



# **Metabolomics Symposium**

**Provence-Alpes-Côte d'Azur, France**

July 3rd and 4th, 2025  
Théâtre du Grand Château  
Parc Valrose, 06108 Nice

UNIVERSITÉ  
**CÔTE D'AZUR**

# Institutional and Industrial Partners

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# Program

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## Metabolomics Symposium Provence-Alpes-Côte d'Azur, France

3rd July 2025

Théâtre du Grand Château, Campus VALROSE, Université Côte d'Azur, 28 avenue de  
Valrose, 06108, Nice

08:30 - 09:00: Welcome and Registration

09:00 - 09:30: Opening Speech

- **Pr. Laurent COUNILLON**

Vice President of Research, Université Côte d'Azur

- Local Organizing Committee

**Pr. Mohamed MEHIRI**, ICN, CNRS, Université Côte d'Azur

**Dr. Delphine DEBAYLE**, IPMC, CNRS, Université Côte d'Azur

**Aurélié SEASSAU**, ISA, INRAE, Université Côte d'Azur

**Dr. Fabien FONTAINE-VIVE**, ICN, CNRS, Université Côte d'Azur

**Dr. Louis-Félix NOTHIAS**, ICN, CNRS, Université Côte d'Azur

### Session 1

09:30 - 10:00: Can internal exposome signatures predict lung cancer risk beyond smoking?  
Evidence from metabolomics and multi-omics studies

**Pr. Sonia DAGNINO**, UMR 4320 PHEN-X, CEA, Université Côte d'Azur

10:00 - 10:15: Search for metabolomic signatures to determine the exposome of honey bees  
(*Apis mellifera*), a colony health indicator

**Safae OUALI**, UMR 4320 PHEN-X, CEA, Université Côte d'Azur

10:15 - 10:30: L'origine environnementale des maladies chroniques et la médecine prédictive  
et préventive autour des biomarqueurs épigénétiques et métaboliques

**Dr. Mohamed BEN AHMED**, TECHINCARE

### 10:30 – 11:00: Coffee Break, Posters, and Discussions

### Session 2

11:00 - 11:45: Leveraging metabolomics to reveal how metabolic alterations drive colorectal  
cancer progression and enable patient stratification.

**Dr. Nathalie LEGRIVE**, Luxembourg Institute of Health, Department of Cancer Research

11:45 - 12:00: The Latest Innovations in Mass Spectrometry: The Orbitrap Astral

**Dr. Marie-Pierre PAVAGEAU**, Thermo Fisher Scientific

## 12:15 - 13:45: Lunch Break and Poster Session

### Session 3

13:45 - 14:15: Metabosmart, a Software for the Interpretation of Metabolomic and Multi-Omics Data Based on Multi-Block Data Segmentation Methods

**Dr. Jean-Charles MARTIN**, C2VN, INRAE, Aix-Marseille Université

**Dr. Abdoulaye SOW**, C2VN, INRAE, Aix-Marseille Université

14:15 - 14:45: High-Throughput Metabolomics and Lipidomics Workflows for Rare Disease and Cancer Research

**Dr. Djawed BENNOUNA**, NIH, NCATS/DPI, USA

14:45 - 15:00: Perspicacité-AI: An LLM-Powered Platform for Accelerating Scientific Research and Education

**Lucas PRADI**, ICN, CNRS, Université Côte d'Azur

## 15:00 - 15:30: Coffee Break and Discussions

### Session 4 – Workshop on Data Analysis

15:30 - 17:30: Data Processing and Spectral Annotation of LC-HRMS/MS Metabolomics Data

**Dr. Corinna BRUNGS**, University of Vienna, Austria

**Dr. Robin SCHMID**,  GmbH, Bremen, Germany

## 17:30: Symposium Closing Session



# Program

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## Metabolomics Symposium Provence-Alpes-Côte d'Azur, France

4<sup>th</sup> July 2025

Théâtre du Grand Château, Campus VALROSE, Université Côte d'Azur, 28 avenue de  
Valrose, 06108, Nice

### 08:30 - 09:00: Welcome and Registration

#### Session 5

09:00 - 9:45: Bioactive Natural Products: From Discovery to Chemo-diversification

**Dr. Emerson F. QUEIROZ**, University of Geneva, Switzerland

09:45 - 10:15: New Strategy in the Discovery of Natural Antituberculosis Agents from French Guiana Flora

**Dr. Elnur GARAYEV**, IMBE, Aix-Marseille Université

10:15 - 10:30: Exploration of the Antitubercular Potential of French Guiana's Flora through a Metabolomic Study

**Célia BREAUD**, IMBE, Aix-Marseille Université

10:30 - 10:45: Development of Innovative Natural Cosmetic Ingredients

**Dr. Camille DUBOIS**, ICN, CNRS, Université Côte d'Azur

10:45 - 11:00: UHPLC-DAD-nano ESI-HRMS/MS-DPPH/ABTS-VIS On-Line coupling for the detection of antioxidant compounds: a novel analytical approach applied to Provence flora species

**Clémentine ACHARD-BACCATI**, Aix-Marseille Université

### 11:00 – 11:15: Coffee Break, Posters, and Discussions

#### Session 6

11:15 - 11:45 : Discovery and quantification of lipoamino acids in bacteria: new partners in microbiote studies?

