Genetic and Mechanic Signaling in Organ Development

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(Datum der Disputation)
# Table of Contents

## Preliminary Content
- Acknowledgements .................................................. 1
- About ................................................................................. 2
- Foreword ............................................................................ 3
- Abbreviations ...................................................................... 4
- Publications ....................................................................... 6

## Zusammenfassung
- Einleitung ........................................................................... 7
- Zielsetzung ......................................................................... 9
- Ergebnisse ........................................................................... 10

## Summary ........................................................................ 13

## Chapter 1: Introduction ................................................... 1

1.1 Development ............................................................... 1
  1.1.1 Cell Types .......................................................... 2
  1.1.2 Cell Signaling ...................................................... 3
  1.1.2 WNT ................................................................. 3
  1.1.2 Fibroblast Growth Factor ....................................... 4
  1.1.2 Notch ................................................................. 4
  1.1.3 Morphogenesis .................................................... 5
  1.1.3 Apical Constriction ............................................... 7
1.2 Model Organism and System ......................................... 8
  1.2.1 Developmental Stages ........................................... 9
  1.2.2 The Lateral Line System ......................................... 9
  1.2.3 Posterior Lateral Line Primordium .......................... 11
  1.2.3 Rosette formation ............................................... 12
  1.2.3 Hair cell specification .......................................... 12
  1.2.4 Shroom3 in the pLLP ........................................... 13
  1.2.4 Recent research .................................................. 13
  1.2.4 Current Model .................................................... 14
1.2.5  \textit{shroom3} mutants ............................................. 15
1.3  Open Questions and Motivation ................................. 16

**Chapter 2: Materials and Methods** .................................. 17

2.1  Materials .......................................................... 17
2.1.1  Chemicals ...................................................... 17
2.1.2  Solutions ....................................................... 18
2.1.3  Antibodies ..................................................... 18
2.1.4  Enzymes ....................................................... 19
2.1.5  Molecular Biology Kits ....................................... 19
2.1.6  Buffers ........................................................ 20
2.1.7  Zebrafish lines ............................................... 20
2.1.8  ISH probes .................................................... 21
2.1.9  Morpholinos ................................................... 21
2.1.10  Hardware ...................................................... 21
   Mounting Stamp .................................................. 21
   Spinning Disc Microscopy ....................................... 22
   Workstation ........................................................ 23
2.1.11  Software ...................................................... 23

2.2  Methods ............................................................ 23
2.2.1  Data Strategy and Analysis .................................... 23
   Sample Preparation ............................................... 24
   Imaging ............................................................. 24
   Data handling .................................................... 25
   Image and Data Analysis ....................................... 25
2.2.2  Zebrafish ....................................................... 25
   Husbandry ........................................................ 25
   Handling and rearing ............................................ 25
   Zebrafish fin clips ............................................... 26
   Adult Genotyping ................................................ 26
   Embryo Genotyping .............................................. 26
   Zebrafish Euthanasia ............................................. 26
   Fixation ............................................................ 26
2.2.3  Wet lab .......................................................... 26
   Sample preparation .............................................. 26
   In Situ Hybridization ........................................... 27
   Immuno staining ................................................ 28
   Mounting ........................................................... 28
2.2.4  Dry lab .......................................................... 29
   Image J macros .................................................... 29
   Proliferation Analysis .......................................... 29
   Apical Index ........................................................ 30

2.3  Ground Truth ....................................................... 36
2.3.1 Cluster Analysis ........................................ 36
Design ..................................................... 36
2.3.2 Morphometric analysis .................................. 38
Design ..................................................... 38
2.4 Image Data Sets ........................................... 39

Chapter 3: Results ............................................ 43
3.1 Quantitative imaging and image analysis ................. 43
  3.1.1 A 3D-printed stamp to standardize sample mounting
        and semi-automatize imaging .......................... 43
        Procedure ........................................... 44
        Summary ............................................. 50
  3.1.2 anaLLzr2D - Automated 2D neuromast analysis and
        nuclei count ......................................... 51
        Image Analysis ...................................... 52
        Code Snippets ....................................... 52
        Data Analysis ....................................... 53
  3.1.3 anaLLzr2DT - Automated 2D pLLP migration and
        neuromast deposition analysis through time ........ 54
  3.1.4 anaLLzr3D - Automated 3D single cell segmentation
        and A.I. analysis in the pLLP ....................... 61
        Image Analysis ...................................... 62
        Code Snippets ....................................... 64
        Data Analysis ....................................... 71
  3.1.5 Rosette Detection .................................... 74
3.2 Shroom3 .................................................. 75
  3.2.1 Phenotype description ................................ 76
  3.2.2 Lateral Line Morphometrics .......................... 76
        Dataset ............................................. 76
        Number and Position of Cell Clusters ............... 76
        Cell Count and Area of CCs ......................... 79
        Temperature Rescue ................................ 80
        Summary ............................................. 81
  3.2.3 Proliferation ........................................ 81
        Datasets ............................................ 82
        Lateral Line Mitoses ................................ 82
        Summary ............................................. 84
  3.2.4 Rosette Formation and Cluster Deposition .......... 84
        Dataset ............................................. 84
        Cluster Deposition ................................ 85
        Registration ........................................ 85
        Rosette Detection .................................. 87
        Correlations ........................................ 89
# List of Tables

2.1 Chemicals ......................................................... 17  
2.2 Solutions ....................................................... 18  
2.3 Antibodies ....................................................... 19  
2.4 Enzymes ......................................................... 19  
2.5 Molecular Biology Kits ........................................ 20  
2.6 Buffers ......................................................... 20  
2.7 Zebrafish lines ................................................. 21  
2.8 ISH probes .................................................... 21  
2.9 Morpholinos ..................................................... 21  
2.10 Materials for production of a standardized mounting stamp 21  
2.11 Spinning Disc system components ............................ 22  
2.12 Available lasers ............................................... 22  
2.13 Available objectives .......................................... 23  
2.14 Workstation hardware components .......................... 23  
2.15 Used software .................................................. 23  
2.16 Proteinase K digestion ........................................ 27  
2.17 Cluster Analysis Model ........................................ 37  
2.18 anaLLzr3D Model ............................................... 38  
2.19 Cell Cluster dataset .......................................... 39  
2.20 A.I. dataset ................................................... 39  
2.21 Proliferation dataset ......................................... 40  
2.22 Detection dataset ............................................. 40  
2.23 Atoh1a dataset ................................................. 41  
2.24 Temperature rescue dataset summary ......................... 81  
2.25 Temperature rescue statistics ................................. 81  
2.26 Rockout rescue dataset summary .............................. 93  
2.27 NMII activation dataset summary .............................. 93  
3.1 Limitations of traditional zebrafish mounting techniques 45  
3.2 Blurring parameter for nuclei count ........................... 54  
3.3 3-D Ground Truth image scaling ................................ 63  
3.4 CNN training data .............................................. 75  
3.5 Temperature rescue dataset summary .......................... 81  
3.6 Temperature rescue statistics .................................. 81  
3.7 Rockout rescue dataset summary ................................ 93  
3.8 NMII activation dataset summary ............................... 93
**List of Figures**

1.1 Waddington’s Classical Epigenetic Landscape ............... 2
1.2 Form and function feedback loops .......................... 6
1.3 Modes of constriction ...................................... 8
1.4 Model organism Danio rerio ................................. 9
1.5 Zebrafish embryonic development ............................ 10
1.6 The lateral line system ...................................... 11
1.7 Neuromast deposition and pattern ............................ 12
1.8 Shroom3 functional domains and mode of operation ........ 13
1.9 Shroom3 in rosette formation ............................... 14
1.10 Shroom3 current model ..................................... 15

2.1 Tracking of mitotic events ................................. 29
2.2 A.I. indeces in the literature ............................... 30
2.3 Different ways to quantify the apical index ................ 31
2.4 Schematic anisotropic AC .................................. 32
2.5 Scheme of ellipsoid measures ............................... 33
2.6 A.I.\textsubscript{Major} / A.I.\textsubscript{Minor} single cell measurements ... 34
2.7 A.I. indices by Harding et al. ............................... 34
2.8 A.I.\textsubscript{Major} / A.I.\textsubscript{Minor} over Major\textsubscript{Angle} ........ 36

3.1 Stamp and micro-well properties ............................ 46
3.2 Mounting results ............................................ 47
3.3 Stamping procedure ......................................... 50
3.4 anaLLzR2D opening dialog .................................. 51
3.5 Registration of 2D data ...................................... 52
3.6 Relative difference of maxima counts ....................... 54
3.7 anaLLzR2DT opening dialog ................................ 55
3.8 anaLLzR3D opening dialog .................................. 62
3.9 Registration of 3D data ...................................... 63
3.10 Image data pLLP segmentation .............................. 64
3.11 anaLLzr3D - Graphical comparison of tested thresholds ... 65
3.12 Relative difference of segment counts .................... 71
Disclosure

Unless otherwise noted, all figures in this manuscript are an expression of my own creative achievement. Similarities between molecule, protein and signalling models exist and are based on well-known textbooks such as Lodish et al., Molecular Cell Biology, 5th Edt. and Scott F. Gilbert et al., Developmental Biology 11th Edt.
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About

Beside a print version, there is also an electronic version\(^1\) of this thesis. In addition, data analysis scripts, figures and tables are stored in a dedicated git repository\(^2\). If you need to have access please contact me.

Print version

- Including Acknowledgements and Preface
- Lateral line primordium *flipbook*
  - right pages (odd numbers) show the wildtype
  - left pages (even numbers) show the mutant

Electronic version

- Download pdf
- Change display settings
- Search contents

\(^1\)https://kleinhansda.github.io/phd_thesis/
\(^2\)https://github.com/KleinhansDa/phd_thesis
Foreword

“The most exciting time in the history of developmental biology is right now. Fueled both by new technologies and by new thought from other fields, we are exploding old notions and opening fantastic new horizons in embryology. […] Next, let’s discuss why developmental biology — both normal and pathological — holds such enduring fascination. I see two intertwined explanations. Obviously, it’s the ultimate personal creation story, telling each of us both where we came from and how we were constructed. Less obvious, but perhaps more tantalizing for us scientists, is the sheer complexity of the process. A single cell with a single genome can somehow create trillions of cells in hundreds of radically different types, and those cells can organize themselves into a specific form. The scale of this self-organization process is mind boggling, surely the most amazing of emergent properties.”
- John B. Wallingford, 2019

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3“We Are All Developmental Biologists,” Developmental Cell, Vol.50, Issue 2, Jul 22, 2019
## Abbreviations

<table>
<thead>
<tr>
<th>Index</th>
<th>Abbreviation</th>
<th>Elaboration</th>
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<tbody>
<tr>
<td>0-9</td>
<td>2-/3-D</td>
<td>Two or Three Dimensional</td>
</tr>
<tr>
<td>A</td>
<td>A.I.</td>
<td>apical index</td>
</tr>
<tr>
<td></td>
<td>AB</td>
<td>Antibody</td>
</tr>
<tr>
<td>C</td>
<td>CC</td>
<td>Cell Cluster</td>
</tr>
<tr>
<td>CNN</td>
<td>Convolutional Neural Network</td>
<td></td>
</tr>
<tr>
<td>CI</td>
<td>confidence interval</td>
<td></td>
</tr>
<tr>
<td>D</td>
<td>dpf</td>
<td>days post fertilization</td>
</tr>
<tr>
<td>DSH</td>
<td>Dishevelled</td>
<td></td>
</tr>
<tr>
<td>E</td>
<td>ECDF</td>
<td>Empirical Cumulative Distribution Function</td>
</tr>
<tr>
<td></td>
<td>EDF</td>
<td>Extended Depth of Field</td>
</tr>
<tr>
<td>F</td>
<td>Fgf</td>
<td>Fibroblast Growth Factor</td>
</tr>
<tr>
<td></td>
<td>Fgfr1</td>
<td>Fgf receptor 1</td>
</tr>
<tr>
<td>FOV</td>
<td>Field of View</td>
<td></td>
</tr>
<tr>
<td>FRZ</td>
<td>Frizzled</td>
<td></td>
</tr>
<tr>
<td>H</td>
<td>hpf</td>
<td>hours post fertilization</td>
</tr>
<tr>
<td>I</td>
<td>IJ</td>
<td>ImageJ</td>
</tr>
<tr>
<td>ISH</td>
<td>In Situ Hybridization</td>
<td></td>
</tr>
<tr>
<td>K</td>
<td>KDE</td>
<td>Kernel Density Estimation</td>
</tr>
<tr>
<td></td>
<td>k.s.</td>
<td>Kolmogorov-Smirnov</td>
</tr>
<tr>
<td>L</td>
<td>l.e.</td>
<td>leading edge</td>
</tr>
<tr>
<td>LL</td>
<td>Lateral Line</td>
<td></td>
</tr>
<tr>
<td>LOESS</td>
<td>Locally Weighted Scatterplot Smoothing</td>
<td></td>
</tr>
<tr>
<td>LSFM</td>
<td>Light sheet fluorescence microscope</td>
<td></td>
</tr>
<tr>
<td>LUT</td>
<td>Lookup Table</td>
<td></td>
</tr>
<tr>
<td>LMPA</td>
<td>low melting point agarose</td>
<td></td>
</tr>
<tr>
<td>M</td>
<td>MaxIP</td>
<td>Maximum Intensity Projection</td>
</tr>
<tr>
<td></td>
<td>MO</td>
<td>Morpholino</td>
</tr>
<tr>
<td></td>
<td>mQ</td>
<td>milli-Q water</td>
</tr>
<tr>
<td>N</td>
<td>N.A.</td>
<td>Numerical Aperture3</td>
</tr>
<tr>
<td>NGS</td>
<td>Normal Goat Serum</td>
<td></td>
</tr>
<tr>
<td>NICD</td>
<td>Notch intracellular domain</td>
<td></td>
</tr>
<tr>
<td>NM</td>
<td>Neuromast</td>
<td></td>
</tr>
<tr>
<td>NMII</td>
<td>Non muscle myosin II</td>
<td></td>
</tr>
<tr>
<td>O</td>
<td>o.n.</td>
<td>over night</td>
</tr>
<tr>
<td>P</td>
<td>p.Hist</td>
<td>phospho-Histone</td>
</tr>
<tr>
<td>PCR</td>
<td>Polymerase chain reaction</td>
<td></td>
</tr>
<tr>
<td>PFA</td>
<td>Paraformaldehyde</td>
<td></td>
</tr>
<tr>
<td>pLLP</td>
<td>Posterior Lateral Line Primordium</td>
<td></td>
</tr>
<tr>
<td>R</td>
<td>Rock</td>
<td>Rho-Kinase</td>
</tr>
<tr>
<td>ROI</td>
<td>Region of Interest</td>
<td></td>
</tr>
<tr>
<td>S</td>
<td>SBD</td>
<td>Shroom binding domain</td>
</tr>
<tr>
<td>SEM</td>
<td>Scanning Electron Microscopy</td>
<td></td>
</tr>
<tr>
<td>SNR</td>
<td>Signal to Noise Ratio</td>
<td></td>
</tr>
<tr>
<td>T</td>
<td>TALEN</td>
<td>Transcription activator-like effector nuclease</td>
</tr>
<tr>
<td>TF</td>
<td>Transcription Factor</td>
<td></td>
</tr>
</tbody>
</table>
Publications

This thesis contains material from the following papers. The rights have been granted by the publisher to include the material in this dissertation. Some passages have been quoted verbatim for the scientific accuracy from the following sources:

Zusammenfassung

Einleitung

neben ihren einzelnen Zytoskelettelementen auf der suprazellulären Ebene über Aktin durch Adherens junctions verbunden. Wenn sich das Gewebe an einer Stelle verformt, werden dadurch auch andere, weiter entfernte Zellen verformt. Für diese Thesis war ich daran interessiert, besser zu verstehen, wie die zelluläre Form und Geometrie die nachfolgende Zell- und Organentwicklung beeinflusst. Was passiert, wenn eine Zelle nicht in eine bestimmte Form übergehen kann? Wie wirkt sich das auf die Gewebestruktur aus? Wie wirkt es sich auf die weitere Entwicklung aus?


**Zielsetzung**


Um die Auswirkungen von Shroom3 auf die Entwicklung des Seitenlinienorgan zu untersuchen, wurde eine Mutanten-Linie generiert und mit verschiedenen transgenen Linien gekreuzt, die fluoreszenzmarkierte Proteine
exprimieren, die sich an Organellen wie der Plasmamembran oder dem Zellkern lokalisieren. Anschließend wurde die Mutante mit ihren fluoreszenten Markierungen unter verschiedenen Bedingungen mikroskopisch abgebildet, um verschiedene zellmorphometrische Merkmale zu quantifizieren und zu analysieren.

Obwohl der Zebrafisch ein beliebter Modellorganismus ist und sich hervorragend für die Entwicklungsbio logie und fortgeschrittene Mikroskopie eignet, gab es bisher keine Methoden, die eine standardisierte und stärker automatisierte Pipeline der Datenerfassung und -verarbeitung ermöglichen würden. Um die morphogenetischen Prozesse, an denen Shroom3 beteiligt ist, genau quantifizieren zu können, wurde ein neues Toolset entwickelt, das meine Arbeit deutlich effizienter gemacht hat. Das Toolset besteht aus (1) einer neuen Methode zur Probennrangement bei der von einem 3D gedruckten Stempel ein Abdruck in einem 3D-Agarosegel hergestellt wird, das die Anzahl der Embryonen die auf einmal montiert und abgebildet werden kön nen deutlich erhöht und den Bildgebungsprozess erheblich beschleunigt. (2) Für die anschließende Bildanalyse wurden vier Programme entwickelt, die die Anzahl der Embryonen die auf einmal montiert und abgebildet werden kön nen deutlich erhöht und den Bildgebungsprozess erheblich beschleunigt. (3) Für die anschließende Bildanalyse wurden vier Programme entwickelt, die die Anzahl der Embryonen die auf einmal montiert und abgebildet werden kön nen deutlich erhöht und den Bildgebungsprozess erheblich beschleunigt. (4) Für die anschließende Bildanalyse wurden vier Programme entwickelt, die die Anzahl der Embryonen die auf einmal montiert und abgebildet werden kön nen deutlich erhöht und den Bildgebungsprozess erheblich beschleunigt.

Ergebnisse

Nach ersten Ergebnissen zur Rolle von Shroom3 im hinteren Seitenlinien-Primordium des Zebrafisches durch Morpholino-injizierte Knockdowns hat eine ehemalige Kollegin im Labor eine Mutante generiert, um die Rolle von Shroom3 während der Morphogenese zu bestätigen und weiter zu untersuchen. Eine acht basen-paar Deletion in der SD2-Domäne wurde isoliert und als stabile Linie erhalten. Diese Mutation führt zu einem vorzeitigen


Für die Probenmontage habe ich eine neue Methode entwickelt, die auf einem 3D-Agarosegel basiert und (1) die Anzahl der Embryonen, die auf einmal montiert und abgebildet werden können, deutlich erhöht (2) durch die optimierte Orientierung weniger häufig belichtet werden und somit schonender für die Embryonen ist und (3) den Bildgebungsprozess durch eine regelmäßige und gleichbleibende Anordnung automatisierbar und somit deutlich beschleunigt. Darüber hinaus habe ich für die anschließende Quantifizierung eine Reihe von Marko-Programmen für die 2D- und 3D-Analyse entwickelt, die den Prozess automatisieren und damit die Ergebnisse reproduzierbarer und die Analyse deutlich effizienter machen. Meine Ergebnisse und meine Methodik zeigen, wie wichtig die Morphologie bei der Steuerung von Entwicklungsprozessen ist und wie schon kleine morphologische Veränderungen auf zellulärer Ebene die weitere Entwicklung erheblich beeinflussen können. Meine Arbeit zeigt auch, wie leistungsfähig die moderne Genetik, Mikroskopie und Bildanalyse sind und wie vielfältig sie in Bezug
Summary

In order to form an organ, cells need to take up specialized functions and tasks. Cellular specialization is guided by an interplay of chemical signals and physical forces, where one influences the other. One aspect in cellular identity is its shape, which e.g. defines how susceptible the cell may be to intercellular signaling or in which section of the cell cycle it is and therefore can tell us about its current state. Shape changes are introduced by motor proteins that are controlled and activated in a locally confined manner. For my thesis, I was interested to understand better how cellular shape and geometry impacts downstream cell and organ development. What happens if a cell cant transition to a specific shape? How does it affect tissue structure? How does it affect further development?

One regulator of motor proteins like non-muscle myosin is Shroom3, which recently has been shown to be expressed and involved in the development of the zebrafish lateral line organ (1). Development of the lateral line occurs through a migrating cluster of initially about 150 cells, the posterior lateral line primordium (pLLP), which migrates from the anterior (head) to the posterior (tail) while depositing cell clusters in a regular pattern. Literature on development of the lateral line suggests that in order for a cell cluster to be deposited from the pLLP, rosette formation is a key requirement. Therefore our expectation from the shroom3 mutant was that the number of clusters deposited was significantly reduced. To our surprise, when we first inspected the end of migration lateral line phenotype we found many individuals with a significant increase in cell clusters deposited. This made us re-think the role of Shroom3 during rosette assembly and the processes its involved in.

To study the effects of Shroom3 on lateral line development, a mutant line was generated and crossed with various transgenic lines which express fluorescently labeled proteins that locate to organelles such as the plasma-membrane or the nucleus. Following, the mutant with its fluorescent labels was microscopically imaged under different conditions to quantify and analyze various cell-morphometric features. Even though the zebrafish is a popular model organism and its perfectly suited for developmental biology
and advanced microscopy, there were no methods that would allow for a standardized and more automated pipeline of data acquisition and processing. Therefore, in order to accurately quantify the morphogenetic processes Shroom3 is involved in, I developed a new toolset that significantly improved and facilitated my research. The toolset consists of (1) a new sample mounting method that is based on a 3D agarose gel that increases the number of embryos that can be mounted and imaged at once and speeds up the imaging process significantly (2) for subsequent image analysis I developed four programs that automate the process and therefore make the results much more reproducible and the analysis much more efficient. The first program is used for end of migration analyses, to deduce the pattern, count and size of Lateral Line cell clusters. The second is used not for end of migration, but for migration analyses (on timelapse recordings). Besides this it also prepares the images for more advanced downstream migration analyses and allows to analyse fluorescence signal on a second channel. The third program is used to analyse the pLLP only at high spatial resolution and to deduce the cell count, 3D cell morphometrics (like the volume) and cell orientation. The fourth program finally is used downstream of the second and third program and is capable of detecting and comparing them with the look of wildtype rosettes.

Here I show that in absence of Shroom3 rosette formation in the migrating pLLP is destabilized leading to facilitated cell cluster deposition and I show how this might be related to traction forces due to a possible interdependence of pLLP acceleration and speed of migration. Furthermore I show that apical constriction and rosette formation is not blocked in Shroom3 deficient embryos, but that larger rosettes are fragmented into many smaller ones. Finally, I give an outlook on how the absence of Shroom3 and hence the absense of morphological changes may deregulate gene transcription by elevating the levels Atoh1a, a transcription factor necessary for hair cell development.

My results and methodology demonstrate the importance of morphology in guiding developmental processes and how rather small morphological changes on the cellular level can impact further development significantly. My work also shows how powerful modern genetics, imaging and image analysis are and how diverse they are in terms of range of questions they are capable of answering. The methods and tools I developed prepare the ground for at least three quarters of the analyses I carried out and together with the documentation and data I provide, they are highly reproducible. In that regard I am especially happy that one of my developments, an improved sample preparation method, is already used by many different labs all over the world helping them to make their results more reproducible.
Chapter 1

Introduction

1.1 Development

**Developmental Biology** studies the continuous process of cells adapting to their ever-changing environment from the fertilized egg to a full organism. For an organism to develop, cells need to proliferate (multiply) and become different from each other. To achieve this, *genes* are utilized that are expressed differently at certain times during development and types of cells. Those proteins again are used for geometrical construction or facilitation of chemical reactions. To illustrate this, the cell can be thought of as a marble rolling down a furrowed landscape (figure 1.1). Physically a cell represents an open system, which is defined as a unit system able for external interactions. Such a system is therefore not self-dependent and self-sustained, but its current conformation is determined by external interactions. In this landscape a hill would be a high energy-, a valley a low energy state. The path the marble will take is determined by the furrows in the landscape, since it would always prefer a valley. However, this landscape is not a *static* structure but one that changes at every instant of time. As cells differentiate they acquire a certain identity - controlled by the genes they express and the proteins they produce - that again influences cell ‘behavior’ and its phenotype which includes cell division, migration and cell shape changes. This allows cells to assemble into tissues, which themselves assemble into organs. For developmental biology, which as a scientific discipline originates from embryology, the central interest is to understand how initially inanimate matter (like single atoms and molecules) is able to organize itself to such complex structures we see in living matter at different levels of organismal hierarchy while ensuring a robust developmental plan. Using the marble analogy, studying developmental biology can be thought of as tracking multiple cells as they roll down the valley while observing
and testing how interactions between them might change their fate. The basic questions arising from this interest are

- How do tissues arise from a population of cells?
- How do organs form from tissues?
- Why do organs form at their particular location?
- How do migrating cells know whether they reached their destination?
- How is growth controlled and how do body axes form?

### 1.1.1 Cell Types

In animals there are two basic types of cells.

1. Epithelial cells, which can form strong adhesions between each other and thereby are able to exert forces upon each other to achieve complex architectures.
2. Mesenchymal cells, which do not adhere as strong with each other and are more independent.

This however describes only the extremes on a continuous scale. A cell is not a binary system but can show characteristics of both extremities, *e.g.* during Epithelial to Mesenchymal transition (EMT), a bidirectional process whereby epithelial cells can gain migratory and invasive properties and *vice versa*.

A cell's identity on this continuum is determined by

- its gene expression profile, which reflects its repertoire of molecular machinery (proteins) and therefore determines its competence to react to internal and external cues.
- its micro-environment, which has a *physical* (forces, energy) and a *chemical* (signaling molecules, diluents) dimension. A change in the
latter usually brings about a reaction in the cell that becomes evident both in expression of genes and morphology.

- its shape and incorporation into a tissue, which may modulate how a cell reacts to its micro-environment (e.g. certain regions of the cell can be more or less exposed, or it can be more or less tightly packed, tuning its susceptibility to forces and signals).

With rising numbers of specialized cells assembling into tissue, a shape and body axes begin to emerge that for the earliest developmental stages is highly similar across certain phyla and only begins to diversify at later developmental stages, which reflects our evolutionary ancestry.

1.1.2 Cell Signaling

To communicate with each other, cells have developed a variety of intercellular communication systems and a complex network of intracellular signal transduction pathways. Some information transfer depends on direct cell-to-cell contact, others rely on freely diffusible ligands that can be sensed by other cells. Each signaling pathway consists of a ligand – receptor pair that determines their main function. In the following, three pathways are introduced that play major roles in embryonic development.

