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What is a Core Facility...

Much, if not most of the research in Life Science depends on the availability of and access to resources like advanced technologies, databases, sophisticated instrumentation, and the expertise for proper operation. The interdisciplinary approach to address current topics in Life Science causes interdependencies, that do not stop at technical, scientific or at institutional boundaries. The complexity of operation has therefore promoted the aggregation of these technologies into Core Facilities.

The Medical Faculty recognizes that Core Facilities require continued and significant investment and support. Several entities exemplify the commitment.

- An active standing faculty committee (Dekanatskommission Core Facilities) that serves as an institutional “board of directors”. The committee meets regularly to advise the Dean on institutional investments of financial resources in the Facilities and reviews facility operations.
- Steering Committees that oversee and address user satisfaction, facility operation and requests for the development of new services in each Core Facility.
- A Core Manager Committee that regularly discusses the practical implementation of recommendations and decisions made by the Dekanatskommission and the Steering Committees and addresses daily life issues concerning the management of Core Facilities.

Core Facilities are evaluated on a yearly basis by the Dekanatskommission to ensure economic efficiency, implementation of strategic decisions, availability of technical equipment and support, technology awareness, scientific outreach, research output and compliance with DFG policies.

... and what is it good for

- Core Facilities provide all researchers access to state of the art instrumentation and methods and assure an easy and structured way to do so. However, cutting edge instrumentation does not guarantee high quality research. Adequate quality controls, system performance tracking, experience with the methods as well as assistance in experimental setup and data interpretation are indispensable components of a Core Facility.

- It is therefore not all about the number of lasers that you have on the machines or the number of base pairs that you can sequence at a given time. It is about people who work in the Facility. What you could or what you should expect from a Facility is people, or, better, experts, that you can talk to and that will listen to you. People that will help you from your idea through your experiments all the way to your publication or next grant proposal.

- They will take care that you’re not starting your project with an error that they have seen many others make before, and they will stay at your side to help with time management. They care about the instrumentation and will provide you with all the tools that you need to do your experiments, so that you can focus on your research. And it doesn’t matter if you are a PhD student or a PI with a European Consolidator Grant.

- Despite the fact that Core Facilities are already established within the faculty, there is a lack of awareness regarding the full potential of currently available resources. The intention of this handbook is to increase the visibility of resources and promote the communication among Core Facilities and their prospective users.

- Enough said about the concept of the Core Facilities at the medical faculty. Enjoy reading about the methods and instrumentation in the Core Facilities and may good ideas come along.

Core Facilities Business and Operations Coordinator

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Expertise

The Core Unit for Bioinformatics Data Analysis (CUBA) provides quantitative and computational analyses, training and consultancy. Increasingly complex and large data sets are characteristic for molecular biology and medical research driven by technology advancements. These high-volume data sets require quality control to filter artefacts, statistical rigor to minimize false positive findings and intelligent experimental design to maximize insights. Data processing pipelines, e.g. for transcriptomic data, are provided as well as guidance on available analysis tools and databases. Analyses are primarily based on freely available open source software essential for reproducible and independent research.

CUBA benefits from shared computing infrastructure and exchange of expertise due to its embedding and liaison with the Institut für Medizinische Biometrie, Informatik und Epidemiologie (IMBIE) and the Institute for Genomic Statistics and Bioinformatics (IGSB). It operates in close collaboration with other core units like the NGS, flow cytometry and mass spectrometry core. Altogether, this facilitates integrative bioinformatics approaches and offers a systems biology perspective.

Moreover, data of different type and technology not only represent the same biological sample but also share characteristics like high dimensionality. Thus, common approaches for clustering, structure preserving visualizations and dimensionality reduction can be applied and synergies across different projects emerge.

Methods

- Our expertise ranges from experimental design and study planning to data processing, bioinformatics, statistical analysis and visualisation. We offer bioinformatics services in:
  - Statistical analysis and machine learning
  - Consulting on experimental design and methods
  - Single cell RNA-seq and RNA-seq (mRNA, miRNA, etc.)
  - Exome and genome sequencing
  - Variant calling and Burden tests
  - Somatic mutation calling
  - Pathway enrichment
  - ATAC-seq and ChIP-seq
  - Mass-Spec based proteomics (LFQ, TMT, DIA, SILAC)
  - Metabolomic and Lipidomics, MS analysis of pull-down proteins
  - Flow cytometry (FACS data)
  - Methylation and copy number variation analysis
  - Microbiome analysis

In addition, CUBA offers trainings in programming and bioinformatics analysis.

