# Tobias **Neumann**

#### BIOINFORMATICIAN · DATA ANALYST

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#### Personal information \_\_\_\_\_

**Date of Birth** December 17, 1986

Nationality Austria

Family status Married, 2 daughters

#### **Experience**

CONTRACTOR

Lexogen GmbH Vienna, Austria

• Lead development of the **SLAMdunk** product backend

- Optimizing and porting established variant-callers for nucleotide-conversion detection
- Containerization of software packages using Docker
- Deployment of Docker containers on the **Bluebee** private cloud platform
- · License assessment and resource benchmarking

#### **IMP Research Institute for Molecular Pathology**

Vienna, Austria

September 2017 - PRESENT

BIOINFORMATICIAN May 2014 - PRESENT

- Lead bioinformatics method and algorithm development for the SLAM-seq sequencing technology
- Design of genome-wide shRNA libraries for pooled RNAi barcode screens in human and mouse
- · Design of genome-wide transspecies sgRNA libraries for simultaneous CRISPR/Cas9 genome-editing in human and mouse
- Analysis and integration of RNA-seq, ChIP-seq, GRO-seq and PRO-seq data
- Software development of *palantir* for integrating, visualizing and querying raw and processed NGS data and metadata utilizing the on-campus sequencing facility API
- Data mining in public databases (COSMIC, TCGA)

Sophia Genetics SA Lausanne, Switzerland

BIOINFORMATICIAN October 2012 - May 2014

- Lead development, maintenance and optimization of a targeted gene sequencing and exome-sequencing pipeline building system forming the bioinformatics backend of the Sophia DDM® SaaS platform
- Tailoring pipelines for diagnostic gene panel kits on various sequencing platforms (Roche 454 / ionTorrent PGM / Illumina MiSeq) for major hospitals and labs in Switzerland and across Europe
- Conducting genetic test kit validation studies with key diagnostic kit developers at European level
- Development of genetic tests with Swiss hospitals
- Exploratory data analysis on prospective gene test data
- · Product presentations for prospective and established customers and on conferences

Max F. Perutz Laboratories Vienna, Austria

BIOINFORMATICIAN April 2012 - September 2012

- De-novo genome assembly of Clunio marinus
  - · Contig assembly, filtering and completeness assessment
  - Setup of the de-novo gene annotation pipeline and genome browser (MAKER, GMOD)
- Scaffold N50 of 1.9 Mb, 98% completeness
- Better assembly quality than honeybee, *Tribolium* or the monarch butterly
- Served as basis to study the genomic basis of circadian and circalunar timing adaptations
- Hosted at **ClunioBase** and published in *Nature*

#### **CIBIV Center for Integrative Bioinformatics Vienna**

Vienna, Austria

PROJECT STUDENT

June 2010 - February 2012

- · Parallelization of sequence alignments on the CPU and cluster
- Software development of an evaluation framework for reference-mapping software
- De-novo transcript assembly of RNA-seq data for *Idiosepius pygmaeus*

#### **SBC Stockholm Bioinformatics Center**

Stockholm, Sweden

 Network analysis of the gene interaction database FunCoup to in-silico identify and evaluate novel neurodegenerative disease candidate genes March 2009 - June 2009

## Education \_\_\_\_\_

RESEARCH INTERN

**Medical University Vienna** Vienna, Austria

DIPLOM-INGENIEUR (M.Sc.ENG.)

October 2009 - April 2012

- Master's program Medical Informatics (major: Bioinformatics)
- · Master's thesis: Challenges of de-novo assembling the transcriptome of Idiosepius pygmaeus for novel gene identification

#### **University for Applied Sciences Hagenberg**

Hagenberg, Austria

October 2006 - July 2009

BACHELOR OF SCIENCE IN ENGINEERING (HONORS)

• Bachelor's program Bioinformatics

**High school** Bad Ischl, Austria

A-LEVELS (DISTINCTION) June 2005

### Skills\_

NGS data Amplicon sequencing, genome-seq, RNA-seq, ChIP-seq, SLAM-seq, GRO-seq, PRO-seq

Sequencers Illumina GA / HiSeq / MiSeq, Ion Torrent PGM, Roche 454 **Programming** C/C++, Java, R / Bioconductor, Python, Perl, Bash, LaTeX

Technologies EJB / CDI, Hibernate / JPA, JSF + PrimeFaces, SSE/AVX, OpenCL, OpenMP, MPI

