Curriculum Vitae

Notarization. I have read the following and certify that this curriculum vitae is a current and accurate statement of my professional record.

The state of the s	September 18, 2016	
Signature	Date	

I. Personal Information

I.A. UID, Last Name, First Name, Middle Name, Contact Information

Last Name: Corrada Bravo First Name: Héctor

3114F Biomolecular Sciences Building #296

University of Maryland College Park, MD 20745 hcorrada@umiacs.umd.edu

http://hcbravo.org UID: 111544240

I.B. Academic Appointments at UMD

- Assistant Professor, Department of Computer Science University of Maryland, College Park, MD July 2010-Present
- Assistant Professor, Institute for Advanced Computer Studies University of Maryland, College Park, MD July 2010-Present
- Affiliate Assistant Professor, Applied Math, Statistics and Scientific Computation University of Maryland, College Park, MD July 2011-Present

I.D. Other Employment

 Postdoctoral Fellow, Department of Biostatistics Johns Hopkins University School of Public Health, Baltimore, MD September 2008-June 2010

- Research and Teaching Assistant, Departments of Computer Science and Statistics University of Wisconsin, Madison, WI September 2003-August 2008
- Resarch Intern, IBM Research, Almaden, CA May 2005-August 2005

I.E. Educational Background

 Ph.D., Computer Science, University of Wisconsin, Madison, WI September 2003 - August 2008
 Dissertation: Graph-based data analysis

Advisor: Grace Wahba and Raghu Ramakrishnan

- D.M.A., Indiana University School of Music, Bloomington, IN (ABD) September 2000 - August 2003
- M.M., Peabody Institute of Music, Baltimore, MD September 1997 - May 1999
- B.M., Peabody Institute of Music, Baltimore, MD September 1993 - May 1997

II. Research, Scholarly, Creative and Professional Activites

Annotation: corresponding authors*, lead authors^, advisee#

II.C. Articles in Refereed Journals

- 1. S. Braid^, K. Okrah[#], A. Shetty, **H. Corrada Bravo*** (2016). Differences in DNA methylation patterns in the cord blood of neonates across gestational age are associated with differences in cell-type composition. *Nursing Research, in press*.
- 2. S. Manimaran^, H.M. Selby, K. Okrah[#], C. Ruberman, J.T. Leek, J. Quackenbush, B. Haibe-Kains, **H. Corrada Bravo**, W.E. Johnson* (2016). BatchQC: interactive software for evaluating sample and batch effects in genomic data. *Bioinformatics, in press*.
- 3. M. Pop^, J.N. Paulson*, S. Chakraborty, I. Astrovskaya, B.R. Lindsay, S. Li, **H. Corrada Bravo**, C. Harro, J. Parkhill, A.W. Walker, R.I. Walker, D.A. Sack, O.C. Stine* (2016). Individual-specific changes in the human gut microbiota after challenge with enterotoxigenic *Escherichia coli* and subsequent ciprofloxacin treatment. *BMC Genomics* 17(1):440, doi:10.1186/s12864-016-27777-0.
- 4. M. Sharmin[#], **H. Corrada bravo**, S. Hannenhalli* (2016). Heterogeneity of TF binding specificity models within and across cell lines. *Genome Research* 26(8):1110-23, doi:10.1101/gr.199166.115.
- 5. M.C. Fernandes^, L.A. Dillon^, A.T. Belew, **H. Corrada Bravo**, D.M. Mosser, N. El-Sayed* (2016). Dual transcriptome profiling of human macrophages infected with *Leishmania* parasites reveals transient reprogramming events followed by a generic phagocytic response. *mBio* 7(3). pii: e00027-16, doi:10.1128/mBio.00027-16.
- 6. Y. Li^, K.L. Caradonna^, K. Okrah^#, J. Choi, P. Padmanabhan, D.M. Ndegwa, S. Shah-Simpson, M.R. Temanni, **H. Corrada Bravo**, N. El-Sayed*, B.A. Burleigh* (2016). Transcriptome Remodeling in Trypanosoma cruzi and Human Cells during Intracellular Infection. *PLOS Pathogens* 12(4):e1005511, doi:10.1371/journal.ppat.1005511.

