

TECHNOLOGY REPORT

Simple and Efficient CRISPR/Cas9-Mediated Targeted Mutagenesis in *Xenopus tropicalis*Takuya Nakayama,¹ Margaret B. Fish,¹ Marilyn Fisher,¹ Jamina Oomen-Hajagos,² Gerald H. Thomsen,² and Robert M. Grainger^{1*}¹Department of Biology, University of Virginia, Charlottesville, Virginia²Department of Biochemistry and Cell Biology, Stony Brook University, Stony Brook, New York

Received 25 September 2013; Revised 9 October 2013; Accepted 10 October 2013

Abstract: We have assessed the efficacy of the recently developed CRISPR/Cas (clustered regularly interspaced short palindromic repeats/CRISPR-associated) system for genome modification in the amphibian *Xenopus tropicalis*. As a model experiment, targeted mutations of the *tyrosinase* gene were verified, showing the expected albinism phenotype in injected embryos. We further tested this technology by interrupting the *six3* gene, which is required for proper eye and brain formation. Expected eye and brain phenotypes were observed when inducing mutations in the *six3* coding regions, as well as when deleting the gene promoter by dual targeting. We describe here a standardized protocol for genome editing using this system. This simple and fast method to edit the genome provides a powerful new reverse genetics tool for *Xenopus* researchers. *genesis* 51:835–843. © 2013 Wiley Periodicals, Inc.

Key words: *tyrosinase*; albinism; *six3*; brain; eye; sgRNA

INTRODUCTION

Xenopus has long been a favored model organism for developmental embryology due to its unique combination of advantageous features, and the recent development of *Xenopus tropicalis* as a new model organism with a diploid genome, short generation time, and sequenced genome information allows researchers to use genetic tools in frogs (Harland and Grainger, 2011). Forward and reverse genetic approaches have identified developmental mutants and their responsible genes (Abu-Daya *et al.*, 2012).

Recent technological advances have allowed researchers to perform rapid targeted gene editing in many organisms.

Two methods, zinc-finger nucleases (ZFNs) and transcription activator-like effector nucleases (TALENs) (Joung and Sander, 2013), have both been successfully applied in *Xenopus* (Ishibashi *et al.*, 2012; Lei *et al.*, 2012; Nakajima *et al.*, 2012, 2013; Suzuki *et al.*, 2013; Young *et al.*, 2011). Another, simpler technology has also emerged: the CRISPR/Cas (clustered regularly interspaced short palindromic repeats/CRISPR-associated) system for genome modification, originally identified as part of the naturally occurring bacterial adaptive defense mechanism. CRISPR/Cas has now been successfully applied in major model organisms such as zebrafish (Hwang *et al.*, 2013), mouse (Wang *et al.*, 2013), fly (Bassett *et al.*, 2013), and worm (Friedland *et al.*, 2013) to effect targeted genome modification. Briefly, the original Cas9 in *Streptococcus pyogenes* is recruited to the target site by two RNAs, a CRISPR RNA (crRNA) that has complementary sequence to the target DNA, and the *trans*-activating CRISPR RNA (tracrRNA) that base pairs with the crRNA. For recognition, the target sequence must be followed by a protospacer adjacent motif (PAM) sequence (nGG, where n can be any nucleotide). For genome editing, the crRNA and tracrRNA can be fused into a single synthetic guide RNA (sgRNA) (Hwang *et al.*, 2013), which functions efficiently with Cas9 to

Additional Supporting Information may be found in the online version of this article.

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Contract grant sponsor: NIH, Contract grant numbers: EY022954, EY018000, and OD010997 (RMG), HD064908 (GHT), T32GM007964 (JOH); Contract grant sponsor: Sharon Stewart Aniridia Trust (RMG)

Published online 8 November 2013 in Wiley Online Library (wileyonlinelibrary.com). DOI: 10.1002/dvg.22720

cause cleavage of the target site (~20 bp), which must be followed by the PAM sequence in the genome (Fig. 1a). A further constraint on the target sequence is due to the utilization of *in vitro* transcription promoters such as T7, T3 or SP6 to produce sgRNAs: since these promoters work optimally with a +1 guanine residue, the genomic target sequence must begin with a "G." As with ZFNs and TALENs, these cleavages are then often imperfectly repaired via nonhomologous end-joining (NHEJ), which can lead to frameshift-causing in-del mutations occurring mosaically throughout the injected embryo. Here, we report the application of CRISPR/Cas system (hereafter called simply CRISPR) to edit the genome of *Xenopus tropicalis*, providing an additional tool for *Xenopus* researchers to achieve simple and efficient targeted mutagenesis.

