Research at a Glance 2014

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Introduction
Four decades ago, the European Molecular Biology Laboratory’s founding nations set out a vision to create a centre of excellence where Europe’s best brains would come together to conduct basic research in molecular biology and related fields.

This year marks EMBL’s 40th anniversary and the laboratory’s founding principles – scientific independence, opportunity for creative young thinkers, interdisciplinarity and collaboration – remain at the heart of our extensive contributions to the life sciences. Over the years EMBL – Europe’s only intergovernmental laboratory in the life sciences – has become the flagship of European molecular biology, spread across five sites in Heidelberg, Grenoble, Hamburg, Hinxton and Monterotondo.

EMBL is continuously ranked as one of the top scientific institutions worldwide but, as well as pursuing research of the highest quality, we also offer a wide range of services to European scientists, most notably in bioinformatics and structural biology; drive research, innovation and progress through technology development and transfer; and have a long-standing tradition of delivering outstanding courses, conferences and workshops to participants from all over the world. These activities would be difficult or even impossible to pursue for national laboratories and are complemented by an extensive outreach programme and significant engagement in discussions of science policy and strategy at the European level. The synergy between our diverse missions and the stringent external quality control to which we submit all our activities regularly, allows EMBL to achieve world-class standards.

In Research at a Glance you will find a concise overview of the work of our research groups and facilities. Science at EMBL covers themes ranging from studies of single molecules to an understanding of how they work together in complex systems to organise cells and organisms. Our research is loosely structured under thematic units, giving scientists the intellectual freedom to pursue the topics that most interest them. But what really distinguishes EMBL is the large number of inter-unit collaborations, bringing people with common interests but distinct skills together to tackle ambitious projects. Increasingly, our young scientists come with physics, chemistry, engineering, mathematics, and computer science backgrounds, contributing new perspectives and the complementary expertise required to unravel the complexity of living systems – a spirit of working together that is also reflected in numerous external collaborations and partnerships.

EMBL combines a critical mass of expertise and resources with organisational flexibility, enabling us to keep pace with today’s biology. Furthermore, the laboratory’s fixed term contract system ensures that the skills our researchers develop in this energetic, collaborative, interdisciplinary, and international environment are exported to our member states when researchers leave to assume key positions in other institutes. Indeed, the impact of the laboratory’s scientific work, the quality of its services, its continued attractiveness to world-leading young scientists, and the continuous support from our member states are testimony to EMBL’s success and show that we are well equipped to tackle the next forty years.

Iain Mattaj
EMBL Director General
## European Molecular Biology Laboratory

### Research topics

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- **Cell signaling and cell differentiation**
- **Cellular organisation and dynamics, and cell division**
- **Chemistry and chemical biology**
- **Computational genomics and metagenomics**
- **Computational modeling of biological systems and processes**
- **Disease mechanisms, pathogens, molecular medicine, stem cells**
- **Evolution**
- **Functional genomics, genetics and gene networks**
- **Gene regulation, transcription, chromatin and epigenetics**
- **Imaging and image analysis**
- **Macromolecular complexes, interaction networks**
- **Neurobiology**
- **Physics and biophysics**
- **Plant biology**
- **Proteomics**
- **RNA metabolism, transport and processing, miRNAs and snoRNAs**
- **Robotics and automation, engineering**
- **Software development and bioinformatics**
- **Tissue morphogenesis, cell polarity and migration**
- **X-ray crystallography, NMR, electron microscopy**
| Research topics | Cell signalling and cell differentiation | Cellular organisation and dynamics, and cell division | Chemistry and chemical biology | Computational genomics and metagenomics | Computational modeling of biological systems and processes | Disease mechanisms, pathogens, molecular medicine, stem cells | Evolution | Functional genomics, genetics and gene networks | Gene regulation, transcription, chromatin and epigenetics | Imaging and image analysis | Macromolecular complexes, interaction networks | Neurobiology | Physics and biophysics | Plant biology | Proteomics | RNA metabolism, transport and processing, ncRNAs and miRNAs | Robotics and automation, engineering | Software development and bioinformatics | Tissue morphogenesis, cell polarity and migration | X-ray crystallography, NMR, electron microscopy |
The European Bioinformatics Institute (EMBL-EBI) is located on the Wellcome Trust Genome Campus in Hinxton, near Cambridge. As a European hub for biomolecular data, EMBL-EBI offers the scientific community access to a variety of bioinformatics services, alongside which a number of active research groups work in areas that complement and extend these services.

Europe’s flagship laboratory for the life sciences

EMBL was founded in 1974 to create a central European laboratory in the emerging field of molecular biology. It remains the only intergovernmental research organisation in Europe that performs research in the molecular life sciences, and is directly supported by 20 member states and one associate member, Australia. EMBL’s goals are:

- Undertaking outstanding life science research: setting trends and pushing the limits of technology.
- Providing world-class research infrastructure and services to the member states.
- Training and inspiring the next generation of scientific leaders.
- Driving research, innovation and progress through technology development, interactions with industry and technology transfer.
- Taking a leading role in the integration of life science research in Europe.

The European Bioinformatics Institute (EMBL-EBI) is located on the Wellcome Trust Genome Campus in Hinxton, near Cambridge. As a European hub for biomolecular data, EMBL-EBI offers the scientific community access to a variety of bioinformatics services, alongside which a number of active research groups work in areas that complement and extend these services.
EMBL Grenoble builds and operates beamlines for macromolecular crystallography, develops instrumentation and techniques, and provides facilities and expertise to visitors in collaboration with its campus partners, the European Synchrotron Radiation Facility (ERSF) and the Institut Laue-Langevin (ILL). The outstation is also part of the Unit of Virus Host Cell Interactions (UVHCI).

EMBL Hamburg develops novel, innovative technologies in structural biology, such as high-throughput crystallisation and data interpretation software, as well as operating cutting-edge synchrotron radiation beamlines and offering world-leading facilities and expertise to the research community. It also has an ambitious research programme for structures of multifunctional proteins and protein complexes of biomedical relevance.

EMBL Heidelberg is home to five research units, central scientific services, the administration, and the laboratory’s technology transfer arm, EMBL Enterprise Management (EMBLEM). Heidelberg is the largest centre for biomedical research in Germany and there are many bilateral links between EMBL scientists and local research institutions.

EMBL Monterotondo, near Rome, focuses on mouse genetics and functional genomics, and offers expertise in mammalian physiology and production of mouse models of human diseases. Researchers form dynamic partnerships with other international research and clinical centres. The outstation shares a campus with Italian national research groups (IBC-CNR) and the headquarters of the the European Mouse Mutant Archive.
Across EMBL’s five sites there are opportunities across the spectrum of life science research for PhD students, postdoctoral fellows, group leaders, and many other professionals, from software development to chemistry and engineering.

**PhD programme**

Training is one of EMBL’s core missions and our International PhD Programme is renowned for offering excellent education to prospective scientists.

Research independence, dedicated mentoring and an international environment are the cornerstones of the programme, in which close to 200 students from all over the world are currently enrolled.

Students have the opportunity to obtain joint PhD degrees between EMBL and one of its partner universities or from a recognised university of their choice.

EMBL recruits PhD students twice a year. For more details please contact predocs@embl.de. www.embl.de/training/eipp

**Postdoctoral fellows**

Postdoctoral fellows at EMBL benefit from the expertise of world class scientists, state-of-the-art scientific equipment, training in career development and an excellent seminar programme.

Our research groups encourage a balance between senior and young scientists, creating the ideal environment to share and discuss research endeavours while supporting junior colleagues to develop and grow into new positions.

The EMBL Interdisciplinary Postdocs (EIPOD) programme builds on highly interactive research between units and is aimed at candidates whose research crosses scientific boundaries.

Please contact group leaders directly to find out if a position is available, or visit www.embl.de/jobs.
EMBL has ongoing opportunities for physicists, computer scientists and electronic engineers, especially early in their careers. Ever-more sophisticated analysis of very large data sets at the European Bioinformatics Institute (EMBL-EBI) draws on a skilled workforce from many disciplines: from scientific expertise in the life sciences to technical know-how in software development. Similarly, qualified technical staff are highly sought after to operate beamlines at EMBL’s outstations in Hamburg and Grenoble.

Other positions include interface development, communications, user support, industry liaison and training. We offer advanced scientific development and vocational training to improve skills in areas such as coaching and communication.

An international, interdisciplinary and collaborative workplace

EMBL’s staff comprises in excess of 1700 people from more than 60 different countries – this internationality creates an atmosphere that is creative, interdisciplinary and collaborative, with an unparalleled breadth of expertise and complementary skills. EMBL is an equal opportunity employer offering attractive conditions and benefits appropriate to an international research organisation. All employees benefit from excellent working conditions, a young and international atmosphere and a high-quality infrastructure of social services. On-site childcare is available at some of EMBL’s locations, helping staff to combine professional and family life.

Group and team leaders

EMBL fosters the pursuit of ambitious and long-term research projects at the highest level. Group and team leaders have the freedom to set their own scientific directions and are encouraged to explore the most challenging research areas.

Support for team and group leaders includes funding for a number of staff, and laboratory space with equipment. Research collaborations between groups are an integral part of EMBL’s scientific culture.

In addition to advanced scientific development, EMBL offers vocational training to improve skills in areas such as coaching, team management and communication. Establishing a good work-life balance is emphasised at every career stage.
The Hentze group combines biochemical- and systems-level approaches to investigate the connections between gene expression, cell metabolism, and their role in human disease. Key goals of the group include collaborative efforts to: understand the basic mechanisms underlying protein synthesis and its regulation by miRNAs and RNA-binding proteins; explore, define and understand REM (RNA, Enzyme, Metabolite) networks; study the mechanisms and circuits that maintain iron homeostasis and its connection to the immune system; and understand more about the molecular pathways of iron overload, iron deficiency, and iron management disorders.

In investigating the mechanisms and forces that determine cell shape in Drosophila, the Leptin group studies two cell types. They look at how the cells at the tips of the fruit fly’s tracheal system rearrange their components as they grow rapidly and branch out to carry air to the animal’s tissues. And at the tissue level, the group investigates how forces generated by single cells give the embryo’s ventral furrow its final shape. The group also studies medaka and zebrafish to understand how signals from damaged cells are recognised by the innate immune system. They are developing methods to assay immune and stress responses in real time as the fish’s cells encounter pathogens and stress signals.
RNA biology, metabolism and molecular medicine

Matthias Hentze
MD 1984, University of Münster.
Postdoctoral training at the NIH, Bethesda.
Group leader at EMBL since 1989.
Senior scientist since 1998.
Co-Director of the EMBL/University of Heidelberg Molecular Medicine Partnership Unit since 2002.
Associate Director of EMBL since 2005.
ERC Advanced Investigator since 2011.
Director of EMBL since 2013.

Previous and current research

Important steps in the control of gene expression are executed in the cytoplasm by regulation of mRNAs via RNA-binding proteins (RBPs) and non-coding regulatory RNAs (e.g. miRNAs). We are elucidating these regulatory mechanisms, combining ‘reductionist’ biochemical and systems level approaches in mammalian, yeast and Drosophila model systems.

We recently developed ‘mRNA interactome capture’ to define ‘all’ RBPs associated with mRNAs in vivo (Castello, et al., 2012). This work offers an ideal starting point for exploration of ‘REM networks’ (Hentze & Preiss, 2010), which we expect to connect cell metabolism and gene expression in previously unrecognised ways (figure 1).

Within the Molecular Medicine Partnership Unit (MMPU), we are investigating the post-transcriptional processes of nonsense-mediated decay (NMD) and 3’ end processing and their importance in genetic diseases (with Andreas Kulozik, University of Heidelberg).

Our second major interest is the biology of mammalian iron metabolism (figure 2). This work includes the definition of the functions of the IRE/IRP regulatory network and its crosstalk with the iron hormone hepcidin. Within the MMPU (together with Martina Muckenthaler, Heidelberg University), we study the molecular basis of genetic and non-genetic diseases of human iron metabolism. Our work employs conditional knockout mouse strains for IRP1 and IRP2 and mouse models of iron metabolism diseases.

Future projects and goals

- To uncover the basic mechanisms underlying protein synthesis and its regulation by miRNAs and RNA-binding proteins in cell metabolism, differentiation, and development.
- To explore, define, and understand REM networks.
- To help elucidate the role of RNA metabolism in disease, and to develop novel diagnostic and therapeutic strategies based on this knowledge.
- To understand the molecular mechanisms and regulatory circuits underlying physiological iron homeostasis.
- To contribute to the elucidation of the molecular pathophysiology of common iron overload (haemochromatosis), iron deficiency (anaemia) and iron management (anaemia, Parkinson’s disease) disorders.

For research themes and projects of the teams in the MMPU, see: www.embl.de/research/partnerships/mmpu.

Selected references


Figure 1: Exploring REM networks.

Figure 2: Systems biology of mammalian iron metabolism.
Cell shape and morphogenesis: subcellular and supracellular mechanisms

The Leptin group studies the mechanisms and forces that determine cell shape in Drosophila and uses the zebrafish to analyse innate immune signalling.

Selected references


Maria Leptin

PhD 1983, Basel Institute for Immunology.

Postdoctoral research then Staff Scientist at the MRC Laboratory for Molecular Biology, Cambridge, UK.

Group leader at the Max Planck Institute for Developmental Biology, Tübingen.

Professor, University of Cologne, Institute of Genetics.

Director of EMBO and group leader at EMBL since 2010.

Cell shape determination during development

The shape of a developing organism is generated by the activities of its constituent cells: growth and proliferation, movements, and shape changes. We are particularly interested in shape changes.

One study concerns an extremely complex single cell, the terminal cell of the Drosophila tracheal system. It is highly branched and carries air to target tissues through an intracellular tube bounded by plasma membrane. During its rapid growth, the cell faces the task of synthesising large amounts of membrane and sorting it correctly to defined membrane domains. Extensive re-organisation of the secretory organelles precedes membrane growth. We are finding out how the cytoskeleton, small GTPases and polarity determinants direct the process, and how membrane trafficking processes contribute to building the tube.

In another project, we try to understand how the forces generated by individual cells are integrated within the supracellular organisation of the whole organism to give the tissue its final shape (see figure 1). We study the formation of the ventral furrow in the early Drosophila embryo. The cells that form the furrow are the major force generators driving invagination but, to allow furrow formation, neighbouring cells must respond and they may contribute. To understand force integration across many cell populations, we use simultaneous time-lapse imaging of multiple-angle views of the gastrulating embryo. We measure the specific shape changes in all the cells of the embryo, as well as the speed and direction of their movements. Genetic and mechanical manipulations reveal the underlying control circuits.

In vivo imaging of innate immune responses

The innate immune system provides rapid defence against pathogens and also deals with non-pathogenic stresses. Macrophages and dendritic cells, two key players in this system, patrol the body and respond to stimuli from damaged cells via extra- and intracellular sensors. We aim to understand how such signals are recognised and how the appropriate subcellular and intercellular responses are triggered. We have discovered that one family of sensors, the cytoplasmic NOD-like receptors (NLRs), are particularly abundant in fish.

Fish model systems allow in vivo observation of physiological processes. Specifically, we watch pathogens and the cells that attack them. We use genetically and chemically engineered in vivo fluorescent reporters to assay immune and stress responses in real time and at high spatial and temporal resolution as the cells of the fish encounter pathogens and stress signals (see figure 2).

Figure 1: A flat projection of the entire surface of a Drosophila embryo in which the position and speed of 6000 cells is followed over a 40 minute period. The head of the embryo is at the top, the center of the image is the ventral midline towards which the lateral cells are moving. Image by Matteo Rauzi.

Figure 2: Zebrafish larvae 24 hours after infection with fluorescent bacteria. Normal fish survive and eventually clear the bacteria (left), but if the interferon signalling pathway is compromised (right) the bacteria proliferate and the fish die. Image by Dirk Sieger.
In this Unit, physicists and chemists work closely together with biologists to elucidate the fundamental rules that govern dynamic cell organisation and function. At the same time, groups are developing new instruments and technologies in order to reach this ambitious goal.

Cells are the smallest autonomous units of life and occupy the midpoint between the molecular and macroscopic scales. In order to understand how living systems are built and function, we need to understand the physical principles that underlie cellular organisation and function.

It is in the cell where we will first understand the basic processes of life at the molecular level in a physiological context. The cell provides the natural coordinate system in space and time onto which we have to map and integrate genomic, transcriptomic, proteomic, structural and biophysical information about the molecules that make up living systems. In short, cell biology has become an integrative hub of much of modern biological research.

This is a time of tremendous opportunity for cell biology, but realising it also represents a formidable challenge and requires new concepts and approaches. Individual cellular processes – such as signalling, membrane trafficking, cytoskeletal dynamics, gene expression or cell division – can no longer be studied in isolation but need to be considered as integrated events. The default situation is that the molecular machinery that performs these functions is complex and combinatorial at the single protein, protein complex, and pathway level. This requires new ways of thinking about cellular functions that use network biology and employing quantitative theoretical methods to generate mechanistic and predictive models that rely on realistic physical principles at the cellular, subcellular and molecular scale. Therefore, cell biology needs to integrate traditionally separate disciplines to realise its potential.

Novel developments in microscopy, computer simulations and chemical biology-based probes are a particular strength of the Unit. We constantly explore new directions and integrate new approaches and disciplines to answer cell biological questions. New correlative light/electron and superresolution imaging methods, as well as mechanistic biochemistry, allow us to directly interface between cell and structural biology to understand molecular mechanisms. Furthermore, advances in live microscopy methods now allow us to carry out cell biology in developing organisms to understand how cell organisation and collective cell behaviour leads to organ formation.

Mechanisms of cellular functions are often best understood when the organisation of the cell changes dramatically to carry out new functions. This is the case when cells divide, or when they change their fate. Both opportunities are exploited in the Unit. As a cell prepares to divide, all the microtubules suddenly depolymerise to reassemble into the mitotic spindle. At the same time, the nucleus is disassembled, mitotic chromosomes are formed, the Golgi complex fragments and membrane traffic ceases. After segregation of the genome is achieved, cellular organisation is re-established. Thus every cell cycle provides the opportunity to study the principles of the biogenesis of cellular compartments. Similarly, when progenitor cells differentiate into new cell types, the genetic programme is changed and a reorganisation of cellular architecture takes place, guided by rules that we begin to unravel. Understanding these rules and principles is our challenge in the years to come.

Jan Ellenberg
Head of the Cell Biology and Biophysics Unit
Previous and current research

The genome of eukaryotic cells is compartmentalised inside the nucleus, delimited by the nuclear envelope (NE) whose double membrane is continuous with the endoplasmic reticulum and perforated by nuclear pore complexes (NPCs). In M-phase, most metazoan cells reversibly disassemble the nucleus. Chromosomes are condensed, attached to cytoplasmic spindle microtubules, faithfully segregated and decondensed, and the nucleus rapidly reassembles. Errors in this beautiful cycle of cell division can lead to severe consequences, such as cancer in somatic cells and infertility in gametes.

The overall aim of our research is therefore to systematically elucidate the mechanisms underlying cell division and nuclear organisation in mitosis and meiosis. To achieve this, we are developing advanced fluorescence microscopy-based methods to systematically identify the required proteins and study their function in living cells. Quantitative imaging is coupled with computerised image processing and simulations to extract biophysical parameters and build mechanistic models. As biological systems, we are using somatic human cells for mitosis and cancer, and mouse oocytes and early embryos for meiosis and infertility.

In the recent past we showed that mitotic nuclear breakdown and reformation is initiated by the ordered dis- and reassembly of NPCs, which interestingly use a different pathway to assemble during interphase. Furthermore, we demonstrated that meiotic spindle assembly and asymmetric positioning rely on novel mechanisms and that meiotic chromosome biorientation is highly error prone (figure 1). We have also identified hundreds of new cell division genes by screening the entire human genome using time-lapse microscopy (figure 2).

Future projects and goals

The objective of our future work is to gain comprehensive mechanistic insight into the division of human mitotic cells, to establish methods for systems analysis of the meiotic and first mitotic divisions of mammalian oocytes and embryos, and to provide a biophysical basis to understand nuclear organisation.

For mitosis, we are aiming to achieve a systems-level understanding of all proteins identified in our phenotypic profiling of the human genome. To this end we are continuously automating advanced fluorescence imaging and single molecule techniques to study protein function in live cells in high throughput.

For oocyte meiosis and early embryonic mitosis, we are pursuing the molecular mechanism of homologous chromosome segregation and are developing new gene silencing methods and light sheet-based imaging systems to establish a physiological model for infertility accessible for systems biology.

For nuclear organisation, we are currently focusing on the structure and assembly of the nuclear pore, the targeting and function of inner nuclear membrane proteins, and chromosome architecture and dynamics.

Selected references


Multicellular Morphogenesis

Using the zebrafish as a model, the Gilmour group takes an integrative, multiscale approach to study how cells collectively migrate and assemble into functional organs.

Selected references

Previous and current research
Collective behaviour lies at the heart of all biological design. Whether it is the assembly of proteins into complexes or the organisation of animal societies, collective interaction creates something much greater than the sum of the parts. A breathtaking example of such behaviour is seen during embryogenesis, when thousands of collectively migrating cells self-organise to form functional tissues and organs. Given the key role played by collective migration in organ formation, wound repair and cancer, it is surprisingly how little we know about how cells organise each other.

We take an integrative, multi-scale approach to study how cells collectively migrate and assemble into functional organs, using the zebrafish lateral line organ as an experimental model. Here, a migrating epithelial primordium comprising of 100 cells, assembles and deposits a series of rosette-like mechanosensory organs across the surface of the embryo. Its superficial migration route, beneath a single transparent cell layer, makes it the dream in vivo sample for quantitative imaging. Moreover, the process can be interrogated using a range of perturbation approaches, such as chemical and optogenetics, and many of the molecular regulators of its migratory behaviour are of general interest due to their role in human disease. For example, the migrating collective is guided by Cxcr4/SDF1 signalling, a chemokine-receptor pair known to control many human cancers.

Future projects and goals
The focus of our group is to use the lateral line to address the general question of how cell behaviours are regulated and coordinated within collectively migrating tissues. We have developed in vivo imaging, analysis and perturbation tools that allow the entire morphogenesis process to be addressed at different spatiotemporal scales. By integrating these data, using statistical methods and modelling, we are aiming to understand the interplay between ‘opposing’ behaviours – namely, cell migration and differentiation. In this way, we hope to move towards a systems-level understanding of how dynamic cell organisation and gene expression are integrated during tissue morphogenesis.

Darren Gilmour
PhD 1996, University of Cambridge.
Postdoctoral research at the Max Planck Institute for Developmental Biology, Tübingen.
Group leader at EMBL since 2004.

The zebrafish migrating lateral line organ allows collective migration to be easily studied in vivo.

Visualising actin dynamics (LifeAct-GFP) within migrating primordium.
EMBL Heidelberg

Previous and current research
Eukaryotic chromosomes undergo enormous changes in structure and organisation over the course of a cell cycle. One of the most fascinating changes is the transformation of interphase chromatin into rod-shaped mitotic chromosomes in preparation for cell division. This process, known as chromosome condensation, is a key step for the successful segregation of chromosomes during mitosis and meiosis. The underlying mechanisms are, however, still poorly understood.

The overall aim of our research is to unravel the action of molecular machines that organise the 3D arrangement of chromosome fibres. Insights into the general working principles behind these machines will be of great importance to our understanding of how cells inherit a complete set of their genomes every time they divide and thereby prevent the emergence of aneuploidies, which are hallmarks of most cancer cells and the leading cause of spontaneous miscarriages in humans.

One of the central players in the formation of mitotic chromosomes is a highly conserved multi-subunit protein complex, known as condensin. We have shown that condensin binds to mitotic chromosomes by encircling chromosomal DNA within a large ring structure formed by its structural maintenance of chromosomes (SMC) and kleisin subunits. Our working hypothesis is that condensin uses this topological principle to tie together loops of chromatin (figure 1), which ensures that chromosome arms clear the site of cell cleavage before cytokinesis.

In an independent project, we use a newly developed time-resolved light microscopy assay to quantitatively measure chromosome condensation in live fission yeast cells in high-throughput (figure 2). This has identified, in addition to condensin, new players that direct the formation of mitotic and meiotic chromosomes.

Future projects and goals
We will continue to use an interdisciplinary approach to advance our understanding of condensin function in yeast and mammalian cells by combining biochemical, molecular, structural, and cell biology methods. In collaboration with other groups, we are taking further advantage of chemical biological techniques as well as single molecular approaches to discover how condensin loads onto chromosomes, how it interacts with other chromosomal components, and how its activity is controlled. In addition, we are further investigating the novel candidates identified in the screen for mitotic chromosome condensation proteins to understand the basis of their functions on mitotic chromosomes.

Selected references
Cuylen, S., et al. (2013). Entrapment of Chromosomes by condensin rings prevents their breakage during cytokinesis. Dev. Cell, 27, 469-78

Figure 1: Model for the organisation of mitotic chromosomes by condensin rings.

Figure 2: Monitoring chromosome structure and segregation in fission yeast cells.

The Häring group aims to understand the molecular machinery that organises chromosomes to allow their correct distribution among daughter cells.

Christian Häring

Chromosome structure and dynamics

The Häring group aims to understand the molecular machinery that organises chromosomes to allow their correct distribution among daughter cells.
Dynamics of cell growth and tissue architecture

The Hufnagel group studies the role of mechanical constraints on processes such as cell growth, programmed cell death, orientation of division, intra-tissue rearrangements and cell differentiation.

Selected references


Previous and current research

Biological processes are highly dynamic and span many temporal and spatial scales. During development, cells must integrate and respond to a multitude of biochemical and biophysical signals: for example, changes in intracellular signalling networks, cytoskeleton remodelling, cell shape changes, long-range signalling and tissue remodelling. A whole-embryo view of morphogenesis with subcellular resolution is essential for unravelling the interconnected dynamics at varying scales of development – from interactions within cells to those acting across the whole embryo. Bridging scales from the submicron to the millimeter range with a temporal resolution of several seconds (combined with a total imaging time of several hours) not only poses tremendous challenges for modern microscopy methods but also requires powerful computational approaches for data handling, processing, and image analysis.

The central question that we are interested in is how a complex multi-cellular tissue or organism is formed from individual cells by spatio-temporal regulation of biophysical and intracellular signalling processes. We address all experimental steps, from innovative transgenic lines and microscope development to systematic image processing and biophysical modelling. This requires a multidisciplinary environment of biologists, physicists and computer scientists working closely together.

In order to address these questions we develop novel imaging techniques based on selective plane illumination microscopy (SPIM). SPIM yields optical sectioning by uncoupling the optical path for sample illumination from emitted photon detection. The illumination branch creates a thin light sheet to illuminate a specimen from the side and the emitted light is collected and imaged onto a high speed and high sensitivity camera by a second objective lens. The unprecedented speed of light sheet-based microscopy poses challenges for data handling and image processing, which we address by developing novel image processing tools.

Currently, we investigate cell shape changes and growth patterns in the Drosophila embryo with emphasis on the role of mechanical constraints on organ formation and tissue differentiation, complemented by mammalian cell culture studies investigating cell cycle response of an epithelial tissue to external and internal mechanical perturbations.

Future projects and goals

Our research interests are focused on the control and regulation of cell proliferation, apoptosis and cellular rearrangement processes in developing tissues, with a specific emphasis on epithelial tissues and the role of mechanical interactions as a regulator. We seek to characterise and quantify the spatio-temporal effects of mechanical stress, deformations and fluid flow-induced shear stress on cell growth, gene expression and cellular polarity in two-dimensional epithelial tissues. To address this issue, we pursue an interdisciplinary approach combining classical biological techniques with detailed modelling methods from various fields, ranging from statistical physics to applied mathematics and computer science. We will continue to not only tailor light-sheet microscopes to match specific biological questions, but also push the boundaries of light-sheet microscopy towards high speed intracellular imaging with extremely thin light sheets, super-resolution techniques, and quantitative in toto imaging.
Dynamics of membrane trafficking

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Previous and current research

Many biological processes at the cellular level are based on complex networks of macromolecular interactions. These networks have a modular organisation, where the different modules form dynamic molecular machines that drive processes such as signalling, cell motility, cytokinesis, and vesicle trafficking. Our group's long-term goal is to contribute to the understanding of the general principles governing the assembly and function of these supramolecular machines.

More specifically, we are interested in the formation of cargo-loaded transport vesicles, such as endocytic vesicles. The formation of the endocytic vesicle is driven by highly dynamic molecular machinery that is composed of more than 50 different protein species and of several thousand individual protein molecules. We aim to understand the processes that regulate the assembly of the endocytic machinery, the recruitment of the cargo molecules, and the selection of the location and timing of endocytic events in the cell.

Our main experimental organism is the budding yeast, Saccharomyces cerevisiae. In our studies we use quantitative live-cell imaging methods (for example particle tracking, FRAP, FCS/FCCS, high-throughput microscopy) in combination with powerful yeast genetics. We also use correlated light and electron microscopy to gain nanometre scale information about the endocytic structures, and biochemistry to characterise protein-protein and protein-lipid interactions.

Future projects and goals

We are interested in the mechanisms that initiate the assembly of the endocytic machinery and regulate the precise timing of the sequential stages of the assembly. The spatial distribution of the endocytic events is tightly coupled to the cell cycle and to the overall polarity of the cell. The spatially regulated initiation of endocytic events is critical for determining the cellular distribution of endocytosis.

We are also studying the mechanisms of selective recruitment of cargo molecules into the endocytic vesicles. The recruitment of cargo proteins is tightly regulated by a family of endocytic adaptors. We want to learn how this adaptor system integrates environmental and intracellular signals in deciding which cargoes to recruit.

Furthermore, we want to understand how actin functions to promote endocytic vesicle budding. In yeast, endocytosis is strictly dependent on actin polymerisation, but the mechanisms by which actin drives vesicle budding are not well understood. We are currently studying the molecular basis of the coupling between the actin cytoskeleton and the endocytic membrane. We have also started to investigate the evolution of the membrane-actin coupling in animals and fungi using a phylogenetic comparative approach.

The core membrane trafficking events, such as the clathrin-mediated endocytosis, are elemental cellular processes that are involved in multiple biological phenomena ranging from cell polarization to neural plasticity. As most of the yeast trafficking proteins are widely conserved in eukaryotes, we believe that mechanisms that we unravel in yeast cells will be applicable to eukaryotes in general.

Selected references


A yeast cell expressing fluorescently labelled endocytic proteins. The first two images show a coat protein Sla1 (green) and an actin-binding protein Abp1 (red). The last image shows both channels merged. The spots at the cell surface reveal the transient accumulation of the proteins at endocytic sites during vesicle budding.
Cytoskeletal dynamics and function in oocytes

Previous and current research
All animal life begins with the fusion of sperm and egg. Our research is focused on the egg cell, specifically investigating how the fertilisable egg develops from the oocyte through meiotic divisions. Oocytes are exceptionally large cells, with diameters up to millimetres in size, because they store large amounts of nutrients to support embryonic development. Therefore, in oocytes and eggs, the cytoskeleton has to transport organelles, separate chromosomes, and organise cellular architecture in a very large cytoplasm. How the cytoskeleton adapts to this unusual size, and how these mechanisms differ from those in small somatic cells, is largely unknown.

We use starfish oocytes as a model system because they are easy to handle, complete meiosis rapidly, develop simply in seawater at room temperature, and are transparent – ideal for high-resolution imaging of cytoskeletal dynamics in live cells. We use confocal microscopy to image live oocytes and employ computational image analysis tools to extract quantitative parameters from these 3D time-lapse datasets. Parameters such as local concentrations or velocities of cellular components provide a quantitative assay for the biological process and, at the same time, serve as inputs for computational models of cytoskeletal dynamics. Model predictions are then tested in perturbation experiments using physical (e.g. laser microsurgery) or molecular (e.g. RNAi knockdown) manipulations. Biochemistry, in combination with the imaging assays, is used to identify the key molecular components in the process.

