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Single-molecule microscopy in zebrafish embryos

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**SINGLE-MOLECULE MICROSCOPY IN
ZEBRAFISH EMBRYOS**

RADOSŁAW JAKUB GÓRA

Single-Molecule Microscopy in Zebrafish Embryos

Cover and layout by **Radosław Jakub Góra**

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Cover image: an artistic representation of trajectories of single molecules diffusing inside a multicellular organism

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SINGLE-MOLECULE MICROSCOPY IN ZEBRAFISH EMBRYOS

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Dr. J.J. Willemse

Keep the ship outside the spray and surge

Homer, the Odyssey

To everyone who joined me on the ship

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