Platforms Linux (Ubuntu, CentOS, SUSE), Mac OS X, AWS, Docker, Windows

Databases Oracle 10g, MySQL, SQL, PL/SQL, XML

**Versioning** Git. Svn Building CMake, Maven

## **Teaching assistance**

**University of Vienna** Vienna, Austria

DARE WINTERSCHOOL TRAINER

February 2012

- · Lecturing on next-generation sequencing data processing
- Setting up and assisting students in practical excercises

**University of Vienna** Vienna, Austria

TEACHING ASSISTANT · Infrastructure setup Summer semester 2011

- · Student program review and testing
- Leading discussions during student presentations

### Languages \_\_\_\_

German Native

**English** Full professional proficiency **Spanish** Elementary proficiency Elementary proficiency French

#### Leisure activities\_

**Sports** Beachvolleyball, Cycling, Bouldering, Slacklining, Badminton

Music Piano, Guitar, Drums

**Others** Reading news, languages, travelling

## List of publications \_\_\_\_\_

- Muhar M, Ebert A, Neumann T, Umkehrer C, Jude J, Wieshofer C, Rescheneder P, Lipp JJ, Herzog VA, Reichholf B, Cisneros DA,

  Hoffmann T, Schlapansky MF, Bhat P, von Haeseler A, Köcher T, Obenauf AC, Popow J, Ameres SL\* & Zuber J: SLAM-seq defines direct gene regulatory functions of the BRD4-MYC axis. Science, 2018. 360(6390), 800–805. http://doi.org/10.1126/science.aao2793
- Herzog VA, Reichholf B, Neumann T, Rescheneder P, Bhat P, Burkard TR, Wlotzka W, von Haeseler A, Zuber J & Ameres SL:

  Thiol-linked alkylation of RNA to assess expression dynamics. *Nature Methods*, 2017. 14(12), 1198–1204. http://doi.org/10.1038/nmeth.4435
- Kaiser T, Poehn P, Szkiba D, Preussner M, Sedlazeck F, Zrim A, <u>Neumann T</u>, Nguyen LT, Betancourt A, Hummel T, Vogel H, Dorner S, Heyd F, Arndt von Haeseler A & Tessmar-Raible K: The genomic basis of circadian and circalunar timing adaptations in a midge. *Nature*, 2016. 540(7631), 69–73. http://doi.org/10.1038/nature20151
- Rathert P\*, Roth M\*, Neumann T, Muerdter F, Roe J-S, Muhar M, Deswal S, Cerny-Reiterer S, Peter B, Jude J, Hoffmann T, Boryn LM,

  Axelsson E, Schweifer N, Tontsch-Grunt U, Dow LE, Gianni D, Pearson M, Valent P, Stark A, Kraut N, Vakoc CR & Zuber J:

  Transcriptional plasticity promotes primary and acquired resistance to BET inhibition. Nature, 2015. 525(7570), 543–547.

  http://doi.org/10.1038/nature14898
- Matsushima W, Herzog VA, Neumann T, Gapp K, Zuber J, Stefan L Ameres SL & Miska EA: SLAM-ITseq: Sequencing cell type-specific transcriptomes without cell sorting. *Development*, 2018. (accepted)
- Fitz J, Neumann T, & Pavri R: Regulation of RNA polymerase II processivity by Spt5 is restricted to a narrow window during elongation. *Embo J*, 2018. http://doi.org/10.15252/embj.201797965
- Chipman AD, Ferrier DEK, Brena C, Qu J, Hughes DST [...], Neumann T et al.: The First Myriapod Genome Sequence Reveals

  Conservative Arthropod Gene Content and Genome Organisation in the Centipede Strigamia maritima. PLoS Biology, 2014. 12(11), e1002005. http://doi.org/10.1371/journal.pbio.1002005

#### Invited talks

## SLAMdunk – a pipeline for analyzing SLAMseq data in established and emerging applications

Orlando, FL, USA

ADVANCES IN GENOME BIOLOGY AND TECHNOLOGY (AGBT)

February 2018

## Conference posters.

## Quantifying experimentally induced nucleotide-conversions in high-throughput sequencing

Orlando, FL, USA

ADVANCES IN GENOME BIOLOGY AND TECHNOLOGY (AGBT)

February 2018

<sup>\*</sup> indicates equal contribution