Introduction

The past decade has witnessed remarkable advances in genome editing. A series of genome editing tools such as ZFNs, TALENs and Cas9 have been developed that enable investigators to manipulate virtually any gene in a wide range of cell types and organisms. Both ZFNs and TALENs have almost the same mode of action with their DNA recognition domains interacting with nucleotides and nuclease catalytic domains producing DNA double strand breaks (DSBs). In the CRISPR/Cas9 system, a Cas9 protein is guided by a synthetic single guide RNA (sgRNA) consisting of a fusion of crRNA and tracrRNA to the target locus to produce cleavage at a PAM motif (NGG) [1]. The induced DSBs can be repaired by the NHEJ system, which in turn generates various mutations including insertion, deletion or even translocation at the target loci [2]. These technologies have been successfully applied in eukaryotic cells and model organisms for the disruption of target genes [3]. However, these genome editing tools are not perfect and about 40–60% of the target loci cannot be efficiently mutated, which calls for sensitive and reliable methods to evaluate the effectiveness of specific targets.

Currently, several assays have been developed to detect the mutations of target loci. The most frequently used assay is the hetero-duplex analysis, which utilizes SURVEYOR nuclease [4], Cel-I [5,6] or T7E1 [7,8,9] to recognize and cut the hetero-duplex of mutant and wild-type DNA fragments containing mismatched base pairs. In addition, the hetero-duplex can be analyzed by high resolution melt curve analysis (HRMA) [10,11,12,13]. Restriction enzyme assay is another frequently used method to evaluate TALEN and Cas9 activity when a target region contains a suitable restriction enzyme site [14,15]. Furthermore, the LacZ recovery/disruption assay has been developed to assess the frame shift mutation resulted in the recovery of lacZ activity by blue/white colony screening [16]. However, current assays have their limitations in that they either require specific sequence in the target sites or are unable to generate sequencing-ready mutant DNA fragments or unable to distinguish induced mutations from natural nucleotide polymorphism.

To overcome these limitations, we have developed a simple PCR-based technique to detect the induced mutations. It is well known that the mutant positions induced by these nucleases are largely predictable, as ZFNs and TALENs induce genomic DSBs in the center of spacer region between their DNA recognition sites and Cas9 induce DSBs in proto-spacer adjacent motif (PAM) [1]. Taking advantage of this, we designed 2 pairs of primers for each TALEN or Cas9 target site, with one putative amplicon extending beyond the putative indel site and the other overlapping it. Thus, the newly introduced mutations will disrupt the amplification of the later amplicon but will not affect that of the first one. By using quantitative PCR (qPCR), we demonstrated that the efficiency of desired mutations could be quantitatively evaluated. And coupled
with a T-vector-based mutant colonies identification assay (T-CIA), it could easily generate single-copy mutant DNA fragments for subsequent sequencing. Thus, our work provides a very simple, efficient and fast assay for detecting induced mutations, which we anticipate will be widely used in the area of genome editing.

**Materials and Methods**

**Ethics Statement**

The treatment and handling of zebrafish was performed under general directives on the protection of animals used for scientific purposes and following operating procedures approved by Sichuan Animal Care and Use Committee, Permit Number: SYXK (Chuan) 2009-119. Detailed treatment of adult zebrafish and embryos were described in the “DNA isolation and template preparation” section.

**Zebrafish Care**

All the Zebrafish used in our experiment were AB strain. Adult fish and embryos were raised and maintained as described Westerfield et al [17,18]. Embryos were cultured in 10 cm petri dishes at 28.5°C. Adult zebrafish were maintained in the standard tank of an automatic fish housing system (ESEN, China). Healthy 3–5 months old adult female and male fish were maintained in separate tanks and mated once a week.

**TALEN Target Site Design and Unit Assembly**

TALEN target sites were designed with an online tool, TALE-NT (https://tale-nt.cac.cornell.edu/). Then, TALEN units were assembled using methods described in Huang et al. [19] and Zu et al. [20]. Briefly, the units were assembled to construct the TALE repeats according to the target sequences by several rounds of digestion/ligation cycles. Each round of digestion/ligation cycle, the recombinant vectors were transformed into *Escherichia coli* Top10 for amplification. The constructed iterative TALE constructed fragments were then digested and inserted into TALEN expression vectors pXt7-TALEN upstream of FokI. The TALEN target sites were listed in Table S1 and Figure S1.

