Rear traction forces drive adherent tissue migration in vivo

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During animal embryogenesis, homeostasis and disease, tissues push and pull on their surroundings to move forward. Although the force-generating machinery is known, it is unknown how tissues exert physical stresses on their substrate to generate motion in vivo. Here, we identify the force transmission machinery, the substrate and the stresses that a tissue, the zebrafish posterior lateral line primordium, generates during its migration. We find that the primordium couples actin flow through integrins to the basement membrane for forward movement. Talin- and integrin-mediated coupling is required for efficient migration, and its loss is partially compensated for by increased actin flow. Using Embryogram, an approach to measure stresses in vivo, we show that the rear of the primordium exerts higher stresses than the front, which suggests that this tissue pushes itself forward with its back. This unexpected strategy probably also underlies the motion of other tissues in animals.

During development, homeostasis and disease, cells and tissues move to form organs, to seal wounds and to hunt pathogens. To move, cells generate force and interact with their surroundings to pull and push themselves forward. Force transmission from the actomyosin network to the surroundings has been molecularly characterized and precisely measured in cultured cells. Cells use integrin-based adhesion complexes to couple the actomyosin network inside the cells to the substrates outside the cells and pull on their surroundings with forces of around 3–30 pN across molecules. Since many processes are altered when cells are removed from their physiological environment and placed in culture, it is largely unclear whether cells in living animals interact with their surrounding in the same manner and pull on their substrate with similar forces. To address these questions, we use the zebrafish posterior lateral line primordium as a model. The primordium is a tissue of about 140 cells that expresses the chemokine receptor Cxcr4b. It migrates directly under the skin from behind the ear to the tip of the tail and follows a gradient of the chemokine Cxcl12a along the body of the embryo.

Results

The primordium migrates on a basement membrane. To learn about the substrate that the primordium uses to push and pull itself forward, we used transmission electron microscopy (TEM) to inspect transverse sections of embryos (30 h post fertilization) at different locations along the migratory route of the primordium. Consistent with previous studies, we found that the two-layered skin is separated from the underlying muscle by a 200-nm thick basement membrane (BM) in front of the migrating primordium (Extended Data Fig. 1a,e). At the position of the primordium, the migrating tissue separates the skin and the BM, such that the basal side of the primordium is juxtaposed to the BM while there is no BM detectable on its apical side (Fig. 1a and Extended Data Fig. 1b–e). We confirmed these observations by inspecting the localization of the core BM component laminin-γ1 tagged with superfolder green fluorescent protein (LamC1–sfGFP) (Extended Data Fig. 1f,g). During its migration, the primordium wedges itself between the skin and the muscle, and pushes the LamC1–sfGFP-labelled BM towards its basal side with little to no LamC1–sfGFP detectable between the primordium and the skin (Extended Data Fig. 1j). This separates the BM from the skin (Fig. 1b). Consistent with this, we found that the basal side of the skin labelled with fluorescently tagged E-cadherin abuts the apical side of the primordium (Fig. 1c). Thus, the primordium migrates on top of the BM and underneath the skin (Fig. 1d).

The primordium requires the BM for migration. If the skin and the BM serve as substrates for the primordium, they should also be required for the migration of the primordium. Indeed, surgical removal of the skin blocks primordium migration. To assess the role of the skin in primordium migration in a less invasive manner, we depleted α-catenin tagged with Citrine (Ctnna1–Citrine) in the primordium using the degron system zGrad expressed from the cxcr4b promoter in the migrating primordium (Extended Data Fig. 2a–d). Ctnna1 links cadherin to the actin cytoskeleton and its depletion should abrogate cadherin-mediated cell–cell adhesion between the skin and the primordium. As previously reported, depletion of Ctnna1–Citrine in the primordium affects sensory organ deposition and directionality of the cells in the primordium (Fig. 2a,b and Supplementary Video 1). However, we found that this does not affect the speed of the cells (Fig. 2c).

To assess the role of the BM in primordium migration, we analysed the migration of the primordium in lamC1 mutant embryos. In such embryos, the BM is disrupted or missing, and inspection of the collagen IV network and the BM confirmed this observation (Fig. 2d,e). Since the lack of LamC1 also impairs the formation of the Cxcl12a-secreting stripe of cells that guides the primordium (Extended Data Fig. 2e), we assessed the ability of the primordium to migrate in lamC1 mutant embryos by generating an ectopic Cxcl12a source in the trunk muscles. The initial location of the primordium is not affected in lamC1 mutant embryos (Extended Data Fig. 2f,g). For this analysis, we also removed endogenous Cxcl12a to avoid competition between endogenous and ectopic chemokine.

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The primordium cells form nascent adhesion-like clusters. To push, pull and exert stresses on the BM, the primordium needs to adhere to the BM. Molecularly, cells can adhere to the BM through focal adhesions. Two core components of these large protein complexes are integrins and talins\(^1\). Integrins bind to specific BM components on the outside the cell and—through talins and other adaptors—to the actin network inside the cell. To test whether the primordium uses focal adhesions to interact with the BM, we first identified the β-integrins and talins that the primordium expresses. Of the 12 β-integrins and 3 talins in zebrafish, integrin-β\(1b\) (itgb1b) and talin1 (tln1) were expressed throughout the primordium (Extended Data Fig. 3a,d). We therefore tagged itgb1b and tln1 with sfGFP and YPet at the endogenous locus and on a bacterial artificial chromosome (BAC) transgene, respectively (Extended Data Fig. 3b,e). Itgb1b–sfGFP and Tln1–YPet recapitulated the endogenous expression pattern (Extended Data Fig. 3c,f) and restored viability in the respective mutant background (viable itgb1b–sfGFP\(^+\) and tln1:tln1–YPet; tln1\(^–/–\) adults, \(n=20\) and \(5\), respectively). While Itgb1b–sfGFP and Tln1–YPet were enriched at the myotendinous junctions of the muscle, itgb1b–sfGFP localized uniformly on the membranes of the primordium cells (Fig. 3a and Supplementary Video 2), and Tln1–YPet was mostly localized in the cytoplasm of the cells of the primordium (Fig. 3b and Supplementary Video 2). Since Itgb1b–sfGFP and Tln1–YPet are also expressed by the surrounding skin and muscle, this expression could mask protein clustering on the membranes of the primordium cells. We therefore used blastomere transplantation to generate embryos in which only a few cells in the primordium express Itgb1b–sfGFP or Tln1–YPet together with membrane-tethered mCherry (mem–mCherry) (Fig. 3c). This analysis revealed that itgb1b–sfGFP and Tln1–YPet formed short-lived clusters with a lifetime of less than 2 min on the basal sides of the cells in the primordium, often within the basal protrusions (Extended Data Fig. 4b,c). We also detected short-lived Itgb1b–sfGFP clusters on the apical side of the superficial cells (Extended Data Fig. 4a)—a layer of thin primordium cells that face the skin\(^1\). These clusters are probably induced by fibronectin or proteoglycans—labelled by chondroitin sulfate—that are expressed around the primordium (Extended Data Fig. 1k,l). Since talin links integrin to F-actin\(^1\), we asked whether clustered Itgb1b and Tln1 colocalized with F-actin. Chimeric analysis showed that this is the case; Itgb1b–sfGFP and Tln1–YPet clusters colocalized with F-tractin-labelled F-actin on the basal sides of primordium cells (Fig. 3d–f and Supplementary Video 3). Control experiments showed that Itgb1b–sfGFP and, to a lesser degree, Tln1–YPet also colocalized with mem–mCherry, as expected for a transmembrane protein and a cytosolic protein, respectively (Extended Data Fig. 4d,e). Thus, the primordium cells form small, transient integrin–talin–F-actin clusters on their basal sides.

We corroborated the transient nature of the integrin clusters by measuring the mobility of integrin in the membrane through fluorescent recovery after photobleaching (FRAP). Ligated integrin couples to the actin network and diffuses more slowly in the membrane than unligated integrin\(^2\). The mobility of integrin is therefore a measure of the degree by which ligated integrins interact with actin. Consistent with integrin function in muscle, the mobility of Itgb1b–sfGFP at the myotendinous junction was increased when ROCK-mediated actin network contractions were blocked (Extended Data Fig. 4f–h). In comparison to the myotendinous junction, the mobility of Itgb1b–sfGFP was higher in the cells of the primordium (Extended Data Fig. 4l–k), which supports the idea that integrin interacts with the actin network only transiently in this migrating tissue. In contrast to migrating cells in vitro\(^3\), these observations suggest that the primordium cells do not form long-lived focal adhesions but rather transient integrin clusters. When placed on a laminin-coated surface, primordium cells formed large integrin and talin clusters along F-actin cables, as observed in cultured cells\(^4\) (Extended Data Fig. 4l–n), which indicates that primordium cells can form focal adhesions and stress fibres ex vivo but do not do so in vivo.

Integrin and talin are required for efficient migration. Next, we asked whether integrin and talin function are required for primordium migration. Since itgb1b and tln1—are and possibly itgb1a, tln2a and tln2b—are expressed in the primordium, we generated mutants in these five genes (Extended Data Figs. 5a and 6a). Phenotypic analysis showed that the primordium was less elongated and migrated more slowly in itgb1b mutant embryos than in wild-type controls (Fig. 4a–d, Extended Data Fig. 5b–f and Supplementary Video 4). Since itgb1a\(^–/–\); itgb1b\(^–/–\) and itgb1a\(^–/–\); itgb1b\(^–/–\) embryos exhibited severe morphogenesis defects (Extended Data Fig. 5b), we could not assess primordium migration in these genetic scenarios. Instead, we assessed whether Itgb1b is required within the tissue for migration by depleting Itgb1b–sfGFP in the primordium

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**Fig. 1** The primordium migrates on top of the BM and directly under the skin. a, TEM images of the skin (s), the primordium (p, purple hue), the muscle (m) and the BM (white arrows). \(n=1\) embryo. b, Optical sections through a primordium in a 31 h.p.f. embryo expressing LamC1–sfGFP. c, Optical section through a primordium labelled with F-tractin–mCherry (mCh) in a live 32 h.p.f. embryo expressing Cdh1–sfGFP. d, A, apical; B, basal. Schematic illustration of the environment around the primordium. Scale bars, 1\(\mu\)m (a), 25\(\mu\)m (c) or 50\(\mu\)m (b).
of itgb1a<sup>−/−</sup>; itgb1b-sfGFP<sup>−/−</sup> embryos using zGrad (Fig. 4e and Extended Data Figs. 2a and 5g,h). In such embryos, primordium migration was slowed to almost the same degree as in itgb1b mutant embryos (Fig. 4f,g), which indicates that Itgb1b is required within the primordium for efficient migration with a minor contribution from Itgb1a.

In talin single-, double- and triple-mutant embryos, the primordium migrated normally and morphogenesis was mostly unaffected (Extended Data Fig. 6b,c). A possible explanation for the mild defects in tln1<sup>−/−</sup>; tln2a<sup>−/−</sup>; tln2b<sup>−/−</sup> embryos could be the maternal contribution (M) of talin mRNA and talin protein to the embryo. To address this possibility, we generated zygotic (Z) Z tln1<sup>−/−</sup>; Z tln2a<sup>−/−</sup>; Z tln2b<sup>−/−</sup> embryos that also lacked the maternal contribution (M) of M tln1 and M tln2b (Extended Data Fig. 6d). However, such embryos had somitogenesis defects and disrupted cxcl12a expression along the migratory route of the primordium (Extended Data Fig. 6e-g), which impeded the analysis of the role of talin in primordium migration. Next, we depleted Tn1–YPet in the primordium by expressing zGrad from the cxcr4b promoter in MZ tln1<sup>−/−</sup>; Z tln2a<sup>−/−</sup> or Z tln2a<sup>−/−</sup>; MZ tln2b<sup>−/−</sup> embryos, whose only source of Tn1–YPet was maternally deposited mRNA and protein (Fig. 4h). zGrad efficiently degraded Tn1–YPet (Extended Data Fig. 6h,i). This analysis showed

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**Fig. 2 | Primordium migration requires an intact BM.**

- **a.** Control and Ctnna1-depleted primordia (arrowheads) in 48 h.p.f. embryos. Close-up of the region is indicated by a dashed square. **b.** Quantification of the migration distance for primordia shown in a. **c.** Speed of Ctnna1-depleted primordium cells. The solid line indicates the median, whereas the dashed line indicates the quartile. n = cell speeds from more than 7 primordia with each more than 100 cells. **d.** Left: TEM images of the ultrastructure of the BM between the skin and the muscle in control (n = 2) and lamC1<sup>−/−</sup> embryos (n = 1). White arrowheads indicate the BM. Right: antibody staining against collagen IV (Col IV) in control and lamC1<sup>−/−</sup> embryos. **e.** Quantification of collagen IV filaments in control and lamC1<sup>−/−</sup> embryos. **f.** Schematic of the strategy used to express Cxcl12a in a few muscle cells in lamC1<sup>−/−</sup> embryos and siblings. **g.** Images of the migrating primordium in cxcl12a<sup>−/−</sup> and cxcl12a<sup>−/−</sup> lamC1<sup>−/−</sup> 32 h.p.f. embryos with clones in the trunk muscle that express mCherry (not shown) or Cxcl12a together with mCherry (not shown). Asterisks indicate the ear and arrowheads the primordium. **h.** Quantification of the distance migrated by the primordium in the indicated experimental conditions at 32 h.p.f. **i.** Number of filaments of acta1a:mCherry or acta1a:cxcl12a clone in muscle. **j.** Images of the migrating primordium in lamC1<sup>−/−</sup> and lamC1<sup>−/−</sup> embryos with clones in the trunk muscle that express mCherry (not shown) or Cxcl12a together with mCherry (not shown). **k.** Quantification of the cxcr4b-Kate to memGFP ratio in the primordia of embryos shown in j. **l.** MemRFP/memGFP ratio. For a, b, c, d, g, h and k, n indicates the number of embryos. For b, e, h and k, individual data points, mean and s.d. are indicated. Scale bars, 2 μm (d, left), 20 μm (j), 50 μm (d, right) or 0.5 mm (a, g).