- We apply a variety of tools with an emphasis on publicly available software like Bioconductor and nf-core. Our own analysis pipelines are developed using R and Python programming languages and executed in a High Performance Computing (HPC) environment. Moving beyond standard analysis, e.g. for transcriptome and proteome data, we are aiming on more integrative multi-omics approaches, network and single-cell analyses.

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Methods

- **The Cell Programming Core Facility employs state-of-the-art cell (re-)programming, genome editing as well as differentiation technologies and is constantly refining its technological portfolio. To promote the establishment of innovative stem cell-based in vitro models, we offer the following services:**

- **Generation and provision of human iPS cell lines**
  As starting material for iPS cell derivation we accept skin fibroblasts or peripheral blood mononuclear cells (PBMCs). Donor cells are reprogrammed using the latest Sendai virus-based reprogramming vectors that guarantee transient introduction of transgenes to avoid the stable integration of reprogramming factors. As a standard procedure, we isolate and expand up to five clonal iPS cell lines per subject for cryopreservation and subsequent shipping to the investigator. Depending on the recipients’ demands, selected clones are subjected to standardized quality control regimens (ICC and gene expression profiling for pluripotency analysis, assessment of transgene removal, STR analysis, SNP genotyping).

- **Genome editing**
  We also offer genome editing services for the generation of isogenic iPS cell lines with KO alleles, introduced and repaired point mutations, respectively, as well as the targeted introduction of transgene expression cassettes into the genomic ‘safe harbor’ locus AAVS1.

- **Neural differentiation of iPS cells**
  The Cell Programming Core Facility has in-depth expertise in the differentiation of iPS cells into neural precursors as well as terminally differentiated neural cells. Currently, we provide the generation and provision of stable neural precursor cells (NPCs), cortical NPCs and NGN2-induced excitatory forebrain neurons, all of which are available in cryopreserved formats. Please contact us for availability of other neuronal cell types.
The Flow Cytometry Core Facility covers a broad spectrum of flow cytometric methods and applications. The multiplexed analysis of immune cell subsets has once again gained importance. 6 to 8 color experiments are now common to precisely identify and separate for example regulatory T-cells, tissue specific naive and memory T-cells, tissue primed T-cells or tumor specific T-cells for ex vivo functional analysis or adoptive transfer. The demand for a multiplexed analysis between 10 and 18 colors simultaneously is no longer extraordinary, and strategies for the isolation of rare cell populations are frequently asked for. Beside these obvious applications of flow cytometry, the “flow cytometry” platform contributed to the analysis and separation of micro vesicles, operating the cytometers at their physical detection limit, and encouraged the interaction of researchers from different disciplines. Furthermore, the flow cytometry platform offers educational activities as stand-alone opportunities or as part of the existing graduate and professional programs, provides technical training, and conducts research that enables development of the next generation of state-of-the-art flow cytometry technology.

Researchers are trained by Core Facility personnel to perform their experiments without assistance of an operator. The Core Facility takes care of eight analyzers from standard three laser, 8 color systems to five laser, 18 color instruments. Users receive detailed advice and support on the choice of dyes and equipment that are best suited to answer their scientific questions. Systems available include: three BD FACSCantoll, one Miltenyi MACSQuant, two BD LSRFortessa.

Cellsorting is mainly performed by experienced operators from the Core Facility. Three high-end high-speed Cellsorters are available: BD FACSaria Fusion (biosafety level S2 approval), BD FACSaria II, BD Influx. For operator free cellsorting of less advanced applications following system is available: BD FACS Melody.

Imaging Flow Cytometry combines the speed, sensitivity, and phenotyping abilities of flow cytometry with the detailed imagery and functional insights of microscopy: Amnis ImageStreamX MarkII. Researchers receive support in experiment design and data analysis by the Core Facility staff.

Multiplexed, bead-based analysis of cytokines, hormones or many other soluble biomolecules can be conducted on one of the Flow Cytometers specially developed for this application: Luminex200, Luminex FlexMap 3D or MAGPIX.