- 7. M. Sharmin^, **H. Corrada Bravo***, S. Hannenhalli* (2016). Characterizing the regulatory landscape of large hypomethylated blocks in colon cancer. *BMC Cancer* 16:88, doi:10.1186/s12885-016-2128-1.
- 8. J. Wagner^#, J. Paulson#, X.S. Wang, B. Bhattacharjee, **H. Corrada Bravo*** (2016). Privacy preserving human microbiome analysis using secure computation. *Bioinformatics*, doi:10.1093/bioinformatics/btw073.
- 9. L.A. Dillon^, R. Suresh, K. Okrah[#], **H. Corrada Bravo**, D.M. Mosser, N.M. El-Sayed* (2015). Simultaneous transcriptional profiling of Leishmania major and its murine macrophage host cell reveals insights into host-pathogen interactions. *BMC Genomics* 16:1108, doi:10.1186/s12864-015-2237-2.
- 10. L. Dillon^, K. Okrah^#, V. Hughitt, R. Suresh, Y. Li, M. Fernandes, A. Belew, **H. Corrada Bravo**, D. Mosser, N. El-Sayed* (2015). Transcriptomic profiling of gene expression and RNA processing during *Leishmania major* differentiation. *Nucleic Acids Research* 43 (14): 6799-6813, doi:10.1093/nar/gkv656.
- 11. W. Dinalankara^#, **H. Corrada Bravo*** (2015). Gene expression signatures based on variability can robustly predict tumor progression and prognosis. *Cancer Informatics* 14:71-81.
- 12. K. Okrah^#, **H. Corrada Bravo*** (2015). Shape analysis for high-throughput transcriptomics experiment data. *Biostatistics* 16(4):627-40. doi:10.1093/biostatistics/kxv018.
- 13. W. Huber[^], V. Carey, R. Gentleman, M. Carlson, B.S. Carvalho, **H. Corrada Bravo**, S. Davis, L. Gatto, T. Girke, R. Gottardo, F. Hahne, K. Hansen, R.A. Irizarry, M. Lawrence, M.I. Love, J. MacDonald, V. Obenchain, A.K. Olés, H. Pagés, P. Shannon, G. Smyth, D. Tenenbaum, L. Waldron, M. Morgan* (2015). Orchestrating high-throughput genomic analysis with Bioconductor. *Nature Methods* 12 115-121, doi:10.1038/nmeth.3252.
- 14. H.S. Parker^, **H. Corrada Bravo**, J.T. Leek* (2014). Removing batch effects for prediction problems with frozen surrogate variable analysis. *PeerJ* 2:e561, doi:10.7717/peerj.561.
- 15. W. Timp^, **H. Corrada Bravo**^, O.G. McDonald, M. Goggins, C. Umbricht, M. Zeiger, A.P. Feinberg*, R.A. Irizarry* (2014). Large hypomethylated blocks related to large heterochromatin regions as a universal defining epigenetic alteration in human solid tumors. *Genome Medicine* 6:61, doi:10.1186/s13073-014-0061-y.
- 16. F. Chelaru^#, L. Smith#, N. Goldstein#, **H. Corrada Bravo*** (2014). Epiviz: interactive visual analytics for epigenomics data. *Nature Methods* 11 938-940, doi:10.1038/nmeth.3038.
- 17. M. Pop^, A.W. Walker^, J.N. Paulson^#, B. Lindsey^, M. Antonio^, M.A. Hossain^, J. Oundo^, B. Tamboura, V. Mai, I. Astrovskaya, **H. Corrada Bravo**, R. Rance, M. Stares, M.M. Levine, S. Panchalingam, K. Kotloff, U.N. Ikumapayi, C. Ebruke, D. Adeyemi, F. Ahmed, M.T. Alam, R. Amin, S. Siddiqui, J.B. Ochieng, E. Ouma, J. Juma, E. Mailu, R. Omore, J.G. Morris, R.F. Breiman, D. Saha, J. Parkhill, O.C. Stine*, J.P. Nataro* (2014). Diarrhea in young children from low-income countries leads to large-scale alterations in intestinal microbiota composition. *Genome Biology* 15, R76.
- 18. M. Aryee^, A. Jaffe, **H. Corrada Bravo**, C. Ladd-Acosta, A. Feinberg, K. Hansen*, R.A. Irizarry* (2014). Minfi: a flexible and comprehensive Bioconductor package for the analysis of Infinium DNA Methylation microarrays. *Bioinformatics* 30 (10):1363-9.
- 19. C. Ye^{^#}, C. Hsiao[#], **H. Corrada Bravo*** (2014). BlindCall: ultra-fast base-calling of second-generation sequencing by blind deconvolution. *Bioinformatics* 30 (9):1214-9.
- 20. N. Akula^, J. Barb, X. Jiang, J. Wendland, K. Choi, S. Sen, L. Hou, D. Chen, G. Laje, K. Johnson, B. Lipska, J. Kleinman, **H. Corrada Bravo**, S. Detera-Wadleigh, P.J. Munson, F.J. McMahon* (2014). RNA-sequencing of brain transcriptome implicates dysregulation of neuroplasticity, circadian rhythms and GTPase binding in bipolar disorder. *Molecular Psychiatry*, doi:10.1038/mp.2013.170.
- 21. E. Alemu^, J.W. Carl, **H. Corrada Bravo***, S. Hannenhalli* (2014). Determinants of expression variability. *Nucleic Acids Research* 42 (6), 3503-14.
- 22. J.N. Paulson[#], O.C. Stein, **H. Corrada Bravo***, M. Pop* (2013). Differential abundance analysis for microbial marker-gene surveys. *Nature Methods* 10 (12):1200-1202, doi:10.1038/nmeth.2658.
- 23. X. He^, R. Chatterjee^, S. John, **H. Corrada Bravo**, B.K. Sathyanarayana, S.C. Biddle, P.C. Fitzgerald, J.A. Stamatoyannopoulos, G.L. Hager, C. Vinson* (2013). Contribution of nucleosome binding preferences and co-occurring DNA sequences to transcription factor binding. *BMC Genomics* 14 (428).

