



**Genome Editing
From Zn finger to CRISPR/CAS9**



NOBELPRISET I KEMI 2020 THE NOBEL PRIZE IN CHEMISTRY 2020



THE ROYAL SWEDISH ACADEMY OF SCIENCES



Photo: Halbauer&Foretti



Emmanuelle Charpentier

Photo: UC Berkeley/Doudna Lab



Jennifer A. Doudna

”för utveckling av en metod för genomeditering”

“for the development of a method for genome editing”

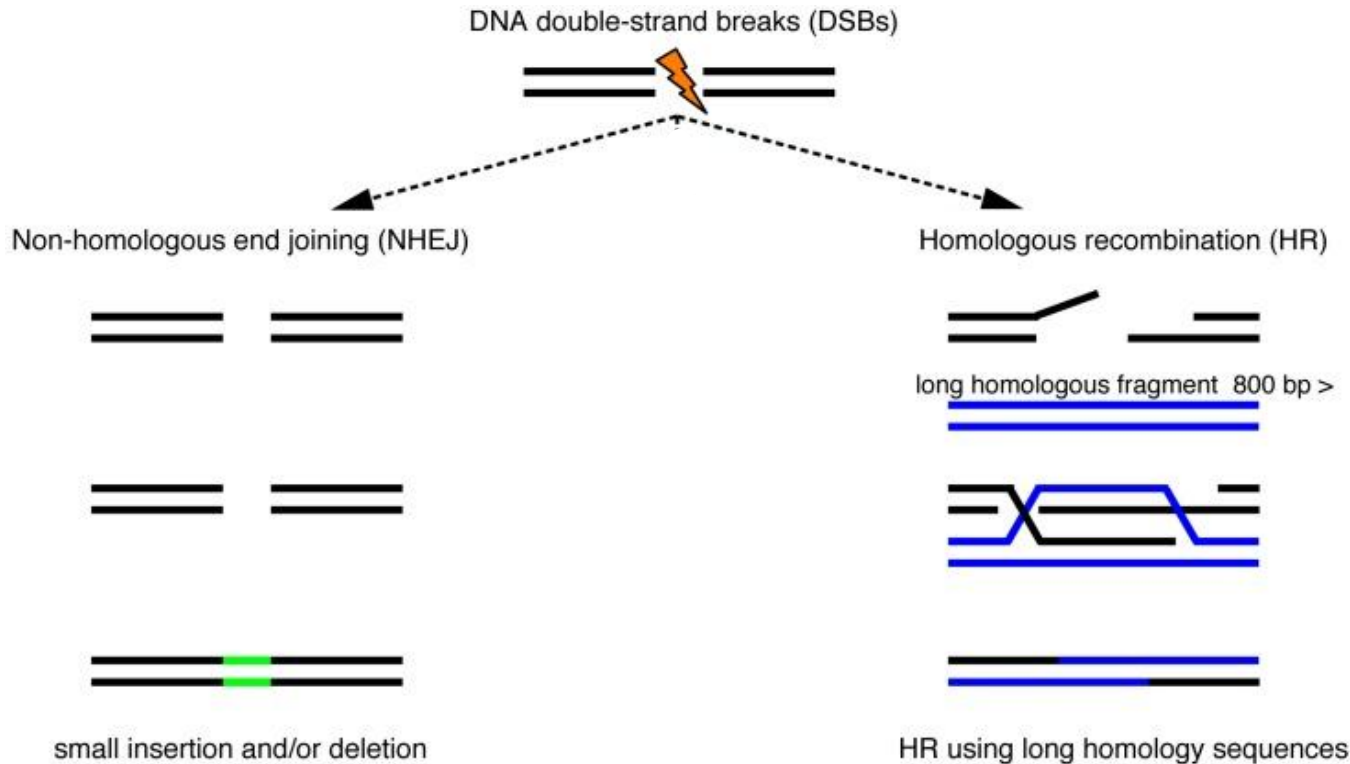
#nobelprize

Registrato con Debut Home Edition. www.nchsoftware.com/capture/it



Targeted genome modification (TGM)

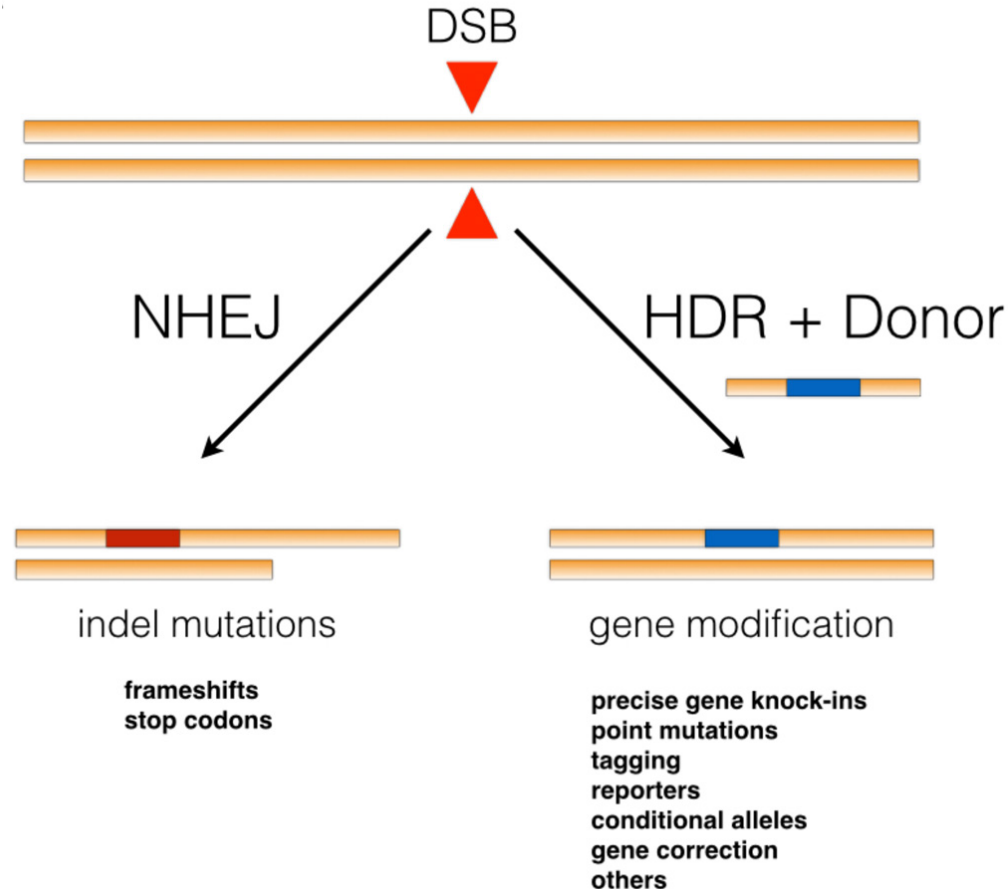
The deployment of engineered nucleases (Zinc finger nucleases, TALENs and the CRISPR/cas system) enabled a significant breakthrough in targeted genome editing and offers an unprecedented way of modifying the genome of organisms in an efficient and cost-effective way.



ZFNs: artificial nucleases that consist of a **synthetic ZFN** domain fused to a **Fok I cleavage domain**