WNT

The word ‘WNT’ is a compound word of Wingless and Int-1, both of which are important genes during development of Drosophila melanogaster (commonly known as fruit fly), where WNT signaling was first studied. WNT is evolutionary conserved with 15 different receptors and co-receptors and plays a major role during embryonic axis formation, body segmentation, organogenesis and stem cell proliferation. Aberrant WNT signaling is involved in diseases like colon cancer, melanoma and neurodegenerative diseases (2).

Beside this, WNT signaling is sub-divided in a canonical (β-catenin\(^1\) dependent) and a non-canonical (β-catenin independent) branch. In canonical signaling, WNT ligand binds, together with co-receptor Lipoprotein Receptor-related Protein 6 (LRP6), to the receptor Frizzled (Frz) – jointly activating protein Dishevelled (Dsh). Dsh again inhibits a protein complex usually degrading β-catenin, leading to the stabilization of β-catenin in the cytoplasm and the nucleus. Within the nucleus, β-catenin forms a complex with LEF/TCF to activate specific target genes (2).

\(^{1}\)catenins are regulators of cell-cell adhesion and gene transcription
CHAPTER 1. INTRODUCTION

Fibroblast Growth Factor

Key roles of Fibroblast growth factor (FGF) signaling is mesoderm\(^2\) patterning in the early embryo, regulation of angiogenesis and wound repair. On a cellular level it is also an important regulator of proliferation and differentiation. The mammalian FGF family comprises 18 ligands and four highly conserved transmembrane tyrosine receptors named FGF receptor 1-4 (FGFR1-4). Aberrant FGF signaling is e.g. associated with tumor growth (3).

Upon ligand binding FGFR dimerizes and undergoes a conformational shift activating the intracellular kinase\(^3\) domain. Subsequent trans-phosphorylation of tyrosine kinase domains serve as docking sites for adaptor proteins. Activated FGFR then phosphorylates FGFR substrate 2 (FRS2), recruiting adaptor protein Son of Sevenless (SoS) and Growth factor Receptor bound 2 (GRb2) to set on a cascade of at least three possible kinase dependent signal transduction pathways eventually leading to activation of target genes involved in the regulation of e.g. proliferation, autophagy and EMT (3).

Notch

In neurobiology the term lateral inhibition describes the process of an excited neuron reducing the activity of its neighbors. The same principle however can be found in other types of cells too where one cell signals its neighbor cell(s) not to acquire a certain fate. For initially equivalent cells to diversify and acquire distinct identities there needs to be a break in symmetry. Often this is accomplished by competition about Notch signaling sources where whichever cell expresses a trait first is able to suppress the same in its neighbors. Once this hierarchy is established, it is maintained in a positive feedback manner (4, 5).

Notch signaling gives cells the ability to self-organize and controls various developmental and homeostatic processes that involve patterning, such as sensory hair cell (HC) formation, branched arterial networks or organ morphogenesis. Notch signaling consists of four components: (1) The extracellular domain of the membrane bound Notch receptor (2) Notch ligands (3) the Notch intracellular domain (NICD) (4) and the \(\gamma\)-Secretase. Upon ligand activation of Notch through Delta, NICD is cleaved and released through \(\gamma\)-Secretase. Subsequently NICD enters the nucleus and together with DNA-binding proteins and co-factors initiates expression of target genes. In contrast to other signaling pathways

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\(^2\)one of the earliest differentiating layer of cells. Cells of the mesoderm will e.g. form the musculature.

\(^3\)enzymes that transfer energy to specific substrates
1. there are no intermediates between membrane signaling and nucleus and therefore no amplification or dampening of the signal occurs
2. signaling requires direct contact between cells, which makes Notch signaling particularly biased by features of cellular morphology and tissue organization (4).

Assuming a homogeneous receptor concentration in a cell membrane and since Notch signaling occurs at sites where cells are in contact, the signal generated is proportional to the contact area. Additionally, the strength of the signal increases further where cells are tightly opposed (4, 6–8).

1.1.3 Morphogenesis

Morphogenesis, from the Greek morphê (shape) and genesis (creation)

“Creation of the shape”

For objects that fulfill a purpose, their form is an expression of their function4. Hence, analyzing the shape of an object can give information about its function. It is therefore an important feature in different scientific disciplines.

Breaking symmetry of daughter cells often results in one of the daughter cells adopting a different fate, which eventually results in diversification of shape (e.g. muscle cells, neural cells, etc.). In order to form a tissue or an organ with a variety of specialized cells, it is important for the single cell to have information about where it is located, what its neighbors are doing, how densely it is packed and what the chemical composition of its surrounding is. This is accomplished by being in constant feedback with its neighbor cells and sensation of its environment. After the information is processed, the cell reacts by adjusting the levels of proteins expressed to undergo proliferation or develop specialized structures like cilia or axons.

“For an isolated cell, cell shape reflects a balance between cortical tension and intracellular pressure.” - Y.Pan et al.(10)

For a cell, its present shape impacts its further specification by defining its ability in perceiving specific signals and the magnitude of signals received and transmitted (8). Shape therefore sets the general framework for cell-cell interaction and follows a

\[
\text{molecular} \rightarrow \text{cellular} \rightarrow \text{tissue}
\]

4 Even though the expression Form follows function (9) is usually found in design and architecture, it formulates the general idea that any objects form is (or in design should) be shaped by the requirements to it.
scale hierarchy, where each scale’s output again feeds back to the others (figure 1.2). E.g. it has been shown that simple changes in cell geometry affect fundamental processes such as cell growth, death, direction of cell divisions and extracellular vesicle cargo (11–22).

Figure 1.2: Form and function feedback loops. On a cellular scale expression of genes may cause a cell to constrict, which again feeds back to the molecular scale (e.g. force transmission) and the tissue scale (e.g. evagination or constriction). All of which standing in a continuous feedback.

Cell shape changes may occur in two different ways...

- active: Each cell has a dynamic skeleton that is composed of a complex matrix of interconnected proteins - the cytoskeleton. It is composed by three basic classes of filaments that have different physical properties. Microtubules (the most rigid), Intermediate and microfilaments (the softest). Motor-proteins like Myosin may adhere to actin micro-filaments and contract them, thereby causing e.g. a narrowing of the cell diameter.
- passive: In a tissue cells are, besides their individual cytoskeletal elements, connected at the supracellular level via actin through Adherens junctions. This way, if the tissue becomes deformed at one place, other more distant cells will get deformed as well (23).
1.1. DEVELOPMENT

Apical Constriction

Apical constriction (AC) is a cell morphogenetic process manifesting by an active narrowing of the apical surface, making an epithelial cell appear bottle or wedge shaped instead of cuboidal. It is usually coordinated by multiple cells within an epithelial layer that raise forces necessary to deform a tissue.

A defining feature of epithelial cells is that they have a basal- (bottom) to apical (top) polarity. At the basal side, the cell is in contact with a specialized extracellular matrix (ECM, a thin sheet-like structure) called the basement membrane, apically the cell forms tight connections to its neighboring cells – a region called the apical junctional complex (AJC). The AJC encompasses three types of junctions: Adherent junctions (AJ) or zonula adherens (ZA), tight junctions (TJ) and desmosomes. Around the ZA dense cables of actomyosin are located that, analog to muscle sarcomeres, are able to contract upon activation of RHO-associated protein kinase (Rock) and Rock-mediated phosphorylation of the motor protein non-muscle myosin II (NMII) \((24)\). The exact mechanism of apical constriction may differ in different organism and organs. E.g. it has been shown by Martin et al. \((25)\) that in *Drosophila* cells are radially organized and RhoA and ROCK have a medioapical focus with RhoA also present at junctions, while chick cells exhibit a planar cell polarity and RhoGEF and ROCK are localized to junctions.

Developmental processes that involve AC are...

- Tissue folding and tube formation
- Single cell ingression and EMT
- Gastrulation
- Healing and sealing of embryonic tissue

Epithelial rosettes are radially organized cell clusters within an epithelial tissue whose vertices interface a common center similar to a garlic bud or a pie cut into pieces along its center. While the mechanisms of cytoskeletal rearrangements seem to be well conserved, the signals that lead to rosette formation are less well understood and more diverse. At least two architectural distinct types of rosettes exist, depending on tissue polarization. First, in a planar polarized tissue, several cells converge at a central apico-basal line where cells go from a square to a more triangular shape to form a cylindrical structure (like a pie cut into \(\geq 4\)) (figure 1.3B). Such rosettes are usually observed during tissue elongation and are rather short-lived. In a second scenario, cells converge to a central apical point through AC (figure 1.3A). This type of rosette is more long-lived and usually does not resolve but already represents a morphologically pre-mature state of the organ to be formed \((26)\).
CHAPTER 1. INTRODUCTION

1.2 Model Organism and System

To study biological phenomena, biologists use a variety of non-human model organisms. While each model organism has its advantages and disadvantages, the choice for a model depends on the scientific question. To study embryonic development the fresh water fish *Danio rerio* (also known as eng: *zebrafish* or ger: *Zebrabärbling* (figure 1.4) has become an important model organism over the recent years. *D. rerio* is a diploid organism with a fully sequenced genome (of the human genes 71.4% have at least one *D. rerio* ortholog, 47% have a one-to-one ortholog (27)). It has a relatively short alternation of generations (12-16 weeks), a regularly large number of embryos (100 / week & female) and is relatively undemanding in terms of space for breeding. Furthermore, it offers well-established methods for mutagenesis, screening, and generation of transgenic lines. Since its embryos are naturally transparent and develop externally, it is an ideal system for microscopic examination using molecular dyes and tags to visualize inter and intra cellular components even deep within the tissue (e.g. cell nuclei or cell membrane fluorescent tags). Together with advances in imaging techniques, this also allows for high-throughput, high-resolution, long-term in vivo imaging. Especially the expression of fluorescent proteins in a tissue- or organ-specific manner in transparent embryos offers enormous possibilities to address interesting and long-standing open questions.

In nature, zebrafish can be found in the shallow waters of the Indian and Pakistan Ganges inflows. It exhibits an oval body shape and can reach a length of up to 5 cm in adulthood. While females are usually more silverish, males have a brownish back and a yellow-whit belly. Laterally it exhibits its name-giving dark-blue iridescent stripes with silver in between.

![Figure 1.3: Modes of constriction. A Cells converge to a central apical point B Cells converge at a central apico-basal line.](image-url)
1.2. MODEL ORGANISM AND SYSTEM

Figure 1.4: Model organism *Danio rerio* (aquarell by Christine Molenda)

1.2.1 Developmental Stages

A single female may lay 100 eggs per week. Each zygote\(^5\) then undergoes the first zygotic cell cycle (at ~ 0.75 h). The following two to seven cell cycles (period: *Cleavage*, at ~ 2.25 h) occur directed and synchronous every ~15 min. Cells in this stage are called blastomeres, are incompletely separated from the yolk and remain interconnected by cytoplasmic bridges. The *Blastula* period (up to 5.25 h) is determined by increasingly asynchronous division, flattening of the blastoderm via cell intercalation and lengthening of the cell cycle. This period is also marked by the onset of *epiboly*, the period when cells in late blastula start to dome while a monolayer of the dome circumference begins to wrap around the yolk (28).

After Blastula and until 10 h the *Gastrula* period takes place, followed by the *Segmentation* period (until 24 h). Both of which are depicted in more detail in figure 1.5. At even later stages the embryo starts to elongate posteriorly, grow and develop organs until it first active muscle are present and it starts to swim (figure 1.5).

1.2.2 The Lateral Line System

The lateral line (LL) system is a mechano-sensory organ that is common to all aquatic vertebrates and evolutionary remnants could even be found in mice (29). It enables the animal to sense water movements and therefore to orient itself, and to detect prey and predators. Fully developed, the lateral line system is comprised of hundreds of neuromasts positioned in an orderly pattern all over the animal’s body (figure 1.6A). Its functional subunits are the neuromasts (NM) (figure 1.6B-C) that, when fully developed, consist of hair-, support- and mantle cells. To sense water movements, each NM projects kino- and stereo-cilia out of the skin that, upon water-induced deflection, generate action potentials that are transduced *via* afferent fibers (30, 31).

Each NM is first deposited as a premature cluster of about 30 cells from a migrating cell-aggregate called the posterior lateral line primordium (pLLP) (figure 1.7A). The pLLP delaminates from the pLL placode, caudal to the

\(^{5}\)first diploid cell after fertilization
**Figure 1.5:** Zebrafish embryonic development. Microscopic images are from a time-lapse where 24 embryos were imaged simultaneously in brightfield and at 488 nm Z-Stacks. Representation shows contrast-enhanced EDFs. Bottom row (30-48 hpf) stages are handmade drawings from live embryos made during the first week of my PhD.
1.2. MODEL ORGANISM AND SYSTEM

Figure 1.6: The lateral line system. A modified after (Ghysen et al., 2012) and A.Bergs, 2016 (student presentation at AK Lecaudey). Development of the lateral line system at embryonic, larval and juvenile stage. B Schematic showing a crossection and organization of a single neuromast. C-C’ SEM images of a single, pre-mature (3 dpf) neuromast.

otic vesicle (figure 1.7B) at around 20 hours post fertilization (hpf) as a group of ~100 cells. After formation of a mesenchymal-like leading region, it starts migrating along a chemokine gradient positioned at the horizontal myoseptum to the tip of the tail (30, 31). To ensure the development of a functional organ, several fundamental biological processes like cell migration, morphogenesis, proliferation and cell polarization need to be integrated into the pLLP. An important breakthrough in LL research has been the development of a transgenic line expressing a membrane-tethered GFP fusion protein (lyn-GFP) that is expressed under the LL specific promotor of cldnb (claudin b6) (32), which allows for a much more detailed view and to observe lateral line development in vivo. An example of the fluorescence signal visible at ~60 hpf can be seen in figure 1.7B.

1.2.3 Posterior Lateral Line Primordium

The pLLP is about 100-150 µm in length (depending on deposition cycle) over which it exhibits a diverse surface topology and cellular morphology. Previous research found that the caudal (also posterior), more mesenchymal cells, are leading the path of migration, while the cranial (also anterior), more epithelial cells, are trailing. Towards the leading region the cells are more flat, towards the trailing region the cells become more columnar and increasingly radially organized into formations called epithelial rosettes. During migration the pLLP typically contains 2-3 rosettes (~25-30 cells each), while the most trailing one will eventually be deposited to further mature to a functional NM (30, 33). Every deposition comes with a loss

6construct name: Tg(-8.0cldnb:lynGFP); ZFIN ID: ZDB-TGCONSTRCT-070117-15
of cells in the pLLP, but this loss is partially compensated by proliferation during migration. While one study concludes a general spatial heterogeneity in distribution of proliferative cells (34), another one suggests a higher proliferative rate specifically near the leading region (35).

**Rosette formation**

The onset of morphological and functional changes is determined by signaling of FGF, which is mostly active in the trailing region. Causal for expression of FGF is a signaling center of WNT in the leading region, which promotes expression of FGF ligands Fgf-3 and -10 (36). Those ligands then diffuse to the trailing domain where binding through FGF receptor 1 (Fgfr1) triggers a signaling cascade through which the cells become more columnar, apically constricted and eventually re-organize into epithelial rosettes (35, 37, 38). Concurrently, cells of the WNT signaling center themselves are not competent to FGF signaling, which is achieved through expression of Sef, an intracellular antagonist of Fgfr1 signaling (39).

It was shown that rosette formation in the lateral line primordium (LLP) is an important morphological feature for a lumen to form on top of the rosette which, filled with FGF, acts as a locally enriched source of FGF signaling (40).

**Hair cell specification**

Downstream of FGF lies the expression of the transcription factor (TF) Atoh1a, which gives cells the potential to become sensory HCs (35).

The current model suggests that:

1. FGF initiates expression of *atoh1a* and *deltaA*, where DeltaA activates Notch in neighboring cells to inhibit expression of *atoh1a* in those.
2. Atoh1a in turn suppresses competence for FGF and initiates expression of atoh1b and deltaD which act synergistically with DeltaA in the center cell of the forming rosette.

3. While the latter acts synergistic with DeltaA, Atoh1b in turn drives expression of atoh1a (41).

4. Once a prospective HC is specified it will itself become a source for Fgf-10. By this process adjacent cells are laterally inhibited and determined as support cells (figure 1.6B), still capable of receiving FGF signals via Fgfr1.

Just before the most trailing rosette is deposited, its prospective HC will undergo a final division to form a doublet of sensory HCs (42, 43).

### 1.2.4 Shroom3 in the pLLP

The Shroom protein family is mostly conserved through animal evolution (figure 3) and involved in contraction of the actomyosin network (e.g. during AC), which has been confirmed in several studies investigating e.g. epithelial planar remodeling, neural tube morphogenesis and Xenopus bottle cells (44–49).

#### Recent research

Shroom proteins have three characteristic domains (1) a PDZ domain close to the N-terminus that interact with other proteins with PDZ-binding domains and (2) two Apx/Shroom domains (ASD-1 and -2), the latter being close to the proteins C-terminus. The Shroom domains may interact with proteins containing a Shroom binding domain (SBD) such as Rock which promotes phosphorylation of NMII (figure 1.8) (50).

![Figure 1.8: Shroom3 functional domains and mode of operation. Sequence of events numbered from 1-4. Approximate scale jumps indicated at arrows.](image-url)
CHAPTER 1. INTRODUCTION

Figure 1.9: Shroom3 in rosette formation (adapted from Ernst et al., 2012) A composite MaxIPs of membrane label and fusion protein showing the localization of Shroom within the pLLP B uninjected control and shroom3 MO injected MaxIPs B' Heat-maps of rosette detector score.

While in all of the above-mentioned studies Shroom3 was the focus of interest, in zebrafish and most other model organisms there are four paralogs. A previous study performed by a colleague done on Shroom3 morphants in *D. rerio* (1) indicates that Shroom3 is also necessary for AC and rosette formation in the migrating pLLP. In summary she was able to show that...

1. *shroom3* is expressed in the pLLP from stages 24 – 48 hpf.
2. *shroom3* is expressed downstream of FGF signaling, which was shown via treatment with an Fgfr inhibiting drug.
3. Shroom3 localizes to rosette centers, which was accomplished by generating a transgenic line expressing a *shroom3-tagRFP* fusion protein under the control of a heat-shock promotor (figure 1.9A).
4. Rosette formation is impaired in MO injected embryos, which was shown quantitatively by using a specifically trained rosette detector (51) to count and weight single rosettes.

Current Model

Based on these and previous results, the current model for apical constriction in the pLLP assumes that (1) expression of *shroom3* is induced by FGF signaling (2) Shroom3 binds Rock and translocates it to the AJC to (3) mediate phosphorylation of NMII which (4) induces contraction of the actin network and AC (figure 1.10). Furthermore, AC is necessary for rosette assembly and subsequent NM deposition. In conclusion the current understanding is that without Shroom3 AC and rosette formation are impaired, leading to a defect in NMs and NM deposition.

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7 Heat shock proteins are enzymes that assist protein folding whose expression is heat activated
1.2.5 \textit{shroom3} mutants

Morpholino injection allows transient gene knockdown in various species which has been broadly used over recent years. Although they have been and are still a useful tool for the zebrafish community, there are a number of limitations.

Those include…

- they need to be injected for each experiment. Even if the person injecting is well trained, there is always some loss in embryos. Furthermore, injection leads to a delay in development
- the induction of non-specific phenotypes due to activation of, among others, p53-mediated cell death.
- the degradation of the morpholino, which allows to analyze the phenotype over a limited period of time only (3 to 5 days after injection)

Because of these limitations, a former colleague in the lab had generated a mutant once it became technically possible using Transcription activator-like effector nuclease (TALEN) to confirm and further study the role of Shroom3 during morphogenesis. An 8 bp deletion in the SD2 domain was isolated and maintained as a stable line. This mutation leads to a premature STOP codon disrupting the SD2 domain, thereby inhibiting Shroom3’s function.

While birth rates follow a distribution of Mendelian inheritance (after genotyping at 3 months of age), \textit{homozygous} mutant adults have a shortened lifespan (~6-9 months). Shroom3 mutants are morphologically similar to their siblings, however their gill flaps seem to be increased in size, swollen, and not exactly streamlined with the body. This is also evident by an increased frequency of gill flap beating. Like the MO injected embryos, in a first qualitative analysis of the phenotype the pLLP exhibits a noticeable defect in rosette assembly. To our surprise however, the number of NMs deposited at the end of migration was significantly increased in the mutants as compared to the controls.
1.3 Open Questions and Motivation

The main objectives of my thesis were...

1. to characterize the newly generated \textit{shroom3} mutant phenotype via quantitative imaging methods
2. to develop a methodology for an automated, more precise, higher throughput and less invasive quantification of the pLLP and LL phenotypes
3. to use the mutant phenotype to better understand the relationship between rosette assembly and NM deposition
4. to analyze the feedback between morphological changes (apical constriction and mediated rosette assembly) and cell fate specification in the pLLP.
Chapter 2

Materials and Methods

2.1 Materials

2.1.1 Chemicals

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CHAPTER 2. MATERIALS AND METHODS

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2.1.2 Solutions

Table 2.2: Solutions

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2.1.3 Antibodies
2.1. MATERIALS

### Table 2.3: Antibodies

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<td>Anti-Digoxigenin</td>
<td>Roche</td>
<td>1:200</td>
<td>11093274910</td>
</tr>
<tr>
<td>Anti-GFP</td>
<td>Torrey Pines</td>
<td>1:200</td>
<td>-</td>
</tr>
<tr>
<td>Anti-TAZ (rabbit)</td>
<td>Cell Signaling Technology</td>
<td>1:200</td>
<td>D24E4</td>
</tr>
<tr>
<td>Anti-ZO1</td>
<td>Zymed</td>
<td>1:200</td>
<td>33-9100</td>
</tr>
<tr>
<td>Alexa Fluor488</td>
<td>Invitrogen</td>
<td>1:500</td>
<td>710369</td>
</tr>
<tr>
<td>Alexa Fluor555</td>
<td>Invitrogen</td>
<td>1:500</td>
<td>Z25005</td>
</tr>
</tbody>
</table>

#### 2.1.4 Enzymes

### Table 2.4: Enzymes

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Company</th>
<th>cat.-no.</th>
</tr>
</thead>
<tbody>
<tr>
<td>BtsCI</td>
<td>NEB</td>
<td>R0647</td>
</tr>
<tr>
<td>DdeI</td>
<td>NEB</td>
<td>R0175</td>
</tr>
<tr>
<td>HaeIII</td>
<td>NEB</td>
<td>R0108</td>
</tr>
<tr>
<td>MnlII</td>
<td>NEB</td>
<td>R0163</td>
</tr>
<tr>
<td>NlaIII</td>
<td>NEB</td>
<td>R0125</td>
</tr>
<tr>
<td>NsiI-HF</td>
<td>NEB</td>
<td>R3127</td>
</tr>
<tr>
<td>Phusion Polymerase</td>
<td>NEB</td>
<td>M0530L</td>
</tr>
<tr>
<td>Pronase</td>
<td>Sigma</td>
<td>P5147</td>
</tr>
<tr>
<td>Ribolock</td>
<td>Thermo</td>
<td>EO0381</td>
</tr>
<tr>
<td>RNase A</td>
<td>Quiagen</td>
<td>1006657</td>
</tr>
<tr>
<td>RNase H</td>
<td>NEB</td>
<td>M0297L</td>
</tr>
<tr>
<td>SP6 RNA Polymerase</td>
<td>Thermo</td>
<td>EP0131</td>
</tr>
<tr>
<td>T4 Ligase</td>
<td>NEB</td>
<td>M0202T</td>
</tr>
<tr>
<td>T7 RNA Polymerase</td>
<td>Thermo</td>
<td>EP0111</td>
</tr>
<tr>
<td>Taq DNA Polymerase</td>
<td>Invitrogen</td>
<td>10342-020</td>
</tr>
<tr>
<td>Taq DNA Polymerase</td>
<td>VWR</td>
<td>733-1301</td>
</tr>
</tbody>
</table>

#### 2.1.5 Molecular Biology Kits
CHAPTER 2. MATERIALS AND METHODS

Table 2.5: Molecular Biology Kits

<table>
<thead>
<tr>
<th>Kit</th>
<th>Company</th>
<th>cat.-no.</th>
</tr>
</thead>
<tbody>
<tr>
<td>EdU Click-iT</td>
<td>Invitrogen</td>
<td>MP 10083</td>
</tr>
<tr>
<td>mMessage mMachine Sp6 Polymerase</td>
<td>Invitrogen</td>
<td>AM1340</td>
</tr>
<tr>
<td>PCR &amp; Gel Clean-Up</td>
<td>Sigma</td>
<td>NA1020</td>
</tr>
<tr>
<td>pGEM-T TA Cloning</td>
<td>Promega</td>
<td>A3600</td>
</tr>
<tr>
<td>Superscript III cDNA Synthesis</td>
<td>Thermo</td>
<td>18080051</td>
</tr>
<tr>
<td>Wizard SV Gel and PCR Clean-Up</td>
<td>Promega</td>
<td>A9282</td>
</tr>
</tbody>
</table>

2.1.6 Buffers

Table 2.6: Buffers

<table>
<thead>
<tr>
<th>Buffer</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Blocking Reagent (BR)</td>
<td>2% BR in maleic buffer + 5% serum</td>
<td></td>
</tr>
<tr>
<td>E3</td>
<td>(52)</td>
<td></td>
</tr>
<tr>
<td>Hybridization buffer</td>
<td>50% Formamide + 25 % 20x SSC + 50mg/mL</td>
<td></td>
</tr>
<tr>
<td>Maleic buffer</td>
<td>250 mM maleic acid + 5M NaCl + 10 % 0.1% Tween-20 + mQ</td>
<td></td>
</tr>
<tr>
<td>NTMT</td>
<td>5 M NaCl + 1 M MgCl₂ + 1 M Tris pH 9.5 + 10% Tween</td>
<td></td>
</tr>
<tr>
<td>PBS</td>
<td>2.7 mM KCl + 12 mM HPO₄²⁻</td>
<td></td>
</tr>
<tr>
<td>PBST</td>
<td>PBS + 0.1% Tween20</td>
<td></td>
</tr>
<tr>
<td>PBDT</td>
<td>PBS + 1% BSA + 1% DMSO + 0.3% Triton</td>
<td></td>
</tr>
<tr>
<td>PFA</td>
<td>4% paraformaldehyde in PBS</td>
<td></td>
</tr>
<tr>
<td>P1</td>
<td>50 mM Tris-HCl pH 8.0 + 10 mM EDTA pH 8.0 + 100 µg/mL RNase</td>
<td></td>
</tr>
<tr>
<td>P2</td>
<td>1M NaOH + 10 % (w/v) SDS</td>
<td></td>
</tr>
<tr>
<td>P3</td>
<td>3M KOAc pH 5.5</td>
<td></td>
</tr>
<tr>
<td>TNT</td>
<td>50 mM Tris-HCl pH 8.0 + 100 mM NaCl + 0.1% Tween-20</td>
<td></td>
</tr>
</tbody>
</table>

