Fish-Specific Duplicated dmrt2b Contributes to a Divergent Function through Hedgehog Pathway and Maintains Left-Right Asymmetry Establishment Function

Sha Liu, Zhi Li, Jian-Fang Gui*

State Key Laboratory of Freshwater Ecology and Biotechnology, Institute of Hydrobiology, Chinese Academy of Sciences, Graduate School of Chinese Academy of Sciences, Wuhan, China

Abstract

Gene duplication is thought to provide raw material for functional divergence and innovation. Fish-specific dmrt2b has been identified as a duplicated gene of the dmrt2a/terra in fish genomes, but its function has remained unclear. Here we reveal that Dmrt2b knockdown zebrafish embryos display a downward tail curvature and have U-shaped somites. Then, we demonstrate that Dmrt2b contributes to a divergent function in somitogenesis through Hedgehog pathway, because Dmrt2b knockdown reduces target gene expression of Hedgehog signaling, and also impairs slow muscle development and neural tube patterning through Hedgehog signaling. Moreover, the Dmrt2b morphants display defects in heart and visceral organ asymmetry, and some lateral-plate mesoderm (LPM) markers expressed in left side are randomized. Together, these data indicate that fish-specific duplicated dmrt2b contributes to a divergent function in somitogenesis through Hedgehog pathway and maintains the common function for left-right asymmetry establishment.

Introduction

Gene duplication is thought to be the primary source of new genes. Many teleost fish, including zebrafish, experience an additional genome-wide duplication event [1]. Since then, many of the duplicated genes have been lost, but a substantial percentage of the duplicates have been retained, and functional divergence has occurred in some duplicates [1,2]. Recently, to identify some differentially expressed genes in early embryogenesis, two kinds of SMART cDNAs were respectively synthesized from the mature eggs and gastrula embryos, and the gastrula embryo SMART cDNAs were respectively synthesized from the mature eggs and gastrula embryos, and the gastrula embryo SMART cDNA library was constructed in Carassius auratus gibelio [3,4,5,6,7]. Following this program, many differentially expressed genes at gastrula stage were screened [5], and some of them were characterized and functionally analyzed [8,9]. Significantly, a fish-specific duplicated gene dmrt2b was identified in dmrt gene family from Carassius auratus gibelio [4].

dmrt2, ddx and mab-3 related transcription factor 2, is a member of a gene family of putative transcription factors. These transcription factors share a highly conserved zinc-finger-like DNA-binding domain (DM domain) which is implicated in sex determination [10]. However, recent studies show that the family genes function not only in sex determination, but also in embryonic development [11]. In zebrafish, dmrt2 was originally called terra [12]. dmrt2b is another duplicated copy of the dmrt2 in the genome, and dmrt2a/terra and dmrt2b have been designated for distinguishing them.

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Results

Molecular characterization and expression pattern of Dmrt2b during embryogenesis

As a new member of dmrt gene family, dmrt2b was firstly cloned in gibel carp (Carassius auratus gibelio) (GenBank accession number:
EF029082). Database searches revealed the closest homologue in zebrafish (Danio rerio) (GenBank accession number: NM_001079976). The DM domain of zebrafish Dmrt2b is 100% identical to that of gibel carp Dmrt2b, and has 94.7% and 93.0% identities to that of zebrafish and gibel carp Dmrt2a respectively. Significantly, the full-length amino acid sequence of zebrafish Dmrt2b has only 31.16% and 30.57% identities to zebrafish and gibel carp Dmrt2a, although it has 75.68% identity to gibel carp Dmrt2b (Figure 1A). The significant sequence divergence between Dmrt2b and Dmrt2a implicates the potential occurrence of functional divergence.

Subsequently, the expression pattern of zebrafish dmrt2b during embryogenesis was analyzed by RT-PCR, Western blotting and whole-mount in situ hybridization. As shown in Figure 1B and Figure 1C, the Dmrt2b transcription is initiated from around shield stage at 6 hpf, and kept at a basic stable level from bud stage at 16 hpf to the hatched larvae stage at 48 hpf. And, the Dmrt2b protein is expressed after bud stage at 10 hpf, and the abundant expression level is detected after 24 hpf of embryogenesis. Figure 1D-G shows in situ mRNA distribution of dmrt2b during embryogenesis. Significantly, dmrt2b is specifically expressed in somites during somitogenesis.

Dmrt2b knockdown leads to defects in somitogenesis and reduces target gene expression of Hedgehog signaling

