**Twist Controls Skeletal Development and Dorsoventral Patterning by Regulating Runx2 in Zebrafish**

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**Abstract**

**Background:** Twist1a and twist1b are the principal components of twists that negatively regulate a number of cellular signaling events. Expression of runx2 and downstream targets is essential for skeletal development and ventral organizer formation and specification in early vertebrate embryos, but what controls ventral activity of maternal runx2 and how twists function in zebrafish embryogenesis still remain unclear.

**Methodology/Principal Findings:** By studying the loss of twist induced by injection of morpholino-oligonucleotide in zebrafish, we found that twist1a and twist1b, but not twist2 or twist3, were required for proper skeletal development and dorsoventral patterning in early embryos. Overexpression of twist1a or twist1b following mRNA injection resulted in deteriorated skeletal development and formation of typical dorsalized embryos, whereas knockdown of twist1a and twist1b led to the formation of abnormal embryos with enhanced skeletal formation and typical ventralized patterning. Overexpression of twist1a or twist1b decreased the expression of runx2b, whereas twist1a and twist1b knockdown increased runx2b expression. We have further demonstrated that phenotypes induced by twist1a and twist1b knockdown were rescued by runx2b knockdown.

**Conclusions/Significance:** Together, these results suggest that twist1a and twist1b control skeletal development and dorsoventral patterning by regulating runx2b in zebrafish and provide potential targets for the treatment of diseases or syndromes associated with decreased skeletal development.


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**Introduction**

The initiation of skeletal development begins with the migration and proliferation of cells from cranial neural crest, sclerotome, and lateral plate mesoderm into mesenchymal condensations that form the template of the future skeleton [1]. Under a precise genetic regulation by a repertoire of transcription factors, chondrocytes or osteoblasts arise from these condensations to form cartilage and bone [2]. Understanding the signaling pathways involved in skeletal development will help to treat diseases associated with abnormal bone formation.

The vertebrate dorsoventral axis establishment represents the earliest event where programmes of induction and cellular commitment are used. The process is controlled by the actions of maternal and zygotic genes, and usually involves cell–cell interactions, cell movements, and spatiotemporally controlled expression of dorsoventral determinants [3,4,5]. The identification of target genes of the signals involved in embryonic axis development will help to establish the genetic network underlying these processes.

Zebrafish was described as ‘the canonical vertebrate’, due to the similarities between zebrafish and mammalian biology. Because of the transparent and continuous visualization of the developmental processes, the use of rapid and transient assays, and the feasibility and affordability of large-scale forward genetic screens in zebrafish [6,7], it has attracted researchers from various fields, such as neuroscience, hematopoiesis or cardiovascular research. These shared features have prompted many laboratories to begin exploiting the unique advantages of the zebrafish system to study human disease, especially in the identification of human disease gene homologs. Thus, the zebrafish is a good model for investigating human development, including skeletal development and dorsoventral axis establishment.

Runx2 (runt-related transcription factor 2, also known as Cbfa1, Osf2 and AML3) is not only a transcription factor essential for skeletal development [8,9], but also an important maternal
determinant of ventral zygotic genes in zebrafish [10]. Previous studies have identified two orthologs of the mammalian Runx2, runx2a and runx2b, in zebrafish. Both genes have type 1 (T1) and type 2 (T2) isoforms and share sequence homology and gene structure with the mammalian genes, and map to regions of the zebrafish genome displaying conserved synteny with the region where the human gene is localized. Although both genes are expressed in developing skeletal elements and skeletal defects appeared following depletion of either runx2a or 2b, the effect of runx2b knockdown on skeletal defects is much more significant than runx2a knockdown [11]. In addition, runx2b has been shown to be able to regulate the expression of osterix and osteocalcin [12–13]; these observations strongly indicate the important role of runx2b during developmental bone formation. Moreover, depletion of maternal runx2b (especially T2 isoform), rather than runx2a, strongly dorsalizes embryos, due to loss of the earliest zygotic expression of ventral genes, resulting in expansion of dorsal gene expression [10,14].

