Leonardo Collado-Torres

Education

- 2011-present **PhD in Biostatistics**, Johns Hopkins Bloomberg School of Public Health, Baltimore, US.
 - 2005-2009 **Bachelor in Genomic Sciences (LCG)**, National Autonomous University of Mexico (UNAM), Cuernavaca, MX, Grade 9.71/10.
 - 2002-2005 High school, ITESM Campus Cuernavaca, Cuernavaca, MX, Grade 97.8/100.

PhD thesis

- title Developing single-base RNA-seq methods and software to understand neuropsychiatric disorders and more.
- advisors Jeffrey T. Leek and Andrew E. Jaffe.
- description The goal is to develop statistical methods and software that enable researchers to differentiate the sources of variation observed in RNA-seq while minimizing the dependance on known annotation. This will allow researchers to correct for technological variation and study the biological variation driving their phenotype of interest. Then apply these methods to further our understanding of neuropsychiatric disorders using the Lieber Institute for Brain Development human brains collection (> 1000 samples).

Honors and awards

- 2011 Awarded CONACyT Mexico scholarship for PhD studies outside Mexico.
- 2009 Summa Cum Laude for bachelor in Genomic Sciences studies at LCG-UNAM.
- 2005 Best high school average (\sim 200 students): awarded ITESM system 90% scholarship for college studies, declined to join LCG-UNAM.

Experience

Vocational

2009-2011 **Scientific executive**, Winter Genomics, Cuernavaca, MX.

Responsible for recruiting and hiring new personnel, overseeing and supervising bioinformaticians, training new employees, writing research reports and presenting them to colleagues, and organizing all scientific projects.

- First scientific staff member at Winter Genomics;
- Projects completed:
 - de novo genome assembly simulations,
 - assembly and annotation of the phiVC8 bacteriophage genome,
 - integrated analysis of more than 20 RNA-seq samples for determination of transcription initiation in *Escherichia coli* reported in Gama-Castro et al., PMID 21051347,
 - de novo assembly of four Escherichia coli strains and lead to Aguilar et al., PMID 22884033;
- o Designed training material for new employees.

Research

2009–2011 Bioinformatician, Enrique Morett lab, IBT-UNAM, Cuernavaca, MX.

Identified transcriptions start sites and transcription units in *Escherichia coli* and *Geobacter sulfurreducens* with RNA-seq data.

Developed the Bacterial Transcription R package.

2007–2009 **Undergraduate research assistant**, *Guillermo Dávila lab*, CCG-UNAM, Cuernavaca, MX.

Determined bacteriophage ecological groups by developing a method based on codon distribution of all phage sequenced genomes.

Joint work with Sur Herrera Paredes.

2007 Undergraduate research assistant, Roberto Kolter lab, Harvard, Boston, US.

Supervisor: Elizabeth Shank.

Carried out screenings to identify bacteria that activate the production of exopolysaccharide through the activation of the gene tasA in *Bacillus subtilis*.

Publications

Peer-reviewed

- 1. **Collado-Torres L**, Jaffe AE and Leek JT. regionReport: Interactive reports for region-based analyses. *F1000Research* 2015, 4:105. doi: 10.12688/f1000research.6379.1.
- 2. Jaffe AE, Shin J, **Collado-Torres L**, Leek JT, et al. Developmental regulation of human cortex transcription and its clinical relevance at single base resolution. *Nat. Neurosci.* 2014. doi: 10.1038/nn.3898.
- 3. Shank EA, Klepac-Ceraj V, **Collado-Torres L**, Powers GE, Losick R, Kolter R. Interspecies interactions that result in Bacillus subtilis forming biofilms are mediated mainly by members of its own genus. *Proc. Natl. Acad. Sci.* U.S.A. 2011 Nov;108(48):E1236–1243. doi: 10.1073/pnas.1103630108.
- Gama-Castro S, Salgado H, Peralta-Gil M, Santos-Zavaleta A, Muñiz-Rascado L, Solano-Lira H, Jimenez-Jacinto V, Weiss V, García-Sotelo JS, López-Fuentes A, Porrón-Sotelo L, Alquicira-Hernández S, Medina-Rivera A, Martínez-Flores I, Alquicira-Hernández K, Martínez-Adame R, 615 N. Wolfe Street, Room E3032 21205-2179 United States

☎ +1 (410) 955 0958 • ☑ lcollado@jhu.edu ¹ http://lcolladotor.github.io/about.html Bonavides-Martínez C, Miranda-Ríos J, Huerta AM, Mendoza-Vargas A, **Collado-Torres L**, Taboada B, Vega-Alvarado L, Olvera M, Olvera L, Grande R, Morett E, Collado-Vides J. RegulonDB version 7.0: transcriptional regulation of Escherichia coli K-12 integrated within genetic sensory response units (Gensor Units). *Nucleic Acids Res.* 2011 Jan;39(Database issue):D98–105. doi: 10.1093/nar/gkq1110.