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**ARTICLES**

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motifs in the cytoplasmic tail of integrin 12. We therefore deleted slower (Extended Data Fig. 6k–m and Supplementary Video 5).

Data Fig. 6j). Compared with controls, such primordia also migrated t-test at the end point). P 0.0001 (two-tailed Mann–Whitney test). For itgb1b–/–, Amino-acid alignment of indicated integrin-

The mean and s.d. values are indicated. ****

m (μm, 500, 0.0001 (two-tailed Mann–Whitney test). For

m (n), mcs.d. values are indicated. ****

and l, b, <

0.0053 (two-tailed

Welch’s t-stack. The primordium is outlined by a dotted yellow line.

z tln1:tln1-YPet

b-stack. The primordium is outlined by a dotted yellow line.

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Fig. 7d–f). Similar to the global and tissue-specific loss of Itgb1b, mobile than Itgb1b–sfGFP in the membrane (Extended Data

Fig. 4l,m). This was also observed for the two NPxY motifs in Itgb1b and tagged Itgb1b\(\Delta\)NPXY with sfGFP at its endogenous locus (Fig. 4k and Extended Data Fig. 7a). Compared with Itgb1b–sfGFP, Itgb1b\(\Delta\)NPXY–sfGFP localized less to the apical and basal sides of the cells in the primordium and its levels were reduced by 16% (Fig. 4l,m). This was also observed for the myotendinous junctions (Extended Data Fig. 7b,c). Consistent with its more uniform distribution, Itgb1b\(\Delta\)NPXY–sfGFP was more mobile than Itgb1b–sfGFP in the membrane (Extended Data Fig. 7d–f). Similar to the global and tissue-specific loss of Itgb1b,
Itgb1bΔNPxY–sfGFP also failed to support efficient primordium migration (Fig. 4n,o). This indicates that the integrin–talin complex is important for the primordium to move along its migratory route at normal speed.

**Integrin-β, slows actin flow in the primordium.** In migrating cells, the integrin–talin complex can couple F-actin flow inside the cell to the BM outside the cell and transduce force. To test whether the cells in the primordium use such a clutch-like mechanism, we measured the speed of F-actin flow on the apical and basal sides of cells in the primordium of wild-type and itgb1b mutant embryos by labelling F-actin with F-tractin–mNeonGreen (F-tractin–mNG) in a few cells of the primordium (Fig. 5a). This analysis showed that F-actin was concentrated in the front of wild-type cells, in the basal cells of the primordium at the basal sides of the cells (Fig. 5b) and in the apically located superficial cells of the primordium on the apical side of the cells (Extended Data Fig. 7g). The F-actin flow was halted or slowed towards the rear of the cells, with a mean speed of 1.5μm.min⁻¹ in the basal cells of the primordium (Fig. 5b,c and Supplementary Video 6), whereas F-actin flowed faster in the superficial cells with a mean speed 7.4μm.min⁻¹ (Fig. 5d and Supplementary Video 6). Removal of Itgb1b function did not significantly affect the pattern or speed of actin flow in the superficial cells (Fig. 5d, Extended Data Fig. 7g and Supplementary Video 6). In contrast, F-actin flow increased to 6.6μm.min⁻¹ in itgb1b mutant basal cells of the primordium, and F-actin formed radial cables...
that have also been observed in talin-depleted cells in culture\(^1\) (Fig. 3b,c and Supplementary Video 6). In basal primordium cells, the actin polymerization rate—the sum of the actin flow rate and the rate of membrane protrusion—also increased from 2.8 \(\mu\)m min\(^{-1}\) in wild-type control cells to 7.6 \(\mu\)m min\(^{-1}\) in \(\text{itgb1b}^{-/-}\) mutant cells (Fig. 5c and Extended Data Fig. 7h,i). Thus, integrins couple force for primordium motility on the basal but not the apical sides of cells in the primordium, and loss of force coupling through integrin results in an increase in the actin polymerization rate on the basal sides of the primordium cells.

The primordium exerts the highest stresses in its rear. If the primordium uses integrin and talin to pull on the BM and push itself forward, the primordium should exert stresses (force per area) on the BM and deform the BM. Such traction stresses have been measured for migrating cells in culture by imaging the displacement of fluorescent beads embedded in elastic surfaces or matrices and the bending of flexible cantilevers—collectively referred to as traction force microscopy\(^2\)\textsuperscript{-5},\textsuperscript{26}. To extend traction force microscopy to living embryos, we created optical landmarks on the BM and assessed how these landmarks are displaced as the primordium moves across them (Fig. 6a). Using a laser, we locally bleached LamC1–sfGFP in an approximately cylindrical volume in the BM (Extended Data Fig. 8a,b and Supplementary Video 7). Minimal LamC1–sfGFP diffused back into the bleached cylinder (Extended Data Fig. 8c,d), and bleached cylinders remained clearly demarcated for 2 h after photobleaching (Supplementary Video 8), whereas untagged, extracellular mCherry rapidly filled bleached cylinders after bleaching (Extended Data Fig. 8c). This indicated that bleaching LamC1–sfGFP is a suitable approach to place local marks on the BM and monitor the deformation of the BM over time. We therefore bleached a hexagonal pattern of marks onto the LamC1–sfGFP-labelled BM in front of the migrating primordium and recorded the position of the marks as the primordium migrated across this pattern (Extended Data Fig. 8a and Supplementary Video 8). To reconstruct the stresses from the displacement of the marks on the BM by the migrating primordium, we developed the analysis pipeline Embryogram\(^3\) (Extended Data Fig. 8e,f; https://zenodo.org/record/5762146#.Ya5X0y-B1QJ), which is inspired by the Cellogram algorithm\(^4\). Embryogram identifies the bleached cylinders in the first frame of the time lapse. This assigns a point in space to each mark in the first frame. The points are connected using a variant of the iterative closest point algorithm to obtain a triangular mesh. This mesh is then deformed to follow the marks on the BM in the subsequent frames of the time lapse, which leads to a time sequence of triangular meshes that captures the deformation of the BM (Extended Data Fig. 8f). The time-varying mesh is used to compute displacements between the marks. To convert the displacements into stresses, we filled an axis-aligned box that contains the sample with a volumetric tetrahedral mesh. This mesh contains the surface mesh that approximates the BM in the first frame of the time lapse. The stresses are then computed by solving an elastic deformation of the volumetric mesh (for details, see Supplementary Note 1). For this conversion, we determined the stiffness, or Young's modulus, of the BM. We removed the skin above the BM and measured the BM stiffness by atomic force microscopy (AFM) (Extended Data Fig. 8g). In agreement with previous in vivo studies that assessed stiffness at the micron scale\(\textsuperscript{18,27}\), this analysis yielded a Young's modulus for the BM of 566 ± 355 Pa (mean ± s.d.), which was reduced to 321 ± 158 Pa (mean ± s.d.) after collagenase treatment (Fig. 6b,c and Extended Data Fig. 8h–k), a value that probably reflects the stiffness of the underlying muscle (Extended Data Fig. 8i,m). Importantly, spontaneous twitches of skin cells contract the underlying BM. This causes the BM to buckle and wrinkle akin to the distortions of the substrate observed around cultured cells\(\textsuperscript{14}\) and in animals\(\textsuperscript{14}\). The wrinkles formed and disappeared in less than 2 min, and tracking the optical marks indicated that the wrinkles do not cause lasting deformation of the BM (Extended Data Fig. 9a–e and Supplementary Video 9). Similarly, repeated probing of the BM at the same location in deskinned embryos by AFM did not alter the stiffness measurements (Extended Data Fig. 8i). These observations suggest that the BM undergoes non-plastic deformations and can be approximated by a linear stress–strain relationship.

Using the Embryogram pipeline together with the stiffness measurements, we found that the front cells of the primordium pull slightly on the BM in random directions. In contrast, the cells in the middle and rear cells of the primordium pull on the BM more strongly and more directional, displacing the BM side-
ways, backwards and downwards (Fig. 6d–f and Supplementary Video 10). The traction stresses (the forces against the plane of the BM) reflected this displacement pattern. In the front, the mean traction stresses averaged 28 ± 22 Pa, and increased to 58 ± 49 Pa and 64 ± 51 Pa (mean ± s.d.) in the middle and rear of the primordium, respectively, with higher traction stresses exerted preferentially along the sides of the primordium and peaking at 600 Pa (Fig. 6g–j and Supplementary Video 10). Similarly, the primordium generated high, mostly rearward-pointing stresses (stresses extracted in the direction of migration) along the sides and towards its rear (Extended Data Fig. 9h), where it also exerted the highest shear stresses on the BM (Extended Data Fig. 9i). Although the primordium moves at relatively constant speed, it does not exert constant stresses on its substrate (Supplementary Video 10), which suggests that its forward motion is the result of the average of the fluctuating stresses across the tissue. At the position where the primordium has passed, the BM returned to its original shape and the marks on the BM snapped back to their original position, which indicates that the BM is not irreversibly deformed by the primordium (Extended Data Fig. 9f-g and Supplementary Video 10). Also, we did not observe such BM displacements and stresses in controls in which we blocked primordium migration through...
the ubiquitous overexpression of Cxcl12a, which is the attractive guidance cue of the primordium (Fig. 6), Extended Data Fig. 9–o and Supplementary Video 10). The observed stress distribution was also reflected in the wrinkling of the BM along the sides of the primordium, with LamC1–sfGFP forming local clusters (Fig. 7a). These LamC1–sfGFP clusters were juxtaposed to F-actin clusters in the primordium (Fig. 7b) and were specifically enriched around the primordium (Fig. 7c), which suggests that actin network contraction in the primordium locally pull on the BM and cause it to wrinkle. Together, these observations indicate that the front cells exert low traction stresses while the rear cells exert high traction stresses. On a tissue level, the primordium moves in a continuous breaststroke-like manner, with its front pushing the BM in random directions while its middle and rear strongly push the BM sideways and backwards. This stress pattern is consistent with theoretical predictions for adherent cell migration.

**Actomyosin activity is highest in the rear of the primordium.** The primordium generates higher stresses in the rear than in the front. This could be because there are more cells in the rear or because the cells in the rear pull stronger on the BM than in the front. Stronger pulling in the rear could be reflected in increased actomyosin activity, slower actin flow rates, increased integrin levels, more engaged integrin, more stable integrin clusters and a greater stress dependence on integrins in the rear than in the front of the primordium.

To test these possibilities, we first assessed the activity of actomyosin across the primordium. Since actomyosin activity correlates with the localization of the actin motor non-muscle myosin II into dots, we generated a myosin II reporter line for the primordium (MyI12.1–mScarlet). Consistent with previous studies that used other contexts17, MyI12.1–mScarlet localized to dots in the cells of the primordium. These dots did not form a cable-like structure around the back of the primordium as in other contexts17 but were mostly localized to the basal sides of the cells and to the lateral sides in the middle to rear of the primordium (Fig. 8a), and enriched in its rear (Fig. 8b). This suggests that the cells in the rear and the sides of the primordium exert more forces than the cells in the front. Actin flow across the primordium was the same (Fig. 8c,d). Itgb1b levels across the primordium were similar (Fig. 8e), and integrin engagement—as judged by the increased clustering of Itgb1b–sfGFP compared with Itgb1bNPxY–sfGFP (Extended Data Fig. 10a,b). This is consistent with the idea that increased clustering of Itgb1b–sfGFP compared with Itgb1bNPxY–sfGFP reflects integrin engagement. In contrast, the levels of phosphorylated paxillin (p-paxillin)—a putative marker for integrin adhesion turnover18—but not total paxillin

**Fig. 7** | The BM wrinkles around the primordium. **a,** F-tractin–mCherry distribution in the primordium (top) and LamC1–sfGFP around the primordium (bottom) in a 32 h.p.f. embryo. LamC1–GFP fluorescence intensity is pseudo-coloured as a heatmap and shown in arbitrary units. The image is a maximum-projected z-stack. **b,** Left, top: a transverse section through the F-tractin–mCherry-expressing primordium and the underlying LamC1–sfGFP-labelled BM. Left, bottom: a corresponding image showing the LamC1–sfGFP fluorescence intensity as a heatmap in arbitrary units. Arrows indicate apposed clusters of F-tractin–mCherry and LamC1–sfGFP. Images are single sections along the yz plane of a z-stack. Right: fluorescence intensity profiles of F-tractin–mCherry and LamC1–sfGFP of image shown in left along the y axis. Arrows indicate the position of the apposed clusters of F-tractin–mCherry and LamC1–sfGFP indicated by arrows in the left panels. **c,** Quantification of the LamC1–sfGFP intensity within 3-μm-wide bands around the perimeter of the primordium (left) and at a distance of 6 μm from the perimeter of the primordium (right). ****P < 0.0001 (two-tailed paired t-test). n indicates the number of embryos. Scale bars, 10 μm (b) or 50 μm (a).