Twist genes code for regulatory bHLH proteins that are essential for embryonic development and are conserved across the metazoan [15]. The zebrafish twist family comprises four genes: twist1α, twist1β—orthologs to mammalian Twist1, twist2—a gene from a new clade that does not exist in mammals [16]. Previous study has demonstrated that the twist developmental patterns, e.g., expression in cephalic neural crest, sclerotome and lateral plate mesoderm, are conserved in tetrapods to the fish [16]. twist is abundantly expressed in invaginating/migrating cells in jellyfish. However, the roles of twist in skeletal development and axis establishment in zebrafish remain to be clarified.

We have demonstrated that TWIST, activated by HIF-1α under hypoxic conditions, inhibited human mesenchymal stem cell (MSC) osteogenesis via direct downregulation of T2 RUNX2, which led to suppression of T1 RUNX2 (DC Yang et al, 2011, PLoS ONE, In press). Since several signaling pathways involved in craniofacial skeletal development in zebrafish are similar to the pathways involved in osteogenic differentiation of mammalian MSCs [6,17], we hypothesized that the expression of T1 and T2 runx2b would also be suppressed by twist in zebrafish under hypoxic conditions. In the current study, we first found that in zebrafish under hypoxic conditions, bone mineralization was inhibited and T1 and T2 runx2b and their downstream targets were downregulated, while hif-1α and twist were upregulated. Inhibition of twist1α and twist1β by morpholino oligonucleotides (MO) increased the expression of T1 and T2 runx2b, and induced ventralized patterning, while microinjecting zebrafish embryos with full length twist1α and twist1β mRNA decreased the expression of T1 and T2 runx2b, and induced dorsalized patterning in zebrafish. Twist1α and twist1β morpholinos also rescued hypoxia-induced decrease in craniofacial skeletal development in zebrafish.

**Results**

**Hypoxia inhibits bone mineralization in zebrafish**

To understand the effects of hypoxia on skeletal development, zebrafish at 2 dpf (day post fertilization) were subjected to the hypoxic environment (5% O2) for 1 day and stained with Alizarin Red S (ARS) for determination of the degree of mineralization at 8 dpf. Hypoxia treatment induced a decrease in ARS staining in the spine area (Fig. 1A) compared to the control group. Similarly, treatment of zebrafish at 2 dpf with DFX, a hypoxia-mimicking agent, for 6 days also induced a decrease in ARS staining in the spine compared to the control group (Fig. 1B). The numbers of positive staining developing centra form ring with ARS were calculated (Fig. 1C). To further visualize normal and defective bone development in zebrafish embryos induced by hypoxia or treatment with DFX, the embryos were labeled with the fluorescent chromophore calcein. Treatment of zebrafish at 2 dpf with hypoxia (Fig. 1D) or DFX (Fig. 1E) and quantitative data (Fig. 1F) for 6 days also induced a decrease in calcein labeling in the spine area compared to the control group. To examine whether decreased bone mineralization was attributed to the reduction in chondrogenesis, Alcian Blue staining was performed at 8 dpf. At this time, no obvious reduction in Alcian Blue staining was noted in zebrafish exposed to hypoxia or DFX treatment compared to the controls (Fig. 1G and Fig. 1H). Collectively, these data suggest hypoxia or DFX treatment inhibited bone mineralization but not chondrogenesis in zebrafish.

**Hypoxia inhibits the expression of type 1 and type 2 runx2b and their downstream targets**

Because MO-based loss-of-function studies revealed the involvement of runx2b, rather than runx2a in skeletal development [11], we chose runx2b for exploring the key molecule that was targeted by hypoxia or DFX to regulate osteogenesis. We found that when zebrafish was exposed to hypoxia or treated with DFX at 48 hpf for 24 h, there was decreased expression of T1 and T2 runx2b (Fig. 2A). Consistent with the effects on upstream transcription factor, hypoxia exposure or DFX treatment also decreased the expression of osterix, collagen 10a1 (coll10a1), alkaline phosphatase (ap) and osteocalcin (oc), (Fig. 2B); Collectively, these data suggest hypoxia or DFX treatment downregulates T1 and T2 runx2b and their downstream targets to inhibit skeletal development in zebrafish.