RESULTS AND DISCUSSION

In order to establish a method for CRISPR-mediated genome editing of *Xenopus tropicalis*, as well as to determine the optimal conditions for inducing mutations, we first chose the *tyrosinase* (*tyr*) gene as a model for targeting because of the readily apparent mutant phenotype, albinism, as seen in previous work using ZFN or TALEN constructs (Ishibashi *et al.*, 2012; Nakajima *et al.*, 2012). The first *tyr* sgRNA was designed to target a sequence near the region that was used effectively with TALEN constructs (Figs. 1b, 2) (Ishibashi *et al.*, 2012).

We initially examined various doses of injected sgRNA (25–200 pg) and Cas9 mRNA (550 pg–2.2 ng). We also tested two Cas9 variants, one using the original bacterial codons (bacCas9 in Fig. 2a) (Hwang *et al.*, 2013) and the other a "humanized" Cas9 (Chang *et al.*, 2013) that instead uses codons optimized for mammalian genes, and which also includes nuclear localization signals at both ends of the protein (humCas9 in Fig. 2a), both of which have been shown to successfully edit the zebrafish genome. In Figure 2a, since the phenotype is hard to score with a quantifiable scale, we show representative embryos as groups to illustrate qualitative differences. Among the doses tested, a combination of the highest doses using humCas9 (i.e., 2.2-ng humCas9 mRNA and 200-pg sgRNA) showed almost complete albinism (Fig. 2a, left bottom panel), whereas at the same doses, bacCas9 showed much less activity, although it still could cause mutations (data not shown) and patchy loss of pigmentation in the retinal pigment epithelium (RPE) (white arrowheads in Fig. 2a). In general, at the doses using more than 100-pg sgRNA and 1.1-ng humCas9, we see the albino phenotype in essentially 100% of injected embryos although the phenotype may vary from the patchy loss of pigmentation in the RPE to almost complete albinism depending on the doses used. As shown in Figure 2b, we developed a quick genotyping method by simple direct sequencing

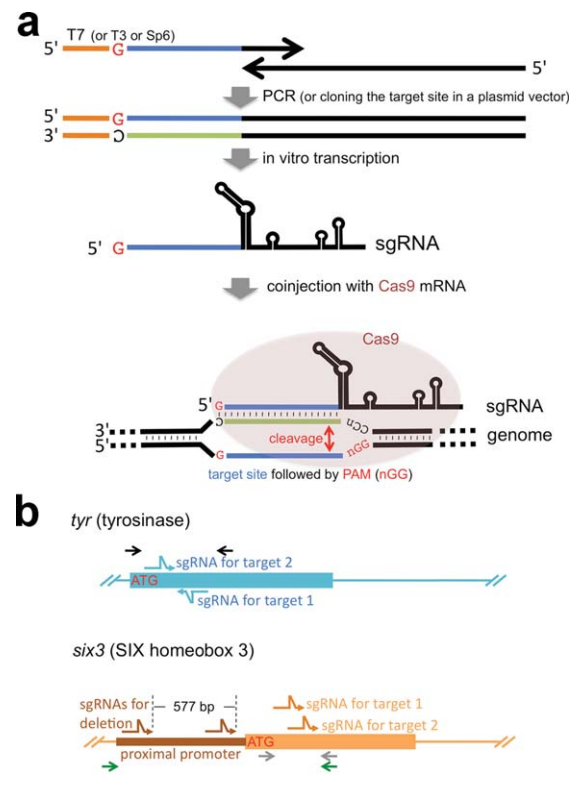


FIG. 1. Strategy for CRISPR/Cas-mediated genome modification. (a) Schematic representation of experimental procedure. We found that the conventional plasmid subcloning method for making sgRNA templates was time-consuming and inconvenient; instead all sgRNAs used in this study are transcribed *in vitro* from double-stranded DNA templates that were made by PCR (except the sgRNA for *tyr* target site 1, which used a cloned template, pDR274-Xt-*tyr*). This PCR strategy uses a 5' oligonucleotide (primer) that begins with the T7 promoter (shown as an orange line in this schematic; alternatively, T3 or SP6 promoters could be used) and contains the genomic target sequence (shown as a blue line in this schematic; note the genomic target sequence must begin with a G for proper transcriptional initiation using the T7 promoter) and a 3' oligonucleotide (primer) that partly overlaps the 5' primer and contains the sgRNA backbone sequence required for proper folding of sgRNAs. This rapid, easy way to make sgRNA templates by PCR takes less than two days to produce sgRNAs to inject once oligonucleotides are received. During the preparation of this manuscript, we have noticed that similar strategies have also been reported [e.g., (Bassett *et al.*, 2013)] and thus this method has general versatility. The resulting sgRNAs were coinjected with Cas9 mRNA into one cell-stage *Xenopus* embryos. The bottom part of the scheme shows how sgRNA and Cas9 work to cleave the target site in the genome. Briefly, the sgRNA forms a complex with the Cas9 protein and identifies the target site via complementary base-pairing. The Cas9 nuclease subsequently cleaves the genomic DNA at the target site, just upstream of the PAM sequence. (b) Targeting strategy for the two genes described in this study. Both *tyr* and *six3* genes were targeted in exon 1 to cause frame-shifts after the translation initiation codon (ATG). Two independent sgRNAs were tested for both genes. The *six3* gene was furthermore targeted in the proximal promoter region with two sgRNAs simultaneously to cause deletion of the promoter region. Bent arrows (blue for *tyr*, orange for *six3* exons and brown for the *six3* promoter) show sgRNA targets. Arrows (black for *tyr*, green and grey for *six3*) indicate genomic PCR primers for mutation analyses. Note that the 3' primer for the *six3* gene is common for both sets of PCR reactions but colored differently for clarity. Drawings are not to scale.

