A CRISPR view of development

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The CRISPR (clustered regularly interspaced short palindromic repeat)–Cas9 (CRISPR-associated nuclease 9) system is poised to transform developmental biology by providing a simple, efficient method to precisely manipulate the genome of virtually any developing organism. This RNA-guided nuclease (RGN)-based approach already has been effectively used to induce targeted mutations in multiple genes simultaneously, create conditional alleles, and generate endogenously tagged proteins. Illustrating the adaptability of RGNS, the genomes of >20 different plant and animal species as well as multiple cell lines and primary cells have been successfully modified. Here we review the current and potential uses of RGNS to investigate genome function during development.

Through the regulated process of development, a single cell divides and differentiates into the multitude of specialized cells that compose a mature organism. This process is controlled in large part by differential gene expression, which generates cells with distinct identities and phenotypes despite nearly identical genomes. Recent advances in genome engineering provide the opportunity to efficiently introduce almost any targeted modification in genomic DNA and, in so doing, the unprecedented ability to probe genome function during development in a diverse array of systems.

Over 25 years ago, homologous recombination (HR)-based gene targeting in mice opened the door to targeted genome engineering for the study of development (Smithies et al. 1985; Thomas et al. 1986; Mansour et al. 1988). HR has been widely used in mice, but its low efficiency requires the selection of rare events in embryonic stem cells, and its implementation in most other organisms has been limited. In contrast, sequence-specific nucleases efficiently catalyze genome editing in a wide variety of organisms. Zinc finger nucleases (ZFNs), transcriptional activator-like effector nucleases (TALENs), and the recently developed two-component CRISPR (clustered regularly interspaced short palindromic repeat)–Cas9 (CRISPR-associated nuclease 9) system can all be programmed to generate targeted double-strand DNA breaks (DSBs) in genomic DNA. Researchers are able to co-opt the endogenous cellular pathways that repair these DSBs to introduce precise changes to the genome. The CRISPR–Cas9 system has propelled genome editing from being a technical possibility to a practical reality for developmental biology studies due to the simplicity with which the Cas9 nuclease is recruited to a specific DNA sequence by a small, easily generated guide RNA (gRNA) that recognizes its genomic target via standard Watson-Crick base-pairing. In this review, we highlight how this transformative technology is being exploited and applied to dissect developmental mechanisms in a wide variety of organisms and cultured cells.

Making the cut: comparing the CRISPR–Cas9 system with ZFNs and TALENs

DSBs induced by sequence-specific nucleases are most commonly repaired by the endogenous cellular DNA repair pathways of nonhomologous end joining (NHEJ) or homology-directed repair (HDR), both of which can be exploited to engineer the genome (Fig. 1). NHEJ is an error-prone process that involves direct ligation of the broken ends and can create disruptive insertions and deletions (indels) at targeted cleavage sites. The HDR pathway uses homologous DNA sequences as templates for repair, and, by supplying an exogenous repair template, HDR can be exploited to precisely edit genomic sequence or insert exogenous DNA.

Cas9 enzymes from type II CRISPR–Cas systems are emerging as the sequence-specific nucleases of choice for genome engineering for several reasons. Most notably, as an RNA-guided nuclease (RGN), Cas9 is guided by a single gRNA that is readily engineered. In the case of the most commonly used Cas9, derived from Streptococcus pyogenes, the gRNA targeting sequence comprises 20 nucleotides (nt) that can be ordered as a pair of oligonucleotides and rapidly cloned. In contrast, generating an effective ZFN or TALEN is labor-intensive [see Box 1]. ZFNs and TALENs are proteins that combine uniquely designed and generated DNA-binding sequences with the FokI
nuclease cleavage domain. FokI is an obligate dimer, necessitating the generation of two novel proteins per editing experiment compared with a single gRNA for CRISPR–Cas9-mediated targeting.

The modular nature of the two-component CRISPR–Cas9 system and small size of the targeting gRNA have the added advantage of being particularly amenable to multiplexing. The use of the common Cas9 nuclease in conjunction with multiple gRNAs to introduce mutations in several genes simultaneously has been carried out in cultured mammalian cells as well as genetic model organisms such as mice, zebrafish, and Arabidopsis (Chang et al. 2013; Cong et al. 2013; Feng et al. 2013; Hwang et al. 2013; Li et al. 2013a; Mali et al. 2013c; Mao et al. 2013; Wang et al. 2013a). More recently, multiplexing was successfully used to introduce mutations in monkeys and silkworms (Wang et al. 2013b; Daimon et al. 2014; Niu et al. 2014b).

Finally, the CRISPR–Cas9 system couples efficiency and high specificity with minimal off-target effects when...
Box: 1. A miniguide to genome engineering techniques

**ZFNs**

ZFNs are fusion proteins comprised of DNA-binding C2H2 zinc fingers fused to the non-specific DNA cleavage domain of the nuclease FokI (for review, see Carroll 2011). Each zinc finger can be engineered to recognize a nucleotide triplet, and multiple (typically three to six) zinc fingers are joined in tandem to target specific genome sequences. Because the FokI cleavage domain must dimerize to be active, two ZFNs are required to create a DSB. This technique, which was first successfully used in fruit flies more than a decade ago (Bibikova et al. 2002), has since been used to modify the genomes of many different organisms, including those that had not previously been developed as genetic model systems.