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The ability to generate genetically modified mouse models significantly contributes to the molecular understanding of human health and disease. The Gene-Editing Core Facility supports researchers in the process of planning and generation of genetically modified mouse models, exploiting a longstanding technical and scientific expertise. A main focus of the Core Facility is put on state-of-the-art gene-editing technologies utilizing CRISPR/Cas9-mediated genome engineering, which has been successfully established since 2015. The application of this versatile genome-editing tool accelerates the generation and establishment of genetically modified mouse models and is about to become the gold-standard for the generation of gene knockouts, conditional mice as well as insertions of small tags or point mutations. These models can be often generated without elaborate and time-consuming cloning of targeting vectors and subsequent recombination in murine embryonic stem cells. Thus, the Core Facility aims to utilize CRISPR/Cas9-mediated approaches for the generation of genetically modified mice whenever possible and is actively working on improvements and refinements of the editing technologies, considering latest publications and methodologies in the field.

Methods

- **Generation of CRISPR/Cas9 gene-edited mouse models**
  The Gene-Editing Core Facility develops and employs customized gene-editing strategies for the generation of genetically modified mouse models. Ubiquitous gene knockouts are generated using a combination of two different guide RNAs in combination with non-homologous end-joining (NHEJ) repair to induce genomic deletions of a few base pairs (bp) up to 5 kbp to disrupt a gene reading frame or to delete larger regions. Precise point mutations or small tags are introduced using the CRISPR/Cas9 system in combination with homology directed repair (HDR) using short single stranded oligonucleotides (ssODNs) acting as repair template. The components are delivered into fertilized oocytes via electroporation allowing for a fast and high-throughput manipulation of oocytes for an efficient and cost-effective generation of gene-edited mice. For conditional strategies we aim to use the Easi-CRISPR approach using 1-2 kbp ssODNs. These constructs are delivered to fertilized oocytes via pronuclear or intracytoplasmatic injection using an inverted DIC (differential interference contrast) microscope equipped with micromanipulators, Piezo-element and an Eppendorf Femtojet injection device. Micromanipulated embryos are transferred into pseudopregnant foster mice by oviduct transfer of 2-cell stage embryos to finally obtain gene-edited founder mice.

- **Blastocyst injections**
  For knock-ins of larger sequences (>1.5 kbp), the CRISPR/Cas9 system is not efficient enough yet and requires „classical“ embryonic stem cell (ES cell) based approaches. If provided with a targeting construct, the Core Facility will perform the ES cell targeting in proven germ-line competent ES cells followed by blastocyst injection. Blastocysts are transferred into pseudopregnant foster mice to generate chimeras.
Expertise

Magnetic resonance imaging is a non-invasive imaging procedure that has become an indispensable part of modern medical research. The advantage of this method is that tissue types of varying density can be displayed in MRI images with different contrasts. This is particularly beneficial in neurological imaging of the brain and is the basis for neurocognitive research.

Via the BOLD effect it is possible to observe neuronal activity in the brain (functional imaging, fMRI). Other techniques include resting state MRI or diffusion tensor imaging (DTI).

The Core Facility “Human 3T MRI” (CF-MRI) provides access to a modern MRI scanner (Siemens Magnetom Trio) for research purposes.

The CF-MRI supports scientists in the implementation of their research project, for example in measurement planning, the creation and optimization of measurement protocols or troubleshooting, but also in the development and programming of paradigms. To enable their users to carry out the investigations independently, CF-MRI offers training courses. The goal is to ensure the highest quality in data recording and the safe execution of the investigations. CF-MRI offers support in accessing and pre-processing the data.

Instrumentation

Scanner

The scanner is equipped with a Spine Matrix Coil and two Body Matrix coils, the standard 12-channel head coil and a 32-channel head coil. Furthermore, a transmit/receive coil (TX/RX) can be used for spectroscopy. The scanner holds sequences to cover a wide range of examinations. Besides the usual sequences for brain imaging (FLAIR, STIR, T2-weighted, T1-MPRAGE), protocols for DTI and functional MRI (EPI) are available. The Core Facility supports its users in setting up the measurement protocols and adjusting the measurement parameters.

Paradigms

The Core Facility provides a workstation for the presentation of paradigms (stimulation in functional MRI). Various software packages like “Presentation” (Neurobehavioral Systems), „matlab toolbox“ or „psychopy“ are available to the user. In addition, the Core Facility also provides the “ScenarioDesigner”, a python-based program developed by the Core Facility itself. It enables the particularly simple programming of paradigms. The Core Facility supports users in the creation of ScenarioDesigner programs.