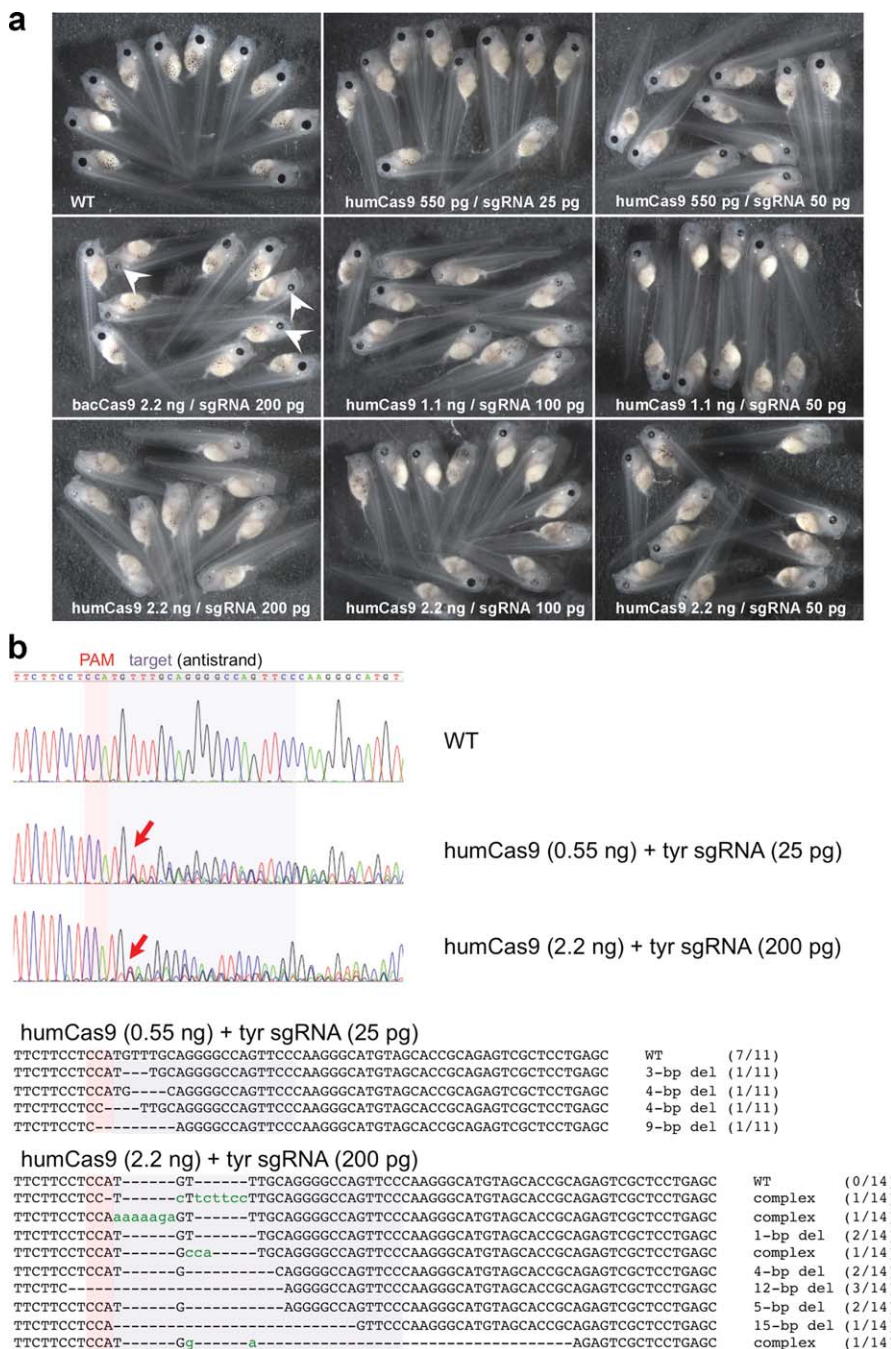


FIG. 2. Successful targeting of the *tyrosinase* gene caused albinism in *Xenopus* embryos. (a) Different dose combinations of Cas9 mRNA and sgRNA were tested. The severity of the phenotype was directly dependent on the amounts of RNAs injected. bacCas9, the original bacterial-codon Cas9 mRNA (which shows examples with a weak phenotype; a patchy loss of pigmentation in the RPE is indicated by white arrowheads). humCas9, the modified Cas9 using humanized-codons. sgRNA, targeting *tyrosinase* gene (first target, see Fig. 3b for location relative to ATG). The toxicity of sgRNA seems to vary depending on its sequence and also on the batch of embryos. For example, in one specific experiment, *tyrosinase* sgRNA (target1) was overall relatively nontoxic, with more than 75% of injected embryos developing normally one day after injection at all tested doses shown here (76.9–97% survivors [n=19–33] as opposed to 84.8% of uninjected embryos [n=46]), whereas in other experiments, survival was between 50–60% (see Supporting Information Table 1). (b) Representative results of conventional sequencing assays (top three panels). Each single embryo injected with indicated RNAs was lysed and the targeted genomic region was PCR-amplified; amplicons were then directly sequenced (DSP assay, see the text). Perturbation of peaks on the 3' side of the PAM region (red arrows) suggests in-del events happen in-between the target sequence (shaded in purple) and the PAM region (shaded in red). PCR amplicons were recloned and sequenced to show the profile of individual mutations found in mosaic individuals (bottom sequence alignments). Dashes (-) indicate gaps. Lowercase green characters indicate insertions or substitutions. The numbers in parentheses indicate the frequency of each mutation pattern seen in total numbers of sequenced clones.

a**b**

| Met | second target | PAM | first target and PAM | | |
