Morphogenetic control of zebrafish cardiac looping by Bmp signaling

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ABSTRACT
Cardiac looping is an essential and highly conserved morphogenetic process that places the different regions of the developing vertebrate heart tube into proximity of their final topographical positions. High-resolution 4D live imaging of mosaically labelled cardiomyocytes reveals distinct cardiomyocyte behaviors that contribute to the deformation of the entire heart tube. Cardiomyocytes acquire a conical cell shape, which is most pronounced at the superior wall of the atrioventricular canal and contributes to S-shaped bending. Torsional deformation close to the outflow tract contributes to a torque-like winding of the entire heart tube between its two poles. Anisotropic growth of cardiomyocytes based on their positions reinforces S-shaping of the heart. During cardiac looping, bone morphogenetic protein pathway signaling is strongest at the future superior wall of the atrioventricular canal. Upon pharmacological or genetic inhibition of bone morphogenetic protein signaling, myocardial cells at the superior wall of the atrioventricular canal maintain cuboidal cell shapes and S-shaped bending is impaired. This description of cellular rearrangements and cardiac looping regulation may also be relevant for understanding the etiology of human congenital heart defects.

KEY WORDS: BMP, Wnt, Cardiac looping, Hemodynamics, Zebrafish

INTRODUCTION
The term ‘cardiac looping’ defines a morphogenetic process during vertebrate heart development that brings the developing heart chambers into an approximation of their definitive topographical relationships. This involves the transformation of an initially straight heart tube into a curved and S-shaped loop that is asymmetrically positioned along the L/R body axis (Männer, 2000, 2009; Patten, 1922). In higher vertebrates, cardiac looping comprises several sub-processes (Männer, 2000, 2009): (1) ventricular bending along the mid-sagittal body plane causes the heart tube to acquire a C-shape with a convexity that is most pronounced along the original ventral midline of the heart tube; (2) ventricular rotation/torsion around the longitudinal axis of the heart tube leads to a helical deformation of the ventricular bend and to dextral heart looping (ventricular D-looping) along the L/R body axis (Männer, 2004); (3) displacements of embryonic heart segments along the cranio-caudal body axis due to a shortening of the distance between the venous and arterial poles, and a displacement of the ventricular bend from its original position cranial to the atria towards its definitive position caudal to the atria; and (4) ‘untwisting’ or ‘terminal repositioning’ results in a partial rewinding of the rightward-most rotation/torsion of the heart loop (Männer, 2009; Singleman and Holtzman, 2012).

Given the morphogenetic complexity of this process, it is not surprising that animal models for human congenital heart defects frequently show abnormal heart looping during early embryonic development (Ramsdell, 2005). Yet our understanding of molecular mechanisms involved in cardiac looping still remains fragmentary. For example, signaling by Nodal (Chen et al., 2010; Grimes and Burdine, 2017) or bone morphogenetic protein (BMP) is essential for asymmetric positioning of the heart loop along the L/R body axis (Breckenridge et al., 2001; Chen et al., 1997; Chochon et al., 2007; Ocaña et al., 2017; Smith et al., 2008; Veerkamp et al., 2013; Zhang and Bradley, 1996), but much less is known about how cardiac L/R asymmetry is determined during looping morphogenesis. In mouse and zebrafish mutants with defective L/R asymmetry due to a loss of Nodal signaling, the transformation of a linear heart tube into an S-shaped heart loop is not affected. However, its orientation along the L/R body axis is randomized, with 50% of mutants displaying the normal ventricular bend (D-loop), while the remainder forms an abnormal L-loop (Baker et al., 2008; Brennan et al., 2002). Similarly, it is unknown whether disturbed Bmp pathway signaling directly causes defective looping morphogenesis or whether such defects are secondary to its earlier roles (Breckenridge et al., 2001).

Both, organ-intrinsic and organ-extrinsic factors have been proposed for regulating cardiac looping morphogenesis (Männer, 2000; Taber, 2006). The strongest evidence for the presence of intrinsic factors stems from the observation that isolated embryonic hearts, explanted at linear heart tube stages and cultured in vitro, still acquire C-shaped and S-shaped morphologies (Bacon, 1945; Butler, 1952; Latacha et al., 2005; Manning and McLachlan, 1990; Noël et al., 2013). Pharmacological inhibition of the actomyosin network suggested that cytoskeleton-driven changes in cell shapes and intrinsic cell chirality may drive intrinsic heart tube bending (Latacha et al., 2005; Noël et al., 2013). External factors that impact cardiac looping could be hemodynamic forces exerted by blood flow. In zebrafish, blood flow triggers the elongation of ventricular cells within the outer curvature (Auman et al., 2007). Similar anisotropic cell growth has also been described during cardiac looping in other vertebrates (Kidokoro et al., 2008; Manasek et al., 1972; Shi et al., 2014a; Soufan et al., 2006; Voronov et al., 2004). However, in lung-breathing vertebrates, cardiac looping is not compromised in the absence of blood flow, indicating that hemodynamic forces do not play a significant role (Abu-Daya 1972; Shi et al., 2014a; Soufan et al., 2006; Voronov et al., 2004).
et al., 2009; Fransen and Lemanski, 1988; Manasek and Monroe, 1972). Another extrinsic factor contributing to cardiac looping could be mechanical forces due to restraints by surrounding tissues. In zebrafish, myocardial chamber size increases mainly due to the accretion of second heart field (SHF)-derived cardiomyocyte progenitor cells to both ends of the heart tube (de Pater et al., 2009; Hami et al., 2011; Zhou et al., 2011). Model simulations suggest that a continuous elongation of the heart tube within a tissue-confined space such as the pericardial cavity could ultimately result in the morphological changes observed during heart looping (Bayraktar and Männner, 2014). To functionally understand the contributions of different morphogenetic processes during cardiac looping morphogenesis, tractable genetic models are required.

Here, we describe the process of cardiac S-looping morphogenesis in the zebrafish embryo and identify three distinct cell- and tissue-scale transformations that all contribute to this process. We recognize BMP signaling as an important tissue-intrinsic regulator of cardiac bending at the AVC region of the heart tube. We also find that BMP signaling has a regional patterning that correlates with bending of the nascent heart tube and is maintained in the absence of Nodal signaling. These results provide novel insights into the molecular control of cardiac looping morphogenesis in the zebrafish, with relevance for a more comprehensive understanding of human congenital heart defects.

