## C. Titus Brown

(As of June 2020.)

EDUCATION

Reed College, Portland, OR; Mathematics; B.A., 1997

California Institute of Technology, Davidson Lab (graduate student);

Developmental Biology; PhD., 2007

California Institute of Technology, Bronner-Fraser Lab (postdoc);

Developmental Biology and Bioinformatics; 2007-2008

APPOINTMENTS

**Assistant Professor**, Microbiology & Molecular Genetics / Computer Science and Engineering Michigan State University, 2008-2014.

**Associate Professor**, Population Health and Reproduction, School of Veterinary Medicine University of California Davis, 2015-present.

Honours and Awards Burroughs-Wellcome Fund Computational Biology Fellowship (1999-2004).

Withrow Award for Teaching Excellence in Computer Science at MSU (2008-2009).

Woods Hole Marine Biological Laboratory Summer Fellow (2013).

Michigan State University / College of Natural Science Teacher-Scholar Award (2013).

Moore Foundation Data Driven Discovery Investigator (2014-2019).

AWARDS

- 20. NSF \$297,922 (100%), 3 years, 8/20-7/23, notified of funding and pending award receipt. "BBSRC-NSF/BIO:Collaborative Research: genomeRxiv: a microbial whole-genome database and diagnostic marker design resource for classification, identification, and data sharing."
- 19. USDA \$498,356 (100%) 3 years, 3/20-2/23. "Tools and resources for cattle pangenomics."
- 18. NIH, \$1,237,582 (100%); 12 months, 1/20-12/20. "The Common Fund Data Ecosystem", 2nd year funding.
- 17. NIH, \$374,645 (100%); 8 months, 4/19-12/19. "The Common Fund Data Ecosystem", 1st year funding.
- 16. NIH, \$1.9m (100%); 18 months, 10/16-4/18. NIH Data Commons Consortium Pilot Phase.
- 15. Department of Homeland Security, \$248,000 (100%); 6/17-9/18. "High Confidence Metagenomics Analysis of Complex Samples."
- 14. Sloan Foundation, \$20,000 (100%); 7/1/16-12/31/16. "A Workshop on Dockerized Notebook Computing."
- 13. Moore Foundation, \$1,750,000 (100%); 12/1/14-12/1/21. "Software to support data driven inquiry in biology."
- 12. USDA, \$129,889 (100%); 10/1/13-9/30/16. Collaborative proposal with WUSTL. "Improvement of the Chicken Reference Genome."
- 11. NSF, \$273,419 (20%); 9/2013-9/2016. co-PI, "Supporting Efficient Discrete Box Queries for Sequence Analysis on Large Scale Genome Databases."

- 10. NIH R25, \$162,000 (100%); 7/2013-7/2016. PI, "Analyzing Next-Generation Sequencing Data." Renewal.
- 9. NIH R01, \$699,231 (100%). 5/2013-5/2016. PI, "BIGDATA: Low-Memory Streaming Prefilters for Biological Sequencing Data."
- 8. USDA, \$2,989,032 (5.8%). 1/2013-1/2017. co-PI, "The Genetics of Johne's Disease."
- 7. NSF OCI Supplement to BEACON STC, \$200k (100%); 1/1/2013-12/31/2013. PI, "Materials and Workshops for Cyberinfrastructure Education in Biology."
- USDA (Grad Fellowships), \$238,000 (15%). 1/2012-11/2017. co-PI, "Integrated genomics training program."
- 5. DOE, \$659,587 (5%). 9/2011-8/2012. co-PI, "Ribosomal Database Project."
- 4. NIH (R25 education), \$104,000 (100%); 7/2011-6/2014. PI, "Analyzing Next-Generation Sequencing Data."
- 3. USDA, \$99,000; 2/2009-12/2011 (100%). PI, "Positional Candidate Genes for Resistance to Marek's Disease by Screening for Marek's Disease Virus Meq-regulated Genes."
- 2. NSF, \$50,000; 9/2009-8/2011 (100%). PI, "RV1: MSB: Collaborative: Symbiont Separation and Investigation of the Novel Heterotrophic Osedax Symbiosis using Comparative Genomics."
- 1. USDA, \$690,000; 12/2009-11/2013 (100%). PI, "Easily accessible Web-based tools for analyzing next-generation sequencing data from agricultural animals."