**Cas9 Target Site Design and Vector Construction**

Cas9 target sites were designed with an online tool, ZIFIT Targeter (http://zifit.partners.org/zifit/Introduction.aspx). The vectors, pCS2-Cas9 and pT7-gRNA, for making Cas9 mRNA and chimera guide RNA were kind gifts from Jingwei Xiong and Bo Zhang (Peking University). The target site sequences were in vitro synthesized and inserted between the T7 promoter and

![Figure 1. Schematic diagrams showing a PCR-based protocol for identifying mutations induced by TALEN and Cas9 in zebrafish.](https://plosone.org/doi/10.1371/journal.pone.0098282.g001)
guide RNA (sgRNA) scaffold sequence of pT7-gRNA [21]. The Cas9 target sites were listed in Table S1 and Figure S1.

mRNA Synthesis
For mRNA synthesis, both the TALEN and Cas9 expression vectors were linearized and then transcribed in vitro with the mMESSAGE mMACHINE T7 ULTRA kit (Ambion) following manufacturer’s instructions. RNAs were purified by LiCl acetate precipitation and re-dissolved in RNase-free water. The RNA quality and concentration were analyzed by electrophoresis and nucleic acid spectrometer respectively.

Microinjection of Zebrafish Embryos
One-cell stage embryos were collected for microinjection by using an electronically regulated air-pressure micro-injector (Harvard Apparatus, NY, PL1-90). For TALEN injection, equal amounts of the Left and Right arm mRNAs were mixed and injected together into the yolk of the embryos. For CRISPR/Cas9 injection, Cas9 mRNA and gRNA were mixed at a ratio of 5:3.

DNA Isolation and Template Preparation
Genomic DNA was extracted using the alkaline lysis method. For adult zebrafish DNA extraction, tails were cut from fish anaesthetized with 0.4 mg/ml tricaine solution (Sigma-Aldrich), and the fish were put back to the tank immediately after tail cutting. For embryonic DNA extraction, 5 embryos at 2 days post fertilization were pooled into each PCR tube. Embryos or tail fins were lysed in 30 µl alkaline lysis buffer (25 mM NaOH, 0.2 mM EDTA) and heated at 95°C for 10 minutes. Then, DNA solution was neutralized by adding 1/10 volume neutralization buffer (40 mM Tris-HCl, pH 8.0). Samples were spun at 10,000 rpm for 5 minutes and the supernatant was transferred into new tubes.

qPCR Analysis for Identifying Mutations
We analyzed a panel of published mutation sites induced by TALEN and Cas9, and found that most mutations affected a 4 to 5 base region around the DSBs. Accordingly, we designed 2 pairs of primers: the first pair of primers were designed to prime outside the DSBs and the other flanked the DSBs at the 3’ most nucleotides. Thus, newly induced mutation will disrupt the amplification of the latter pair of primers. Before performing qPCR, we ran an ordinary PCR to obtain high quality DNA fragments using the outside primers, and these fragments were quantified and adjusted to equal amounts for subsequent qPCR. qPCR was performed with SsoAdvanced SYBR Green Supermix (Bio-Rad) by using a Bio-Rad CFX96 Real-Time system. All the primers were listed in Table S2. The qPCR conditions were programmed as follows: 40 cycles of 10 s at 95°C and 20 s at 60°C after initial denaturing for 30 s at 95°C.

Isolation of Monocolonies Harboring Induced Mutations
The products from outside primers of qPCR were directly inserted into T-clone vector pMD19-T (Takara) and transformed into Escherichia coli Top10 for white-blue screening. Then the white
colonies were analyzed by PCR using primers as described in Table S2. Those colonies that could not generate inside PCR fragments were the putative mutant ones which were further analyzed by DNA sequencing.

Results and Discussion

Mutations caused by TALEN or Cas9 are largely predictable as the induced mutant loci were produced at a fixed target sequence. Most of the indel mutations occurred at the DSBs sites, referred as mutational hot spot regions (MHS). An analysis of several hundreds of the published mutant sequences revealed that most TALEN induced mutations affected the middle 4–5 base pairs of the spacer regions and Cas9 induced mutations affected the 4 base pairs upstream of the PAM ( counting 5' to 3' from the PAM). Specifically, in TALEN-induced mutations, 70.18% affected all the 4–5 base pairs, 11.47% affected 3 of them, 7.34% affected 2 of them, and 6.57% affected 1 of them (n = 654) (Document S1). In Cas9 induced mutations, 42.67% affected all the 4 base pairs, 17.48% affected 3 of them, 14.91% affected 2 of them, and 20.31% affected 1 of them (n = 389) (Document S2).

