



# RBC: RNA Bioinformatics Center: Fkz 031A538C BIMSB The Berlin Institute for Medical Systems Biology

Bora Uyar, Dilmurat Yusuf\*, Ricardo Wurmus, Dan Munteanu, Lorenzo Calviello, Altuna Akalin, Nikolaus Rajewsky, Uwe Ohler The Berlin Institute for Medical Systems Biology, Max-Delbrueck Center for Molecular Medicine, Berlin, Germany \*Current affiliation: Bioinformatics Group, Department of Computer Science, Albert-Ludwigs-University Freiburg, Freiburg, Germany

## Short description of the project

We offer tools, services and training for the analysis of RNA-binding proteins and post-transcriptional regulation.

de.NBI services

|                 | Service          | Description   | Stats until April '17 |
|-----------------|------------------|---|-----------------------|
| Data-<br>bases  | Galaxy<br>Server | Galaxy server at MDC  | 82 unique users (UU)  |
|                 | doRiNA           | A database of RNA interactions  | 15680 UU              |
|                 | circBase         | A database to explore and discover circular RNAs                      | 34215 UU              |
| RNA (Seq) Tools | RCAS             | RNA-centric annotation system   | 652 downloads (Ds)    |
|                 | NASTIseq         | identifies cis-Natural Antisense Transcripts from RNA-seq data        | 417 Ds                |
|                 | RiboTaper        | Ribo-Seq analysis pipeline  | 1257 Ds               |
| miRNA<br>Tools  | PIPmiR           | identifies novel plant<br>miRNA genes from deep<br>sequencing data    | 586 Ds                |
|                 |                  | Web resource for microRNA targets                                     | 66158 UU              |
|                 | miRDeep          | microRNA detection tool in deep sequencing data                       | 5644 Ds               |
|                 | miReduce         | discovers motifs in mRNAs and correlates with gene expression changes | 34 Ds                 |
| RBP Tools       | PARalyzer        | PAR-CLIP sequencing analysis  | 1201 Ds               |
|                 | micro-<br>MUMMIE | microRNA target-site<br>prediction in PAR-CLIP<br>data                | added later           |
|                 | cERMIT           | finds optimal motifs in high-<br>throughput ChIP or RIP<br>datasets   | added later           |

#### General information on the project

- No. of staff paid from de.NBI grant: 2
- Other staff involved: 2

## **Progress report**

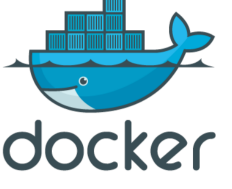
• Developed and maintained tools & databases made accessible via many channels











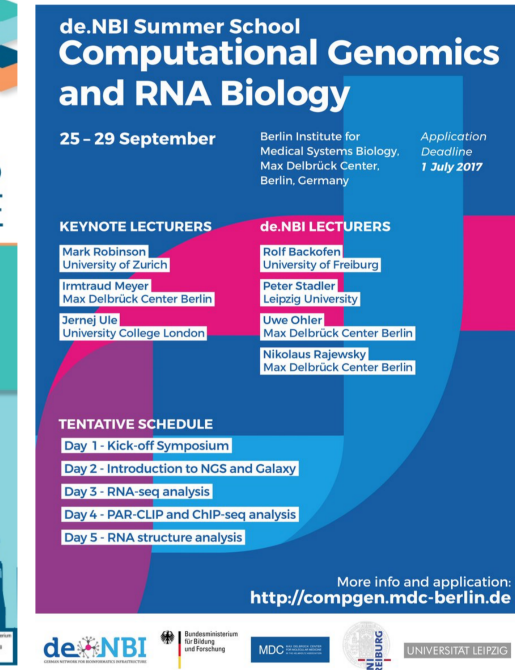
- Organised three summer schools
- o Organised a bioinformatics hackathon
- Organised three Galaxy Workshops
- o Participated in deNBI related events

## de.NBI Training and education

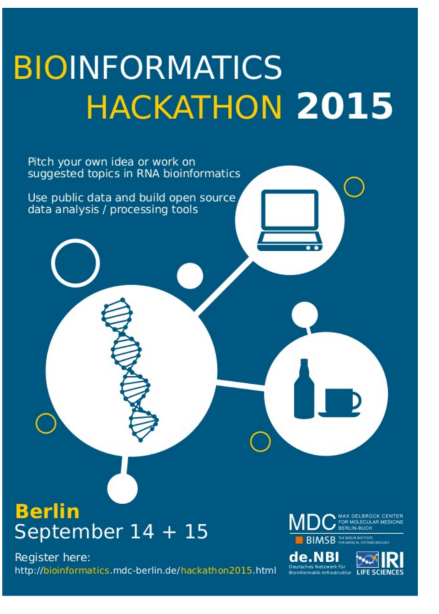
Summer schools (Total: 60 participants)







#### Bioinformatics Hackathon



15 participants

Also, offered
three workshops for
RNA-seq analysis
using Galaxy



Total: 55 participants

# **Selected Publications**

- O Calviello, Lorenzo, Neelanjan Mukherjee, Emanuel Wyler, Henrik Zauber, Antje Hirsekorn, Matthias Selbach, Markus Landthaler, Benedikt Obermayer, and Uwe Ohler. 2015. "Detecting Actively Translated Open Reading Frames in Ribosome Profiling Data." Nature Methods 13 (2). Nature Research: 165–70.
- O Uyar, Bora, Dilmurat Yusuf, Ricardo Wurmus, Nikolaus Rajewsky, Uwe Ohler, and Altuna Akalin. 2017. "RCAS: An RNA Centric Annotation System for Transcriptome-Wide Regions of Interest." Nucleic Acids Research, February. doi:10.1093/nar/gkx120.
- O Blin K, Dieterich C, Wurmus R, Nikolaus Rajewsky, Markus Landthaler, Akalin A. DoRiNA 2.0—upgrading the doRiNA database of RNA interactions in post-transcriptional regulation. Nucleic Acid Research, 2014





