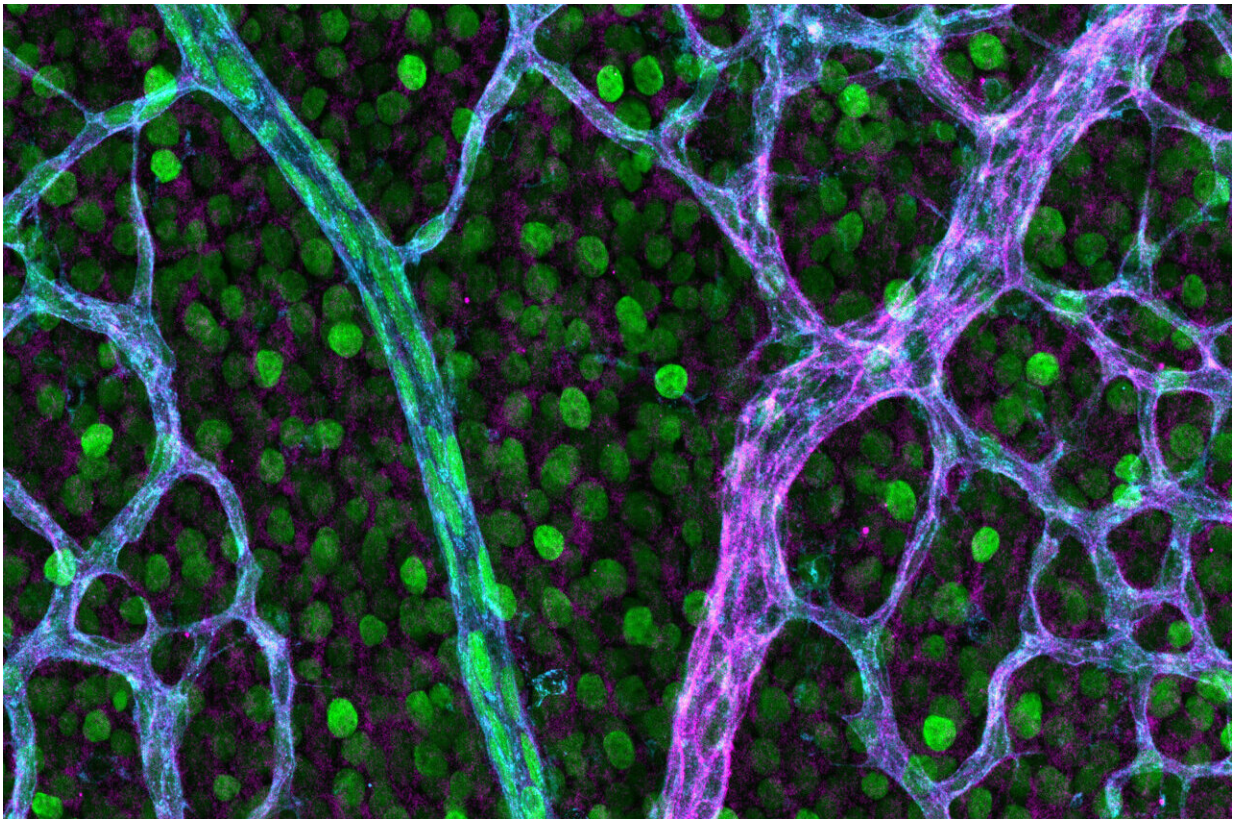


A balancing act: EphB4 and ephrin-B2 regulate artery formation

April 3 2024, by Jeanine Müller-Keuker



EphB4 and ephrin-B2 expression in the retinal vasculature. Credit: MPI for Molecular Biomedicine / Jonas Stewen

Conditions affecting the arterial vasculature present pressing challenges in global health. Yet, the complex mechanisms underlying artery

formation remain elusive, impeding the development of new treatments.

A team of researchers from the Max Planck Institute for Molecular Biomedicine, led by Mara Pitulescu and Ralf Adams, have uncovered new insights into how arteries form. Building on their earlier discovery of how Notch-signaling directs "tip" cells to become arteries, their latest study [published](#) in *Nature Communications* reveals the crucial role of EphB4 and ephrin-B2 in shaping the fate of tip cells.

At the heart of this discovery lie two key molecules: EphB4 and ephrin-B2. Traditionally, EphB4, a transmembrane receptor protein, is linked to endothelial cells of the venous system, while ephrin-B2, a transmembrane ligand protein, dominates in arterial endothelial cells.