|-------------------|---------------------------------------|--------------------------|----------------------|--------|--|
| ATGGAAGGAACATGG | TCCCTCTGGCATTCTGCTGCCTGTTCTTCTTCTCC | TCCATGTTTGCAGGGGCCAGTTCC | WT | (0/7) | |
| ATGGAAGGAACATGG | CTCTGGCATTCTGCTGCCTGTTCTTCTTCC | TCCATGTTTGCAGGGGCCAGTTCC | 3-bp del | (1/7) | |
| ATGGAAGGAACATGGTC | TGGCATTCTGCTGCCTGTTCTTCTTCTCC | TCCATGTTTGCAGGGGCCAGTTCC | 4-bp del | (5/7) | |
| ATGGAAGGAACACGG | TCTGGCATTCTGCTGCCTGTTCTTCTTCTCC | TCCATGTTTGCAGGGGCCAGTTCC | complex | (1/7) | |
| | | | | | |
| ATGGAAGGAACATGG | TCCCTCTGGCATTCTGCTGCCTGTTCTTCTTCTCC | TCCATGTTTGCAGGGGCCAGTTCC | WT | (0/10) | |
| ATGGAAGGAACATGG | TC--TCTGGCATTCTGCTGCCTGTTCTTCTTCTCC | TCCATGTTTGCAGGGGCCAGTTCC | 2-bp del | (3/10) | |
| ATGGAAGGAACATGG | CATTCTGCTGCCTGTTCTTCTTCTTCTCC | TCCATGTTTGCAGGGGCCAGTTCC | 9-bp del | (1/10) | |
| ATGGAAGGAACA | GCATTCTGCTGCCTGTTCTTCTTCTTCTCC | TCCATGTTTGCAGGGGCCAGTTCC | 11-bp del | (1/10) | |
| ATGGAAGGAACATG | TGCTGCCTGTTCTTCTTCTTCTTCTCC | TCCATGTTTGCAGGGGCCAGTTCC | 15-bp del | (1/10) | |
| ATGGAAGGAACATGGaa | ca-TGGCATTCTGCTGCCTGTTCTTCTTCTCC | TCCATGTTTGCAGGGGCCAGTTCC | complex | (1/10) | |
| ATGGAAGGAACATGG | cccatggCTGTCATTCTGCTGCCTGTTCTTCTTCTCC | TCCATGTTTGCAGGGGCCAGTTCC | complex | (2/10) | |
| ATGGAAGGAACATGG | ctcatggCTGTCATTCTGCTGCCTGTTCTTCTTCTCC | TCCATGTTTGCAGGGGCCAGTTCC | complex | (1/10) | |

FIG. 3. The mutation of a second *tyr* gene target site also caused albinism. **(a)** Phenotype of injected embryos (bottom five embryos; top five embryos are uninjected sibling controls) with Cas9 mRNA (2.2 ng) and sgRNA (200 pg) for the second target site. **(b)** Representative mutation profiles from two embryos sequenced after cloning. The second target sequence (shaded in purple) and the PAM sequence (in red) are shown together with first target and PAM sequences (in gray; see Fig. 2) relative to the ATG (Met) site. The alignment shown is labeled as described in Fig. 2b.

