Tiffany M. Delhomme

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Looking for a postdoctoral position in computational cancer genomics

Education

International Agency for Research on Cancer (IARC)

Lyon, France

Nov. 2015 - currently

Ph.D. IN COMPUTATIONAL CANCER GENOMICS

• Thesis: Dealing with NGS errors to produce an efficient variant calling. Application to early cancer detection.

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

Lyon, France

M.Sc. IN THEORETICAL COMPUTER SCIENCE

Universite Claude Bernard

Sept. 2014 - July. 2015

• Major in complex system modelisation, minor in algorithms

Lyon, France

M.Sc. in Bioinformatics and Evolutionary Biology

Sept. 2009 - 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

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

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

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

Identification of Circulating Tumor DNA for the Early Detection of Small-cell Lung Cancer

EBioMedicine

Lynnette Fernandez-Cuesta*, Sandra Perdomo*, Patrice H.Avogbe*, Noemie Leblay, Tiffany M.Delhomme et

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.

link

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

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

Oncotarget

Oct. 2016 link

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

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

Nature Communications

Mar 2018

link

SUBMITTED

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

Maria Zvereva, Gabriel Roberti, Geoffroy Durand, Catherine Voegele, MinhDao Nguyen, Matthieu Foll, **Tiffany M. Delhomme** *et al.*

Clinical Chemistry

sub. Nov 2018

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

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.*

Nature Communications

sub. Nov 2018

IN PREPARATION

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

sequencing data
Tiffany M.Delhomme et al.

Contribution: development of the algorithm and implementation (mostly in R, bash and nextflow). needlestack is based on a robust negative binomial regression and is implemented in nextflow to emphasize scientific reproducibility and efficient computations. Analyses on real and simulated tumor data to evaluate the performance of needlestack in case of germline, somatic and low allelic fraction somatic mutations.

expected sub. Dec. 2018

link

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.

Contribution: application of needlestack on two new cohorts and development of variant filtering steps in order to increase the specificity obtained in Fernandez-Cuesta *et al.*

expected sub. Dec. 2018

Communications

POSTERS

needlestack: an ultrasensitive variant caller for multisample deep next generation sequencing data

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

June 2015

needlestack: an ultrasensitive variant caller for multisample deep next generation sequencing data

RECOMB-Computational Cancer Biology, Paris, France

Apr. 2018

TALKS

needlestack: an ultrasensitive 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

Ph.D. Fellowship, La Ligue Nationale Contre le Cancer

2015-2018

Referees

Matthieu Foll

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

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James McKay

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