Towards 3D in silico modeling of the sea urchin embryonic development

Barbara Rizzi · Nadine Peyrieras

Abstract Embryogenesis is a dynamic process with an intrinsic variability whose understanding requires the integration of molecular, genetic, and cellular dynamics. Biological circuits function over time at the level of single cells and require a precise analysis of the topology, temporality, and probability of events. Integrative developmental biology is currently looking for the appropriate strategies to capture the intrinsic properties of biological systems. The “–omic” approaches require disruption of the function of the biological circuit; they provide static information, with low temporal resolution and usually with population averaging that masks fast or variable features at the cellular scale and in a single individual. This data should be correlated with cell behavior as cells are the integrators of biological activity. Cellular dynamics are captured by the in vivo microscopy observation of live organisms. This can be used to reconstruct the 3D + time cell lineage tree to serve as the basis for modeling the organism’s multiscale dynamics. We discuss here the progress that has been made in this direction, starting with the reconstruction over time of three-dimensional digital embryos from in toto time-lapse imaging. Digital specimens provide the means for a quantitative description of the development of model organisms that can be stored, shared, and compared. They open the way to in silico experimentation and to a more theoretical approach to biological processes. We show, with some unpublished results, how the proposed methodology can be applied to sea urchin species that have been model organisms in the field of classical embryology and modern developmental biology for over a century.

Keywords Morphogenesis · Sea urchin · Cellular dynamics · In vivo imaging · Multiscale digital specimens · In silico modeling

The phenomenological and theoretical reconstruction of multiscale dynamics in animal embryogenesis: a general perspective

In an ideal scenario, morphogenetic events could be understood through a circular path integrating quantitative reconstructions, data analysis, and modeling (depicted in Fig. 1). The bottom-up digital reconstruction and analysis from the sub-cellular scale to the whole embryo level should be combined with a top-down approach. The latter includes the effect of high-level phenotypic features, such as tissue biomechanics on biochemical processes and on the regulation of gene expression. Events at the subcellular level modify the properties of cells and extracellular matrices, and the latter, in turn, modulates the biochemical activity of the cell [13, 74].

Among all possible genetically and epigenetically encoded processes, cells divide, differentiate, and migrate in the context of their self-produced external environment [67, 102]. Cell features over time can be described with a set of parameters including the cell position, cell lineage, volume, surface, shape indexes, local convexity or concavity, nuclear/cytoplasmic ratio, cell neighborhood and surface of contact, cell polarization assessed by the asymmetric distribution of sub-cellular structures, intrinsic motility, and directionality. At the mesoscopic level, tissue patterning depends on cell displacements, cell–cell adhesion, and cell division characteristics. The emergence of the most macroscopic features shapes the whole embryo and provides global mechanical constraints.
Changes in cell and tissue properties can be described by biomechanical models whose behavior depends on initial and boundary conditions given by the cellular environment [135]. A number of recent publications on different model organisms have focused on the mechanical properties of cells and their emergence from the interaction of cells with their environment [41, 46, 65, 66] and the interplay between mechanical constraints and gene expression patterning [13, 74, 114]. The biomechanical approach readily leads to the integration of the genetic and molecular basis of cell activities into the macroscopic deformation of tissues that shape the embryo [20].

Building multiscale models, integrating the biomechanics, and the molecular and genetic dynamics of embryonic development requires the quantitative description of cell behaviors. We review here how the state of the art reflects this paradigm, focusing on the reconstruction of digital specimens from in vivo and in toto imaging.

**From in vivo imaging to modeling to understand animal embryogenesis: the premises**

State of the art imaging techniques combined with the expression of fluorescent proteins provide time lapse recordings of the embryonic development of small transparent animal model organisms with resolution at the cellular level [69, 89]. The tridimensional images can be processed to reconstruct digital specimens composed of labeled volumes or surfaces corresponding to subcellular structures such as cell nuclei and membranes over time, and encompassing the clonal history of each
cell. The digital reconstruction provides many advantages compared with raw 3D + time data. The visualization of biological structures (e.g., nuclei and membranes) in the digital specimen is simpler and can be combined with different color maps to highlight various features, select cells, and assess their clonal history (examples are shown for the sea urchin in Figs. 2 and 3). Even more importantly, digital embryos can be analyzed to quantify features at all scales.

A number of studies in different model organisms have more or less explicitly tackled the reconstruction of digital specimens from live observation. Pioneering work in the late 1970s [23, 128] led to the full reconstruction of the cell lineage tree of Caenorhabditis elegans [129]. This was obtained from the manual annotation of different embryos observed through Nomarski optics and was made possible thanks to the lineage invariance in Caenorhabditis elegans. Then, in the late 1990s, advances in microscopy and computation to handle high-dimensional data brought the first computer-aided cell tracking [124]. The authors used a new 3D multifocal time lapse system [55] to observe the early embryogenesis of Caenorhabditis elegans and the software Simi BioCell provided means for annotating and comparing the lineage trees of different embryos. The synergistic efforts of different disciplines brought further major advances. Developmental biology benefited from breakthroughs in fluorescent protein engineering [14, 16, 84, 85], microscopy imaging systems

Fig. 2 Paracentrotus lividus embryo reconstruction at the 370-cell stage. Raw and reconstructed data displayed with the Mov-It interactive visualization software. Raw data with cell membranes (a) and nuclei (b) staining represented in volume rendering according to the color map displayed bottom right. Colored dots indicate the approximate nucleus center for each cell. Orthoslices in the x y plane for raw membranes (c) and nuclei (d), colored dots as in a,b; in the inset, orthoslices of raw data and their position in the volume indicated by blue lines. e Same as in c, with a cut of the segmented surfaces in addition. f Isosurfaces of segmented membranes. The different cell populations (i.e., small and large micro-meres, Veg2 and Veg1 macromeres, and mesomeres) are displayed in different colors, color map described on the right. a–f Referential displayed bottom left. Scale bar 20 μm (Images from Duloquin, L., and Rizzi B., unpublished)
[34, 60, 107, 127, 130], image processing methods for cell segmentation and cell tracking [69, 90–92], the development of software for computer-aided data processing and visualization [26], and computer hardware for processing high dimensional datasets on computing grids [115].