We have recently shown that meiotic chromosomes scattered in the large oocyte nucleus are collected by an actin meshwork and transported to the spindle, whose short microtubules cannot reach the chromosomes directly, as they do in somatic cells. This novel actin-based chromosome transport system forms as the nuclear envelope breaks down and fills the nuclear space with an actin meshwork, physically entrapping chromosomes. We were able to show that the actin meshwork contracts homogeneously; however, because it is mechanically anchored to the cell cortex, this homogeneous contraction is translated into directional transport towards the cortex where the spindle forms. By understanding the mechanism of chromosome transport essential to oocyte division and fertility, our studies revealed a novel design principle for a cytoskeletal ‘transport machine’ that is very different from previously known mechanisms of actin-driven intracellular transport.

Future projects and goals
Immediate goals include determining the detailed structure of the F-actin meshwork, understanding the molecular mechanisms of meshwork contraction, and identifying the mechanisms by which chromosomes attach to the meshwork. We will employ high-resolution imaging methods, including electron tomography and super-resolution light microscopy, to resolve single actin-filaments and to identify, localise and perturb molecules regulating actin filament dynamics that, combined with high-resolution imaging, will allow us to address the underlying molecular mechanisms. Longer term, we are interested in related cytoskeletal processes that occur in oocytes, eggs and early embryos, with the overall goal to mechanistically understand the organisational principles of the actin and microtubule cytoskeleton.

Selected references

Using starfish as a model organism, the Lénárt group combines molecular methods with imaging assays to investigate how the fertilisable egg cell develops from the oocyte.

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Group leader since 2011.

The actin filament network (grey) embedding the chromosomes (red).

Transparent starfish oocytes are uniquely suited for imaging meiotic divisions.
Previous and current research

Modern microscopy has demonstrated the dynamic nature of biological organisation. The mitotic spindle, for example, is a stable and solid cellular structure: in a given cell type, it has a precise symmetry and very reproducible dimensions. Yet, except for the chromosomes, all the components of a spindle — polar filaments called microtubules and associated proteins — are in rapid turnover. Microtubules grow, shrink and disappear in less than a minute and their associated proteins continuously and stochastically bind and unbind even faster. The resulting assembly, although highly dynamic, is remarkably precise: it can remain steady for hours waiting for the right signal, to eventually apply the balanced forces necessary to position and segregate the chromosomes exactly. The spindle is thus a fascinating structure that illustrates a central question in biology: how can the uncoordinated and inevitably imperfect actions of proteins and other molecules collectively fulfill the biological needs with the required accuracy?

Today, understanding biological phenomena from their multiple biological components seems within our reach, as testified by the rise of systems biology. Yet, collective behaviours in biology require more than statistical averages. They are challenging us for many reasons: 1) the diversity of molecular players is enormous; 2) their interactions are often dynamic and out-of-equilibrium; and 3) the properties of the constituents have been selected by natural evolution.

We approach this topic in practical terms by developing in vitro experiments and modelling tools. This approach allows us to reduce the number of components in the system: we can either remove specific proteins, or start from scratch by mixing purified components. Modelling allows us to recapitulate the process of protein organisation in a framework in which all the interactions are known exactly and can even be specified at will. We have developed an advanced simulation engine — called Cytosim — to simulate ensembles of multiple polar fibres and associated proteins. Cytosim can simulate various problems involving microtubules, actin filaments or both. Simulations are often used to validate or refute existing ideas, but we also try to use them in a more creative way: one can generate systematically various properties for the molecules and automatically test their ability to form stable structures. The analysis of successful scenarios leads to the formulation of new hypotheses.

Future projects and goals

We will study systems in which experiments and theory can be synergistically combined. We currently focus on Xenopus egg extracts, an experimental system in which many aspects of mitosis can be recapitulated. We are also generally interested in modelling cellular processes in which the cytoskeleton is a major player, such as the different stages of mitosis, the generation of cell shape in S. pombe, or the generation of asymmetry during cell division.
Systems biology of stem cell differentiation

The Neveu group takes an integrated systems biology approach to investigate the molecular changes that determine what a stem cell becomes.

Selected references

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Previous and current research
Pluripotent cells have the dual ability to self-renew and differentiate. Therefore, in pluripotent cells, the expression of hundreds of genes should be stable in the self-renewal case, but gene expression can also be directed in a coordinated manner towards particular states upon external signalling cues (lineage commitment towards terminal differentiation). Deciphering this complex problem has garnered much attention at the systems level.

Tackling this challenge requires good characterisation of the pluripotent state. miRNAs are suitable marker candidates because they are excellent classifiers of tissue types or cellular states and they also play a crucial role in differentiation. By profiling miRNA expression in human cells, we have previously shown that pluripotency surprisingly emerges as a much more diverse state than previously believed: variability in miRNA expression is comparable to that found in differentiated cells and cancer cells. We have also shown that it is possible to dramatically reduce the complexity of miRNA expression patterns to a few meaningful dimensions. This reductionist approach still allows us to quantitatively and robustly discriminate pluripotency, cancer and lineage commitment. More importantly, it suggests that complex processes of the stem cell system, such as differentiation and reprogramming, can be mapped quantitatively.

Currently, we are employing a dynamic approach at the single cell level to resolve the dynamics of differentiation and the different molecular and cellular processes at play during fate determination. Indeed, differentiation is intrinsically a dynamic process, where individual cells have to change from one state to another. Having developed fluorescent reporters to assess miRNA expression in single cells, we are characterising mouse embryonic stem cell (ESC) self-renewal using single-cell live imaging.

Future projects and goals
We plan to study the dynamics of differentiation at the single-cell level both in vitro in mouse embryonic stem cells and in vivo. The ultimate goal is to dissect the transcriptional regulation and gene networks and the associated cellular changes underlying stem cell differentiation. We are taking an integrated systems biology approach that combines single-cell live imaging of miRNA expression, image processing, perturbation approaches, and mathematical modelling.

We wish to address the following questions:
- How dynamic is the pluripotent state?
- What are the in vitro dynamics of differentiation of mouse ESCs?
- How do in vitro findings compare to in vivo differentiation behaviour?

Molecular cartography of stem cells: miRNA expression classifies pluripotent cells, cancer cells and differentiated cells. This map allows us to quantitatively follow changes in cell identity, such as differentiation and reprogramming. It reveals that reprogramming involves a cancer-like behaviour.
Membrane traffic and organelle biogenesis

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Team leader at EMBL since 1998.
Senior scientist since 2012.

Previous and current research
Our research goal is to comprehensively understand the temporal and spatial organisation of the early secretory pathway and organelle morphogenesis at the molecular level.

While many of the core components of the secretory machinery have been identified and characterised to some detail in the past decades, still little is known about how all components function together and how they are regulated in response to extracellular stimuli, stress or differentiation. Transport of material from one organelle to the other involves several steps, which have to occur sequentially and thus require a high degree of control at the molecular level (see figure 1). In order to understand such regulation in the physiological system that contains all possible components involved in the intact cell, we have developed and applied microscopy-based approaches to systematically identify components that regulate the early secretory pathway and the biogenesis and maintenance of the Golgi complex, down to the genome level. We have also developed and applied high-throughput microscopy techniques to quantitatively image genetic or physical interactions of the components we identified.

Network analyses of the components identified in our large-scale screens revealed links between early secretory pathway function, small GTP-binding protein regulation, actin and microtubule cytoskeleton organisation and growth factor mediated signalling. It provides a basis for understanding the global cellular organisation and regulation of the secretory pathway.

In order to investigate the mechanisms of Golgi biogenesis we have developed an approach, using laser nanosurgery, to deplete living cells from their Golgi complex and subsequently analyse the ‘Golgi-less’ karyoplast by time-lapse light and electron microscopy (figure 2). With this approach we are able to show that Golgi biogenesis in mammalian cells occurs de novo from ER derived membranes by a self-organising mechanism that integrates Golgi biogenesis, ER-exit sites biogenesis and the organisation of the microtubule network.

Future projects and goals
We will study the complement of components that our genome-wide screens identified as being involved in the early secretory pathway in further detail. An important question in this context will be if and how they participate in the temporal and spatial organisation of ER-exit sites and their function, and the biogenesis of the Golgi complex.

Ultimately, we hope to be able to define and understand the molecular network(s) underlying trafficking at the ER/Golgi boundary and Golgi function, also considering their relationship to other cellular processes such as transcriptional control, lipid or general metabolism, or signalling and thus contribute towards a global molecular understanding of the living cell.

Figure 1: The four steps involved in ER to Golgi transport in mammalian cells. (I): Biogenesis of COPII coated vesicles occurs at specialised ER exit sites of the ER. (II): COPII vesicles homotypically fuse to form larger vesicular tubular transport carriers (VTCs) that are transported to the Golgi complex along microtubules. (III): VTCs arrive at the Golgi complex and fuse to it to deliver their cargo. (IV): Transport machinery and misrouted proteins are return back to the ER by a distinct class of carriers.

Figure 2: (A) Cells are cut by laser nano-surgery to generate a Golgi-less karyoplast and Golgi containing Golgiplasts (arrowhead). Karyoplasts are the followed by time-lapse microscopy to monitor de novo Golgi biogenesis in living cells (B). The arrowhead points to the Golgi-like structure reforming after nano-surgery in karyoplasts.

Selected references
Cellular nanoscopy

The Ries group develops cutting-edge superresolution microscopy methods, such as automated localisation microscopy for proteome-wide superresolution imaging.

Selected references

Previous and current research
The resolution of optical microscopy is limited by diffraction to about 200 nm, which is much larger than the relevant length-scales in cell biology, defined for instance by the size of organelles or supramolecular complexes. Single molecule localisation-based superresolution microscopy methods, such as photoactivated localisation microscopy (PALM), rely on the stochastic activation and subsequent localisation of individual fluorophores. They reach a 10-fold higher resolution, which is optimal for the study of intracellular structures. Until now however, these techniques required special fluorescent proteins to be cloned or high-affinity antibodies to be generated for specific labelling. On the other hand, many laboratories have most of their constructs in green fluorescent protein (GFP) form and entire genomes are available as functional GFP-fusion proteins. We recently developed a labelling scheme to make all these constructs immediately available for superresolution microscopy by targeting them with small antibodies labelled with bright organic dyes. This opens the door to high-throughput localisation analysis of entire genomes at the nanoscopic level in cells.

Our current research efforts are threefold. First, collaborating closely with other groups, we are establishing state-of-the-art superresolution microscopy to answer exciting questions in cell biology, which have only now become accessible due to greatly improved resolution. Second, we are working on automating single-molecule localisation microscopy with the aim of proteome-wide imaging – such superresolution localisation maps of proteins will be an invaluable resource for the life science community. Third, we are developing novel detection schemes for localisation microscopy. In one project we intend to use the principle of surface-generated fluorescence to improve the axial resolution. In another we aim to measure the 3D orientation of single molecules, in addition to their position by polarised detection, in order to resolve the structure of multi-molecular complexes.

Future projects and goals
Our goal is to establish cutting-edge superresolution microscopy and to apply it to biological systems. We will implement the newly developed detection schemes in a powerful microscope and combine this with advanced data analysis and our expertise in sample preparation and labelling. Using automated single-molecule localisation microscopy we are planning to image the whole proteome of budding yeast with a resolution of ~20 nm in dual-color and 3D. The combination of optical superresolution microscopy with dynamic microscopy techniques such as fluorescence correlation spectroscopy (FCS) or single particle tracking (SPT) bears great potential in relating structure, localisation and function, as does the combination with electron microscopy to add molecular specificity to the ultra structure.
Chemical cell biology

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Group leader at EMBL since 2001.
Senior scientist since 2008.

Previous and current research

Past projects: Before joining EMBL in 2001, our research focused on finding novel ways to stimulate chloride and water secretion of epithelial cells in understanding the genetic disease cystic fibrosis (CF). Our compounds helped to investigate some of the underlying intracellular signalling pathways and provided drug candidates to eventually treat CF patients. Of particular significance was the development of chemical methods to convert highly polar signalling molecules like cyclic nucleotides, inositol phosphates and phosphoinositides to membrane-permeant, bioactivatable derivatives (‘prodrugs’) (Schultz, 2003; Laketa et al., 2009), and lately peptides (Cobos-Correa et al., 2012).

Current projects: Our interest in CF has shifted to the development of lung emphysema (the ultimate cause of death in the patient). In a truly translational collaboration with the Mall group (Molecular Medicine Partnership Unit & German Center of Lung Research), we develop FRET reporters to sense enzyme activities detrimental to lung tissue, such as macrophage and neutrophil elastases. In ex vivo experiments, we are now able to monitor these enzyme activities on cells from both mouse models and patients (Cobos et al., 2009; Gehrig et al., 2012). At the cell biology level, our interest focuses on signalling networks regulated by G-protein-coupled and growth factor receptors. We developed a wide range of fluorescent reporter molecules, either genetically encoded (Piljic & Schultz, 2011) or as small molecule fluorescent probes (see figure). We hope to provide a more complete picture of the signalling network and to help find compounds beneficial in unravelling basic principles in signal transduction and, ultimately, in ion and enzyme secretion relevant to CF patients. In addition, we prepared a large number of tools to manipulate signalling networks and are able to locally activate the important messenger such as PIP3 and DAG with a light flash in subcellular resolution in living cells (Mentel et al., 2011; Nadler et al. 2013). Alternatively, we switch on enzymes such as single G-proteins by translocating them to their site of action with the help of a chemical dimeriser (Putyrski et al., 2011).

Hot projects: Currently, we are very excited about performing bioorthogonal chemistry inside living cells. In collaboration with the Lemke group (page 62), we developed a new set of amino acids that can be site-specifically incorporated into a protein of interest by amber stop codon suppression and then labelled with bioorthogonal chemistry inside living cells. In collaboration with the Häring group (page 20), we developed a new method to cross-link proteins in a protein-protein interaction-dependent fashion in living cells by using FlAsH technology (Rutkowska et al., 2012).

Future projects and goals

In 2014, we will focus predominantly on lipid signalling and lipid-controlled cell biology and examine the effect of sphingo- and phospholipids on endocytosis, lipid trafficking, and secretion. Most projects rely on organic chemistry and the group has a significant number of preparative chemists at the graduate student and postdoc level. The symbiosis of chemistry, biochemistry and cell biology opens new doors and grants novel insights into how cells function.

The Schultz group develops tools for imaging and for manipulating cellular enzyme activities, with a particular emphasis on lipid signalling and the hereditary disease cystic fibrosis.

Selected references


Several reporter and modulator molecules have been developed in our lab, including: small molecule sensors for lipases and proteases; genetically encoded reporters for kinase and phosphatase activities; membrane-permeant and photoactivatable lipid molecules; and lipid derivatives that can be fluorescently labelled in living cells.
Volume correlative light and electron microscopy

The Schwab team is interested in developing tools for the 3D correlation of data generated by fluorescent imaging and by electron microscopy.

Selected references

Previous and current research
Correlative light and electron microscopy (CLEM) is a set of techniques that allow data acquisition with both imaging modalities on a single object. It is a growing field that now includes a large variety of strategies, and one that reaches a high degree of precision even in complex biological models. Before joining EMBL, we were developing tools and protocols to track rare objects or dynamic phenomena on cultured cells and bulk specimen such as nematodes and murine tissues.

One common challenge when trying to combine imaging modalities on the same sample is to identify space cues (external or internal) to track single objects when switching from light microscopy (LM) to electron microscopy (EM). On adherent cultured cells, we have previously developed specific substrates with coordinates to precisely record the position of cells (Spiegelhalter et al., 2009; Gibbings et al., 2011). Currently, we are exploiting these approaches to develop new workflows allowing the study of a higher number of cells.

On more complex specimens, such as multicellular organisms, this targeting is even more critical as systematic EM acquisition of their entire volume would otherwise involve tedious and extremely long processes. For this reason, we are developing new methods to map the region of interest (ROI) within large living specimens, taking advantage of structural hallmarks in the sample that are visible with both LM and EM. The position of the ROI is mapped in 3D by confocal or multiphoton microscopy and then tracked at the EM level by targeted ultramicrotomy (Kolotuev et al., 2009, 2012; Goetz et al., 2014). Relying on structural features of the sample as anchor points, the cell or structure of interest can then be retrieved with sub-micrometric precision.

Future projects and goals
In parallel to the fast evolution of CLEM techniques over the past decade, acquisition methods in electron microscopes have significantly evolved with special breakthroughs in the volume analysis of cells by transmission electron microscopy (TEM) and scanning electron microscopy (SEM) tomography. Our team, in collaboration with other research teams at EMBL, will now combine these advanced techniques to perform CLEM in the 3D space of complex model specimens for cell and developmental biology. We aim to develop new techniques and software to facilitate and automate the correlation and acquisition of large amounts and volumes of sample. By automating these tedious procedures, we intend to enhance the throughput of data collection.

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Yannick Schwab

Figure 1: CLEM on cultured cells. A & B: the Golgi apparatus is tagged with a GFP marker and imaged by light microscopy (in collaboration with the Pepperkok team). Using a CLEM workflow, the same cell is tracked and the ROI studied by electron microscopy.

Figure 2: CLEM on cancer cells. Intravital imaging and ultrastructure. A: Fluorescent cancer cells were injected in mouse skin and imaged with multiphoton microscopy (J. Goetz and L. Mercier, Inserm, France), enabling the visualisation of both the invasive cells (green) and the vasculature (red). B: Following EM sample preparation, the ROI (boxed in A) can be retrieved and imaged at high-resolution with TEM and electron tomography, revealing the complex organisation of the cell membrane in 3D (C and D).
A fundamental question in developmental biology is the mechanism by which symmetry is broken and cells with distinct fates are specified. Researchers in the Unit are studying a number of related research areas, including the mechanisms underlying cell polarisation, mRNA transport, and translational control in *Drosophila*; how auxin specifies different cell types in *Arabidopsis*; and a systems-level understanding of the symmetry breaking processes operating in the early mouse embryo.

During development, progenitor cells divide and differentiate into tissues of characteristic shape and function. Another aim is to elucidate how cells in the early *Drosophila* embryo reorganise their content in response to the expression of key developmental transcription factors and, specifically, how tissue-specific gene expression controls protein and membrane trafficking, and how this trafficking regulates cell fate and behaviour.

Elucidating the temporal organisation of embryonic development is a further goal. Using the mouse model, the mechanisms controlling overall developmental rate at an organisational level, as well as the timing of individual patterning processes and the dynamics of underlying signalling pathways, are being investigated. Analysis of novel mouse reporter lines using real-time imaging techniques allows visualisation of the activity and dynamics of signalling pathways in the context of a developing embryo.

The marine annelid Platynereis is an ideal model for exploring the evolution of cell types. Large-scale expression profiling at cellular resolution has revealed the evolutionary origin of the vertebrate hypothalamus. Using this model, research in the Unit aims at solving one of the major remaining mysteries in animal evolution: the evolution of the central nervous system.

Several groups seek to understand both normal development and its deviations in disease. During brain development, vast numbers of neurons are targeted for death and are cleared rapidly and efficiently by a resident lineage of phagocytes, the microglia. Combining live imaging and genetic approaches, the dynamic relationship between neurons and microglia in zebrafish is actively investigated.

Re-shuffling of regulatory inputs after chromosomal rearrangements is the likely cause of several human genetic disorders. Focusing on the regulatory architecture of key developmental loci, another goal in the Unit is to understand the molecular mechanisms that control functional interactions between genes and remote *cis*-regulatory elements, and to determine how they contribute to phenotypic variations during vertebrate evolution and in humans.

Anne Ephrussi
Head of the Developmental Biology Unit
RNA localisation and regulation in development

Anne Ephrussi
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Group leader at EMBL since 1992.
Coordinator of EICAT since 2005.
Head of Developmental Biology Unit since 2007.

Previous and current research
Intracellular RNA transport coupled with localised translation is a powerful and widespread mechanism that promotes the functional polarisation of cells, from yeast to man. Asymmetric localisation of messenger RNAs within cells has key roles in cell fate decisions, cell migration, cell morphology and function. mRNA targeting is particularly evident in large cells, such as eggs and neurons, where it allows rapid and localised deployment of protein activities in response to extrinsic signals.

An ideal model for the study of RNA transport is the large Drosophila oocyte, in which asymmetrically localised cell fate determinants specify the body axes and pattern the future embryo. During oogenesis, mRNAs encoding these embryonic axis determinants are transported to specific sites within the oocyte, where they are anchored and locally translated, thus ensuring spatial restriction of their protein products. A polarised cytoskeleton and specific motor proteins mediate mRNA transport and anchoring within the cell. We use these RNAs as models to understand how mRNAs localisation and translational control are regulated in space and time.

One RNA of particular interest is oskar, which encodes the posterior determinant of Drosophila. Oskar protein is uniquely endowed with the capacity to induce germ cell formation in the embryo, which it does by nucleating formation of the germ plasm and its germline determining RNP complexes, called polar granules. How oskar mRNA is transported and anchored at the posterior pole of the oocyte and its translation regulated is one focus of research in the lab.

We are also investigating the roles of other classes of RNAs, including long non-coding RNAs and piRNAs, and of non-canonical RNA binding proteins, in Drosophila embryonic development and neurogenesis. Drosophila, with its exceptional genetic tools, is also well suited to biochemical and cell biological investigation, including live imaging, of the processes of cell polarisation, mRNA localisation and translational control.

Future projects and goals
We combine genetics, biochemistry and a broad spectrum of cell biological and imaging approaches to study:

- Polariation of the cytoskeleton.
- The roles and regulation of cytoskeletal motors in RNA localisation.
- Assembly of transport-competent RNPs: the cis-acting RNA targeting elements and interacting proteins, how they assemble and associate with motor proteins.
- Translational regulation of localised mRNAs.
- Germ plasm assembly and function.

Selected references

The Ephrussi group aims to understand the mechanisms underlying RNA transport and localised translation – fundamental processes that promote the functional polarisation of cells during development.

Right: oskar mRNA on the move. Time projection of a squash of ooplasm from a stage 9 oocyte, imaged with TIRF microscopy. oskar mRNA (labelled with MS2-MCPGFP, shown in rainbow colours) utilises microtubules (labelled with mCherry-a1-tubulin and EB1-Cherry, shown in grey with cyan tips, indicating plus ends) to take fast, long linear runs.

Above: A Drosophila egg-chamber, showing colocalisation of oskar mRNA, Staufen protein and a microtubule polarity marker at the posterior of the oocyte.
Evolution of the nervous system in bilateria

As a ‘living fossil’, Platynereis represents an ideal connecting link between vertebrates and the fast evolving protostome models, Drosophila and Caenorhabditis.

Selected references

Previous and current research
We are intrigued by one of the remaining great mysteries in animal evolution: how did our central nervous system (CNS) come into existence? What did it look like at first and how did it function? We are especially interested in the CNS of an extinct animal known as Uribilatera, the last common ancestor of humans, flies and most other ‘higher’ animals that live today, which lived some 600 million years ago in the ocean.

Our lab has chosen to investigate a new molecular animal model, the marine annelid Platynereis dumerilii. As a ‘living fossil’, Platynereis represents an ideal connecting link between vertebrates and the fast evolving protostome models, Drosophila and Caenorhabditis. Genomic resources and molecular techniques have been generated that make it a model marine invertebrate for ocean biology and for organisational systems biology. Platynereis is amenable to high throughput imaging techniques and functional interference approaches, for example the first genetic knock-out lines have been generated. With the recent development of the PrImR (Profiling by Image Registration) resource, Platynereis has become the first animal model for which gene expression profiling data can be obtained in cellular resolution for the whole organism. Building on these resources, we have discovered that their brains harbour sensory-associative parts and a neurosecretory centre that corresponds to the vertebrate pallium and hypothalamus, respectively. These findings revolutionise our current understanding of brain evolution – a clear picture is emerging that the Platynereis brain harbours many cell types so far known only for vertebrates, but in a much more simple and different overall arrangement.

In an attempt to broaden our comparative approach, we have introduced two new model species to the lab, the lancelet amphioxus and the sea anemone Nematostella, representing distinct divisions of the animal kingdom: chordates and cnidarians. Amphioxus has a very simple brain uniting invertebrate- and vertebrate-like features. The Nematostella nervous system is very simple and is a good proxy for an early stage of nervous system evolution.

Future projects and goals
Our aim is to gain a systems view of the Platynereis brain and nervous system and to track the evolutionary history of all constituent cell types by identifying and investigating their evolutionary counterparts in sea anemone and amphioxus. This will involve investigations of cell type-specific gene regulatory networks in all species as well as neurobiological and behavioural approaches. In 2012, we began an ERC-funded project called BrainEvoDevo, which aims at generating a neuron-type atlas of the annelid larval brain. This atlas will combine neuronal morphologies, axonal projections and cellular expression profiling for an entire bilaterian brain. In order to generate the first cellular resolution expression atlas for a whole animal nervous system, involving early developmental as well as differentiation stages, we are working in collaboration with the Marioni group (page 87), the Keller lab (Janelia Farm Research Centre) and the Tomanacak lab (MPI for Molecular Cell Biology and Genetics). We are combining a full tracking of the developmental lineage for all cells at several developmental stages, with cellular resolution expression profiling by image registration and single-cell sequencing. Building on the Atlas, we will dissect Platynereis chemosensory-motor forebrain circuits by laser ablation of GFP-labelled single neurons, gene knockout studies, and behavioural assays based on microfluidics to explore duplication, divergence and expansion of neural circuits in central nervous system development and evolution. We are also interested in exploring population genetics and the variability of development and differentiation in different habitats and conditions. To this end, we are collecting strains of Platynereis and amphioxus as part of the TARA Oceans expedition and as an active member of the EMBL Oceans Team.

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Timing of mammalian embryogenesis

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Previous and current research
During an embryo’s journey from a single cell to a complex organism, countless patterning processes unfold with remarkable precision, spatially but also in respect to temporal sequence, or timing. This temporal aspect of embryonic development is the focus of our research. How is time measured during embryonic development and what extrinsic and intrinsic signals control this timing? How are embryonic oscillators/clocks employed during patterning? What are the dynamics of signalling pathways?

To approach these questions, novel methodologies are required (see video 1). We are generating novel real-time reporter mouse lines using knock-in technology that enables visualisation and quantification of temporal dynamics at different levels in the context of mouse embryonic development. Using in vivo imaging, we are focusing on the somite segmentation clock, an oscillatory system that is thought to control the formation of the pre-vertebrae that form periodically in a head-to-tail sequence within the paraxial mesoderm. In mouse embryos this clock, with a periodicity of around two hours, drives oscillatory activity of several signalling pathways (Wnt, Notch and Fgf signalling) in the developing mesoderm. We recently developed an ex vivo assay that, in combination with real-time imaging reporters, has become instrumental for our approach: the assay recapitulates mesoderm patterning, including segment formation and spatio-temporally controlled oscillatory signalling activities, within the simplified context of a monolayer of primary mesoderm cells put in culture (see figure & video 2).

Scaling and phase-shifted oscillators: One fundamental property of vertebrate segment formation is its ability to maintain proportions even when overall embryo size is experimentally altered, a process termed scaling. Intriguingly, scaling behaviour can be observed in the ex vivo assay system as well. This enabled us to identify a novel scaling mechanism employing phase-shifted oscillatory activity (Lauschke et al, 2013). How temporal devices, or oscillators, mechanistically encode spatial information for patterning constitutes a major interest in the lab. We are particularly interested in studying these problems at an integrated, higher-order level, so as to reveal emergent properties, incorporating mathematical modelling into our approach.

Oscillatory Wnt-signalling: This signalling pathway serves a multitude of evolutionarily conserved functions during development and has been shown to play an essential role during somite formation. Our novel real-time reporter system is designed to reflect oscillatory Wnt-signalling activity both at gene activity and at protein levels. This will enable us to determine how the striking oscillations of Wnt-signalling activity are generated in the first place and, moreover, to functionally test their role in embryonic patterning. We are particularly interested in identifying the intrinsic and extrinsic factors that are responsible for controlling these oscillations within the segmentation process.

Future projects and goals
- Quantitative (imaging) approach to understand the role of dynamic oscillatory signalling during patterning and scaling.
- Study of emergent properties of coupled oscillator populations.
- Discovery of oscillatory signalling phenomena during embryogenesis.

Selected references

Video links
http://s.embl.org/Vg1

Ex vivo cell culture model for mesoderm patterning and oscillations. a, b) Primary mesoderm cells retain undifferentiated PSM identity in the centre of the culture, before initiating a differentiation program in the periphery. c) Snapshot of time-series using fluorescent lunatic fringe reporter mouse line (LuVeLu), overlaid with time-projection of activity patterns (in green) seen during the time-lapse recordings (see video 2). d) Raw photon counts (measured in quadrant shown in c) demonstrating robust oscillatory activity for extended recording times. e) Time-space kymograph along dashed arrow in c). From this quantification, critical oscillation parameters, wave speed and phase-distributions can be calculated.
Cell dynamics and signalling during morphogenesis

Cell shape changes are of fundamental importance during embryonic development – how cells form and change shape during morphogenesis are the key questions addressed by the De Renzis group.

Selected references


Previous and current research

Tissue morphogenesis is triggered by shape changes in single cells or groups of cells. This remodelling depends on a complex interaction between cortical forces exerted by the actin cytoskeleton and membrane homeostasis (i.e. vesicular trafficking and lipid metabolism). We want to understand how membrane trafficking and cytoskeletal dynamics are regulated during morphogenesis and how this, in turn, impacts on specific cell and tissue behaviour. To this end, we combine high-resolution imaging methods with genetics and biochemistry using the early Drosophila embryo as model system (see figure 1 and video link).

We have recently developed a modified form of total internal reflection fluorescence (TIRF) microscopy to follow apical surface dynamics in live embryos with unprecedented spatio-temporal resolution. Using this approach we have identified a novel endocytic pathway controlling the morphology of the apical surface during epithelial morphogenesis (figure 2), thus demonstrating for the first time that endocytosis directly controls cell and tissue shape. We are now using similar high-resolution imaging methods in combination with electron tomography to study the involvement of endocytosis in the regulation of cell signalling and membrane remodelling during tissue morphogenesis.

We are also interested in characterising the impact of lipid metabolism during morphogenesis. Using a chemical genetic approach we uncovered an exciting link between so-called ‘lipid induced phenotypes’ and developmental gene activities underlying the regulation of cell and tissue shape.

Finally, we are developing new optogenetic tools to control protein activity with light during tissue morphogenesis with high spatio-temporal precision.

Future projects and goals

Using a combination of imaging, genetics and optogenetic approaches we wish to elucidate how machineries controlling intracellular trafficking re-organise during differentiation and how this in turn impacts on global changes in tissue morphology.

Figure 1: Cross-section of a Drosophila embryo during late cellularisation (left panel) and ventral furrow formation (right panel) stained with antibodies against b-catenin (red), Notch (green) and Delta (blue). Embryos are oriented with the ventral side facing down and dorsal up. Cells on the ventral side are elongated along the apico-basal axis (left panel arrowhead) compared to their dorso-lateral neighbours. Endocytosis of Notch and Delta is specifically up-regulated in ventral cells during invagination.

Figure 2: Application of TIRF-M imaging to early Drosophila embryo allowed uncovering a prominent endocytic pathway controlling the morphology of the apical surface during epithelial development. Rab5 endosomes (purple), plasma membrane (green).
Previous and current research

In addition to providing us with the air we breathe, the food we eat and much of the energy and materials we use, plants exhibit a unique beauty associated with their strikingly symmetrical patterns of development. Multicellularity also evolved independently in plants giving us an opportunity to compare and contrast the developmental strategies used in different kingdoms.

Lateral organ formation in the model plant species *Arabidopsis thaliana* provides an ideal system for investigating such questions, since organ formation involves the coordination of cell polarity, gene expression and morphogenesis. Our recent work reveals that patterns of cell polarity control both morphogenesis at the cellular level as well as at the tissue level. This integration occurs through the co-alignment of microtubule arrays with the polar localisation patterns of the auxin efflux carrier PIN1. The microtubule cytoskeleton regulates growth direction at the cellular level, while PIN1 works to concentrate the hormone auxin at the tissue level to localise growth. Our data so far suggests a role for mechanical stresses in orienting these factors and we are further investigating this possibility. Interestingly, we have also found that the patterns of cell polarity associated with organogenesis correlate spatially with particular patterns of gene expression normally associated with the dorsal and ventral cell types of lateral organs. This raises the question of whether these expression domains play a causal role in organising cell polarity patterns and, in turn, whether these polarity patterns influence dorsiventral gene expression. This rich interplay is one of our prime focuses.

