MAXIME BORRY | Curriculum Vitae

SIIMMARY

Bioinformatician - Life Data Scientist - Postdoctoral researcher at the Max Planck Institute for Evolutionary Anthropology and the Leibniz Institute for Natural Product Research and Infection Biology Hans Knöll Institute - Working on ancient DNA metagenomics

Enucation _____

FRIEDRICH-SCHILLER-UNIVERSITÄT

since 2018

PhD candidate in Bioinformatics

Jena, Germany

Software development and statistical data analysis with a focus in ancient DNA metagenomics

UNIVERSITÉ PARIS DIDEROT

2016 - 2018

MSc. Biology-Computer Science/Bioinformatics

Paris, France

Valedictorian

Python programming - R programming - Web programming - Databases - Algorithms -Genomics - Structural Biology - Structural Bioinformatics - Data analysis - Machine learning

UNIVERSITÉ GRENOBLE ALPES

2014 - 2015

MSc. Biology, Ecology, Environment

Grenoble, France

Populations genetic - Evolution - Biostatistics - Metabarcoding

UNIVERSITÉ PARIS DESCARTES

2011 - 2014 Paris, France

BSc. Interdisciplinary approaches to life sciences

Biology - Chemistry - Physics - Dynamic system Modelling - Perl programming - Research oriented projects - Phylosophy of Science - Research ethics - Editor of a scientific journalism publication

EXPERIENCE

POSTDOCTORAL RESEARCHER

Since 04/2023

Jena & Leipzig, Germany

Leibniz Institute for Natural Product Research and Infection Biology

Developing bioinformatics software and analyzing ancient metagenomics sequencing data of ancient fermentation samples.

DOCTORAL RESEARCHER 09/2021-03/2023

Max Planck Institute for Evolutionary Anthropology

Leipzig, Germany

I continue to develop and publish bioinformatics tools and pipelines in Python and Nextflow for ancient DNA metagenomics data, as well as my work on analysing ancient DNA metagenomics data.

DOCTORAL RESEARCHER 09/2018 - 08/2021

Max Planck Institute for the Science of Human History

Jena, Germany

I developed and published bioinformatics tools in Python and Nextflow for ancient DNA metagenomics data, as well as performed ancient DNA metagenomics data analysis that led to publications in peer-reviewed journals.

GRADUATE RESEARCH INTERN

01/2018 - 07/2018

Paris, France

Muséum national d'Histoire naturelle

bronze-age dental calculus samples.

Bioinformatics development and analysis for ancient DNA metagenomics. Nextflow pipeline development for dietary and microbial composition from metagenomics data of

LAB TECHNICIAN 10/2017 - 05/2018

Learning Planet Institute

Paris, France

Setting up a Lab Inventory Management System (LIMS) - Inventory management, ordering, and supplier contact point - Growing media preparation

GRADUATE RESEARCH INTERN 03/2017 - 05/2017 Institut Pasteur Paris, France

Development of ensemble methods for comparative gene expression analysis, with a R package and an interactive dashboard with Shiny

GRADUATE RESEARCH ASSISTANT 06/2016 - 08/2016

Centre for Genomic Regulation

Exploration of the oral mycobiome with Gabaldon's comparative genomics research group.

Barcelona, Spain

MASTER STUDENT INTERN 12/2015 - 02/2016

Max Planck Institute for Evolutionary Biology

Detection and functional study of gene duplication between Neanderthal and Mordern

Leipzig, Germany

GUEST RESEARCHER 05/2015 - 10/2015

Nature History, University of Oslo

Development of new markers and metagenomics software for plant identification. De novo and mapping assembly of Illumina data, phylogeny of the Anacyclus genus, and metabarcoding visualization software development.

Oslo, Norway

Gdansk, Poland

Bodø, Norway

RESEARCH INTERN 02/2014 - 06/2014

University of Gdansk

RESEARCH INTERN

Molecular biology internship: Expression of PIK3CA protein in insect cell lines using the BacToBac system.

09/2013 - 12/2013

Universitetet i Nordland

Benchmarking de novo assemblers on copepod's transcriptome sequencing data

RESEARCH INTERN 10/2012 - 11/2012

INRA Jouy en Josas, France

Investigating new recombinase families in Mycobacteriophages

SKILLS_

Meet-U

PROGRAMMING LANGUAGE **Experienced**: Python, Nextflow Familiar: R | JavaScript

LIBRARIES pandas | scikit-learn | statsmodels | scikit-bio | pysam | biopython LANGUAGES Native: French | Fluent: English (C2) | Conversational: German

HONORS & AWARDS

BEST POSTER PRIZE 01/2019

Société d'Anthropologie de Paris conference

"Analysis of the human periodontal microbiome from ancient DNA." Poster based on the research I conducted at the "Musee de l'Homme" in Paris in the Éco-anthropologie et Ethnobiology Lab during my master Thesis.

