Curriculum Vitae

KATHRIN M. SEIBT

PERSONAL INFORMATION

Full Name: Kathrin Martina Seibt

Date of Birth: 02.07.1984

(Görlitz, Germany)

Marital Status: unmarried

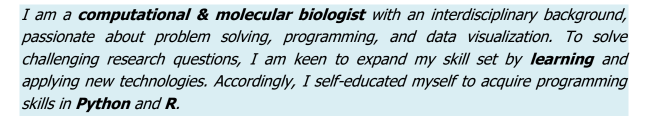
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GitHub: https://github.com/KathSe1984

Google Scholar: https://scholar.google.com/citations?user=5tWrgxgAAAA]



WORK EXPERIENCE

04/2012 - 10/2020

Technische Universität Dresden, Plant Cell & Molecular Biology (Prof. Thomas Schmidt/Prof. Stefan Wanke)

Topics

- BMBF projects "AVIMA" and "MASEPA" development of resistance-linked molecular markers for marker-assisted potato breeding
- transposable element genomics in Solanaceae & higher plants

Methods

- bioinformatics: comparative genomics, sequence analyses, development of analysis workflows in Python scripting, statistical analyses and data visualization in R, sequence analyses
- molecular biology: molecular markers, cloning, DNA sequencing

02/2009 – 03/2012 **potato breeder NORIKA GmbH**

Topics

• BMBF project "RetroKartoffel" – development of a marker system for potato cultivar differentiation

Methods

- molecular biology: plant DNA isolation, molecular markers, molecular cloning
- bioinformatics: sequence analyses, establishment of a fingerprint database, genotype clustering

11/2007 - 10/2008

Freiberg University of Mining & Technology,

Diploma thesis

Environmental Microbiology

(Prof. Michael Schlömann)

Topic

• "Cultivation and molecular characterization of sulfate-reducing bacteria from mining sites" (grade 1,2)

Methods

 molecular/microbiology: cultivation of anaerobic bacteria, characterization of substrate utilization, DNA isolation, molecular cloning, molecular markers, Sanger DNA sequencing

03/2007 – 11/2007 **Student project**

Freiberg University of Mining & Technology,

Environmental Microbiology (Prof. Michael Schlömann)

Topic

• "Bacterial diversity of a water sample from the uranium mine Königstein (Germany)"

Methods

 molecular/microbiology: DNA isolation, molecular cloning, molecular markers, Sanger DNA sequencing

08/2006, 08/2007

Medizinisches Labor Ostsachsen, Görlitz

Internship

(Medical Laboratory East Saxony; Dr. med. Roger Hillert)

Methods

- molecular biology: routine microbiological and molecular biological diagnostics like ELISA, PCR, agglutination tests
- microbiological routine techniques

EDUCATION

10/2008	Diploma degree Diplom-Naturwissenschaftler (diploma natural scientist, grade 1.4)
10/2003 – 09/2008	Course of studies "Angewandte Naturwissenschaft" (applied natural science) at Freiberg University of Mining & Technology (TUBAF; Germany) Specialization: biotechnology, environmental microbiology
2003	Abitur (grade 1.6) at Gymnasium 2 Annenschule, Görlitz (Germany) Specialization: mathematics, biology

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FURTHER EDUCATION AND PROFESSIONAL DEVELOPMENT

2019 MOOC on **open science** by Technische Universität Dresden

Science 2.0 and open research methods (Moving MOOC)

since 2013 **Bioinformatics and programming** MOOCs on Coursera platform

Bioinformatics algorithms [part 1] (University of California, USA; 14 weeks)

Bioinformatics: Introduction and methods (Peking University, China; 15 weeks)

Command line tools for genomic data science (John Hopkins University, USA; 4

weeks)

Computational molecular evolution (Technical University of Denmark,

Denmark; 6 weeks)

Introduction to interactive programming in Python

(Rice University, USA; 8 weeks)

Principles of computing [part 1 and 2] (Rice University, USA; 5 weeks each)

Learn to program: The fundamentals (University of Toronto, Canada; 10

weeks)

since 2013 MOOCs on data analysis und writing on Coursera platform

Data analysis and statistical inference (Duke University, USA; 8 weeks)

Pattern discovery in data mining (University of Illinois at Urbana-

Champaign, USA; 4 weeks)

Crafting an effective writer: Tools of the trade (Mt. San Jacinto College, USA; 6 weeks)

Creativity, innovation and change (Penn State University, USA; 4 weeks)

