*mafba* and *mafbb* differentially regulate lymphatic endothelial cell migration in topographically distinct manners

**Graphical abstract**

**Highlights**

- *mafba* and *mafbb* are required for facial lymphatic development and function
- *mafbb* is dispensable for lymphatic development in the trunk
- *mafba* and *mafbb* regulate directionality of lymphatic endothelial cell migration
- Differential signaling regulates *mafba* and *mafbb* facial lymphangiogenesis

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**In brief**

Arnold et al. investigate the role of transcription factors Mafba and Mafbb during lymphatic development. Using zebrafish mutants and live imaging in conjunction with molecular analysis, they identify topographically distinct requirements of these transcription factors. In the face, *mafba* and *mafbb* steer the direction of lymphatic endothelial cell migration.
**mafba** and **mafbb** differentially regulate lymphatic endothelial cell migration in topographically distinct manners

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**SUMMARY**

Lymphangiogenesis, formation of lymphatic vessels from pre-existing vessels, is a dynamic process that requires cell migration. Regardless of location, migrating lymphatic endothelial cell (LEC) progenitors probe their surroundings to form the lymphatic network. Lymphatic-development regulation requires the transcription factor MAFB in different species. Zebrafish Mafba, expressed in LEC progenitors, is essential for their migration in the trunk. However, the transcriptional mechanism that orchestrates LEC migration in different lymphatic endothelial beds remains elusive. Here, we uncover topographically different requirements of the two paralogs, Mafba and Mafbb, for LEC migration. Both mafba and mafbb are necessary for facial lymphatic development, but mafbb is dispensable for trunk lymphatic development. On the molecular level, we demonstrate a regulatory network where Vegfc-Vegfd-SoxF-Mafba-Mafbb is essential in facial lymphangiogenesis. We identify that mafba and mafbb tune the directionality of LEC migration and vessel morphogenesis that is ultimately necessary for lymphatic function.

**INTRODUCTION**

Lymphatic vascular networks are found in the majority of tissues and organs, and their functions include immune surveillance, body fluid homeostasis, and fat absorption (Koltowska et al., 2013). The diverse functionality of lymphatic vessels is achieved by distinct morphological features across the network. Despite these differences, the current dogma states that common molecular mechanisms drive lymphatic development.

Both in zebrafish and mouse, lymphatic vessels originate from different sources. Lineage-tracing studies in mice have shown that lymphatic endothelial cells (LECs) originate from veins (Karkkainen et al., 2004; Wigle and Oliver, 1999), but also in heart, skin, and mesentery, they can be derived from a non-venous origin (Klotz et al., 2015; Martinez-Corral et al., 2015; Pichol-Thievend et al., 2018; Stanczuk et al., 2015). Zebrafish lymphatic development is studied primarily in two beds, face and trunk, which are morphologically distinct. Developing trunk lymphatics originate solely from the posterior cardinal vein (PCV) and form the trunk lymphatic network, consisting of the thoracic duct (TD), the dorsal longitudinal lymphatic vessel (DLLV), and the intersegmental lymphatic vessels (ISLVs) (Küchler et al., 2006; Yaniv et al., 2006). The facial lymphatics arise from venous origins, the common cardinal vein (CCV) and the primary head sinus (PHS), and from a non-venous origin, the ventral aorta lymphangioblast (VA-L) (Eng et al., 2019; Okuda et al., 2012). The facial LEC progenitors come together to form the lateral facial lymphatic (LFL), the medial facial lymphatic (MFL), the lymphatic branchial arches (LAAs), and the otolithic lymphatic vessel (OLV) (Okuda et al., 2012). Noticeably, regardless of the origin, lymphatic- vessel identity is marked by the transcription factor Prox1 (Koltowska et al., 2015a; Martinez-Corral et al., 2015; Pichol-Thievend et al., 2018; Shin et al., 2016; Wigle and Oliver, 1999). Moreover, the Vegfc-Vegfr3 signaling cascade is required for LEC sprouting, proliferation, and migration (Karaman et al., 2018; Koltowska et al., 2013). Thus, common molecular mechanisms regulate main lymphangiogenesis in different species.

Recent work in zebrafish has uncovered signaling nuances in the regulation of face and trunk lymphatic development. Experiments disrupting Vegfc and Vegfr3 signaling have revealed their differential requirements during lymphangiogenesis (Astin et al., 2014; Bower et al., 2017a, 2017b; Shin et al., 2016). Moreover,
facial lymphangiogenesis is dependent on both Vegfc and Vegfrd signaling via their receptors Kdr and Flt4 (orthologs of Vegfr2 and Vegfr3) (Bower et al., 2017a, 2017b; Vogrin et al., 2019). In contrast, trunk lymphangiogenesis is regulated by Vegfc signaling through Flt4. Thus, a ligand’s ability to compensate for the other is location dependent. It remains unclear if these topographical requirements for different ligands and receptors are precursors to unique downstream transcriptional outputs.