RESULTS
Zebrafish cardiac morphogenesis involves early and advanced S-looping phases
In zebrafish, cardiac looping morphogenesis becomes apparent between 30 and 54 h post fertilization (hpf). To analyze cardiac morphologies during looping stages, we used the transgenic reporter line Tg(myl7:EGFP)^pxu34, which labels myocardial cells, and extracted hearts at 30, 36, 42, 48 and 54 hpf (Fig. 1A-E). Based on two-dimensional projections of confocal z-scans of these hearts (Fig. 1A-E), cardiac looping morphologies can be divided into an ‘early S-looping’ phase, between 30 and 42 hpf (Fig. 1A-C) and an ‘advanced S-looping’ phase, between 42 and 54 hpf (Fig. 1D,E). Early S-looping involves ventricular D-bending along the outer curvature and atrial bending with an outer curvature towards the left. Advanced S-looping involves a cranial shift of the atrium and, as a consequence, the two heart chambers are positioned side-by-side and their convex curvatures become prominent (Fig. 1E). This coincides with an angulation of the heart at the AVC, which is a useful measure for the extent of S-looping (see below). Simultaneously with advanced S-looping, the heart chamber volumes expand anisotropically along their outer curvatures in a process referred to as cardiac ballooning (Bakkers, 2011; Christoffels et al., 2000). In addition, while cardiac looping progresses, the heart shifts from a left-sided body position at heart tube stages (around 30 hpf) to a cranial-ventral position at advanced S-looping stages (>48 hpf). At ballooning stages (54 hpf), the two heart chambers, ventricle on the right and atrium on the left, are positioned in a single plane perpendicular to the mid-sagittal body plane (Fig. 1F). Hence, the deformation of the linear heart tube into a S-shaped loop resembles ‘planar bending’ of the two heart chambers into opposite directions.

Cardiac S-looping coincides with planar bending, torsional winding and anisotropic ballooning
These findings suggest that the S-shaped morphology of the embryonic zebrafish heart results from at least two independent processes: planar bending and anisotropic ballooning of both chambers. To further characterize these morphogenetic processes, we developed quantitative methods to: (1) precisely measure the extent of planar bending; (2) analyze regional changes in shape, size and position of myocardial cells that may contribute to S-looping; and (3) quantify the process of cardiac torsion. This analysis would also clarify whether torsion of the heart, similar to cardiac looping in lung-breathing vertebrates, may also occur in zebrafish. First, we analyzed the extent of cardiac planar bending during S-looping stages based on two independent morphometric measurements. Shown is the looping angle (α) (as defined in the Materials and Methods section) (Fig. 1G). Using this method, we measured a looping angle of 85±4° for wild-type hearts at 54 hpf, which shows that the midline of the heart bends almost at a right angle (Fig. 1G,H; Table S1). As a second parameter for cardiac bending, we compared the distance between atrial and ventricular apex (b) with the entire length of the heart (a) (see scheme in Fig. II for a more detailed explanation). The index value (a/b) decreases inversely proportional to the extent of chamber overlap. Thus, the index value (a/b) decreases while S-looping progresses. By this method, we measured a looping overlap of 2.2±0.2 for wild-type hearts at 54 hpf (Fig. 1J; Table S1). Hence, the index value (a/b) correlates with the shortening of the total length of the heart (a) relative to the increasing region of chamber overlap (b).

In lung-breathing vertebrates, cardiac looping involves a rotation/torsion of the heart tube along its longitudinal axis, which is responsible for asymmetric positioning of the ventricular bend along the L/R body axis and for its helical deformation (Männer, 2000, 2009; Taber et al., 2010; Voronov et al., 2004). Torsion may also contribute to looping of the zebrafish heart (Weber et al., 2017). Hence, we performed multi-color mosaic labeling of cardiomyocytes combined with high-resolution 4D confocal microscopy to assess the exact three-dimensional changes that occur during zebrafish cardiac looping morphogenesis. To unambiguously identify cardiomyocytes, fertilized eggs carrying the transgenic Tg(myl7:EGFP)^pxu34 reporter were injected with a mixture of myl7:TagRFP-T and myl7:TagBFP constructs (Staudt et al., 2014). This resulted in a mosaic expression of fluorescently colored cardiomyocytes due to random combinations and varying expression levels of the three fluorescent proteins. Based on this approach, we identified cell positions and morphologies of individual cardiomyocytes, which were monitored in vivo during cardiac advanced S-looping stages (40-54 hpf) (Fig. 1K, top panel). To quantify torsional winding of the heart tube, we assessed whether mosiacally labelled cross-section planes derived from high-resolution reconstructions of stacks of confocal z-scans were angularly shifted relative to other section planes over time along the heart tube (Fig. 1K, bottom panel; see color-coded arrowheads and dotted lines, top panel). In total, three individual section planes with an even distribution along the heart were selected in each chamber (Fig. 1K, top panel). First, all cross-section planes at 40 and 54 hpf stages were normalized according to their respective geometric orientation at 48 hpf (Fig. S1A); this provided the necessary spatial reference for determining angular shifts (ρ) within each section plane. Within the selected section planes, we identified individual cells at different time points (40, 48 and 54 hpf) by taking advantage of their mosaic color labeling (Fig. S1B) and determined the angular position of each of these uniquely labeled cells relative to the midline (Fig. S1C) (cells that were not identified at each of the three time points were not considered in this analysis). Then angular shifts for a specific time window (40-48 hpf, 48-54 hpf and 40-54 hpf) were calculated by subtracting the angular positions (θ) of all cells for t2-t1 (with t1=earlier stage) (Fig. S1D; Fig. 1L with a representative angular shift measurement of cross-section plane 1 between 40 and 54 hpf).
Finally, we compared the angular shifts between individual section planes to determine whether a torsional winding had occurred within the heart tube. Between 40 and 48 hpf, the angular shifts within all section planes were less than ±10°, indicating only minor torsional shifts within the heart at the beginning of advanced S-looping phase [Fig. 1M; mean±s.d., sample size (Table S2A) and statistical analysis (Table S2B)]. However, at the end of advanced S-looping phase between 48 and 54 hpf, a strong rightward shift relative to the rest of the heart tube of −34.7° and −51° occurred mainly in the ventricular region close to the outflow tract (planes 5 and 6, respectively) (Fig. 1N,P, red arrows; Table S2A,B). In comparison, section planes at the lower atrium (planes 1, 2) had only a minor leftward torsional movement, with positive angular shifts of +3.5° and +8.8°, respectively. Similarly, section planes closest to the AVC (planes 3, 4) had only low negative angular shifts of −4.6° and −1.5°, respectively. Fig. 1O shows the angular shifts between 40 and 54 hpf (Fig. 1O; Table S2A,B). Hence, between 48 and 54 hpf, zebrafish S-looping involves a torsional winding within the heart tube that is particularly strong within the region of the ventricle closest to the outflow tract.
In order to assess the impact of cellular dynamics and morphologies on S-looping morphology, we combined the multi-color mosaic-labeling approach with high-resolution live imaging at 32, 40, 48 and 54 hpf (Fig. 2A-D; Fig. S2). This approach allowed us to analyze cell morphologies and positions of individual cardiomyocytes during these cardiac looping stages. In contrast to a more conventional subdivision of the heart chambers into an inner curvature (IC) comprising the regions neighboring the AVC and an outer curvature (OC) comprising the residual regions of atrium and ventricle, we used a more detailed subdivision of the heart to characterize regional myocardial cell morphologies (Fig. 2E). Accordingly, heart chambers were subdivided along their R-L and cranial-caudal axes, which resulted in nine regions (Fig. 2E-L). To assess cell morphology changes, myocardial cell surface area and circularity were quantified using Fiji software. An extensive comparison of these morphological parameters revealed highly dynamic changes of cell morphologies with regional characteristics during cardiac looping morphogenesis (Fig. 2M,N; statistical analysis in Tables S3 and S4, respectively). Ventricular cardiomyocytes were generally smaller and more circular, with fewer regional differences than atrial cardiomyocytes. In general, regional differences, both between chambers and within the atrium, were more significant from 48 hpf. At 48 and 54 hpf, cells located on the left side of the atrium (LLA+LUA) were larger and more circular than cells located within the corresponding right atrial regions (RLA+RUA). Myocardial cells close to the inflow tract (regions RLA+LLA) acquired highly elongated shapes, whereas cells in the left upper atrium (LUA) acquired large and rounded shapes. These dynamic regional morphologies correspond with regionally anisotropic ballooning of the heart chambers, which contributes to the S-looped heart morphology.

