Mathias VANDENBOGAERT

Life Sciences Bio-Informatics – Scientific Data A.I. Analyst

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Plinkedin.com/in/mathiasvandenbogaert

EXPERIENCE

Bio-informatics Scientist

BIOLIZARD N.V.

Bioinformatics Scientist & Te

♀ Ghent, Belgium.

Bioinformatics Scientist & Technology Consultant as part of a data science consultancy company, involved in computational bulk and single-cell RNA, (meta-) genomics and proteomics data analyses.

• Bio-informatics Scientist

INSTITUT CURIE

2020 – 2023

Paris, France

Translational Immunotherapy Research. Integrative Functional Genomics of Cancer. Single-cell *scRNA-seq* & *scATAC-seq* data analysis. Proteomics: *immuno-peptidomics* mass-spectrometry data analysis.

• Bio-informatics Analyst

INSTITUT PASTEUR

2012 – 2020

Paris, France.

Pôle de Génotypage des Pathogènes (PGP) – Cellule d'Intervention de Biologie d'Urgence (CIBU) ; Unité de Recherche et Expertise « Environnement et Risques Infectieux » (URE-ERI). Microbial *meta-genomics* NGS data analysis.

Bio-informatics Analyst

INSTITUT PASTEUR

2006 -2012

Paris, France.

Unité de Biologie Systémique – Proteomics: Analysis of LC-MS/MS high-throughput data; Genomics: DNA-chip data analysis.

• Post-doc Computational Biology

ECOLE NORMALE SUPÉRIEURE

2005 –2006

Paris, France

Dynamics and Organization of Genomes -- Center for Computational Biology: Comparative Genomics to Identify Gene Transcription Regulation.

Post-doc Computational Biology

UNIVERSITY BASEL

2004 –2005

Pasel, Schweiz.

Computational Systems Biology -- Biozentrum & Swiss Institute of Bioinformatics: Identification of Genome-Wide Functional Regulatory Networks.

EDUCATION

Ph.D. Computational Biology

2000 - 2004

Laboratoire bordelais de Recherche en Informatique (LABRI, Université de Bordeaux I, France), and Institut National de Recherche en Informatique et Automatique (I.N.R.I.A. Rocquencourt, France).

 M.Sc. Computer Science for Biology Institut Pasteur, Paris, France. 2000

M.Sc. Biochemistry,

2000

Laboratory of "Protein Biochemistry and Protein Engineering" University Ghent, Belgium.

B.Sc., Biology

University Ghent, Belgium.

=== 1997



ACADEMIC & INDUSTRY EXPERTISE

Linux/Unix High-Performance Computing (HPC) & Data analysis & visualization

- Environments: Linux RedHat (CentOS), Unix FreeBSD 10.3/11.x, SunOS/OpenIndiana b151a9+.
- Programming languages: C, C++, Bash, Java, Perl, Python, Awk, R, GPU parallel programming with C/CUDA, OpenMPI.
- Cluster Management tools: PBS/SGE/SLURM, Pvfs2, Google cloud.
- Containers & Virtualization: Docker, Singularity.
- Software Design and development:
 Eclipse Cvs, Svn, Git MySql DB –
 Jupyter Snakemake / NextFlow (from scratch & Nf-Core)

Single-cell data analysis, 10X Genomics.

- scRNA-seq: Seurat, Scanpy, Scirpy
- scATAC-seq: ArchR, Signac
- single-cell lineage tracing: monocle, Slingshot

High-throughput data analysis

- Meta-genomics: Illumina, Ion Torrent Proton, Oxford Nanopore – workflow & pipeline development to production-level instances.
- Bulk RNA-seq data analysis: R universe & python 3.12 – bio-statistics for client-intimate consultancy commitments.

 $Proteomics \ / \ mass-spectrometry \ data \ analysis$

 DB Search engines: Mascot, Comet, MGSF+, MaxQuant Nf-core

Multi-omics data analysis

- 10X genomics: scRNA-seq / scATAC-seq
- Smart-seq2

A.I. Scientific data analysis:

 Large Language Models (LLM): Nucleotide Transformer (NT), DNABERT-2/S, HyenaDNA.

MOST SIGNIFICANT PUBLICATIONS IN THE LAST 5 YEARS

Hamy, Anne-Sophie, Judith Abécassis, Keltouma Driouch, Lauren Darrigues, <u>Mathias Vandenbogaert</u>, Cecile Laurent, Francois Zaccarini, et al. "Evolution of Synchronous Female Bilateral Breast Cancers and Response to Treatment." Nature Medicine 29, no. 3 (2023): 646–655. https://www.nature.com/articles/s41591-023-02216-8.

Balière, Charlotte, Elodie Calvez, Jean-Michel Thiberge, Somphavanh Somlor, Mathias Vandenbogaert, Marc Grandadam, and Valérie Caro. "A Six Years (2010–2016) Longitudinal Survey of the Four Serotypes of Dengue Viruses in Lao PDR." Microorganisms 11, no. 2 (January 18, 2023): 243. https://www.mdpi.com/2076-2607/11/2/243.

<u>Vandenbogaert M.</u> Kwasiborski A, Gonofio E, Descorps-Declère S, Selekon B, Nkili Meyong AA, Ouilibona RS, Gessain A, Manuguerra JC, Caro V, Nakoune E, Berthet N. Nanopore sequencing of a monkeypox virus strain isolated from a pustular lesion in the Central African Republic. Sci Rep. 2022 Jun 24;12(1):10768. doi: 10.1038/s41598-022-15073-1. PMID: 35750759; PMCID: PMC9232561.

Virginie Sauvage, Johanna Gomez, Laure Boizeau, Mathias Vandenbogaert, Léa Barbier, Claude Tayou Tagny, Aimée Olivat Rakoto Alson, Pascal Bizimana, Sekou Oumar Coulibaly, Mohamed Abdallahi Boullahi, Hadiza Soumana, Guy Mbensa, Valérie Caro, Syria Laperche. New insights into Human Pegivirus-1 (HPgV-1) genotypes diversity in sub-Saharan Africa. Infection, Genetics and Evolution, Volume 94, p. 104995, Oct 2021.

Ian Nunes Valença, Rafael dos Santos Bezerra, Kamila Chagas Peronni, Virginie Sauvage, <u>Mathias Vandenbogaert</u>, Valérie Caro, Wilson Araújo da Silva Junior, Dimas Tadeu Covas, Ana Cristina Silva-Pinto, Syria Laperche, Simone Kashima, Svetoslav Nanev Slavov. Deep sequencing applied to the analysis of viromes in patients with beta-thalassemia. Revista do Instituto de Medicina Tropical de São Paulo (vol. 63), May 2021.

Sauvage V, Boizeau L, Candotti D, <u>Vandenbogaert M</u>, Servant-Delmas A, Caro V, Laperche S. Early MinION™ nanopore single-molecule sequencing technology enables the characterization of hepatitis B virus genetic complexity in clinical samples. PLoS One. 2018 Mar 22;13(3):e0194366. doi: 10.1371/journal.pone.0194366. eCollection 2018.