ABSTRACT

CRISPR/Cas9-based tissue-specific knockout techniques are essential for probing the functions of genes in embryonic development and disease using zebrafish. However, the lack of capacity to perform gene-specific rescue or live imaging in the tissue-specific knockout background has limited the utility of this approach. Here, we report a robust and flexible gateway system for tissue-specific gene inactivation in neutrophils. Using a transgenic fish line with neutrophil-restricted expression of Cas9 and ubiquitous expression of single guide (sg)RNAs targeting rac2, specific disruption of the rac2 gene in neutrophils is achieved. Transient expression of sgRNAs targeting rac2 or cdk2 in the neutrophil-restricted Cas9 line also results in significantly decreased cell motility. Re-expressing sgRNA-resistant rac2 or cdk2 genes restores neutrophil motility in the corresponding knockout background. Moreover, active Rac and force-bearing F-actins localize to both the cell front and the contracting tail during neutrophil interstitial migration in an oscillating fashion that is disrupted when rac2 is knocked out. Together, our work provides a potent tool that can be used to advance the utility of zebrafish in identifying and characterizing gene functions in a tissue-specific manner.

KEYWORDS: Actin stress, Cell migration, Leukocytes, Live imaging, Rac2

INTRODUCTION

Over the past decade, zebrafish (Danio rerio) has gained popularity as a vertebrate model organism for biological and biomedical studies, including neutrophil biology (Deng and Huttenlocher, 2012). Transparent embryos, a short life cycle (Driever et al., 1994), a highly conserved innate immune system (Lieschke and Trede, 2012), as well as ease of genetic manipulation (Lawson and Wolfe, 2011), allow for the dissection of mechanisms regulating neutrophil migration using both genetics and non-invasive high-resolution intravital imaging approaches.

In general, generating tissue-specific knockouts is essential to delineate the function of genes-of-interest in different cells and tissues. However, this technique is not sufficiently developed in the zebrafish model. The Cre/loxP site-specific recombination technology is one of the earliest conditional gene modification approaches (Hoess and Abremski, 1985) and is widely applied in mice (Branda and Dymecki, 2004). The first Cre/loxP system in zebrafish was developed in 2004, by injecting Cre RNA into embryos of a floxed gfp transgenic line (Dong and Stuart, 2004). Indeed, several studies demonstrated the feasibility of using the Cre/loxP system for tissue-specific gene inactivation in zebrafish (Cantu et al., 2018; Hans et al., 2009; Langenau et al., 2005; Pan et al., 2005; Thummel et al., 2005; Xiong et al., 2013). However, making floxed alleles at the endogenous loci is technically challenging and time consuming. A recent advance in generating the conditional allele could expand its utility in the future (Li et al., 2019a). Meanwhile, the gene-silencing approach using RNAi has only been shown to be successful in limited circumstances (de Rienzo et al., 2012; Dong et al., 2009; Kelly and Hurlstone, 2011), possibly due to a lack of reliable methods to express small interfering RNAs in zebrafish tissues (Oates et al., 2000; Wang et al., 2010; Zhao et al., 2001).

Over the past decade, the prokaryotic clustered regularly interspaced short palindromic repeats (CRISPR) and the CRISPR-associated system (Cas) (CRISPR/Cas system)-based technology has been successfully used in zebrafish to efficiently generate insertions and indels to disrupt gene function (Varshney et al., 2015). The Zon group designed a CRISPR-based vector system that enabled tissue-specific gene inactivation in zebrafish (Ablain et al., 2015). They later incorporated this system into the MiniCoopR (Ceol et al., 2011) vector and developed the CRISPR MiniCoopR vector for melanocyte-specific gene disruption and validated Spred1 loss as a driver of mucosal melanoma in zebrafish (Ablain et al., 2018). A similar approach has been used to disrupt the androgen receptor gene in zebrafish liver to determine its contribution in hepatocellular carcinoma (Li et al., 2019b). di Donato et al. (2016) used the Gal4-UAS system to control the expression of Cas9 and achieved tissue-specific gene disruption. The Chen group used different U6 promoters for multiplex single guide (sg)RNA expression and also demonstrated tissue-specific disruption (Yin et al., 2015). Our group incorporated the Zon and Chen methods into a gateway system to express multiple sgRNAs for gene disruption in neutrophils in zebrafish (Zhou et al., 2018). Although high efficiency is achieved, a major limitation is also observed: knockout efficiency reduces significantly when the knockout line is crossed with other fish lines that use neutrophil-specific promoters. This limitation created two
problems: (1) without a sgRNA-resistant rescue construct, the specificity of the sgRNAs and the related phenotype cannot be concluded; and (2) incorporating biosensors into the knockout lines requires generating additional Cas9-2a-sensor lines, thereby limiting the flexibility of incorporating additional genetically encoded probes as projects evolve. We also found, to the best of our knowledge, that tissue-specific gene rescue and biosensor imaging in a tissue-specific knockout background have not yet been achieved in any previous work in zebrafish.