2015 – 2017 Graduate Academy courses at Technische Universität Dresden

Research data managements

Fundamentals in R

Reading strategies and techniques

Writing papers and theses in the life and natural sciences

Time and self-management for doctoral candidates

Enhance your visibility: Creating and developing your professional profile

2009 – 2015 **Bioinformatics and biology** lectures at Technische Universität Dresden

Medical microbiology (grade 1.0) Probability, statistics and programming

Clinical genetics (grade 1.3) for computational biology

Interaction bacteria plants (grade 1.0) Introduction to mathematical biology

Applied bioinformatics Genes and genomes

Programming for bioinformatics

2015/2018 **Good Manufacturing Practice**

Good Manufacturing Practice (GMP) (TU Dresden, Germany; 1 week)

GMP Statistics Course (Universität Leipzig, Germany; 3 days)

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BIOINFORMATICS SKILLS

I quickly learn to apply, interpret, and integrate new software into custom pipelines. Here, I am only listing tools I used frequently.

programming languages Python, R, RMarkDown (advanced)

UNIX tools, bash scripting (intermediate)

SQL, HTML (basic)

general sequence analysis Geneious, Staden package, EMBOSS tools, Galaxy platform

alignment and mapping Muscle, ClustalW, MAFFT, T-Coffee, Bowtie2

clustering MEGA, SplitsTree (network analysis), CAFE (k-mer based

clustering), SiLiX (similarity-based clustering), STRUCTURE

(population structure analysis)

sequence search BLAST, FASTA, nhmmer

NGS data analysis FastQC, Trimmomatic, TrimGalore!, samtools, bedtools,

SegMonk, edgeR

transposable element

annotation

RepeatExplorer, RepeatMasker, LTR harvest, LTR Finder,

SINE-Finder, and others

visualization MapChart, IGV, FigTree, FlexiDot, ggplot2

others handling various biological file formats (fastq, bed, gff,

sam, vcf,...)

GENERAL COMPUTATIONAL SKILLS

operating systems Windows, Linux

administrator at the TU Dresden lab responsible for setup

and maintenance of dual-boot computers

databases GelComparII (molecular marker fingerprint manager)

ARB (16S rDNA sequence database and analysis package)

Microsoft SQL Server

general software Microsoft Office

RStudio, Sublime, GitHub

Endnote, Mendeley

Adobe Photoshop, Inkscape

EXPERIMENTAL SKILLS

molecular biology DNA isolation, PCR, electrophoresis, molecular cloning,

molecular markers (ARDRA, T-RFLP, ISAP, STS), Southern hybridization, Sanger sequencing, preparation of plant chromosome spreads, fluorescent *in situ* hybridization

other biological techniques plant in vitro culture, grafting of potato plants, ELISA assay

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LANGUAGES

German native language

English fluent (Unicert III)

Italian conversational (Unicert I, Level B1)

Spanish basic

Latin Latinum (Abitur, grade 1,0)

OTHER SKILLS

drivers license German class B GxP (good scientific practice) basic knowledge

PUBLISHED SOFTWARE

<u>Seibt</u> KM, Schmidt T & Heitkam T (2018) *FlexiDot: Highly customizable, ambiguity-aware dotplots for visual sequence analyses.* **Bioinformatics** 34:3575-3577

GitHub: https://github.com/molbio-dresden/flexidot

RESEARCH PUBLICATIONS

Reiche B, Kögler A, Morgenstern K, Brückner M, Weber B, Heikam T, <u>Seibt</u> KM, Tröber U, Meyer M, Wolf H, Schmidt T & Krabel D: *Application of retrotransposon-based inter-SINE amplified polymorphism (ISAP) markers for the differentiation of common poplar genotypes.* Canadian Journal of Forest Research (accepted)

Maiwald S, Weber B, <u>Seibt</u> KM, Schmidt T & Heitkam T (2020) *The Cassandra retrotransposon landscape in sugar beet (*Beta vulgaris*) and related Amaranthaceae: Recombination and re-shuffling lead to a high structural variability.* **Annals of Botany** (doi: 10.1093/aob/mcaa176)

Seibt KM, Schmidt T & Heitkam T (2020) *The conserved 3' Angio-domain defines a superfamily of short interspersed nuclear elements (SINEs) in higher plants.* **Plant Journal** 101(3):681-699

Kögler A, **Seibt** KM, Heitkam T, Morgenstern K, Reiche B, Brückner M, Wolf H, Krabel D & Schmidt T (2020) *Divergence of 3' ends as a driver of short interspersed nuclear element (SINE) evolution in the Salicaceae*. **Plant Journal** doi:101111/tpj.14721