Current projects and goals

Establishment and function of dorsiventral boundaries (ERC funded): Previously we developed confocal-based methods for imaging growing plant tissues, enabling us to obtain dynamic high-resolution data for protein localisation and gene expression (making full use of the different GFP spectral variants). By incorporating such data directly into mathematical models we aim to develop an explicit understanding of the complexity underlying patterning processes associated with dorsiventral cell-type specification. Our main questions include: How do dorsiventral gene expression boundaries regulate organ morphogenesis and positioning (e.g. cell polarity patterns)? How are dorsiventral gene expression boundaries established and regulated?

Like animals, plants can also re-pattern their tissues in response to wounding. Wounding also causes dramatic changes to dorsiventral patterning, although the mechanisms by which this occurs remain unknown. Our recent results show that cell polarity patterns respond dramatically to wounds, suggesting this cellular response may play an important role in tissue reorganisation. We aim to investigate this possibility using two-photon induced ablation and DSLM microscopy.

Confocal projection showing polar localisation of the auxin efflux carrier PIN1 fused to GFP. At organ inception PIN1 polarities are directed away from adjacent organ sites and towards the new site.
Systems-level understanding of early mammalian development

Looking at the molecular, cellular and systems levels, the Hiragi group studies how, early in mammal development, the embryo is shaped from a spherical mass of cells.

Selected references


Motosugi, N., et al. (2005). Polarity of the mouse embryo is established at blastocyst and is not prepatterned. Genes Dev., 19, 1081-92

Previous and current research

A fundamental question in biology is the mechanism by which the embryonic asymmetry is established during development. In contrast to many organisms in which embryonic development is driven by determinants localised asymmetrically in the egg, mammalian eggs lack polarity and thus symmetry has to be broken during early embryogenesis. This symmetry breaking process in mammalian embryos results in the formation of the blastocyst, composed of the inner cell mass surrounded by the trophectoderm. Despite its importance, the molecular mechanism of blastocyst patterning has long been elusive. How is the symmetry broken in the mammalian embryo? How is the definitive embryonic pattern established?

We have developed a live-imaging system for mouse pre-implantation embryos, demonstrating unexpectedly high dynamicity, stochasticity and molecular heterogeneity (figure 1) during early embryogenesis. Our recent study also demonstrated that centriole is generated de novo during the gradual transition from meiosis to mitosis at the pre-implantation stage (figure 2). Taken together, and in light of its highly regulative capacity, early mammalian embryo may be viewed as a self-organising system, patterning through stochastic processes in a particular structural context. These features suggest that, in order to fully understand the mechanisms of early mammalian development, it will be essential to address how the diverse inputs acting on individual cells are integrated in the embryo at the systems level. Thus we have recently established necessary tools and multi-disciplinary strategies, including fluorescence gene-trap mice that allow quantitative characterisation of gene-expression dynamics, and mapping of cellular mechanical properties during morphogenesis. Furthermore, our recent single-cell transcriptome analysis led us to propose a new model for symmetry breaking during embryogenesis, in which stochastic cell-to-cell gene expression variability followed by signal reinforcement progressively and antagonistically segregate lineages within initially equivalent cells.

Overall we aim at understanding principles and robustness underlying early mammalian development.

Future projects and goals

We adopt a wide variety of experimental strategies including embryology, molecular genetics, live-imaging, cell physics and modelling in order to address fundamental questions in development and cell biology at a molecular, cellular and systems level. Our goals include:

- identification of the symmetry-breaking cue in the mouse embryo;
- investigating the significance of molecular heterogeneity in lineage segregation;
- understanding the role of mechanical cues in embryogenesis;
- identification of the trigger and mechanism for centriole biogenesis in vivo.

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Figure 1: Unprecedented molecular heterogeneity during mouse blastocyst patterning. Cells expressing Nanog (green), Gata6 (red) or Serpinh1 (blue).

Figure 2: Microtubules (green) self-assemble into a mitotic spindle from multiple microtubule-organising centres (red) in the absence of centriole, during progressive transition from meiosis to mitosis in early mouse development.
Previous and current research

During brain development, neurons are generated in great excess and only those that make functional connections survive, while the majority is eliminated via apoptosis. Such huge numbers of dying cells pose a problem to the embryo, as leaking cell contents damages the surrounding environment. Therefore, the clearance of dying cells must be fast and efficient and is performed by a resident lineage of ‘professional’ phagocytes, the microglia. These cells patrol the entire vertebrate brain and sense the presence of apoptotic and damaged neurons. The coupling between the death of neurons and their phagocytosis by microglia is striking: every time we observe dead neurons we find them already inside the microglia. This remarkable correlation suggests a fast acting communication between the two cell types, such that microglia are forewarned of the coming problem. It is even possible that microglia promote the controlled death of neurons during brain development. Despite the importance of microglia in several neuronal pathologies, the mechanism underlying their degradation of neurons remains elusive.

The zebrafish Danio rerio is an ideal model system to study complex cell-cell interactions in vivo. As the embryo is optically transparent, the role of molecular regulators identified in large-scale forward and reverse genetic screens can be studied in vivo. Moreover, a key advantage of the system is that zebrafish microglia are extremely large, dynamic cells that form a non-overlapping network within the small transparent fish brain. Labelling microglia, neurons and organelles of the microglial phagocytic pathway simultaneously in the living zebrafish embryos allows us to image, for the first time, the entire microglial population in order to study the interaction between neurons and microglia.

Future projects and goals

Despite the importance of microglia in several neuronal pathologies, many fundamental questions concerning microglial-neuronal interactions remain unaddressed. How these cells collectively ensure that the entire brain is surveyed and how they react to damage with high precision is still entirely unknown. Recent findings suggest that diffusible molecules such as lipids and nucleotides could attract microglia in response to neuronal apoptosis and injury, respectively. While these molecules can trigger dynamic changes in microglia motility in vitro, elucidating how their activity is controlled within the intact brain, both in space and time, remains the most important challenge in understanding this fascinating biological problem. We aim to further exploit the massive imaging potential of the transparent zebrafish embryo for studying microglial biology in vivo. By combining forward and reverse genetic approaches with quantitative imaging technology, we will directly address the mechanisms underlying the attraction of microglia towards apoptotic, sick and injured neurons. By applying cutting-edge microscopy technology, such as the SPIM/DSLM (Selective Plane Illumination Microscopy), we will image all interactions between neurons and microglia and derive from this time-lapse analysis real quantitative data in a spatiotemporal manner.

Selected references

Gene regulation and genome architecture

Previous and current research
The patterning of the embryo and the specification of its different cell types are driven by the implementation of cell-specific gene expression programs. In vertebrates, the cis-acting elements that regulate transcription can be located hundreds of kilobases away from the genes they control, particularly for genes with important functions during development. Because of this, the genome appears to be composed of intermingled arrays of unrelated genes and cis-regulatory elements. Therefore, the mechanisms that regulate enhancer-promoter interactions are essential to transform this apparent genomic and regulatory conundrum into gene- and tissue-specific expression programs. Recent data reveal that genomic loci adopt specific chromatin structures and conformations in the nuclei of different cell types, correlating with differential gene activity. Yet, the cis-acting genomic elements that determine how a genomic locus folds into specific structural and regulatory architectures, and the precise roles of the chromatin, protein-complexes and non-coding RNAs suggested to contribute to this process, are still unclear.

Our lab has developed several experimental approaches to explore the regulatory architecture of the mouse genome and characterise functionally the mechanisms that organise it. Towards this aim, we have established an efficient in vivo system that, through the combined used of transposases and recombinases, allows the re-engineering, in a systematic manner, of the mouse genome. With this approach, we generated a unique resource comprising hundreds of mouse strains carrying regulatory sensors throughout the genome, and series of specific chromosomal rearrangements in selected loci. This genomic resource enables us to dissect functionally the genomic information and the mechanisms that organise a linear genome into structurally distinct domains and chromatin loops, so as to implement long-range specific regulatory interactions.

Future projects and goals
Structural conformation and regulatory organisation of the genome: By combining advanced genomic engineering with chromatin profiling (ChIP-Seq) and conformation analyses (4C-chromatin conformation capture, super-high resolution FISH), we aim to learn how the genomic organisation of a locus determines the specific chromatin structures and conformations that it adopts in the nucleus, and determines their functional significance in the context of a developing embryo.

Regulatory architecture, disease and evolution: Our mouse models provide insights into the consequences of structural variations or chromosomal aneuploidies found in humans, both at the phenotypic and molecular level. Comparison of the regulatory architecture of developmental gene loci between different species can reveal how large-scale changes in chromosomal organisation may have contributed to evolution of body forms.

Abnormal skull development in mice with a deletion of distal enhancers engineered by in vivo recombination (see Marinic et al., 2013).
In eukaryotes, many steps of gene expression, such as transcription and RNA processing, take place in the structurally complex environment of the nucleus and often involve remodelling of chromatin into active and inactive states. Messenger RNAs, once exported from the nucleus, undergo additional regulatory steps. Their translation results in the production of proteins, whose functions define the characteristics of different cell types, or cellular phenotypes. Not all RNAs are translated, however. In recent years, multiple types of non-coding RNAs have been discovered that display diverse functionality. Genetic variation in non-coding and protein-coding genes alike, as well as the regulatory elements that govern their expression, can adversely affect the function of these genes, leading to diseases such as cancer. Groups within the Unit are investigating various aspects of genome biology in order to understand these processes leading from genotype to phenotype.

A notable strength of the Unit is its ability to address questions at different scales, ranging from detailed mechanistic studies (using biochemistry, genetics, microfluidics and chemistry) to genome-wide studies (using functional genomic, proteomic and computational approaches), often by developing new enabling technologies. For example, the development and integration of chemistry and microfluidic devices with the recent advances in next-generation sequencing will facilitate major advances in these areas in the coming years. Global, dynamic and quantitative measurements of biological molecules at all levels (DNA, RNA, proteins, cells, organisms, etc) as well as the integration of hypothesis and discovery-driven research characterise the Unit. The synergy between computational and wet-lab groups provides a very interactive and collaborative environment to yield unprecedented insights into how genetic information is ‘read’ and mediates phenotype through molecular networks.

Eileen Furlong
*Head of the Genome Biology Unit*
Regulatory networks driving cell fate decisions: Dissecting the logic

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Previous and current research
Development is driven by the establishment of complex patterns of gene expression at precise times and spatial locations. Although a number of mechanisms fine-tune expression states, it is initially established through the integration of signalling and transcriptional networks converging on enhancer elements, or cis-regulatory modules (CRMs). Understanding how CRMs function is therefore central to understanding metazoan development and evolutionary change. Although there has been extensive progress in deciphering the function of individual regulatory elements, how these modules are integrated to regulate more global cis-regulatory networks remains a key challenge. Even in the extensively studied model organism Drosophila, there are no predictive models for a transcriptional network leading to cell fate specification.

Our research includes studies of the mechanism of enhancer function (figure 1) and the interplay of transcription factors and chromatin state (figure 2), as well as studies of how gene regulatory networks control development and how network perturbations lead to specific phenotypes. To address this we integrate functional genomic, genetic and computational approaches to make predictive models of transcription and developmental progression. We use Drosophila mesoderm specification into different muscle primordia as a model system. The relative simplicity of the fly mesoderm, in addition to the number of essential and conserved transcription factors already identified, make it an ideal model to understand cell fate decisions at a systems level.

Future projects and goals
Chromatin remodelling during cell fate decisions: We have developed a very accurate and sensitive method to investigate cell type-specific changes in chromatin status and chromatin binding protein occupancy in the context of a multicellular embryo’s development (figure 2). We are currently using this method to examine the interplay between changes in chromatin remodelling with dynamic changes in transcription factor occupancy and developmental transitions.

Variation and plasticity in cis-regulatory networks: Variation in cis-regulatory elements can affect gene expression and account for individual differences in phenotypes, like taste sensation and olfactory sensitivity. However, little is known about how much variation in gene expression or transcription factor function can be tolerated for essential developmental processes during embryonic development. We plan to investigate this by extending our current knowledge of the transcriptional network regulating cell fate choices during mesoderm development to many Drosophila individuals (isogenic Drosophila strains) whose genomes have been fully sequenced.

Predictive models of embryonic development: Our previous work demonstrated that only using information on combinatorial occupancy of transcription factors is sufficient to predict spatio-temporal cis-regulatory activity (Nature, 2009) and that information on chromatin state and RNA polymerase II occupancy on enhancers can predict the precise timing and location of active enhancer elements de novo (Nature Genetics, 2012). We plan to build on this by predicting a gene’s expression. Our ultimate goal is to use this systems-level approach to make predictive models of embryonic development and the effect of genetic perturbations. Working in Drosophila allows us to readily test the predicted outcome of network perturbations on embryonic development.

The Furlong group aims to understand fundamental principles of transcription, focusing on the processes that determine what a cell becomes during embryonic development.

Selected references

Figure 1: Enhancers can function by highly cooperative transcription factor occupancy using very flexible motif content and organisation (Junion, Spivakov, et al. 2012).

Figure 2: Chromatin state and RNA polymerase II occupancy on enhancers (yellow) is highly predictive of enhancers’ activity and is very dynamic, mirroring that of dynamic enhancer usage during development (Bonn, Zinzen, Girardot, et al. 2012).
The Steinmetz group bridges diverse domains of genome science, from deciphering the structure and function of genomes to the application of these insights in understanding diseases.

Selected references

Previous and current research
One of the most daunting challenges in medicine is the complex nature of most common diseases (including cancer, diabetes, and heart disease) due to interactions between multiple genetic variants and environmental influences. Our research is directed at understanding such complex traits; to do so, we develop novel genomic approaches to investigate the molecular processes that link genotype to phenotype, identify the underlying factors, and quantify their contributions. We investigate variation at the level of the genome, transcriptome, and proteome, which we integrate with higher-level phenotypes. Our projects are mainly in the following areas:

Function and mechanisms of transcription: We have developed several technologies to characterise pervasive transcription at the genome-wide level as well as its functional impact. We are interested in the function and regulation of non-coding RNAs, antisense transcription, and the molecular phenotypes that arise from transcriptome complexity. Recently, we discovered extensive variation in the start and end sites of transcript molecules produced by each gene by developing a novel technique to map full-length transcript isoforms genome-wide (figure 1).

Systems genetics: We have worked extensively on dissecting the genetic basis of complex phenotypes. We are interested in studying the network-level consequences of genetic variation and learning to predict phenotype from genotype. Recently, by studying genetic interactions with the environment, we have defined new experimental and statistical techniques that facilitate the distinction of genes that play a causal role in mediating genetic effects on phenotype (figure 2).

Mitochondria: Using an array of systematic and biochemical approaches, we investigate the effects of genetically perturbing mitochondrial function and strategies to rescue these perturbations. We also study mitochondrial genetics, including recombination of mitochondrial DNA in yeast and the relocation of mitochondrial genes to the nucleus.

Future projects and goals
We are integrating multiple layers of molecular data in order to understand how the genome is read for function. Using novel algorithms, intervention points can be identified from such data that can be targeted to modulate phenotypes of interest. We are also following up on our studies of transcriptional regulation through targeted investigations of the interplay between epigenetics and transcription, the functional consequences of complex transcriptome architecture, and its contribution to single-cell heterogeneity. Ultimately, by integrating genetics, genomics, systems biology, and computational modelling, we aim to develop approaches that unravel disease mechanisms and predict effective therapeutics, enabling personalised and preventive medicine.
**Computational biology and genomics**

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**Previous and current research**

Our aim is to understand biological systems through large-scale data acquisition and quantitative models. Our main tool is statistics – the science of computing with uncertainty, drawing conclusions from incomplete and noisy data. Together with its sister discipline, machine learning, it helps humans to discover patterns in large datasets and to infer underlying mechanisms, and predictive and causal relationships.

Our projects have a mix of two objectives: method development and biological discovery – to understand genetic and phenotypic variation between individuals on a genome-wide scale. We have projects in the areas of gene expression and regulation, in the genetics of complex phenotypes and genetic interactions, in cell division and cell migration, and in cancer genetics.

The group brings together expertise from quantitative disciplines – mathematics, statistics, physics and computer science – and from different areas of biology to design and analyse genomic experiments and their biological interpretation. Computational and statistical methods are at the heart of systematic, large-scale experimental approaches. Our aim is to develop high-quality methods of general applicability that can be widely used in genomic research. We regard the publication of scientific software as an integral part of the publication of new methodical approaches and contribute to the Bioconductor Project, an open source software initiative to provide tools for the analysis and study of high-throughput genomic data.

**Future projects and goals**

An emphasis of the group’s work is on project-oriented collaborations with experimenters. We aim to develop the computational techniques needed to make new types of experiments feasible and to turn the data into biology. Among our current projects are:

- RNA-Seq analysis.
- 3D nuclear organisation by HiC-related techniques.
- Cancer genomics and precision oncology.
- Image analysis for systems biology: measuring the dynamics of cell cycle and of migration of individual cells under normal conditions and many different perturbations (RNAi, drugs).
- Systematic mapping of molecular interactions and life cycles within single cells.
- Open source software for genomics, high-throughput phenotyping and statistical bioinformatics, to support reproducible research and wide dissemination of state-of-the-art methods.

**Selected references**


Phosphatase chemistry and biology

The Köhn group combines molecular biology, biochemistry and synthetic chemistry to develop new approaches to study phosphatases, which can play a major role in cancer.

Selected references

Previous and current research

Within intracellular signalling networks, phosphatases are counter players of kinases and play crucial roles in health and disease. The investigation of phosphatases is challenging, which is also due to the lack of tools to selectively study particular phosphatases. Understanding of phosphatase function, regulation and substrate interaction is therefore still quite limited. Our main interest is thus to control and investigate phosphatases using interdisciplinary approaches.

Specifically, we are interested in the phosphatase of regenerating liver (PRL) enzymes, in particular PRL-3. We study biological pathways and roles of this family using biochemical and molecular cell biology approaches, and we aim to design inhibitors for PRL members. We have observed phosphoinositide-phosphatase activity in vitro for one member, PRL-3 (McParland et al., 2011). In this regard, we developed a solid phase synthesis strategy that accelerates access to phosphoinositides and their analogues (figure 1a, Bru et al., 2012). We aim to obtain a detailed picture of substrate specificities of lipid phosphatases in biochemical structure–activity relationship (SAR) studies using a library of phosphoinositide analogues to support designing specific inhibitors of lipid phosphatases.

Another interest is the tool development for protein phosphatase-1 (PP1), a ubiquitous phosphatase that is responsible for a majority of all dephosphorylation reactions on Ser/Thr inside cells and is involved in many processes such as mitosis and cell cycle regulation. We have developed a selective peptide-based PP1 activator (figure 1b), which is an enabling tool to gain novel insights into PP1 biology (Chatterjee et al., Angew. Chem. Int. Ed. 2012). We follow up several strategies for the refinement of this activator and for the development of further PP1-directed chemical tools.

To support the research on phosphatase-kinase-substrate networks, in collaboration with the Wilmanns (page 112) and Thornton (page 78) groups, we have created the human DEPhOlyphosphorylation Database: DEPOD (figure 2). We have applied DEPOD data to re-classify the human phosphatome and to analyse phosphatase substrate specificities and their relation to kinases (Li et al., Sci. Signal. 2013).

Future projects and goals

Studying PRL biology will remain a focus of the laboratory in the future, with the aim to understand the underlying mechanisms of oncogenesis caused by these phosphatases. We continue to develop chemical methods to enable applying peptides and inositides as phosphatase modulators inside cells. Designing modulators for the highly complex serine/threonine phosphatases is another long-term goal. Furthermore, we will continue to develop, extend and maintain DEPOD.

The lab consists of an equal number of molecular biologists and organic chemists on the graduate student and postdoctoral level. The combination of biology and chemistry not only opens up new ways to approach the challenging phosphatase research, but also broadens views and skills of every lab member.

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Figure 1: (A) Solid phase synthesis of phosphoinositides for the preparation of libraries for SAR studies with lipid phosphatases (Bru et al., Chem. Sci. 2012). (B) Selective activators of PP1 in cells enable us to gain new insights into PP1 biology (Angew. Chem. Int. Ed. 2012, Chem. Biol. 2013).

Figure 2: DEPOD, the human dephosphorylation database (Li et al., 2013) – see www.depod.org.
Previous and current research

Genetic variation is a key reason why we differ from one another and can be used as starting point to unravel disease mechanisms. Recent advances in DNA sequencing technology have facilitated the characterisation of genetic variation at genome-wide scale. Our group is devoting efforts to characterising the extent, origin, and functional consequences of DNA variation, with a particular focus on genomic structural variants (SVs) such as deletions, duplications, inversions and translocations – the most consequential type of heritable genetic variation in humans in terms of the sheer number of nucleotides affected. Germline and somatic SV classes have been linked to numerous heritable diseases and different cancer types. We are pursuing research using both laboratory and computational techniques, a ‘hybrid’ approach that allows us to combine data generation and analysis with hypothesis generation and testing in experimental model systems.

A cancer genome study that we recently performed revealed that the development of medulloblastoma, the most common malignant brain tumour in children, frequently involves a remarkable process known as chromothripsis, where localised chromosomal shattering and repair occur in a one-off massive DNA rearrangement event (see figure). We also recently made progress in understanding the etiology of early onset prostate cancer, the initiation of which we found to be largely driven by androgen-mediated somatic SVs. Our group also participates in genome research consortia such as the 1000 Genomes Project, with the aim of contributing to the generation of fine-resolution genetic variation maps in humans that can later be related to functional genomics data. We also investigate SVs in non-human species to pursue genetic variation studies in the context of evolution.

Within the Whole Genome Sequencing (WGS) Pan-Cancer Analysis Initiative of the International Cancer Genome Consortium (ICGC) we have begun investigating whole genome, DNA methylome, and transcriptome sequencing data of ~2000 cancer patients. Using integrative analyses we aim to unravel commonalities and discrepancies between cancer types at the molecular level, to facilitate the molecular classification of malignancies with a potential impact on diagnostics and treatment.

Future projects and goals

- Uncovering genetic determinants for the development and progression of cancer in humans, and studying commonalities and differences between tumour types.
- Development of wet-lab and in silico approaches for deciphering the molecular origin and function of SVs in humans and model organisms.
- Constructing near-complete human genome variation maps using second and third generation sequencing technologies.
- Deciphering the mechanistic basis of chromothripsis, an SV process particularly abundant in highly aggressive malignancies.

Selected references

Functional proteomics

The Krijgsfeld team uses biochemical and mass spectrometric approaches to understand the dynamics of protein expression and interaction in the context of cellular differentiation and stress response.

Selected references

Previous and current research
Proteins fulfill most of the functions that are crucial in establishing cellular phenotypes. In addition, it is becoming increasingly clear that proteins rarely act alone, but that they constitute intricate networks, both among themselves and with other biomolecules. This system is both robust and dynamic, allowing a cell to respond to external cues, or to develop from an embryonic to a mature state. Our interest is in understanding cellular properties from this perspective, realising that one needs to study proteins collectively rather than in isolation, and dynamically rather than under static conditions.

Our research is centred on quantitative proteomics, combining biochemistry, mass spectrometry, analytical chemistry, and bioinformatics, applied to various biological systems (yeast, Drosophila, mammalian cells). Our main interest is to understand how changes in protein expression, localisation and interaction underlie processes in stress-response, differentiation and reprogramming. For instance, large-scale proteomic experiments enable us to characterise the proteomes of highly purified mouse hematopoietic stem cells and progenitor populations obtained by fluorescence-activated cell sorting, generating novel insights in the initial steps of hematopoiesis in vivo. Furthermore, we have performed time course analyses quantifying the proteome changes in fibroblasts during their reprogramming to induced pluripotent stem cells (iPSCs), identifying and functionally validating proteins that are key in the gain of pluripotency. Apart from these large-scale analyses of intracellular proteomes, we have developed new tools to study secretory proteins and their role in cell signalling and communication. Furthermore, we are interested in regulatory principles of transcriptional activation and protein turnover in the face of developmental processes or response to stress. We are therefore developing novel techniques to identify proteins that interact with regulatory domains in the genome, both in vivo and in vitro. In doing so, we aim to identify proteins that drive (or inhibit) transcription in a gene- and condition-specific manner, for example to understand how transcription of developmentally important genes is controlled. To further explore the link between genome regulation and protein output, we study protein turnover taking yeast as a model system. By determining protein synthesis and degradation proteome-wide and across a range of growth conditions, we aim to construct models of how protein homeostasis is maintained.

Future projects and goals
- Develop new tools to study protein-DNA and protein-RNA interactions to identify and functionally characterise proteins that regulate transcription and translation.
- Integration of proteomics and next-generation sequencing to understand the molecular basis of protein homeostasis.
- Study cellular communication via secretory proteins.

Changes in protein expression during reprogramming of fibroblasts (A, B) leading to the formation of induced pluripotent stem cells (C).
Previos and current research

Working on the micro-scale offers some unique advantages:

- Drastically increased throughput (processing up to a million samples an hour).
- Superb spatio-temporal resolution (assays can be carried out on micro-metre length scales and sub-millisecond timescales).
- Low material consumption, enabling single-organism, single-cell, or even single-molecule assays.

During the past couple of years we have developed powerful microfluidic platforms for cell-based and (bio)chemical assays. In our laboratory, we perform all steps ranging from the design and manufacturing of microfluidic chips and detection systems to the cultivation and study of human cells and multicellular organisms (prior knowledge in microfluidics is not obligatory for joining the group). Furthermore, we are interested in combinatorial chemistry, perform computational fluid dynamics simulations, and develop novel software controlling our microfluidic systems.

For many applications we use two-phase microfluidics, in which aqueous droplets within an immiscible oil phase serve as miniaturised reaction vessels. As they can be generated at kilohertz frequencies, they are of particular interest for high-throughput screens. Furthermore, the small assay volumes (pico- to nanoliters) facilitate the obtainment of high concentrations of nucleic acids (mRNA, DNA) or proteins (for instance, secreted antibodies) from individually encapsulated cells, paving the way for single cell assays. We also use continuous-phase microfluidics to generate laminar flow patterns, where we expose cells and organisms (or even small parts thereof) to different chemical environments. Amongst other applications, this allows the analysis of signalling events in developing embryos.

Future projects and goals

Having a comprehensive microfluidic toolbox at hand (and expanding it continuously), we are now focusing on applications in three different research fields:

Biomedical applications: Droplet-based microfluidics enables functional antibody screening at very high throughput (several hundred thousand antibodies per experiment). We want to use this technology to identify therapeutic antibodies, starting with primary plasma cells from immunised mice or even human disease (such as HIV and HCV) survivors. In parallel, we exploit a reversed experimental setup to derive potential HIV vaccine candidates (in close collaboration with the International AIDS Vaccine Initiative). Furthermore, we are actively developing microfluidic approaches for personalised cancer therapy.

Cell biology: We will perform large-scale chemical perturbations to identify pathway interactions in stem cell differentiation and carcinogenesis. These microfluidic chemical genetics approaches require only minute numbers of cells and are hence compatible with primary cells or even patient biopsies.

Genomics: We are developing microfluidic modules for single-cell barcoding and sequencing. Furthermore, we are setting up integrated microfluidic ChiPseq platforms allowing for the analysis of less than 5000 cells. Notably, some of our modules (such as our microfluidic sonicator for the shearing of gDNA and cross-linked chromatin) have already been commercialised.
Selected references


Previous and current research

Chromatin, the faithful association of genomic DNA with histone proteins, exists as the physiological form of our genome and the substrate for processes that regulate cellular gene expression. An increasing body of evidence suggests that epigenetic mechanisms influence gene expression profiles with far-reaching implications for human biology, health and disease. Recent advances in genome sequencing have pinpointed genetic mutations associated with human disease at base pair resolution and further afford us new opportunities to study disease mechanisms at the molecular level. Numerous diseases are associated with mutations in genes that encode for chromatin-binding and/or chromatin-modifying enzymes, which together act as epigenetic regulators. Combining neurobiology and chromatin biology, we aim to study the molecular mechanisms that link genetic mutations encoded in epigenetic regulators to the widespread chromatin alterations associated with brain diseases. A central question grounding our research is how the chromatin modification network engages in brain development, function and disease.

Previously, colleagues and I studied a cellular pathway of ischemia-induced neuronal death, and showed that the transcriptional repressor REST causes epigenetic remodeling and repression of multiple target genes including AMPA receptor in postischemic neurons. We further demonstrated that REST knockdown prevents neuronal death in a clinically relevant in vivo model of ischemia. During the past few years, we uncovered the localisation and function of a histone H3 variant, H3.3. We found that H3.3, guided by distinct chaperone systems, marks the genomic regions of histone turnover. We mapped the genome-wide localisation of H3.3 in mouse embryonic stem cells (mESCs) and neuronal precursor cells, and further expanded to terminally differentiated neurons for studying its functional role in promoting neuronal plasticity. In addition, we revealed the molecular mechanisms that underlie DNA methylation, specifically, the interplay between histone post-translational modifications and DNA methylation, and identified the biological function of this interaction in cell lineage specification.

Future projects and goals

We aim to study chromatin regulation, its interpretation during brain development, and its misinterpretation in relation with brain cognitive and developmental diseases. We will utilise differentiating neurons from mESCs and human induced pluripotent cells (hiPSCs) to model developmental stages and facilitate necessary genetic manipulation/engineering. Collectively, defining the ‘epigenetic landscape’ – both in normal and abnormal brain cells – will help provide novel targets for therapeutic intervention for cognitive and developmental diseases of the brain.

Our research projects are to:

- Determine combinatorial histone modifications that link de novo DNA methyltransferase location/function during neuronal lineage commitment.
- Identify the location/function of mutated histones and epigenetic regulators specific to cognitive deficits, and explore alterations of the epigenetic landscape in developing neurons.
- Investigate PHD (Plant Homeo Domain)-containing epigenetic regulators that integrate specific signalling pathways into developmental transcription programs.
Previous and current research

The recent explosion of genomic sequence information provides a first step towards better understanding diverse bacteria, but also makes it crucial to develop large-scale phenotyping approaches to characterise functions of novel genes and to map them within pathways. Research pioneered in yeast indicates that some of the most powerful global phenotyping approaches are based on scaling up long-standing concepts in biology. Epistasis genetic interactions, for instance, assess how the function of one gene depends on the function of another, while chemical genetic interactions measure how the function of one gene affects cellular responses to chemical stress. We have developed quantitative high-throughput versions of these approaches in E. coli, and used them to obtain a mechanistic understanding of key aspects of its envelope function and architecture.

The bacterial envelope is sandwiched between two membranes in gram-negative bacteria and constitutes the interface of the bacterium to the environment. This compartment acts as a protective barrier, but also allows bacteria to sense changes in, and communicate with, their environment and is vital for pathogenesis, cell morphogenesis and cell developmental programs. Although many envelope structural components have been characterised, we often have limited information on how their biosynthesis and transport are interconnected, regulated, or linked to the overall status of the cell, how they sense perturbations, and how signals are transduced to achieve homeostasis.

Working at the intersection between systems genomic biology and mechanistic molecular biology, we have discovered key missing players of major envelope components, uncovered niche-specific regulation of conserved envelope processes, identified linking proteins that allow coordination between processes and mapped network rewiring under different stresses. Specific examples of the biological stories we have uncovered include: i) identification of the first regulators controlling peptidoglycan synthesis outside of the cell, thereby providing an opportunity for outer membrane status to influence the fundamental shape forming structure in bacterial cells; ii) discovery of a new player that orchestrates outer membrane constriction during cell division; iii) a mechanism to transduce information about cell size shape to the transcriptional machinery; iv) a mechanism to control the activity of lipoproteins.

Future projects and goals

Our main goal is to expand our efforts in two directions. First, we want to develop analogous high-throughput methods for other bacteria. This will enable us to generate comprehensive interaction datasets in different bacteria, enrich them with other available genome-wide resources, and ultimately use them to create inferences about common and individual characteristics of biology in those organisms. Second, we plan to expand this technology to report on more social characteristics of bacteria. This will enable us to probe interspecies interactions and interactions with the host, shedding light on different mechanistic aspects of these interactions. Our specific aims include: i) generation of global genetic and chemical genetic interaction profiles for different pathogen and commensal bacteria and integration with other available global phenotyping data; ii) cross-species comparisons to map key features of divergence in the organisation of the bacterial envelope; iii) development of high-throughput approaches to study bacterial interspecies interactions and bacterial-host interactions.