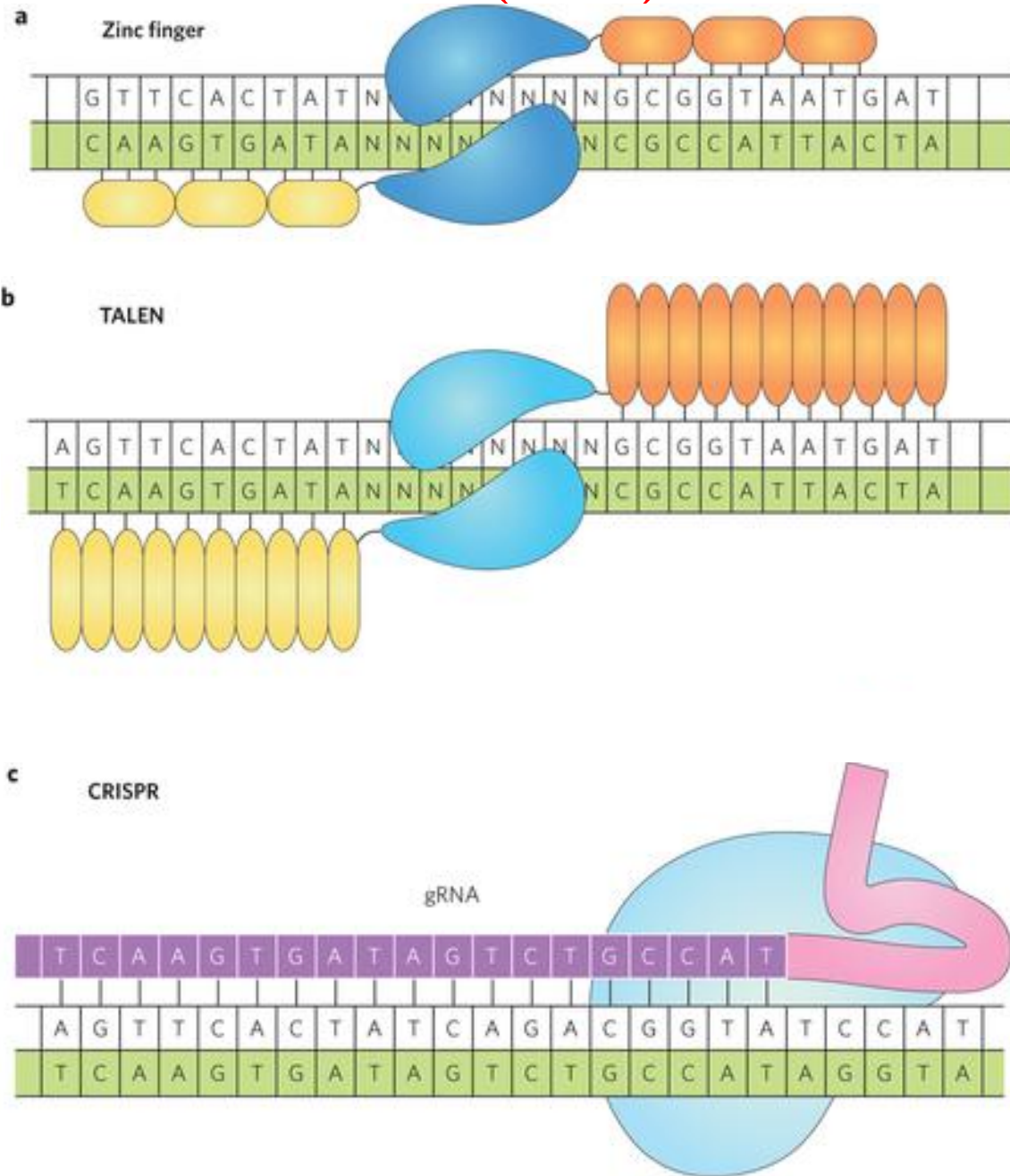
TALENs: transcription activator-like effector nucleases, consisting of an **engineered specific** (TALE) DNA binding domain and a **Fok I cleavage domain**

A crucial first step for performing targeted genome editing is **the creation of a DNA double-stranded break (DSB) at the genomic locus to be modified**. Nuclease-induced DSBs can be repaired by: **Non-Homologous End-Joining (NHEJ)** and **Homology-Directed Repair (HDR)**. **NHEJ can lead to the efficient introduction of insertion/deletion mutations (indels) of various lengths. HDR-mediated repair can be used to introduce specific point mutations or to insert desired sequences through recombination of the target locus with exogenously supplied DNA ‘donor templates’.** With targeted nuclease-induced DSBs, the frequencies of these alterations are typically **greater than 1% and, in some cases, over 50%.**



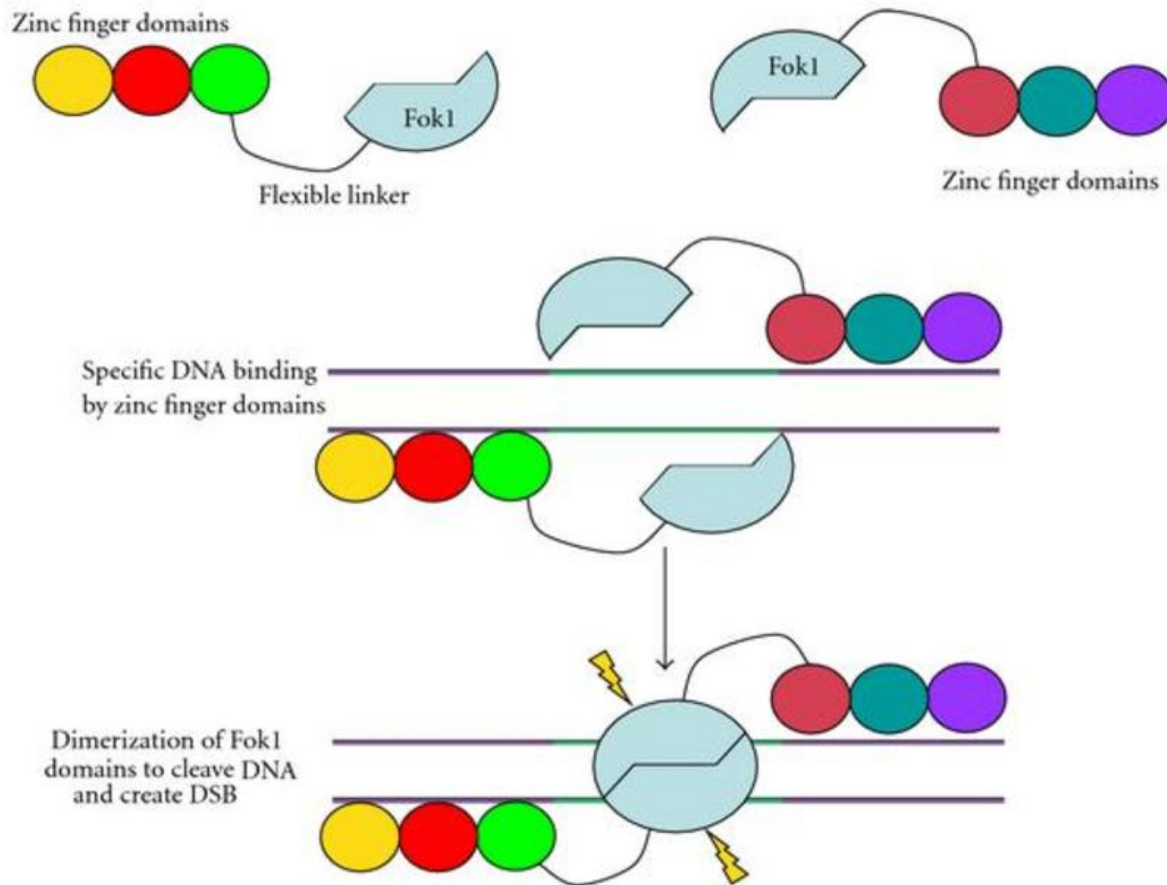
Targeted genome modification (TGM)

Site-directed nucleases. a–c, DNA nucleases bind to and cut DNA at specific locations. Each nuclease comprises a DNA-cutting domain (depicted in blue) and a DNA-targeting domain. **Zinc finger nucleases** and **Transcription Activator-Like Effector Nucleases** (TALENs) possess **protein-based DNA recognition domains** (depicted by yellow and orange ovals). **Clustered Regularly Interspaced Short Palindromic Repeats (CRISPR) nucleases** (c) rely on short-guide RNAs (gRNA) to locate the target DNA. The DNA recognition domain of all three nucleases can be engineered to target pre-determined sites in the genome for the purposes of genome editing.



The enzyme **FokI**, naturally found in *Flavobacterium okeanoikoites*, is a bacterial restriction endonuclease consisting of an N-terminal DNA-binding domain and a **non-specific DNA cleavage domain at the C-terminal**.

FokI domain requires dimerization to function.