**From in vivo imaging to modeling to understand animal embryogenesis: reconstructing multiscale digital specimens**

However, the reconstruction of animal embryogenesis from 3D + time imaging data remains a challenging approach, suffering from intrinsic limitations mainly related to the quality of the original data and the simultaneous requirement of automation and precision. The field of image processing has only recently turned to biological images, first dealing with 2D images, then 3D data, and then addressing one more level of complexity with 3D + time processing [93]. But even the most sophisticated algorithms are currently unable to distinguish objects or track them where the human eye cannot provide an accurate benchmark. So, the next breakthroughs in the field will have to originate from the biological side. In vivo 3D + time imaging has to deal with the trade-off between conflicting requirements including spatial and temporal resolution, signal-to-noise ratio, and photo damage. High spatial resolution and signal-to-noise ratio are needed to capture cell position and shape, but high temporal resolution is necessary to achieve single cell tracking. Further constraints are imposed by photo bleaching and photo damage, which limit the acquisition rate and overall imaging duration. In addition, fluorescent staining of subcellular structures in live model organisms is often non-homogeneous and adds artifacts in data acquisition, limiting the quality of the reconstruction, despite the development of sophisticated pre-processing methods [72, 134].

We can expect further improvements in fluorescent staining with more stable and brighter protein variants. Revealing contrasts intrinsic to the tissues such as the generation of harmonics by multiphoton illumination, has also proved to be useful [101]. When the cell density becomes too high, e.g., at late developmental stages, strategies to decrease the image complexity by providing color combinations and mosaic staining should be further developed. On the side of microscopy techniques, there is still room for improvement in depth imaging, spatial and temporal resolution, and signal-to-noise ratio. The recently developed single-plane illumination microscopy seems a valuable alternative to laser point-scanning microscopy, as it definitely improves the temporal resolution and the signal-to-noise ratio. But various imaging artifacts have not been solved yet and single-plane 2-photon excitation, which looks very promising, is still under development by physicists [62].

![Fig. 3 Cell clonal analysis in Paracentrotus lividus digital embryo. Segmentation of cell membranes represented as isosurfaces. a 32-cell stage, b 201-cell stage, c 334-cell stage, and d 545-cell stage. Scale bar 20 μm. e Flat representation of the cell lineage tree. Line segments between branching points represent the cell life time, branching points indicate mitoses. Color map (a–e) given by a set of colors associated with each cell at the 32-cell stage, color propagated along the cell lineage tree and thus revealing cell clones. This representation highlights the fact that very little cell dispersion occurs through the first 10 cell divisions (Reconstructions from live imaging from Rizzi B., unpublished)](image-url)
Except with ideal image data sets, e.g., highly contrasted objects at low density and moving slowly enough, state of the art image processing strategies are still far from providing error-free, fully automated processing of large 3D + time images. Every cell tracking error propagates to its progeny and impairs the reconstruction of cell clonal history. Similarly, shape segmentation inaccuracy adversely affects further analysis of the cells’ characteristic features. Consequently, although protocols for 3D + time imaging of development based either on 2-photon or confocal laser scanning or on selective plane illumination microscopy have been described for several model organisms—Caenorhabditis elegans [44, 97], Ascidians [116–118], Drosophila melanogaster [105, 131], zebrafish [88], zebrafish and Drosophila [64], mouse [37, 100], and avian [73]—the most reliable results still come from manual reconstructions or extensively supervised and corrected data. In this context, the best results have been obtained for Nematoda and Ascidiacea embryos, thanks to their small cell number and largely invariant cell lineage that allows the matching of different specimens with precision at the single cell level (Table 1). Drosophila [35, 36, 68, 71, 80, 87, 109, 131] and zebrafish [63, 82, 101, 137] are more difficult to handle and cell tracking results obtained from state of the art 3D + time imaging data of developing embryos are suitable for average analysis but lack precision for cell clonal analysis. We review below the main results obtained for Nematodases which provides an example of error-free reconstruction of the cell lineage, and for Ascidians which gives an example of the digital representation of cell shapes combined with a quantitative description of cell interactions.

The digital reconstruction of Nematodases

The reconstruction of the cell lineage tree of Caenorhabditis elegans led to the development of dedicated software applications. The first available tools were designed to perform manual tracking and annotations. Worth mentioning, as it was pioneering in the field, is the development of 3D-DIASemb [50] for performing manual tracking, nuclear and membrane segmentation, and cell motion analysis from Nomarski video microscopy imaging of Caenorhabditis elegans up to the 28-cell stage. Simi BioCell, first mentioned in 1997, was used for manual tracking and annotation in Nematoda [58, 59], for combining the lineage tree with gene expression [96], and even lately for investigating cell lineages in the early mouse embryo [94]. Semiautomatic tracking with StarryNite [11] and AceTree [97] (available at http://waterston.gs.washington.edu/) provided the cell lineage tree of the Caenorhabditis elegans embryo up to 350 cells [8]. A digital atlas of the first larval stage (L1) with single-cell resolution was obtained from confocal images of 15 individual worms [78]. Gene expression and cell fate were then investigated [77] with the volume-object annotation system [108]. More recently, NucleiTracker4D (available at http://sourceforge.net/projects/nucleitracker4d/) was used for the semiautomatic tracking of nuclei imaged by laser scanning confocal microscopy of Caenorhabditis elegans development [39]. Further details concerning the cell clonal analysis in Caenorhabditis elegans can be found in [38, 44] and all the available data has been collected by the Nematode community on http://www.wormatlas.org/. It should be noted that the digital reconstruction of Nematodases

