# dr. philippbayer

plant bioinformatician

### contact

philippbay@gmail.com github.com/philippbayer twitter.com/philippbayer

### languages

German: mother tongue English: fluent French & Japanese: advanced

## programming

Python Go, Perl, Bash, Java Ruby on Rails, HTML

# education

2012-2015 **PhD** Applied Bioinformatics

University of Queensland, Brisbane

Working in the Applied Bioinformatics group on the use of genotyping by sequencing to improve the genome assembly of canola. Thesis submission data: 23rd Septem-

ber 2015. Date of acceptance: 4th May 2016

2010-2012 Master of IT Bond University, Gold Coast

Graduated with Honours

2006-2009 **Bachelor of Science Biology**  University of Münster, Germany

Thesis: Analysis of splicing in two populations of marine plants using bioinformatic

approaches

# **experience**

### 2017–Current Hacky Hour Founder

UWA, Perth

Founded the Hacky Hour at UWA, a weekly get-together of researchers and staff working with programming and data, doubles as a help-desk for students with programming problems.

### 2017-Current Mozilla Open Science Leadership mentor

UWA, Perth

Mentoring open source programmers and researchers on how to streamline and grow open source and open science projects under the umbrella of Mozilla.

### 2016-Current EMBL-ABR Head of Nodes member, Open Science Special Interest Group mem-UWA, Perth

EMBL-ABR is an Australian-wide network supporting the technical needs of life sciences researchers. Members of the group of Head of Nodes meet monthly to discuss the way forward for the organisation. The Open Science Special Interest Group meets bimonthly to discuss how EMBL-ABR can advance open science in Australia.

### 2016–Current **Postdoctoral researcher**

UWA Perth

Researching the genetics of complex plants with a focus on canola and wheat. Working closely with industry partners to improve their breeding programs. Preparing, writing, and publishing research. Currently supervising two interns, co-supervising four PhD students and one MSc student. Supervising the local computational infrastructure and data management. Assisting other researchers. Started and continue to run the group's journal club.

### 2013-Current Software Carpentry and Data Carpentry instructor

Australia

Certified Software Carpentry and Data Carpentry instructor. Software Carpentry is a Mozilla/Alfred P. Sloan Foundation funded non-profit organization which teaches best programming practices (structured programming, reproducible research, version control etc.) in bootcamps to scientists around the world. Data Carpentry is a sister-organisation that focuses on teaching best data management practices.

2012-2012 Research exchange at Bayer CropScience Ghent, Belgium

For 4 weeks, worked on the assembly of the *Brassica napus* genome. Learned to work in a corporate science environment. Still regularly involved with Bayer CropScience and continue to collaborate with the company on many projects.

2011-Now Co-founder openSNP.org Germany/Australia

A project for costumers of genotyping companies like 23andMe to share their data with scientists around the world, for free.

Responsibilities:

- Partially wrote and still maintain the site's Ruby on Rails code-base
- Interact and manage with the community of 5000 users
- System administration of the site's servers

### awards

2014 **GRDC Travel Award** GRDC \$4000 travel cost scholarship 2014 **SAFS Travel Award** University of Queensland \$2500 travel cost scholarship 2011-2014 Two postgraduate scholarships University of Queensland For the work on genotyping by sequencing, covers tuition and living costs. First place in PLOS/Mendeley Binary Challenge 2012 Won with openSNP.org Won first price in a competition aimed towards the advancement of open science 2009-2011 **Master IT** Bond University 5x Top of class, 3x Vice-Chancellor's List for Academic Excellence, 1x IT Award Aca-

demic Excellence. Graduated with honours. Recipient of John Oglethorpe Medal for highest GPA of all IT students graduating that semester

# teaching

2016

2017	<b>Teaching Software Carpentry</b> Introduction to data manipulation using Python	Research Bazaar, Curtin University, Perth
2016	<b>Teaching and hosting Data Carpentry</b> UWA, Perth Hosted, planned, and set up the first Data Carpentry workshop at UWA, taught best data management practices	
2016, 2017	<b>University teaching</b> Co-taught and co-supervised SCIE4002, compubiomedical MSc students. Set up the computatio ticals.	, 0,

2016 **Teaching Software Carpentry** Research Bazaar, Murdoch University, Perth Taught an introduction to Python

**Software Carpentry** UQ, Brisbane Hosted, planned, and set up the first Software Carpentry workshop at UQ. Taught introduction to programming.

2013-2014 **Software Carpentry** Adelaide/Melbourne Assisted Software Carpentry bootcamp in Adelaide, taught basic to intermediate

Python as well as documentation and assisted at bootcamp in Melbourne.