Note: ▶ highlights the most important papers, in my opinion (personal contribution/impact).

Whole genome assembly of Culex tarsalis. BJ Main, CT Brown, M Marcantonio, CM Barker. bioRxiv.

Draft genome assemblies using sequencing reads from Oxford Nanopore Technology and Illumina platforms for four species of North American killifish from the Fundulus genus, LK Johnson, R Sahasrabudhe, T Gill, J Roach, L Froenicke, CT Brown, A Whitehead. BioRxiv.

Microbial contamination in the genome of the domesticated olive. T Reiter, CT Brown. bioRxiv.

▶ A Reference-Free Algorithm for Computational Normalization of Shotgun Sequencing Data. Brown CT, Howe AC, Zhang Q, Pyrkosz AB, Brom TH. 2012. arXiv:1203.4802.

Crossing the streams: a framework for streaming analysis of short DNA sequencing reads. QP Zhang, S Awad, CT Brown. 2015. https://peerj.com/preprints/890/.

Evaluating Metagenome Assembly on a Simple Defined Community with Many Strain Variants. S Awad, LC Irber, CT Brown. 2017. https://www.biorxiv.org/content/early/2017/07/03/155358.

Preprints

PEER REVIEWED PUBLICATIONS

Full publication list at: http://scholar.google.com/citations?user=O4rYanMAAAAJ

▷ Exploring neighborhoods in large metagenome assembly graphs reveals hidden sequence diversity.
CT Brown, D Moritz, M O'Brien, F Reidl, T Reiter, B Sullivan. ACCEPTED, Genome Biology, May 2020. (Available on biorxiv.)

A phylogenetically novel cyanobacterium most closely related to Gloeobacter. CL Grettenberger et al. The ISME Journal, 1-11

Whole genome sequencing for mutation discovery in a single case of lysosomal storage disease (MPS type 1) in the dog. TA Mansour et al. Scientific Reports 10 (1), 1-10

Genomic and phenotypic analyses of six offspring of a genome-edited hornless bull. AE Young et al. Nature biotechnology 38 (2), 225-232

FAIRshake: Toolkit to Evaluate the FAIRness of Research Digital Resources. DJB Clarke et al. Cell systems 9 (5), 417-421

Sequencing data discovery with MetaSeek. A Hoarfrost, N Brown, CT Brown, C Arnosti. Bioinformatics 35 (22), 4857-4859

Kevlar: a mapping-free framework for accurate discovery of de novo variants. DS Standage, CT Brown, F Hormozdiari. iScience 18, 28-36

Reproducible, interactive, scalable and extensible microbiome data science using QIIME 2. E Bolyen et al. Nature biotechnology 37 (8), 852-857

*Large-scale sequence comparisons with sourmash.* NT Pierce, L Irber, T Reiter, P Brooks, **CT Brown**. F1000Research 8, 1006.

Re-assembly, quality evaluation, and annotation of 678 microbial eukaryotic reference transcriptomes. LK Johnson, H Alexander, CT Brown. GigaScience 8 (4), giy158.

Review, evaluation and directions for gene-targeted assembly for ecologic analyses of metagenomes. J Guo, J Quensen, Y Sun, Q Wang, **CT Brown**, JR Cole, JM Tiedje. Frontiers in Genetics 10, 957.

VA-Store: A Virtual Approximate Store Approach to Supporting Repetitive Big Data in Genome Sequence Analyses. X Liu, Q Zhu, S Pramanik, CT Brown, G Qian. IEEE Transactions on Knowledge and Data Engineering

Whole genome variant association across 100 dogs identifies a frame shift mutation in DISHEV-ELLED 2 which contributes to Robinow-like syndrome in Bulldogs and related screw tail dog breeds. TA Mansour et al. PLoS genetics 14 (12), e1007850.

A Missense Mutation in the Vacuolar Protein Sorting 11 (VPS11) Gene Is Associated with Neuroaxonal Dystrophy in Rottweiler Dogs. KL Lucot et al. G3: Genes, Genomes, Genetics 8 (8), 2773-2780.

Sustainable computational science: the ReScience initiative. NP Rougier et al, PeerJ Computer Science 3, e142.