**TALENs**

Similar to ZFNs, TALENs are chimeric proteins comprised of a programmable DNA-binding domain fused to the FokI nuclease domain (for review, see Joung and Sander 2013). TALEs are naturally occurring proteins that are secreted by the bacteria *Xanthomonas* and bind to sequences in the host plant genome, activating transcription. The TALE DNA-binding domain is composed of multiple repeats, each of which are 33–35 amino acids long. Each repeat recognizes a single nucleotide in the target DNA sequence. Nucleotide specificity is conferred by a two-amino-acid hypervariable region present in each repeat. Sequence-specific TALENs are generated by modifying the two residues in the hypervariable region and concatenating multiple TALE repeats together. Because the TALE DNA-binding domain is fused to Fok1, TALENs, like ZFNs, must also be used as dimers to generate DSBs.

**The CRISPR–Cas9 system**

The CRISPR–Cas9 genome-editing method is derived from a prokaryotic RNA-guided defense system (Gasiusnas et al. 2012, Jinek et al. 2012, 2013; Cong et al. 2013; Mali et al. 2013c). CRISPR repeats were first discovered in the *Escherichia coli* genome as an unusual repeat locus (Ishino et al. 1987). The significance of this structure was appreciated later when investigators realized that phage and plasmid sequences are similar to the spacer sequences in CRISPR loci (Bolotin et al. 2005, Mojica et al. 2005, Pourcel et al. 2005). Soon afterward, it was shown that spacers are derived from viral genomic sequence (Barrangou et al. 2007).

In the CRISPR–Cas system, short sequences (referred to as “protospacers”) from an invading viral genome are copied as “spacers” between repetitive sequences in the CRISPR locus of the host genome. The CRISPR locus is transcribed and processed into short CRISPR RNAs (crRNAs) that guide the Cas to the complementary genomic target sequence. There are at least eleven different CRISPR–Cas systems, which have been grouped into three major types (I–III). In the type I and II systems, nucleotides adjacent to the protospacer in the targeted genome comprise the protospacer adjacent motif (PAM). The PAM is essential for Cas to cleave its target DNA, enabling the CRISPR–Cas system to differentiate between the invading viral genome and the CRISPR locus in the host genome, which does not incorporate the PAM. For additional details on this fascinating prokaryotic adaptive immune response, see recent reviews (Sorek et al. 2013, Terns and Terns 2014).

Type II CRISPR–Cas systems have been adapted as a genome-engineering tool. In this system, crRNA teams up with a second RNA, called *trans*-acting CRISPR RNA (tracrRNA), which is critical for crRNA maturation and recruiting the Cas9 nuclease to DNA (Deltcheva et al. 2011; Jinek et al. 2012). The RNA that guides Cas9 uses a short (12–20 nt) sequence to identify its genomic target. This three-component system was simplified by fusing together crRNA and tracrRNA, creating a single chimeric “guide” RNA (abbreviated as sgRNA or simply gRNA) (Gasiusnas et al. 2012, Jinek et al. 2012). While some early experiments indicated that a gRNA may not cleave a subset of targets as efficiently as a crRNA in combination with tracrRNA (Mali et al. 2013c), the ease of using a single RNA has led to the widespread adoption of gRNAs for genome engineering. A number of resources for designing experiments using the CRISPR–Cas9 system are freely available online. (A comprehensive list is available at http://www.gewisc.wisc.edu.)

gRNAs are well designed. This was evident in whole-genome analysis of engineered human stem cells from multiple laboratories, which uncovered few off-target effects (Duan et al. 2014; Kiskinis et al. 2014; Smith et al. 2014; Suzuki et al. 2014; Veres et al. 2014). Recent work has shown that truncating the 5′ end of the gRNA, where mismatches with the genomic target sequence are tolerated, further increases specificity (Fu et al. 2014). The potential for off-target cleavage can also be reduced by using a mutant nickase version of Cas9 (Cong et al.
In this approach, pairs of Cas9 nickases are targeted to generate single-strand breaks on opposite strands of the genomic target DNA. While the frequency of off-target effects is also low for TALENs, off-target cleavage has been a significant concern for ZFNs (for review, see Carroll 2014). The coassembly of multiple zinc fingers in a single ZFN can alter how the zinc finger and DNA interact, reducing specificity and posing challenges for optimal ZFN design. It is of interest to note that rare-cleaving meganucleases (also known as homing endonucleases) hold the promise of even greater specificity than TALENs or RGNs, which makes them of great interest for gene therapy applications. A drawback of meganucleases, however, has been in the difficulty of programming the nuclease to recognize a desired target. To circumvent this issue, a recent study has generated a meganuclease–TALE chimera [megaTAL], which uses the TALE DNA-binding module to target a meganuclease to the desired sequence in the genome (Boissel et al. 2014). Analogous efforts to combine the precision of meganucleases with the flexibility and ease of targeting of RGNs may further revolutionize genome engineering.

Practical considerations of using RGNs in developing organisms

RGNs hold great potential for dissecting how the genome functions during development. Since the CRISPR–Cas9 system has been recently described in detail elsewhere (Hsu et al. 2014; Sander and Joung 2014), we provide just a brief overview of the system (Box 1, Fig. 1A–C) and focus here on a few practical considerations for using RGNs to edit the genome of a developing organism.