2.1.7 Zebrafish lines
2.1. MATERIALS

### Table 2.7: Zebrafish lines

<table>
<thead>
<tr>
<th>Allele</th>
<th>name</th>
<th>zfin</th>
</tr>
</thead>
<tbody>
<tr>
<td>zf106Tg</td>
<td>cldnb:lyn-gfp</td>
<td>Tg(-8.0cldnb:LY-EGFP)</td>
</tr>
<tr>
<td>fu13Tg</td>
<td>exer4b(BAC):H2BRFP</td>
<td>TgBAC(exer4b:Hsa.HIST1H2BJ-RFP)</td>
</tr>
<tr>
<td>nns8Tg</td>
<td>atoh1a:Tom</td>
<td>Tg(atoh1a:dTomato)nns8</td>
</tr>
<tr>
<td>fu50</td>
<td>shroom3</td>
<td>-</td>
</tr>
<tr>
<td>m1274Tg</td>
<td>hsp70:shr3v1FL-taqRFP</td>
<td>Tg1(hsp70l:shroom3-TagRFP)</td>
</tr>
</tbody>
</table>

#### 2.1.8 ISH probes

### Table 2.8: ISH probes

<table>
<thead>
<tr>
<th>Probe</th>
<th>Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>atoh1a</td>
<td>see (38)</td>
</tr>
<tr>
<td>deltaD</td>
<td>see (53)</td>
</tr>
</tbody>
</table>

#### 2.1.9 Morpholinos

### Table 2.9: Morpholinos

<table>
<thead>
<tr>
<th>Probe</th>
<th>Sequence</th>
<th>Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>MoAtoh1a</td>
<td>see (38, 54)</td>
<td>0.4 ng/mM</td>
</tr>
<tr>
<td>p53</td>
<td>see (55)</td>
<td>2 ng/mM</td>
</tr>
</tbody>
</table>

#### 2.1.10 Hardware

##### Mounting Stamp

An stl file for 3D printing can be found at github.com/KleinhansDa/3DModels

### Table 2.10: Materials for production of a standardized mounting stamp

<table>
<thead>
<tr>
<th>Component</th>
<th>Company</th>
<th>cat.-no.</th>
</tr>
</thead>
<tbody>
<tr>
<td>µ dish</td>
<td>Ibidi</td>
<td>81,218–200</td>
</tr>
<tr>
<td>Stamp</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Preparation needles</td>
<td>VWR</td>
<td>USBE5470</td>
</tr>
<tr>
<td>Pasteur Pipettes</td>
<td>Roth</td>
<td>4518</td>
</tr>
<tr>
<td>Rubber / Silicone bulb</td>
<td>VWR</td>
<td>612-2327</td>
</tr>
</tbody>
</table>
CHAPTER 2. MATERIALS AND METHODS

<table>
<thead>
<tr>
<th>Component</th>
<th>Company</th>
<th>cat.-no.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Microtubes 2 mL</td>
<td>Sarstedt</td>
<td>2691</td>
</tr>
<tr>
<td>Heating block</td>
<td>PeqLab</td>
<td>HX2</td>
</tr>
<tr>
<td>Microwave oven</td>
<td>Severin</td>
<td>MW7849</td>
</tr>
<tr>
<td>Stereo microscope</td>
<td>Leica</td>
<td>M165FC</td>
</tr>
<tr>
<td>Transmitted Light Base</td>
<td>Leica</td>
<td>MDG36</td>
</tr>
<tr>
<td>Countersunk screw DIN7991, 8 × 20 mm</td>
<td>Dresselhaus</td>
<td>7662389</td>
</tr>
<tr>
<td></td>
<td>(Hornbach)</td>
<td></td>
</tr>
<tr>
<td>Superglue</td>
<td>UHU</td>
<td>509141</td>
</tr>
</tbody>
</table>

Spinning Disc Microscopy

Table 2.11: Spinning Disc system components

<table>
<thead>
<tr>
<th>Component</th>
<th>Company</th>
<th>Product</th>
<th>Specs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Microscope</td>
<td>Nikon</td>
<td>Eclipse Ti-E</td>
<td>fully motorized</td>
</tr>
<tr>
<td>PFS</td>
<td>Nikon</td>
<td>Perfect focus system</td>
<td>Z repositioning</td>
</tr>
<tr>
<td>XY-table</td>
<td>Merzhaeuser</td>
<td>XY motorized table</td>
<td>1 µm accuracy</td>
</tr>
<tr>
<td>Piezo</td>
<td>Yokogawa</td>
<td>CSU-W1</td>
<td>50 µm pattern</td>
</tr>
<tr>
<td>SD system</td>
<td>Laser Combiner</td>
<td>see table 2.12</td>
<td></td>
</tr>
<tr>
<td>FRAPPA</td>
<td>Revolution</td>
<td>FRAPPA</td>
<td>-</td>
</tr>
<tr>
<td>Borealis</td>
<td>Borealis</td>
<td>Borealis</td>
<td>flat field correction</td>
</tr>
<tr>
<td>sCMOS</td>
<td>Andor</td>
<td>ZYLA PLUS</td>
<td>4.2Mpix; 82%QE</td>
</tr>
<tr>
<td>Immersion</td>
<td>Merzhaeuser</td>
<td>Liquid Dispenser</td>
<td>-</td>
</tr>
</tbody>
</table>

Table 2.12: Available lasers

<table>
<thead>
<tr>
<th>Lasers</th>
<th>Type</th>
<th>Power</th>
</tr>
</thead>
<tbody>
<tr>
<td>405 nm</td>
<td>diode</td>
<td>100 mW</td>
</tr>
<tr>
<td>445 nm</td>
<td>diode</td>
<td>80 mW</td>
</tr>
<tr>
<td>488 nm</td>
<td>DPSS</td>
<td>100 mW</td>
</tr>
<tr>
<td>561 nm</td>
<td>DPSS</td>
<td>100 mW</td>
</tr>
<tr>
<td>640 nm</td>
<td>diode</td>
<td>100 mW</td>
</tr>
</tbody>
</table>
2.2. METHODS

Table 2.13: Available objectives

<table>
<thead>
<tr>
<th>Objective</th>
<th>Company</th>
<th>Type</th>
<th>Immersion</th>
<th>N.A.</th>
<th>Working distance</th>
</tr>
</thead>
<tbody>
<tr>
<td>10x</td>
<td>Nikon</td>
<td>CFI APO</td>
<td>air</td>
<td>0.45</td>
<td>4.00 mm</td>
</tr>
<tr>
<td>20X</td>
<td>Nikon</td>
<td>CFI APO</td>
<td>water</td>
<td>0.95</td>
<td>0.95 mm</td>
</tr>
<tr>
<td>40X</td>
<td>Nikon</td>
<td>CFI APO</td>
<td>water</td>
<td>1.15</td>
<td>0.60 mm</td>
</tr>
<tr>
<td>60X</td>
<td>Nikon</td>
<td>CFI APO</td>
<td>water</td>
<td>1.20</td>
<td>0.30 mm</td>
</tr>
</tbody>
</table>

Workstation

Statistical computation and image analysis were done on a Fujitsu Siemens (FS) Workstation CELSIUSM740 with the following hardware components...

Table 2.14: Workstation hardware components

<table>
<thead>
<tr>
<th>Component</th>
<th>Company</th>
<th>Product</th>
<th>Specs</th>
</tr>
</thead>
<tbody>
<tr>
<td>CPU</td>
<td>Intel</td>
<td>Xeon E5-1660v4</td>
<td>3.2 GHz, 20MB, 8cores</td>
</tr>
<tr>
<td>RAM</td>
<td>Fujitsu</td>
<td>-</td>
<td>4x16GB DDR4-2400</td>
</tr>
<tr>
<td>Graphics</td>
<td>NVIDIA</td>
<td>Quadro M4000</td>
<td>8GB RAM</td>
</tr>
</tbody>
</table>

2.1.11 Software

Table 2.15: Used software

<table>
<thead>
<tr>
<th>Software</th>
<th>Version</th>
<th>web</th>
</tr>
</thead>
<tbody>
<tr>
<td>Imagej FIJI</td>
<td>1.48</td>
<td><a href="https://fiji.sc/">https://fiji.sc/</a></td>
</tr>
<tr>
<td>R</td>
<td>3.6.1</td>
<td><a href="https://cran.r-project.org/">https://cran.r-project.org/</a></td>
</tr>
<tr>
<td>RStudio</td>
<td>1.0.153</td>
<td><a href="https://www.rstudio.com/">https://www.rstudio.com/</a></td>
</tr>
<tr>
<td>Ubuntu</td>
<td>17.1</td>
<td><a href="https://www.ubuntu.com/">https://www.ubuntu.com/</a></td>
</tr>
<tr>
<td>Windows 10 Pro</td>
<td>10.0.16299</td>
<td></td>
</tr>
<tr>
<td>Total Commander</td>
<td>9.0</td>
<td><a href="http://ghisler.com/">http://ghisler.com/</a></td>
</tr>
</tbody>
</table>

2.2 Methods

2.2.1 Data Strategy and Analysis

Due to the history of Developmental Biology and the complexity of biological processes per se, the field heavily relies on image data. Since
the advent of electronic imaging techniques\textsuperscript{1} scientific image data can be processed and analyzed \textit{in silico}. To take advantage of

1. live imaging, which (as compared to fixation techniques) conserves the cellular integrity and morphology while also offers the possibility of recording time-lapses
2. the optically clear specimen and
3. high throughput image analysis and state-of-the-art data science using algorithmic implementations,

the three following points were paid special attention to.

\textbf{Sample Preparation}

For fluorescence microscopy zebrafish embryos are usually immersed in a 1\% solution of low melting-point agarose (LMPA) and then oriented on an optical cover slip manually until the LMPA has solidified. This process allows to mount\textsuperscript{2} eight to ten embryos \textit{per} dish. To make use of the high number of offspring, a single zebrafish female may lay, which rapidly leads to a sample number of more than 300, a new sample preparation technique was designed that allows for (1) a four to five - time increase in samples per dish (2) a facilitated navigation \textit{via} a grid-like orientation through the samples and (3) an improved spatial orientation where the embryos body axes are aligned parallel to the optical Z-sections of the confocal microscope. For details, see Materials and Methods section 2.2.3 and Kleinhans \textit{et al.}, 2019 (\textsuperscript{56}).

\textbf{Imaging}

Technically, speed and sensitivity are most important for live imaging. Considering these two parameters a light-sheet (\textsuperscript{57}) fluorescence microscope (LSFM) would be the best fit. However, LSFM\textsuperscript{s} also have several limitations. First, due to the sample preparation methods available, the number of samples that can be imaged at a time is highly restricted. Second, for subcellular resolution a high magnification is required, which is limited by working distances and third - for optimal image analysis a high signal-to-noise ratio (SNR) and numerical aperture (N.A.) is preferable. Therefore a spinning disc (\textsuperscript{58}) system was chosen for most of the imaging. The system makes use of (1) an extra-large field of view (FOV) ideal for large specimen, (2) the possibility of a high degree of automation with state-of-the-art software and (3) a water dispensing system for long-term water immersion imaging. For details about the system see Materials section 2.1.10.

\textsuperscript{1}e.g. photomultipliers or charge-coupled devices
\textsuperscript{2}the process of embedding the samples in agarose
2.2. METHODS

Data handling

After data acquisition and pre-processing, the image data was transferred from the microscope system to the lab’s main workstation. To uniquely identify each file and have them appear in a structured manner, a filename-naming system was established after the following structure

\[ \text{[stage]}\text{[group]}\text{[id]}\text{[date]} \]

Where \text{stage} would e.g. be 32hpf, \text{group} would be a genotype or drug treatment, \text{id} would be a positional identifier on an imaging dish like B1P01\(^3\) and \text{date} would be a date in the form of YYMMDD.

Image and Data Analysis

In order to be as objective and as high throughput as possible, almost all of the analyses performed for this study was solved either algorithmically or using convolutional neural networks (CNNs). Furthermore, to meet the terms and conditions of open science\(^4\) standards, all pipelines were implemented in open source software frameworks such as Fiji is just image J (FIJI) and R. For further information about training datasets, algorithms and versions used see Materials section 2.1.11.

2.2.2 Zebrafish

Husbandry

Zebrafish husbandry was maintained at the University of Frankfurt am Main. All legal procedures were followed while handling and maintaining zebrafish husbandry. All zebrafish lines used in and generated for this study are listed in Materials section 2.1.7.

Handling and rearing

In the afternoon preceding the embryo collection, 1 male and female were set up in crossing cages, physically separated by a transparent separator. Next day, before noon, separators were removed allowing fertilization. Fertilized eggs were then collected, sorted and reared in the well-defined culture medium E3 (Kimmel et.al. 1995, section 2.1.6) at 25°, 28.5°, or 30°C depending on the experimental condition required. To grow larvae to the adulthood, they were transferred to the system on day 5. Till day 12, larvae were fed Vinegar Eels, Paramecia, and caviar powder.

\(^3\)Where B stands for a batch, that is if multiple dishes were imaged and P stands for the position within a single batch

\(^4\)“movement to make scientific research […] and its dissemination accessible to all levels of an inquiring society” – Wikipedia/en/Open_science
CHAPTER 2. MATERIALS AND METHODS

After the 12th day, water supply was started and fish were fed Brine Shrimp, Artemia, Paramecia and Vinegar Eels. Adult fish (>1 Month) were fed Artemia and the dry flakes.

Zebrasfin fin clips

Adult fish were anesthetized with buffered Tricaine (1X, see section 2.1.6) until loss of motion. About 1/3 of the caudal fin was cut with a sterile scalpel in a sterile Petri Dish. Immediately the dissected fin was transferred to 100 µL of 50 mM NaOH. Fish were returned to system water and kept in 1L system water in single tanks with 200 µL of 0.01% Methylene Blue.

Adult Genotyping

The clipped fins were digested for 1 h at 95°C and neutralized subsequently with 10 µL of 1M Tris-HCl of (pH 9).

Embryo Genotyping

Single fixed/live embryos were denatured at 95°C in 20 µL of 50 mM NaOH for 1 hour and neutralized by adding 2.5 µL of 1 M Tris-HCl (pH 9).

Zebrasfin Euthanasia

Adult zebrafish were euthanised by an overdose of Tricaine in ice cold water so as to sacrifice them by hypothermia.

Fixation

Embryos and dechorionated larvae were fixed in 2 mL of 4% PFA in 1X PBS overnight at 4°C.

2.2.3 Wet lab

Sample preparation

For samples older than 24 hpf, embryos were grown in 1X PTU till desired stage and treated with 150 µL per 10 mL of 0.1 mg/mL Pronase for ~30 min.. Choria were removed by gentle pipetting with a 2 mL plastic pasteur pipette. To replace the Pronase solution with fresh E3, embryos were immobilized by anesthesia and collected in the center of the dish by gentle rotational movement. Then the medium was decanted by collecting the embryos at the corner bottom of the dish while taking care not to loose any. After, the dish was filled with fresh E3. This process was repeated three times.
For samples **younger than 18s stage**, embryos were treated with 150 μL per 10 mL of 0.1 mg/mL Pronase directly. Choria were removed the same way as for > 24 hpf embryos but when pouring away the Pronase solution, the dish was simultaneously and very carefully filled with fresh E3 again. Since younger embryos are more fragile and to avoid damage, they must be kept in solution constantly.

Fixation started at the desired stage in 4% PFA in 0.1% PBST in 2 mL tubes at 4°C o.n.. The next day, samples were rinsed 3 times for ~5 min. in PBST and passed through a MeOH series of 25% → 50% → 75% → 100% MeOH/PBST (V/V)). For permanent storage, samples were stored in 100% MeOH at -20°C.

**In Situ Hybridization**

Samples were prepared after the method described in section 2.2.3.

1. **Permeabilisation & Probe Hybridization**  
   Permeabilisation → without shaking

Samples were rehydrated in an inverse MeOH series of 75% → 50% → 25% → 0% PBST and washed again for five min. two times in pure PBST. Finally, samples were digested in 10 μg/mL Proteinase K according to table 2.16. Samples were rinsed again two times in PBST and post-fixated in 4% PFA at 4°C for > 30 min. Samples were washed again for 5 min. three times in PBST.

**Probe Hybridisation** → all steps at 60°C, except stated differently

Samples were pre-hybridized in 350 μL of hybridization buffer (section 2.1.6) for 1 - 8 h. Just before detection probe treatment, the probe was denatured at 80°C in 1:200 of hybridization buffer. Subsequently, hybridization buffer was taken off the samples and replaced with the heated probe. Finally, samples were incubated o.n. at a desired temperature (around 65°C).

2. **Probe removal & Antibody incubation**  
The next day the probe got collected and stored at -20°C for re-use. Washing took place at the
same temperature as hybridization (from step 1) To keep solutions at temperature a Thermo-Block was used. For the washing series the samples were first washed one time for 20 min. in hybridization buffer, then two times for 30 min. in 50% Formamide and one time for 20 min. in 25% Formamide. Then two times for 15 min. in 2X SSCT and two times for 30 min. in 0.2X SSCT. Finally, one time for 5 min. in TNT.

To reduce noise and increase specific signal strength, the samples were treated with blocking solution (section 2.1.6) for 1 - 8 h in 350 µL of 2% BR/TNT at room temperature (RT). Afterwards the samples were incubated in 100 µL Anti-Digoxigenin diluted in NTMT buffer (1/4000 (V/V)) in 2% BR/TNT for 2 h at RT or o.n. at 4°C.

3. Probe detection First, the samples were washed six times for ~20 min. (or one wash o.n.) in TNT and two times for ~5min. in NTMT. After washing, color staining was performed with 4.5 NBT µL + 3.5 µL BCIP per mL NTMT in the dark and at RT without shaking (in a drawer) for 2 - 8 h, regularly checking the progression of the reaction. As soon as an appropriate degree of color intensity on the target site was achieved (up to two days), the samples were again washed three times in PBST.

Afterwards the samples were either prepared for immunostaining or imaging. For permanent storage samples were kept in 50% Glycerol at 4°C.

Immuno staining

Samples were prepared after the method described in section 2.2.3.

First, samples were blocked in 2% Goat Serum / PBDT (V/V) for 30 min.. For protein target site detection, a primary antibody (150 µL of 2% NGS / PBDT (V/V)) was incubated for ~2 h at RT or o.n. at 4°C. Samples were washed for 2 h in PBDT while changing the solution 5 - 6 times. To stain the now bound primary antibody, a secondary antibody (150 µL of 2% NGS / PBDT (V/V)) was incubated for 2 h at RT or o.n. at 4°C. Samples were washed for 2 h in PBDT while changing the solution 5 - 6 times.

Mounting

For live microscopy zebrafish embryos are usually immersed in a 1% solution of low melting-point Agarose (LMPA) solution and then oriented on an optical cover slip manually until the LMPA has solidified. This process allows to mount eight to ten embryos per dish.

To take advantage of the high number of offspring a single zebrafish female may lay, a new sample preparation technique was designed that allows for

1. a four to five times increase in samples per dish

28
2.2. METHODS

2. a facilitated navigation via a grid-like orientation through the samples
and
3. an improved spatial orientation where the embryos body axes are
aligned parallel to the optical Z-sections of the confocal microscope.

A detailed protocol can be found under section 3.1.1

2.2.4 Dry lab

Image J macros

Three IJ macros have been developed to facilitate image analysis and make
results more reproducible. Each of them is specifically designed for input
of LL and pLLP images of the cldnb:lyn-gfp transgenic line.
Hence, they are called *anaLLzr* . . .

- **2D** - analysis of Z-projected images of the LL at end of migration
- **2DT** - analysis of Z-projected images of the LL during migration
- **3D** - analysis of 3D image stacks of the pLLP at a given timepoint

Since their development was an integral part of my PhD work, the de-
scription of the macros can be found in the results part in section section
3.1.

Proliferation Analysis

The basic principle is based on work done by Laguerre *et al.*, 2009 (34).
For registration of mitotic events an IJ manual tracking tool was used that
allows to track an image feature through a stack of images creating tracks
as it progresses through volume / time (‘MTrackJ’(59)).

For mounting the embryo, the procedure described in section 2.2.3 was
used. Nuclei were visualized in a *TgBAC(cxcr4b:H2B-RFP)* transgenic line.
After Z-projection of volumetric timelapses, mitotic events were tracked in
each CC and the pLLP. Afterwards the data was exported as one table
per embryo and processed by counting mitoses per pLLP / CC / total CC
mitoses. Figure 2.1 shows an exemplary track for the data analyzed.

![Figure 2.1: Tracking of mitotic events. T1-T3 show consequetive timepoints of a single nucleus.](image-url)
CHAPTER 2. MATERIALS AND METHODS

Apical Index

Rationale The earliest attempt found for indexing AC can be found in a study published by Lee et al (60) where they were interested in the ‘apical index’ (A.I.) of bottle cells during X.laevis gastrulation (figure 2.2 Lee). Another example for measuring AC is the apical constriction index (A.C.I., figure 2.2 Harding) for the cells of the D.rerio lateral line primordium (pLLP), which can be found in a study from 2012 where it was shown that FGFR-Ras-MAPK signaling is required for Rock2a localization and AC (61, 62).

In these two publications, the way they measure A.I. (60) and ACI (62) respectively, does not differ and is the ratio of lateral height over apical width.

\[
\text{ACI} = \frac{\text{lateral height [\mu m]}}{\text{apical width [\mu m]}}
\]

We found two principal weaknesses of applying this ratio to the cells of the pLLP. First, it does not respect the independence of lateral height to AC. Second, it does not differentiate between constriction along the anterio-posterior (AP) or the dorso-ventral (DV) axis. Third, it actually represents the A.I. rather than the apical constriction index.

Parameter definition To obtain a precise and biologically meaningful way to quantify AC, first a couple of definitions had to be made.

Definition 2.1 (AC is independent of orientation). In a 3D space a cell can have any orientation and still be apically constricted. Therefore, before measuring one should make sure orientation between embryos is aligned and
also consider taking measurements along two different directions. Since apical constriction is not necessarily isotropic, it is important to consider constriction along 2 perpendicular axes (AP and DV axis of the embryo/pLLP).

**Definition 2.2** (AC is independent of lateral height). Lateral height can be described as the distance of the two farthest points on the surface area of a cell. Two cells with different lateral heights can be equally apically constricted.

**Definition 2.3** (AC is independent of cell volume). The volume of a cell represents its size. A large cell can be equally constricted as a small cell.

**Adaption for variation in lateral height** To test different A.I. conditions, an apically constricted cell can be approximated by modeling a tetrahedron. For example, shrinking or enlarging a cell symmetrically should not affect the A.I. As described by Harding (2014) (62), the apical width of a cell is measured first by manual 3-D reconstruction, second manual re-orientation, and third by going 1 µm from the apical tip into the cell (from now on referred to as Δap, 2.3B). Finally, apical width is the total width of the 2D object in the respective volume. If Δap is a constant, the A.I. in a symmetrically enlarged cell increases from e.g. ~15 to ~23, since apical width stays the same but lateral height increases (compare figure 2.3A to A’). On the contrary, if Δap is adjusted relative to a cells lateral height, e.g. by percentage, the A.I. in a symmetrically enlarged cell stays the same (compare figure 2.3A to A’’).

![Figure 2.3: Different ways to quantify the apical index. A-A’’ A.I. Cell Models. A’ and A’’ show cells that are symmetrically increased versions of A. While in A’, constant delta was used, in A’’ delta is proportional to the lateral height. B Illustrating delta ap. (left) apically constricted cells volume rendered in XY (top) and as a lateral cross-section in X-Z (bottom). (right) 2-D area as seen at Δap of 1 or 2.5 µm.](image)

Therefore the measurement for apical width has to be relative to lateral height.
\[ \text{ACI} = \frac{\text{lateral height [\(\mu m\)]}}{\text{relative apical width [\(\mu m\)]}} \]

**Adaption for tissue polarization**  Organs develop in a 3-D space and are polarized along each axis. AC usually describes a 2-D morphogenetic movement towards a center along the X-Y axes. However, the constriction movements along X and Y might be independent of one another. This could mean that they happen at different speeds, or that one is absent. As a result, the tissue would look less radially (figure 2.4B) constricted, but more constricted along one particular axis (anisotropic). In order to separate those two AC dimensions, the A.I. can be calculated for the *anterio-posterior* and for the *dorso-ventral* axis (figure 2.4, horizontal vs. vertical).

![Figure 2.4: Schematic AC along the A-P and D-V axis. A shows a A-P and D-V constricted cluster of cells. B shows a D-V constricted cluster of cells.](image)

By fitting an ellipsoid to the area taken at \(\Delta ap\), one will obtain the following parameters (figure 2.5).

1. **Length of Major axis**
   - indicates the level of constriction along the less constricted axis
2. **Length of Minor axis**
   - indicates the level of constriction along the most constricted axis
3. **Angle of Major from 0°**
   - indicates the orientation of the long, less-constricted axis: If the angle is close to 0°, the long axis of the apical area is parallel to the AP axis (the cell is constricted along the DV axis). If the angle is close to 90°, the long axis of the apical area is parallel to the DV axis (the cell is constricted along the AP axis).
2.2. METHODS

Figure 2.5: Scheme of ellipsoid measures. A shows the major axis as apical width and the minor axis as apical height. B shows the angular displacement from the horizon in steps of 30°.

**Measurement definition**  The two dimensions of A.I. indices can therefore be defined as the following ratios...

**Definition 2.4 (A.I. Major).**

\[
\text{A.I. Major} = \frac{lateral \ height \ [\mu m]}{ellipsoid \ major \ axis \ at \ relative \ \Delta ap \ [\mu m]}
\]

**Definition 2.5 (A.I. Minor).**

\[
\text{A.I. Minor} = \frac{lateral \ height \ [\mu m]}{ellipsoid \ minor \ axis \ at \ relative \ \Delta ap \ [\mu m]}
\]

**Definition 2.6 (Angle Major).**

\[
\text{Angle Major} = \angle = \Delta \ from \ horizon \ [0 - 90°]
\]

**Measurements**  As a proof of principle of the definitions stated in the previous section we compare our results to previously published results from Harding et al. (61) who, as a control, measured apical constriction in embryos treated with an FGF inhibitor (SU5402) and their DMSO controls.