Work from Koltowska et al. (2015b) uncovered that the transcription factor Mafba is part of the Vegfc-SoxF pathway and is essential for trunk LEC migration. Intriguingly, the facial lymphatic phenotype in mafba mutants is mild, with only a slight reduction in the LFL, MLF, and OLV, suggesting that other transcription factors are required for the formation of this lymphatic bed. Similarly, in mouse, Mafb regulates morphogenesis of a subset of lymphatic beds, including the skin and diaphragm (Dieterich et al., 2020; Rondon-Galeano et al., 2020), further supporting that a lymphatic-bed-specific transcriptional code may exist. As a consequence of the Teleost whole-genome duplication, Mafb and mafb, mafbb remains expressed in mafba mutants and therefore serves as a promising candidate to explore in lymphatic-bed-specific development.

Here, we exploit the transcription factors Mafba and Mafbb as a model to uncover molecular mechanisms orchestrating lymphatic development in a topographically dependent manner. We uncovered a unique transcriptional requirement for the development of different lymphatic beds, with mafba and mafbb together directing facial lymphatic development. We show that, mechanistically, Mafba and Mafbb are required for directional migration of LECs in the facial lymphatics but not for their motility. The differential lymphangiogenic requirement of mafba and mafbb in the face and trunk creates possibilities for understanding how diversity is generated between lymphatic beds.

RESULTS AND DISCUSSION

mafbb has a compensatory role in lymphatic development in the facial lymphatic bed

We have previously shown that mafbb is expressed in mafba mutant endothelium (Koltowska et al., 2015b). To assess the role of mafbb in lymphatic development, we generated mafbb mutants using CRISPR-Cas9 technology. We introduced a 10 base pair (bp) deletion in the critical basic region leucine zipper (BRLZ) (Figure S1A), creating a premature stop codon resulting in a predicted loss-of-function allele. Phenotypic analysis of Tg(-5.2lyve1b:DsRed2) and Tg(fli1a:EGFP) transgenic zebrafish lines at 5 days post-fertilization (dpf) revealed morphologically intact facial lymphatics in mafbb mutants with a small decrease in cell number compared with mafbb heterozygotes (Figures 1A and 1C). Trunk lymphatic development and cell number was unperturbed in mafbb mutants (Figures 1B and 1C). mafbb mutants have normal gross morphology (Figure 1G), growing into viable adults (Figure S1E). To test for maternal contribution, we crossed mafbb mutant females with mafbb heterozygous males, observing no additional phenotypes (data not shown). Therefore, mafbb loss is likely compensated by its paralog, mafba.

To investigate the facial LEC phenotype and address compensation, we generated mafba;mafbb double mutants and found more severe facial lymphatic phenotypes. mafba;mafbb mutants have a 35.9% LEC reduction compared with siblings (Figures 1F and S1B) and disrupted morphology (Figure 1D) compared with mafba mutants with a 25% reduction in face LECs (Figures 1F and S1B) with relatively unaffected morphology (Figure 1D). In addition, mafba;mafbb mutants have weaker Tg(-5.2lyve1b: DsRed2) expression than siblings or single mutants (Figure S1F). mafba;mafbb double-mutant trunk lymphatics show similar LEC numbers compared with mafba mutants at 5 dpf (Figures 1E, 1F, and 1C). This suggests that there are topographical differences in mafba and mafbb requirements across lymphatic vessels.

To investigate differences in other LEC derivatives, we have focused on mural LECs (muLECs), also known as fluorescent granular perihelial cells or brain LECs (Bower et al., 2017a; Galanternik et al., 2017; van Lessen et al., 2017). This revealed that muLEC development was unaffected in mafba, mafbb, or double mutants (Figures 1F and S1G), indicating that muLEC migration is independent of Mafba and Mafbb regulation. Survival severity is comparable between mafba;mafbb double mutants and mafba mutants, and both genotypes show otolithic vesicle (OV) enlargement and absent swim bladders (Figures 1G and S1E). Quantitative real-time PCR did not reveal a striking upregulation of mafba in mafbb mutants, or vice versa (Figure S1D). Thus, we speculate that these two genes play specific roles during facial lymphatic-vessel development rather than follow the mechanisms of genetic compensation. Our findings reveal different
regulatory requirements of lymphatic-vessel formation: mafba is needed for trunk lymphatic development, while both mafbb and mafba regulate facial lymphatic development.

**LEC progenitors are specified independently of mafba and mafbb**

LEC progenitors are still specified in the trunk of mafba mutants and make their way to the horizontal myoseptum (Koltowska et al., 2015b). We asked if the phenotypic severity in mafba-mafbb double mutants originates from LEC-progenitor specification. Using Tg(fli1a:EGFP) and immunostaining for Prox1 at 56 h post-fertilization (hpf) (Figures 2A and 2B), we found specified LECs in both beds for all mutant backgrounds (Figures 2A and 2B). Quantification of double-positive cells in the trunk and face revealed no difference in LEC numbers across all genotypes (Figure 2C), even when normalized to the total number of Tg(fli1a:EGFP)-positive cells within the CCV and facial lymphatic sprout (FLS) (Figure S2A). Defects in proliferation in mafba;mafbb double mutants were ruled out by EdU staining (Figure S2B). Thus, mafba and mafbb are required for lymphatic development after LEC-progenitor specification.

