

Jean MONLONG

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Education

- 2012 – 2018** [MCGILL UNIVERSITY](#), Montreal, Canada.
PhD in the Human Genetics Department in [Guillaume Bourque's Lab](#).
- 2010 – 2011** [UNIVERSITAT POLITÈCNICA DE CATALUNYA](#), Barcelona, Spain.
Master in Statistic and Operations Research as an exchange student in the Faculty of Mathematics and Statistics.
- 2008 – 2010** [ENSIMAG](#), Grenoble, France.
Computer Science and Mathematics course with specialization in **Bioinformatics**.
- 2006 – 2008** [LYCÉE MICHEL MONTAIGNE](#), Bordeaux, France.
Preparatory classes for entrance to the Grandes Écoles. Mathematics and Physics.

Professional Experiences and Projects

- 2018 –** [UNIVERSITY OF CALIFORNIA SANTA CRUZ](#), Santa Cruz, USA.
Postdoctoral researcher with Benedict Paten in the [Computational Genomics Lab](#), working on [variation graphs](#).
- 2012 – 2018** [BOURQUE LAB](#), [McGill Genome Centre](#), Montreal, Canada.
PhD project. Implementation of a [CNV detection method](#) for whole-genome sequencing that can deal with repeat-rich regions. Application to large cohorts of normal and disease genomes.
- 2017** [CENTER FOR GENOMIC MEDICINE](#), Kyoto, Japan.
(3 months) As part of an exchange program, I visited Dr. Matsuda's group and contributed to their ongoing sequencing projects to characterize Japanese genomes.
- 2017** [GENOMICS ENGLAND](#), London, UK.
(1.5 month) Visiting the bioinformatics team lead by Dr. Rendon, I contributed to the curation of structural variant calls for their genome analysis pipeline.
- 2011 – 2012** [CENTER FOR GENOMIC REGULATION\(CRG\)](#), Barcelona, Spain.
(1 year) Graduation project in [Roderic Guigó's group](#). Comparison of splicing activity and [detection of splicing QTLs](#) from RNA-Seq experiments. Participation in the [Geuvadis](#) and [GTEx](#) projects.
- 2011** [UNIVERSITAT POLITÈCNICA DE CATALUNYA](#), Barcelona, Spain.
(3 months) Study of the regularization of the generalized canonical correlation analysis.
- 2010** [NEOMADES](#)(Mobile software development), Bidart, France.
(3 months) Implementation of a Java module of their principal product.
- 2010** [ENSIMAG](#), Grenoble, France.
(1 month) Breast cancer modelization and Bayesian estimation of the overdiagnosis rate. Collaboration with La Tronche's hospital (Grenoble).

Expertise

R, Java, C, Bash, Python, AWK, Perl, D3/Javascript/HTML/CSS.
Git, Linux, High Performance Computing, LaTeX, R/Markdown, Emacs.
Bioinformatics, Genomics, Transcriptomics.

Scholarships and Awards

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| 2018 | Department Activity Award. Human Genetics Dept, McGill University. |
| 2017 | Student Exchange Support Program. JASSO, Japan. |
| 2017 | Graduate Mobility Award. Graduate and Postdoctoral Studies, McGill University. |
| 2017 | Oral Presentation, Honorable Mention. Human Genetics Dept's Research Day, McGill University. |
| 2017 | Excellence Award. Human Genetics Dept, McGill University. |
| 2013-2016 | Graduate Research Enhancement and Travel Award funding attendance to five international conferences. Human Genetics Dept, McGill University. |
| 2015 | Excellence Award. Human Genetics Dept, McGill University. |
| 2014 | Best oral presentation. Human Genetics Dept's Research Day, McGill University. |