Selected references


Dissecting bacterial lifestyle and interspecies interactions with systems approaches

The Typas group develops and utilises high-throughput methods to study the cellular networks of different species of bacteria, and how these bacteria interact with the environment and with each other.

Two novel niche-specific lipoproteins control peptidoglycan synthesis from the outside of the sacculus (Typas et al., Cell, 2010).
The Unit pursues an ambitious research programme with a strong basis in integrated structural systems biology and a far-reaching computational component that bridges into various areas of biology. A wide spectrum of expertise allows the Unit to tackle problems at different ranges of spatial resolution, connecting atomic structures and dynamic information obtained by X-ray crystallography and NMR with medium-range resolution from single particle electron microscopy, and cellular imaging obtained by electron tomography and light microscopy. Dedicated large scale biochemistry, proteomics, chemical biology, biophysics, and cell biology approaches complement the structural biology activities and, in conjunction with a wide range of innovative computational biology activities, are integrated into a comprehensive description of biological function.

Within the Unit, there is a continuing interplay between groups with expertise in different methodologies. This reflects our belief that a combination of structural and functional studies is the most rewarding route to an understanding of the molecular basis of biological function, and that computational biology is essential to integrate the variety of tools and heterogeneous data into a comprehensive spatial and temporal description of biological processes. Along those lines, groups in the Unit pursue a few common large projects. One example is the comprehensive structural and temporal description of an entire cell at almost molecular resolution. It goes hand in hand with the application of and integration of various ‘omics’ approaches to the small bacterium *Mycoplasma pneumoniae*, by characterising its dynamic protein organisation and merging this molecular information to cellular, high-resolution tomograms. In the thermophilic fungus *Chaetomium thermophilum* spatial and temporal networks will be deduced using multidisciplinary approaches including structural studies, large scale biochemistry and computational biology. Together, they will provide insight into eukaryotic thermophily at the molecular and cellular level.

Currently, the Unit consists of twelve research groups covering a broad methodological spectrum. The core technologies include electron microscopy (three groups), X-ray crystallography (two groups), NMR (one group), chemical biology (two groups) and computational biology (four groups). However, each group reaches out into different areas, for example, there is considerable expertise in proteomics, metabolomics and next generation sequencing. In addition, several groups based in other Units have shared appointments with the Unit.

The Unit is very well equipped for experimental and computational work. Experimental facilities include: a crystallisation robot and automated crystal visualisation; rotating anode and image plate detector for the collection of X-ray diffraction data; 800 MHz, 700 MHz, 600 MHz and 500 MHz NMR spectrometers; and several transmission electron microscopes, including a high-throughput Titan Krios microscope for single particle cryo-electron microscopy and cryo-electron tomography. The Unit also has facilities for single-molecule light microscopy, isothermal calorimetry, circular dichroism, static and dynamic light scattering and analytical ultracentrifugation, as well as for large-scale growth of prokaryotic and eukaryotic cells. The computing environment offers access to around 3000 CPU cores, whereby large central clusters and separate workstations are conveniently networked.

Peer Bork and Christoph Müller
*Joint Heads of the Structural and Computational Biology Unit*
Deciphering function and evolution of biological systems

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Previous and current research

The group currently works on three different spatial scales, but with common underlying methodological frameworks:

- genes, proteins and small molecules;
- molecular and cellular networks;
- microbial communities.

All of these biological entities we try to analyse in the context of their respective environments. We often develop tools and resources – our research is usually in new or emerging areas. Past highlights include work on the Human Genome Project (Lander et al., 2001); as well as being amongst the first to study protein interaction networks (von Mering et al., 2002), microbial communities using metagenomics (Tringe et al., 2005), and drug-target interactions using global readouts such as side effects (Campillos et al., 2007). Other exploratory projects have targeted, for instance, genome evolution, dynamic aspects of networks, post-translational modifications, and data integration for species of interest.

Our current focus is the human gut where, on one hand we employ metagenomics to uncover principles of microbial communities in the human population and in the context of disease, and on the other we use systems pharmacology to understand and predict the impact of drugs on humans at the molecular level. We develop analytic frameworks (e.g. see Qin et al., 2010; Sunagawa et al., 2013) and study stratification of the human population – we found three principle community compositions in developed countries across the world, dubbed enterotypes (Arumugam et al., 2011). We also explore variation at the nucleotide level, where each human appears to carry individual strains (Schloissnig et al., 2013). In the context of such global observations, we hope to find microbial markers for a number of diseases such as obesity (Le Chatelier et al., 2013) and colon cancer.

Furthermore, the human gut does not only harbour hundreds of microbial species, but its environment also impacts the efficacy of orally administered drugs. We try to repurpose existing drugs and to understand more about human biology using large-scale integration of various molecular and phenotypic datasets (e.g. Kuhn et al., 2013; Işkar et al., 2013).

Future projects and goals

We aim to develop community-wide population genetics to understand how microbial communities are transmitted or evolve within ourselves and under which constraints. This might help us to answer many long-standing questions, for instance how many pathogens do we tolerate in us as phenotypically healthy individuals? How is antibiotic resistance developing? For which diseases or phenotypes can we identify microbial biomarkers or can we develop a formula that will replace faecal transplantations? In the future we hope to connect microbiomics with diet, host interactions and drug intake. In this regard, we will continue to explore networks between proteins and chemicals such as lipids or carbohydrates and link them to phenotypic data such as disease status, side effects or toxicology. Several current exploratory projects study, for example, systematically the interactions between post-translational modifications and their evolution.

While the human microbiome will remain our main focus, we will also take part in collaborations exploring other habitats such as the ocean to better understand biodiversity – for example, we are involved in the TARA Oceans project (Karsenti et al., 2011). To foster translational research, the group is partially associated with the Max Delbruck Center for Molecular Medicine in Berlin and with the Molecular Medicine Partnership Unit at the University of Heidelberg.

Selected references


Multiple roles for different microbial communities in the human gut (modified from the German newspaper Zeit covering the work of the group, original designed by J. Schievink). Metagenomic data from thousands of individuals from all over the world are analysed. For example, three stratifying gut microbial community types (enterotypes) have been discovered in the human population (Arumugam et al., 2011); shown are 1000 individuals clustered by their gut microbial composition. Each individual is a dot, coloured by enterotype.
Molecular mechanisms of transcriptional regulation in eukaryotes

Selected references


Previous and current research

In the context of chromatin, we are interested how sequence-specific transcription factors assemble on DNA and how these factors interact with co-activators and general transcription factors to recruit RNA polymerases to the transcription start site. We are also studying the overall structure, architecture and inner workings of large molecular machines like RNA polymerases or chromatin modifying complexes involved in the transcription process. Finally, we would like to gain insight into how DNA sequence information and epigenetic modifications work together to regulate gene transcription.

To achieve these goals, we use structural information mainly obtained by X-ray crystallography and electron microscopy combined with other biophysical and biochemical approaches. Systems currently under investigation include multi-protein complexes involved in chromatin targeting, remodelling and histone modifications, yeast RNA polymerases I and III, and the Elongator complex.

Chromatin modifying complexes: The accessibility of chromatin in eukaryotes is regulated by ATP-dependent chromatin remodelling factors and histone modifying enzymes. Both classes of enzymes use similar domains like bromodomains, chromodomains, MBT domains, PHD fingers and SANT domains for the controlled access to defined genomic regions. We try to understand the molecular architecture of large chromatin modifying complexes, such as polycomb repressive complexes (PRCs), by which mechanisms they are recruited, how they interact with the nucleosome, and how their activities are regulated.

RNA polymerase I and III transcription: RNA polymerase I (Pol I) and III (Pol III) consist of 14 and 17 subunits, respectively. Whereas Pol I is responsible for the biosynthesis of ribosomal RNA, Pol III synthesises small RNAs like tRNA and 5S RNA. Misregulation of Pol I and Pol III has been associated with different types of cancer. Our research aims to understand the overall architecture of the Pol I and Pol III enzymes and the architecture of their pre-initiation machineries using a broad and interdisciplinary approach that combines integrated structural biology with in vitro and in vivo functional analysis.

Elongator: The 6-subunit Elongator complex was initially identified as a transcriptional regulator associated with elongating RNA polymerase II. However, recent results suggest that Elongator is involved in the specific modification of uridines at the wobble base position of tRNAs. Our group recently solved the Elp456 subcomplex that forms a ring-like heterohexameric structure resembling hexameric RecA-like ATPases. We are now pursuing the structural and functional analysis of the entire Elongator complex to gain further insight into its molecular function.

Future projects and goals

- Molecular insights into the recruitment of transcriptional regulators through the combination of DNA sequence-specific recognition and epigenetic modifications.
- Structural and functional analysis of macromolecular machines involved in transcription regulation, chromatin remodelling and chromatin modification.
- Contributing to a better mechanistic understanding of eukaryotic transcription and epigenetics using integrated structural biology combined with biochemical and cell biology approaches.
Previous and current research

Our research focuses on transposons, a class of mobile genetic elements that can autonomously move from one location to another in the genome. They drive genetic diversity and evolution and constitute about half of the human genome. However, the physiological roles of transposons are just starting to be unravelled. Recent studies show that they have key functions in gene regulation, development, immunity, and neurogenesis (Beck et al., 2011). In addition, these ‘jumping’ DNA elements offer attractive tools for genetics and human gene therapy.

To better understand transposition and facilitate the development of transposon-based genetic tools, we investigate the molecular mechanisms of their movement and regulation using structural biology (mainly X-ray crystallography), molecular biology, biochemistry, biophysics, microbiology, and cell biology approaches. We strive to understand the structure of functional transposition complexes, the chemistry they use to cut and paste DNA, their target-site selection and their regulation in the cell.

Sleeping Beauty: This resurrected transposon provides a prime tool in vertebrate genetics with applications spanning from forward mutagenesis screens to chromosomal engineering and gene therapy (Ivics et al., 2009). We study the structure and mechanisms of this transposon and, in collaboration with the Gavin (page 60) and Beck (page 57) groups, we also investigate how it interacts with other components of the human host cells.

Target site-specific transposons: One of the main obstacles in gene therapy is integration of the therapeutic gene at unwanted locations. We seek tools that integrate to specific genomic sequences. Our mechanistic work revealed that the IS608 transposon uses a short sequence in the transposon DNA to guide its integration to a specific sequence via base pairing (Barabas et al., 2008). The site of insertion can also be altered by making point mutations in the transposon (Guynet et al., 2009). We are now testing if this target recognition can be extended to select unique genomic sites.

Antibiotic resistance carrying elements: The spread of antibiotic resistance is one of today’s biggest public health concerns. Conjugative transposons provide a major mechanism to transfer resistance between bacteria. To shed light on their mechanism of transfer, we study two conjugative transposons from Helicobacter and Enterococcus, respectively.

Transposon regulation: To avoid deleterious outcomes, cells must keep their transposons under control. One major control mechanism is provided by small RNAs. In collaboration with the Carlomagno (page 59) and Pillai (page 108) groups, we investigate these processes in prokaryotes and eukaryotes. Our recent work on the eukaryotic piRNA pathway has revealed the structure and function of a novel component, a piRNA biogenesis factor called Zucchini.

Future projects and goals

- Develop novel genetic engineering tools and explore their applications in transgenesis and synthetic biology.
- Study the mechanism and regulation of a class of ‘beneficial’ transposons that are involved in the development of ciliated protists.
- Explore the movement of retrotransposons.
Structure and function of large macromolecular assemblies

Research in the Beck group combines biochemical approaches, proteomics and cryo-electron microscopy to study large macromolecular assemblies.

Selected references


Previous and current research

Integrated structure determination approaches: Research in our laboratory combines biochemical approaches, proteomics and cryo-electron microscopy to study the structure and function of large macromolecular assemblies. Cryo-electron tomography is the ideal tool to observe molecular machines at work in their native environment (figure 1). Since the attainable resolution of the tomograms is moderate, the challenge ahead is to integrate information provided by complementary approaches in order to bridge the resolution gap towards high-resolution techniques (NMR, X-ray crystallography). Mass spectrometry approaches can provide the auxiliary information that is necessary to tackle this challenge. Targeted mass spectrometry can handle complex protein mixtures and, in combination with heavy labelled reference peptides, provides quantitative information about protein stoichiometries. Using this together with cross-linking techniques can reveal protein interfaces. The spatial information obtained in this way facilitates the fitting of high-resolution structures into cryo-EM maps in order to build pseudo-atomic models of entire molecular machines (figure 2).

Large macromolecular assemblies: Megadalton protein complexes are involved in a number of fundamental cellular processes such as cell division, vesicular trafficking and nucleocytoplasmic exchange. In most cases such molecular machines consist of a multitude of different proteins that occur in several copies within an individual assembly. Their function is often fine-tuned towards context specific needs by compositional remodelling across different cell-types. Structural variations occur through stoichiometric changes, subunit switches or competing protein interfaces. Studying the structure and function of Megadalton protein complexes is a challenging task, not only due to their compositional complexity but also because of their sheer size, which makes them inaccessible to biochemical purification.

Future projects and goals

- To develop integrated workflows for structure determination of large macromolecular assemblies such as the nuclear pore complex (figure 2).
- To reveal the function of cell-type specific variations of macromolecular assemblies.

Figure 1: Cryo-electron tomogram of a fraction of the cytoplasm of a human cell. Microtubules are coloured in orange, stress fibres in grey, protein complexes in green, membranes in cyan and vesicular contents in yellow.

Figure 2: Model of the scaffold arrangement of the human Nuclear Pore Complex revealed by an integrated approach consisting of cryo-electron tomography, single particle EM, cross-linking MS and structural modelling (Bui, von Appen et al., Cell, 2013).
Viruses and vesicles – cryo-electron microscopy and tomography

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Group Leader at EMBL since 2006.
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Senior Scientist since 2013.

Previous and current research

We aim to understand the structure and molecular assembly mechanisms of important pathogenic enveloped viruses (e.g. HIV and Influenza virus), and of cellular trafficking vesicles (e.g. clathrin, COPI or COPII coated vesicles). To do this we need to understand how the protein components of the virus or vesicle interact with one another, how the cargo of the virus or vesicle is collected, and how the proteins interact with the lipid bilayer to reshape it and form a free virus or vesicle. The level of understanding we aim to achieve could be imagined as a 3D, functionally annotated movie – with molecular resolution – of the assembly and budding process. We apply methods including cryo-electron microscopy and tomography, correlated light and electron microscopy, and computational image processing. The unique power of cryo-electron microscopy and tomography methods is that they can provide detailed structural information under close-to-native conditions, even within cells.

For many problems we take a step-by-step approach. Correlative fluorescence and electron microscopy methods can be used to locate and characterise features of interest. 3D reconstructions of these features can be obtained using electron tomography of the biological system in its native state. These reconstructions can be better interpreted by comparison with data collected from in vitro reconstituted systems. A detailed view is obtained by carrying out image processing using sub-tomogram averaging or single-particle reconstruction. As well as applying existing methods, we develop and apply new approaches, in particular correlative fluorescence and electron microscopy techniques, and image processing protocols for high-resolution sub-tomogram averaging. Members of the group have varied and complementary skills, including biochemistry, cell biology, physics, engineering and computer sciences.

A particular emphasis of our research is the structure and life-cycle of asymmetric membrane viruses such as HIV. The structure and assembly of HIV particles offers insights into general features of membrane budding. Further details on our research into the structure and inhibition of HIV are available on our Molecular Medicine Partnership Unit website: www.embl.de/mmpu/mmpu/research_groups/hiv.

Future projects and goals

Our goal is to understand the interplay between protein complexes, membrane shape and virus/vesicle structure. What kind of protein-protein interactions can drive virus assembly while maintaining structural flexibility? How do structural switches allow viruses and vesicles that have completed the assembly pathway to start disassembling? How do proteins induce the distortion of cellular membranes into vesicles of different dimensions? What are the similarities and differences between the variety of cellular budding events? How do viruses hijack cellular systems for their own use? How does the curvature of a membrane influence its interaction with particular proteins? We are developing and applying novel microscopy and image processing approaches to address these questions.

Selected references


From cells to structure: using correlative light and electron microscopy, fluorescent images can be used to locate sites for tomography. Subtomogram averaging can be used to obtain higher-resolution structures of repeating features. These can be interpreted by comparison with in vitro cryo-EM or crystallography.
Functional mechanisms of complex enzymes involved in RNA metabolism and methodology development for drug design

The Carlomagno group uses NMR spectroscopy in combination with biochemical and biophysical techniques to study the structure and dynamics of biomolecular complexes.

Previous and current research

Our group focuses on studying: i) structure-activity relationships of RNP complexes involved in RNA processing; and ii) the interaction of small drugs with cellular receptors.

Our work aims at describing the features of RNA-protein recognition in RNP complexes and at characterising the structural basis for their function. Recently, we investigated the nucleolar multimeric Box C/D RNP complex responsible for the methylation of the 2'-O-position in rRNA. During the biosynthesis and processing of the pre-rRNA transcripts, post-transcriptional modifications of ribonucleotides occur in functionally important regions, such as at intersubunit interfaces, decoding and peptidyltransferase centers. Among the possible modifications, 2'-O-ribose methylation was shown to protect RNA from ribonucleolytic cleavage, stabilise single base pairs, serve as chaperone, and impact folding at high temperatures. We solved the structures of the 400 kDa enzyme in solution. A large conformational change is detected upon substrate binding, revealing an unexpected 3D organisation of the catalytic RNP (figure 1). In addition, the structure revealed an unsuspected mechanism of sequentially controlled methylation at dual sites of the rRNA, which might have important implications for ribosome biogenesis.

Conformational switches occur in macromolecular receptors at all cellular levels, dependent on the presence of small organic molecules that are able to trigger or inhibit specific cellular processes. In a second area of research, we develop both computational and experimental tools to access the structure of large receptors in complex with function regulators. We are the developers of INPHARMA, a novel approach to structure-based drug design that does not require crystallographic structures of the receptor-drug complex (figure 2). We apply our methods to study the functional mechanisms of anti-cancer drug-leads, designed as inhibitors of kinases, proteasome and membrane receptors.

Future projects and goals

My team uses a multidisciplinary approach combining nuclear magnetic resonance spectroscopy (NMR), and biochemical, biophysical and computational methods. Our philosophy is to tackle the structure of high molecular weight complexes, whose large size impedes a detailed structural description by NMR only, with an array of different complementary methodologies, such as segmental and specific labelling of both proteins and RNAs, small angle scattering (SAS), electron microscopy (EM), electron paramagnetic resonance (EPR), fluorescence resonance energy transfer (FRET), mutational analysis and biochemical experiments (e.g. cross-link). With our complementary approach it is possible to examine RNP particles in solution, in their native environment, where they preserve both their structure and dynamic properties.
Biomolecular networks

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Previous and current research

The rules that govern the behaviour of biological systems are the focus of intense research in the field of systems biology. The resulting models are expected to be predictive of different healthy and pathological conditions and might provide the general principles for the (re)engineering of biological systems. Our group has pioneered biochemical methods, coupled to quantitative mass-spectrometry, with the aim of systematically linking dynamic protein interaction networks to various phenotypes in model organisms, human cells and human pathogens. Long term, we aim to advance network biology and medicine through the integration of quantitative biochemistry, proteomics and structural biology, and define system-wide hypotheses explaining complex phenotypes and human diseases. We will contribute new strategies for the targeting of human pathologies and provide insight into fundamental principles and rules guiding biomolecular recognition.

Charting biological networks: The way biological systems organise themselves in dynamic, functional assemblies with varying levels of complexity remains largely elusive. One of our main focuses is on deciphering the molecular mechanisms of cell function or dysfunction, which relies to a large extent on tracing the multitude of physical interactions between the cell’s many components. We apply a range of biochemical and quantitative mass spectrometry approaches to organisms including yeast, a human pathogen and human somatic stem cells. The datasets guide the identification of drug targets and help us understand the mechanisms and side-effects of therapeutic compounds. Incorporation of structural models, single-particle electron microscopy, and cellular electron tomograms (collaboration with structural groups at EMBL) provide supporting details for the proteome organisation.

Development of new methods for charting new types of biological networks: While protein–protein and protein–DNA networks currently produce spectacular results, other critically important cellular components – metabolites – have rarely been studied in systematic interaction screens and remain best known for their housekeeping, metabolic functions. We currently focus on lipids and have developed new technologies with the capacity to produce systematic datasets measuring protein-lipid interactions. We designed miniaturised arrays of artificial membranes on a small footprint, coupled to microfluidic systems. We have also combined protein fractionation and lipidomics to characterise soluble protein-lipid complexes. We aim to extend the analyses to the entire proteome and lipidome and develop more generic approaches measuring all protein-metabolite interactions.

Future projects and goals

- Development of chemical biology methods based on affinity purification to monitor protein-metabolite interactions.
- Global screen aiming at the systematic charting of the interactions taking place between the proteome and the metabolome in the model organism *Saccharomyces cerevisiae* and in human.
- Development of new and existing collaborations to tackle the structural and functional aspects of biomolecular recognition.

Selected references


The group studies diverse organisms: the yeasts *Saccharomyces cerevisiae*, *Chaetomium thermophilum* (thermophilic eukaryote), the human pathogen *Mycoplasma pneumoniae*, and human somatic stem cells (MMPU group and EU-funded SyStemAge), with datasets contributing detailed cartographies of biological processes relevant to human health or disease. Another major goal is the generation of organism-wide, systematic datasets of protein-metabolite regulatory circuits, and hypotheses or models concerning the consequences of dysfunction in human diseases.
Previous and current research

Regulatory decisions during eukaryotic cell signalling are made within large dynamic protein complexes. Cell regulation is networked, redundant and, above all, cooperative. Decisions are made by in-complex molecular switching (see Van Roey et al., 2012, 2013). The deeply misleading ‘kinase cascade’ metaphor needs to be retired and the sooner, the better. Regulatory proteins make remarkable numbers of interactions, with the corollary that they also have highly modular architectures.

We and collaborators develop and deploy the Eukaryotic Linear Motif (ELM) resource for investigating functional sites in modular protein sequences. Linear motifs (LMs) are short functional sites used for the dynamic assembly and regulation of large cellular protein complexes and their characterisation is essential if we are to understand cell signalling. So-called ‘hub’ proteins that make many contacts in interaction networks are being found to have abundant LMs in large segments of IUP (intrinsically unstructured protein segments). Viral proteomes are rich in LMs that are used for hijacking cell systems required for viral production (see figure). We have now created a new resource – called switches.ELM – to capture motif-based regulatory switching mechanisms (see figure). ELM data are now being used by many bioinformatics groups to develop and benchmark LM predictors. We are now actively hunting for new LM candidates and we look to collaborate with groups undertaking validation experiments – for example, in a recent interdisciplinary collaboration we performed bioinformatics analyses of the SxIP motif that is critical for the regulation of microtubule ends.

We also undertake more general computational analyses of biological macromolecules. Where possible, we contribute to multidisciplinary projects involving structural and experimental groups at EMBL and elsewhere. Our collaborators Des Higgins (Dublin) and Julie Thompson (Strasbourg) have released Clustal Omega, a major update to the widely used multiple sequence alignment software. Rodrigo Lopez provides a Clustal Omega web server at EMBL-EBI.

Future projects and goals

We will continue to hunt for regulatory motifs and we will undertake proteome surveys when we have specific questions to answer. Protein interaction networks are anticipated to become increasingly important to our work. Molecular evolution is also one of the group’s interests, especially when it has practical applications. With our collaborators, we will look to build up protein architecture tools, especially the unique ELM resource, taking them to a new level of power and applicability. We will apply these tools to investigate modular protein function and may deploy them in proteome and protein network analysis pipelines. We are working to improve the way that bioinformatics standards represent cooperative molecular interactions (with the Hermjakob team, page 95). As part of the consortia DiGtoP, SyBoSS and SYSCILIA we are looking at interaction networks and systems in stem cells and primary cilia.
Previous and current research

Research in our laboratory combines advanced fluorescence and single molecule techniques with modern chemical biology methods to elucidate the nature of protein plasticity and disorder in biological systems and disease mechanisms.

Currently, more than 50,000 protein structures with atomic resolution are available from the protein databank. However, even if all 3D protein structures were available, our view of the molecular building blocks of cellular function would still be incomplete, as we now know that many proteins are intrinsically disordered, which means that they are unfolded in their native state. Interestingly, the estimated percentage of intrinsically disordered proteins (IDPs) grows with the complexity of the organism (eukaryotes ≈ 50%). In a modern view of systems biology, these disordered proteins are believed to be multi-functional signalling hubs and their ability to adopt multiple conformations is considered a major driving force behind their evolution and enrichment in eukaryotes.

While the importance of IDPs in biology is now well established, many common strategies for probing protein structure are incompatible with molecular disorder and the highly dynamic nature of those systems. In contrast, single molecule and super-resolution techniques, which directly probe the distribution of molecular events, can reveal important mechanisms that otherwise remain obscured. In particular, highly time resolved advanced fluorescence tools allow probing of molecular structures and dynamics at near atomic scale down to picosecond resolution. While such experiments are possible in the natural environment of the entire cell, single molecule fluorescence studies require labelling with special fluorescent dyes, which still hampers the broad application of this technique. In our group we are utilising a large spectrum of chemical biology and state-of-the-art protein engineering tools to overcome this limitation. Furthermore, microfluidics – and its potential to miniaturise lab efforts and increase throughput of single molecule science – is an area we also explore. With a focus on studying biological questions, we are continuing to develop new methods and recruit techniques from other disciplines whenever they promise to assist our overall goal of improving biological understanding.

Future projects and goals

Recent studies have shown that even the building blocks of some of the most complex and precise machines with an absolute critical role to survival of the cell are largely built from IDPs. For example, many nucleoporins are known to have central roles in the nuclear pore complex, but also in chromatin organisation, epigenetic mechanisms, transcription, and oncogenesis. How this multifunctionality can be encoded into protein disorder is a central question in biology. We aim to explore the physical and molecular rationale behind the fundamental role of IDPs by combining molecular biology and protein engineering tools with single molecule biophysics.

Selected references


Our long-term goal is to interface a large set of tools with our home-built, highly sensitive single molecule and super-resolution equipment to study structure and dynamics of heterogeneous biological systems, such as nuclear pore complexes, chromatin and transcription in 4D.
Architecture and regulation of metabolic networks

The Patil group uses a combination of modelling, bioinformatics, and experimental approaches to study metabolic networks and how they are controlled.

Selected references

Previous and current research

Metabolism is a fundamental cellular process that provides molecular building blocks and energy for growth and maintenance. In order to optimise the use of resources and to maximise fitness, cells respond to environmental or genetic perturbations through a highly coordinated regulation of metabolism. The research in our group focuses on understanding the basic principles of operation and regulation of metabolic networks. We are particularly interested in developing models connecting genotype to the metabolic phenotype (metabolic fluxes and metabolite concentrations) in cell factories and in microbial communities.

With a foundation in genome-scale metabolic modelling, optimisation methods and statistics, we develop novel computational algorithms that are driven by mechanistic insights. For example, we have previously shown that the transcriptional changes in metabolic networks are organised around key metabolites that are crucial for responding to the underlying perturbations (see figure). We complement our computational analyses with experimental activities carried out within our group (microbial physiology and genetics) and in close collaboration with other groups at EMBL and elsewhere (high-throughput phenotyping, metabolomics, proteomics, and more). This combination of computational and experimental approaches has previously enabled us to improve yeast cell factories producing vanillin – a popular flavouring agent. Currently we are developing novel tools, concepts and applications in the following research areas:

i) Metabolic interactions in microbial communities: Microbial communities are ubiquitous in nature and have a large impact on ecological processes and human health. A major focus of our current activities is the development of computational and experimental tools for mapping competitive and cooperative metabolic interactions in natural as well as in synthetic microbial communities. With the help of these tools, we aim at uncovering the role of inter-species interactions in shaping the diversity and stability of complex microbial communities.

ii) Computer-aided design of cell factories: Cell factories, such as yeast and CHO cells, are at the heart of biotechnological processes for sustainable production of various chemicals and pharmaceuticals. We are using modelling and bioinformatics tools to identify genetic redesign strategies towards improving the productivity of cell factories. These strategies guide our experimental implementation, which in turn help us to further improve the design algorithms in an iterative fashion.

Future projects and goals

We are keenly interested in expanding the scope of our computational and experimental models to gain mechanistic insight into following biological processes: i) xenobiotic metabolism in microbial communities; ii) crosstalk between metabolism and gene regulatory networks; and iii) metabolic changes during developmental processes. To this end, we are actively seeking collaborative projects within EMBL and elsewhere.
Previous and current research

The molecular hallmark of neurodegenerative diseases, such as Alzheimer’s and Parkinson’s, is the formation of large protein aggregates called amyloid fibrils. The net build-up of these fibrillar aggregates is a result of an imbalance in the cellular production and clearance of misfolded polypeptides. In our group, we are investigating the molecular structures involved in these aberrant processes as they provide fundamental insights in our understanding of ageing and neuronal dysfunction.

We are visualising the molecules by electron cryo-microscopy (cryo-EM), because large macromolecular structures and multi-protein complexes can be studied in their near-native environment without the need for crystals. Small amounts of material are sufficient to obtain ‘snapshots’ of single particles in the electron cryo-microscope and subsequent computer-aided image processing enables 3D image reconstruction. To realise the promising potential of the technique, the scientific community is still in great need of hardware-based improvements and software enhancements. Therefore, we are also interested in developing techniques, including sample preparation and data processing, to ultimately increase the resolution of single-particle cryo-EM. We would like to make it a routine tool for structural biology studies of large macromolecules.

Future projects and goals

Autophagy (from Greek, meaning ‘to eat oneself’) is the cell’s housekeeping mechanism to engulf and degrade large protein aggregates, damaged organelles and even microbes in double-membrane vesicles called autophagosomes. Multiprotein complexes are essential mediators in the events leading to autophagy. On the structural level, little is known about their 3D architecture and thus fundamental questions on the nature of these complexes need to be addressed:

- How are protein deposits structurally linked to autophagy?
- What are the shapes of these multiprotein assemblies at the membrane?
- How do they give rise to the cellular structure of the autophagosome?
Personalised genomics to study genetic basis of complex diseases

The Zaugg group investigates the variation of molecular phenotypes among individuals along with their genetic variation with the aim of better understanding the molecular basis of complex genetic diseases and inter-individual differences in drug response.

Selected references


Previous and current research

One of the continuing challenges in biomedical research, in particular in translating personalised molecular medicine to the clinic, is to understand the contribution of genetic variation to hereditary traits and diseases. Genome-wide association studies have revealed thousands of associations between genetic variants and complex diseases. However, since most of these variants lie in non-coding parts of the genome, our understanding of the molecular mechanisms underlying these associations is lagging far behind the number of known associations.

To gain a better mechanistic insight into potential causes of known genotype-disease associations our lab is investigating the variability of molecular phenotypes among individuals and trying to link them to genetic variation. In addition, since many of the disease-associated SNPs are located in regulatory elements, we have a general interest in understanding gene regulatory mechanisms.

Our recent findings suggest a genetic basis of chromatin states, challenging the traditional view of chromatin being an epigenetic mark. Interestingly, there is a dramatic discrepancy in variability among individuals between enhancer elements (most variable) and gene expression (least variable). We further found that regulatory elements that are variable among individuals are enriched for SNPs that have previously been found to associate with complex traits or diseases, highlighting the functional significance of studying inter-individual variation of molecular phenotypes. We are currently investigating potential mechanisms, such as enhancer compensation models as well as transcript isoform variation, to understand the complex relationship between gene expression and regulatory elements.

Future projects and goals

In the future we will expand our efforts to contributing to the understanding of complex traits and diseases along three lines of research:

- We will apply our models to current genome-wide association studies to increase our power of understanding known associations between genetic variants and complex diseases.
- We will expand our models to include more downstream molecular phenotypes, such as protein levels and complex composition, to estimate the impact of genetic variation on biological pathway activity.
- We will use drug response as a model to investigate the role of chromatin in mediating genotype-environment interactions across individuals.

Judith Zaugg

PhD 2011, EMBL-EBI and Cambridge University.

Postdoctoral research fellow, Stanford University.

Group leader at EMBL (from September 2014).