Because a highly complementary 3-prime end of the PCR primer is critical for amplification, mismatches in the 3’ end will greatly reduce the amplification efficiency. This phenomenon is widely used in PCR-based genotyping methods for detecting genetic polymorphism or point mutation [22,23]. Theoretically, the induced indel mutations will cause complete or partial loss of the PCR amplification if the 3’ end of primers were designed to exactly cover MHSs. In view of this hypothesis, we designed a simple PCR based protocol to detect the expected mutations (Figure 1). As shown in Figure 1A and Table S2, one pair of primers were designed to prime outside the MHS, referred as Primer outside (Po), and the other flanked the MHS at the 3’ most nucleotides, referred as Primer flank (Pf). Thus, Po can amplify both wild-type and mutant alleles, while Pf can only amplify wild-type alleles, and any newly arising mutation will disrupt the amplification of the Pf. Technically, mutation frequency can be easily calculated by the decreased ratio between Pf amplicons and Po amplicons.

As a first step to test the hypothesis, we examined whether this method could identify and quantify an already established 4 bp deletion mutation in the zebrafish ldlr gene. We diluted the mutant and wild-type templates to equal concentration and mixed them in various ratios ranging from 0% to 100%, herein referred as putative ratios. Then we used ldlr-Po primers that are mutation insensitive as an internal control and ldlr-Pf primers that are mutation sensitive as a test. As expected, the amplification efficiencies of ldlr-Pf were gradually reduced along with the

Figure 3. Identification of mutations induced by ldlr TALEN. (A) A schematic diagram showing primers for detecting mutations in TALEN target sites of ldlr gene. (B) The relative amplification efficiency of ldlr Pf primers to Po primers. ldlr TALEN mRNAs injection dramatically reduced the amplification efficiency of ldlr Pf primers. 5 embryos from the wild-type group or the ldlr-TALEN-injected group were pooled and analyzed by qPCR, and 3 pools of each group were analyzed in an independent experiment. Similar results were obtained in three independent experiments. (C) Identification of a mutation in a single allele with T-CIA. Upper panel was an agarose electrophoresis result of PCR products amplified with ldlr Po and Pf primers. Lower panel listed the sequences of mutant colonies identified in upper panel.

doi:10.1371/journal.pone.0098282.g003
increasing ratios of mutant templates in the mixtures (Figure 2A). More importantly, these data showed that qPCR could detect at least 10% mutation in the template which is much lower than the efficiency of most reported TALEN and Cas9 induced mutations. However, the amplification efficiencies of Po primers were gradually increased along with the increasing ratios of mutant templates in the mixtures (Figure 2A), and the calculated ratios of mutant fragments by qPCR were greater than the putative ratios (Figure 2B). The deviation could be caused by inaccurate measurement of the concentration of original templates or from errors during dilution and mixing. Nevertheless, this is not an issue when using this method in detecting induced mutations, because we do not need to mix the wild-type and the putative mutant templates in such situations.

Next, we tested this method in detecting new mutations at several designed TALEN or Cas9 target sites. We injected one cell-stage zebrafish embryos with either TALEN mRNAs or an RNA mixture of Cas9 mRNA and sgRNAs. Genomic DNA was prepared from a pool of 5 injected or wild-type embryos and analyzed by qPCR. As shown in figure 3B, ldr2 TALEN mRNA injection greatly reduced the amplification of the Pf primers but not that of the Po primers. The normalized efficiency of ldr2 TALENs was about 91.6%. Figure 4B showed the qPCR results of a Cas9 targeting the ldr2 gene, in which the normalized efficiency was about 44.1%. We also used this method to successfully test 2 additional TALEN or Cas9 target sites, apoeb and nsd3 (Figure S2B and S1B). At the same time, we also confirmed the presence of indel mutations in ldr2 and nsd3 loci with restriction endonuclease assays, and observed similar mutant efficiency (Figure S3). Thus, these results established that the qPCR assay can reliably calculate the efficiency of TALEN or Cas9 induced mutations.