Taken together, S-looping morphogenesis in zebrafish coincides with planar bending of the heart tube, a torsional deformation that is particularly strong within the proximal part of the ventricle, and an anisotropic ballooning of the heart chambers corresponding with regionally confined changes in cardiomyocyte morphology.

Fig. 2. Anisotropic growth of myocardial cells contributes to asymmetric ballooning of the heart chambers. (A–D) Reconstructions of confocal z-scan images taken from a time series (32-54 hpf). Individual cell size and shape changes during cardiac looping stages were quantified based on in vivo multi-color mosaic-labeling by 1- to 4-cell stage plasmid injections into Tg(myl7:EGFP)twu34 transgenic zebrafish (false-colored red). (E) Schematic of different heart chamber regions at 54 hpf. RUV, right upper ventricle; LUV, left upper ventricle; RLV, right lower ventricle; LLV, left lower ventricle; AVC, atrio-ventricular canal; RUA, right upper atrium; LUA, left upper atrium; RLA, right lower atrium; LLA, left lower atrium. (F–L) Regional differences in myocardial cell surface areas and morphologies at 54 hpf. (M,N) Quantifications of dynamic changes in (M) myocardial cell surface area (values are shown relative to the RLV region at 32 hpf) and (N) myocardial cell circularity between 32 and 54 hpf. Kruskal–Wallis test was used for multiple comparisons between the different cardiac regions (statistical analysis in Tables S4 and S5). Data are mean±s.d.; *P≤0.05; **P≤0.01; ***P≤0.001; ****P≤0.0001. Scale bars: 50 μm in A-D; 10 μm in F-L.
Bmp signaling activity is asymmetric during S-looping morphogenesis

To elucidate molecular regulatory mechanisms that control cardiac S-looping, we first focused on the role of Bmp signaling due to its well-established role in controlling cell morphological changes that contribute to organ morphogenesis (Jidigam et al., 2015; Widmann and Dahmann, 2009). First, we analyzed the spatial-temporal activity of Bmp signaling within hearts between 24 and 54 hpf using the transgenic \( Tg(BRE-AAVmlp:dmKO2)^{mw40} \) Bmp reporter line (Collery and Link, 2011) [referred to as \( Tg(BRE:dmKO2)^{mw40} \)]. This reporter expresses destabilized monomeric Kusabira-Orange 2 (dmKO2) under the control of a Bmp-responsive element (BRE), which contains multiple binding sites for phosphorylated Smad (P-Smad) proteins. A C-terminal PEST domain mediates rapid degradation of dmKO2, which enables a precise monitoring of Smad-dependent Bmp pathway activity. For a time course analysis of Bmp signaling activity during cardiac looping stages, we imaged double-transgenic \( Tg(BRE:dmKO2)^{mw40} \) (false-colored gray) and \( Tg(kdrl:EGFP)s843 \) (false-colored red) hearts at 24, 30, 36, 42, 48 and 54 hpf (Fig. 3A-F; Fig. S3). At early heart stages (24 and 30 hpf), Bmp signaling activity was higher in endocardium compared with myocardium (Fig. 3A,B; Fig. S3A-B).
asterisks; Fig. S3C-F.

Throughout cardiac looping stages (36-54 hpf), Bmp activity was more pronounced within the superior compared with the inferior AVC, which created an asymmetric pattern of activity (Fig. 3C-F, asterisks; Fig. S3C-F’). These observations raised the intriguing possibility that asymmetric Bmp activity might play a role in the regulation of zebrafish S-looping. The asymmetry of Bmp signaling activity within the superior AVC myocardial tissue was verified by immunolabeling against P-Smad1/5/8 (Fig. S4A,C).

However, with respect to its onset, an asymmetric P-Smad1/5/8 immunostaining at the future superior AVC region within myocardial tissue was observed as early as at the linear heart tube stage (30 hpf) (Fig. S4A), which preceded the asymmetric expression of the Bmp reporter activity by 6 h (Fig. 3C). To evaluate the specificity of Bmp signaling activity indicated by the P-Smad1/5/8 immunostaining, we carried out pharmacological inhibition with 20 μM LDN193189 treatment between 24 and 30 hpf (Fig. S4B) and 24 and 54 hpf (Fig. S4D), and found that the immunostaining signal strongly decreased compared with the 0.2% DMSO-treated control hearts (Fig. S4A,C). Hence, Bmp signaling activity is stronger at the superior compared with the inferior AVC during S-looping stages.