Here, we report an updated CRISPR/Cas9 system for a robust and flexible neutrophil-restricted knockout in zebrafish. We successfully disrupted different genes in neutrophils and applied live imaging using various biosensors in the knockout background. As a proof-of-principle, we inactivated the rac2 gene. Rac2 is essential for actin polymerization, cell migration and intracellular signaling. Loss of Rac2 activity leads to defects in neutrophil motility and chemotaxis in zebrafish (Deng et al., 2011; Rosowski et al., 2016). However, it remains unknown whether Rac activation is restricted to the front of the cell during neutrophil migration in vivo. The specific functions of Rac2, in comparison to its homologue Rac1 that is also expressed in neutrophils, are also not clear. Using multiple biosensors, we observed that Rac activation and force bearing actin structures are localized to the cell front and back in zebrafish neutrophils in an oscillating fashion, and the localizations are dependent on Rac2. Together, our system here provides a robust tool for discovering and characterizing genes that regulate neutrophil migration in vivo.

RESULTS

A Gateway cloning system for a tissue-specific knockout

Our previous strategy was to express Cas9 tagged with mCherry specifically in neutrophils and the sgRNA ubiquitously to achieve a neutrophil-specific knockout (Zhou et al., 2018). We speculate that the presence of another construct driven by neutrophil-specific promoters in the genome may compete with the transcriptional factors for Cas9-2A-mCherry expression and reduces Cas9 protein to a level that is not sufficient for an efficient knockout. On the contrary, several studies incorporated the same tissue-specific promoter multiple times and still achieved sufficient target gene disruption using untagged Cas9 (Ablain et al., 2015; Ceol et al., 2011; Li et al., 2019b). Thus, we decided to use the untagged Cas9. We designed two different plasmids to express the Cas9 and sgRNA in two separate lines. To generate a final plasmid construct for cell-specific Cas9 expression, we used three entry plasmids, containing, respectively, the neutrophil-specific promoter lysozyme C (lyzC), Cas9 with nuclear localization sequences, SV40 polyA and a destination Tol2 vector with a GFP reporter gene driven by the α-crystallin (cry, also known as cryaa) promoter. GFP+ lenses enable the selection of zebrafish with stable genomic integration (Fig. 1A). To introduce ubiquitous sgRNA expression, we used a plasmid harboring a GFP reporter gene controlled by the lyzC promoter, and two gene-specific sgRNAs driven by the zebrafish RNA polymerase (RNAP) III-dependent U6 promoters (U6a and U6c) (Fig. 1B). The successful incorporation of sgRNA sequences into the zebrafish genome can be visualized by GFP expression in neutrophils.