**Fig. 8** | The primordium generates larger forces in the rear. **a,** Top: MyI12.1–mScarlet distribution in the primordium. Middle: MyI12.1–mScarlet in the entire primordium. Bottom: MyI12.1–mScarlet on the basal side of the primordium pseudo-coloured. Dotted line indicates the outline of the primordium. **b,** Quantification of the basal MyI12.1–mScarlet intensity at indicated positions in the primordium. Data points, the mean and s.d. values are indicated (one-way ANOVA followed by Tukey’s multiple comparison test). **c,** Top: images of F-tractin–mNG at the basal sides of primordium cells. Arrows indicate the direction of migration. Bottom: Kymographs along the dotted line indicated in the top images. **d,** Actin flow rates in basal primordium cells. n indicates the number of cells pooled from more than five primordia. Data points, the mean and s.d. values are indicated (two-tailed Mann–Whitney test). **e,** Distribution (top) and quantification (bottom) of Itgb1b–sfGFP on the cell membranes of the primordium. Itgb1b–sfGFP intensity is pseudo-coloured. Mean and s.d. are indicated. **f,** Left: fluorescence intensity ratio images of Itgb1b–tdTomato to Itgb1b–sfGFP (top) and Itgb1bNPxY–sfGFP (bottom) in the membrane of primordium cells. The dotted lines indicate the outline of the primordium. Right: quantification of red/green (R/G) fluorescence intensity ratios. Ratios are shown in pseudo-colour. **g,** BM displacement around the primordium (grey) in itgb1bΔ embryos shown as a vector field along the x, y and z axes (Supplementary Video 10). The vector magnitude is indicated as a colour map. The x,y view is from the basal side of the primordium. **h,** Traction stresses on the BM indicated as a colour map (Supplementary Video 10). **i,** Pooled traction stresses exerted by the primordium on the BM in itgb1bΔ embryos. Median (thick line) and quartile (thin line) are shown. The wild-type control is the same as shown in Fig. 6j (two-tailed Mann–Whitney test). N indicates the number of embryos, n indicates the number of bleached cylinders. w was used for statistics. **j,** Left: staining against paxillin (top) and p-paxillin (bottom) protein in primordia shown as a heatmap. Right: quantification of the fluorescence intensity on the primordium membrane for paxillin (top) and p-paxillin (bottom). Mean and s.d. are indicated. **k,** Model for primordium motility. For b, e, f and j, n indicates the number of embryos. Scale bars, 2 μm (c), 20 μm (a, e, f, j) or 25 μm (g, h). Coloured bars in a, e, f indicate fluorescence intensities in arbitrary units.
were lower in the rear than the front of the primordium (Fig. 8i), which suggests that the rear of the primordium forms longer-lived integrin-mediated adhesions. Consistent with this idea, the traction stresses in the plane of migration (xy plane)—but not the traction stresses against the BM (z axis)—decreased more in the rear than in the front of primordia in embryos lacking Itgb1b compared with wild-type embryos but did not become equal in magnitude (Fig. 8g–i and Supplementary Video 10). Thus, Itgb1b couples more force to the BM in the rear of the primordium than in its front and is dispensable for force transmission against BM, but does not account for all the difference in stresses between the front and the rear of the primordium. Together, these observations suggest that the cells in the rear of the primordium generate more force than the cell in its front, which—probably together with the higher cell number in the rear—accounts for the high stresses exerted by the rear of the primordium on the BM.
Another migrating tissue that is guided by Cxcl12 and propelled forward by its rear is the cranial neural crest in *Xenopus*39. The front cells of the cranial neural crest are attracted by Cxcl12 and the rear cells form a supraacellular cable around the back of the tissue. This cable contracts periodically and pushes rear cells forward while front cells become displaced to the side and then to the back (Extended Data Fig. 10e). In contrast, the primordium is attracted by a self-generated Cxcl12a gradient that extends almost across the entire tissue46,47 (Extended Data Fig. 10c,d). It does not form a supraacellular actomyosin cable across adherens junctions around its back (Fig. 8a, and Extended Data Fig. 10h,i), does not periodically contract its rear (Extended Data Fig. 10f), and its cells do not intercalate but remain next to their neighbours during migration11 (Extended Data Fig. 10g and Supplementary Video 1). Thus, although both tissues generate highest forces in their rear, the underlying guidance and propulsion mechanisms are different and might represent two solutions to the same problem (Extended Data Fig. 10e): how to propel a tissue through an animal.

**Discussion**

Together, this work elucidates how a tissue moves through a live animal. It provides three major insights. First, primordium cells link the force-generating actomyosin network to the BM through integrin clusters on their basal sides. These integrin clusters are less than 2 μm in size and form and disassemble in less than 2 min. This is in contrast to the larger and longer-lived focal adhesions that migrating cells in culture use to pull themselves forward46,47 and is more reminiscent of nascent adhesions that form at the edge of protrusions and underneath spreading cells in culture46–48. This suggests that tissues in animals rely on transient rather than prolonged cell–substrate interactions for movement. Second, the primordium cells pull and deform the BM on their outside with maximal stresses around 600 Pa. This is comparable to the average stress that migrating cells in culture exert on their substrates26–28, which suggests that stresses in this range are inherent to adherent migration in simplified and physiological scenarios. Consistent with this notion, retrograde actin flow is slowed or stalled in primordium cells, which suggests that most of the flow is converted into forward movement. Intriguingly, disrupting the coupling of actin flow across integrin results in increased flow. Such a compensatory response has also been observed in cultured dendritic cells and macrophages49. These cells also increase actin polymerization in response to decreased force coupling to the substrate to maintain forward movement—probably by coupling through other integrins and nonspecific adhesion. Third, the primordium moves similar to a continuous breaststroke by pushing the BM downward, sideways and backwards. Counterintuitively, the rear cells of the primordium generate higher traction stresses than the front cells, probably because the rear cells need to overcome greater resistance in their environment than the front cells. One possible reason for the greater resistance is the front cells, which the rear cells might need to push to move forward. Consistent with this idea, the front cells pull and push the BM in random directions (Fig. 6i), which suggests that they contribute little to directed force generation in the tissue. Also, when severed from the rear cells, the front cells cease to move forward46. It therefore seems that the rear pushes the front to propel the primordium forward, a scenario that is akin to a rear-engine-like design (Fig. 8k). Intriguingly, the two collectively migrating heart cells in *Ciona* and crawling *Dictyostelium* slugs, banana slugs and garden snails also push stronger with their rears than their fronts on their surroundings47–49 which indicates that this propulsion design is conserved across different length scales to drive tissue and animal movement.

**Online content**

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at [https://doi.org/10.1038/s41556-022-00844-9](https://doi.org/10.1038/s41556-022-00844-9).

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Methods

Zebrafish husbandry. This study was performed in accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. All of the animals were handled according to approved institutional animal care and use committee protocols (I16-00788_AMM2012000320) of the NYU Grossman School of Medicine.

Zebrafish strains. Embryos were staged as previously described. The cxxla1a:exon2 allele has been previously described. The lamC2a allele was obtained from the Zebrafish International Resource Center (https://zfin.org) and contains a nonsense mutation that results in a premature stop codon. Homozygous lamC1a:exon4a mutants were identified by their shortened body axis or by PCR-based genotyping. Primers used for genotyping of digested amplicons on a 3% agarose gel with the above genotyping protocol.

AM1344) using the linearized plasmid MEGAscript T7 Transcription kit (Thermo Fisher Scientific, AM1334) to obtain Purification kit (Qiagen, 28106) and subjected to in vitro transcription using a gRNA backbone primer were annealed, filled-in by Taq polymerase and amplified sequence, the target sequence without the PAM site and an overhang for the Ensembl genome browser (GRCz10). gRNA sequences were identified Otherwise, we designed gRNAs targeting the available coding sequence in (ref.41) and (ref. 35), (ref. 11), (ref. 57), (ref. 53), (refs. 55,56), (ref. 9), (ref. 54).

Generation and genotyping of the tln1d4 mutant. The gRNAs used to target tlb1a are listed in Table 1. The isolated mutant contains a 34-bp deletion of tlb1a exon 13 that results in a frame shift that causes a premature stop codon. tlb1a mutants can be kept as homozygous adult fish. The tlb1a allele was genotyped by PCR. The primers used for genotyping by PCR are listed in Supplementary Table 1.

Generation and genotyping of the itgb1bi mutant. The gRNAs used to target itgb1a are listed in Supplementary Table 1. The isolated mutant contains a 70-bp insertion that results in a frameshift that introduces two successive premature stop codons. The predicted mutant protein comprises only the first 320 amino acids of the total 806 amino acids. itgb1b homzygous mutant embryos display a shorter body axis (Extended Data Fig. 3b–d) and die as embryos, which is consistent with previous reports. The itgb1b allele was genotyped by PCR. The primers used for genotyping by PCR are listed in Supplementary Table 1.

Generation of transgenic lines. To generate BAC-mediated transgenes, we modified the BAC clone of interest with gRNA-mediated BAC recombinasing amino acids, as previously described. In brief, we first modified the BAC clone to add sequences complementary to the target region with a linker sequence to include a transgene marker and the tol2 sequences in the BAC backbone. Next, we modified the protein-coding sequence of interest on the BAC to add a fluorochrome protein or to express the coding sequence under the regulation of a specific promoter. The final BAC was characterized by EcoRI digestion and PCR followed by sequencing around the modified sequences. The final BAC was purified with a Nucleobond BAC 100 kit (Takara Bio, 740579) and co-injected with 1 nl of 40 ng/g DNA tol2 mRNA into 1-cell-stage embryos. The stable transgenic line was established by out-crossing the adult fish injected with the BAC transgene and raising transgenic embryos.

TgBAC(lamC1:lamC1-gGFP). For the lamC1:lamC1-gGFP transgene, we used the BAC clone CHORI-211-19441, which was obtained from BACPAC Resources, Children’s Hospital Oakland Research Institute, CA, USA. This BAC spans 192,491 bp of genomic DNA and contains the lamC1 locus with about 50 kb of the genomic sequence upstream of lamC1 exon 1 and about 35 kb of the genomic sequence downstream of lamC1 exon 1. As a transgenesis marker is cyrrus/Cerulean. This transgene expresses full-length laminin-γ1 fused to sfGFP from the lamC1 promoter. The full name of this transgenic line is TgBAC(lamC1:lamC1-gGFP).1 Note that this transgenic recapitulates the recently reported expression pattern of lamC1 mRNA expression,1 partially resuces lamC1 homozygous mutant embryos (Extended Data Fig. 1h) and does not affect primordium migration (Extended Data Fig. 1f).

TgBAC(lamC1:lamC1-gGFP). For the lamC1:lamC1-gGFP transgene, we used the BAC clone DKEY-42J10, which was obtained from imaGenes. The BAC clone spans 194,108 bp of genomic DNA. This includes 40 kb of the genomic sequence upstream of the beginning of lamC1 exon 1, lamC1 exon 1 to 44, but lacks lamC1 exon 45 to 56. To include the complete coding sequence of lamC1 on the BAC, we inserted the sequence of the missing lamC1 exons (exons 45 to 56) directly downstream of exon 44. This design was guided by the annotated lamC1 transcript lamC1-202 (ENSDBT00000167799.2, Ensembl). We inserted the coding sequence for Ypet between the head and the rod domains of lamC1 (ref. 18), which is located in lamC1 exon 13. As a transgenesis marker, we inserted cryrrus/Red in the BAC backbone. A cassette comprising lamC1 exons 45–56 and the kanamycin-resistant gene KanR was inserted. The amino acid sequence around the Ypet insertion is Gly-Ser-Val-X-Ala-Leu-Pro, which is consistent with previous reports.1 The lamC1 allele was genotyped by PCR. The primers used for genotyping by PCR are listed in Supplementary Table 1.

Generation and genotyping of the tln1d4 mutant. The gRNAs used to target tln1a are listed in Supplementary Table 1. The isolated mutant fish harbour a 4-bp deletion in tln1 exon 2 that results in a frame shift that introduces a premature stop codon. tln1a mutant fish can be kept as homozygous adult fish. The tln1d4 mutant allele was established by out-crossing the adult fish injected with the BAC transgene and raising transgenic embryos.

Generation and genotyping of the tln1d4 mutant. The gRNAs used to target tln1a are listed in Supplementary Table 1. The isolated mutant contains a 70-bp insertion that results in a frame shift that introduces two successive premature stop codons. The predicted mutant protein comprises only the first 320 amino acids of the total 806 amino acids. itgb1b homzygous mutant embryos display a shorter body axis (Extended Data Fig. 3b–d) and die as embryos, which is consistent with previous reports. The itgb1b allele was genotyped by PCR. The primers used for genotyping by PCR are listed in Supplementary Table 1.

Generation and genotyping of the tln1d4 mutant. The gRNAs used to target tln1a are listed in Supplementary Table 1. The isolated mutant contains a 70-bp insertion that results in a frame shift that introduces two successive premature stop codons. The predicted mutant protein comprises only the first 320 amino acids of the total 806 amino acids. itgb1b homzygous mutant embryos display a shorter body axis (Extended Data Fig. 3b–d) and die as embryos, which is consistent with previous reports. The itgb1b allele was genotyped by PCR. The primers used for genotyping by PCR are listed in Supplementary Table 1.

Generation and genotyping of the tln1d4 mutant. The gRNAs used to target tln1a are listed in Supplementary Table 1. The isolated mutant contains a 70-bp insertion that results in a frame shift that introduces two successive premature stop codons. The predicted mutant protein comprises only the first 320 amino acids of the total 806 amino acids. itgb1b homzygous mutant embryos display a shorter body axis (Extended Data Fig. 3b–d) and die as embryos, which is consistent with previous reports. The itgb1b allele was genotyped by PCR. The primers used for genotyping by PCR are listed in Supplementary Table 1.

Generation and genotyping of the tln1d4 mutant. The gRNAs used to target tln1a are listed in Supplementary Table 1. The isolated mutant contains a 70-bp insertion that results in a frame shift that introduces two successive premature stop codons. The predicted mutant protein comprises only the first 320 amino acids of the total 806 amino acids. itgb1b homzygous mutant embryos display a shorter body axis (Extended Data Fig. 3b–d) and die as embryos, which is consistent with previous reports. The itgb1b allele was genotyped by PCR. The primers used for genotyping by PCR are listed in Supplementary Table 1.
For constructing cxcr4b:EGFP-CAAX plasmids, we used the BAC clone DKEY-169F10 described above. We identified one correct knock-in event by sequencing PCR amplicons spanning the genomic insertion site.