### Table 1 Model organisms with their digital representation

| Animal species | Outline | Software utilities | In totto | In vivo | References |
|----------------|---------|--------------------|----------|---------|------------|
| *C. elegans*   | Manual lineage tree extraction and comparison of different embryos. | SIMIBiocell | Yes | Yes | [124] |
|                | Semiautomatic tracking and annotations at the single cell resolution up to 350 cells | StarryNite and AceTree | Yes | Yes | [8, 11, 97] |
|                | Automated analysis of reporter gene expression in *Caenorhabditis elegans* with cellular resolution | StarryNite and AceTree | Yes | Yes | [96] |
|                | Digital nuclear atlas of first larval stage (L1) at single-cell resolution from confocal imaging of 15 individual worms | VANO | Yes | No | [78, 108] |
|                | Analysis of cell fate from single-cell gene expression profiles | VANO | Yes | No | [77] |
|                | Tracking and analysis at the single cell resolution | NucleiTracker4D | Yes | Yes | [39] |
| *C. intestinalis* and *H. roretzi* | Quantitative reconstruction and analysis of digital data at the cellular scale | Imaris and Amira | Yes | No | [132, 133] |
| *C. intestinalis* | Interactive developmental table | – | Yes | No | [57] |
| *Drosophila melanogaster* | Manual segmentation of the tailbud embryo and analysis of its anatomy | Amira | Yes | No | [98] |
| *Zebrafish*    | 3D reconstruction and analysis of gastrulation movement | Imaris | No | Yes | [87, 131] |
|                | Imaging and reconstruction of the first 24 h of development from DLSLM imaging | Matlab | Yes | Yes | [63] |
|                | Imaging and reconstruction of the first 10 cleavages from multiharmonic microscopy imaging | Mov-It | Yes | Yes | [101] |
reconstruction performed so far on Nematoda has focused on displaying the gene regulatory network data according to cell position, but little integration has been achieved with cell behavior, and to our best knowledge, the assessment of cell shape through the segmentation of membrane images has not been addressed.

The digital reconstruction of Asciidiacea

As for the Nematodes, Ascidians exhibit a largely invariant cell lineage and digital embryos have been reconstructed from confocal images of fixed specimens at different stages of development. The most remarkable work has been collected in the Ascidian Network for In Situ Expression and Embryological Data (ANISEED) database [133] and the 3D Virtual Embryo software for visualization and interaction [132], both available through http://aniseed-ibdm.univ-mrs.fr/virtual_embryo.php. The 3D Virtual Embryo has been introduced as a tool for quantifying the geometry of cells and cell–cell contacts in an interactive three-dimensional environment and provides cell volumes, mathematical descriptors of cell shape, and cell–cell contacts. Initially used to describe the first developmental stages of Ciona intestinalis and a late 32-cell stage of Halocynthia roretzi, the concept has been extended to other Ascidian species. The ANISEED database integrating molecular, embryological, and anatomical data in the virtual embryo provides a new paradigm in the field of developmental biology. It should, however, be noted that so far, ANISEED has been based on the segmentation of fixed embryos. Further extending the concept will require the segmentation of live specimens. The Ascidian community has also contributed another interesting initiative. At first, dealing with raw data only, the Four-dimensional Ascidian Body Atlas, with interactive visualization of the development of Ciona intestinalis [57] is available online through http://chordate.bpnli.bio.keio.ac.jp/faba/1.4/top.html. More recently, Hotta et al. achieved the manual segmentation of Ciona intestinalis tailbud embryos, and the interactive visualization of the data is available through 3D pdf files [98].

Reconstructing the multiscale dynamics of the sea urchin embryogenesis

The sea urchin as an animal model in experimental biology: historical perspective

The importance of the sea urchin as a model organism in biological investigations dates back to the nineteenth century and its characteristic features led to cornerstone discoveries in the history of embryology and cell biology. The transparency of the embryo allowed for the first time to observe the fusion between oocyte and sperm during the process of fertilization [52]. In 1892, blastomeres isolated at the 2-cell stage were shown to produce complete sea urchin larvae [24]. Such experimental possibilities made the sea urchin a very valuable model for investigating the components of individual variation. Pioneering studies on teratogenesis carried out by changing the chemical content of the sea urchin embryo helped to found the concept of embryonic induction [51]. The manipulation of double fertilized eggs [9] inspired the chromosome theory of inheritance and opened the way to the understanding of cancer as a genetic disease (English translation in [10]). Further work opened the way to modern Developmental Biology with the exploration of early development including gastrulation [56], cell mechanics underlying morphogenetic movements [18], and the mechanical properties of the sea urchin embryo at early stages of development [53, 54]. During the same period, Gustafson and Kinnander developed a technique allowing the gastrulation of the sea urchin Psammechinus miliaris to be analyzed for the first time, through time-lapse cinematography [42]. A few years later, the sea urchin was selected as a suitable animal model for large-scale analysis of gene expression regulation during early development [12]. More recently (2001), the discovery of cyclin and its role in cell cycle progression in the sea urchin [33] resulted in the award of the Nobel Prize in Physiology and Medicine to Tim Hunt (jointly with Leland H. Hartwell and Paul M. Nurse). Further information about the contribution of the sea urchin animal model to the foundation of paradigms in embryology and developmental biology can be found in [27].