The current methods of producing the CRISPR–Cas9 components provide great flexibility in terms of expression and delivery, and biologists can exploit these options to control when and where DSBs are generated in an organism. To introduce DSBs and generate modifications early in development, the CRISPR–Cas9 components can be injected as DNA, RNA, or protein into most developing organisms. This approach, which has been widely used, generates mosaic organisms for analysis. To gain control over which tissues are affected, a plasmid expressing Cas9 under the control of tissue-specific enhancers can be used. Since each cell has a choice of whether to repair a break through NHEJ or HDR, a variety of different repair events will be present in the injected organism (and in individual cells). The frequency at which both alleles of a gene are affected has been reported to be high enough to visualize null phenotypes in developing mice and zebrafish (Jao et al. 2013; Wang et al. 2013a; Yasue et al. 2014; Yen et al. 2014). For example, zebrafish and mice injected with a gRNA targeting the tyrosinase gene resulted in embryos displaying mosaic pigmentation (Jao et al. 2013; Yen et al. 2014). This indicates that it is possible to generate homozygous mutant tissue in developing animals through injection of the CRISPR–Cas9 components.

Genetic mosaicism, however, may not be desirable for all experiments. In this case, it is necessary to first generate a strain that has the desired modifications. Thus, the targeted modification must occur in the germ cells of the injected organism and be compatible with viable progeny. In fruit flies, this process is facilitated by using transgenic animals that selectively express Cas9 in the germline (Kondo and Ueda 2013; Ren et al. 2013; Gratz et al. 2014; Sebo et al. 2014; Xue et al. 2014). The frequency of targeted events is increased using Cas9 transgenic animals, most likely because more consistent levels of Cas9 are achieved with a stably integrated transgene than with injected plasmid, mRNA, or protein. Transgenic expression of the gRNA has also been demonstrated to increase the frequency of targeted events in fruit flies (Kondo and Ueda 2013; Port et al. 2014; Xue et al. 2014). The expression of Cas9 can be restricted to the germline by placing it under the control of tissue-specific regulatory sequences. In contrast, gRNAs are expressed using polymerase III [Pol III] promoters, which are used to prevent gRNAs from being unduly modified and exported from the nucleus. However, Pol III promoters are typically constitutively active and lead to the ubiquitous expression of gRNAs. To provide spatial and temporal control of gRNA expression, strategies to place gRNAs under the control of Pol II promoters are being developed (Gao and Zhao 2014; Nissim et al. 2014). One approach includes nesting gRNA sequence between ribozymes, thus liberating the gRNA from a Pol II transcript. Other approaches take advantage of the endoribonuclease Csy4, a CRISPR-associated enzyme also known as Cas6, which processes CRISPR repeats and can free an appropriately sized gRNA from cellular transcripts (Nissim et al. 2014; Tsai et al. 2014). These approaches will allow greater control over gRNA expression in developing organisms and in cultured cells and would also enable multiple gRNAs to be generated from a single transcript.

Genome engineering with RGNs enables the direct manipulation of nearly any sequence in the genome to determine its role in development. The major limitation as to which genomic loci can be targeted is the requirement of a specific protospacer adjacent motif (PAM). The PAM is a short DNA motif adjacent to the Cas9 recognition sequence in the target DNA and is essential for cleavage. The most commonly used S. pyogenes Cas9 requires the PAM sequence 5′-NGG (in cell lines, other PAMs are recognized, including 5′-NAG, but at a lower frequency) (Jinek et al. 2012; Esvelt et al. 2013; Hsu et al. 2013; Jiang et al. 2013a; Zhang et al. 2014). The PAM is critical for cleavage and increases target specificity but, conversely, can also make some segments of the genome refractory to Cas9 cleavage. For example, AT-rich genomic sequences may contain fewer PAM sites that would be recognized and cleaved by S. pyogenes Cas9. Thus, some poly[dA-dT] tracts, which are implicated in nucleosome positioning (for review, see Struhl and Segal 2013), may be difficult to manipulate using S. pyogenes Cas9. However, Cas9 from other bacteria recognizes different PAM sequences, and some have already been
adapted for genome engineering in human cells. This includes Cas9 from Neisseria meningitides (5'-NNNNGATT), Streptococcus thermophilus (5'-NNNAGAAW), and Treponema denticola (5'-NAAAAC) (Cong et al. 2013; Esvelt et al. 2013; Hou et al. 2013). These nucleases extend the catalog of possible target sequences, opening up different parts of a genome to analysis. Furthermore, Cas9 from different bacteria can be used in combination, increasing the multiplex capabilities of the CRISPR–Cas9 system.

Catching a break with RGNs to introduce genome modifications

With RGNs, a variety of genomic manipulations are brought within reach of developmental biologists studying a diversity of organisms (Table 1). This approach also makes it possible to readily generate mutations in different genetic strains, making it easier to control genetic background and eliminating the need to carry out multigenerational mating schemes to bring different mutations together in the same animal. While the CRISPR–Cas9 system has been widely used to introduce indels and deletions, HDR makes it possible to introduce more precise gene mutations, deletions, and exogenous sequences, such as loxP sites and green fluorescent protein (GFP).