Generation of plasmid constructs. Plasmids were generated by Gibson cloning. To construct pDEST-to2-ta2a1-cxcl12a-t2a-mCherry and pDEST-to2-ta2a1-cxcr4a-F-tractin-mNeonGreen plasmids, the following plasmids were used as PCR templates: the plasmid backbone including pDEST-tol2-hsp70l-zGrad-t2a-mNeonGreen (ref. 9) and the pCS2-cxcl12a (ref. 9), respectively. To construct pCS2(+)-YPEt-ZF1, pCS2(+)-mNeonGreen-ZF1 and pCS2(+)-mCherry-ZF1, the following plasmids were used as templates: the plasmid backbone, the YPetr, the mNeonGreen and the mCherry sequences were amplified from pCS2(+)-tGFp-ZF1 (ref. 7) or pCS2(+)-tGFp-ZF1, pUC19-t1n-5arm-YPet-t3arm, the Addgene plasmid 98886 (ref. 10) and pDEST-to2-hop70l-scElc-mCherry-SV40pA (ref. 11). To construct pDEST-to2-hop70l-scElc-mCherry-SV40pA (ref. 11), the transgenetics marker cxcr4b:EGFP-CaaX was amplified from zebrafish cDNA by PCR. This transgene expresses the first five amino acids from cxcr4bΔN20 with PCR primers that also included the SP6 promoter sequence. mRNAs were synthesized on plasmids using primers containing a SP6 promoter sequence. mRNAs were co-injected into zebrafish embryos. The cxcr4b:EGFP-CaaX promoter drives high expression only in the posterior region except that the BAC was modified to contain the cxcr4b:F-tractin-mNeonGreen expression driven from the cxcr4b promoter.

Antibody staining and quantification of paxillin and p-paxillin levels. To stain for F-actin, phospho-MLC, fibronectin, chondroitin sulfate, paxillin, and p-paxillin (p-Y118) and the membrane of the primordium cells, we fixed 32h.p.i. TgBAC(cxcr4b-EGFP-CaaX) embryos in 4% PFA/0.1% Triton X-100 in PBS for 20 min at room temperature before blocking. The embryos were permeabilized in 1% BSA/PBS and blocked in 1% BSA/PBS at 1:1,000 in 0.5% Triton X-100. Embryos were immunostained overnight with the following secondary antibodies at a 1:1,000 dilution at room temperature for 1 h: donkey anti-actin (Jackson ImmunoResearch, 711-546-152, lot no. 109010). The posterior lateral line cells were identified based on F-tractin–mCherry expression and co-stained with a semi-automated manner with a custom-written macro in Fiji. In brief, the primordium region was manually cropped. The macro generated a binary mask based on the EGFP–CaaX signal (autothresholding with the Iso-Data method). The mask was applied to the paxillin or p-paxillin channel, and the signal on the primordium membrane was extracted. Next, three consecutive optical slices were acquired with a sequential scan setting. Quantification of filamin and p-paxillin intensities were performed in a semi-automated manner with a custom-written macro in Fiji. In brief, the region of interest of 10-μm wide and 10-μm high was manually defined from the tip of the primordium, and the signal intensity profile was obtained along the front-to-rear axis of the primordium. We did not include the membrane at the leading edge of the primordium because the signal in this region could not be separated from the signal of the skin. Finally, the signal was normalized to the average intensity for each embryo and plotted.

Quantification of filamentous collagen IV by immunofluorescence staining. To stain for collagen IV and the membrane of the primordium cells in lamC1 mutant and control embryos, we in-crossed cldBlyn:GFP, lamC1−/− fish and sorted for cldBlyn:GFP, lamC1−/− and cldBlyn:GFP, lamC1+/− or cldBlyn:GFP, lamC1−/− embryos. The lamC1−/− larvae were incubated in anti-collagen IV (1:200, ab5686, Abcam) and goat anti-GFP (1:500, Covance, custom-made antibody) at 4°C. Embryos were washed four times with PBS and incubated with donkey anti-rabbit-Cy3 (1:500, Jackson ImmunoResearch, 711-165-152, lot no. 102215) or goat anti-mouse Cy3 (1:500, Jackson ImmunoResearch, 115-165-003) together with donkey anti-goat Alexa488 (1:500, Jackson Immunolrodes, 711-546-152, lot no. 1108949), or rabbit anti-paxillin(pY118) (1:500, Novus, NBP1-24459, lot no. 031327 42D0P2) together with goat anti-GFP (1:500, Covance, custom-made antibody) overnight at 4°C. Embryos were washed twice with PBS and incubated with goat anti-rabbit Cy3 (1:500, Jackson ImmunoResearch, 711-165-152, lot no. 102215) or goat anti-mouse Cy3 (1:500, Jackson ImmunoResearch, 115-165-003) together with donkey anti-actin Alexa488 (1:500, Jackson Immunolrodes, 711-546-152, lot no. 1108949), or rabbit anti-paxillin(pY118) (1:500, Novus, NBP1-24459, lot no. 031327 42D0P2). The embryos were mounted in PBS. Images were taken on a Leica SP8 confocal microscope equipped with Hyd detectors (Leica Microsystems) using a x40 (NA 1.1) objective with a sequential scan setting.

Quantification of filamin and p-paxillin intensities were performed in a semi-automated manner with a custom-written macro in Fiji. In brief, the region of interest of 10-μm wide and 10-μm high was manually defined from the tip of the primordium, and the signal intensity profile was obtained along the front-to-rear axis of the primordium. We did not include the membrane at the leading edge of the primordium because the signal in this region could not be separated from the signal of the skin. Finally, the signal was normalized to the average intensity for each embryo and plotted.
overnight at 4°C. Embryos were washed four times with PRST and incubated with donkey anti-rabbit Cy3 (1:500, Jackson ImmunoResearch, 711-165-152, lot no. 102215) and donkey anti-goat Alexa488 (1:500, Jackson ImmunoResearch, 709-546-148) for 1 h. The samples were then incubated in 0.5% low-melt agarose/Ringer’s solution. Images were taken on a spinning disk confocal Nikon W1 microscope. The number of filament structures along the primordium was quantified using a custom-written macro in Fiji. In brief, a 150 × 150 μm ROI that contained the entire primordium was manually defined. The red fluorescent channel of the z-stack representing the collagen IV signal was sum-projected, and only the fluorescence values above 1.25 the mean fluorescence intensity of the image were kept. Filamentous structures were extracted using the Tubeless filter (https://www.longair.net/edinburgh/imagej/tubeless/) in Fiji with sigma set to 0.7. Then, the image was thresholded using “Otsu” method and the number of filaments was counted with the Analyze Particles command in Fiji (settings: limiting size = 50–infinity, circularity = 0.0–0.3).

Whole-mount in situ hybridization. The procedures for RNA probe synthesis and zebrafish embryo whole-mount in situ hybridization were performed as previously described47. The RNA probe against cxcl12a has been previously described48 and no new synthetic procedures were performed. The collagenase (collagenase, MS-222 anaesthetic). The skin of the embryos was peeled off under a Leica inverted microscope (settings: limiting size 50–infinity, circularity 0.0–0.3).

Dissection of embryos and collagenase treatment. The head and the yolk of 28 h.p.f. live zebrafish embryos were removed in dissection medium supplemented with 0.4 mg ml⁻¹ MS-222 anaesthetic. Dissected embryos were transferred into dissection medium (Ca²⁺-free Ringer’s solution, 50 mM EDTA and 0.4 mg ml⁻¹ MS-222 anaesthetic). The skin of the embryos was peeled off under a Leica dissecting scope (Leica, Wild M420 with light stand) using forceps. The deskinned embryos were then transferred into Leibovitz L-15 medium (Fisher, 11415064) supplemented with 0.4 mg ml⁻¹ MS-222 anaesthetic. The collagenase (collagenase, purified, 4 kU, Worthington, LS005275) stock was prepared in Leibovitz L-15 medium at a concentration of 1,000 U ml⁻¹. The deskinned embryos were soaked in the collagenase solution (900 U ml⁻¹) supplemented with 0.4 mg ml⁻¹ MS-222 anaesthetic at room temperature for 30 min. After the treatment, the collagenase was washed out using Leibovitz L-15 medium supplemented with 0.4 mg ml⁻¹ MS-222 anaesthetic, the embryos were quickly mounted for AFM or confocal microscopy.

AFM measurements and data analysis. Deskinned embryos were glued to FluoroDish dishes (World Precision Instruments, FD5040-100) using CELL-TAK (Corning, 354240) and imersed in Leibovitz L-15 medium supplemented with 0.4 mg ml⁻¹ of the anaesthetic MS-222 (Sigma Aldrich, A5040-25g). All AFM measurements were carried out within 90 min after skin removal. The AFM measurements were performed on an Asylum Research MFP-3D-BIO atomic force microscope using the Asylum Research software package vIX (AR software) as previously described49. The AR software was used for cantilever calibration, force mapping, data export and data visualization (Extended Data Fig. 8–n).

We used a spherical borosilicate glass bead probe with a 2.5 μm radius, a spring constant of 0.07 N m⁻¹, a Young’s modulus of 68.0 GPa and a Poisson ratio of 0.19 (Novascan, PT-G5). In contrast to pyramidal probes, which probe structures at the point defined by the user to only fit a select part of the curve to the Hertz model (contact point to end of force curve/point of maximum deformation or the end point defined by the user). These statistical parameters were used as quality control criteria to include or reject force curves. For the calculation of Young’s modulus, the sample Poisson ratio was assumed to be 0.45. The reduced Young’s modulus $E'$ was obtained by fitting the first 200 nm of the approach curve past the contact point to the Hertz model as follows:

$$F = \frac{1}{2}E' r^2 \delta^2,$$

where $F$ is the loading force, $E'$ is the reduced Young's modulus, $r$ is the radius of the spherical probe used and $\delta$ is the indentation. The sample Young's modulus $E$ was calculated using

$$\frac{1}{E'} = \frac{1}{E} - \frac{1}{E} \frac{\delta^2}{r^2} + \frac{1}{E} \frac{\delta^2}{r^2} E.$$

where $\nu_i$ is the indenter’s Poisson ratio, $\nu_i$ is the sample Poisson ratio and $E$ is the indenter’s Young's modulus.

Owing to debris after skin removal and the curvature of the BM above the muscle, some force curves were of low quality. To automatically select high-quality force curves, we applied three criteria. The first criterion was the slope of the approach curve (between the approach part of the force curve to contact point) to a linear equation. The second criterion was the residuals between the measured and fitted curve of the baseline of the approach curve to a linear equation. The third criterion was the $P$ value of the fit of the approach curve to the Hertz model from the contact point to 200 nm or 500 nm into the sample. Only force curves with a baseline slope between −10 and $+10\,\text{pN}/\text{μm}$, baseline sum-of-squared-residuals smaller than 0.1 $\text{nN}$² and $P$ values smaller than 1.0 × 10⁻¹⁰ were included to determine the overall stiffness of the BM. These criteria select for force curves that have a flat baseline approach curve and a clearly defined contact point. Extended Data Fig. 8n shows representative force curves that meet and do not meet these criteria.

Electron microscopy. lamC1⁻⁻ embryos were generated by in-crossing lamC1⁻⁻ fish. Embryos (30 h.p.f.) were fixed in electron microscopy fixative containing 2% PFA, 2% glutaraldehyde in 0.1 M sodium cacodylate buffer at room temperature for 2 h and then overnight at 4°C. Fixed embryos were rinsed with 0.1 M sodium cacodylate buffer and post-fixed with 1% OsO₄ in 0.1 M cacodylate buffer, followed by block-staining with 1% uranyl acetate aqueous solution overnight at 4°C. The samples were rinsed with water, dehydrated in a graded series of ethanol, infiltrated with propylene oxide/Epon mixtures and finally embedded in EMbed812 (Electron Microscopy Sciences). Sections (70 nm) were cut and mounted on 200 mesh copper grids and carbon coated. Ultrathin section imaging was performed on a Talos120C transmission electron microscope (Thermo Fisher Scientific) with a Gatan (4k × 4k) OneView Camera (Gatan). The primordium cells were pseude-coloured using Adobe Illustrator 2020 (Adobe).

Ectopic expression of Cxcl12a from the trunk muscle cells. A total of 1 nl of 10 ng ml⁻¹ PDEST-tol2-acta1a-cxcl12a-t2a-mCherry plasmid DNA or PDEST-tol2-acta1a-cxcl12a-t2a-mCherry plasmid DNA was injected together with 40 ng ml⁻¹ tol2 mRNA into 1-cell-stage embryos obtained from the following crosses: lamC1⁻⁻; cxcl12a⁻⁻; cdna3B-gfp, gfp in-cross, lamC1⁻⁻; cxcl12a⁻⁻; cdna3B-gfp, gfp in-cross, wild-type in-cross and lamC1⁻⁻; cxcl12a⁻⁻; cxcr5c-vat3x-Kate2-iRES-GFP-CaaX in-cross. lamC1⁻⁻ mutant embryos were identified by morphology.

Live imaging with a Leica SP8 confocal system and image analysis. The live imaging experiments detailed below were performed on a Leica SP8 confocal microscope system equipped with a x40 (NA 1.1), a x20 (NA 0.7) and a x20 (NA 0.5) objectives. Samples were kept at 28°C with a heated stage (Warner Instruments, Quick Exchange Heated Base, QE-1) for prolonged time-lapse imaging. The power of the laser lines was calibrated using a power meter (X-Cite Power Meter Model, Lumen Dynamics, XR2200). For quantifying the fluorescence intensity, image acquisition was carried out using the photon-counting mode. Embryos were mounted in 0.5% low-melt agarose (National Diagnostics, EC-205)/ Ringer’s solution supplemented with 0.4 mg ml⁻¹ MS-222 anaesthetic on a coverslip or a plastic dish. 

Itgb1b mutant and control embryos were generated by crossing itgb1b-itgb1b-sGFp; prpemc-mcherry fish to itgb1b-itgb1b-sGFp; fish. Such embryos were imaged for 10 h with the following settings: z-step size of 5.0 μm, time interval of 30 min and duration of 9.5 h. Mutants were identified by

$$\text{itgb1b}.$$
the absence of Itgb1b-sfGFP expression. Note that itgb1b mutants produced by this cross showed a slightly stronger overall morphological defect than mutants generated by in-crossing itgb1b+/− fish, which indicated that maternal itgb1b rescues the loss of zygotic itgb1b slightly better than maternal itgb1b-sfGFP mRNA and protein.