**Facial lymphatic-vessel morphology is disrupted in mafba;mafbb mutants**

To dissect how regulation by mafba and mafbb is translated into tissue organization, we measured different morphological features of the developing facial lymphatic vessels. Surface rendering of Tg(-5.2lyve1b:DsRed2) revealed that the facial lymphatic network was disrupted, with thinner and mis-shaped vessels in mafba;mafbb mutants (Figure 2D). The facial lymphatic vessels are shorter and globular with volume and area reduced by half compared with siblings; however, sphericity was unaltered across the mutant backgrounds (Figure 2E; Video S1). Together, these findings reveal that mafba and mafbb are required for craniofacial lymphatic-vessel morphogenesis and proper vessel organization.

**Lymphatic function is disrupted in mafba;mafbb mutants**

We speculated that the dysmorphic vessels in mafba;mafbb mutants would impact lymphatic function. To test functionality, we injected fluorescein isothiocyanate (FITC)-dextran into the tissue ventral to the LFL and in between the LAA at 5 dpf. We found no difference in FITC-dextran uptake after 30 min between the different genotypes. However, after 3 h, the uptake in the double mutants was significantly reduced compared with the other genotypes (Figures 2F–2H and S2C). Moreover, while fluid uptake in siblings, mafba mutants, and mafbb mutants significantly increased from 30 min to 3 h, there was no change in uptake over time in mafba;mafbb mutants (Figure 2F). The uptake defects were independent of the reduction in vessel size in mafba;mafbb mutants (Figure S2D). These results were confirmed by injection of Qtracker 655 Vascular Labels (hereafter referred to as Qtracker) at 5 dpf (Figures S2E and S2F). These data suggest that the reduced volume and altered morphology of mafba;mafbb mutants impacts their ability to remove fluid from the surroundings at the appropriate rate.

**mafba and mafbb regulate lymphatic-vessel maturation**

Defective vessel morphology in mafba;mafbb mutants was further validated with the transgenic line TgBAC(prox1a:KlaTA4-4xUAS-ADV.E1b:TagRFP) (hereafter referred to as Tg(prox1a:RFP), reporting prox1a expression in the developing lymphatic vasculature. prox1a is expressed in all backgrounds at 5 dpf, and the facial lymphatic phenotypes recapitulated those observed in Tg(-5.2lyve1b:DsRed2) transgenic fish (Figure 3A). A newly discovered morphological feature of facial lymphatics is a valve (Shin et al., 2019). Tg(prox1a:RFP) is expressed in the developing valve at 5 dpf, which appeared to be mis-patterned with 55.1% length reduction in mafba;mafbb mutants compared with siblings (Figures 3A and 3B). To analyze valve morphology, we generated average phenotype images by computationally overlaying single images from siblings and mafba, mafba;mafbb mutants (Figures 3C, 3D, S2G, and S2H). We observed a gradient of phenotype from siblings and mafbb mutants (narrow valve) to mafba;mafbb mutants (unclear valve), in which mafba mutants (broad valve) are a hybrid of the two phenotypes, with increasing phenotypic variability across genotypes (Figures 3C, 3D, and S2G; Video S2). To test valve functionality in the different genetic backgrounds, we injected Qtracker into the facial lymphatic vessels (FLVs) of anaesthetized embryos and imaged 5 min post-injection (Figure 3E). We observed a 70.6%
**A**

*Tg(fli1a:nEGFP)*\(^{+}\);*Tg(prox1a:RFP)*\(^{+}\) overlay,

- **sibling**
- 5dpf
- *mafbba\(^{gfp}\)*
- *mafbba\(^{gfp}\), *mafbbb\(^{gfp}\)*

**B**

Valve length

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<td>*mafbba(^{gfp}), <em>mafbbb(^{gfp})</em></td>
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**C**

*Tg(prox1a:RFP)*\(^{+}\) overlay,

- **sibling**
- 5dpf
- *mafbba\(^{gfp}\)*
- *mafbba\(^{gfp}\), *mafbbb\(^{gfp}\)*

**D**

Rendered *Tg(prox1a:RFP)*\(^{+}\)

- Frontal
- Lateral

**E**

*Tg(fli1a:nEGFP)*\(^{+}\);*Tg(prox1a:RFP)*\(^{+}\);*QTracker™ 655*

- **siblings**
- *mafbba\(^{gfp}\)*
- *mafbba\(^{gfp}\), *mafbbb\(^{gfp}\)*

**F**

Valve leakage

- **Embryo number (%)**
- **ns**
- **ns**

(legend on next page)
increase of embryos with leakage from the FLV to facial collecting lymphatic vessel (FCLV) in mafb;mafbb mutants compared with siblings (Figure 3f). These data suggest that mafb and mafbb are required for the formation and function of the valve in the facial lymphatics and that in their absence, lymph flow cannot be correctly channeled.

**Topographical differences in Vegfc/d-SoxF downstream effectors during lymphatic-vessel formation**

Recent reports have revealed differential requirements of Vegf ligands and receptors during lymphangiogenesis (Astin et al., 2014; Bower et al., 2017a, 2017b; Shin et al., 2016; Vogrin et al., 2019). We have previously shown that mafb expression is dependent on Vegfc-SoxF (Koltowska et al., 2015b). We hypothesize that mafb and mafbb also act downstream of the Vegf-SoxF axis. First, we confirmed that MO-vegfc and -vegfd recapitulate the published mutant phenotypes, while standard control MO did not induce lymphatic phenotypes (Figures S4A and S4B) (Astin et al., 2014; Hogan et al., 2009a, 2009b). We investigated the facial lymphatic phenotype in the previously published sox7 and sox18 mutants at 3 dpf, before the onset of significant edema due to PCV-DA shunts (Herpers et al., 2008) (Figure 4A). The single sox7 or sox18 mutants did not show lymphatic phenotypes, but sox7+/-sox18+/- and sox7-/-sox18+/- embryos showed a reduction in FLS (Figure 4A), recapitulating the sox7 double morpholino phenotype (Figures S4A and S4B).