- 24. S. Boca^, **H. Corrada Bravo**, B. Caffo, J.T. Leek*, G. Parmigiani (2013). A decision-theory approach to interpretable set analysis for high-dimensional data. *Biometrics* 69 (3):614-23, doi:10.1111/biom.12060.
- 25. M.L. Nickerson[^], K.M. Im, K.J. Misner, W. Tan, H. Lou, B. Gold, D.W. Wells, **H. Corrada Bravo**, K.M. Fredrikson, T.T. Harkins, P. Milos, B. Zbar, W.M. Linehan, M. Yeager, T. Andresson, M. Dean*, G.S. Bova* (2013). Somatic alterations contributing to metastasis of a castration-resistant prostate cancer. *Human Mutation* 34 (9): 1231-41, doi:10.1002/humu.22346.
- 26. W. Shi^, G. Wahba, R.A. Irizarry, **H. Corrada Bravo**, S.J. Wright* (2012). The Partitioned LASSO-Patternsearch Algorithm with Application to Gene Expression Data. *BMC Bioinformatics* 13:98, doi:10.1186/1471-2105-13-98.
- 27. **H. Corrada Bravo**^, V. Pihur, M. McCall, R.A. Irizarry, J.T. Leek* (2012). Gene expression anti-profiles as a basis for cancer diagnostics. *BMC Bioinformatics* 13:272, doi:10.1186/1471-2105-13-272.
- 28. T.S. Niranjan^, A. Adamczyk^, **H. Corrada Bravo**^, M. Taub, S.J. Wheelan, R.A. Irizarry, T. Wang* (2011). Effective detection of rare variants in pooled DNA samples using Srfim and cross-pool tail-curve analysis. *Genome Biology* 12 (9):R93.
- 29. K. Hansen^, W. Timp^, **H. Corrada Bravo**^, S. Sabunciyan^, B. Langmead^, O.G. McDonald, B. Wen, H. Wu, D. Diep, E. Briem, K. Zhang, R.A. Irizarry*, A.P. Feinberg* (2011). Increased methylation variation in epigenetic domains across cancer types. *Nature Genetics* 43 (8):768-75.
- 30. A. Rivas^, K. Bohane, **H. Corrada Bravo**^, M. Tan, R. Tamargo, H.W. Francis (2011). A model for early prediction of facial nerve recovery after vestibular schwannoma surgery. *Otology & Neurotology* 32 (5):826-33.
- 31. L. Shan^, H.C. Yang, S.A. Rabi, **H. Corrada Bravo**, J.D. Siliciano, R.A. Irizarry, H. Zhang, J. Margolick, R.F. Siliciano* (2011). Influence of host gene transcription level and orientation on HIV-1 latency in a primary cell model. *Journal of Virology* 85 (11):5384-93.
- 32. M. Taub^, **H. Corrada Bravo**, R.A. Irizarry* (2010). Overcoming bias and systematic errors in next generation sequencing data. *Genome Medicine* 2 (12):87.
- 33. J.T. Leek^, R. Scharpf, **H. Corrada Bravo**, D. Simcha, B. Langmead, W.E. Johnson, D. Geman, K. Baggerly, R.A. Irizarry* (2010). Tackling the widespread and critical impact of batch effects in high-throughput data. *Nature Reviews Genetics* 11 (10):733-739.
- 34. H. Wu, R.A. Irizarry*, **H. Corrada Bravo**^(2010). Intensity normalization improves color calling in SOLiD sequencing. *Nature Methods* 7:336-337.
- 35. M. Acevedo^, T.M. Aide*, L. J. Villanueva-Rivera, **H. Corrada Bravo**^, C. J. Corrada-Bravo* (2009). Automated classification of bird and amphibian calls using machine learning: a comparison of methods. *Ecological Informatics* 4 (4):206-214.
- 36. **H. Corrada Bravo**^, R.A. Irizarry* (2009). Model-based quality assessment and base-calling for second-generation sequencing data. *Biometrics* 66(3):665-74, doi:10.1111/j.1541-0420.2009.01353.x.
- 37. **H. Corrada Bravo**^, K.E. Lee, B.E.K. Klein, R. Klein, S.K. Iyengar, G. Wahba* (2009). Examining the relative influence of familial, genetic and environmental covariate information in flexible risk models. *Proceedings of the National Academy of Science* 106 (20): 8128-8133.
- 38. K. H. Eng^, **H. Corrada Bravo**^, S. Keles* (2009). A phylogenetic mixture model for the evolution of gene expression. *Molecular Biology and Evolution* 26 (10):2363-2372.
- 39. A. T. Evan, R. Bennartz, V. Bennington, **H. Corrada Bravo**, A. K. Heidinger, N. M. Mahowald, C. S. Velde, G. Myhre, J. P. Kossin (2008). Ocean temperature forcing by aerosols across the Atlantic tropical cyclone development region. *Geochem, Geophys. Geosyst.* 9:Q05V04, doi:10.1029/2007GC001774.
- 40. C. Kuang, P. McMurry, A. McCormick, F. Eisele, S.H. Lee, L.H. Young, D.R. Benson, et al. (2007). A system for operational aerosol optical depth data assimilation over global oceans. *J. Geophys. Res* 113(D10):D10208.

II.D. Published Conference Proceedings

II.D.1. Refereed Conference Proceedings

- 1. F. Dorri^#, L. Mendelowitz, **H. Corrada Bravo*** (2015). methylFlow: cell-specific methylation pattern reconstruction from high-throughput bisulfite-converted DNA sequencing. *HitSEQ 2015; Bioinformatics, in press.*
- 2. F. Chelaru^**, **H. Corrada Bravo*** (2015). Epiviz: a view inside the design of an integrated visual analysis software for genomics. *BioVis 2015; BMC Bioinformatics 16(Supl 11):S4*.
- 3. **H. Corrada Bravo**^, K. Eng, S. Keles, G. Wahba, S. Wright (2009). Estimating tree-structured covariance matrices via mixed integer programming. *Twelfth International Conference on Artificial Intelligence and Statistics (AISTATS '09); Journal of Machine Learning Research Workshop and Conference Proceedings*, 533:40.
- 4. **H. Corrada Bravo**^, R. Ramakrishnan (2007). Optimizing MPF queries: decision support and probabilistic inference. *26th ACM SIGMOD Intl. Conf. on Management of Data*701:712.
- 5. **H. Corrada Bravo**^, D. Page, R. Ramakrishnan, J. Shavlik, V. Santos Costa (2005). A framework for set-oriented computation in inductive logic programming and its application in generalizing inverse entailment. *15th ILP Conf*.69:86.