To test the gene knockout efficiency, we injected the F2 embryos of the newly generated Tg(lyzC:cas9, cry:GFP)pu26 line with the plasmids carrying rac2-targeting sgRNAs or control (ctrl) sgRNAs for transient gene inactivation. The sequences of the sgRNAs are described in Fig. 1C,D. A longer sequence with no predicted binding sites in the zebrafish genome was used as a non-targeting control (Fig. 1D). As expected, we observed significantly decreased neutrophil motility in larvae of Tg(lyzC:cas9, cry:GFP)pu26 fish transiently expressing sgRNAs targeting rac2 (Fig. 1E,F; Movie 1), consistent with a functional disruption of the rac2 gene. To test the knockout efficiency, we generated two transgenic lines, Tg(U6a/c: ctrl sgRNAs, lyzC:GFP)pu27 and Tg(U6a/c: rac2 sgRNAs, lyzC:GFP)pu28. The F1 fish were crossed with
Tg(lyzC:cas9, cry:GFP)mu26, and the velocity of neutrophils in the head mesenchyme was quantified in embryos at 3 days post fertilization (dpf). As expected, a significant decrease of motility was observed in the neutrophils expressing Cas9 protein and the rac2 sgRNAs (Fig. 2A,B; Movie 2). To make sure that the sgRNA expression alone does not influence neutrophil motility, we compared neutrophil motility in the transgenic lines Tg(U6a/c: ctrl sgRNAs, lyzC:GFP)mu22 or Tg(U6a/c: rac2 sgRNAs, lyzC:GFP)mu26 with that in Tg(lyzC:GFP) (Hall et al., 2007). All lines displayed similar neutrophil motility, indicating that the migration defects are dependent on the expression of Cas9 in neutrophils (Fig. 2C,D; Movie 3). The lyzC is a well-characterized promoter for driving gene expression in neutrophils (Kitaguchi et al., 2009), albeit with a much lower expression level in macrophages. As Rac2 is also required for macrophage migration in tissue (Rosowski et al., 2016), we measured the speed of macrophage migration to infer the function of rac2 in macrophages. To confirm that the Cas9-mediated depletion was neutrophil-specific, we bred in a macrophage reporter line Tg(mpeg:mcherry-H2B) (Davis et al., 2016) into the two lines described in Fig. 2A,B. Although neutrophil motility remained deficient (Fig. 2E,F), macrophage migration was intact (Fig. 2E,G; Movie 4), indicating that rac2 inactivation in macrophages was not significant.

To demonstrate the editing of the rac2 locus in neutrophils, we crossed Tg(LyzC:Cas9, Cry:GFP)mu26 with Tg(U6a/c: ctrl sgRNAs, lyzC:GFP)mu27 or Tg(U6a/c: rac2 sgRNAs, LyzC:GFP)mu28, and the neutrophils from the 3 dpf larvae were enriched using fluorescence-activated cell sorting. The rac2 sgRNA1 and sgRNA2 target sites were amplified and the PCR products were treated with T7 endonuclease 1 (T7E1). In neutrophils sorted from the rac2 sgRNA expressing lines, sufficient genome editing was detected at the sgRNA1 targeting site, but not at the sgRNA2 targeting site (Fig. 2H). No editing was detected in the sorted neutrophils expressing control sgRNAs. To further demonstrate the specific gene disruption in neutrophils, we used a T7E1 assay to check the sgRNA target sites in the GFP+ non-neutrophil population. We did not observe significant editing (Fig. 2I), indicating that the rac2 disruption is primarily restricted to neutrophils. We further deep sequenced these loci and detected a mutation efficiency of 34.73% at the sgRNA1 target site and 17.16% at the sgRNA2 target site (Fig. 2J), whereas in the GFP+ non-neutrophil population, a similar level of editing was not detected.


tools and resources

Expression of sgRNA-resistant rac2 rescued the neutrophil migration defect

A gold standard to confirm that a particular phenotype results from a specific gene disruption is to perform a gene-specific rescue. We therefore replaced the GFP gene with a sgRNA-resistant rac2 gene in the PME entry plasmid described in Fig. 1B and constructed the final plasmid to allow co-expression of the rescue gene and the sgRNAs (Fig. 3A). Restoring the expression of wild-type Rac2 rescued the neutrophil motility defect seen in embryos with neutrophil-specific rac2 disruption (Fig. 3B,C; Movie 5), indicating that the motility