Abundant expression of dmrt2b in somites implies its significant functions in zebrafish embryo development. To assess the functions, we undertook loss-of-function experiments in zebrafish by morpholinos. We utilized two non-overlapping antisense morpholino oligonucleotides (MOs), Dmrt2b-MO1 and Dmrt2b-MO2, which designed to block translation by binding to 25 bases of the 5' UTR of the Dmrt2b mRNA. Because the translation-blocking morpholino was demonstrated to be a powerful tool for studying the effects of near-total loss of function during early stages of development [16], Western blot detection showed a predicted size protein band in the Cont-MO embryos at 24 hpf, whereas the corresponding protein band was notably reduced in the embryos injected with Dmrt2b-MO1 (5 ng per embryo) and Dmrt2b-MO2 (5 ng per embryo) respectively, and no any signal was observed in the embryos co-injected with Dmrt2b-MO1 (2.5 ng) and Dmrt2b-MO2 (2.5 ng) (Supplementary Information, Figure S1A). Because the translation-blocking morpholino was demonstrated to be a powerful tool for studying the effects of near-total loss of function during early stages of development [16], Western blot detection showed a predicted size protein band in the Cont-MO embryos at 24 hpf, whereas the corresponding protein band was notably reduced in the embryos injected with Dmrt2b-MO1 (5 ng per embryo) and Dmrt2b-MO2 (5 ng per embryo) respectively, and no any signal was observed in the embryos co-injected with Dmrt2b-MO1 (2.5 ng) and Dmrt2b-MO2 (2.5 ng) (Supplementary Information, Figure S1A). Morphological observation displayed obvious embryonic development defects in the Dmrt2b morphants. As shown in Figure 2, in comparison with normal embryos injected with 5 ng Cont-MO (Figure 2A), the morphant embryos injected with 5 ng Dmrt2b-MO1 or 5 ng Dmrt2b-MO2 exhibit an abnormal ventrally curved body shape, short trunk and U-shaped somites at 24 hpf (Figure 2B,C), and when the two non-overlapping antisense MOs (2.5 ng each, totally 5 ng) were jointly injected, the morphant embryos exhibit similar phenotypes and even more severe defects (Figure 2D). Because co-injection of multiple targeting MOs was found to be effective in mediating p53 activation, and p53-MO can attenuate the off-targeting effects [20,21], we also checked the embryonic effects of Dmrt2b-MO and p53-MO co-injection. In comparison with normal embryos injected with p53-MO (Figure 2F), no any signal was observed in the embryos co-injected with Dmrt2b-MO and p53-MO (Figure 2G). To find further evidence that Dmrt2b is involved in Hedgehog signal pathway, we first checked the expression of patched1 (ptc1), a direct transcriptional target of the Hh pathway [28]. In comparison with high expression in somites of Cont-MO embryos (Figure 2N), an obvious down-regulation of ptc1 transcript was observed in the Dmrt2b morphants (23/27 embryos) at 24 hpf (Figure 2O). In embryos co-injected with Dmrt2b-MO and dmrt2b mRNA, low expression of ptc1 was rescued (21/25 embryos) at 24 hpf (Figure 2P). To reveal the signal point at which Dmrt2b exerts its activity on the Hh signaling cascade, we also examined the expression of the Hh target gene nkx2.2a in the developing brain [29]. As shown in Figure 2Q-S, Dmrt2b-MO causes a notable reduction of nkx2.2a in the brain (22/25 embryos), and co-injection with dmrt2b mRNA could rescue the phenotype (19/24 embryos). Quantification analysis through qPCR further confirmed the down-regulated changes of nkx2.2a and ptc1 expression (Figure 2T).

Considering sequence similarity and functional redundancy of Dmrt2a and Dmrt2b, we additionally tested the ability whether Dmrt2a could rescue Dmrt2b knockdown and vice versa. As shown in Figure 3, both Dmrt2a and Dmrt2b morphant defects (Figure 1A, B) could not be rescued by Dmrt2b mRNA (Figure 3C) and dmrt2a mRNA respectively (Figure 3D). Then we analyzed the protein level of Dmrt2b in the Dmrt2a-MO injected embryos, and no any notable change of Dmrt2b expression was observed in the Dmrt2a morphants (Figure 3E). We also examined the expression of dmrt2a in the Dmrt2b morphants by whole-mount in situ hybridization, and no any change was not detected (Figure 3F and G). The data indicate that the loss-of-function of Dmrt2b leads to notable defects in somitogenesis and reduces target gene expression of Hedgehog signaling. And, the defects do not affect the expression of dmrt2a and Dmrt2a and Dmrt2b can not compensate the function for each other when they are knocked down.