II.E. Conferences, Workshops and Talks

II.E.2. Invited Talks

- 1. What is Biomedical Data Science?. *Symposium of Health Informatics in Latin America and the Caribbean*, San Juan, PR. November 2015.
- 2. Visualization, Statistical Modeling and Discovery in Computational Epigenomics. *Computer Science Colloquium Series*, College Park, MD. September 2015.
- 3. Interactive and exploratory visualization of epigenome-wide data. *BioIT World Conference*, Boston, MA. April 2015.
- 4. Interactive and exploratory visualization of epigenome-wide data. *UMD Campus Visualization Partnership Lecture Series*, College Park, MD. February 2015.
- 5. Interactive and exploratory visualization of epigenome-wide data. *Epigenomics in Disease, Molecular Medicine Tri-Con*, San Francisco, CA. February 2015.
- 6. Exploring tumor epigenetic heterogeneity by cell-specific methylation pattern reconstruction. *Department of Biostatistics and Computational Biology, Johns Hopkins Cancer Center,* Baltimore, MD. November 2014.
- 7. Exploring tumor epigenetic heterogeneity by cell-specific methylation pattern reconstruction. *CMU-Pitt Ph.D. Program in Computational Biology Seminar Series*, Pittsburgh, PA. April 2014.
- 8. Cell-specific methylation pattern reconstruction using minimum cost network flow algorithms. *Department of Mathematics, George Mason University,* Fairfax, VA. February 2014.
- 9. Gene expression anti-profiles as a basis for accurate universal cancer signatures. *Greenbaum Cancer Center, University of Maryland School of Medicine, Baltimore, MD. September 2013.*
- 10. Gene expression anti-profiles as a basis for accurate universal cancer signatures. *Institute for Genome Sciences, University of Maryland School of Medicine, Baltimore, MD. March 2013.*

- 11. Gene expression anti-profiles as a basis for accurate universal cancer signatures. *Department of Bioinformatics and Computational Biology, Genentech, Inc.*, South San Franciso, CA. February 2013.
- 12. Gene expression anti-profiles as a basis for accurate universal cancer signatures. *Innovation Center for Biomedical Informatics, Georgetown University,* Washington, DB. December 2012.
- 13. Statistical and computational methods for the analysis of pooled, targeted, second-generation re-sequencing data. *Biostatistics Department, University of Alabama*, Birmingham, AL. April, 2012.
- 14. Modeling gene expression variability for prediction in disease populations. *Department of Biostatistics, Columbia University School of Public Health,* New York, NY. December 2010.
- 15. Modeling gene expression variability for prediction in disease populations. *Johns Hopkins University School of Medicine*, Baltimore, MD. December 2010.
- 16. Gene expression variability in disease populations. *National Cancer Institute*, Bethesda, MD. October 2010.
- 17. Modeling uncertainty in second-generation sequencing data. *Dept. of Biostatistics, Harvard School of Public Health*, Boston, MA. November 2009.
- 18. Model-based quality assessment and base-calling for second-generation sequencing data. *University of Wisconsin-Milwaukee*, Milwaukee, WI. October 2009.
- 19. Modeling and managing uncertainty in second-generation sequencing data. *Dept. of Computer Science, University of Maryland,* College Park, MD. October 2009.
- 20. Model-based quality assessment and base-calling for second-generation sequencing data. *University of Manchester*, Manchester, England. October 2009.
- 21. Model-based quality assessment and base-calling for second-generation sequencing data. *University of Wisconsin*, Madison, WI. May 2009.
- 22. Kernel methods for examining the relative influence of familial, genetic and environmental covariate information in risk models: results and (more importantly) extensions. *University of Wisconsin*, Madison, WI. May 2009.
- 23. Model-based quality assessment and base-calling for second-generation sequencing data. *Case Western Reserve University*, Cleveland, OH. April 2009.
- 24. Data analysis at the computational/statistical sciences borderland: two examples from genomics. *Johns Hopkins Bloomberg School of Public Health*, Baltimore, MD. February 2009.
- 25. Estimating tree-structured covariance matrices via mixed-integer programming. *Johns Hopkins School of Public Health,* Baltimore, MD. January 2008.

II.E.3. Refereed Presentations

- 1. Differential abundance analysis of metagenomic whole-genome sequencing. *Joint Statistical Meetings*, Seattle, WA. August 2015.
- 2. Addressing reproducibility in genomic signatures by characterizing variance and estimation stability. *Joint Statistical Meetings*, Seattle, WA. August 2015.
- 3. Epiviz: a view inside the design of an integrated visual analysis software for genomics. *BioVis*, Dublin, Ireland. July 2015.

- 4. Interactive and exploratory visual analytics of epigenome-wide data. ISMB, Dublin, Ireland. July 2015.
- 5. methylFlow: cell-specific methylation pattern reconstruction from high-throughput bisulfite-converted DNA sequencing. *HiTSeq*, Dublin, Ireland. July 2015.
- 6. Interactive and exploratory visualization of epigenome-wide data. *Joint Statistical Meetings*, Boston, MA. July 2014.
- 7. Interactive, Exploratory Visualization and Statistical Analysis of Genome-Scale Data. *International Biomteric Society ENAR Meeting*, Baltimore, MD. March 2014.
- 8. Gene expression anti-profiles as a basis for accurate universal cancer signatures. *ISMB '13*, Berlin, Germany. July 2013.
- 9. Srfim2: using basecalling model parameter estimates to understand sequencing bias. *2012 Joint Statistical Meetings*, San Diego, CA. August 2012.
- 10. Increased methylation variation in epigenetic domains across cancer types. *16th Annual International Conference on Research in Computational Molecular Biology (RECOMB)*, Barcelona, Spain. April 2012.
- 11. Statistical and computational methods for the analysis of pooled, targeted, second-generation re-sequencing data. *2011 Joint Statistical Meetings*, Miami Beach, FL.. August 2011.
- 12. Model-based quality assessment and base-calling for second-generation sequencing data. *WNAR/IMS annual meeting*, Portland, OR. June 2009.
- 13. Tuning regularized kernel estimation parameters for prediction. *SIAM Conference on Optimization*, Boston, MA. May 2008.
- 14. Optimizing MPF queries: decision support and probabilistic inference. *26th ACM SIGMOD Intl. Conf. on Management of Data*, Beijing, China. June 2007.
- 15. A framework for set-oriented computation in inductive logic programming and its application in generalizing inverse entailment.. *15th ILP Conf.*, Bonn, Germany. August 2005.