To uncover the potential direct regulators of mafb, we investigated the presence of cis-regulatory elements (CREs). We identified conserved non-coding DNA sequences in the regions upstream and downstream of mafb using mVista (Dubchak et al., 2000; Frazer et al., 2004; Mayor et al., 2000) and found a 151 bp conserved region 6 kb downstream of mafb (Figure S3A). This element is part of a previously described mafb enhancer driving reporter expression in the venous and lymphatic endothelium at 55 hpf (Quillen et al., 2017). We generated a stable transgenic line, Tg(mafbE1bas:EGFP), which recapitulates endogenous mafb expression (Figures S3C and S3D). We found no changes in the expression of Tg(mafbbE1bas:EGFP) in MO-vegfc, MO-vegfd, MO-sox7, MO-sox18, or double morpholino treatments (Figures 4B and S3F–S3I). We used MEMESuite (Bailey et al., 2009) to identify conserved motifs in the mafb CREs and found 10 predicted transcription-binding sites, none of them Sox7 or Sox18 (Figures S3B and S3E). Together, these observations suggest that this mafb CRE is not regulated by SoxF.

To investigate mafb and mafbb expression in the facial lymphatics in response to Vegfc/d-SoxF signaling, we took a quantitative real-time PCR approach. We isolated LECs and venous endothelial cells (VECs) from dissected heads by sorting for cells double positive for Tg(5.2lyve1b:Venus) and Tg(fli1a:H2B-mCherry) (Figure S4C) and found a trend of mafb and mafbb downregulation in the craniofacial VECs and LECs depleted of Vegfc-Vegfd signaling or SoxF (Figure S4D). Next, we determined if the level of expression of mafb and mafbb is downregulated specifically in LECs. We isolated facial LECs by sorting cells double positive for Tg(prox1a:RFP) and Tg(fli1a:nEGFP) from dissected heads (Figure S4C). We compared expression levels of mafb and mafbb in wild-type embryos and did not observe differences, confirming that both genes are present in developing LECs at similar levels (Figure 4C). We observed that mafb and mafbb expression is reduced in vegfc morphants and that there is a trend of reduced expression in vegfd and vegfc-vegfd morphants (Figures 4D and 4E). We found a significant difference in mafb, but not mafbb, expression in sox7/18 morphants (Figures 4D and 4E). This revealed an unexpected difference in mafb and mafbb regulation in facial lymphatics downstream of Vegfc-Vegfd and SoxF. This shows that the two paralogs are differentially activated downstream of Vegf-SoxF signaling and that mafbb is constitutively expressed and regulated in facial LECs and is required for proper facial lymphatic formation.

**mafbb and mafba are required for face LEC migration**

Our previous work has shown that mafb is essential for LEC migration in the trunk (Koltowska et al., 2015b). To identify if mafb and mafbb play a similar role in facial LEC migration, we used time-lapse imaging. By combining Tg(5.2lyve1b:Venus) and Tg(fli1a:H2B-mCherry) transgenic lines, we documented in real-time the LEC trajectories in all genotypes.

From 40 hpf, we observed initial migration of the FLS from the CCV in an anterior-ventral direction (Figures 4F and S4E;
Tracking the FLS migratory tip or individual cells, we found that in mafb;mafbb mutants, the FLS tip migratory distance was reduced compared with siblings, yet the FLS cells’ velocity was not affected at 40–45 hpf (Figure 4G). In addition, there was no significant difference in the leading and following cells’ velocity within the FLS in sibling and mafb; mafbb double mutants (Figure S4F), suggesting that cell motility was not affected. In siblings, the FLS tip advances over time, as revealed by the increasing or constant mean-squared displacement (MSD) (Figure 4H). However, in mafb; mafbb mutants, this value fluctuates, indicating that the tip progression rate is disrupted, resulting in stalling or changes in the migratory route (Figure 4H).

To establish if the migration defects are caused by an altered trajectory, we quantified cell-movement directionality (Figures 4I and S4H). In siblings, FLS cells collectively migrate in an anterior-ventral direction, predominantly between 120° and 150°, whereas this shifts ventrally to between 150° and 180° in mafb; mafbb double mutants (Figure 4I). To understand these directional differences, we separated leading cells from following cells and repeated the analysis. We observed that sibling leading cells are moving in various directions and sense their environment, with 0°–30°, 90°–150°, and 180°–210° being the most common cell-migration directions (Figure 4I). In mafb; mafbb double mutants, leading cells have instead a strong directional preference for the ventral-posterior direction (180°–210°) (Figure 4I), potentially due to loss of the ability to sense the environment. To further determine how the change of behavior in leading cells impacts the following LECs, we analyzed the followers’ trajectory indices. We found that sibling following cells migrate toward 0°–30°, 90°–120°, and 180°–210°; while in mutants, the following-cell migration is more randomized. Further, while in siblings, the migratory direction of leading and following cells is remarkably similar, in mafb; mafbb double mutants, leading and following LECs have different directional preferences (Figure 4I).