Dmrt2b is involved in slow muscle development through Hedgehog signaling

To assess the role of Dmrt2b in slow muscle development, we further analyzed the expression affect of Dmrt2b knockdown on
Figure 1. Molecular characterization and expression pattern of zebrafish Dmrt2b during embryogenesis. (A) Amino acid alignment of zebrafish Dmrt2b with gibel carp Dmrt2b, gibel carp Dmrt2a and zebrafish Dmrt2a. Similar and identical amino acids are highlighted in grey and black boxes. The line indicates the DM domain. (B) RT-PCR detection of dmrt2b in zebrafish embryonic development stages, and beta-actin mRNA as the control. (C) Western blot detection of Dmrt2b during the zebrafish embryo development, and Tubulin was used for the control. (D-H) Whole-mount in situ hybridization detection of dmrt2b on somitogenesis embryos as indicated stage. The arrows indicate positive signals in the somites. doi:10.1371/journal.pone.0007261.g001
Figure 2. Dmrt2b morphants display defects in somitogenesis and Hedgehog signaling. (A–G) Morphology of 24 hpf embryos injected with Cont-MO (A), Dmrt2b-MO1 (B), Dmrt2b-MO2 (C), Dmrt2b-MO (D), Dmrt2b-MO+dmrt2b mRNA (E), p53-MO (F), Dmrt2b-MO+p53-MO (G). The arrows indicate the off-target cell death in Dmrt2b-MO morphant and the reduced cell death in the Dmrt2b-MO+p53-MO morphant. (H) The statistical data of three independent experiments on Dmrt2b knockdown, dmrt2b mRNA rescue and p53 MO co-injection. Results are represented as mean±SD of three separate experiments. (I) Western blot detection of Dmrt2b knockdown during embryogenesis. The protein extracts from embryos (16 hpf and 24 hpf) were analyzed by Western blot using the polyclonal anti-Dmrt2b antibody. A band of about 41 KD was not detected in Dmrt2b morphants. The picture represents typical result from three separate experiments. (J–M) Dmrt2b morphant exhibits U-shape somites. Morphology of embryos injected with Cont-MO display the typical 'chevron' shape (J, K). Morphology of embryos injected with Dmrt2b-MO display the U-shape (L, M). Whole-mount in situ hybridization of ptc1(N, O, P) (Anterior is left) and nkx2.2a(Q, R, S) (Anterior is top) in embryos injected with Cont-MO (N, Q) or Dmrt2b-MO (O, R) and embryos co-injected with Dmrt2b-MO with dmrt2b mRNA (P, S) at 24 hpf. (T) qPCR analysis of the expression changes of ptc1 and nkx2.2a in 24 hpf embryos injected with Cont-MO or Dmrt2b-MO. Results represent mean±SD of three separate experiments.
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Rescued by co-injection of dmrt2b embryos at 26 hpf. Significantly, the perturbed phenotypes can be muscle pioneers are inhibited (Figure 4D) in the morphant embryos. Especially, the numbers of superficial slow muscle fibers is severely perturbed in the morphant embryos. As shown in Figure 4, in comparison with normal PKA previously [31,32], when the dominant negative segmentation, resulting in significant impairment in slow muscle fibers and muscle pioneers cannot be increased (Figure 4N). In these experiments, a total of 20 to 30 embryos were contributed to each analysis. Figure 4O and 4P respectively show the quantitative data of slow muscle fibers and muscle pioneer cells per somite. The data indicate that Dmrt2b knockdown does appear to affect slow muscle fibers, and therefore, Dmrt2b is involved in slow muscle development.

Dmrt2b is involved in neural tube patterning through Hedgehog signaling

Hh signaling has been identified as a key morphogen in patterning of the ventral neural tube [34]. To assess the contribution of Dmrt2b to Hh-mediated signaling in the neural tube, we analyzed the expression of nks2.2a. In non-injected and control-injected embryos, nks2.2a is expressed in lateral floor plate of the neural tube at 24 hpf (Figure 3A). In the Dmrt2b morphant embryos, the nks2.2a expression is severely suppressed (22/25 embryos) (Figure 3B). Moreover, we observed that dnPKA mRNA injection could lead to ectopic expression of nks2.2a in the neural tube of Cont-MO embryo (19/25 embryos) (Figure 3C) as described previously [31,55]. Consistently with the epistatic analysis using slow muscle fiber and muscle pioneer markers (Figure 4I, J), co-injection of Dmrt2b-MO with dnPKA normalizes the ectopic expression of nks2.2a (18/26 embryos) (Figure 3D). These data indicate a requirement of Dmrt2b during the development of neural tube for the appropriate cellular response to the Hh signals. To determine whether Dmrt2b is required for transcription of Hedgehog genes, we further examined the expression of three Hedgehog genes shha, shhb and shhc in Dmrt2b morphant embryos and control embryos. Whole mount in situ hybridization analysis showed that shha, shhb and shhc expression appeared normal in Dmrt2b morphant embryos at 10 hpf (bud stage) and at 24 hpf (Supplementary Figure S2). These data suggest that Dmrt2b should act downstream of Hedgehog genes and upstream of Gli protein, the core components of the Hh pathway [36], because three Hedgehog genes shha, shhb and shhc are not affected in Dmrt2b morphants, but dnPKA and Su(fu)-MO can counteract the Dmrt2b morphant phenotypes [22,31].

Dmrt2b morphants display defects in heart and visceral organs asymmetry

As described previously, Dmrt2a is essential for establishment of LR asymmetry in zebrafish development [14], and shh regulates the establishment of LR asymmetry in zebrafish development [37]. We also examined heart asymmetry defects in Dmrt2b morphant embryos by using specific marker of heart, cardiac myosin light chain 2 (cmlc2) [38]. In embryos injected with Cont-MO, 100% of the embryos show the heart tube on the left side, at 48 hpf (Figure 6A, I). By contrast, in the Dmrt2b morphant embryos, 24.9% of the heart tubes shift rightward and 19% remain in the middle (Figure 6B, C, D and I) (n = 86 to 113 in three separate experiments).

