# Tiffany M. Delhomme

PH.D

2 Rue Garibaldi, 69006 Lyon, France

# **Experience**

#### Postdoctoral researcher (Juan de la Cierva fellow)

IRB Barcelona, Spain

GENOME DATA SCIENCE LAB (PI: PR. SUPEK)

Jan. 2020 - present

• Scientific projects: mining genomics data in order to explore how exogenous and endogenous factors can impact mutational processes in the

#### **Ph.D. in Computational Cancer Genomics**

Lyon, France

INTERNATIONAL AGENCY FOR RESEARCH ON CANCER (IARC)

Nov. 2015 - July 2019

Thesis: Using the systematic nature of errors in NGS data in order to efficiently detect mutations. Computational methods and application to
early cancer detection.

# **Education**

### **Ph.D. in Computational Cancer Genomics**

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 Thesis: Using the systematic nature of errors in NGS data in order to efficiently detect mutations. Computational methods and application to early cancer detection.

#### M.Sc. in Theoretical Computer Science

Lyon, France

ENS Lyon, IXXI (Rhone-Alpes Complex Systems Institute)

Sept. 2014 - July. 2015

• Major in complex system modelisation, minor in algorithms

#### M.Sc. in Bioinformatics and Evolutionary Biology

Lyon, France

Universite Claude Bernard

Sept. 2012 - June. 2014

· Major in Bioinformatics and biostatistics, minor in evolution and biometry

## Skills\_

**Omics** NGS, somatic and germline variant calling and filtering, RNA-seq analysis, TCGA data

**Cancer Evolution** Neutral tumor evolution, repeated tumor evolution **Cancer Biomarkers** Circulating tumor DNA for early cancer detection

**Machine Learning** Supervised learning and semi-supervised learning (random forest)

Unsupervised learning (cluster-then-label)

Statistics Hypothesis testing and probability theory, bayesian inference, (generalized) linear models

**Discrete Mathematics** Graph theory, clustering, biological networks

**Programming** Bash, Python, cloud computing (SevenBridges Genomics plateform)

Statistical Programming Languages F

**Scientific Reproducibility** Docker, DockerHub, Singularity, SingularityHub, Git, GitHub

Workflow writing and data format Nextflow, CWL, JSON, YAML

**Language** French: native, English: good writing and communication, Spanish: notions

# **Publications**\_

# The PI3K/mTOR pathway is targeted by rare germline variants in patients with both melanoma and renal cell carcinoma

Cancers

Jean-Noël Hubert, Voreak Suybeng, Maxime Vallée, Tiffany M Delhomme et al.

May 2021

link

TP53 Targeted Deep Sequencing of Cell-Free DNA in Esophageal Squamous Cell Carcinoma Using Low-Quality Serum: Concordance with Tumor Mutation

Int. J. Mol. Sci.

Dariush Nasrollahzadeh, Gholamreza Roshandel, Tiffany M Delhomme et al.

May 2021

link

# The PI3K/mTOR Pathway Is Targeted by Rare Germline Variants in Patients with Both Melanoma and Renal Cell Carcinoma Jean-Noël Hubert, Voreak Suybeng, Maxime Vallée, Tiffany M Delhomme et al.

Cancers

2021 link

Development of sensitive droplet digital PCR assays for detecting urinary TERT promoter mutations as non-invasive biomarkers for detection of urothelial cancer

Ismail Hosen, Nathalie Forey, Geoffroy Durand, Catherine Voegele, Selin Bilici, Patrice H. Avogbe, Tiffany M

Cancers

2020 link

needlestack: an ultra-sensitive variant caller for multi-sample deep next generation sequencing data

Nucleic Acid Res. - Genomics and Bioinformatics

June 2020

**Tiffany M.Delhomme**, Patrice H. Avogbe, Aurelie AA. Gabriel *et al.* 

**Delhomme** et al.

link

Integrative and comparative genomic analyses identify clinically relevant groups of pulmonary carcinoids and unveil the existence of supra-carcinoids

Nat. Commun.