**Single cell measurements**  Each geometric object has a centroid coordinate in X and Y (and Z) which is represented as the mean of all X or Y coordinates within the object. In figure 2.3, centroid coordinates in X and Y are used to plot the cells as points in the X-Y plane. Additionally, each point is colored for the A.I. value (high values are dark red - red, middle values are green, low values are cyan - blue).
Harding et al. (62) were using a constant $\Delta a_p$ to measure the apical width, which we have shown to be incorrect in certain cases. In their study they found that certain mean A.C.I. values in the DMSO go as high as 15 (figure 2.7), which might be related to this (see figure 2.3). By measuring apical width at a relative $\Delta a_p$ and taking into account all pLLP cells of the two exemplary pLLPs shown in figure 2.3, we measure a mean difference in the Major of 0.53 and 1.11 in the Minor.

Figure 2.6: A.I. $\text{Major} / \text{Minor}$ single cell measurements

Figure 2.7: A.I. indices by Harding et al. E-G’ 3-D reconstructions of the highlighted cell. H A.C.I.s for embryos treated with DMSO, SU5402, PD0325901 or following induction of $hsp70:dn-Ras$. (n = 180 cells / N = 6 embryos).
2.2. METHODS

**Angle densities**  To check whether there is a bias in orientation of the apical width, the angle measurements 2.5 can be shown as a function of density along X (figure 2.8A).

Interestingly the results indicate that there is less of a difference for the Major Angle at angles bigger than 15-20°. This would mean that the apical surface of the cells in SU5402 treated embryos is more strongly oriented along the horizontal *antero-posterior* axis.

**ACI magnitude at different angles**  Now, to get an idea of the magnitude of constriction relative to the orientation of the cell (angle to the horizontal), the A.I. Major/Minor can be shown as a function of the Major Angle (figure 2.8B-B').

Since AC is a 3-D morphogenetic process and since cells in a wild type pLLP are mostly radially organized, it does make sense to look at AC from more than just one perspective. Here we propose to separate the A.I. into an *antero-posterior* and a *dorso-ventral* dimension.

1. While for the control (DMSO treated) embryo the distribution of the cells Major Angles seem to be mostly uniform, for the SU5402 treated embryo there is an accumulation of lower Major Angles. This means that cells in SU5402 treated embryos are more oriented along the horizontal (anterior - posterior) axis.

2. Interestingly there does not seem to be much of a difference in A.I. Major (figure 2.8B), which can also be shown by the mean values which are at 2.6 for the DMSO control and at 2.1 for the SU5402 treated condition.

3. For the A.I. Minor (figure 2.8B') the means are 4.7 for the DMSO control and 3.6 for SU5402. The base constriction for both, DMSO and SU5402 is at around 3.6, however there is a peak at around 40 - 60° in the DMSO control where cells are most constricted having a maximum A.I. at 15.8. This indicates that for the Minor, cells in that range of angles are more constricted than cells oriented in different directions.
2.3 Ground Truth

Analyzing images and extracting quantitative measurements can be a tedious task, especially when the analysis becomes more complex. Fortunately, there are ways to automate image analysis by using either machine learning approaches or by tailoring hand-crafted algorithms in an image analysis software tool like e.g. ImageJ(63). The main advantages of doing so are to...

- be independent of confirmation bias
- make the analysis more robust against oversight
- increase the statistical power by increasing the number of data points(64)
- increase effect size by increasing the measurement accuracy(64)

However, to ensure the measurements taken by a tailored or trained algorithm are meaningful, they must be compared to a ground truth dataset which again describes a general measure of algorithmic quality performance(65).

2.3.1 Cluster Analysis

The \textit{anaLLzR2D} algorithm was designed for semi-automatic cell cluster detection in the \textit{cldnb:lyn-gfp} transgenic line and optional nuclei counting in a second DAPI-labeled channel within the Regions of Interest (ROIs) derived from the cell cluster detection.

Design

To assess the quality of the \textit{anaLLzR2D} algorithm for nuclei detection the ground truth was designed as follows.

Model
2.3. GROUND TRUTH

- each Cell Cluster (CC) consists of a number of objects (cells)
- each object is part of the respective CC and defines one cell entity
- each object is determined via a fluorescent nucleus label
- embryos are mounted within a 3D mold (section 2.2.3) to reduce noise

Table 2.17: Cluster Analysis Model

<table>
<thead>
<tr>
<th>XY scale</th>
<th>0.32 px/µm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Z-spacing</td>
<td>4 µm</td>
</tr>
<tr>
<td>Camera</td>
<td>Rolera</td>
</tr>
</tbody>
</table>
**Training & test data** The training set consists of three randomly picked wildtype pLLPs. For each the algorithm was run with standard parameters. Cell cluster ROIs and nuclei multi-point labels were edited manually. To test the algorithm, it is run at different nuclei detection thresholds on the same image data.

### 2.3.2 Morphometric analysis

The *anaLLzr3D* algorithm was designed for fully automated, single cell volume segmentation in the *cldnb:lyn-gfp* transgenic line. In addition to 3D cellular metrics, it offers Apical Constriction measurement of each cell.

**Design**

To assess the quality of the anaLLzr3D algorithm the ground truth was modeled as follows.

**Model**

- each pLLP consists of a number of objects (cells)
- each object is part of the pLLP and defines one cell entity
- cell boundaries are determined via the transgene *cldnb:lyn-gfp* +/+ 
- embryos are imaged live to conserve signal and membrane integrity 
- embryos are mounted within a 3D mold for improved 3D alignment

**Table 2.18:** anaLLzr3D Model

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Specification</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exposure time</td>
<td>100 ms</td>
</tr>
<tr>
<td>Laser intensity</td>
<td>100% / 9.3 mW</td>
</tr>
<tr>
<td>Objective</td>
<td>40X; CFI APO LWD WI; N.A. = 1.15, W.D. = 0.60 mm</td>
</tr>
<tr>
<td>Tube lens</td>
<td>1.0X</td>
</tr>
<tr>
<td>Z-spacing</td>
<td>0.4 - 0.5 μm</td>
</tr>
<tr>
<td>Camera</td>
<td>sCMOS; 4.2 M.Pix; 82% Q.E.</td>
</tr>
<tr>
<td>SD system</td>
<td>Yokogawa CSU - W1; 50 μm pattern</td>
</tr>
<tr>
<td>Piezo</td>
<td>Piezo Z-table; 300 μm scan range</td>
</tr>
</tbody>
</table>

**Training & test data** The training set consists of three randomly picked wildtype pLLPs. For each the algorithm is run with no filters (X, Y, Z border objects, size) and a minimum segmentation threshold. Afterwards the segmentation result is corrected manually for over- and under-segmentation and objects that are not part of the pLLP.
To test the algorithm it is run at different segmentation thresholds on the same image data.

2.4 Image Data Sets

Summaries of Image datasets. *Pairs* describe the number of parent pairs I harvested eggs from. *Stage* describes the time I waited for the parent pairs to mate and lay eggs. Since pair #1 might have laid their eggs earlier than pair #2, those batches would be some time apart in their staging. *Stamp* describes the version of the stamp I used, where the main difference between version 4 and 5 are more wells added and some minor upgrades in well-design.

Table 2.19: Cell Cluster dataset

<table>
<thead>
<tr>
<th>Crossings</th>
<th>Pairs</th>
<th>4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Transgenes</td>
<td>cldnb:lyn-gfp +/-</td>
<td></td>
</tr>
<tr>
<td>Mutation</td>
<td>shroom3</td>
<td></td>
</tr>
<tr>
<td>Staging</td>
<td>60 min.</td>
<td></td>
</tr>
<tr>
<td>Mounting</td>
<td>Fixation</td>
<td>4% PFA o.N.</td>
</tr>
<tr>
<td>Agarose</td>
<td>1% LMPA</td>
<td></td>
</tr>
<tr>
<td>Imaging</td>
<td>Magnification</td>
<td>25X WI + 1.0x zoom</td>
</tr>
<tr>
<td>Channels</td>
<td>488 nm</td>
<td></td>
</tr>
<tr>
<td>Z-Stack</td>
<td>2.5 µm Z-spacing; 110 µm stack size; 12*X large image</td>
<td></td>
</tr>
</tbody>
</table>

Table 2.20: A.I. dataset

<table>
<thead>
<tr>
<th>Crossings</th>
<th>Pairs</th>
<th>4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Transgenes</td>
<td>cldnb:lyn-gfp +/-</td>
<td></td>
</tr>
<tr>
<td>Mutation</td>
<td>shroom3</td>
<td></td>
</tr>
<tr>
<td>Staging</td>
<td>30 min.</td>
<td></td>
</tr>
<tr>
<td>Mounting</td>
<td>Protocol</td>
<td>Kleinhaus et al., 2019</td>
</tr>
<tr>
<td>Agarose</td>
<td>0.5% LMPA + 20% Tricaine (V/V%)</td>
<td></td>
</tr>
<tr>
<td>stamp</td>
<td>version 4A</td>
<td></td>
</tr>
<tr>
<td>Imaging</td>
<td>Magnification</td>
<td>40X objective + 1.0x zoom</td>
</tr>
<tr>
<td>Camera</td>
<td>Binning 1x1; Gain 1; Exposure 100 ms</td>
<td></td>
</tr>
<tr>
<td>Channels</td>
<td>488 nm (100%)</td>
<td></td>
</tr>
<tr>
<td>Z-Stack</td>
<td>0.4 µm Z-spacing</td>
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### Table 2.21: Proliferation dataset

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</thead>
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<td></td>
</tr>
<tr>
<td>Mutation</td>
<td>shroom3</td>
<td></td>
</tr>
<tr>
<td>Staging</td>
<td>30 min.</td>
<td></td>
</tr>
<tr>
<td>Mounting</td>
<td>Protocol</td>
<td>Kleinhans et al., 2019</td>
</tr>
<tr>
<td>Imaging</td>
<td>Agarose</td>
<td>0.3% LMPA + 20% Tricaine (V/V%)</td>
</tr>
<tr>
<td></td>
<td>stamp</td>
<td>version 4A</td>
</tr>
<tr>
<td></td>
<td>Magnification</td>
<td>20X + 1.5x zoom</td>
</tr>
<tr>
<td>Camera</td>
<td>Binning 2x2; Gain 4; Exposure 35 ms; full FOV*150 µm</td>
<td></td>
</tr>
<tr>
<td>Channels</td>
<td>651 nm (25%)</td>
<td></td>
</tr>
<tr>
<td>Z-Stack</td>
<td>2.5 µm Z-spacing; 110 µm stack size; 2*X large image</td>
<td></td>
</tr>
<tr>
<td>Time</td>
<td>20 h / 7 min. interval / start 2 p.m.</td>
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</tbody>
</table>

### Table 2.22: Detection dataset

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<td>Transgenes</td>
<td>cldnb:lyn-gfp +/?</td>
<td></td>
</tr>
<tr>
<td>Mutation</td>
<td>shroom3</td>
<td></td>
</tr>
<tr>
<td>Staging</td>
<td>30 min.</td>
<td></td>
</tr>
<tr>
<td>Mounting</td>
<td>Protocol</td>
<td>Kleinhans et al., 2019</td>
</tr>
<tr>
<td>Imaging</td>
<td>Agarose</td>
<td>0.3% LMPA + 20% Tricaine (V/V%)</td>
</tr>
<tr>
<td></td>
<td>stamp</td>
<td>version 5A</td>
</tr>
<tr>
<td></td>
<td>Magnification</td>
<td>20X + 1.5x zoom</td>
</tr>
<tr>
<td>Camera</td>
<td>Binning 2x2; Gain 4; Exposure 20 ms; full FOV*150 µm</td>
<td></td>
</tr>
<tr>
<td>Channels</td>
<td>488 nm (25%)</td>
<td></td>
</tr>
<tr>
<td>Positions</td>
<td>36</td>
<td></td>
</tr>
<tr>
<td>Z-Stack</td>
<td>3 µm Z-spacing; 100 µm stack size; 3*X large image</td>
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</tr>
<tr>
<td>Time</td>
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### Table 2.23: Atoh1a dataset

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</tr>
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<td>cldnb:lyn-gfp +/?; atoh1a:Tom +/0</td>
<td></td>
</tr>
<tr>
<td>Mutation</td>
<td>shroom3</td>
<td></td>
</tr>
<tr>
<td>Staging</td>
<td>30 min.</td>
<td></td>
</tr>
<tr>
<td>Mounting</td>
<td>Protocol</td>
<td>Kleinhans <em>et al.</em>, 2019</td>
</tr>
<tr>
<td>Agarose</td>
<td>0.3% LMPA + 20% Tricaine (V/V%)</td>
<td></td>
</tr>
<tr>
<td>stamp</td>
<td>version 5A</td>
<td></td>
</tr>
<tr>
<td>Imaging</td>
<td>Magnification</td>
<td>20X + 1.0x zoom</td>
</tr>
<tr>
<td>Camera</td>
<td>Binning 2x2; Gain 4; full FOV*150 µm</td>
<td></td>
</tr>
<tr>
<td>Channels</td>
<td>488 nm (Int: 20%; Exp: 25 ms)</td>
<td></td>
</tr>
<tr>
<td>Channels</td>
<td>561 nm (Int: 30%; Exp: 50 ms)</td>
<td></td>
</tr>
<tr>
<td>Positions</td>
<td>36</td>
<td></td>
</tr>
<tr>
<td>Z-Stack</td>
<td>2.5 µm Z-spacing; 100 µm stack size; 2*X large image</td>
<td></td>
</tr>
<tr>
<td>Time</td>
<td>20 h / 20 min. interval / start 2 p.m.</td>
<td></td>
</tr>
</tbody>
</table>
Chapter 3

Results

3.1 Quantitative imaging and image analysis

3.1.1 A 3D-printed stamp to standardize sample mounting and semi-automatize imaging

NOTE

Most of Chapter 3.1.1 has been published as an article in the Journal BMC Biotechnology (Kleinhans and Lecaudey, BMC Biotechnology (2019) 19:68). The author contribution was described in the paper as follows: DSK designed the study, carried out the experiments and analysis of the results. VL provided the infrastructure and funding for performing the experiments. DSK and VL wrote the final manuscript. All authors read and approved the final manuscript. Some passages in this section 3.1.1 have been quoted verbatim from the above-mentioned article for the scientific accuracy of the terms used.

Even though on a macroscopic scale development is a remarkably similar and synchronized process between zebrafish embryos, a single biological process on a microscopic scale even in sibling embryos can look drastically different. Given the noisy and variable character of biological systems, it is important to record a sufficient number of samples to obtain a quantitative and representative view of a biological process. Furthermore, to process biological samples of whole organisms in a high-content manner it is important to have a standardized way of sample mounting, data acquisition, data processing and analysis.
However, imaging a high number of samples and generating large datasets to date is still largely limited by the classical way developmental biologists mount embryos for imaging. A number of factors limiting the standardization are summarized in table 3.1. Therefore, especially for 3D segmentation and 2D tracking experiments where an exact staging and orientation of the embryo is necessary, there is a need for methods to standardize sample mounting and image acquisition of multiple embryos.

The protocol we describe here was designed to be used with XY scanning universal sample holders that usually come with any motorized-stage inverted microscope. Similar to previous approaches (66–69), it uses a 3D-printed stamping device to produce an Agarose imprint with a diameter of 20 mm on the cover glass of a 35 mm μ-dish. The imprint consists of 44 equally spaced μ-wells, which are designed to fit the average morphology of a zebrafish embryo between 24 and 96 hours-post-fertilization (hpf).

The aim was to develop a standardized mounting method allowing us to: (i) mount many samples in parallel in a 2D coordinate system of rows and columns, (ii) reduce the acquisition time and thus photo-bleaching and photo-toxicity during imaging, (iii) semi-automatize the acquisition, (iv) reduce the post-processing steps, and (v) facilitate subsequent processing such as genotyping due to a 1:1 correlation between image data and specimen arrangement sequence.

For mounting, an improved 3D specimen preparation and well-plate-like sample navigation for zebrafish larvae confocal microscopy was developed with which lateral line development can be recorded over more than 20 h, in up to 44 positions, in a confocal Z-stack of less than 120 μm and a time interval of 5–10 min. (depending on the number of channels and exposure time). The stamp was designed to be used for embryos between 24 and 96 hpf. For a tailor-made well, embryos were fixed and imaged in toto to measure the dimensions in X, Y and Z of different (whole embryo, trunk, yolk) embryonic structures (figure 3.1B-B'). Using Microsoft 3D-Builder a well was assembled from basic shapes like cube, sphere and wedge. After, the well was duplicated a couple of times and put in a grid-like arrangement to fit on a disc base 20 mm in diameter. Printing was performed on a Formlabs extrusion printer (figure 3.1C).

Procedure

**Preparation of the agarose cast** To prepare the agarose cast, the stamp is first cleaned from dust and remnants with tissue soaked in 70% Ethanol and pressured air (figure 3.3 IA). To prepare the casting medium a 1% Agarose (w/v) solution is prepared in an autoclaved 100 mL bluecap bottle by dissolving 200 mg of agarose in 20 mL of E3 using a microwave oven. From the ready solution 650 μL are applied to the 20 mm cover-
### 3.1. QUANTITATIVE IMAGING AND IMAGE ANALYSIS

**Table 3.1:** Limitations of traditional zebrafish mounting techniques

<table>
<thead>
<tr>
<th>Standard</th>
<th>Limitations</th>
<th>Solutions</th>
<th>Improvement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mounting in 1% LMPA</td>
<td>• Polymerization speed limits the number of embryos that can be mounted in parallel</td>
<td>• Use lower concentration of LMPA</td>
<td>• Mounting time extended</td>
</tr>
<tr>
<td></td>
<td>• Embryo growth is restricted by high-percentage LMPA</td>
<td>• Use heating device to keep the LMPA liquid longer</td>
<td>• Growth and change in shape of the embryo allowed</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Sample size increased</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Embryo retrieval facilitated</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Possibility to grow embryos to adulthood afterwards</td>
</tr>
<tr>
<td>No pre-defined positions</td>
<td>• Positioning and alignment in XY are neither standardized nor reproducible</td>
<td>• Mount embryos at pre-defined identically oriented and equidistant positions</td>
<td>• Relative positions of embryos identical in all experiments</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Easier setup of multi-dimensional imaging experiments</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Easier navigation between different XY locations</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Possibility for semi-automated imaging</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Identification of individual embryo facilitated for downstream experiments (genotyping)</td>
</tr>
<tr>
<td>No μ-wells that model the average</td>
<td>• Time-consuming orientation of the embryos</td>
<td>• Use μ-wells that model the average embryo shape</td>
<td>• Orientation of individual embryos during mounting in LMPA much faster</td>
</tr>
<tr>
<td>embryoshape at a defined</td>
<td>• Body axes not aligned to optical sectioning in Z due to huge yolk sac</td>
<td></td>
<td>• No need to re-orient the embryos in XY plane post-imaging</td>
</tr>
<tr>
<td>developmental stage</td>
<td></td>
<td></td>
<td>• Aligned morphological shapes in Z projections</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Reduced stack and file size</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Reduced photo-bleaching and toxicity- Reduced post-processing</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Increased scanning speed</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Improved signal-to-noise ratio</td>
</tr>
</tbody>
</table>
slip of a 35 mm imaging dish (materials in table 2.10). Subsequently the clean stamp is gently placed onto the placed solution and adjusted to the center. The dish is then rotated to distribute excess agarose over the entire dish surface to stabilize the imprint once polymerized. After about 30 min. the stamp is removed by first slipping a clean preparation needle between the stamp and the polymer and then lifting it from the cast (figure 3.3IB). If necessary, air bubbles appearing between the cover glass and the polymer are eliminated by punctuation with a preparation needle. The mounting cast may be used immediately or stored at 4°C for several days (with lid closed).

**Preparation of mounting media** Two solutions of low-melting point Agarose (LMPA) are prepared in autoclaved 100 mL bluecap bottles by dissolving 60 and 100 mg LMPA in 16 mL of E3 in a microwave oven - yielding 0.375 and 0.625 % respectively. Per stamped cast, 2 aliquots of 1.6 mL are prepared in 2 mL tubes for each LMPA concentration and placed in a heating block adjusted to 41°C. For live imaging, 400 µL of 4.2 mg/mL Tricaine (25X) are added to keep the embryos anesthetized during imaging. Final concentrations of LMPA solution are therefore 0.3 and 0.5 %, respectively. LMPA solutions containing Tricaine were prepared fresh for each mounting session. The LMPA solution and the mounting cast have almost equal refractive indices. Therefore, when adding the LMPA solution the cast becomes invisible. To still be able to locate the μ-wells and
3.1. QUANTITATIVE IMAGING AND IMAGE ANALYSIS

**Figure 3.2: Mounting Results.**

A - A' Maximum intensity projection of a 50 hpf embryo mounted on its side in XY and XZ. (right) Color scale indicates depth encoding.

C Multi-position (36), multi-channel (2) time-lapse recording (13 h duration; 15 min. interval).

D Multi-channel (2) Extended Depth of Focus (EDF) projections from widefield Z-stacks (recorded with 20x Objective). Scale Bar = 1mm

E Multipoint coordinates in X, Y and Z (recorded with 40x Objective). The offset describes the distance of each point from the mean of all points in X, Y and Z. Panel 1–3 (top to bottom) show dimensions X, Y and Z in comparison for the pLLP, the eye and the otic vesicle. The red line indicates the median, the blue line indicates zero offset, error bars indicate mean ± s.d.. Numeric values indicate the variance in each group.

F-F' Systematic retrieval for genotyping. F Mounted embryos in a 2-D coordinate system of rows (A-M) and columns (1–3). F' Imaging Sequence in a snake-by-column fashion. In a time-lapse setting, it starts at point 1 (P01) again to initiate the next timepoint. F'F' After imaging, the embryos are retrieved in the same sequence as they were imaged (snake by column, left panel). F'F' Each genotyping result on the electrophoresis gel is easily correlated to one imaging dataset with defined X-Y coordinates.
to position the embryos accordingly, the illumination contrast and mirror angle of a transmitted light base are adjusted to make the \( \mu \)-wells visible again (figure 3.3IIA-C).

**Mounting procedure**  In case of live imaging the embryos are first anesthetized in a Petri dish with 4 to 5 drops of 4.2 mg/mL Tricaine (40 \( \mu \)g/mL in E3) added 4 to 5 min. before usage.

For mounting, the cast is first gently filled from the border (figure 3.3II A3) with 500 \( \mu \)L of 0.3\% LMPA solution. Then, 44 embryos (one for each well) are collected from their Petri dish with a glass Pasteur pipette. To minimize the amount of liquid added to the LMPA, the embryos are allowed to sink to the air – liquid interface and immediately added in one drop to the liquid LMPA solution in the stamped cast. Next, each embryo is moved to a separate \( \mu \)-well with a preparation needle by positioning the yolk within the hemi-spherical structure of each well and the tail aligned horizontally with the shape of the \( \mu \)-well (figure 3.3III C-D). The LMPA was allowed to polymerize for about 40 min. For time-lapse recording longer than 1 h, 1 mL of 0.5\% LMPA was added on top and allowed to polymerize for another 10 min. to construct an Agarose *sandwich* to stabilize the structure. Since the 0.3\% LMPA will still be very fragile, the 0.5\% LMPA should be added to the outer well first, carefully raising the level. Since Agarose polymerization speed depends on temperature, for mounting the temperature of the room should not be less than 23\(^\circ\)C to give sufficient time. For indefinite time of embryo orientation, a higher room temperature or a 5 V terrarium heating mat (at maximum temperature ca. 38\(^\circ\)C) can be used. For the latter, a hole with the diameter of an imaging dish should first be cut in the middle of the heating mat. For mounting, the mat should then be placed and fixed on the stereo-microscope stage with the dish in the hole.

**Imaging setup**  The dish is placed onto the sample holder of an inverted confocal spinning disc microscope so that the embryos are aligned to the Y axis of the microscope stage. The stage is then moved to place the embryo at Position 01 (P01, top-left position) right above the objective (figure 3.3III A).

**Define embryo positions**  Since the embryos are mounted in a 3-D grid with well defined dimensions, all positions can be defined *via* a pre-defined points list that is loaded into the microscope software. For our system we use the ‘Nikon Imaging Software’ where we move the stage to P01, define a multi-point list with distance X / distance Y = \( \frac{3450}{1280} \) \( \mu \)m, bring P01 into focus and offset all points in Z. The list can also be saved for re-use in
3.1. QUANTITATIVE IMAGING AND IMAGE ANALYSIS

Refine Positions  Even though the mounting method allows for a precise positioning, each embryo physiology is still a bit different resulting in differences between positions but same structures of up to 100 µm in X, Y and Z (figure 3.2E). Therefore, before starting an experiment each position needs to be refined.

Retrieval  For further experiments such as genotyping, the embryos are retrieved from the agarose in the same sequence as they were imaged (figure 3.2F'). To do so, a glass pipette is inserted into the agarose and directed to the head region of an embryo. By applying a gentle underpressure the embryo is then sucked into the glass pipette.

To lyse the embryos and extract the genomic DNA, each embryo is placed in a single tube of an 8-tube PCR strip. Since 8-tube PCR strips are designed to work with multichannel pipettes, the genotyping PCR is performed and analysed by gel electrophoresis using an 8x-multichannel pipette. When using a 34-well comb, the pipette tips will reach every second well of the agarose gel. Filling the wells staggered (offset by 1), one can load $4 \times 8$ wells in one row (figure 3.2F''). Since each embryo has a defined position, it is straightforward to associate each genotype to the corresponding image data (figure 3.2F' - F'''). Since a single mismatch would mess up the entire experiment by resulting in a frameshift of the one-to-one correspondence, this is a very important feature. The imaging dish can be reused several times. For cleaning, the agarose bed is removed from the dish using a small scoop or preparation needle and wiped gently with a lint-free tissue soaked in Ethanol.
Figure 3.3: Stamping procedure Agarose Cast (A) clean stamp surface (B) preparation of the stamp before lifting (C) ready-for-use agarose imprint. Mounting (A) without LMPA (B and C) with LMPA, while the latter shows the imprint with light coming from a different angle, making the chambers visible again. (D) Horizontal alignment of embryos. Imaging (A) Positioning of the μ-well (B) Alignment in Brightfield and (C) Definition of a custom well plate.

Summary

The major improvements introduced by this method are (1) using a low percentage LMPA, which extends the timespan for mounting which is nec-
3.1. QUANTITATIVE IMAGING AND IMAGE ANALYSIS

It is essential to align a higher number of embryos. It also gives the embryo more freedom to grow during longer time-lapse imaging and facilitates retrieval afterwards. (2) using a stamped cast, which allows for standardized and reproducible positions of the embryo as shown for the lateral line primordium, the eye and the otic vesicle (figure 3.2E). A significant increase of number of embryos that can be imaged during a single experiment (figure 3.2C-D). A significant reduction of the Z-stack size and therefore of the illumination of the samples (figure 3.2A).