Overall, we see no significant difference in the migratory distance of the FLS or the directionality of collective cell migration in mafb or mafbb single mutants when compared with siblings (Figures 4G and S4H). In mafb mutants, the MSD is the same as for siblings (Figure S4G), while the direction of migration is disrupted in leading cells (Figure S4H), whereas the reverse is found in mafbb mutants where leading cells show the same directional preference as siblings (Figure S4H), but the MSD is similar to mafb; mafbb mutants (Figure S4G). In both mafb and mafbb single mutants, we see a significant increase in velocity (Figures 4G and S4F). As the vessel morphology and function are largely normal in these mutants, we speculate that the velocity increase ensures that the cells migrate the same distance. These data indicate that mafb and mafbb together are necessary for directional LEC migration.

In summary, we have found that facial lymphatic development differentially requires Mafb and Mafbb downstream of Vegfc/d-SoxF to promote the directionality of LEC migration. Ultimately, this work defined that Mafb and Mafbb are required to regulate LEC migration in the face, while Mafba alone regulates this process in the trunk. On the contrary, muLECs develop in a Mafba- and Mafbb-independent manner. This highlights the important differences in regulatory mechanisms driving lymphatic development across the zebrafish embryo. Together, our findings uncovered that selective transcription-factor combinatorial levels drive lymphatic development in a topographically distinct manner. This spatially distinct regulation of lymphangiogenesis suggests that despite the highly conserved developmental programs in lymphatic development, progenitor niches present key differences in transcriptional code, which might change the way we view lymphatic-vessel development on a whole-organism level.

**Figure 4. mafb and mafbb regulate facial LEC migration downstream of Vegfc-Vegfd-SoxF**

(A) (Left) Confocal images of Tg(flt1ABAC;mCitrine) expression in facial lymphatics in sibling, sox7+/+, sox18+/+, sox18+/−, and sox7+/−; sox18+/− embryos at 3 dpf. *: reduced sprout length. Scale bars, 100 μm. (Right) Quantification of facial lymphatic sprout length at 3 dpf in sox7 and sox18 mutant backgrounds.

(B) (Top) Heatmap of Tg(mafbbE1bas:EGFP) signal intensity from confocal projection at 2.5 dpf in face LECs. Gray value intensity scale: 0–113 for control and MO-vegfc/mvegfd, 0–147 for control and MO-vegfc, 0–147 for control and MO-vegfd, 0–210 for control and MO-sox7+MO-sox18. White bracket: quantification area. Scale bar, 50 μm. (Bottom) Quantification of Tg(mafbbE1bas:EGFP) intensity. (Left) Control n = 7; MO-vegfd and MO-vegfc n = 9; MO-vegfc + MO-vegfd n = 8; (right) control n = 7; MO-sox7 n = 9; MO-sox18 n = 14; MO-sox7 + MO-sox18 n = 10. Kruskal-Wallis test: ns p > 0.2372.

(C) Quantitative real-time PCR of mafb and mafbb expression relative to cdh5 in facial LECs at 48 hpf. 12 replicates. Wilcoxon test: ns p = 0.3013.

(D and E) Quantitative real-time PCR of mafb and mafbb expression in face LECs at 48 hpf. (D, left) 6 replicates for control and MO-vegfc and 5 for MO-vegfd and MO-vegfc + vegfd; expression relative to cdh5. (D, right) 6 replicates for control, MO-sox7, 5 for MO-sox18, and 4 for MO-sox7+MO-sox18; expression relative to B actin. (E, left) 6 replicates for control, MO-vegfd, MO-vegfc, and MO-vegfc + vegfd; expression relative to cdh5. (E, right) 6 replicates for control, 5 for MO-sox7 and MO-sox18, and 4 for MO-sox7+MO-sox18; expression relative to B actin. Kruskal-Wallis: p values reported on the graph.

(F) Confocal projections from time-lapse imaging of LEC migration in siblings and double mutants labeled by Tg(5.25yi2e1b:Venus) from 40–48 hpf. Arrows: leading cells in FLS. Scale bars, 50 μm.

(G) (Left) FLS migratory distance (from F) in siblings (n = 4) and double mutants (n = 4). One-way ANOVA: ***p < 0.0001 for siblings versus double mutants; ns for all other comparisons.

(H) FLS tip mean-squared displacement (MSD) (from F) in siblings (n = 4) and double mutants (n = 4). One-way ANOVA: ***p < 0.0001 for siblings versus double mutants; ns for all other comparisons.

See also Figures S3 and S4 and Videos S3, S4, S5, and S6.
Limitations of the study
Due to technical limitations, this study has not identified the mechanisms by which mafba and mafbb are differentially regulated directly downstream of VegF and SoxF signaling. It is also limited in the mechanistic insight by which mafba and mafbb regulate the direction of LEC migration. Future work is needed to further identify the direct downstream targets of Mafba and Mafbb, which orchestrate the cellular processes involved in shaping functional lymphatic vessels.

