### WEDNESDAY, SEPTEMBER 19TH, 2018

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<tr>
<th>Time</th>
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<th>Event</th>
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</thead>
<tbody>
<tr>
<td>9:00-12:00</td>
<td>Institute of Immunology</td>
<td>Presentation and Demonstration of Flow Cytometers</td>
</tr>
<tr>
<td>10:00-12:00</td>
<td>Center for Applied Research</td>
<td>Raman Microspectroscopy – a new Tool for Cultivation-free Pathogen Identification</td>
</tr>
<tr>
<td>12:00-13:00</td>
<td>Foyer</td>
<td>Welcome &amp; Registration</td>
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<tr>
<td>13:00-14:30</td>
<td>Lecture Hall 3</td>
<td>Nanotechnology Session</td>
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<td>19:00-22:00</td>
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<tr>
<td>9:00-10:30</td>
<td>Lecture Hall 3</td>
<td>NEW! Core Facility Session: Reproducibility in Flow Cytometry</td>
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<tr>
<td>10:30-11:00</td>
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<tr>
<td>11:00-12:30</td>
<td>Lecture Hall 3</td>
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<td>17:00-18:00</td>
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<tr>
<td>19:30-23:00</td>
<td>ZEISS PLANETARIUM</td>
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### FRIDAY, SEPTEMBER 21ST, 2018

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<th>Time</th>
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<th>Event</th>
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<tbody>
<tr>
<td>9:30-11:00</td>
<td>Lecture Hall 3</td>
<td>Microbiology Session</td>
</tr>
<tr>
<td>11:00-11:30</td>
<td>Foyer</td>
<td>COFFEE BREAK</td>
</tr>
<tr>
<td>11:30-13:00</td>
<td>Lecture Hall 3</td>
<td>Microscopy Session</td>
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<tr>
<td>13:00-14:00</td>
<td>Foyer</td>
<td>FAREWELL &amp; SNACKS</td>
</tr>
</tbody>
</table>

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**Life in Focus**

Dear Cytometrists,

welcome to the annual DGfZ meeting 2018!

The meeting will cover various aspects of cytometry, in scientific sessions such as on nanotechnology, microscopy or microbiological applications, in a newly established core facility session, on posters, and in the industrial exhibition [Thanks to the sponsors for making this meeting possible!]. Technology innovations will be presented in the Cutting Edge session as well as in the company product slam. This years European Guest Session will introduce our Hungarian partner organization, The Cell Analysis Section of the Hungarian Biophysical Society.

Jena has a unique tradition of fruitful interactions between technology and science aimed at bringing "Life in Focus", as demonstrated in the work of people like Carl Zeiss and Ernst Abbe, Matthias Schleiden and Hans Knöll. This special spirit reflects the motivation of our society, and shall therefore also guide our meeting.

At this point I would like to thank all funding bodies and industrial partners for their generous support without which this meeting would not have been possible. I am looking forward to three days of scientific exchange in fascinating presentations and interesting discussions with colleagues, to meet old and to make new friends – let’s enjoy the various aspects of the exciting world of cytometry here in Jena!

Wolfgang Fritzsche – President DGfZ
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General Information

VENUE ADDRESS
Friedrich Schiller University // Carl-Zeiss-Straße 3 // 07743 Jena // Germany

CONFERENCE CHAIR
(APL) Prof. Wolfgang Fritzsche // wolfgang.fritzsche@leibniz-ipht.de

CONFERENCE ORGANIZERS
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Daniel Siegesmund // daniel.siegesmund@leibniz-ipht.de
Felicitas Deckert // felicitas.deckert@leibniz-ipht.de

OPENING HOURS
Wednesday, September 19 // 12:00 – 22:30
Thursday, September 20 // 8:30 – 19:30
Friday, September 21 // 9:00 – 14:00

WIFI-PASSWORD
Information and assistance provided at the registration desk
## Program

### WEDNESDAY, 19 / SEP / 2018

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<th>Time</th>
<th>Session</th>
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</thead>
<tbody>
<tr>
<td>9:00-12:00</td>
<td>Institute of Immunology // Tutorial &quot;Presentation and Demonstration of Flow Cytometers&quot; // Nico Andreas // Institute of Immunology / Jena, Germany</td>
</tr>
<tr>
<td>10:00-12:00</td>
<td>Center for Applied Research // Tutorial &quot;Raman Microscopy – a New Tool for Cultivation-free Pathogen Identification&quot; // Petra Rösch // Institute for Physical Chemistry / Jena, Germany</td>
</tr>
<tr>
<td>12:00-13:00</td>
<td>FOYER // WELCOME &amp; REGISTRATION</td>
</tr>
<tr>
<td>13:00-14:30</td>
<td>Lecture Hall 3 // Nanotechnology Session // Chair: Wolfgang Fritzsche, Chair: Ulrike Taylor</td>
</tr>
<tr>
<td>13:00-13:15</td>
<td>Opening // Wolfgang Fritzsche // Leibniz Institute of Photonic Technology / Germany</td>
</tr>
<tr>
<td>13:15-14:00</td>
<td>Molecular Imaging of Cancer Cells with Ultra-small 5 nm Gold Nanoparticles // Konstantin Sokolov // M. D. Anderson Cancer Center / United States of America</td>
</tr>
<tr>
<td>14:00-14:30</td>
<td>How to Re-activate Antibiotic Efficacy in Biofilm-associated Lung Infections by Nanoparticles // Julia Ernst // Friedrich Schiller University Jena / Germany</td>
</tr>
<tr>
<td>14:30-15:30</td>
<td>Lecture Hall 3 // Product Slam // Chair: Frank Schildberg, Chair: Elmar Endl</td>
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<tr>
<td>15:30-16:30</td>
<td>FOYER // COFFEE BREAK &amp; INDUSTRY SESSION</td>
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<tr>
<td>16:30-18:00</td>
<td>Lecture Hall 3 // Cutting Edge Session // Chair: Hyun-Dong Chang</td>
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<tr>
<td>16:30-17:00</td>
<td>Light Sheet for the Masses // Emmanuel Reynaud // UCD Centre for Biomedical Engineering / Ireland</td>
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<tr>
<td>17:00-17:20</td>
<td>Luminescence Lifetime Encoding in Flow Cytometry – First Steps and Assessment // Daniel Kage // Bundesanstalt für Materialforschung und -prüfung (BAM)</td>
</tr>
<tr>
<td>17:20-17:40</td>
<td>Advanced Imaging Flow Cytometry // Andreas Kleiber // Leibniz Institute of Photonic Technology / Germany</td>
</tr>
<tr>
<td>17:40-18:00</td>
<td>Opto Biolabs – Combining Optogenetics with Flow Cytometry // Kathrin Brenker^2 // 1: Opto Biolabs, Germany, 2: Albert-Ludwigs-Universität Freiburg</td>
</tr>
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### 18:00-19:00 | Lecture Hall 3 // Keynote // Sepsis – Changing the Paradigm // Michael Bauer // Jena University Hospital, Germany // Chair: Thomas Kamradt |

### 19:00-22:00 | FOYER // WELCOME RECEPTION AT INDUSTRY EXHIBITION |

### 20:00-21:00 | Seminar Room 131 // Tutorial "Scientific Publication with Special Aspects for Cytometry Part A" // Attila Tärnok // University of Leipzig / Germany |

### 20:00-22:00 | Seminar Room 113 // Core Facility Networking Event // Chair: Desiree Kunkel, Chair: Steffen Schmitt |

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<tr>
<th>Time</th>
<th>Session</th>
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<tbody>
<tr>
<td>9:00-10:30</td>
<td>Lecture Hall 3 // NEW! Core Facility Session: Reproducibility in Flow Cytometry // Chair: Desiree Kunkel, Chair: Frank Schildberg</td>
</tr>
<tr>
<td>9:00-9:30</td>
<td>Validation and Standardization of Flow Cytometry Based Immune Monitoring in Clinical Trials // Mathias Streitz // Institute for Medical Immunology, Charité / Germany</td>
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<tr>
<td>9:30-10:00</td>
<td>A Novel Method for Flow Cytometer Characterization by Determination of Detector Background, Signal-to-noise, and Dynamic Range // Claudia Giesecke // Deutsches Rheuma-Forschungszentrum Berlin / Germany</td>
</tr>
<tr>
<td>10:00-10:15</td>
<td>Minimal Requirements for the Presentation of Flow Cytometry Data // Steffen Schmitt // German Cancer Research Center (DKFZ) / Germany</td>
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<tr>
<td>10:15-10:30</td>
<td>cytometry.de – the Communication Platform of the DGfZ // Elmar Endl // University Bonn / Germany</td>
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### 10:30-11:00 | FOYER // COFFEE BREAK & POSTER SESSION |

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\(^{1}\) Opto Biolabs: Combining Optogenetics with Flow Cytometry

\(^{2}\) Kathrin Brenker: Co-author of the paper "Opto Biolabs – Combining Optogenetics with Flow Cytometry"
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<tr>
<td>11:00-11:30</td>
<td>Immunophenotyping of Treatment Naive Patients with Systemic Autoimmune Diseases by Single Cell Mass Cytometry // Gábor Szebeni // Hungarian Academy of Sciences / Hungary</td>
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<tr>
<td>11:10-12:00</td>
<td>Targeting the HER2 Oncoprotein: Lessons Learned from Quantitative Cytometry // György Veréb // University of Debrecen / Hungary</td>
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<tr>
<td>12:00-12:30</td>
<td>The Effect of Microenvironmental Factors on the Development of Myeloma Cells // Gábor Barna // Semmelweis University, Hungary / Hungary</td>
</tr>
<tr>
<td>12:30-13:00</td>
<td>Lecture Hall 3 // Poster Pitch &amp; Poster Session</td>
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<tr>
<td>13:30-14:30</td>
<td>FOYER // LUNCH &amp; POSTER SESSION</td>
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<tr>
<td>14:30-16:00</td>
<td>Lecture Hall 3 // Klaus-Goerttler-Session // Chair: Julia Reinhardt, Chair: Wolfgang Fritzsche</td>
</tr>
<tr>
<td>14:30-15:00</td>
<td>Deformability Cytometry and 1D Fluorescence Imaging in Real-time // Philipp Rosendahl // TU Dresden / Germany</td>
</tr>
<tr>
<td>15:00-15:20</td>
<td>NKT Cells Promote Alternative Cross-priming in two Distinct Phases // Christoph Heuser // Technical University of Munich / Germany</td>
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<tr>
<td>15:20-15:40</td>
<td>Alternative Splicing: An Alternative Level of NLRP3 Inflammasome Regulation // Florian Hoss // University of Bonn / Germany</td>
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<tr>
<td>15:40-16:00</td>
<td>Metabolic Regulation of ILC Mediated Barrier Protection and Pathology // Christoph Wilhelm // University Hospital Bonn / Germany</td>
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<tr>
<td>16:00-17:00</td>
<td>FOYER // Poster Session</td>
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<tr>
<td>17:00-18:00</td>
<td>Lecture Hall 3 // Guest Lecture – Early Precursors of Fluorescence Cytometry: Foundation of Cell Theory and Fluorescence Microscopy // Timo Mappes // Deutsches Optisches Museum / Germany // Chair: Wolfgang Fritzsche</td>
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<tr>
<td>18:00-19:30</td>
<td>Lecture Hall 3 // Members Assembly</td>
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<td>19:30-22:30</td>
<td>Zeiss Planetarium // Social Event &amp; Conference Dinner</td>
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<th>Time</th>
<th>Session</th>
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<tbody>
<tr>
<td>9:30-10:00</td>
<td>Molecular Diagnostic in the Age of Multiresistant Gram-negative Bacteria // Oliwia Makarewicz // University Hospital Jena / Germany</td>
</tr>
<tr>
<td>10:00-10:20</td>
<td>Raman Spectroscopic Cytometry for the Detection and Characterization of Bacterial Infection // Astrid Tannert // Leibniz IPHT / Germany</td>
</tr>
<tr>
<td>10:20-10:40</td>
<td>Who am I and if Yes, How many? A new Way of Phytoplankton Species Identification by Combination of Image Cytometry and Deep Learning // Susanne Dunker // Helmholtz-Centre for Environmental Research – UFZ / Germany</td>
</tr>
<tr>
<td>10:40-11:00</td>
<td>Microfluidic System for Single-cell Analysis in Picoliter-sized Batch Bioreactors // Eugen Kaganovitch // Forschungszentrum Jülich GmbH / Germany</td>
</tr>
<tr>
<td>11:00-11:30</td>
<td>FOYER // COFFEE BREAK</td>
</tr>
<tr>
<td>11:30-12:00</td>
<td>Lecture Hall 3 // Microscopy Session // Chair: Anja Hauser, Chair: Raluca Niesner</td>
</tr>
<tr>
<td>11:30-12:00</td>
<td>histoCAT: Analysis of Cell Phenotypes and Interactions in Multiplex Image Cytometry Data // Denis Schapiro // University of Zurich / Switzerland</td>
</tr>
<tr>
<td>12:00-12:30</td>
<td>Super-resolution Microscopy: Challenges and Potentials in Biomedical Research // Christian Eggeling // Leibniz Institute of Photonic Technology / Germany</td>
</tr>
<tr>
<td>12:30-12:45</td>
<td>Histo Cytometry using Multi Epitope Ligand Cartography (MELC) // Ralf Köhler // Immunodynamics, German Rheumatism Research Center, Berlin / Germany</td>
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<tr>
<td>12:45-13:00</td>
<td>Lecture Hall 3 // uC2 – An Open-Source Optical Toolbox for Multi-Modal Imaging in the Incubator // Benedict Diederich // Leibniz IPHT / Germany</td>
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<tr>
<td>13:00-14:00</td>
<td>FOYER // FAREWELL &amp; SNACKS</td>
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Jena – City of Light