In comparison to existing methods we provide a solution that is easy and in-expensive even for non-specialized labs. Also, while similar methods are well suited for high throughput and lower resolutions, ours may also be used for long time-lapse and high resolution imaging.

3.1.2 anaLLzar2D - Automated 2D neuromast analysis and nuclei count

For LL analysis I developed a custom IJ macro script that segments individual cell clusters and the pLLP. From the opening dialog (figure 3.4) the user can choose to count nuclei and / or sort ROIs. If nuclei count is chosen, the macro expects a dual-channel (Ch1: cldnb:lyn-gfp; Ch2: a nuclei label) tiff-file as input. If ROI sorting is selected, segmented CCs are numbered and sorted from left to right instead of top to bottom (IJ’s native sorting method).

- **Membrane label blur** controls the detail of pLLP and CC segmentation, where lower values result in more detail
- **Closing filter** controls how harsh objects are separated from each other
- **Nuclei label blur** controls the details of single nuclei and is evaluated after ground truth data described in section 2.3

![anaLLzar2D opening dialog](image)

**Figure 3.4:** anaLLzar2D opening dialog **checkboxes** Choose to count nuclei and whether ROIs sorting should be applied. **sliders** Choose filter and blurring levels.
CHAPTER 3. RESULTS

Image Analysis

Using ROIs as masks, the nuclei within each ROI are counted with a 2-D maxima finder. However, in their unprocessed form the images are too noisy to get meaningful results. The images therefore have to be smoothened with a blurring filter. To detect the right amount of nuclei, it is necessary to evaluate the distance over which the blurring should be applied. A typical nucleus in the pLLP is about 5 µm in diameter. To determine the right blurring value, a range of 4-6 µm in steps of 0.5 was tested. Figure 3.5 shows a registered maximum Z projected lateral line used for ground truth evaluation.

Figure 3.5: Registration of 2D data. Cluster Segments Registered, MaxIP data with cell cluster segments of the lyn-GFP signal laid upon the DAPI signal. Cell Labels Magenta dots represent the maxima found within each ROI and hence the nuclei labels.

Code Snippets

The [...] symbol indicates code re-use from an earlier instance.

Segmentation The membrane label image is segmented based on optimized filter parameters that were derived from trial and error. After, the macro halts for manual correction.

```plaintext
# Background subtraction
run("Subtract Background...", "rolling = 50");
run("Morphological Filters",
    "operation = Opening element = Disk radius = 20");
run("Gaussian Blur...", "sigma = 5 scaled");

# Thresholding
setAutoThreshold("Moments dark");
setOption("BlackBackground", false);
run("Convert to Mask");

# Particle analysis
run("Analyze Particles...", "size = 250-Infinity exclude add");
roiManager("Show All without labels");
run("Enhance Contrast", "saturated = 0.35");
waitForUser("Check ROIs, correct if necessary");
if (sort) {
    sortROIs();
}
```

52
3.1. QUANTITATIVE IMAGING AND IMAGE ANALYSIS

**Sorting** To sort manually corrected ROIs from left to right, each ROI's position in calculated relatively to total image width.

```plaintext
# Sort ROIs from left to right
function sortROIs() {
  run("Set Measurements...", "centroid redirect = None decimal = 0");
  for (j = 0 ; j < roiManager("count"); j++) {
    roiManager("select", j);
    roiManager("measure");
    x = getResult("X", 0);
    w = getWidth();
    a = x/w;
    roiManager("rename", a);
    run("Clear Results");
  }
  setBatchMode(false);
  roiManager("sort");
  for (j = 0 ; j < roiManager("count"); j++) {
    roiManager("select", j);
    roiManager("rename", j);
    run("Clear Results");
  }
  for (j = 0 ; j < roiManager("count"); j++) {
    roiManager("select", j);
    roiManager("rename", j+1);
    run("Clear Results");
  }
}
```

**Count nuclei** After smoothing the nuclei signal in the DAPI labeled channel, maxima are detected only within each ROI.

```plaintext
# count nuclei within segmented cell clusters
# steps performed on DAPI channel
# count nuclei
run("Gaussian Blur...", "sigma = 0.6 scaled");
rcount = roiManager("count");
for (j = 0 ; j < rcount; j++) {
  roiManager("open", datdir + filename + ".ROIset.zip");
  roiManager("select", j);
  run("Duplicate...", " ");
  run("Enhance Contrast", "saturated = 0.35");
  run("Find Maxima...", "noise = 0 output = [Point Selection]");
  run("Capture Image");
  roiManager("select", j);
  run("Find Maxima...", "noise = 0 output = Count");
  NC = getResult("Count");
  setResult("Nuclei", j, NC);
  setResult("Pos", j, j + 1);
}
```

**Data Analysis**

Comparing the maxima counts of each Gaussian parameter with the Ground Truth gives an indication for false -positives resp. -negatives. In figure 3.6 the relative numbers for each blurring parameter can be seen in
CHAPTER 3. RESULTS

Table 3.2: Blurring parameter for nuclei count

<table>
<thead>
<tr>
<th></th>
<th>4.5</th>
<th>5.0</th>
<th>5.5</th>
<th>6.0</th>
<th>7.0</th>
</tr>
</thead>
<tbody>
<tr>
<td>mean</td>
<td>1.30</td>
<td>1.14</td>
<td>1.04</td>
<td>0.95</td>
<td>0.86</td>
</tr>
<tr>
<td>ratio</td>
<td>0.23</td>
<td>0.17</td>
<td>0.15</td>
<td>0.14</td>
<td>0.14</td>
</tr>
<tr>
<td>std.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*column headers*
blurring parameters

percentage above or below the mean cell count of the ground truth (blue horizon). The red area represents false negatives, the green false positives.

Figure 3.6: Relative difference of maxima counts

To estimate the quality of nuclei detection for each parameter, the ratio of automatically detected and ground truth objects count can be calculated and compared (table 3.2). The closer it is to 1, the better. In summary, maximum performance is achieved at a scaled parameter of 6 \( \mu \text{m} \), with a ratio of 0.95 for the count objects and a standard deviation of 0.14.

3.1.3 anaLLzr2DT - Automated 2D pLLP migration and neuromast deposition analysis through time

For the analysis of the migratory behavior (speed, acceleration) and shape (area, roundness) of the pLLP, as well as for the formation and deposition of the pro-Neuromasts through time, I developed a custom IJ macro script that segments the migrating pLLP and individual cell clusters on each frame of a timelapse. Upon macro execution the opening dialog is presented which is divided in three sections (figure 3.7A).

In the processing section the user may choose which modules are executed.

- **Segmentation** controls whether the images are segmented before measurement. If de-selected the user has to provide segmentation masks separately.
3.1. QUANTITATIVE IMAGING AND IMAGE ANALYSIS

- *Include Cell Clusters* controls whether Cell Clusters should be included in the analysis. If de-selected, only the pLLP will be considered.
- *Registration* controls whether the pLLP should be captured in time and space and saved in a separate stack.
- *Multichannel* controls whether a second channel summary statistics from each ROI at each timepoint should be taken.
  - measurements taken are the mean, standard deviation, minimum and maximum intensity

In the **options** section the user may choose whether the macro should be run in *headless mode* (without showing every single action), whether the input images are timeseries and whether all other windows should be closed upon start of processing.

After confirmation, the user has to enter a date of experiment as an identifier in a second dialog. Furthermore the user is presented the images physical properties pre-filled where the idea here is just a review since this is a major source of mistakes (figure 3.7B). Finally, a third dialog is presented to the user giving an approximate duration and basic instructions (figure 3.7C).

![Figure 3.7: anaLLzR2DT opening dialog](image)

**A** Opening dialog: Main functionality **B** Opening Dialog: metadata setup options **C** Time approximation dialog and basic instructions.

Code-snippets describing the main functionality are described in the next couple of sub-sections. The `[...]` symbol indicates code re-use from an earlier instance.

**Registration** The first module of the macro is pLLP registration in X, Y and cropping in Z. For better segmentation results, first the SNR is enhanced by background subtraction. To rotate the image the pLLPs migrational path is approximated by the position of the first and the last segment, then the image is cropped to a fixed height.
# subtract background
run("Z Project...", "projection = [Average Intensity]");
ZPAVG = getTitle();
if (reg) {
  print("Calculating registration parameters...");
  setSlice(n);
  run("Duplicate...", " ");
  DORG = getTitle();
  imageCalculator("Subtract create", DORG, ZPAVG);
  close(DORG);
  run("Morphological Filters",
  "operation = Closing element=Disk radius = 15");
  REG = getTitle();
  run("Gaussian Blur...", "sigma = 6 scaled");
  run("Enhance Contrast...", "saturated = 0.3 normalize");
  run("8-bit");
  setAutoThreshold("MaxEntropy dark");
  run("Convert to Mask");
}

# analyze segments
run("Analyze Particles...", "size = 150-10000 include exclude add");
rmcount = roiManager("count")-1;
print("rois: "+ rmcount);  
if(roiManager("count") == 1) {
  print("Angle: "+ Angle);
  if (Angle < 0) {Angle = Angle * (-1);}
  if (Angle > 90) {Angle = (180-Angle) * (-1);}
} else {
  print("Angle: "+ Angle);
  run("Rotate... ",
  "angle = " + Angle + " grid = 1 interpolation = Bilinear");
  ZPAVG = getTitle();
  selectWindow(REG);
  run("Rotate... ",
  "angle = " + Angle + " grid = 1 interpolation = Bilinear");
  run("Make Binary");
  REG = getTitle();
}

# cropping
roiManager("reset");
run("Analyze Particles...",
  "size = 150 - 10000 include add");
roiManager("select", 0);
List.setMeasurements;
3.1. QUANTITATIVE IMAGING AND IMAGE ANALYSIS

```python
XRect = List.getValue("X");
YRect = List.getValue("Y");
selectWindow(REG);
getDimensions(width, height, channels, slices, frames);
List.setMeasurements;
height = 120 / sizeX;  # change height of rect here
toUnscaled(YRect);
YRect = YRect - (height / 2);
print("YRectcor: " + YRect);
}

define measures

XRect = List.getValue("X");
YRect = List.getValue("Y");
selectWindow(REG);
getDimensions(width, height, channels, slices, frames);
List.setMeasurements;
height = 120 / sizeX;  # change height of rect here
toUnscaled(YRect);
YRect = YRect - (height / 2);
print("YRectcor: " + YRect);
}

# register
resetMinAndMax();
if (dual) {
    # C1
    selectWindow(ORG);
    if (reg) {
        print(" Registering " + embryoID + "...");
        run("Rotate... ",
            "angle = " + Angle + " grid = 1 interpolation = Bilinear stack");
        makeRectangle(0, YRect, width, height);
        run("Crop");
    }
    # C2
    open(dualdir + dualdirlist[q]);
    dualname = replace(dualdirlist[q], ".tif", ";
    if (reg) {
        run("Rotate... ",
            "angle = " + Angle + " grid = 1 interpolation = Bilinear stack");
        makeRectangle(0, YRect, width, height);
        run("Crop");
    }
    close();
} else {
    selectWindow(ORG);
    if (reg) {
        run("Rotate... ",
            "angle = " + Angle + " grid = 1 interpolation = Bilinear stack");
        makeRectangle(0, YRect, width, height);
        run("Crop");
    } else {
    }
}

ORG = getTitle();
# crop ZPAVG for image calc
selectWindow(ZPAVG);
if (reg) {
    makeRectangle(0, YRect, width, height);
    run("Crop");
}

# background correction
print(" Segmenting " + embryoID + ".RC...”);
getDimensions(width, height, channels, slices, frames);
imageCalculator("Subtract create stack", ORG, ZPAVG);
IC = getTitle();
selectWindow(IC);
print("Bleach correction...");
```

**Segmentation**  After registration of the image follows segmentation. For this we again improve SNR by background correction, following by disconnecting loosely joint segments.
run("Bleach Correction",
    "correction = [Simple Ratio] background = 0");
nslbc = nSlices();
for (j = 1; j < nslbc; j++) {
    setSlice(j);
    run("Morphological Filters",
        "operation = Closing element = Disk radius = 15");
}
run("Images to Stack", "name ="+ ORG +" title = [] use");

# segmentation
selectWindow(MC);
run("Gaussian Blur...", "sigma = 5.5 scaled stack");

# disconnect segments
run("Enhance Contrast...", "saturated = 0.5 normalize process_all");
setSlice(n);
resetThreshold();
setAutoThreshold("MaxEntropy dark");
run("Convert to Mask",
    "method = MaxEntropy background = Dark black");
run("Invert LUT");
run("Fill Holes", "stack");
run("Options...", 
    "iterations = 2 count = 1 pad do = Erode stack");
run("Options...", 
    "iterations = 2 count = 1 pad do = Open stack");
run("Options...", 
    "iterations = 1 count = 1 pad do = Dilate stack");
waitForKeyUser("Check Segmentations");

Analysis  Finally we measure and save the results in the defined names
and directories

for (b = 0; b < orgdirlist.length; b++) {
    # get genotypes and embryoIDs from arrays
    type = types[b];
    embryoID = embryoIDs[b];
    orgname = replace(orgdirlist[b], ".tif", ");
    embryodir = output + File.separator + orgname + File.separator;
    File.makeDirectory(embryodir);

    # open and define binary
    open(bindir+bindirlist[b]);
    BIN = getTitle();

    # open and define original
    if (dual) {
        open(rcdir1 + rcdirc1list[b]);
    } else {
        open(rcdir + rcdirlst[b]);
    }
    RC = getTitle();
    dotIndex = indexOf(RC, ".");
    title = substring(RC, 0, dotIndex);

    # enter 2nd loop to increment over each slice of the time-series
    selectWindow(BIN);  # select binary
    pangles = newArray(nSlices() + 1);
    for (i = 1 ; i <= nSlices(); i++) {
3.1. QUANTITATIVE IMAGING AND IMAGE ANALYSIS

```plaintext
s = nSlices();
setSlice(i);
if (ccs) {
    run("Analyze Particles...",
        "size = 150-10000 include add");
} else {
    run("Analyze Particles...",
        "size = 750-10000 include add");
}

# loop through ROI List
for (j = 0 ; j < roiManager("count"); j++) {
    roimanager("select", j);
    run("Set Scale...",
        "distance = 1 known = 0.0005 pixel = 1 unit = micron");
    List.setMeasurements;
    x = List.getValue("X");
    roiManager("rename", x);
}
run("Properties...",
    "channels = 1 slices = 1 frames = [s] unit = micron pixel_width = [xs]

# Sort ROIs and select last one
roiManager("Sort");
for (j = 0 ; j < roiManager("count"); j++) {
    ccn = roiManager("count")+j;
    if (ccn == roiManager("count")) {
        ccn = "prim";
        roiselect = roiManager("count")-1;
    } else {
        ccn = "CC"+j;
        roiselect = j-1;
    }
    roiManager("select", roiselect);
    roiManager("rename", ccn);
}
rmc = roiManager("count");
m = rmc-1;
selectWindow(Rec);

# Prim registration
run("Select None");
roiManager("Select", m);
sln = getSliceNumber();
run("Enlarge...", "enlarge=6");
run("Fit Ellipse");
run("Duplicate...", "use");
rename(sln);
resetMinAndMax();

# Rotate
List.setMeasurements;
A = List.getValue("Angle");
run("Select None");
if (A < 10) {
    A = A;
} else {
    A = 180-A;
    A = A*(-1);
}
pangles[i] = A;
run("Rotate... ", "angle = [A] grid = 1 interpolation = Bilinear slice");
run("Flip Horizontally");
```
# Measure and save segmented Mask ROI

\[ \text{pLLProis = embryodir + File.separator + "ROIs" + File.separator;} \]
\[ \text{File.makeDirectory(pLLProis);} \]
\[ \text{pLLPxy = embryodir + File.separator + "ROIsXY" + File.separator;} \]
\[ \text{File.makeDirectory(pLLPxy);} \]

roiManager("show none"); # suppress roimanager popping up

roiManager("Select", m);

# Save ROIs and XY coordinates

if (i < 10) {
    slice = d2s(0,0) + d2s(i,0);
    roiManager("save", pLLProis + "s" + slice + ".zip");
    saveAs("XY Coordinates", pLLPxy + "s" + slice + ".txt");
} else {
    roiManager("save", pLLProis + "s" + i + ".zip");
    saveAs("XY Coordinates", pLLPxy + "s" + i + ".txt");
}

# Measure

run("Set Measurements...",
    "area centroid bounding fit shape feret"/
    "/quotesingle.ts1
ds stack redirect = None decimal = 2");

roiManager("measure");

roiManager("reset");

run("Select None");

# Calculate additional variables based on measurements

n = nResults();
r = n-1; # actual RowNumber
r2 = n-2; # RowNumber -1
if (i == 1) {
    X0 = getResult("X");
    Y0 = getResult("Y");
} else {
    X1 = getResult("X", r2);
    X2 = getResult("X", r);
    Y1 = getResult("Y", r2);
    Y2 = getResult("Y", r);
}

# Width of bounding rectangle

W = getResult("Width");

# Calculations (XN = normalized X; LE = Leading Edge)

# Euclidian Distance of X + normalized to offspring 'zero'

if (i == 1) {
    XED = 0;
    XN = 0;
} else {
    XED = sqrt((X2-X1)*(X2-X1)+(Y2-Y1)*(Y2-Y1));
    XN = (X2 - X0) + XED;
}

LE = XN + (W/2); # Leading Edge

T = time * r; # Time interval

setResult("embryo", r, orgname); # set Results

setResult("group", r, type);

setResult("time", r, T);

setResult("deg", r, A);

setResult("X_ED", r, XED);

setResult("X_N", r, XN);

updateResults();

close(BIN, RC);

# Merge registered prim timepoints

setBatchMode("exit and display");

run("Images to Stack", "method = [Copy (top-left)] name = Stack title = [] use");
3.1. QUANTITATIVE IMAGING AND IMAGE ANALYSIS

```plaintext
run("Properties...");
"channels = 1 slices = 1 frames = [s] unit = micron pixel_width = [xs]
run("Flip Horizontally", "stack");
if (dual) {
    # save C1
    saveAs("Tiff", pLLPdir + orgname + "+C01.tif");
    close();
    # open C2
    open(rcdir2 + rcdirc2list[b]);
    resetMinAndMax();
    RC = getTitle();
    dotIndex = indexOf(RC, ".");
    title = substring(RC, 0, dotIndex);
    for (i = 1 ; i <= nSlices(); i++) {
        s = i;
        setSlice(i);
    # Prim registration
        if (i < 10) {
            slice = d2s(0, 0) + d2s(i, 0);
            roiManager("open", pLLProis + "s" + slice + "+.zip");
        } else {
            roiManager("open", pLLProis + "s" + i + "+.zip");
        }
        rmc = roiManager("count");
        m = rmc-1;
        roiManager("Select", m);
        selectWindow(RC);
        run("Enlarge...", "enlarge=6");
        run("Fit Ellipse");
        run("Duplicate...", "use");
        rename(sln);
    # Rotate
        A = pangles[s];
        run("Select None");
        run("Rotate...", "angle = [A] grid = 1 interpolation = Bilinear slice");
        run("Flip Horizontally");
    # select & deselect to remove selected ROIs
        selectWindow(RC);
        run("Select None");
        roiManager("reset");
    }
    # close and merge individual pLLP images into one stack
    close(RC);
    run("Images to Stack", "method = [Copy (top-left)] name = Stack title = [] use");
    run("Properties...", "channels = 1 slices = 1 frames = [s] unit = micron
    run("Flip Horizontally", "stack");
    roiManager("reset");
    # Save Results Table
    run("Input/Output...", "jpeg = 100 gif = -1 file = .txt use_file copy_column copy_row save_column");
    saveAs("results", embryodir + orgname + "+_Results" + "+.txt");
```

3.1.4 anaLLzr3D - Automated 3D single cell segmentation and A.I. analysis in the pLLP

For measurement of the apical index (section 2.2.4) I developed a custom IJ macro script that segments and analyses each cell of a pLLP in 3D. Upon
CHAPTER 3. RESULTS

macro execution the opening dialog is presented which is divided in three sections (figure 3.8A).

In the **processing** section the user has to define the input format as well as in which direction the Z-stack was recorded. Furthermore, the user may choose to have the pLLP registered, save intermediate steps for debugging and to have objects segmented without any restrictions and manual ROI correction.

In the **thresholds** section the user may fine tune segmentation and filter thresholds:

- **Segmentation** controls the segmentation threshold at which the membrane signal is detected and therefore the cell volumes are separated from each other (section 2.3)
- **Min. volume** controls the minimum volume, below which objects are discarded
- **Max. volume** controls the maximum volume, above which objects are discarded

In the **measurements** section, the user may choose whether apical constriction measurement should be applied or not. In case apical constriction measurement is selected, the user may choose from a second dialog box whether A.I. should be measured at an absolute- or relative- distance from the tip. Furthermore, the user has the option to measure from a fit ellipsoid (as done for the A.I. measurement described in section 2.2.4) or rectangle.

**Image Analysis**

For pLLP analyses I developed a custom IJ macro script that recognizes cell boundaries via the fluorescence signal emitted by a membrane tethered eGFP which expression is controlled by the *claudinB* lateral line specific promotor (32). The central IJ tool used to do this is the MorphoLibJ’s(70) **Morphological Segmentation** plugin. The plugin however requires to choose
3.1. QUANTITATIVE IMAGING AND IMAGE ANALYSIS

Figure 3.9: Registration of 3D data. (unregistered) location and orientation of the
unregistered MaxIPs in XY. The red line indicates the angle in degrees from the horizontal
midline. The blue oval indicates registration ROI as determined by the macro. (registered)
pLLPs after XY transformation took place. red circles indicate rosette centers as
detected by the macro based on maximum signal intensity.

for a ‘segmentation threshold’ that determines the quality and the quantity
of segmented objects. This parameter therefore plays an essential role in
the reliability of the analysis results.

Registration The first module of the macro is the registration of the
pLLP in X, Y and cropping in Z. This is accomplished by an initial max-
imum Z projection and blurring of the image, 2-D segmentation using a
minimum threshold and lastly by rotating the segment through the angle
formed by the long axis of the ellipsoid (see section 2.2.4 for more informa-
tion) and the horizon (at 0°). After rotation the image is cropped according
to the obtained ROI, as described before. Additionally, the centers of the
most constricting areas are detected via an intensity based dynamic thresh-
hold and highlighted as magenta circles in figure 3.9.

<table>
<thead>
<tr>
<th>Table 3.3: 3-D Ground Truth image scaling</th>
</tr>
</thead>
<tbody>
<tr>
<td>XY resolution</td>
</tr>
<tr>
<td>Z resolution</td>
</tr>
<tr>
<td>Time difference between Z-planes in Z</td>
</tr>
</tbody>
</table>

Image data In figure 3.10 the fluorescence signal of the three pLLPs used
for the Ground-Truth is shown in a single central cross-section along the
dorso-ventral and the apico-basal axis.
To compare the results between the Ground Truth segments and the seg-
ments obtained from different threshold levels graphically, for a single pLLP
Figure 3.10: cldob:lyn-gfp fluorescence signal in a cross-section of the pLLP (Obj.: 40X APO, scale bar = 100 µm)

the Ground Truth and threshold levels are shown as a composite color image in figure 3.11. By using a green lookup-table (LUT) for the Ground Truth and a magenta LUT for the threshold level, one can readily detect overlapping objects (white), over segmentation (magenta) and under segmentation (green). False positive segments are cells that are not part of the Ground Truth, including those cells that would distort our dataset. False negative segments are cells that are part of the Ground Truth but were not detected by the macro (green cells), excluding those cells impacts the cell count. As one can see, the green cells are randomly distributed, therefore, when averaging a variable for all cells, those cells are less likely to distort our dataset. Only if under segmentation becomes too high will it impact the distribution of values. At Threshold Level 2 (T02) we have the least green cells, but the most magenta cells. At Threshold Level 4 (T04) we have no magenta, but some more green cells. Therefore, to be on the safe side in terms of cell parameters and dataset integrity, for the definition of our Ground Truth T04 would be the best pick.

Code Snippets

The [...] symbol indicates code re-use from an earlier instance.

Registration  First we need to get parameters angle and height for registration. All steps are performed on Z-projected data.

```plaintext
1 # 2D segmentation mask
2 run("Z Project...", "projection = [Max Intensity]");
3 run("Gaussian Blur...", "sigma = 8 scaled");
4 setAutoThreshold("Minimum dark");
5 run("Convert to Mask");
6 run("Select None");
```
3.1. QUANTITATIVE IMAGING AND IMAGE ANALYSIS

Figure 3.11: Graphical comparison of the thresholds tested. Volume renderings have been done with IJ's VolumeViewer

```plaintext
# angle from horizontal midline
run("Analyze Particles...", "include add");
rmcount = roiManager("count")-1;
if(roiManager("count") == 1) {
    roiManager("select", 0);
    run("Fit Ellipse");
    List.setMeasurements;
    Angle = List.getValue("FeretAngle");
    if (Angle < 0) {Angle = Angle * (-1);}
    if (Angle > 90) {Angle = (180 - Angle) * (-1);}
} else {
    roiManager("select", 0);
    roiManager("update");
    List.setMeasurements;
    X1Line = List.getValue("X");
    Y1Line = List.getValue("Y");
    roiManager("select", rmcount);
    List.setMeasurements;
    X2Line = List.getValue("X");
    Y2Line = List.getValue("Y");
    makeLine(X1Line, Y1Line, X2Line, Y2Line);
    List.setMeasurements;
    Angle = List.getValue("Angle");
    if (Angle < 0) {Angle = Angle*(-1);}
    if (Angle > 90) {Angle = (180-Angle)*(-1);}
}
run("Select None");
run("Rotate... ", "angle = " + Angle + " grid = 1 interpolation = Bilinear");

# height to crop image to
roiManager("reset"); # the image was rotated, so we need to get the ROIs again
run("Select None");
run("Make Binary");
run("Erode");
run("Analyze Particles...", "size = 150-10000 include exclude add");
rmcount = roiManager("count") - 1;
if(roiManager("count") == 1) {
```

CHAPTER 3. RESULTS

Transformation  Next we transform our 3D data based on the registration parameters derived from the previous step.