To further analyze the involvement of Bmp signaling in heart looping, we used the two specific small molecule inhibitors [LDN193189 (Cuny et al., 2008; Dietrich et al., 2014) and K02288 (Sanvitale et al., 2013)] between 30 and 54 hpf (Fig. 3G-I). Both compounds interfere with Bmp signaling by preventing the activation of the Smad-dependent pathway via Bmp type I receptors. Indeed, both inhibitors caused a significant reduction of the relative dmKO2 fluorescence intensity levels within the superior or inferior myocardial AVC when compared with DMSO-treated control hearts (superior AVC: control, 100±32%; LDN193189, 71±25%; K02288, 57±28%; inferior AVC: control, 74±32%; LDN193189, 45±11%; K02288, 42±15%) (Fig. 3F; sample size and statistical analysis in Table S5). Next, we assessed the extent of S-looping after Bmp signaling inhibition by quantifying the looping angle (α) (LDN193189, 105±20°; K02288, 98±16°) and index a/b (LDN193189, 4.3±2.6; K02288, 3.3±1.7) compared with control hearts (control, 77±12°; control, 2.2±0.7, respectively) (Fig. 3K,L; sample size and statistical analysis in Table S1). In comparison with DMSO-treated control hearts, the looping angles in LDN193189- or K02288-treated hearts were significantly higher, which shows that these Bmp signaling-inhibited hearts were more linear (Fig. 3H,I,K; Table S1; Fig. S4D). Consistent with this finding, the index a/b was significantly increased in LDN193189- or K02288-treated hearts, which also implies a reduced lateral overlap of the heart chambers and thus a more linear heart morphology (Fig. 3H,IL; Table S1; Fig. S4D). Thus, abnormal S-looping may be affected by abnormal planar bending.

The essential role of Bmp in this morphogenetic process was demonstrated by higher Bmp reporter signaling at the superior AVC myocardium that coincided with the onset of S-looping and by a failure of correct S-looping upon pharmacological Bmp signaling inhibition. This also suggested an involvement of Bmp activity in planar bending of the heart tube, as indicated by the more linear morphology of Bmp signaling-inhibited hearts. To analyze whether Bmp promotes planar bending by driving cell morphology changes, we used an antibody against the cell-adhesion protein ALCAM to outline cell borders and closely examined the cellular morphologies of superior and inferior AVC myocardial cells in control embryos and in embryos treated with LDN193189 (Fig. 3M-N’). Single confocal z-scan images at the AVC revealed that 54 hpf myocardial cells at the superior AVC have a conical shape and are bottlenecked (Fig. 3M, M’), which may contribute to the constriction of the AVC and facilitate bending of the heart tube. In contrast, hearts treated with 20 μM LDN193189 between 30 and 54 hpf had more cuboidal shapes (Fig. 3N, N’), which is similar to myocardial cell shapes present at earlier stages (Fig. S5A,A’).

At 54 hpf, myocardial cells at the inferior AVC had cuboidal shapes in both control (Fig. 3M,M’) and LDN193189-treated hearts (Fig. 3N,N’). Hence, Bmp inhibition caused reduced bending of the heart and resulted in more cuboidal myocardial cell morphologies at the superior AVC that resemble those cell morphologies present at the onset of S-looping when the Bmp signaling asymmetry initiates at the superior AVC.

In a complementary approach, we assayed whether misregulation of Bmp signaling during S-looping morphogenesis causes any planar-bending defects (Fig. 3O-T). To downregulate Bmp signaling, we clonally overexpressed the secreted Bmp antagonist Noggin3 within myocardial tissue (Fig. 3O) or overexpressed Noggin3 from the stable Tg(hsp70I:Nog3)1/14 (Chocron et al., 2007) transgenic line of zebrafish by heat shock at 32 hpf (Fig. 3P). We found that these conditions had a stronger effect on planar bending compared with the pharmacological inhibition of Bmp signaling. The stronger effect of Noggin3 clonal overexpression (Fig. 3O) in comparison with pharmacological (Fig. 3H,I) or genetic (Fig. 3P) inhibition of Bmp signaling could be due to the fact that myocardial clones are expressing Noggin3 from the onset of cardiac morphogenesis. Instead, the timing of the Bmp inhibition treatment between 30 and 54 hpf, and of the heat shock at 32 hpf, may have had less of an impact on the process of cardiac looping. Overexpression of the Bmp2b ligand was induced from the stable heat-shock promoter-driven transgene Tg(hsp70: Bmp2b)1/13 (Chocron et al., 2007) at 32 hpf (Fig. 3Q). In all cases, hearts analyzed at 54 hpf were abnormally looped (Fig. 3O-Q), as quantified by S-looping parameters (Fig. 3S,T). For clonally overexpressed Noggin3, it was also possible to measure the relative dmKO2 fluorescence intensities showing that Bmp signaling was reduced at both superior and inferior AVCs (Fig. 3R; sample size and statistical analysis in Table S5) (superior AVC: control, 100±34%; Nog3 clonal overexpression, 91±9%; inferior AVC: control, 66±32%; Nog3 clonal overexpression, 10±8%). Looping angles increased in all three conditions (Nog3 clonal overexpression, 149±8°; hs:Noggin3, 137±4°; hs:Bmp2b, 130±4°) in comparison with control hearts (73±10°) (Fig. 3S; sample size and statistical analysis in Table S1). The index a/b increased in all conditions when compared with control hearts, which also suggested defective bending and a more linear heart upon loss of Bmp signaling (Nog3 clonal overexpression, 14±8; hs:Noggin3, 9±8; hs:Bmp2b, 5.9±0.8; control, 1.9±0.4) (Fig. 3T; sample size and statistical analysis in Table S1). A closer examination of the AVC regions revealed that these cardiomyocytes had abnormal cell shapes under conditions of downregulation or overactivation of Bmp signaling (Fig. SS5B-C’), which is consistent with its role in endocardial cell proliferation and morphogenesis.
Establishment of asymmetric Bmp signaling within the AVC is not dependent on Nodal or Wnt signaling