To image the localization of itgb1b-sfGFP in the primordium cells, 34 h.p.f. itgb1b:itgb1b-sfGFP; primem-m-Cherry and itgb1b:itgb1b-sfGFP; primem-m Cherry control embryos were mounted. The z-step size was 0.42 μm. The power of the 488-nm and 594-nm laser lines was calibrated to 95 μW and 29 μW, respectively. The GFP-to-m-Cherry fluorescence intensity ratio in the basal-to-apical axis was obtained by a custom-written macro in Fiji. In brief, the macro generated a cell-membrane mask based on the m-Cherry channel using the “Default” thresholding method in Fiji. This mask was applied to the GFP and the m-Cherry channels, and the signal intensities were obtained for each z-slice. To analyse the ratio of itgb1b-sfGFP in the primordium cells using custom-written macros in Fiji, in brief, we first manually aligned the primordium along the x, y and z axes such that the three central slices in the z-stack (= 1.2 μm) neither contained the apical constrictions nor the lateral nerve but contained the tip and the rear of the primordium. Next, the macro generated a binary mask for the cell membrane using mem-m-Cherry fluorescence and the “Default” thresholding method in Fiji. The mask was applied to the GFP channel and the three central slices were average-projected. Next, the masked GFP channel was resliced from the tip of the primordium to the rear, and the intensity profile along this line was obtained.

To image the localization of F-tractin–mCherry in the primordium with skin cells labelled with Cdh1-sfGFP, 33 h.p.f. TgBAC(cxcr4b:H2A–mCherry) embryos were mounted. The z-step size was 0.42 μm. The power of the 488 nm and 594 nm laser lines were calibrated to 68 μW and 14 μW, respectively. Two channels were sequentially scanned. To analyse the ratio of Cxcl12a–EGFP in the primordium cells using custom-written macros in Fiji, in brief, we first aligned the primordium along the x, y and z axes. The macro includes only the bottom half of the z-stack such that the stack did not contain the apical constrictions. Then, the stack was maximum-projected. The primordium area was masked by a binary mask generated as follows: the EGFP–CaaX channel was maximum-projected, thresholded using the “Huang” method in Fiji and the holes were filled. Then, 1.7 μm was eroded to eliminate the Myl12.1–mScarlet signal localized at the periphery of the primordium. The value (mean ± s.d.) of Myl12.1–mScarlet was subtracted from the Myl12.1–mScarlet intensity to highlight only accumulated myosin II punctae. The total intensity of Myl12.1–mScarlet punctae in the front, middle and rear regions of the primordium was manually measured. The total intensity of Myl12.1–mScarlet punctae was normalized by the area of the front, middle and rear of the primordium. Finally, these values were divided by the mean for normalization and plotted.

To image the localization of Ctnna1–citrine in the primordium cells, TgBAC(cxcr4b:myl12.1–mScarlet); TgBAC(cxcr4b:EGFP–CaaX) embryos were mounted at 32 h.p.f. The z-step size was 0.42 μm with a pinhole of 1.0 Airy units. The power of the 488-nm and 546-nm laser lines were calibrated to 105 μW and 112 μW, respectively. Two channels were sequentially scanned. Basally localized Myl12.1–mScarlet was then subtracted from the EGFP and GFP channels. In brief, we first aligned the primordium with the x, y and z axes. The macro includes only the bottom half of the z-stack such that the stack did not contain the apical constrictions. Then, the stack was maximum-projected. The primordium area was masked by a binary mask generated as follows: the EGFP–CaaX channel was maximum-projected, thresholded using the “Huang” method in Fiji and the holes were filled. Then, 1.7 μm was eroded to eliminate the Myl12.1–mScarlet signal localized at the periphery of the primordium. The value (mean ± s.d.) of Myl12.1–mScarlet was subtracted from the Myl12.1–mScarlet intensity to highlight only accumulated myosin II punctae. The total intensity of Myl12.1–mScarlet punctae in the front, middle and rear regions of the primordium was manually measured. The total intensity of Myl12.1–mScarlet punctae was normalized by the area of the front, middle and rear of the primordium. Finally, these values were divided by the mean for normalization and plotted.

To observe the localization of TgBAC(cxcr4b:myl12.1–mScarlet); TgBAC(cxcr4b:EGFP–CaaX) embryos were mounted at 32 h.p.f. The z-step size was 0.42 μm with a pinhole of 1.0 Airy units. The power of the 488-nm and 546-nm laser lines were calibrated to 105 μW and 112 μW, respectively. Two channels were sequentially scanned. Basally localized Myl12.1–mScarlet was then subtracted from the EGFP and GFP channels. In brief, we first aligned the primordium with the x, y and z axes. The macro includes only the bottom half of the z-stack such that the stack did not contain the apical constrictions. Then, the stack was maximum-projected. The primordium area was masked by a binary mask generated as follows: the EGFP–CaaX channel was maximum-projected, thresholded using the “Huang” method in Fiji and the holes were filled. Then, 1.7 μm was eroded to eliminate the Myl12.1–mScarlet signal localized at the periphery of the primordium. The value (mean ± s.d.) of Myl12.1–mScarlet was subtracted from the Myl12.1–mScarlet intensity to highlight only accumulated myosin II punctae. The total intensity of Myl12.1–mScarlet punctae in the front, middle and rear regions of the primordium was manually measured. The total intensity of Myl12.1–mScarlet punctae was normalized by the area of the front, middle and rear of the primordium. Finally, these values were divided by the mean for normalization and plotted.

To observe receptor internalization induced by ectopic Cxcl12a expression, the "Spot" tool in the Imaris software was used with a specified cell diameter of 3.78 μm and the default background subtraction was activated. The histogram of the spot quality was adjusted such that about 70–80% of the cells were selected. The cell nuclei were tracked using the "Autoregressive motion" method with a maximum distance that an object can move between two consecutive time points set to 20 μm for the red channel (H2A–mCherry). The maximum gap size—the maximum number of consecutive time points that are allowed to be missing to join track fragments—was set to two. To correct for stage-drift, a muscle cell nucleus was tracked and chosen to track fragments that do not belong to the primordium or represent dying cells and to correct or delete track segments. For dividing cells, only one daughter cell was tracked. To correct for stage-drift, a muscle cell nucleus was tracked and chosen as a reference point. The x, y and z values and the track IDs were exported and the position of the reference cells was subtracted from all tracked cells at each time point. With the exported data, the speed of each cell in the indicated genotype was pooled.

To observe receptor internalization induced by ectopic Cxcl12a secreted from the muscle cells, 25–26 h.p.f. embryos were imaged. The power of the 488-nm and 591-nm laser lines was calibrated to 152 μW and 110 μW, respectively. The z-step size was 0.42 μm. We manually cropped the z-stack such that it only contained the primordium we then generated a binary mask based on the GFP channel and applied it to the RFP and GFP channels. The RFP and GFP channels were then summed-projected, the mean intensity of each channel was measured and the RFP mean intensity was divided by the GFP mean intensity using a custom-written macro in Fiji.
The tagged protein expression patterns of TgBAC(lamC1:lamC1-sfGFP), TgBAC(thb:thb1n1-Ypet), thbg1b-thbg1b-sfGFP and thbg1b-thbg1b-mCherry were imaged in 28-h.p.f. embryos. The z-step size was set to 1.5μm. The images were stitched together using the auto-tiling feature in the LAS X Life Science microscopy software (Leica) and sum-projected.

Live imaging with a Nikon W1 spinning disk confocal system and image analysis. The live imaging experiments detailed below were performed using a Nikon W1 spinning disk confocal microscope that was equipped with an Apo LWD ×40 NA 1.15 objective lens (Nikon, MRD77410) and a SR HP Plan Apo ×100 NA 1.45 objective lens (Nikon, MRD01905). The x100 objective was used only for the actin flow experiments. All other experiments were performed using the x40 objective. The samples for live time-lapse imaging experiments were kept at 30°C using a Tokai Hit incubation system STXG-TIZWX-SET.

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The live imaging experiments detailed below were performed using a Leica TCS SP8 confocal laser scanning microscope. The embryos were mounted in 0.5% low-melt agarose (National Diagnostics, 555553-10MG) in 1% dimethylsulfoxide (DMSO) or control-treated with 1% DMSO for 3 h.

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The embryos were mounted in 0.5% low-melt agarose (National Diagnostics, 555553-10MG) on a glass-bottom dish. The embryos were digested and genotyped for itgb1b–/–; TgBAC(tln1:tln1-YPet) or itgb1b–/–; TgBAC(lamC1:lamC1-sfGFP; prim:mem-mCherry) TgBAC(cxcr4b;F-tratin-mCherry) embryos. The green and the red channels were sequentially imaged to prevent fluorescent bleaching.

To quantify the LamC1–sfGFP intensity around the edge of the primordium and beyond the primordium, we used a semi-automated custom-written macro in Fiji. Z-stacks were maximum-projected. The outer circumference of the primordium was manually traced using the primordium-specific mCherry signal as a guide. The encircled area was filled and thresholded. The thresholded image was duplicated. The first duplicate was masked and the outer circumference of the primordium as a binary mask. We plotted the mean intensity of the LamC1–sfGFP signal in each region from individual embryos.

Analysis of actin flow. To image actin flow in single primordium superficial and basal cells, we co-injected BAC DNA coding for cxcr4b;F-tratin-mNG with 1 nl of 40ng μl–1 tol2 mRNA into 1-cell-stage wild-type and thbg1b–/– embryos that were transgenic for prim:mem-mCherry. The thbg1b–/– embryos were generated by crossing thbg1bthbg1b-sfGFP; prim:mem-mCherry to thbg1bthbg1b-sfGFP and sorting for embryos lacking thbg1b–sfGFP expression at 24 h.p.f. The embryos were mounted at 32–34 h.p.f. Clones in the primordium were identified based on mCherry and mNG expression and imaged on their basal side, collecting single planes every 2 s for 3 min.

Actin flow analysis was performed by generating kymographs using Fiji. In brief, a 1 pixel–wide 20 μm ROI line was drawn manually from the centre of the cell outwards across the protrusion on the basal side. The kymograph was generated using the KymoResliceWide plugin provided by E. Katrukha and L. Young (https://imagej.net/KymoResliceWide). In singly labelled cells, the front of the actin flow and the front of the protrusion were visually identified, manually traced and the actin flow rate and protrusion rate were extracted from the slopes of the trace lines. The actin polymerization rate was calculated by subtracting the actin flow rate from the protrusion rate. To image and analyse actin flow in single primordium basal cells across the front-to-rear axis of the primordium, we prepared injected embryos as described above. To identify the location of F–tratin–mNG expressing cells, we first imaged a z-stack of the whole primordium. Then switched to the x-y plane. To image the edge of the primordium, the actin flow was later identified based on the entire primordium z-stack image. To image F-actin retrograde flow in the tip cells, we used TgBAC(cxcr4b;F-tratin-mCherry); TgBAC(cxcr4b;EGFP-CaaX) embryos. The imaging conditions and analysis were identical.

Blasteomere transplantation and imaging of primordia with clones expressing thbg1b–sfGFP and Tln1–Ypet. We transplanted 20–50 cells from donor embryos at the 1,000–8,000 cell stage into host embryos of the same stage. All host embryos were wild type. Donor embryos were transgenic for TgBAC(thb:thb1n1-Ypet) or thbg1bthbg1b-sfGFP and Tg(primmem-mCherry) or TgBAC(cxcr4b; F-tratin-mCherry). At 28 h.p.f., we isolated embryos that contained donor cells in the primordium based on the expression of the Tg(primmem-mCherry) or TgBAC(cxcr4b;F-tratin-mCherry) transgenes. To observe clones in the superficial cells of the primordium, we used thbg1bthbg1b-sfGFP; TgBAC(cxcr4b;F-tratin-mCherry) as donor embryos. Chimeric embryos were mounted at 32–34 h.p.f. Images were collected every 30 s for 10 min as z-stacks with a z-step size of 1.0μm. The green and mNG fluorescent channels were sequentially scanned to prevent fluorescent bleed-through.

Blasteomeric transplantation and imaging of primordia with clones in talin activity. We transplanted 20–50 cells from donor embryos at the 1,000–8,000 cell stage into embryos of the same stage. Donor and recipient embryos were transgenic for thbg1bthbg1b-sfGFP; thbg1bthbg1b-mCherry and TgCd34cd34;F-tratin-mCherry. Donor embryos were transgenic for Tg(primmem-mCherry). Talin-depleted donor embryos were obtained from an in-cross of thbg1bthbg1b-sfGFP; thbg1bthbg1b-mCherry; TgCd34cd34;F-tratin-mCherry; Tg(primmem-mCherry) fish. Wild-type and talin-depleted donor embryos were injected with 50ng μl–1 zGrad mRNA at the 1-cell stage. Embryos (32–34 h.p.f.) were kept at 30°C using a Tokai Hit incubation system STXG-TIZWX-SET.

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The embryos were mounted at 30°C using a Tokai Hit incubation system STXG-TIZWX-SET.

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The embryos were mounted at 30°C using a Tokai Hit incubation system STXG-TIZWX-SET.
with chimeric primordia were mounted. Images were collected every 5 min for 2 h as z-stacks with a z-step size of 1.0 μm. The green and red fluorescent channels were sequentially scanned to prevent fluorescent bleed-through. Cumulative migration distance was quantified by manually tracking the tip of the primordium using the “Manual tracking” plugin in Fiji. Kymographs were drawn using the “KymoResliceWide” plugin in Fiji.