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**Fig. 2. Neutrophil-specific knockout of rac2 in stable transgenic lines.** Stable lines were generated by crossing Tg(LyzC:Cas9, Cry:GFP)mu26 with Tg(U6a/c: ctrl guide, LyzC:GFP)mu22 or Tg(U6a/c: rac2 guides, LyzC:GFP)mu28. (A,B) Representative images (A) and quantification (B) of neutrophil motility in the head mesenchyme of 3 dpf larvae, n=55 for control from four different larvae and n=60 for rac2 knockouts from five different larvae. P<0.0001 (Mann-Whitney test). (C,D) Representative images (C) and quantification (D) of neutrophil motility in the head mesenchyme of 3 dpf larvae, n=41 for Tg(lyzC:GFP), n=38 for Tg(U6a/c: ctrl sgRNA, lyzC:GFP)mu21 and n=47 for Tg(U6a/c: rac2 sgRNA, lyzC:GFP)mu28 from three different larvae. P=0.9282 and P=0.4747 (one-way ANOVA). (E-G) Tg(LyzC:Cas9, Cry:GFP)mu26 was crossed with Tg(U6a/c: ctrl guide, LyzC:GFP, mpeg:mcherry-H2B) or Tg(U6a/c: rac2 guides, LyzC:GFP, mpeg:mcherry-H2B). Representative images (E) and quantification (F) of macrophage motility (F) and neutrophil motility (G) in the head mesenchyme of 3 dpf larvae. The macrophage tracks are red, and the neutrophil tracks are yellow. n=80 for macrophages in the control group and n=65 for macrophages in rac2 knockouts. n=34 for neutrophils in the control group and n=32 for macrophages in rac2 knockouts from four different larvae. P=0.0735 and P=0.0001 (Mann-Whitney test). In B,D,F and G, the individual points are mean speeds for individual cells. The red and green lines indicate the mean velocity in each group. (H,I) Adults from three separate founders in Tg(LyzC:Cas9, Cry:GFP)mu26 were crossed with Tg(U6a/c: ctrl guide, LyzC:GFP)mu21 or Tg(U6a/c: rac2 guides, LyzC:GFP)mu28. Representative gel images of the T7E1-treated PCR products amplifying the sgRNA1 and sgRNA2 targeting sites (H) or non-neutrophil cells (I) from 3 dpf embryos. The white arrows indicate the expected cleavage bands. (J) Deep sequencing of the rac2 loci described in H. The sequences on the top are wild-type sequences, and the five most frequent types of mutations are shown. Point mutations, deletions and insertions are all observed. See also Movies 2, 3, 4. Scale bars: 100 µm.
Fig. 3. Re-expression of \( \text{rac2} \) rescued neutrophil motility defects in neutrophil-specific \( \text{rac2} \) knockout fish. (A) Schematics of the plasmids for neutrophil-specific rescue. The sgRNA-resistant \( \text{rac2} \) gene of wild type (\( \text{rac2-R-WT} \)), dominant-negative (\( \text{rac2-R-DN} \)) or constitutively active (\( \text{rac2-R-CA} \)), along with the mCherry reporter gene, were cloned into the plasmid carrying \( \text{rac2} \) sgRNAs. (B,C) Representative tracks (B) and quantification (C) of neutrophil motility in the head mesenchyme of 3 dpf larvae injected with plasmids encoding \( \text{rac2-R (WT)} \), \( \text{rac2-R-D57N (DN)} \) or \( \text{rac2-R-Q61L (CA)} \). (D) Schematics of the plasmids used to generate stable \( \text{Tg[lyzC:rac2-2A-mcherry]} \) or \( \text{Tg[lyzC:rac2-Q61L-2A-mcherry]} \) lines. Representative tracks (E) and quantification (F) of neutrophil motility in the head mesenchyme of 3 dpf larvae injected with plasmids encoding \( \text{rac2-R-WT} \), \( \text{rac2-R-DN} \) and \( \text{rac2-R-CA} \) from four or five different larvae. \( p<0.0001 \) (one-way ANOVA). See also Movies 5, 6. Scale bars: 100 µm.

Defect is a result of the loss of Rac2 function in neutrophils. As controls, supplementing \( \text{rac2} \)-deficient neutrophils with either a dominant-negative D57N (Deng et al., 2011), or a constitutively active Q61L version of \( \text{rac2} \) (Gu et al., 2001), failed to restore cell motility, indicating that spatially or temporally regulated \( \text{Rac2} \) activity is required to drive neutrophil migration.

The \( \text{Rac2} \) Q61L induces abnormal cell proliferation in human cells (Gu et al., 2001). However, whether this mutation impacts neutrophil migration is yet to be determined in both human and zebrafish. On this end, we generated a transgenic zebrafish line, \( \text{Tg[lyzC:mcherry-2A-Rac2CA]} \), overexpressing the Rac2 Q61L in neutrophils (Fig. 3D). Interestingly, in the \( \text{Tg[lyzC:mcherry-2A-Rac2CA]} \) stable line, no significant change in neutrophil motility was observed compared to the \( \text{Tg[lyzC:mcherry-2A-Rac2WT]} \) stable line, which overexpresses wild-type \( \text{Rac2} \) (Fig. 3E,F, Movie 6). Together, the \( \text{Rac2 Q61L} \) mutant does not have a dominant function but only impacts neutrophil mobility in the \( \text{rac2} \) knockout background.