The sea urchin in modern developmental biology

The sea urchin remains a leading model organism in developmental biology [4–6, 32], and the sea urchin community made a major breakthrough in building the architecture of the gene regulatory networks (GRN) underlying the embryo patterning and functional differentiation [86, 110]. The corresponding resources including protocols, database, and data annotation are publicly available (Table 2). However, cell lineage and biomechanics have only been scarcely addressed. The fate of embryonic cells was identified by means of fluorescent dyes [113, 120, 121] and uncaging methods [112], but very little work has been done to achieve in vivo and in toto 3D digital reconstructions from time-lapse imaging, and the resulting data have not yet permitted 3D + time quantitative analysis of cell behavior. Regarding the early development of Paracentrotus lividus, Luengo et al. [81] modeled cell geometries with a Voronoi diagram and compared results with membrane shapes obtained with a viscous watershed segmentation method [136]. Pastor et al. [106] designed a cell tracking method based on the introduction of a 4D tube-structuring element. Rubio et al. [119] merged lineage trees encompassing different periods of the sea urchin development.
using an error-tolerant graph matching technique. So far, however, these methods and the corresponding data have not led to new biological insights or any quantitative assessment of cell behavior. A quantitative 2D analysis of the mitotic gradient observed in different sea urchin species [1, 2, 103, 104] has been performed in the sand dollar *Dendraster excentricus* [25]. The quantitative assessment of cell behavior from time-lapse video microscopy imaging suggested a possible dependency of cell division timing and position on the cell volume between the fourth and the tenth cell cleavage. In this work, cell volumes were calculated indirectly from the measurement of cell diameter under the assumption of a spherical cell shape. Conclusions derived from 2D analysis might, however, be revisited at some point with more precise 3D data. This paradigm has been explored—although the corresponding data did not have single cell resolution—for the 3D reconstruction of adult sea urchin soft tissue from magnetic resonance imaging [138]. The method proposed for the inference of phylogenetic relationships was described together with a simple and attractive way to explore 3D structures interactively in digital specimens embedded in pdf files.

Reconstructing the digital sea urchin from 3D + time in toto imaging

The sea urchin has a number of desirable characteristics, making it a model of choice for in toto time-lapse microscopy imaging of the embryonic and larval stages. The embryo is easily accessible, relatively transparent and small, develops fast, grows to a fairly small number of cells and can be engineered to express fluorescent proteins. After overcoming the problem of the immobilization of the swimming blastula, the embryo can be imaged in toto for extended periods of time by two-photon laser scanning or selective plane illumination microscopy. The sea urchin is the ideal model for making a proof of concept for the reconstruction of the multiscale dynamics of a developing model organism from Deuterostomia, exploiting the unique data available in terms of the GRN architecture and dynamics.

We should, however, emphasize here that the task of long-term 3D + time imaging of the developing sea urchin and automated image processing to achieve cell tracking and shape segmentation is still subject to a number of limitations. These limitations are mainly on the biological side: highlighting structures without ectopic staining, e.g., staining membranes or nuclei without spurious cytoplasmic staining, immobilizing the specimen without mechanical deformation, compensating for light absorption and dispersion and thus loss of signal at depth, resolving individual nuclei throughout the entire volume including at late developmental stages, keeping a temporal resolution high enough to optimize the cell tracking outcome, e.g., ideally keeping the time step smaller than 2 min. Both 2-photon laser scanning and selective plane microscopy imaging are suitable, although they are based on quite different compromises, for the long-term 3D + time imaging of sea urchin embryos. It may soon be demonstrated that for specimens as small as sea urchin embryos, SPIM/DSLM offers the best conditions, provided that specimen mounting is improved compared with what is currently achieved [62].

Over the past few years, the BioEmergences platform (http://www.bioemergences.eu) and its interdisciplinary team have been exploring this paradigm, starting with the systematic reconstruction of the cell lineage tree of *Paracentrotus lividus* from in toto imaging and automated algorithmic processing. The reconstructed digital embryo consists of cell positions and shapes over time and cell clonal history. The corresponding data can be displayed and interactively explored, validated, and annotated with the visualization interface, Mov-IT, [101] (Figs. 2 and 3). The lineages of the different cell populations described in the early embryo according to [19] were highlighted from the 32-cell stage until blastula stages. These reconstructions provide precise measurements hitherto unavailable to model the dynamics at the microscopic scale and reconstruct the overall dynamics of the morphogenetic process at the macroscopic scale as well as possible. This is a most promising approach for the systematic identification of symmetry breaking or intra-individual and inter-individual variation. It would be a major paradigm change in the field of developmental biology, as it uses new theoretical frameworks to revisit classical embryology concepts.

### Table 2 Some online resources from the sea urchin community

| Resource | Web site |
|----------|----------|
| Sea urchin embryology tutorial, the web site is designed to provide material for teaching developmental biology to undergraduates. | http://worms.zoology.wisc.edu/urchins/SUIntro.html |
| Interactive tutorial developed by the Stanford University with support from the National Science Foundation. | http://virtualurchin.stanford.edu/microscope.htm |
| Protocols to study the sea urchin development from the Stanford University | http://www.stanford.edu/group/Urchin/contents.html |
| Protocols from the Swarthmore College | http://www.swarthmore.edu/NatSci/sgilber1/DB_lab/Urchin/urchin_protocols.html |
| The web site of E. Davidson’s laboratory providing an updated version of the BioTapestry Interactive Network Viewer [79] for the GRN of the endomesoderm in *Strongylocentrotus purpuratus* | http://sugp.caltech.edu/endomes/ |
When completed up to late developmental stages, the precise reconstruction of the cell's clonal history with associated shape, volume, and contact changes will contribute to our understanding of morphogenesis and differentiation processes [47–49] and answer a number of open questions. Digital specimens and the corresponding raw data should be made available to the entire community and scientists should all work to enrich a common database and contribute to the validation and correction of cell trajectories and cell shape. By sharing this 3D + time quantitative data for cohorts of individuals in defined genetic and environmental conditions, scientists will have the tools to bring a new perspective to classical questions of embryology.

First, a series of questions would be settled with the analysis of cell clonal history and cell fate such as for ectoderm/endoderm boundaries in the veg1 territory (reviewed by [32]) or the clonal origin of larval structures such as coelomic pouches. A second series of questions concerns the biomechanics of generic morphogenetic movements. The thickening of the vegetal plate during the ingression of the primary mesenchyme blastula stage [22] revealed that the extracellular matrix of the blastula wall was considerably stiffer than the cell layer. The authors then suggested that neither apical constriction nor annular ring contraction could drive invagination while all other hypotheses remained plausible.