II.E.7. Non-Refereed Presentations

- 1. Does Healthcare Insurance Matter? Establishing Insurance Status as a Risk Factor for Trauma Mortality. *INFORMS 2014*, San Francisco, CA. November 2014.
- 2. Epiviz(r): turning a genome browser into a display device. *Bioconductor conference*, Boston, MA. July 2014.
- 3. Statistical and computational methods for the analysis of pooled, targeted, second-generation re-sequencing data. *8th International Chinese Statistical Association (ICSA) International Conference*, Guangzhou, China. December 2010.
- 4. Model-based quality assessment and base-calling for second-generation sequencing data. *Conference on Next-Generation Sequencing*, Barcelona, Spain. October 2009.

II.E.11. Symposia

1. Gene expression network anti-profiles. *UMD/NIST Network Science Mini-Symposium*, College Park, MD. January, 2014.

- 2. Increased methylation variation in epigenetic domains across cancer types. *Omics Day, University of Maryland,* Shady Grove, MD. May 2012.
- 3. Model-based quality assessment and base-calling for second-generation sequencing data. *Third Annual Young Investigators Symposium on Genomics and Bioinformatics*, Baltimore, MD. September 2009.

II.E.12. Workshops

- 1. Gene expression network anti-profiles. *UMD/NIST Network Science Mini-Symposium*, College Park, MD. January, 2014.
- 2. Increased methylation variation in epigenetic domains across cancer types. *Omics Day, University of Maryland,* Shady Grove, MD. May 2012.
- 3. Model-based quality assessment and base-calling for second-generation sequencing data. *Third Annual Young Investigators Symposium on Genomics and Bioinformatics*, Baltimore, MD. September 2009.

II.F. Professional Publications

II.F.2. Pre-print/Working Paper (Not Work in Progress)

- 1. J. Wagner^{^#}, J.N. Paulson[#], X.S. Wang, B. Bhattacharjee, **H. Corrada Bravo*** (2015). Privacy-Preserving Microbiome Analysis Using Secure Computation. *biorxiv* http://dx.doi.org/10.1101/025999.
- 2. M. Sharmin^#, **H. Corrada Bravo***, S. Hannenhalli* (2015). Distinct genomic and epigenomic features demarcate hypomethylated blocks in colon cancer. *biorxiv* http://dx.doi.org/10.1101/028803.
- 3. M. Sharmin[#], **H. Corrada Bravo**, S. Hannenhalli* (2015). Heterogeneity of Transcription Factor binding specificity models within and across cell lines. *biorxiv* http://dx.doi.org/10.1101/028787.
- 4. H.S. Parker^, **H. Corrada Bravo**, J.T. Leek (2013). Removing batch effects for prediction problems with frozen surrogate variable analysis. *arXiv preprint server* arXiv:1031.3947 [stat.ME].
- 5. W. Dinalankara^{*}, **H. Corrada Bravo** (2013). Anomaly classification with the anti-profile support vector machine. *arXiv preprint server* arXiv:1301.3514 [stat.ML].
- 6. S. Boca^, **H. Corrada Bravo**, B. Caffo, J.T. Leek, G. Parmigiani (2010). A decision-theory approach to interpretable set analysis for high-dimensional data. *Johns Hopkins University, Dept. of Biostatistics Working Papers* Working paper 211. http://biostats.bepress.com/jhubiostat/paper211.
- 7. R.A. Irizarry, **H. Corrada Bravo**^(2009). Model-based quality assessment and base-calling for second-generation sequencing data. *Johns Hopkins University, Dept. of Biostatistics Working Papers* Working paper 184. http://biostats.bepress.com/jhubiostat/paper184.

II.H. Completed Creative Works

II.H.8. Software and Applications

- 1. Epiviz: Interactive visualization for genomics data. Released June 2013. http://github.com/epiviz.
- 2. Epivizr: interactive visualization of genomics data in R/Bioconductor. Released June 2013. http://bioconductor.org/packages/release/bioc/html/epivizr.html.
- 3. antiProfiles: Gene expression anti-profiles as a basis for accurate universal cancer signatures. Released April 2013. http://bioconductor.org/packages/release/bioc/html/antiProfiles.html.

- 4. Healthvis: Interactive visualization in health. Released April 2013. http://healthvis.org.
- 5. Minfi: Analyze Illumina's 450k methylation arrays. Released Febraury 2013. http://bioconductor.org/packages/release/bioc/html/minfi.html.
- 6. metagenomeSeq: Statistical analysis for sparse high-throughput sequencing. Released February 2013. http://bioconductor.org/packages/release/bioc/html/metagenomeSeq.html.
- 7. bumphunter: Tools for finding bumps in genomic data.
 Released February 2013. http://bioconductor.org/packages/release/bioc/html/bumphunter.html.
- 8. Servic4e: Effective detection of rare variants in pooled DNA samples. Released November 2011. http://www.cbcb.umd.edu/~hcorrada/secgen.
- 9. Rsolid: Intensity normalization for SOLiD sequencing. Released April 2010. http://www.cbcb.umd.edu/~hcorrada/secgen.
- 10. Srfim: Model-based base-calling and quality assessment for second-generation sequencing. Released April 2010. http://www.cbcb.umd.edu/~hcorrada/secgen.
- 11. Rcsdp: An interface to the CSDP semidefinite programming library for R. Released Dec. 2008. http://cran.r-project.org/web/packages/Rcsdp/index.html.
- 12. Rcplex: An interface to the CPLEX optimization engine for R. Released Jan. 2008. http://cran.r-project.org/web/packages/Rcplex/index.html.