In addition to the heart, morphological LR asymmetry of the visceral organs was also observed in zebrafish embryos [37]. To determine if LR asymmetry of the visceral organs was affected in Dmrt2b knockdown embryos, we checked the expression of foxa3 which marks the liver. As expected, 100% of embryos injected...
Figure 4. Dmrt2b is involved in slow muscle development through Hedgehog signaling. Analysis of slow muscle development by whole-mount immunostaining of 26 hpf embryos with antibodies labeling slow MyHC (F59) (A, C, E, G, I, K, M) and engrailed muscle pioneer cells (4D9) (B, D, F, H, J, L, N). Lateral view of embryos injected with Cont-MO (A, B), Dmrt2b-MO (C, D), embryos co-injected with Dmrt2b-MO+dmrt2b (E, F), Cont-MO+dnPKA (G, H), Dmrt2b-MO+dnPKA (I, J), Cont-MO+Su(fu)-MO (K, L), Dmrt2b-MO+Su(fu)-MO (M, N). White arrows in 4d9 panel indicate reduction of engrailed staining in the knockdown embryos compared to controls. All images show the somites over the yolk extension. Anterior is left in all images. (O) Quantitative analysis of slow muscle fiber number per somite in embryos at 26 hpf, injected as indicated. Data represent average ± SD. *** indicates significance of p < 0.0001 for Dmrt2b-MO vs. Cont-MO, Dmrt2b-MO+dmrt2b, Dmrt2b-MO+dnPKA and Dmrt2b-MO+Su(fu)-MO. n = 20 to 38 embryos per condition. (P) Quantitative analysis of muscle pioneer cells number per somite in embryos at 26 hpf, injected as indicated. Data represent average ± SD. *** indicates significance of p < 0.0001 for Dmrt2b-MO vs. Cont-MO, Dmrt2b-MO+dmrt2b and Dmrt2b-MO+dnPKA. n = 20 to 38 embryos per condition. doi:10.1371/journal.pone.0007261.g004
with Cont-MO exhibit a leftward budding of the liver, at 48 hpf (Figure 6E, J). By contrast, 24.0% of the liver shift rightward, and 20.7% remain in the middle in the Dmrt2b marphant embryos (Figure 6F, G, H and J) (n = 83 to 100 in three separate experiments).

Dmrt2b activity is important for establishment of left-right asymmetry in lateral plate mesoderm

Left-sided expression of several genes such as lefty1, spaw and pitx2c in the lateral plate mesoderm (LPM) is important to heart and visceral organs asymmetry [39,40,41]. To determine whether the randomized heart and visceral organs asymmetry is preceded by alteration of underlying molecular cues, we analyzed these LR markers.

*lefty1* is a Nodal signaling antagonist, which expresses in the left dorsal diencephalon and left LPM at 22 somites stage (overlapping the prospective heart field) (Figure 7A, E). However only 33.3% of the Dmrt2b-MO injected embryos show the normal *lefty1* patterns, the rest exhibit right-sided, bilateral, or absent expression (Figure 7B–D, F–H and Figure 7Q) (n = 36).

*spaw*, which encodes a nodal-related protein in zebrafish, is normally expressed in the left LPM at 20 somites stage and regulates left-right asymmetry (Figure 7I). However, the later asymmetric *spaw* expression in the LPM is dramatically altered in Dmrt2b morphant embryos with only 38.1% showing left-sided expression, compared with embryos injected with Cont-MO (Figure 7F–I and Figure 7K) (n = 63). *pitx2c* encoding a bicoid-related transcription factor is also expressed in the left LPM at 22 somite stage (Figure 7M). As expected, in Dmrt2b-MO injected embryos (Figure 7M–P), only 38.6% showing normal left-sided expression compared with embryos injected with Cont-MO (Figure 7Q) (n = 70).

Since defects in midline integrity can also lead to abnormal LR asymmetry. We examined the midline structure of these embryos by using *in situ* hybridization of *ntl*. Expression of the *ntl* was not affected in embryos injected with Dmrt2b-MO (Figure 7T and 7U). The above data demonstrate *dmrt2b* importance in the establishment of left-right asymmetry of the body plan.

**Dmrt2a does not contribute to Hedgehog pathway during zebrafish somitogenesis**

As previously reported, Dmrt2a is also involved in somitogenesis in zebrafish [12]. To examine whether Dmrt2a is contribute to hedgehog pathway as Dmrt2b, we performed comparative studies on two marker genes between Dmrt2a and Dmrt2b. As shown in Figure 8, in comparison with normal expression in Cont-MO embryos at 24 hpf (Figure 8A), the down-regulation of *ptc1* transcripts in Dmrt2b morphant embryos (Figure 8B) is not observed in Dmrt2a-MO injected embryos (Figure 8C). Similar results are also observed in checking *nkx2.2a* expression. In contrast with the severely suppression in Dmrt2b morphant embryos (Figure 8E), the expression level of *nkx2.2a* does not reduce in the Dmrt2a morphant embryos (Figure 8F). Moreover, we also detected abnormal *spaw* expression in the Dmrt2a morphant embryos as described previously [14]. Figure 8G–J show the *spaw* expression alteration in left LPM in the 43 Dmrt2a-MO injected embryos with only 39.5% left-sided expression, 20.9% bilateral expression, 23.3% right-sided expression and 16.3% absent expression. The data indicate that Dmrt2a and Dmrt2b play similar rules in the establishment of left-right asymmetry of the body plan, but Dmrt2a does not contribute to Hedgehog pathway during zebrafish somitogenesis.