Rosette detection  To analyze cells within rosettes resp. within a certain radius of rosettes we first need to know where the rosettes are. At rosette centers we observe an increase in signal intensity since here the membranes of many cells come together in a very small area. This effect we can use to utilize a maximum finder algorithm together with a threshold that is defined individually for each image.
3.1. QUANTITATIVE IMAGING AND IMAGE ANALYSIS

pointthresh = mean/2.5;
pointthresh = round(pointthresh);
run("Find Maxima...", "noise = " + pointthresh + " output = [Point Selection]");
run("Point Tool...", "type = Dot color = Green size = [Extra Large] label counter = 0");
getSelectionCoordinates(xpoints, ypoints);
roiManager("Add");

# measure intensities along horizontal midline
# [...] registration
RX = lengthOf(xpoints); # collect Arrays
RX = Array.sort(xpoints); # put xpoints in right order
RX = Array.invert(RX);

# fill ypoints with mean values of all y coordinates
Array.getStatistics(ypoints, min, max, mean, stdDev);
meanline = mean;
Array.fill(ypoints, meanline);
RY = ypoints;
getDimensions(width, height, channels, slices, frames);
makeLine(0, meanline, width, meanline, 1);
run("Clear Results");
profile = getProfile();
for (a = 0; a < profile.length; a++) {
    setResult("Value", a, profile[a]);
    updateResults();
}

Segmentation  For image segmentation we use the MorphoLibJ’s Morphological Segmentation plugin. Function calls and arguments are defined in the publication documentation.

# 3D gaussian blur
run("Gaussian Blur 3D...", "x=2 y=2 z=0.5");
resetMinAndMax();

# run segmentation
run("Morphological Segmentation");
selectWindow("Morphological Segmentation");
call("inra.ijpb.plugins.MorphologicalSegmentation.setInputImageType", "border");
call("inra.ijpb.plugins.MorphologicalSegmentation.segment", "tolerance = " + tol + ",
    "calculateDams = true", "connectivity = 6");

# wait till segmentation is done
initTime = getTime();
oldTime = initTime;
while (isOpen("Morphological Segmentation")) {
    elapsedTime = getTime() - initTime;
    newTime = getTime() - oldTime;
    if (newTime > 10000) {
        oldTime = getTime();
    }
    if (newTime == 0) {
        loginfo = getInfo("log");
        loginfo = split(loginfo, " ");
        loginfo = Array.reverse(loginfo);
        loginfo = Array.trim(loginfo, 5);
        loginfo = Array.reverse(loginfo);
        loginfo = split(loginfo[0], ".");
        loginfo = Array.reverse(loginfo);
        loginfo = Array.reverse(loginfo);
        loginfo = loginfo[0];
        if (loginfo == "Whole") {
            call("inra.ijpb.plugins.MorphologicalSegmentation.setDisplayFormat",
                "Catchment basins");
        }
CHAPTER 3. RESULTS

```javascript
32 call("inra.ijpb.plugins.MorphologicalSegmentation.createResultImage");
33 run("Grays");
34 selectWindow("Morphological Segmentation");
35 close();
36 }
37 }
38 run("Properties...", "channels = 1 slices = " + n + " frames = 1 unit =
39 microns pixel_width = " + sizeX + " pixel_height = " + sizeY + " voxel_depth = " + sizeZ);

Filter and clearing  Remove segments below a certain volume threshold defined in the startup dialog and clear blank slices in Z.

1 # erase objects V < vmin and V > vmax
2 run("3D Manager Options", "volume surface compactness fit_ellipse 3d_moments
3 feret centroid (pix) centroid (unit) distance_to_surface centre_of_mass (unit)
4 bounding_box radial_distance surface_contact closest exclude_objects_on_edges_xy
5 sync distance_between_centers = 10 distance_max_contact = 1.80");
6 run("3D Manager");
7 selectWindow(OMap);
8 Ext.Manager3D_AddImage();
9
10 # get number of objects
11 Ext.Manager3D_Count(nb);
12 Ext.Manager3D_MultiSelect();
13
14 # loop through all the objects and erase by filter settings
15 for(k = 0; k < nb; k++) {
16   showStatus("Processing " + k + "+/" + nb);
17   Ext.Manager3D_Measure3D(k, "Vol", V);
18   if (V < vmin) {
19     Ext.Manager3D_Select(k);
20     Ext.Manager3D_Erase();
21     if (V > vmax) {
22       Ext.Manager3D_Select(k);
23       Ext.Manager3D_Erase();
24     }
25   }
26 }
27
28 # clean blank slices from bottom and top
29 getDimensions(width, height, channels, slices, frames);
30 var done = false;
31 for(l = 1; l < slices &!done; l++) {
32   setSlice(l);
33   getStatistics(area, mean, min, max, std, histogram);
34   if(max > 0) {
35     amax = l-1;
36     run("Slice Remover", "first = 1 last = " + amax + " increment = 1");
37     run("Reverse");
38     getDimensions(width, height, channels, slices, frames);
39   for(l = 1; l < slices &!done; l++) {
40     setSlice(l);
41     getStatistics(area, mean, min, max, std, histogram);
42     if(max > 0) {
43       bmax = l-1;
44       run("Slice Remover", "first = 1 last = " + bmax + " increment = 1");
45       run("Reverse");
46     done = true;
47   }
48 }
49 }
50 ```
Apical Constriction  Next we navigate to the calculated relative distance from the apical site and take measurements.

```plaintext
# define Z values
if (rosAC) {
    cellum = rosum;  # I copy pasted the code, so rosette um would be cell um
    Zslice = rosum/sizeZ;
    round(Zslice);
}

# cell Constriction
if (cellAC) {
    aciza = cellum / sizeZ;
    Zslice = aciza;
    round(Zslice);
    round(aciza);
    if (fixAC||symAC) {
        # crop single objects and measure ----------------------
        selectWindow(GMap);
        run("3D Manager Options", "volume surface compactness fit_ellipse 3d_moments
        feret centroid_(pix) centroid_(unit) distance_to_surface centre_of_mass_(unit)
        bounding_box radial_distance surface_contact closest exclude_objects_on_edges_xy
        sync distance_between_centers = 10 distance_max_contact = 1.80");
        run("3D Manager");
        Ext.Manager3D_AddImage();
        # get number of objects
        Ext.Manager3D_Count(nb);
        # create arrays to fill with measurements
        objlabelArray = newArray(nb);
        MajorAngle = newArray(nb);
        ACIMajor = newArray(nb);
        ACIMinor = newArray(nb);
        Dap = newArray(nb);
        Ext.Manager3D_MultiSelect();
        for (k = 0; k < nb; k++) {
            if (k > 0) {
                Ext.Manager3D_AddImage();
            }
            Ext.Manager3D_GetName(k, obj);
            Ext.Manager3D_Centroid3D(k, cx, cy, cz);
            # measure feret
            if (symAC) {
                Ext.Manager3D_Measure3D(k, "Feret", ferr);
                da = ferr * cellum;
                da = round(da);
                if (da == 0) {
                    da = 1;
                }
            }
            Dap[k] = da;
        }
        toString(obj);
        objlabelArray[k] = obj;
        # erase all objects except current(k)
        Ext.Manager3D_SelectAll();
        Ext.Manager3D_Select(k);
        Ext.Manager3D_Erase();
        run("Enhance Contrast...", "saturated = 0.3 equalize process_all");
        run("8-bit");
        run("Crop Label", "label=255 border=5");
        # clear cell stack in Z
        getDimensions(width, height, channels, slices, frames);
        var done = false;
        for(l = 1; l < slices && !done; l++) {
            setSlice(l);
        }
```
getStatistics(area, mean, min, max, std, histogram);
if(max > 0) { // from apical
  smax = l-1;
  run("Slice Remover", "first=1 last="+smax+" increment=1");
  run("Reverse");
getDimensions(width, height, channels, slices, frames);
for(l = 1; l < slices &&!done; l++) { // from basal
  setSlice(l);
  getStatistics(area, mean, min, max, std, histogram);
  if(max > 0) {
    smax = l-1;
    run("Slice Remover", "first = 1 last = "+ smax +" increment = 1");
    run("Reverse");
    done = true;
  }
}
}
}
}
}
}
}
}
}
}
# if fixed ACI
if(symAC) {
  aciza = da / sizeZ;
  db = naci - da;
}
# measure apical
run("Make Binary", "method = Default background=Default calculate black");
setSlice(aciza);
run("Set Measurements...", "area centroid bounding fit feret ts1 redirect = None decimal = 2");
run("Analyze Particles...", "display slice");
MajorAngle[k] = getResult("FeretAngle", 0);
if(MajorAngle[k] > 90) {
  MajorAngle[k] = 90 - (MajorAngle[k] - 90);
}
# get min feret / major / minor
resultsArray = newArray(nResults());
for(p = 0; p < nResults(); p++) {
  resultsArray[p] = getResult("Minor", p);
}
total = 0;
for(p = 0; p < nResults(); p++) {
  total = total + resultsArray[p];
}

Single cell measurement  Get 3D measurements from each object using the 3D Manager plugin
run("3D Manager Options", "volume surface compactness fit_ellipse 3d_moments feret centroid_(pix) centroid_(unit) distance_to_surface centre_of_mass_(unit) bounding_box radial_distance surface_contact closest exclude_objects_on_edges_xy sync distance_between_centers=10 distance_max_contact = 1.80");
run("3D Manager");
Ext.Manager3D_AddImage();
Ext.Manager3D_DeselectAll();
Ext.Manager3D_Measure();
Ext.Manager3D_SaveResult("M", datcelldir + name + ".csv");
Ext.Manager3D_CloseResult("M");
Functions