Cardiac looping morphogenesis can occur in zebrafish embryos with defective L/R asymmetry (Burdine and Schier, 2000). This raises the issue of whether and how asymmetric Bmp signaling at the AVC is initially established. We therefore analyzed developmental pathways that determine the general body plan of the embryo, including Nodal and Wnt signaling. First, we downregulated Nodal signaling by injection of an antisense oligonucleotide morpholino (MO) against the primary cardiac Nodal ligand Southpaw (Spaw), which is a key regulator of cardiac L/R asymmetry (Long et al., 2003; Rebagliati et al., 1998; Smith et al., 2011; Veerkamp et al., 2013). Knockdown of Nodal signaling caused a randomized looping of the heart in the right-left (R-L) embryonic body axis with D-looped (54%) (Fig. 4A) and L-looped (27%) (Fig. 4B) hearts, and an abnormal looping along the dorsal-ventral (D-V) embryonic body axis with V-looped (19%) hearts (Fig. 4C) (n=83 embryos). Bmp signaling activity was in an asymmetric pattern that strictly correlated with the orientation of S-looping (Fig. 4D,E) when hearts were looping along the R-L axis (D- and L-looped hearts). Despite a lack of Nodal signaling, these hearts looped to a degree comparable with wild type (D-looped heart, 2.3±0.4; L-looped heart, 2.4±0.4) (Fig. 4G; Table S1). For the third group of hearts that were looping within the D-V axis (Fig. 4C,F), Bmp activity was present either on both sides of the midline or with an D/V asymmetry (Fig. 4F,G; Table S1; V-looped heart index a/b, 5.4±0.7). Taken together, the asymmetric expression of Bmp within the embryonic heart is independent of Nodal activity and strictly correlates with the looping orientation of D- and L-looped hearts.

To downregulate the Wnt signaling pathway, which is essential for the establishment of the embryonic cranial-caudal body axis (Yamaguchi, 2001), embryos were treated between 30 and 54 hpf with 10 µM of the Wnt antagonist IWR-1, which stabilizes Axin1 (thereby promoting the degradation of β-catenin and inhibiting the activation of Wnt target genes) (Nusse, 2005) (Fig. 4H-L). As a proof of efficacy, during cardiac valve leaflet formation at 72 hpf, 10 µM IWR-1 was sufficient to inhibit Wnt signaling, which was observed in transgenic embryos of the Wnt reporter line Tg(7xTCF-Xla.Siam:nls mCherry)ia5 (data not shown). Subsequently, the cardiac Tg(BRE:dmKO2)mvo40 Bmp signaling reporter activity was quantified at 54 hpf (Fig. 4L). Upon Wnt signaling inhibition, cardiac S-looping at 54 hpf was not affected (Fig. 4L-K). Quantifications of planar bending in IWR-1-treated hearts based on the looping angle (α) or the index a/b did not reveal significant differences in comparison with DMSO-treated controls (Fig. 4J,K; mean±s.d, sample size and statistical analysis in Table S1). In addition, upon IWR-1 treatment, Bmp signaling based on dmKO2 fluorescence intensity levels showed no significant changes within superior AVC myocardial cells and only a slight reduction within inferior AVC myocardium (P≤0.05) (Fig. 4L; mean±s.d, sample size and statistical analysis in Table S5). Altogether, we conclude that downregulation of Wnt signaling by IWR-1 treatment from 30-54 hpf did not affect S-looping morphogenesis or the asymmetry of Bmp signaling activation within the superior AVC.

Establishment of asymmetric Bmp signaling within the AVC is dependent on cardiac contractility

Because the onset of a heart beat and blood flow coincides with the initial stages of cardiac looping, we next analyzed whether changes in hemodynamic forces or cardiac contractility affect cardiac looping. Both Gata1 and Gata2 are transcription factors required for hematopoiesis in zebrafish and their knockdown results in abnormal hemodynamic forces (Galloway et al., 2005). Knockdown of Gata1 shifts erythropoiesis towards myelopoiesis (Galloway...
et al., 2005), which results in a complete absence of red blood cells (Lyons et al., 2002; Vermot et al., 2009) and reduces blood viscosity by 90% (Vermot et al., 2009). Loss of Gata2 causes a reduction in blood cell numbers (Dietrich et al., 2014; Galloway et al., 2005; Tsai and Orkin, 1997; Vermot et al., 2009). As a consequence, blood viscosity in zebrafish embryos is reduced by 70% and the retrograde flow fraction is strongly reduced (Vermot et al., 2009). Combined loss of Gata1 and Gata2 causes a complete loss of red blood cells but a milder reduction of the retrograde flow fraction when compared with gata2 morphants (Dietrich et al., 2014; Vermot et al., 2009). Cardiac TroponinT2a (Tnnt2a) is essential for sarcomere assembly and cardiac contractility (Huang et al., 2009; Sehnert et al., 2002). To assess the impact of reduced shear stress and retrograde flow fraction, or of cardiac contractility, on cardiac S-looping and on the asymmetric activation of Bmp signaling within the superior AVC, we carried out individual or combined injections of MOs against Gata1, Gata2, Gata1/2 or Tnnt2a (Fig. 5A–E). Quantification of the relative dmKO2 intensity at 54 hpf within both superior and inferior AVC myocardium revealed significant differences in Bmp reporter activity only for the condition with a lack of Tnnt2a (Fig. 5F); mean±s.d., sample size and statistical analysis in Table S5; superior AVC: control, 100±30%; MO Gata1, 59±27%; MO Gata2, 79±21%; MO Gata1/2, 60±7%; MO Tnnt2a, 47±12%; inferior AVC: control, 56±4%; MO Gata1, 31±13%; MO Gata2, 39±11%; MO Gata1/2, 34±7%; MO Tnnt2a, 42±14%). Hence, Bmp asymmetric activity within the AVC is significantly reduced in the absence of heart contractility and not simply by a reduction of shear stress or retrograde flow patterns. Quantification of cardiac S-looping parameters, revealed a significant increase of the looping angle (α) (Fig. 5G; sample size and statistical analysis in Table S1; control, 82±5°; MO Gata1, 95±10°; MO Gata2, 103±13°; MO Gata1/2, 112±19°; MO Tnnt2a, 135±24°) and of the index a/b (Fig. 5H; sample size analyzed and statistical analysis are in Table S1; control, 2.1±0.2; MO Gata1, 3.1±0.3; MO Gata2, 3.4±0.7; MO Gata1/2, 4.6±1.6; MO Tnnt2a, 8.0±4.2) under conditions of loss of Gata1/2 or Tnnt2a. This also correlated with changes in S-looping morphology (Fig. 5D,E). The effects of a lack of cardiac contractility were further tested in TropomysinT2a (tnnt2a)β/10 mutations (Sehnert et al., 2002). At 54 hpf, mutant hearts were not looped and P-Smad1/5/8 levels were reduced in myocardial AVC cells (Fig. S6B,B’). This correlated with an increase in the looping angle (α) (tnnt2a mutant, 125±15°; control, 93±8°) and in the index a/b (tnnt2a mutant, 9.6±3.5; control, 2.1±0.3) (Fig. S6D,E; sample size analyzed and statistical analysis are in Table S1). This also correlated with a loss of asymmetric BMP signaling activity within the superior AVC (Fig. S6C, sample size analyzed and statistical analysis are in Table S6). In summary, alterations of retrograde flow patterns result in S-looping defects that are not caused by obvious changes in Bmp signaling at the AVC. However, as BMP is only weakly and homogeneously active in tnnt2a morphant and mutant hearts, cardiac contractility is apparently an important modifier of its activity.