Joint appointment with Genome Biology.
EMBL Heidelberg

Core Facilities

The EMBL model for Core Facilities has developed a first-rate reputation in the European life sciences community. The Core Facilities contribute significantly to internal and external training courses and workshops, often in collaboration with industrial partners. Moreover, institutions in member states frequently seek our advice and guidance in setting up their own core facilities and services to enhance the efficiency and effectiveness of their scientific research.

EMBL's Core Facilities play a crucial role in enabling scientists to achieve ambitious research goals in a cost effective way. Following the establishment of a small set of facilities in 2001, the support of EMBL Council has enabled significant expansion, with the development of a number of high-level support teams that help focus diverse sets of expertise and multiple expensive technologies on specific biological problems. Currently, facilities cover the following areas: Advanced Light Microscopy, Chemical Biology, Electron Microscopy, Flow Cytometry, Genomics, Protein Expression and Purification, and Proteomics. In line with EMBL’s mission to provide services to Member States, Core Facilities are open to both internal and external scientists, who benefit significantly from our contributions and advice and are able to conduct research at and beyond normal state-of-the-art.

Core Facilities are staffed by technology experts who focus entirely on service provision, delivering technologies to be used in research projects designed and run by others. Each is run by a Head of Facility who is responsible for daily operations and ensuring high user satisfaction. Close attention is given to the delivery of quality services, fast reaction times to user demands, affordable prices and the complete integration of Core Facilities with the scientific objectives of EMBL.

Such attributes are enhanced by a user committee, which consists of representatives of EMBL’s research units. The committee helps to ensure that support activities are tailored to the demands of the research community, supports the introduction of new services, helps to define future strategies and provides valuable feedback on current operations.

Christian Boulin
Director, Core Facilities and Services
Advanced Light Microscopy Core Facility

Rainer Pepperkok
PhD 1992, University Kaiserslautern.
Postdoctoral research at University of Geneva.
Lab head at the Imperial Cancer Research Fund, London.

The facility was set up as a cooperation between EMBL and industry to improve communication between users and producers of high-end microscopy technology and to support in-house scientists and visitors in using light microscopy methods for their research. The ALMF also regularly organises in-house and international courses to teach basic and advanced light microscopy methods.

Major projects and accomplishments
- The ALMF presently manages 18 top-of-the-line microscope systems and eight high-content screening microscopes from leading industrial companies, as well as four image analysis workstations.
- More than 50 visitors per year come to carry out their own experiments in the ALMF or to evaluate microscopy equipment.
- The ALMF was the seed for the European Light Microscopy Initiative (ELMI) that establishes links with facilities throughout Europe.
- Usage of the facility exceeds 50,000 hours per year.

Services provided
- Assisting all EMBL groups and visitors through the microscopic imaging process: project planning, sample preparation, staining, microscope selection and use, image processing, presentation, data transfer and storage.
- Developing accessory software and microscopy equipment; co-developments with industrial partners; pre-evaluation of commercial equipment.
- Supporting all aspects of automated microscopy and high-content screening projects, including RNAi technology.

Technology partners
The ALMF presently has collaborations with the following companies:
- Applied Precision*
- Bitplane*
- Carl Zeiss*
- Eppendorf*
- Lambert Instruments*
- Leica Microsystems*
- Olympus*
- Eppendorf*
- Perkin Elmer*
- PicoQuant
- Scientific Volume Imaging (SVI)

Selected references
The facility assists groups in developing primary and secondary assays for screening against the in-house compound library and guide them in developing tool compounds for their specific target.

**Selected references**


Small molecules play essential roles in many areas of basic research and are often used to address important biological questions. Our aim is to enable research groups to address biological questions by identifying and developing ‘biotool’ compounds against novel targets. We can assist groups in the development of primary and secondary assays for screening against our in-house compound library and guide them through the process of developing tool compounds for their specific target. Chemical optimisation projects can be done in collaboration with our chemistry partners. The facility is a collaboration between EMBL, the German Cancer Research Center (DKFZ), and the University of Heidelberg (since February 2012) to provide the infrastructure and expertise to open up small molecule development to research groups at these institutions.

**Major projects and accomplishments**

The facility was established at the beginning of 2004. We have a very strong pipeline of projects from all three institutes covering biochemical- and cell-based targets. At the end of 2009 we established computational chemistry as part of the facility offering. Elara Pharmaceuticals GmbH and Savira Pharmaceuticals GmbH have been founded to further develop and commercialise active compounds identified in the facility, targeting specific cancer cell signalling pathways and the influenza virus respectively.

**Services provided**

Our screening library is composed of around 80 000 compounds. The selection focused on compound catalogues from three leading vendors in the field. Each vendor offers access to significantly larger collections, with low redundancy and highly competitive prices, coupled with attractive options for resupply and follow-up synthesis services. Selected compounds were checked for drug-likeness, structural and shape diversity, novelty, and compliance with medicinal chemistry requirements.

Individual compound selection was done by picking representative compounds around selected scaffolds. A scaffold-based selection offers the advantage of high information screening: as the structural space around each scaffold is covered appropriately, any hit compounds from a high throughput screen can be rapidly followed up by selecting similar compounds to enable initial structure-activity relationships to be discerned. This will help in the prioritisation of the hit compounds for further medicinal chemistry optimisation.

**Further services include:**

- Selection of appropriate assay technology platforms.
- Developing assays for medium throughput screening.
- Assisting in the design of secondary specificity assays.
- Compound characterisation.
- Managing compound acquisition through our chemistry partners.
- Computational screening using ligand-based and structure-based design strategies.

**Partners**

- Technology partners: Perkin Elmer, IDBS, Certara, GE, TTP Labtech.
- Chemistry partners: ChemDiv, Chembridge and Enamine.
The EMCF activities cover a large spectrum of EM techniques with a major focus on sample preparation, immuno-localisation of proteins, ultrastructural analysis in 2D and 3D, and data processing. Staff in the facility can help you to define optimal experimental conditions for your project – we have experience spanning virtually the full spectrum of biological specimens, with high-level resources for both research and training.

**Major projects and accomplishments**

**Advanced equipment:** Besides conventional methods for sample preparation, we offer access to a set of high-pressure freezing machines that are routinely used to vitrify biological samples. Strong expertise has been developed in yeast cells, adherent cultured cells, Drosophila embryos, nematodes, zebrafish embryos, and mouse tissues. A new microwave-assisted sample processor for chemical fixation greatly reduces time spent preparing the samples (from days to hours). Our electron tomography equipment includes a transmission electron microscope (a FEI 30 300kV microscope with a field emission gun and Eagle FEI 4K camera) and computing set-up with programs for 3D reconstruction and cellular modelling. Specialised EM engineers have expertise in tomography data acquisition and processing.

**The Electron microscopist ‘savoir faire’:** We are deeply involved in method development and training. A recent example in correlative light and electron microscopy (CLEM) is the implementation of a technique developed by the Briggs (page 58) and Kaksonen (page 22) groups, which tracks the signal of fluorescent proteins in resin sections with high precision.

**The future in perspective:** We are in the process of implementing FIB-SEM technology that will allow, among other applications, automated serial image acquisitions. The technology presents new opportunities for the understanding of the cellular fine architecture as it offers high-resolution 3D imaging (5 nm, isotropic) of volumes containing up to several cells. A Crossbeam Auriga 60 from Zeiss will be fully operational in spring 2014.

**Services provided**

**Techniques**
- Sample preparation for single particle analysis.
- Chemical fixation, high pressure freezing of cells and multi-cellular specimens.
- Resin embedding.
- Ultramicrotomy and cryo-ultramicrotomy (Tokuyasu technique).
- Immuno-labelling and TEM imaging.
- Electron tomography.
- Correlative light and electron microscopy.
- Image analysis and 3D cellular modelling.

**Teaching and training**
- Training of the users on all the accessible techniques.
- Organising courses and lectures on EM methods in cell biology.

**Technology partners**
FEI Company (our Transmission Electron Microscopes), Carl Zeiss (the Crossbeam), Leica Microsystems (ultramicrotomes, high pressure freezing and freeze substitution), AbraFluid (high pressure freezing).

Selected references


Immungold labeling is a routine technique offered by the facility. In this work, performed by Charlotte Funaya, an EGFP tagged protein targeted to the trans-Golgi network was stained with an anti-GFP antibody revealed with 10nm gold particles (Teresa Alonso, IBGM, Valladolid, Spain).
The goal of the facility is to proactively introduce flow cytometric methods into new research areas while supporting and extending current research.

Alexis Pérez González
PhD 2003, Center of Molecular Immunology & University of Havana, Cuba.
Postdoctoral research and cytometry lab manager at Gulbenkian Institute of Science, Oeiras, Portugal.
At EMBL since 2006. Facility Manager since 2012.

We offer a wide range of flow cytometric techniques. Our equipment adds flexibility in the preparation and execution of experiments, allowing different approaches to addressing scientific problems. Our facility strives to meet researchers’ needs and enable the highest possible resolution in terms of analysis and product.

Major projects and accomplishments
- Cell cloning by Darwinian selection, requiring several rounds of single cell sorting of target populations into 96-well plates, agar plates or other culture vessels. Examples of how this technique is being used include the generation of homogeneous cell lines with stably genome-integrated transfected genes and the selection of clones carrying improved fluorescent protein variants out of genetically engineered libraries.
- High-resolution analysis through photo saturation of dimly fluorescent bistable states in reworked bacterial signalling cascades.
- Evaluation of FRET probes in a novel assay for the evaluation of apoptosis.
- High throughput sorting of low frequency tissue-specific fly embryo nuclei carrying dim fluorescent signals as a preparative step in the analysis of chromatin modifications with regulatory function.
- Chromosome karyotyping and sorting for DNA sequencing and proteomics studies.

Services provided
- Complex multi-colour analysis of cell populations based on light scatter, fluorescent probes content and light intensities (including polarisation).
- Sorting of rare populations out of a heterogeneous particle mix. Cell cloning, particle enrichment and high purity bulk sorts.
- Providing EMBL scientific staff with expertise in flow cytometric techniques required in their research projects.
- Providing our researchers with advice and training in the use of flow cytometry, instrument operation and post-acquisition data analysis.
- Developing novel flow cytometric techniques to meet EMBL’s diverse scientific needs.

Technology partners
We work with equipment from Beckman Coulter, Cytopeia Inc., Becton Dickinson, Union Biometrica, Coherent Inc., and Miltienyi Biotec. We are open to testing new technological developments to best serve the needs of the scientific community.

The facility provides key services, such as sorting heterogeneous cell populations into homogeneous populations based on their fluorescence.
GeneCore is the in-house genomics service centre at EMBL equipped with state-of-the-art EMBL equipped with state-of-the-art facilities required for functional genomics analyses and operated by highly qualified staff.

Selected references


Bustin, S.A., et al. (2013). The need for transparency and good practices in the qPCR literature. Nat. Methods, 10, 1063-7

Genomics Core Facility

Vladimír Beneš
PhD 1994, Czech Academy of Sciences, Prague.
Postdoctoral research at EMBL.
Facility head since 2001.

Major projects and accomplishments

GeneCore provides its services to a broad range of users ranging from small research groups to large consortia (the International Cancer Genome Consortium, for example). The implementation of new-generation sequencing technology was a vital step to ensure EMBL remains at the forefront of European research. Since 2010, our massively parallel sequencing (MPS) suite has been upgraded and expanded significantly and now includes HiSeq2000 and cBot instruments, as well as MiSeq and Ion Torrent sequencers. Preparation of MPS libraries for various applications is supported by a robust instrumentation infrastructure (e.g., Covaris, Bioanalyzer, AAT Fragment Analyzer, Qubit, and more). To deal with increasing numbers of incoming samples, we recently reinforced our instrumentation infrastructure through acquisition of Beckman FX liquid handling robots.

GeneCore provides the following analyses in a single- or a pair-end sequencing mode, including multiplexing and mate-pair libraries:

- Genome-wide location analysis of nucleic acid-protein interactions – ChIP-Seq, CLIP-Seq.
- Transcriptome sequencing: RNA-Seq (including strand-specific libraries).
- Discovery of small non-coding RNAs: ncRNA-Seq.
- Genome-wide DNA methylation analysis: Methyl/BS-Seq.
- De novo sequencing & re-sequencing of genomic DNA.
- Targeted enrichment (sequence capture) in solution coupled with MPS.

GeneCore continues to establish new protocols enabling the processing of challenging samples such as low input or metagenomics samples. For analysis of MPS data, we work intensively with EMBL’s bioinformatics community on the development of terabases of MPS sequence data for its users.

Training is another crucial aspect of our work – GeneCore staff tutor individual researchers and also organise practical courses on corresponding subjects.

Services provided

- MPS sequencing, microarrays (homemade, commercial).
- miRNA qPCR profiling, Bioanalyzer, liquid handling robotics.
- Access to instruments and complete support: qPCR, NanoDrop, PCR cyclers.

We offer processing of samples for a range of microarray applications (mRNA, miRNA and other ncRNA expression profiling, comparative genome hybridisation) available from Affymetrix and Agilent platforms and, upon demand, spotting of customised arrays. In addition to three qPCR instruments managed by GeneCore, our qPCR capacity has been considerably enhanced by a Fluidigm Biomark HD instrument – a device capable of quantitation of transcripts on a single cell level.

Technology partners

MPS is still a very dynamic and rapidly evolving technology. We collaborate with several companies involved in developing MPS-related products, for instance testing them in our workflows. GeneCore is a member of the early-access program of Illumina, Agilent, NuGEN and Beckman Coulter.

Services provided

- MPS sequencing, microarrays (homemade, commercial).
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Protein Expression and Purification Core Facility

The facility produces and purifies proteins from E. coli, insect, mammalian cells and sera, using a variety of chromatographic methods and provides support for biophysical characterisation of purified proteins.

Selected references

Following each purification, we can perform biophysical analyses to ensure the quality of the purified sample in terms of correct folding and stability. Our facility also develops or evaluates new techniques and advanced protocols for protein production and purification and there is significant focus on developing time-saving solutions for these activities. Moreover, we are keeping stocks of a large number of expression vectors and bacterial strains for the users as well as preparing a collection of frequently used proteins for general use, which helps to considerably reduce the expenses of our users.

Major projects and accomplishments
We have evaluated new variants of our pETM-series expression vectors for E. coli that can now be used for sequence and ligation-independent cloning (SLIC). We have adapted vectors for insect and mammalian cells for the same cloning protocol. Using a single PCR product with the gene of interest, it is possible to integrate the insert into all of the vectors due to the universal overlaps that are present in the linearised vectors and the PCR product. A lethal ccdB gene in the original template vectors inhibits the growth of false positive colonies, which reduces the number of clones to test for the correct insert. With this new vector set, one can test the expression of a gene in different expression systems in parallel and avoid the redesigning of inserts for restriction-based cloning.

We have established a generic protocol for expression of fusion protein based on the small SUMO proteins and their highly specific protease SenP2. In most of our expressions, SUMO-fusion proteins showed high expression yields. In cases of insoluble product, we developed a protocol for proteolytic cleavage of the urea-denatured fusion protein, with the robust protease under conditions where other proteases show a poor performance. We can obtain pure, untagged proteins that are otherwise difficult to express or purify which include cytokines for cell culture or antigens for immunisation.

Services provided
- Expression and purification: proteins in E. coli, insect and mammalian cells.
- Scientific and technical advice to users at EMBL and external researchers.
- Preparing injection material for immunisations and purification of antibodies from serum and hybridoma supernatants.
- Providing quality analysis and biophysical characterisation of purified proteins (ITC, analytical ultracentrifugation, CD).
- Maintaining collections of expression vectors and bacterial strains.
- Producing frequently used enzymes and protein molecular weight markers for general use within EMBL.
- Developing and testing new vectors and protocols.
- Access to protocols and vector sequence information on the website.
- Caring for equipment for protein production and analysis.

Technology partners
We are open to collaborations with academic or industrial partners to evaluate new products or technological developments. Furthermore, we have initiated a network of protein facilities across Europe called P4EU (Protein Production and Purification Partnership in Europe) to improve information exchange and evaluation of new technologies.

Hüseyin Besir
PhD 2001, Max Planck Institute of Biochemistry, Munich.
Postdoctoral research at Roche Diagnostics, Penzberg, and the Max Planck Institute of Biochemistry, Munich.
Facility head at EMBL since 2006.
Infrastructure in the Proteomics Core Facility is centered around state-of-the-art mass spectrometry for MS and LC-MSMS experiments. This is complemented by chromatographic and electrophoretic systems for protein and peptide separation.

**Major projects and accomplishments**
- Molecular weight determination of intact proteins.
- Routine identification of proteins from coomassie and silver-stained gels.
- Identification of post-translational modifications.
- Nano flow liquid chromatography coupled to high resolution mass spectrometry; (LC-MSMS) for the identification of proteins in complex mixtures.
- Protein quantification by stable-isotope labelling (e.g. SILAC).

**Services provided**

**Proteomics:**
- Protein identification from gel or in solution.
- High resolution and high mass-accuracy MS, MSMS, and LC-MSMS (Thermo Orbitrap Velos Pro and Q-Exactive) for identification and quantification of proteins in complex mixtures.
- Ion trap (Bruker HCT) LC-MSMS for routine identification of proteins from coomassie and silver-stained gels.
- Triple-quad mass spectrometry (Thermo Vantage) for targeted protein analysis.
- Protein quantification by stable-isotope labeling (SILAC, TMT and dimethyl labelling).
- Identification of post-translational modifications.
- Enrichment of phosphopeptides (TIO2 and IMAC).
- Multi-dimensional peptide separation (isoelectric focusing and liquid chromatography).

**Analysis of intact proteins:**
- Molecular weight determination of intact proteins by ESI mass spectrometry.
- Determination of N- and C-termini of proteins and products of limited proteolysis.
- Verification of incorporation of non-natural amino acids.

**Technology partner**
- BIO-RAD

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**Selected references**

Yokoyama, H., et al. (2013). CHD4 is a RanGTP-dependent MAP that stabilizes microtubules and regulates bipolar spindle formation. *Curr. Biol.*, 23, 2443–51


EMBL-EBI Hinxton

European Bioinformatics Institute

EMBL-EBI is probably best known worldwide for its provision of biological information and bioinformatics services. However, about 20% of the institute is devoted to curiosity-driven research using computational approaches to unravel the secrets of life.

The development of new technologies provides a constant driver for innovative research into processing and analysing the data generated. For example, the wide uptake of next-generation sequencing by life scientists has led to unprecedented growth in sequence data. These data require novel algorithms to turn them into reliable information, and perhaps even more challenging is to use these new data to obtain novel insights into biological processes.

Research at EMBL-EBI is carried out both in groups devoted solely to research and in some of the larger service teams that have associated research activities. All researchers have computational approaches as their major focus, but most also collaborate closely with experimentalists and often generate experimental data themselves. Our research is highly collaborative within EMBL as well as with many external colleagues. We are highly interdisciplinary; our faculty comprises scientists who originally trained in biology, physics, chemistry, engineering, medicine and mathematics. We develop novel algorithms and protocols for handling data, such as checking the quality of the data; interpret data; and integrate data to generate new knowledge. We use this information to develop novel hypotheses about the basic molecular processes of life.

Although we are united in using computers, the biological questions we address and the algorithms we develop and use are very diverse. We explore biological questions spanning genome evolution, transcripational regulation and systems modelling of basic biological processes and disease. For example, different groups are investigating the molecular basis of ageing; the differentiation of stem cells; the basis for neuronal plasticity; and the early development of brain structure. Others are exploring regulation through epigenetics or RNA processing; how phenotype is related to genotype both in mice and humans; and how new enzyme reactions appear during evolution. All our discoveries are published in peer-reviewed journals but in addition, as part of these studies, our researchers often develop novel bioinformatics services, which are usually made freely available for all users so that our work helps facilitate new discoveries throughout the global scientific community.

Increasingly, much of our work is related to problems of direct medical significance, and with the emergence of personal genomes we are very conscious of the need to contribute to the translation of this new knowledge into medicine and the environment. This process is just beginning and will provide many challenges to computational biologists over the coming years.

Janet Thornton  
Director, EMBL-EBI
Previous and current research
The goal of our research is to understand more about how biology works at the molecular level, with a particular focus on proteins and their 3D structure and evolution. We explore how enzymes perform catalysis by gathering relevant data from the literature and developing novel software tools, which allow us to characterise enzyme mechanisms and navigate the catalytic and substrate space. In parallel, we investigate the evolution of these enzymes to discover how they can evolve new mechanisms and specificities. This involves integrating heterogeneous data with phylogenetic relationships within protein families, which are based on protein structure classification data derived by colleagues at University College London (UCL). The practical goal of this research is to improve the prediction of function from sequence and structure and to enable the design of new proteins or small molecules with novel functions.

We also explore sequence variation between individuals in different contexts and for different species. To understand more about the molecular basis of ageing in different organisms, we participate in a strong collaboration with experimental biologists at UCL. Our role is to analyse functional genomics data from flies, worms and mice – and by developing new software tools relate these observations to effects on life span.

Future projects and goals
Our work on understanding enzymes and their mechanisms using structural and chemical information will include a study of how enzymes, their families and pathways have evolved. We will study sequence variation in different individuals, including humans, flies and bacteria, and explore how genetic variations impact on the structure and function of a protein and sometimes cause disease. We will seek a better understanding of reaction space and its impact on pathways, and to use this new knowledge to improve chemistry queries across our databases. Using evolutionary approaches, we hope to improve our prediction of protein function from sequence and structure. We will also improve our analyses of survival curves and combine data with network analysis for flies, worms and mice in order to compare the different pathways and ultimately explore effects related to human variation and age.

Selected references
de Beer, T.A.P., et al. (2013). Amino acid changes in disease-associated variants differ radically from variants observed in the 1000 Genomes Project dataset. PLoS Comp. Biol. 9, e1003382
Analysis of protein and RNA sequence

The Bateman group endeavors to classify proteins and certain RNAs into functional families with a view to producing a ‘periodic table’ of these molecules.

Selected references
Buljan, M., Frankish, A., & Bateman, A. (2010). Quantifying the mechanisms of domain gain in animal proteins. Genome Biol., 11, R74

Previous and current research
Our work has centred around the idea that there are a finite number of families of protein and RNA genes. We wish to enumerate all of these families to gain an understanding of how complex biological processes have evolved from a relatively small number of components. We have produced a number of widely used biological database resources such as Pfam, Rfam, TreeFam and MEROPS to collect and analyse these families of molecules. Over the years we have published a large number of novel protein domains and families of particularly high interest. For example, we discovered the Paz and Piwi domains, which allowed us to identify the Dicer proteins as having an important role in RNAi several months before this was experimentally verified. More recently, we showed that the scramblase genes may act as membrane-tethered transcription factors.

Our research interests focus on how proteins and non-coding RNAs interact with each other and how these interaction networks can be rewired due to disease mutations or natural variation. We are interested in how proteins have evolved through the gain and loss of new protein domains. Recently we have been involved in using Wikipedia for collecting community annotation and other biological information for biological databases. Wikipedia provides an enormous opportunity for public engagement in science and we have been encouraging scientists in a number of ways to edit it. Current research is looking at identification of non-coding RNAs and understanding the function through computational analysis.

Future projects and goals
We will continue to develop tools and databases to understand the function and evolution of RNA and proteins. Using this data and computational analyses we aim to investigate interaction networks in two directions. Firstly, we will investigate the plasticity of the protein interaction network between individuals. To do this we will identify natural human variation such as SNPs and CNVs that rewire the protein interaction network.

We will also explore the large and growing set of important molecular interactions involving RNA that are currently dispersed among diverse databases and experimental studies. By bringing this data together we wish to uncover the extent and evolution of the RNA interaction network compared to the protein interaction network. In another strand of our research we will develop automated techniques to identify spurious protein predictions that are polluting sequence databases. We have collected thousands of examples of proteins that are unlikely to be translated. These examples will form a good training set for machine learning techniques to identify further suspicious proteins.

The flow of information, from sequence to knowledge of function through classification and hypothesis generation, used in the Bateman group.

Alex Bateman
PhD 1997, University of Cambridge.
Postdoctoral work at the Sanger Centre.
Group leader at Wellcome Trust Sanger Institute 2001-2012.
Head of Protein Sequence Resources at EMBL-EBI since 2012.
Evolution of cellular networks

Pedro Beltrão
PhD 2007, University of Aveiro (research conducted at EMBL Heidelberg).
Postdoctoral research at the University of California San Francisco, USA.
Group leader at EMBL-EBI since 2013.

Previous and current research

Our group is interested in understanding how novel cellular functions arise and diverge during evolution. We study the molecular sources of phenotypic novelties, exploring how genetic variability that is introduced at the DNA level is propagated through protein structures and interaction networks to give rise to phenotypic variability. Within the broad scope of this evolutionary problem, we focus on the function and evolution of post-translational regulatory networks and the evolution of genetic and chemical-genetic interactions. Looking beyond evolutionary processes, we also seek to understand the genomic differences between individuals and improve our capacity to devise therapeutic strategies.

In collaboration with mass-spectrometry groups, we develop a resource of experimentally derived, post-translational modifications (PTMs) for different species to study the evolutionary dynamics and functional importance of post-translational regulatory networks. We use these data to create novel computational methods to predict PTM function and regulatory interactions. Our goal is to gain insights into the relationship between genetic variation and changes in PTM interactions and function.

Changes in cellular interaction networks underpin variation in cellular responses and sensitivity to environmental perturbations or small molecules. As we model and study the evolution of cellular interaction networks, we begin to see how different individuals or species diverge in their response to drugs. Understanding this relationship will enable us to develop methods to predict how genetic changes result in specific sensitivity to drug combinations.

Future projects and goals

In 2014 group will continue to study the evolution of cellular interactions networks with a specific focus on post-translational regulatory networks, genetic and chemical-genetic networks. We will also continue to study the evolution of function, specificity and conditional regulation of PTM signalling as well as the evolution and conditional regulation of genetic interactions networks. In addition, we are developing methods to study how the genetic variability observed in different individuals of the same species might impact on the response to environmental conditions or drugs.

Selected references


Evolution of cellular networks

The Beltrão group studies the molecular impacts of genetic variability on phenotypic variability in order to understand better the function and evolution of cellular networks.

Functional role of post-translational modifications. PTMs act to change the activity of proteins through different mechanisms and in response to different conditions. (A) Different mechanisms used by PTMs to regulate protein activity. (B) Example of conditional regulation of phosphorylation sites. (C) Mechanisms of cross-regulation between different PTM types. (Beltrao et al., 2013).
Pluripotency, reprogramming and differentiation

The Bertone group investigates the cellular and molecular processes underlying mammalian stem cell differentiation and induced pluripotency.

Previous and current research

We investigate the cellular and molecular attributes of embryonic and tissue-specific stem cells using a combination of experimental and computational methods. We develop and apply genomic technologies to the analysis of stem cell function to address fundamental aspects of development and disease. Embryonic stem (ES) cells are similar to the transient population of self-renewing cells within the inner cell mass of the pre-implantation blastocyst (epiblast), which are capable of pluripotential differentiation to all specialised cell types comprising the adult organism. These cells undergo continuous self-renewal to produce identical daughter cells, or can develop into specialised progenitors and terminally differentiated cells. Each regenerative or differentiative cell division involves a decision whereby an individual stem cell remains in self-renewal or commits to a particular lineage. The properties of proliferation, differentiation and lineage specialisation are fundamental to cellular diversification and growth patterning during organismal development, as well as the initiation of cellular repair processes throughout life.

The fundamental processes that regulate cell differentiation are not well understood and are likely to be misregulated in cancer. A second focus in the lab is the study of neural cancer stem cells derived from human glioblastoma multiforme tumours. Using neural stem (NS) cell derivation protocols, it is possible to expand tumour-initiating, glioblastoma-derived neural stem (GNS) cells continuously in vitro. Although the normal and disease-related counterparts are highly similar in morphology and lineage marker expression, GNS cells harbour genetic mutations typical of gliomas and give rise to authentic tumours following orthotopic xenotransplantation. We apply genomic technologies to determine transcriptional changes and chromosomal architecture of patient-derived GNS cell lines and their individual genetic variants. These data provide a unified framework for the genomic analysis of stem cell populations that drive cancer progression, and contribute to the molecular understanding of tumourigenesis.

Future projects and goals

We have in place the most robust and stable systems for stem cell derivation and propagation, where controlled experiments can be performed in well-defined conditions. These assets are particularly valuable for studying cell populations that would normally be inaccessible in the developing embryo. To realise the potential of ES cells in species other than mouse, precise knowledge is needed of the biological state of these cells, and particularly of the molecular processes that maintain pluripotency and direct differentiation. We are working to translate the knowledge and methods that have been successful in mouse ES cell biology to other mammalian species. This involves the characterisation of germline-competent ES cells from the laboratory rat, along with the production of pluripotent human iPS cells using alternative reprogramming strategies. Through deep transcriptome sequencing we have shown a broad equivalence in self-renewal capacity and cellular state, albeit with intriguing species-specific differences. Thus, while ground-state pluripotency can be captured and maintained in several species, the mechanisms used to repress lineage differentiation may be fundamentally different.

Selected references


Sequence algorithms and intra-species variation

Ewan Birney
PhD 2000, Wellcome Trust Sanger Institute.
At EMBL-EBI since 2000. Joint Associate Director since 2012.

Previous and current research
Our group has a long-standing interest in developing sequencing algorithms. Over the past four years a considerable focus has been on compression, with theoretical and now practical implementations of compression techniques. Our ‘blue skies’ research includes collaborating with the Goldman group (page 86) on a method to store digital data in DNA molecules. We continue to be involved in this area as new opportunities arise – including the application of new sequencing technologies.

We are also interested in the interplay of natural DNA sequence variation with cellular assays and basic biology. During the past five years there has been a tremendous increase in the use of genome-wide association to study human diseases. However, this approach is very general and need not be restricted to the human disease arena. Association analysis can be applied to nearly any measurable phenotype in a cellular or organismal system where an accessible, outbred population is available. We are pursuing association analysis for a number of both molecular (e.g. RNA expression levels and chromatin levels) and basic biology traits in a number of species where favourable populations are available including human, and Drosophila. We hope to expand this to a variety of other basic biological phenotypes in other species, including establishing the first vertebrate near-isogenic wild panel in Japanese Rice Paddy fish (Medaka, Oryzias latipes).

Future projects and goals
In 2014 we will continue to work on sequence algorithms and intra-species variation. Our work with human data will focus on molecular phenotypes in an induced pluripotent stem cell (iPSC) panel generated as part of the HipSci consortium, and on a project based on normal human cardiac data. Our work in Drosophila will investigate multi-time-point developmental biology measures. We will also assess the near isogenic panel in Japanese Rice Paddy fish for a number of molecular and whole body phenotypes.

Example of differential phenotypes of inbred Medaka fish. (A) Four lateral (L1-L4) and five dorsal (D1-D5) morphometric distances were extracted and analysed for each inbred strain. (B) The proportion of variance explained by the difference between strains as a fraction of the total variance. The variables are the measurements corrected by the appropriate body length measurement (L4 and D5, respectively). (C) Substantial differences in lateral length and eye diameter between fish from the two Southern inbred strains HdrR and Icab. (D) The distribution of eye diameter, lateral length and the ratio between eye diameter and lateral length for HdrR and Icab. Eye diameter and lateral length: y-axis = value in pixel. HdrR (n=70), Icab (n=78).

Selected references
Functional genomics research

Research in the Brazma team focuses on analysis of gene, transcript and protein expression, on cancer genomics and proteomics, and on integrative analysis of functional genomics data.

Selected references

Gonzàlez-Porta, M., et al. (2013). Transcriptome analysis of human tissues and cell lines reveals one dominant transcript per gene. Genome Biol., 14, R70

Previous and current research

Our research group complements the Functional Genomics service team, and focuses on developing new methods and algorithms and integrating new types of data across multiple platforms. We are particularly interested in cancer genomics and transcript isoform usage. We collaborate closely with the Marioni group (page 87) and others throughout EMBL.

As a part of our participation in the GEUVADIS project (funded by the European Commission’s Seventh Framework Programme), we analysed mRNA and small RNA from lymphoblastoid cell lines of 465 individuals who participated in the 1000 Genomes Project. Our group led the analysis of transcript isoform use and fusion gene discovery. By integrating RNA and DNA sequencing data, we were able to link gene expression and genetic variation, and to characterise mRNA and miRNA variation in several human populations. All of the data generated in the project are available through ArrayExpress.