Jena is one of the most popular university and college cities in Central Germany and has an outstanding reputation as a high-tech center. It is the second largest city in Thuringia and attractively situated in the picturesque landscape of the Saale Valley. The city surprises visitors with gratifyingly short distances, an almost Mediterranean flair and a contemporary and open-minded atmosphere. Jena’s nickname “City of Light” stands for its many scientific institutions, world-renowned high-technology companies and young start-up businesses in the field of optics and photonics. Essentially, the name “City of Light” is inspired by Jena’s bright minds. Among researchers, Jena is one of the best-known places in Germany. One quarter of the 108,000 inhabitants are students and 4500 scientist work at the universities and research institutes. For many entrepreneurs, the city with one of the highest number of patent applications provides the perfect environment to realize their ideas.

Friedrich Schiller University – Thinking without Limits

The FSU Jena still maintains the sense of innovation which characterized its early period, and today it views itself as a forward-looking hub for new research fields and emerging generations of scientists. As a result, the University traditionally thinks and operates on an interdisciplinary and international basis. Over 18,000 students currently study at the University in Jena, and a large proportion of these students have an international background. Moreover, Jena’s character has been fundamentally shaped by the embedding of the University within the city’s historically rich and diverse cultural landscape, as well as the emergence of the region as a hotbed for innovative and economically successful high-tech companies. As an active initiator in the fields of science, culture, and business, the University is committed to promoting the development of the city and region as a whole.
The Society of Cytometry (Gesellschaft fuer Zytometrie, GZ) was founded in 1989 in Heidelberg (Germany) by the Foundation Council represented by Cess Cornelisse, Georg Feichter, Wolfgang Goehde, Klaus Goerttler, Holger Hoehn, Andreas Radbruch, Peter Schwarzmann, and Günter Valet. An association was born dedicated to provide an interdisciplinary platform for interested scientists basically in the field of flow and image cytometry. Founding members were scientists whose personal scientific development was and is still closely interlinked with the development of cytometric technologies in Europe.

**Council**

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**Dr. Frank Schmidt // University of Greifswald Functional Genomics**

**Dr. Gergely Toldy // Birmingham Women's Hospital, Birmingham, UK**
Excellent Plus Sponsors

Excellent Sponsors

Funded by

Floorplan Exhibition Hall
Tutorials

It is a good tradition of the Annual Conference of the German Society for Cytometry to offer preconference tutorials. The tutorials are a great opportunity to gain deeper insights and practical skills with a special focus on cytometry related content. Therefore the tutorials are recommended especially, but not solely for students. The tutorials are free of charge for registered participants of the conference.

Tutorial 1

9:00-12:00  Presentation and Demonstration of Flow Cytometers // Nico Andreas // University Hospital Jena / Institute of Immunology

Tutorial 2

10:00-12:00  Raman Microspectroscopy – a new Tool for Cultivation-free Pathogen Identification // Petra Rösch // Center for Applied Research Jena

Tutorial 3

20:00-21:00  Scientific Publication with Special Aspects for Cytometry Part A // Attila Tärnk // Seminar Room 131 / Friedrich Schiller University
Molecular Imaging of Cancer Cells with Ultra-small 5 nm Gold Nanoparticles.

This talk will present biologically-induced plasmonic effects in nanoscale spherical gold nanoparticles that can be exploited for multi-modal imaging including dark-field microscopy, reflectance, photoacoustic and two-photon luminescence imaging. Previously, we have demonstrated a strong spectral shift of > 100 nm and color change of 20 and 40 nm gold nanoparticles targeted to epidermal growth factor receptor (EGFR) after their receptor mediated uptake by cancer cells [1]. We showed that this effect is associated with plasmon resonance coupling of gold nanoparticles in cellular endosomal compartments [2] and it allows achieving an unprecedented sensitivity in detection of metastatic foci of ca. 30 cells at depth in vivo in animal models of head and neck cancer [3].

Currently, we are exploring multimodal nature of ultra-small 5 nm gold nanoparticles in detection of cancer cells. This size domain is comparable (or even smaller) to large biomolecules such as antibodies and it can offer a number of advantages including better intratumoral distribution of nanoparticles, improved delivery and pharmacokinetics. Furthermore, nanoparticles with sizes below than 10 nm can undergo efficient body clearance that can facilitate future clinical translation. Our results show that labeling of cancer cells with EGFR-targeted 5nm gold nanoparticles results in a very strong contrast in dark-field and photoacoustic imaging with near-infrared (NIR) excitation that is very similar to the much bigger 20-40 nm particles. Confocal and multi-photon microscopy studies of nanoparticles trafficking in live cells indicate that this strong signal is associated with nanoparticles’ coating degradation inside cells. Interestingly, molecular specific intracellular uptake of 5nm gold nanoparticles results in a very strong two-photon luminescence, which is characterized by broad emission and relatively narrow excitation spectral properties and very short lifetimes. The dual photoacoustic (due to absorption) and luminescence signal from nanoparticles’ labeled cells can be highly advantageous in combining a lower resolution in depth tissue imaging using photoacoustic with a high-resolution two-photon microscopy that we are exploring in a number of application ranging from cell tracking to sensitive detection of molecular targets and cancer cells in vivo.

References:
How to Re-activate Antibiotic Efficacy in Biofilm-associated Lung Infections by Nanoparticles

Bacteria, such as Pseudomonas aeruginosa and Burkholderia cepacia, are a major cause of chronic lung infections in cystic fibrosis (CF) patients. The ability of the bacteria to form biofilms and the presence of a thick and stagnant mucus in the airways of CF patients lead to antibiotic therapy failure and request innovative antibiotic delivery systems to improve the antibiotics' effectiveness in the CF environment [1]. Biodegradable nanoparticles (NP) as carriers for antimicrobials are promising to break through the mucus and biofilm barrier but are demanding in pulmonary delivery because of particle aggregation and exhalation [2]. In the present study, differently sized nanoparticles made of fluorescently labeled poly(lactic-co-glycolic acid) (PLGA) and poly(ethylene glycol)-grafted PLGA (PEG-PLGA) were compared and evaluated as lung applicable carriers to re-activate the efficacy of the antibiotic tobramycin (Tb) in biofilms of P. aeruginosa and B. cepacia.

NPs loaded with Tb were prepared by a double-emulsion evaporation method [3] and exhibited mean hydrodynamic diameters of 200 nm (Tb-NP200) and 900 nm (Tb-NP900), respectively, with narrow size distributions measured by dynamic light scattering (Zetasizer Nano ZS, Malvern Instruments, Herrenberg, Germany) and confirmed by scanning electron microscopy. Tb incorporation increased the zeta potentials to about -10 mV compared to drug-free particles of about 30 mV. The drug content was 1.7 µg Tb per mg NP analyzed by HPLC. To determine the in vitro mucus permeation of NPs, artificial mucus (AM) was filled in ThinCerts™ layered by fluorescently labeled NPs and placed in simulated lung fluid. By measuring the fluorescence intensity in the upper compartment over time, a size and surface potential dependent permeation was found as Tb-NP200 permeated faster compared to Tb-NP900 and Tb-loaded NPs faster than the corresponding drug-free NPs. In vitro biocompatibility studies showed that no cytotoxicity in A-549 cells were observed for all particle types up to a concentration of 1 mg/mL and 24 h incubation time. For biofilm experiments, bacteria were cultivated in AM-containing chamber slides to allow the formation of a biofilm close to those of CF patients or in a microfluidic device to imitate the physiological shear flow in the body. The excellent penetration abilities of Tb-loaded particles through AM and biofilms and the remarkable antimicrobial efficacy in comparison to the free drug was confirmed by confocal laser scanning microscopy of live/dead stained biofilms.

Nebulization experiments were performed by air-jet (PARI boy, LC SPRINT, PARI, Germany) or vibrating-mesh technology (IH 50, Beurer, Germany). First, the re-suspension of the NPs in therapeutically relevant inhalation solution (0.9% NaCl, 3% NaCl and 7% NaCl) was tested and resulted in slightly increased hydrodynamic diameters compared to the NPs in deionized water. Physiological NaCl was identified as the most suitable inhalation solution due to minor alteration of the particle characteristics and was therefore chosen for the further nebulization experiments. Laser diffraction (HELOS, Sympatec, Germany) was used to analyze the aerodynamic characteristics of nebulized formulations over 10 min to determine the mean median aerosol diameter (MMAD), the geometric standard deviation (GSD) and the fine particle fraction (FPF). All nebulized formulations demonstrated appropriate output and aerodynamic characteristics for peripheral lung delivery. The MMAD increased only slightly due to the incorporation of NPs in aerosols, e.g. for the vibrating mesh nebulizer, the MMAD was found to be 5.4 µm for the Tb-NP200 formulation compared to 4.3 µm measured in the NP-free set-up. Despite a large output rate of the vibrating-mesh nebulizer, the air-jet technology was beneficial for deep lung delivery indicated by the larger amount of aerosolized particles smaller than 5 µm (85% FPF) and an MMAD between 2.1 µm and 2.3 µm.

In conclusion, we demonstrated that PLGA and PEG-PLGA based NPs displayed excellent properties as biocompatible, mucus-penetrating delivery systems for antibiotics with improved deposition and bacterial killing of biofilm-embedded and mucus covered pathogens. The nebulization of NPs, especially via air-jet technique, offers a highly suitable approach to deliver differently sized NPs efficient to the deep lungs.

Acknowledgments:
The authors thank the German Research Foundation (DFG) for financial support (FI 899/4-1, FI 899/4-2, PL 320/3-1, PL 320/3-2).