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SUPPLEMENTAL INFORMATION
Supplemental information can be found online at https://doi.org/10.1016/j.celrep.2022.110982.

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AUTHOR CONTRIBUTIONS
H.A., V.P., and K.K. conceptualized the project, performed experiments, analyzed data, and co-wrote the manuscript. M.H., B.F.-G., R.S., M.G., and S.S.-M. performed experiments and analyzed data. P.R. and A.A. analyzed experiments. B.H. provided unpublished reagents.

DECLARATION OF INTERESTS
The authors declare no competing interests.

REFERENCES


## STAR METHODS

### KEY RESOURCES TABLE

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Software and algorithms

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RESOURCE AVAILABILITY

Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Katarzyna Koltowska (kaska.koltowska@igp.uu.se).

Materials availability
Transgenic lines, gRNA primers and RT-qPCR primers will be made available on request.

Data and code availability
Custom code written for this project has been made available on GitHub and is publicly available. The DOI and URL for the GitHub repository can be found in the Key resources table. Further information required to reanalyse the data generated in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS
Zebrafish work was carried out under ethical approval from the Swedish Board of Agriculture (5.8.18–10590/2018). Zebrafish were housed at the Genome Engineering Zebrafish National Facility (SciLifeLab, Uppsala Sweden) and adults and embryos were housed according to the standard laboratory procedure (Åleström et al., 2020). At 24 hpf embryos were transferred to 200 μM 1-phenyl 2-thiourea for imaging. Published zebrafish lines used were Tg(fli1a:nEGFP)y7 (Lawson and Weinstein, 2002), Tg(-5.2lyve1b:DsRed2)y7107 (Okuda et al., 2012), TgBAC(prox1a:KaiTA4-4xUAS-ADV.E1b:TagRFPtnms5 (Dunworth et al., 2014; van Impel et al., 2014), Tg(kdr-l:ras-Cherry)s916 (Hogan et al., 2009a; 2009b), Tgfltl4BAC:mCitrine)b7135 (van Impel et al., 2014) and Tgfli1a:H2B-mCherry)yq37bh (Baek et al., 2019). The transgenic lines Tg(-5.2lyve1b:Venus)yq42k and Tg(mafbbE1bas:EGFP)yq2kk were generated for this study.

METHOD DETAILS

Mutant generation and identification
mafbb mutants were generated by CRISPR/Cas9 genome editing as described in Gagnon et al. (2014). The guide was designed in the BRLZ domain of mafbb. Zebrafish embryos were injected at one cell stage with the 70–140 ng/μL gRNA and 200 ng/μL Cas9 mRNA. Founder identification was carried out by Fragment Length Analysis (FLA) as described (Carrington et al., 2015). Primers used to generate guides and for FLA are listed in the Key resources table (the target site of the gRNA can be found in capitals for mafbb_g21 primer).

Genotyping for mafbaq42h and mafbbq47bh was carried out using KASP assay according to the manufacturer’s instructions (LGC Biosearch Technologies) and using a OneStepPlus Real-Time PCR System (Thermo Fisher). Primers used are listed the Key resources table.

mafbb CRE identification
Actinopterygian sequences of the complete upstream and downstream non-coding regions of mafbb were aligned using the LAGAN alignment program (Brudno et al., 2003) and peaks of conservation were identified using mVISTA (Dubchak et al., 2000; Frazer et al., 2004; Mayor et al., 2000).

Generating a cis-regulatory transgenic line
We established the transgenic line Tg(mafbbE1bas:EGFP)yq2kk by injecting the construct pT2-cryR;mafbbCE1-basEgfp (Addgene plasmid #90137), at 20 ng/μL with 100 ng/μL tol2 transposase mRNA into the cell of wild type zebrafish embryos at the one-cell stage.

Morpholino-mediated knockdown, genotyping and genome editing
Morpholinos (MOs) used have been previously published: MO-vegfc and MO-vegfd by (Astin et al., 2014), MO-sox7 and MO-sox18 by (Herpers et al., 2008). Standard control morpholino (Gene Tools) was used to test toxicity. 1 nL of MO solutions were injected into the yolk at the one-cell stage. The MO-vegfcSB has been used in the RT-qPCR experiment, while the MO-vegfcT was used for the intensity quantifications. Details of each MO are listed in the Key resources table.

Dye injections
For functional experiments embryos were injected with 1 nL of 5 mg/mL FITC-dextran 10 kDa MW (Sigma) in 0.2 M KCl in ventral to the facial lymphatics. Embryos were imaged on a Leica TCS SP8 DLS microscope 30 min and 3 h post-injection. Embryos were also injected with 1nL of Qtracker™ 655 Vascular labels (Thermo Fisher) at 5 dpf into the region dorsal and ventral to the LFL for uptake experiments and imaged on a Leica TCS SP8 DLS microscope 3 h post-injection.
Embryos anaesthetised with Tricane were injected with 1 nL of Qtracker™ 655 Vascular labels (Thermo Fisher) at 5 dpf in the FLV for leakage experiments and imaged on a Leica TCS SP8 DLS microscope immediately post-injection, approximately 5 min post-injection a Leica TCS SP8 DLS microscope.