**Dr. Justine BERTRAND MICHEL**, MTH\_MetaToul, I2M, Toulouse

11:45- 12:15: Lipidomics with a broad coverage of lipid isomers

**Dr. Takeshi HARAYAMA**, IPMC, CNRS, Université Côte d'Azur

12:15 - 12:30: Latest Sciex Advances in Metabolomics and Lipidomics

**Heather CHASSAING**, Sciex

12:45 - 14:30: Inauguration of the CPER METABOLOME –  
Lunch

### Session 7

14:30 - 15:00: Characterizing marine benthic exometabolites from species to ecosystems  
**Dr. Charlotte SIMMLER**, IMBE, CNRS, IRD, Aix Marseille Université

15:00 - 15:30: Coffee Break and Discussions

### Session 8 – Workshop on Data Analysis

15:30 - 17:00: Comprehensive Analysis of Small Molecule MS/MS Data with SIRIUS  
**Dr. Kai DÜHRKOP**, Bright Giant GmbH

17:00 - 17:30: Symposium Closing Session

# Can internal exposome signatures predict lung cancer risk beyond smoking? Evidence from metabolomics and multi-omics studies

**Prof. Sonia DAGNINO**

UMR 4320 PHEN-X Physiopathologie, ENvironnement & eXposome  
Team EARTH, Directrice Scientifique de la Multi-Omiques, B. Rossi  
Université Côte d'Azur, CEA. Nice, France.



Lung cancer is the leading cause of cancer-related mortality worldwide, with over 1.8 million deaths annually. While tobacco smoking remains the main recognized risk factor, a significant proportion of lung cancer cases occur in non-smokers, especially among women. Environmental exposures beyond smoking remain insufficiently characterized, and internal exposome profiling offers a novel approach to detect early biological responses to environmental insults. In this work we investigate whether molecular signatures of internal exposome (proteomic, metabolomic, adductomics and epigenomic markers) are associated with lung cancer risk independently of smoking, and whether these signatures can serve as early biomarkers for precision prevention strategies. This work integrates data from two nested case-control studies within large prospective cohorts. Blood samples were analyzed using: (i) an inflammatory proteomics panel (92 proteins), (ii) genome-wide DNA methylation (485,512 CpG sites), and (iii) untargeted metabolomics (16,778 features). Multi-omics data were analyzed using univariate models, directed acyclic graphs (DAGs), and integrated network-based approaches to disentangle the contribution of smoking-mediated and independent molecular alterations.

We identified multiple omics markers significantly associated with lung cancer risk years before diagnosis. Only a subset of these biomarkers were associated with smoking, suggesting distinct exposome-driven pathways. Our findings support the presence of robust, early molecular fingerprints of environmental exposure linked to lung cancer development, independent of tobacco use. These exposome-informed biomarkers offer promising avenues for early detection and prevention, particularly in non-smokers.

## Search for metabolomic signatures to determine the exposome of honey bees (*Apis mellifera*), a colony health indicator

**Safae OUALI**

PhD in bioinformatics,  
Université Cote d'Azur (PHEN-X UMR 4320, Team EARTH)  
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Bees play a fundamental role in the balance of ecosystems, contributing to biodiversity and serving as bioindicators of environmental quality. However, in recent years, a worrying decline in bee populations has been observed worldwide. This phenomenon is attributed to various stress factors likely to weaken colonies and increase their mortality, including pesticides and pathogens. The links between these factors and the deterioration in bee health have been established, but the underlying molecular mechanisms remain poorly understood. Metabolomics offers great potential for the discovery of biomarkers and for enhancing our understanding of physiological disruptions at the molecular level. To better characterize the effects of pesticides exposure, controlled laboratory experiments and field exposures under natural conditions have been carried out. The aim of these studies is to identify biomarkers of bee metabolome capable of specifically signaling the cause of mortality, thereby adding to the targeted analyses. To achieve this, an analytical approach based on high-resolution mass spectrometry (HRMS) is being developed. Statistical analyses, including univariate and multivariate methods, revealed distinct metabolic signatures associated with pesticide exposure. Several biological pathways were significantly altered, highlighting key molecular disruptions linked to chemical stress. The results highlight the utility of metabolomics in identifying candidate biomarkers indicative of pesticide-induced stress in bees. This approach not only contributes to monitor and protect bee populations, but also to take advantage of their role as sentinel species, providing valuable information on the health of the ecosystem.