Publications

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| 2019 | G Hickey*, D Heller*, J Monlong* , JA Sibbesen, J Siren, J Eizenga, E Dawson, E Garrison, A Novak, and B Paten. Genotyping structural variants in pangenome graphs using the vg toolkit. <i>bioRxiv</i> , 2019. |
| 2018 | J Monlong , P Cossette, C Meloche, G Rouleau, SL Girard, and G Bourque. Human copy number variants are enriched in regions of low mappability. <i>Nucleic Acids Research</i> , 2018. |
| 2018 | J Monlong* , SL Girard*, ..., JL Michaud, G Rouleau, BA Minassian, G Bourque, and P Cossette. Global characterization of copy number variants in epilepsy patients from whole genome sequencing. <i>PLOS Genetics</i> , 2018. |
| 2017 | M Arseneault*, J Monlong* , ..., M Lathrop, G Bourque, and Y Riazalhosseini. Loss of chromosome Y leads to down regulation of KDM5D and KDM6C epigenetic modifiers in clear cell renal cell carcinoma. <i>Scientific Reports</i> , 2017. |
| 2017 | F Hamdan, CT Myers, P Cossette, P Lemay, D Spiegelman, A Dionne-Laporte, C Nassif, O Diallo, J Monlong , ..., BA Minassian, and JL Michaud. High Rate of Recurrent De Novo Mutations in Developmental and Epileptic Encephalopathies. <i>American journal of human genetics</i> , 2017. |
| 2017 | The GTEx Consortium . Genetic effects on gene expression across human tissues. <i>Nature</i> , 2017. |
| 2017 | A Saha, et al., and The GTEx Consortium . Co-expression networks reveal the tissue-specific regulation of transcription and splicing. <i>Genome research</i> , 2017. |
| 2017 | F Yang, et al., and The GTEx Consortium . Identifying cis-mediators for trans-eQTLs across many human tissues using genomic mediation analysis. <i>Genome research</i> , 2017. |
| 2017 | X Li, et al., and The GTEx Consortium . The impact of rare variation on gene expression across tissues. <i>Nature</i> , 2017. |
| 2017 | MH Tan, et al., and The GTEx Consortium . Dynamic landscape and regulation of RNA editing in mammals. <i>Nature</i> , 2017. |
| 2017 | T Tukiainen, et al., and The GTEx Consortium . Landscape of X chromosome inactivation across human tissues. <i>Nature</i> , 2017. |
| 2015 | M Melé, PG Ferreira, F Reverter, DS DeLuca, J Monlong , M Sammeth, ..., and The GTEx Consortium . The human transcriptome across tissues and individuals. <i>Science</i> , 2015. |
| 2015 | The GTEx Consortium . The Genotype-Tissue Expression (GTEx) pilot analysis: Multitissue gene regulation in humans. <i>Science</i> , 2015. |

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| 2015 | D Pervouchine, S Djebali, A Breschi, CA Davis, PP Barja, A Dobin, A Tanzer, J Lagarde, C Zaleski, L See, M Fastuca, J Drenkow, H Wang, G Bussotti, B Pei, S Balasubramanian, J Monlong , ..., C Notredame, R Guigo, and TR Gingeras. Enhanced transcriptome maps from multiple mouse tissues reveal evolutionary constraint in gene expression. <i>Nature Communications</i> , 2015. |
| 2014 | J Monlong , M Calvo, PG Ferreira, and R Guigó. Identification of genetic variants associated with alternative splicing using sQTLseeker. <i>Nature Communications</i> , 2014. |
| 2014 | PG Ferreira, P Jares, D Rico, G Gomez-Lopez, A Martinez-Trillos, N Villamor, S Ecker, A Gonzalez-Perez, DG Knowles, J Monlong , ..., E Campo, and R Guigo. Transcriptome characterization by rna sequencing identifies a major molecular and clinical subdivision in chronic lymphocytic leukemia. <i>Genome research</i> , 2014. |
| 2014 | L Greger, J Su, J Rung, PG Ferreira, The Geuvadis Consortium , T Lappalainen, ET Dermitzakis, and A Brazma. Tandem RNA chimeras contribute to transcriptome diversity in human population and are associated with intronic genetic variants. <i>PloS one</i> , 2014. |
| 2013 | T Lappalainen, M Sammeth*, MR Friedlander*, PA t Hoen*, J Monlong* , MA Rivas*, and et al. Transcriptome and genome sequencing uncovers functional variation in humans. <i>Nature</i> , 2013. |
| 2013 | PC t Hoen, et al., and The Geuvadis Consortium . Reproducibility of high-throughput mRNA and small RNA sequencing across laboratories. <i>Nature biotechnology</i> , 2013. |
| 2013 | The GTEx Consortium . The Genotype-Tissue Expression (GTEx) project. <i>Nat Genet</i> , 2013. |
| 2011 | A Seigneurin, O Francois, J Labarere, P Oudeville, J Monlong , and M Colonna. Overdiagnosis from non-progressive cancer detected by screening mammography: stochastic simulation study with calibration to population based registry data. <i>BMJ</i> , 2011. |