In order to further confirm the reliability of this method and to generate single-copy mutant DNA fragments for subsequent sequencing, we cloned the PCR products of the injected embryos into T-cloning vectors, pMD19-T. About 10 colonies of each target site were examined by ordinary PCR using the corresponding Po and Pf primers. As shown in Figure 3C, 4C and Figure S1C, all the white colonies can be amplified by the Po primers while some of them cannot be amplified by the Pf primers, indicating that these colonies harbored mutant DNA fragments. To confirm this, we selected several colonies that possess only Po fragment for sequencing and found that almost all the positive colonies had indel mutations in the target sites (Figure 3C and 4C). Together, our data suggested that this PCR-based assay is indeed
affective and sensitive for both determining mutation efficiency and identifying mutant sequences.

Then, we tested if this method could be utilized to detect germ line transmitted mutations. TALEN or Cas9 mRNA injected embryos were raised to adulthood (referred as F0) and outcrossed with wild-type fish. Pools of 5 F1 embryos were lysed and subjected to qPCR analysis. In Figure S4 A and C, we identified one out of four fish that had germ line mutation in the nsd2 gene, and all 3 F0 fish examined had germ line mutation in ldlr gene. Again, the offspring of these F1 fish were raised to adulthood and were genotyped by qPCR using DNA extracted from tail fins (Fig. S4 B and D). Interestingly, all the F1 fish analyzed were heterozygous mutant for nsd2 or ldlr gene. Then the mutant allele of each F1 fish was analyzed by T-CIA (data not shown). Thus, this method is practical for detecting germ line transmitted mutations as well.

In summary, we have developed a simple PCR-based method for detecting indel mutations induced by TALEN and Cas9 in zebrafish. We demonstrated that qPCR assay could specifically detect indel mutations arising at designed target sites, and results from this assay should not be disturbed by natural polymorphism in the genome. In addition, this assay does not require a restriction site, thus it does not bias the choice of a target site. Furthermore, we also provided a T-vector-based assay for identifying mutant sequences of the target loci. However, theoretically, this assay is less efficient in detecting mutations in regions containing repeated nucleotides, especially mono-nucleotide repeats. It is also not efficient in detecting mutations that do not happen in the predicted MHS. But besides these situations, this assay is highly efficient. Considering that it could be used in the entire procedure of nuclease-mediated genome editing, including the evaluation of the efficiency of a given TALEN or Cas9, and the identification of founder or F1 fishes, we hope this assay will be widely used in the field of in genome editing technologies.

Supporting Information

Figure S1 The designs of TALENs and Cas9 targeting regions in zebrafish. (A) and (B) were using TALENs to disrupt ldlr and apoeb genes respectively. (C) and (D) were using Cas9 to disrupt nsd2 and nsd3 genes respectively.

Figure S2 qPCR identify mutations induced by NSD3 Cas9. (A) A chemic diagram showing primers for detecting mutations in Cas9 target site of nsd3 gene. (B) The relative amplification efficiency of nsd3 PI primers to Po primers. nsd3 cas9 mRNAs injection dramatically reduced the amplification efficiency of nsd3 PI primers. (C) Identification of mutation in single allele with T-CIA. Upper panel was agarose electrophoresis result of PCR products amplified with nsd3 Po and PI primers. Lower panel listed the sequences of mutant colonies identified in upper panel. Every 5 embryos served as a pool for qPCR. The Data in B were obtained from 3 independent experiments, with 3 replicates for each pool.

Figure S3 qPCR identify mutations induced by TALEN. (A) A chemic diagram showing primers for detecting mutations in TALEN target site of apoeb gene. (B) The relative amplification efficiency of apoeb PI primers to Po primers. apoeb TALEN mRNAs injection dramatically reduced the amplification efficiency of nsd3 PI primers. (C) A list of the mutant sequences of TALEN target site in apoeb gene. Every 5 embryos served as a pool for qPCR. The Data in B were obtained from 3 independent experiments, with 3 replicates for each pool.