Disruption of \( \text{cdk2} \) using the tissue-specific knockout system also suppressed neutrophil motility

Our previous study revealed an unexpected and critical role of Cdk2 in neutrophil migration and chemotaxis (Hsu et al., 2019). To ensure that the neutrophil-specific knockout system is feasible for disrupting other genes, we injected plasmids carrying \( \text{cdk2} \) targeting sgRNAs into \( \text{Tg[lyzC:cas9, cry:GFP]} \) embryos. Two sgRNAs targeting exon 4, which encodes the Cdk2 catalytic domain were selected (Fig. 4A). Neutrophil motility was significantly reduced by the tissue-specific \( \text{cdk2} \) disruption (Fig. 4B,C; Movie 7), recapitulating the phenotypes observed in the stable lines overexpressing a kinase-dead dominant-negative form of Cdk2, D145N (DN) (Hsu et al., 2019). To confirm that neutrophil migration defects did indeed result from \( \text{cdk2} \) disruption, we again performed rescue experiments. Restoring wild type, but not the dominant-negative, Cdk2 expression partially rescued the neutrophil migration defects caused by \( \text{cdk2} \) gene disruption (Fig. 4D-F; Movie 8). Our results indicate that the Cdk2 kinase activity is required for neutrophil motility in zebrafish, and that the neutrophil-specific knockout system is a robust tool to reach this conclusion.

Neutrophil-specific \( \text{rac2} \) knockout disrupts Rac activation

To observe alternations of Rac activation resulting from \( \text{rac2} \) disruption, we used a Rac-binding domain of PAK1 fused with GFP (PBD-GFP) (Benink and Bement, 2005) to mark the location of active Rac in neutrophils. This reporter has been used in previous studies in human neutrophil-like HL-60 cells and revealed that active Rac localizes to the cell front during migration in vitro (Benard et al., 1999; Srinivasan et al., 2003; Peng et al., 2011). Again, we cloned

Fig. 4. Neutrophil-specific knockout of \( \text{cdk2} \) reduced neutrophil motility. (A) Schematic of the gene structure of the zebrafish \( \text{cdk2} \) gene. The two target sites of the sgRNAs are in exon 4. (B,C) Representative tracks (B) and quantification (C) of neutrophil motility in the head mesenchyme of 3 dpf \( \text{Tg[lyzC:Cas9, cry:GFP]} \) larvae injected with plasmids carrying \( \text{cdk2} \) sgRNAs. (D) Schematic diagrams of the plasmids used to rescue Cdk2 expression. (E,F) Representative tracks (E) and quantification (F) of neutrophil motility in the head mesenchyme of 3 dpf \( \text{Tg[lyzC:Cas9, cry:GFP]} \) larvae injected with plasmids carrying \( \text{cdk2-R-WT} \) or \( \text{cdk2-R-D145N (DN)} \). \( n=20 \) for each group from three different larvae. \( p<0.0001 \) (Mann–Whitney test). In C and F, the individual points are mean speeds for individual neutrophils. The red and green lines indicate the mean velocity in each group. See also Movies 7, 8. Scale bars: 100 µm.
this construct in the pME entry vector described in Fig. 1B and obtained the plasmid that allowed expression of the biosensor in the control or rac2 knockout background (Fig. 5A). Here, in zebrafish, the PBD-GFP probe was enriched at both the front and rear in the migrating neutrophils in the control. When neutrophils migrate, Rac activity oscillated: active Rac first concentrated on the cell front and later shifted to the back. No discernible enrichment of Rac activity was detected in rac2-deficient neutrophils (Fig. 5B,C; Movie 9).

Forster resonance energy transfer (FRET)-based biosensors are used to detect protein conformational changes and interactions. To determine the subcellular location of Rac activation using a second approach, the ‘Raichu’ (Ras superfamily and interacting protein chimeric unit) Rac1-FRET probe developed by the Matsuda group (Itoh et al., 2002) was cloned into our sgRNA plasmids (Fig. 5D). When Rac is activated, the binding of Rac-GTP to PAK1-CRIB (CDC42/Rac interactive binding domain) increases FRET and the YFP/CFP fluorescence ratio. In neutrophils isolated from the ‘Raichu’ reporter mouse strain, active Rac localizes at both the front and back during chemotaxis (Johnsson et al., 2014). Consistent with this report, in zebrafish, Rac activity is higher at the cell periphery and oscillated between the front and back of migrating neutrophils expressing control sgRNAs. The rac2-defective neutrophils lost the ability to polarize and protrude, and did not display proper RAC activity (Fig. 5E,F; Movie 10), indicating that Rac2 function is required for the spatially and temporally coordinated Rac activation during neutrophil interstitial migration.