2009-2011 **Tutoring** Tutored students in Intro to Programming (Java), Database Management (Ora-

cle/MySQL) and Networks & Applications, held several all-day refresher courses be-

fore exams

# public speaking

2017	<b>Presentation</b> Title: Towards better plant breeding at UW	COMBINE event, Perth
2017	<b>Presentation</b> Title: Improving Plant Breeding using KNet	Plant And Animal Genome conference, San Diego Miner
2016	<b>Presentation</b> Title: Towards a canola pan-genome: caut	CCDM, Curtin University ionary tales from the assembly bench
2015	<b>Presentation</b> Plant And Animal Genome conference, San Diego Title: Assessing and validating the amphidiploid genome of <i>Brassica napus</i> using genotyping by sequencing	
2014	<b>Presentation</b> Title: Assembling and validating the genome genotyping by sequencing	University of Queensland, GenGen Seminar Series ne of the <i>Brassica napus</i> using skim-based
2013	<b>Poster</b> Title: Genome Assembly Validation and Traing by Sequencing in Canola	Plant And Animal Genome conference, San Diego it Association using Skim Based Genotyp-
2012	<b>Presentation</b> Presented the work on openSNP, talked ab privacy implications	28th Chaos Communication Congress, Berlin pout the future of personal genomics and

# **publications**

- **Bayer, Philipp E**, Bhavna Hurgobin, Agnieszka A Golicz, Chon-Kit Kenneth Chan, Yuxuan Yuan, HueyTyng Lee, Michael Renton, Jinling Meng, Ruiyuan Li, Yan Long, et al. "Assembly and comparison of two closely related Brassica napus genomes" *Plant biotechnology journal (2017)*
- Conceived of the study, performed the majority of analysis steps, wrote the publication.
- Yuxuan Yuan, **Bayer, Philipp E**, Jacqueline Batley, and David Edwards "Improvements in Genomic Technologies: Application to Crop Genomics" *Trends in Biotechnology (2017). Elsevier Current Trends*
- Added several citations, added paragraphs, proof-read and error-corrected.
- Juan D Montenegro, Agnieszka A Golicz, **Bayer, Philipp E**, Bhavna Hurgobin, HueyTyng Lee, Chon-Kit Kenneth Chan, Paul Visendi, Kaitao Lai, Jaroslav Doležel, Jacqueline Batley, et al. "The pangenome of hexaploid bread wheat" *The Plant Journal (2017)*
- Streamlined the annotation pipeline.
- Parwinder Kaur, **Bayer, Philipp E**, Zbyněk Milec, Jan Vrána, Yuxuan Yuan, Rudi Appels, David Edwards, Jacqueline Batley, Phillip Nichols, William Erskine, et al. "An advanced reference genome of Trifolium subterraneum L. reveals genes related to agronomic performance" *Plant biotechnology journal (2017)*
- Created the advanced ordering of assembly, performed new gene prediction, analysed cultivar differences.
- Katarzyna Gacek, **Bayer, Philipp E**, Iwona Bartkowiak-Broda, Laurencja Szala, Jan Bocianowski, David Edwards, and Jacqueline Batley "Genome-wide association study of genetic control of seed fatty acid biosynthesis in Brassica napus" *Frontiers in Plant Science 7 (2016). Frontiers Media SA*
- Ran the genome-wide association study, interpreted results.

- Agnieszka A Golicz, **Bayer, Philipp E**, Guy C Barker, Patrick P Edger, HyeRan Kim, Paula A Martinez, Chon Kit Kenneth Chan, Anita Severn-Ellis, W Richard McCombie, Isobel AP Parkin, et al. "The pangenome of an agronomically important crop plant Brassica oleracea" *Nature Communications 7 (2016). Nature Publishing Group*
- Ran different orders of the *B. oleracea* assembly steps, ran annotation, helped with biological analysis and plotting.
- James K Hane, Yao Ming, Lars G Kamphuis, Matthew N Nelson, Gagan Garg, Craig A Atkins, **Bayer, Philipp E**, Armando Bravo, Scott Bringans, Steven Cannon, et al. "A comprehensive draft genome sequence for lupin (Lupinus angustifolius), an emerging health food: insights into plant–microbe interactions and legume evolution" *Plant biotechnology journal 15.3 (2017) pp. 318–330*
- Helped with setup of annotation and repeat pipeline.
- Mark Barash, **Bayer, Philipp E**, and Angela van Daal "Candidate gene scan for Single Nucleotide Polymorphisms involved in the determination of normal variability in human craniofacial morphology" bioRxiv (2016) p. 060814. Cold Spring Harbor Labs Journals
- Ran the entire genome-wide assocation study, interpreted results.
- HueyTyng Lee, Agnieszka A Golicz, **Bayer, Philipp E**, Yuannian Jiao, Haibao Tang, Andrew H Paterson, Gaurav Sablok, Rahul R Krishnaraj, Chon-Kit Kenneth Chan, Jacqueline Batley, et al. "The genome of a southern hemisphere seagrass species (Zostera muelleri)" *Plant Physiology (2016) pp–00868. American Society of Plant Biologists*
- Supervised part of the analysis,