Draft genome of tule elk Cervus canadensis nannodes. JE Mizzi, ZT Lounsberry, CT Brown, BN Sacks. F1000Research 6.

Identification of long non-coding RNA in the horse transcriptome EY Scott et al., CJ Finno. BMC genomics 18 (1), 511.

Central nervous system transcriptome of Biomphalaria alexandrina, an intermediate host for schistosomiasis TA Mansour et al., MW Miller. BMC research notes 10 (1), 729.

FGF4 retrogene on CFA12 is responsible for chondrodystrophy and intervertebral disc disease in dogs. EA Brown et al., DL Bannasch. PNAS 2017/10/11.

Evolutionary loss of melanogenesis in the tunicate Molgula occulta. C Racioppi et al., F. Ristoratore. EvoDevo 2017.

sourmash: a library for MinHash sketching of DNA. CT Brown and LC Irber. Journal of Open Source Software, 1(5), 27, doi:10.21105/joss.00027.

Centromere location in Arabidopsis is unaltered by extreme divergence in CENH3 protein sequence. S Maheshwari, T Ishii, CT Brown, A Houben, L Comai. Genome Research 27 (3), 471-478. 2017.

khmer release v2.1: software for biological sequence analysis. Standage et al., CT Brown. Journal of Open Source Software, 2(15), 272, doi:10.21105/joss.00272

Tissue resolved, gene structure refined equine transcriptome. A Mansour, EY Scott, CJ Finno, RR Bellone, MJ Mienaltowski, MC Penedo, PJ Ross, SJ Valberg, JD Murray, CT Brown. BMC Genomics 18 (1) p103. 2017.

A new chicken genome assembly provides insight into avian genome structure. WC Warren et al. G3 7 (1) 109-117. 2017.

Transcriptome of the Caribbean stony coral Porites astreoides from three developmental stages. Mansour TC, Rosenthal JJC, Brown CT, Roberson LM. GigaScience 5 (1), 33, August 2016.

How open science helps researchers succeed. McKiernan EC et al., eLife 5, e16800, July 2016.

Haplotype-phased synthetic long reads from short-read sequencing. JA Stapleton, et al., PLOS One 11 (1), e0147229, 2016.

Microbial community analysis with ribosomal gene fragments from shotgun metagenomes. Guo J, Cole JC, Zhang Q, Brown CT, Tiedje JM. Applied and environmental microbiology. Jan 2016.

Haplotype-phased synthetic long reads from short-read sequencing. Stapleton JA, Kim J, Hamilton JP, Wu M, Irber LC, Maddamsetti R, Briney B, Newton L, Burton DR, Brown CT, Chan C, Buell CR, Whitehead TA. PLOS One. Jan 2016.

Hsp90 and hepatobiliary transformation during sea lamprey metamorphosis. Chung-Davidson YW, Yeh CY, Bussy U, Li K, Davidson PJ, Nanlohy KG, Brown CT, Whyard S, Li W. BMC Developmental Biology. Dec 2015.

A new method for DNA sequencing error verification and correction via an on-disk index tree. Gu Y, Liu X, Zhu Q, Dong Y, Brown CT, Pramanik S. Proceedings of the 6th ACM Conference on Bioinformatics, Computational Biology and Health Informatics. Sep 2015.

Differentially-Expressed Pseudogenes in HIV-1 Infection. Gupta A, Brown CT, Zheng YH, Adami C. Viruses. Sep 2015.

The khmer software package: enabling efficient nucleotide sequence analysis. Crusoe MR et al., Brown CT. F1000Research. Sep 2015.

Xander: employing a novel method for efficient gene-targeted metagenomic assembly. Wang Q, Fish JA, Gilman M, Sun Y, Brown CT, Tiedje JM, Cole JR. Microbiome. August 2015.

Transcriptome variation in response to Marek's disease virus acute infection. Preeyanon L, Brown CT, Cheng H. Cytogenetics. July 2015.

Phylogeny and phylogeography of functional genes shared among seven terrestrial subsurface metagenomes reveal N-cycling and microbial evolutionary relationships. Lau MCY et al., Onstott TC. Frontiers in Microbiology. Oct 2014.

