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Téo Lemane

Education

2022 **Ph.D in computer science**, *University of Rennes 1, France*.

2019 Master's Degree in bioinformatic, with high honours, University of Rennes 1, France. Subjects: Algorithmic, sequence analysis, programming, semantic web, combinatorial optimization, machine learning, biological networks, structural biology

2017 **Bachelor's Degree: biology of organisms**, *University of Rennes 1, France*.

2013 High School Diploma, scientific option, Lycée André Malraux, Allonnes, France.

Research experiences

Postdoc Large-scale sequence abundances indexing.

Since May 2023 Supervisor: Eric Pelletier

CEA/Genoscope Development of bioinformatic methods and tools for large-scale indexing of biological sequences with abundances, LAGE team including powerful interfaces for end-users.

Ph.D thesis Indexing and analyses of large sequencing collections using k-mer matrices.

2019 - 2022 Supervisors: Pierre Peterlongo, Rayan Chikhi

Inria Rennes Development of bioinformatic methods for constructing k-mer matrices from large sequencing collections. This project GenScale team focused on making the computational tools necessary to scale up and enable indexing and downstream analyses of

these collections.

MSc2 Internship Search engine for genomic sequencing data.

January - July 2019 Supervisor: Pierre Peterlongo

> Inria Rennes Study and development of algorithms and methods for indexing sequencing data.

GenScale team Key words: Algorithmic, probabilistic data structures, indexing, C++ implementation, metagenomic

MSc1 Internship Curation of the metabolic network of *Tisochrysis lutea* using heterogeneous methods.

April - July 2018 Supervisor: Jeanne Got

> Inria Rennes Improvement of the metabolic network quality of Tisochrysis lutea, especially the metabolic pathways related to vitamin

Dyliss team

Key words: GEMs (Genome-scale metabolic models), flux balance analysis, topological analysis

Publications

2024 [preprint] Lucas Elliott, Frédéric Boyer, Téo Lemane, Inger Alsos, and Eric Coissac. wholeskim: Utilizing genome skims for taxonomically annotating ancient DNA metagenomes. September 2024 [preprint].

Beatriz Willink, Kalle Tunström, Sofie Nilén, Rayan Chikhi, Téo Lemane, Michihiko Takahashi, Yuma Takahashi, Erik I. Svensson, and Christopher West Wheat. The genomics and evolution of inter-sexual mimicry and female-limited polymorphisms in damselflies. Nature Ecology & Evolution, volume 8, pages 83-97, January 2024. Publisher: Nature Publishing Group.

2024 Téo Lemane, Nolan Lezzoche, Julien Lecubin, Eric Pelletier, Magali Lescot, Rayan Chikhi, and Pierre Peterlongo. Indexing and real-time user-friendly queries in terabyte-sized complex genomic datasets with kmindex and ORA. Nature Computational Science, volume 4, pages 104-109, February 2024. Publisher: Nature Publishing Group.

2023 Camila Duitama González, Riccardo Vicedomini, **Téo Lemane**, Nicolas Rascovan, Hugues Richard, and Rayan Chikhi. decOM: similarity-based microbial source tracking of ancient oral samples using k-mer-based methods. Microbiome, volume 11, page 243, November 2023.

2022 **Téo Lemane**, Paul Medvedev, Rayan Chikhi, and Pierre Peterlongo. kmtricks: efficient and flexible construction of Bloom filters for large sequencing data collections. Bioinformatics Advances, volume 2, page vbac029, January 2022.

- **Téo Lemane**, Rayan Chikhi, and Pierre Peterlongo. kmdiff, large-scale and user-friendly differential k-mer analyses. *Bioinformatics*, volume 38, pages 5443–5445, December 2022.
- 2022 Yoann Dufresne, **Téo Lemane**, Pierre Marijon, Pierre Peterlongo, Amatur Rahman, Marek Kokot, Paul Medvedev, Sebastian Deorowicz, and Rayan Chikhi. The K-mer File Format: a standardized and compact disk representation of sets of k-mers. *Bioinformatics*, volume 38, pages 4423–4425, September 2022.
- Grégoire Siekaniec, Emeline Roux, **Téo Lemane**, Eric Guédon, and Jacques Nicolas. Identification of isolated or mixed strains from long reads: a challenge met on Streptococcus thermophilus using a MinION sequencer. *Microbial Genomics*, volume 7, page 000654, 2021. Publisher: Microbiology Society,.