**Immunostaining**

Prox1 and GFP were detected by immunofluorescence staining as previously described (Le Guen et al., 2014; Shin et al., 2016), with the addition of a 25 min Proteinase K treatment at RT as described in (Koltowska et al., 2015b). To view 5-ethynyl-2'-deoxyuridin (EdU) incorporation into newly synthesised DNA we followed the method previously described in (Koltowska et al., 2021), with the adaptation that embryos were treated in EdU at 54 hpf and fixed at 56 hpf. Tg(-5.2lyve1b:Venus)ss11kh staining was carried out using the method described above. The details of the antibodies used are listed in the Key resources table.

**In situ hybridization**

The coding region of mafb was cloned by In-Fusion® HD Cloning (Takara Bio) using the primers listed in the Key resources table into a PCS2+ vector linearized with XbaI and EcoRI. The probe was synthesised using the MAXIscript T7 Transcription Kit (Invitrogen) on the vector after linearization with BbvCI. Primers for probe synthesis can be found in the Key resources table. In situ hybridization against mafb was performed as previously described (Kartopawiro et al., 2014). For imaging, embryos were embedded in 8% agarose and sectioned in a Microm HM 650V microtome (Thermo-Fisher) with a slice thickness of 60 μm. Sections were mounted in Mowiol and imaged on a DMI8 Leica microscope with a N Plan L 20× objective (Leica).

**Image acquisition**

Transgenic fluorescent embryos were mounted laterally in 1% low-melting agarose and imaged in the face or trunk using a Leica TCS SP8 DLS microscope. The following objectives were used: Fluotar VISR 25× water objective (objective number: 11504416), HC PL APO CS2 40× water objective (objective number: 11506360) and HC PL Fluotar 20× dry objective (objective number: 11506519). For time-lapse imaging, embryos were imaged overnight from 40 hpf acquiring a frame approximately every 10 min. Brightfield images were taken using a Leica Fluorescent Stereo Microscope M165 FC and 1x objective (objective number: 10450028). Embryos were mounted in 3% methylcellulose. Images were processed using ImageJ 2.0.0 (Schindelin et al., 2012).

**VAST system imaging**

mafbuq4bh, mafbbuq47bh, mafbauq4bh, mafbbuq47bh and sibling embryos were imaged at 5 dpf using the Vertebrate Automated Screening Technology (VAST) system (Almstedt et al., 2020; Gudmundsson et al., 2019). Fluorescent images of the facial lymphatics, muLECs loops and trunk lymphatics were acquired using the Tg(-5.2lyve1b:Venus)ss11kh line and used in the quantifications in Figure S1G.

**Fluorescence-activated cell sorting (FACS) and gene expression analysis**

Whole embryos were harvested at 3 dpf from an incross of mafba+/uq4bh;mafbb+/u or mafba+/uq4bh;mafbb+/uq47bh and selected based on phenotype. Embryos were harvested at 48 hpf and the heads were dissected by making an incision posterior of the otolithic vesicle. Isolation of cells was performed as previously described (Kartopawiro et al., 2014). Double positive cells were sorted for either transgenes Tg(prox1a:RFP)ss13b and Tg(fli1a:nEGFP)y7, or Tg(-5.2lyve1b:Venus)ss11kh and Tg(fli1a:H2B-mCherry)ss13bh directly into 300 μL TRizol™ LS Reagent (Thermo Fisher). RNA was extracted using the Quick-RNA Microprep kit (Cultura Bioscience) according to manufacturers’ instructions. cDNA synthesis was performed with the SuperScript™ VILO™ cDNA Synthesis Kit (Thermo Fisher) according to manufacturer’s instructions with a longer incubation at 42°C of 120 min cDNA was amplified using the SsoAdvanced™ PreAmp Supermix (Bio-Rad) according to manufacturer instructions. Each sample was amplified in two replicates. RT-qPCR was carried out using the SsoAdvanced Universal SYBR Green Supermix (Bio-Rad) according to manufacturer’s instructions on CFX384 Touch Real-Time PCR Detection System (Bio-Rad). Data were analysed using the CFX Maestro Software (Bio-Rad). Changes in gene expression were calculated relative to the expression of a control gene, and normalized to the corresponding controls. Primers are listed in the Key resources table.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

**LEC number quantification**

Image quantification was performed using z-stacks and manual counting in ImageJ 2.0.0. For LECs at 5 dpf, double-positive cells for Tg(-5.2lyve1b:DsRed2)ss11kh and Tg(Fli1a:nEGFP)y7 in face and trunk lymphatic vessels were scored. For LEC specification at 2 dpf double-positive cells for Prox1 and Tg(Fli1a:nEGFP)y7 we scored. For image representation, trunk images were rendered using Imaris v9.3.0 with a surface detail of 1μm for Tg(Fli1a:nEGFP)y7. Both channels, for Tg(Fli1a:nEGFP)y7 and for Prox1, were then masked and these channels used for representative images. Final images were processed using ImageJ 2.0.0 for both face and trunk.
Image rendering
For volume, area and sphericity quantifications, surface rendering of the \( Tg(-5.2lyve1b:DsRed2)^{az101} \) channel was carried out using Imaris v8.3.0. Surfaces were rendered with a surface detail of 2 \( \mu m \), and vessels that were not part of the facial lymphatic network were removed (Figure 2D). Measurements were acquired using the statistics tool.