**Analysis of rear contraction and migration.** To analyse rear contractions in the primordium, TgBAC(lamC1:lamC1-sfGFP) embryos were mounted at 33 h.p.f. Images were collected every 30 s for 30 min as z-stacks with a z-step size of 1.0 μm. As a proxy for the rear cells, we tracked the apical constriction site located in the rear of the primordium (>65 μm from the tip) using the sum-projected EGFP-CaaX signal and the “TrackMate” plugin in Fiji. The contraction of the rear cells was analysed as follows. First, we duplicated the middle nine slices (=8.0 μm of the z-stack) and summed them next. Then, we subtracted the background from the projected image using 5 pixels in Fiji to enhance the membrane signal. Third, we applied the “Huang” thresholding method to this image and binarized it. Finally, we manually identified two vertices, which were located three cells apart from each other, located at the periphery of the rear region of the primordium, and manually tracked their locations over time.

**Live imaging with a Leica 165M FC fluorescence stereo microscope.** The experiments detailed below were performed using a Leica 165M FC fluorescence stereo microscope equipped with a Leica DFC345 FX camera.

To assess the migration distance of the primordium in embryos in different genetic scenarios (cldnB:lysGFP; lamC1–/– and cldnB:lysGFP embryos with clones in the muscle expressing mCherry from the acta1α promoter, TgBAC(lamC1:lamC1-sfGFP), 26 h.p.f. (initial stage), 32 h.p.f. (assessment of migration) and 48 h.p.f. embryos, respectively, were imaged. The TgBAC(lamC1:lamC1-sfGFP) copy number was determined based on the intensity of the sfGFP fluorescence. Migration measurements were performed using Fiji.

To measure the migration distance of the primordium in itgb1b–/– fish, 54 h.p.f. embryos were generated by crossing itgb1b-itgb1b-sfGFP; prim:mem-mCherry fish with itgb1b-itgb1b-sfGFP; fish. Itgb1b–/– embryos were identified by the absence of sfGFP expression. The primordium migration distance was measured using Fiji. Note that the primordium reaches the tip of the tail by 42 h.p.f. in wild-type embryos as well.

To analyse the distance that the primordium migrated in various mutant backgrounds at 48 h.p.f., the following crosses were set up to obtain embryos of the required genotypes. At 48 h.p.f., the following crosses were set up to obtain embryos of the required genotypes.

- To quantify the circularity of the primordium migration distance, time-lapse image sequences were performed using an Axiocam (Zeiss) and a ×5 (NA 0.25) objective.

**Documentation of the phenotype of lamC1–/–; TgBAC(lamC1:lamC1-sfGFP) embryos.** The TgBAC(lamC1:lamC1-sfGFP) was identified based on sfGFP fluorescence. The lamC1 mutant embryos were identified and scored based on the morphological defects at 48 h.p.f.

**Primordium migration distance quantification.** To quantify the cumulative migration distance of the primordium, time-lapse image sequences were maximum- or sum-projected, and the tip of the primordium was tracked using the “Manual tracking” plugin provided by Fabrice Cordelieres in Fiji (https://imagej.nih.gov/ij/plugins/manual-tracking.html).

**Circularity analysis of the primordium.** To quantify the circularity of the primordium, we limited the length of the primordium to the first 100 μm from the tip of the primordium. Z-stacks were sum-projected. Using Fiji, the primordium region was manually cropped based on the intensity of Lyn–/– sfGFP fluorescence. Then, a median filter with a radius of 5 pixels was applied and the background was subtracted using a rolling ball radius of 100 pixels. Images were rendered binary using the “Huang” thresholding algorithm to obtain a clear outline of the primordium. Finally, we quantified the circularity of the primordium for each time point using the “Analyze particles” macro in Fiji. The circularity C is defined as

\[ C = \frac{4 \pi A}{P^2} \]

where A is the area of the primordium and P is the perimeter.

**Analysis of itgb1b–/–sfGFP and Tln1–YPet localization in primordial cells.** We analysed the spatial distribution of itgb1b–/–sfGFP and Tln1–YPet with respect to F-tractin–mCherry and mem-mCherry at the basal side of small clones of labelled primordial cells in two ways. First, we chose a single slice at the basal side of the cells in a clone. The clone contour was manually selected based on the F-tractin–mCherry or mem-mCherry fluorescence using a 5 pixel-wide (=1.625 μm) segmented line to obtain the fluorescence intensity profile at each point for the green and red channels in Fiji. Second, we assessed the colocalization of the fluorescently tagged proteins using the “coloc2” plugin in Fiji. Before applying the coloc2 plugin, we processed the images as follows: due to photobleaching, only the first ten time points were included for analysis. Images were rotated such that the direction of migration aligned with the x-axis. Then the images were sliced to obtain transverse sections in the x plane. This resulted in images with the basal sides of the cells pointing upwards (z axis of imaging) and the direction of migration pointing to the right. Next, a median filter with 1 pixel width was applied and the images were rotated again such that the basal sides of the cells were horizontally aligned. An ROI with a width of 3.25 μm from the basal membrane inward was manually defined as the basal region of cells and used for the colocal2 analysis in Fiji with a custom-written macro. The macro computes the individual ROIs of the basal region from each time point, creates a mask based on the sum intensities of the green and red channels using the “Default” method in Fiji. The individual ROIs were also compiled into a single image for each channel as a montage as shown in (Extended Data Fig. 4d). The macro then calculates the degree of colocalization, the FICQ value, using the coloco2 function in Fiji. Li’s ICQ is calculated as follows. For each pixel in the ROI, the product of the difference of intensity and the mean intensity for each channel is calculated ((Ch1 – mean(Ch1)) × (Ch2 – mean(Ch2))). Then, the number of pixels with a positive product are normalized to the total pixel number and 0.5 is subtracted. Therefore, Li’s ICQ ranges from –0.5 (signals perfectly segregated) to 0.5 (signals perfectly overlapped). We performed the calculation for three consecutive x/z slices per clone and pooled the data for display in the panel.

**Analysis of skin-generated BM wrinkles and traction.** A 30×30 μm ROI centred on a transient increase in LamC1–/–sfGFP fluorescence intensity at the third time point using the “Analyze particles” macro in Fiji. The circularity C is defined as

\[ C = \frac{4 \pi A}{P^2} \]
point in a four-dimensional stack spanning five time points with a 1 min interval was manually defined. The third point was set to 0 min. To visualize the LamC1–sfGFP increase as a graph, the stack was maximum-projected along the z-axis. The fluorescence intensity profile across the LamC1–sfGFP increase at the 0 min time point. The same fluorescence intensity profiles along the same line ROI were obtained for the 1 min and +1 min time points. The fluorescence intensity was normalized to the mean of the −1 min time point. Traction was analysed using Embryogram and calculated and visualized with ParaView (ParaView-5.8.1, Kitware). For the traction stress calculation, a Young’s modulus of 566.7 Pa was used for the BM based on the AFM measurements (Fig. 6c) and a Poisson ratio of 0.45. Traction was obtained using the "PIVlab" plugin (v.2.38 by W. Thielicke) in Matlab (v.9.9, MathWorks).

Analysis of primordium-generated traction stresses and angles of BM deformation.

Four-dimensional confocal 2-stacks were denoted by applying a median filter (width: 3 pixels) and analysed in Embryogram. In Embryogram, an area containing a well-defined bleach pattern that was 1 μm line ROI across the z-axis was manually defined. The third point was set to 0 min. The spatial displacement of the membrane from traction stresses measured at the three marks closest to the LamC1–sfGFP increase. Particle image velocimetry (PIV) analysis of the basal skin cells was used for detecting the displacement of the membrane from −1 min to 0 min using the "PIVlab" plugin (v.2.38 by W. Thielicke) in Matlab (v.9.9, MathWorks). Every second cell was visualized on the cell membrane image at the −1 min time point. The displacement of the BM was obtained by Embryogram. The three-dimensional quiver plots of the displacement of the BM from −1 min to 0 min with the scale factor 3 were drawn using ParaView (ParaView-5.8.1, Kitware).

The magnitude of traction stress in the direction of primordium migration in three-dimensions was manually obtained in Fiji. Half-circle polar diagrams were drawn using Matlab. The cosines of these angles are shown in Fig. 6i. The average SD of the traction stress values was obtained using the R environment. In brief, the cosines between the directions of two given cells at a given time frame was obtained. We performed this analysis for all the combinations of two cells for all the time frames and pooled data. The trajectories of the primordium cells were also drawn using R.

Data availability

Previously published genome assemblies as either GRCz10 or GRCz11 that were used here for the design of gRNA constructs are available for tln1, tln2a, tln2b, itgb1a and itgb1b under accession codes ENSDARG00000100729, ENSDARG0000117901, ENSDARG000011973, ENSDARG000017863 and ENSDARG000014484, respectively. All other data supporting the findings of this study are available from the corresponding authors on reasonable request. Source data are provided with this paper.

Code availability

The code for Embryogram software has been deposited at Zenodo (https://zenodo.org/record/5762146#.Y63X0y-B1QIY). The codes for image analysis using ImageJ and R are provided as a zip file.

