 enzymes belonging to the type II and type V families that are widely deployed for genome editing applications. Cas1, one of the most divergent in sequence of the phage-encoded type V enzymes identified in this study, was found to have robust biochemical activity as an RNA-guided double-stranded DNA (dsDNA) cutter. Its cryoelectron microscopy (cryo-EM)-determined molecular structure explains its use of a natural single-guide RNA for DNA binding, and cell-based experiments demonstrated robust endogenous genome editing activity in plant and human cells. The compact architecture of Cas1 and other phage-encoded CRISPR-Cas proteins holds significant promise for vector-based and direct delivery into cells for wide-ranging biotechnological applications.

RESULTS

A wide diversity of phages across many bacterial phyla encode divergent CRISPR-Cas systems

Using genome-resolved metagenomics, we analyzed over 660 giga base pairs (bp) of assembled genomic DNA from both environmental and animal-associated microbiomes to reveal a surprising diversity of over 6000 CRISPR-encoding phages (Figure 1B). Our analysis of publicly available phage genomes revealed that CRISPR-Cas systems occur in only 0.4% of phages, making them exceptionally rare compared to their abundance in prokaryotic genomes where they occur in 40% of bacteria and 85% of archaea. CRISPR-encoding phages, rather than being limited to specific phylogenetic clusters, are found within many diverse phage subtypes (Figure 1B). This finding is consistent with previous work that determined phage phylogeny from protein-clustering analysis. At least two phages harboring CRISPR arrays were alternatively coded such that the TAG stop codon was recoded to glutamine. Although circularized CRISPR-encoding phages included huge phages such as a >620 kbp megaphage (Figure 1C), most had a genome size close to the average of 52 kbp. Notably, however, relatively few phages encode complete CRISPR-Cas systems. Fewer than 10% of CRISPR-encoding phages were found to contain machinery for the acquisition of new spacer sequences into their CRISPR arrays, consistent with observations in huge phages. Many phages encode CRISPR arrays, but few of these (~6%) include Cas effectors encoded nearby (Figure 1D). In such situations, phages may produce their own guide RNAs but hijack the Cas effectors provided by their hosts. Consistent with this possibility, ~1% of phages encode only the Cas1-Cas2 integrase used for the acquisition of new spacers, but no other Cas enzymes. In some cases, phage-encoded Cas1 contained a fusion to another protein such as reverse transcriptase, suggesting the possibility of the acquisition of RNA protospacers into the phage array. Notably, only 27 of the thousands of phage-encoded CRISPR-Cas loci identified in this study target RNA and can be classified as new homologs of previously described RNA-targeting systems. Thus, the vast majority of phage-encoded CRISPR systems target DNA.

Phage-encoded CRISPR-Cas systems include all six known types but with phage-specific properties

We found that all six known types of CRISPR-Cas systems occur in phages, and relative to host-encoded systems, they have various unique properties associated with their existence within phage genomes. These include missing sequence integration or targeting machinery as mentioned above, modified type III and VI systems that mitigate the abortive infection mechanism, and spacers that target other mobile genetic elements. For example, some of the rare phage-encoded type III systems are associated with CRISPR arrays targeting vital or highly abundant RNA transcripts of other mobile elements, such as phage tail proteins or transposases (Figure 2). In well-studied type III systems, the Cas10 protein converts ATP into a cyclic oligoadenylate (cOA) product, which allosterically activates an auxiliary Csm6 ribonuclease. The activated Csm6 amplifies the immune response by degrading RNA transcripts indiscriminately, thereby destroying the invasive transcriptome or inducing host cell dormancy or death, aborting the phage infectious cycle. Interestingly, in phage-encoded type III systems, the Cas10 subunit contains multiple mutations, hinting at an inability to produce cOA (Figure S1), and Csm6 or a related CARF-domain ribonuclease is absent, similarly to archaeal Borg elements. Notably, the key residues for DNA cleavage in the Cas10 HD domain, and for RNA cleavage in Cas7, remain intact (Figure S1). Unless the cOA production and Csm6 ribonuclease functionalities are complemented by orthogonal type III systems from the host genome, this suggests that the type III phage systems may be capable of cleaving key RNA transcripts and genomic DNA of competing mobile elements to interfere with their infectious cycle without activating abortive infection in which cOA signaling triggers trans-cleavage of transcripts in the host cell.

In addition to type III systems, we found the first examples of phage-encoded type VI (Cas13) ribonucleases, most of which belong to the Cas13b and the relatively small Cas13d super-families (Figure 3). Analogous to the findings above with type III systems in abortive infection, the lack of signature Csx27 and Csx28 proteins, which are transmembrane factors that enhance abortive infection mechanisms, may indicate the absence of an abortive infection pathway unless supplemented by the host.

Miniature single-effector CRISPR-Cas systems are enriched in phage genomes

Class 2 CRISPR-Cas systems, including types II, V, and VI, generally employ single-subunit RNA-guided, nucleic acid-targeting interference enzymes. In addition to new Cas9 (a, b, c) and Cas12 (a, b, c, f, i) enzyme variants, we identified miniature CRISPR-associated nucleases in phages harboring both HNH and RuvC catalytic domains characteristic of Cas9.
miniature nucleases constitute phylogenetically distinct clades denoted as types II-x, -y, and -z (Figure 3A). These systems lack the Cas1, Cas2, or Csn2 sequence acquisition machinery (Figures 1D and 3A) and have distinct domain organizations compared to previously studied Cas9 orthologs, with significant deletions across the proteins in comparison (Figure S2). The phylogenetic analyses indicated multiple evolutionary origins of the type II CRISPR-Cas systems in viruses, the evolutionary relatedness of virally and bacterially encoded CRISPR-Cas systems suggesting that those encoded by viruses were obtained from their host during prior infection.

Furthermore, we observed that bacteriophage genomes harbor an unusual enrichment of hypercompact type V effectors (Figures 1D and 3B) compared to abundance in bacteria, including hundreds of variants comprising 44 protein families that are evolutionarily distant from previously reported and experimentally validated miniature type V CRISPR-Cas nucleases, including Cas12f and CasF12,13 (Figure 3B). Evolutionary analysis suggests that distinct type V nuclease subtypes may have evolved multiple times from separate transposon-encoded TnpB families, which have recently been shown to be RNA-guided nucleases themselves,14 and we observe that TnpB is also widely encoded on phages. CRISPR arrays associated with the type V families contained spacer sequences targeting competing dsDNA-based extrachromosomal elements that are predicted to infect the same host (Figure 2). We found that in multiple related Biggiephages, miniature type V families including Casm and CasF co-occurred with a type I system that we term type I-X, of which we only had one example previously,5 bearing similarities to type I-C CRISPR systems but featuring a distinct helicase in place of the processive nuclease Cas3. Biggiephage genomes were recovered over a four-year time span, and remained identical save for their CRISPR arrays, which only exhibited minor differences (Figures S2C and S2D). Though we were unable to validate DNA cleavage by this system, it is possible that dsDNA binding silences the expression of target genes. In some cases, the arrays of the type I-X system target the same circular extrachromosomal element, albeit with distinct spacers, as the array associated with co-occurring type V systems. One such cryptic element harbored restriction enzymes and retron-based anti-phage defense systems that could limit Biggiephage infectivity, underscoring the dynamic nature of the evolutionary arms race between mobile elements in competition for host resources.
We also found the first type IV systems encoded in lytic phage genomes. Type IV systems are predominantly found on plasmids, where their mechanisms of action are poorly understood and they sometimes lack a CRISPR array. We report a type IV subtype that lacks the DinG hallmark gene and encodes in its place a CysH-like protein bearing limited similarity to non-CRISPR associated CysH phosphoadenosine 5'-phosphosulfate reductases. Remarkably, the CRISPR array associated with this type IV-F system and a neighboring type V targets the type V Cas gene encoded in a competing cyanophage (Figure 2).