Akiyama et al. developed an ad hoc mathematical model to reproduce the first four cleavages of the sea urchin egg, including the asymmetric divisions giving rise to micromeres [3]. The model established the planes of mitosis by assuming a chemotactic motion of centrosomes based on the diffusion of a chemical substance with repellent effects at the animal pole and attractive activity at the vegetal pole.

Ciliberto and Tyson [17] modified a previously developed mathematical model [99] based on molecular interactions underlying early embryonic cell-cycle control to investigate the hypothesis of a mitotic gradient along the animal-vegetal axis. Their results exhibited a good match with experimental data from [83] on Temnopleurus toreumaticus. They concluded that the mitotic wave was primarily attributable to differences in interdivision times in different parts of the embryo, although some subtle details of the wave were probably due to diffusive coupling between neighboring cells.

The transformation of the well-established GRN architecture into a predictive, dynamic Boolean computational model was a major recent breakthrough by the team led by Eric Davidson. This Boolean model computes spatial and temporal gene expression according to the GRN structure established for the embryonic development in the sea urchin. Direct comparison with experimental observations showed that the model predictively computed known gene expression patterns with remarkable spatial and temporal accuracy, thus validating the GRN architecture established so far.

All these models are intended to be both predictive and explanatory and to provide a formal basis to support or refute the intuition. They also tackle specific and limited scales and components of the system. Their integration in a multiscale modeling of morphogenesis will require a huge interdisciplinary and synergistic effort.

Modeling morphogenetic processes in sea urchin development

The long-term goal of integrated modeling of the multiscale dynamics of sea urchin development will take advantage of the partial models established at different scales. The tentative modeling strategies available so far are scarce and much remains to be done in terms of comparing the models with quantitative data. A careful bibliographic search has brought to light four papers, all addressing early developmental steps and dealing respectively with: the biomechanical modeling of the primary invagination [21], the mechanical modeling of division asymmetry [3], the molecular modeling of cell proliferation [17] and the Boolean modeling of GRN dynamics [111].

Finite element modeling was used to discriminate between the possible mechanisms accounting for the primary invagination leading to the formation of the archenteron [21]. Hypothetical mechanisms driving invagination included at the time: apical constriction [7, 31], annular ring contraction [15, 31], cell tracting [15], gel swelling [75], and apicobasal contraction [43]. But each of these mechanisms required different properties of the composite epithelial sheet. Further studies on Strongylocentrotus purpuratus embryos at the mesenchyme blastula stage [22] revealed that the extracellular matrix of the blastula wall was considerably stiffer than the cell layer. The authors then suggested that neither apical constriction nor annular ring contraction could drive invagination while all other hypotheses remained plausible.
interdisciplinary community coming from different perspectives and sharing the same goals is progressing fast. However, the overall conceptual and experimental landscape is still disparate, resembling a huge puzzle whose pieces do not yet fit together. Synergistic efforts by the scientific community are producing public databases of genome and GRN components for many model organisms. State-of-the-art microscopy imaging has made it possible to observe embryo development in small transparent organisms in toto, in vivo, with resolution at the cellular level. Software tools are being developed for visualizing, reconstructing and analyzing 3D + time image data. Computer hardware enables the processing of high-dimensional datasets on computing grids. Digital embryos can be directly manipulated and visualized in 3D to inspect and analyze cell clonal history and cell shape changes and cell–cell interactions over time. Much effort is being dedicated to the development of mathematical models of morphogenesis, based on either GRN architecture or the mechanical properties of tissue, for testing biological hypotheses and predicting the system’s behavior. A new generation of digital resources anticipates the need to integrate multimodal and multiscale data and to provide tools for in silico experimentation. The availability of multimodal and multiscale quantitative data from the in vivo observation and reconstruction of cohorts of individuals should revolutionize practices in the field of developmental biology. Some major bottlenecks remain, related to the current limitations of long-term in vivo imaging and their consequences in terms of automated image processing. Imaging artifacts make 3D + time data from developing organisms difficult to process automatically. And the need for human supervision and time-consuming manual validation and correction is a major issue. The next breakthroughs may come from imaging techniques and may also rely on crowd sourcing for validating and correcting data. We also need new concepts in data analysis and theoretical modeling based on new representations of biological systems, faithful to biological observations but derived from formal descriptions to be shared by a new interdisciplinary community. In this context, the sea urchin is a model organism of choice for unprecedented achievements.

Acknowledgments We thank Louise Duloquin for the time-lapse imaging of live sea urchin embryos and our collaborators at the BioEmergences platform for their contribution. This work was supported by the Centre National de la Recherche Scientifique, the Agence Nationale de la Recherche, the European Commission and the Region Ile de France. We thank Paul Bourigne for helpful discussions and Yannick Kergosien for critical reading of the manuscript. We apologize to the many authors whose work we were unable to cite owing to space constraints.

Open Access This article is distributed under the terms of the Creative Commons Attribution License which permits any use, distribution, and reproduction in any medium, provided the original author(s) and the source are credited.