Visual System

The “Visual System” from NNL (Nordic Neuro Lab) consisting of video goggles and response buttons is available. The video goggles create the impression of a large image (like a cinema screen) and shield lateral influences. This allows the respondent to immerse himself more intensively in the scene and is less distracted. A correction of defective vision is directly integrated into the goggles.

LCD Monitor

Alternatively, the paradigms can be displayed on an LCD screen placed at the head-side magnetic tunnel entrance. The subjects see the image through a mirror attached to the head coil. A set of lenses is available to correct defective vision.

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**Expertise**

The Core Facility Mass Spectrometry offers services for a broad range of protein analyses. We apply a variety of mass spectrometric approaches to measure intact protein masses, identify proteins after proteolytic digestion (e.g. from polyacrylamide gels) or analyze post translational modifications. Modern mass spectrometers with high resolution and sensitivity allow analysis of complex samples such as cell lysates. These machines are also used for quantitative analyses with or without stable isotope labeling strategies like SILAC and TMT. Targeted quantification can be done for example if good antibodies are not available (multiplexed “next-generation Western Blot”). The data we provide can be very informative but also challenging, in particular the quantitative characterization of proteomes. Therefore, early consultation with us is essential to improve the chances for a successful mass spectrometric analysis. We need to discuss the strategy, scope, and pitfalls of the experiments. We collaborate closely with the Core Unit Bioinformatics in order to assess the data quality and apply statistical methods. Together, we will help you in finding biological knowledge in your proteomics datasets.

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**Instrumentation**

- **Thermo Orbitrap Lumos with ETD**
  The Orbitrap Lumos is a modern spectrometer that is extremely versatile: Classical measurements can be done with high sensitivity; label-based quantification is improved. It is also capable of better intact protein analyses, targeted measurements for selected proteins and the modern quantitative approach of Data Independent Acquisitions.

- **Hybrid instrument Thermo Orbitrap Velos**
  The Orbitrap Velos has two detectors: a fast low-resolution linear ion trap and a high resolution, high accuracy Orbitrap detector which can be used in parallel. This instrument is a workhorse for proteome analyses for medium complexity.

- Additionally, we use an *Autoflex III MALDI-TOF* and an *HCTultra ion trap* (both from Bruker) for special applications.

- **nanoLC systems**
  Three ultra-high performance liquid chromatographs for nanoflow (~300 nl/min) are used for coupling with mass spectrometers. These systems perform chromatographic separations very reproducibly and assure a stable delivery of analytes to the mass spectrometers. An Advion Triversa Nanomate nanospray robot can be used for automated delivery of small sample amounts for direct infusion.
Expertise

The Microscopy Core Facility provides services for imaging microscopy techniques for live and fixed cells and tissue sections. We also provide scientific and technical assistance for researchers to design experiments and to facilitate image acquisition and analysis. We also offer full service sample preparations and imaging for electron microscopy related projects.

The Core Facility is equipped with five wide-field microscopes, five confocal systems, and an electron microscope. We also have five analysis workstations.

The Core Facility has two sites, one in BMZ 2 and the other at the Neurocenter, with the instruments evenly distributed between the sites.

The light microscopes can be booked for an hourly fee after sufficient training. We are also happy to perform the measurements for you, if required. The electron microscope is operated by staff members as full service.

Instrumentation

Wide-field microscopes
- Color- and fluorescence-based automated whole slide scanning
- Automated high-content screening of multi-well plates
- Live-cell imaging
- Dual-camera ratiometric imaging
- STORM super-resolution

Confocal microscopes
- High-resolution live-cell imaging
- Whole-organ / Cleared tissue imaging
- Large-tissue section imaging
- FLIM and FCS measurements
- Two-photon imaging
- On-the-fly deconvolution super-resolution
- STED super-resolution

Electron microscopy
- Scanning EM (SEM)
- Scanning transmission EM (STEM)
- FIB-SEM 3D nanotomography
- Sample preparation
- Ultramicrotomy
- Immunogold labeling
- Pre- & post-embedding correlative light and electron microscopy (CLEM)

Image processing and analysis
- Fiji and CellProfiler free software
- Licensed Zeiss, Leica, and Visitron software
- Huygens and AutoQuant deconvolution
- Imaris 4D image visualization and analysis
- Neurolucida analysis software

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Expertise

The Core Facility Nanobodies at the University of Bonn offers diverse services based on alpaca or llama variable domains of heavy chain-only antibodies (VHHs), also called nanobodies, and is the first of its kind in Germany.