Multiplexing advantages

Genes that have essential roles in development are often functionally redundant, and thus the effects of mutating a single gene can be masked by the presence of another gene. Due to the ease and efficiency with which gRNAs can be generated, multiple gRNAs can be used in a single experiment to simultaneously mutate multiple genes, overcoming issues of redundancy. Recent technical innovations now make it possible to express multiple gRNAs from a single transcript (Nissim et al. 2014; Tsai et al. 2014), making RGN multiplexing experiments even easier to carry out. Such multiplexing experiments will also facilitate multifaceted experiments, including epistasis tests and manipulating genes that are physically very close together in the genome. Multiplexing has already been used successfully to simultaneously disrupt both Tet1 and Tet2 in developing mice following injection into zygotes (Wang et al. 2013a). The CRISPR–Cas9 system has also been used to eliminate two genes in monkeys [Niu et al. 2014b], demonstrating the potential

Table 1. Organisms that have been modified using the CRISPR–Cas9 system

<table>
<thead>
<tr>
<th>Organism</th>
<th>Mutations created in</th>
<th>Alleles generated by</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Cultured cells (heritable?)</td>
<td>NHEJ</td>
<td>HDR</td>
</tr>
<tr>
<td>Vertebrates</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Axolotl</td>
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<td></td>
<td></td>
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<tr>
<td>Human</td>
<td>[Yes]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mouse</td>
<td>[Yes]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Monkey</td>
<td>[Yes]</td>
<td></td>
<td></td>
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<tr>
<td>Rat</td>
<td>[Yes]</td>
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<tr>
<td>Tilapia</td>
<td>[Yes]</td>
<td></td>
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<tr>
<td>Zebradfish</td>
<td>[Yes]</td>
<td></td>
<td></td>
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<tr>
<td>Invertebrates</td>
<td></td>
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<tr>
<td>Freshwater flea</td>
<td>[Yes]</td>
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<tr>
<td>Fruit fly</td>
<td>[Yes]</td>
<td></td>
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<tr>
<td>Roundworm</td>
<td>[Yes]</td>
<td></td>
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<tr>
<td>Plants</td>
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<td></td>
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<tr>
<td>Corn</td>
<td>[Yes]</td>
<td></td>
<td></td>
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<tr>
<td>Liverwort</td>
<td>[Yes]</td>
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<tr>
<td>Rice</td>
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<td></td>
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<tr>
<td>Sorghum</td>
<td>[Yes]</td>
<td></td>
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<tr>
<td>Sweet orange</td>
<td>[Yes]</td>
<td></td>
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<tr>
<td>Thale cress</td>
<td>[Yes]</td>
<td></td>
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<tr>
<td>Tobacco</td>
<td>[Yes]</td>
<td></td>
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<tr>
<td>Wheat</td>
<td>[Yes]</td>
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</table>

We limited our list to those organisms that provide platforms for the study of development (as indicated for some organisms, only cells derived from the organism have been modified to date). Blanks indicate “not tested.” For the organisms in which the CRISPR–Cas9 system has been used extensively, see recent reviews. Given the rapid advances in the field, we apologize for any organisms or references that were inadvertently not included.
of RGN-mediated mutagenesis to create primate models of multigenic human disorders. Multiplexing can be further extended by taking advantage of a catalytically inactive Cas9 that has been adapted as a programmable RNA-guided platform to regulate gene expression [see below] and by using Cas9 from different bacteria in combination.

Conditional alleles

Many gene products of interest to developmental biologists are essential early in development, and mutations in these genes are lethal to an animal before it reaches later developmental stages. Conditional alleles provide spatial and temporal control over gene inactivation and therefore have been invaluable tools for working with genes that cause early lethality. Conditional alleles have also been used to determine where and when a gene is acting during development. The utility of exerting conditional control over gene activity is widely recognized, and an international consortium is currently working to create a library over gene activity is widely recognized, and an international consortium is currently working to create a library of conditional alleles for ~20,000 genes in the mouse genome (Skarnes et al. 2011). While conditional alleles are typically created using HR, in the past year, RGNs have been used to rapidly generate conditional alleles in mice, rats, and flies (Yang et al. 2013; Gratz et al. 2014; Ma et al. 2014d). RGNs will also be valuable in expanding the repertoire of recombinase-expressing lines. Since the expression of the conditional allele reflects the expression pattern of the recombinase, it is advantageous to have a variety of lines that express recombinase in specific tissues or at discrete developmental stages. The CRISPR–Cas9 system was recently used to generate two different Cre recombinase-expressing lines in rats [Ma et al. 2014b]. Thus, RGNs are being used to rapidly generate the tools necessary to probe gene function in a tissue- and time-dependent manner.

Introducing tags to illuminate and manipulate development

Simply observing an organism as it develops can yield valuable insight into mechanisms that regulate processes such as cell proliferation, differentiation, and morphogenesis and tissue growth. At the cell and molecular level, a broad selection of molecular tags has enabled researchers to track the localization of proteins and RNAs in fixed and live tissue during development. While endogenous genes can be tagged using HR or transposable elements (TEs), these approaches are generally technically challenging and time-consuming. Thus, localization experiments often use transgenes to ectopically express tagged proteins or RNAs. However, a drawback of this approach is that transgenes are typically expressed using exogenous promoters and/or enhancers and are integrated in the genome in a nonnative chromatin environment. Therefore, transgenes are often misexpressed relative to the wild-type gene, and this difference in expression pattern or level can result in altered localization and/or activity of the tagged protein or RNA. A key advantage of tagging an endogenous gene is eliminating expression artifacts. A limitation of this approach is that natively expressed protein or RNA may not be easily detected if it is weakly expressed and/or dispersed within cells. To overcome difficulties resulting from sparse expression, approaches such as tag multimerization may be necessary to effectively visualize endogenous proteins and RNAs.

RGNs open the door to quickly and easily tagging endogenous genes for developmental studies. Furthermore, because the CRISPR–Cas9 system is amenable to multiplexing, tags could be added simultaneously to multiple genes or different splice isoforms of a single gene. There is an ever-growing number of genetically encoded molecular tags that can be used for functional analysis, protein purification, or protein and RNA localization studies.