Nicolas Alcala\*, Noemie Leblay\*, Aurelie Gabriel\*, Lise Mangiante, David Hervas Marin, Theo Giffon, Anne-Sophie Sertier, Anthony Ferrari, Jules Derks, Akram Ghantous, **Tiffany M.Delhomme** *et al.* 

Aug. 2019

Urinary TERT promoter mutations as non-invasive biomarkers for the comprehensive detection of urothelial cancer

**EBioMedicine** 

Patrice H. Avogbe, Arnaud Manel, Emmanuel Vian, Geoffroy Durand, Nathalie Forey, Catherine Voegele, Maria Zvereva, Ismail Hosen, Sonia Meziani, Berengere De Tilly, Gilles Polo, Olesia Lole, Pauline Francois, **Tiffany**M. Delhomme et al.

June 2019

link

Integrative genomic profiling of large-cell neuroendocrine carcinomas reveals distinct subtypes of high-grade neuroendocrine lung tumors

Nature Communications

Julie George, Vonn Walter, Martin Peifer, Ludmil B. Alexandrov, Danila Seidel, Frauke Leenders, Lukas Maas, Christian Müller, Ilona Dahmen, **Tiffany M.Delhomme** *et al.* 

Mar. 2018

link

link

link

KRAS mutations in blood circulating cell-free DNA: a pancreatic cancer case-control study

Oncotarget

Oct. 2016

Florence Le Calvez-Kelm, Matthieu Foll, Magdalena B. Wozniak, **Tiffany M.Delhomme** et al.

FRioMedicine

**Identification of Circulating Tumor DNA for the Early Detection of Small-cell Lung Cancer** Lynnette Fernandez-Cuesta\*, Sandra Perdomo\*, Patrice H.Avogbe\*, Noemie Leblay, **Tiffany M.Delhomme** *et al.* 

Aug. 2016

Contribution: development of needlestack, an highly sensitive variant caller that can detect low abundance mutations such as tumor-derived mutations from circulating tumor DNA data. In this study, we applied needlestack to Small-cell Lung Cancer cases and matched controls in order to use circulating tumor DNA as an early cancer biomarker.

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SUBMITTED

Disentangling heterogeneity of Malignant Pleural Mesothelioma through deep integrative omics analyses

Lise Mangiante\*, Nicolas Alcala\*, Alex Di Genova\*, Alexandra Sexton-Oates\*, [+14], **Tiffany M. Delhomme** *et* 

Nat. Commun. 2022

IN PREPARATION

Proton and alpha radiation-induced mutational profiles in human cells

Tiffany M. Delhomme, Manuela Buonanno, Veljko Grilj, Josep Biayna, Fran Supek

To be submitted in July 2022

al.

# Assessment of the diagnostic value of circulating RB1 and TP53 mutations for early detection of small cell lung cancers

Patrice H. Avogbe\*, **Tiffany M.Delhomme**\* et al.

expected sub. Dec. 2022

# **Communications**

#### **POSTERS**

# needlestack: an ultra-sensitive variant caller for multisample deep next generation sequencing data

Journees Ouvertes Biologie, Informatique et Mathematiques (JOBIM), Clermont Ferrand, France

June 2015

# needlestack: an ultra-sensitive variant caller for multisample deep next generation sequencing data

RECOMB-Computational Cancer Biology, Paris, France

Apr. 2018

#### **TALKS**

# needlestack: an ultra-sensitive variant caller for multisample deep next generation sequencing data

Journees Ouvertes Biologie, Informatique et Mathematiques (JOBIM), Lyon, France

Aug. 2016

## IARC nextflow pipelines: toward efficient cancer genomics analyses

Nextflow workshop, invited speaker, Barcelona, Spain

Nov. 2018

# Honors & Awards

#### **FELLOWSHIPS**

**Postdoctoral Fellowship**, "Juan de la Cierva", Spanish Ministry of Science **Ph.D. Fellowship**, La Ligue Nationale Contre le Cancer

2020-2022 2015-2018

## Referees

### Dr. Matthieu Foll

GEN/GCS group International Agency for Research on Cancer Lyon, France

FollM@iarc.fr +334-72-73-85-37 Dr. James McKay

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## Pr. Fran Supek

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