Nanobodies are highly specific single domain antibodies (ca. 15 kDa) that recognize their targets with affinities comparable to conventional antibodies. We can identify suitable nanobodies by phage display, produce them in bacteria, or express them in the cytosol of mammalian cells to visualize targets or alter protein function.

The ease of modification by genetic fusion, chemical or enzymatic conjugation makes Nanobodies versatile tools used in the clinic, for diagnostics or for research. They have been used for perturbation studies, microscopy, flow cytometry, mass cytometry, mass spectrometry, or non-invasive imaging methods such as immuno-PET. In structural biology Nanobodies are appreciated as chaperones facilitating protein crystallization.

We offer the customized immunization of alpacas or llamas with provided protein, as well as the identification and validation of specific nanobodies against your desired target. We also produce and (site-specifically) modify nanobodies to adjust them to your needs.

Methods

- **Scientific and technical advice for successful generation and usage of nanobodies**
  Contact us and make an appointment. We are happy to help get your nanobody project started!

- **Generation of custom-made nanobodies**
  Immunization of camelids with your antigen of choice and selection of binders by phage display

- **Validation of custom-made nanobodies**
  *In vitro* by ELISA or *in situ* by LUMIER assay

- **Recombinant expression of nanobodies**
  Production at multiple scales in bacteria and in eukaryotic cells (with human/mouse Fc part)

- **Chemical and chemo-enzymatic modification of nanobodies**
  Amine-reactive (NHS ester) or sulfhydryl-reactive (maleimide chemistry) labeling or site-specific Sortase A-mediated labeling by transpeptidation („sortagging“)

- **Custom services involving development, selection or use of nanobodies**
  Ask us, we will gladly discuss your specific idea involving nanobodies upon request!
Next Generation Sequencing

Expertise

Massive parallel sequencing, called Next Generation Sequencing (NGS), is becoming widely applied in many research projects. However, investment in maintenance of NGS systems as well as establishing different applications is expensive and time consuming. Therefore, the NGS Core Facility at the Institute of Human Genetics is dedicated to provide the research community in Bonn with access to state-of-the-art sequencing technologies and to help facilitate cutting-edge genomics research. The service comprises the following library preparation protocols:

- Whole-Genome Sequencing
- Exome Sequencing
- Whole-Transcriptome Sequencing (mRNA and total RNA-Seq)
- Gene Expression Profiling (3´-mRNA-Seq)
- Single Cell Sequencing (10X Genomics)
- Small RNA sequencing (miRNA-Seq)
- Targeted Sequencing (Amplicon)
- 16S Metagenomics
- ChIP Sequencing

The service is all-encompassing and covers all phases of a research project, from experimental design to data generation. The facility’s modern technical equipment enables sample preparation and sequencing for in-depth analysis of some few samples to high throughput omics-profiling of larger cohorts and guarantees an optimal balance between data output and costs.

Since 2019 the NGS Core Facility is the production site Bonn of the DFG-funded West German Genome Center.

Instrumentation

- Sample Preparation & Quality Control
  To ensure a high quality of NGS libraries modern technical devices are available for quantification and/or size selection (Tapestation 4200, Agilent; Qubit/Quant-iT, Thermo Fisher; Fragment Analyzer, Agilent; BluePippin, Sage Science), sample preparation (LE-220 plus, Covaris; Chromium Controller, 10X Genomics), and liquid handling (Biomek i7, Beckman Coulter).

- Sequencing Devices
  Due to a comprehensive equipment of sequencing devices, the NGS Core Facility is able to offer the most suitable sequencer for each project in terms of required data volume and costs. Currently, most of the short-read sequencing systems from Illumina (NovaSeq 6000, NextSeq 550, MiSeq, MiSeq and iSeq 100) are available to the scientific community at the Bonn Campus.

- Data Analysis
  Upon run completion, raw data are converted into standard-format FastQ and provided per download link. Quality control and comprehensive secondary data analysis is ensured through a strong collaboration with the Bioinformatics Core Facility Bonn.
Expertise

The Viral Core Facility (VCF) offers scientists of the Bonn Medical Faculty and external academic scientists (“non-profit organizations”) services for the generation of viral particles. Using these particles functions and properties of specific proteins can be studied by overexpression or down-regulation in various cell types.

For the production of our viruses we use HEK cell cultures, helper plasmids and viral DNA vectors. Depending on the application the viruses will be purified for in vitro or in vivo experiments.