**Hypoxia downregulates type 1 and type 2 runx2b via the hif-1α-twist pathway**

It has been demonstrated that HIF-1α directly downregulated Twist under hypoxic conditions in mammalian cells [18]. To examine the molecular mechanism involved in hypoxia-mediated inhibition of osteogenesis, we first demonstrated that the expression of hif1α and twist1/2 (Fig. 3) was increased when zebrafish was treated with DFX or hypoxia at 48 hpf for 24 h. These data suggest hypoxia inhibits skeletal development and activates the hif1α-twist pathway in zebrafish.

**twist1α and twist 1β knockdown independently inhibits runx2b expression and induces ventralized embryos in zebrafish**

Since runx2b also controls the dorsoventral patterning in the early zebrafish embryos [10,14], we therefore examined the involvement of the twist isoforms in zebrafish axis establishment [16]. Interestingly, microinjection of twist1α and twist1β atgMOs but not scrambled MO (MO-SC), twist2 and twist3 atgMOs dose-dependently induced an increase in Class 3 (V3) and 4 (V4) ventralized embryos [19] (Fig. 4A–C, Fig. S1A), in which anterior forebrain was deficient, and eyes and notochord were completely lost. Because runx2b upregulates ventral genes [10,14], these results suggested that knockdown of twist1α and twist1β increased the expression of runx2b. Consistently, quantitative RT-PCR revealed twist1α and twist1β atgMOs increased T1 and T2 runx2b expressions compared to MO-SC embryos at 3, 14, and 48 hpf (Fig. 4D, 4E), while MO-SC, twist2 and twist3 atgMOs failed to induce any increase in runx2b expression (Fig. S1B).

More importantly, twist1α or twist1β atgMO-induced ventralized embryos and increase in runx2b gene expression were rescued by co-injection with twist1α or twist1β mRNA (Fig. 4C–4E).
Further, injection of twist1a or twist1b mRNA alone induced the appearance of dorsalized embryos (Fig. S1C), in which ventral tail vein and fin were missing, blood circulation was impaired [20], and the expressions of T1 and T2 runx2b were inhibited (Fig. S1D). It has been reported that knockdown of T2 runx2b with T2 runx2b atgMO induces the appearance of dorsalized embryos [10]. Similarly, twist1a or twist1b atgMO-induced increase of the ventralized embryos were inhibited by co-injection with T2 runx2b atgMO (Fig. 4C), suggesting the involvement of T2 runx2b in determining the twist1a or twist1b atgMOs-induced ventralized patterning.

Whole-mount in situ hybridization further demonstrated that no injection or MO-SC embryos expressed endogenous runx2b on all enveloping layer in a faint manner at 8 and 14 hpf, respectively (Fig. 5A, 5B). In contrast, the expression of runx2b at 8 hpf was strongly induced by twist1a atgMO and twist1b atgMO in the enveloping layer on the future ventral region (Fig. 5A). At 14 hpf, runx2b expression was induced by twist1a atgMO mainly in the neural crest of the forebrain and by twist1b atgMO both in the neural crest of the forebrain and somites (Fig. 5B). At 48 hpf, twist1a atgMO increased runx2b expression in the whole brain and produced new signals in the neural tube region of the V5/no defect embryos compared to no injection zebrafish embryos injected with MO-SC, twist2 or twist3 atgMOs (Fig. S2), which only showed runx2b expression in ceratobranchial 1–5 (Cb1-5) and cleithrum (Cl); twist1b atgMO also increased runx2b expression in the whole brain and produced a new signal in the tail of the V5/no defect embryos (Fig. 5C).