References:
1.) M. Klinger-Strobel, Expert Opin Drug Deliv, 2015, 12, 1351-74.
Industrial partners of the conference will get the chance to present their newest innovations and products within three minutes in this latest event. This type of appetizer is a fantastic marketing opportunity supplying talking points for later conversations. As bonus, there will be a prize for the best performance — chosen by vote of the audience.
Abstracts

October 4th // 16:30-18:00

Session 3: Cutting Edge Session

Chair: Hyung-Dong Chang

Cutting-edge research not only evolves from newly invented and optimized technologies and strategies, but may arise from combining well-known technologies in a new and optimized way. The combinations of flow cytometry and Next Generation Sequencing or microscopy and mass spectrometry are just recent examples. New technologies not only facilitate or allow high-throughput of cellular and biochemical analyses but also lead to the discovery of novel biological phenomena and processes. This session will introduce some of these novel developments and showcase the high dynamics of the field of cytometry.

Abstract – Session 3: Cutting Edge Session

16:30-17:00 Emmanuel Reynaud // UCD Centre for Biomedical Engineering / Dublin / Ireland

Light Sheet for the Masses

More than a hundred years ago in Jena a physicist and a chemist combining their efforts created the first light sheet for imaging. Their efforts won them the Nobel prize in 1925 but their idea of uncoupling illumination and detection was lost. But with better laser illumination and electronic devices, this basic principle has become a revolution in biological imaging since its revival in 1993. It is now available for histology, pathology, developmental biology, plant biology, cell biology as well as cytometry. So far more than 80 acronyms described at least one type of light sheet microscope, but is it that amazing? The talk will walk you through history and the jungle of variations around the same theme and how this technology will affect biological imaging as well as cytometry in the coming decade.

Biography:
Dr. Emmanuel G. Reynaud, MSc, BSc, PhD, is a Lecturer in Integrative Biology at the UCD School of Biomolecular & Biomedical Science. Prior to joining UCD, he was a Researcher and Postdoctoral Fellow at the European Molecular Biology Laboratory, where his research focused on development of new imaging methods (e.g. Light Sheet Microscopy) and optical micromanipulations in Cell Biology (e.g. laser nanosurgery), in the laboratory of Prof. Ernst H.K Stelzer and Dr Rainer Pepperkok. He is also the co-founder of the Light Sheet Microscopy community alongside Dr Pavel Tomancak. He also built and coordinated for 2.5 years a unique imaging platform during the circumnavigation of the Earth as part of the Tara Oceans (2009-2012). He has been awarded a Knight of Palmes academiques for his educational works.
Luminescence Lifetime Encoding in Flow Cytometry – First Steps and Assessment

Numerous analytical techniques in biomedical research rely on multiparametric analyses [1,2]. This often implies encoding or labeling by means of easily distinguishable properties like different luminescence features. Optical encoding is frequently combined with high-throughput optical-spectroscopic methods such as flow cytometry, one of the most widespread techniques in the life sciences [3,4]. Typically, luminescence encoding schemes rely on spectral (color) and/or intensity codes which can be prone to spectral crosstalk limiting the number of distinguishable codes [5]. A promising alternative is to exploit the characteristic parameter luminescence lifetime for species identification and encoding [6]. This has been suggested for flow cytometry already several decades ago [7]. However, up to now there are only few reports on the challenges of lifetime measurements within the short interaction times in flow cytometry. Moreover, most reports focus on scanning techniques, specialized applications without a general assessment, and on measurements in the frequency domain [8-12].

Here, we present first results from flow cytometry measurements in the time-domain using a custom-designed instrument with luminescence lifetime analysis capability. In this respect, major challenges of time-resolved flow cytometry and requirements imposed on temporal resolution and discrimination are discussed that originate from the short interaction time per code and the resulting weak signal-to-background ratio. To that end, we studied a set of lifetime-encoded polymer microbeads loaded with luminophores with varying luminescence decay kinetics. The experimental results were supplemented by numerical simulations of the decay kinetics and underlying photon statistics.

Our results show that lifetime encoding in flow cytometry may serve two current trends [2]: simplified instrument setups for cost-effective high-throughput methods on the one hand, and highly multiplexed analyses by introducing the additional encoding parameter luminescence lifetime on the other hand.

Acknowledgments:
We acknowledge financial support from the Federal Ministry of Education and Research 13N13357 (Forschungsförderung Photonik, VDI Technologiezentrum GmbH). We would like to thank the Laboratory of Nano-Bioengineering, National Research Nuclear University (Moscow Engineering Physics Institute, Russian Federation) for providing encoded polymer microbeads.

References:
2) V. V. Tuchin: Advanced Optical Flow Cytometry: Methods and Disease Diagnoses. 2011, Wiley-VCH, 1st Ed.
9) A. V. Gohar, R. Cao, P. Jenkins, W. Li, J. P. Houston and K. D. Houston, Biomed Opt Express, 2013, 4, 1390-400.

Advanced Imaging Flow Cytometry

Imaging flow cytometry (IFC) is a hybrid technology which extends conventional flow cytometry with additional high resolution morphological information. The objective of our work is to develop a microfluidic system for conventional and tomographic IFC.

Both, conventional and tomographic IFC is realized by advanced 3D hydrodynamic focusing which automatically aligns all particles as a sheet at a controllable z-position. Tomographic imaging Flow Cytometry extends conventional imaging flow cytometry for the image based measurement of 3D-geometrical features of cells. The required multidirectional views are generated by rotating all cells while passing the imaging window of the developed microfluidic chip. Rotation is implemented by guiding them at a shear flow position of the parabolic velocity profile. All cells pass the detection chamber as a two dimensional sheet under controlled rotation where each cell is imaged multiple times.

Experimental results show a strong focusing quality even under flow velocities below 1 mm/s. For the tomographic IFC, white blood cells with fluorescent stained nuclei are been recorded in parallel for the bright field and the fluorescence channel. Different subtypes of white blood cells can be distinguished by the shape of its nucleus. The experiments show that the multidirectional imaging enhances the identification of these subtypes compare to a single 2D view. Ongoing experiments are focusing on a label free classification of a mixed population of eight allergic pollen types using a convolutional neuronal network (CNN). The
whole process requires a high effort in data-processing containing algorithms for object
detection, particle tracking and mapping (multi-channel applications) and a CNN-model for
the particle classification.

In our work we report on a microfluidic system and method for tomographic imaging flow
cytometry, where the angular velocity of a rotating cell is controlled by its z-position in the
parabolic velocity profile of a carrier fluid. We also show the need of advanced data-process-
ing tools for image analysis.

Acknowledgments:
We acknowledge the microsystem group and the cleanroom staff at the IPHT for the devel-
opment and realization of the microfluidic units. The funding from WaterChip (EU Era-NET-
DLR 01DQ16009A) is gratefully acknowledged.

17:40-18:00 Kathrin Brenker // Opto Biolabs / Freiburg / Germany

Opto Biolabs – Combining Optogenetics with Flow Cytometry

Optogenetic tools allow isolated, functional investigations of almost any signaling molecule
within complex signaling pathways. A major obstacle is the controlled delivery of light to the
cell sample and hence the most popular tools for optogenetic studies are microscopy-based
cell analyses and in vitro experiments. The flow cytometer has major advantages over a mi-
croscope, including the ability to rapidly measure thousands of cells at single cell resolution.
However, it is not yet widely used in optogenetics.

Opto Biolabs is an EXIST-funded SpinOff from the University of Freiburg and builds custom-
ized illumination devices for optogenetic flow cytometry. Opto Biolabs’ pxONE illuminates
cells at specific wavelengths, light intensities and temperatures during flow cytometric
measurements. To demonstrate the utility of the pxONE, we characterized the photoswitch-
ing kinetics of Dronpa proteins and performed calcium flux experiments. This protocol can be
adapted to almost all optically controlled substances and substantially expands the set of
possible experiments. More importantly, it will greatly simplify the discovery and develop-
ment of new optogenetic tools and accelerate the screening for novel drugs.
Abstracts

September 19th // 18:00-19:00

Keynote – Michael Bauer

Chair: Thomas Kamradt

Michael Bauer is professor and chair for Anesthesiology and Critical Care Medicine as well as spokesman of the Center for Sepsis Control and Care (CSCC) at the Jena University Hospital.

He worked as a post-doc at Johns Hopkins University addressing molecular mechanisms of organ failure. He serves in the board of directors in research programs by the German Research Foundation (DFG) and the Federal Ministry of Education and Research (BMBF), such as “PolyTarget” or “InfectoGnostics” and was part of the task force to redefine sepsis (“Sepsis-3”).

Abstract – Keynote

18:00-19:00 Michael Bauer // Jena University Hospital / Germany

Sepsis – Changing the Paradigm

Sepsis is the most common preventable cause of death in hospitals. The adoption of the new sepsis definition last year shifted the interest from systemic inflammation (SIRS) to organ dysfunction as the clinical hallmark of an inappropriate host response. Based on a better understanding of the molecular mechanisms, the focus of the new definition is no longer the inflammatory response, but rather the impairment of organ function which results not exclusively from inflammation but also from e.g. metabolic dysfunction. The paradigm thus moves away from the infection and the systemic inflammatory response, and toward that which makes sepsis so dangerous in terms of both disease dynamics and outcome: organ dysfunction. This change of perspective requires novel diagnostic tools to enable early recognition of infected patients with an increased risk of developing sepsis in clinical routine, even outside of the intensive care unit. The new definition also promotes development of new treatment strategies with improved ability to treat sepsis causally. Diagnostic uncertainty is the main driver for delays in therapy, the mis- and overuse of antibiotics, and the failure to identify patients who might benefit from adjunctive therapies. There is a need for new sepsis biomarkers that can aid in therapeutic decision making and add information about screening, diagnosis, risk stratification, and monitoring of the response to therapy. The most promising novel approaches for diagnosis of infection and the ensuing host response consist of light-based (photonic) tools as well as on transcriptomic, proteomic, or metabolic profiling. Novel approaches to sepsis diagnostics promise to transform sepsis from a single syndrome into a group of distinct ‘endophenotypes’ and help in the development of better diagnostic tools and effective adjunctive sepsis therapies.
Abstract – Core Facility Networking Event

September 19th // 20:00-22:00
Core Facility Networking Event

Chair: Desiree Kunkel
Chair: Steffen Schmitt

This event should be an opportunity to meet and share your experiences and challenges working in a core facility. We will have three short presentations and hopefully lots of ideas to discuss afterwards in an informal atmosphere among colleagues.

We hope to spend a wonderful evening with you at the DGfZ meeting 2018 in Jena.

Abstracts

20:00-20:15 Claudia Dumrese // University of Zurich / Switzerland

Users Sorting on High Speed Sorters in a Shared Resource Environment: Effects of a Well Balanced Sorter Training

The Cytometry Facility of University Zurich provides dedicated sorter training on high speed cell sorters to its user base for 9 years already. The effects on sorter performance, as well as user behavior in a highly supportive shared resource environment are presented along with a training schedule and usage rules as an example for successful operator-free system.

20:15-20:30 Stefanie Bürger // Institute of Molecular Biology, Mainz / Germany

Open IRIS – an Open Source Resource Management Tool Implemented at the University of Mainz

Open IRIS is a free non-commercial resource management tool that was developed at the Friedrich Miescher Institute in Basel. This presentation should give you an overview about the major functions of the software and what it takes to implement Open IRIS as a campus-wide booking system as we did at the University of Mainz.