Divergent mechanisms regulate conserved cardiopharyngeal development and gene expression in distantly related ascidians. Stolfi A, Lowe EK, Racioppi C, Ristoratore F, Brown CT, Swalla BJ, Christiaen L. eLife. Sep 2014.

These are not the k-mers you are looking for: efficient online k-mer counting using a probabilistic data structure. Zhang Q, Pell J, Canino-Koning R, Howe AC, **Brown CT**. PLoS One. July 2014.

▶ Tackling soil diversity with the assembly of large, complex metagenomes. Howe AC, Jansson J, Malfatti SA, Tringe SG, Tiedje JM, **Brown CT**. Accepted at PNAS, 2/2014. preprint arXiv:1212.2832.

The Ribosomal Database Project: Data and Tools for High Throughput rRNA Analysis. Cole JR, Wang Q, Fish J, Chai B, McGarrell D, Sun Y, Brown CT, Porras-Alfaro A, Kuske C, Tiedje JM. Accepted, Nucleic Acid Res, Nov 2013.

Genomic versatility and functional variation between two dominant heterotrophic symbionts of deep-sea Osedax worms. Goffredi S, Yi H, Zhang Q, Klann J, Struve I, Vrijenhoek RC, Brown CT. Accepted, ISME Journal, October 2013.

Best practices for scientific computing. Wilson GV et al. preprint arXiv:1210.0530. Accepted PLoS Biology, October 2013.

FunGene: the Functional Gene Pipeline and Repository. Fish JA, Chai B, Wang Q, Sun Y, **Brown** C, Tiedje JM and Cole JR (2013). Front. Microbiol. 4:291. doi: 10.3389/fmicb.2013.00291

The genome and developmental transcriptome of the strongylid nematode Haemonchus contortus. Schwarz EM, Korhonen PK, Campbell BE, Young ND, Jex AR, Jabbar A, Hall RS, Mondal A, Howe AC, Pell J, Hofmann A, Boag PR, Zhu XQ, Gregory TR, Loukas A, Williams BA, Antoshechkin I, **Brown CT**, Sternberg PW, Gasser RB. Genome Biol. 2013 Aug 28;14(8):R89.

The sea lamprey has a primordial accessory offactory system. Chang S, Chung-Davidson YW, Libants SV, Nanlohy KG, Kiupel M, **Brown C.T.**, Li W. BMC Evol Biol. 2013 Aug 17;13(1):172. doi: 10.1186/1471-2148-13-172.

Integrated Analyses of Genome-Wide DNA Occupancy and Expression Profiling Identify Key Genes and Pathways Involved in Cellular Transformation by a Marek's Disease Virus Oncoprotein, Meq. Subramaniam S, Johnston J, Preeyanon L, **Brown CT**, Kung HJ, Cheng HH. J Virol. 2013 Aug;87(16):9016-29. doi: 10.1128/JVI.01163-13.

A thermogenic secondary sexual character in male sea lamprey. Chung-Davidson, Y.-W., Priess, M.C., Yeh, C.-Y., Brant, C.O., Johnson, N.S., Li, K., Nanlohy, K.G., Bryan, M.B., **Brown, C.T.**, Choi, J., Li, W. Journal of Experimental Biology. 2013 Jul 15;216(Pt 14):2702-12. doi: 10.1242/jeb.085746.

Sequencing of the sea lamprey (Petromyzon marinus) genome provides insights into vertebrate evolution. Smith JJ et al. Nature Genetics, Apr;45(4):415-21, 421e1-2. doi: 10.1038/ng.2568. Epub 2013 Feb 24.

Draft Genome Sequences of two Campylobacter jejuni Clinical Isolates, NW and D2600. Jerome JP, Klahn BD, Bell JA, Barrick JE, **Brown CT**, Mansfield LS. Journal of Bacteriology, 194 (20), 5707-5708.

Standing Genetic Variation in Contingency Loci Drives the Rapid Adaptation of Campylobacter jejuni to a Novel Host Jerome JP, Bell JA, Plovanich-Jones AE, Barrick JE, **Brown CT**, Mansfield LS. PLoS One 6 (1), e16399, Jan 24 2011.