**twist1a and twist1b knockdown independently enhances bone formation in zebrafish**

We then examined whether microinjection of twist1a and twist1b atgMOs upregulated the expression of other osteoblast markers. Quantitative RT-PCR revealed both twist1a and twist1b atgMOs enhanced the expression of osteirx and col10a1 at 14 and 48 hpf, while no significant change was noted at 8 hpf (Fig. 5D). Finally, we determined whether microinjection of twist1a and twist1b atgMOs promoted functional mineralization both under normoxia and hypoxia. The embryos had normal morphology and survived up to 8 dpf were assayed for the degree of mineralization. These embryos showed increase in bone mineralization, as evident by the apparent increase of ARS staining in the spine area (Fig. 5E) in whole embryos studies. Similarly, calcein labeling also revealed...
an increase in bone mineralization in embryos microinjected with twist1a and twist1b atgMOs both in the absence or presence of DFX treatment (Fig. 5F). Together, these data suggest that the knockdown of twist1a or twist1b in zebrafish enhanced runx2b transcription, induced ventralized patterning, and promoted bone formation both under normoxic and hypoxic conditions.

**Discussion**

Twist is known to trigger epithelial-mesenchymal transition (EMT) mechanisms and increase cells with migratory ability. Although Twist is constantly expressed in various cells including osteoblasts, its roles in skeleton development is seldom, if ever, investigated [21]. Previous studies have shown that TWIST silencing enhanced osteoblast gene expression and matrix mineralization [22,23]. In this current study, we demonstrate that twist plays an essential role in skeleton development and axis establishment by regulating runx2b in zebrafish. Interestingly, only TWIST in human MSCs and twist1a and twist1b (orthologs of mammalian Twist), but not twist2 and twist3, in zebrafish possess these functions. Moreover, RUNX2 in human and runx2b in zebrafish are mainly involved in skeleton development or axis establishment, suggesting the conservation of Twist-Runx2 pathway in mammalian cells and zebrafish.

Previously, Twist has been demonstrated to inhibit DNA binding and gene activation by Runx2, while Runx2 expression was not affected in mice carrying Twist heterozygosity [24]. The current study found overexpression and knockdown of Twist increased and decreased the expression of Runx2 both at the mRNA and protein levels, respectively. The discrepancy between our study and the previous one [24] may be because the suppression of Twist expression by Twist heterozygosity in mice is not sufficient to downregulate Runx2, as is the minimal effect of the low dose of twist1b atgMO to increase runx2b expression (Fig. S3).

Twist has been reported to have a synergistic effect with dorsal and snail in integrating diverse dorsoventral patterning in the Drosophila embryo [25,26]. Runx2b, a maternal and zygotic mediator, has been reported to induce the expression of ventral gene such as ved, vent and vox in the earliest embryo of zebrafish (4 hpf) [10]; however the upstream molecule regulating the expression of runx2b has not been discovered. Knockdown of twist1a and twist1b inducing the expression of runx2b was observed as early as 8 hpf and 14 hpf, suggesting the early involvement of twist in controlling the dorsoventral patterning. The phenotypes of the knockdown of twist1a and twist1b include abnormalities in eyes, fusion of fore/midbrain and hindbrain, notochord, trunk, and other skeleton deformity, which are normally observed in the...
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S-stained zebrafish were photographed or preserved in solution made by 4 parts 100% glycerol plus 1 part 95% ethanol (v/v).

Calcein immersion

The procedures of calcein immersion are described by Du et al. [33] and Chen et al. [34] with some modifications. Briefly, 0.2% (w/v) of calcein (Sigma) solution was prepared and adjusted to pH 7.0 with 0.5 N NaOH. Zebrafish embryos from 8 day were immersed in the calcein solution in petri dishes for 5 min. Calcein-immersed embryos were rinsed several times with tap water, and then immersed in one liter of tap water for 10 min to allow the excess calcein to diffuse out of the tissues. The embryos were then euthanized in tricaine–methanesulfonate (MS 222), mounted on glass slides with methylcellulose (3%), and observed under a fluorescent microscopy.