of the PCR-amplified targeted genome region (hereafter, called DSP assay), namely, a single embryo was subjected to genomic PCR to amplify the target region and the resultant amplicon was directly sequenced without cloning. If mutations occur at relatively high frequency, the resultant PCR amplicon is a mixture of heterogeneous fragments with different mutations, and the reading of sequences becomes perturbed in the mutated region. Typical results of samples from a group injected with the lowest doses (humCas9 550 pg/sgRNA 25pg) or from a group injected with the highest doses (2.2 ng/200 pg) compared with wild-type sample are shown (Fig. 2b). Each sample was also subsequently recloned and individual clones per embryo were sequenced as shown in Figure 2b (bottom sequencing alignments). Not surprisingly, since we saw almost completely albino embryos, indicative of highly efficient mutagenesis, the sample from the highest doses showed perturbation of peaks in the targeted region (Fig. 2b, the bottom chromatogram, arrow), suggesting the presence of in-del mutations in this region. Interestingly, a sample from the lowest doses also clearly showed perturbation of peaks in the targeted region (Fig. 2b, middle chromatogram) although more than 60% (7 out of 11 clones sequenced) of recovered sequences from an embryo were not mutated (Fig. 2b, top sequence alignment), suggesting that this method (DSP assay) is sensitive enough for quick screening of functional sgRNAs that

have ability to cause mutations at the molecular level, even if relatively infrequent. Given that the phenotype at this dose was very mild, the DSP assay is also suitable for screening of putative carriers of potentially lethal mutations where a low mutation rate is necessary to be able to raise carriers. Although, more sensitive screening methods could detect less efficiently mutated carriers, such carriers might not be practical for making mutant line(s) since one would need to screen so many offspring to identify heterozygotes in the next generation.

Since it has shorter recognition sequences compared to ZFNs or TALENs, CRISPR may potentially create more off-target mutations, and to overcome this problem several approaches are being used (Carroll, 2013). In order to rule out the possibility that off-target effects are causing observed phenotypes, we have established a standard method for performing F0 embryo assays using *Xenopus tropicalis*: (1) target at least two independent regions of the gene of interest to see reproducibility of the phenotype (including its variations), and, if possible, (2) perform rescue to confirm the phenotype is specific to a particular gene. However, it remains valuable to consider the issue of off-target effects, and the bioinformatics approaches described in the accompanying paper (Blitz et al., 2013), and below, can be used to identify putative sites and then evaluate the degree of mutagenesis in such off-target sites as performed by Blitz et al. (2013).

To pursue the two-target strategy, as shown in Figure 3, we designed a second sgRNA for a different region of the *tyr* gene (Fig. 1b, 3b). Hereafter, we only used hum-Cas9 (therefore simply called Cas9 below). Using the combination of the highest doses of RNAs as used earlier, the second sgRNA also showed the expected phenotype in 100% of embryos injected in multiple experiments although the severity varied from patchy loss of pigmentation to almost complete albinism (Fig. 3a). Two independent sgRNAs targeting different regions of the gene showed the same phenotype, sug-

gesting that the phenotype is likely due to inactivation of the targeted gene, and not due to off-target mutations. However, we failed to rescue the phenotype by coinjection of *tyr* mRNA (up to 200 pg) under conditions we tested (data not shown). This is likely due to the timing of expression of the *tyr* gene, which begins at late tailbud stages when injected mRNA is likely to be already degraded.

To demonstrate the general applicability of this approach, we chose a second gene, *six3*, of direct interest to ongoing studies of eye determination in the