Dmrt2a has been described to be involved in synchronizing left-right somitogenesis in zebrafish. To clarify whether Dmrt2b has similar role for synchronizing left-right somitogenesis, we further analyzed expression changes of the presomatic mesoderm marker genes *her1* and *deltaC* in the Dmrt2b morphants. As shown in Supplementary Figure S3, the expression of *her1* and *deltaC* are not affected in the Dmrt2b-MO injected embryos. The data confirm that Dmrt2b is not involved in synchronizing left-right somitogenesis in zebrafish.
Discussion

Gene duplication provides raw material for functional divergence and innovation [42], and neofunctionalization and subfunctionalization hypotheses have been proposed. The neofunctionalization hypothesis argues that after duplication one daughter gene retains the ancestral function while the other acquires new functions. In contrast, the subfunctionalization hypothesis asserts that the functions of the ancestral gene are partitioned between the duplicated genes [43]. Recent advances have suggested a new model termed subneofunctionalization, because neither neofunctionalization nor subfunctionalization can alone be explained in a large proportion of duplicate genes, and rapid subfunctionalization is often accompanied by prolonged and substantial neofunctionalization. This new model proposes that a large part of duplicated genes might play both overlapping functions and divergent functions [44].

In this study, we found significant sequence divergence between the fish-specific duplicated gene dmrt2b and dmrt2a, observed severe defects in somitogenesis, slow muscle development and neural tube patterning in the Dmrt2b loss-of-function zebrafish embryos by using morpholino-mediated knockdown strategy, and revealed the underlying mechanism that is involved in Hedgehog signaling pathway. Moreover, we also found the overlapping functions between dmrt2b and dmrt2a in LR asymmetry establishments, including heart and visceral organ asymmetry establishment and lateral plate mesoderm asymmetry establishment. Additionally, we further verified that Dmrt2a and Dmrt2b play common overlapping functions in the establishment of left-right asymmetry of the body plan, but Dmrt2a does not contribute to Hedgehog pathway (Figure 9A). Therefore, the current study clarifies exact function of the fish-specific duplicated gene dmrt2b, and provides a significant and interesting case for functional divergence of duplicate genes.

Another significant advance in this study was to find similar defect phenotypes in the Dmrt2b morphants to that in the so called “you-type mutants” which are demonstrated to be inactivated in particular genes in the Hedgehog signaling pathway [22,23,24,25,26,27,45]. The defects include the curled tail and the U-shaped somite boundaries. Subsequently, we have confirmed that the expression of Hedgehog target genes ptc1 and nck2.2a are severely inhibited in the Dmrt2b morphants, and both slow muscle development and neural tube development are also disrupted by Hedgehog pathway impairment. Moreover, we have observed normal expression of three Hedgehog genes shha, shhb and shhb in Dmrt2b morphants, indicating that Dmrt2b mophant defects are in the response to Hedgehog signals not in Hedgehog transcript expression at low level. The data suggest that Dmrt2b might act downstream of Hedgehog gene transcription. To assess what level in the pathway Dmrt2b is active, we further used two rescue experiments. Co-injection of dmPKA and Su(fu)-MO could rescue some defects of the Dmrt2b morphants in slow muscle development, implicating that Dmrt2b should act downstream of Hedgehog genes and upstream of Gli protein, the core components of the Hh

Figure 6. Dmrt2b morphant embryos display defects in heart and visceral organs asymmetry. (A–D) Expression of cmilc2 in 48 hpf embryos detected by whole mount in situ hybridization with antisense RNA probe. (E–H) Expression of foxa3 in 48 hpf embryos detected by whole mount in situ hybridization. Arrows in foxa3 panel indicate the liver. Graphical representation of the percentage of embryos exhibiting the expression patterns for cmilc2 (I) and for foxa3 (J). Results are represented as mean±SD of three separate experiments.

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Divergent Function of *dmrt2b*

Figure 7. *Dmrt2b* knockdown disrupts L–R identity in lateral plate mesoderm. (A–D) *lefty1* normal expression in left dorsal diencephalon at 22 somites stage were disrupted embryos injected with *Dmrt2b-MO*. (E–H) *lefty1* normal expression in left LPM at 22 somites stage were disrupted embryos injected with *Dmrt2b-MO*. (Q) Graphical representation of the percentage of embryos exhibiting the expression patterns for *lefty1* in *Dmrt2b* knockdown embryos and control embryos. (I–L) *spaw* normal expression in left LPM at 20 somites stage were disrupted embryos injected with *Dmrt2b-MO*. (R) Graphical representation of the percentage of embryos exhibiting the expression patterns for *spaw* in *Dmrt2b* knockdown embryos and control embryos. (M–P) *pitx2c* normal expression in left LPM at 22 somites stage were disrupted embryos injected with *Dmrt2b-MO*. (S) Graphical representation of the percentage of embryos exhibiting the expression patterns for *pitx2c* in *Dmrt2b* knockdown embryos and control embryos. (T) Lateral view of the embryos at 22 somites stage. Expression of the *ntl* in embryos injected with *Dmrt2b-MO* is similar to expression of *ntl* in control embryos. Results are represented as mean±SD of three separate experiments. (U) qPCR analysis of the expression changes of *ntl* in 24 hpf embryos injected with *Cont-MO* or *Dmrt2b-MO*. The data represents mean±SD of three separate experiments.