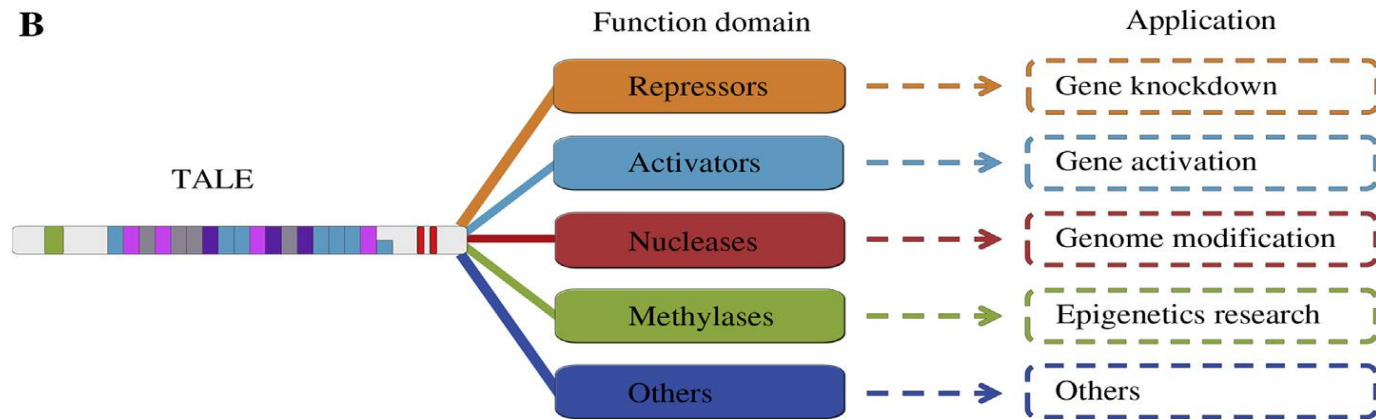
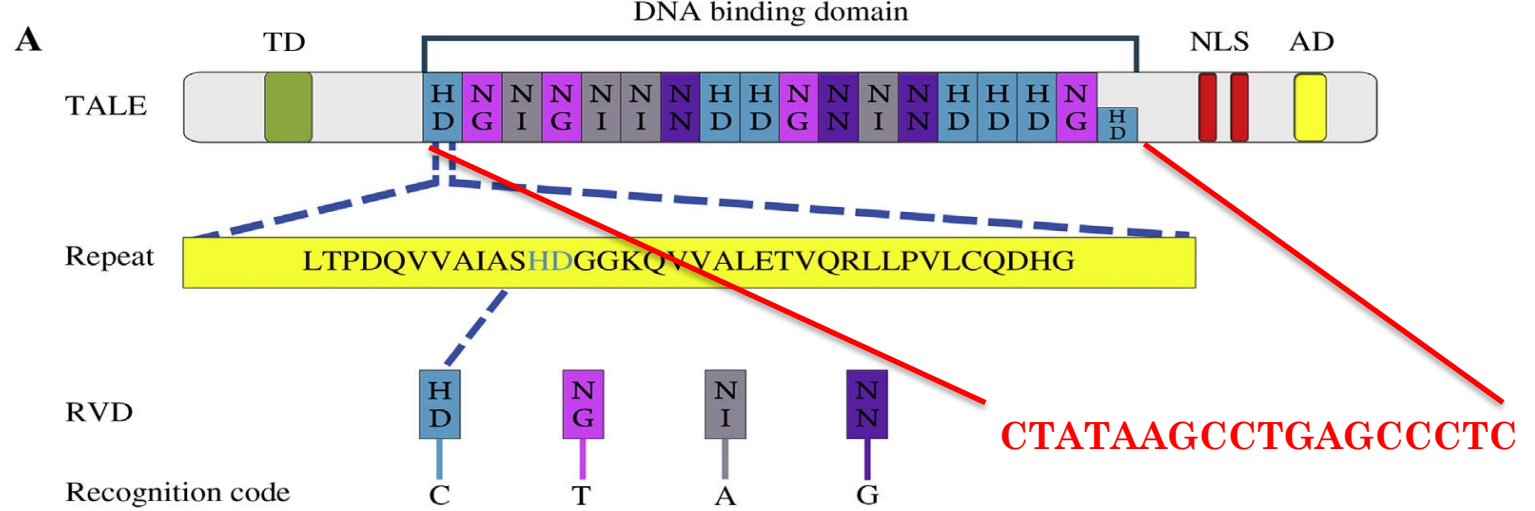
The customizable **TALE DNA binding domain**, composed of several nearly identical tandem repeat arrays, can target any given sequence according to a simple **repeat variable di-residue (RVD)**-nucleotide recognition code

- TALEs are major **virulence factors secreted by the plant pathogenic bacterial genus *Xanthomonas***. They are injected into host cells through the secretion system and interfere with cellular activities by activating the transcription of specific target genes

- They have specific structural features, including secretion and translocation signals in the N-terminal region, nuclear localization signals (**NLS**) and an acidic transcription-activation domain (**AD**) in the C-terminal region and a central DNA binding domain (**DBD**) with **33-35 nearly identical long amino acid repeats**, followed by the last module which contains only 20 amino acids (“half repeat”)

- The **Repeat Variable Di-residue (RVD)** at positions 12 and 13 of each repeat **dictates the specificity of repeat binding to a nucleotide in the DNA target**

- The DNA binding specificity of a TALE is determined by its repeat number and the sequence of the RVD: the repeat number determines the length of the target sequence, while the RVD corresponds directly to the nucleotide in the target site.



A: TALE structure and DNA recognition code: a typical TALE structure (top) comprises an N-terminal translocation domain (TD), a central DNA binding domain (DBD), two nuclear localization signals (NLS) and a transcriptional activation domain (AD) in the C-terminal region. The DBD is composed of several tandem repeats which end with a half repeat. **Each repeat consists of 34 nearly identical amino acids except for the central repeat variable di-residue (RVD) in positions 12 and 13.** The one-to-one RVD-nucleotide code of the four common RVDs used for DNA specific targeting in TALENs is shown below. **B:** TALE proteins: an optimal TALE scaffold including the central DBD can be fused to some functional domain, such as repressor, activator, nuclease or methylase, to generate TALE-based designer proteins for site-specific modification of the genome.

1987 a Japanese group identifies **clustered regularly interspaced palindromic repeats** (CRISPRs) as a series of short direct repeats interspaced with short sequences in the genome of *Escherichia coli*

CRISPRs were later detected in numerous bacteria and archaea, and predictions were made about their possible roles in DNA repair or gene regulation

```

TGAAAATGGGAGGGAGTTCTACCGCAGAGGCGGGGAACTCCAAGTGATATCCATCATCGCATCCAGTGCGCC (1,451)
(1,452) CGGTTTATCCCCGCTGATGCGGGGAACACCAGCGTCAGGCGTGAAATTCACCGTCGTTGC (1,512)
(1,513) CGGTTTATCCCTGCTGGCGCGGGGAACTCTCGGTTCAAGCGTTGCAAACCTGGCTACCGGG (1,573)
(1,574) CGGTTTATCCCCGCTAACGCGGGGAACTCGTAGTCCATCATCCACCTATGTCTGAACTCC (1,634)
(1,635) CGGTTTATCCCCGCTGGCGCGGGGAACTCG (1,664)

consensus: CGGTTTATCCCCGCTGGAACGCGGGGAACTC

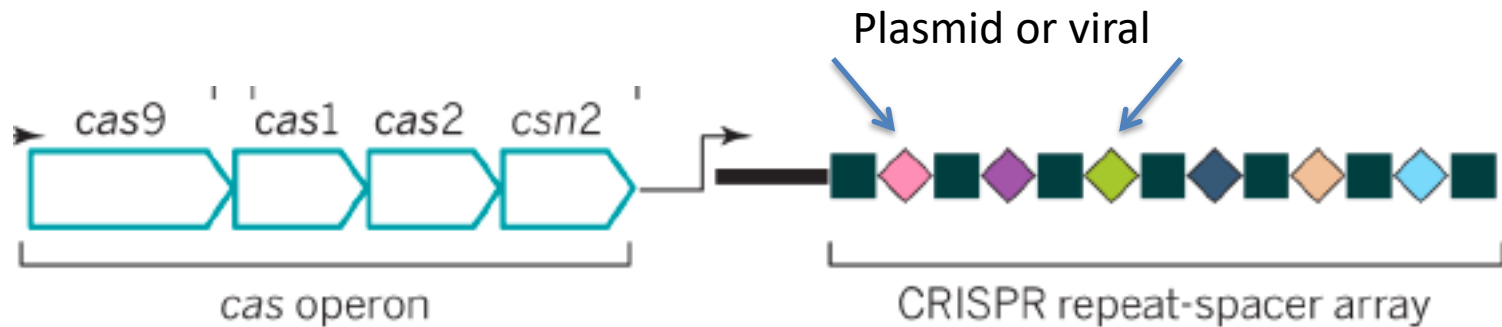
```

FIG. 5. Comparison of direct-repeat sequences consisting of 61 base pairs in the 3'-end flanking region of *iap*. The 29 highly conserved nucleotides, which contain a dyad symmetry of 14 base pairs (underlined), are shown at the bottom. Homologous nucleotides found in at least two DNA segments are shown in boldface type. The second translational termination codon is boxed. The nucleotide numbers are in parentheses.



2005: sequence analyses indicate that many spacer sequences within CRISPRs derive from **plasmid and viral origins**.

CAS (CRISP-Associated) genes:



Identification of 86 candidates for small non-messenger RNAs from the archaeon *Archaeoglobus fulgidus*

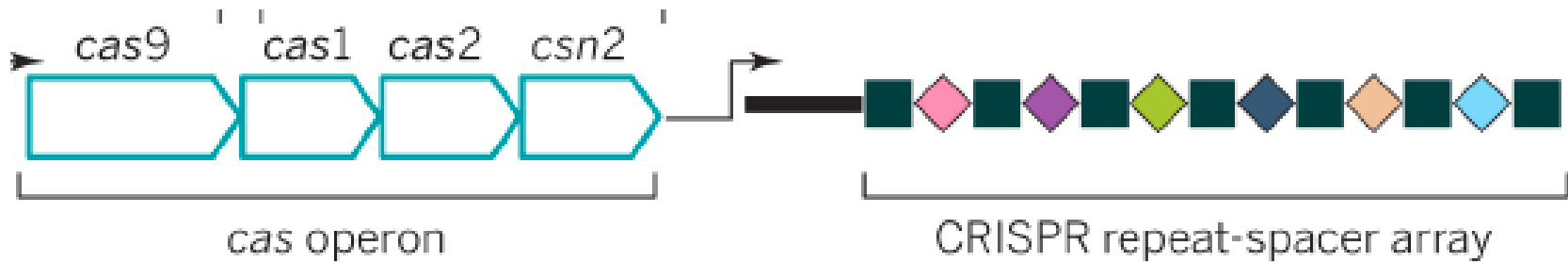
Thean-Hock Tang*, Jean-Pierre Bachelierie†‡, Timofey Rozhdestvensky*, Marie-Line Bortolin†, Mario Drungowski¶, Thorsten Elge||, Jürgen Brosius*, and Alexander Hüttenhofer**