Tagging proteins and RNAs to visualize their expression in vivo is likely to be a common application of RGNs. Indeed, groups working with flies, mice, and rats have already used epitope tags and fluorescent proteins to label endogenous proteins and generated gene expression reporters [Yang et al. 2013; Ma et al. 2014b; Yu et al. 2014]. In flies, a histone acetyltransferase protein encoded by the gene chameau was C-terminally tagged with GFP, and myc was used to tag an uncharacterized gene, CG4221 [Yu et al. 2013]. In mice, the Sox2 gene was tagged with the V5 epitope [Yang et al. 2013]. Additionally, two different fluorescent reporters were generated for the genes nanog and Oct4 [Yang et al. 2013]. These reporters used either the viral 2A peptide or an internal ribosome entry site (IRES) to express fluorescent proteins with the same expression pattern as the endogenous gene but not fused to the protein product. While these groups used standard fluorescent proteins, a spectrum of fluorescent proteins of different colors and with diverse functions are available [for review, see Dean and Palmer 2014].

In addition to tags for visualizing protein localization, there are also genetically encoded tags to manipulate other aspects of protein function. The CRISPR–Cas9 system was recently used to introduce a small destabilization domain tag into the Treacher Collins-Franceschetti syndrome 1 [TCOF1] gene in human 293T cells [Park et al. 2014]. This FKBP protein-based destabilization tag makes the protein to which it is attached susceptible to degradation in the absence of an exogenously supplied chemical, enabling precise post-transcriptional control over protein expression [Banaszynski et al. 2006]. As the RGN-mediated tagging of endogenous proteins becomes more widely adapted, it will be interesting to see how this will drive an expansion of the current repertoire of genetically encoded fluorescent proteins and tags.

Dishing up cultured cells with modified genomes

Cultured cells are an important and widely used system for dissecting cellular, molecular, and biochemical mechanisms of development. Moreover, human-derived cells are essential for characterizing the function of human genes that lack orthologs in other organisms. The first reports of genome engineering using RGNs were carried out in cultured mammalian cells [Cong et al. 2013; Jinek et al. 2013; Mali et al. 2013c], and, in the short time since
these initial reports, RGNs have been used successfully in a large number of cell lines. Mammalian cell lines, such as HEK293 cells, are relatively easy to manipulate and have been extensively used as a quick and straightforward system to characterize and develop rapid improvements in the technology. Because there have been a number of recent reviews covering the use of RGNs [Mali et al. 2013b; Hsu et al. 2014; Wilkinson and Wiedenheft 2014], we focus here on applications of particular interest for the study of developmental mechanisms, specifically the ability to engineer the genomes of pluripotent stem cells.

Human pluripotent stem cells have been difficult to engineer using classic gene targeting strategies (Capecci 2005; Hockemeyer and Jaenisch 2010). New avenues of genetic manipulation of these cell types have been opened up by RGNs, which can be readily programmed with different gRNAs. One of the first reports of the use of RGNs for genome engineering demonstrated success in induced pluripotent stem cells [iPSCs] with a frequency of between 2% and 4% when assayed by deep sequencing of bulk culture [Mali et al. 2013c]. Recovery of engineered cells is increased when Cas9-expressing cells are marked with a fluorescent marker and selected by cell sorting (Ding et al. 2013). Using this strategy, it was reported that clones containing at least one mutant allele could be isolated at frequencies between 51% and 79%. In comparison, TALENs designed against the same set of genes resulted in between 0% and 34% of clones containing at least one mutant allele. This use of RGNs resulted in a considerable improvement over ZFNs and TALENs in the efficiency of genome engineering in pluripotent cells.

To further increase the frequency of editing events and eliminate the necessity for cell sorting, three human PSC [hPSC] lines and one iPSC line were generated that express doxycycline-inducible Cas9 iCRISPR [Gonzalez et al. 2014]. Because gRNAs are relatively small (~100 nt), these RNAs can be transfected into cell types that are poorly transfectable, such as hPSCs. This system has been used for multiplexing and to generate biallelic knockouts for six genes with efficiencies ranging from 17% to 67%. Mutations could be generated after inducing differentiation, allowing for studies of tissue-specific effects. Pluripotent lines have also been made that enable the simultaneous induction of Cas9 and gRNA expression. These inducible pluripotent cell lines will enable the study of gene knockdown following differentiation into a wide variety of cell types.

As discussed below, the development of additional Cas9-mediated tools for regulating gene expression in combination with lentiviral delivery systems has enabled the modulation of the differentiation status of hPSCs [Kearns et al. 2014]. This continuously expanding Cas9-mediated toolbox will advance the study of genetic contributions to differentiation and disease mechanisms using pluripotent cells and has potential for use in gene therapy.

Disease modeling and treatment with RGNs

Gene editing provided by RGNs has offered and will continue to offer important advancements in genetic studies of disease. The ability to introduce deletions as well as point mutations in a wide variety of cell types allows for the generation of cell lines with human disease-causing mutations. A large number of genes in a diversity of human cell types have already been modified [for review, see Niu et al. 2014a], providing the capacity to study mechanistic effects of these mutations. The relative ease of generating mutant animals will yield many additional animal models of disease and supply a means of testing whether specific polymorphisms are the proximal cause of disease in vivo. Additionally, the CRISPR–Cas9 system is amenable to application in organisms not widely used for genetic studies. Organisms that may be better suited to mimic human disease can now be more easily used to generate disease models. For example, mouse models of the bleeding disorder von Willebrand disease fail to fully recapitulate the human disease. To address this, RGNs were used in pigs to engineer a knockout of the vWF gene, whose deficiency causes the human disease [Hai et al. 2014]. Because the organ size and vasculature of pigs more closely resemble those of humans as compared with the more commonly used mouse models, it is likely that studies in this model will better recapitulate the human disease state.