# L'origine environnementale des maladies chroniques et la médecine prédictive et préventive autour des biomarqueurs épigénétiques et métabolomiques

**Dr. Mohamed BENAHMED**

TECHiNCARE, 149, Promenade des anglais, Nice, 06000

TECHiNCARE est une Société de Biotechnologies dans les domaines de la Bonne Santé et de la Maladie. Elle est née des grandes ruptures conceptuelles et technologiques apparues au cours des vingt dernières années sur l'origine des Maladies Chroniques. En effet, l'apparition des Maladies Chroniques (Maladies cardiovasculaires, métaboliques, cancers, auto-immunes, neurodéveloppementale/neurodégénératives, les maladies de la Reproduction, ...) est liée à notre environnement et à notre mode de vie, comme les déséquilibres nutritionnels, l'exposition aux toxiques (eau, air, alimentation, ...) et au stress (psycho-social et psycho-émotionnel). Le lien entre les désordres de notre mode de vie et les modifications de l'expression de notre génome allant dans le sens de la bonne santé ou de la maladies est sous-tendu par des mécanismes épigénétiques. C'est ainsi que le séquençage du génome humain dans les années 2000-2003 est venu confirmé l'origine majoritairement (de l'ordre de 80%) épigénétique (modifications de l'«habillage» du génome) plutôt que génétique (modifications de la structure du génome) des maladies chroniques.

Ces ruptures conceptuelles ont été accompagnées de ruptures méthodologiques et technologiques qui permettaient pour la première fois de mesurer et de quantifier l'impact de l'environnement, au sens des modes de vie, sur notre génome.

La Société s'est ainsi fixé comme objectifs : (i) la production et l'utilisation d'une nouvelle génération de biomarqueurs diagnostiques, prédictifs, pronostiques et théranostiques, en particulier d'origine épigénétiques et métabolomiques dans le domaine de la Médecine 4P (Prédictive, Préventive, Personnalisée, Participative) et Intégrative (car intégrant en particulier l'approche nutritionnelle) et (ii) l'identification de nouvelles cibles épigénétiques des nutriments (Nutrition de Précision) en prévention de la Maladie mais aussi au cours de sa prise en charge.

# Leveraging metabolomics to reveal how metabolic alterations drive colorectal cancer progression and enable patient stratification

**Dr. Nathalie LEGRAVE**

Head of Metabolomics Platform, Metabolomics Platform  
Luxembourg Institute of Health  
Department of Cancer Research  
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Luxembourg



Colorectal cancer (CRC) is projected to have the highest global incidence and mortality by 2040, with rising rates of therapy resistance and an alarming increase among younger individuals. Despite advances in targeted and immune therapies, CRC's heterogeneity continues to limit universal treatment strategies. To address this, a deeper understanding of the mechanisms driving CRC progression is essential for improving patient stratification and enabling personalised care. Our recent work highlights metabolic rewiring as a key driver of disease evolution, with formate emerging as a metabolite of particular interest. Here, we present the implementation of a robust metabolomics strategy for large-scale cohort analysis applied to a well-characterised CRC cohort. This includes tumour and matched normal tissues, as well as longitudinal plasma samples from CRC patients and healthy controls. Our ongoing approach integrates clinical data, transcriptomic analysis and cytokine profiling with in-depth metabolomic and lipidomic analyses, with a particular focus on formate metabolism. This comprehensive multi-omics workflow is currently being developed to minimise technical variability, enhance analytical sensitivity, and uncover meaningful biological trends beyond statistical significance—particularly in longitudinal data. Additionally, our approach enables a multi-compartment analysis that expands biomarker discovery and provides mechanistic insight into metabolic reprogramming in cancer. With this strategy, we aim to demonstrate that the observable metabolic reprogramming in CRC, and the dysregulated formate metabolism, can serve as a powerful tool for patient stratification, predicting disease trajectories, and discovering novel therapeutic targets and strategies. We are also working toward developing predictive models based on combined molecular and metabolic features. The workflow we are developing in the context of CRC is broadly applicable to other disease cohorts, and we believe it holds real potential for advancing precision medicine.

## Advancing metabolomics with the novel Orbitrap Astral zoom<sup>™</sup> mass spectrometer

**Dr. Marie-Pierre PAVAGEAU**

Application Scientist Omics, South, Thermo Fisher Scientific



The Orbitrap Astral<sup>™</sup> is a powerful platform for metabolomics and lipidomics, combining speed, sensitivity, and depth of coverage. It is particularly suited for high-throughput biomarker discovery, systems biology, and clinical omics studies.

Untargeted metabolomics often struggles with accurate metabolite identification and quantification due to the lack of internal standards and comprehensive libraries. This limitation can hinder study design and data interpretation, leading researchers to focus on a small subset of analytes, potentially overlooking biologically relevant compounds.

To address this, the SQUAD (Single-injection Simultaneous Quantitation and Discovery) workflow combines accurate targeted quantitation with untargeted allowing the discovery of biologically unknown compounds.

The faster MS<sup>2</sup> scanning of the high-resolution MS Orbitrap Astral<sup>™</sup> enables sensitive quantitation of numerous metabolites through Astral PRM, simultaneously, ensuring robust unknown annotation and comprehensive sample coverage.