20:30-20:45 Christian Kukat // Max Planck Institute for Biology of Ageing, Cologne / Germany

Messenger Apps Facilitate Group Communication: Introduction to Slack

Slack is a workplace messaging app and team collaboration tool which is used widely in science labs (alternatives are Rocket.Chat and Mattermost). We present how we use it in our core facility and how it has improved our team communication, compared to using email.
Abstract – Session 4: NEW! Core Facility Session: Reproducibility in Flow Cytometry

9:00-9:30 Mathias Streitz // Institute for Medical Immunology, Charité Berlin / Germany

Validation and Standardization of Flow Cytometry Based Immune Monitoring in Clinical Trials

Flow cytometry is an important tool within immunological research and immune diagnostic. The current technical developments improve the handling and precision of flow cytometers but also increase the complexity of cytometry systems with respect to the numbers of variables that need to be controlled. In addition the lack of reference material and the insufficient comparison with other methods hinders the determination of the accuracy of flow cytometry data. Therefore the data quality is directly linked to the description of the imprecision and the stability of a flow cytometry based immune monitoring. At the moment, increasing interest in the effects of therapeutics on the immune system results in an increase of flow cytometry based immune monitoring in clinical studies. Therefore, quality of data becomes more important and harmonization, standardization and validation come more to the focus of immune monitoring. On the other hand this competes with the material costs and the consumption of lab resources. In addition, even if a number of guidelines report the different technical aspects of flow cytometry and provide protocols for the phenotyping of major players of the immune system, there are only few agreements about minimal standards for the determination of the imprecision and stability of a flow cytometry based diagnostic tests or immune monitoring via flow cytometry. Here we describe the necessary procedures for the validation and standardization based on the experiences within clinical diagnostic and the validation of multi-center clinical trials.

9:30-10:00 Claudia Giesecke-Thiel // Max Planck Institute for Molecular Genetics, Berlin / Germany

A Novel Method for Flow Cytometer Characterization by Determination of Detector Background, Signal-to-noise, and Dynamic Range

Accurate flow cytometer setup is fundamental to empower best experimental results. Regardless of the instrument used, maximum resolution of the populations of interest is the primary goal, meaning that ideally negative populations should be above noise or background, positive populations should be below the upper range limit and separation of the populations should be at maximum. In engineering sciences specific metrics are assigned to evaluate the lower detection limit, sensitivity and the upper detection limit at once, i.e. signal-to-noise ratio (SNR)
and the dynamic range (DNR). Recent introduction of the quantiFlash®, a pulsed precision LED light source, now enables a new way of determination of these performance metrics for flow cytometry, independent of sample or bead preparation and instrumental factors not related to signal intensity. We used the quantiFlash® to characterize an instrument's response to a stable input light signal over the entire PMT gain range. As a consequence, we propose a method to determine a flow cytometer's SNR and DNR. This allows the selection of a voltage to optimize the signals delivered by the PMTs with respect to the background consisting of scattered laser light and electronic noise. Both contributions to the background vary depending on PMT voltage and are now ascertainable and distinguishable with the here proposed method. Such knowledge further allows for separation of technical and biological background which can help with experiment design. In conclusion, we introduce a new practical method for instrument sensitivity characterization, show how the optimal PMT voltage can be defined with respect to the SNR and DNR and discuss practical implications.

10:00-10:15 Steffen Schmitt // German Cancer Research Center (DKFZ), Heidelberg / Germany

Minimal Requirements for the Presentation of Flow Cytometry Data

As a growing number of publications is indicating, reproducibility of experimental data is widely discussed in many biomedical and preclinical research fields. The correct interpretation of flow cytometric data - a relative measure of cellular properties - is strongly dependent on the context in which they have been generated. Therefore, minimal requirements for presenting this type of data have been published a while ago. These requirements could serve as a kind of standard, helping to improve the quality of flow cytometric data and will be briefly introduced during this session.

10:15-10:30 Elmar Endl // University Bonn / Germany

cytometry.de – the Communication Platform of the DGfZ

cytometry.de has been developed by a group of core facility managers to serve as a communication platform of the german society for cytometry (DGfZ). This communication platform and the connected task force have been officially integrated into the society at the member assembly 2017. The idea is to give core facility staff and researchers working in flow cytometry the opportunity to present themselves and to implement tools to communicate in an easy and personal way.

The platform along with its history will be introduced and the features that have been added so far will be discussed.
Abstract – Session 5: European Guest Session: Hungary

11:00-11:30 Gábor Szebeni // Hungarian Academy of Sciences, Budapest / Hungary

Immunophenotyping of Treatment Naive Patients with Systemic Autoimmune Diseases by Single Cell Mass Cytometry

INTRODUCTION. Epidemiological data highlights the rising incidence of rheumatoid arthritis (RA), systemic lupus erythematosus (SLE) and systemic sclerosis (SSC) in the developed world over the last decades. Currently available treatments are palliative reducing the symptoms and supporting patient’s wealth. Therefore, studies are much needed to deeply reveal cellular features responsible for pathologies or to identify early diagnostic and prognostic markers associated with RA, SLE and SSC, respectively.

METHODS. We focus on the multiparametric and functional characterization of RA, SLE and SSC using single cell mass cytometry. Treatment-naive patients suffering either in RA, SLE or SSC who had not received non-steroidal anti-inflammatory drugs, disease modifying anti-rheumatic drugs and glucocorticoids until the time of blood sampling are enrolled aligned with age and gender matched healthy individuals. Healthy controls have a negative history of rheumatic symptoms, negative status upon detailed physical and laboratory examination. Human subjects are informed and handled in line with the Declaration of Helsinki and relevant Directives of the EU. PBMCs are purified by Ficoll density gradient centrifugation. CD4+ T-cells are purified by magnetic bead separation for whole transcriptome analysis. PBMCs are labeled by antibodies to detect markers at single cell resolution by mass cytometry. In mass cytometry detection is based on antibodies which are labeled by stable metal isotopes. Thus, autofluorescence and spectral overlapping are eliminated. Markers identified by the transcriptome analysis associated with either RA, SLE or SSC are screened for validation by mass cytometry at protein level.

RESULTS. After gating out the calibration beads and cellular doublets (191Ir+/193Ir+ heterogeneous) live cells (195Pt cisplatin negative) are determined. Using 26 antibodies in one single tube the following sub-populations have been defined within single living CD45+ cells: B-cells, monocytes, CD16+ myeloid dendritic cells, NK-cells, CD4Th1, CD4Th2, CD4Th0, CD8Th1, CD8Th0, CD8Tcm, CD8Tnaive, CD8Tc1, CD8Tc2, CD8Tc0. In order to characterize the phenotype of the above mentioned cell types the following markers have been investigated on each subtype: CD5, CD7, CD9, CD28, CD40, CD161, CXXR3, CD25, CD69, CD27, CD57, CD127. Stochastic neighbor embedding (viSNE) analysis dissects the different immunophenotype of RA, SSC and SLE samples based on the multidimensional cellular relatedness of 26 marker expression at single cell resolution in each disease.

CONCLUSION. Mass cytometry deeply reveals cellular heterogeneity on the basis of highly multiplex phenotypical and functional characterization in RA, SLE and SSC.
Breast cancers, and malignant tumors of several other organs (gastric, ovary, brain) often overexpress HER2 (neu, ErbB2), a receptor tyrosine kinase of the epidermal growth factor receptor family. This is often correlated with rapid clinical progression, resistance to radio or chemotherapy, and, overall, bad prognosis. Since HER2 is not, or weakly expressed in healthy human tissues, it can be considered an ideal target of antitumor therapy. In coherency with this, the first humanized antibody against solid tumors was trastuzumab (Herceptin), targeting HER2. Although trastuzumab treatment of HER2 positive tumors has shown considerable success, resistance occurs in a significant fraction of patients, both from the start of treatment and evolving during treatment. Earlier we have shown that an important cause of this resistance is the massive extracellular matrix evolved by tumors. Considering that the antibody can mainly act by down-regulating HER2 or by recruiting antibody dependent cellular cytotoxicity (ADCC), various biochemical and immunological approaches can be proposed for decreasing or obviating the resistance against trastuzumab treatment.

Since HSP-90 serves to retain HER2 in a well-folded, but inactive conformation, the geldanamycin derivative 17-AAG can be used to enhance HER2 dimerization, consequential down-regulation, and thus to decrease proliferation of trastuzumab resistant breast cancer cells. However, this comes at a price, since HER2 activation is also increased transiently, and side effects in the clinic can be significant.

Recently, the mycobacterial antibiotic archazolid, a potent inhibitor of the lysosomal V-ATPase has been produced synthetically, which enabled widespread testing. It appears that archazolid interferes with the regular recirculation of HER2, results in its retention in phagolysosomes and autophagosomes, decreases HER2 phosphorylation, and inhibits the growth of xenograft tumors. Yet, its bioavailability needs to be improved to make it a viable option for adjuvant therapy.

Trastuzumab can also be combined with other HER2-targeted antibodies, such as pertuzumab, which increases the resistance-free period in tumor-bearing mice. The beneficial, additive effect of the combination is related to the fact that maximal approved clinical doses of either antibody alone do not saturate ADCC. Thus, it is recommended that both in the adjuvant and neoadjuvant setting the two antibodies are applied in combination from the start.

Finally, it is possible to build a chimeric antigen receptor (CAR) containing trastuzumab scFv for MHC non-restricted target recognition, and important motifs of the T cell receptor and co-signaling molecules. This, when transduced into naive T cells, enables the destruction of HER2 positive tumors by the CAR T cells that perform active reconnaissance, even when the therapeutic antibody cannot anymore penetrate the extracellular matrix with passive diffusion.
Database-Guided Flow-Cytometry: Simplification and Reproducibility in Flow Cytometry

Multiparameter immunophenotyping by flow cytometry is indispensable for the management of hematological malignancies. Despite the significant benefits provided by this technique that fulfills the requirements for high speed and broad applicability for diagnosis and follow-up of a wide range of lymphoid disorders, there are still drawbacks for myeloid disorders. The difficulty is related to the lack of consensus for panels of antibodies with high specificity and sensitivity for diagnostic assessment and prognostic stratification. Alongside, standardization of sample preparation, instrument setup protocols, fluorescence compensations and data interpretation across institutions is needed.

The aim of this study is to evaluate the potential of an immunophenotypic myeloid database for identification of surrogate markers for genetic aberrations frequently observed during AML diagnosis and which could guide the clinical protocols. In addition, evaluation of myeloid cell maturation abnormalities against an immunophenotypic myeloid database allow identification of the most relevant phenotypic abnormalities related to dysplastic changes in BM myeloid cells.

The database is an archive of cell-surface antigen expression profiles representative for normal and malignant hematopoiesis. The database of normal hematopoiesis was constructed with normal bone marrows from healthy donors of allogeneic bone marrow transplant or individuals with no evidence of a hematopoietic disease. The data were acquired on five different centers from France under the standardized conditions set by France Flow Clinical Cytometry Group. Immunostaining was performed after erythrocyte lysis using the 8-color antibody panel according to the EuroFlow consortium’s guideline (1). As already referred, the strategy of data analysis consists in separate analysis of the CD34+/-CD117+ immature committed precursors and of more mature myeloid cells (2-4). Data from distinct subtypes of acute myeloid leukemia (AML) were included in a database allowing to directly comparing antigen expression of malignant cells with those of their closest normal counterpart. Furthermore, the comparison of patterns of expression was performed in bulk bone marrows of the three cytogenetic subgroups of AML: AML t(8;21), AML t(15;17) and AML inv(16)/t(16;16). The Compass database was constructed in Infinicyt™ software. In addition, the Maturation databases were built to help evaluation of phenotypic abnormalities in cases with myelodysplastic syndromes (MDS) or suspicions of MDS.

The Myeloid Database is a comprehensive visualization interface that can make it useful as a daily procedure for pathologists and cancer researchers for AML/MDS assessment. The reduction of investigator subjectivity in data analysis is an important advantage of this method.
Inclusion of an increased number of data from healthy donors in the database improves the robustness of the analysis. Undoubtedly information stored in database will have an impact on the diagnosis, prognosis, and treatment of myeloid malignancies in the future.

Acknowledgments:
The antibodies used in this study were provided by BD Biosciences. The authors are thankful for the clinician hematologists for their interest and involvement in this study and for the patients and healthy donors for their agreement to participate in this study.