**Casλ** is a divergent phage-specific CRISPR-Cas enzyme with a unique guide RNA

A distinctive phage-encoded enzyme family, Casλ, exists within huge bacteriophages that are evolutionarily linked to the recently reported Mahaphage clade. Named using Greek nomenclature to indicate its phage origin, this family of 55 compact systems exhibited such sequence divergence that it had negligible sequence identity (<5%) to, and clustered separately from, type V and type II enzymes (Figure S3A). In addition, Casλ sequences have low similarity to these enzymes (<10%) but are phylogenetically closest to Cas14J5 (Figure 3B). The protein is not encoded along with any other Cas proteins, and the RuvC nuclease was not immediately identifiable from the sequence. Difficulty in aligning this system to reported enzymes via remote homology (Figure S6) further suggested that a direct evolutionary relationship with known Cas superfamilies was questionable. CRISPR arrays associated with Casλ contain spacer sequences complementary to dsDNA-based extrachromosomal elements predicted to infect the same Bacteroidetes host (Figure 2). These observations implied that
Casλ may be targeting dsDNA in native contexts of the host similar to Cas9 or Cas12 systems.

In any CRISPR-Cas system, processing of CRISPR array transcripts, consisting of repeats and spacer sequences acquired from previously encountered mobile genetic elements (MGEs), is essential to generating mature crRNAs that guide Cas proteins to destroy foreign viruses. Analogous to the distinct nature of the protein, the Casλ crRNA is predicted to form an elongated hairpin secondary structure not previously observed in guide RNAs associated with Cas12 (Figure 4A).

Despite their divergent nucleotide sequences, crRNAs retain a similar predicted hairpin structure across the protein family (Figure S3B). Furthermore, Casλ crRNAs contain conserved sequences at their 5' and 3' ends and in the center of the RNA (Figure 4B). The overall sequence divergence of the protein, its putative RuvC domain, and the encoded crRNA prompted us to further analyze this protein family.

RuvC-mediated crRNA processing in the spacer region and dsDNA cutting by Casλ

The lack of a detectable tracrRNA encoded within the genomic locus begged the question of how this aberrant RNA, akin to a naturally occurring crRNA-tracrRNA hybrid (Figure 4A), may be processed by the CRISPR-Cas system or host factors to produce mature crRNA.

Using radiolabeled precursor crRNAs (pre-crRNAs) as substrates, we first tested whether purified Casλ protein catalyzes RNA cleavage. Surprisingly, analytical denaturing gel electrophoresis showed that pre-crRNAs are cut by Casλ in the spacer region as opposed to the 5' end of the RNA, where cutting has been observed in nearly all self-processing single-effector systems analyzed previously (Figures 4C, 4D, and S3D), with the exception of a type V-C system.18,19 The Casλ-induced pre-crRNA processing yields a crRNA spacer sequence that is complementary to DNA target sites 14–17 nucleotides (nt) in length.

The fact that Casλ can process its own pre-crRNA obviates the need for Ribonuclease III or other host factors required for the function of most known Cas9 and Cas12 family members. Although some CRISPR-Cas proteins process pre-crRNAs using
an internal active site distinct from the RuvC domain or by recruiting Ribonuclease III to cleave a pre-crRNA:tracrRNA duplex, we wondered whether phage-encoded Casλ, like phage-encoded CasΦ, processes pre-crRNA using its RuvC active site. We thus tested Mg²⁺ dependence and showed that Casλ is indeed reliant on the presence of Mg²⁺ and thus, by extension, the RuvC active site for crRNA maturation (Figure 4D).

CRISPR–Cas systems target DNA sequences following or preceding a 2–5 bp Protospacer Adjacent Motif (PAM) for self-versus-non-self discrimination. We determined the sequence requirements for DNA targeting by Casλ using a plasmid deletion assay in which targeting a library of putative PAM sites revealed sequence specificity. This assay demonstrated the ability of crRNA-guided Casλ to cleave dsDNA, without requirement for additional RNA components, and a putative TTR PAM sequence specificity (Figure S4A). Casλ with host genome-targeting guides showed a reduction in colony-forming units (as a proxy for cell viability) of multiple orders of magnitude, in comparison to negative control of Casλ with a non-targeting guide (Figure 4E).

In vitro incubation of purified Casλ with crRNAs, along with a linear dsDNA target substrate, generated cleavage products with surprisingly pronounced staggered 5’-overhangs of 11–16 nt (Figures 4F and 4G). Type V CRISPR-Cas enzymes such as Cas12a have also been observed to generate staggered overhangs, albeit smaller at only 5 nt. Furthermore, the non-target strand (NTS) was cleaved faster than the target strand (TS) within the RuvC active site over a 2 h time period (Figure 4H).

Casλ ribonucleoproteins induce genome editing in endogenous genes in human and plant cells

The development of single-effector CRISPR-Cas systems for editing eukaryotic cells has revolutionized genome engineering. However, the large sizes of Cas9 and Cas12a enzymes can inhibit delivery into many cell types, for which hypercompact genome editors with favorable kinetics imply great promise as an alternative. We conducted a head-to-head comparison of insertion and deletion efficiencies using Casλ and Cas12a ribonucleoproteins (RNPs) with identical guide RNA spacers targeting sequences recognizing Vascular Endothelial Growth Factor A (VEGFA) and Empty Spiracles Homeobox1 (EMX1) genes in HEK293T cells. Despite their miniature size, Casλ RNPs generated promising genome-editing outcomes compared to Cas12a, and in at least one case exceeded Cas12a insertion-deletion (indel) percentages (Figure 5A). Extending these experiments to Arabidopsis thaliana, we confirmed that Casλ exhibited editing efficiencies of up to 18% at the endogenous PDS3 gene (Figure 5B), notably higher than what we observed previously using CasΦ. The efficiency of editing was highly dependent on temperature, with no editing occurring at 23 °C, an intermediate level of editing occurring at 28 °C, and the highest level of editing occurring at 32 °C. Furthermore, we were able to achieve editing in the endogenous disease resistance gene Snm5 in hexaploid wheat protoplasts, where a genome ~5x larger than the human genome poses a scanning challenge to achieve successful editing at the target site (Figure 5C). Next-generation sequencing for both human and plant cells revealed indel profiles with large deletions (Figures 5D and S5C), consistent with the staggered cuts observed in vitro at the PAM distal region.

Casλ protein structure explains interference mechanism

To explore the mechanism by which Casλ achieves RNA-guided DNA recognition, we generated a purified Casλ-crRNA-dsDNA ternary complex suitable for analysis by cryo-EM. CryoEM maps of this complex revealed a bilobed architecture analogous to Cas9 and Cas12 enzymes, despite the divergence in both sequence and size between Casλ and these much larger enzymes (Figures 6A–6C, S6, and S7). The 3 Å resolution structure revealed the shape and domain organization of Casλ and the unique structure of the guide RNA (Figures 6A–6C, S6, and S7). Notably, the RuvC domain of Casλ is split into four parts across the C-terminal half of the protein, likely hindering reliable alignment and clustering with reported Cas12 systems (Figure 6D).

The REC I and REC II domains are also segmented in the protein sequence, with the PAM-interacting domain wedged between REC I as opposed to the N terminus of the protein as seen in CasΦ, but similar to Cas12i. In contrast to CasΦ, Casλ contains a Target Strand Loading (TSL) domain that likely functions to load the single-stranded DNA (ssDNA) substrate, in a position analogous to the “Nuc” domain that was incorrectly hypothesized in other type V CRISPR-Cas enzymes to be a second nuclease domain responsible for DNA cleavage. Casλ also exhibits a distinct structure in the REC I domain compared to CasΦ (Figure S6D).

The crRNA assumes an unexpected shape that blankets the protein, with a divergent recognition lobe in Casλ that binds to distinct sequences and structural features of the guide RNA (Figures 6C, and S6A–S6C). Specifically, we observed possible interactions between primarily polar or charged residues within the REC II domain in Casλ with the conserved motifs of the crRNA hairpin (Figures 4B and S6C). These residues are conserved across the protein family and likely interact either directly with the RNA nucleobases (Q452, N510), or with the RNA phosphate backbone to stabilize the guide (S451, K496, E444, N445, K503, Y619) (Figures 6C and S6C). We did not observe protein contacts to the unpaired nucleobases A9, A30, A31 motif in the middle of the guide RNA stem-loop (Figure 3A), which is further supported by the lack of sequence conservation. However, interestingly, the non-complementarity is conserved, which is likely important for the hairpin kink geometry.