The human transcriptome contains in excess of 100 000 different transcripts. We analysed transcript composition in 16 human tissues and five cell lines to show that, in a given condition, most protein coding genes have one major transcript expressed at significantly higher level than others, and that in human tissues the major transcripts contribute almost 85% to the total mRNA. We also found that often the same major transcript is expressed in many tissues. These observations can help prioritise candidate targets in proteomics research and to predict the functional impact of the detected changes in variation studies.

Future projects and goals

Large-scale data integration and systems biology will remain the focus of our research. We will work to develop methods for RNA-seq data analysis and processing, and apply these to address important biological questions such as the role of alternative splicing and splicing mechanisms. With our collaborators from the International Cancer Genome Consortium we will be seeking new insights into cancer genomes and their impacts on functional changes in cancer development, as well as discovery and analysis of fusion genes and their role in cancer development.

Alvis Brazma

MSc in Mathematics 1982, University of Latvia.
PhD in Computer Science 1987, Moscow State University.
At EMBL-EBI since 1997.

A combined human and mouse gene expression data matrix (principal components 1 and 3). Each dot represents a sample, which is labelled by (a) species and (b) tissue type.
Functional genomics and analysis of small RNA function

Anton Enright
PhD 2003, University of Cambridge.
Postdoctoral research at Memorial Sloan-Kettering Cancer Center, New York.
At EMBL-EBI since 2008.

Previous and current research
Complete genome sequencing projects are generating enormous amounts of data. Although progress has been rapid, a significant proportion of genes in any given genome are either un-annotated or possess a poorly characterised function. Our group aims to predict and describe the functions of genes, proteins and regulatory RNAs as well as their interactions in living organisms. Regulatory RNAs have recently entered the limelight, as the roles of a number of novel classes of non-coding RNAs have been uncovered. Our work involves the development of algorithms, protocols and datasets for functional genomics. We focus on determining the functions of regulatory RNAs including microRNAs, piwiRNAs and long non-coding RNAs. We collaborate extensively with experimental laboratories on commissioning experiments and analysing experimental data. Some laboratory members take advantage of these close collaborations to gain hands-on experience in the wet lab.

Future projects and goals
Our long-term goal is to combine regulatory RNA target prediction, secondary effects and upstream regulation into complex regulatory networks. In 2014 we will continue to build an accurate database of piRNA loci in animals and explore the importance and evolution of these molecules. We are extremely interested in the evolution of regulatory RNAs and developing phylogenetic techniques appropriate for short non-coding RNA. We are equally interested in the analysis of IncRNAs and how they fit into the non-coding RNA landscape. We will continue to build strong links with experimental laboratories that work on miRNAs in different systems, as this will allow us to build better datasets with which to train and validate our computational approaches. The use of visualisation techniques to assist with the interpretation and display of complex, multi-dimensional data will continue to be an important parallel aspect of our work.

Evolutionary distribution of miRNA families.
Phylogenetic tree representing miRNA family gains and losses. Branch width represents the number of miRNA families present among leaves of the branch, while the colour represents significant miRNA family loss (blue) or gain (red). For each of 408 miRNA families present at multiple loci at least two species, we also build a graphical ‘glyph’. This glyph can be used to quickly assess presence, absence or expansion of families between clades. Each square represents a specific miRNA family. Squares are coloured as follows: white, indicates that this species does not contain a particular family, black indicates that this species contains at least 10 copies of miRNAs within that family. Copies between 1 and 10 are indicated as a rainbow gradient (red through violet). Groups of species are labelled according to the name of the evolutionary branch preceding them.

Selected references

The Enright group focuses on small non-coding RNAs and develops computational tools, systems, and algorithms to predict the function and interactions of small RNAs.
Evolution of transcriptional regulation

The Flicek team develops large-scale bioinformatics infrastructure, explores the evolution of transcriptional regulation and develops algorithms to study epigenomic data.

Selected references


Evolutionarily conserved transcription factor binding around the Factor IX gene promoter in five species (human, macaque, mouse, rat and dog). Mutations known to cause haemophilia are indicated by triangles about the genome sequence and are consistently found at sites of conserved transcription factor binding.

Paul Flicek

DSc 2004, Washington University.
Honorary Faculty Member, Wellcome Trust Sanger Institute since 2008.
At EMBL-EBI since 2005.
Team leader since 2008.
Senior Scientist since 2011.

Previous and current research

Our team creates and maintains the genomic resources of the Ensembl project and is responsible for data management for a number of large-scale international projects, including the 1000 Genomes Project and, in collaboration with the Brazma team, the International Mouse Phenotyping Consortium. We also collaborate on the development of EMBL-EBI’s major variation databases, including the European Genome-phenome Archive (EGA) and the DGVs database of copy number and structural variation. All of these resources are publicly available and are widely used by the scientific community and by the team itself as part of our research into evolution, epigenetics and transcriptional regulation.

Our specific research projects focus on the evolution of transcriptional regulation. Recently we have expanded ‘comparative regulatory genomics’ techniques including mapping the same DNA–protein interactions in matched tissues in multiple species to understand how gene regulation has evolved while the tissue-level functions are largely conserved. We are also interested in the role of chromatin conformation in tissue-specific gene regulation and have investigated both the CTCF and cohesin complex in this context.

Future projects and goals

With the issue of major datasets from the EU-funded Blueprint project and the NIH-funded KOMP2 project, we continue to play an end-to-end role in major genomics projects from raw-data management for the project to summary-data presentation to the wider scientific community. Our research projects are expanding in number of species, tissues and specific DNA–protein interactions. We will also focus on understanding the differentiation process and components of cell- and tissue-specific regulation. We will address these questions both in the context of our established collaborative projects with the Odom group at the University of Cambridge and as part of other collaborations, including larger EU-funded projects.
Digital information encoding in DNA. Digital information (a, in blue), here binary digits holding the ASCII codes for part of Shakespeare’s sonnet 18, was converted to base-3 (b, red) using a Huffman code that replaces each byte with five or six base-3 digits (trits). This in turn was converted in silico to our DNA code (c, green) by replacement of each trit with one of the three nucleotides different from the previous one used, ensuring no homopolymers were generated. This formed the basis for a large number of overlapping segments of length 100 bases with overlap of 75 bases, creating fourfold redundancy (d, green and, with alternate segments reverse complemented for added data security, violet). Indexing DNA codes were added (yellow), also encoded as non-repeating DNA nucleotides (figure courtesy of Nature).
Classifying genes by their regulatory function. We used RNA-seq data generated from F0 mice and their F1 hybrids to classify genes into sets depending upon their regulatory mechanism (Goncalves et al., 2012).

### Future projects and goals

In 2014 and beyond, our lab will continue to focus on developing computational tools for understanding the regulation of gene expression levels. An especially strong focus will be on the development of methods for analysing single-cell RNA-sequencing data (in conjunction with other members of the Sanger-EBI Single-Cell Genomics Centre), which has the potential to reveal novel insights into cell type identity and tumourigenesis.
Previous and current research

Although great progress has been made in developing biological drugs, synthetic small molecule and natural product-derived drugs still form the majority of novel, life-saving medicines. The process complexity and costs of discovering new drugs is so high that conventional industrial R&D efforts are being complemented with public-private partnerships. Central to this public-sector activity is data sharing and availability of structure, binding, functional and ADMET data for large sets of compounds. The ChEMBL database, which our group maintains, stores curated, two-dimensional chemical structures and abstracted quantitative bioactivity data alongside calculated molecular properties.

Our research interests centre on mining ChEMBL for data that can be applied to drug-discovery challenges, and finding new ways to increase the database in a qualitative and quantitative manner. ChEMBL data is mainly derived by manual abstraction and curation from the primary scientific literature, and cover a significant fraction of the structure-activity relationship (SAR) data for the discovery of modern drugs. We complement this data with depositions, and selected data from other public resources, such as PubChem.

Our work on differential expression during development focuses on identifying molecular differences between humans of different ages in order to better understand age-related drug differential responses reported in the literature, and to identify how these changes translate to species commonly used in drug development. We also work on chemogenomic characterisation of allosteric modulators, using resources in ChEMBL to develop predictive models that can quantitatively predict the likelihood of compounds modulating a certain target. We are also developing the Functional Therapeutic Chemical Classification System, a resource formalising the mode and mechanism of action of drugs using semantic technologies. This classification has been designed to support drug re-purposing and can be seen as a toolbox designed to fix dysfunctional biological systems.

Future projects and goals

We will extend our work on translational and safety biology to areas including the study of the impact of human genetic and physiological variation of drug efficacy and safety.

ChEMBL: a database of bioactive drug-like small molecules
Julio Saez-Rodriguez
PhD 2007, University of Magdeburg.
Postdoctoral work at Harvard Medical School and MIT.
At EMBL-EBI since 2010.
Joint appointment, Genome Biology Unit.

Previous and current research

Human cells are equipped with complex signalling networks that allow them to receive and process the information encoded in myriad extracellular stimuli. Understanding how these networks function is a compelling scientific challenge and has practical applications, as alteration in the functioning of cellular networks underlies the development of diseases such as cancer and diabetes. Considerable effort has been devoted to identifying proteins that can be targeted to reverse this deregulation; however, their benefit is often unexpected. It is hard to assess their influence on the signalling network as a whole and thus their net effect on the behaviour of the diseased cell. Such a global understanding can only be achieved by a combination of experimental and computational analysis.

Our research is hypothesis-driven and tailored towards producing mathematical models that integrate diverse data sources. To this end, we collaborate closely with experimental groups. Our models integrate a range of data, from genomic to biochemical, with various sources of prior knowledge, with an emphasis on providing both predictive power of new experiments and insights into the functioning of the signalling network. We combine statistical methods with models describing the mechanisms of signal transduction, either as logical or physico-chemical systems. For this, we develop tools and integrate them with existing resources. We then use these models to better understand how signalling is altered in human disease and predict effective therapeutic targets.

Future projects and goals

In 2014, we will continue to develop methods and tools to understand signal transduction in human cells and its potential to yield insights of medical relevance. Our main focus will be on modelling signalling networks using phospho-proteomics data with our software tool CellNOpt, and finding ways to employ different proteomics technologies and sources of information about pathways. We will also develop methods to infer drug mode of action and drug repurposing by integrating genomic and transcriptomic data with drug screening data. Using these methods we will address questions such as: What are the origins of the profound differences in signal transduction between healthy and diseased cells and, in the context of cancer, between normal and transformed cells? What are the differences in signal transduction among cancer types? Can we use these differences to predict disease progression? Do these differences reveal valuable targets for drug development? Can we study the side effects of drugs using these models?
Statistical genomics and systems genetics

Oliver Stegle

PhD 2009, University of Cambridge.
Postdoctoral fellow, Max Planck Institutes Tübingen 2009-2012.
Research group leader at EMBL-EBI since 2012.

Previous and current research

Our interest lies in computational approaches to unravel the genotype–phenotype map on a genome-wide scale. How do genetic background and environment jointly shape phenotypic traits or causes diseases? How are genetic and external factors integrated at different molecular layers such as transcription and translation?

To make accurate inferences from high-dimensional ‘omics’ datasets, it is essential to account for biological and technical noise and to propagate evidence strength between different steps in the analysis. With this in mind, we develop statistical analysis methods in the areas of gene regulation, genome-wide association studies (GWAS) and causal discovery in molecular systems. Our methodological work is tied in with experimental collaborations, in which we study the variability of molecular phenotypes in different systems including yeast models, plants genomics and human genetics. Success include: the development and application of new statistical tools to integrate molecular phenotype data and global phenotypes in systems-level models; joining the Human Induced Pluripotent Stem Cell Initiative (HipSci); running a new EMBO training course on genotype-phenotype mapping.

In 2013 we continued to develop and apply methods for linking genetic variation data and molecular phenotypes. We contributed to major studies mapping the genetic component of gene expression variation in humans, including contributions to an initial publication of the GEUVA-DS consortium (Lappalainen et al., 2013).

To fully explain differences in gene expression between individuals, it is necessary to consider genetic effects in combination with other sources of variation. Together with colleagues from Sheffield (Fusi et al., 2013) we devised new statistical approaches to study genetic effects on gene expression in the context of different environmental exposures. The genotype–environment interactions uncovered by our new approach shed light on the interplay of genetic signals and external stimuli.

In addition to studying genetic effects on molecular phenotypes in isolation, we developed and applied statistical tools for tying together multiple layers of molecular readouts with global phenotypes. In a collaborative effort with the Steinmetz group (page 43) and the Gagneur group at Ludwig Maximilian University Munich, we demonstrated how these methods allow for teasing apart cause–consequence relationships when studying gene expression levels in the context of genotype-phenotype relationships of fitness traits (Gagneur et al., 2013). This work laid the groundwork for other projects in human genetics in which we exploit rich molecular phenotype data to dissect the GWAS loci.

The group also conducted research in the field of machine learning – collaborating with colleagues at MPI Tübingen, we developed new models that allow for jointly analysing statistical associations of multiple response variables in a single model (Rakitsch et al., 2013). These theoretical developments form the basis of integrative genetic approaches to study high-dimensional phenotypes.

Future projects and goals

In 2014 we will continue to devise statistical methods to model and analyse data from high-throughput genetic and molecular profiling experiments. Technically, the development of new approaches for tying together quantitative readouts across multiple molecular layers will increase in importance. To this end, we develop causal inference methods to deduce functional relationships from the wealth of correlative ‘omics datasets being generated. We are particularly interested in applying these methods to data from HipSci.

Selected references


Small molecule metabolism in biological systems

The Steinbeck group studies small molecule metabolism in biological systems, including reconstruction of metabolic networks from genomic data, and structure elucidation and identification of metabolites.

Selected references

Previous and current research
Our research is dedicated to the elucidation of metabolomes, Computer-Assisted Structure Elucidation (CASE), the reconstruction of metabolic networks, biomedical and biochemical ontologies, and algorithm development in chem- and bioinformatics.

Part of our research comprises the development and implementation of methods to analyse spectroscopic data in metabolomics. The chemical diversity of the metabolome and a lack of accepted reporting standards currently make analysis challenging and time consuming. Typical mass spectrometry-based studies, for instance, generate complex data where the signals of interest are obscured by systematic and random noise. Proper data pre-processing and consequent peak detection and extraction is essential for compound identification.

CASE methods developed by us provide a means to determine the structure of metabolites by stochastic screening of large candidate spaces based on spectroscopic methods. Our SENECA system is based on a stochastic structure generator, which is guided by a spectroscopy-based scoring function. Simulated Annealing and Evolutionary Algorithms are at the core of the structure generation process, allowing us to explore the structural space of isomers.

Another of our activities involves employing machine-learning methods to correlate graph-based molecular descriptors with database knowledge. We use the resulting prediction engines as judges in our SENECA scoring function. In order to narrow down our search space during the structure-determination process, we employ Natural Product likeness as a filter.

Future projects and goals
Our focus is efficient methods and algorithms for the assembly, analysis and dissemination of information on small molecules of relevance for biological systems. This includes information about primary and secondary metabolites, and also on xenobiotics and other molecules of relevance, such as epitopes. To this end, we will continue our work in various related areas of ontology development, research on the computational representation of related data, inference of metabolomes from all types of available information, processing of metabolic and metabolomics information, and reconstruction of metabolic networks. We select these projects with an emphasis on applicability in our service foci. Here, our focus is on the extension of the ChEBI database towards greater usability for metabolism and natural products research, improving the curation efficiency (which requires work on curation tools and narrow AI in the area of on-the-fly information retrieval) and the extension and establishment of our metabolomics database, MetaboLights.

Christoph Steinbeck
PhD 1995, Rheinische Friedrich-Wilhelm-Universität, Bonn.
Postdoctoral research at Tufts University, Boston, 1996-1997.
Lecturer in Cheminformatics, University of Tübingen, 2007.
At EMBL-EBI since 2008.

PCA plot from metabolomics LC-MS data for four genotypes of tomato (Solanum lycopersicum). Metabolomics snapshots were studied at different stages of ripening (days indicated by data point size). The plot reveals distinctly different ripening trajectories for each genotype.
Gene expression regulation and protein complex assembly

Sarah Teichmann

PhD 2000, University of Cambridge and MRC Laboratory of Molecular Biology,
MRC Career Track Programme Leader, MRC Laboratory of Molecular Biology, 2001-5 and MRC Programme Leader, 2006-12.
Fellow and Director of Studies, Trinity College, since 2005.
Group leader at EMBL-EBI and Sanger Institute since 2013.

Previous and current research

Our group seeks to elucidate general principles of gene expression and protein complex assembly. We study protein complexes in terms of their 3D structure, structural evolution and the principles underlying protein-complex formation and organisation. We also explore the regulation of gene expression during switches in cell state, and use mouse T-helper cells as a model of cell differentiation. We combine computational and wet-lab approaches at both EMBL-EBI and the Wellcome Trust Sanger Institute.

The wealth of genome-scale data now available for sequences, structures and interactions provides an unprecedented opportunity to investigate systematically principles of gene and protein interactions. We focus on the evolution and dynamics of regulatory and physical interaction networks, combining computational and mathematical approaches with genome-wide and gene/protein experiments. Our two main areas are transcription factors and the regulation of gene expression; and physical protein–protein interactions and protein complexes.

Differences in genes and their spatio-temporal expression patterns determine the physiology of an organism: its development, differentiation and behaviour. Transcription factors regulate this process by decoding DNA elements and binding to DNA in a sequence-specific manner. Our group has developed a prediction pipeline (transcriptionfactor.org) that identifies repertoires of transcription factors in genomes.

We are very interested in elucidating transcriptional regulatory networks that orchestrate T-helper-cell differentiation and plasticity. Using the T helper cell system, we explore the hierarchy and kinetics of molecular events that contribute to changes in gene expression, and whether the kinetics of these interactions graded or switch-like.

Our group also investigates the principles that govern the folding and assembly of protein complexes. Using the informative power of genomic, proteomic and structural data, we capture the critical changes in sequence and structure that distinguish protein-complex formation from the sea of functionally neutral changes. The 3DComplex.org database is a research tool for our work in this area. Our in silico, phylogeny-based methods predict critical ancestral mutations involved in changing protein complexes, and we test these using wet-lab biophysical and biochemical techniques.

Future projects and goals

We will continue our projects in structural bioinformatics of protein complex assembly and expand our programme in genomics of gene expression. A major thrust will be single cell transcriptomics of the dynamics of immune responses to pathogens. This will reveal the full spectrum of CD4+ T cell types, and the evolution of the T cell response during an infection time course. These in vivo experiments, together with in vitro T-cell and ES cell experiments, will inform us about the cellular circuitry and decision-making in switching from one cell type to another. To gain more insight into cellular switches, we will work towards integrating high-throughput high-content imaging with single-cell RNA sequencing.

Methods development in single cell bioinformatics approaches is a further strand of research that we will pursue energetically over the next few years. This is an exciting field still in its infancy, and there are many open questions that require new statistical and computational techniques. Together with the Marioni (page 87) and Stegle (page 90) groups at EMBL-EBI, we are keen to find new ways to dissect technical from biological cell-to-cell variation in gene expression, predict regulatory relationships, gene expression modules and cell states from the new flood of single cell RNA-sequencing data.

Selected references

Bioinformatics Services

Service teams at EMBL-EBI focus on gathering, adding value to and presenting important collections of biological and chemical information for the benefit of the larger research community. Their work is enhanced by the input of basic researchers, some of whom are embedded in service teams. Master’s students and other visiting scientists who conduct their research at EMBL-EBI have the opportunity to work in a unique environment, exploring how we can use an incredibly diverse range of information to understand life on a fundamental level.

Building on more than 20 years’ experience in bioinformatics, EMBL-EBI maintains the world’s most comprehensive range of molecular databases. We are the European node for globally coordinated efforts to collect and disseminate biological data. Many of our databases are household names to biologists – they include EMBL-Bank (DNA and RNA sequences), Ensembl (genomes), ArrayExpress (microarray-based gene-expression data), UniProt (protein sequences), InterPro (protein families, domains and motifs) and PDB (macromolecular structures). Others, such as IntAct (protein–protein interactions), Reactome (pathways), ChEBI and ChEMBL (small molecules), help researchers understand not only the molecular parts that go towards constructing an organism, but how these parts combine to create systems. The details of each database vary, but they all uphold the same principles of service provision: accessibility, compatibility, comprehensive datasets, portability, and quality.

European Nucleotide Archive

Guy Cochrane
PhD 1999, University of East Anglia.
At EMBL-EBI since 2002.
Team leader since 2009.

The European Nucleotide Archive (ENA) provides globally comprehensive primary data repositories for nucleotide sequencing information. ENA content spans the spectrum of data from raw sequence reads through assembly and alignment information to functional annotation of assembled sequences and genomes. Services for data providers include interactive and programmatic submission tools and curation support. Data consumers are offered a palette of services including sequence similarity search, text search, browsing, rich integration with data resources beyond ENA, provided both over the web and through an increasingly sophisticated programmatic interface. These services are focused towards users who approach ENA data and services directly, and those who provide secondary services, such as UniProt, Ensembl, Ensembl Genomes and ArrayExpress, that build on ENA content. Reflecting the centrality of nucleotide sequencing in the life sciences and the emerging importance of the technologies in applied areas such as healthcare, environmental and food sciences, ENA data and services form a core foundation upon which scientific understanding of biological systems has been assembled and our exploitation of these systems will develop. With an ongoing focus on data presentation, integration within ENA and with resources beyond it, tools and services development, the team’s commitment is to the utility of ENA content and the broadest reach of sequencing applications.
Proteomics services

Henning Hermjakob
Dipl. Inf (MSc) in Bioinformatics 1996, University of Bielefeld.
Research assistant at the National Centre for Biotechnology (GBF), Braunschweig, Transfac Database team.
At EMBL-EBI since 1997.

Our team develops tools and resources for the representation, deposition, distribution and analysis of proteomics and systems biology data. We follow an open-source, open-data approach: all of the resources we develop are freely available. The team is a major contributor to community standards, in particular the Proteomics Standards Initiative (PSI) of the international Human Proteome Organisation (HUPO), and systems biology standards (COMBINE Network). We provide public databases as reference implementations for community standards: the PRIDE proteomics identifications database, the IntAct molecular interaction database, the Reactome pathway database and BioModels Database, a repository of computational models of biological systems. As a result of long-term engagement with the community, journal editors and funding organisations, data deposition in our standards-compliant data resources is becoming a strongly recommended part of the publishing process. Our curation teams ensure consistency and appropriate annotation of all data to provide the community with high-quality reference datasets. We also contribute to the development of data integration technologies, using protocols like the PSI Common Query Interface (PSICQUIC) and Semantic Web technologies, and provide stable identifiers for life science entities through Identifiers.org.

Non-vertebrate genomics

Paul Kersey
PhD 1995, University of Edinburgh.
At EMBL since 1999.
Team leader since 2008.

The Ensembl Genomes team provides services based on the genomes of non-vertebrate species. The falling costs of DNA sequencing have led to an explosion of reference genome sequences and genome-wide measurements and interpretation. Ensembl Genomes (Kersey et al., 2010) provides portals for bacteria, protists, fungi, plants and invertebrate metazoa, offering access to these data through a set of programmatic and interactive interfaces, exploiting developments originating in the vertebrate-focused Ensembl project. Collectively, the two projects span the taxonomic space.

The development of next-generation sequencing technologies has led to the performance of complex and highly data-generative experiments, now performed even in species studied only by small communities with little informatics infrastructure. Through collaborating with EMBL-EBI and re-using our established toolset, such small communities can store, analyse and disseminate data more cheaply and powerfully than if they develop their own tools. Our leading collaborators include VectorBase (Megy et al., 2011), a resource focused on the annotation of invertebrate vectors; WormBase (Yook et al., 2011), a resource for nematode biology; and PomBase (Wood et al., 2011), focused on the fission yeast Schizosaccharomyces pombe. In the plant domain, we collaborate closely with Gramene in the US and with a range of European groups in the transPLANT project. Our major areas of interest include broad-range comparative genomics and the visualisation and interpretation of genomic variation, which is being increasingly studied in species throughout the taxonomy. We have developed a new portal for plant pathogen data, PhytoPath (launched in early 2012), and are involved in the development of Microme, a new resource for bacterial metabolic pathways.
The Protein Data Bank in Europe

Gerard Kleywegt

PhD 1991, University of Utrecht.
Postdoctoral research, University of Uppsala
Coordinator, then programme director of the Swedish Structural Biology Network, 1996-2009
Professor of Structural Molecular Biology, University of Uppsala, 2009.
At EMBL-EBI since 2009.

The Protein Data Bank in Europe (PDBe) is the European partner in the Worldwide Protein Data Bank organisation (wwPDB), which maintains the single international archive for biomacromolecular structure data. The other wwPDB partners are the RCSB and BMRB in the United States and PDBj in Japan. PDBe is a deposition and annotation site for the Protein Data Bank (PDB) and the Electron Microscopy Data Bank (EMDB).

The major goal of PDBe is to provide integrated structural data resources that evolve with the needs of biologists. To that end, our team endeavours to: handle deposition and annotation of structural data expertly; provide an integrated resource of high-quality macromolecular structures and related data; and maintain in-house expertise in all the major structure-determination techniques (e.g. X-ray crystallography, Nuclear Magnetic Resonance spectroscopy and 3D electron microscopy). Our specific focus areas are: advanced services, ligands, integration, validation and experimental data.

UniProt development

Maria J. Martin

BSc in Veterinary Medicine, Universidad Complutense, Madrid, 1990.
PhD in Molecular Biology (Bioinformatics) 2003, Universidad Autonoma, Madrid.
At EMBL-EBI since 1996.
Technical team leader since 2009.

Our team provides the bioinformatics infrastructure for the databases and services of the Universal Protein Resource (UniProt). The team consists of software engineers and bioinformaticians responsible for the UniProt software and database development, and the study of novel automatic methods for protein annotation and representation. A user-experience analyst in the team coordinates the user-request gathering process for the design and development of the website. The team is also responsible for the maintenance and development of tools for UniProt curation. We work in a fully complementary fashion with Claire O’Donovan’s UniProt Content group (see page 97) to provide essential resources to the biological community so that our databases can remain an integral part of the tools researchers use regularly for their work.
Literature services

Johanna McEntyre

PhD in Plant Biotechnology 1990, Manchester Metropolitan University.


Staff scientist, NCBI, National Library of Medicine, NIH, USA. 1997-2009.

At EMBL-EBI since 2009.

Direct access to the scientific literature and the data that underlie it have become increasingly important as data-driven science continues to trend upwards. The Literature Services team addresses this in a number of ways, supporting the wider scientific research community and our data-provider colleagues at EMBL-EBI by providing valuable, multi-layer functionality in Europe PubMed Central. Europe PubMed Central, now the sole literature database offered by EMBL-EBI, contains over 26 million abstracts and 2.7 million full text articles. The abstracts component includes all of PubMed, agricultural abstracts from Agricola and patents from the European Patent Office. About 700,000 of the full-text articles are open access, so they are free to read and to reuse in ways such as text mining. Europe PubMed Central is funded by 24 European funding organisations, whose commitment supports their own Open Access mandates. Our goal is to provide fast and powerful access to the literature, as well as features and tools that place the article narratives in the wider context of related data and credit systems such as article citations. One of the key approaches we employ to meet this goal is to engage with individual scientists, text miners and database managers to understand how layers of value can be built upon the basic article content. We provide the infrastructure that enables the enrichment of the literature by individuals and computational methods developed throughout the community, and publish the results to maximise the usefulness of the core content and allow their widest possible reuse.

UniProt content

Claire O’Donovan

BSc (Hons) in Biochemistry, 1992, University College Cork, Ireland.

Diploma in Computer Science, 1993, University College Cork, Ireland.

At EMBL since 1993.

At EMBL-EBI since 1994.

Technical team leader since 2009.

The UniProt Content Team comprises biocurators and bioinformaticians working on the Universal Protein Resource (UniProt). Biocuration involves the translation and integration of information relevant to biology into a database or resource that enables integration of the scientific literature as well as large data sets. Accurate and comprehensive representation of biological knowledge, as well as easy access to this data for working scientists and a basis for computational analysis, are the primary goals of biocuration. The goals of biocuration are achieved thanks to the convergent endeavors of biocurators and software developers and our team works in a fully complementary fashion with Maria Jesus Martin’s UniProt Development group (see page 96) to provide essential resources to the biological community such that databases have become an integral part of the tools researchers use on a daily basis for their work.
The Protein Data Bank in Europe (PDBe) is one of six core databases located at EMBL-EBI and is also a partner in the Worldwide Protein Data Bank organisation (wwPDB) along with the RCSB and BMRB in the United States and PDBj in Japan. The PDBe team manages two production systems: the weekly update of deposited data, and the weekly increment of new released data. These production data systems are managed within multiple Oracle® databases and support a large number of integrated web resources to collect data and disseminate information to the wider community. With the future provision of new services based on structure validation data there is an emphasis to extend the core databases and infrastructure. This will require an optimisation of the loading tools to manage both the increased amount of depositions expected and also the breadth of data required to support new services.

Our team comprises the Gene Ontology Editorial Office, Mouse Informatics and the former Functional Genomics Production teams. We provide ontology resources such as the Gene Ontology and Experimental Factor Ontology, and sample/phenotype resources such as the BioSamples database, Infrafrontier and the International Mouse Phenotyping Consortium (IMPC), among others. The team works with external collaborators on projects funded by the European Commission, the BBSRC, the Wellcome Trust, the National Institutes of Health and the National Science Foundation. Examples include PhenImageShare, which addresses the indexing of image annotation in the context of genomic data, so that images are accessible and queryable with biomolecular datasets. Another of our projects, Embryonic Phenotyping, captures and integrates mouse embryonic images with genomic and phenotypic data. DIACHRON addresses both the use of RDF technology to represent scientific data and strategies for exploring data life cycles in the interests of enabling data preservation. The inception of these projects expands our efforts in RDF generation and application building, ontology delivery and provision of community access to data.

In addition to microRNA data analysis, our research activities have focused on analyses (in R) of RNA-seq data in collaboration with the Geuvadis consortium and genotype imputation for individuals from the 1000 Genomes Project. Semantic web projects include development of an RDF representation of Gene Expression data from the Gene Expression Atlas and ArrayExpress.
Our team develops software for ArrayExpress, a core EMBL-EBI resource. A new project we started in 2013 is the BioStudies database, a resource for biological datasets that do not have a dedicated home within EMBL-EBI services. We also contribute to the development of the EBI BioSamples database, which centralises biological sample data.

Together with the Samples, Phenotypes and Ontologies team (formerly the Functional Genomics Production team), we build and maintain data management tools, user interfaces, programmatic interfaces, and annotation and data submission systems for functional genomics resources. Our team participates in several pilot projects in a data management role. We believe that being close to large consortia that generate different types of high-throughput data places us in a better position to fulfil our main objective of developing ArrayExpress and BioSamples infrastructures. Our participation in diXa, a toxicogenomics data management project, will enable us to build better links between BioSamples, ArrayExpress and other EMBL-EBI assay data resources. We are also partners on EU-AIMS, a project devoted to autism research, which will contribute to a deeper understanding of ways to manage complex endophenotype data, such as imaging data.

The Protein Data Bank in Europe (PDB) and is a founding member of the Worldwide Protein Data Bank (wwPDB). We accept and annotate worldwide depositions of biomacromolecular structures determined using X-ray crystallography, Nuclear Magnetic Resonance (NMR) spectroscopy, 3D Electron Microscopy (EM) and other structure determination methods. PDBe is also a founding member of EMDataBank, which manages the deposition and annotation of electron microscopy data in EMDB.

Our goal is to ensure that PDBe truly serves the needs of the biomedical community. As part of that effort, we are constantly improving the web interface for existing tools and services and designing new tools to make structural data available to all. In the context of the SIFTS project, we integrate structural data with other biological data in the interests of facilitating discovery. These integrated data form the basis for many query interfaces that allow biomacromolecular structure data to be presented in its biological context. Our specific focus areas are: data integrity, data quality, integration and data dissemination to the non-expert biomedical community.
EMBL Grenoble

Structural Biology

EMBL Grenoble is a key player in the Partnership for Structural Biology (PSB) which integrates the activities of several Grenoble institutes involved with structural biology and thus is able to provide a uniquely comprehensive range of platforms for both inhouse research and external users.