II.H.9. Websites

1. Epiviz: Interactive visualization for genomics data. Released May 2014. http://epiviz.cbcb.umd.edu/help.

II.J. Sponsored Research

II.J.1. Grants

1. Title: R01: Overcoming bias and unwanted variability in next generation sequencing

Funding Agency: NIH HG005220

Recipient Institution: Dana Farber Cancer Institute, Rafael A. Irizarry (PI)

Amount Awarded: \$397,903 subcontracted to UMCP

Dates: March 2015-February 2019

Role: PI

2. Title: R01: Integrative visual and computational exploratory analysis of genomics data

Funding Agency: NIH GM114267

Recipient Institution: University of Maryland, College Park

Amount Awarded: \$1,800,421

Dates: September 2015-August 2019

Role: PI

3. Title: U54: Longitudinal multiomics microbial profiling in healthy and disease individuals

Funding Agency: NIH

Recipient Institution: Stanford University, Michael P. Snyder (PI)

Amount Awarded: \$100,702 subcontracted to UMCP

Dates: September 2015-December 2016

Role: PI

4. Title: NRT-DESE; Network biology: from data to information to insights

Funding Agency: NSF

Recipient Institution: University of Maryland, College Park

Amount Awarded: \$2,999,847 Dates: September 2016-August 2021

Role: co-PI

5. Title: R01: Analysis tools and software for second generation sequencing

Funding Agency: NIH HG005220

Recipient Institution: Johns Hopkins University, Rafael A. Irizarry (PI)

Amount Awarded: \$380,400 subcontracted to UMCP (\$1,230,000 total awarded by NIH)

Dates: August 2010-May 2013

Role: PI

6. Title: R01: Alignment software for second generation sequencing

Funding Agency: NIH HG006102

Recipient Institution: Johns Hopkins University, Steven Salzberg (PI)

Amount Awarded: \$432,259.00 subcontracted to UMCP (\$3,585,436 total awarded by NIH)

Dates: May 2011-April 2014

Role: PI

II.K. Fellowships, Gifts and Other Funded Research

II.K.1. Fellowships

- 1. Ford Fellowship, National Academies of Science.
- 2. Advanced Opportunity Fellowship, University of Wisconsin-Madison.

II.L. Submissions and Works in Progress

II.L.1. Current Grant Applications

II.L.2. Manuscripts in Preparation

II.L.3. Manuscripts under Review

1. J.N. Paulson^#, H. Talukder^#, **H. Corrada Bravo**. Longitudinal differential abundance analysis of microbial marker-gene surveys using smoothing splines. *Statistics in Medicine, submitted*.

III. Teaching, Advising and Mentoring

III.A. Courses Taught

Term	Course	Enrollment	Description
Spring 2016	CMSC 320	50	Introduction to Data Science

Term	Course	Enrollment	Description
Spring 2016	CMSC 898	1	Pre-candidacy Research (Individual instruction)
Spring 2016		1	Doctoral Dissertation Research (Individual instruction)
Fall 2015	CMSC 898	3	Pre-candidacy research (Individual instruction)
Spring 2015	CMSC 498T	32	Introduction to Data Science II
Fall 2015	CMSC 423	49	Bioinformatics Databases, Tools and Algorithms
Spring 2015	CMSC 899	2	Doctoral dissertation research (Individual instruction)
Spring 2015	CMSC 898	1	Pre-candidacy research (Individual instruction)
Fall 2015	CMSC 499A	1	Independent Undergraduate Research (Individual instruction)
Spring 2015	AMSC 899	2	Doctoral dissertation research (Individual instruction)
Fall 2014	CMSC 423	49	Bioinformatics Databases, Tools and Algorithms
Spring 2014	CMSC 702	30	Computational Systems Biology and Functional Genomics
Fall 2014	CMSC 423	35	Bioinformatics Databases, Tools and Algorithms
Spring 2014	CMSC 899	1	Doctoral dissertation research (Individual instruction)
Spring 2014	AMSC 899	3	Doctoral dissertation research (Individual instruction)
Fall 2014	AMSC 899	2	Doctoral dissertation research (Individual instruction)
Fall 2014	CMSC 798	1	Graduate seminar (Individual instruction)
Fall 2014	AMSC 760	1	Applied Statistics Practicum (Individual instruction)
Fall 2014	CMSC 899	2	Doctoral dissertation research (Individual instruction)
Spring 2014	CMSC 898	3	Pre-candidacy research (Individual instruction)
Fall 2014	CMSC 898	1	Pre-candidacy research (Individual instruction)
Spring 2013	CMSC 702	35	Computational Systems Biology and Functional Genomics
Spring 2013	CMSC 798	1	Graduate seminar (Individual instruction)
Spring 2013	AMSC 899	1	Pre-candidacy research (Individual instruction)
Spring 2013	AMSC 898	3	Pre-candidacy research (Individual instruction)
Spring 2013	CMSC 898	2	Pre-candidacy research (Individual instruction)
Fall 2013	AMSC 689	1	Research Interactions: Regularized Regression Methods
Fall 2013	CMSC 423	48	Bioinformatics Databases, Tools and Algorithms
Fall 2013	AMSC 899	3	Doctoral dissertation research (Individual instruction)
Fall 2013	CMSC 898	4	Pre-candidacy research (Individual instruction)
Fall 2013	AMSC 898	1	Pre-candidacy research (Individual instruction)
Fall 2012	CMSC 726	48	Machine Learning
Fall 2012	CMSC 898	1	Pre-candidacy research (Individual instruction)
Fall 2012	AMSC 899	1	Doctoral dissertation research (Individual instruction)
Fall 2012	AMSC 898	3	Pre-candidacy research (Individual instruction)
Spring 2012	CMSC 351	90	Introduction to Algorithms
Spring 2012	CMSC 858B	26	Computational Systems Biology and Functional Genomics
	CMSC 898	1	Pre-candidacy research (Individual instruction)
Spring 2012	AMSC 898		· · · · · · · · · · · · · · · · · · ·
Spring 2012		4	Pre-candidacy research (Individual instruction)
Spring 2011	CMSC 858P	17	Computational Methods for High-Throughput Analysis
Spring 2011	AMSC 900	1	of Biological Systems Pro-candidacy research (Individual instruction)
Spring 2011	AMSC 898	1 12	Pre-candidacy research (Individual instruction) Team tought graduate 'Programming for Piologists' course
Fall 2011	CBMG 688P	12	Team-taught graduate 'Programming for Biologists' course
Fall 2011	AMSC 898	3	Pre-candidacy research (Individual instruction)
3rd Term 2010	140.644	16	Practical Machine Learning (Johns Hopkins University
F-11 2010	CDMC coop	10	School of Public Health, Department of Biostatistics)
Fall 2010	CBMG 688P	12	Team-taught graduate 'Programming for Biologists' course