CRISPR–Cas proteins initiate the unwinding of target dsDNA following PAM recognition. In Casλ, this recognition is achieved via interactions with the oligonucleotide-binding domain (OBD), REC I, and a five α-helical bundle referred to as the PAM-interacting domain (PID). Residues within the three domains interact with the sugar-phosphate backbone of the target DNA (Figure S6B) and, in some cases such as residue N102, interact directly with the nucleobases. The interaction between N102 and nucleobase G(-1) may explain the preference for purines in this position as opposed to pyrimidines, since a pyrimidine substitution would result in a base that is too distant from the interacting asparagine (Figure S6B). In examining the aftermath of cis-cleavage of DNA, we found that Casλ had a very low level of ssDNA or ssRNA cleavage in trans upon DNA recognition in cis (Figure S4B). Incubation of the Casλ protein with non-cognate guides from other orthologs within the protein family replicated the ssDNA trans cleavage effect despite differences...
in their sequence (Figure 6E), suggesting that guides within the Casλ family may be interchangeable, unlike Cas9. These findings warrant further investigation to examine the determinants of guide RNA binding by Casλ. Single mismatches across the ssDNA target revealed that the seed region of the target DNA (1–5) and the region extending from bases 7–13 are required to match the spacer sequence of the guide RNA for efficient cleavage (Figure 6F). Investigation of positions that possibly interact with the DNA in these regions (Figure S6D) or the corresponding RNA revealed conserved residues in REC, OBD, PID, and RuvC domains that may account for the complex’s intolerance to target mismatches, and, therefore, the possibility of relatively

Figure 5. Casλ RNPs are functional for editing endogenous genes in human, Arabidopsis, and wheat cells with large deletion profiles

(A) Indel efficiency using Casλ and Cas12a RNPs with identical spacers targeting VEGF, and Casλ RNPs targeting EMX1 genes in HEK293T cells, and a schematic of the in vitro model of DNA cleavage outcomes following DNA cleavage by Casλ.

(B and C) Indel efficiencies in Arabidopsis thaliana protoplasts show significantly higher levels of editing than previously achieved by Casλ for the same PDS3 gene, and (C) in wheat protoplasts targeting the disease resistance gene Snn5.

(D) Indel profiles generated by Casλ RNP administration show primarily large deletions, and little change without Casλ.
high fidelity in the context of genome editing. Overall, more domains within Casλ, such as the RECI, RECI, OBD, RuvC, and TSL domains, exhibit segmentation and rearrangement compared to known type V systems.

Figure 6. Structure of Casλ-gRNA-DNA complex
(A) Schematic representation of the Casλ-gRNA-DNA complex. Disordered linkers are shown as dotted lines. Insets for protein-DNA interactions are shown in Figure S5.
(B) Cryo-EM maps of the Casλ-guide-RNA-DNA complex. The target strand is shown in cyan and the non-target strand is shown in magenta.
(C) Cryo-EM-based model of guide RNA-target DNA complex.
(D) Schematic of the domain organization and secondary structure of Casλ.
(E) Hierarchical clustering dendrogram of different repeats with their predicted secondary structures. Casλ can still cleave ssDNA in trans with guide RNAs consisting of non-cognate repeats that are divergent at the sequence level.
(F) Fluorescence output using oligonucleotide activators with mismatches at each respective position along the target DNA. “0” indicates no mismatches (control). Insets relating to protein-DNA interactions are shown in Figure S6.
See also Figures S5, S6, and S7.

DISCUSSION
CRISPR-Cas systems occur rarely in bacteriophage genomes, but large-scale analysis of all known and novel phages
uncovered thousands of viruses encoding an unexpected diversity, distribution, and potential function of RNA-guided proteins across the biosphere. CRISPR-encoding bacteriophages are predicted to infect a remarkable diversity of bacterial phyla, and the spacer sequences associated with their CRISPR arrays show complementarity to dsDNA-based viruses and extrachromosomal elements predicted to infect the same hosts. These observations underscore the general strategy of phages to protect their hosts against superinfection by competing elements. We find that phage genomes harbor CRISPR-Cas systems corresponding to all six known types, and we also note differences observed bioinformatically or biochemically between the viral CRISPR-Cas systems and their bacterial counterparts. These differences hint that phages have co-opted CRISPR anti-viral systems with modifications that tune RNA-guided pathways to their needs.

At least two aspects of phage-encoded CRISPR systems differ notably from cellular systems, highlighting the versatility of these pathways and also the potential for phage-mediated functional evolution. First, some RNA-targeting type III and VI systems recognize abundant or essential transcripts of competing phages, and type III systems retain the catalytic residues required to cleave ssDNA and enable nicking of the target DNA, but lack components used for non-specific transcript cleavage following RNA target recognition. The absence of these components, which trigger abortive infection by analogous host-encoded systems, suggests that some phages prefer to avoid self-destruction of transcripts or induction of a dormant state in the host, both of which may be disadvantageous to the phage life cycle. Consistent with this idea, the minimal trans- cutting of ssDNA and RNA observed for Casλ implies limited ability to target single-stranded replication intermediates of MGEs (Figures S4 and S8). A second important difference between phage- and phage-encoded CRISPR systems is the absence of a processive nuclease, such as Cas3, in some phage type I systems. Together with the presence of CysH, which may be recruited as a putative effector in type IV systems, these observations suggest alternate outcomes to nucleic acid-targeting by these phage-encoded pathways. In particular, the lack of a Cas3 nuclease in type II systems targeting plasmid-like elements suggests a gene silencing mechanism that precludes DNA cutting. We speculate that such phage-based type I systems could assist the activity of the co-occurring Casϕ systems that are found in the same genome. Because the targeted plasmid-like elements harbor restriction enzymes and retron-based anti-phage defense systems that could limit the infectivity of the CRISPR-encoding phage, coordinated activities of orthogonal CRISPR systems could assist competition between mobile elements.

This study demonstrates that phage genomes are a natural reservoir of miniature single-effector CRISPR-Cas systems, including DNA targeting type II and type V enzymes belonging to the Cas9 and Cas12 superfamilies. We use Greek nomenclature to indicate the phage origins of Casϕ, Casω, and Casλ, extending the naming convention established by phage-encoded Casϕ. In contrast to the prevalence of multi-subunit type I and type III CRISPR systems in prokaryotic genomes, the notable abundance of miniature Cas12-family enzymes in phages may reflect the size restriction of many phage genomes. Because phages evolve quickly, they serve as important sources of new, divergent, or hypercompact CRISPR systems. Some of these, such as Casλ, bear sufficient sequence-level divergence to cluster separately from Cas12 and Cas9 systems and obscure a direct evolutionary relationship with known Cas superfamilies. Nonetheless, Casλ’s structure, domain composition, and biochemical mechanism are similar to other type V enzymes. This finding implies that within phage genomes, distinct type V nucleases may have evolved multiple times from ancestral transposon-encoded TnpB families, which also function as RNA-guided nucleases. Despite being from different clades of phages and having divergent sequences and domain organizations, we observe a convergent evolution of Cas12-like architecture in the Casλ and Casϕ protein structures. In addition, both can process their own pre-crRNA and rely on the same RuvC active site used for DNA cleavage for this activity. This extreme compression of enzymatic activities within one active site has not been observed for bacterially encoded CRISPR-Cas proteins. Nonetheless, the phage-encoded enzymes diverge functionally from one another in other ways, including guide RNA structure and maturation process. This may reflect the interplay between rapid phage evolution, which generates diversity, and selective pressure to maintain CRISPR competency in a variety of host environments over time, which favors pathway conservation. In both cases, phages that encode their own Cas variants that do not rely on host factors may eliminate the possibility that the ongoing evolution of essential host proteins or cofactors will result in incompatibility with phage-encoded anti-viral systems.

The molecular structure of the Casλ-crRNA-dsDNA complex reported in this study illustrates possible convergent evolution of RNA-guided effectors, despite extreme sequence divergence and distinct ancestral protein origins. The domain architecture of Casλ exhibits more segmentation and likely structural rearrangements than have been seen in other Cas12-family enzymes, with multiple functional domains split at the sequence level into separate segments that assemble during protein folding. This unique domain organization may explain the difficulty in accurately aligning Casλ to previously reported enzymes, despite overall structural similarity. Remarkably, however, this segmented domain composition does not compromise genome editing activity as observed for Casλ-based editing of human, Arabidopsis, and wheat cells. The finding that Casλ can induce efficient genome editing of endogenous genes in these diverse cell types, in some cases exceeding the efficacy of Cas12a-mediated genome editing, shows that there isn’t necessarily a tradeoff between Cas effector size and function. This result, together with the compact size of phage-encoded CRISPR-Cas proteins that is advantageous for vector-based cellular delivery (Figure S7), shows that nature’s phage reservoir is an important future source of enzymes useful for genome editing in heterologous cell types.