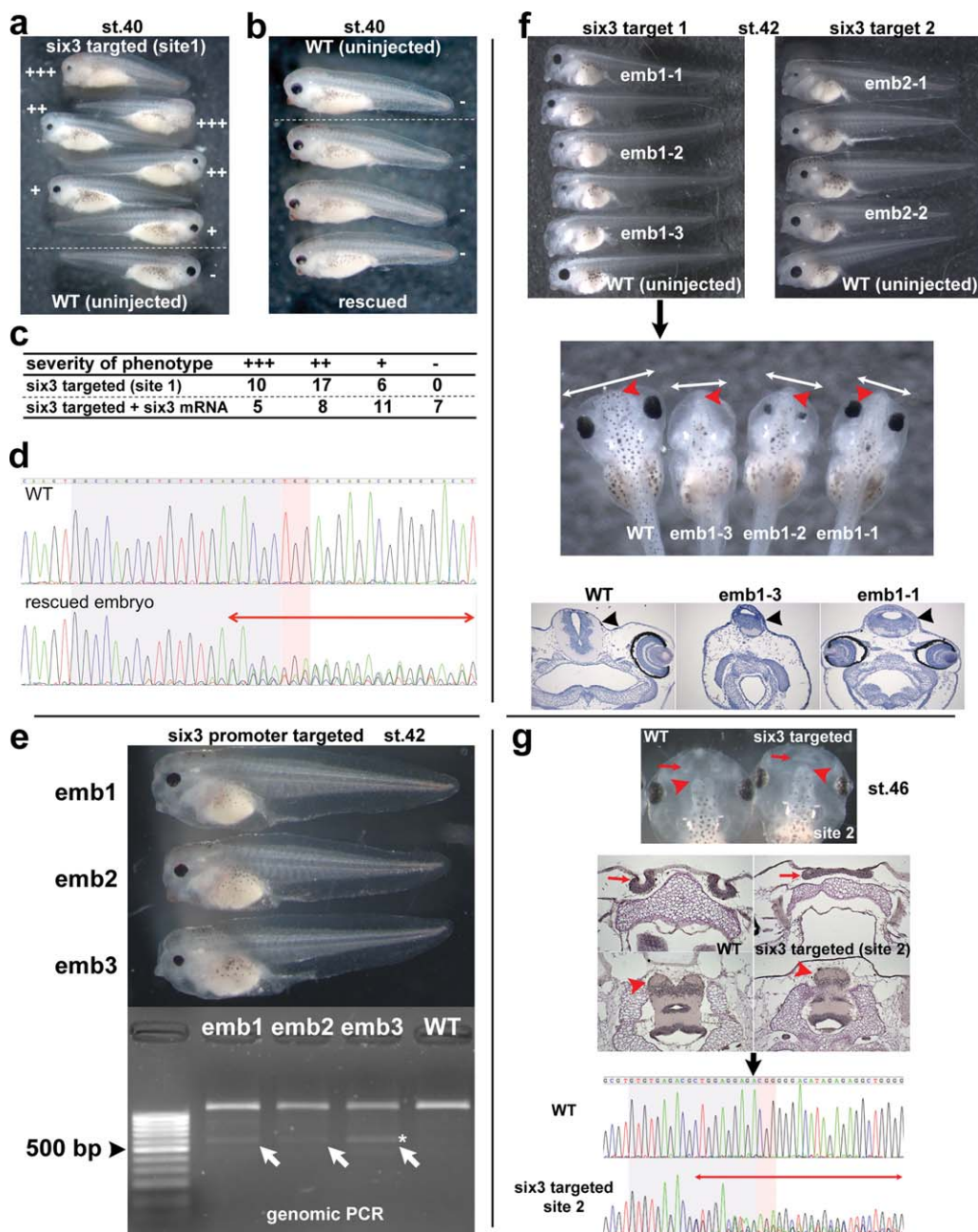


FIG. .4.

Grainger lab. *six3* is one of a group of eye-field transcription factors expressed in the anterior region of the vertebrate neural plate and is essential for eye formation (Zuber *et al.*, 2003). We found that embryos injected with the first *six3* targeted CRISPR (for target site1) showed consistent abnormalities that were similar to known loss-of-function phenotypes of *six3* protein. In *X. laevis*, loss-of-function of *six3* using a dominant-negative construct or morpholino (MO) knock-down was reported to show eye and anterior head defects (Gestri *et al.*, 2005). In humans, mutations in the homeodomain of *SIX3* gene are known to cause holoprosencephaly (HPE) (Wallis *et al.*, 1999), and haploinsufficiency of *Six3* in the mouse causes HPE (Geng *et al.*, 2008). Complete inactivation of *Six3* results in the absence of the eyes and nose in mouse (Lagutin *et al.*, 2003). What we have observed in *X. tropicalis* mutant embryos here is consistent with those reports (see Fig. 4).

As described earlier, we tested two sgRNAs targeting different coding regions and one targeting the proximal promoter (Fig. 1b, Supporting Information Fig. S1). Although, the efficiency of each sgRNA differed, all sgRNAs caused a similar phenotype, suggesting the phenotype is likely due to the mutations of *six3* gene but not due to mutations in off-target sites. We used 3.2-ng Cas9 mRNA and 200-pg sgRNA for F0 mutant characterization for single sgRNAs. This dose combination induced reproducible phenotypes as described in detail below in essentially all embryos injected in multiple experiments although the severity varied from one embryo to another. In another set of experiments, we attempted to delete an approximately 600bp *cis*-regulatory element, the *six3* proximal promoter, by designing CRISPR target sites flanking the region (Fig. 1b). When

CRISPR-mediated cleavage occurs at both sites surrounding this region, NHEJ may result in a deletion of the flanked region. For this experiment, 4.7 or 5.7 ng Cas9 mRNA and 200 pg of each sgRNA were injected, showing a consistent phenotype in up to 50% of the survivors (Fig. 4e). We could confirm that a deletion of approximately 600bp occurred in some cells of injected embryos as determined by PCR assay and subsequent cloning and sequencing of the targeted region (Fig. 4e and Supporting Information Fig. S1).