DISCUSSION

As discussed by Männner (2000, 2009) and Le Garrec and colleagues (2017), the definition of cardiac looping has differed in a number of studies (Le Garrec et al., 2017; Männner, 2000; Männner, 2009).

Fig. 5. Cardiac contractility is required for Bmp signaling at the superior AVC. (A–E) Reconstructions of confocal z-stacks of hearts at 54 hpf in (A) wild-type, (B) gata1 morphant, (C) gata2 morphant, (D) gata1/2 double morphant or (E) tnnt2a morphant embryos. Myocardial tissue is marked by expression from the Tg(my7:EGFP)Tub09 reporter (false-colored red) and Bmp reporter Tg(BRE:dmKO2)Mw40 (false-colored gray). (A–E) Bmp reporter. (A–E) Bmp reporter Tg(BRE:dmKO2)Mw40 expression. The outline of the heart is delineated by dotted lines. Asymmetric Bmp signaling activity within the superior AVC is marked by yellow asterisks. (F–H) Quantifications of (F) relative dmKO2 intensities at the superior (Sup.) and inferior (Inf.) AVC (data in Table S5), (G) looping angle (α) (data in Table S1), and (H) index a/b (data in Table S1) were measured in gata1, gata2, gata1/2 and tnnt2a morphant versus wild-type hearts. Under abnormal hemodynamic conditions (gata1, gata2 and gata1/2 morphant hearts) the relative dmKO2 intensity is not significantly affected. However, asymmetric Bmp signaling activity is lost in the absence of cardiac contractility. The looping angle (α) (G) and the index a/b (H) are significantly increased in gata1/2 and tnnt2a morphant versus control hearts (statistical analysis in Table S1). In G,H, the limits of the boxes indicate the range between the first quartile (25th percentile) and the third quartile (75th percentile). The line inside box indicate the median value. The error bars indicate the maximum and minimum values. Data are mean±s.d. in F; ns, not significant; *P≤0.05; **P≤0.01; ***P≤0.001. Scale bar: 50 μm.
Therefore, a clear description of the morphological changes and the cellular mechanisms that contribute to zebrafish cardiac looping was needed. Here, we have characterized S-looping morphogenesis in the zebrafish and describe three cell- or tissue-scale morphogenetic processes that all contribute to the deformation of the linear heart tube (Fig. 6): (1) zebrafish S-looping involves a torque-like winding deformation mainly at the arterial pole of the early heart tube; (2) anisotropic growth of ventricle and atrium along their outer curvatures contributes to the emergence of the S-shaped morphology of the zebrafish heart; and (3) Bmp signaling is an intrinsic signaling pathway that controls cell shape changes at the superior myocardial wall of the AVC, which determines bending of the heart tube.

Some of these cellular processes are remarkably similar to morphogenetic processes within higher vertebrates. Rotation of the outflow tract region has been observed in the chick (Thompson et al., 1987) and mouse embryo (Meilhac et al., 2004; Bajolle et al., 2006). Anisotropic growth of cardiomyocytes during cardiac looping has also been described in higher vertebrates (Kidokoro et al., 2008; Manasek et al., 1972; Meilhac et al., 2004; Shi et al., 2014a; Soufan et al., 2006; Voronov et al., 2004). In zebrafish, blood flow is an important trigger that causes the elongation of ventricular cardiomyocytes within the outer curvature. Heart contractility has some role in restricting the extent of their elongation (Auman et al., 2007). This balanced interplay between contractility as a cell-intrinsic force and blood flow as an extrinsic force demonstrates the interdependence of these forces in shaping the developing heart (Auman et al., 2007; Deacon et al., 2010). Our data extend these previous studies and provide a dynamic multi-color mosaic labeling-based view of this process over the course of cardiac looping. This analysis also reveals that regionally confined anisotropic growth of cardiomyocytes contributes to chamber morphogenesis in a way that contributes to the cardiac S-shaped morphology.

Several lines of evidence suggest that Bmp signaling acts as an intrinsic signaling pathway during cardiac S-looping morphogenesis. Unlike Nodal, which is not expressed within the mouse heart tube at looping stages (Vincent et al., 2004), we find that, in zebrafish, Bmp signaling has an asymmetric activity within the heart that strongly correlates with looping orientation and is not dependent on the Nodal ligand Spaw. Similar patterns of Bmp expression are also present in the hearts of higher vertebrates (Jiao et al., 2003; Somi et al., 2004) and bmp4 is asymmetrically expressed within the zebrafish heart at the cone stage prior to the formation of a heart tube (Chen et al., 1997; Smith et al., 2008; Somi et al., 2004; Veerkamp et al., 2013). However, to date the role of Bmps in cardiac looping morphogenesis beyond their well-established role in L/R asymmetry within the embryo (Breckenridge et al., 2001; Chen et al., 1997; Chocron et al., 2017; Smith et al., 2008; Veerkamp et al., 2013; Zhang and Bradley, 1996) and in the formation of the AVC region.