# Metabosmart, a Software for the Interpretation of Metabolomic and Multi-Omics Data Based on Multi-Block Data Segmentation Methods

**Dr. Jean-Charles MARTIN**, C2VN, INRAE, Aix-Marseille Université

**Dr. Abdoulaye SOW**, C2VN, INRAE, Aix-Marseille Université



Metabolomics generates complex data that can sometimes be difficult to interpret. We have developed software that enables the stratification of metabolic information into blocks (functional or statistical), thereby reducing dimensionality and facilitating the interpretation and representation of metabolic systems. The software uses R scripts and the R Shiny user interface. Annotated metabolites are grouped by the user into blocks according to functional ontology criteria of their choice, or into statistical clusters (using the WGCNA method). For each observation, a combined value (score) is calculated for each block using a hierarchical PCA, PLS, or OPLS method. The score matrices generated by each model are compared to determine which best matches the PCA of the initial data. Finally, classification techniques—including PLS-DA, OPLS-DA, logistic regression, and random forest—are applied to the selected matrix to identify the biological functions that are most determinant in the classification of individuals. Interactions between functions (blocks) can be visualized as an interaction network. The tool aims to optimize the interpretation of metabolomic data by leveraging advanced statistical modeling approaches. This method is also

applicable to lipidomics and multi-scale data grouped according to functional criteria, allowing them to be represented in the same normalized space, assembled, and compared.

# High-Throughput Metabolomics and Lipidomics Workflow for Rare Disease and Cancer Research

**Dr. Djawed BENNOUNA**

Research Scientist, Mathé Lab

National Center for Advancing Translational Sciences (NCATS) National Institutes of Health (NIH) U.S. Department of Health and Human Services (HHS) 9800 Medical Center Dr. Rockville, Maryland



Characterizing metabolic alterations in diseases such as cancer is critical for uncovering disease mechanisms and identifying biomarkers. To support this goal, our lab at the NIH has developed a high-throughput approach for metabolomics and lipidomics that combines automation, speed, and reproducibility. Our focus has been on building an automated and reproducible sample preparation and LC-MS workflow capable of processing hundreds to thousands of biological samples. Using plasma as a model matrix, we have optimized a biphasic extraction protocol for simultaneous recovery of polar and non-polar metabolites, preserving the advantages of biphasic extraction in terms of metabolic coverage, sensitivity, and reduced matrix effects. This protocol has been implemented in a 384-well format using automated liquid handling, allowing us to minimize variability and enable consistent, and large-scale sample processing. To improve analytical efficiency while maintaining resolution, we developed rapid LC-MS methods for both metabolomics and lipidomics. These methods showed robust performance across multiple plasma sources and analytical runs, with high retention time stability across timepoints and columns, ensuring reliable analysis across extended study durations and large sample sets. Beyond analytical development, we have also focused on improving metabolite annotation and biological interpretation. We introduced metScribeR, a user-friendly R/Shiny tool for building in-house compound libraries that combine retention time and m/z information, enhancing identification confidence by integrating publicly available MS/MS data without acquiring them experimentally. The tool also highlights potential identification conflicts—such as compounds sharing similar m/z and retention times—and incorporates the latest identification probability scoring method to prioritize the most likely compound assignments. Following metabolite identification, we also incorporate to our workflow RaMP-DB, a comprehensive relational database that integrates metabolite and gene annotations from multiple sources, to perform pathway and chemical class enrichment analyses. This step helps interpret metabolomic data by linking metabolite changes to relevant biological pathways and functions. This workflow—from automated extraction to compound annotation and pathway analysis—offers a robust platform for large-scale, reproducible metabolomics studies in cancer and other biologically complex conditions.

**Acknowledgement:** This research was supported in part by the Intramural/Extramural research program of the NCATS, NIH.



# Perspicacité-AI: An AI-Pipeline for Accelerating Scientific Education and Research

**Lucas PRADI**<sup>1</sup>, Tao Jiang<sup>1,2</sup>, Matthieu Ferraud<sup>1,2</sup>, Madina Bekbergenova<sup>1,2</sup>, Louis-Félix Nothias<sup>1,2\*</sup>

1 — Université Côte d'Azur, CNRS, ICN, Nice, France.

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Recently, emerging tools such as LLM have gained attention in scientific research and education. However, their limitations — notably outdated knowledge and untraceable provenance — hinder their ability to keep pace with rapid advancements in scientific fields, like metabolomics. Some of these challenges can be addressed by coupling LLM with Retrieval-Augmented Generation (RAG) techniques, which combine relevant data retrieval with generative AI. Here, we developed Perspicacité-AI, a free and open-source agentic literature assistant with advanced RAG capabilities designed for scientific research and education.

Perspicacité-AI required the development of the Bibt2KB python package, which seamlessly transforms open-access bibliographic references into a structured and searchable Facebook AI Similarity Search (FAISS) knowledge base (KB). The KB generated is used by the pipeline's framework to deploy an AI-assistant enhanced by RAG, ensuring that each question asked is grounded in relevant literature and documents. The framework features different advanced modes of document search and question-answering, including Perspicacité-Profound, an iterative, structured reasoning workflow suitable for answering complex scientific questions with the trackable facts and sources. Furthermore, the pipeline supports major open source and proprietary LLM providers, broadening access for different users.