Bayer, Philipp E "Genomics of Salinity" (2016) pp. 179–194. Springer New York

- Paul Visendi, Paul J Berkman, Satomi Hayashi, Agnieszka A Golicz, **Bayer, Philipp E**, Pradeep Ruperao, Bhavna Hurgobin, Juan Montenegro, Chon-Kit Kenneth Chan, Helena Staňková, et al. "An efficient approach to BAC based assembly of complex genomes" *Plant methods* 12.1 (2016) p. 2. BioMed Central
- Annaliese S Mason, Mathieu Rousseau-Gueutin, Jérôme Morice, **Bayer, Philipp E**, Naghmeh Besharat, Anouska Cousin, Aneeta Pradhan, Isobel AP Parkin, Anne-Marie Chèvre, Jacqueline Batley, et al. "Centromere locations in brassica A and C genomes revealed through half-tetrad analysis" *Genetics* 202.2 (2016) pp. 513–523. Genetics
- Analysed differential
- **Bayer, Philipp E** "Skim-Based Genotyping by Sequencing Using a Double Haploid Population to Call SNPs, Infer Gene Conversions, and Improve Genome Assemblies" *Plant Bioinformatics: Methods and Protocols* (2016) pp. 285–292. Springer New York
- **Bayer, Philipp E**, Pradeep Ruperao, Annaliese S Mason, Jiri Stiller, Chon-Kit Kenneth Chan, Satomi Hayashi, Yan Long, Jinling Meng, Tim Sutton, Paul Visendi, et al. "High-resolution skim genotyping by sequencing reveals the distribution of crossovers and gene conversions in Cicer arietinum and Brassica napus" *Theoretical and Applied Genetics* 128.6 (2015) pp. 1039–1047. Springer Berlin Heidelberg
- Agnieszka A Golicz, **Bayer, Philipp E**, and David Edwards "Skim-based genotyping by sequencing" *Plant Genotyping: Methods and Protocols (2015) pp. 257–270. Springer New York*

- Kaitao Lai, Michał T Lorenc, Hong Ching Lee, Paul J Berkman, **Bayer, Philipp E**, Paul Visendi, Pradeep Ruperao, Timothy L Fitzgerald, Manuel Zander, Chon-Kit Kenneth Chan, et al. "Identification and characterization of more than 4 million intervarietal SNPs across the group 7 chromosomes of bread wheat" *Plant biotechnology journal* 13.1 (2015) pp. 97–104
- Helped with
- Boulos Chalhoub, France Denoeud, Shengyi Liu, Isobel AP Parkin, Haibao Tang, Xiyin Wang, Julien Chiquet, Harry Belcram, Chaobo Tong, Birgit Samans, et al. "Early allopolyploid evolution in the post-Neolithic Brassica napus oilseed genome" Science 345.6199 (2014) pp. 950–953. American Association for the Advancement of Science
- Carried out genotyping by sequencing analysis, manually validated contig placement, compiled report of misassemblies.
- Annaliese S Mason, Jacqueline Batley, **Bayer, Philipp E**, Alice Hayward, Wallace A Cowling, and Matthew N Nelson "High-resolution molecular karyotyping uncovers pairing between ancestrally related Brassica chromosomes" *New Phytologist* 202.3 (2014) pp. 964–974. Wiley Online Library
- Performed the technical analysis and wrote the R-script.
- Bastian Greshake, **Bayer, Philipp E**, Helge Rausch, and Julia Reda "OpenSNP–a crowdsourced web resource for personal genomics" *PLoS One 9.3 (2014) e89204. Public Library of Science*
- Built large parts of the website, co-wrote the publication.
- Emanuela Dattolo, Jenny Gu, **Bayer, Philipp E**, Silvia Mazzuca, Ilia Anna Serra, Antonia Spadafora, Letizia Bernardo, Lucia Natali, Andrea Cavallini, and Gabriele Procaccini "Acclimation to different depths by the marine angiosperm Posidonia oceanica: transcriptomic and proteomic profiles" *Frontiers in plant science* 4 (2013) p. 195. Frontiers
- Ran EST-based differential expression analysis of seagrasses grown in different depths.