References

1. Agrell I (1964) Natural division synchrony and mitotic gradients in metazoan tissues. In: Zeuthen E (ed) Synchroly Cell Div. Growth. Interscience, New York, pp 32–67
2. Agrell I (1956) A mitotic gradient as the cause of the early differentiation in the sea urchin embryo. In: Wingstrand K (ed) Zoological Papers in Honour of B. Hanstrom, Lund, pp 27–34
3. Akiyama M, Tero A, Kobayashi R (2010) A mathematical model of cell movement. J Theor Biol 264:84–94. doi:10.1016/j.jtbi.2009.12.016
4. Angerer LM, Angerer RC (2003) Patternning the sea urchin embryo: gene regulatory networks, signaling pathways, and cellular interactions. Curr Top Dev Biol 53:159–198
5. Angerer LM, Angerer RC (2000) Animal-vegetal axis patternning mechanisms in the early sea urchin embryo. Dev Biol 218:1–12. doi:10.1006/dbio.1999.9553
6. Angerer LM, Oleksyn DW, Levine a M et al (2001) Sea urchin goosecoid function links fate specification along the animal-vegetal and oral-aboral embryonic axes. Development 128:439–440
7. Anstrom JA (1992) Microfilaments, cell shape changes, and the formation of primary mesenchyme in sea urchin embryos. J Exp Zool 264:312–322
8. Bao Z, Murray JI, Boyle T et al (2006) Automated cell lineage tracing in Caenorhabditis elegans. Proc Natl Acad Sci U S A 103:2707–2712. doi:10.1073/pnas.0511111103
9. Boveri T (1904) Ergebnisse über die Konstitution der chromatischen Substanz des Zellkerns. Verlag von Gustav Fischer, Jena
10. Boveri T (2008) Concerning the origin of malignant tumours by Theodor Boveri. Translated and annotated by Henry Harris. J Cell Sci 121:1–84
11. Boyle TJ, Bao Z, Murray JI et al (2006) AceTree: a tool for visual analysis of Caenorhabditis elegans embryogenesis. BMC Bioinforma 7:275. doi:10.1186/1471-2105-7-275
12. Britten RJ, Davidson EH (1969) Gene regulation for higher cells: a theory. Science 165:349–357
13. Brouzès E, Farge E (2004) Interplay of mechanical deformation and patterned gene expression in developing embryos. Dev Cell 21:394
14. Buckingham ME, Melilhac SM (2011) Tracing cells for tracking cell lineage and clonal behavior. Dev Cell 21:394–409. doi:10.1016/j.devcel.2011.07.019
15. Burke RD, Myers RL, Sexton TL, Jackson C (1991) Cell movements during the initial phase of gastrulation in the sea urchin embryo. Dev Biol 146:542–557
16. Chudakov DM, Matz MV, Lukyanov S, Lukyanov KA (2010) Fluorescent proteins and their applications in imaging living cells and tissues. Physiol Rev 90:1103–1163. doi:10.1152/physrev.00038.2009
17. Cliberto A, Tyson JJ (2000) Mathematical model for early development of the sea urchin embryo. Bull Math Biol 62:37–59. doi:10.1006/bulm.1999.0129
18. Dan K, Okazaki K (1956) Cyto-embryological studies of sea urchins III. Role of the secondary mesenchime cells in the formation of the primitive gut in the sea urchin larvae. Biol Bull 110:29–42
19. Davidson EH (1989) Lineage-specific gene expression and the regulative capacities of the sea urchin embryo: a proposed mechanism. Development 105:421–445
20. Davidson LA, Von Dassow M, Zhou J (2009) Multi-scale mechanics from molecules to morphogenesis. Int J Biochem Cell Biol 41:2147–2162. doi:10.1016/j.biocel.2009.04.015
21. Davidson LA, Koehl MA, Keller R, Oster GF (1995) How do sea urchins invaginate? Using biomechanics to distinguish between mechanisms of primary invagination. Development 121:2005–2018
22. Davidson LA, Oster GF, Keller RE, Koehl MAR (1999) Measurements of mechanical properties of the blastula wall reveal which hypothesized mechanisms of primary invagination are physically
plausible in the sea urchin Strongylocentrotus purpuratus. Dev Biol 209:221–238
23. Deppe U, Schierenberg E, Cole T et al (1978) Cell lineages of the embryo of the nematode Caenorhabditis elegans. Prot Natl Acad Sci U S A 75:376–380
24. Driesch H (1892) The potency of the first two cleavage cells in echinoderm development. Experimental production of partial and double formations. In: Willer BH, Oopenheimer JM (eds) Foundations of experimental embryology. Hafner, New York
25. Duncan REL, Whiteley AH (2011) The echinoid mitotic gradient: effect of cell size on the micromere cleavage cycle. Mol Reprod Dev 78:686–878. doi:10.1002/mrd.21573
26. Elieci K, Berthold MR, Goldberg IG et al (2012) Biological imaging software tools. Nat Methods 9:697–710. doi:10.1038/nmeth.2084
27. Ernst SG (2011) Offerings from an urchin. Dev Biol 358:285–294. doi:10.1016/j.ydbio.2011.06.021
28. Ettensohn CA (1985) Mechanisms of epithelial invagination. Q Rev Biol 60(3):289–307
29. Ettensohn CA (1984) Primary invagination of the vegetal plate during sea urchin gastrulation. Am Zool 24:571–588. doi:10.1093/icb/24.3.571
30. Ettensohn CA (1985) Gastrulation in the sea urchin embryo is accompanied by the rearrangement of invaginating epithelial cells. Dev Biol 112:383–390. doi:10.1016/0012-1606(85)90410-5
31. Ettensohn CA (1984) An analysis of invagination during sea urchin gastrulation. Yale University
32. Ettensohn CA, Sweet HC (2000) Patterning the early sea urchin embryo. Curr Top Dev Biol 50:1–44
33. Evans T, Rosenthal ET, Youngblom J et al (1983) Cyclin: a protein specified by maternal mRNA in sea urchin eggs that is destroyed at each cleavage division. Cell 33:389–396