Overall, the discovery of thousands of viruses encoding CRISPR systems representing all six CRISPR-Cas types highlights the sparsity but broad diversity of RNA-guided systems within viruses. Genome-resolved metagenomics and bioinformatics-enabled phylogenetic insights enabled us to analyze these systems from uncultivated viruses and infer their mechanisms of action within their biological contexts. We investigated...
hundreds of novel hypercompact and divergent CRISPR-Cas systems, with particular focus on the unique Casλ family. We report the utility of Casλ as a valuable tool for genome editing in plant and human cells. In addition, our data show how the structural compaction of this protein family preserves robust biochemical and cell-based functionality essential to both natural activities and biotechnological applications. Finally, our structural understanding of Casλ proteins provides a starting point for the future design of variants with expanded genome editing capabilities that combine the advantages of a small protein with the versatility of a robust RNA-guided DNA recognition machine.

**Limitations of the study**

Some mechanistic aspects of Casλ remain to be defined, including the possible interchangeability of different variant guide RNAs to support catalytic activity. In addition, details of target binding, including the seed region within the target binding site as well as target recognition accuracy, remain to be established. Potential interactions between phage-encoded and host-encoded CRISPR systems have not been explored and will be fascinating to understand. Future research is needed to identify the functions of phage-encoded CRISPR systems that may extend beyond anti-phage defense and to further test the applications of Casλ as a genome editing tool in plants, other organisms, and human cells.

**STAR★METHODS**

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  - Plant genome editing
  - Ternary complex reconstitution for cryo-EM
  - Electron microscopy grid preparation and data collection
  - Single-particle cryo-EM data processing and 3D volume reconstruction

**SUPPLEMENTAL INFORMATION**

Supplemental information can be found online at https://doi.org/10.1016/j.cell.2022.10.020.

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**AUTHOR CONTRIBUTIONS**

B.A.-S. conceived of and designed the study with guidance from J.F.B. and J.A.D. B.A.-S. and J.F.B. designed and implemented the computational searches, host matching, phylogenetic analyses, and classification of phages and CRISPR-Cas systems. B.A.-S. performed PAM and cell viability assays, radiolabeling, and DNA cleavage biochemical work and analyzed the data, with guidance from P.P. and B.F.C. C.J.H. performed the crRNA processing assay, B.A.-S. and D.S. performed protein purifications and trans-cleavage assays, B.A.-S., P.S., and K.M.S. prepared cryoEM grids and collected cryoEM dataset. B.A.-S. generated the initial protein model. P.S. and B.A.-S. processed cryoEM data with guidance from K.M.S. B.A.-S. designed guide RNAs and purified protein for eukaryotic genome editing experiments that were performed by Z.L., E.C.S., E.G., and S.E.J. with oversight from S.E.J., D.F.S. and B.S. B.A.-S. and A.R.E. performed phage-type I-X analyses. J.A.D. and J.F.B. provided advisory support and funding. J.A.D. and B.A.-S. wrote the manuscript with input from all authors.

**DECLARATION OF INTERESTS**

The Regents of the University of California have filed patents related to this work on which the authors are inventors. J.A.D. is a co-founder of Caribou Biosciences, Editas Medicine, Scribe Therapeutics, Intellia Therapeutics, and Mammmbio Biosciences. J.A.D. is a scientific advisory board member of Vertex, Caribou Biosciences, Intellia Therapeutics, Scribe Therapeutics, Mammmbio Biosciences, Algen Biotechnologies, Felix Biosciences, The Column Group, and Inari Agriculture. J.A.D. is Chief Science Advisor to Sixth Street, a Director at Johnson & Johnson, and Altos and Tempus, and has research projects sponsored by Apple Tree Partners and Roche. J.F.B. is a co-founder of Meta-genomi. S.E.J. is a scientific co-founder of Inari Agriculture, and a science advisory board member to Invaio Sciences, Sendia Biosciences, FL70, and Sixth Street.

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## STAR+METHODS

### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Bacterial and virus strains** | | |
| *E. coli* BL21(DE3) | New England Biolabs | C2527H |
| *E. coli* BL21(DE3)-Star | QB3-Macrolab, UC Berkeley | N/A |
| *E. coli* 5-alpha Competent cells | New England Biolabs | C2987H |
| **Chemicals, peptides, and recombinant proteins** | | |
| 384 well flat bottom black polystyrene assay plate | Corning | 3820 |
| 2xYT media | Sigma-Aldrich | Y2377 |
| Acrylamide/Bis 19:1, 40% solution | Fisher | 187,328 |
| Agarose | Omnipur | 2125 |
| Alkaline Hydrolysis Buffer | Invitrogen | AM2283 |
| Alt-R Cas12a (Cpf1) Electroporation Enhancer | IDT | 1,076,301 |
| ATP, [γ-32P]- 3000 Ci/mmol | Perkin Elmer | BLU002A001MC |
| DNase Alert | IDT | 11-02-01-04 |
| Dulbecco’s Modified Eagle Medium (DMEM) | Corning | 10-017-CV |
| EDTA | Fisher Chemical | S311500 |
| fetal bovine serum (FBS) | VWR | 1500-500 |
| Glycerol | Fisher bioreagents | BP229 |
| HEPES | Sigma | PHG0001 |
| Imidazole | Sigma-Aldrich | IX0005 |
| isopropyl-β-D-thiogalactopyranoside (IPTG) | Goldbio | I-902 |
| Luria-Bertani Broth (LB) | USBiological Life Sciences | L1505 |
| Magnesium chloride | Sigma-Aldrich | M2670 |
| penicillin-streptomycin | Gibco | 10,378,016 |
| Potassium chloride | Sigma-Aldrich | P9541 |
| Q5 High-Fidelity DNA Polymerase | NEB | M0491 |
| Quick Extract | Lucigen Corporation | QE09050 |
| Recovery Medium | Lucigen Corporation | NC9996840 |
| RNase Alert | IDT | 11-02-01-02 |
| SF nucleofection buffer | Lonza | S50-642 |
| SOC media | NEB | B9020 |
| Sodium chloride | JT Baker | 4058 |
| T4 PNK | NEB | M0236S |
| Terrific Broth (TB) | VWR | 97,063–420 |
| Tris (2-carboxyethyl) phosphine hydrochloride (TCEP) | Sigma-Aldrich | C4706 |
| UREA | Sigma | u5378 |

### Critical commercial assays

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| DNA Clean and Concentrator kit | Zymo Research | D4033 |
| HiSpeed Plasmid Maxi Kit | QIAGEN | 12,663 |
| QIAprep Spin Miniprep kit | QIAGEN | Cat#27106 |

### Deposited data

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Cryo-EM density map of the Casλ-crRNA-dsDNA ternary complex | This paper | EMDB: EMD-27320 |
| Coordinates of the Casλ-crRNA-dsDNA ternary complex | This paper | PDB: 8DC2 |

(Continued on next page)
### REAGENT or RESOURCE SOURCE IDENTIFIER

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Continuation**    |        |            |

| Experimental models: Cell lines | | |
|---|---|---|
| HEK293T | University of California Berkeley Cell Culture Facility | N/A |

| Oligonucleotides | | |
|---|---|---|
| rBAS80 ssRNA oligo | AUUGUUGUAAACUUAUUUU GUAUGGAGUAACACUACCAUCACC | This paper N/A |
| dBAS608 ssDNA oligo | CTTGAACGTTTGTACTGATG TGAAGTGGGAGATAGTAATGTAATG | This paper N/A |
| dBAS609 ssDNA oligo | CATAACATTACTAAGAGGG TGAAGGATGTGCTAAAAACGGGTCAAG | This paper N/A |

| Recombinant DNA | | |
|---|---|---|
| Cas12L1 Non-Targeting array and Cas12L1 ORF | This paper | pBAS12 |
| pBAS12 with GFP targeting spacer | This paper | pBAS18 |
| L1_orf L2 guide 19 targeting GFP | This paper | pBAS40 |
| L1_orf L1 guide non-targeting GFP | This paper | pBAS41 |
| L1_orf L1 guide non-targeting GFP | This paper | pBAS42 |
| L1_orf L1 guide non-targeting GFP | This paper | pBAS43 |
| L1_orf L1 guide 19 targeting GFP | This paper | pBAS44 |