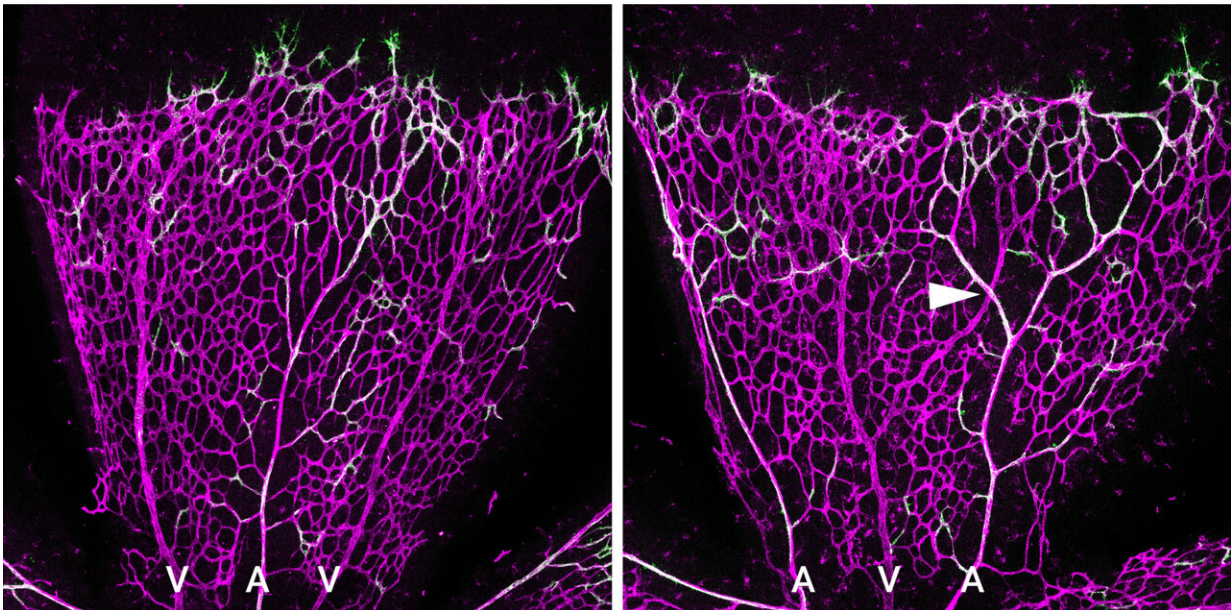
Interestingly, recent studies have revealed their co-expression in tip cells, the frontier navigators of vascular expansion. "Imagine the developing body's vascular system as an expanding network of roads, with arteries acting as life-sustaining highways for oxygen and nutrients. At the network's edge are tip cells, mediating expansion into new territories," explains Mara Pitulescu, senior scientist in the department of Ralf Adams.

Pitulescu's earlier research showed the remarkable ability of tip cells to shift towards the arterial fate. Migrating against [blood flow](#), these cells serve as building blocks that incorporate into arteries, mediating their growth.

In search of the molecular mechanisms underlying the fate specification of tip cells, Pitulescu and Ph.D. student Jonas Stewen analyzed the retinal vasculature of mouse mutants as well as cell culture models where EphB4 or ephrin-B2 signaling is impaired or enhanced. "We found that EphB4 and ephrin-B2 inhibit each other: When EphB4 is low and/or ephrin-B2 is high, the arterial program in endothelial cells is switched

on," Stewen says.

In addition, the researchers found intricate feedback loops from EphB4 and ephrin-B2 activating or inhibiting Notch and [vascular endothelial growth factor](#) (VEGF) signaling, known players involved in arterial fate acquisition. "The tight balance of EphB4 and ephrin-B2 thus orchestrates a well-wired molecular network governing arterial fate specification," Stewen adds.



EphB4 deletion enhances tip cell-mediated artery formation. Credit: MPI for Molecular Biomedicine / Jonas Stewen

Strikingly, disrupting the balance of EphB4 and ephrin-B2 in tip [cells](#) has profound effects on whole vascular plexus development. "We found that deletion of EphB4 causes so-called arteriovenous crossings, a retinal vascular pathology found in [human patients](#) where an artery lies on top

of the vein," Pitulescu says. This can lead to arteriovenous nicking and further retinal vein occlusion—a phenomenon increasingly linked to heightened risks of stroke formation.

"Our findings could shed light on the underlying mechanisms of certain vascular diseases, potentially paving the way for the development of novel therapeutic interventions," explains Ralf Adams, head of the Tissue Morphogenesis department at the Max Planck Institute for Molecular Biomedicine.

More information: Jonas Stewen et al, Eph-ephrin signaling couples endothelial cell sorting and arterial specification, *Nature Communications* (2024). [DOI: 10.1038/s41467-024-46300-0](https://doi.org/10.1038/s41467-024-46300-0)

Provided by Max Planck Society

Citation: A balancing act: EphB4 and ephrin-B2 regulate artery formation (2024, April 3) retrieved 16 May 2024 from <https://phys.org/news/2024-04-ephb4-ephrin-b2-artery-formation.html>

<p>This document is subject to copyright. Apart from any fair dealing for the purpose of private study or research, no part may be reproduced without the written permission. The content is provided for information purposes only.</p>
--