Perspicacité-AI also incorporates a novel reranking function for document selection, Sigmoid Weighted RRF (SW-RRF). The benchmarking of SW-RRF showed improvement over the traditional Reciprocal Reranking Function (RRF) for all tested metrics: Recall, Precision, F1, Mean Reciprocal Rank (MRR), Average Precision (AP) and Normalized Discounted Cumulative Gain (NDCG) at 1, 3, 5 and 10 documents retrieved ( $p < 0.05$ ) at no computational expenses.

Initial applications of Perspicacité-AI in computational metabolomics [metaboguide.holobiomicslab.eu] demonstrated effective chatbot responses with accurate citations, and while still ongoing, initial evaluations of the pipeline show promising results, not only mitigating some of the limitations of standalone LLMs but empowering scientific literature with generative AI.

**Acknowledgement:** This work was supported by the French government through the France 2030 investment plan managed by the National Research Agency (ANR), as part of the Initiative of Excellence Université Côte d'Azur under reference number ANR-15-IDEX-01. Lucas Pradi is Supported by the Fonds National de la Recherche, Luxembourg, project 17994255.

# Computational Mass Spectrometry in **mzmine** - Hands-on Training



**Dr. Corinna BRUNGS**, University of Vienna, Austria  
Pharmacist and Analytical Chemist focused on reference data and plant metabolism  
Reference Data Project Lead [cbrungs1789@gmail.com](mailto:cbrungs1789@gmail.com)  
[GitHub](#) | [Twitter](#) | [LinkedIn](#)



**Dr. Robin SCHMID**, **mzIO** GmbH, Bremen, Germany  
Food Chemist and Analytical Chemist focused on computational mass spectrometry  
Lead Architect of mzmine <https://robinschmid.github.io>

This workshop will introduce non-target LC-MS<sup>2</sup> data processing workflows in mzmine. You will integrate feature detection, compound annotation, molecular networking, and statistical analysis. The new interactive molecular networking in mzmine clusters MS<sup>2</sup> fragmentation spectra by similarity reflecting the structural similarity of their underlying compounds. All the results from these workflows can be exported for downstream analysis in other popular tools like GNPS, SIRIUS, and statistical pipelines. The mzwizard aids in setting up workflows for various instruments and methods, e.g., for mass spectral reference library generation. We encourage you to bring your laptops for the best hands-on experience, but you can also take it as a live demo. Please download the provided dataset and install mzmine before the workshop.

## Dataset

The dataset is constituted of HRMS/MS data acquired over *Pseudomonas* strains. It contains wild type or mutant strains together with medium blanks and QCs. More background on the dataset is available at <https://doi.org/10.1111/1462-2920.15139>.

## Downloads

- Download the **dataset zip** file ~500 MB and unpack it.  
<https://drive.google.com/drive/folders/12X1fGBcTpkcfsAijlpNXQxjU484xFL6Y?usp=sharing>

- Open mzmine and register a **new user** for free or download the workshop user:  
<https://drive.google.com/file/d/1H03wfwFCZEvIFSRiBggcvfFkUIYNTrSy/view?usp=sharing>
- **Spectral library:**  
The Google Drive dataset **already** contains the MoNA (LC-MS<sup>2</sup> pos) library  
[Alternative link](#) if downloading from MassIVE

## Installation

- **mzmine:** Download and install the latest version - Current **mzmine 4.2.0**.  
<https://github.com/mzmine/mzmine/releases/latest>  
There are platform-specific installers for Windows, Mac, and Linux.  
There is **NO** need to install any other tool or Java Virtual Machine (JVM):  
Refer to the documentation for installation instructions and post issues on GitHub:  
<https://mzmine.org/documentation/>
1. **mzmine paper:** <https://www.nature.com/articles/s41587-023-01690-2>
  2. Development: <https://github.com/mzmine/mzmine/>
  3. Documentation: <https://mzmine.org/documentation/>
  4. Protocol: <https://doi.org/10.26434/chemrxiv-2023-98n6q-v2>
  5. YouTube: <https://www.youtube.com/@mzmineproject/playlists>

# Bioactive Natural Products: From Discovery to Chemo-diversification

Plinio Scapozza<sup>1,2</sup>, Marion Zwingelstein,<sup>3</sup> Robin Huber<sup>1,2</sup>, Laurence Marcourt<sup>1,2</sup>, Emilie Michellod<sup>3</sup>, Jean-Luc Wolfender<sup>1,2</sup>, Katia Gindro<sup>3</sup>, and **Emerson F. QUEIROZ**<sup>1,2</sup>

<sup>1</sup> School of Pharmaceutical Sciences, University of Geneva, CMU - Rue Michel-Servet 1, 1211 Geneva 4, Switzerland.

<sup>2</sup> Institute of Pharmaceutical Sciences of Western Switzerland (ISPSO), University of Geneva, CMU – Rue Michel Servet 1, 1211 Geneva 4, Switzerland.