Pre-prints

- Nellore A, Collado-Torres L, Jaffe AE, Morton J, Pritt J, Alquicira-Hernández J, Leek JT, Langmead B. Rail-RNA: Scalable analysis of RNA-seq splicing and coverage. bioRxiv 019067 (2015). doi:10.1101/019067.
- 2. **Collado-Torres L**, Jaffe AE, Leek JT. regionReport: Interactive reports for region-based analyses. *bioRxiv* 016659 (2015). doi:10.1101/016659.
- 3. **Collado-Torres L**, Frazee AC, Love MI, Irizarry RA, Jaffe AE, Leek JT. derfinder: Software for annotation-agnostic RNA-seq differential expression analysis. *bioRxiv* 015370 (2015). doi:10.1101/015370.

Books

 Frazee AC, Collado-Torres L, Jaffe AE, Langmead B, Leek JT. Measurement, Summary, and Methodological Variation in RNA-sequencing in Statistical Analysis of Next Generation Sequencing Data, Springer, 2014, 115-128.

Google scholar

https://scholar.google.com/citations?user=h57-MykAAAAJ

ORCID

http://orcid.org/0000-0003-2140-308X

LinkedIn

https://www.linkedin.com/in/lcollado

Epernicus

http://www.epernicus.com/lc40

GitHub

https://github.com/lcolladotor

Referee for

Since 2015 Bioinformatics

Since 2013 Biostatistics

Presentations

Talks at conferences

2015 Collado-Torres L, Frazee AC, Love MI, Irizarry RA, Jaffe AE, Leek JT. Annotation-agnostic differential expression analysis, Genomics and Bioinformatics Symposium, Center for Computational Genomics, Hopkins, Baltimore – US.

- 2015 Jaffe AE, Shin J, **Collado-Torres L**, Leek JT, et al. Dissecting human brain development at high resolution using RNA-seq, *ENAR*, Miami US.
- 2014 Jaffe AE, Shin J, **Collado-Torres L**, Leek JT, et al. Developmental regulation of human cortex transcription at base-pair resolution, *is3b*: 1st International Summer Symposium on Systems Biology, INMEGEN, Mexico City MX.
- 2014 **Collado-Torres L**, Frazee AC, Love MI, Irizarry RA, Jaffe AE, Leek JT. Fast differential expression analysis annotation-agnostic across groups with biological replicates, LCG 10 year anniversary, LCG-UNAM, Cuernavaca MX.
- 2013 **Collado-Torres L**, Frazee AC, Irizarry RA, Jaffe AE, Leek JT. Differential expression analysis of RNA-seq data at base-pair resolution in multiple biological replicates, *useR2013*, Albacete Spain.
- 2010 Collado-Torres L, Reyes-Quiroz A, Cuéllar-Partida G, Moreno-Mayar V, Vargas-Chávez C, Collado-Vides J. BacterialTranscription: a R package to identify Transcription Start Sites and Transcription Units, *Bioconductor Developer Meeting*, EMBL, Heidelberg Germany.

Posters

- 2015 **Collado-Torres L**, Frazee AC, Love MI, Irizarry RA, Jaffe AE, Leek JT. Annotationagnostic RNA-seq differential expression analysis software, *ASHG2015* and *IDIES2015*, Baltimore US.
- 2014 **Collado-Torres L**, Frazee AC, Love MI, Irizarry RA, Jaffe AE, Leek JT. Fast annotation-agnostic differential expression analysis, *ENAR* and *Delta Omega Poster Competition* (*JHSPH*), Baltimore US.
- 2013 **Collado-Torres L**, Jaffe AE, Leek JT. Fast annotation-agnostic differential expression analysis, *Genomics and Bioinformatics Symposium*, Center for Computational Genomics, Hopkins, Baltimore US.
- 2010 Collado-Torres L, Reyes-Quiroz A, Cuéllar-Partida A, Moreno-Mayar V, Taboada B, Vega-Alvarado L, Jiménez-Jacinto V, Mendoza-Vargas A, Grande R, Olvera L, Olvera M, Vargas-Chávez C, Júarez K, Collado-Vides J, Morett E. Global Analysis of Transcription Start Sites and Transcription Units in Bacterial Genomes, From Functional Genomics to Systems Biology, EMBL, Heidelberg Germany.
- 2010 Collado-Torres L, Reyes-Quiroz A, Cuéllar-Partida A, Moreno-Mayar V, Taboada B, Vega-Alvarado L, Jiménez-Jacinto V, Mendoza-Vargas A, Grande R, Olvera L, Olvera M, Vargas-Chávez C, Júarez K, Collado-Vides J, Morett E. Global Analysis of Transcription Start Sites and Transcription Units in Bacterial Genomes, *BioC2010*, FHCRC, Seattle US.