Diekmann K, <u>Seibt</u> KM, Muders K, Wenke T, Junghans H, Schmidt T & Dehmer KJ (2017) Diversity studies in genetic resources of Solanum spp. (section Petota) by comparative application of ISAP markers. **Genetic Resources and Crop Evolution** 64:1937-1953

Tomlekova N, Spasova-Apostolova V, Nacheva E, Stoyanova M, Teneva A, Petrov N, <u>Seibt</u> KM & Schmidt T (2017) *Genotyping of Bulgarian potato varieties by SINE-based ISAP markers*. **Comptes Rendus de l'Académie Bulgare des Sciences** 70:63-72

Seibt KM, Wenke T, Muders K, Truberg B & Schmidt T (2016) *Short interspersed nuclear elements (SINEs) are abundant in Solanaceae and have a family-specific impact on gene structure and genome organization.* **Plant Journal** 86:268-285

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- Schwichtenberg K, Wenke T, Zakrzewski F, <u>Seibt</u> KM, Minoche A, Dohm JC, Weisshaar B, Himmelbauer H & Schmidt T (2016) *Diversification, evolution and methylation of short interspersed nuclear element families in sugar beet and related Amaranthaceae species*. **Plant Journal** 85:229-244
- Menzel G, Heitkam T, **Seibt** KM, Nouroz F, Müller-Stoermer M, Heslop-Harrison JS & Schmidt T (2014) *The diversification and activity of hAT transposons in Musa genomes*. **Chromosome Research** 22:559-571
- <u>Seibt</u> KM, Wenke T, Wollrab C, Junghans H, Muders K, Dehmer KJ, Diekmann K & Schmidt T (2012) *Development and application of SINE-based markers for genotyping of potato varieties.* **Theoretical and Applied Genetics** 125:185-196

BOOK CHAPTERS

- Reiche B, Kögler A, Morgenstern K, Brückner M, Weber B, Heitkam T, **Seibt** KM, Tröber U, Meyer M, Wolf H, Schmidt T, & Krabel D (2020): "Anwendung des SINE-basierten Markersystems ISAP zur Identifizierung von Pappelklonen." Chapter in Thünen Report 76: Forstpflanzenzüchtung für die Praxis, M. Liesebach (ed.), pp 144-154
- Wenke T, <u>Seibt</u> KM, Döbel T, Muders K & Schmidt T (2015) *Inter-SINE amplified polymorphism (ISAP) for rapid and robust plant genotyping*. In: Batley J (ed) Plant Genotyping: Methods and Protocols. Springer Science+Business Media, New York, pp 183-192
- Seifert J, Erler B, <u>Seibt</u> K, Rohrbach N, Arnold J, Schlömann M, Kassahun A & Jenk U (2008) *Characterization of the microbial diversity in the abandoned uranium mine Königstein*. In: Merkel B & Hasche-Berger A (ed) Uranium, Mining and Hydrogeology. Springer, Berlin, pp 733-742

PUBLICATIONS IN PREPARATION

- Hartig N, <u>Seibt</u> KM, Schmidt T & Heitkam T: *The impact of allopolyploidization on RTE long interspersed nuclear elements (LINEs) in* Nicotiana.
- Schmidt N, Weber B, **Seibt** KM, Schwarzacher T, Schmidt T & Heitkam T: *The genomic organisation and environment of caulimoviruses in* Beta vulgaris *shed a new light on the endopararetrovirus ability to outlast silencing.*

GRANTS

2018

Conference travel grant by the German Academic Exchange Office (DAAD) for the "*Transposable elements meeting*" in Cold Spring Harbor, USA

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CONFERENCE TALKS

- **RepeatExplorer Meeting** 2019 (workshop on the RepeatExplorer pipeline for repetitive DNA analyses; České Budějovice, Czech Republic) "*The Angio-SINE superfamily with a conserved 3' domain is widely distributed across the Angiosperms and frequently associated with genes"*
- **Biopolis** 2018 (PhD conference; Dresden, Germany) "The Angio-SINE superfamily with a conserved 3' domain is widely distributed across the Angiosperms"
- iJaDe 2016 (conference for German-Japanese cooperation; Dresden, Germany)
 "Solanaceae short interspersed nuclear elements and their impact on gene structure
 and genome organization."