Exploring the future of bioinformatics data sharing and mining with Pygr and Worldbase Lee C, Alekseyenko A, **Brown CT**. in Proceedings of the 8th Python in Science conference (SciPy 2009), G Varoquaux, S van der Walt, J Millman (Eds.), pp. 62-67.

Diverse syntrophic partnerships from deep-sea methane vents revealed by direct cell capture and metagenomics.

Pernthaler A, Dekas AE, Brown CT, Goffredi SK, Embaye T, Orphan VJ.

Proc Natl Acad Sci U S A. 2008 May 13;105(19):7052-7. Epub 2008 May 8. PMID: 18467493.

The genome of the sea urchin Strongylocentrotus purpuratus.

Sea Urchin Genome Sequencing Consortium.

Science. 2006 Nov 10;314(5801):941-52. PMID: 17095691.

High regulatory gene use in sea urchin embryogenesis: Implications for bilaterian development and evolution.

Howard-Ashby M, Materna SC, **Brown CT**, Tu Q, Oliveri P, Cameron RA, Davidson EH. Dev Biol. 2006 Dec 1;300(1):27-34. Epub 2006 Oct 18. PMID: 17101125.

Gene families encoding transcription factors expressed in early development of Strongylocentrotus purpuratus. Howard-Ashby M, Materna SC, **Brown CT**, Chen L, Cameron RA, Davidson EH. Dev Bio 2006 300 (1), 90-107.

Identification and characterization of homeobox transcription factor genes in Strongylocentrotus purpuratus, and their expression in embryonic development Howard-Ashby M, Materna C, **Brown CT**, Chen L, Cameron RA, Davidson EH. 2006 Dev Bio 300 (1), 74-89.

Sea urchin Forkhead gene family: phylogeny and embryonic expression Tu Q, **Brown CT**, Davidson EH, Oliveri P. 2006 Dev Bio 300 (1), 49-62.

Paircomp, FamilyRelationsII and Cartwheel: tools for interspecific sequence comparison. Brown CT, Xie Y, Davidson EH, Cameron RA.

BMC Bioinformatics. 2005 Mar 24;6:70. PMID: 15790396.

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Anaerobic regulation by an atypical Arc system in Shewanella oneidensis.

Gralnick JA, Brown CT, Newman DK.

Mol Microbiol. 2005 Jun;56(5):1347-57. PMID: 15882425.

▶ Evolutionary comparisons suggest many novel cAMP response protein binding sites in E. coli. Brown CT, Callan CG Jr.

Proc Natl Acad Sci U S A. 2004 Feb 24;101(8):2404-9. PMID: 14983022.

Patchy interspecific sequence similarities efficiently identify positive cis-regulatory elements in the sea urchin. Yuh CH, **Brown CT**, Livi CB, Rowen L, Clarke PJC, Davidson EH. 2002 Dev Bio 246 (1), 148-161.

New computational approaches for analysis of cis-regulatory networks.

**Brown CT**, Rust AG, Clarke PJ, Pan Z, Schilstra MJ, De Buysscher T, Griffin G, Wold BJ, Cameron RA, Davidson EH, Bolouri H.

Dev Biol. 2002 Jun 1;246(1):86-102. PMID: 12027436.

A genomic regulatory network for development.

Davidson EH, Rast JP, Oliveri P, Ransick A, Calestani C, Yuh CH, Minokawa T, Amore G, Hinman V, Arenas-Mena C, Otim O, **Brown CT**, Livi CB, Lee PY, Revilla R, Rust AG, Pan Z, Schilstra MJ, Clarke PJ, Arnone MI, Rowen L, Cameron RA, McClay DR, Hood L, Bolouri H. Science. 2002 Mar 1;295(5560):1669-78. PMID: 11872831.

A provisional regulatory gene network for specification of endomesoderm in the sea urchin embryo. Davidson EH, Rast JP, Oliveri P, Ransick A, Calestani C, Yuh CH, Minokawa T, Amore G, Hinman V, Arenas-Mena C, Otim O, Brown CT, Livi CB, Lee PY, Revilla R, Schilstra MJ, Clarke PJ, Rust AG, Pan Z, Arnone MI, Rowen L, Cameron RA, McClay DR, Hood L, Bolouri H. Dev Biol. 2002 Jun 1;246(1):162-90. PMID: 12027441.