References:
2.) S Matarazz et al., The immunophenotype of different immature, myeloid and B-cell lineage-committed CD34+ hematopoietic cells allows discrimination between normal/reactive and myelodysplastic syndrome precursors. Leukemia (2008) 22, 1175–1183.

P02 Anna Bajnok // Semmelweis University, Budapest / Hungary

Distinct Cytokine Patterns May Regulate the Severity of Neonatal Asphyxia – an Observational Study

Background: Neuroinflammation and a systemic inflammatory reaction are important features of perinatal asphyxia. Neuroinflammation may have dual aspects being a hindrance, but also a significant help in the recovery of the CNS. We aimed to assess intracellular cytokine levels of T-lymphocytes and plasma cytokine levels in moderate and severe asphyxia in order to identify players of the inflammatory response that may influence patient outcome.

The effect of cytokines is primarily exerted at a microenvironmental level, but we generally measure the level of cytokines at the macroenvironmental level, from the serum. The relationship between cellular cytokine production and serum cytokine levels is undefined, cytokines in the serum come from various different sources and show a less stable kinetic in time. The advantage of analysing cytokine levels intracellularly by flow cytometry is that this methodology opens up the opportunity for precise characterization of the function and cytokine production of each cell type in a physiological, which could be of great value in identifying key cellular players of various inflammatory conditions.

Methods: We analyzed data of 28 term neonates requiring moderate systemic hypothermia in a single-centre observational study. Neonates were divided into a moderate (n=17) and a severe (n=11) group based on neuroradiological and aEEG characteristics. Blood samples were collected between 3-6h of life, at 24h, 72h, 1 week and 1 month of life. Whole blood samples were stimulated for 6h, then intracellular cytokine levels were determined using flow cytometry. Cytokine plasma levels were measured using Bioplex immunoassays.

Results: The prevalence and extravasation of IL-1β+ CD4 cells was higher in severe than in moderate asphyxia at 6h. Based on ROC analysis, the assessment of the prevalence of CD4+ IL-1β+ and CD4+ IL-1β+ CD49d+ cells at 6h appears to be able to predict severity of the insult at an early stage in asphyxia. Intracellular levels of TNF-α in CD4 cells were increased at all time points compared to 6h in both groups. At 1 mo, intracellular levels of TNF-α were higher in the severe group. Plasma IL-6 levels were higher at 1wk in the severe group and decreased by mo in the moderate group. Intracellular levels of IL-6 peaked at 24h in both groups. Intracellular TGF-β levels were increased from 24h onwards in the moderate group.

Conclusions: IL-1β and IL-6 appear to play a key role in the early events of the inflammatory response, while TNF-α seems to be responsible for prolonged neuroinflammation, potentially contributing to a worse outcome. TGF-β has a compensatory role in decreasing inflammation. These results provide novel data on the role of distinct cytokines in shaping the inflammatory response in perinatal asphyxia and influencing patient outcome.

P03 Sabine Baumgart // Deutsches Rheuma-Forschungszentrum Berlin / Germany

Reference Sample Quality Control in Mass Cytometry: Is it Necessary for Clinical Applications?

OBJECTIVE: Mass cytometry (MC) has become an established technology for cell diversity studies and in-depth immune cell profiling of patient samples. Several techniques have been implemented to secure the quality of mass cytometry data such as cell sample banking, standardized instrument setup protocols, signal normalization according to bead standards, sample barcoding, antibody cocktail freezing, signal spill-over correction, and algorithm-based data analysis. However, large sample numbers e.g. from patient cohorts require sample processing and measurement cycles on multiple days, illustrating the need to consider and monitor day-to-day variation in the data. To this end, the inclusion of a standardized reference sample into each sample processing/measurement cycle has been suggested. Based on including such a reference sample into our mass cytometry workflow, we have investigated day-to-day data variation of 19 assay/measurement cycles performed across 3 months.

METHOD: A 43-parameter mass cytometry immunophenotyping study was performed on 190 fixed and cryopreserved patient and control whole blood samples. Reference sample aliquots were processed once from a single control donor. For one sample processing/measurement...
cycle, 10 study samples were Pd-barcoded and pooled, and one aliquot of the reference sample was labeled with a separate barcode (mDOTA-103Rh) was added. All antibody staining, washing, fixation etc. steps were carried out on the sample pool ensuring identical conditions for study and reference samples. Data were acquired on a Helios mass cytometer connected to a supersampler for approx. 3 hours per pool. After normalization and debarcoding, reference sample data were clustered using the FlowSOM algorithm and manual gating was performed by using FlowJo.

RESULTS: First, signals elicited by normalization beads were analyzed as a reference. As expected, the inter-pool coefficients of variation (CV) of the mean signal intensities of the bead elements (Ce, Eu, Ho, Lu) after data normalization were low (0.5 – 1.2 %), while CV of signals resulting from staining reference sample cells with metal-conjugated cell lineage-specific antibodies ranged between 15 % for 196Pt channel (CD14, monocyte marker) and 62 % for 175Lu channel (CD7, NK and T cell marker). This documents that despite employing consistent sample banking, standardized instrument setup and signal correction according to bead standards, some day-to-day variation of data remained that could not be explained by individual sample properties and was not fully eliminated by bead-based data normalization. Despite this signal variation, both manual gating and FlowSOM clustering delivered consistent results, indicating that signal variability had no major impact on cell subset identification. Coherently, CV of frequencies of major manually gated leukocyte subsets or FlowSOM cluster among total CD45+ leukocytes showed low variability in the reference sample across the 19 days. Here, the lowest inter-pool CV was detected for neutrophils, the most abundant cell type in leukocytes (11 %). Frequencies of B cells making up about only 2 % of leukocytes, showed a CV of 59 %.

Moreover, the intra-pool analyses of post-normalization reference sample data showed a time-dependent decrease of mean signal intensity in several mass channels to varying degree, not apparently related to a specific isotope or a defined mass range, revealing that bead-based data normalization does not fully eliminate time-dependent signal drift observed for cellular signals.

SUMMARY & CONCLUSION: Analyzing reference sample data from multi-day measurements helps monitoring the technical day-to-day variability in mass cytometry data, which can be considered when analyzing and interpreting study data. Reference sample data can be used to back up the robustness of mass cytometric studies, and to disclose any artifacts possibly caused during sample processing or data acquisition. The use of reference samples will be specifically useful for monitoring assay and data consistency in multi-center studies. In addition, an error analysis for the entire process of data acquisition to estimate/calculate accuracy and precision of our mass cytometry approach can be performed based on reference sample data.

We conclude that the inclusion of reference samples in multi-day mass cytometry studies is suitable to assess the robustness of the data obtained, and should be considered a valuable additional quality control in mass cytometry.

OBJECTIVE: Crohn’s disease (CD) is a chronic inflammatory disease affecting the entire gastrointestinal tract, whereby inflammation often occurs in patches and spreads into deeper layers of the intestinal tissue. Unlike CD, in ulcerative colitis (UC) the innermost mucosal lining of the colon or rectum becomes chronically inflamed. The hallmark of both active CD and UC is the infiltration of the intestinal mucosa by innate and adaptive immune cells. To characterize CD and UC-related immune cell abnormalities, we analyzed peripheral blood leukocytes from active CD and UC patients and compared them to a group of sex- and age-matched healthy controls (HC) by mass cytometry.

METHOD: We applied a 43-parameter mass cytometric immune phenotyping panel established to be compatible with “Smart Tube” whole blood fixation system. In total, blood samples from 27 IBD patients stratified according to the disease activity scores (Harvey-Bradshaw Index, HBI, for CD and Mayo score for UC), into 11 active CD (HBI > = 5), as well as 16 active UC patients (MAYO > = 3), and 30 HC were analyzed. Samples were barcoded with Pd isotopes, processed and analyzed on a mass cytometer (Helios) equipped with a super sampler. Normalized and manually debarcoded data were analyzed using a bioinformatic pipeline based on the FlowSOM algorithm. Global data were randomly subsampled to 30,000 cells per donor and clustered altogether by FlowSOM yielding a total of 40 cell population clusters. Frequencies of clusters were statistically analyzed for differences between CD, UC and HC using an unpaired T test.

RESULTS: As expected, FlowSOM clustering yielded clusters indicative of known T and B cell subsets, monocytes, dendritic cells etc. Both, CD and UC patients showed significantly aberrant leukocyte composition compared to controls. In both diseases, frequencies of peripheral blood NK cell, B cell, eosinophil, and TCRgd+ CD4-CD8- T cell clusters were significantly decreased in comparison to controls. While frequencies of CD4+ central memory T cells with a CD161+ phenotype were increased only in active CD patients, frequencies of monocytes and regulatory T cells showed opposite deviations from controls in CD and UC, with increased frequencies in CD and decreased frequencies in UC patients. Within the T cell compartment of active UC patients, frequencies of CD4 and CD8 naive T cells were lower in comparison to the control group.

SUMMARY & CONCLUSION: Our analysis reveals multiple shared and non-shared immunophenotypic aberrations of patients with active CD and UC. While shared abnormalities may point towards immune cell subset regulation as a result of general gut inflammation, disease-specific differences observed may reflect rather specific immunopathogenic pathways active in CD and UC. In conclusion, our study demonstrates the power of mass cytometry paired with whole blood preservation of patient blood samples to provide a comprehensive overview of changes in global immune cell profiles in patients suffering from inflammatory bowel diseases.
Osmium-labeled Microspheres for Bead-based Assays in Mass Cytometry

Polystyrene microspheres are broadly applied in flow cytometry for instrument setup and monitoring instrument stability, for assessing fluorescent spillover and in various cytometric assays e.g. for absolute quantification of cellular receptors and multi-analyte profiling. The implementation of bead-based assays in mass cytometry for the same purposes is strongly desired but hampered by the lack of functionalized beads associated with sufficient amounts of heavy metal allowing for the unequivocal detection of these by the mass cytometer.

We here introduce osmium tetroxide labeling for polystyrene microspheres as a simple, quick, and universal approach to produce various kinds of functionalized beads applicable in mass cytometry. Osmium detection does not interfere with any existing reagents routinely used in mass cytometric assays, and osmium labeling of various commercially available antibody capture beads resulted in stably and uniformly labeled beads, while retaining their antibody-capture functionality. Osmium-labeled antibody capture beads retained both their functionality and osmium signal for at least two weeks when stored at -80 °C.

We show that osmium-labeled antibody capture beads can be employed for uncompromised characterization of metal-antibody conjugates, and for signal spillover assessment in complex mass cytometric data sets resulting from e.g. minor isotopic impurities of metal labels. Furthermore, osmium-labeled beads facilitate the absolute quantification of cell-surface receptors in mass cytometry.

Osmium labeling of polystyrene beads permits robust implementation of beads-based assays in mass cytometry, broadening the applicability of mass cytometry in biomedical and basic research, and increasing the quality and quantity of information retrievable in mass cytometric studies.

Microbial Flow Cytometry – a High Dynamic Resolution Method for Monitoring Ecological Stabilities of Microbial Communities.

Microbial communities drive many processes which affect human well-being directly as the human microbiome or indirectly as in natural environments or in biotechnological applications. Their complexity and their dynamics are difficult to monitor, and currently, sequence-based approaches are limited with respect to the temporal resolution. However, in order to eventually control microbial community dynamics, monitoring schemes of high temporal resolution are required. Flow cytometry provides single-cell based data in the required temporal resolution and we here use such data to compute ecological stability properties of microbial communities, such as resistance, resilience, displacement speed and elasticity. For resilience, we additionally introduce an on-line methodology which is suitable for continuous community monitoring.