<sup>3</sup> Agroscope, Plant Protection Research Division, Route de Duillier 50, P.O. Box 1012, 1260 Nyon, Switzerland



Over the past decade, we have witnessed a revolution in the methodologies applied to natural product research. Current approaches combine powerful metabolite profiling methods for compound annotation and prioritization. Targeted isolation is performed using high-resolution chromatographic methods that closely match those obtained for analytical profiling [1]. Thanks to these tools, minor bioactive compounds were identified. However, in plant extracts, the major compounds are generally common structures, apparently irrelevant for drug discovery. In this context, biotransformation could be an alternative to valorize them. For this proposal, we employ two biotech approaches: reactions using fungal secretomes (mixture of enzymes), and biotransformation using living organisms (whole-cell biotransformation). Saprophytic fungi, such as *Botrytis cinerea* and *Trametes versicolor*, were used as a source of enzymes [2]. Stilbenes, chalcones, phenylpropanoids, and terpenes from the Swiss flora were used as substrates. Biotransformations were first conducted at an analytical scale and monitored by UHPLC-PDA-ELSD-HRMS for the detection of unusual features. Promising reactions were scale-up at the gram scale, and high-resolution preparative chromatography combined with dryload was used for their purification [1]. Enantiomers were purified by chiral chromatography. HRMS, 2D NMR, and ECD were used for structural elucidation. Compounds were evaluated for their antibacterial and antiviral activity against relevant targets. A library of over 280 compounds was generated at the mg scale from common NPs. Some of the compounds obtained presented unique scaffolds and potent biological activities [3]. In most cases, it was possible to propose the enzymes and mechanisms involved in the synthesis of each compound [4]. The applications, possibilities, and limitations of these latest technologies will be illustrated with recent investigations carried out in our laboratory.

[1] Queiroz EF, Guilleme D, Wolfender JL. Advanced high-resolution chromatographic strategies for efficient isolation of natural products from complex biological matrices: from metabolite profiling to pure chemical entities. *Phytochem Rev* 2024; 23: 1415-1442; DOI; 10.1007/s11101-024-09928-w.

[2] Huber R, Marcourt L, Koval A, Schnee S, Righi D, Michellod E, Katanaev VL, Wolfender JL, Gindro K, Queiroz EF. Chemoenzymatic synthesis of complex phenylpropanoid derivatives by the *Botrytis cinerea* secretome and evaluation of their wnt inhibition activity. *Frontiers in Plant Science* 2021; 12: 805610; DOI; 10.3389/fpls.2021.805610.

[3] Huber R, Marcourt L, Heritier M, Luscher A, Guebey L, Schnee S, Michellod E, Guerrier S, Wolfender JL, Scapozza L, Kohler T, Gindro K, Queiroz EF. Generation of potent antibacterial compounds through enzymatic and chemical modifications of the *trans*-delta-viniferin scaffold. *Scientific Reports* 2023; 13,: DOI; 10.1038/s41598-023-43000-5.

[4] Huber R, Marcourt L, Félix F, Tardy S, Michellod E, Scapozza L, Wolfender JL, Gindro K, Queiroz EF. Study of phenoxy radical couplings using the enzymatic secretome of *Botrytis cinerea*. *Front Chem* 2024; 12: DOI; 10.3389/fchem.2024.1390066.

# New Strategy in the Discovery of Natural Antituberculosis Agents from French Guiana Flora

**Dr. Elnur GARAYEV**

Associate Professor

Aix Marseille Univ, CNRS 7263, IRD 237, Avignon Université, IMBE, 27 Blvd Jean Moulin,  
Service of Pharmacognosy, Faculty of Pharmacy, 13385 Marseille, France



Nowadays, approximately 60% of the world's population uses herbal medicines, and plants are recognized as a source of therapeutically effective drugs. Natural products are an important source of new structural scaffolds and historically have always been a privileged source of inspiration in drug discovery. Tuberculosis (TB) is one of the top 10 world's leading causes of death despite 100 years of vaccination and 60 years of antibiotherapy. The region the most affected of France is French Guiana with 3-times more notification rate compared to the whole country. As French Guiana is also one of the world's biodiversity spots, the local people in French Guiana have long used plants to treat the TB symptoms.

In this context, combination of the traditional ethnopharmacology with new metabolomics and bio/chemoinformatics approaches are used to foster the detection of bioactive compounds and select the appropriate samples from which further isolation/purification will be performed.

The ethnopharmacological approach represents the first filtering step, focusing only on the plants with already reported anti *Mycobacterium tuberculosis* activity and/or used in traditional medicine in the treatment of TB symptoms. Bio/cheminformatics approaches by molecular network and *in silico* pharmacophore screening represent the second and the third filtering steps, allowing to decrease the number of compounds to purify and avoid unnecessary and time-wasting purification.