The Earthshine Project: update on photometric and spectroscopic measurements. E. Palle, P. M. Rodriguez, P. R. Goode, J. Qiu, V. Yurchyshyn, J. Hickey, M.-C. Chu, E. Kolbe, C. T. Brown, and S. E. Koonin. Solar Variability and Climate Change Advances in Space Research 34, 288 (2004).

The earthshine spectrum P. M. Rodriguez, E. Palle, P. R. Goode, J. Hickey, J. Qiu, V. Yurchyshyn, M.-C. Chu, E. Kolbe, C. T. Brown, and S. E. Koonin. Solar Variability and Climate Change Advances in Space Research 34, 293 (2004).

Sunshine, Earthshine and Climate Change: II. Solar Origins of Variations in the Earth's Albedo. P.R. Goode, E. Palli, V. Yurchyshyn, J. Qiu, J. Hickey, P. Montaqis-Rodriguez, M.-C. Chu, E. Kolbe, C.T. Brown, S.E. Koonin. Journal of the Korean Astronomical Society, 35, 1 (2003).

Earthshine and the Earth's albedo: 1. Earthshine observations and measurements of the lunar phase function for accurate measurements of the Earth's Bond albedo J. Qiu, P. R. Goode, E. Palle, V. Yurchyshyn, J. Hickey, P. M. Rodriguez, M.-C. Chu, E. Kolbe, C. T. Brown, and S. E. Koonin. J. of Geophys. Res.-Atmospheres 108, 4709 (2003).

Earthshine and the Earth's albedo: 2. Observations and simulations over three years E. Palle, P. R. Goode, V. Yurchyshyn, J. Qiu, J. Hickey, P. M. Rodriguez, M.-C. Chu, E. Kolbe, C. T. Brown, and S. E. Koonin. J. of Geophys. Res.-Atmospheres 108, 4710 (2003).

Earthshine observations of the earth's reflectance P. R. Goode, J. Qiu, V. Yurchyshyn, J. Hickey, M.-C. Chu, E. Kolbe, C. T. Brown, and S. E. Koonin Geophys. Res. Lett. 28, 1671 (2001).

Visualizing Evolutionary Activity of Genotypes in Evita; with M. Bedau. Adaptive Systems, 1998.

A Comparison of Evolutionary Activity in Artificial Living Systems and in the Biosphere; Snyder E, Brown CT, Bedau M, Packard N. in the Proceedings of the 4th Europ. Conf. on Artificial Life, July, 1997.

Abundance Distributions in Artificial Life and Stochastic Models: "Age and Area" revisited, Adami, C., Brown, C.T., Haggerty, M.R. Proc. of 3rd Europ. Conf. on Artificial Life, June 4-6, 1995, Granada, Spain, Lecture Notes in Computer Science, Springer Verlag (1995), p.503.

Evolutionary Learning in the 2D Artificial Life System "Avida" Adami C, Brown CT. Proc. of "Artificial Life IV", MIT Press, p. 377-381 (1994).

## Invited Articles and Reviews

Strain recovery from metagenomes. Brown CT. Nature Biotechnology 33 (10), 1041-1043, Oct 2015.

Metagenomics: the paths forward. Brown CT and Tiedje JM. Handbook of Molecular Microbiology II: Metagenomics in Different Habitats. Wiley-Blackwell 10 Nov 2011.

Computational approaches to finding and analyzing cis-regulatory elements. Brown CT. Methods Cell Biol. 2008;87:337-65. Review. PMID: 18485306.

Reproducible Bioinformatics Research for Biologists. Preeyanon L, Pyrkosz AB, and Brown CT. Chapter in Implementing Reproducible Computational Research, V. Stodden and R. Peng, ed. Forthcoming in Dec 2013.

## REPORTS AND EDITORIALS

The Architecture of the LISA Science Analysis. SA Teukolsky et al. Keck Institute for Space Studies

The principles of tomorrow's university. DS Katz et al. F1000Research 7

Keeping it light:(Re) analyzing community-wide datasets without major infrastructure. H Alexander, LK Johnson, CT Brown. GigaScience.

Cephalopod genomics: A plan of strategies and organization. Albertin et al., Standards in Genomic Sciences 7 (1), 175.