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pathway [36], and therefore contribute to somitogenesis, slow muscle development and neural tube patterning by inhibiting the expression of Hh target genes (Figure 9B). In addition, we found that Dmrt2a expression was not affected in the Dmrt2b morphants (Figure 3F and G), and overexpression of dmrt2a was previously observed to induce rapid apoptosis in the mesoderm [12]. Moreover, no any Hedgehog signals responding defects were observed in zebrafish Dmrt2a morphants in our experiment (Figure 8). Therefore, the new function of Dmrt2b during somitogenesis and the underlying mechanism that is involved in Hedgehog signaling pathway are different from the function of Dmrt2a during somitogenesis which is linked to apoptosis.

Additionally, an interesting aspect of the Dmrt2b morphants is that it encompasses only a subset of defects observed from other zebrafish Hedgehog pathway mutants. For example, pectoral fins, which are disrupted in mutants syu [22] and smu [24], appear normal in Dmrt2b knockdown embryos (some Dmrt2b morphant embryos have smaller pectoral fins). We also observed incomplete absence of the slow muscle phenotype. These phenotypic characteristics may be a result of the functionally redundant components in Hh pathway. Indeed, similar varying levels of slow muscle development defects have been described in the different Hh pathway genetic mutants [46], potentially indicating compensatory effects by other genes in the Hedgehog pathway.

In conclusion, we have endeavored to elucidate the roles of Dmrt2b in the developing zebrafish embryo, and compared the Dmrt2b function to the Dmrt2a function. Through morpholino loss-of-function approach, we demonstrated that Dmrt2b contributes to multiple developmental processes, including zebrafish somitogenesis through Hedgehog pathway and establishment of left-right asymmetry. We have uncovered important new roles for Dmrt2b in zebrafish, which were unanticipated by studies of this molecule’s duplicated gene Dmrt2a in zebrafish and orthologous gene Dmrt2 in mouse. The data suggest fish-specific duplicated genes Dmrt2b and Dmrt2a play overlapping roles in establishment of Left-Right asymmetry and divergent functions in somitogenesis, which Dmrt2b contributes to Hedgehog pathway.

Materials and Methods

Maintenance of zebrafish
A breeding colony of zebrafish (Danio rerio) were maintained at 28.5°C on a 14 h light/10 h dark cycle [47]. All embryos used were collected by natural spawning and staged according to standard procedures [48].

Antisense morpholino and mRNA microinjection
Two non-overlapping translation-blocking morpholino oligonucleotides (MOs) [16] Dmrt2b-MO1 and Dmrt2b-MO2, standard control MO against the zebrafish β-globin intron, p53 MO, and Su(fu) MO, were obtained from Gene Tools, LLC. The morpholino sequences were as follows: Dmrt2b-MO1, 5’-TTCTCACG-AACCCACGACGCTCATC-3’; Dmrt2b-MO2, 5’-CCGCTTTAGTGAGAGATTTGAGCT-3’; stand control MO(Cont-MO), 5’-CCTCCTACCTCAGTTACAATTTATA-3’; p53 MO

Figure 8. Dmrt2a does not contribute to Hedgehog pathway during zebrafish somitogenesis. (A–C) The expression of ptc1 in embryos injected with Cont-MO, Dmrt2b-MO and Dmrt2a-MO at 24 hpf. (D–F) The expression of nkk2.2a in embryos injected with Cont-MO, Dmrt2b-MO and Dmrt2a-MO at 24 hpf. (G–J) spaw normal expression in left LPM at 20 somites stage were disrupted embryos injected with Dmrt2b-MO.

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Figure 9. A schematic diagram illustrating the duplication and subsequent functional diversification of zebrafish Dmrt2a and Dmrt2b (A), and a hypothesized acting position that Dmrt2b is involved in Hedgehog pathway (B).

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essentially as previously described [21]. Embryos were fixed at the stages indicated and processed respectively (kind gifts from Dr. Randall Moon, University of Washington). Embryos were incubated at 28.5°C in a microcentrifuge to remove insoluble particles, samples were loaded on a 12% SDS gel. Western blot analysis was performed according to the previous report [50] using the anti-Dmrt2b antibody and anti-tubulin antibody.

Whole-mount in situ hybridization

Antisense probes for pitx2c [35], nkx2.2a [29], pax6a [52], lefty1 [41], spaw [39], en [38], foxa2 [53] and ntl [5] were prepared as described previously. Probes for pitx2c and echidna Hedgehog (shh) were generated directly from PCR products that included T7 or T3 RNA polymerase binding sequence at the 3’ end. Primer sequences used were: pitx2c forward, 5’-ACTGGCCGCAAATCTTGCACTA-3’; reverse, 5’-CATTAACCTCATAAAGGAAGTTGGCTTGGCTTAGCTTCATA CGC-3’. shh forward, 5’-ATGAGACTCTCGACGGGGCCG-3’; reverse, 5’-CATTAACCTCATAAAGGAAGTTGGCTTGGCTTAGCTTC ATAGGC-3’. Probes for sonic Hedgehog (shha) and tiggywinkle Hedgehog (shhb) were prepared using shhb-pT7Ts and shhb-pT7Ts respectively (kind gifts from Dr. Randall Moon, University of Washington). Embryos were fixed at the stages indicated and processed essentially as described previously [21].