**Neutrophil-specific rac2 knockout alters actin cytoskeletal dynamics**

To observe the alternations in the actin cytoskeleton resulting from rac2 disruption, we used the calponin-homology domain of Utrophin (Utr-CH)-GFP (Burkel et al., 2007; Barros-Becker et al., 2017; Lam et al., 2014) to label stable F-actin in zebrafish neutrophils as described previously (Barros-Becker et al., 2017; Lam et al., 2014) (Fig. 6A). In Tg(lyzC:cas9, cry:GFP)pu26 larvae transiently expressing control sgRNAs, stable F-actin was enriched at the rear of migrating neutrophils. On the contrary, (Utr-CH)-GFP was not enriched at any specific subcellular locations in rac2 knockout neutrophils (Fig. 6B,C; Movie 11), indicating a loss of stable actin and cell polarity.

An actin stress probe, actin–cpstFRET–actin (AcpA), was recently developed to report forces within F-actin filaments (Johnsson et al., 2014). The sensor consists of a FRET pair flanking two linked β-actin monomers. When incorporated into F-actin filaments, the mechanical force in the filaments twists AcpA and decreases FRET efficiency. Thus, actin stress can be inferred from CFP/YFP ratio (Fig. 6D). We expressed the sensor and control or rac2 sgRNAs in Tg(lyzC:cas9, cry:GFP)pu26 embryos. Actin stress was relatively higher at the front and rear of neutrophils in control cells during migration. In contrast, rac2 knockout neutrophils showed decreased actin stress (Fig. 6E,F; Movie 12), suggesting that Rac2 is required for actin polymerization and force generation. Taken together, the ease of incorporating various biosensors into the neutrophil-specific knockout system allows live imaging of dynamic signaling events during cell migration.
migration in the knockout background, providing a flexible tool to interrogate gene function and determine mechanism.

Ribozyme-mediated gRNA generation for a neutrophil-specific knockout

We evaluated another gateway system for neutrophil-specific gene modification, in which the Cas9 protein is ubiquitously expressed and the sgRNA is processed by ribozymes and expressed in a neutrophil-restricted manner (Walton, 2018). This strategy was adapted from a previous study using an all-in-one plasmid containing a universal promoter (Lee et al., 2016). Here, we separated the Cas9 and sgRNA into two plasmids (Fig. 7A). The sgRNA is processed by the hammerhead and hepatitis delta virus ribozymes at the 5′ and 3′ ends, respectively. The noncoding RNA, MALAT1, forming a triple helical structure at the 3′ end (Wilusz et al., 2012), is incorporated to stabilize the tdTomato mRNA. The sgRNA and the tdTomato reporter gene is expressed as a single transcript driven by the lyzC promoter. We generated a transgenic zebrafish line, Tg(ubb:cas9, cry:GFP)\textsuperscript{lyt2}, and cloned the same control and the rac2 sgRNA1 into the ribozyme-mediated knockout system. The sgRNA plasmids carrying rac2 sgRNA or ctrl sgRNA were injected into the F2 embryos of the Tg(ubb:cas9, cry:GFP)\textsuperscript{lyt2} line. Neutrophil motility was significantly decreased in the zebrafish larvae carrying rac2 sgRNA, indicating sufficient gene disruption (Fig. 7B,C; Movie 13). Notably, we observed a slight decrease in neutrophil motility when we transiently injected the control sgRNA plasmids into the wild-type background (Fig. 7D,E; Movie 14), raising some slight concerns regarding expressing the sgRNA using ribozyme-mediated processing machinery in neutrophils.

**DISCUSSION**

Here, we report a robust and flexible neutrophil-specific knockout system in zebrafish. Using the lens-restricted GFP expression, we can easily select the positive fish carrying the Cas9 transgene. Transient expression of sgRNAs and visualization of edited cells can be achieved by injecting plasmids containing sgRNAs and GFP into the Cas9-expressing embryos. We also demonstrated the knockout efficiency using sgRNAs targeting rac2 and cdk2 using neutrophil motility as a proxy. To establish a causal relationship between the phenotype and the sgRNA-mediated genome editing, we rescued the migration phenotype by expressing sgRNA-resistant rac2 or cdk2. The stable lines expressing Cas9 protein and rac2 sgRNAs constructed here showed an inheritable ability to generate rac2 knockout lines at the F2 generation. We expect that the Cas9-driver line can be maintained and crossed with different reporter lines with different sgRNAs to achieve a multiplexed knockout. With three lyzC promoter-driven constructs in one neutrophil, we were still able to observe the expected phenotypes (Fig. 5B, Fig. 6B).