Apart from point mutations and gene deletions, large chromosomal rearrangements can drive specific cancers. By simultaneously introducing gRNAs targeting two different chromosomes or two widely separated regions of the same chromosome, RGNs have been used to introduce targeted inversions and translocations into otherwise wild-type human cells [Choi and Meyerson 2014; Torres et al. 2014]. These engineered cells will ultimately allow for studies of the causative role of these gene fusions in cancer progression. Translocations that drive lung adenocarcinoma [Choi and Meyerson 2014], acute myeloid leukemia, and Ewing’s sarcoma [Torres et al. 2014] have been generated in both HEK293 cells and more physiologically relevant cell types [nontransformed immortalized lung epithelial cells and human mesenchymal stem cells]. Additionally, cell lines harboring chromosomal inversions found in lung adenocarcinoma have also been created [Choi and Meyerson 2014].

As genome editing with RGNs becomes increasingly efficient and precise, it provides exciting opportunities for gene therapy. Previous work using ZFNs and TALENs has demonstrated the promise suggested by genome editing for gene therapy [for review, see Lisa Li et al. 2014]. Proof of principle for the utility of the CRISPR–Cas9 system has recently been shown for a number of well-characterized disease genes. The CRISPR–Cas9 system has been used to correct a mutation that causes cystic fibrosis in both patient-derived primary cultured small intestinal cells and large intestinal stem cells. When assayed in organoid culture, disease-associated defects were rescued in these engineered cells [Schwank et al. 2013]. As discussed above, RGNs can also be used to edit the genomes of pluripotent cells such as iPSCs, allowing for the generation of cell type-specific disease models. Furthermore, such studies suggest that, in the future, it may be possible to generate iPSCs from patients, correct the causative
mutation, and reintroduce these cells (or the differentiated derivatives of these cells) back into the patient to provide rescuing wild-type function.

Corrections of disease loci have also been generated in animals. RGNs were used in mouse embryos to correct a dominant mutation in the Crygc gene, which causes cataracts, and those mice with the corrected locus were free of cataracts [Wu et al. 2013]. Providing additional promise for the potential of gene therapy in adults, delivery of the CRISPR–Cas9 system and a single-stranded donor template by hydrodynamic injection into mice was able to produce edited hepatocytes that were corrected for a point mutation in Fah, a gene that, when mutated, causes hereditary tyrosinemia. These hepatocytes were rescued for function, as indicated by the fact that resulting animals had substantially decreased liver damage as compared with Fah mutant mice [Yin et al. 2014]. The adaptability of the CRISPR–Cas9 system and its high efficiency will result in significant progress in the mechanistic studies of disease and provide promise for advancements in gene therapies.

**Bringing surgical precision to genetic screens**

Advances in developmental biology have been propelled by genetic screens carried out in a variety of organisms and cultured cells. Genetic screens have used a number of different approaches to introduce mutations into the genome or knock down gene activity. These methods include the use of chemicals [e.g., alkylating agents such as EMS and ENU], TEs, and RNAi. Each of these methods has advantages and disadvantages; for example, although chemical mutagenesis generates unbiased mutations, these mutations are typically single base-pair changes and can be difficult (or sometimes impossible) to specifically identify. In contrast, TE-induced mutations are easier to map [for review, see Friedel and Soriano 2010; Ammar et al. 2012; St Johnston 2013]. However, TEs often preferentially insert at particular sites, leading to TE “hot spots” and the lack of mutations in other genomic regions. This uneven distribution of TE-induced mutations can be somewhat addressed by using multiple TEs with different insertional preferences. Another caveat is that TEs can insert and excise multiple times before mapping. This can result in “hit-and-run” mutations for TEs that excise imprecisely, and these mutations can be difficult to identify. More recently, RNAi has been widely used for large-scale genetic screens in a variety of organisms and cultured cells [for review, see Mohr et al. 2010]. Since base-pair interactions mediate target identification, the gene product targeted by RNAi is easy to determine based on sequence. Moreover, RNAi can be controlled temporally and spatially during development, a boon for characterizing genes that might cause early lethality and identifying the cells in which a gene is active. In addition, unlike conditional alleles, RNAi is reversible, allowing for screens to specifically target distinct cell populations at discrete developmental time points. Although RNAi has been a powerful tool for both forward and reverse genetic screens, there are some drawbacks to this approach: Off-target effects are common, targets are generally limited to mRNAs, and, since the target gene is not directly affected, expression is knocked down but not entirely eliminated.

RGN-based screens have the potential to combine the strengths of chemical and TE mutagenesis with the advantages of RNAi. Like RNAi, RGN targets are identified via standard base-pair interactions, making it relatively straightforward to predict which genes will be affected by a given gRNA. Because RGNs directly target genomic DNA, it is possible to target nearly every part of the genome, including noncoding RNAs that are difficult to target using RNAi. RGNs can also be designed to target short sequence motifs that are impossible to target with precision using TEs or chemical mutagenesis. Thus, RGN-based screens can be designed to systematically probe elements that, for example, regulate gene expression [promoters and enhancers] as well as motifs that direct the post-transcriptional processing of a transcript [splice site sequences and mRNA 5' and 3' untranslated regions]. As described elsewhere in this review, a variety of different alleles can be generated using RGNs, and, furthermore, a catalytically inactive Cas9 can be used as a programmable platform for manipulating gene expression levels, expanding the repertoire of screening approaches.