| Software and algorithms | | |
|---|---|---|
| AlphaFold2 | Jumper et al. | https://github.com/deepmind/alphafold |
| ChimeraX version 1.2.5 | Pettersen et al. | https://www.cgl.ucsf.edu/chimerax/download.html |
| Coot version 0.9.4.1 | Emsley et al. | https://www2.mrc-lmb.cam.ac.uk/personal/pemsley/coot/ |
| CRISPResso2 | Clement et al. | https://github.com/pinellolab/CRISPResso2 |
| cryoSPARC version 3.2.0 | Punjani et al. | https://cryosparc.com/download |
| ImageQuant TL 8.1 | Cytiva Life Sciences | https://www.cytivafsciences.com/en/us/shop/molecular-biology/nucleic-acid-electrophoresis blotting-and-detection/molecular-imaging-for nucleic-acids/imagequant-tl-10-2-analysis software-p-28619 |
| IQTREE version 1.6.6 | Nguyen et al. | https://github.com/Cibiv/IQ-TREE/releases/tag/v1.6.6 |
| LocSpiral | Kaur et al. | https://github.com/1aviervargas/LocSpiral LocBSharpen-LocBFactor-LocOccupancy |
| MAFFT version 7 | Katoh and Standley | https://mafft.cbrc.jp/alignment/software/ |
| MinCED | Bland et al. | https://github.com/ctSkennerton/minced |
| Mtriage | Afonine et al. | https://phenix-online.org/documentation/reference/mtriage.html |
| Phenix version 1.19.2–4158 | Liebschner et al. | https://phenix-online.org/download/ |
| SerialEM version 3.8.7 | Mastronarde | https://bio3d.colorado.edu/SeriaEM/download.html |
| Topaz | Bepler et al. | https://github.com/tbepler/topaz#installation |
| ViennaRNA | Lorenz et al. | https://github.com/ViennaRNA/ViennaRNA |

| Other | | |
|---|---|---|
| 1.2/1.3 300 mesh UltrAuFoil gold grids | Mitigen | Q350AR13A |
| 10 kDa MWCO concentrator | Amicon Ultra, Merck | UFC9010 |
| 4D-nucleofector | Lonza | N/A |
| Amersham Typhoon phosphorimager | GE Healthcare | N/A |
| Cytation 5 plate reader | BioTek | N/A |

(Continued on next page)
RESOURCE AVAILABILITY

Lead contact
Requests for additional information and reagents should be directed to the lead contact, Jennifer Doudna (doudna@berkeley.edu).

Materials availability
Reagents generated in this study are available, upon reasonable request, from the lead contact with a Materials Transfer Agreement.

Data and code availability
- Structural data were deposited in EMDB and PDB with accession numbers reported in the Key Resources Table.
- This paper does not report original code. Analyses were performed as previously described in Al-Shayeb et al. 2020.
- Any additional information required to reanalyze the data reported in this work is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Mammalian models
Mammalian gene-editing experiments were performed in HEK293T cells obtained from University of California Berkeley Cell Culture Facility. HEK293T cells were female in origin and grown in DMEM media (Corning) containing 10% fetal bovine serum (VWR) and 100U/mL of penicillin-streptomycin (Gibco) at 37°C with 5% CO₂.

Plant models
PDS3 gene-editing was tested in A. thaliana protoplasts isolated from the leaves of 4-week-old plants. Following RNP screening experiments, protoplasts were incubated in W5 solution (4 mM MES pH 5.7, 0.5 M mannitol, 20 mM KCl) at RT for 12 h, then moved to 37°C for 2.5 h, followed by a final incubation at room temperature for 48 h.

Additional experiments in wheat (Triticum aestivum, L cv. Fielder) were performed using protoplasts extracted from leaves of 2-week-old plantlets. Edited cells were incubated in darkness with W5 solution at 30°C for 24 h.

METHOD DETAILS

Phylogenetic analysis
For analysis of publicly available phage genomes, we analyzed Genbank-recorded phages, complete RefSeq-recorded phages, IMG-VR-recorded phages. Cas protein sequences and representatives from the TnpB superfamily were collected from literature.2,5,42–45 The resulting set was clustered at 90% amino acid identity to reduce redundancy. A new alignment of Casλ with the resulting sequence set was generated using MAFFT with 1000 iterations and filtered to remove columns composed of gaps in 95% of sequences. The phylogenetic tree was inferred using IQTREE v1.6.6 using automatic model selection and 1000 bootstraps.

crRNA sequence analysis
CRISPR-RNA (crRNA) repeats from Phage-encoded CRISPR loci were identified using MinCED (github.com/ctSkennerton/minced). The repeats were compared by generating pairwise similarity scores using the Needleman-Wunsch algorithm. A heatmap was built using the similarity score matrix and hierarchical clustering produced dendrograms that were overlaid onto the heatmap to delineate different clusters of repeats. The RNA structures were predicted with ViennaRNA.41

PAM depletion analysis
PAM depletion assays were performed with plasmids containing the casλ protein coding sequence as derived from metagenomics and a mini CRISPR targeting guide (pBAS18), or with plasmids that contained only the casλ gene and a non-targeting guide (pBAS12). Assays were performed as three individual biological replicates. Plasmids containing casλ and mini CRISPRs were transformed into E. coli BL21(DE3) (NEB). Subsequently, electrocompetent cells were prepared by ice-cold H₂O and 10% glycerol washing. A plasmid
library was constructed with 8 randomized nucleotides upstream of the (5') end of the target sequence. Competent cells were transformed in triplicate by electroporation with 200 ng library plasmids (0.1 mm electroporation cuvettes (Bio-Rad) on a Micropulser electro- porator (Bio-Rad)). After a 2 h recovery period, cells were plated on selective media and colony forming units were determined to ensure appropriate coverage of all possible combinations of the randomized 5’ PAM region. Strains were grown at 25°C for 48 h on media containing appropriate antibiotics (either 100 μg/mL carbenicillin and 34 μg/mL chloramphenicol, or 100 μg/mL carbenicillin and 50 μg/mL kanamycin) and 0.05 mM isopropyl-β-D-thiogalactopyranoside (IPTG), or 200 nM anhydrotetracycline (aTc), depending on the vector to ensure propagation of plasmids and Casλ effector production. Subsequently, propagated plasmids were isolated using a QiAprep Spin Miniprep Kit (Qiagen).

PAM depletion sequencing analysis

Amplion sequencing of the targeted plasmid was used to identify PAM motifs that are preferentially depleted. Sequencing reads were mapped to the respective plasmids and PAM randomized regions were extracted. The abundance of each possible 8 nucleotide combination was counted from the aligned reads and normalized to the total reads for each sample. Enriched PAMs were computed by calculating the log ratio compared to the abundance in the control plasmids, and were used to produce sequence logos.

Programmable DNA targeting

A flip recombination assay was performed in E. coli to eliminate the Kanamycin resistance cassette from E. coli strains that contain GFP and RFP expression cassettes integrated into the genome. Individual colonies of the E. coli ΔKan were picked to inoculate three 5 mL (LB) starter cultures to prepare electrocompetent cells the following day. 100 mL (LB) main cultures were inoculated from the starter cultures and grown vigorously shaking at 37°C to an OD600 of 0.6–0.7 before preparation of electrocompetent cells by repeated ice-cold H2O and 10% glycerol washes. Cells were resuspended in 10% glycerol and 50 μL aliquots were flash frozen in liquid nitrogen and stored at −80°C. Casλ vectors were generated containing codon optimized cas:1 gene and a guide comprised of its cognate repeat element and selections of spacers targeting the GFP DNA within the resulting E. coli ΔKan strain (pBAS41, pBAS42, pBAS43, pBAS44) were subcloned from pBAS12. Casλ vectors containing Casλ1 and a guide composed of a non-cognate repeat unit from cas2/2 and a GFP-targeting spacer (TAGCATCACCTTCACCCTCTCCAGGACAG) guide were also subcloned to form pBAS40. The Casλ vectors and Casλ vectors with a non-targeting guide control plasmid were transformed into 25 μL of electro- competent cells with 100 ng of plasmid via electroporation in 0.1 mm electroporation cuvettes (Bio-Rad) on a Micropulser electro- porator (Bio-Rad), cells were recovered in 1 mL recovery medium (Lucigen) shaking at 37°C for 1 h 10-fold dilution series were then prepared and 3.5 μL of the respective dilutions were spot-plated on LB-Agar containing the appropriate antibiotics and IPTG inducer. Plates were incubated overnight at 37°C and colonies were counted the following day to determine the transformation efficiency. To assess the transformation efficiency, the mean and standard deviations were calculated from the cell forming units per ng transformed plasmids for the electroporation triplicates. The experiment showed marked reduction of GFP E. coli using Casλ vectors with their cognate guides (pBAS44) in comparison to the non-targeting control, indicating a dsDNA break at the target region. The growth of primarily RFP-positive/GFP-negative colonies under blue light further supports the ability to confer targeted programmable genome editing to result in strains lacking GFP production. Growth inhibition using Casλ vectors with guides from a separate Casλ ortholog (pBAS40), with observed expressing primarily RFP and no GFP, also indicate that Casλ orthologs may function using guides from related CRISPR-Cas systems to confer editing in cells, with a precise ablation of GFP production. This can be further expanded to HEK293T mammalian cells with integrated GFP, which indicate activity in mammalian cells. The sickly phenotype of E. coli colonies that have grown in both cases even in undiluted samples is also indicative of possible trans-cleavage of nucleic acids (RNA or DNA), which can be used for diagnostic purposes by providing a sample containing the target nucleic acid with the Casλ RNP and a ssDNA fluorophore-quencher (ssDNA-FQ) reporter or RNA fluorophore-quencher (ssRNA-FQ) reporter molecule, generating a strong fluorescence signal in the presence of the target nucleic acid compared to a markedly lower fluorescence signal in its absence.