Due to the workflow’s immanent ability to support high temporal sample densities, even in generation-time level, such monitoring tools will allow for a fast, continuous, and cost effective screening of ecological stable behaviour of microbes. The proposed workflow was tested on a long-term continuous reactor experiment employing both an artificial and a natural microbial community which were exposed to identical pulse disturbances. 16S rRNA amplicon sequencing of sorted cells was used to verify the findings. The computed stability properties uncovered superior stabilities of the natural community, and demonstrated the global applicability of the protocol to microbial communities in various environments, like the diverse localities of the human body or in surface water and in bioreactors, and will contribute to the development of managing schemes to manipulate microbial community structures and performances.
Circulating platelets are essential players in haemostasis and thrombosis, as they “survey” the integrity of the vascular system. Upon vascular injury, platelets rapidly adhere to the exposed extracellular matrix (ECM). After adhesion, they spread by forming protrusions like lamellipodia and filopodia and form a haemostatic plug to seal the wound. This process is associated with generation of contractile forces through the coordination of physicochemical interactions between platelet cytoskeletal components (e.g. actin, myosin and tubulin) and surface receptors (e.g. glycoprotein Ib-V-IX complex, integrins α2β1 and αIIbβ3) that recognize the exposed ECM. However, under in vivo physiological conditions and in blood flow platelets encounter a complex and dynamically changing biophysical microenvironment. So far, the biomechanics of force generation by platelet filopodia and lamellipodia and the role of cytoskeletal components in optimally sensing extracellular biophysical environment to accomplish haemostasis are poorly understood. Recent advances in microfabrication engineering have enabled the development of elastic vascular tissue-mimetic niches with a microfluidic environment for platelet function analysis.

Here we present a rapid prototyping of 3D printed deformable microstructures using two-photon lithography to assess biomechanics during fundamental biological events during platelet adhesion, spreading and activation using novel surface topographies.

**P08 Henrik Mei // German Rheumatism Research Centre Berlin / Germany**

**Immune Cell Profiling of Rheumatoid Arthritis Patients undergoing Helminth Therapy by Mass Cytometry**

Based on the hygiene hypothesis, the treatment of rheumatoid arthritis (RA) by immune modulation induced by infection with eggs of the intestinal helminth Trichuris suis (TSO) promises amelioration of the disease, associated with fewer side effects as compared to currently approved therapies. In order to gain insight in the mechanism of action of TSO therapy and to facilitate precision medicine in the treatment of RA, we studied the composition and activation state of PBMC of RA patients undergoing targeted and timely controlled TSO treatment in a placebo-controlled trial by mass cytometry.

We developed a mass cytometric antibody panel comprising 44 markers for complex immune profiling focused on T and B cell subsets and their activation status. A high degree of assay standardization was achieved by novel beta-2-microglobulin-based live-cell barcoding in conjunction with a new preservation method for metal-labeled antibody cocktails.

The comparison of 31 age- and gender-matched healthy controls with 36 RA patients prior to TSO therapy revealed numerous RA-related phenotypical disturbances in the T cell, B cell and monocyte lineages. RA patients showed diminished frequencies of MAIT cells, significantly lower frequencies of IgG- and IgM-memory B cells and plasmablasts, while a subset of CD14hi CD16lo monocytes was increased. Interestingly, we identified significantly lower expression of the inflammatory chemokine receptor CXCR3 in RA patients’ T, B and NK cell subsets. After treatment with TSO, RA patients exhibited a transient increase of CD23+ naïve B cells and γδ T cells. Ongoing work aims at further delineating trajectories of immune cell subsets over the treatment course and correlating these with the serological response to TSO and the clinical improvement of RA in the patients.

This study demonstrates the utility of massively high-parametric immune profiling by mass cytometry in chronic inflammatory conditions to identify immune cell aberrations for further consideration in immunopathogenesis research and therapy.

**P10 Vikram Srinivasa Raghavan // Indian Institute of Science, Bangalore / India**

**Validating Effectiveness of HIV-ART Using Superparamagnetic Iron Oxide Nanoparticles (SPIONs)-conjugated CD4 Cell Count**

Antiretroviral therapy (ART) for HIV infected patients includes combination of antiretroviral (ARV) drugs to suppress the HIV virus and stop the progression of HIV disease. Patients under ART medication are monitored to ensure treatment efficacy. Viral load and CD4 cell counts are being performed to determine the prognosis and for monitoring the response to therapy. CD4 cell counts are measured at the time of diagnosis and every 3-6 months. Currently, expensive and bulky flow cytometers are used for monitoring the ART efficacy. There is a need for a novel, simple and inexpensive technique to test for counting CD4 cells in resource limited settings in developing and underdeveloped countries. We developed a technique to quantify CD4 cells in whole-blood; using magnetic nanoparticles and widely used automated hematology analyzer. Magnetic field assisted cell separation is an interesting technique, in which magnetic particles are bound over target cells and separated using an external magnet. Targeted binding of nanoparticle to particular type of cell is achieved by conjugating antibodies to nanoparticles.

Magnetic assisted cell sorting technique is utilized for this purpose using Superparamagnetic Iron Oxide Nanoparticles (SPIONs). The nanoparticles were synthesized using conventional co-precipitation; the particle size was controlled by tuning the molar ratio of Fe2+/Fe3+ ions and the magnetic properties were studied. Among the different size of SPIONs, 20nm size ones were found most appropriate according to their magnetophoretic force from theoretical calculation. These SPIONs were surface functionalized to covalently link antibodies specific to CD4...
CD4 cells. The CD4-tagged SPIONs (CD4-SPIONs) had effective magnetization of 60 emu / g and average hydrodynamic diameter of 100 nm.

Here, we demonstrate an indirect method of counting CD4 cells in blood using CD4-SPIONs and Automated Hematology Analyzer. To test the concept of magnetic separation of cells; we initially used pure isolated white blood cells (WBCs) and mixed with CD4-SPIONs to separate CD4 cells from the total WBCs. The performance of cell separation is highly efficient, but isolation of WBCs is a tedious task. Hence, we experimented with whole blood and diluted blood. In whole blood, the separation efficiency was very low as the interference due to very high cells concentration. Different dilution ratios were experimented and at 1:26 ratio of whole blood and saline; optimum cell separation was achieved. Fig. 1 shows the separated CD4 cells count from total blood cells. The concentration of CD4-SPIONs, incubation time, separation time were optimized with respect to the dilution ratio of 1:26. The volume of SPIONs-CD4 reagent required for optimum separation is between 60-80 microliters for a reaction volume of 100 µl.

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Further, for more accurate and direct count of target cells, a system with imaging capability and cell sorting in flow need to be developed. This can be achieved by integration of microfluidics and microscopy system. Currently, we are performing the above Off-chip experiments in an On-chip microfluidic device (MFD) which we have customized for this application. This MFD will be incorporated with an inexpensive portable imaging flow cytometry (IFC) system for absolute CD4 counts with simple image processing algorithms.

Application of Flow Cytometry Analysis for Studying Toxicological Effects of Nanoparticles on Human Caco-2 Cells

Flow cytometry is a fast, reliable, and low-cost high-throughput screening tool for studying the toxicological properties of hazardous substances on human cells. Nanotoxicological research has also been able to benefit from the advantages of this technique in order to estimate potential risks of engineered nanomaterials. However, in addition to the advantages of toxicological analysis by flow cytometry, unintentional interactions of nanoparticles with the used measurement systems have been discovered, leading to partial false-positive or false-negative results.

In our recent work, we have therefore studied induction of apoptosis and influence on the cell cycle in human intestinal cells (Caco-2) after exposure with three metal oxide nanoparticles: titanium dioxide (TiO2), copper oxide (CuO), and zinc oxide (ZnO). Shortly, cells were grown for 48 h and treated with nanoparticle concentrations from 1-100 µg / ml for 24 h. After harvesting and washing the cells, they were stained with Annexin-FITC and propidium iodide (PI) for analysis of apoptosis and a PI staining solution (0.1 % Triton X-100, 100 µg / ml DNase-free RNase, 10 µg / ml PI) for the detection of cell cycle phases. Additionally, the interaction of the used nanoparticles (100 µg / ml in double distilled water) with the assay components Annexin-FITC and PI was investigated in a cell-free solution.

While our results show that TiO2 did not influence the induction of apoptosis, an increase of apoptotic and dead cells at 100 µg / ml was measured for CuO (8 % apoptotic cells; 38 % dead cells) and ZnO (26 % apoptotic cells, 50 % dead cells). The cell cycle analysis showed that the proportion of cells in the G1 phase increased for TiO2, and ZnO and decreased in the G2 phase. No influence of nanoparticles with PI was observed, while an unspecified binding of Annexin-FITC with TiO2, and ZnO led to an increase of FITC signal in the cell-free system.

As a conclusion, toxicological effects of nanoparticles on cells can be detected by flow cytometry, whereby the non-specific influence of these small particles on the test system is not negligible and must be excluded in preliminary experiments.

Assessing Subset-specific Leukocyte Signaling Aberrations in Chronic Inflammatory Diseases: Paving the Way for Precision Medicine in Chronic Inflammation

Objective: Aberrant signaling in leukocytes contributes to the establishment and maintenance of chronic inflammation and autoimmunity. While the activity of signaling cascades and their abnormalities are known to be cell subset-specific, we lack a systems-level understanding of which immune cells signal aberrantly in different chronic inflammatory conditions and through which pathways.

Method: A new sample collection workflow involving whole blood fixation and cryopreservation was established to arrest the ex vivo or stimulation-induced activation of leukocytes within 30 minutes after blood collection, providing access to cellular activation states that are otherwise subject to time- or environment-dependent degradation. Mass cytometry, a multiplex analysis tool offering more than 50 cytometric channels, is applied to these samples for simultaneous in depth phenotyping of peripheral blood leukocyte subsets and interrogation of cell type-specific intracellular signaling on the levels of abundance and/or phosphorylation status of intracellular signal transducers.

Results: In a proof-of-concept study, the workflow was evaluated by monitoring protein phosphorylation in ex vivo-stimulated whole blood samples from healthy donors in a time-dependent manner. Short-term kinetics for responses to IFN-α, TNF-α, IL-6 and...
PMA/ionomycin were recorded by simultaneously measuring the activation of intracellular signal transducers like STAT1,3,5, Syk/Zap70, PLCγ2 or MAP kinases in neutrophils, monocytes, B- and T-lymphocytes. These data were used to determine stimulus-specific reference response signatures in a cell-type specific manner. Preliminary data from a small set of patient samples suggests that aberrant pathway activation between diseased and healthy individuals can already be detected in ex vivo samples, without performing re-stimulation.

Summary and Outlook: By transferring this approach to patient blood samples and combining it with computational data analysis, this study aims to identify cellular activation patterns that are either commonly or specifically dysregulated in different chronic inflammatory conditions. It can be expected that these data will provide valuable new insights into disease pathogenesis and single cell-based signaling processes targeted by a new class of small molecule therapeutics in the field of chronic inflammatory diseases. Altogether, this information will contribute to a better understanding and transfer of therapeutic concepts between different disciplines and provide a new basis for precision medicine to stratify individual patients with respect to the best-fitting therapy option.

References:
1.) Baumgart et al., OMIP-034: Comprehensive immune phenotyping of human peripheral leukocytes by mass cytometry for monitoring immunomodulatory therapies. Cytometry A (2017)

COPASTM Vision Flow Cytometer Sorts and Captures Images

Union Biometrica has developed instrumentation for large particle flow cytometry that can capture images in flow. Adding imaging capability to flow cytometry greatly enhances the phenotyping of samples by providing morphological and spatial information of the sample constituents not collected by conventional flow cytometers. Traditional measurements of size, optical density, and fluorescence, as well as Profiler data, are also collected, and these measurements are used for making sorting/dispensing decisions. The collected images and flow cytometry measurements are synchronized so that objects dispensed to wells of multiwell plates can be traced back to their corresponding image. Our COPASTM technology platform is designed for large particles making it ideally suitable for large single cells, cell clusters, and small model organisms. The COPASTM Vision instrument is based on this platform and is ideally suited for samples made up of particles of varying sizes and shapes. Our data from the COPAS Vision shows proof-of-principle support for increased level of phenotyping of these types of samples, including the small model organism C. elegans and marine meiofauna samples.