The small size of gRNAs and the ease with which they are generated make it feasible to generate gRNA libraries for large-scale genetic screens. In fact, the first RGN-based genetic screens were recently carried out in cultured mammalian cells [Koike-Yusa et al. 2014; Shalem et al. 2014; Wang et al. 2014; Zhou et al. 2014]. When carrying out such a screen, it is important to consider both the number of genes targeted by the library and the degree of coverage of each gene. The largest library reported to date is comprised of 90,000 gRNAs designed to target 19,000 genes, which equates to about four to five gRNAs per targeted gene [Koike-Yusa et al. 2014]. The screens identified targets affecting the DNA mismatch repair pathway [Koike-Yusa et al. 2014; Wang et al. 2014], resistance to bacterial and chemical toxins [Koike-Yusa et al. 2014; Wang et al. 2014; Zhou et al. 2014], and cell survival and proliferation [Shalem et al. 2014; Wang et al. 2014]. The Zheng group [Shalem et al. 2014] also compared the results of their screen for genes involved in resistance to a drug that inhibits B-Raf with a prior RNAi screen that used the same cell line and drug. This comparison revealed that gRNAs identified targets that could be validated more consistently and efficiently than shRNAs, pointing to the potential advantages of using gRNAs to knock out, rather than knock down, gene function in genetic screens.

While these screens in cultured cells demonstrated the power of a CRISPR–Cas9-based approach, the question remains whether similar screens can be performed in a developing organism. Excitingly, two recent proof-of-principle studies using worms and mice indicate that RGNs will likely be useful for in vivo genetic screens, including unbiased forward genetic screens [Liu et al. 2014a; Mashiko et al. 2014]. The study in worms revealed...
that gRNA-expressing bacteria could be fed to transgenic worms expressing Cas9 to generate targeted mutations in somatic tissue and the germline, the latter giving rise to heritable mutations [Liu et al. 2014a]. This approach is based on a similar method that has been widely used for RNAi-based screens in worms [Fraser et al. 2000]. The study carried out in mice aimed simply to determine the frequency at which RGN-induced mutations could be recovered [Mashiko et al. 2014]. This approach initially used a fluorescence reconstitution assay in HEK293T cells to eliminate ineffective gRNAs. This screening step eliminated ~20% of the gRNAs, which had been randomly selected. These results point to the utility of using a cell culture-based prescreening step to rapidly enrich for gRNAs that can cleave the target site. Since the gRNAs that did not result in GFP reconstitution were not injected into mice, it is not known how performance in this in vitro assay correlates with targeting frequency in vivo. The 238 validated gRNAs, targeting 32 genes, were injected into mouse zygotes. Of the nearly 200 mice recovered, approximately half contained a targeted mutation, and 40% of the autosomal genes that were targeted had biallelic mutations. The high frequency of biallelic mutations suggests that it would be possible to screen the injected mosaic animals for phenotypes, which would preclude the necessity of generating mutant strains. Together, these studies establish the feasibility of carrying out RGN-based genetic screens in developing organisms, setting the stage for screens designed to identify new players and pathways in development.

The CRISPR–Cas9 system as a platform for genome analysis

The CRISPR–Cas9 system also provides a set of readily programmable tools to probe and manipulate the genome without altering the underlying genetic sequence. By fusing different effector domains or proteins to nuclease-inactive Cas9 (dCas9), the CRISPR–Cas9 system can be exploited to interrogate a spectrum of dynamic genome properties (Fig. 1D). The ability to use a short RNA sequence to target different dCas9 fusion proteins to a specific location within the genome has many potential applications. To date, dCas9 chimeras have been used to activate as well as repress gene expression and follow the subnuclear localization of genes and DNA sequences. Most recently, the dCas9 platform was co-opted to create another RNA-guided system for genome engineering by fusing FokI nuclease to dCas, creating an RNA-guided FokI nuclease [RFN] [Tas et al. 2014]. With the Cas9 crystal structure now in hand to provide additional insight into how Cas9 interacts with the gRNA and PAM [Jinek et al. 2014; Nishimasu et al. 2014], even more modifications and applications of this system are likely.

Catalytically inactive dCas9 was initially used as an RNA-guided platform to disrupt gene expression [Qi et al. 2013]. This approach, called CRISPR interference (CRISPRi), revealed that Cas9 by itself can disrupt gene expression, most likely by sterically interfering with the transcriptional machinery. Following this work, several groups then fused different transcription activation domains, such as VP16 and VP48, to dCas9 and demonstrated that these fusion proteins can activate transgenes as well as endogenous genes [Bikard et al. 2013; Cheng et al. 2013; Maeder et al. 2013; Perez-Pinera et al. 2013; Kears et al. 2014]. Similarly, in an extension of the original CRISPRi experiments, effector domains that inhibit gene expression, such as Krüppel-associated box [KRAB] domain, have also been attached to dCas9 [Gilbert et al. 2013]. While these approaches to manipulate gene expression use Cas9, it is worth pointing out that the Cas RAMP module complex [Cmr] has been developed as a tool to target mRNA for cleavage [Hale et al. 2009, 2012]. Cmr belongs to the type III-B CRISPR–Cas system, which is the only CRISPR–Cas system to target RNA rather than DNA. Thus, a number of different CRISPR–Cas-based approaches are being developed to regulate gene expression.