Immunohistochemistry

Embryos were grown to the indicated stages and processed essentially as described previously. Immunostainings of whole zebrafish embryos were performed following standard protocols. The 4d9 antibody used 1:100 dilution, which recognizes the engrailed protein in the nuclei of muscle pioneer cells, the F59 antibody used 1:100 dilution, which labels slow myosin heavy chain (MyHC) in zebrafish [30], were obtained from the Developmental Studies Hybridoma Bank. Fluorescent secondary antibodies against mouse used for detection were FITC or Cy3 conjugated. Stained embryos were photographed on fluorescence optics of Confocal Microscope LEICA DMIRE2 (Leica, Germany).

Quantification of slow muscle defects

As previous report [33], 26 hpf embryos were fixed and stained using F59 antibody. Numbers of slow muscle fibers were counted in 5 somites over the yolk extension per embryo.

Supporting Information

Figure S1 Dmrt2b translation is blocked by non-overlapping morpholinos. (A) The Dmrt2b-MO1 and Dmrt2b-MO2 target sequences are shown in relation to the 5’ UTR region of the Dmrt2b mRNA sequence. (B) Western blot assay showing Dmrt2b translation in the embryos injected with Cont-MO, Dmrt2b-MO1, Dmrt2b-MO2 and Dmrt2b-MO (Dmrt2b-MO1+Dmrt2b-MO2). The signal of Dmrt2b protein were significant reduced in Dmrt2b-MO1, Dmrt2b-MO2 and Dmrt2b-MO injected embryos. Found at: doi:10.1371/journal.pone.0007261.s001 (0.72 MB TIF)

Figure S2 Dmrt2b is not required for shha, ihhb and shhb transcription. Dorsal views of embryos at the bud stage (10 hpf) (A, C, E, G and I). Lateral views of embryos at 24 hpf (B, D, F, H and J). Expression of shha in embryos injected with Dmrt2b-MO (C and D) is similar to expression of shha in control embryos (A and B). Expression of ihhb in embryos injected with Dmrt2b-MO (G and H) is similar to expression of ihhb in control embryos (E and F). Expression of shhb in embryos injected with Dmrt2b-MO (K and L) is similar to expression of shhb in control embryos (I and J). Found at: doi:10.1371/journal.pone.0007261.s002 (3.61 MB TIF)

Figure S3 Whole-mount in situ hybridization of persomotic mesoderm genes her1 and deltaC in the Dmrt2b morphants. Expression patterns of her1 in the Dmrt2b-MO (A) and Cont-MO (B) embryos. Expression patterns of deltaC in the Dmrt2b-MO (C) and Cont-MO (D) embryos. All the embryos are at 10 somites stage. Panels show dorsal views, anterior to the top. Found at: doi:10.1371/journal.pone.0007261.s003 (4.94 MB TIF)

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Author Contributions

Conceived and designed the experiments: SL, JFG. Performed the experiments: SL, ZL. Analyzed the data: SL. Contributed reagents/materials/analysis tools: SL, JFG. Wrote the paper: SL, JFG.