A cornerstone of the PSB is the close interaction of EMBL Grenoble with European Synchrotron Radiation Facility (ESRF), which involves collaboration on building and operating beamlines for macromolecular crystallography and small-angle scattering, developing instrumentation and techniques, and providing expert help to visitors. The highly automated ESRF crystallography beamlines are equipped with EMBL-designed high-precision micro-diffractometers and frozen crystal sample changers. A new X-ray small-angle scattering instrument built by ESRF and EMBL is now operational with a custom designed small-volume automatic sample changer.

High throughput methods have also been introduced in other steps of the structure determination process. These include a very successful robotic system for nanovolume crystallisation and a novel high throughput screening method, ESPRIT, which enables soluble protein domains to be identified in otherwise badly expressed or insoluble proteins. More recently, a Eukaryotic Expression Facility (EEF) has been established specialising in the expression of multi-subunit complexes in insect cells, building on and further developing the MultiBac method. All these platforms are now available to external users under the EU funded BioStruct-X projects.

A strong tradition at the outstation is the study of systems involving protein-nucleic acid complexes and viruses. Structural work on aminocycl-tRNA synthetases is particularly well known and has recently focussed on elucidation of the mode of action of novel boron-containing antibiotics, which target leucyl-tRNA synthetase. Projects involving protein-RNA interactions also include cryo-EM studies of the signal recognition particle and holo-translocon with the ribosome and other proteins and complexes involved in RNA processing, transport and degradation, such as the nonsense-mediated decay (NMD) pathway.

Other important areas include the analysis of mechanisms of transcriptional regulation, including at the epigenetic level: groups are working on the structural analysis of eukaryotic transcription factor and chromatin-modification complexes as well as elucidation of the mechanisms by which piRNAs (small non-coding RNAs) protect the genome. Another focus is the study of segmented RNA viruses, particularly influenza and bunyaviruses, with the aim of understanding how their polymerases replicate and transcribe the viral genome. Complementary to these are studies on the innate immune receptors which detect the presence viral RNA in infected cells and activate interferon production. The first crystal structures of domains of the influenza virus polymerase, determined in collaboration with other members of the CNRS-Grenoble University-EMBL Unit of Virus Host Cell Interactions (UVHCI), which is situated next to the outstation, have led to an important programme of anti-influenza drug design in collaboration with pharmaceutical companies.

Scientists at EMBL Grenoble have access to a wide range of techniques, including molecular biology and biophysical techniques, cryo-electron microscopy, isotope labelling, NMR, neutron scattering, X-ray crystallography and small angle scattering. A confocal microscope with facilities for cross-correlation spectroscopy is available for the study of complex formation in cells, as well as a top-end Polara electron microscope with direct detector and cryo-tomography capability.

Stephen Cusack
Head of EMBL Grenoble
Structural biology of RNA-protein complexes in gene expression and host-pathogen interactions

Stephen Cusack
PhD 1976, Imperial College, London, UK.
Postdoctoral work at EMBL Grenoble.
Group Leader, Senior Scientist and Head of Outstation since 1989.
Director of EMBL-Grenoble University-CNRS Unit for Virus Host Cell Interactions since 2007.
ERC Advanced Investigator.

Previous and current research
Aminoacyl-tRNA synthetases play an essential role in protein synthesis by charging specifically their cognate tRNA(s) with the correct amino acid. Structural information can help understand the catalytic mechanism of the enzymes and their substrate specificity for ATP, cognate amino acid and tRNA. In recent years we have focussed on leucyl-tRNA synthetase which contains a large inserted ‘editing’ domain able to recognise and hydrolyse mischarged amino acids – a proof-reading activity that is essential for maintaining translational fidelity. We have structurally characterised the large conformational changes required to switch from the aminoacylation to the editing configurations. We collaborated in the elucidation of the mechanisms of a new boron-containing anti-fungal compound (now in Phase III clinical trials) that targets the editing site of leucyl-tRNA synthetase and helped design related anti-bacterial compounds that are active against multi-drug resistant strains, including tuberculosis.

The nuclear cap-binding complex (CBC) binds to m7Gppp cap at the 5’ end of Pol II transcripts and mediates interaction with nuclear RNA processing machinery for splicing, polyadenylation and transport. We determined the structure of cap-bound human CBC, a 90 kDa heterodimeric protein, and have studied several other proteins involved in cap-dependent processes. Once in the cytoplasm, mRNAs are subject to a quality control check to detect premature stop-codons. In all eukaryotic organisms studied, this process, known as nonsense-mediated decay (NMD), crucially depends on the three conserved Upf proteins (Upf1, Upf2 and Upf3). In mammals, it is linked to splicing. We obtained the first structural information on binary complexes of these three proteins whose ternary complex formation triggers decay.

Future projects and goals
Ongoing projects related to RNA metabolism include continued studies on PHAX and ARS2, both of which bind CBC and are linked to the metabolism of small RNAs. A major focus is structure determination of the influenza virus RNA-dependent RNA polymerase, the viral replication machine. We have determined the structure of four distinct domains from the polymerase, including the two key domains involved in the ‘cap-snatching’ process of viral mRNA transcription: the cap-binding site in PB2 and the endonuclease in PA. These results give some insight into the polymerase mutations required to adapt an avian virus to be able infect humans and permit structure-based antiviral drug design. To pursue this we have co-founded a Vienna-based company called SAVIRA. This work is now being extended to the polymerases of other segmented RNA viruses such as bunyaviruses, which also perform cap-snatching. In collaboration with the Ellenberg (page 18) and Briggs (page 58) groups we are engaged in confocal and cross-correlation fluorescence studies as well as correlative EM microscopy of the assembly and trafficking of the influenza polymerase and RNPs in living, infected cells. Another major project is innate immune pattern recognition receptors such as NOD proteins and RIG-I like helicases. In 2011 we published the first structure-based mechanism of activation of RIG-I and are continuing to study this signalling pathway. Finally we have new projects on epigenetic complexes involving the histone acetylase MOF, such as the X-chromosome dosage compensation or male-specific lethal complex, which in Drosophila contains an essential non-coding RNA and the more recently discovered NLS complex (with the Akhtar group, MPI, Freiburg).

Selected references

Model of the activated state of RIG-I with bound dsRNA (centre) and ATP (top-right). The helicase domains (green and cyan), the insertion domain (yellow) and the C-terminal domain (gold) all contribute to RNA binding, which displaces the CARD domains thus allowing downstream signalling and interferon expression.
We develop and utilise advanced, automated technologies to produce eukaryotic multiprotein complexes for structural and functional analysis by a variety of methods including X-ray crystallography.

**Previous and current research**

Human gene transcription requires the controlled step-wise assembly of the pre-initiation complex (PIC), comprising a large ensemble of proteins and protein complexes including RNA Pol II and the general transcription factors. TFIIID is the first general transcription factor to bind to gene promoters, and is a cornerstone of PIC assembly. Understanding of TFIIID and its crucial role in transcription regulation is hampered by a lack of detailed knowledge of its molecular architecture, assembly in the cell, and interactions with chromatin and other factors. The paucity and heterogeneity of TFIIID in endogenous cells impede its extraction from cells for high-resolution structural and functional studies. Our lab addressed this challenge by creating new technologies for recombinant production of TFIIID and other complex protein machines. Notably, our MultiBac system – a modular, baculovirus-based technology specifically designed for eukaryotic multi-protein expression – is now used in many labs worldwide, in areas including structural biology, vaccine development and gene therapy vectors. Recently, we determined the architecture of the 700 kDa heterodecameric human TFIIID core complex by hybrid methods, combining MultiBac-based production, cryo-EM, X-ray crystal structures, homology models, and proteomics data. It is thought to represent a central scaffold that nucleates holo-TFIIID formation and provided first impressions on how the functional holo-TFIIID complex is assembled in the nucleus.

We collaborate with groups from academia and industry for technology development. We are striving to automate labour-intensive steps in the multiprotein complex structure determination process, and have harnessed homologous and site-specific recombination methods for assembling multigene expression plasmids. We have implemented a full robotics setup by developing ACEMBL, a proprietary automated suite for multigene recombineering on our TECAN Evoli platform. By using our technology, we produced numerous large multiprotein assemblies for structural studies, including multicomponent membrane protein complexes and the 1.6 MDa human TFID holo-complex and expanded our multiprotein expression strategies to prokaryotic and mammalian hosts.

**Future projects and goals**

We continue to advance our expression technologies to entirely automate and standardise the process of production for eukaryotic gene regulatory multiprotein complexes including the entire human TFIIID holocomplex, its various isoforms and other components of the pre-initiation complex. In collaboration with the Schaffitzel Team (page 109) and the Schultz Group (IGBMC Strasbourg), we subject the complex specimens produced to electron microscopic analyses. We use homogenous complexes thus identified for X-ray crystallography, aim to understand physiological function, and explore and challenge our findings by *in vitro* and *in vivo* biochemical analysis.

Using state-of-the-art mass spectrometric methods from systems biology, we are developing MultiTRAQ, a new technology addressing a further bottleneck in complex crystallography, namely the challenge of defining crystallisable core assemblies of multiprotein complexes in a reasonable time frame (a collaboration with ETH Zürich and Lund University). Another recent project line in our lab exploits synthetic biology techniques for genome engineering, with the aim of creating disruptive platforms for recombinant protein production, for both academic and industrial applications.
Previous and current research
The core activity of our team is to develop instruments and methods for X-ray scattering experiments in collaboration with the ESRF Structural Biology group, EMBL@PETRA3 and the McCarthy (page 106) and Márquez (page 105) teams. Strongly involving EMBLEM, the EMBL partner for technology transfer, an important part of our mission is to make our instruments available to the scientific community worldwide. The MD3 Kappa diffractometer developed in collaboration with EMBL@PETRA3 is now available to the users of the P14 beamline (led by Gleb Bourenkov, see: www.embl-hamburg.de/facilities/beamlines/P14). The combination of a small and intense beam with a fast X-ray detector and MD3 is ideal for collecting diffraction data from micron-sized needles and batches of micro-crystals using 4D scans or serial rastered data collection methods.

CrystalDirect™ (CD), a system jointly conceived with the Márquez team, is now available to users through the high throughput crystallisation (HTX) laboratory (see page xx). Crystals grown in CD plates can be marked remotely using CRIMS, the crystallisation laboratory information management system of the HTX lab, and automatically harvested and frozen in the CD lab. The automated removal of the solution surrounding the crystals is often sufficient to safely freeze the crystals, but a cryo-protective treatment can also be automatically applied before harvesting if necessary. More than 1500 crystals from 50 different proteins have been successfully harvested so far.

A basic setup to diffuse chemical solutions in crystallisation drops was recently integrated and the first ligand soaking experiment has been very encouraging. The intrinsic low background of the CD plates makes them ideal for in situ X-ray data collection. Specific goniometer supports were developed to process CD plates on the MD2 and MD3 diffractometers. In situ data collection is now offered on request at the EMBL/ESRF BM14 beamline. The first in-situ rastered data collections were made on the MD3 of P14, with batches of 5-20 micron-sized crystals of model proteins (see figure). Several hundred crystals in the same crystallisation drop were shot, producing thousands of diffraction images, from which high resolution structures were obtained. Similar experiments were successfully performed with crystals harvested in batches using CD and frozen, showing both the potential of serial data collection and the benefit of CD for this method. In parallel, studies are ongoing to develop a compact and precise sample holder for frozen crystals. Prototypes are available and a robotised test setup is installed on BM14. This collaborative project called ‘NewPin’ is part-supported by BioStruct-X.

Future projects and goals
A second CD harvester with extended crystal treatment capabilities will soon be built for projects involving ligands or derivatives. Our long-term ambition is to bridge the HTX lab and the BM14 beamline via CD as a pilot project for an automated and integrated MX facility. In the context of the ESRF MASSIF upgrade programme, we will equip the ESRF ID30B beamline with a MD2 diffractometer and new 6-axis robotic sample changing solution. The ultimate goal is to offer the ESRF user community a serial in situ and frozen data collection environment in a flexible robotised manner ready for the next generation of sample holders.

Selected references
High-throughput crystallisation laboratory

The Márquez team develops low volume, high-throughput, techniques to optimise protein crystallisation and uses them to study the structure of sensing and signalling molecules.

Selected references

José A. Márquez
PhD 1997, University of Valencia, Spain.
Postdoctoral research at EMBL.
Staff scientist at EMBL Grenoble since 2003.
Team leader since 2005.
Head of Crystallisation Facility since 2012

Previous and current research
The HTX lab is one of the major facilities for high-throughput nanovolume crystallisation screening in Europe and one of the major resources of Grenoble’s Partnership for Structural Biology, open to scientists working in European academic institutions through the EC-funded P-CUBE and BioStructX projects. The lab has offered services to hundreds of scientists, performing several million experiments. We are involved in the development of data management resources and new crystallisation techniques.

The Crystallisation Information Management System (CRIMS): CRIMS tracks experiments and makes results available to users via the web in real-time, along with all experimental parameters. It has been licensed to 10 other laboratories in Europe, three of them at synchrotron sites. Recently, data mining through CRIMS has allowed us to develop a new method to determine the crystallisation likelihood of a protein sample based on a simple assay measuring thermal stability (Dupeux et al., 2011).

Integration of crystallisation and synchrotron data collection facilities: While both highly automated platforms, recovering crystals and mounting them on supports remains a difficult and time-consuming manual process. In collaboration with the Cipriani Team (page 104), we developed Crystal Direct®, an approach that enables full automation of the crystal harvesting process (see figure). Crystals are grown on an ultrathin film in a vapour-diffusion crystallisation plate and recovered through laser-induced photo ablation. Advantages include: elimination of crystal fishing with X-ray data collection. The first prototype is now in operation and could benefit many projects.

Molecular mechanisms in sensing and signalling: Our research focus is on understanding the mechanisms of sensing and signalling at a structural level. Recently, we have obtained the structure of the receptor for abscisic acid (ABA), a hormone regulating the response to environmental stress in plants and shown how receptor dimerisation modulates ligand binding affinity leading to differential sensitivities towards the hormone (Dupeux et al., 2011). This provides a framework for understanding the ABA signalling pathway and activation of the stress response in plants, and illustrates how receptor oligomerisation can modulate ligand binding affinity by influencing the thermodynamics of the overall activation reaction.

Future projects and goals
In collaboration with the McCarthy (page 106) and Cipriani teams we will link the crystallisation screening service with the automated evaluation of crystals in situ with X-rays. New interfaces will be added to CRIMS to allow remote operation and we aim (with collaborators) to integrate CRIMS with synchrotron data management systems. We will develop vapour diffusion and microfluidic devices for crystal optimisation experiments. We aim to establish CrystalDirect for routine use in macromolecular crystallography and a series of pilot projects will be selected to establish standard protocols and develop new applications. Prototypes of the new plates will be distributed to other labs as part of the INSTRUCT project, and we will work towards the development of promising new approaches for crystal processing, including for cryo-protection, crystal soaking and crystal freezing. Towards this goal, an advanced version of the CrystalDirect harvester will be designed in collaboration with the Cipriani team.

The CrystalDirect technique could benefit challenging structural biology projects, such as studies of membrane proteins or multi-protein complexes.
Synchrotron crystallography team

Andrew McCarthy
PhD 1997, National University of Ireland, Galway.
Research associate, Utrecht University.
Postdoctoral research at Massey University and Auckland University.
Staff scientist at EMBL Grenoble.
Team leader since 2007.

Previous and current research

The Synchrotron Crystallography Team works in close collaboration with the Structural Biology Group of the European Synchrotron Radiation Facility (ESRF) in the design, construction and operation of macromolecular crystallography (MX) and biological X-ray scattering (bioSAXS) beamlines. We are currently responsible for the microfocus MX beamline on ID23-2 and the bioSAXS beamline at BM29 as well as the commissioning of MASSIF-1 on ID30A and the new tuneable MX beamline on ID30B. The team also manages the operation of BM14, which is run as a partnership with the ESRF and the Indian government. The structural biology beamlines at the ESRF are currently undergoing an exciting transformation following the closure of the ID14 suite last year and the start of commissioning on ID30. On BM14, we are working in close collaboration with the Cipriani Team (page 104) and are currently developing a new type of sample changer based on a 6-axis robot. We are also actively developing the use of workflows in the design and implementation of complex MX experiments for advanced sample screening and data collection.

In addition we study proteins involved in neuronal development, particularly the Slit-Robo signalling complex, and proteins involved in the synthesis of plant secondary metabolites. Meanwhile, the BM14 group is actively involved in the structural studies of proteins involved in the Toxoplasma gondii epigenetic machinery in collaboration with Mohamed-Ali Hakimi (Grenoble Medical University).

Future projects and goals

This year will be exciting with the completion of the next phase of the UPBL10 project with our ESRF colleagues, the first user operation of the new suite of MX-beamlines on ID30A (MASSIF) and ID30B. This ambitious project is part of the ESRF upgrade programme and will ensure that European users will have continued access to state-of-the-art structural biology beamlines for the next decade. On BM14, we will continue to develop a new sample changing robotic system and optimise the in situ screening of crystals in plates (see figure). We will continue to develop automated data screening, collection and analysis schemes, phasing methods using UV induced damage, and new workflow routines and display software using Eclipse-RCM technology as part of the DAWN and BioStruct-X collaborations respectively. We will further develop the highly automated BioSAXS beamline in collaboration with the ESRF, the Cipriani team and EMBL Hamburg, including the full integration of an online high-performance liquid chromatography system with additional biophysical characterisation features. We hope that our combined efforts will push the boundaries of structural biology to better understand how complex biological systems work at the molecular level.

In the laboratory we will continue our research on the Slit-Robo signalling complex by trying to decipher how exactly Slit activates Robo on the cell surface, and we will extend work on phosphoryl transfer into human kinase signal cascades. The BM14 group will over-express several Toxoplasma proteins in E. Coli to be structurally and biochemically characterised using the PSB facilities. Lastly, the initial work on a novel dequibiquitinase will be continued.

Selected references


BM14 in an in situ plate screening setup mode.
The Panne group looks to understand important signalling processing pathways in the cell, which could help in the discovery of anti-viral drugs.

**Selected references**


**Previous and current research**

Cellular control logic is ultimately embedded in the molecular architecture of the molecular machines that make up the living cell. Many molecular machines, especially complexes involved in cellular signalling, are transient, with a variety of states and a succession of structures. For example, transcription factors such as NFkB, IRF3, IRF7 and ATF2/c-Jun bind as complexes to enhancers (‘enhanceosomes’) in a combinatorial and dynamic fashion to regulate expression of genes (figure 1). Enhanceosome assembly is mediated by a set of IKK kinases such as TBK1 that regulate NFkB and IRF3/IRF7 activation. These signalling components interact with each other and with other molecules in highly structured but complex ways.

Understanding such transient and dynamic complexes of the cellular machinery is one of the most important challenges in biology today. One important first step toward characterising such dynamic processes is to determine the molecular architecture of essential components. We are using a combination of biophysical techniques including X-ray crystallography, electron microscopy, native mass spectrometry, and more to address the following questions: What is the architecture of signalling complexes that direct innate immune responses? How do these signalling pathways lead to assembly of higher-order regulatory complexes? How does assembly of such transcription factor complexes ultimately lead to chromatin modification? How does chromatin modification direct nucleosome remodelling and gene regulation?

**Future projects and goals**

Cellular signalling ultimately results in assembly of transcriptional regulatory complexes that direct chromatin modification, remodelling and gene expression. The enhanceosome has served as a paradigm for understanding signal integration on higher eukaryotic enhancers (figure 2). Assembly of the enhanceosome results in recruitment of enzymes such as CBP/p300 that acetylate chromatin. We aim to understand how recruitment of CBP/p300 allows cellular signal integration and chromatin acetylation. We ask how chromatin acetylation changes the structure of inhibitory nucleosomes and leads to a more permissive chromatin structure for gene expression. We also ask how CBP/p300 read out histone modification patterns, and how chromatin recognition and modification are coupled. Answers to some of these questions are likely to contribute to our understanding of epigenetic gene regulation and dysregulation in disease. This is not only of fundamental importance for cellular signalling, but also opens up opportunities for pharmacological targeting.
Regulation of gene expression by non-coding RNAs

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Previous and current research

Past invasion events from mobile genetic elements have left eukaryotic genomes littered with repeats and other transposon sequences. Much of these are inactive fossils, but some still retain the potential to get activated and cause genome instability. Protection from transposons is achieved by silencing them in the germline, which is then maintained throughout the life of the individual. Animal germ cells express a specialised class of ~30 nt small non-coding RNAs called piwi-interacting RNAs (piRNAs), which are implicated in guiding this silencing. Indeed, one universal feature of piRNAs in all animals is their origin from transposon-rich genomic regions. In mammals, they are believed to recruit DNA methyltransferases to transposon sequences. In Drosophila, maternally produced piRNAs are deposited in the egg and they contribute to protection from new transposons brought in by the paternal genome. Thus, piRNAs constitute an epigenetic component of the genome defence mechanism in animals.

Our lab is interested in understanding the molecular mechanisms involved in piRNA biogenesis and function. A striking feature of piRNAs is their clustered genomic origins. It is believed that a long single-stranded transcript arising from a cluster is processed into thousands of piRNAs. The mechanism of this processing and the identity of factors involved are unknown. We have taken a biochemical approach to identify these factors by isolating mouse Piwi-associated proteins. This led to the identification of Tudor domain-containing protein 1 (Tdrd1), which interacts by recognising symmetrical dimethyl arginine modification marks on Piwi proteins. Another factor is the putative helicase Mov10l, which is an essential piRNA biogenesis factor, as piRNAs fail to accumulate in mutant mice. In all these studies, we have used a variety of techniques ranging from protein biochemistry, cellular imaging, small RNA bioinformatics, and mouse mutants. We are now setting up insect cell culture lines that have an active piRNA pathway, paving the way for potential mechanistic insight into the function of the identified factors. To deepen our understanding, we collaborate with structural biologists to obtain atomic resolution images of the identified pathway components. Recently, this effort resulted in a structure describing the recognition of the 2’-O-methyl mark on piRNAs by the PAZ domain of a Piwi protein.

Future projects and goals

We will continue to analyse additional factors identified in our complex purifications. Another goal is to understand the features that define genomic regions as piRNA clusters, and whether there is a link between transcription from the clusters and piRNA biogenesis. We also hope to use live cell imaging techniques to study assembly of small RNPs in vivo and define the contribution of the individual constituents of the complex to this process. It is our desire to intensify the collaborative work on structural biology of Piwi complexes, adding another dimension to our understanding of germline small RNAs. In addition to small RNAs, our cells express longer non-coding RNAs (ncRNAs), which are implicated in a variety of gene regulatory functions, usually in epigenetic roles. We wish to apply biochemical methods to identify protein components of long ncRNPs to understand their contribution to the molecular function of the RNA.

Selected references


Localisation of a tagged insect Piwi protein to perinuclear cytoplasmic granules in insect cell cultures. These are putative piRNA biogenesis sites, similar to the nuage in germ cells.
Ribosomal complexes: targeting, translocation and quality control

The Schaffitzel team combines molecular biology, biochemistry and cryo-electron microscopy to study large macromolecular complexes in protein targeting, secretion and membrane protein integration.

Selected references

Previous and current research
Research in our laboratory combines molecular biology, in vitro and in vivo biochemistry, and single-particle cryo-electron microscopy (cryo-EM) to study the structure and function of ribosomal complexes. Synthesis and folding of proteins require concerted interactions of the translating ribosome with translation factors, regulatory factors, molecular chaperones, and factors involved in the export of proteins. Structures of translating ribosomes in complex with these factors provide critical insight into the interaction networks, stoichiometry, and molecular mechanisms of these megadalton-size complexes. Using cryo-EM, we can study the multi-component translation machinery at close to physiological conditions. By using state-of-the-art electron microscopes and by image processing of large data sets, EM structures of prokaryotic and eukaryotic ribosomes have been obtained at subnanomolar resolution, demonstrating the power of this method.

A prerequisite for our functional and structural studies is the production of large amounts of homogenous, stable complexes in the quantity and quality required for interaction assays, mass spectrometry and single-particle cryo-EM. We have established bacterial and eukaryotic cell-free translation systems for the in vitro generation of ribosomes that display homogenous nascent polypeptide chains or have stalled at a defined step in translation. We reconstitute the ribosomal complexes along the pathways of co-translational targeting and translocation, and mRNA quality control. This approach was successfully applied in the case of the cryo-EM structures of the complex of the ribosome with the translocation machinery SecYEG (figure 1), of the translating ribosome-signal recognition particle (SRP) complex, and of the ribosome in complex with SRP and SRP receptor (figure 2). The data from intermediate resolution structures derived from cryo-EM, in conjunction with high-resolution structures of the ribosome and of the isolated factors, were combined in a hybrid approach to generate quasi-atomic models of the ribosomal complexes involved. The structural data, supported by biochemical data, provide important and detailed snapshots of the mechanisms underlying these cellular processes, ensuring correct folding, targeting and translocation of nascent proteins.

Future projects and goals
We study ribosomal complexes involved in targeting, membrane protein integration, folding, and assembly. We analyse the membrane protein complexes biochemically, by cross-linking/mass spectrometry (collaboration with Juri Rappisilber, TU Berlin) and single-particle cryo-electron microscopy. In collaboration with the Casack (page 102), Hentze (page 14) and Kulozik (Molecular Medicine Partnership Unit) groups, we study mammalian ribosomal complexes involved in nonsense-mediated mRNA decay. We produce the eukaryotic factors involved using advanced recombinant eukaryotic technologies in collaboration with the Berger group (page 103). Finally, our team collaborates with a number of groups to solve the structures of large macromolecular complexes in transcription, epigenetics and cellular signalling.

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Figure 1: EM reconstruction of the ribosome nascent chain complex and the translocon.

Figure 2: Cryo-EM structure of the ribosome (yellow/blue) bound to the signal recognition particle (SRP) and the SRP receptor (both in red). Below it is an atomic model of SRP (green-yellow/orange) and its receptor (pink).
EMBL Hamburg

Structural Biology

Activities at EMBL Hamburg focus on state-of-the-art structural biology methods using synchrotron radiation, combining cutting-edge technology with an ambitious research programme for structures of multifunctional proteins and protein complexes of biomedical relevance.

EMBL Hamburg’s laboratories are on the German Synchrotron Research Centre (DESY) campus, with synchrotron radiation (PETRA III) and laser (FLASH) facilities available. In addition, a powerful X-ray free electron laser is under construction. EMBL operates a new integrated facility, called EMBL@PETRA3, for applications in structural biology at the PETRA III ring. It comprises three state-of-the-art beamlines for macromolecular X-ray crystallography and small angle X-ray scattering of biological samples, complemented by facilities for sample preparation and characterisation, and data evaluation. EMBL Hamburg is also one of the main partners in the future Centre for Structural Systems Biology (CSSB) on the DESY campus.

EMBL Hamburg has set up an ambitious research programme for structures of multifunctional proteins and protein complexes of biomedical relevance. Present research interests of group leaders include cell surface receptors, protein assemblies in muscle cells, protein kinases, protein translocation into peroxisomes, and several projects relating to tuberculosis. Common to all projects is the goal to make optimum use of on-site high-brilliance synchrotron radiation and to explore novel opportunities of the X-ray Free Electron Laser. Beyond the tools in structural biology that are available on-site, EMBL Hamburg groups are engaged in many interdisciplinary collaborations with colleagues from other EMBL units, enabling access to a large variety of in vitro and in vivo functional techniques, including cellular imaging techniques.

EMBL Hamburg also has a well-established record for the development of novel, innovative technologies in structural biology. Leading software packages for the automation of data interpretation have been developed here and are used in a large number of projects across the world’s research community. One example is the ARP/wARP package that allows automatic X-ray structure determination. Another package, ATSAS, allows the automatic interpretation of small angle X-ray scattering data for structural shape determination. Finally, there are two groups that focus on the development and construction of new equipment for experimental stations in structural biology, using synchrotron radiation. Present efforts focus on the installation of new robotics that allow automatic placement of biological samples into specialised synchrotron experiment facilities.

Matthias Wilmanns
Head of EMBL Hamburg
Structure and function of protein complexes in biological systems

The architecture of the protein interactome in sarcomeric muscle cells: Many proteins found in muscle cells, when dysfunctional, are associated with cardiovascular diseases. We investigate how large protein filament systems forming the overall architecture of ‘sarcomeric units’ in muscle cells – such as actin, myosin, nebulin, titin, myomesin and obscurin – are connected and interact with each other, frequently mediated via small scaffold proteins. We have determined the structure and function of some key complexes, including telethonin-mediated assembly of the N-terminus of titin (Zou et al., 2006) and the overall architecture of the elastic filament protein myomesin (Pinotsis et al., 2008; Pinotsis et al., 2012). Our future focus will be on novel protein interactions within the sarcomeric Z-disk and M-line region, and novel signalling functions of the protein partners involved.

Activity regulation in protein kinases: The human kinome – the part of the genome that expresses protein kinases – comprises about 600 genes. About 70 protein kinases share a common C-terminal autoregulatory domain. To investigate the mechanism of activity regulation in these kinases, we first determined the structure of the kinase domain from the giant filament protein titin, in the inhibited apo-conformation (Mayans et al., 1998). More recently, we have unravelled the structure of the apoptotic Death Associated Protein Kinase-1, in the presence of the regulatory scaffold calcium/calmodulin (CaM) (figure 1). This structure provides insight into how CaM binding leads to kinase activation by withdrawing the autoregulatory domain from the kinase active site. Our goal is to complement ongoing structural studies by in vitro and in vivo functional studies, to decipher underlying, general molecular mechanisms that regulate the activity of members of the CaM-dependent protein kinase family and, ultimately, to promote drug discovery against those kinases involved in cancer formation and progression.

The architecture of the translocon of peroxisomes: Peroxisomes are cell organelles that allow sequestered metabolic processes that would interfere with other processes in the cytosol. Proteins involved in these processes are generally translocated as active and folded targets. For the first time, we have unravelled the mechanism involved in the recognition of peroxisome protein targets by the peroxisome import receptor Pex5p, by determining the structure of the cargo-binding domain of the receptor in the absence and presence of the cargo protein sterol carrier protein 2 (Stanley et al., 2006) and alanine-glyoxylate aminotransferase (figure 2). Our goal, with support of two national and international research networks, is to provide insight into the overall architecture of the peroxisomal translocon, using a broad range of structural biology and imaging methods, complemented by genetic and cell biology-oriented approaches.

Structural systems biology in M. tuberculosis: We have determined the X-ray structures of a number of protein targets, some of them of known function and others unknown. For instance, we were able to identity Rv2217 as a novel cysteine/lysine dyad acyltransferase, which allows activation of several important protein complexes by lipoylation (Ma et al., 2006). Using available structural data and with support of European research network systeMTB, we aim to use systems biology-oriented approaches (such as proteomics, metabolomics, lipidomics and transcriptomics) to investigate functional processes in living mycobacteria, with the aim of making data available to promote the development of new drugs, vaccines and diagnostic markers.
Synchrotron instrumentation for structural biology beamlines at PETRA III

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**Selected references**


**Previous and current research**

EMBL has designed, built and operates three beamlines for structural biology at the PETRA III synchrotron radiation source on the DESY campus in Hamburg. Beamline facilities are dedicated to the leading techniques for X-ray-based structural research of biological samples: small angle X-ray scattering and macromolecular crystallography. Our team provides expertise in X-ray optics, precision mechanical engineering, robotic, control software and electronics and is in charge of the X-ray optical elements, experimental endstations, vacuum system, cryogenic system, control system, data acquisition system, technical infrastructure and parts of the civil engineering.

**Recent major projects:** All three beamlines reached regular user operation. In preparation for this, it was necessary to install and commission adaptive focusing optics at all beamlines, develop a multi-degree-of-freedom heavy-duty detector stage for the large-area pixel detectors at the MX beamlines (P13 instrument installed) and to develop and install white beam X-ray monitor systems for all beamlines. A cryogenic supply system for the beamline endstations has been installed and is in operation and a two-floor control hutch area with computing infrastructure was set-up in order to allow beamline control and users to perform experiments in a comfortable manner.