![Fig. 6. Model of zebrafish cardiac looping.](image)

(A) Summary of cellular processes that contribute to zebrafish S-looping morphogenesis: (1) BMP-dependent planar bending (white line at heart midline) of the heart tube due to cuboidal-to-conical cell shape changes of superior AVC cardiomyocytes (yellow triangles); (2) myocardial anisotropic ballooning due to region-specific cell morphology changes causes an expansion of the two cardiac chambers (white arrowheads); and (3) ventricular rightward-torsional winding (twisted arrows at ventricular outflow tract) is most pronounced at the cranial pole of the heart. Bmp signaling (false-colored gray) with strong activation at myocardial superior AVC, myocardium (yellow dotted lines) and endocardium (false-colored red). (B) Summary of cell-intrinsic and -extrinsic factors involved in cardiac planar bending of the heart tube. Cardiac contractility is essential for a strong Bmp signaling pathway activation within myocardial cells at the superior AVC region. This Bmp activation induces myocardial changes from cubical to conical cell shapes and results in an S-planar bending of the heart tube. In addition, hemodynamic forces (shear stress and retrograde flow fraction) affect Bmp signaling-independent planar bending. (C) During zebrafish cardiac looping morphogenesis, two separate processes occur in parallel: (1) planar S-looping along the mid-sagittal body axis and (2) lateral looping during which the ventricle is positioned towards the right and the atrium towards the left (D-looping). The mid-sagittal plane is in grey; original right and left sides of the heart are marked in blue and yellow, respectively. Under some experimental conditions, inverted L-looping can also occur. In comparison with lung-looping during which the ventricle is positioned towards the right and the atrium towards the left (D-looping). The mid-sagittal plane is in grey; original right and left sides of the heart are marked in blue and yellow, respectively. Under some experimental conditions, inverted L-looping can also occur.
principle be driven by the same fundamental biophysical forces in lower and higher vertebrates. Thus, understanding cardiac development in zebrafish, at a cellular resolution and with high molecular precision, will also be essential for a better understanding of human cardiac heart defects.

**MATERIALS AND METHODS**

**Zebrafish lines**

Zebrafish were kept according to standard laboratory procedures (Westerfield, 2007). Handling of zebrafish was carried out in compliance with German and Berlin state law, carefully monitored by the local authority for animal protection (LaGeSo, Berlin-Brandenburg, Germany and LANUV, Lower-Saxony, Germany). We used the following zebrafish mutant and transgenic lines: tnnt2a-lacZ (Sehnert et al., 2002), Tg(kdr:EGFP)jsw1 (Jin et al., 2005), Tg(myl7:EGFP)sm4 (Huang et al., 2003), Tg(BR-B;ATG)tnnt2a3 (Collary and Link, 2011), Tg(hsp70l:tdTomT)icos (Chocron et al., 2007) and Tg(hsp70:Bmp2b)jsw1 (Chocron et al., 2007).

**Antisense oligonucleotide morpholinos**

Knockdown studies were performed by MO injection as described previously (Nasevicius and Ekker, 2000). The following antisense oligonucleotide morpholinos (Gene Tools were used: gata1a_ATG MO, 5′-CTGCAAGTGTGAATGAGATGTC-3′ (Galloway et al., 2005); gata2a_3ex_in MO, 5′-CATCCTACTACCTGCTCCTTTCG-3′ (Galloway et al., 2005); spaw_ATG MO, 5′-GACGCTATGCTAGCTGCTGATGG-3′ (Long et al., 2009); and MO1-tnnt2a, 5′-CATGTTGCTGCTGCTGACACCGA-3′ (Sehnert et al., 2002).

Morpholino solution (1 nl) was injected into the yolk at the following concentrations: 1 mM MO spaw, 1 mM MO gata1, 0.2 mM MO gata2 and 0.1 mM MO-tnnt2a.

**Pharmacological treatments**

For pharmacological treatments, embryos were incubated in egg-water supplemented with 20 μM LDN193189 trichloride (Axon Medchem, Axon 1509), 20 μM K02288 (Sigma-Aldrich, SML1307) or 10 μM 1WR-1 (Sigma-Aldrich, I0161). If not stated otherwise, embryos were treated between 30 and 54 hpf at 28.5°C. Control embryos were incubated in egg-water containing an equal amount of DMSO.

**Gene overexpression via heat shock**

For heat-shock-induced overexpression of a gene of choice, transgenic embryos carrying the respective gene under control of the heat-shock promoter hsp70l were transferred to a 2 ml reaction tube. Heat-shock was induced at 32 hpf by addition of pre-warmed egg-water of 38°C and incubated at 38°C with shaking at 300 rpm for 45 min. After heat-shock, embryos were transferred to petri dishes filled with egg-water and incubated at 28.5°C until 54 hpf. Efficacy of the heat-shock-induced overexpression was monitored by expression of GFP within the entire embryo.

**Immunohistochemistry**

Embryos of the desired developmental stage were dechorionated, anesthetized with 0.16 mg/ml tricaine and fixed in 4% paraformaldehyde overnight at 4°C. Embryos were rinsed four times with phosphate-buffered saline with Tween 20 (PBST) for 10 min and blocked for 2 h at room temperature in PBST supplemented with 10% normal goat serum (NGS). Subsequently, embryos were incubated with the primary antibody mouse anti-Alcam (1:100); Developmental Studies Hybridoma Bank, zm-8) at 4°C overnight. After washing with PBST containing 5% NGS, fluorescently conjugated secondary antibody goat anti-mouse DyLight649 (1:200; Jackson Immuno Research, 115-495-003) was incubated overnight at 4°C and washed as described before. The pSmad-1/5/8 labeling was performed as described by Lenhart et al. (2013) using the following antibodies: rabbit anti-pSmad-1/5/8 (1:100; Cell signaling Technology, 9511), chicken anti-GFP (1:600; Aves labs, GFP-1020), goat anti-rabbit DyLight 649 (1:200, Jackson Immuno Research, 111-495-003) and goat anti-chicken FITC (1:200; Aves labs, F-1005). Dissected hearts were mounted in SlowFade Gold (Invitrogen, S36936) and imaged on a Leica TCS SP8 or a Zeiss LSM
510 META NLO confocal microscope with a 40× objective. Images were analyzed using Volocity 3D Image Analysis Software (Perkin Elmer) or Imaris software (Bitplane).

**Clonal overexpression and placid constructs**

For clonal Noggin3 overexpression within myocardial cells, the Tol2 plasmid myl7:nog3-IRESpGF was co-injected with transposase mRNA in one-cell transgenic Tg(BRE-Azmmlp:dmKO2)5mo1 Bmp reporter embryos. The myl7:nog3-IRESpGF expression vector was Gateway cloning (Invitrogen, 12537-103) using the following primers for nog3 (GenBank Accession Number AL935268; nog3_1 Thr1_fwd primer, 5'-GGGACCAAAGTGTACACAAAAAACGAGGCTCATTGATGAAACATCCCGATATTCC-3' and nog3_1 Thr2_rev primer, 5'-GGGGAGCACCTTTGTCACAGAAGAGCT-GGGTTCAGTCCGCGGAGGC3').