34. Fischer RS, Wu Y, Kanchanawong P et al (2011) Microscopy in 3D: a biologist’s toolbox. Trends Cell Biol 21:682–691. doi:10.1016/j.tcb.2011.09.008. Microscopy
35. Fowlkes CC, Eckernode KB, Bragdon MD et al (2011) A conserved developmental patterning network produces quantitatively different output in multiple species of Drosophila. PLoS Genet 7:e1002346. doi:10.1371/journal.pgen.1002346
36. Fowlkes CC, Hendriks CLL, Keränen SVE et al (2008) A quantitative spatiotemporal atlas of gene expression in the Drosophila blastoderm. Cell 133:364–374. doi:10.1016/j.cell.2008.01.053
37. Garcia MD, Udan RS, Hadjantonakis AK, Dickinson ME (2011) Live Imaging of Mouse Embryos. Cold Spring Harb Protoc 2011; pdb.top104. doi:10.1010/pdb.top104
38. Giurumescu CA, Chisholm AD (2011) Cell identification and cell lineage analysis. Meth Cell Biol 106:325–341. doi:10.1016/B978-0-12-454172-8.00012-8
39. Giurumescu CA, Kang S, Planchon TA et al (2012) Quantitative semi-automated analysis of morphogenesis with single-cell resolution in complex embryos. Development 139:4279–4279. doi:10.1242/dev.0986256
40. Glickman NS (2003) Shaping the zebrafish notochord. Development 130:873–887. doi:10.1242/dev.00314
41. Gorflinkiel N, Schamburg S, Blanchard GB (2011) Integrative approaches to morphogenesis: lessons from dorsal closure. Genesis 49:522–533. doi:10.1002/dvg.20704
42. Gustafson T, Kinnander H (1956) Microaquaria for time-lapse cinematographic studies of morphogenesis in swimming larvae and observations on sea urchin gastrulation. Exp Cell Res 11:36–51. doi:10.1016/0014-4827(56)90188-4
43. Gustafson T, Wolpert L (1963) The cellular basis of morphogenesis and sea urchin development. Int Rev Cytol 15:139–214
44. Hardin J (2011) Imaging embryonic morphogenesis in C. elegans. Methods Cell Biol 106:377–412. doi:10.1016/B978-0-12-454172-8.00014-1
45. Hardin J (1989) Local shifts in position and polarized motility drive cell rearrangement during sea urchin gastrulation. Dev Biol 136:430–445. doi:10.1016/0012-1606(89)90268-6
46. Hardin J, Walston T (2004) Models of morphogenesis: the mechanisms and mechanics of cell rearrangement. Curr Opin Genet Dev 14:399–406. doi:10.1016/j.gde.2004.06.008
47. Hardin JD (1987) Archenteron elongation in the sea urchin embryo is a microtubule-independent process. Dev Biol 121:253–262
48. Hardin JD (1987b) The cellular mechanisms and mechanics of archenteron elongation in the sea urchin embryo. University of California
49. Hardin JD, Cheng LY (1986) The mechanisms and mechanics of archenteron elongation during sea urchin gastrulation. Dev Biol 115:490–501
50. Heid PJ, Voss E, Soll DR (2002) 3D-DIA Semb: a computer-assisted system for reconstructing and motion analyzing in 4D every cell and nucleus in a developing embryo. Dev Biol 245:329–347. doi:10.1006/dbio.2002.0631
51. Herbst C (1893) Experimentelle Untersuchungen über den Einfluss der veränderten chemischen Zusammensetzung des umgebenden Mediums auf die Entwicklung der Thiere. II. Wierteres über die morphologische Wirkung der Lithiumsalze und ihre theoretische Bedeutung. Mitt D Zool Station Neapel 11:136–220
52. Hertwig O (1876) Beiträge zur Kenntniss des Bildung und Theilung des tierischen Eies. Morphologisches Jahrbücher 1:349–434
53. Hiramoto Y (1963) Mechanical properties of sea urchin eggs. I. Surface force and elastic modulus of the cell membrane. Exp Cell Res 32:59–75
54. Hiramoto Y (1963) Mechanical properties of sea urchin eggs. II. Changes in mechanical properties from fertilization to cleavage. Exp Cell Res 32:76–89
55. Hirst SN, White JG (1993) Cortical and cytoplasmic fluid flow in early embryonic cells of Caenorhabditis elegans. J Cell Biol 121:1343–1355
56. Horstadius S (1953) Vegetalization of the sea-urchin egg by dimithrophenol and animalization by trypsin and ficin. J Embryol Exp Morphol 1:327–348
57. Hotta K, Mitsuhara K, Takahashi H et al (2007) A web-based interactive developmental table for the Ascidian Ciona intestinalis. Exp Cell Res 324:380–390. doi:10.1016/j.gde.2004.06.008
58. Houthoofd W, Borgonie G (2007) The embryonic cell lineage of the nematode Halicephalobus gingivalis (Nematoda: Cephalobina: Panagrolaimoidea). Nematology 9:573–584
59. Houthoofd W, Jacobsen K, Mertens C et al (2003) Embryonic cell lineage of the marine nematode Pseudolittorina marina. Dev Biol 258:57–69. doi:10.1016/S0012-1606(03)00101-5
60. Huisken J, Swoger J, Del Bene F et al (2004) Optical sectioning deep inside live embryos by selective plane illumination microscopy. Science 305:1007–1009. doi:10.1126/science.1100035
61. Irvine KD, Wieschaus E (1994) Cell intercalation during Drosophila germband extension and its regulation by pair-rule segmentation genes. Development 120:827–841
62. Keller PJ, Schmidt AD, Wittbrodt J, Stelzer EHK (2008) Reconstruction of Zebrafish early embryonic development by scanned light sheet microscopy. Science 322:1065–1069
63. Keller PJ, Schmidt AD, Wittbrodt J, Stelzer EHK (2011) Digital scanned laser light-sheet fluorescence microscopy (DSLM) of zebrafish and Drosophila embryonic development. Cold Spring Harb Protoc 2011:1235–1243. doi:10.1010/pdb.prot065839
reconstruction by 4D-tubular structuring elements. EMBC 2009:970–973. doi:10.1109/EMBS.2009.5333756