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Author contributions

N.Y., D.P. and H.K. conceptualized the study and designed the experiments. N.Y. performed all the zebrafish experiments with support from H.K., except for the AFM measurements, which were performed by B.W. with samples prepared by N.Y. The Embryogram software was developed by Z.Z., T.S. and D.P. (with inputs from N.Y. and H.K.). N.Y. analysed most of the data with help from Z.Z. and T.S. for the traction stress analysis and from Z.Z. and B.W. for the AFM data analysis. N.Y. and H.K. wrote the main manuscript (with input from Z.Z., T.S. and D.P.). Z.Z., T.S. and D.P. wrote Supplementary Note 1 (with input from N.Y. and H.K.). All authors approved of and contributed to the final version of the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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Extended Data Fig. 1 | Ultrastructure of the basement membrane along the migratory route of the primordium and the characterization of the TgBAC(lamC1:lamC1-sfGFP) line. a, Overview of the primordium. Dotted lines indicate the location of the cross-sections shown in (b–e). b–e, TEM images of cross-sections at the level of the most recently deposited neuromast (b), at the level of the primordium’s rear (c), at the level of the primordium’s front (d), and in front of the primordium (e). Scale bars = 10 μm. b’–e’, Magnification of area outlined by a dotted line in (b–e). Scale bars = 1 μm. The skin (s), primordium (p, purple hue), the muscle (m), and the BM (arrows) are indicated. n = 1 embryo. f, Schematic of the TgBAC(lamC1:lamC1-sfGFP) transgene. g, Image of the expression of LamC1-sfGFP from TgBAC(lamC1:lamC1-sfGFP) transgene in a 28 hpf embryo. The image is a sum-projected z-stack. Scale bar = 0.5 mm. h, The TgBAC(lamC1:lamC1-sfGFP) transgene partly rescues the lamC1 mutant phenotype. Crosses from lamC1-/-; lamC1:lamC1-sfGFP to lamC1-/- fish resulted in embryos with three different phenotypes shown on the left. Quantification of the phenotypic categories from these crosses for non-transgenic embryos and embryos expressing LamC1-sfGFP are shown on the right. Note that the mild phenotype correlates with the presence of LamC1-sfGFP and the severe phenotype represents the lamC1 mutant phenotype. Scale bars = 0.5 mm. i, Quantification of the primordium migration in the presence of different copy numbers of the TgBAC(lamC1:lamC1-sfGFP) transgene. Data points, means, and SD are indicated. n.s.: p = 0.6514 (non-transgenic vs. lamC1:lamC1-sfGFP/+), p = 0.7842 (non-transgenic vs. lamC1:lamC1-sfGFP/lamC1:lamC1-sfGFP) (two-tailed Mann-Whitney test). j, Cross-section along apical-basal axis of a primordium (dotted line in top panel) of embryos expressing LamC1-sfGFP (BM) and the Cdh1-tdTomato (skin). The LamC1-sfGFP signal is enhanced to saturated levels (bottom panel). Scale bars = 25 μm. k, Images of slices from a z-stack of 32 hpf TgBAC(cxcr4b:EGFP-CaaX) embryos stained for Fibronectin and GFP. Orthogonal views are shown. l, Images of slices from a z-stack of 32 hpf TgBAC(cxcr4b:EGFP-CaaX) embryos stained for Chondroitin sulfate and GFP. Orthogonal views are shown. For h, i, n = number of embryos.
Extended Data Fig. 2 | Depletion of Ctnna1-Citrine by zGrad and characterization of the lamC1 mutants. a, Principle of zGrad-mediated protein degradation. b, Left: 8 hpf embryos injected with sfGFp-ZF1 mRNA and mCherry-ZF1 mRNA with or without co-injected zGrad mRNA. Middle: 8 hpf embryos injected with YPet-ZF1 mRNA and mCherry-ZF1 mRNA with or without co-injected zGrad mRNA. Right: 8 hpf embryos injected with mNeonGreen-ZF1 mRNA and mCherry-ZF1 mRNA with or without co-injected zGrad mRNA. n ≥ 20 embryos. Scale bar: 1 mm. c, Single confocal slices of primordia in prim:mem-mCherry; ctnna1:ctnna1-citrine control (left) and prim:mem-mCherry; ctnna1:ctnna1-citrine; cxc4b:zGrad 32 hpf embryos (right). Lower panels show the Ctnna1-Citrine fluorescence as a heat map. Scale bar = 20 μm. d, Quantification of the Ctnna1-Citrine fluorescence intensity in control and zGrad-expressing embryos at 32 hpf. Data points, means, and SD are indicated. ****: p < 0.0001 (two-tailed Welch’s t-test). e, Expression of cxcl12a in control (wild-type or lamC1-/-) and lamC1 mutant 30 hpf embryos. Bracket indicates the location of interrupted cxcl12a expression domain. Scale bar = 0.5 mm. f, mCherry-expressing clones in muscle of 26 hpf control (wild-type or lamC1-/-) and lamC1 mutant embryos also transgenic for cldnB:lyn2GFP. Arrowheads indicate the position of primordium. Scale bar = 0.5 mm. g, Quantification of the distance from the ear to the first somite in the indicated genotypes at 26 hpf. Data points, means, and SD are indicated. n.s.: p = 0.5516 (two-tailed Mann-Whitney test). h, Images of the primordium in wild-type and cxcl12a-/-; 32 hpf embryos with clones in the trunk muscle that express Cxcl12a together with mCherry (not shown) (Left). Asterisks indicate the ear and arrowheads the primordium. Scale bar = 0.5 mm. i, Cross-sectional images of the Cxcl12a sensor in primordia of cxcl12a-/- and cxcl12a-/-; lamC1-/- embryos with clones in the muscle of the trunk that express mCherry or Cxcl12a. Quantification shown in Fig. 2k. Scale bar = 20 μm. For d, e, g, h, n = number of embryos.
Extended Data Fig. 3 | β-integrin and talin expression analysis. a, Expression analysis of β-integrins in the migrating primordium by in situ hybridization on 32 hpf embryos. Note that itgb8 could not be amplified from embryonic cDNA. Arrows indicate expression in the primordium. Scale bar = 0.5 mm. b, Schematics of the itgb1b locus, the itgb1b targeting cassette, and the modified itgb1b locus. c, Itgb1b-sfGFP expression in a 28 hpf itgb1b:itgb1b-sfGFP embryo. The image is a sum-projected z-stack. Scale bar = 0.5 mm. d, in situ hybridization against the three zebrafish talin genes on 32 hpf embryos. Arrow indicates enriched talin expression in the primordium. Scale bar = 0.5 mm. e, Schematic of the TgBAC(tln1:tln1-YPet) transgene and its protein product. f, Tln1-YPet expression in a 28 hpf tln1:tln1-YPet embryo. The image is a sum-projected z-stack. Scale bar = 0.5 mm.
Extended Data Fig. 4 | See next page for caption.
Extended Data Fig. 4 | Integrin-β1b and Talin1 dynamics in cells of the primordium. a, Localization of Itgb1b-sfGFP and F-tractin-mCherry at the apical side of superficial cells in the primordium. The images are single optical slices. Arrowheads indicate Itgb1b-sfGFP clustering. Scale bar = 10 μm. b, Localization of Itgb1b-sfGFP (top) and Tln1-YPet (bottom) with membrane-mCherry at the basal sides of cells in clones in the primordium imaged over time from Supplementary Video 3. The images are single optical slices. Arrowheads indicate Itgb1b-sfGFP and Tln1-YPet clustering. Scale bar = 10 μm. c, Intensity profiles of Itgb1b-sfGFP (left) and Tln1-YPet (right) together with membrane-tethered mCherry along the contours of clones at indicated times taken from Supplementary Video 3. Arrows indicate Itgb1b-sfGFP and Tln1-YPet clusters that do not coincide with membrane-tethered mCherry clustering. Representative profile of 5 or more imaged cells. d, Montage of 10 consecutive images of the basal sides of the clones. The images are single transverse sections from a time lapse video. Scale bar = 10 μm. e, Quantification of co-localization of Itgb1b-sfGFP and Tln1-YPet with F-tractin-mCherry and membrane tethered mCherry. Li’s ICQ co-localization indices of 0.5 and −0.5 indicate perfectly co-localized and perfectly anti-co-localized signals, respectively. n = number of cells. Data points, means, and SD are indicated. Three data points were analyzed from the same embryo. **: p = 0.0015 (two-tailed t-test). f, Images from time-lapse video after photo-bleaching of Itgb1b-sfGFP at the myotendinous junction of embryos treated with DMSO or 50 μM Rockout. GFP intensities are pseudo-colored as a heat map. Scale bars = 10 μm. g, Graph of Itgb1b-sfGFP fluorescence intensity over time before and after photo-bleaching in embryos treated with DMSO or 50 μM Rockout. The fluorescence intensities are normalized to the minimal intensities after photo-bleaching. Dots indicate mean intensities and error bars are SD. n = number of experiments, N = number of embryos. h, Plot of the percent recovery of Itgb1b-sfGFP fluorescence intensity at 28 sec after photo-bleaching in embryos treated with DMSO or 50 μM Rockout. Data points, means, and SD are indicated. ****: p < 0.0001 (two-tailed Welch’s t-test). **: p = 0.0015 (two-tailed t-test). i, Images from time-lapse video after photo-bleaching of Itgb1b-sfGFP in the primordium (left) and at the myotendinous junction (right). Fluorescence intensities are pseudo-colored as a heat map. Scale bars = 10 μm. j, Graph of Itgb1b-sfGFP fluorescence intensity over time before and after photo-bleaching in the primordium and at the myotendinous junction. The fluorescence intensities are normalized to the minimal intensities after photo-bleaching. Dots indicate mean intensities and error bars are SD. n = number of experiments, N = number of embryos. k, Plot of the percent recovery of Itgb1b-sfGFP fluorescence intensity at 27 sec after photo-bleaching in the primordium and at the myotendinous junction. n = number of experiments (used for statistical test), N = number of embryos. Data points, means, and SD are indicated. ****: p < 0.0001 (two-tailed Welch’s t-test). l, Experimental design to culture primordium cells. m, Antibody staining against Itgb1b-GFP and F-tractin-mCherry on cultured primordium cells. Arrowheads indicate actin stress fibers (m’) with Itgb1b-GFP clusters (arrows in m”) in the cell center and in protrusions (m’’’). Scale bars = 20 μm (m-m”) and 1 μm (m’’’). n, Antibody staining against Tln1-YPet and F-tractin-mCherry on cultured primordium cells. Arrowheads indicate actin stress fibers (n’) with Tln1-YPet clusters (arrows in n”) in the cell center and in protrusions (n”’). Scale bars = 20 μm (n-n”) and 1 μm (n”’). Images are max-projected z-stacks. Close-ups (right panels) are magnifications of the regions indicated by dotted squares in the middle panels.
Extended Data Fig. 5 | β1-integrin mutational analysis. **a**, Schematic of the itgb1a and itgb1b alleles. d and i denote deletion and insertion, respectively. Arrows indicate the position of the primordium. Scale bar = 0.5 mm. **b**, Quantification of the primordium migration distance (left), the body length (middle), and primordium migration distance normalized to body length (right) in 48 hpf itgb1 mutant embryos. n = number of embryos. Data points, means, and SD are indicated. ****: p < 0.0001 (two-tailed Welch’s t-test). **: p = 0.0026 (two-tailed Welch’s t-test, middle plot). d, Overall morphology of control (wild-type or itgb1b-/+ ) and itgb1b mutant embryos at 24 hpf. Scale bar = 0.5 mm. e, Primordium migration in control (wild-type or itgb1b-/+ ) and itgb1b mutant embryos at 54 hpf. Scale bar = 0.5 mm. The arrows indicate the position of the primordium and the arrowheads indicate the position of the tip of the tail. f, Quantification of primordium migration in control (wild-type or itgb1b-/+ ) and itgb1b mutant embryos at 54 hpf. Data points, means, and SD are indicated. n = number of embryos. ****: p < 0.0001 (two-tailed Welch’s t-test). g, quantification of the Itgb1b-sfGFP fluorescence intensity in control and zGrad-expressing embryos at 33 hpf. Data points, means, and SD are indicated. ****: p < 0.0001 (two-tailed Welch’s t-test). n = number of embryos.
Extended Data Fig. 6 | See next page for caption.
Extended Data Fig. 6 | Generation and characterization of talin mutant and analysis of primordium migration in embryos or primordia with depleted Talin activity. a, Schematic of the tln1, tln2a and tln2b mutant alleles. The sequence around the deletions (d) and insertions (i) are shown. The start codons are indicated for tln1d4 and tln2a22; and the premature stop codon for tln2a22. b, Primordium migration distance in 48 hpf embryos with different levels of talin activity. Scale bar = 0.5 mm. c, Quantification of the primordium migration distance (left), the body length (middle), and primordium migration distance normalized to body length (right) in tln mutants at 48 hpf. Data points, means, and SD are indicated. ****: p<0.0001 (one-way ANOVA with Tukey's multiple comparisons test, left plot). ***: p=0.0002 (two-tailed Welch's t-test, middle plot). n = number of embryos. d, Crosses to generate embryos with depleted Talin activity. e, in situ hybridization against cxcl12a mRNA on 28 hpf wild-type (top) and Talin-depleted (bottom) embryos injected with zGrad mRNA. Scale bar = 0.5 mm. f, Quantification of the percentage of control and Talin-depleted embryos with perturbed cxcl12a expression along the horizontal myoseptum in 28 hpf embryos. n = number of embryos. g, Quantification of the body length in control and Talin-depleted 28 hpf embryos. n = number of embryos. Data points, means, and SD are indicated. ***: p=0.0063. ****: p<0.0001, n.s.: p=0.1610 (two-tailed Mann-Whitney test). h, prim:mem-mCherry; tln1:tln1-YPet control (left) and prim:mem-mCherry; tln1:tln1-YPet; cxcr4b:zGrad 33 hpf embryos (right). Lower panels show the Tln1-YPet fluorescence as a heat map. Scale bar = 25 μm. Images are single confocal slices from a z-stack. i, Quantification of the Tln1-YPet fluorescence intensity in control and zGrad-expressing embryos at 33 hpf. Data points, means, and SD are indicated. ****: p<0.0001 (two-tailed t-test). n = number of embryos. j, Experimental strategy to generate embryos with Talin-depleted clones in the primordium. k, Migration of wild-type primordia with clones of control cells (top) and Talin-depleted cells (bottom). Images are maximum-projected z-stacks from Supplementary Video 5. The dotted lines indicate the location of primordium tip. Scale bar = 20 μm. l, Kymographs of migrating chimeric primordia shown in (k) and Supplementary Video 5. m, Quantification of the cumulative migration distance of primordia with clones of control cells and Talin-depleted cells. Dots are means, error bars are SD. *: p=0.03131, p=0.03046 and p=0.04856 (45, 50 and 55 min in the graph) (two-tailed t-test). n = number of embryos.
Extended Data Fig. 7 | Generation and characterization of itgb1b:itgb1b\(^{ΔNPxY}\)-sfGFP mutant knock-in line and F-actin retrograde flow analysis.