*Institute of Experimental Pathology, Von-Esmarch-Strasse 56, 48149 Münster, Germany; †Laboratoire de Biologie Moléculaire de la Recherche Scientifique, Université Paul-Sabatier, 31062 Toulouse, France; ‡Lehrstuhl für Mikrobiologie, Universität F 93053 Regensburg, Germany; ¶Max-Planck-Institut für Molekulare Genetik, Harnackstrasse 23, 14195 Berlin, Germany; an Primärdatenbank Deutsches Ressourcenzentrum für Genomforschung GmbH, Heubnerweg 6, 14059 Berlin, Germany

2002

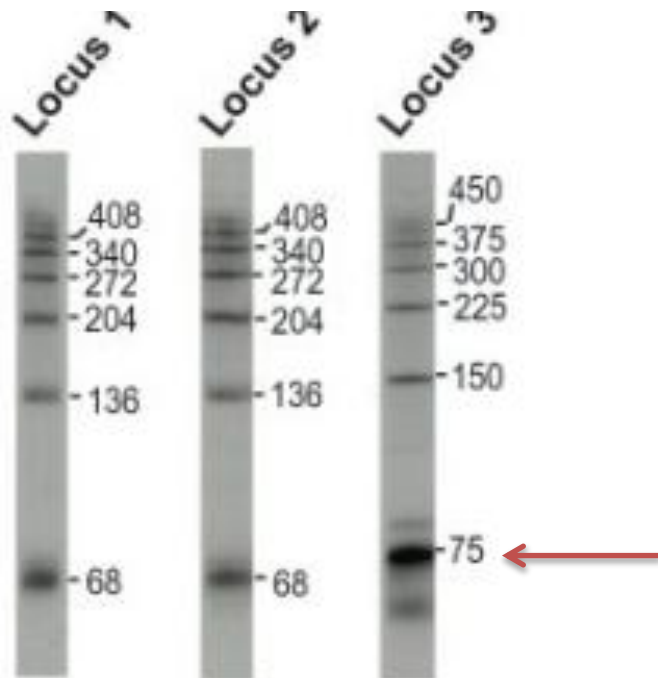
Goal: Identify small non-mRNA in the thermophilic sulfur-metabolizing *Archaeoglobus fulgidus*

Methodology: Select small RNA (from ≈50 to ≈500 nt) and prepare a cDNA library from *Archaeoglobus fulgidus* -> sequencing and confirmation of expression by northern blotting -> **they find CRISP repeats!**



TRANSCRIBED!!!

Northern blot analysis of repeated sequences. For each locus, an oligonucleotide derived from the respective repeat motif was used as a probe



mature CRISPR RNAs (crRNAs)

Cas (CRISPR-associated)

genes encode proteins with **putative nuclease and helicase domains**

CRISPR-Cas is an adaptive defense system that might use antisense RNAs as **memory signatures of past invasions**. Alignment of the amino acid sequences of putative Cas proteins 1–4 from published genomes. The number indicates the last amino acid residue of the alignment. Only matching residues of identical or chemically related amino acid residues are given. **The boxed regions in the Cas3 homologues are characteristic of helicases.**

Cas1 homologues

A. pernix	..R-----NA.L.G.S.LY...L...L.P.LG.H...SL.LDA.E.FR..IVD...L...	270
A. aeolicus	..R-----NA.I.G.S.Y...I...L.P.WGYLH...R.SL.LDV.E.FR..VD...LI...	237
A. fulgidus (Af187)	..R-----NA.L.G.S.S.L...V...L.P.AGFLN...R.SL.LDE.E.FR..VVD...LI...	271
A. fulgidus (Af2435)	..R-----NA.I.G.S.LY...I...L.P.ISYLN...R.SL.LDI.E.FR..VVD...LV...	245
B. halodurans	..R-----NA.L.G.S.LY...L...L.P.WGF.H...R.SL.LDE.E.R...D...LI...	266
C. jejuni	..R-----N.L.G...V...L.P.WG.H...L.DE.E.FR...VD...L...	239
E. coli	..R-----N.I...S.LY...I...P.ISFVN...S...DE.D...VV...I...	238
E. coli O157	..R-----N.I...S.LY...I...P.ISFIN...S...DE.D...VV...I...	238
M. thermoautotrophicum	..R-----NA.I.G.S.LY...L...L.P.ISYLN...R.SL.LDE.E.FR..LID...LI...	258
M. jannaschii	..R-----NA.I...S.LY...L...L.P.WGYLH...R.SL.LDE.E.FR..I.D...LV...	246
M. tuberculosis	..R-----NS.V.G.S.LY...I...L...IGFLN...L...DE.E.WK...IID...LI...	252
N. meningitidis	..R-----NA.L...L...L...L.P.LG.H...L...D.E.R...D...L...	235
P. multocida	..R-----NA.L...L...L...L.P.LG.H...L...D.E.R...D...L...	274
P. horikoshii (Ph0173)	..R-----NA.I.G.S.LY...I...L.P.I.YLN...R.SL.LDE.E.FR..I.D...LV...	246
P. horikoshii (Ph1245)	..R-----NA.I...S.LY...I...L.P.I.YLN...R.SL.LDE.E.FR..VV...LV...	242
S. pyogenes (Spy1047)	..R-----NA.L.G.S.V...L...G.H...L...D.E.R...D...L...	236
S. pyogenes (Spy1562)	..R-----NA.L.G.S.L...L...L.P.WG.H...R.SL.LDE.E.FR..IVD...LI...	265
S. solfataricus	..R-----N.L.G.L...V...L.P.ISFLN...R.SL...DE.E.FR...VD...L...	306
T. volcanium	..R-----NA.I.G.S.LY...I...L.P.ISFLN...R.SL.LDI.E.FR..IVE...LV...	248
T. maritima	..R-----N.I.G.S.LY...I...L.P.IGYLN...R.SL.LDI.E.FR..VVD...LV...	246

Cas2 homologues

A. aeolicus	..VILVYDV...R..K.K...-I..VQ.SVFEGET...L...I...D.V.IY...-D	84
A. fulgidus (Af1876)	..LWVYDI...R..RL.K.L...L..VQNSAF.GEL...L...V.K.V...D.I.L...GVE	84
A. fulgidus (Af2434)	YVIVAYDV...R..RV.K.L...L..VQNSLPEGELS...V...L...I...D.V.IY...GIE	82
B. halodurans	Y.V...R..KV.K...L...L...VQNSVFE...V...L...I...D.L.IY...GIE	85
M. thermoautotrophicum	YLLIVYDV...R..RV...L...L..VQNSVFEGETV...L...L.R.I...D.V.IY...GLE	81
M. jannaschii	YVIVYDV...R..KI...L...L..VQNSVFEGETV...I...I.R.I...D.V.IY...GLE	104
M. tuberculosis	..VLVIYDI...R..L.K.L...L..VQ.SAFE.LT...L...I.R.A...D.I.IY...G-	103
P. horikoshii (Phrep02)	YIVVYDI...R..KV.K.L...L..VQNSVFEGETV...L...L.K.I...D.V.IY...GIE	77
P. horikoshii (B2)	YIVVYDV...R..R.K.L...L...QNSVFEGELS...L...L.R.V...D.V.IY...G.D	77
S. pyogenes	YDV...R..V.K...L...L..VQNSLPEGELS...I...L...I...D.I.Y...G...	85
S. solfataricus	YLI...YDI...R..RV...L...L..IQ.SVP.GDL...V...L...I...E.I...I...	90
T. maritima	YVI.VYDV...R..KI.K.A...L..VQNSVFE.VT...L...V.R.I...D.V.Y...GVE	79

Cas3 homologues

A. pernix	...I.V.TQVLE.VDL...TE..ID.VIGR.GR..R	385
A. aeolicus	...V.VTQIAE.LDL...TE..ID.LIGR.GR..R	556
A. fulgidus	...V.VTQVIE.VDI...TE..LIGR.GR..R	382
B. halodurans	...I.V.TQVIE.VDV...LD.I.Q.GR..R	600
E. coli	...I.V.TQVVE.LDV...T...D.L.GR.GR..R	691
E. coli O157	...V.I.TQVLE.VD...VD.LIGR.GR..R	695
M. thermoautotrophicum	...I...TQVIE.VDI...TE..ID...GR.GR..R	653
M. jannaschii	...V.V.TQVIE.LD...SE...D.LIGR.GR..R	565
P. horikoshii (Ph0176)	...I.V.TQVVE.LDI...TE..ID.LIGR.GR..R	535
P. horikoshii (Ph1246)	...V.TQVIE.LDI...TE..LD.LIGR.GR..R	571
S. pyogenes	...I.L.TQVIE.VDV...TE..ID.IVQ.GR..R	624
S. solfataricus	...I.I.TQVIE.V.I...TE..I.IVGR.GR..R	400
T. maritima	...L.V.TQVVEA.VDI...D...VD.IVQ.GR..R	565

Cas4 homologues

A. pernix	...I...L...G...I...K...V...PP.P...-C...C...C...	212
A. aeolicus	...QV.YTL...L...-GV...G.I.YPK...VEL...I...PP...-C...C.YYE.C...	177
A. fulgidus (Af2436)	...QL.YTL...YL...-GV...G.I.YPK...VEL...L...P...-C...C.YYE.C...	167

CRISPR Provides Acquired Resistance Against Viruses in Prokaryotes

Rodolphe Barrangou,¹ Christophe Fremaux,² H el ene Deveau,³ Melissa Richards,¹ Patrick Boyaval,² Sylvain Moineau,³ Dennis A. Romero,¹ Philippe Horvath^{2*}

Clustered regularly interspaced short palindromic repeats (CRISPR) are a distinctive feature of the genomes of most Bacteria and Archaea and are thought to be involved in resistance to bacteriophages. We found that, after viral challenge, bacteria integrated new spacers derived from phage genomic sequences. Removal or addition of particular spacers modified the phage-resistance phenotype of the cell. Thus, CRISPR, together with associated *cas* genes, provided resistance against phages, and resistance specificity is determined by spacer-phage sequence similarity.