By st. 40, the majority of embryos injected with sgRNAs targeting *six3* coding regions (essentially 100% in multiple experiments for target 1, and 60–100% for target 2) showed reduction of eye size, although the severity varied (Fig. 4a, c). Importantly, this phenotype was partially rescued by coinjection of *X. laevis six3* mRNA (20 pg) as evidenced by shift of severity toward a milder phenotype (Fig. 4b, c). We confirmed that the rescued embryos were mutated by the DSP assay (Fig. 4d). By st. 42, the heads of mutants had a more angular and flattened shape from a lateral view (Fig. 4e,f, top panels) and narrower heads with smaller brain regions from the dorsal view (Fig. 4f, middle panel) or in a frontal section (Fig. 4f, bottom panel, black arrowhead) than wild-type embryos. When using the sgRNA for target 2, by st. 46 we also noticed a milder phenotype where the nasal pits were fused or shifted medially (not shown), and the telencephalon was smaller and fused in the mutant (Fig. 4g). Mutation profiles of mutant embryos described earlier are shown in Supporting Information Figure S1. In summary, multiple sgRNAs showed common phenotypes and the phenotypes could be significantly rescued by coinjection of mRNA, suggesting that the observed phenotypes are due to

FIG. 4. Summary of phenotype caused by mutations of *six3* gene. All targeted mutations showed a similar phenotype. (a–d) Phenotype variations seen in st.40 embryos targeted with the sgRNA for coding region site 1. By this stage, the obvious phenotype is significantly reduced eye size, but a brain defect (see below) is not yet obvious. The phenotype is *six3*-specific because it was partially rescued by coinjection of *six3* mRNA (b, c). Severity of phenotype was scored as +++ (most severe, no eye or tiny piece of eye), ++ (severe, small and malformed eye), + (modest, small relatively normal looking eye), and - (no or little phenotype) (a–c). As observed with the *tyr* target injections, toxicity of injected RNAs varied greatly depending on batch of embryos. For example, in different experiments, the percentage of normal embryos recovered after the same RNA doses were injected (*six3* target 1) varied from 43% to 87% (see Supporting Information Table S1). (d) Chromatograms of the DSP assay from an uninjected embryo (WT) and one of rescued embryos shown in (b). Perturbation of peaks is seen around the PAM region (two-headed red arrow), suggesting the presence of mutated sequences. At later stages, the brain phenotype becomes obvious (e–g). (e) Examples of the phenotype of embryos (emb1 to emb3) targeted to delete the proximal promoter region at st.42. Using the primer as schematically shown in Fig. 1b (green arrows), genomic PCR of sibling uninjected wild-type embryo showed an expected ~ 1.2 kbp band, whereas embryos injected with sgRNAs had an extra ~ 0.6 kbp band (emb1–3, white arrows). One band (marked by *) was re-cloned and sequenced; the results are shown in Supporting Information Fig. S1. The promoter deletion injections were on average more toxic, perhaps due to higher total amounts of RNAs being injected, but survival rates as high as 68% were observed in individual experiments. (f) Phenotype examples of st. 42 embryos injected with sgRNA targeting coding region site 1 (left panel) and site 2 (right panel), lateral views. Compared to wild-type embryos (bottom, WT, round shaped head), mutant embryo heads had a more angular and flattened shape. The dorsal view (second panel) and frontal sections (third panel) of the same embryos (emb1-1, 1-2, 1-3) in first left panel clearly show narrow heads (compare two-headed white arrows between WT and mutants) and smaller brains (black arrowheads in frontal sections), especially the forebrain (red arrowheads in second panel, see also Fig. 4g) in the mutant embryos. The sequencing summary of mutated loci of emb1-1 and 1-3 in the top left panel and emb2-1, 2-2 in the top right panel are shown in Supporting Information Fig. S1. (g) A mild phenotype seen in embryos (target site 2) at st. 46. Top panel shows a dorsal view of embryos. Second panel shows horizontal sections of equivalent embryos shown in the top panel. At early stages, these embryos could be scored as wild type. Later observation of mutant embryos reveals that the nasal pits (red arrows) are fused (shown here) or shifted medially (not shown), and the telencephalon (red arrowheads) is smaller and fused. The last panel shows chromatograms of the DSP assay from one of the sectioned embryos shown above to confirm genomic mutations as evidenced by perturbation of peaks around PAM region (two-headed red arrow) in the mutant embryo.

partial inactivation of the *six3* gene but not due to mutations of off-target sites. Also, importantly, we successfully showed that by simultaneously targeting two sites surrounding the promoter, the flanked region (up to 600 bp in this experiment) could be deleted. This strategy is particularly useful for studying promoter and enhancer regions of a gene.