Optimized 7-Color REAfinity™ Antibody Cocktail for Immunophenotyping

Introduction: Immunophenotyping by multicolor flow cytometry is a powerful diagnostic tool to assess the status of a patient’s immune system, e.g. for immune monitoring and complementary research in clinical trials or the detection of minimal residual disease after treatment for leukemia. Other possible applications include the quality control of cellular therapy products or preclinical research, e.g. mouse xenograft models. Having reliable, consistently performing antibodies and dyes is crucial to all the aforementioned applications. Here we present an improved ready-to-use seven color immunophenotyping cocktail, made up solely of our Miltenyi Biotec REAfinity™ antibodies, for the differentiation of nine leukocyte subsets. REAfinity™ are recombinant antibodies of human IgG1 isotype with a mutated region within the Fc portion, which prevents Fcγ receptor binding. Furthermore, they are characterized by high antigen specificity and consistent quality compared to conventional hybridoma-derived monoclonal antibodies. Here we show their superior performance in a multicolor flow cytometry application.

Methods: Whole blood of healthy human donors was stained with a cocktail containing antibodies recognizing the following antigens: CD3, CD4, CD8, CD14, CD16, CD19, CD45, and CD56. The cocktail either consisted of hybridoma-derived or REAfinity™ antibodies. Optimization of the REAfinity™ cocktail allowed for dead cell exclusion using 7-AAD. After red blood cell lysis stained samples were acquired on the MACSQuant Analyzer 10. This instrument supports the enumeration of the identified cell subsets via volumetric cell count determination. For further validation of the REAfinity™ cocktail additional cell materials were used, e.g. peripheral blood mononuclear cells (PBMCs), leukapheresis or cultured chimeric antigen receptor (CAR) T cells.

Results: Both REAfinity™ and hybridoma-derived cocktails enabled the identification and enumeration of B cells, CD4+ and CD8+ T cells, NK cells, NKT cells, monocytes, neutrophils, and eosinophils. Labeling with the REAfinity™ cocktail resulted in improved separation of the target populations when compared to the hybridoma-derived cocktail. For whole blood, this was in large part due to reduced unspecific background caused by Fc receptor binding on granulocytes when using the REAfinity™ cocktail. For the other cell materials tested, the cocktail showed consistent and reproducible results.

Conclusion: Results presented here demonstrate improved performance of the new REAfinity™ immunophenotyping kit in comparison with hybridoma-derived antibodies, due to high antigen specificity and lower unspecific background. Together with an upcoming new Express Mode (automated gating and analysis tool), the kit allows for rapid and highly reproducible immunophenotyping of various cell materials.
The Domain of Image Based, Fluorescence Analysers in the Generation and Monitoring of CAR-T Cell Therapy

There is a new therapeutic concept, based on a creation of modified CAR-T (chimeric antigen receptor T-lymphocyte) cells, dispensed to patients by a one-time intravenous infusion. As the CAR-T cells are the essence of this novel medicine, the monitoring of their status over the whole production process, as well as the monitoring of patient’s blood samples, is an indispensable part of the quality assurance of the therapy concept.

We will present innovative, image based methods, using a fluorescent and bright field cell culture analyser. The Countstar Fluorescence (FL) is an ideal system to combine not only cell concentrations, and viabilities measurements, but additionally documents various cell-specific characteristics of CAR-T cells, and their stability during the consecutive steps of a QbD designed manufacturing process. We were able to detect, and to quantify changes of cells during their cultivation process. The Countstar FL demonstrated to be fast, accurate, and sensitive analyser.

Advanced Applications with CCD Based Flow Cytometry

Beside our traditional CCD based imaging flow cytometers (ImageStream® and FlowSight®), Merck has launched this year the new CellStream® benchtop flow cytometry system, a highly-customizable and compact flow cytometer that is the first to use a camera for detection. Its unique optics system and design provides researchers with unparalleled sensitivity and flexibility when analyzing cells and submicron particles. Within the CellStream® system, the Amnis® time delay integration (TDI) and camera technology rapidly captures low resolution cell images and transforms them into high-throughput intensity data. Researchers acquire the intensity data they are accustomed to from traditional flow cytometers, but with greater fluorescence sensitivity. Here we will take a deeper look inside the instrument design, system performance and software of this exciting new flow cytometer.

Maximizing Human Immune Monitoring Studies with Mass Cytometry

Immune monitoring is an essential method for quantifying changes in immune cell populations in chronic inflammation, infectious disease, autoimmune disease and cancer studies. The extreme heterogeneity of immune cells demands a high-parameter approach to more fully and efficiently quantify the immune response in health and disease. Mass cytometry is an ideal solution, enabling the simultaneous detection of over 40 phenotypic and functional markers in a single tube of sample. We report development of a 29-marker panel for mass cytometry based on the Human ImmunoPhenotyping Consortium (HIPC) consensus panel [Maecker et al. Nature Reviews Immunology (2012)], expanded to allow identification of additional leukocyte subsets, particularly T cells. Automated data analysis with Verity Software House GemStone™ software has been developed specifically for data collected with the panel. Extensive panel testing for repeatability, reproducibility and agreement of full versus partial panel population identification was performed. Repeatability was tested with a single PBMC sample stained by a single technician in two technical replicates and acquired in triplicate on two Helios™ mass cytometers. SDs for percent of parent were 1% or less for 16 identified populations. Reproducibility was tested by determining the variability in measurements of five PBMC lots stained by five technicians and collected on two Helios instruments. CVs on mean percent of 13 populations were under 15% for all but three of 130 measurements. Lastly, R2 values for agreement of percent parent populations using the full 29-marker panel compared to a 10-marker panel for T cell populations were 0.94 or higher. We conclude that this panel kit can provide consistent immune population identification and enumeration for any given lot of PBMC.
Deformability Cytometry and 1D Fluorescence Imaging in Real-time

During the last decades, tools for rapid characterization of large cell quantities have become indispensable not only for basic research but also clinical diagnostics. The gold standard for cell characterization is flow cytometry. Its success is closely tied to the availability of fluorescent labels. But what if there is no molecular marker known for the cells of interest? Or if the label changes cell function? Or cells shall be used for transplantation? As an attractive alternative, deformability cytometry exploits cell mechanics as a sensitive, inherent, label-free functional marker, but lacks the specificity provided by a fluorescent signal. Here we present real-time fluorescence and deformability cytometry (RT-FDC), the ideal, combined system. It facilitates fluorescence detection as in conventional flow cytometry, extended by 1D analysis of spatial information encoded in the fluorescence pulse shape, and adds bright field imaging for mechanical phenotyping of single cells — all in real-time at rates of 100 cells/s. We show utility of RT-FDC for the most common fluorescent labels: Fluorescent surface markers (CD34) are used to separate human hematopoietic stem and progenitor cells (HSPCs) from an unpurified apheresis sample as harvested for bone marrow transplantations. Membrane permeant dyes identify reticulocytes by their ribonucleic acid (RNA) content in a blood sample. And endogenously expressed fluorescent proteins (FUCCI) reveal cell cycle phases in an unsynchronized sample of retinal pigment epithelial cells (RPE1). In addition, we can now also directly correlate mechanical characteristics with fluorescence intensity and localization for each single cell to improve correct classification. In future, this combined approach could establish mechanical phenotyping as equivalent to fluorescent labeling, or even identify subpopulations invisible to molecular labels.
i.e. T helper-cell mediated, cross-priming, it is believed that a brief licensing interaction can result in a licensed state of dendritic cells that wanes over time. In consequence, dendritic cells, T helper cells, and CTL may hypothetically interact sequentially. The in vivo dynamics of alternative (i.e. NKT-cell mediated) cross-priming are unknown. Using a novel approach to conditionally deplete NKT cells in vivo, we compared the sequence of cellular interactions in classical and alternative cross-priming. Strikingly, in classical cross-priming, CTL expansion was blunted when they were not present simultaneously with T helper cells early after immunization. In alternative cross-priming, NKT cells activated DCs during this time, but their simultaneous presence with CTL only had a weak effect on the CTL response. NKT cells provided productive CD40L signaling later, when they had relocated to the splenic white pulp. Depletion of NKT cells after the initial activation of DCs, therefore, shortened the CTL expansion phase and compromised their differentiation into effector phenotype CTL. In contrast, depletion of T helper cells at this time did not alter the quantity or quality of the CTL response.

These data suggest that NKT cells unfold their effect on the primary expansion of CTL in two phases: They activate DCs in the marginal zone upon which both cell types relocate to the white pulp. There, NKT cells shape the CTL response with the help of CD40L, ensuring optimal CTL generation. T helper cells, on the contrary, provide CD40L signaling early after immunization and lose their influence on the CTL response thereafter. These results provide novel insights into the dynamic three-cell interactions during the cross-priming of CTL.

Some cytosolic pattern recognition receptors (e.g. NLRP3) can recruit multi-protein signaling platforms, termed inflammasomes, which recognize diverse danger signals derived from either pathogens or sterile cell damage.

Inflammasomes are powerful inducers of caspase-1-dependent pyroptosis and maturation of IL-1β and IL-18. These cytokines are potent pro-inflammatory mediators implicated in numerous infectious, metabolic, and autoimmune diseases. Therefore, activation of the inflammasome is tightly regulated.

During inflammasome activation, all ASC molecules of a cell are recruited into a single cluster, resulting in a dramatic molecular redistribution. Using flow cytometry, this relocalization event can be monitored by pulse shape analysis.

Intriguingly, we observed that the percentage of NLRP3-activatable human primary cells is strongly reduced in comparison to murine cells. In the following, we could show, that human but not murine, NLRP3 mRNA is subject to alternative splicing. The major alternative NLRP3 isoform lacks the first two canonical leucine-rich repeats, is not activated by commonly used NLRP3 stimuli and does not induce inflammasome formation. Consequently, cells expressing the NLRP3 splice variant are protected from pyroptosis. We could link this loss of function of the alternative splice variant to the inability to interact with NEK7. Furthermore, we could show differential expression of the different isoforms on single cell level, potentially rendering some sub-fractions of a cell pool inert to activation and pyroptosis.

We propose that alternative splicing represents an additional, not yet described level of NLRP3 regulation and it is very likely that similar regulatory mechanisms are conserved across most NLR proteins.
Abstract – Guest Lecture

17:00-18:00 Timo Mappes // Deutsches Optisches Museum / Germany

Early Precursors of Fluorescence Cytometry: Foundation of Cell Theory and Fluorescence Microscopy

Fluorescence based cytometry is eventually based on (1) understanding of cells, and (2) the principle of fluorescence. The foundation of both topics is closely connected with the site of Jena.

The cell theory was formulated in 1839 by Theodor Schwann (1810-1882) and Matthias Jakob Schleiden (1804-1881). Schleiden performed the major part of his research in Jena, closely collaborating with the microscope makers of his time. The developments in optics enabled him to microscopically see and characterize the cellular structures and eventually to postulate: (A) All living organisms are composed of one or more cells, and (B) the cell is the most basic unit of life.

Pushing the resolution in microscopy has been a major driver of development since establishing the Abbe equation: \( d = \frac{\lambda}{2 \text{NA}} \). Reducing the wavelength into the deep Ultraviolet and thus improving the resolution, August Köhler (1866-1948) and Moritz von Rohr (1868-1940) designed dedicated optics. In 1903 the team presented a microscope's beam path designed entirely for 275 nm, enabling wide-field immersion microscope-photography with magnifications up to 2500 x. While doing so they were the first to observe autofluorescence in the microscopic sample.
Enjoy the captivating movement of the celestial body come rain or shine. The Zeiss-Planetarium was the fourth big planetarium to open worldwide. Since March 1926 people are able to study the stars on a projection screen of nearly 800 m². After the destructions during WW II, Jena’s planetarium remained as the oldest one in the world. In 1984/85 extensive reconstructions led to many advancements. The installation of computer-operated technology and the spatial expansion led to a new orientation in program offers and visitor service.