The optimization of the CRISPR/Cas9 vectors from our previous research lies in: (1) removing the 2A-mcherry tag and (2) separating the Cas9 and sgRNA elements into two constructs. Although the knockout efficiency is acceptable with multiple genes in our previous system, gene-specific rescue or biosensor imaging in the knockout background was challenging. The updates described here overcome previous limitations and significantly increase the efficiency of making genetic changes in zebrafish, and understanding the resulting changes in cell structural and signaling molecules. An example of how this work changes our ability to probe mechanisms comes from our own previous experience. In 2017, we obtained a neutrophil-specific mfn2 knockout line that displayed strong neutrophil adhesion defects. However, at that time, we encountered significant problems performing the rescue experiment or conducting a mechanistic study in the TSKO background. We could not observe mitochondrial morphology or related molecular/metabolic changes in any neutrophils with gene knockouts, and our work in zebrafish stopped with phenotypic observation. We only used zebrafish data for initial discovery, and the rest of the work was completed in human cells (Zhou et al., 2020). With the recent advance described here, we could have performed more
Fig. 7. Neutrophil-specific expression of ribozyme-processed rac2-targeting sgRNAs reduced neutrophil motility. (A) Schematic of the design of the plasmids for a second neutrophil-specific knockout system. Cas9 is expressed ubiquitously, whereas an sgRNA is expressed only in neutrophils. (B,C) Representative tracks (B) and quantification (C) of neutrophil motility in the head mesenchyme of 3 dpf Tg(uub:Cas9, cry:GFP)Δ466 larvae injected with plasmids carrying sgRNAs of control (ctrl) or rac2. n=29 for control and n=30 for rac2 transient knockouts from three different larvae. P=0.0001 (Mann–Whitney test). (D,E) Representative tracks (D) and quantification (E) of neutrophil motility in the head mesenchyme of 3 dpf wild-type AB zebrafish larvae injected with Tol2-lyzC-RFP plasmid or plasmids carrying sgRNAs of control (ctrl) or rac2. n=32 for no sgRNA control from three different larvae, n=29 for control sgRNA and n=28 for rac2 sgRNA from four different larvae. P=0.0018 and P=0.0111 (one-way ANOVA). In C and E, the individual points are mean speeds for individual neutrophils. The red and green lines indicate the mean velocity in each group. See also Movies 13, 14. Scale bars: 100 µm.
processed in zebrafish (Lee et al., 2016). Our results indicate that this approach is efficient enough to induce a phenotype. However, we observed some reduction in cell speed when the ribozyme-flanking sgRNA was expressed alone. This effect could be tissue specific and will require careful evaluation in the future. Following this work, our aim is to expand functionality to allow multiplexed sgRNA expression. One direction to pursue is to evaluate both microRNA-based and tRNA-based processing machinery to process multiple sgRNAs from one transcript for tissue-specific gene disruption in zebrafish (Port and Bullock, 2016; Wang et al., 2015; Xie et al., 2015). The Cas12a family proteins have recently been demonstrated to mediate highly efficient genome editing in zebrafish, and will likely also enable tissue-specific disruption (Liu et al., 2019; Moreno-Mateos et al., 2017).

In summary, we developed a robust and flexible neutrophil-specific knockout system in zebrafish. Using this system, we gained insights into the role of Rac2 in regulating the actin cytoskeleton and the subcellular location of Rac activation in zebrafish neutrophils. Our system is suitable for various genetic studies and screens, which can be achieved by injecting plasmid encoding different sgRNAs into the Cas9-expression fish embryos. We also expect that our system can be adapted for gene function studies in various tissues using other tissue-specific promoters.

MATERIALS AND METHODS

Animals

The zebrafish experiment was conducted in accordance with internationally accepted standards. The Animal Care and Use Protocol was approved by the Purdue Animal Care and Use Committee, adhering to the Guidelines for Use of the National Institutes of Health Intramural Research Program (protocol number 1401001018). MATLAB and the samplesizewsp function was used to calculate the sample sizes required for each experiment based on conservative estimates for the variability in the controls for each type of experiments, with a power of 0.9 significance level of 0.05) in two sample experiments. Data were quantified blindly by an investigator not involved in data collection. To generate transgenic zebrafish lines, plasmids with the Tol2 backbone were co-injected with Tol2 transposase mRNA into embryos of the AB strain at the one-cell stage as described previously (Deng et al., 2011).