III.B. Teaching Innovations

III.B.5. Course or Curriculum Development

CMSC 498 - Introduction to Data Science II, Exploring, Modeling and Communicating with Data

This is the second of two new courses covering the practice of data science. This course focuses on exploratory and statistical data analysis, data and information visualization, and the presentation and communication of analysis

results. It is heavily assignment-based and draws extensively from applications.

CMSC 320 - Introduction to Data Science II

Developed a new undergraduate course covering the practice of data science, and a required course on a new Data Science specialization within the undergraduate program of the Department of Computer Science. This course focuses on data management, exploratory and statistical data analysis, data and information visualization. It is heavily assignment-based and draws extensively from applications.

CMSC 702 - Computational Systems Biology and Functional Genomics

Developed this new advanced graduate-level course concentrating on the use of Statistical Learning methods and algorithms in the analysis of large high-throughput biomedical assays. Core course in the Computational Biology concentration area.

CMSC 858P - Computational Methods for High-Throughput Analysis of Biological Systems

Developed this new advanced graduate-level course concentrating on the use of Statistical Learning methods and algorithms in the analysis of large high-throughput biomedical assays. First offering attracted 21 students, including students from ECE, AMSC and BISI. This course was merged with other content to become CMSC 702 (Computational Systems Biology) to be offered as a regular course in the CS department starting in Spring 2012.

140.644 - Practical Machine Learning

Developed this new advanced graduate-level course as postdoc at the Johns Hopkins University School of Public Health. The course provided students from a variety of backgrounds with practical knowledge of Machine Learning models and applications.

III.C. Advising

III.C.1. Undergraduate

Name	Department	Stage	Role
Daniel Konecki	CS	Completed Baylor College of Medicine	advisor
Mihai Sirbu	CS	Current	advisor

III.C.2. Master's

III.C.3. Doctoral

Name	Department	Stage	Role
Faezeh Dorri	CS	Pre-candidacy	advisor
Justin Wagner	CS	Candidacy	advisor
Senthil Muthiah	CBMG	Candidacy	co-advisor
Mohammed Gunady	CS	Pre-candidacy	advisor
Mahfuza Sharmin	CS	Candidacy	co-advisor
Nathanael Olson	CBMG	Pre-candidacy	advisor
Florin Chelaru	CS	Completed	advisor
		Postdoc, MIT Broad Institute	
Chiao-Wen Hsiao	AMSC	Completed	advisor
		Postdoc, University of Chicago	
Kwame Okrah	AMSC	Completed	advisor
		Biostatistician, Genentech Inc.	
Wikum Dinalankara	CS	Completed	advisor
		Postdoc, Johns Hopkins University Medicine	

Name	Department	Stage	Role
Hisham Talukder	AMSC	Completed	advisor
		Data Scientist, Dow Jones	
Joseph Paulson	AMSC	Completed	co-advisor
		Postdoc, Harvard/Dana-Farber Cancer Institute	
Chengxi Ye	CS	Pre-candidacy	committee member
Viet-An Nguyen	CS	Completed	committee member
		Data Scientist, Facebook	
Justin Malin	CBMG	Completed	committee member
		Postdoc, National Institutes of Health	
Ted Gibbons	CBMG	Candidacy	committee member
Steven Smith	CBMG	Candidacy	committee member
Cody Buntain	CS	Candidacy	committee member
David Kelley	CS	Completed	committee member
		Postdoc, Broad Institute	
Guillaume Marcais	CS	Completed	committee member
		Research Scientist, University of Maryland	
Bhargav Kanagal	CS	Completed	committee member
Yuan Li	CBMG	Completed	committee member
Mohammadreza Ghodsi	CS	Completed	
		Software Engineer, Facebook	
Ginger Houston-Ludlam	CBMG	Candidacy	committee member
Louis Licamele	CS	Completed	committee member
Benjamin Langmead	CS	Completed	committee member
		Assistant Professor, Johns Hopkins University	
Rob Partro	CS	Completed	committee member
		Postdoc, Carnegie Mellon University	
Bo Liu	CS	Completed	committee member
		Software Engineer, Google	
Charles Dawkins	CBMG	Pre-candidacy	committee member
Daehwan Kim	CS	Completed	committee member
		Postdoc, Johns Hopkins University	
Laura Dillon	CBMG	Completed	committee member
Shane Squires	Physics	Completed	committee member
Srutii Sarda	CBMG	Candidacy	committee member
Kun Wang	CBMG	Candidacy	committee member
Vahid Liaghat	CS	Completed	committee member
Adam Bazinet	CS	Completed	committee member
		Research Scientist, University of Maryland	
Sameh Khamis	CS	Completed	committee member
Snighda Chaturvedi	CS	Candidacy	committee member
Alex Malozemoff	CS	Candidacy	committee member
Chris Hill	CS	Completed	committee member
		Postdoc, University of Washington	
Jose Carrillo	ANSC	Candidacy	committee member
Stephen Xi Chen	CS	Completed	committee member
Yang Shen	Chemical Physics	Candidacy	committee member
Jay Pujara	CS	Completed	committee member
Jayanta Mondal	CS	Completed	committee member
Mossaab Bagdouri	CS	Candidacy	committee member
Avinash Sahu	CS	Completed	committee member
Mohamed Rastegari	CS	Completed	committee member
Sana Malek	CS	Candidacy	committee member