Selected Scientific Presentations

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| 2017 | QUEBEC, CANADA, Quebec. Oral presentation, <i>6th Annual Canadian Human and Statistical Genetics Meeting</i> . |
| 2016 | JAPAN, Kyoto. Oral presentation, <i>13th International Congress of Human Genetics</i> . |
| 2015 | BC, CANADA, Vancouver. Oral presentation, <i>4rd Annual Canadian Human and Statistical Genetics Meeting</i> . |
| 2014 | UK, Cambridge. Oral presentation, <i>Genome Informatics</i> conference. |

Other Scientific Presentations

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| 2019 | TX, USA, Houston. Oral presentation, <i>GRC-GIAB Workshop</i> at ASHG. |
| 2019 | CA, USA, UC Davis. Oral presentation, <i>Northern California Computational Biology Symposium</i> . |
| 2017 | QUEBEC, CANADA, Montreal. Oral presentation, Human Genetics Dept. Research Day. |
| 2016 | SPAIN, Barcelona. Poster presentation, <i>European Society of Human Genetics</i> conference. Best Poster Award candidate. |
| 2015 | SCOTLAND, Glasgow. Poster presentation, <i>European Society of Human Genetics</i> conference. |

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| 2014 | QUEBEC, CANADA, Montreal. Oral presentation, Human Genetics Dept. Research Day. |
| 2014 | BRITISH COLUMBIA, CANADA, Victoria. Poster presentation at the <i>3rd Annual Canadian Human and Statistical Genetics Meeting</i> . |
| 2014 | NEW MEXICO, USA, Santa Fe. Poster presentation at the <i>Mobile Genetic Elements and Genome Evolution</i> Keystone conference. |
| 2013 | QUEBEC, CANADA, Esterel. Poster presentation at the <i>2nd Annual Canadian Human and Statistical Genetics Meeting</i> . |
| 2013 | MONTANA, USA, Big Sky. Poster presentation at the <i>Mobile DNA in Mammalian Genomes</i> FASEB conference. |
| 2011 | SPAIN, Barcelona. Poster presentation at the <i>XIIIth Spanish Biometry Conference and 3rd Ibero-American Biometry Meeting</i> . |

Other Activities

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| 2018 | IRCM, MONTREAL. Gave a short seminar about R Markdown at the Montreal Bioinformatics User Group . |
| 2013-2018 | MCGILL UNIVERSITY. Organizer and instructor of eight full-day <i>Introduction to R and Bioinformatics</i> workshops for Human Genetics graduate students. |
| 2014-2017 | MCGILL UNIVERSITY. VP Communications at the Human Genetics Student Society . |
| 2017 | UNIVERSITÉ DE MONTREAL. Participated to a Bio-Hackaton. Our 24h project was to detect extending filaments from 3D cell imaging. |
| 2017 | IRCM, MONTREAL. Gave a short seminar about the Gviz package at the Montreal Bioinformatics User Group . |
| 2016 | KYOTO, JAPAN. Instructor at the <i>4th Kyoto Course and Symposium on Bioinformatics for Next Generation Sequencing with Applications in Human Genetics</i> . |
| 2015 | UNIVERSITÉ DE MONTREAL. Participated to a Bio-Hackaton. Our 24h project was to develop interactive visualization of gene expression in cancer samples. |
| 2014 | MCGILL UNIVERSITY. Teaching Assistant for the <i>Advanced Statistical Concepts in Genetic and Genomic Analysis</i> course. |
| 2014 | MCGILL UNIVERSITY. Organizer of a four-days <i>R and Bioinformatics</i> workshop for System Biology graduate students. |