Protein purification

Casλ overexpression vectors containing a His-Tag were transformed into chemically competent E. coli BL21(DE3)-Star (QB3-Macrolab, UC Berkeley) and incubated overnight at 37°C on LB-Kan agar plates (50 μg/mL Kanamycin). Single colonies were picked to inoculate 50 mL (LB, Kanamycin 50 μg/mL) starter cultures which were incubated at 37°C shaking vigorously overnight. The following day, 1.5 L TB-Kan media (50 μg/mL Kanamycin) were inoculated with 40 mL starter culture and grown at 37°C to an OD600 of 0.6, cooled down on ice, and gene expression was subsequently induced with 0.5 mM IPTG followed by incubation overnight at 16°C. The cells were harvested by centrifugation and resuspended in wash buffer (50 mM HEPES-Na pH 7.5 RT, 500 mM NaCl, 20 mM Imidazole, 5% glycerol and 0.5 mM TCEP), and then subsequently lysed by sonication. The soluble fraction was loaded on a 5 mL Ni-NTA Superflow Cartridge (Qiagen) which had been pre-equilibrated in the same wash buffer. Bound proteins were washed with 20 column volumes (CV) wash buffer and subsequently eluted in 5 CV elution buffer (50 mM HEPES-Na pH 7.5 RT, 500 mM NaCl, 5% glycerol, and 0.5 mM TCEP). The eluted proteins were concentrated to 1 mL before injection into a HiLoad 16/600 Superdex 200pg column (GE Healthcare) pre-equilibrated in size-exclusion chromatography buffer (20 mM HEPES-Na pH 7.5 RT, 500 mM NaCl, 5% glycerol, and 0.5 mM TCEP). Peak fractions were concentrated to 1 mL and concentrations were determined using a NanoDrop 8000 Spectrophotometer (Thermo Scientific). Proteins were purified at a constant temperature of
4°C and concentrated proteins were kept on ice to prevent aggregation, snap-frozen in liquid nitrogen, and stored at –80°C. SDS-PAGE gel electrophoresis of Casλ at varying stages of protein purification showed a protein size in line with computationally predicted values of 85 kDa.

**Pre-crRNA processing assays**

The reactions were carried out in RNA cleavage buffer containing 20 mM Tris-Cl (pH 7.5 at 37°C), 150 mM KCl, 5 mM MgCl₂, 1 mM TCEP, and 5% (v/v) glycerol. Pre-crRNA substrates were 5’S-radiolabeled with T4 PNK (NEB) in the presence of γ-32P-ATP. In a typical pre-crRNA processing reaction, the concentrations of Casλ and γ-32P-labeled pre-crRNA substrates were 100 and 3 nM, respectively. Reactions were incubated at 37°C, and an aliquot of each reaction was quenched with 2x Quench Buffer (94% (v/v) formamide, 30 mM EDTA, 400 μg/mL heparin, 0.2% SDS, and 0.025% (w/v) bromophenol blue) at 0, 1, 3, 5, 15, 30, and 60 min. RNA hydrolysis ladders were prepared by incubating RNA probes in 1X RNA Alkaline Hydrolysis Buffer (Invitrogen) at 95°C before the addition of 2x Quench Buffer. Quenched reactions were incubated at 95°C for 3 min, and products were then resolved by denaturing PAGE (10% or 20% acrylamide:bis-acrylamide 19:1, 7 M urea, 1X TBE). Gels were dried (3 h, 80°C) on a Model 583 Gel Dryer (Bio-Rad) and exposed to a phosphor screen. Phosphor screens were imaged on an Amersham Typhoon phosphorimager (GE Healthcare). For assays in an EDTA-containing buffer, 25 mM EDTA was substituted for 5 mM MgCl₂.

**In vitro cleavage assays - Radiolabeled nucleic acids**

crRNA oligonucleotides were synthesized by IDT and dissolved in DEPC-treated ddH₂O to a concentration of 0.5 mM. Subsequently, the crRNA was heated to 65°C for 3 min and allowed to cool down to room temperature. Casλ RNP complexes were reconstituted at a concentration of 10 μM by incubation of 10 μM Casλ and 12 μM crRNA for 10 min at RT in 2x cleavage buffer (20 mM Hepes-Na pH 7.5, 300 mM KCl, 10 mM MgCl₂, 20% glycerol, 1 mM TCEP). RNPs were aliquoted to a volume of 10 μL flash-frozen in liquid nitrogen, and stored at –80°C. RNPs aliquots were thawed on ice before experimental use. Substrates were 5’S-end-labelled using T4-PNK (NEB) in the presence of γ-32P-ATP. Oligonucleotide-duplex targets were generated by combining γ-32P-labeled and unlabelled complementary oligonucleotides in a 1:1.5 M ratio. Oligos were hybridized to a DNA-duplex concentration of 50 nM in hybridization buffer (10 mM Hepes-Na pH 7.5 RT, 150 mM NaCl), by heating for 5 min to 95°C and a slow cool down to RT in a heating block. Cleavage reactions were initiated by combining 200 nM RNP with 2 nM substrate in CB buffer and subsequently incubated at 37°C. Reactions were stopped by the addition of two volumes of formamide loading buffer (96% formamide, 100 μg/mL bromophenol blue, 50 μg/mL xylene cyanol, 10 mM EDTA, 50 μg/mL heparin), heated to 95°C for 5 min, and cooled down on ice before separation on a 12.5% denaturing urea-PAGE. Gels were dried for 4 h at 80°C before phospho-imaging visualization using an Amersham Typhoon scanner, v2.0.0.6 firmware version 208 (GE Healthcare). Bands were quantified using ImageQuant TL 8.1 (Cytiva) and the cleaved fraction was calculated as the product intensity sum divided by the combined substrate and product intensity sum. Curves were fitted to a One-Phase-Decay model to derive the rate of cleavage.

**Fluorophore quencher and DNA mismatch tolerance assay**

DNA oligo activators were ordered from IDT to contain mismatches at each respective position, (A->C, T->G, C->A, G->T). Casλ RNPs were prepared as described above. Reactions were started by combining 100 nM RNP (100 nM Casλ, 120 nM crRNA), 100 nM DNase Alert (IDT) FQ probe, with and without activator ssDNA and with the addition of a non-targeting guide or activator control in 1X RNA Alkaline Hydrolysis Buffer (Invitrogen) at 95°C before phosphor-imaging visualization using an Amersham Typhoon scanner, v2.0.0.6 firmware version 208 (GE Healthcare). Bands were quantified using ImageQuant TL 8.1 (Cytiva) and the cleaved fraction was calculated as the product intensity sum divided by the combined substrate and product intensity sum. Curves were fitted to a One-Phase-Decay model to derive the rate of cleavage.

**I-X PAM binding assay**

The PAM binding assay was conducted using NEB 5-alpha Competent *E. coli* cells. Plasmids containing the type I-X system included a targeting or non-targeting guide downstream of T7 promoters. PAM library plasmids contained sfGFP under the control of an araBAD promoter. Downstream of the promoter was a six-nucleotide variable region of potential PAM sequences, resulting in loss of sfGFP fluorescence for a successful PAM binding event. All cultures used 2xYT media and were supplemented with kanamycin and ampicillin as needed for plasmid maintenance. Cell densities were maintained at greater than 100x library coverage throughout the assay.