Join us for the DGfZ Networking & Get-Together at the Zeiss Planetarium on Thursday evening. There will be a buffet, drinks and samples of various Planetarium shows. Furthermore the Klaus Goerttler prize will be awarded, as well as the poster prize and the prize for the best product slam presentation.

How to get there:
The Planetarium is within walking distance from the city center. From the conference venue: 9 min walk (700 m)
Molecular Diagnostic in the Age of Multiresistant Gram-negative Bacteria

Currently, the spread of multiresistant Gram-negative bacteria (MRGN) producing extended-spectrum \(\beta\)-lactamases (ESBL) and carbapenemases is a global threat to public health. Bloodstream infections with MRGN are associated with increased mortality, which is primarily due to delayed appropriate treatment resulting in clinical failure. As most sepsis guidelines, the German guideline suggest an empiric treatment with a \(\beta\)-lactam with antipseudomonal activity (i.e. piperacillin/tazobactam, ceftazidim, cefepim or a carbapenem) in an optional combination with an aminoglycoside or a fluoroquinolone, which does not cover ESBL-producers except for carbapenems (last resort antibiotics). With up to 72 hours to result, the standard blood culture-based routine diagnostic is highly sensitive for detection of bloodstream infection, but takes often too long for critically ill or septic patients.

In attempts to improve and accelerate diagnostic procedures, numerous molecular methods have been developed. Commercially available molecular approaches focus on species identification and cover only a limited number of resistance genes and are therefore of minor use for clinical decisions regarding antibiotic treatment. Whereas a simple PCR based approach maybe suitable for multi-drug resistant Gram-positives with ‘monogenetic’ resistance, most of these tests fail to accurately detect MRGN. This is explained by the highly genetically and phenotypically diverse superfamily of \(\beta\)-lactamases containing more than 1500 variants subdivided in at least four purely related classes mediating resistance to penicillins, monobactams, cephalosporins (ESBL) and/or carbapenemases. The resistance genes are usually encoded on highly mobile multi-resistance plasmids driving a fast intra- and intra-species spread of those in Gram-negatives. Moreover, other factors such as reduced influx and increased efflux, or changes in expression levels, influence the phenotypic resistance profiles, but these factors are not covered by molecular tests. Thus, an one-fits-all assay remains unlikely to be developed covering the species, all \(\beta\)-lactamases and other resistance mechanisms. Therefore, we need to better understand the dynamics of resistance genes in MRGN to find appropriate targets for molecular diagnostics. In the meantime, blood-culture based diagnostic remains the gold standard.
Raman Spectroscopic Cytometry for the Detection and Characterization of Bacterial Infection

Infectious diseases are one of the leading causes for deaths worldwide. In order to efficiently treat an infection, physicians need to know which pathogen is causing the infection and—in case of a bacterial infection—the pathogen’s antibiotic susceptibility. Established microbiological methods used in routine clinical diagnostics are mainly based on cultivation to detect the bacteria and to generate enough biological material for subsequent analysis, such as antibiotic susceptibility testing. Thus, they need at least a full day to provide the result. Faster methods are urgently needed to administer tailored antibiotic therapy already early on.

In this presentation, a new Raman spectroscopy-based approach is presented which holds the potential to dramatically reduce diagnosis times. By means of Raman spectroscopy (individual) cells can be characterized without the need of any external label making sample preparation very easy. The inelastic scattered light provides highly specific information of the overall molecular composition of the investigated cells yielding a so-called “spectroscopic fingerprint”. This spectroscopic fingerprint in combination with multivariate statistical data analysis can be used to characterize immune cells from the host, but also to characterize the pathogen causing the infection.

Here, we will focus on the characterization of the bacterial pathogen and present how Raman spectroscopy can be used to distinguish different pathogens. This will exemplarily be shown for bacteria from patient’s urine samples. Furthermore, Raman spectroscopy can be applied to differentiate the bacteria’s response to antibiotic treatment. From this response valuable information on the pathogen’s antibiotic susceptibility can be extracted. This can be done in a qualitative manner to classify the bacteria as sensitive or resistant; as well as in a quantitative manner yielding the antibiotic’s minimal inhibitory concentration (MIC).

Acknowledgements:
Financial support by the BMBF via the Integrated Research and Treatment Center “Center for Sepsis Control and Care” (CSC, FKZ 01E01502) and via the Research Campus InfectoGnostics (FKZ 13GW0096F) is highly acknowledged. Furthermore, the project was supported by the Free State of Thuringia (FKZ 2015 FGI 0011 and 2016 FGI 0010) with cofinancing from the European Union within the European Regional Development Fund (EFRE). The work is supported by the COST Action “Raman-based applications for clinical diagnostics — Raman-4Clinics” (BM 1401) and the DFG via the Jena Biophotonic and Imaging Laboratory (JBIL).

Who am I and if Yes, How any? A new Way of Phytoplankton Species Identification by Combination of Image Cytometry and Deep Learning

Phytoplankton species composition is directly linked to water quality. Especially drinking water reservoirs, bathing and ballast water need to be regularly monitored for harmful species. Microscopic investigations are the common standard of phytoplankton monitoring, guaranteeing high quality of analyses. However, by using microscopy, there are severe limitations like taxonomic expert requirement, difficulties in providing standardized taxonomic knowledge and time consumption (limiting the samples which can be processed in total). Image cytometry allows to collect thousands of images per minute. This technique allows to separate first phytoplankton functional groups based on fluorescence images and subsequently using brightfield images for species identification. Images of reference species can ideally be used for training of a deep learning network. Deep learning is an advanced form of machine learning, avoiding subjective feature selection. A training and validation of a dataset with nine phytoplankton species at three different life cycle stages (young, well developed and senescent cells) revealed a high accuracy (97%) of correct species identification and life cycle assignment. The innovative combination of high throughput image cytometry and deep learning is therefore suggested as a promising tool to monitor water quality in future.
Microfluidic System for Single-cell Analysis in Picoliter-sized Batch Bioreactors

In this contribution we demonstrate a polydimethylsiloxane (PDMS)-based device for the cultivation and analysis of bacterial cells under batch conditions. Microfluidic cultivation devices, in combination with time-lapse microscopy, provide simple but capable tools to study biotechnologically relevant parameters like cell growth and productivity of individual cells [1]. In comparison to other cell analysis tools, these devices facilitate a high level of environmental control combined with time-resolved observation of living cells.

So far, most microfluidic cultivation systems feature chambers for cell growth, which are connected to continuous medium perfusion, thereby facilitating a continuous nutrient supply and removal of side products [2], leading to continuous cell growth. However, most conventional analyses and cultivations are performed in batch mode. Results obtained from continuously perfused systems can therefore hardly be transferred to applications conducted in batch mode.

We present a device featuring six channels, each containing 49 picoliter-sized bioreactors that can be reversibly isolated from continuous medium supply (Fig. 1). Therefore the cultivation medium is removed by a continuous flow of pressurized air, preserving a fixed volume of cultivation medium inside the bioreactors.

We show the application of our device for the analysis of survival strategies of Bacillus subtilis in case of nutrient deficiency. Therefore, the cells were cultivated in batch mode in our device until stationary phase. The live-cell imaging revealed a heterogeneous production of a toxin which is assumed to be responsible for cannibalism within the cell population during nutrient limitations (Fig. 2). The physiological relevance and the consequences of the heterogeneous toxin production are still under investigation. We were also able to observe cell growth and spore formation when the majority of the population died, indicating that Bacillus cells exhibit a range of physiological states during the stationary phase in order to increase the chances for the population to survive adverse environmental conditions.
Abstracts

September 21st // 11:30-13:00
Session 8: Microscopy Session

Chair: Anja Hauser
Chair: Raluca Niesner

The present session gives insight into new developments such as multiplexing in histology and its analysis, an emerging field known as histocytometry. We will also get an overview on how super-resolution microscopy can be used to image at a molecular level in biomedical research. Finally, we will learn about innovative ways to make cutting-edge high resolution microscopy available to a large community.

11:30-12:00 Denis Schapiro // University of Zurich / Switzerland

histoCAT: Analysis of Cell Phenotypes and Interactions in Multiplex Image Cytometry Data

Single-cell, spatially resolved ‘omics analysis of tissues is poised to transform biomedical research and clinical practice. We have developed a computational histology topography cytometry analysis toolbox (histoCAT) to enable the interactive, quantitative, and comprehensive exploration of phenotypes of individual cells, cell-to-cell interactions, microenvironment, and morphological structures within intact tissues. histoCAT will be useful in all areas of tissue-based research. We highlight the unique abilities of histoCAT by analysis of highly multiplexed mass cytometry images of human breast cancer tissues.

12:00-12:30 Christian Eggeling // Leibniz Institute of Photonic Technology / Germany

Super-resolution Microscopy: Challenges and Potentials in Biomedical Research

Understanding the complex interactions of molecular processes underlying the efficient functioning of the human body is one of the main objectives of biomedical research. Scientifically, it is important that the applied observation methods do not influence the biological system during observation. A suitable tool that can cover all of this is optical far-field fluorescence microscopy. Yet, biomedical applications often demand coverage of a large range of spatial and temporal scales, and/or long acquisition times, which can so far not all be covered by a single microscope and puts some challenges on microscope infrastructure. Taking immune cell responses and plasma membrane organization as examples, we outline these challenges but also give new insights into possible solutions and the potentials of these advanced microscopy techniques, e.g. for solving long-standing questions such as of lipid membrane rafts.

Key words: Super-resolution microscopy, STED microscopy, STED-FCS, T-cell activation, plasma membrane organization, lipid rafts
Histo Cytometry using Multi Epitope Ligand Cartography (MELC)

Bone marrow stromal cells take a crucial part in controlling of immune responses by influencing the generation of lymphocytes, the induction of central tolerance, the response to antigen and the maintenance of immunological memory. They support differentiation and survival of hematopoietic stem and progenitor cells, and provide crucial survival signals to certain subsets of memory cells, such as long lived plasma cells and T cells. Bone marrow stroma cells are thought exert their function by forming tissue niches, which are multi-component entities on a cellular and molecular level. The analysis of single niches needs a multiplexed analysis in situ. Here, we demonstrate the use of Multi Epitope Ligand Cartography (MELC) to gain multiplexed information about spatial distribution of cells in histological sections.

MELC allows us to co-map and investigate a large number of histological markers on a single tissue section. Its principle is based on a sequential cycle of staining with an antibody coupled to a fluorophore, washing, and imaging of the sample with subsequent fluorescence removal via photo-bleaching.

The acquired images are transformed to signal-to-noise ratio (SNR) images. This normalization of the images enables the investigation of the single staining quality and the comparison of different stainings. Transformed images are then segmented within CellProfiler to extract the contained information on a single cell level. Cells, i.e. objects, obtained in this way are further used to measure object-based properties like spatial distribution, co-localization or protein expression levels. These analysis steps are processed within Matlab and a self-written ImageJ/Fiji Plugin.

We developed a Plugin in ImageJ/Fiji to analyze stroma cells subsets from MELC image data sets. In addition to the amount of co-localized markers, the spatial distribution of the chosen markers indicates heterogeneous distribution in the bone marrow. A marker-specified searching approach separates classified objects inside the segmented images, to analyze and visualize co-expressed markers in specific combinations. We further developed scripts in Matlab to investigate the neighborhood behavior of these cells.

Conclusion: Using Multi Epitope Ligand Cartography in combination with self-developed analyzing tools allows us to visualize the heterogeneity of cellular subsets in tissue. We highlight the power of our evaluation tools by studying the heterogeneity of stromal cells in the bone marrow.
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