VALEDICTORIAN 07/2018

Université Paris Diderot

Valedictorian of the 2018 promotion of the BIB bioinformatics master

Paris, France

BIOINFORMATICS COMPETITION - WINNING TEAM

01/2018 Paris, France

Jena, Germany

Paris. France

Structural bioinformatics competition: conceive, design, develop, test and validate a computational program to predict the three-dimensional structure of a protein complex,

given the structures of the two monomeric partners.

XTRACURRICULAR ACTIVITIES _

PHD REPRESENTATIVE Max Planck Institute for the Science of Human History

06/2020 - 06/2021 PhD representative of the doctoral researchers of the Max Planck Institute for the Science

of Human History. I organized social events, and represented them at the instances of the Max Planck society.

BOARD MEMBER Paris Montagne Paris, France

10/2011 - 11/2012Member of the board and volunteer for this scientific popularization NGO offering research lab internship placements to high-school students

Teaching experience

MENTOR Since 05/2022

Seqera Labs Remote

Mentor for the nexflow bioinformatics mentoring program from the nf-coreorganization/Seqera Labs, with support from the CZI Science fundation.

LECTURER 08/2021

MPI-SHH IMPRS
Teaching a course on microbiome data analysis with a focus on microbial community

ecology

TEACHING ASSISTANT 09/2015

Petnica Science Center Petnica, Serbia

Teaching assistant for a molecular biology and ecology workshop for bachelor students.

TEACHING ASSISTANT 04/2015

Université Paris Descartes Paris, France

Teaching assistant on a workshop about thermodynamics for bachelor students.

PUBLICATIONS

- Martin Klapper, Alexander Hübner, Anan Ibrahim, Ina Wasmuth, Maxime Borry, Veit G. Haensch, Shuaibing Zhang, Walid K. Al-Jammal, Harikumar Suma, James A. Fellows Yates, Jasmin Frangenberg, Irina M. Velsko, Somak Chowdhury, Rosa Herbst, Evgeni V. Bratovanov, Hans-Martin Dahse, Therese Horch, Christian Hertweck, Manuel Ramon González Morales, Lawrence Guy Straus, Ivan Vilotijevic, Christina Warinner, Pierre Stallforth. Natural Products from Reconstructed Bacterial Genomes of the Middle and Upper Paleolithic. Science, 2023
- Maxime Borry, Alexander Hübner, Christina Warinner. sam2lca: Lowest Common Ancestor for SAM/BAM/CRAM alignment files. Journal of Open Source Software, 2022
- Maxime Borry, Alexander Hübner, Adam B. Rohrlach, Christina Warinner. PyDamage: automated ancient damage identification and estimation for contigs in ancient DNA de novo assembly. PeerJ, 2021
- Marsha C. Wibowo, Zhen Yang, Maxime Borry, Alexander Hübner, Kun D. Huang, Braden T. Tierney, Samuel Zimmerman, Francisco Barajas-Olmos, Cecilia Contreras-Cubas, Humberto García-Ortiz, Angélica Martínez-Hernández, Jacob M. Luber, Philipp Kirstahler, Tre Blohm, Francis E. Smiley, Richard Arnold, Sonia A. Ballal, Sünje Johanna Pamp, Julia Russ, Frank Maixner, Omar Rota-Stabelli, Nicola Segata, Karl Reinhard, Lorena Orozco, Christina Warinner, Meradeth Snow, Steven LeBlanc, Aleksandar D. Kostic. Reconstruction of ancient microbial genomes from the human gut. Nature, 2021.
- James A. Fellows Yates, Thiseas C. Lamnidis, Maxime Borry, Aida Andrades Valtueña, Zandra Fagernäs, Stephen Clayton, Maxime U. Garcia, Judith Neukamm, Alexander Peltzer. Reproducible, portable, and efficient ancient genome reconstruction with nf-core/eager. PeerJ, 2021.
- James A. Fellows Yates, Aida Andrades Valtueña, Åshild J. Vågene, Becky Cribdon, Irina M. Velsko, Maxime Borry, Miriam J. Bravo-Lopez, Antonio Fernandez-Guerra, Eleanor J. Green, Shreya L. Ramachandran, Peter D. Heintzman, Maria A. Spyrou, Alexander Hübner, Abigail S. Gancz, Jessica Hider, Aurora F. Allshouse, Valentina Zaro, Christina Warinner. Community-curated and standardised metadata of published ancient metagenomic samples with AncientMetagenomeDir. Scientific Data, 2021.
- Maxime Borry et al. CoproID predicts the source of coprolites and paleofeces using microbiome composition and host DNA content. PeerJ, 2020.
- Maxime Borry. Sourcepredict: Prediction of metagenomic sample sources using dimension reduction followed by machine learning classification. Journal of Open Source Software, 2019.