# Exploration of the Antitubercular Potential of French Guiana's Flora through a Metabolomic Study

**Célia BREAUD**

PhD student in Natural Products Chemistry

Aix Marseille Univ, CNRS 7263, IRD 237, Avignon Université, IMBE, 27 Blvd Jean Moulin,  
Service of Pharmacognosy, Faculty of Pharmacy, 13385 Marseille, France



Tuberculosis, a bacterial infection caused by *Mycobacterium tuberculosis*, is still one of the world's deadliest infectious diseases. With the rise of multidrug-resistant strains, the need for new therapeutic options is more urgent than ever. Natural products remain a key source of bioactive molecules, but their chemical complexity often makes it difficult to identify the compounds responsible for biological activity.

In this work, we applied an integrative metabolomics strategy, to explore the chemical composition and antitubercular properties from 80 extracts and fractions, obtained from seven medicinal plants collected in French Guiana. We combined high-resolution LC-MS/MS, bioactivity-guided Feature-Based Molecular Networking, and *in vitro* screening against *Mycobacterium tuberculosis* H37Ra. The most active samples were non polar extracts from *Zingiber zerumbet*, *Tetradenia riparia*, and *Indigofera suffruticosa*. By integrating spectral data, taxonomic information, and antitubercular screening results into a single molecular network, we could quickly associate specific metabolite families with the observed bioactivity. Methoxylated flavonoids and one sesquiterpene emerged as the main contributors, and their predicted activity was validated with commercial standards, showing micromolar MICs (Minimum inhibitory concentration).

This approach allowed us to distinguish active compounds from structurally similar, inactive analogues, thus saving time and avoiding redundant isolations. This study highlights the power of combining untargeted metabolomics with molecular networking to accelerate the discovery of natural antitubercular agents.

# Development of Innovative Natural Cosmetic Ingredients

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Cosmetics is a key sector of the French industry actually, strongly influenced by the pursuit of innovation, the "natural" movement, and the growing trend toward more responsible consumption [1]. The diversity of plant biodiversity has consistently been a source of inspiration for the development of cosmetic ingredients, aligning with the current trend toward a "green" and sustainable approach. Nissactive, in collaboration with the MVBV group of the Institute of Chemistry of Nice, is specialized in the development of cosmetic actives derived from Mediterranean natural sources and/or upcycled materials [2,3].



Over the years, various resources derived from Mediterranean biodiversity and/or upcycling have been investigated for their potential in cosmetic applications. Among them, the strawberry tree (*Arbutus unedo*) is a representative example of Mediterranean resources, identified as a potential source of cosmetic active compounds [4]. Among the various parts of the plant (buds, bark, leaves, flowers, etc.), two extracts—one from the leaves and one from the bark—were studied in greater detail to identify the active compounds. The experimental conditions for eco-extraction techniques were optimized to obtain the most active and promising extracts for the cosmetic industry [5].

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## Discovery and quantification of lipoamino acids in bacteria: new partners in microbiote studies?

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Improving knowledge about metabolites produced by the microbiota is a key point to understand its role in human health and disease. In a first study we characterized lipoamino acid (LpAA) containing asparagine and their derivatives by a semi targeted approach. We demonstrated that these bacterial metabolites could have an impact on the host and could have a potential analgesic effect. We will present here the second study where our aim was to extend the characterization of this family. We developed a semi-targeted workflow to identify and quantify new candidates. First, the sample preparation and analytical conditions using liquid chromatography (LC) coupled to high resolution mass spectrometry (HRMS) were optimized. Using a theoretical homemade database, HRMS raw data were manually queried. This strategy allowed us to find 25 new LpAA conjugated to Asn, Gln, Asp, Glu, His, Leu, Ileu, Pro, Lys, Phe, Trp and Val amino acids. These metabolites were then fully characterized by MS<sup>2</sup>, and compared to the pure synthesized standards to validate annotation. Finally, a quantitative method was developed by LC coupled to a triple quadrupole instrument, and linearity and limit of quantification were determined. 14 new LpAA were quantified in gram positive bacteria, *Lactobacillus animalis*, and 12 LpAA in *Escherichia coli* strain Nissle 1917. First data about the quantification of these metabolites, completed by other metabolites (short chain fatty acid, bile acids) in a model of irritable bowel syndrome in prenatally stressed mice and in patients will be presented.

## Lipidomics with a broad coverage of lipid isomers

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Lipidomics is used to detect and quantify lipids comprehensively. It is an essential tool for fundamental lipid biology as well as applied research to find biomarkers or novel disease mechanisms. Lipidomics remains a method that requires a high degree of expertise, as the interpretation of chromatographic peaks and MS/MS spectra done by lipidomics software contains multiple errors that require extensive data curation. Via the extensive analysis of lipid standards artificially generated in live cells, we are proposing approaches to solve existing issues, improving the accuracy of lipidomics for a broader usership. I will describe the current developments of our approaches, as well as its future directions.