**CRISPR-Cas is an adaptive defense system
that might use antisense RNAs
as memory signatures of past invasions**

Small CRISPR RNAs Guide Antiviral Defense in Prokaryotes

Stan J. J. Brouns,^{1*} Matthijs M. Jore,^{1*} Magnus Lundgren,¹ Edze R. Westra,¹
Rik J. H. Slijkhuis,¹ Ambrosius P. L. Snijders,² Mark J. Dickman,² Kira S. Makarova,³
Eugene V. Koonin,³ John van der Oost^{1†}

Prokaryotes acquire virus resistance by integrating short fragments of viral nucleic acid into clusters of regularly interspaced short palindromic repeats (CRISPRs). Here we show how virus-derived sequences contained in CRISPRs are used by CRISPR-associated (Cas) proteins from the host to mediate an antiviral response that counteracts infection. After transcription of the CRISPR, a complex of Cas proteins termed Cascade cleaves a CRISPR RNA precursor in each repeat and retains the cleavage products containing the virus-derived sequence. Assisted by the helicase Cas3, these mature CRISPR RNAs then serve as small guide RNAs that enable Cascade to interfere with virus proliferation. Our results demonstrate that the formation of mature guide RNAs by the CRISPR RNA endonuclease subunit of Cascade is a mechanistic requirement for antiviral defense.

1. J. A. Hoffmann, *Nature* **426**, 33 (2003).
2. P. Schmid-Hempel, *Annu. Rev. Entomol.* **50**, 529 (2005).
3. Y. Moret, M. T. Siva-Jothy, *Proc. Biol. Sci.* **270**, 2475 (2003).
4. M. J. Brown, Y. Moret, P. Schmid-Hempel, *Parasitology* **126**, 253 (2003).

Materials and Methods
Figs. S1 to S8
Table S1

9 April 2010; accepted 22 July 2010
10.1126/science.1190689

Sequence- and Structure-Specific RNA Processing by a CRISPR Endonuclease

Rachel E. Haurwitz,^{1*} Martin Jinek,^{1*} Blake Wiedenheft,^{1,2} Kaihong Zhou,^{1,2} Jennifer A. Doudna^{1,2,3,4}†

Many bacteria and archaea contain clustered regularly interspaced short palindromic repeats (CRISPRs) that confer resistance to invasive genetic elements. Central to this immune system is the production of CRISPR-derived RNAs (crRNAs) after transcription of the CRISPR locus. Here, we identify the endoribonuclease (Csy4) responsible for CRISPR transcript (pre-crRNA) processing in *Haemophilus influenzae*. A 1.8 angstrom crystal structure of Csy4 bound to its cognate RNA reveals sequence-specific interactions in the major groove of the crRNA repeat stem-loop. Electrostatic contacts to the phosphate backbone, these enable Csy4 to bind select pre-crRNAs using phylogenetically conserved serine and histidine residues in the recognition mechanism identified here explains sequence- and structure-specific processing of CRISPR-specific endoribonucleases.

In prokaryotes, fragments of foreign DNA are integrated into clustered regularly interspaced short palindromic repeat (CRISPR) loci that are transcribed as long RNAs containing a repetitive sequence element derived from the host (*1–6*). These CRISPR transcripts posttranscriptionally process that serve as homing oligonucleotides for the propagation of invading cognate sequence

ARTICLE

doi:10.1038/nature09886

CRISPR RNA maturation by *trans*-encoded small RNA and host factor RNase III

Elitza Deltcheva^{1,2}, Krzysztof Chylinski^{1,2*}, Cynthia M. Sharma^{3*}, Karine Gonzales², Yanjie Chao^{3,4}, Zaid A. Pirzada², Maria R. Eckert², Jörg Vogel^{3,4} & Emmanuelle Charpentier^{1,2}

CRISPR/Cas systems constitute a widespread class of immunity systems that protect bacteria and archaea against phages and plasmids, and commonly use repeat/spacer-derived short crRNAs to silence foreign nucleic acids in a sequence-specific manner. Although the maturation of crRNAs represents a key event in CRISPR activation, the responsible endoribonucleases (CasE, Cas6, Csy4) are missing in many CRISPR/Cas subtypes. Here, differential RNA sequencing of the human pathogen *Streptococcus pyogenes* uncovered tracrRNA, a *trans*-encoded small RNA with 24-nucleotide complementarity to the repeat regions of crRNA precursor transcripts. We show that tracrRNA directs the maturation of crRNAs by the activities of the widely conserved endogenous RNase III and the CRISPR-associated CsnI protein; all these components are essential to protect *S. pyogenes* against prophage-derived DNA. Our study reveals a novel pathway of small guide RNA maturation and the first example of a host factor (RNase III) required for bacterial RNA-mediated immunity against invaders.

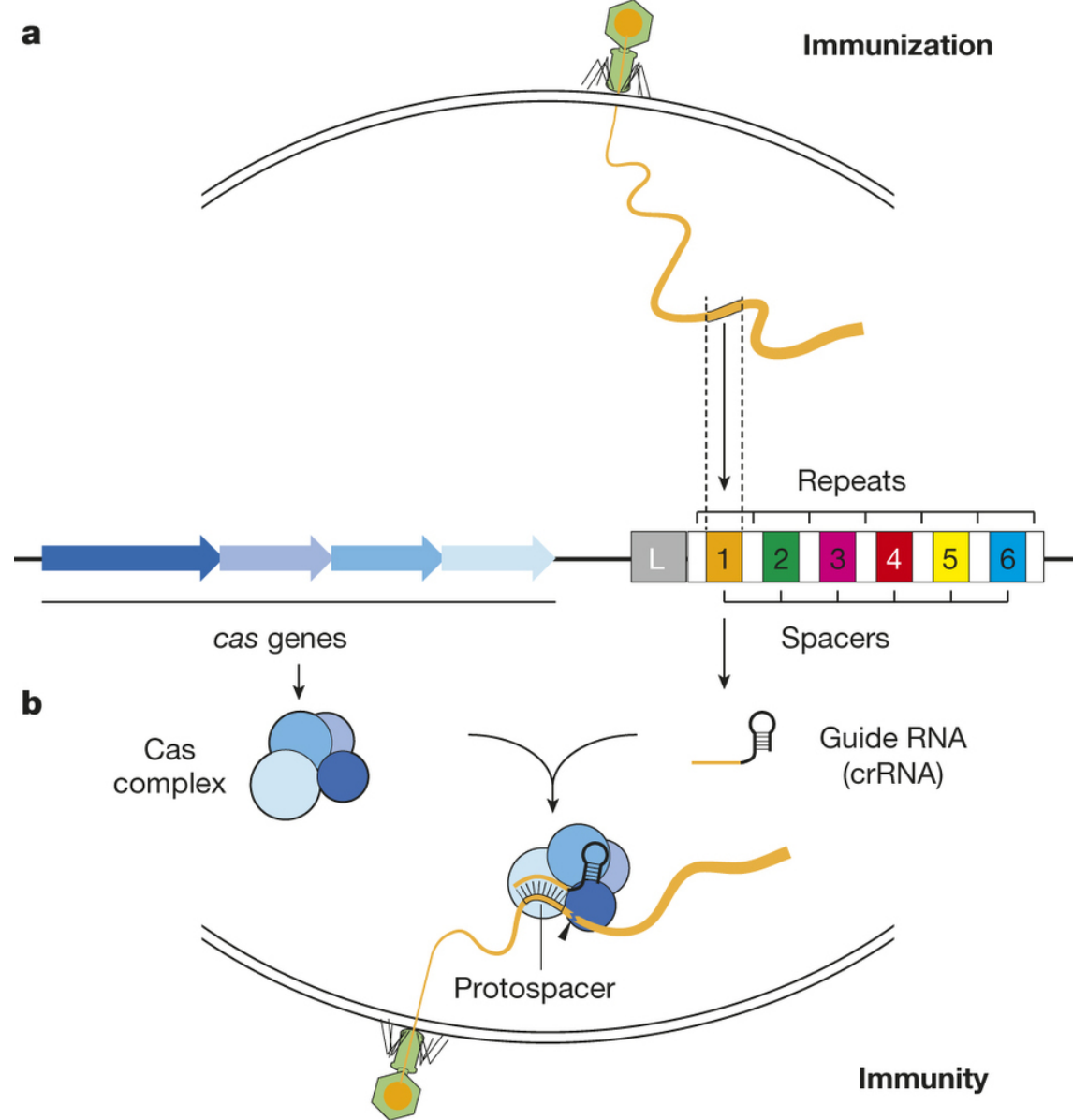
Organisms of all kingdoms of life have evolved RNA-guided immunity mechanisms to protect themselves against genome invaders^{1–6}. In bacteria and archaea, CRISPR/Cas (clustered, regularly interspaced short palindromic repeats/CRISPR-associated proteins) constitutes