**Multilayer optics:** An ongoing in-house development is the construction of a double multilayer monochromator that is intended to boost the flux density at the P12 SAXS beamline – this enables time-resolved solution scattering experiments in the microsecond range. This instrument has been further developed from a prototype installed at a beamline of the former DORIS storage ring that can preserve the highly increased coherence of the radiation delivered by the PETRA III synchrotron. We are also working on the development of downstream experimental instrumentation with the level of synchronisation necessary for dynamic measurements.

**Automation:** For all beamline facilities, it is important to develop capabilities such as automatic operation and remote access. In this context, a robotic sample mounting system named MARVIN is being developed for the MX applications (see figure). This is characterised by high sample storage capacity, high sample mounting speed and flexibility. It is integrated, like all beamline elements, into a software-based control system which allows for a heterogeneous control environment and provides distributed access. The prototype has been in user operation on the BW7b beamline at DORIS and adapted versions are undergoing installation on the MX beamlines at PETRA.

**Future projects and goals**

- Positional and intensity feedback and (active) vibrational stabilisation of the monochromatising optics.
- Automatic tuning of adaptive focusing mirror optics to vary beam size/shape.
- Integration of beamline elements into a global instrument protection system.
- Further automation of alignment and data acquisition and integration with sample preparation/crystallisation.
- Improved sample observation and positioning at or beyond the optical resolution limit for microcrystallographic investigations.
- Exploring preparative or bridging developments for the X-ray free electron laser with state-of-the-art synchrotron beamlines.

*Design of the P14 experimental endstation with sample mounting system MARVIN.*
To fully understand the function of biological systems, accurate structures of their components - DNA, RNA, proteins, macromolecular complexes and assemblies - are required. We apply and develop cutting-edge computational methods and experimental approaches for sample quality control, experimentation and data interpretation in macromolecular crystallography and with the use of X-ray free-electron lasers that may also have potential use in electron microscopy.

Previous and current research

Methods for biological structure determination: We develop a comprehensive range of algorithms for protein/ligand/DNA/RNA X-ray crystal structure determination and new procedures for dealing with challenging problems (Hattne & Lamzin, 2011). We exploit inherent properties of macromolecular structures (Wiegels & Lamzin, 2012) and integrate additional information derived from a priori knowledge and dedicated databases. The group’s main methodological focus is the ARP/wARP software project (Langer et al., 2008) for macromolecular crystallography, which is based on the use of pattern-recognition methods. An intuitive and user-friendly molecular viewer – the ArpNavigator (Langer et al., 2013) – enables user control of the model building process and provides easy access to a range of methods for quality assessment and model completion.

Structure-based drug design: We make use of various novel algorithms and, through their combination (Langer et al., 2012), develop new tools for drug discovery. The VICI software, available for use through an online webservice (www.embl-hamburg.de/vici), allows for the in silico screening of known ligands to provide new leads for drug design. Our interest in this direction is stimulated by our research into the biology of pathogenic species associated with human morbidity and mortality, and is focused on the probing of bacterial antibiotic resistance.

Biological imaging with Free-Electron Lasers (FEL): Breathtaking results from initial diffractive imaging experiments using coherent FEL radiation (Siebert et al., 2011) show the potential for imaging cellular organelles and understanding dynamics of complex formation. In order to exploit the numerous novel and unique opportunities for structural biology that will be provided by the European X-ray FEL source near DESY (www.xfel.eu), we are developing protocols for handling of biological samples for FEL experiments as well as novel computational methods for the interpretation of measured data (Mancusi et al., 2012). We succeeded in 3D imaging of frozen-hydrated S. cerevisiae cells using ptychographic coherent X-ray diffractive imaging.

Other targets of biomedical interest: We integrate X-ray crystallography, lower resolution imaging, biochemistry, computational biology and biophysical methods in order to investigate targets of biomedical interest. These include the nuclear pore complex and hydrophobins (Kallio & Rovinen, 2011), which may be used in drug delivery to solubilise hydrophobic pharmaceuticals. We also investigate the pathway of amyloid fibril formation via class I hydrophobins and fragments of human gelsolin.

Future projects and goals

The group will continue to focus on crystallographic software development driven by general academic interest, provision of state-of-the-art beamline facilities at PETRA III in Hamburg, future applications of such developments to FEL-based diffraction, and by the potential use of such developments in projects of medical or biotechnological importance. Together with our international collaborators, we will undertake novel pilot projects aiming at interpretation of structural data obtained from various sources.

Selected references


A selection of model-viewing options in ArpNavigator. Shown clockwise from the top left are a stick representation in solid electron density, a ball-and-stick representation in planar density, a skeleton representation of the electron density shown as a mesh and the protein in cartoon representation in planar density.

A known inhibitor in green, aligned with two hits from the VICI software, in the binding pocket of beta lactamase. Important interactions maintained are highlighted with red circles and those created for exploration with full red spots.
Structural and dynamic insights into nutrient uptake systems

Using biophysical and biochemical methods, the Löw group aim to enhance understanding of the structural basis for substrate recognition in peptide transporters.

Selected references

Previous and current research
Cell membranes compartmentalise metabolic processes and present a selective barrier for permeation. Therefore, nutrient transport through the plasma membrane is essential to maintain homeostasis within the cell. Many proton-coupled secondary active transporters of the major facilitator superfamily (MFS) are involved in the accumulation of nutrients above extracellular levels in the cell. Structural and functional analyses of MFS transporters suggest an alternating-access mechanism for the transport of substrates across the membrane. Here the transporter adopts different conformational states, allowing the substrate binding site to face either side of the membrane. A full transport cycle involves at least three different conformational states (inward open, occluded and outward open), with each of them in a ligand-bound and ligand-free state.

Proton coupled oligopeptide transporters of the PepT family (also known as the POT family) are responsible for the uptake of a range of different di- and tripeptides, derived from the digestion of dietary proteins, and are highly conserved in all kingdoms of life. The best studied members of this family include the two human peptide transporters, PepT1 and PepT2. Besides their role in uptake of short-chain peptides, PepT1 and PepT2 are also of great pharmacological and pharmaceutical interest since they also accept a number of drugs and amino acid-conjugated pro-drugs as substrates. A detailed understanding of the structural basis for substrate recognition can therefore help to convert pharmacologically active compounds into substrates for PepT1 and PepT2 and thus improve their absorption in the small intestine and subsequent distribution in the body. To this end we will study the proton-dependent oligopeptide transporter (POT) family using a combination of biochemical and biophysical methods. POTs share the canonical fold of MFS transporters with 12 predicted transmembrane helices each. There are currently no crystal structures available for any of the human PepT transporters, but the first bacterial PepT structures have recently been reported (including one from our lab).

Future projects and goals
- Structural and functional studies of the reaction cycle of bacterial POTs using X-ray crystallography.
- Structural and dynamic insights into the binding mode of POTs to peptides, drugs, and inhibitors.
- Functional expression, purification, biochemical characterisation and crystallisation of eukaryotic POTs.

Integral membrane proteins are a challenging class of proteins in terms of their structural and functional characterisation. Over the years we have developed and established new tools and a workflow for protein production and quality control of membrane proteins including functional assays (in whole cells or in reconstituted systems) with the major focus on nutrient uptake systems. We will make use of the newest synchrotron radiation at PETRA III, where EMBL Hamburg operates two beamlines for macromolecular crystallography. These beamlines are integrated into advanced facilities for biological sample preparation, characterisation and crystallisation as well as for X-ray data processing and evaluation.
Previous and current research

Intercellular interactions occur through supramolecular clusters that form asymmetric kinapses when a cell is scanning the environment, and symmetric synapses when cell-cell recognition is established. This ancient recognition process has been observed in as diverse systems as predatory amoeba, lymphocytes scanning for antigens, and nerve cells seeking connections to form a brain map. At the centre of these interactions are very specific molecular recognition events that trigger a reorganisation of the cluster on the cell surface, amplifying the recognition event, resulting in the activation of a signalling cascade within the cell that leads to physiological changes.

Immune receptors: T cell receptors (TCR) on lymphocytes are the central recognition unit of a supramolecular complex that scans for pathogenic peptides loaded on MHC molecules at the surface of host cells. We have studied the molecular basis for the dominant response of the adaptive immune system to certain influenza peptides, which provides insights that should help in the design of a peptide vaccine against influenza (Meijers et al., 2005). We have also studied the interaction between the T cell receptor/MHC complex and CD4, a cell surface receptor that acts as a co-receptor in T cell recognition (Wang et al., 2001). CD4 is a prime fusion target of the HIV virus, and the structure between CD4 and the MHC class II molecule I-Ak shows that the viral envelope protein gp120 and the MHC molecule bind the same region of CD4. The structure confirmed that immunodeficiency is caused in part by the disruption of the binding of MHC class II molecules to CD4, an essential coreceptor in the supramolecular TCR complex.

Neuronal receptors: The nervous system consists of neuronal circuits, and it is thought that the individual neurons find their targets and establish synaptic connections within the circuit following a specific developmental program. Some of the molecules that guide the neurons to their targets are cell surface receptors that provide a unique identity tag to each neuron. The Down syndrome cell adhesion molecule (Dscam) from Drosophila was identified as an axon guidance receptor that has the potential to provide a large number of unique identity tags. Dscam contains three variable extracellular immunoglobulin domains, which can provide 19 008 unique cell surface receptor identity tags through splicing. We have studied the structural basis for the remarkable specificity of the receptor, which seems to interact only with receptors that consist of identical isoforms (Meijers et al., 2007). The structure shows how variable regions of the Dscam receptor interact through a palindromic hydrogen bonding network that is unique for each isoform.

Future projects and goals

Viruses and bacterial pathogens use cell surface receptors to invade host cells, but they also deregulate the order established in the supramolecular cluster to jam the recognition machinery of the host. The same (de)regulation mechanism is used by the immune system itself, which employs internal regulators that act on the synapse (such as hormones and cytokines) to tune the response of the immune system. We are using molecular fragments of pathogenic and self-regulating factors to gauge the interactions with and within the supramolecular complex. A detailed understanding of these interactions will allow us to tinker with cell surface receptors in order to manipulate the behaviour of certain individual cells.

The identification tag of the Dscam receptor is encoded in a palindromic hydrogen bonding network. Two receptors that contain the same code bind together, but a slight difference in amino acid sequence prevents binding. The Drosophila Dscam gene is spliced to generate 19 008 different receptors that provide each neuron in the brain with a unique identity tag. This helps the neurons to orient themselves, and to decide where to form a connection with another neuron.
Selected references


Previous and current research

During the past six years, we built three beamlines at the new PETRA III synchrotron in close collaboration with the Cipriani (page 104) and Fiedler (page 113) teams. Since 2012, the beamlines for small angle X-ray scattering on solutions (SAXS) and X-ray crystallography on crystals of biological macromolecules (MX) are open for users. The beamlines are embedded in an Integrated Facility for Structural Biology that supports non-specialists in taking a project from producing a suitable sample to determination of the structure by SAXS and/or MX.

Partly due to the enormous progress in synchrotron radiation-based structural biology, structural data on biological macromolecules are produced at an ever-increasing rate, creating the need to develop tools for efficient mining of structural data. We are developing tools for which the central concept is to use coordinate errors throughout all calculations. The necessity of this approach becomes clear when one considers that in contrast to sequence data, where a nucleotide entry can only be right or wrong, the precision in the location of an atom in a crystal structure can vary over several orders of magnitude. While the position of an atom in a rigid region of a protein giving diffraction data to high resolution may be known to within 0.01 Å, for an atom in a flexible region of a poorly diffracting protein the coordinate error may reach more than 1.0 Å.

From a technical point of view, extracting information from large amounts of raw structural data (as many as hundreds of structures containing thousands of atoms each) is a very complex task and requires sophisticated algorithms, both for the analysis and for the presentation and 3D visualisation of the results.

Future projects and goals

We will continue the commissioning work on the new beamlines and strengthen the user programme to full capacity. The two beamlines for macromolecular crystallography will offer excellent conditions for data collection on small (micron-sized) crystals, crystallographic phasing, and the development of new experimental protocols for challenging systems.

On the computational side, we will work on improving the error models underlying our methods and on expanding our computational framework using genetic- and graph-based algorithms. We also plan to use recurrent structural fragments extracted from ensembles of structures as search models in molecular replacement and for the interpretation of low-resolution electron density maps. In fact, this aspect of our computational work will be very helpful in the interpretation of experimentally phased electron density maps obtained on the PETRA III beamlines.

For further information, see: www.embl-hamburg.de/facilities/mx.
Small-angle X-ray scattering from macromolecular solutions

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At EMBL since 1991, Group leader since 2003, Senior scientist since 2011.

Previous and current research

Small-angle X-ray scattering (SAXS) reveals low-resolution (1-2 nm) structures of biological macromolecules and functional complexes in solution. Recent experimental and methodical developments have significantly enhanced the resolution and reliability of the SAXS-based structural models, and the last decade saw a renaissance of biological SAXS worldwide.

Our group leads the development of novel computational methods for constructing structural models from the scattering data. Special attention is given to the joint use of SAXS with other methods including crystallography, NMR, electron microscopy and bioinformatics. We developed the world’s most used program package, ATSAS, employed by more than 8000 users from more than 50 countries.

Our group runs a dedicated high brilliance synchrotron beamline P12 at DESY’s third generation storage ring, PETRA III. P12 has a robotic sample changer for rapid automated experiments, and possesses a data analysis pipeline for building structural models online. The beamline offers FedEx-style and remote data access options, as well as an in-line purification and biophysical characterisation setup using size exclusion chromatography (Malvern).

In collaborative projects, group members help users not only with data collection, but also with structural modelling. SAXS is employed to study overall structural organisation of macromolecules, conformational transitions such as upon ligand binding (figure 1), and also to quantitatively characterise oligomeric mixtures (figure 2), intrinsically unfolded proteins, hierarchical systems and other objects of high biological and medical importance.

Future projects and goals

- Further methods development for the reconstruction of macromolecular structure from X-ray and neutron scattering.
- Hybrid applications of SAXS with crystallography, NMR, electron microscopy and bioinformatics to construct and validate structural models.
- Participation in collaborative SAXS projects at the P12 beamline.
- Further extension of P12 including time-resolved and anomalous scattering approaches.

Selected references


Figure 1: A spectacular quaternary structure transition of a protein complex upon tRNA binding (Koehler et al., 2013)

Figure 2: Model of estrogen-related receptor ERα in complex with IRE1 inverted repeat DNA obtained from polydisperse SAXS data (Petoukhov et al., 2013).
Dynamic partnerships and exchanges with other international academic research and clinical centres, and participation in multiple EU-wide mouse research and informatics initiatives, are integral parts of our discovery process into genetics and genomics, cell biology and pathology.

The continued refinement and diversification of methodologies for introducing genetic and epigenetic perturbations of cellular and physiological function is allowing the outstation to generate ever more accurate mouse models of human disease and multigenic disorders. The use of engineered nucleases in particular is allowing the ‘democratisation’ of experimental perturbation to virtually any mouse strain, allowing much more sophisticated approaches to be applied to understanding the complex interactions occurring between genes and across the genome—interactions which will likely have both epigenetic and environmental components.

Research groups at EMBL Monterotondo are using these powerful tools to investigate wide-ranging aspects of mammalian biology, including gene expression during early embryogenesis, haemopoietic development and differentiation, cancer and regeneration, behaviour and sensory perception.

A state-of-art animal facility provides a full range of mouse transgenic and gene knock-out production, embryo redervation and cryopreservation services, together with a specialised phenotyping suite. Other centralised facilities include histology, confocal microscopy and flow cytometry. The many interactions with groups on other EMBL campuses including through the Interdisciplinary Postdoctoral (EIPOD) Fellowship Programme provides exciting and continuing prospects for joint projects. Ongoing international collaborations include those with groups at the University of Florida, the MPI Freiburg, the EPFL in Lausanne, the Gordon Institute in Cambridge, the Institut Pasteur in Paris, and the CNIO in Madrid Spain.

Philip Avner  
Head of EMBL Monterotondo
Dynamics of epigenetic regulation

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Previous and current research

The genetic material of the cell is not all equally available for transcription and this availability, which varies with cell type and developmental stage, is mediated largely by epigenetic modifications to the genome playing out mainly at the level of the chromatin. The double focus of our research has been on mouse genetics and epigenetics, with a particular emphasis on the interface between genetics and epigenetics, as revealed using X-inactivation as an experimental paradigm. X-inactivation, which occurs early during development in female mammalian embryos, ensures the dosage compensation between females carrying two copies of the X chromosome and males with a single copy of the X. Parts of the process in the mouse can be modelled ex vivo using female ES cells. We have been at the forefront of research into the characterisation and functional analysis of the different components of the X-inactivation centre, the key complex on the X chromosome for the initiation of X-inactivation. Ongoing research involves, for instance, the study of the Xce locus (X-controlling element), a genetic locus existing in different forms, which appears to influence which of the two X chromosomes will be chosen to undergo X-inactivation, and studies on the basis of the differences in the stability of the X-inactivation process occurring in the different extra-embryonic lineages of the mouse.

By carrying out this research we are not only discovering the multiplicity and plasticity of mechanisms that feed into this process of epigenetic control but also providing insights into the links between epigenetic processes and development. Our approach involves a combination of genetics, genomics, biochemistry and cell biology and both ex vivo and in vivo experimental approaches.

Future projects and goals

Objectives of our future work will be to gain further mechanistic insights into variations in the X-inactivation process that occur in different cell lineages and the sensitivity of the process to trans-acting genetic factors. We will use our knowledge of the X-chromosome to explore more widely the contribution of epigenetic regulation to allele-specific epistasis, the process where the effects of one gene are modified in an allele-specific manner by one or several other genes.

The Avner group combines genetics, genomics, biochemistry, and cell biology to study the nature of the key complex on the X chromosome responsible for X-inactivation.

Selected references

Chureau, C., et al. (2011). Ftx is a non-coding RNA which affects Xist expression and chromatin structure within the X-inactivation center region. Human Molecular Genetics, 20, 705-18

Mouse female trophoectoderm stem cells: The XIST non-coding RNA (green) shows partial overlap with the repressive H3K27Me3 histone mark on the inactive X chromosome (Morey et al.).
Developmental programming of behaviour

The Gross group uses pharmacological, histochemical, electrophysiological and behavioural genetic approaches to study the neural circuits underlying behaviour in mice.

Selected references

Previous and current research
The laboratory is interested in understanding at a molecular and neural circuit level how early life events influence brain development in order to establish behavioural traits in adulthood, with a particular focus on fear and anxiety. We are currently pursuing two areas of research:

Neural circuits encoding fear and anxiety: Fear is a mental state that is elicited by exposure to threats or cues that signal those threats. Fear is part of an organism’s natural defense mechanism and the accompanying behavioural and physiological responses are essential for it to cope with potential bodily harm. However, in its pathological form, fear can become excessive or inappropriate – features associated with anxiety disorders. It is accepted that the amygdala plays a central role in processing fear. However, it is less widely appreciated that distinct amygdala outputs and downstream circuits are recruited in response to different types of fear (Gross & Canteras, 2012). We have recently shown that fear responses to painful stimuli, predators, and aggressive members of the same species, depend on distinct neural circuits that involve the amygdala, medial hypothalamus, and periaqueductal gray (Silva et al., 2013). These data demonstrate that independent fear circuits exist to respond to different classes of threat and imply that pathological fear may come in different flavours and be amenable to selective therapeutic treatment. Current work in the lab combines molecular genetic, electrophysiological, and genetically encoded neural manipulation tools (figure 1) with behavioural methods in mice to understand how amygdala, hypothalamic, and brainstem circuits support and adapt fear responses to diverse threats.

Developmental programming of brain wiring by microglia: Microglia are non-neuronal cells of the hematopoetic lineage that infiltrate the brain during development and are thought to play a role in brain surveillance. Recent studies from our group and others have shown that microglia are particularly abundant during the period of postnatal brain development when synapses are formed (figure 2) and that they play a key role in the elimination of synapses during this period, a phenomenon called ‘synaptic pruning’ (Paolicelli et al., 2011). Mice with deficient synaptic pruning show weak functional brain connectivity, poor social behaviour, and increased repetitive behaviour – all hallmarks of autism – suggesting that some features of this neurodevelopmental disorder may depend on a deficit in synaptic pruning (Yang, Paolicelli et al., 2014). We are currently using a variety of tools to identify the ‘eat me’ and ‘spare me’ signals that regulate pruning and understand the mechanisms by which synapse elimination remodels neural circuits during development.

Future projects and goals
Together these approaches are aimed at discovering the neural circuits and molecular mechanisms that support individual differences in behavioural traits in health and disease. A better understanding of the molecular signals that influence the formation and remodelling of these circuits will allow us to form specific hypotheses about how human behaviour is determined and lead to improved diagnostic and therapeutic tools for mental illness.
Molecular physiology of somatosensation

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Previous and current research

Somatosensation is the process by which we sense touch and pain. It is dependent upon specialised sensory neurons which extend from the skin to the spinal cord and are tuned to detect mechanical, thermal and chemical stimuli. Surprisingly, the mechanisms that transduce these forces into electrical signals at the peripheral endings of sensory neurons are not well understood. Our research focuses on identifying and characterising these transduction components and exploring how they are altered during chronic pain states.

We use a combination of molecular, imaging and electrophysiological techniques to examine functional properties of sensory neurons at their peripheral and central terminals. At the molecular level, we are interested in mechanisms of touch sensitivity of sensory neurons. Normal mechanical sensitivity is dependent upon a complex of proteins that are localised at the peripheral endings of sensory neurons. Evidence supports a central role for the cytoskeleton in regulating the composition and function of this complex. Using cellular, electrophysiological and molecular imaging techniques we are characterising the contribution of the cytoskeleton, in particular microtubules to mechanotransduction.

Another focus of the group is to understand the biophysical properties of ion channels involved in sensory transduction. Much of our work has concentrated on the ion channel TRPA1, a member of the Transient Receptor Potential (TRP) family of channels. In mammals, TRPA1 is expressed by nociceptors and plays a key role in detecting noxious chemicals. We demonstrated that intracellular calcium ions directly activate TRPA1 via an EF-hand domain in the N-terminus of the protein and that calcium is essential for normal activation of the channel by noxious chemicals. We are now interested in how TRPA channels have evolved to sense diverse stimuli across different phyla – for example, in snakes and insects TRPA1 orthologues are activated by warm temperatures. Using a combination of molecular and electrophysiological techniques we have mapped the regions in Drosophila TRPA1 that are responsible for sensing temperature and described how single TRPA1 channels are activated by heat.

Future projects and goals

- Identification of novel genes involved in touch and pain.
- Mutagenesis of transduction channels and associated proteins to determine their mechanism of action.
- Tissue-specific and conditional mutagenesis of sensory-related genes in defined subpopulations of sensory neurons.
- Development of new techniques to measure functional properties of sensory neurons at their terminals.

The major focus of the laboratory is to correlate cellular studies on somatosensation with observations made at the physiological level. To this end we are developing genetic approaches that, combined with electrophysiological and molecular imaging techniques, will enable us to characterise sensory neurons in situ. A better understanding of sensory neuron function may ultimately lead to improved therapies for the treatment of chronic pain.

Selected references

The Jechlinger group uses a 3D culture system of primary mouse mammary epithelial cells to study cancer-initiating oncogenes.

Selected references

Previous and current research
Extensive evidence now supports the concept of oncogene addiction (the dependence of tumour cells on their initiating lesion for survival). In patients and mouse models interference with the activity of cancer-initiating oncogenes can result in tumour regression. However, novel therapies that target the products of mutant alleles in human cancers are only partly successful, since maintenance of remission requires long-term treatment and relapse often occurs in the presence of therapeutic agents. Hence, a better understanding of drug resistance and tumour recurrence is needed for the design of more successful anti-cancer strategies.

Transgenic mice carrying regulatable transgenes represent tractable systems for studying the mechanisms of oncogene dependence, the response and resistance to targeted drugs and tumour recurrence. In a complementary approach, we have developed a 3D culture system of primary mouse mammary epithelial cells to study detailed responses to the induction and de-induction of oncogenes (mimicking treatment with an ideally targeted drug). This 3D system produced phenotypic changes similar to those observed in the mammary glands of the transgenic mice from which the cultures were derived. In addition, this new approach identified and isolated cells that had survived oncogene withdrawal, exhibited characteristics of mammary gland progenitors and could efficiently re-populate the mammary fat pads of immunodeficient mice. The successful isolation of a pure population of surviving cells after oncogene withdrawal will allow us to characterise these residual ‘dormant’ tumour cells in detail.

Future projects and goals
- Determine at which point during tumourigenesis cells acquire the ability to survive oncogene withdrawal.
- Identify the molecular properties that distinguish surviving-residual cells, from naïve cells.
- Interfere with the mechanisms important for survival of residual ‘dormant’ cells.

(A) One cell in anaphase divides with the sets of chromosomes perpendicular to the apical surface, while keeping ZO1 localised to the luminal membrane.
(B) Confocal microscopy (5mm projection through the middle) shows epithelial cell polarity. E-Cadherin, (adherens junctions, lateral); ZO1, (tight junctions, apical); Integrin a6 (basolateral).
(C) Doxycycline exposure causes loss of epithelial polarity and filling of the lumen; Removal of doxycycline results in survival of a re-polarised cell layer, that acquires the ability to exclude Hoechst 33342. Left panels: Confocal microscopy (5mm projection through the middle) shows Dapi, GM130 (apical), ZO1 (tight junction), Integrin a6 (basal) at indicated times. Middle panels: Bright-field pictures show: (top) small, hollow acini; (middle) filled, irregular shaped spheres (bottom) hollow, irregular shaped spheres that show debris of internal cells. Right panels: Exclusion of Hoechst 33342 (1 hour incubation) at indicated times.
The haemogenic endothelium: a key stage in the generation of the first blood cells

Previous and current research

The continuous generation of blood cells throughout life relies on the existence of haematopoietic stem cells (HSC) generated during embryogenesis. They have the ability to self-renew and to generate all types of blood cells. Any pathology affecting these cells could lead to the development of serious diseases such as leukaemia and anaemia. That is why understanding how HSC and haematopoietic progenitors are produced during embryonic life is so important.

The origin of blood cells has been the subject of an intense scientific debate during the last decade. It has been proposed that during embryonic development, haematopoietic cells arise from a mesodermal progenitor with smooth muscle, endothelial, and haematopoietic potential called the haemangioblast. However, a conflicting theory instead associates the first haematopoietic cells with a phenotypically differentiated endothelial cell with haematopoietic potential (i.e. a haemogenic endothelium).

To investigate the cellular origin of blood cells, we used a model of early haematopoiesis based on the differentiation potential of the mouse embryonic stem cells (ESC) in vitro. These cells are derived from the inner cell mass of the blastocyst – an early-stage mouse embryo – and have the capacity to generate any cell types. Using this system coupled with time-lapse microscopy, clonogenic assays and flow cytometry analysis, we have demonstrated that the haemangioblast generates haematopoietic progenitors through the formation of a haemogenic endothelium stage, providing the first direct link between these two precursor populations. Together our results merge the two a priori conflicting theories on the origin of haematopoietic development into a single linear developmental process. This finding allowed us to identify the haemogenic endothelium as the immediate precursor of blood cells (figures 1 and 2).

Future projects and goals

Recently, the generation of the ESC-like induced pluripotent stem cells (iPSC) from fully differentiated cell types, such as skin fibroblast, provided a major breakthrough in the field of regenerative medicine. Indeed iPSC offer a great opportunity to implement replacement therapy by bypassing the use of human embryos to generate ESC, therefore decreasing ethical concerns. However, important work has to be done to differentiate efficiently iPSC or ESC toward specific cell types including blood cell progenitors such as HSC.

Consequently, in order to better understand the development of the haematopoietic system, the focus of our research is to unravel the mechanisms underlying the generation of haemogenic endothelium from its precursor, the haemangioblast, and its subsequent commitment to haematopoiesis. Combining genomics, time-lapse microscopy, and loss and gain of function experiments in vitro and in vivo, we plan to identify and study the genes responsible for the generation of the first blood progenitors during embryonic life. Our research will bring a further understanding of the mechanisms of cell fate decisions leading to the production of the first haematopoietic cells and enable the development of new strategies to improve methods of blood cell generation from ESC or iPSC for regenerative medicine.

Selected references


The Lancrin group studies the haematopoietic system and looks to develop strategies to improve methods for generating blood cells from stem cells.
Non-coding RNA function and RNA modification in germ/stem cell biology

The O’Carroll group studies mouse blood cell formation, embryology and germ cell development using state-of-the-art genetic strategies and high-throughput sequencing approaches.

Selected references

Previous and current research
The goal of my laboratory is to explore the contribution of non-coding RNA as well as RNA modification pathways to tissue development and homeostasis. We have garnered much experimental expertise in both haematopoiesis and the germline. Within the laboratory, an emphasis (non-exclusive) is now placed on studying the immortal lineage. The integrity of the genome transmitted to the next generation intrinsically relies on cells of the germline. Processes that ensure germ cell development, genomic stability, and reproductive lifespan are essential for the long-term success of a species. We tackle fundamental questions regarding the mammalian male germline and heredity from an RNA perspective. Specifically, our research explores the contribution of non-coding RNA (miRNA, piRNA and lncRNA) as well as RNA modification pathways within germ cell development as well as testicular homeostasis and regeneration.

The precise identity of the spermatogonial stem cell (SSC) in vivo that supports spermatogenesis throughout life remains unknown. Capitalising on the fact that SSC maintenance is dependent upon several RNA-binding proteins, we hope that investigation into these pathways may reveal the identity of this stem cell in vivo. The maturation of RNA sequencing techniques, in combination with refined genetic approaches, now renders the identification and functional evaluation of non-coding RNAs and RNA modifications in vivo within the realm of experimental feasibility. Our research objectives focus on the contribution of these emerging pathways on the underlying circuitry of self-renewal that underpins the SSC, as well as the coordination of the various cellular/differentiation processes of spermatogenesis.

The acquisition of both pluripotency and totipotency is associated with the deregulation of transposable elements – our goal is understand the mechanisms by which the germ cells manage this formidable threat to gametes, and thus transgenerational genome stability. Specifically, transposon silencing in the germline by the Piwi-interacting RNA (piRNA) pathway as well as epigenetic mechanisms will be extended upon from our previous findings.

Future projects and goals
- The identification and characterisation of the spermatogonial stem cell populations.
- Dissection of the pathways required for spermatogonial stem cell self-renewal and testicular regeneration.
- Post-transcriptional RNA modification in germ cell and hematopoietic development.
- Long non-coding RNA function in spermatogenesis.
- Establishment and maintenance of epigenetic transposon silencing in the male germ line.
Mitotic chromosomal instability and oncogene dependence

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Previous and current research
Chromosomal instability (CIN), the inability to correctly segregate sister chromatids during mitosis, is a hallmark of cancer cells. Overexpression of the mitotic checkpoint protein Mad2, commonly found in human tumours, leads to CIN and the development of aneuploid tumours in mouse models. Moreover, CIN can facilitate escape from oncogene addiction (the dependence of tumour cells on their initiating lesion for survival) and may be responsible for tumour relapse after targeted therapies. Very little is known about the mechanism of how and when CIN promotes tumour relapse. Our lab focuses on understanding the molecular mechanisms that lead to CIN and the consequences it may have in tumour initiation, suppression and relapse. We hope that the genes or proteins identified could be targeted therapeutically. We will use a combination of mouse genetics and highly innovative 3D in vitro culture systems.

Future projects and goals
- To study the dependence of tumour cells on the mitotic checkpoint in vivo and evaluate the potential for therapeutic interference with mitotic checkpoint genes.
- To study the effects of tumour regression and recurrence in chromosomally unstable tissues.
- To analyse the dual role of chromosome instability (tumour initiating and tumour suppressive) dependent on levels of aneuploidy, tissue type and molecular nature of the cooperating lesion(s).

Selected references

Primary mammary cells from TetO-Mad2/TetO-Myc/MMTV-rtTA mice grown in 3D culture. Left panel shows never induced cells that are beginning to form a polarized acinus. Right panel: loss of epithelial cell polarity in an acinus grown from tritransgenic cells and exposed to doxycycline for 36h. White arrows show abundant mitotic cells and lagging chromosomes after Mad2 overexpression.
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