## Latest SCIEX Advances in Metabolomics and Lipidomics: Unleashing the Power of the ZenoTOF 8600

**Dr. Heather CHASSAING**

Sciex, Luisant, France



Metabolomics and lipidomics continue to push the boundaries of biological discovery, demanding ever-greater sensitivity, robustness, and data quality from analytical platforms. In this talk, we will explore the latest advances from SCIEX, focusing on the groundbreaking capabilities of the newly launched **ZenoTOF 8600** system, introduced at ASMS 2025. Built upon the proven architecture of the ZenoTOF platform, the 8600 represents a leap forward in performance—delivering **over 10x increased sensitivity** compared to the ZenoTOF 7600, enabling the detection of low-

abundance metabolites and lipids with unprecedented clarity and confidence.

Key highlights include the integration of **ZTScan DIA**, a novel data-independent acquisition (DIA) workflow designed for improved deconvolution of spectra from non-targeted data acquisition. ZTScan DIA provides broad coverage with exceptional quantitative reproducibility, empowering researchers to profile complex biological matrices with greater depth and reliability. Moreover, the ZenoTOF 8600 exhibits exceptional **robustness and stability**, supporting high-throughput studies with minimal downtime and consistent performance across large sample cohorts.

This presentation will showcase real-world applications and comparative performance data, demonstrating how the ZenoTOF 8600 is redefining the capabilities of mass spectrometry in metabolomics and lipidomics research. Join us to discover how this next-generation platform is accelerating the path to new biological insights and translational breakthroughs.



# UHPLC-DAD-nano ESI-HRMS/MS-DPPH/ABTS-VIS On-Line coupling for the detection of antioxidant compounds: a novel analytical approach applied to Provence flora species

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Conventional antioxidant screening methods, such as well-plates assays using DPPH or ABTS reagents, provide a global evaluation of antioxidant activity through IC<sub>50</sub> determination, but do not allow the identification of individual bioactive compounds. This study presents an on-line coupling methodology that integrates UHPLC-DAD-HRMS/MS with real-time DPPH/ABTS-VIS detection, enabling simultaneous chromatographic separation, high-resolution mass spectrometric compound identification, and antioxidant activity profiling, within a single run. By eliminating the need for time-consuming fractionation and post-fractionation bioassays, this method enhances the efficiency to identify antioxidant components in a complex mixture, and considerably reduces analysis time.

This approach was applied in a comparative study of antioxidant properties and phytochemical profiles of three pairs of plant species, each composed of a widely used medicinal plant and its regional counterparts from Provence with the same traditional indications:

- *Arnica montana* with *Pentanema montanum* (formerly known as *Inula montana*)
- *Helichrysum italicum* with *Helichrysum stoechas*
- *Satureja hortensis* with *Satureja montana*.

Phytochemical characterization using UHPLC-HRMS/MS and molecular networking revealed chemical profiles dominated by phenylpropanoids and flavonoids. According to DPPH and ABTS well-plate assays, all three Provence species demonstrated antioxidant activities comparable or even superior to their conventional counterparts. The on-line system successfully mapped antioxidants activities to 34 individual phenolic compounds, especially mono- and di-caffeoylquinic acid derivatives.

This work—recently published as “*Can Provence Flora Offer Effective Alternatives to Widely Used Medicinal Plants?*”—establishes this on-line coupling technique as a powerful tool for high-throughput bioactive compound discovery and demonstrates the potential of Provence flora as sustainable alternative to over-harvested medicinal plants for cosmetic or pharmaceutical applications.

# Characterizing marine benthic exometabolites from species to ecosystems

**Dr. Charlotte SIMMLER (CR CNRS)**

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Sponges are aquatic sessile invertebrates known to produce a wide range of specialized metabolites, which are extensively explored in drug discovery contexts.<sup>1</sup> These metabolites are also thought to be part of the sponge's defense and communication toolkit. However, in contrast to their pharmacological activities, the ecological functions of these specialized metabolites are generally more difficult to assess and therefore remain less well understood.<sup>2</sup> One way to address this challenge is to further study the metabolic or chemical output of sponges within their ecosystems. Through filter-feeding and metabolic processes, sponges recycle organic matter and subsequently release various metabolites (*i.e* exometabolites), including their specialized ones. These exometabolites may participate to the cycle of energy and nutrients in the benthos, and could also act as chemical cues involved in species interactions, thereby influencing the structure of marine biodiversity.<sup>3</sup> This presentation will showcase MS-based metabolomic results obtained from exometabolite captures performed with I-SMEL (In situ Marine moleculeE logger)<sup>4</sup> in sponge-dominated Mediterranean ecosystems.

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# Comprehensive Analysis of Small Molecule MS/MS Data with SIRIUS

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SIRIUS is a powerful tool for the automated analysis of tandem mass spectrometry (MS/MS) data, enabling the identification and annotation of small molecules with high confidence. In this workshop, we will guide participants through the entire SIRIUS data analysis pipeline, from loading LC-MS/MS data to feature detection, alignment, and molecular formula annotation. We will demonstrate how to search structure databases efficiently and validate results by integrating in-silico annotations, analog spectral library searches, and combinatorial fragmentation techniques. Additionally, we will explore strategies for processing large datasets, including classifying thousands of metabolites by chemical class and leveraging confidence scores to identify relevant compound annotations. This hands-on session will equip participants with practical skills for enhancing their MS/MS data analysis workflows.