Transformations of type I-X systems with guide and library plasmids were conducted consecutively. Type I-X systems with guides were transformed into NEB 5-alpha Competent *E. coli* cells following manufacturer’s instructions. Individual colonies were incubated at 37°C overnight at 250 RPM. Non-transformed cultures were included for library only and no plasmid controls. Cells were back diluted 1000x and cultured to 0.6 OD, pelleted, washed 3 times with sterile water, and resuspended in 10% glycerol to make them electrocompetent. Type I-X systems with guide, and non-transformed cultures were electroporated at 1800V with 100 ng of PAM library stock and recovered for 1 h in SOC media. Recovered cells were plated with appropriate antibiotics and incubated overnight. Plates were scraped, resuspended, and incubated at 37°C 250 RPM for 3 h. 25% glycerol stocks were stored at –80°C.
In preparation for fluorescence-activated cell sorting, stocks were back-diluted 100x and cultured to ABS$_{300}$ \( \sim 1 \) OD. To induce proteins and guides, cultures were back diluted 1000x, supplemented with 0.5 mM IPTG and 1% arabinose. Strains were cultured overnight at 30°C until targeting and non-targeting strains reached an ABS$_{300}$ \( \sim 1 \) OD. Prior to sorting on the Sony SH800 Cell Sorter, cells were pelleted and resuspended in PBS to 0.6 OD. The non-targeting strain was used to set forward scatter (Gate 1) and singlets (Gate 2) gates and to detect any decrease in fluorescence in the targeting strain. For the targeting strain, at least 270,000 events were sorted for the lowest \( \sim 0.13 \% \) of the fluorescent cell population (Gate 3) and 500,000 events for the next \( \sim 0.13 \% \) to 1.23% lowest fluorescent cells (Gate 4). Sorted cells were grown overnight on plates containing appropriate antibiotics along with non-targeting, library only, and no plasmid controls.

Targeting, non-targeting, and library only strains were individually prepared for next generation sequencing by first purifying plasmid DNA using a Qiagen HiSpeed Plasmid Maxi Kit. Plates were gently scraped and resuspended in \( \sim 50 \) mL 2xYT prior to pelleting. Concentrations were determined with a Nanodrop. In conjunction with the original naive PAM library stock control, PAM sequences were amplified using primers containing the 5′ stub sequence GCTCTTCGATCT. Samples were submitted to the Innovative Genomics Core for completion of library preparation and iSeq sequencing at greater than 100x library coverage.

**Mammalian genome editing**

RNPs were formed in the SF nucleofection buffer (Lonza) with 100pmol protein & 120pmol crRNA at 10μM concentration for 10’ at RT. 78 pmol (1μL) of IDT Cas12a electroporation enhancer was then added. HEK293T cells (University of California Berkeley Cell Culture Facility) were added in a 10μL SF nucleofection buffer at 200,000 cells per nucleofection. 21μL reactions were loaded into cuvettes and electroporated with pulse code DS-150 in a 4D-nucleofector (Lonza). Nucleofections were performed in triplicate for each guide RNA tested. Cells were grown in duplicate in DMEM media (Corning) containing 10% fetal bovine serum (VWR) and 100U/mL of penicillin-streptomycin (Gibco) from each nucleofection in 24-well plates at 37°C with 5% CO$_2$. gDNA was collected after 72 h in Quick Extract (Lucigen) by heating at 65°C for 20 min followed by 95°C for 20 min. PCR1 was performed followed by bead clean-up to remove primers and submitted for PCR2, bead clean-up, and iSeq (illumina) at the IGIC Center for Translational Genomics. Approximately 20,000 reads per sample (2 x 150bp) were analyzed for genome editing using CRISPResso2 (https://crispresso.pinellolab.partners.org/login).

**Plant genome editing**

Guides were designed to target the PDS3 gene in *A. thaliana* protoplasts, incubated with protein as described for *in vitro* assays, and 26μL of 4μM RNP was transferred onto Arabidopsis protoplasts as previously described.\(^{48}\) Leaves were cut into 0.5mm strips perpendicular to the leaf midrib in 0.6 M mannitol solution, then placed in freshly exposed to 6 h low light (\( \sim 100 \) μE m⁻² s⁻¹). Protoplasts from 2-week-old plantlets were isolated from leaf tissue as previously described.\(^{46}\) Leaves were cut into 0.5mm strips perpendicular to the leaf midrib in 0.6 M mannitol solution, then placed in freshly prepared enzyme solution (20 mM MES pH 5.7, 0.6 M mannitol, 10 mM KCl, 1.5% cellulase R10). Leaf strips in solution were vacuum infiltrated for 30 min in darkness and then incubated for 6 h shaking at 70 rpm. After the incubation, the enzyme/protoplast solution was placed with equal volume of W5 solution (2 mM MES pH 5.7, 154 mM NaCl, 125 mM CaCl$_2$, 5 mM KCl) and filtered through 40 μm cell strainers. Protoplasts were spun down at 80g for 3 min, then resuspended in 15mL W5 solution and left to aggregate in ice for 60 min. Supernatant was removed and protoplasts were resuspended in MMG solution (4mM MES pH 5.7, 0.4 M mannitol, 15mM MgCl$_2$ at 2.5x10$^5$ cells/mL. Cas$_{9\lambda}$ RNP complexes were reconstituted with 6 μM Cas$_{9\lambda}$ protein, purified as described, and 10 μM guideRNA assembled in RNP reconstitution buffer (20mM Hepes-Na pH 7.5, 300 mM KCl, 10mM MgCl$_2$, 20% glycerol, 1mM TCEP) and incubated for 20 min at 37°C. 25 μL of 6 μM assembled RNP were added to a 1.5 mL tube, then mixed with 200 μL protoplasts. After flicking to mix, 220 μL PEG-CaCl$_2$ solution (40% PEG 4000, 0.2 M mannitol, 100 mM CaCl$_2$) was added to tubes, and samples were mixed thoroughly by slowly inverting the Cryo-EM tube until streaks of PEG were no longer visible. Samples were incubated for 15 min in darkness at RT, then 880 μL of W5 solution was added and mixed by inverting. Protoplasts were harvested by centrifugation at 80g for 3 min, resuspended in 1.2 mL W5 solution (4 mM MES pH 5.7, 0.5 M mannitol, 20 mM KCl), and plated into 12-well plates. Plate edges were sealed with parafilm and cells were incubated for 24 h in darkness at 30°C. At the end of the incubation period, protoplasts were collected in 1.5 mL tubes, pelleted at 8000 g, and flash frozen. gDNA was extracted from protoplasts using 2X CTAB\(^{47}\) and suspended in 50 μL DEPC-treated H$_2$O. 150–200 bp amplicons for amplicon sequencing were obtained using 30 cycles of Q5 High Fidelity DNA Polymerase, cleaned using the Zymo DNA Clean and Concentrator kit, then subsequently amplified using 15 cycles of PrimeSTAR GXL DNA Polymerase to add the requisite indices for sequencing. Amplicons were sequenced via paired-end 150 bp amplicon sequencing using an Illumina iSeq 100.

**Ternary complex reconstitution for cryo-EM**

Cas$_{9\lambda}$ was produced as described above. crRNA (rBAS80) was ordered as a synthetic RNA oligonucleotide from IDT and dissolved in DEPC-treated ddH$_2$O to a concentration of 0.5mM. Subsequently, the crRNA was heated to 65°C for 3 min and cooled down to RT to allow for hairpin formation. DNA oligonucleotides (dBAS608, dBAS609) were designed to contain a non complementary protospacer segment to produce ‘bubbled’ substrates and facilitate rapid R-loop formation during ternary complex reconstitution. Oligonucleotides were ordered from and synthesized by IDT. DNA oligonucleotides were combined in a 1:1.2 M ratio (target strand: non-target strand) and annealed to form a DNA duplex in hybridization buffer (10mM Heps-Na pH 7.5 RT, 150mM NaCl) by heating for 5 min at 95°C and a subsequent slow cool down in a thermocycler.
Prior to reconstitution, thawed Casλ protein was incubated with crRNA in a 1:1.1 ratio for 10 min at room temperature, and the DNA duplex was added. The ternary complex was reconstituted with the final Casλ: crRNA: TS: NTS strands stoichiometry of 1:1.1:1.2:1.4, for another 10 min at RT, and further injected into a Superdex 200 prep grade 10/300 column (GE Healthcare) pre-equilibrated in low salt buffer (10mM Hepes-Na pH 7.5, 150mM NaCl) at 4°C to separate complexes from excess nucleic acids. Peak fractions were pooled and concentrated down to ~20 μM with a centrifugal filter device (Millipore 10 kDa Mw cutoff), as measured by absorbance at 260 nm with a NanoDrop 8000 Spectrophotometer (Thermo Scientific), and kept on ice before plunge-freezing.