In our experiments, as noted in Figure legends 2 and 4, there is some impact of injecting CRISPR RNAs on embryo survival, and we consider possibilities here to explain these results to help other investigators who may use this technology. Three obvious possibilities are: egg quality, toxicity of RNAs, and off-target effects of CRISPR RNAs. We are certain that egg quality issues come into play, since the same constructs are found to result in very different survival rates with eggs from different females (see Supporting Information Table S1). In injections involving all target sequences used in this work where there were low survival rates in one experiment (e.g., as low as 50%), there were other experiments where survival was significantly higher (e.g., 80%). To examine whether individual RNAs are themselves toxic, we injected either Cas9 or sgRNA's alone at doses used in targeting experiments, and there we found significant mortality (up to 50%) in some egg batches where higher doses were used, suggesting that RNA toxicity is present to a significant degree. In addition to our multiple target and rescue strategies to minimize the chance of off target effects, both egg batch and single RNA injection results argue against off-target effects as an explanation for the lethality we see, since the latter is variable (and can be quite low) and can be elicited without biologically active CRISPR function (i.e., embryos injected with Cas9 or sgRNA alone).

As shown in the accompanying paper (Blitz *et al.*, 2013), one can bioinformatically identify and systematically evaluate off-target effects experimentally; for the *tyr* target studied, these are not significant. The same *tyr* target was used in our study (target 1), along with a second target that also induced albinism. In our experiments off-target effects were studied in two ways. First, possible target sites were evaluated bioinformatically during the design process for both *tyr* and *six3* (see Methods) to eliminate any target that might have identical sequence or a single base mismatch in sequence elsewhere in the genome. Second, although our data strongly suggests we are not seeing off-target effects in our experiments for reasons noted earlier, we also searched for off-target sequences with up to four base mismatches in all of the targets used in our studies of *six3*, and have found none in coding sequences in genomic sequence (Supporting Information Table S2).

Here, we have demonstrated for two genes that CRISPR-mediated mutagenesis is useful for quickly analyzing loss-of-function phenotypes in genes of interest

that function at later stages of development when conventional MO-type loss-of-function assays are not feasible. Because of mosaicism of mutant and wild-type cells in a whole embryo, we are probably seeing hypomorphic phenotype(s), which is actually an advantage of using F0 animals because we might not see subtle gene effects in the case of complete inactivation of a gene in an established mutant line. Conversely, because of the mosaicism, phenotype variation could be significant from one animal to another. Therefore, we need to be careful in concluding that observed phenotypes are due to inactivation of the gene of interest. For detailed, precise analysis of loss-of-function phenotypes, mutant lines should be established, for which CRISPR methodology will be a very powerful and efficient tool. For the creation of mutant lines that may cause embryonic lethal phenotypes in F0 animals with high mutation rates, the use of lower doses of RNAs or use of the less efficient, bacterial Cas9, may be advantageous. Furthermore, the sensitivity of the DSP assay is sufficient to identify nonphenotypic putative founders, and can later be used to genotype heterozygous offspring (e.g., see Supporting information Fig. S2). Maintenance of such lines through out-crossing also serves to breed out putative off-target mutations. In the future, more detailed analysis of *six3* null mutant(s) will be performed using established mutant line(s) where the value of consistent, heritable phenotypes provides additional advantages over the more variable phenotypes (and issues like toxicity of injected embryos) that one sees in F0-injected embryos.

We should keep in mind that the availability of CRISPR target sequence is somewhat constrained and thus there will be situations where one needs TALENs for gene editing, since they have more flexibility in selection of target sequences. Additionally, the increased possibility of off-target effects when using CRISPR to study F0 phenotypes may be problematic; however, the guidelines presented here, that is, targeting different regions of the same gene as well as performing rescue experiments when they are feasible, should help address these concerns. Also, as has been shown by Blitz *et al.* (2013), the off-target sites may not be problematic in *Xenopus* for some target sequences, since they could not find evidence for mutations of possible off-target sites for the targets site 1 of *tyr* sgRNA. Furthermore, as suggested earlier, over the course of the creation of stable mutant lines, any off-target mutations will be bred out, and such lines will be powerful tools for studying gene function. The clear advantages of the CRISPR-mediated genome editing method, including the ability to easily and quickly design oligonucleotides to create sgRNA templates, as well as effect targeted deletions of larger regions of regulatory DNA such as promoters and enhancers, makes this method an important part of the growing *Xenopus* genetic-manipulation toolkit.

during the 2013 Cold Spring Harbor Laboratory *Xenopus* Course, which authors would like to acknowledge. Authors also thank Ira Blitz and Ken Cho for sharing information before publication. The authors would also like to gratefully acknowledge the support of the National *Xenopus* Resource and its staff. Due to the rapid accumulation of reports using the same methodologies and space limitations, authors apologize for not being able to cite all relevant references. The plasmid pDR274-Xt-tyr described in the study can be obtained from GHT (gerald.h.thomsen@stonybrook.edu) and RMG (rmg9p@virginia.edu). pCS2-Cas9 is available from RMG.

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