a, itgb1b:itgb1b\(^{ΔNPxY}\)-sfGFP expression in a 28 hpf itgb1b\(^{ΔNPxY}\)-sfGFP embryo. Image is sum-projected z-stacks. Scale bars = 0.5 mm.
b, Distribution of Itgb1b-sfGFP and Itgb1b\(^{ΔNPxY}\)-sfGFP in muscle of 33 hpf embryos. Images are single z-slices through muscle at the myotendinous junction imaged and scaled identically. The GFP intensity is pseudo-colored as a heat map. Scale bars = 20 \(\mu\)m.
c, Quantification of the Itgb1b-sfGFP and Itgb1b\(^{ΔNPxY}\)-sfGFP fluorescence intensities at the myotendinous junction (MTJ). Data points, means, and SD are indicated. ****: p<0.0001 (two-tailed Mann-Whitney test). n = number of experiments (used for statistical analysis), N = number of embryos.
d, Images from time-lapse video after photo-bleaching of Itgb1b-sfGFP and Itgb1b\(^{ΔNPxY}\)-sfGFP at the myotendinous junction. GFP intensities are pseudo-colored as heat maps. Scale bars = 10 \(\mu\)m.
e, Graph of Itgb1b-sfGFP and Itgb1b\(^{ΔNPxY}\)-sfGFP fluorescence intensities over time before and after photo-bleaching. The GFP fluorescence intensities are normalized to the minimal intensities after photo-bleaching. Dots indicate mean intensities and error bars are SD. n = number of experiments, N = number of embryos.
f, Plot of the percent recovery of Itgb1b-sfGFP and Itgb1b\(^{ΔNPxY}\)-sfGFP fluorescence intensities at 1 min after photo-bleaching shown in e. n = number of experiments (used for statistical analysis), N = number of embryos. Data points, means, and SD are indicated. ****: p<0.0001 (two-tailed t-test).
g, Images of F-tractin-mNeonGreen localization at the apical sides of wild-type and itgb1b\(^{-/-}\) primordium superficial cells (top). White arrows indicate the direction of migration. Scale bar = 2 \(\mu\)m. Images are single optical sections from Supplementary Video 6. Kymographs of Supplementary Video 6 along the dotted line indicated in top images (bottom). The dotted cyan line indicates the rate of actin flow.
h, Protrusion rates in wild-type and itgb1b\(^{-/-}\) mutant primordium basal cells. Data points, means, and SD are indicated. n.s.: p=0.3167 (two-tailed t-test). n = number of cells.
i, Plot of the protrusion rate versus the actin flow rate in individual primordium basal cells. n = number of cells.
Extended Data Fig. 8 | See next page for caption.
Extended Data Fig. 8 | LamC1-sfGFP mobility, Embryogram workflow, and basement membrane stiffness measurements. a, Optical sections along the indicated planes of a z-stack of the primordium and the BM labeled with LamC1-sfGFP from Supplementary Video 7. LamC1-sfGFP was bleached in front of the primordium in a hexagonal pattern. Scale bar = 50 μm. b, Hexagonal bleach pattern on LamC1-sfGFP-labeled BM underneath the migrating primordium (left). Dotted line indicates location of intensity profile shown on right. Scale bar = 5 μm. The image is a maximum-projected z-stack. c, FRAP analysis of LamC1-sfGFP and extracellular mCherry in heat-shocked hsp70I:sec-mCherry; lamC1:lamC1-sfGFP embryos. Images from the time course are shown on the left and quantification of fluorescence recovery is shown on the right. Scale bar = 10 μm, error bars = SD, n = measurements from N embryos, dots = means, n was used for statistical analysis. d, Extended FRAP analysis of LamC1-sfGFP over 50 min in lamC1:lamC1-sfGFP embryos. Images from time course are shown on the left and quantification of fluorescence recovery is shown on the right. Scale bar = 10 μm, error bars = SD, dots = means, n = measurements from N embryos. n was used for statistical analysis. e, Image of Embryogram application user interface. f, In Embryogram, candidate locations for the bleached markers are identified by a grid search (1), clustered in the XY-plane (2), and then along the Z-axis (3). We match these candidates with a regular hexagonal grid using the iterative closest point algorithm (4). Markers are tracked in subsequent frames using optical flow and numerical optimization. The user can manually offset rigid body motions caused by the movement of the microscope, sample movement or sample growth (6). The displacement of each dot is calculated using the mesh for the first time frame as the relaxed reference (7). To perform finite element analysis (FEA), the user constructs a volumetric tetrahedral mesh above, below or both (8) and inputs the Young’s modulus and the Poisson ratio of the material. The results of the FEA can be exported and visualized in other software packages such as ParaView (9). For detail see Supplementary Note 1. g, 28 hpf embryos with labeled skin and BM before and after surgical skin removal. Images are maximum-projected z-stacks. Scale bar = 100 μm. h, Deskinned and collagenase-treated lamC1:lamC1-sfGFP, cdh1:cdh1-TagRFP embryo. Image is a maximum-projected z-stack. Scale bar = 50 μm. i, TEM-image of the BM underneath the primordium. The semi-transparent yellow line traces the BM and the black lines indicate the thickness of the BM. Scale bar = 1 μm. j, Bright-field image of a deskinned embryo tail with the cantilever during an AFM measurement (left). A grid of 8x8 squares (20 μm by x 20 μm) on the BM was probed for its stiffness (square in left image) and the resultant stiffness map is shown on the right. Scale bar = 1 mm. k, Representative force curves showing the approach (red) and retraction (blue) curves for a deskinned embryo (top) and a collagenase-treated deskinned embryo (bottom). Cross-hairs indicate contact point position and force. Red dots on the approach curves indicate the first 200 nm from the contact point. The fit to the baseline and the Hertz model is indicated by a dotted black line. l, Analysis of the effect of repeated probing of the same area by AFM. The left image is a montage of the stiffness values obtained for the same location after measurements 1 to 100. The order of the measurements is indicated by the arrows. The force curves for the first and 100th measurements are shown on the right. The fit to the Hertz model is indicated in cyan. m, Quantification of the stiffness of the BM of deskinned embryos when fitting the first 500 nm after the contract point to the Hertz model. Data points, mean and SD are shown. Values for the fit of the first 200 nm to the Hertz model are shown for comparison. ****: p<0.0001 (two-tailed Mann-Whitney test). n, Representative force curves that meet (left) and do not meet (right) the indicated quality criteria.
Extended Data Fig. 9 | See next page for caption.
Extended Data Fig. 9 | Distribution of stresses under the skin, under the primordium, and in the absence of the primordium. a, Images of LamC1-sfGFP (left) and basal skin cell membranes (middle) from Supplementary Video 9. The LamC1-sfGFP intensity is pseudo-colored as a heat map. The area outlined by a dotted line was analyzed using Embryogram to calculate the traction stresses (right) pseudo-colored as temperature map (right). The arrowhead indicates a spot of transient accumulation of LamC1-sfGFP. Images are maximum-projected z-stacks. Scale bar = 5 μm. b, Quiver plots of the BM displacement at 0 min in the XY- and XZ-planes. The XZ-plane quiver plot shows a subset of the vector field outlined by the orange rectangle. The magnitude of the vectors was increased by a factor of 3 for visualization purposes. Scale bar = 5 μm. c, Image of cell membrane at −1 min with arrows indicating the direction of movement from time point −1 min to 0 min as determined by PIV. Vector magnitudes are magnified three-times. Scale bar = 5 μm. d, Quantification of traction stresses. Traction stresses at the three vertices closest to a given wrinkle were averaged. Individual data points are shown. Individual data points are indicated. **: p = 0.0027 (−1 min vs. 0 min), p = 0.0043 (0 min vs. 2 min) and n.s.: p = 0.1479 (−1 min vs. 2 min) (two-tailed paired t-test). n = number of measured cells, N = number of embryos, n was used for statistical analysis. e, Quantification of LamC1-sfGFP accumulation during BM wrinkling. Intensity profiles were obtained from a line plot across the BM wrinkle at 0 min indicated by the arrowhead in a, and from line plots at the same location of the images at the time points −1 min and 1 min. Intensities were normalized to the mean intensities at time point −1 min. Mean and SD are shown. n = number of measurements, N = number of embryos. f, Deformation of the BM before, during, and after primordium (magenta) migration. Images are from Supplementary Video 8. The white arrow indicates the direction of migration. Scale bar = 5 μm. g, Quantification of the displacement of bleached marks (yellow circles 1–4 in f) relative to control bleached marks (cyan circles in f). h, Quiver plot of the stresses in the direction of primordium migration. The magnitude of the stress vectors is color-coded. i, Distribution of the tensile and shear stresses around the migrating primordium outlined by a dotted line. The value of each unique component of the stress tensor is colored as a temperature map. The X and Y direction are indicated. The Z direction is orthogonal to the image plane. j, Experimental design of the stress analysis with blocked primordium migration. k, Images of a heat-shocked control embryo at 0 min and 80 min of Supplementary Video 8. The dotted line indicates the region used for the analysis. Images are maximum-projected z-stacks. Scale bar = 50 μm. l, Quiver plot of the displacement vectors shown along the Z, Y and X axes. The magnitude of the displacement vectors is color coded. Scale bar = 10 μm. m, Quiver plot of the displacement vectors projected in the XY-plane. The magnitude of the vectors was increased twofold. Scale bar = 10 μm. n, Distribution of the traction stress magnitudes color-coded using a temperature map. Scale bar = 10 μm. o, Quiver plot of the stresses in the direction of horizontal myoseptum. The magnitude of the stress vectors is color-coded. Scale bar = 10 μm. (l–o) Data correspond to the at the 80 min time point of Supplementary Video 10.
Extended Data Fig. 10 | See next page for caption.
Extended Data Fig. 10 | The primordium is a continuously migrating tissue. a, Itgb1b-tomato-to-Itgb1b-sfGFP (left) and Itgb1b-tomato-to-Itgb1bΔNPxY-sfGFP (right) ratio images in trunk muscle cells. Images are single optical slices from z-stacks. Ratios are color-coded as indicated. Scale bar = 25 μm. b, Quantification of ratios Itgb1b-tomato to Itgb1b-sfGFP and Itgb1b-tomato to Itgb1bΔNPxY-sfGFP at the myotendinous junction and lateral sides of muscle cells. Data points, means, and SD are indicated. ****: p<0.0001 (two-tailed Mann-Whitney test). n = number of measurements at indicated locations, N = number of embryos, n was used for statistical analysis. c, Image of Cxcr4b-EGFP and membrane-tethered Kate2 expressed from the Cxcl12a sensor in the primordium. Image is a maximum-projection of a z-stack. Scale bar = 25 μm. d, Quantification of the Cxcr4b-EGFP/Kate2 ratio across the primordium. Mean (black line) and SD (gray lines) are shown. n = number of embryos. e, Illustrations and predictions for two models of tissue migration. f, Quantification of junction length (left) and the cumulative migration distance over time for three primordia. g, Trajectories of individual primordium cells (left) and frequency plots for angles between any two given cell velocity vectors (right). h, Localization of Cadherin-2-mCherry and membrane-tethered EGFP in the primordium (left). Cdh2-mCherry fluorescence intensity pseudo-colored as a temperature map (middle) and on the primordium's rear at higher magnification (right). Images are single confocal slice from the z-stack. Scale bars = 25 μm (left) and 10 μm (right). i, Images of slices from a z-stack of 32 hpf TgBAC(cxcr4b:EGFP:CaaX) embryos stained for F-actin (left) or phospho-MLC (right) and GFP. Orthogonal views are shown.
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*Give P values as exact values whenever suitable.*

☐  For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
☐  For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
☐  Estimates of effect sizes (e.g. Cohen’s d, Pearson’s r), indicating how they were calculated

*Our web collection on statistics for biologists contains articles on many of the points above.*

Software and code

Policy information about availability of computer code

Data collection  No software was used for data collection.

Data analysis  All images were analyzed using Fiji (Image J v1.0 and v2.0). Custom codes were written and included in this article. All statistical analyses were performed using GraphPad Prism 9. The atomic force microscopy measurements were analyzed with a custom written macro (attached on the manuscript) based on Rasylum version 0.4.0. ([https://github.com/Instone8/Rasylum](https://github.com/Instone8/Rasylum) and [https://doi.org/10.1016/j.rasinfo.2019.05.001](https://doi.org/10.1016/j.rasinfo.2019.05.001)). Stresses were analyzed using the software embroygram which is included in the submission ([https://zenodo.org/record/5752146#Ya5X0y81Qq](https://zenodo.org/record/5752146#Ya5X0y81Qq)). The atomic force microscope was controlled using the Asylum Research software package Version IX (AR Software). Cell tracking was partly done using Imaris Version 8.0 (Bitplane, Oxford Instruments). Confocal image collection and tiled images were stitched using the LAS X Life Science Microscope Software (Leica). Stresses were visualized using ParaView (ParaView-5.8.1, Kitware). Polar diagrams were drawn using MATLAB (MathWorks). Spinning disk confocal microscopy images were collected using NIS-Elements (Nikon).

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Data

Policy information about availability of data

All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:
- Accession codes, unique identifiers, or web links for publicly available datasets
- A list of figures that have associated raw data
- A description of any restrictions on data availability

The source data for Figs. 1–8 and Extended Data Figs. 1–10 are provided as separate excel files. The code for the software embryogram is deposited with Zenodo (https://zenodo.org/record/5762146#.Ya5X0y-B1QJ). The codes for image analysis are provided as a zip file. The Ensembl accession numbers for the genes modified in this study are:

- Itgb1b: ENSDARG00000104484 (ZFIN:Acc:ZDB-GENE-030909-10)
- Itgb1a: ENSDARG00000071863 (ZFIN:Acc:ZDB-GENE-060803-2)
- Tn1: ENSDARG000000100729 (ZFIN:Acc:ZDB-GENE-031002-48)
- Tn2a: ENSDARG000000117901 (ZFIN:Acc:ZDB-GENE-040724-263)
- Tn2b: ENSDARG00000110913 (ZFIN:Acc:ZDB-GENE-130325-1)

All other data supporting the findings of this study are available from the corresponding authors on reasonable request.

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Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

- **Sample size**
  No sample size calculation was performed. The sample size is chosen based on work of other groups using zebrafish embryos as a model system.

- **Data exclusions**
  No data were excluded from the analysis.

- **Replication**
  All attempts at replication were successful. Special cases are clearly stated in the manuscript.

- **Randomization**
  No randomization methods were used to allocate samples into experimental groups.

- **Blinding**
  Blinding is not relevant to this study. The experiments require the investigators to group the data between control and testing condition to quantify differences. However, embryos of different genotypes were imaged and grouped based on genotype after image collection for data analysis. Thus, at the time of data collection the genotype was unknown.

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| Materials & experimental systems | Methods |
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| ✔ Eukaryotic cell lines | ✔ Flow cytometry |
| ✔ Palaeontology and archaeology | ✔ MRI-based neuroimaging |
| ✔ Animals and other organisms | | |
| ✔ Human research participants | | |
| ✔ Clinical data | | |
| ✔ Dual use research of concern | | |

**Antibodies**

- Antibodies used: rabbit anti-collagen IV (Abcam, Cambridge UK, cat no. ab6586)
- rabbit anti-Phospho-Myosin Light Chain 2 (Ser19) (Cell Signaling Technology, cat. no. 3671, lot no. 6)
Validation
Collagen IV: https://www.abcam.com/collagen-iv-antibody-ab6586.html and Maaike C.W. van den Berg et al., 2019, Cell Report
pMLC2(Ser19): https://www.cellsignal.com/products/primary-antibodies/phospho-myosin-light-chain-2-ser19-antibody/3671 and Sandra Ernst et al., 2012, Development, Jaydeep Sidhaye and Caren Norden, 2017, eLife
Fibronectin: https://www.sigmaaldrich.com/GB/en/product/sigma/F3648 and Jaydeep Sidhaye and Caren Norden, 2017, eLife
Chondroitin sulfate: https://www.thermofisher.com/antibody/product/Chondroitin-Sulfate-antibody-clone-CS-56-Monoclonal/MA1-83055
Same clone from the different manufacture was used in Jaydeep Sidhaye and Caren Norden, 2017, eLife
Paxillin: https://www.bdbiosciences.com/en-us/products/reagents/microscopy-imaging-reagents/immunofluorescence-reagents/purified-mouse-anti-paxillin-610051
Sofia Hirth et al., 2016, PlosOne
pPaxillin(pTyr118): https://www.novusbio.com/products/paxillin-antibody_nb2-24459
Hannah M. Olson and Alex V. Nechiporuk, 2021, Developmental Biology
GFP (Torrey Pines Biolab): https://www.labome.com/product/Torrey-Pines-Biolab/T401.html
GFP (Covance): Gayatri Venkiteswaran et al., 2013, Cell [https://doi.org/10.1016/j.cell.2013.09.046]
mCherry (Covance): John Wang et al., 2018, Developmental Cell [https://doi.org/10.1016/j.devcel.2018.07.015]
anti-rabbit Cy3: https://www.jacksonimmuno.com/catalog/products/711-165-162
anti-mouse Cy3: https://www.jacksonimmuno.com/catalog/products/115-165-003
anti-goat Alexa488: https://www.jacksonimmuno.com/catalog/products/705-546-147
anti-DIG antibody coupled to alkaline phosphatase: https://www.sigmaaldrich.com/GB/en/product/roche/11093274910
AlexaFluor594 Phalloidin: https://www.thermofisher.com/order/catalog/product/A12381

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For laboratory work with field-collected samples, describe all relevant parameters such as housing, maintenance, temperature, photoperiod and end-of-experiment protocol OR state that the study did not involve samples collected from the field.

Ethics oversight
This study was performed in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. All of the animals were handled according to approved institutional animal care and use committee [IACUC] protocols [IA16-00788, AMEND20200320] of the NYU School of Medicine.

Note that full information on the approval of the study protocol must also be provided in the manuscript.