CONFERENCE POSTERS (LAST 5 YEARS)

- Maiwald M, Weber B, <u>Seibt</u> KM, Schmidt T & Heitkam T (2020) "*The highly recombinated landscape of Cassandra retrotransposons in* Beta vulgaris" Biopolis (Dresden, Germany) [presented by first author]
- **Seibt** KM, Schmidt T & Heitkam T (2019) "FlexiDot: Customize your dotplots for visual sequence analyses" Meeting of the GPZ group Cytogenetics (Dresden, Germany)
- **Seibt** KM, Schmidt T & Heitkam T (2019) "The conserved 3' Angio-domain defines a novel superfamily of short interspersed nuclear elements (SINEs) in higher plants" Meeting of the GPZ group Cytogenetics (Dresden, Germany)
- Schmidt N, Weber B, **Seibt** KM, Schwarzacher T, Schmidt T & Heitkam T (2019) "Endogenous pararetroviruses in the genome of sugar beet (Beta vulgaris)" Meeting of the GPZ group Cytogenetics (Dresden, Germany) [presented by first author]
- Heitkam T, Weber B, <u>Seibt</u> KM, Hoffmann J, Badstübner M & Schmidt T (2019) "*Taking the retro-ride: Long terminal repeat retrotransposons carry tandem repeats and disperse them through plant genomes*" Meeting of the GPZ group Cytogenetics (Dresden, Germany) [presented by first author]
- **Seibt** KM, Heitkam T & Schmidt T (2019) "*The conserved 3' Angio-domain defines a novel superfamily of SINEs in higher plants"* Crossroads between transposons and gene regulation (London, United Kingdom)
- **Seibt** KM, Schmidt T & Heitkam T (2019) "FlexiDot: Customize your dotplots for visual sequence analyses" Biopolis (Dresden, Germany)
- **Seibt** KM, Heitkam T & Schmidt T (2018) "The Angio-SINE superfamily with its conserved 3' domain is widely distributed across the Angiosperms" Transposable elements meeting (Cold Spring Harbor, USA)
- **Seibt** KM, Schmidt T & Heitkam T (2018) "SINEs contribute to gene evolution, regulation and genome rearrangement in Solanaceae plants" EMBO Workshop plant genome stability and change (IPK Gatersleben, Germany)
- Hübler N, **Seibt** KM, Schmidt T & Heitkam T (2018) "*The impact of allopolyploidization on retrotransposable elements in Nicotiana"*, Plant Science Student Conference (IPK Gatersleben, Germany) [presented by first author]

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- **Seibt** KM, Wenke T, Muders K, Truberg B & Schmidt T (2017) "SINEs in Solanaceae gene association and use as molecular markers" EMBL symposium: The mobile genome: Genetic and physiological impacts of transposable elements (Heidelberg, Germany)
- Weber B, **Seibt** KM, Hoffmann J, Ha HB, Bannack E & Schmidt T, Heitkam T (2017) "Where it starts and how it ends the evolution of tandem repeats within LTR retrotransposons" EMBL symposium: The mobile genome: Genetic and physiological impacts of transposable elements (Heidelberg, Germany) [presented by first author]
- **Seibt** KM, Wenke T, Muders K, Truberg B, Heitkam T & Schmidt T (2017) "SINEs in Solanaceae chromosomal localization, gene association and utilization as molecular markers" XIX international botanical congress (Shenzhen, China)
- **Seibt** KM, Wenke T, Muders K, Truberg B & Schmidt T (2017) "SINEs in Solanaceae genome organization, evolution and application as molecular markers" Biopolis (Dresden, Germany)
- <u>Seibt</u> KM, Wenke T, Muders K, Truberg B & Schmidt T (2015) "*Informative retrotransposon-based markers are facilitated by recent amplificational bursts and a dispersed genomic organization"* Plant Science Student Conference (IPB Halle, Germany)

TEACHING

2017	Supervision of a Bachelor thesis entitled: "New ISAP markers based on the widespread plant SINE family Au and their evaluation using reference potato genotypes"
2016 – 2020	Teaching during the practical course "DNA sequencing": Introduction to bioinformatics (theoretical background of BLAST, file formats, Galaxy platform, trimming of NGS data)
2012	Supervision of a school thesis "Besondere Lernleistung" (9 months) entitled: "Mobile DNA sequences as markers in crop breeding"
since 2011	Supervision of visiting students (high-school students, university students, guest students from USA)

PERSONAL INTEREST

Travelling, Board Games, Photography, Pilates, E-Learning, listening to Blinkist

Dresden, 23.11.2020

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