References

1. Postlethwait J, Amores A, Cresko W, Singer A, Yan YL (2004) Subfunction partitioning, the telost radiation and the annotation of the human genome. Trends Genet 20: 481–490.
2. Taylor JS, Braasch I, Pickney T, Meyer A, Van de Peer Y (2003) Genome duplication, a trait shared by 22000 species of ray-finned fish. Genome Res 13: 382–390.
3. Dong CH, Yang ST, Yang ZA, Zhang L, Gui JF (2004) A C-type lectin associated and translocated with cortical granules during oocyte maturation and egg fertilization in fish. Dev Biol 265: 341–354.
4. Liu J, Shi YH, Yin J, Gui JF (2005) Screen of differentially expressed genes between gastrula embryos and tail bud embryos in genomic gilber carp (Carassius auratus gibelio). J Genet Genomics 32: 253–263.
5. Liu JX, Shi YH, Gui JF (2005) Screen of differentially expressed genes at gastrula stage during embryogenesis of gilber carp. Acta Hydrobiologica Sinica 29: 359–365.
6. Xia J, Wen JJ, Chen B, Gui JF (2001) Differential gene expression in fully-grown oocytes between gynogenetic and gonochoristic crucian carps. Gene 271: 109–116.
7. Zhou L, Wang Y, Gui JF (2000) Genetic evidence for gonochoristic reproduction in gynogenetic silver crucian carp (Carassius auratus gibelio) as revealed by RAPD assays. J Mol Evol 51: 498–506.
8. Xia JH, Liu JX, Zhou L, Li Z, Gui JF (2000) Apo-14 is required for digestive system organogenesis during fish embryogenesis and larval development. Int J Dev Biol 45: 1089-1098.
9. Yin J, Xia JH, Du ZX, Liu J, Zhou L, et al. (2007) Developmental expression of CagMdkb during gibel carp embryogenesis. Int J Dev Biol 51: 761–769.
10. Raymond CS, Shamu CE, Shen MM, Seifert KJ, Hirsch R, et al. (1998) Evidence for evolutionary conservation of sex-determining genes. Nature 391: 691–695.
11. Hong C, Park B, Saint-Jeannet J (2007) The function of Dnmt genes in vertebrate development: it is not just about sex. Dev Biol 310: 1–9.
12. Meng A, Moore B, Tang H, Yuan B, Lin S (1999) A Drosophila doublesex-related gene, terra, is involved in somitogenesis in vertebrates. Development 126: 1259–1268.
13. Zhou X, Li Q, Lu H, Chen H, Guo Y, et al. (2008) Fish specific duplication of Dnmt2: characterization of zebrfish Dnmt2b. Biochimie 90: 878–877.
14. Saide L, Lourenço R, Gonçalves A, Palmeirim I (2005) terra is a left-right asymmetry gene required for left-right synchronization of the segmentation clock. Nat Cell Biol 7: 918–920.
15. Luo C, Wang Y, Kobuho H, Kettlewell J, Zarkower D, et al. (2006) Targeted disruption of the DM domain containing transcription factor Dnmt2 reveals an essential role in somite patterning. Dev Biol 290: 280–290.
16. Nasvisci A, Ekker SC (2000) Effective targeted gene 'knockdown' in zebrfish. Nat Genet 26: 216–220.
17. Etker SC (2000) Morphants: a new systematic vertebrate functional genomics approach. Yeast 17: 392–396.
18. Etker S, Larsson J (2001) Morphant technology in model developmental systems. Genesis 30: 89–93.
19. Schlueter PJ, Royer T, Farah MH, Laser B, Chan SJ, et al. (2006) Gene activation by knockdown technologies. PLoS Genet 3: e78.
20. Mei J, Zhang QY, Li Z, Lin S, Gui JF (2008) C1q-like inhibits p53-mediated apoptosis and controls normal hematopoiesis during zebrfish embryogenesis. Dev Biol 319: 273–284.
21. Barrese MJ, Sticken HL, Devoto SH (2000) The zebrfish slow-muscle-omitted gene product is required for Hedgehog signal transduction and the development of slow muscle identity. Development 127: 2189–2199.
22. CagMdkb during gibel carp embryogenesis. Int J Dev Biol 51: 761–769.
23. Westerfield M (2000) The zebrafish book. Eugene: Univ. of Oregon Press.
24. Wolff C, Roy S, Lewis KE, Schauerte H, Joerg-Rauch G, et al. (2004) iguana mutations in the zebrafish hedgehog and protein kinase A in neural tube and somite patterning. Development 122: 2835–2846.
25. Ruiz i Altaba A (1999) GlI proteins and Hedgehog signaling: development and cancer. Trends Genet 15: 418–425.
26. Schilling TF, Conkordet JP, Ingham PW (1999) Regulation of left-right asymmetries in the zebrfish by Shh and BMP4. Dev Biol 210: 277–287.
27. Etker S, Larsson J (2001) Morphant technology in model developmental systems. Genesis 30: 89–93.
28. He X, Zhang J (2005) Rapid subfunctionalization accompanied by prolonged and substantial neofunctionalization in duplicate gene evolution. Genetics 169: 1157–1164.
29. van Eeden F, Granato M, Schach U, Brand M, Furutani-Seiki M, et al. (1996) Mutations affecting somite formation and patterning in the zebrfish, Danio rerio. Development 123: 153–164.
30. Wollf C, Roy S, Lewis KE, Schauerte H, Joerg-Rauch G, et al. (2004) iguana encodes a novel zinc-finger protein with helix-coil domains essential for Hedgehog signal transduction in the zebrfish embryo. Genes Dev 18: 1565–1576.
31. Westerfield M (2000) The zebrafish book. Eugene: Univ. of Oregon Press.
32. Kimmel CB, Ballard WW, Kimmel SR, Ullmann B, Schilling TF (1995) Stages of embryonic development of the zebrafish. Dev Dyn 203: 155–310.
33. van Eeden F, Granato M, Schach U, Brand M, Furutani-Seiki M, et al. (1996) Mutations affecting somite formation and patterning in the zebrfish, Danio rerio. Development 123: 153–164.
34. Wang Y, Zhou L, Gui JF (2000) Differential and spermatogenic cell-specific expression of DMRT1 during sex reversal in protogynous hermaphroditic groupers. Mol Cell Endocrinol 263: 156–172.
35. Link V, Shevelevko A, Heisenberg C (2006) Proteomes of early zebrafish embryos. BMC Dev Biol 6: 1.
36. Krauss S, Johansen T, Korzh V, Moens C, Ericson J, et al. (1991) Zebrafish pax2a: a paired box-containing gene expressed in the neural tube. EMBO J 10: 3609–3619.
37. Edental J, Naslein-Vollhard C (1998) fork head domain genes in zebrfish. Dev Genes Evol 208: 245–250.