dCas9 has also been co-opted to visualize the subnuclear localization of genomic sequences. By attaching a fluorescent protein to dCas9, it is possible to follow the position of repetitive telomeric sequences, satellite repeats, and individual genes within nuclei of fixed and living cells [Chen et al. 2013; Antin et al. 2014]. This method is analogous to the zinc finger- and TALE-based approaches in which a fluorescent protein is joined to a programmable DNA-binding domain [Lindhout et al. 2007; Miyanari et al. 2013; Thanisch et al. 2014]. By following fluorescently tagged dCas9 [or TALE or zinc finger] in living cells, it is possible to explore how genomic architecture changes as cells divide and differentiate. These approaches can also be applied to determine whether there is a functional relationship between the position of a gene [or other genetic sequence] within the nucleus and gene expression. Moreover, the TALE-based method is sensitive enough to distinguish individual single-nucleotide polymorphisms [SNPs] [Miyanari et al. 2013], allowing for allele-specific analysis.

As mentioned above, TALEs have also been used as a programmable platform. One particularly notable study created a two-component system to regulate gene expression in response to light [Kornermann et al. 2013]. This method, called LITE [light-inducible transcriptional effectors] capitalizes on a blue light-sensitive interaction between cryptochrome 2 [Cry2] and cryptochrome-interacting basic helix-loop–helix [CIB1]. In this approach, CIB1 is fused to different transcriptional regulators, and Cry2 is targeted to specific genomic locations by attaching the TALE DNA-binding domain. LITE has been shown to effectively control gene expression in a light-dependent manner in both cultured neurons and the mouse brain. Given that the designs of the TALE and Cas9 platforms are similar, it is likely that techniques developed with one platform can be easily adapted for the other, creating an expanding number of tools to dissect gene function. One TALE-based method that has already been adapted to the dCas9 platform is the engineered DNA-binding, molecule-mediated chromatin immuno-precipitation method [or, more simply, enChIP] [Fujita
and Fujii et al. 2013]. In enChIP, a Flag-tagged TALE DNA-binding domain or dCas9 is used in combination with an anti-Flag antibody to pull down the targeted genomic sequence and associated proteins and RNA. This approach has been used to isolate telomere-binding proteins and RNAs [Fujita et al. 2013].

While these studies have demonstrated the great versatility of dCas9 and TALE platforms for genome analysis, there are several considerations that a developmental biologist should take into account before planning experiments. First, with just a few exceptions, the experiments using the dCas9 and TALE platforms have been carried out in bacteria and cultured cells. Thus, it remains to be determined how effective many of these techniques will be in dissecting genome dynamics in a developing organism. Additionally, dCas9-based approaches generally require multiple gRNAs for maximum effect; in a somewhat extreme example, it is estimated that ~25–35 gRNAs will be required to visualize the subnuclear position of a typical gene that lacks repetitive sequences (Chen et al. 2013). In addition, off-target effects are an essential consideration for both TALE and dCas9 platforms, especially given recent work revealing that dCas9 can bind up to several thousand off-target sites, many of which are in genes [Kuscu et al. 2014; Wu et al. 2014]. Although these sites are infrequently cleaved by active Cas9, it is unclear how stably Cas9 associates with off-target sequences and how these interactions might affect the use of dCas9 as a programmable platform. Also, in regards to knocking down gene expression, it remains to be determined how effective CRISPRi and dCas9 chimeras are in comparison with RNAi. Notably, CRISPRi and the dCas9 chimeras designed to inhibit gene expression are reportedly less effective in cultured mammalian cells than in bacteria [Gilbert et al. 2013]. Nonetheless, given the ease with which dCas9 and TALE platforms can be programmed and their versatility, the potential application of these approaches to investigating genome dynamics in vivo is enticing to consider.

Concluding remarks

RGNs have made it possible to precisely modify the genomes of a great variety of organisms and cultured cells with unprecedented ease. The rapid pace of improvements, new applications, and adoption for use in diverse organisms makes the CRISPR–Cas9 system an exciting and significant technical leap forward for developmental biology studies. Additional methodological advances will undoubtedly further enhance the use of RGNs. Currently, the majority of RGN-editing experiments have taken advantage of NHEJ to create small indels and larger deletions, which are useful for disrupting gene expression. However, to introduce specific mutations or other tailored modifications [e.g., genetically encoded tags], the HDR pathway must be activated. In most eukaryotic cells, DSBs are repaired more frequently through NHEJ than HDR [for review, see Lieber et al. 2003; Carroll 2014]. Developing effective methods to bias DSB repair toward HDR to integrate a designed mutation or other modification will be of great utility for many developmental biology applications. Additional improvements in the delivery and/or expression of CRISPR–Cas9 components in different systems as well as in the reduction of off-target effects will further increase the efficient use of RGNs. The CRISPR–Cas9 system holds the potential to revolutionize developmental biology by making it possible to probe with exquisite control the interplay between genome activity and developmental events such as cell proliferation, differentiation, and morphogenesis.

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References


Cross-references and annotations for the gene targeting strategies in *C. elegans*, *Drosophila melanogaster*, and *Zebrafish* are included, along with the relevant experimental setups and methodologies.


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