RNA extraction and quantitative RT-PCR

Total RNA was prepared using the TriZol reagent (Invitrogen) according to the manufacturer’s specifications. cDNA was synthesized from 2 μg RNA and using Superscript III (Invitrogen), random primers (Invitrogen), 10 mM DTT (Invitrogen), and RNaseOUT ribonuclease RNase inhibitor (Invitrogen). cDNA was then 1/20 diluted with dH2O (Final cDNA concentration: 5 ng/μl). The quantitative RT-PCR was performed using 40 ng cDNA as the template in a 20 μl reaction mixture containing FastStart SYBR Green Master (Roche Applied Science) and a specific primer pair of each cDNA according to the published sequences which listed in the Table S1. Analysis of the results were carried out using the software supplied with the ABI Step One Real-Time PCR System machine and each expression was calculated relative to the zebrafish β-Actin (delta CT) and then relative to controls (delta delta CT) using the fluorescence threshold of the amplification reaction and the comparative CT method.

Preparation and microinjection of morpholino or RNA

Twists Morpholino antisense oligonucleotides (twist-MOs) were obtained from Gene Tools (Philomath, OR) with the sequences: zebrafish twist1α atgMO (5'-CTTAAGCGAGATGCCGCAAGGACCC-3'); twist1β atgMO (5'-GATGCGGCAAAGCGCGGAGGAC-3'); twist2 atgMO (5'-GAATTAATGGAAGAGGTTTCAATGCTC-3'); twist3 atgMO (5'-CATTGGAGAGGAACAGACTTGTGGA-3') and T2 runx2b atgMO (5'-CATTGCTGCCACATTGCGTCCTCCAAA-3') targets the sequence at the translation start site of twist mRNAs. As a control experiment, Scrambled control (MO-SC, 5'-GGGCTCTTTCGGGAGATCCTCGAA-3') recommended by Gene Tools was used. All of the above were prepared at stock concentrations of 1 mM and 2.3 nl per embryo was injected. Synthetic capped-mRNAs were prepared using mMessage mMachine (Ambion) from pGEMT-twist1α or pGEMT-twist1β, which containing coding sequences of zebrafish twist1α and twist1β, respectively. Embryos were collected by natural spawning and microinjected with MO or mRNA (overexpression) or MO/mRNA mixtures (rescue experiments) at 1- or 8-cell stage using a Narishige micromanipulator. For constructing plasmids pGEMT-twist1α or pGEMT-twist1β, the pDRN-LIB-twist1α and pME18s-FL3-twist1β were purchased from Open Biosystems and zebrafish twist1α or twist1β mRNAs were amplified by PCR using the primer pairs: twist1α-rescue-F: 5'-GATGTTTTAGGAGGAACAGC-3'; twist1α-rescue-R: TTTCCCTGAGGAGGTATC-3'; twist1β-rescue-F: GAGATGCGGAGAGGACACCTTGTGGA-3' and T2 runx2b-rescue-R: ATGGTGTTTGAATGATGATGAC-3' targets the sequence in cDNA of twist1α and twist1β, respectively. Embryos were collected by natural spawning and microinjected with MO or mRNA (overexpression) or MO/mRNA mixtures (rescue experiments) at 1- or 8-cell stage using a Narishige micromanipulator. For constructing plasmids pGEMT-twist1α or pGEMT-twist1β, the pDRN-LIB-twist1α and pME18s-FL3-twist1β were purchased from Open Biosystems and zebrafish twist1α or twist1β mRNAs were amplified by PCR using the primer pairs: twist1α-rescue-F: 5'-GATGTTTTAGGAGGAACAGC-3'; twist1α-rescue-R: TTTCCCTGAGGAGGTATC-3'; twist1β-rescue-F: GAGATGCGGAGAGGACACCTTGTGGA-3' and T2 runx2b-rescue-R: ATGGTGTTTGAATGATGATGAC-3' targets the sequence in cDNA of twist1α and twist1β, respectively.