Services

- **Viral vector service – Cloning**
  Adeno-associated virus (AAV) and Lentivirus (LV) can be generated and cloned upon request. We use standard cloning techniques; restriction-ligation approaches and Infusion Cloning. All newly made plasmid DNAs will be sequenced and tested for fluorescence activity.

- **Viral vector service – Purification AAV**
  For AAV replication we use helper plasmids that provide all the necessary adenoviral genes. Depending on the target cell type different serotypes of rAAV can be purified (serotype 1,2,5,8,DJ8,9). For the purification of new AAVs we use heparin columns or iodixanol gradient centrifugation and FPLC. We routinely test purified rAAV viruses for their physical and infective titer. On request we can also perform qPCR to determine the genomic titer of the virus.

- **Methods:**
  - Generation of endotoxinfree plasmid DNA for viral production
  - Cloning of new AAV or LV vector (simple or complex cloning strategies)
  - AAV purification (Heparin columns or gradient centrifugation and FPLC)
  - SDS-PAGE Coomassie staining and viral transduction test in primary neurons
  - Viral titer determination using qPCR

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Expertise

Zebrafish (danio rerio) and especially zebrafish larvae has become an important vertebrate model organism within the last 30 years. Now they became even more challenging with the invention of the CRISPR/Cas technology in the last few years. There are several reasons to use zebrafish as a vertebrate model: mainly the fast ex utero development of the small nearly transparent larvae fish, which makes them very approachable for whole mount (fluorescence) in vivo microscopy, as well as the fact that they can be raised in large numbers on relatively small amount of space and therefore costs.

Thousands of mutant and transgenic zebrafish (reporter) lines have been established and described already worldwide and are easily available to everyone free of charge (Small handling fees for shipping of the fish-eggs might apply). Furthermore it is easy for everybody to transiently manipulate the larvae fish by injections of RNA, Morpholino® antisense knock-down constructs or overexpressing plasmid DNA which can also be used to raise stable transgenic fishlines via germ line insertion within the next generation.

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Methods

The idea of our zebrafish core facility within the medical faculty of the University of Bonn is twofold:

- To give easy access, possibilities and help all researchers raise, maintain, bread and initially screen and genotype their zebrafish lines of interest within the fish facility (fishroom) of the newly established core facility.

- To provide expertise, equipment and help to all researchers who plan to do (or have done for them) transient injection experiments into fertilized fish eggs as well as larvae fish for generating and screening transient larvae fish research models as well as generating transgenic zebrafish lines (after application).

For this purpose our zebrafish core facility is equipped with:

- Freshwater fish-racks providing space for 200 three-liter fish tanks (~ 20 fish each)
- Breeding tanks for daily matings of up to 50 fish-pairs
- Wild type line zebrafish for matings to get fish eggs
- Two manual fully equipped air pressure fish egg injection setups (with stereo microscopes)
- Temperature and light controlled incubator for raising of fish embryos
- Fluorescence stereomicroscope for screening of embryos and larvae fish
Data Storage

Registration
Once you are registered for the Bonn Technology Campus and you are located in the network of the university or the Forschungsnetz at the Venusberg Campus you get...

Groups
Group Leaders can ask for additional storage, where the group can share data among the corresponding members.

One Terabyte of personal storage on our central Petabyte Storage cluster.

Cloud
and an additional 128 Gigabyte on our Cloud Installation.
https://btc-cloud.ukb.uni-bonn.de

Need Information?
Find more on our INTRANET pages:
https://community.ukb.uni-bonn.de

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Your Acknowledgement Matters

An acknowledgement in your publication is more than just a nice way to say „Thank you!“.

IT HELPS US TO MAINTAIN OPERATIONS
Showing to be part of the scientific community is one of the important measures to maintain funding and investments for core facilities.

IT IS ABOUT PEOPLE IN SCIENCE TOO
The core facility personnel may have contributed significantly to your research project. Authorship may also be appropriate.

Projektergebnisse, die aus mit DFG-Mitteln finanzierten Projekten resultieren, müssen in geeigneter Art ... einen Hinweis auf die DFG-Förderung (sog. „Funding Acknowledgement“) ... enthalten. [DFG-Vordruck 2.00 – 01/21, Verwendungsrichtlinien]

As an example: „We would like to thank the ... Core Facility of the Medical Faculty at the University of Bonn for providing support and instrumentation funded by the Deutsche Forschungsgemeinschaft (DFG, German Research Foundation) – Projektnummer …“
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