107. Pawley J (2006) Handbook of biological confocal microscopy. Springer, New York

108. Peng H, Long F, Myers EW (2009) VANO: a volume-object image annotation system. Bioinformatics 25:695–697. doi:10.1093/bioinformatics/btp046

109. Peng H, Long F, Zhou J et al (2007) Automatic image analysis for gene expression patterns of fly embryos. BMC Cell Biol 8(1):S7. doi:10.1186/1471-2121-8-S1-S7

110. Peter IS, Davidson EH (2011) A gene regulatory network controlling the embryonic specification of endoderm. Nature 474:635–639. doi:10.1038/nature10100

111. Peter IS, Faure E, Davidson EH (2012) Predictive computation of genomic logic processing functions in embryonic development. Proc Natl Acad Sci U S A 109:16434–16442. doi:10.1073/pnas.1207852109

112. Piston DW, Summers RG, Knobel SM, Morril JB (1998) Characterization of involution during sea urchin gastrulation using two-photon excited photorelease and confocal microscopy. Microsc Microanal 4:404–414

113. Ransick A, Davidson EH (1998) Late specification of Veg1 lineages contributing to mesenchyme cell fates in the sea urchin embryo. Dev Biol 160:285–297. doi:10.1006/dbio.1997.8509

114. Rembold M, Looshi F, Adams RJ, Wittbrodt J (2006) Individual cell migration serves as the driving force for optic vesicle evagination. Development 133:1130–1134. doi:10.1126/science.1127144

115. Reuillon R, Chuffart F, Leclaire M, et al. (2010) Declarative task delegation in OpenMOLE. High Performance Computing and Simulation (HPCS), 2010 International Conference on. pp 55–62

116. Robin FB, Dauga D, Tassy O, et al. (2011a) Creating 3D digital replicas of Ascidian embryos from stacks of confocal images. Cold Spring Harb Protoc 2011: pdb–prot065862.

117. Robin FB, Dauga D, Tassy O, et al. (2011b) Time-lapse imaging of live Phallusia embryos for creating 3D digital replicas. Cold Spring Harb Protoc 2011: pdb–prot065847.

118. Robin FB, Dauga D, Tassy O, et al. (2011c) Imaging of fixed Ciona embryos for creating 3D digital replicas. Cold Spring Harb Protoc 2011: pdb–prot065854.

119. Rubio-Guivernau JL, Luengo-Oroz MA, Dulosoquin L et al (2009) Combining sea urchin embryo cell lineages by error-tolerant graph matching. EMBC 2009(2009):5918–5921. doi:10.1109/EMBS.2009.5334851

120. Ruffins SW, Ettensohn CA (1993) A clonal analysis of secondary mesenchyme cell fates in the sea urchin embryo. Dev Biol 160:285–288. doi:10.1006/dbio.1993.1306

121. Ruffins SW, Ettensohn CA (1996) A fate map of the vegetal plate of the sea urchin (Lytechinus variegatus) mesenchyme blastula. Development 122:253–263

122. Sausedo RA, Schoenwolf GC (1993) Cell behaviors underlying notochord formation and extension in avian embryos: quantitative and immunocytochemical studies. Anat Rec 237:58–70. doi:10.1002/ar.1092370107

123. Sausedo RA, Schoenwolf GC (1994) Quantitative analyses of cell behaviors underlying notochord formation and extension in mouse embryos. Anat Rec 239:103–112. doi:10.1002/ar.1092390112

124. Schnabel R, Hutter H, Moerman D, Schnabel H (1997) Assessing normal embryogenesis in Caenorhabditis elegans using a 4D microscopy: variability of development and regional specification. Dev Biol 184:234–265. doi:10.1006/dbio.1997.8509

125. Schoenwolf GC, Alvarez JS (1989) Roles of neuroepithelial cell rearrangement and division in shaping of the avian neural plate. Development 106:427–439

126. Shook D, Keller R (2003) Mechanisms, mechanics and function of epithelial–mesenchymal transitions in early development. Mech Dev 120:1351–1383. doi:10.1016/j.mod.2003.06.005

127. So PTC, Dong CY, Masters BR, Berland KM (2000) Two-photon excitation fluorescence microscopy. Annu Rev Biomed Eng 2:399–429

128. Sulston JE, Horvitz HR (1977) Post-embryonic cell lineages of the nematode, Caenorhabditis elegans. Dev Biol 56:110–156

129. Sulston JE, Schierenberg E, White JD, Thomson JN (1983) The embryonic cell lineage of the nematode Caenorhabditis elegans. Dev Biol 100:64–119

130. Sun C-K, Chu S-W, Chen S-Y et al (2004) Higher harmonic generation from 3-D Images from light-sheet-based fluorescence microscopy. Nat Protoc 4:1397–1412. doi:10.1038/nprot.2009.130

131. Supatto W, McMahon A, Fraser SE, Stathopoulos A (2009) Quantitative imaging of collective cell migration during Drosophila gastrulation: multiphoton microscopy and computational analysis. Nat Protoc 4:1397–1412. doi:10.1038/nprot.2009.130

132. Tassy O, Daiana F, Hudson C et al (2006) A quantitative approach to the study of cell shapes and interactions during early chordate embryogenesis. Curr Biol 16:345–358. doi:10.1016/j.cub.2005.12.044

133. Tassy O, Dauga D, Diagn A et al (2010) The ANISEED database: digital representation, formalization, and elucidation of a chordate developmental program. Genome Res 20(10):1459–1568

134. Temeirac-Ott M, Ronneberger O, Ochs P et al (2012) Multiview deblurring for 3-D Images from light-sheet-based fluorescence microscopy. IEEE T Image Process 21:1863–1873

135. Trier SM, Davidson LA (2011) Quantitative microscopy and imaging tools for the mechanical analysis of morphogenesis. Curr Opin Genet Dev 21:664–670. doi:10.1016/j.gde.2011.08.005

136. Vachier C, Math CDM (2005) The viscous watershed transform. J Math Imaging Vis 22:251–267

137. Zanella C, Campana M, Rizzi B et al (2010) Cells segmentation from 3-D confocal images of early zebrafish embryogenesis. IEEE T Image Process 19:770–781. doi:10.1109/TIP.2009.2033629

138. Ziegler A, Faber C, Mueller S, Bartolomaus T (2008) Systematic comparison and reconstruction of sea urchin (Echinoidea) internal anatomy: a novel approach using magnetic resonance imaging. BMC Biol 6:33. doi:10.1186/1741-7007-6-33