Other talks

- 2015 Easy parallel computing with BiocParallel and HTML reports with knitrBootstrap, *Biostatistics Computing Club*, JHSPH, Baltimore US.
- 2015 Does mapping simulated RNA-seq reads provide information?, *Joint Genomic Meeting*, JHSPH, Baltimore US.

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- 2014 Git for research, Biostatistics Computing Club, JHSPH, Baltimore US.
- 2013 Introduction to ggbio, Genomics for Students, JHSPH, Baltimore US.
- 2013 Introduction to knitr, Biostatistics Computing Club, JHSPH, Baltimore US.
- 2013 Introduction to High-Throughput Sequencing and RNA-seq, *Genomics for Students*, JHSPH, Baltimore US.
- 2012 DEXSeq paper discussion, Genomics for Students, JHSPH, Baltimore US.
- 2012 Introduction to R and Biostatistics, LCG-UNAM via Skype.
- 2012 Introducing Git while making your academic webpage, *Biostatistics Computing Club*, JHSPH, Baltimore US.
- 2011 Introducing Biostatistics to first year LCG students, LCG-UNAM via Skype.
- 2010 Introduction to using Bioconductor for High Throughput Sequencing Analysis, *National Bioinformatics Week*, CCG-UNAM, Cuernavaca MX.
- 2009 Bacteriophages: analyzing their diversity, *LCG third generation symposium*, CCG-UNAM, Cuernavaca MX.

Courses and Meetings Attendance

- 2015 ENAR, Miami US.
- 2014 is3b, INMEGEN, Mexico City MX.
- 2014 BioC2014, Harvard, Boston US.
- 2014 IDIES2014, Johns Hopkins University, Baltimore US.
- 2014 ENAR, Baltimore US.
- 2014 Delta Omega Poster Competition, Johns Hopkins University, Baltimore US.
- 2014 LCG 10 year anniversary, LCG-UNAM, Cuernavaca MX.
- 2013 Genomics and Bioinformatics Symposium, Johns Hopkins University, Baltimore US.
- 2013 useR2013, Albacete Spain.
- 2011 BioC2011, FHCRC, Seattle US.
- 2010 From Functional Genomics to Systems Biology, EMBL, Heidelberg Germany.
- 2010 BioC2010, FHCRC, Seattle US.
- 2009 BioC2009, FHCRC, Seattle US.
- 2009 Course on Oral Communication taught by the master Rafael Popoca, CCG-UNAM, Cuernavaca MX.
- 2008 BioC2008, FHCRC, Seattle US.
- 2008 A Short R/Bioconductor Course by James Bullard from UC Berkeley, LCG-UNAM, Cuernavaca MX.
- 2007 Boston Bacterial Meeting, Boston US.
- 2007 Retreat of the Department of Microbiology and Molecular Genetics Harvard, Boston US.

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- 2006 Winter School in Genomics, CCG-UNAM, Cuernavaca MX.
- 2005 HUGO 2005, Kyoto Japan.

Software

Bioconductor - main author

- 2014 derfinder: Annotation-agnostic differential expression analysis of RNA-seq data at base-pair resolution via the DER Finder approach 2963 downloads in last 12 months.
- 2014 derfinderPlot: plotting functions for derfinder results 2372 downloads in past 12 months.
- 2014 regionReport: generate HTML reports for exploring a set of regions 2292 downloads in last 12 months.
- 2014 derfinderHelper: helper functions for derfinder package 2756 downloads in last 12 months.
- 2014 derfinderData: data for derfinder examples 600 downloads in past 12 months.

 Bioconductor contributor role
 - . . .
- 2015 bumphunter
- 2014 ballgown

Other R packages

- 2014 dots: simplify function calls https://github.com/lcolladotor/dots.
- 2013 fitbitR: visualize your FitBit data https://github.com/lcolladotor/fitbitR.
- 2011 BacterialTranscription: identify TSSs and TUs from RNA-seq data.
- shiny applications
- 2014-2015 MPH capstone TA office hours sign up https://github.com/lcolladotor/MPHcapstoneTA
 - 2014 Simple mortgage calculator https://github.com/lcolladotor/mortgage

Computer skills

all-purpose **R**

Ranked 138/5506 in the US and 366/52549 worldwide by http://github-awards.com/users/lcolladotor as of November 04, 2015. Does not take into account contributions at https://github.com/leekgroup.