Plasmids

All plasmid constructions were generated by gateway cloning using LR Clonase II Plus enzyme (Invitrogen). The pME-GFP, p3E-polyA and the destination vector for the sgRNA plasmid, pDesTol2pA2, were obtained from the Tol2Kit (Kwan et al., 2007). To design sgRNAs, CRISPRscan (www.crisprscan.org) was used. SgRNAs with the highest score and without any off-targets were selected. The destination vector pDestTol2pAcryFP (Addgene, 64022) was used to generate the final Cas9 plasmids. The p3E-U6a-U6c plasmids containing rac2 sgRNAs or control sgRNAs (Addgene 107591, 107600), and p5E-lyzC (Addgene 107591) were described as described previously (Deng et al., 2011). At least two founders (F0) for each line were obtained. Experiments were performed with F2 larvae produced by F1 fish derived from multiple founders. For transient knockout, only plasmids were injected into the Tg(lyzC:cas9, cry:GFP)pu26 transgenic line.
performed at temperatures between 26 and 28°C. The cell velocity was quantified using ImageJ with the MTrackJ plug-in and plotted in Prism 6.0 (GraphPad). The fluorescence intensity quantification was performed using an algorithm written in our lab (https://github.com/tomato990/subcellular-intensity-reader). The kymograph was generated using Helm 1.0 (Deng et al., 2014).

T7 endonuclease I assay
T7 endonuclease I (New England Biolabs, M0302) was used to detect mutations caused by CRISPR/Cas9. Neutrophils and non-neutrophil cells were sorted based on GFP expression as described previously (Hsu et al., 2019). Genomic DNA containing sgRNA recognition sites was amplified by PCR from sorted cells. PCR products were purified with a PCR purification kit (Clontech) and re-annealed in a thermocycler using the following conditions: 95°C for 5 min; 95-85°C with a ramp rate of −0.3°C/s; and 85-25°C with a ramp rate of −0.1°C/s. Re-annealed PCR products were incubated with T7 endonuclease I at 37°C for 1 h, followed by agarose gel electrophoresis. Primers used for this assay are described below.

**Mutational efficiency quantification**
To determine the mutation efficiency in Tg(LyzC:Cas9, Cry:GFP, U6a/c: rac2 guides, LyzC-GFP), the rac2 loci around the sgRNA binding site was amplified by PCR using the following primers: Rac2-sgRNA1-F, 5’-GT-GAGTATCACTACATAGAAGG-3’; Rac2-sgRNA1-R, 5’-GTCGCCACT-GAAGTCTTGATG-3’; Rac2-sgRNA2-F, 5’-GGCTGTATCTAGTC-AAGAGATAG-3’; and Rac2-sgRNA2-R, 5’-GGCATCATTGTTGCATGA-CAC-3’. Amplification was followed by wide sequencing at the sequencing center at Purdue University. Mutational efficiency was calculated as the ratio of sequence reads containing the CRISPR-endonuclease-generated mutations to total sequence reads obtained from the sequencing center.

**Statistical analysis**
Statistical analysis was performed using Prism 6 (GraphPad). An unpaired two-tailed Student’s t-test or one-way ANOVA was used to determine the statistical significance of differences between groups. *P*<0.05 was considered statistically significant. Individual *P* values are indicated in the figures, with no data points excluded from statistical analysis. One representative experiment of at least three independent repeats is shown.

**Acknowledgements**
We thank Dr David Umulis (Purdue University) for critical reading of our manuscript and insightful comments. Raichu-Rac1 was a generous gift from Dr Miki Matsuda (Kyoto University, Japan).

**Competing interests**
The authors declare no competing or financial interests.

**Author contributions**

**Funding**
This work was supported by the National Institutes of Health (R3SGM119787 to D.Q.; AI125517, AI130236 and AI127115 to D.T.; and R56GM124913 to G.Z.); and the Purdue Center for Cancer Research (P30CA232168, for shared resources). A.Y.H. was supported by a Cagiantas Fellowship, Purdue University. Deposited in PMC for release after 12 months.

**Data availability**
Raw sequencing reads have been deposited to the NCBI Sequence Read Archive under BioProject accession PRJNA706779.

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