Changing computational research. The challenges ahead. Neylon, C., Aerts, J., Brown, C.T., Lemire, D., Millman, J., Murray-Rust, P., Perez, F., Saunders, N., Smith, A., Varoquaux, G. Source Code for Biology and Medicine 7 (1), 2 (2012).

Meeting report: the terabase metagenomics workshop and the vision of an Earth microbiome project. Gilbert, J.A., Meyer, F., Antonopoulos, D., Balaji, P., Brown, C.T., Brown, C.T., Desai, N., Eisen, J.A., Evers, D., Field, D. Standards in genomic sciences, 3(3) 243 (2010).

ONLINE COMMENTARIES, BLOGS, AND SOCIAL MEDIA Personal/professional blog at: ivory.idyll.org/blog/. A few selected posts (click on links): "Our approach to replication in computational science", "Thoughts on Assemblathon 2", "The future of khmer (2013)".

Twitter: @ctitusbrown

BioMedCentral invited blog post: "Version control for scientific research"

SELECTED INVITATIONS AND MEETINGS April 2013 - NIH NHGRI Education and Training committee.

March 2013 - Invited speaker at National Center for Atmospheric Research Software Engineering Assembly.

March 2013 - NSF/Moore Foundation meeting on Cyberinfrastructure for Marine 'Omics.

September 2012 - Invited speaker at Extremely Large Databases 2012 (XLDB 2012).

June 2012 - NSF BIO Centers meeting on Cyberinfrastructure Needs in BIO.

Professional Activities iPlant Scientific Advisory Board member.

Software Carpentry Scientific Advisory Board member.

Accredited Software Carpentry Instructor.

NIH Committee Member for Cloud Computing and the Human Microbiome.

Cephalopod Genome Sequencing Consortium Steering Committee, 2012-present.

Member of the Editorial Board for Open Research Computation, Frontiers in Livestock Genomics.

Xconomist.com, invited member, Advisory Board (Michigan chapter).

BEACON NSF STC, Thrust Group co-leader (responsible for reviewing proposals, organizing activities), 2010-2014.

Course director, 2010-2015, Next-Generation Sequence Analysis for Biologists, KBS, MSU.

Director, Data Intensive Biology Summer Institute 2017, UC Davis.

Course faculty, 2006-2008, Embryology Course, Woods Hole Marine Biological Laboratory.

Founder, Caltech Bioinformatics Journal Club; biology-in-python mailing list.

Faculty advisor, Metagenomics Journal Club at MSU.

Development and maintenance of several open source bioinformatics tools, including Cartwheel server for comparative sequence analysis, khmer k-mer software, and screed; github.com/ctb/.

Active in open source testing community: twill, figleaf, pony-build.

Reviewer for National Science Foundation; Developmental Biology, BMC Bioinformatics, BMC Genomics, Genome Biology, Bioinformatics, PLoS One.

TEACHING AND WORKSHOPS Open Problems in Bioinformatics, CSE/MMG graduate seminar course (2008-2009)

Database-Backed Web Development, CSE 4xx (2008-, yearly), at MSU.

Introduction to Computational Science for Evolutionary Biologists, CSE 801 (was 891) (2010-2014), at MSU.

Analyzing Next-Generation Sequencing Data, research workshop (2010-, yearly).

Software Carpentry workshops: Scripps Research Institute (11/2012), U. Arizona (4/2013)

Instructor at Marine Biological Laboratory course on Strategies and Techniques for Analyzing Microbial Population Structures, 2012 and 2013.

Co-instructor for Workshop on Microbial Bioinformatics, 10/2013, Caltech.

Lead instructor for Workshop on mRNAseq for Biologists, and Workshop for Advanced Bioinfor-

matics Developers, 11/2013, The Centre For Genome Analysis, Norwich, UK. Intro to Genetics and Genomics lab, GGG 201(b), UC Davis, 2017-present.

Tools to Support Data Intensive Research, GGG 298, UC Davis, 2020.

References

(Contact details available upon request.)

Dr. Ewan Birney, Associate Director of the EMBL-EBI.

Professor Jonathan Eisen, University of California Davis.

Professor Paul W. Sternberg, California Institute of Technology.

Professor Billie J. Swalla, University of Washington at Seattle.