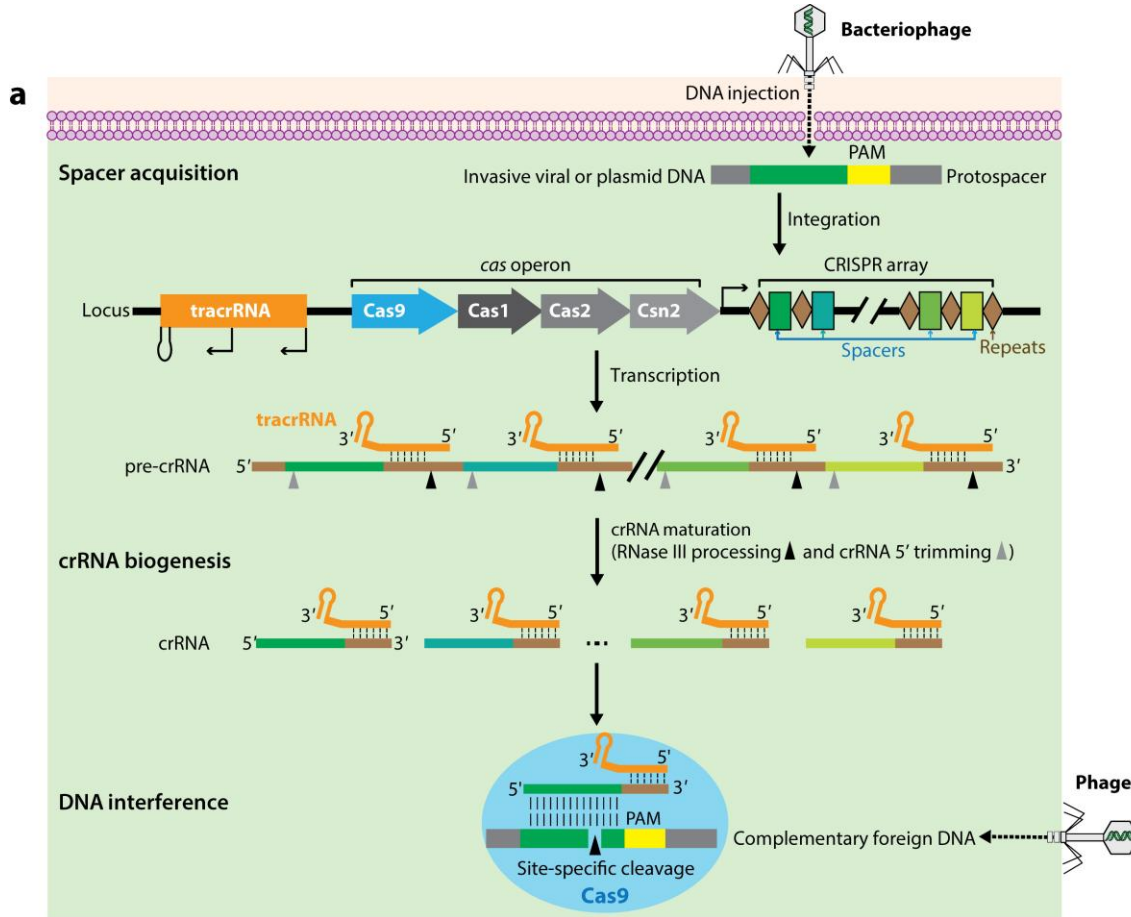
To examine the *in vivo* expression of CRISPR01 and CRISPR02, we analysed *S. pyogenes* strain SF370 (M1 serotype) by differential RNA sequencing (dRNA-seq)⁷. The most abundantly recovered small RNA species were CRISPR01 crRNAs originating from a ~511-nucleotide

CRISPR loci are a cluster of short DNA repeats (white boxes) separated by **equally short spacer sequences of phage and plasmid origin**. This repeat/spacer array is **flanked by an operon of CRISPR-associated (cas) genes** that encode the machinery for the immunization and immunity stages of the system. The CRISPR array is preceded by a leader sequence (grey box) containing the promoter for its expression.

Immunization stage: spacer sequences are captured upon entry of the foreign DNA into the cell and integrated into the first position of the CRISPR array.

Immunity stage: spacer used to target invading DNA that carries a cognate sequence for destruction. **Spacers are transcribed and processed into small CRISPR RNAs (crRNAs)**. These small RNAs act as **antisense guides for Cas RNA-guided nucleases** that cleave the target sequence (black arrowhead) in the invader's genome





CRISPR–Cas system comprises an array of repeat interspaced by short stretches of nonrepetitive sequences (spacers), and a set of CRISPR-associated (cas) genes. Preceding the cas operon is the **trans-activating CRISPR RNA (*tracrRNA*) gene, which encodes a unique noncoding RNA homologous to the repeat sequences.**

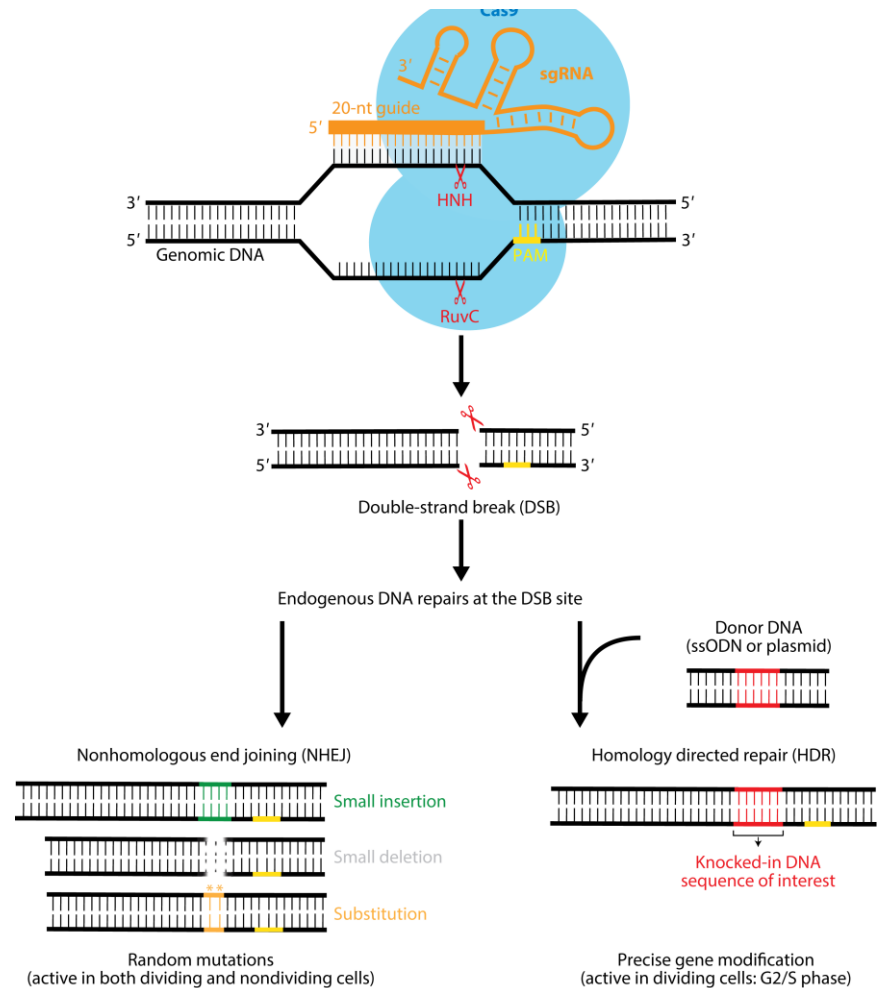
Upon phage infection, a new spacer derived from the phage is incorporated into the CRISPR array by the acquisition machinery (Cas1, Cas2, and Csn2). Then, it is cotranscribed with all spacers into a precursor CRISPR RNA (pre-crRNA). **The *tracrRNA* is transcribed separately and anneals to the pre-crRNA repeats for crRNA maturation by RNase III cleavage.**

During interference, the mature crRNA–*tracrRNA* structure engages Cas9 endonuclease and further directs it to cleave foreign DNA containing a 20-nt crRNA complementary sequence preceding the PAM sequence (protospacer adjacent motif).

CRISPR–Cas9–mediated genome engineering.