**Multicolor mosaic-labeling approach**

To unambiguously identify cardiomyocytes, myl7:TagRFP-T and myl7:TagBFP constructs (kindly provided by D. Stainier (Staudt et al., 2014), Max Planck Institute for Heart and Lung Research, Bad Nauheim, Germany) together with transposase mRNA were co-injected into 1- to 4-cell-stage embryos of the transgenic Tg(myl7:EGFP)m144 line. Tol2-mediated transgenesis (Kawakami, 2007) resulted in a clonal expression pattern of these constructs, creating numerous differently colored cardiomyocytes throughout the entire myocardium by random combination as well as different cytoplasmic expression levels of the three fluorescent proteins. High-resolution live imaging of each single heart was performed at regular intervals during cardiac looping development. Before imaging, each live embryo was incubated with 0.16 mg/ml tricaine (3-amino benzoic acid ethyl ester; Sigma-Aldrich, A-5040) and 4 mg/ml of the myosin ATPase inhibitor BDM (2,3 butanediol monoxime; Sigma-Aldrich, B0753) to anesthetize and suppress the heartbeat, and then mounted in 1% low-melting agarose (Lonza, 50081) supplemented with tricaine and BDM. To ensure physiological heart development, embryos were extracted from agarose after imaging each time point to ensure physiological heart development, which results in a change of heart orientation for the next round of imaging. In order to keep the quantification consistent at all time-points, it is important to first orient the images acquired from the hearts before performing any analysis. A customized Matlab-based software was used for orienting the heart at different time points. Briefly, to normalize changes in orientation of the heart at different stages, we selected a reference cross-section plane of the 40 hpf atrium (section plane 2). For each stage, plane number 2 was rotated gradually and we calculated the Pearson correlation coefficient between the 40 hpf reference plane and the rotated reference plane at all other stages. The rotation angle, at which this correlation coefficient is maximum, was selected to adjust all stage-matched section planes.

In the oriented hearts, the angle (θi) of each cell (ci) at the polar coordinate (with the reference point being the cross-section of the centerline of the heart tube with the plane and the reference direction being the right side) was calculated for all the cells contained in each of the section planes (pi) at each time-point (ti). A custom Matlab-based software was designed for this purpose.

The angular shift for each cell is defined as:

\[ \rho_{ij} = \theta_{ji} - \theta_{ij}, \]

where ρ is the angular shift, θ is the angle, ci is the jth cell, pi is the jth plane and ti is the kth time point. A value of ρ = 0 indicates a counterclockwise (leftward) rotation of the jth cell of the kth plane at time ti. If all of the cells of the jth plane have a positive or negative angular shift, it means that the jth ring is rotated.

We define torsion as:

\[ \tau_{nm} = \frac{\sum_{j=1}^{n \cdot m} \rho_{ij} \cdot I_p}{I_p} - \frac{\sum_{j=1}^{n-1} \rho_{ij} \cdot I_p}{I_p}, \]

where \( \tau_{nm} \) is the torsion between planes m and n, \( I_p \) is the number of cells in the nth plane, and \( I_p \) is the number of cells in the nth plane.

**Quantifications of myocardial cell surface area and shape**

To allow assessment of cell morphologies, myocardial cell borders were outlined with an antibody against the cell adhesion protein ALCAM (see Immunohistochemistry section) or measured from multicolor mosaic-labelled hearts. Fiji software (Schindelin et al., 2012) was used to quantify the morphological parameters. Cell surface area and shape were calculated as was described by Dietrich et al. (2014). Cell surface areas are shown relative to the RLV region at 32 hpf. Cell shape values are from 0.1 to 1, where a value of 0.1 characterizes an elongated cell and a value of 1.0 corresponds with a perfect circle.

**Heart torsion calculations**

We selected six cross-section planes of fluorescence projection images (Imaris software, Bitplane) perpendicular to the mid-sagittal plane of the mosaically labelled 48 hpf heart with an even distribution throughout ventricle and atrium. Uniquely identifiable cardiomyocytes within selected cross-section planes were then matched to cross-section planes of earlier and later stages (40 and 54 hpf). The matching rings were selected in a way that they were perpendicular to the centerline of the heart and contained the maximum number of tracked cells. Embryos were extracted from agarose after imaging each time point to ensure physiological heart development, which results in a change of heart orientation for the next round of imaging. In order to keep the quantification consistent at all time-points, it is important to first orient the images acquired from the hearts before performing any analysis. A customized Matlab-based software was used for orienting the heart at different time points. Briefly, to normalize changes in orientation of the heart at different stages, we selected a reference cross-section plane of the 40 hpf atrium (section plane 2). For each stage, plane number 2 was rotated gradually and we calculated the Pearson correlation coefficient between the 40 hpf reference plane and the rotated reference plane at all other stages. The rotation angle, at which this correlation coefficient is maximum, was selected to adjust all stage-matched section planes.

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We define torsion as:

\[ \tau_{nm} = \frac{\sum_{j=1}^{n \cdot m} \rho_{ij} \cdot I_p}{I_p} - \frac{\sum_{j=1}^{n-1} \rho_{ij} \cdot I_p}{I_p}, \]

where \( \tau_{nm} \) is the torsion between planes m and n, \( I_p \) is the number of cells in the nth plane, and \( I_p \) is the number of cells in the nth plane.

**Statistical analysis**

All experiments were performed in at least three independent biological replicates. All data are presented as mean±standard deviation (s.d.). Statistical analysis was carried out using PRISM Graph Pad software. D’Agostino & Pearson omnibus normality test (p=0.05) was used to check for parametric or non-parametric distribution. For parametric distribution, a one-way ANOVA Holm-Sidak test was used. For non-parametric distribution, Kruskal–Wallis or Dunn’s test were used. Significance levels for multiple comparisons were assigned by PRISM based on mean rank differences or mean differences, respectively. ns, not significant; *P≤0.05, **P≤0.01, ***P≤0.001 and ****P≤0.0001.
Morphogenetic control of zebrafish cardiac looping by Bmp signaling


Lenhart, K. F., Holtzman, N. G., Williams, J. R. and Burdine, R. D. (2013). Integration of nodal and BMP signals in the heart requires FoxH1 to create left-right differences in cell migration rates that direct cardiac asymmetricality. PLoS Genet. 9, e1003109. doi:10.1371/journal.pgen.1003109