Methods

Zebrafish maintenance and histological staining: Alcian blue and ARS staining

The procedures of ARS stain are described by Walker and Kimmel [32]. Zebrafish (Danio rerio) embryos were maintained in Embryo Medium at 28.5°C. To visualize developing bone at 8dpf, the embryos were immobilized in ice water, and then fixed for 24 h in buffered 4% paraformaldehyde. After washing twice with PBS, they were dehydrated in 50% and 100% ethanol each for 24 h. They were then placed in 0.05 mg/ml Alizarin Red S in 1% KOH for 1 h and cleared in 20% glycerol in 1% KOH for 40 min. Alizarin Red II stained zebrafish were photographed or preserved in 4% formaldehyde and 50% glycerol. S-stained zebrafish were photographed or preserved in solution made by 4 parts 100% glycerol plus 1 part 95% ethanol (v/v). Zebrafish at 48 hpf were treated with or without 21% O2, 5% O2, or indicated concentration of DFX for 1 day and the expression of hif-1α, twist1α, twist1b, twist2 and twist3 genes was assayed by quantitative RT-PCR. Results are shown as the relative expression to β-actin (mean ± SD) and significance was determined by Student’s t-test. (* p<0.05 and ** p<0.01 versus 21% O2).

Figure 3. Hypoxia increases the expression of hif1α and twist in zebrafish. Zebrafish at 48 hpf were treated with or without 21% O2, 5% O2, or indicated concentration of DFX for 1 day and the expression of hif1α, twist1α, twist1b, twist2 and twist3 genes was assayed by quantitative RT-PCR. Results are shown as the relative expression to β-actin (mean ± SD) and significance was determined by Student’s t-test. (* p<0.05 and ** p<0.01 versus 21% O2).

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ventralized embryos induced by mutation or knockdown of dorsal-specific genes [19] or overexpression of ventral-specific genes [27]. Together with Runx2, a number of molecules involved in dorsoventral patterning also participate in bone formation, including BMPs, Chordin, Noggin, Wnt/β-catenin, transforming growth factor-β (TGF-β) and fibroblast growth factors (FGFs) [28,29]. Runx1 and Runx3 also interact with similar molecules in haematopoiesis and gastric epithelial maintenance, respectively [30,31]. Whether the expressions of these genes are also regulated by Twist has not been clarified and necessitates future investigation. Future exploration of the twist signaling pathways may help in developing strategies to control skeleton development and dorsoventral patterning through suppressing runx2b.

In conclusion, these data provide convincing evidences for the important roles of Twist in controlling dorsoventral patterning, skeleton development and bone mineralization. Further exploration of the mechanism involved in Twist-mediate regulation of skeleton development and regeneration may provide new strategies for treating these diseases.

Preparation and microinjection of morpholino or RNA
Figure 4. Morpholino knockdown of twist1a and twist1b induces the appearance of ventralized embryos and increases the expression of runx2b in zebrafish. 

A, Antisense morpholinos (MO) for twist1a (twist1a atgMO), twist1b (twist1b atgMO), twist2 (twist2 atgMO) and twist3 (twist3 atgMO), were designed against the 5' UTR and ATG regions, which blocked translation of each transcript. Each atgMO was microinjected into 1-cell to 4-cell embryo and the percentage of each dorsoventral patterning was calculated at 36–48 hpf. B, Representative pictures of induced Class 1–5 ventralized (V1-5/no defect) and dorsalized (D) embryos. V and D phenotype annotations were described in ref. 20 and 21, respectively. C, Table summarizing the ventralized or dorsalized embryo features of zebrafish microinjected with indicated concentration of each atgMO with or without twist1a/1b mRNA or T2 runx2b atgMO. Zebrafish were microinjected with MO-SC, (D) twist1a atgMO or (E) twist1b atgMO with or without (D) twist1a mRNA or (E) twist1b mRNA and quantitative RT-PCR for T1 runx2b and T2 runx2b were performed at 8, 14 and 48 hpf (n = 3). Results are shown as the relative expression to β-actin (mean ± SD) and significance was determined by Student’s t-test. (* p<0.05 and ** p<0.01 versus MO-SC; # p<0.05 and ## p<0.01 versus twist1a/1b atgMO). doi:10.1371/journal.pone.0027324.g004
Figure 5. Morpholino knockdown of twist1a and twist1b induces runx2b expression and promotes bone formation in zebrafish. (A to D) Zebrafish were microinjected without (No injection, NI) or with control MO (MO-SC), twist1a or twist1b atgMOs and runx2b expression was analyzed by in situ hybridization at 8 hpf (A), 14 hpf (B), and 48 hpf (C) with runx2b probe. Pictures of lateral (L), animal pole (AP), and head region dorsal (HD) views are displayed. Runx2b expression was induced by twist1a or twist1b atgMO compared to MO-SC at 8 hpf, 14 hpf and 48 hpf. (arrow, ventral region; arrowhead, dorsal region; fb, forebrain area; hb, hindbrain area; ceratobranchial 1–5, Cb1–5; cleithrum, Cl). D, Quantitative RT-PCR for osterix and col10a1 was performed at 8, 14 and 48 hpf (n = 3). Results are shown as the relative expression to β-actin (mean ± SD) and significance was determined by Student’s t-test. (* p<0.05 and ** p<0.01 versus MO-SC). E, F, Zebrafish survived after microinjection with MO-SC or twist1a or twist1b atgMO were cultured with or without 100 μM DFX and bone mineralization was analyzed by (E) ARS staining and (F) calcein labeling at 8 dpf. The number of positive staining developing centra form ring with ARS or calcein labeling were calculated.