The sgRNA or crRNA–tracrRNA structure directs a Cas9 endonuclease to almost arbitrary DNA sequence in the genome through a user-defined 20-nt guide RNA sequence and further guides Cas9 to introduce a double-strand break (DSB) in targeted genomic DNA. The DSB generated by two distinct Cas9 nuclease domains is repaired by host-mediated DNA repair mechanisms. In the absence of a repair template, the prevalent error-prone nonhomologous end joining (NHEJ) pathway is activated and causes random indels or even substitutions at the DSB site, frequently resulting in the disruption of gene function. In the presence of a donor template containing a sequence of interest flanked by homology arms, the error-free homology directed repair (HDR) pathway can be initiated to create desired mutations through homologous recombination, which provides the basis for performing precise gene modification, such as gene knock-in, deletion, correction, or mutagenesis.

CRISPR–Cas9 DNA targeting can be uncoupled from cleavage activity by mutating the catalytic residues in the HNH and RuvC nuclease domains, making it a versatile platform for many other applications beyond genome editing.



Error-prone NHEJ

- Indel mutations
- Frameshift mutations
- Large deletions or inversions using two adjacent DSBs
- Loss-of-function screens
- Genomic rearrangements
- NHEJ-mediated homology-independent knock-in

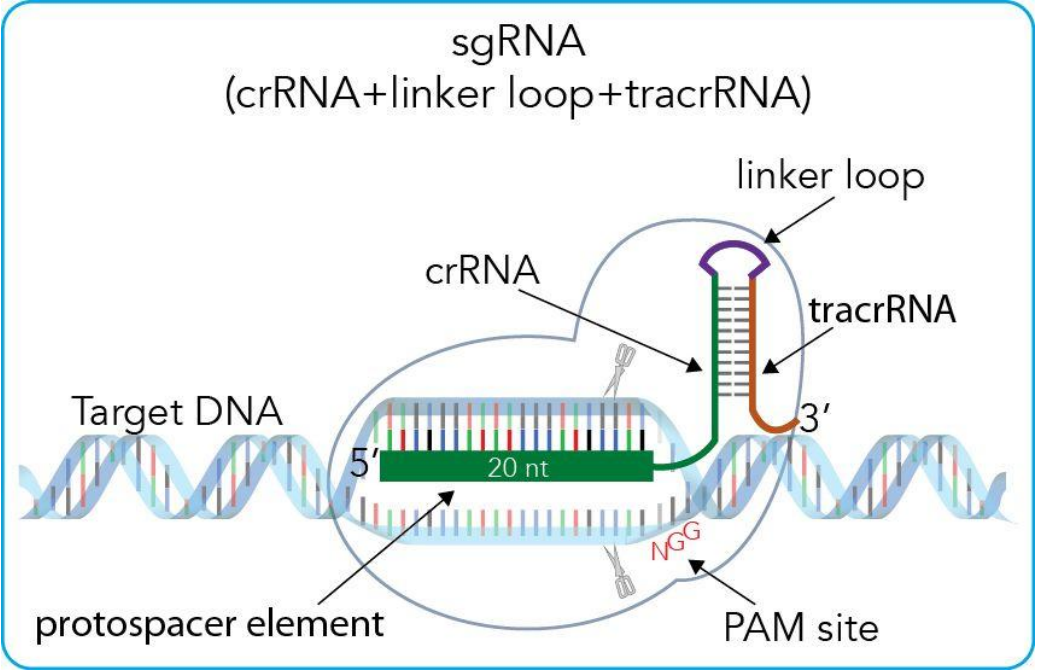
Other applications

- Gene targeting (dCas9–effector):
 - transcriptional regulation
 - epigenetic modification
 - live-cell imaging
 - nucleotide editing
- Genetic screens/drug screens
- Ligation-mediated gene editing by double Cas9 nickases (D10A)

High-fidelity HDR

- Gene insertion/correction/replacement
- Precise point mutations
- Precise gene knockout
- Conditional alleles (Cre–loxP, etc.)
- Introduction of tags, reporters, etc.
- Gain-of-function mutations

A CRISPR/Cas9 nuclease system requires two components: a **Cas enzyme** for cutting the target sequence and a **single guide RNA (sgRNA)**, which binds to the target sequence of 20-base pair (bp). The target sequence (complementary to the sgRNA sequence) is followed by two cytosine nucleotides because the sgRNA binds best when the opposite DNA strand is comprised of any nucleotide followed by two guanines (NGG). This sequence is called a **Protospacer Adjacent Motif (PAM)** sequence. The PAM varies depending on the origin of Cas9.



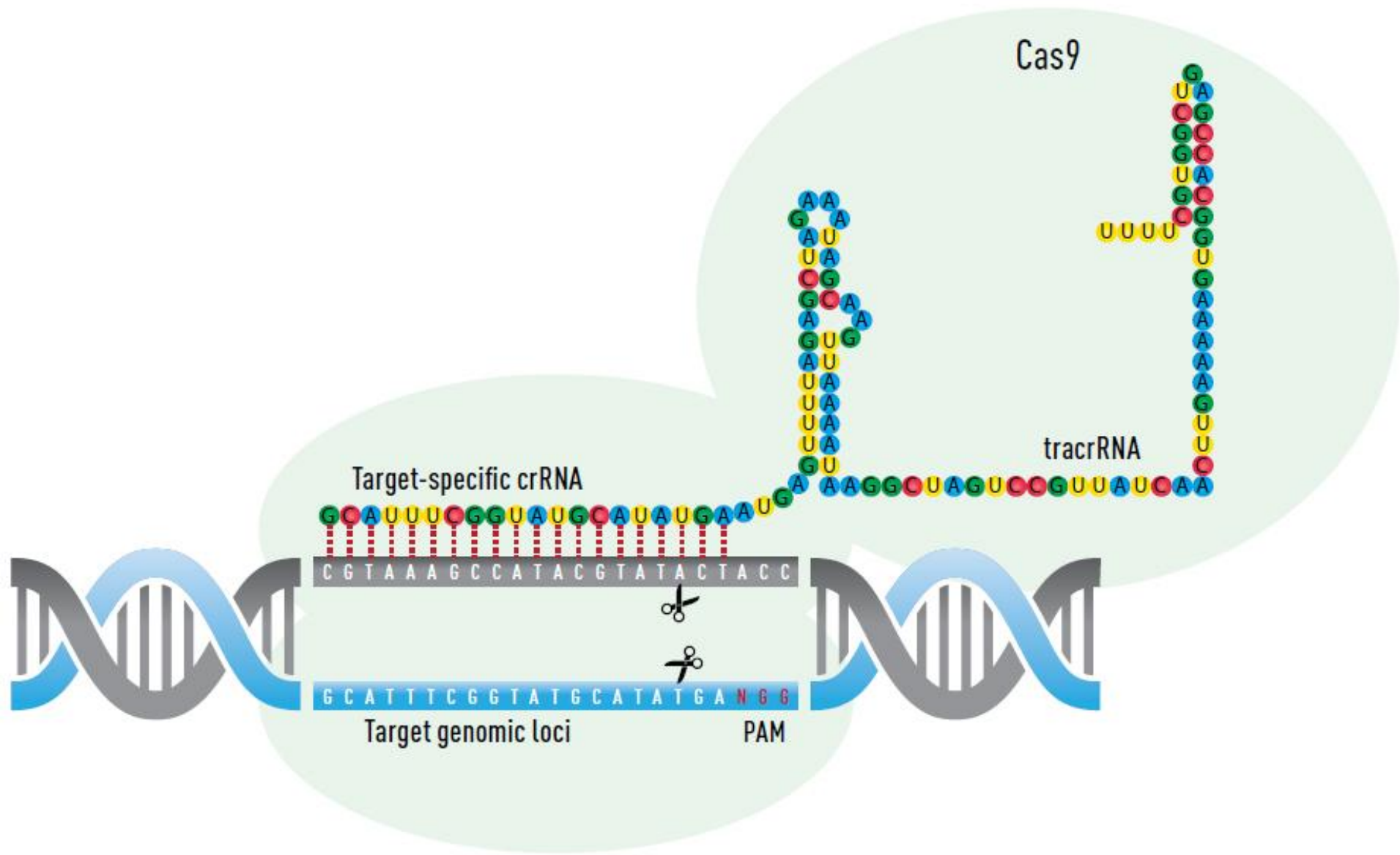
crRNA (CRISPR RNA)—contains both the 20 base protospacer element and additional nucleotides which are complementary to the tracrRNA.

tracrRNA (transactivating RNA)—hybridizes to the crRNA and binds to the CAS9 protein activating the complex to creating double-stranded breaks at specific sites within genomic sequence.