Talks

- 2022 **Téo Lemane**, Paul Medvedev, Rayan Chikhi, and Pierre Peterlongo. kmtricks: creating bloom filters for indexing large sequencing data collections. JOBIM, 2022.
- 2021 **Téo Lemane**, Rayan Chikhi, and Pierre Peterlongo. Advances in k-mer matrix construction for analysis of large sequencing collections. SeqBIM, 2021.
- 2020 **Téo Lemane**, Paul Medvedev, Rayan Chikhi, and Pierre Peterlongo. kmtricks: Modular k-mer count matrix and Bloom filter construction for large read collections. SeqBIM, 2020.
- 2020 **Téo Lemane**, Rayan Chikhi, and Pierre Peterlongo. HowDeSBT and Simka wedding: what has been done and what we plan to do? DSB, 2020.

Softwares

kmtricks Modular k-mer count matrix and Bloom filter construction for large read collections.

kmtricks is a generic tool suite along with a library for the construction of k-mer matrices and Bloom filters from large sequencing collections. It provides also a C++ plugin system to extend its features. Availability: $\mathbf{\Omega}$ tlemane/kmtricks

${\it kmdiff}$ Large-scale and user-friendly differential $k{\it -mer}$ analyses.

kmdiff is a tool allowing differential k-mer analysis on large cohorts (hundreds of individuals). It uses the concept of k-mer matrix to find differentially represented k-mers between two cohorts, opening the door to large-scale downstream analysis like structural variant detection.

Availability: O tlemane/kmdiff

kmindex Real-time k-mers query in TB-sized banks.

kmindex is a tool for real-time querying of sequencing samples indexed using kmtricks. It allows to compute the percentage of shared k-mers between a query and each indexed sample. It is the indexing backend of the ORA (https://ocean-read-atlas.mio.osupytheas.fr) web service.

Availability: **O** tlemane/kmindex

kmviz A generic web interface for exploring biological sequence indexes.

kmviz is a generic web interface and a REST API for interacting with sequence indexes. It connects to multiple, local or distant, sequence indexes to performs sequence queries. Results and metadata can then be explored through interactive tables, plots, sequence views, or maps.

Availability: tlemane/kmviz

Teaching

2019 - 2022 String algorithms and bioinformatics, MSc1 Software Engineering, Istic Univ. Rennes 1, France.

Computer skills

Systems GNU/Linux (CentOS, Arch, Debian, NixOS), basic knowledge of macOS

Programming Advanced: C, C++ and Python, Intermediate: JavaScript, Nix and Java

Perf Engineering Microbenchmarking (gbench, nanobench, hyperfine), Profiling (perf, gprofng, flamegraph), SIMD

CI/CD Git, Github Actions, GitLab CI, Jenkins

Packaging RPM, Nix, Conda, Poetry

Containerization Docker, Singularity, Apptainer, Applmage

Orchestration Docker Swarm, basic knowledge of Kubernetes

Workflow Nextflow, Snakemake

HPC Slurm, SGE

Cloud Microsoft Azure (Blob, Batch), basic knowledge of AWS

Languages

French Native English Advanced

Popular science

2021 Organization of "Sciences en cour[t]s".

http://sciences-en-courts.fr

2020 Production of an animated short film for "Sciences en cour[t]s".

http://sciences-en-courts.fr

Referees

Dr. Pierre Peterlongo

Research Associate, Head of GenScale team
Inria, Rennes, France

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Dr. Eric Pelletier

Research Associate
Genoscope, Évry-Courcouronnes, France

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Dr. Rayan Chikhi

Research Associate, Head of Sequence Bioinformatics team Institut Pasteur, Paris, France

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