References

1. Van Der Oost J (2013) New Tool for Genome Surgery. Science 339: 768–770.
2. Haber JE (2000) Recombination: a frank view of exchanges and vice versa. Curr Opin Cell Biol 12: 216–292.
3. Gaj T, Gersbach CA, Barbas III CF (2013) ZFN, TALEN, and CRISPR/Cas-based methods for genome engineering. Trends Biotechnol 31(7): 397–405.
4. Cong L, Ran FA, Cox D, Lin S, Barretto R, et al. (2013) Multiplex Genome Engineering Using CRISPR/Cas Systems. Science 339: 819–823.
5. Liu X, Wang YS, Guo W, Chang B, Liu J, et al. (2013) Zinc-finger nickase-mediated insertion of the lysostaphin gene into the beta-casein locus in cloned cows. Nat Commun 4: 2363.
6. Lo TW, Pickle CS, Lin S, Ralston EJ, Gurung M, et al. (2013) Precise and Heritable Genome Editing in Evolutionarily Diverse Nematodes Using TALENs and CRISPR/Cas9 to Engineer Insertions and Deletions. Genetics 195: 331–348.

7. Kim HJ, Lee HJ, Kim H, Cho SW, Kim JS (2009) Targeted genome editing in human cells with zinc finger nucleases constructed via modular assembly. Genome Res 19: 1279-1288.

8. Kim H, Um E, Cho SR, Jung C, Kim H, et al. (2011) Surrogate reporters for enrichment of cells with nuclease-induced mutations. Nat Methods 8: 941–943.

9. Cho SW, Kim S, Kim JM, Kim JS (2013) Targeted genome engineering in human cells with the Cas9 RNA-guided endonuclease. Nat Biotechnol 31(3): 230–232.

10. Dahlem TJ, Hoshijima K, Jurynec MJ, Gunther D, Starker CG, et al. (2012) Simple Methods for Generating and Detecting Locus-Specific Mutations Induced with TALENs in the Zebrafish Genome. PLoS Genet 8(8): e1002861.

11. Xing L, Hoshijima K, Gruenwald DJ, Fujimoto E, Quist TS, et al. (2012) Zebrafish foxP2 Zinc Finger Nuclease Mutant Has Normal Axon Pathfinding. PLoS ONE 7(8): e43968.

12. Bassett AR, Tibbit C, Ponting CP, Liu JL. (2013) Highly Efficient Targeted Mutagenesis of Drosophila with the CRISPR/Cas9 System. Cell Rep 4(1): 220–228.

13. Ota S, Hisano Y, Muraki M, Hoshijima K, Dahlem TJ, et al. (2013) Efficient identification of TALEN-mediated genome modifications using heteroduplex mobility assays. GENES CELLS 18(6): 450–458.

14. Nekrasov V, Staskawicz B, Weigel D, Jones JDG, Kamoun S (2013) Targeted mutagenesis in the model plant Nicotiana benthamiana using Cas9 RNA-guided endonuclease. Nat Biotechnol 31(5): 691–693.

15. Liang Z, Zhang K, Chen K, Gao C (2013) Targeted Mutagenesis in Zea mays Using TALENs and the CRISPR/Cas System. J Genet Genomics 41(2): 63–68.

16. Hisano Y, Ota S, Arakawa K, Muraki M, Kono N, et al. (2013). Quantitative assay for TALEN activity at endogenous genomic loci. Biology Open 2: 363–367.

17. Kimmel CB, Ballard WW, Kimmeld SR, Ullmann B, Schilling TF (1995) Stages of embryonic development of the zebrafish. Dev Dyn 203: 233–310.

18. Westerfield M (2000) The Zebrafish Book: A guide for the laboratory use of zebrafish (Danio rerio). 4th ed. Eugene: Univ of Oregon Press.

19. Huang P, Xiao A, Zhen M, Zhu L, Lin S, et al. (2011) Heritable gene targeting in zebrafish using customized TALENs. Nat Biotechnol 29: 699–700.

20. Zu Y, Tong X, Wang Z, Liu D, Pan R, et al. (2013) TALEN-mediated precise genome modification by homologous recombination in zebrafish. Nat Methods 10: 329–333.

21. Chang N, Sun C, Gao L, Zhu D, Xiong JW, et al. (2013) Genome editing with RNA-guided Cas9 nuclease in Zebrafish embryos. Cell Res 23: 465–472.

22. Rigat B, Hubert C, Corvol P and Soubrier F (1992) PCR detection of the insertion/deletion polymorphism of the human angiotensin converting enzyme gene (DCP1) (dipeptidyl carboxypeptidase 1). Nucl Acids Res 20(6): 1433.

23. Orum H, Nielsen PE, Egholm M, Berg RH, Buchhardt O, et al. (1993) Single base pair mutation analysis by PNA directed PCR clamping. Nucl Acids Res 21(23): 5332–5336.