```javascript
function sliceclear() {
  for (j = 1; j <= n; j++) {
    roiManager("reset");
    setSlice(j);
    run("Analyze Particles...", "include add slice");
    if (roiManager("count") == 0) {
      setSlice(j);
      makeRectangle(0, 0, width, height);
      run("Cut");
    } else if (roiManager("count") == 1) {
      setSlice(j);
      allroi();
      wait(200);
      run("Clear Outside", "slice");
    } else if (roiManager("count") > 1) {
      setSlice(j);
      allroi();
      roiManager("Combine");
      run("Clear Outside", "slice");
    }
  }
}
```

Data Analysis

**Cell Count**  In figure 3.12 the relative numbers for each threshold level can be seen in percentage above or below the mean cell count of the ground truth (blue horizon). Analogous to the graphical inspection, the magenta area represents over segmentation, the green under segmentation.

![Figure 3.12: Relative difference of segment counts](image)

**Apical Constriction**  For a proof of concept of segmentation results and concept around measuring Apical Constriction Index, analog to Harding(2013)(62), 13 DMSO and 15 SU5402 treated embryos were imaged, segmented and processed. The results (figure 3.13) show a strong significant
difference between DMSO und SU5402 treated embryos in both A.I.\textsubscript{Major} and A.I.\textsubscript{Minor}, validating the general concept.

Cell Morphology

*There is always a precision / recall trade-off in detection tasks,\textsuperscript{51} e.g., when we set a lower threshold \( [. . .] \), we can get a higher recall with a lower precision \( [. . .] \), but meanwhile we also get more false-positives in the results*.\textsuperscript{(19)}

Inspecting the cell count unfortunately does not directly tell us how well the cell morphology is conserved at different threshold levels, since at higher threshold levels the cell boundaries are differently determined and eventually not even recognized as such anymore. The volume of a cell is a very robust morphological feature with respect to changes in pose and topology \textsuperscript{(19)}. Therefore, if its volume does not differ significantly at a given threshold level we consider its morphology to be conserved.

In figure 3.14 the cell volumes and their distribution across the different threshold levels can be inspected. The closer the slopes are to the Ground Truth, the stronger they are conserved. The upper graph of figure 3.14 shows the full distribution, where the major differences seem to appear within the 0.4 quantile (red line). To get a more detailed impression, in the lower graph of figure 3.14 only the values *within* the 0.4 quantile are shown.

**Confidence Level** To statistically check how closely related each threshold level (section 3.1.4) sample distribution is compared to the Ground Truth, a Kolmogorov-Smirnov-Test (ks.test) was performed. The ks.test is
3.1. QUANTITATIVE IMAGING AND IMAGE ANALYSIS

Figure 3.14: upper Cell volumes full distribution (red line = .4 Quantile) lower Cell volumes distribution within .4 Quantile
a nonparametric test whose null hypothesis is that the both groups compared were sampled from populations with identical attributes. Therefore, the closer the p-value is to 1, the more similar the tested sample distribution would be to the Ground Truth (figure 3.15).

![Figure 3.15: Volumetric conservation (red line = 5%)](image)

In comparison to the Mann-Whitney test, the ks.test is more sensitive to detect changes in the shape of the distribution than to detect a shift of the median(71). Since we have no reason to assume a significant change in the average cell volume, it would not make much sense to use a test that requires averages. Instead we want to test if there is a change in the shape or skew of the whole distribution, which allows to compare each group on the single cell level (figure 3.14). In summary, the test results suggest a larger deviance in shape of distribution between the Ground Truth and the automatically segmented cells for Threshold levels T02 and T10. For T03 we are already close to rejection of the \( H_0 \) hypothesis (assuming a 5% threshold). Only for T04 and T05 are well within the 5% rejection threshold. However, since we want to be as close to the Ground Truth as possible, we settled for T04.

### 3.1.5 Rosette Detection

The method used for rosette detection is based on a convolutional neuronal network (CNN) and was modified from the “rosette detector” algorithm previously used in the lab and described in (1, 51). Since the former method was technically outdated and since we had new data in which we needed to detect rosettes, we updated the former method to a state-of-the-art CNN using Caffe(72) as a backend. Network configuration and training was done by our collaborators at the institute for Informatics, Albert-Ludwigs-University Freiburg.

The training dataset consists of 17 DMSO- and SU5402- treated embryos each. SU5402 is an inhibitor of FGF receptors and embryos treated with these inhibitors show a strongly reduced number of rosettes (38). In order to train the network, the data had to be labeled manually, which was done in ImageJ by placing multipoint ROIs at the center of the rosettes. The data was then further permuted (rotated and changed in size) to artificially increase the amount of training data and make the detection more robust.
3.2. SHROOM3

Figure 3.16: Example for rosette detection on the training data. left Maximum Z-projected input data. middle heatmap of scores. blue indicates a low score, red a high one. right score map projected onto the input data.

against different kinds of input. Further parameters about the training data is listed in Table 3.4. One example for each is shown in figure 3.16. The main advantages for using a neural network in a task like this are...

1. Objectivity
   • A computer model is not biased in a way that it prefers one outcome over the other. It evaluates based on what it was trained to.
   • Unlike the human brain, once a CNN is trained it is static and does not keep on learning. This promotes reproducibility.

2. Degree of rosette registration
   • The output data are continuous rational numbers (Q) instead of integers (Z) which does not only tell if a rosette is there or not, but also for ‘how much’ (50-100%) it is there.

3. Training is done relatively quick

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3.2 Shroom3

In the previous section a number of methods were presented I developed specifically to perform the analyses on the shroom3 mutant phenotype
which are described in the following section.

3.2.1 Phenotype description

While birth rates follow a distribution of Mendelian inheritance (after genotyping at 3 months of age), homozygous mutant adults seem to be more sensitive to mechanical stress and have a shortened lifespan (~6-9 months). In body shape, shroom3 mutants are not smaller or possess any other striking phenotype (figure 3.17C), however their gill flaps seem to be increased in size, swollen, and not exactly streamlined with the body. This is also evident by an increased frequency of gill flap beating. When looking at the pLLP at different stages (figure 3.17D), they exhibit the same phenotype as the MO injected embryos (I).

Shroom3 heterozygous zebrafish show no phenotypic abnormality. When incrossed, genotyping of two to five days post fertilization (dpf) embryos results in a Mendelian distribution. However, stock-fish that are regen-erated and genotyped by finclip at about 3 months of age show only a rate of 5-10 % of shroom3 homozygous mutants. After another 3 – 6 months those would usually decease too. Therefore, the loss of Shroom3 activity leads to increased mortality for reasons we have not uncovered yet.

3.2.2 Lateral Line Morphometrics

Since in the previous study performed on shroom3 morphants we found a significantly reduced number of rosettes in the pLLP, a reduced number of CCs was expected to be deposited at the end of pLLP migration as well. Instead we found more CCs deposited (figure 3.17E).

Dataset

To analyze this observation quantitatively a dataset was put together consisting of 83 zebrafish embryos fixed at the end of pLLP migration, derived from four different parent pairs. After genotyping, this gave us 26 wildtypes, 31 heterozygous and 26 homozygous mutants for statistical tests. In addition to the number and position of deposited NMs, we also measured the area and the number of nuclei in each deposited CC.

Number and Position of Cell Clusters

Although the embryos were closely staged (± 20 min.), there are always some individual variations in the distance migrated by the pLLP of each. To compare the number of deposited CCs independently of LL length the ratio of LL length over CC count (3.18B) was calculated. Fixation with
3.2. SHROOM3

Figure 3.17: shroom3 mutant phenotype (A–A′) mutation strategy (A) Talen arms (black blocks) bordering a sequence within SD1 including a restriction site for NsiI (indicated back arrows and grey background) (A′) wildtype and mutant allele with 8 bp deletion (B–B′) Amino acid code and protein schematic with functional domains and stop codon (indicated by asterisk) (C–C′) Adult phenotype with closeup to gill flaps (D) pLLP phenotype for three different stages (columns) and different manifestations (rows) (40X WI objective). Arrows indicate epithelial rosettes. (E) LL phenotype at end of migration (10X air objective). Scale bars = 1 mm. MaxIPs for D–E.
PFA may introduce a slight bending of the embryo and mounting introduces a different tilt for each embryo, to correct for uneven LL paths and irregularities in mounting, CC distances are calculated in Euclidean space rather than solely in dimension X. While \textit{shroom3}\textsuperscript{+/+} embryos deposit 6 ± 0.7 CCs, \textit{shroom3}\textsuperscript{-/-} embryos deposit 8 ± 0.9 CCs (figure 3.18A). This difference stays true also when normalizing against length (figure 3.18B).

Even though CC position in individual \textit{shroom3}\textsuperscript{-/-} embryos seems more random, the position of the first deposited CC is mostly conserved (3.19A), \( p \) for difference in position = 0.2). Similarly, the position of the pLLP isn’t significantly changed confirming that Shroom3 activity is not required for migration, as was already shown in \textit{shroom3} morphants (1). While for the remaining CCs an average lag of -50.4 \( \mu \text{m} \) as compared to \textit{shroom3}\textsuperscript{+/+} CC positions is observed, it also seems to increase with later CC positions.

For \textit{shroom3}\textsuperscript{+/+} embryos CC position is mostly conserved through development, however it remains elusive if this is true for \textit{shroom3}\textsuperscript{-/-} embryos too. Figure 3.19A’, shows the kernel density distribution for the CC position without grouping individual positions. At a binwidth of 50 \( \mu \text{m} \), the distribution curves show a high and narrow density at around 350 \( \mu \text{m} \), which is the average location of CC1 (3.19 2A). For the remaining CCs the kernel distribution does, neither for \textit{shroom3}\textsuperscript{+/+} nor for \textit{shroom3}\textsuperscript{-/-} embryos, reveal a precise location. In contrast, if based on CC sequential identity the mean position and standard deviation is calculated, a more explicit pattern emerges which clearly shows the increased count and average frequency (3.19 2A).

This shows that in the absence of Shroom3 activity, more NMs form. This is particularly surprising since the lab had previously shown that \textit{shroom3} knock-down leads to a defect in rosette formation and these rosettes pre-figure the deposited NM.
3.2. **SHROOM3**

Figure 3.19: Cell Cluster positions. **A** Exemplary *shroom3*+/+ and *shroom3*−/− embryos with CCs highlighted. CC0 marks the reference location to compare individual embryos. Scale bar = 1 mm. **A’** Kernel Density Estimate without (KDE) grouping (*n*++ = 162, *n*− = 206). **A’’** Dots = mean positions, bars = standard deviation (*n*max for both ++ and −− = 26).

### Cell Count and Area of CCs

The CC count and position analysis clearly shows that more CCs are deposited in *shroom3* mutant embryos. In order to determine whether these additional clusters are comprised of as many cells as their wild-type counterparts or less, I quantified the number of cells and the size of each clusters in *shroom3*−/− embryos, the cell count per CC was determined by counting DAPI stained cells within CC segments derived from the *cldnb:lyn-gfp* membrane signal (section 2.3). Both the CC cell number and the CC area were reduced by about 6% in *shroom3*−/− embryos so that the density remains unchanged (3.20A). Interestingly, the total number of cells in CC per embryo is increased by 9% (3.20B). Both the sum of CC cells and total LL cells (CC+pLLP) are significantly increased in *shroom3*−/− embryos, while the count of only pLLP cells remains unchanged.
CHAPTER 3. RESULTS

Temperature Rescue

Increasing temperature usually leads to an acceleration in thermodynamics and embryonic development, while lower temperatures slow down development (73). Slowing down development also leads to a reduction in speed of migration and therefore to a reduction in forces that act on cells, cell-cell junctions and organ structures.

For this experiment the hypothesis to test was that it should be possible to at least partially restore the shroom3\(^{+/+}\) number of CCs by giving rosettes a longer time to develop properly. Since embryos develop at different speeds, it is more challenging to exactly stage match them at end of migration. To account for this the CC count was normalized to LL length.

The question was if we could restore wildtype CC counts in shroom3\(^{-/-}\)
3.2. SHROOM3

Table 3.5: Temperature rescue dataset summary

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Table 3.6: Temperature rescue statistics

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embryos when incubating at lower temperatures. Therefore, we need to compare CC counts between T25-- and T30++ respectively p-values between the pairs T30-- vs T30++ (the original) and T25-- vs T30++ (the actual hypothesis). Even though the CC count T25-- cannot be completely restored (~3.2 to ~2.8 in median normalized CC count), the difference between tested pairs shrunk from 0.00000 (T30-- vs T30++) to 0.00210 (T25-- vs T30++) (figure 4D and 3.21B), which is much closer to a rejection of $H_0$. At this point, the question remains open but seems probable that at even lower temperatures the wildtype CC count could be rescued in shroom3−/− embryos.

Summary

Two additional CCs are deposited in shroom3−/− embryos. While each CC is comprised of less cells, in sum there are more cells in embryos deficient for Shroom3 leading to a net increase in cell count of ~9% (31 cells). Lowering the temperature by 5° C was sufficient to partially rescue the number of NMs suggesting that Shroom3 might be required to make the system more robust against stress conditions.

3.2.3 Proliferation

LL morphometric analysis revealed that deposited CCs in shroom3−/− embryos were on average slightly smaller and had fewer cells incorporated. However, due to the additional two cell clusters deposited, the net count is ~9% increased at the end of migration. To test whether this is due to higher proliferative activity a dataset of time-lapse movies (12 h / ΔT = 6 min.)
was generated to count the number of mitoses in a \textit{cxcr4b(BAC):H2BRFP} transgenic background similar to previous proliferation studies in the pLLP (\textit{34, 74}).

\textbf{Datasets}

The live imaging dataset consists of 12 \textit{shroom3}+/+ and 18 \textit{shroom3}−/− embryos with about 75 timepoints for each pLLP and 150 cell clusters. For counting, tracks were created for each proliferative cell on MaxIPs (section 2.2.4). Figure 3.22A shows a \textit{shroom3}+/+ and a \textit{shroom3}−/− LL after 12 h of imaging where each mitotic track is individually colored and overlaid to the image data.

To support and validate this data two more datasets from fixed embryos were generated, one with a EdU (\textit{N = 3, n++ = 26, n-- = 17}) and one with a phospho-Histone marker (\textit{N = 3, n++ = 22, n-- = 29}).

\textbf{Lateral Line Mitoses}

Tracking mitoses in the pLLP does not reveal any difference in proliferation from start (~28 hpf) till about mid of migration 12 hours later (3.22B). For confirmation, this finding was also validated \textit{via} two more additional methods. During the cell cycle genetic material is replicated in S-phase, while in metaphase of mitosis histones are found to be heavily phosphorylated (\textit{75}). With an 5-Ethynyl-2'-deoxyuridine (EdU) assay (\textit{76}), cells in S-phase can be detected (3.22B'). Using a specific phospho-Histone (p-Hist.) antibody cells in meta-phase can be detected. When comparing the EdU index (EdU positive cells over total pLLP cell number) or the mitotic index (p-Hist. positive cells over total pLLP cell number) I could confirm that there is no difference in proliferation in the migrating pLLP. Still, at the end of migration the LL system in \textit{shroom3}−/− embryos does contain 9\% more cells (figure 3.20).

Since individual \textit{shroom3}−/− CCs are smaller, we wondered if there could be compensation mechanisms activated that increases proliferation to restore wildtype CC size once they are deposited. To verify this, CC mitoses were tracked on the same data as before. Other than the pLLP, which can be observed throughout the time-lapse, CCs only begin to \textit{exist} when they are deposited. To normalize for individual CC lifetimes, the number of mitoses per CC is divided by the duration of the total time-lapse minus the time span to when it first appeared.

\textbf{Definition 3.1} (Normalized mitotic rate).

\[
\text{mitoses per CC} \ [n] \over \text{total time} - \text{time of deposition} \ [T]
\]
3.2. SHROOM3

**Figure 3.22:** pLLP proliferation A Tracks of Mitosis. (Image data) Last timepoint of Z-projected time-lapse movie. (Labels) each dots marks a mitotic events. Each color represents one mitosis. Each connection between two dots represents 6 min. B-B’ Mitoses in the pLLP. (B) Count of Mitoses through time (mean±sd) (B’) Proliferation Index at 36 hpf for EdU and phospho Histone labeled pLLPs (B’’) Examples of EdU staining (scale bar = 100 µm) C Mitoses in CCs. B’ and C, the colored bar indicated the median, bars indicate the 95% CI.
CHAPTER 3. RESULTS

This clearly shows that the additional cells in \textit{shroom3}\textsuperscript{−/−} embryos are derived from CC proliferation, rather than pLLP proliferation (3.22C).

Summary

These results show that the additional cells at the end of migration observed in the LL analysis do not stem from an increase in proliferation in the pLLP, but instead from increased proliferation after CCs are deposited.

3.2.4 Rosette Formation and Cluster Deposition

Having shown that the additional CC deposition in \textit{shroom3}\textsuperscript{−/−} embryos is not caused by an over-proliferation in the migrating pLLP, the next step was to have a closer look at the dynamics of rosette formation, in relation to pLLP morphometrics and CC deposition. To test dependencies between different observations and developmental dynamics, variables can be correlated. To do this and to have a coherent dataset to work with, the data of three different analyses on a single set of image data were joined. 

\textbf{First}, to get to know the exact timing of CC deposition, a manual tracking tool (59) was used (figure 3.23, tracking). \textbf{Second}, to deduce pLLP morphometrics an IJ macro was developed (anaLLzr2DT, section 3.1.3) for spatiotemporal registration of the pLLP and to yield information about its speed, area, roundness, etc. (3.23, Registration). \textbf{Third}, to detect rosettes and quantify their weights, a CNN (77) was used on the registered pLLP output from the before mentioned IJ macro (section 3.1.5; figure 3.23 - Detection). Finally, all three datasets were joined by embryo id and timepoint.

Dataset

The image data set analyzed consists of 20 time-lapse movies (11 \textit{shroom3}\textsuperscript{−/−}, 9 \textit{shroom3}\textsuperscript{+/+}). Each time-lapse has a duration of ~20 h (~8 min. interval)\textsuperscript{1}, summing up to ~1650 \textit{shroom3}\textsuperscript{−/−}, and ~1350 \textit{shroom3}\textsuperscript{+/+} timepoints.

\textsuperscript{1}for further details about dataset acquisition see section 2.4 - Proliferation dataset
3.2. **SHROOM3**

**Figure 3.23:** Rosette Formation Joined Datasets. **Tracking** pLLP is marked as no.1. The rest of the CCs is numbered sequentially as they appear. **Registration** The black outline marks the region of interest (ROI) that is the pLLP as it is detected by the anaLLzr2DT. The red line highlights the pLLPs leading edge. **Detection** Each square highlights a detected rosette by the CNN, colors represent rosette weights.

### Cluster Deposition

Figure 3.24A shows a montage of a shroom3\(^{+/+}\) and a shroom3\(^{-/-}\) scenario in cluster deposition (3.5 h / ~20 min. interval). For shroom3\(^{+/+}\) the rosette structure seems tight and two depositions occur in a regular manner. In the shroom3\(^{-/-}\) on the other hand, rosette structure are more fragile with less pronounced rosette centers and four observed depositions. Interestingly, based on the cldn6:lyn-gfp signal the area of constriction seems to be less radially organized but more oriented towards the horizontal midline. Furthermore, the trailing rosettes are significantly smaller and do not seem to separate as clean from the migrating primordium. In addition, for L3 it first seems like two CCs could be deposited, until they merge again to a single CC about 1.5 h later.

On average, as it was shown in section 3.2.2, there’s a significant increase in clusters deposited (3.24B). Also, neither shroom3\(^{+/+}\) nor shroom3\(^{-/-}\) CCs drastically change their position once they are deposited.

### Registration

As the pLLP migrates it moves through space and time. To measure velocity and acceleration, one needs to detect where in space (XY coordinates) the pLLP is located at multiple timepoints. When duplicating the pLLP based on a detection ROI from the whole image, the pLLP is registered in time and space (figure 3.25A).

As shown in figure 3.25B-B’ neither speed nor acceleration drastically differ throughout the complete course of migration. Speed drops from an initial ~75 \(\mu\)m/h to about ~30 \(\mu\)m/h for both shroom3\(^{+/+}\) and shroom3\(^{-/-}\) 17 h later (figure 3.3 A). Similarly, while there is a positive acceleration of almost ~2 \(\mu\)m/h peaking after ~2-3 hours, the remaining two peaks progressively get smaller (3.25A’). While for the area no difference can be
Figure 3.24: Cluster Deposition. **A** *shrm3<sup>+/+</sup>* and *shrm3<sup>−/−</sup>* LL development in comparison. L1 - L4 are deposited CCs. Arrows indicate deposition events. Dotted lines are tracks of rosette to CC transition. **B** Statistics of deposited cluster counts. **C** Change of CC position through time. Each line represents the locally weighted scatterplot smoothing (LOESS) of all CCn positions observed. See section 3.2.4 for a dataset description.
detected (3.25C), interestingly the roundness is on average significantly reduced in shroom3−/− pLLPs (3.25D). This is also evident from the montage in figure 3.24A.

**Rosette Detection**

To quantify the maturity of rosettes within the pLLP a CNN was used which allows to detect objects within images (Region Proposal based CNN). Inherent to object detection and classification by a neural network is that the network not only tells us if an object was detected (of numbers room $x \in [0, \infty] \subset \mathbb{N}$), but also how secure it is - the probability - that the object detected is the right one (of numbers room $x \in [0, 1] \subset \mathbb{R}$) - the weighted detection. Furthermore, the weighted detection was capped a score $\leq 0.5$, since below 0.5 the probability would be to insecure. In our case the network was trained to detect wildtype rosettes, while also presenting pLLPs of embryos treated with a compound that inhibits rosette formation (SU5402) to refine learning. Therefore, if the network detected 1 rosette with a
probability of 0.99, it is very likely that this is a wildtype rosette. However, at different developmental stages and genetic background (e.g. mutants with impaired rosette formation) the pLLP exhibits a different count of rosettes. Therefore it would be wrong to compare just the sum of detections or weighted detections per pLLP, but it is necessary to normalize for the different count of rosettes to get a global figure of pLLP rosettization - the rosettiness.

**Definition 3.2 (rosettiness).**

\[
\frac{\sum \text{Weighted Detections}}{\sum \text{Detections}} = \text{Normalized Weight}
\]

**Figure 3.26:** Quantification of rosettes. A Model of detection and weighted detection in shroom3\(^{+/+}\) and shroom3\(^{-/-}\) embryos B Actual detection in pLLPs of shroom3\(^{+/+}\) and shroom3\(^{-/-}\) embryos at six different timepoints and highlighted detection weight.

The kymographs\(^2\) in figure 3.27A-A’ were generated from pLLPs that were previously registered with the anaLLzr2DT IJ macro (section 3.1.3). After registration, a line was drawn along the horizontal midline (3.25A). Finally, recorded kymographs were turned to false color (blue = low intensity, red = high intensity) for better visual interpretation. In shroom3\(^{+/+}\) the higher intensities are highly concentrated to two to three distinct regions within the migrating pLLP, while in the shroom3\(^{-/-}\) pLLP the higher intensities are more fragmented and overall reduced.

The numbers reveal two very interesting things. First, the medians show that rosette count (3.25B’) is only reduced for the first six hours, while rosettiness is reduced almost throughout the entire 20 hours. While the median rosettiness of shroom3\(^{+/+}\) is at a constant level of about 0.9, shroom3\(^{-/-}\) rosettiness starts out at about 0.5 but then linearly grows until it also reaches levels of about 0.9 at 16-20 hours of migration. These results, together with the finding that we could reduce the number of CCs deposited

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\(^2\)a kymograph is a tool to record position over time
when lowering the temperature, suggests that there should be an interdependence with speed of migration. As for rosette detection, variance is much higher.

Second, while for rosette counts comprised between 2 and 6, half or more of the pLLPs are \textit{shroom3}^+/+, for counts below two and above six there are more \textit{shroom3}^-/- pLLPs (3.25B). Furthermore figure 3.27C shows that at a rosettiness of ~0.7 there are about as many \textit{shroom3}^+/+ as there are \textit{shroom3}^-/- pLLPs, marking this point as a threshold were pLLPs above are more likely to be \textit{shroom3}^+/+ and below more likely to be \textit{shroom3}^-/- . At ~0.1 rosettiness there are almost only \textit{shroom3}^-/- pLLPs, while at ~0.9 rosettiness there are almost only \textit{shroom3}^+/+ pLLPs. In addition, the data distributions shown on the right (3.27B'' and C'') reveals that for the rosettiness the distribution did not actually shift to a lower number, but rather flattened and variance is increased.

\textbf{Correlations}

In the previous sections I have shown that in embryos deficient for Shroom3 more NMs are deposited and that this is not caused by an overproliferation in the pLLP. In this section my goal was to collect detailed information about LL morphometrics and rosette formation to find out what is causing the additional CCs to be deposited.

To do this I combined the results of three different analyses to be able to detect relationships between variables. For my question the most interesting variable to find correlations with is rosettiness. As mentioned in section 3.2.1 when we had a first closer look at the pLLP phenotype we found that some of the clearly as \textit{shroom3}^-/- identified embryos still were reminiscent of the \textit{shroom3}^+/+ phenotype. However, what surprised us the most (due to previous results indicating an important role during rosette assembly for Shroom3), was that a fraction had significantly more CCs deposited. To test the interdependence between rosettiness and CCs deposited quantitatively we set up two hypotheses.

First, when we had a closer look at timelapse movies of the migrating pLLP at a high temporal resolution we had the impression that while in \textit{shroom3}^+/+ embryos the pLLP stalled for a moment during CC deposition, this was not the case in \textit{shroom3}^-/- embryos. Therefore the hypothesis to test was that higher velocity or acceleration lead to a lower degree of rosettiness and \textit{vice versa} (figure 3.28A-B). While velocity is the rate at which an object moves in a certain direction, acceleration tells us the rate at which velocity changes. While most of the \textit{shroom3}^+/+ pLLPs nicely cluster in the top right corner (high rosettiness and velocity, green dots in figure 3.28A), for the \textit{shroom3}^-/- pLLPs there is more of a spread. However, there seems to be a weak relationship between velocity and rosettiness.
CHAPTER 3. RESULTS

Figure 3.27: Rosette counts and weights. A-A’ pLLP kymographs. Length in A’ = 100 µm. (A) shows the signal and line drawn. (A’) shows the signal through time. B- B’’ Rosette detections as whole numbers (Z). (B) Relative shows the ratio of detections in shroom3\(^{+/+}\) and shroom3\(^{-/-}\) (x axis) at each count of detections per pLLP (y axis) throughout time. (B’) Medians shows the median ± standard deviation of detections (y axis) as dots in time intervals of 2 h (x axis). (B’’) KDE shows the density in the distribution of the data. C-C’’ Rosettiness as rational numbers (Q). (B) Relative shows the ratio of rosettiness in shroom3\(^{+/+}\) and shroom3\(^{-/-}\) (x axis) at level of rosettiness per pLLP (y axis) throughout time. At ~70% rosettiness the ratio of shroom3\(^{+/+}\) over shroom3\(^{-/-}\) is equally distributed (B’). Medians shows the median ± standard deviation of rosettiness (y axis) as dots in time intervals of 2 h (x axis). (B’’) KDE shows the density in the distribution of the data. See section 3.2.4 for a dataset description.
which is also reflected by the correlation coefficient. The relationship between acceleration and rosettiness on the other hand seems to be stronger. Again, *shroom3*+/+ pLLPs nicely cluster in the lower right corner (high rosettiness and negative acceleration, green dots in figure 3.28B). Since over the course of migration the pLLP gets slower (3.25B), the average acceleration is negative. For the *shroom3*−/− pLLPs there is also more of a spread, but the trend is more clearly visible which is also reflected by the correlation coefficient.

Second, since we know Shroom3 plays a role in rosette assembly and that embryos deficient for Shroom3 have on average more CCs deposited, we wanted to test if there is a direct link between the two and have statistical certainty. Therefore, the hypothesis to test for figure 3.28B was if lower rosettiness leads to a higher number of CCs deposited and *vice versa*. Since here the end of migration cluster count is one-dimensional (one number per embryo), we had to correlate it to a per embryos summary statistic rosettiness. Here the correlation coefficient is rather high, suggesting there might be an interdependency between pLLP rosettiness and the final pattern of NM deposition. Additionally, the clusters determined are nicely separated from each other and the *shroom3*+/+ and *shroom3*−/− pLLPs seem to form two independent groups.

**Figure 3.28:** Correlation between rosettiness, speed, acceleration and final pattern of NMs. Single dots in the scatterplots represent the variables scatter in x and y, the red line shows a linear model through the point cloud of both groups *shroom3*+/+ and *shroom3*−/−, red digits indicate the correlation coefficient. Black spots with grey circles indicate the centers for clusters calculated on these two sets of data. Clustering was performed hierarchical A Correlation between pLLP rosettiness and speed of migration B Correlation between pLLP rosettiness and acceleration C Correlation between pLLP rosettiness and number of deposited clusters normalized to the length of the lateral line.

### Mechanics

It is very well known that Rho kinases are necessary for mechanic force transmission in various developmental processes (26, 44, 49, 78, 79). Since Rock is also thought to be a key player of the Shroom force mediation pathway in the pLLP, the idea for a mechanic rescue was to titrate dif-
different concentrations of a Rock inhibitor (Rockout, section 2.1.1) and a non-muscle myosin II (NMII) activator (Calyculin, section 2.1.1) to see if this way can confirm the correlation between rosettiness and CC number in an independent manner.

Rock inhibition  Rock2a phosphorylates NMII which is necessary for contraction of the actin network and apical constriction (section 1.2.4). Here we wanted to mimic the \textit{shroom3}^{−/−} phenotype in \textit{shroom3}^{+/+} embryos by inhibiting Rock activity via small molecule Rockout drug treatment. Embryos were incubated at distinct (figure 3.29 legend) concentrations from 24 hpf till end of migration (about 60 hpf). At this stage embryos incubated at higher concentrations displayed a significant reduction in overall body length (3.29A’). Therefore, to see if there is a specific effect on length, LL length was normalized to body length.

Figure 3.29: Titration of Rock inhibitor. A LL lengths are reduced with increased concentrations of Rock inhibitor and B groupwise comparison of the length normalized cluster count per concentration of Rock inhibitor. Rockout treatment at 10 µM does not lead to a significant reduction in LL length but does significantly increase the number of CCs deposited (figure 4B and figure 3.29B), confirming a possible link between lower actomyosin contraction in the p LLP and increased number of deposited NMs.

NMII activation  Downstream of Shroom3 NMII, as a motor protein, is necessary for contraction of the actin network and apical constriction (section 1.2.4). The idea for this experiment was to rescue the \textit{shroom3}^{−/−} phenotype by over-activation of NMII. Unfortunately, even when incubation at 100 nM (the maximum solubility), no significant differences can be detected in LL length (figure 3.30A), cluster count (figure 3.30B) or cluster position (figure 4A).
Table 3.7: Rockout rescue dataset summary

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conc.
concentrations in 0.1% DMSO solution

Table 3.8: NMII activation dataset summary

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conc.
concentrations in 0.1% DMSO solution

Figure 3.30: NMII activation. A LL lengths and B groupwise comparison of the length normalized cluster count.

Summary

I could (1) demonstrate, from fixed embryos (figure 3.18B) as well as here on live imaging data (figure 3.24B), that in shroom3 heterozygous mutants significantly more CCs are deposited, gaining more insight in the deposition process which shows smaller and more fragmented CCs and a more chaotic deposition process overall (figure 3.24A) (2) show a possible interdependence with velocity and acceleration (figure 3.28A-B). Also the area of the pLLP is not affected, but roundness, confirming our impression that the
pLLP in shroom3−/− embryos is more elongated (figure 3.25C-D) (3) show that while the integer number of rosettes in the pLLP is only significantly increased during the first 6-8 hours of migration, the overall rosettiness (rosette integrity per pLLP) is significantly reduced over the whole path of migration (figure 3.27C”). Rosettiness is strongly correlated to the final number of CCs deposited (3.28C), concluding that rosettiness is the key parameter for the future outcome of CC pattern. Interestingly however this conclusion contradicts the previous, and intuitive, assumption that embryos deficient for Shroom3 should not deposit CCs at all. However, in the light of this data Shroom3 is important to strengthen rosette integrity in order to have CCs of the right size deposited. Still, the question is: What determines rosette weight? Velocity and rosettiness are correlated weaker than acceleration and rosettiness are which indicates that the rate at which velocity changes has more of an impact than velocity itself. This finding is also confirmed by an experiment where the CC count of shroom3−/− can be rescued when incubated at lower temperature (section 3.2.2).

3.2.5 Apical Constriction

In the previous analysis I have shown a direct relationship between rosette integrity and CCs deposited, where lower rosette integrity leads to a fragmented rosettes and therefore to an increase in CC deposition frequency. To preserve rosette integrity and for proper CC maturation cells need to apically constrict. To gain further insight into Shroom3 contribution to apical constriction, it is appropriate to have a closer look at rosette conformation on a cellular scale. To do so, first a set of high spatial resolution, volumetric image data was generated (XY = 0.164 µm / pixel; Z = 0.4 µm) that allowed for...

1. a more rigorous inspection of morphological differences, and
2. automated and unbiased single cell 3D reconstruction (section 2.2.4) using a newly developed IJ macro (section 3.1.4). Secondly, to investigate the contribution of various single cell morphometrics to rosette formation an app was developed3 that allows for a convenient handling of large amounts of retrieved data.

3LLMapR - dskleinhans.shinyapps.io/LLmapR
Inhibition of FGFR1 via drug treatment (SU5402, section 2.1.1) results in a concentration dependent loss of morphogenesis and rosette assembly in the pLLP (38, 40, 61). As a proof of principle and to validate previous studies, data of drug treated embryos (20 µM) and appropriate DMSO (0.1% DMSO / E3) controls were included.

**Dataset**

The dataset generated consists of three pLLP stages (3.32), four groups (DMSO, SU5402, shrm3++ and shroom3−/−), 267 pLLPs and 33,163 single cells.

**Shape and Arrangement**

Figure 3.32 illustrates the three developmental stages used for this analysis. Figure 3.33 shows for each of the 3 stages a representative example of pLLP in shroom3+/+, shroom3−/− and SU5402-treated embryo. For shroom3+/+ pLLPs two (3.33A, 32 hpf) to four (3.33A, 40 hpf) areas of constricting regions can be observed (XZ pane), in shroom3−/− its three (3.33A, 36 hpf) to seven (3.33A, 40 hpf), confirming the results of rosette

---

4: Embryo Medium 3, section 2.1.6
Table 3.9: Apical Constriction dataset summary

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*Note: When looking at the central rosette along the width (indicated by arrows in 3.33), YZ panes, it seems like there is no considerable difference between shroom3+++ and shroom3+++ rosettes. In SU5402 treated embryos, neither in XZ nor in YZ constriction can be observed and the cells seem to be mostly columnar in shape. In addition, while constriction seems isotropic in the shroom3+++ rosette center, constriction in the shroom3+++ rosette center cells appears anisotropic.*
3.2. SHROOM3

Figure 3.33: Apical Constriction in 3D. **Stages are in columns, the groups are in rows.** Each panel shows a MaxIP (top left), and orthogonal views along the length (XZ, bottom left) and the width (YZ, right). Arrow annotations in YZ panels indicate points of constriction. Bar annotations in XZ indicate the spread of region of constriction.

**Height**

Earlier attempts for indexing AC include measuring the ‘apical index’ (A.I.) of bottle cells during *X.laevis* gastrulation (80) or for cells of the pLLP (61). For both, the metric describes the ratio of lateral height over apical width which, depending on the situation, is a very coarse and inaccurate description. In the pLLP cells of different forms, volumes and heights can be found. When measuring apical width at a fix distance from the apical site cells with different height but otherwise equal degree of AC turn out to have a very different A.I. (figure 2.3A and A’). If however apical width is measured at a distance relative to the cells height, cells with an equal degree of AC turn out to have the same A.I. (figure 2.3A and A’’). In addition, it remains questionable exactly how to define *apical width* in a two-dimensional cellular cross-section. By introducing apical width measurement along both, the major and the minor axis (section 2.2.4), the improved model for AC and method for automated and unbiased cellular segmentation therefore represents a major advancement.

For another demonstration of the method, figure 3.34 shows a heatmap of cell height, how its distributed in the pLLP and under different conditions. At stages 32 and 36 hpf, height in *shroom3−/−* cells is not as strongly reduced as in SU5402 treated ones and cells in the central region of the pLLP are higher than in the leading, trailing and lateral regions. Therefore this representation also gives an idea which cells are most affected.
Figure 3.34: pLLP terrain maps. To get a more detailed impression about how variables (Height in this case) are distributed within the cells of the pLLP, single cell measurements of pLLPs that are normalized in orientation can be superimposed in a two-dimensional map of organ length and width. Next, cell coordinates are grouped into bins of hexagons and the summary statistic is represented in color. (bottom right) link to the LLMapR web-app.

Apical Index