III.F. Professional and Extension Education

III.F.3. Workshops

- 1. Big Data in Social Sciences: Introduction to Data Science with R. *Monterrey Technological College,* Mexico City, Mexico. June 2015.
- 2. Computational and statistical challenges in second-generation sequencing data analysis. *Emerging Biotechnologies, Georgetown University Masters in Biotechnology program,* Washington, DC. February and March 2012.
- 3. Topics in modern biology: omics: new tools and new areas. *Guest lecturer, Department of Biology, University of Puerto Rico*, San Juan, PR. March 2011.
- 4. Computational and statistical challenges in second-generation sequencing data analysis. *Emerging Biotechnologies, Georgetown University Masters in Biotechnology program,* Washington, DC. March 2011.
- 5. Model-based base-calling and quality assessment for second-generation sequencing data. *Integrative Statistical Analysis of Genome Scale Data. Cold Spring Harbor Laboratory,* Cold Spring Harbor, NY. June 17, 2010.
- 6. Introduction to R and Bioconductor for Microarray Analysis. PASI, Guanajuato, México. May 4-5, 2010.
- 7. Clustering and Classification of Gene Expression Data. *Workshop. Johns Hopkins University Center for Computational Genomics*, Baltimore, MD. May 14, 2010.

IV. Service and Outreach

IV.A. Editorships, Editorial Boards, and Reviewing Activities

IV.A.1. Editorships

1. Special Issue Editor, Proceedings IEEE. 2013-2015

IV.A.3. Reviewing Activities for Journals and Presses

- 1. Bioinformatics
- 2. Annals of Applied Statistics
- 3. Journal of Machine Learning Research
- 4. Biostatistics
- 5. Journal of Artificial Intelligence Research
- 6. Genome Biology
- 7. Journal of the Royal Statistical Society (Series C)
- 8. IEEE Transactions on Computational Biology and Bioinformatics
- 9. BMC Genomics
- 10. Biometrics
- 11. BMC Bioinformatics

- 12. Genome Research
- 13. Nucleic Acids Research
- 14. Nature Communications
- 15. Nature Methods
- 16. New England Journal of Medicine
- 17. Nature Scientific Reports
- 18. Manning Publications Press
- 19. Chapman & amp; Hall/CRC

IV.A.4. Reviewing Activities for Agencies and Foundations

- 1. NSF review panel 2011
- 2. NIH ad-hoc review panel 2014
- 3. Austrian Science Fund
- 4. NSF review panel 2016
- 5. NCI review panel 2016

IV.A.5. Reviewing Activities for Conferences

- 1. ISMB/ECCB 2013
- 2. ISMB 2014
- 3. ISMB/ECCB 2015
- 4. ISMB 2016
- 5. ACM-BCB 2014
- 6. AMC-BCB 2015

IV.B. Committees, Professional and Campus Service

IV.B.1. Campus Service - Department

- Computational Biology Field Committee. 2010-
- UMIACS APT Committee. 2013-2015
- Computer Science Recruitment Committee. 2014-2015
- CBBG Graduate Admissions Committee, 2012-2014
- Center for Women in Computing Advisory Board. 2014-2015
- Strategic Planning Subcommittee on Diversity and Outreach. 2014-2015

IV.B.2. Campus Service - College

• Computer Science Chair Search Committee. December 2011-March 2012

IV.B.7. Leadership Roles in Meetings and Conferences

• Organizing Committee, Biovis Workshop. 2015-2016

IV.C. External Service and Consulting

IV.C.1. Community Engagements, Local, State, National, International

- City of Hyattsville Council Educational Advisory Committee. September 2014-August 2016
- Outreach: computer science education after-school club, Hyattsville Elementary School. March-April 2015

IV.C.5. Consultancies

• Visualization in Bioconductor. *Genentech*, September 2013-December 2014.

V. Awards, Honors and Recognition

V.D. Recognition in Media

- Researchers develop tool to better visualize, analyze human genomic data, *Genetics Times*, 11 Aug 2014.
- New Tool To Better Visualize, Analyze Human Genomic Data, Counsel and Heal, 04 Aug 2014.
- UMD Researchers Develop Tool to Better Visualize, Analyze Human Genomic Data, *Science Newsline*, 04 Aug 2014.
- UMD Researchers Develop Tool to Better Visualize, Analyze Human Genomic Data, *Technobahn*, 04 Aug 2014.
- New web-based tool to better analyse human genomic data, Vancouver Verdesi, 04 Aug 2014.
- New web-based tool to better analyse human genomic data, Yahoo! News India, 04 Aug 2014.
- New web-based tool to better analyse human genomic data, Business Standard, 04 Aug 2014.
- New web-based tool enables researchers to quickly visualize genomic information, *The Medical News*, 04 Aug 2014.
- UMD researchers develop tool to better visualize, analyze human genomic data, *Innovations Report*, 04 Aug 2014.
- Researchers Develop Tool to Analyze Human Genomic Data, *Medindia*, 04 Aug 2014.
- Tool to better visualize, analyze human genomic data developed, Science Daily, 03 Aug 2014.
- Team develops tool to better visualize, analyze human genomic data, MedicalXpress, 03 Aug 2014.
- UMD researchers develop tool to better visualize, analyze human genomic data, EurekAlert!, 03 Aug 2014.