**Electron microscopy grid preparation and data collection**

The resulting sample was frozen using FEI Vitrobot Mark IV, cooled to 8°C at 100% humidity. 1.2/1.3 300 mesh UltrAuFoil gold grids (Electron Microscopy Sciences #Q350AR13A), were glow discharged at 15 mA for 25 s using PELCO easyGLOW. Total volume of 4 μL sample was applied to the grid and immediately blotted for 5 s with a blot force of 8 units. Micrographs were collected on a Talos Arctica operated at 200 kV and ×36,000 magnification (1.115 Å pixel size), in the super-resolution setting of K3 Direct Electron Detector. Cryo-EM data was collected using SerialEM v.3.8.7 software. Images were obtained in a series of exposures generated by the microscope stage and beam shifts.

**Single-particle cryo-EM data processing and 3D volume reconstruction**

In total, 2795 movies were collected with a defocus range of −0.8 to −2.2 μm and a 20° tilt. Data processing was further performed in cryoSPARC v3.2.0. Movies were corrected for beam-induced motion using patch motion, and CTF parameters were calculated using patch CTF. Two rounds of Topaz training were applied to the data to enrich the amounts of Casλ ternary complex particles picked as follows. In the first round, as a result of initial curation, a subset of 562 micrographs with seemingly best ice quality and CTF fit were selected. Further, 3931 particles were manually picked and submitted to Topaz particle training. The resulting Topaz model was used to pick particles from the micrographs, and a total of 153,537 particles were extracted with bin factor 2, and applied to 2D classification. Following the selection of the best classes, 113,638 particles were used for *ab initio* reconstruction with three classes. The 55,587 particles constituting the best class in terms of resolution and resemblance to an RNP were subject to non-uniform map refinement, and an initial complex map was obtained. In the second round, the latter particles were used to train a new Topaz model. Following the second round of curation, a total of 1931 micrographs were selected, and the new Topaz model was applied to pick and extract the particles. In total, 884,595 particles were subject to a round of heterogeneous refinement. Particles constituting the best class in terms of resolution were subject to the remove duplicates procedure, and further to non-uniform map refinement. As a result, a 2.99 Å map reconstructed from a total of 369,389 particles was obtained. Half-maps from this refinement were used to generate the final LocSpiral map with improved weaker density regions.

**Model building and refinement**

The initial model of the Casλ protein was obtained with the AlphaFold2 program. The predicted model was split into two parts (eventually constituting REC and Nuc lobes), and each was docked independently into the map with the fitmap tool in UCSF ChimeraX v1.2.5. The dsDNA and crRNA models were built de novo. The combined ternary complex model was refined using the real-space refinement and rigid body fit tools in Coot v0.9.4. Finally, the model was subject to a round of real_space_refine tool in Phenix v1.19.2–4158, using secondary structure, Ramachandran, and rotamer restraints.

**Data deposition and figure preparation**

Cryo-EM maps and model coordinates were deposited to the EMDB (code EMD-27320) and PDB (code 8DC2). The structure figures were generated in UCSF ChimeraX v1.2.5. Cryo-EM map FSC levels were calculated as: map level/root-mean-square deviation from zero. The orientation distribution plots were either obtained from CryoSPARC or generated using pyem csparc2star.py and star2bild.py programs. Map versus model Fourier shell correlation (FSC) graphs were calculated in Mtriage, as implemented in Phenix. Gold standard FSC plot was generated in cryoSPARC.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

Computational analyses and figures were prepared as described in Method Details. Where applicable, the SEM was graphed to depict variability across replicates.
**Figure S1. Sequence similarity of phage-encoded CRISPR-Cas systems, related to Figure 3**

(A) Alignment of Cas10 effectors from Huge Phages with those sourced from bacterial genomes. Phage type III Cas10’s are predicted to cleave DNA via Cas10 HD nuclease, but lack the residues required for the Palm domain to generate cyclic oligonucleotide signaling molecules.

(B) Alignment of Cas7 proteins from phages with those sourced from bacterial genomes. Phage type III Cas7’s have conserved motifs that are predicted to cleave RNA.
Figure S2. Structure of phage-encoded Cas9-like systems and comparison of type I-X CRISPR arrays, related to Figures 1 and 2

(A) Predicted domain organization for hypercompact phage-encoded Cas9-like systems.

(B) Predicted models for Cas9-like phage-encoded systems.

(C) Comparison of type I-X and Cas9-encoded Biggiephages recovered across a four-year time frame using Mauve, with the CRISPR repeat locations denoted in blue. Identical sequences at the nucleotide level are shown in green, with differences shown in brown or red.

(D) Phage type I-X CRISPR arrays from metagenomes sampled from the same site over the span of four years show remarkably stable arrays.
Figure S3. Divergent properties of Casλ, related to Figure 4
(A) Casλ remote homolog searches across public databases led to poor hits and no similarity to known CRISPR-Cas proteins, where only poor hits (green-black) were observed in one RuvC motif.
(B) Comparison of crRNA repeat similarity across orthologs.
(C) Comparison of protein similarity across orthologs.
(D) A time-series experiment incubating Casλ with 5' radiolabeled crRNAs with the product run on a 20% Urea PAGE gel supports the finding that Casλ uniquely processes its own single crRNA in the spacer region (or 3' end).
Figure S4. Casλ PAM specificity and comparison with other Cas ortholog trans-cleavage and indel profiles, related to Figure 4
(A) The most depleted 5' PAMs resulting from the PAM depletion assay, indicating DNA recognition and cleavage preferences for Casλ1. (B) DNase alert trans-cleavage assay with the same molarities of Cas12a, Casλ, and CasΦ targeting the same ssDNA activator. (C) Casλ indel profile in HEK293T cells compared to AsCas12a. Guide 107 targets the antisense strand, while guide 109 targets the sense strand of VEGFa.
Figure S5. Cryo-EM workflow, related to Figure 6

(A) Map generation pipeline in cryoSPARC.

(B–D) Representative 2D class averages of the final set of particles, (C) the corresponding 3D maps resulting from \textit{ab initio} reconstruction, and further (D) from heterogeneous refinement.

(E) Local resolution map as calculated in cryoSPARC v3.3.

(F) Orientation distribution of the final set of refined particles.

(G and H) gold standard, and (H) map versus model FSC curves of the model refined to the LocSpiral map and plotted with the final cryoSPARC sharp experimental map.
Figure S6. Structure of Casλ ternary complex, related to Figure 6

(A) Cryo-EM maps of the Casλ-guide RNA-DNA complex in two 90°-rotated orientations.
(B) Cartoon representation of the Casλ-gRNA-DNA complex. Insets highlight residues N102, S253, N254 predicted to be responsible for PAM recognition. Hydrogen bonds are shown as dashed lines.
(C) Model of guide RNA-target DNA complex, with insets highlighting residues conserved across the protein family that are predicted to be interacting with the RNA.
(D) Close-up views of the residues predicted to be responsible for recognition of the seed and low mismatch tolerance regions observed in (Figure 6F).
(E) Direct comparison of Casλ and CasΦ (PDB-ID: 7LYS) with a dashed bubble highlighting the Casλ TSL domain. Differences in RecI (Blue) can also be observed between the two proteins.
Figure S7. Structural comparison of Cas12 orthologs, related to Figure 6
Structural comparison of all DNA-targeting Cas12’s in order of increasing RNP size: Cas6 (7LYS), CasX (6NY3), Cas12i (6W5C), Cas12a (5XUS), Cas12b (5WTI), Cas12f (7C7L).
Figure S8. Trans-cleavage assay, related to Figure S4

(A–F) Trans-cleavage assays conducting with RNase Alert reporter substrate at decreasing RNP concentrations (A–C) for binary and ternary complexes of Casλ, and with (D) PolyU RNA reporter substrates, and testing cell viability assays with cells expressing Casλ in conjunction with (E) targeting and (F) non-targeting guides.