**Welcome**

Emma Faulkner has joined Drs Steve Thomas and Natalie Poulter’s team as a Research Fellow at the University of Birmingham on a BHF funded project with COMPARE researchers, Steve Watson, Jeremey Pike, Rob Neely and Iain Styles. Emma will be working on a project applying expansion and super resolution microscopy to look at the subcellular organisation of individual platelets within aggregates under normal and perturbed situations. The aim of this project is to provide detailed analysis of heterogeneity of platelet organisation within thrombi to better understand healthy and diseased situations to potentially identify how we can target this in disease.

**Congratulations**

Josh Bourne, University of Birmingham, won the Best Oral Presentation at the UK Cell Adhesion Society meeting; he also gave a talk at the 1st Platelet Society Meeting in Cambridge which had wonderful feedback. Josh is a PhD student working with Dr Julie Raye’s, and has just published his first review in RPTH on ‘The dual role of platelet-innate immune cell interactions in thrombo-inflammation’ https://onlinelibrary.wiley.com/doi/full/10.1002/rth2.12266

Joanne Clarke, COMPARE PhD student, with Steve Watson and Steve Hill, based at the University of Birmingham, has had her STORM microscopy image featured on the front cover of the July issue of Thrombosis and Haemostasis. The image shows a cluster plot generated by a DBSCAN algorithm showing the localisation of GPVI receptor clustering in platelets spread on collagen obtained by STORM microscopy. This image is included in her paper Adenosine and Forskolin Inhibit Platelet Aggregation by Collagen but not the Proximal Signalling Events.

birmingham-nottingham.ac.uk/compare

Follow us on Twitter
@compare_uobuon
@uobuonpartners

If you have any items for the next newsletter please send to:
compare@birmingham-nottingham.ac.uk
Applications for the 2020 summer placements are now open. Team Science funding will be used to provide up to 4 summer placements in 2020. The placements will be supervised by a COMPARE Post Doc/Research Fellow for a maximum of 6 weeks, providing up to £200/week stipend for the student and £300 consumables. Applications involving interdisciplinary projects or undergraduate students from the physical sciences are encouraged. If you have a current project that would benefit from a summer student and would like to develop your supervisory skills, please complete the application form which has been circulated via email, in conjunction with your COMPARE PI and student and return to compare@birmingham-nottingham.ac.uk.

Closing date for applications is Friday 28th February 2020. Applicants will be informed of the result by the mid-March and for successful applicants final reports on the project will need to be submitted by Monday 7th September 2020.

Applications are invited for the Team Science Grants. Applications for collaborative projects will be considered (can be between Birmingham and Nottingham and/or external collaborations). Funding can support collaborative projects or visits to a partner laboratory with a maximum application value of £2,000, to support cross University research projects or visits to learn new techniques and skills that are within the scope of COMPARE. The money can be used to cover travel, consumables and equipment costs occurred during the visit. A post-award report and submission of an abstract and poster at the COMPARE Annual Research Symposium 2020, will be required from all successfully funded applicants.

Closing date 5pm on Friday 31st January 2020. Applications will be reviewed by the Team Science Committee and Team Science leads and results will be provided by mid-February.

Please email compare@birmingham-nottingham.ac.uk to request an application form (if not received via email) or submit completed forms.

The cardiac research dream team visits Birmingham

Alexander (Sasha) Kondrashov and Nguyen (Kate) T N Vo from Chris Denning’s group at the University of Nottingham are experts in genome editing and phenotyping of pluripotent stem cell-derived cardiomyocytes, respectively. Kate and Sasha visited the University of Birmingham and presented their work as part of the COMPARE Team Science seminar series in July and October. During their visit, they exchanged science with researchers from both within and outside of COMPARE, from the Institute of Cardiovascular Sciences and Cells and Molecules Research Theme in the School of Biosciences. Thank you, Kate and Sasha!
Below is a summary of the microscopes in COMPARE. For further information including training and costings please contact Dee Kavanagh (D.M.Kavanagh@bham.ac.uk) or Joelle Goulding (Joelle.Goulding@nottingham.ac.uk).

Please note the equipment is not maintained to GCP in the Laboratory standard and therefore cannot be used in the analysis of clinical trial (CTIMP or non-CTIMP) samples which contribute to the primary, secondary or exploratory outcomes of such trials.

**COMPARE Microscopes**

**The UltramicroscopeII**
The UltramicroscopeII is a fluorescent microscope used for 3D imaging of large samples such as whole mouse organs and small tissue samples (essentially 3-D histology). The microscope is suitable for imaging multi-colour fluorescently labelled cell types and vasculature e.g. renal, cardiac and lymphatic, in samples that have been made optically clear using established protocols.

**Marianas diSPIM microscope**
The Marianas captures two alternating views of a sample. The microscope is suitable for imaging multi-colour fluorescent cellular structures in spheroids and organoids. These views are then merged together to give an even 3D image without any distortions. The Marianas microscope can be used for long time lapse imaging of living samples (hours) such as formation of platelets from megakaryocytes.

**Lattice Light Sheet**
The Lattice microscope is the leading fluorescent microscope for imaging live cellular processes in single cells (hours). Using this microscope you will be able to gain precise information on protein and organelle dynamics, with the highest and most accurate time resolution currently available in a fluorescent imaging system.