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Whole-Mount In Situ Hybridization and Sections

Whole mount in situ hybridization for the runx2b gene was performed at 68°C as described by Flores et al., 2008[10]. Samples were imaged using a Leica MZ16FA stereomicroscope with a Leica DC490 camera and the associated software. The designation of developmental stage of zebrafish was following those of Kimmel et al [35].

Supporting Information

Figure S1 Morpholino knockdown of twist1a and twist1b, but not twist2 and twist3, induces the appearance of ventralized embryos and increases the expression of runx2b in zebrafish. A, The table summarizing the ventralized embryo features (V1-5/no defect) of zebrafish injected with indicated concentration of each atgMO. B, Zebrafish were microinjected with twist2 or twist3 atgMO and quantitative RT-PCR for T1 runx2b and T2 runx2b were performed at 48 hpf (n = 3). Results are shown as the relative expression to β-actin (mean ± SD). C, D, Microinjection of twist1a and twist1b mRNA induces the appearance of dorsalized embryos and decreases the expression of runx2b. C, The table summarizing the dorsalized embryo features (D1-5/no defect) of zebrafish injected with each mRNA. Dorsalized (D) phenotype annotations were described in ref. 21. D, Zebrafish were microinjected with or without (WT) twist1a or twist1b mRNA and quantitative RT-PCR for T1 runx2b and T2 runx2b was performed at 48 hpf (n = 3). Results are shown as the relative expression to β-actin (mean ± SD) and significance was determined by Student’s t-test. (* p<0.05 and ** p<0.01 versus no injection [NI]). Results are shown as the relative expression to β-actin (mean ± SD) and significance was determined by Student’s t-test. (* p<0.05 and ** p<0.01 versus MO-SC).

Figure S2 Morpholino knockdown of twist2 or twist3 induces no changes in runx2b transcription. Zebrafish were microinjected with twist2 and twist3 atgMOs and runx2b expression was analyzed by in situ hybridization at 48 hpf (n = 40 for each). Pictures of lateral (L) and dorsal views (D) are displayed. (TIF)

Figure S3 Morpholino knockdown of twist1b increases runx2b transcription. Morpholino knockdown of twist1b increases runx2b expression in a dose-dependent manner. Zebrafish were microinjected with indicated amount of MO-SC or twist1b atgMO and quantitative RT-PCR for T1 runx2b and T2 runx2b was performed at 48 hpf (n = 3). Results are shown as the relative expression to β-actin (mean ± SD) and significance was determined by Student’s t-test. (* p<0.05 and ** p<0.01 versus MO-SC).

(TIF)

Table S1 PCR primer list. Primers designed to perform quantitative RT-PCR for this study are listed.

(PDF)

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Author Contributions

Conceived and designed the experiments: DCY YHC SCH. Performed the experiments: DCY CCT HJT YFL TFH YHC SCH. Analyzed the data: DCY CCT HJT YFL TFH. Contributed reagents/materials/analysis tools: YHC SCH. Wrote the paper: DCY CCT YHC SCH.

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