statistics Stata

scripting bash, Perl

markup LaTeX, markdown

OS Linux

cluster queue Sun Grid Engine

Teaching Experience

Instructor

- o PDCB-UNAM, Cuernavaca, MX
 - 2011 Invited instructor for the course Introduction to R and Biostatistics http://lcolladotor.github.io/courses/PDCB-Biostats.html ~ 10 enrollment.
 - 2010 Analysis of High-Throughput Sequencing data with Bioconductor for Biomedical Sciences PhD Program students http://lcolladotor.github.io/courses/PDCB-HTS.html ~ 10 enrollment.
- o CCG-UNAM, Cuernavaca, MX
 - 2010 Introduction to Using Bioconductor for High-Throughput Sequencing Analysis practice lab at the National Bioinformatics Week ~ 40 enrollment.
- o IBT-UNAM, Cuernavaca, MX
 - 2010 Introduction to R and plotting with R course for Morett's lab ~ 10 enrollment.
 - Organized and gave a lecture for the course on Statistical Methods and Analysis of Genomic Data http://lcolladotor.github.io/courses/MEyAdDG.html ~ 20 enrollment.
 - 2009 Organized the course Introduction to Bioinformatics for Morett's lab and served as instructor for the Introduction to R and plotting with R module http://lcolladotor.github.io/courses/mIntroR.html ~ 10 enrollment.
- LCG-UNAM, Cuernavaca, MX
 - 2009 Seminar III: R/Bioconductor. In-depth Bioconductor course http://www.lcg.unam. mx/~lcollado/B/ ~ 30 enrollment.

Guest lecturer

- o JHSPH, Baltimore, US
 - 2015 Introduction to R for Public Health Researchers: Reproducible research module ~ 20 enrollment.
- LCG-UNAM, Cuernavaca, MX
 - 2012 Introduction to R and Biostatistics lecture for Seminar 1: Introduction to Bioinformatics course ~ 30 enrollment.
 - 2011 Introduction to R and Biostatistics lecture for Seminar 1: Introduction to Bioinformatics course ~ 30 enrollment.

Lead teaching assistant

- o JHSPH, Baltimore, US
 - 2014–2015 Statistical Methods in Public Health I and II ~ 550 enrollment.

Teaching assistant

- o JHSPH, Baltimore, US
 - 2015–2016 Statistical Methods in Public Health I ~ 550 enrollment.
 - 2015 Introduction to R for Public Health Researchers ~ 20 enrollment.

- 2015 MPH capstone project: 30 min one-on-one consulting sessions (biostatistics, Stata coding) ~ 500 enrollment.
- 2014 MPH capstone project: 30 min one-on-one consulting sessions (biostatistics, Stata coding) ~ 500 enrollment.
- 2013–2014 Statistical Methods in Public Health I and II ~ 550 enrollment.
- 2012–2013 Statistical Methods in Public Health I, II, III, and $IV \sim 550$ enrollment.
- LCG-UNAM, Cuernavaca, MX
 - 2009 Principles of Statistics. Basic R http://www.lcg.unam.mx/~lcollado/E/ ~ 30 enrollment.
 - 2008 Bioinformatics and Statistics I. R and Bioconductor overview http://www.lcg.unam. mx/~lcollado/R/ ~ 40 enrollment.

Mentoring

- 2015 Mentored 1 LCG-UNAM student visiting Jeff Leek's group at JHSPH: Alquicira-Hernández J.
- 2009–2011 Advised and trained 13 LCG-UNAM students and alumni while working at *Winter Genomics*: Riveros-McKay F, Vargas-Chávez C, Dulanto-Acevedo V, Romero-Martínez S, Samaniego-Castruita J, Zepeda-Mendoza L, Vargas-Velázquez A, Noé-González M, Soto Jiménez LM, López Moyado I, Medina-Abarca H., Izquierdo-Rangel E, and Berrocal-Quezada NA.
 - 2009 Trained 3 LCG-UNAM students to take over the R/Bioconductor course: Reyes-Quiroz A, Moreno-Mayar V, and Reyes-López J.

Other

- 2012-2014 Organized the *Genomics for Students* group http://lcolladotor.github.io/courses/GenomicsForStudents.html
- 2012–2014 Organized *Cultural Mixer* events for the Department of Biostatistics (JHSPH) with Amanda Mejia.
- 2009–2011 Organized a Genomics Journal Club at IBT-UNAM.
- 2008–2009 Elected class representative for the LCG Academic Committee.
- 2008–2009 Class representative for Administration Unit for Technology Information committee.
 - 2008 Helped start the National Node of Bioinformatics online forum http://foros.nnb.unam.mx/.

Languages

Native Spanish

Bilingual English

Basic French

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