**Structured Illumination Microscope (SIM)**
SIM allows the observation of live and fixed fluorescent samples at a resolution (100x100nm) greater than a standard fluorescent microscope (300x300nm). Imaging at this resolution can reveal structural organisation that can be otherwise masked.

**Single Molecule Stochastic Optimal Reconstruction Microscope (STORM)**
The single molecule microscope is used to identify the position of single fluorescent protein and lipid molecules. The molecule positions can then be used to extract information on protein organisation at a higher level using mathematical algorithms.

**Fast multicolour single-molecule microscope (4-camera TIRF)**
This custom 4-camera TIRF is capable of simultaneously imaging up to four different molecules (labelled with different colours) with single-molecule sensitivity, permitting analysis of individual molecules as they diffuse and interact in living cells with a spatial resolution of approximately 20 nm and a temporal resolution of 20 ms. This provides quantitative information on complex populations of molecules, including their nanoscale arrangement, diffusion and interactions.

**Ultrafast FRET/calcium imaging microscope**
This custom microscope is capable of performing ultrafast FRET and calcium imaging measurements in living cells with up to sub-millisecond temporal resolution. It is equipped with a fast perfusion system for rapid stimulation, which allows to exchange the solution received by the perfused cell in a few milliseconds. Combined with a series of FRET sensors and indicators, it allows monitoring the kinetics of cell signalling events such as receptor activation or the production of second messengers such as calcium or cyclic AMP in living cells with high spatiotemporal resolution.

**Computational resources and specialist software (Birmingham)**
There are six high-spec dedicated analysis workstations, as well as preferential access to the University’s central high performance computing (HPC) resources. The workstations and HPC resources are free to use.

We have in-house expertise in a range of leading open-source software and can provide consultation and advice on image analysis applications. There are two commercial software packages; Arivis Vision4D and SVI Huygens. Vision4D enables user friendly analysis of large volumetric data and facilities impressive and fast 3D volume rendering for visualisation and inspection of data. Huygens is specialist deconvolution software which improves image quality by removing out of focus light from fluorescent microscopy datasets. Commercial software is chargeable either by annual fee, or at an hourly rate.
Bioluminescence – Olympus LV200
The Olympus LV200 is capable of imaging bioluminescence within an environmentally controlled setting. Bioluminescence is the chemically induced release of light which does not need laser or LED excitation light (as with fluorescence) and as such does not suffer from the side-effects of photo-toxicity and photo-bleaching. The microscope can be used to image bioluminescence in cell monolayers, and for proximity assays using bioluminescence as donor (Bioluminescence Resonance Energy Transfer, BRET).

Ptychography – PhaseFocus Livecyte
The PhaseFocus Livecyte can measure the change in the phase of transmitted light as it passes through a sample and can reconstruct an image which contains information on cell volume, shape, size etc. Cell movement can be tracked and cell splitting and lineage monitored. No lasers are used and as such photosensitive cells can be imaged for several days, within multi-well plates. The microscope has temperature, humidity and gas control which allows automated imaging long term within the environmentally controlled casing. Potential assays include cell migration, mobility, proliferation, wound healing and angiogenesis (formation of blood vessels).

Confocal Microscopy – Zeiss 710
This microscope allows fluorescence confocal microscopy. By the addition of a pinhole within the laser beam path, out-of-focus light from your sample can be discarded allowing the imaging of thin confocal slices and if desired 3D reconstruction of your sample. This is perfect for localisation and quantification of your fluorescent target and can be coupled with more advanced techniques such as FRET (to look at protein-protein interactions) and FRAP (recovery after photo-bleaching to study diffusion and immobile fractions).

Fluorescence Correlation Spectroscopy (FCS) – Zeiss 880 & Zeiss 510NLO
These two confocal microscopes are equipped with single photon sensitive detectors which allow fluorescence fluctuation spectroscopy techniques such as FCS. FCS allows quantification of particle number, diffusion speed and brightness within a very small volume (0.25 femtolitre) and can be applied to samples both in solution or expressed within live cells. Within solution, important pharmacological parameters such as drug binding affinity can be determined. Within live cells the organisation of membrane proteins can be studied and in addition the effects of drug treatment examined. This technique is ideal for low expression systems such as those observed physiologically.

Total Internal Reflection Fluorescence (TIRF) – Nikon Ti2
TIRF microscopy produces a very thin field of illumination above the sample chamber (100-200nm) which allows imaging of the lower membrane of a cell or a layer of immobilised protein. Time-lapse capture can be amenable for tracking the movement of membrane proteins and methods can be optimised to enable the study of single molecule dynamics and interactions.

High-content plate readers – MD Image Express Ultra and Micro
High-content plate readers enable automated, multi-parameter image capture across various plate formats. Both live and fixed cells can be imaged for fluorescence in 96- or 384-well format. The microscope moves the low-throughput quantification and localisation microscopy into a high-throughput scheme. Analysis packages are available to quantify receptor internalisation, cellular structures and fluorescence intensity/per cell or region. These can subsequently be used to determine important parameters used in screening such as drug binding affinity and potency.