To get a hold on AC anisotropy we measured the length of the minor and major axis of an ellipsoid fitted to the imaging Z-plane at a certain distance from the cell apex (as described in section 2.2.4 and illustrated in figure 3.35A). The angle formed by the major axis of the ellipsoid and the horizontal midline indicate the orientation of AC (figure 3.35A'). If more cells are constricted along the DV axis, we expect angles closer to 0°. If more cells are constricted along the AP axis, we expect angles closer to 90°. For angle measurements, all angles are normalized to a 90° range, where 0° is along the horizontal midline.

When comparing the ratio of cells in intervals of 15° along a range of 90° it can be shown that there are more $shroom3^{-/-}$ than $shroom3^{+/+}$ cells within the 0-15° interval through all three stages (3.32). Even more, it shows that there are more $shroom3^{+/+}$ than $shroom3^{-/-}$ cells within the remaining intervals – confirming the model of anisotropic constriction along the DV axis.

While reduction in A.I. Major is stronger than the A.I. Minor in $shroom3^{-/-}$ cells (3.35C-C'), both are significantly reduced throughout time – again confirming the increase in constriction an-isotropy. For all three timepoints the A.I. for SU5402 treated embryos is not only the smallest, but also stays at the same level (~2 in the A.I. Major, ~3.4 in the A.I. Minor), therefore it can be seen as a lower threshold. Interestingly, the A.I.s for DMSO, $shroom3^{+/+}$ and $shroom3^{-/-}$ are progressively converging this threshold from earlier to later stages.
3.2. SHROOM3

Summary

In the previous chapter I provided strong indication that the main reason for the additional CCs deposited in Embryos deficient for Shroom3 is reduced rosettiness. Here I wanted to get a more detailed view on how rosette configuration and cell morphology compares between wildtype and mutant. The method developed to quantify AC allows measurement along two orthogonal axes, the major (the longer axis) and a minor (the shorter axis), of an ellipse fit to a cellular cross-section at a distinct distance from the apical site. In addition, the cells orientation in degrees from the horizontal midline is obtained (3.35A-A’).

In summary, AC is significantly reduced in \textit{shroom3}^{-/-} embryos. The results show an increased axial an-isotropy, meaning that AC is more reduced along the major than along the minor axis. In conclusion this means that due to less tight rosette packing and apical constriction the migrational pulling forces are probably accounting for the an-isotropy. In terms of cell shape this means that rosette cells go from a pyramidal geometry, more to a prism geometry (figure 3.36). This is also confirmed by angular measurement which show that in \textit{shroom3}^{-/-} pLLPs there are more cells in the range of 0-15° while in \textit{shroom3}^{+/+} pLLPs more cells are in the range of 30-45°.

Zona occludens-1 (ZO-1) is a peripheral membrane, tight-junction associated protein. Within the apical region of pLLP rosettes FGF filled luminal
structures are developing that are hubs for locally confined morphogen signaling. All cells of a rosette make up and are connected to a single lumen. When visualizing the lumen via an antibody for ZO-1, the lumen appears like a buckyball where each side is the connection to a single cell (40). While in \textit{shroom3}^{+/+} rosettes those structures appear circular in the trailing rosette, in \textit{shroom3}^{−/−} rosettes they appear more oval (supplement figure 2, which supports the theory of an-isotropy. Furthermore, in more leading regions the signal is more fragmented, which supports the theory of micro-rosette structures.

### 3.2.6 Haircell Specification

In the analyses presented so far I have shown that embryos deficient for Shroom3 deposit more CCs, that this is caused by a fragmentation of rosettes (not by overproliferation) and that AC, which is important for CC maturation, is only partially diminished (more in A-P than in D-V axis). Since we also observe a shift in cell geometry from a pyramidal to a more prism geometry, we wanted to know if this had downstream consequences on hair-cell (HC) specification.

As described in section 1.1.3, the shape of a cell has direct impact on its ability to perceive and react to external cues. Here we were interested to test if the reduction in AC (change in cell shape, section 2.2.4) in the pLLP had an impact on HC specification and therefore if the LLs function as a sensory system would still be intact. Atoh1a (section 1.2.3, a TF that gives cells the potential to become sensory HCs), in the pLLP is locally expressed between leader and follower cells and is finally restricted to a single center cell via lateral inhibition (section 1.1.2). Since till today it was not shown why the center cell becomes the HC - here our hypothesis was that rosette assembly (radial organization) respectively AC could play a role.

First, to check if \textit{atoh1a} was still expressed in \textit{shroom3}^{−/−} cells and if there...
were further implications in feedback loops with Notch signaling, an *In Situ* Hybridization (ISH, section 2.2.3) experiment was conducted (probes in section 2.1.8). For 36 and 40 hpf the count of *atoh1a* and *deltaD* expressing cells (3.37) seemed to be increased. Furthermore, especially at 40 hpf, the signal seems to be a lot more fragmented and fuzzier.

Traditional ISH allows for detection of RNA transcripts on a whole embryo scale via a hybridizing anti-sense probe. However, since embryos are fixed, it only allows to analyze single time points. Furthermore, since images are taken in brightfield, quantification of signal intensity is difficult and since what is visualized is a precipitate rather than a signal there is not a clear linear relationship to quantify the amount of expression.

To get access to developmental dynamics we made use of an additional, besides cldnb:lyn-gfp, transgenic construct where tdTomato (81) is expressed under direct control of the *atoh1a* promotor. By simultaneous observation of both fluorophores in a time-lapse setup (3.38B), the strategy was to follow up the dynamics of tdTomato to quantify the expression of *atoh1a* in cell count and intensity of expression.

**Dataset**

For each group, *shroom3*+/+ and *shroom3*−/−, and timepoint the dataset was split into pLLP and CC (table 3.10). Each time-lapse has a duration of 18-20 hours, consists of about 54 timepoints and two channels (488 & 561 nm).
### Table 3.10: Hair-cell specification dataset summary

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#### Atoh1a in the LL

While in both $shroom3^{+/-}$ and $shroom3^{-/-}$ tdTomato signal intensities are rising as soon as CCs are deposited (figure 3.38A), in $shroom3^{-/-}$ pLLPs intensities are on the rise already in the first four hours of migration. Comparing signal intensities in deposited CCs for both, $shroom3^{+/-}$ and $shroom3^{-/-}$ show a similar pattern of increasing tdTomato signal intensity over the course of development but decreasing tdTomato signal intensity at the timepoint of deposition with progressive CC depositions (figure 3.38B). Similarly, when looking at ungrouped CC signal intensity (not per CC, but as one group), no difference can be detected (figure 3.38C). While there is no difference in relative numbers of $atoh1a$ expressing cells within the migrating pLLP throughout time (figure 3.38D), when comparing mean $atoh1a$ expressing cell counts at specific intervals (about the deposition frequency) the $shroom3^{-/-}$ line is always on or above the $shroom3^{+/-}$ line (figure 3.38D’), indicating significantly more $atoh1a$ expressing cells in $shroom3^{-/-}$ pLLPs.

The extrusions to the left in the Kymographs in figure 3.38D are representative to the elongating pLLP during the process of CC deposition. Magenta curves represent expression of $atoh1a$. For the $shroom3^{-/-}$ pLLP, premature expression of $atoh1a$ can be detected. The diagonal lines in the kymographs in figure 3.38D’ represent the moving pLLP along the LL. Vertical curves represent deposited CCs. As for the registered pLLP, Magenta curves represent expression of $atoh1a$.

Deposition and position of the first CC is more independent of Shroom3 (section 3.2.2) than for the remaining CCs. Interestingly during the first four hours, when the first CC is deposited, analysis of hair-cell specification shows that not only the promotor for $atoh1a$ is more active in $shroom3^{-/-}$ embryos (figure 3.38 A and B), but also that more cells are differentiating to become HCs (figure 3.38C and C’). Together with my previous results these results suggest that the reduction in AC and rosettiness in $shroom3^{-/-}$ embryos correlate with a premature activation of $atoh1a$ expression in the pLLP which, in turn, suggests that changes in cell shape might impact cell
Figure 3.38: Hair-cell specification in the LL. **A** Sample of images analyzed. *Upper panel* shows the cldnb:lyn-gfp transgene with segmentation masks highlighted in red. Each structure is labeled with the corresponding label in the tabular data. *Lower panel* shows the atoh1a:Tom transgene with segmentation mask transferred from upper panel. **B and C** Maximum tdTomato intensities in the pLLP and CCs. All CC lifetimes were normalized to the timepoint of deposition (timepoint zero, highlighted by striped vertical line). pLLP intensities are shown to the right (black curve), CC intensities are shown on a negative scale to the left (colored curves). **D** Relative numbers of atoh1a expressing cells to number of timepoints and **D’** mean counts of atoh1a expressing cells. **D-D’** Kymographs along the horizontal midline showing nascent signal of Tom (in magenta) **E** two individual pLLPs that were registered in time and to the leading edge. And **E’** Tom signal during LL development.
fate decision in the pLLP.

**Morphogen Rescue**

As stated in section 1.1.2 and 1.2.4, FGF plays a key role during LL development and, additionally, more recent research shows that levels of FGF activity have direct impact on the number of CCs deposited (40).

Since in embryos deficient for Shroom3 there are significantly more CCs deposited and since we observe a pre-mature activation of Atoh1a (which activates FGF signaling), we wanted to test whether the increase in CCs deposited can be explained by an increase in FGF activity. Inhibition of FGF also reduces LL length (38, 40). Therefore, a titration series experiment with an FGF inhibitor (SU5402) was planned to find a concentration that would not affect the length of the LL but still restore shroom3+/+ levels of CC count.

**Figure 3.39:** Titration of FGF inhibitor. **A** LL lengths are reduced with increased concentrations of FGF inhibitor and **B** groupwise comparison of the length normalized cluster count per concentration of FGF inhibitor.

Concentrations 0.10 and 0.20 \( \mu \text{M} \) do not show a significant reduction in LL length (figure 3.39A), but they also do not exhibit any difference in CC count (figure 4C and figure 3.39B). While the titration worked out well, we reach wildtype levels of CC count only at a concentration of 1 \( \mu \text{M} \) of SU5402, however at this concentration the LL length is also reduced by \( \sim 1 \) mm. Since FGF plays a crucial role in many developmental processes and since SU5402 affects all of these, \( H_0 \) can at this point neither be rejected nor accepted. More precise methods need to be applied.

**atoh1a knockdown**

Since expression of Atoh1a depends on Fgf signaling (40), the hypothesis to test was if premature CC deposition in shroom3+/− embryos is determined by a premature rise in Atoh1a signaling and if Fgf was also sufficient for CC deposition.
Table 3.11: Morphogen rescue dataset summary

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conc.
concentrations in 0.1% DMSO solution

To test this hypothesis, I injected shroom3−/− embryos with an anti-sense morpholino (MO) that had previously been shown to inhibit atoh1a mRNA translation. Furthermore, since many MOs have been shown to induce unspecific, p53-mediated cell death (82), in a second group I co-injected embryos with an MO for p53 (section 2.1.9 for details about MOs). Interestingly injection of atoh1a MO lead to a loss of the first CC, usually deposited at around 400 µm distance from the otic vesicle (figure 3.40A). Furthermore, while median CC counts in atoh1a MO injected shroom3−/− embryos are actually restored to shroom3+/+ levels, CC counts in shroom3+/+ embryos are significantly reduced as well.

Figure 3.40: atoh1a knockdown. A CC mean positions and B groupwise comparison of the length normalized cluster count.

In the expression analysis of atoh1a in shroom3−/− embryos I could show a premature expression of atoh1a in the pLLP. When trying to rescue the phenotype by knocking down atoh1a, neither in shroom3−/− nor in shroom3+/+ embryos the first NM is deposited anymore. Both of these results would in theory argue for an Atoh1a dependent mechanism of CC deposition, however this result has not been reported before and would
Table 3.12: Atoh rescue dataset summary

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Table 3.13: Atoh1a MO rescue statistics

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need further validation. Furthermore, even though an MO for p53 was co-injected, MO injection has been shown to potentially have many more unspecific effects due to the high concentrations usually injected (83). For this reason double mutants (shroom3−/−; atoh1a−−) were generated - which at this timepoints however are not analyzed yet.

Summary

Spatiotemporal expression analysis of atoh1a in shroom3−/− revealed that expression of atoh1a is initiated earlier in shroom3−/− embryos (figure 3.38B-C) and that the number of cells expressing atoh1a within the migrating pLLP is also increased (figure 3.38D'). Reduction of atoh1a expression by inhibition of FGF did not work since inhibition of FGF also significantly affects LL length. Rescue by morpholino injection resulted in a complete absence of the location of the first neuromasts - a result which does not allow for concise conclusion at this point and needs to be repeated in vivo with shroom3; atoh1a double mutants.
Chapter 4

Discussion

4.1 Methods

Due to the pLLP’s relative simplicity (~100 cells) and excellent accessibility for advanced light-microscopes (~1 cell layer beneath the skin), it promises an in toto understanding and complete model of factors influencing its development.

To create an accurate and robust model of a developmental process, it is necessary to have precise, meaningful and accurate data. Furthermore, to test a hypothesis thoroughly, it is beneficial to be able to analyze a single set of image data in different ways which gives the advantage to directly link records in the separate datasets to each other. E.g. with zebrafish you have a problem of exact stage matching. Even if at one point, when starting to live image, the embryos are stage-aligned, developmental speed may differ between embryos which results in a stage mis-alignment at a later point in time. Therefore you need biological and technical replicates to statistically test your hypothesis. This is true if you want to test for an effect in a single feature like the area of an organ. However, if you also want to know about other features that require a different kind of analysis you will have to record another set of image data, again with biological and technical replicates to say that at a given timepoint feature a and b relate to each other in a certain way. Doing so comes at cost of data consistency since (1) the samples are in fact different ones (2) they have to be prepared in a different way and (3) the measuring tool is a different one.

One way to overcome this is standardization and automation which, if accomplished, would allow to generate large datasets of precise measurements. The idea here is to record a set of image data at a sufficient resolution and number of dimensions necessary to conduct the analyses, allowing to connect datapoints from a certain embryo and timepoint. Thereby different
features don’t need to be compared by their means, but directly, which reduces complexity and increases precision. The tools and methods developed for this work have proven easy to adopt even by novice users and very useful in extracting large scale quantitative information.

4.1.1 Standardized mounting

Although efforts have been made to standardize zebrafish embryo mounting and imaging, most protocols were designed for high throughput widefield screening (67–69, 84) rather than high-resolution confocal microscopy (85–87). In recent years a couple of studies tackled specifically this issue by developing standardized mounting methods using 3D printed stamps (66, 67, 88, 89). The solution provided here offers a much more efficient way of mounting zebrafish embryos for high resolution confocal microscopy.

Most methods concerned with specimen mounting are developed by microscopy specialized labs that are working with state-of-the-art microscopes. Usually, the most successful of these developments are those which find a collaboration partner that can provide a specimen that fits the microscope use case. Still, for most labs the microscope technology and mounting technique is not in reach since they are usually expensive and are targeted towards very specific scientific questions. The mounting method described in this thesis targets a gap that previously had not been filled. Today, the most useful and most spread microscope within the developmental biology community is the spinning disc microscope. It allows to record relatively high-resolution time-lapse movies (depending on the size of the sample and number of channels) as well as 3-D imaging of multiple specimen at once (58). Sample preparation for spinning disc microscopy does not differ from preparing samples for standard brightfield imaging. Therefore, for many it is a door opener into advanced microscopy.

For zebrafish, for many years the standard mounting technique was to mount zebrafish embryos on their side without assistance on a flat surface in a rather fast solidifying mounting medium. Since the embryos are not flat on their side they tend to fall over and have to be re-positioned frequently. Furthermore, the researcher needs to be quite skilled to know how to handle the embryo so it moves in a desired direction. Taken together, this limits the number of embryos that can be mounted simultaneously tremendously. Since one can harvest around a hundred eggs from a zebrafish female from a single crossing, for live imaging most of them could not be used or it would be extremely time consuming to mount and image all of them. Therefore, most of the eggs / larvae are not used and are discarded after some time. Since the method is easy to adopt and uses already existing microscopy resources, it increases the data output and time efficiency by a multitude
4.1. METHODS

without investment in new hardware and excessive training. I chose to mount in round dishes because (1) they are used classically and (2) the selection with the manufacturers is more suited for this application. However, round dishes require a manual alignment of the embryonic body axes with the microscope stage axes - which may compromise the use with pre-set positions if the dish is not placed accurately onto the microscope sample holder. By using rectangular dishes together with a rectangular stamp it would be possible to minimize the angular degrees of freedom.

4.1.2 Automated image analysis

Humans depend on vision to orient themselves in their environment. Even though, from a technical perspective, eyesight is not at all a perfect system (blind spot, foveal centralis, farsightedness, shortsightedness, ...), humans usually don’t perceive these shortcomings. This is because vision is mostly just a heavily processed and augmented version of the faulty and blurry data coming from the eye, processed by the brain. The brain on the other hand is a very advanced pattern recognition system (90).

For humans, analyzing image data is a tedious and tiring process. First, one has to be trained to detect certain features in images, then one has to recognize those features in many different images (from different angles, different sizes, ...), sometimes going into the thousands. Furthermore, even though humans can orient themselves in a 3-D space, for most people recognizing image features in 3-D is a spacial challenge. Whether analyzing images in 2-D or 3-D, it demands a high degree of attention which, in order to ensure high quality data, in turn requires the analyst to have regular breaks. Computers are made to process data. Therefore, to overcome the shortcomings of a human image analyst it is instead possible to train a computer program to specifically detect image features.

2D Analysis

The anallyzr2D and anallyzr2DT detect rather simple image features - the deposited Neuromasts, the pLLP and the nuclei within each structure. While this could, even for large datasets, still be done by a human analyst, one also needs to take into account human confirmation bias during an analysis session. Since the algorithm is not subject to fatigue, it is also less error prone.

1Assuming the selection of images analysed is randomized and the analyst doesn’t know what he is looking at, the analyst could still think to be looking at a certain mutant or wildtype and therefore draw a line about a shape more or less generously. The algorithm on the other hand, depending on how its programmed, usually is not biased. Therefore, it generates more neutral and trustworthy measurements.
In addition to the analytical improvement, the anallzr2DT processes and prepares the images for further downstream processing in a way pLLP timelapses could not be analyzed before. In a timelapse of the LL, the pLLP migrates from anterior to posterior, while the embryo still grows. Since this means a shift of the position of the pLLP from timepoint to timepoint, it is hard to detect and analyze cellular dynamics within the pLLP. During processing of the anallzr2DT the migratory dimension as well as sideward movements of the pLLP are eliminated, making it possible to investigate single cell behaviour during migration more thoroughly. Furthermore, while CC count and CC position had been analysed before (40, 41), using this technique allowed us to conduct the most thoroughly and most accurate analysis conducted on cell cluster deposition so far.

### 3D Analysis

The anallyzr3D is able to segment and analyze cells within the pLLP in 3D and measure a cells apical area at a certain distance from the apex. Since the fluorescence signal in some areas of the cell can be very faint (e.g. in basal regions) or very high (e.g. in rosette centers, where many cell apices come together) it is almost impossible for a human analyst to track and detect the cells boundaries correctly. While manual 3D cell segmentation in the pLLP has been done before (62), here I provided a method that allows for segmentation of all the cells of the pLLP and large datasets. Furthermore, for certain measurements and when comparing positions in 3D, a normalized orientation of cells and between specimen is necessary, which is taken care of by the standardized mounting method.

For computational image analysis, the researcher has different options in methodology where each have their pros and cons. (1) For segmentation, traditionally one would use thresholding or a watershed algorithm. For the latter, the image is treated like a topological map, assuming higher gray values represent something topologically high and lower gray values something topologically low, the watershed algorithm then ‘fills’ the lower regions stating from the peak of each maximum until it comes in touch with another filled region (91). (2) Alternatively, one can also train a computer to detect cell boundaries. With this technique, an artificial neural network (part of Machine- and Deep - Learning) is trained to detect image features on a training dataset where the features (cell membranes, nuclei, ...) were already marked by a human analyst. When the network has reached a certain precision through learning, it is asked to find the same features on a test dataset (92) to validate the training results. For the segmentation results, for the moment, neither option has a clear advantage over the other. For my analyses, I wanted to make sure they were usable and reproducible by other people in the field. While there are people working on making
4.1. METHODS

Machine Learning approaches like Deep Learning more approachable to non-technical people (like most biologists are), they still require a very good understanding and knowledge of the subject and its respective implementation (typically programming languages like Python or R). Meanwhile, many biologists are familiar with the image analysis software ImageJ respectively FIJI, which has a rich catalog of actively developed and maintained libraries or update sites. ImageJ again has the ability to record, parameterize and execute macro code, where it is possible to not only incorporate the core features, but also many of the update site plugins. This way, it is possible to compose a macro program that automates even challenging image analysis pipelines and to easily distribute it to a wide audience - instantly able to reproduce the analyses. In addition, this method is much more transparent in terms of how the features are actually extracted and may be forked to develop it further or adjust it to the users own needs.

In addition to visualization, large multidimensional datasets offer the possibility for advanced computational methods such as Machine- and Deep Learning. For example one could label cells of a dataset as either leading, trailing, rosette, lateral, ... etc. to train an Machine learning model on. The model could then be used on unlabeled data to assign the previous learned labels to cells that fit the right parameters, a strategy that has recently been published by J.Hartmann et al. Yet another application example would be to use the Ground Truth image data generated for my studies to train a CNN that potentially would be more robust to data of different quality / resolution. For the future it would be interesting to apply this method on timeplapse movies too.

4.1.3 Rosette Detection

While the anallzr3D actually detects and counts pLLP rosette centers using traditional image processing techniques, it does not quantify the rosettes maturity respectively to which degree it actually resembles a wild-type rosette. An inherent feature of object detection via neural networks is that they tell us how safe it is for an object detected to actually be what it was trained to detect. In other words, neural networks will also detect objects which do not specifically resemble what they know from the training data - but with a lower detection score. While the interpretation of neural network results should be taken cautiously, for my analyses I interpreted the detection score as the rosettiness, or how much on average the pLLP is rosettized.

For classification tasks one can use discriminative or generative models. The main difference here is that (1) generative models will model the posi-

\[\text{\footnotesize 2 provided the necessary computing power} \]

\[\text{\footnotesize 3 to open a parallel branch of development} \]
tives (the distribution or label to detect) as well as the negatives (the part of the data that is not labeled). When asking a generative model to classify data it will look for a boundary where one model becomes more plausible than the other. Hence they are probabilistic. (2) Discriminative models on the other hand focus on the boundary that separates the labeled from the unlabeled data. Hence they are not probabilistic.

The model used for rosette detection is a discriminative one. During the course of training the model is optimized to reach a softmax of 1 which later allows to make a statement if the pattern is more similar to a pattern of one class seen in the training data than to patterns of the other class. The gradation however is not necessarily linear, which makes a statement like ‘the mutant rosette is 50% wildtype’ impossible. However, even for generative models we have the problem of choosing the right set of training data that covers all the aspects and expressions of possibly occurring real patterns. In addition there is the problem of modelling multi-modal distributions. Many questions that cannot be answered conclusively, but do not prevent us from using these models in the hope that they will do something useful and keep us busy for many years to come.

4.2 Shroom3

4.2.1 Lateral Line

Till the end of migration, on average two additional CCs are deposited in shroom3−/− embryos. Proliferation is not increased in the pLLP - but in CCs once they are deposited (section 3.2.3) reaching at ~ 2 dpf wild-type levels of cells per area. Therefore the increase in CC count is unrelated to the amount of proliferation in the pLLP. While I did not quantify the size of CCs directly after deposition, it seems likely that the observation of an increase in proliferation after CCs are deposited is due to a compensatory effect. Interestingly, there are reports showing an inexhaustible hair cell generation (95) which seems to be triggered by an interaction between WNT and Notch Pathways (96–98). The cell count and area per CC show an average reduction of 6% respectively and no difference in density (section 3.2.2).

Conjecture 4.1. The increase in deposited cell clusters is independent of proliferation

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4An important metric in object detection is the softmax-score (also “detection score,” the final result of all weights of the NN). It is a metric that tells about the security of the network how safe it is in its prediction.
4.2. SHROOM3

4.2.2 Rosette Formation

One of the most important and most interesting results of my work is the correlation between rosettiness and CC count as it clearly shows the interdependence between a high median rosettiness and low number of deposited CCs and vice versa - therefore highlighting the role of Shroom3 as a stabilizing factor during NM formation.

**Conjecture 4.2.** The increase in deposited cell clusters is due to a reduction in rosettiness.

As explained in section 1.1.1, cells in the leading region have a more mesenchymal (more loose and more migrating) while cells in the trailing region have a more epithelial character. The cells in the trailing region therefore are more adhesive to their substrate, the basement membrane, while at the same time they are pulled by the more mesenchymal leading cells towards the direction of migration. This results in traction forces in the trailing cells, becoming stronger at times of acceleration and towards the more trailing part, which potentially destabilizes the migrating cluster. To resist these forces, the cells within the cluster need to adhere strongly, which is made sure by Shroom3. Interestingly this also means that there are no external factors regulating the pattern of LL development, but that it is completely controlled by internal factors.

**Conjecture 4.3.** The pattern of deposited Neuromasts is controlled by internal factors of the posterior Lateral Line Primordium.

While the CC count can be derived from images of the LL at end of migration, the rosettiness at this timepoint can not be measured anymore since the pLLP disperses into three terminal NMs. Likewise, the rosettiness can be measured in images during migration - but the final CC count is only revealed at end of migration. One possible solution to this is to first take single timepoint images of the pLLP during migration, leave the embryos embedded in agarose while incubating them till end of migration, and then image them again at end of migration to derive the CC count. While this way one can image more embryos, one only obtains a single measurement for a single timepoint - making the statistics much poorer. This is mostly due to the measurement of rosettiness. As explained, the result of the weight from the rosette detector does not exactly depict a linear relationship between 0 - no rosette and 1 - perfect rosette. Therefore the variance in measurements may be quite high. Taking however the median of a series of rosettiness measurements from the same pLLP reduces this error. Additionally this kind of procedure produces a number of other useful measurements like speed and acceleration, pLLP area and roundness that may be subject to investigate possible interdependencies.
As figure 3.25B-B’ shows, the method allows to measure velocity and acceleration quite accurately (as indicated by the standard error). Especially for that time scale, there are no reports where researchers were able to successfully derive these measurements - even though it was long known that phases of acceleration and deceleration existed (31). One of our hypothesis was that the higher number of CCs may be related to phases of acceleration, since here cells within CCs would get stretched more which in absence of Shroom3 may lead to a more easy and therefore more frequent deposition of CCs. Even though I have shown a there is a relationship between velocity / acceleration and rosettiness I think the approach is not flawless. As shown in figure 3.25B’ acceleration has a wave-like pattern of acceleration and deceleration through time. Therefore, to correlate the measures acceleration and rosettiness precisely it would be necessary to compare acceleration and rosettiness at a certain timepoint and to replicate this wave-like pattern in the correlation coefficient. However, the dataset recorded does not allow for such an analysis. To calculate a correlation one needs at least two data-points that are precisely stage-matched for each timepoint. For a convincing correlation analysis it is necessary to have much more than two datapoints. In general, the more, the better. However, even in wild-type embryos the cluster deposition cycle is not exactly the same throughout time and between embryos. To make things easier it might be sufficient to stage-match just a single deposition cycle, but even here it would be necessary to record a multiple of embryos I recorded for this analysis to find similar deposition cycle patterns.

4.2.3 Apical Constriction

The main learning from the quantification of the apical index is that it is far from easy not only to accurately quantify a single cell 3D morphometric feature, but also to interpret it.

Apical Index

The main findings from this study are that the difference in A.I. between the cells of shroom3+/+ and shroom3−/− pLLPs is more pronounced along the major than the minor. For recap, the A.I. is lateral height over the respective fit ellipsoid axis of a cells cross-section at a relative distance from the apex (section 2.2.4). If the lateral height was 5 µm and major / minor were 2 / 1 µm, then the A.I. major was 2.5 and the A.I. minor 5. The inversion of values represents the reality, where the minor axis is more constricted than the major axis. The axes however do not reflect a specific body axis (anterior - posterior or dorsal - ventral), but essentially the longer (major) and the shorter (minor) axis of the ellipsoid. If the reduction in apical constriction in shroom3−/− embryos would occur
isotropically we would expect a similar increase in axis length (major and minor) as compared to shroom3\(^{+/+}\) controls. At 32 hpf we see about a 10% reduction in both the major and the minor, while at the later stages of 36 and 40 hpf we see a stronger reduction in the major only (section 3.2.5). A main difference between the earliest stage and the later ones is the speed of migration, which reaches its maximum at around 36 - 40 hpf (section 3.2.4). Therefore one possible explanation could be that at lower speed of migration apical constriction is reduced less, while at higher speeds the cells are stretched more strongly along the axis parallel to the direction of migration. Interestingly, this explanation would also concur with a study by Kozlovskaja-Gumbriené et al. (99), who report Shroom3 independent rosette formation at early stages and active Notch signaling.

**Conjecture 4.4.** The reduction in apical constriction is anisotropic. The anisotropcity may be related to speed of migration.

**Cell orientation**

The second finding is the difference in cell radial orientation. A rosette can be divided in four quadrants of 0 - 90°, where 0° represents the anterior - posterior axis (section 3.2.4A'). The measurement taken represents the angle between the ellipse major axis and the horizon, which is oriented along the pLLP's horizontal midline. Interestingly, the results show an increase in cell count in cells oriented along the horizontal midline in shroom3\(^{-/-}\) embryos which suggests a shift of organ morphology from a radial to a more wedge like shape (section 3.2.5). Since a wedge is also anisotropically apically constricted, this is also in line with the first results. In the range of 30 - 45° we see the opposite ratio, where now the count of cells in shroom3\(^{-/-}\) embryos is reduced, suggesting that those cells are under more stress to apically bind to the rosette center. Again, this result is in favor of the theory that Shroom3 acts as a stabilizer against cell stretching and maintenance of radial cell arrangements under extreme conditions.

**Conjecture 4.5.** Organ morphology shifts from a radial, to a more wedge-like shape.

**4.2.4 Hair Cell specification**

In the pLLP, Notch signaling is important for selection and specification of hair-cells (section 1.1.2). For our observation this could mean that (1) either expression of atoh1a is up-regulated due to a compensation mechanism to ensure CC deposition or (2) that the cellular rearrangements lead to a bias in Notch signaling. A phenomenon that in fact has been reported (8). For the latter, the proposed model would be that in each micro-rosette the HC is in contact with less cells due to a more wedge-like organ morphology, but
the amount of Notch ligand stays the same. Therefore, lateral inhibition and feedback is stronger, leading to an increase in expression of \textit{atoh1a} and ultimately to premature CC deposition.

**Conjecture 4.6.** \textit{Expression of atoh1a appears pre-mature and more frequent}

Even though the dataset for hair-cell specification provides a solid base for hypothesis testing, the count of Atoh1a expressing cells shows only a relatively low significance difference. This could be attributed to an imprecision in measurements. However, the same dataset also shows an earlier activity of the \textit{atoh1a} promotor 3.38B-C, which is independent of cell count and makes the results more trustworthy. Furthermore, the control experiment of rescuing the \textit{shroom3}^{-/-} phenotype of more deposited CCs via an \textit{atoh1a} knockdown shows that neither in \textit{shroom3}^{-/-} nor in \textit{shroom3}^{+/+} embryos the first NM is deposited anymore revealing a possibly novel role for Atoh1a that has not been reported before. Arguing - in theory - for an Atoh1a dependent mechanism of CC deposition. Furthermore, even though an MO for p53 was co-injected, MO injection has been shown to potentially have many more unspecific effects due to the high concentrations usually injected (83). For this reason double mutants ( \textit{shroom3}^{-/-}; \textit{atoh1a}--) were generated - which at this timepoints are not analyzed yet.

### 4.2.5 Model

Based on my results the phenotype of the \textit{shroom3}^{-/-} pLLP is defined by a greater amount of rosettes that are smaller and are less pronounced. In this model Shroom3 stabilizes the cell clusters as more cells are integrated and more traction forces are built up. Without this stabilizing force the cell clusters could not aggregate above a certain threshold, which results in 'micro-rosettes.' In those micro-rosettes the cells are more wedge-like shaped and are less densely packed, which results in pre-mature hair cell specification and deposition. The results are summarized and graphically modeled in figure 4.1.
Figure 4.1: Shroom3 dependent rosette formation. First morphogen FGF binds to FGFR1, leading to expression of shroom3, interaction with Rock and actin network contraction through phosphorylated NMII. Without Shroom3 there’s only a partial contraction of the action network which leads to smaller and more rosettes. Altered cellular morphology then leads to a premature expression of atoh1a and hair-cell specification.
Supplement

**Figure 2:** Luminal signaling. The pLLP is indicated by a dark line. Arrows indicate tight-junctions and possibly luminal structures on the apical side. Nuclei are visualized via a TgBAC(cxcr4b:H2B-RFP) transgenic line (section 2.1.7. ZO-1 is made visible via Immunostaining (section 2.1.3. Scalebar = 100 µm

**Figure 3:** Shroom Ortho- and Paralogs. Groups on y-axis. Amino acid chain length indicated on x-axis. Data derived from Uniprot.
**Figure 4:** Collection of rescue experiment CC positions A Calyculin treatment (section 3.2.4) B Rock inhibition (section 3.2.4) C SU5402 treatment (section 3.2.4) D Temperature rescue (section 3.2.4)
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Publications
Statutory Declaration

I herewith declare that I have composed the present thesis myself and without use of any other than the cited sources and aids. Sentences or parts of sentences quoted literally are marked as such; other references with regard to the statement and scope are indicated by full details of the publications concerned. The thesis in the same or similar form has not been submitted to any examination body and has not been published. This thesis was not yet, even in part, used in another examination or as a course performance.

26.10.21, Frankfurt am Main

Date, Place

David Kleinhans
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