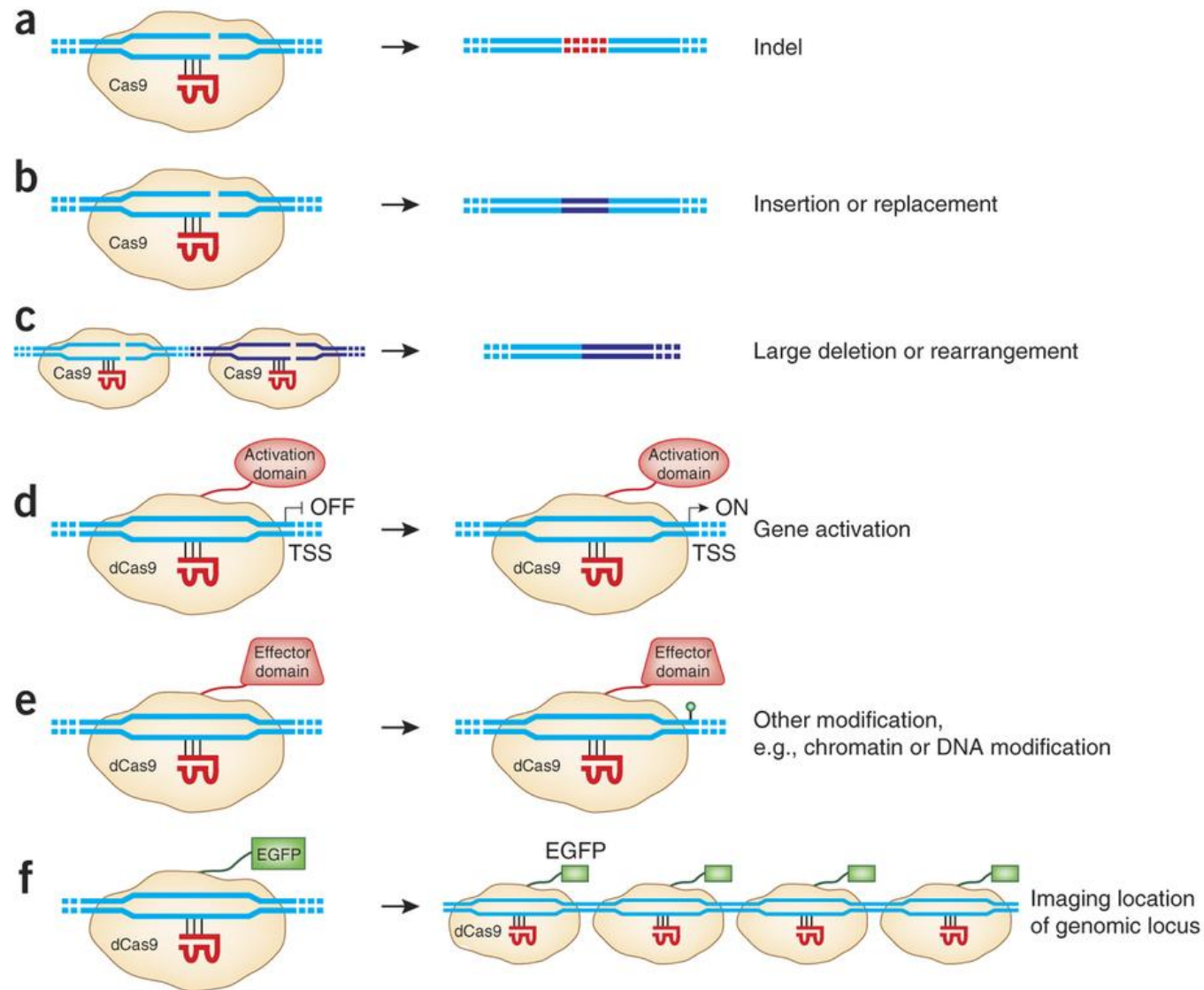
sgRNA (single-guide RNA)—combines the tracrRNA and crRNA, which are separate molecules in the native CRISPR/Cas9 system in *S. pyogenes*, into a single RNA construct, simplifying the components needed to use CRISPR/Cas9 for genome editing. A linker loop sequence is included between the two.

protospacer element — the portion of the crRNA (or sgRNA) that is complementary to the genomic DNA target sequence; usually 20 nucleotides in length.

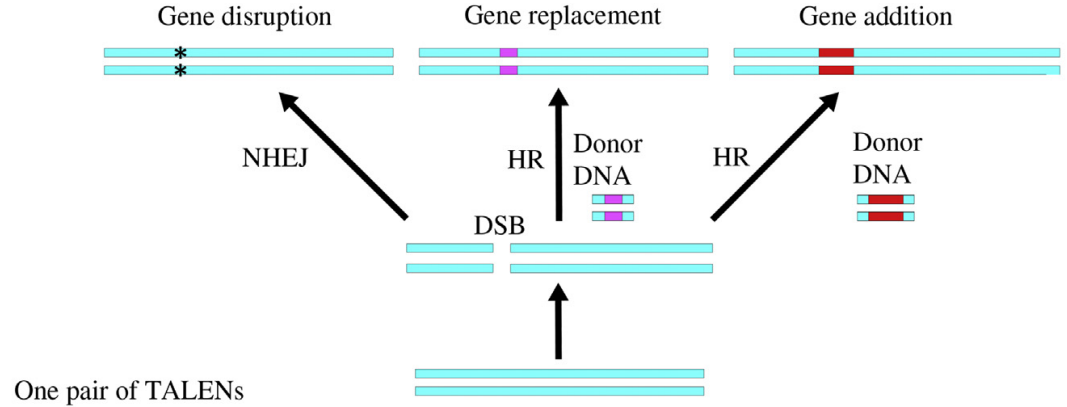
RNA trigger—a generic term for the RNAs that activate the CRISPR/CAS9 complex. They can be sgRNA or crRNA/tracrRNA.



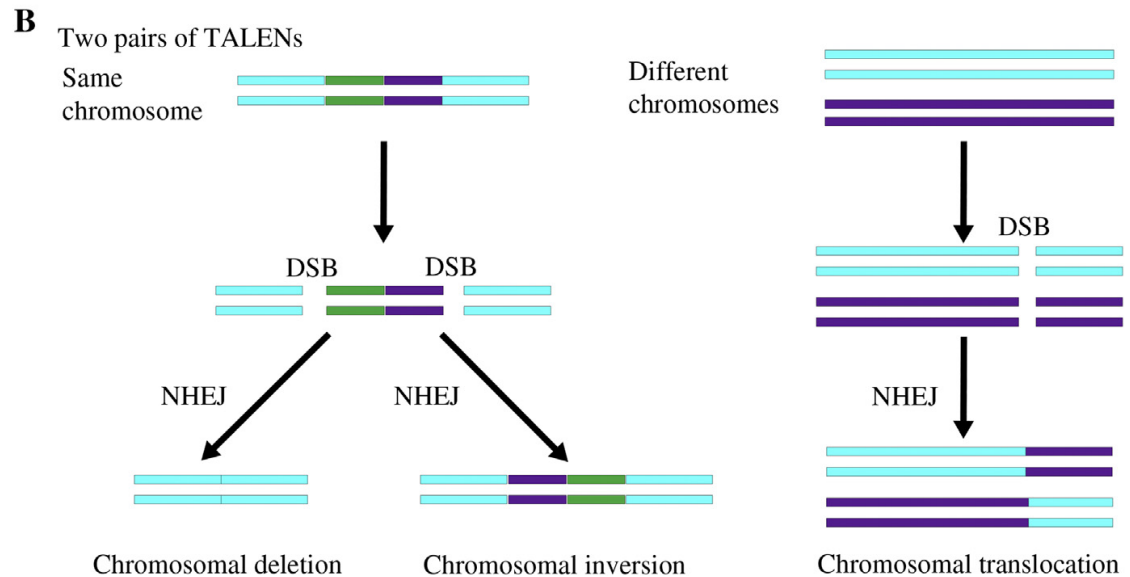
Overview of **various Cas9-based applications**. (a,b) gRNA-directed Cas9 nuclease can induce indel mutations (a) or specific sequence replacement or insertion (b). (c) Pairs of gRNA-directed Cas9 nucleases can stimulate large deletions or genomic rearrangements (e.g., inversions or translocations). (d–f) gRNA-directed dCas9 can be fused to **activation domains** (d) to mediate upregulation of specific endogenous genes, heterologous effector domains (e) to alter histone modifications or DNA methylation, or fluorescent proteins (f) to enable imaging of specific genomic loci.



A: genome modification with one pair of TALENs. Double-strand breaks (DSBs) induced by one pair of TALENs will activate DNA repair mechanisms either by **non-homologous end joining (NHEJ)** or **homologous recombination (HR)**. Through NHEJ, DSBs lead to frame-shift mutations with small insertions or deletions (**indels**) and result in gene disruption (left panel); HR, stimulated by the homologous DNA template, leads to **gene replacement** (middle panel) or **gene insertion** (right panel).



B: genome modification with two pairs of TALENs. The introduction of **two pairs of TALENs targeted at two sites on the same chromosome can cause large deletions** or inversions (left panel). The introduction of two nuclease-induced DSBs **on different chromosomes can lead to translocations** or multiple gene disruptions (right panel).



Since Fok I functions as a dimer, TALENs designed as pairs **allow two monomers to bind at two adjacent sites separated by a DNA spacer that allows Fok I to form dimers to cleave the given target sites**. Consequently, DSBs induced by TALENs will activate DNA repair either by non-homologous end joining (NHEJ) or homologous recombination (HR). If DSBs are repaired by NHEJ, an error-prone pathway which often simply rejoins broken DNAs imprecisely, this will lead to frame-shift mutations with small insertions or deletions (indels) at or near the DSB, so this repair pathway can be used to create KO mutants. Alternatively, if DSBs are repaired by HR, which is stimulated by the homologous DNA template surrounding the DSBs, HR can be employed to create site-specific sequence modifications or insertions.