Valve surface rendering of \( Tg(prox1a:RFP)^{vms5} \) was carried out using Imaris v9.3.0. Surfaces were rendered with a surface detail of 0.1 \( \mu m \), and the valve region extracted from the vessel (Figure 3D).

Lymphatic functional analysis
The lymphatic function analysis by dextran uptake was measured as a ratio of inside:outside average pixel value of the LFL. Image analysis was carried out using ImageJ 2.0.0. ROIs were drawn on individual z-slices which included the main vessel of the LFL (inside) or ventral to the LFL (outside), using the \( Tg(-5.2lyve1b:DsRed2)^{az101} \) channel. The average pixel value was then taken for the FITC-dextran channel for the ROI on each z-slice and the mean pixel value was calculated across the z-slices, which varied between 5-9 slices depending on the thickness of the vessel. The average pixel value at 30 min and 3 h post-injection was normalised to the level from before FITC-dextran injection by taking the average of the z-slices of the LFL. To visualise the FITC-dextran distribution of the images in Figure 3G, a fluorescent profile was taken through the LFL at the anterior branching point in a z-stack of the slices spanning the LFL. A line was drawn intersecting the LFL and including the region dorsal and ventral to the LFL. The grey values for FITC-dextran and \( Tg(-5.2lyve1b:DsRed2)^{az101} \) were plotted against the position along the line to show the relationship between the LFL and dextran absorbance.

For lymphatic function analysis using Qtracker\textsuperscript{TM} 655 Vascular Labels uptake was measured inside the LFL at 3 h post-injection. Image analysis was carried out using ImageJ 2.0.0. ROIs were drawn on individual z-slices which included the main vessel of the LFL using the \( Tg(prox1a:RFP)^{vms5} \) channel. The mean pixel value calculated across the z-slices, which varied between 5-9 slices depending on the thickness of the vessel. This method differed from that of the FITC-dextran uptake experiment as Qtracker\textsuperscript{TM} 655 Vascular Labels was more stable within the tissue and was not cleared at 3 h post-injection despite uptake into the tissue for some genotypic backgrounds.

Fluorescent intensity quantification
For fluorescent intensity quantifications, images used in Figure 2B were rendered in Imaris v9.3.0 with a surface detail of 0.75 \( \mu m \) using \( Tg(-5.2lyve1b:DsRed2)^{az101} \). Both channels were then masked and used for representative images. In the head, the distal section of the developing lymphatic sprout was isolated by hand using the Cut Surface tool. Strongly GFP-positive cells with a morphology distinct from endothelial (Figure S3G) were removed through rendering of \( Tg(mafbE1bas:EGFP)^{u2kk} \). In the trunk, the PCV was rendered using \( Tg(-5.2lyve1b:DsRed2)^{az101} \) and the sprouting vessels were trimmed manually. The average intensity of the trimmed rendered surfaces was calculated for both channels using the Statistics tool in Imaris v8.3.0. \( Tg(-5.2lyve1b:DsRed2)^{az101} \) was confirmed not to change in intensity between the controls and MO-injected \( Tg(-5.2lyve1b:DsRed2)^{az101};Tg(mafbE1bas:EGFP)^{u2kk} \) embryos (Figure S3H), he normalised GFP intensity value for each image was calculated by dividing the mean intensity value of the GFP channel by the mean intensity value of the DsRed channel in the rendered lymphatic surfaces. The normalized GFP values of every imaging session were then divided by the median of the controls of that session.

Average image analysis
Average image overlays were generated using MATLAB and elastix (Klein et al., 2010), with the method previously described (Allalou et al., 2017). Embryos of each genotypic group were aligned to generate an average pattern using Iterative Shape Averaging (ISA) algorithm (Rohlfing et al., 2001), and iteratively improved using an affine transformation and non-linear transformation.

Cell migration analysis
Movies were prepared for cell tracking in ImageJ using the Registration tool to account for drift during the course of imaging, then tracking was carried out using the Manual Tracking tool.

The Mean Squared Displacement (MSD) is computed by calculating the MSD for each frame as the sum of squared distances to the tip position for that frame, this is then divided by the number of objects.

Cell directionality was quantified by comparing the angle of each track to the first position of that track. This is then represented as an angle in the polar histogram. The angle interval for each sector (=bins) is set to 30 degrees, and the distance to the origin corresponds to the number of positions within that angle interval.

Statistical analysis
Normality of all numerical datasets was tested with a Shapiro-Wilk test. When the number of datapoints was lower than 5, the data were assumed not to be normally distributed.

For pair-wise comparison, an unpaired two-tailed Student’s \( t \)-test was run on normally distributed data, while a Mann-Whitney test was run if normality was not confirmed. A Wilcoxon test was run on paired data.
For multiple comparisons, a one-way ANOVA was run on normally distributed data, while a Kruskal-Wallis test was run if normality was not confirmed. Both were run together with Dunn’s multiple comparison test. When two independent variables were present, a two-way ANOVA with Sidak’s multiple comparison test was used.

For qualitative data, a Chi-square test was used. For survival curves, a Gehan-Breslow-Wilcoxon test was used.