The people of the 24th annual Pacific Symposium on Biocomputing in Hawaii

This manuscript (permalink) was automatically generated from dhimmel/psb-manuscript@c3fd119 on January 5, 2019.

Authors

Weixuan Fu

© 0000-0002-6434-5468 • • weixuanfu • weixuanfu

Department of Biostatistics, Epidemiology and Informatics, Institute for Biomedical Informatics, University of Pennsylvania

Casey Greene

Department of Systems Pharmacology & Translational Therapeutics, University of Pennsylvania; Childhood Cancer Data Lab, Alex's Lemonade Stand Foundation

• Daniel Himmelstein

□ 0000-0002-3012-7446 **□** dhimmel **■** dhimmel

Department of Systems Pharmacology & Translational Therapeutics, University of Pennsylvania

Lawrence Hunter

University of Colorado Anschutz School of Medicine; University of Colorado Boulder

• Trang Le

D 0000-0003-3737-6565 **○** trang1618 **У** trang1618

University of Pennsylvania

Luca Pinello

● 0000-0003-1109-3823 • pinellolab • lucapinello

Massachusetts General Hospital/ Harvard Medical School

· Jaclyn Taroni

Childhood Cancer Data Lab, Alex's Lemonade Stand Foundation

• Ryan Whaley

Stanford University

Abstract

Manubot is an open source tool for writing manuscripts on GitHub in markdown format. Manubot applies the git-based software workflow to scholarly writing, enabling enhanced transparency, collaboration, automation, and reproducibility.

This manuscript is the result of a *special working group* at the 2019 Pacific Symposium on Biocomputing that will introduce attendees to collaborative writing with Manubot. Each conference attendee is invited to write a small blurb on themselves and their research, by submitting a pull request to the manuscript repository at https://github.com/dhimmel/psb-manuscript.

The working group also covers how to write your next manuscript using Manubot and what features of Manubot can help biomedical researchers document and publish their computational research. For example, Manubot enables citation by persistent identifier to automate bibliographic metadata retrieval and formatting as well as allowing templating so results can be directly inserted from the analyses that produced them.

Methods

In this section, PSB 2019 attendees are asked to contribute a bit about themselves and their research. As part of the special working group, we thought this would be a helpful activity to introduce biocomputational scientists to writing with Manubot. For inspiration, here are some prompts:

- Introduce yourself briefly.
- What do you research? Include any relevant links to your lab or personal website.
- What is your favorite study from your career or what study was your biggest discovery?
- What was your first scholarly publication?
- Add a code snippet or mathematical equation.
- Add a figure with a caption. This could be a picture of you in Hawaii or a figure from your previous work if the license is permissive enough to allow reuse.

Self-citations are explicitly encouraged, since one goal of this activity is to introduce attendees to the concept of citation by persistent identifier. By having attendees cite their existing works, we hope to show researchers how references can be created from just persistent identifiers, and how this can assist with collaborative and transparent authoring.

The markdown manuscript source has a section for each PSB 2019 attendee, generated from the online attendee list. Names are ordered alphabetically by last name. If you'd like to contribute, but are not already in the list, please insert your section at the appropriate alphabetical location.

For questions on how to use Manubot, see the usage documentation. More information on the tool and its inception is available in the project manuscript [1].

Attendees

Weixuan Fu

Aloha, I'm in the Institute for Biomedical Informatics (IBI) at the University of Pennsylvania and the developer of TPOT and PennAI [2].

My main interest of research is developing automated machine learning tools for the analysis of large scale biomedical/sequencing data. Besides that, I am working on optimizing analysis pipeline of predicting neoantigen specifically presented in tumor cells using DNA and RNA sequencing data, for designing personalized neoantigen vaccines in cancer immunotherapies.

Casey Greene

I run an integrative genomics research lab at the University of Pennsylvania, and I direct the Childhood Cancer Data Lab for Alex's Lemonade Stand Foundation. The lab at Penn develops methods to integrate large-scale public datasets, primarily from transcriptomic assays, and applies these methods to a broad set of biological questions. We've studied numerous systems, and we currently have active research projects in the application areas of microbial systems [3,4], cancers [5,6,7,8], and rare diseases [9]. At this PSB, a postdoc from the group will present a paper describing Continental Breakfast Included (CBI) effect in the final talk of the final session of this year's meeting [10].

I'm also interested in technologies that enhance the future of scientific communication. Our lab has studied Sci-Hub [11]. We've led a large collaborative review of deep learning in biology and medicine [12]. Members of the lab have developed tools like manubot [1], which you are using now. More publications are available on our lab website.

Daniel Himmelstein

Greetings, I'm in the Greene Lab at the University of Pennsylvania and am the lead developer of the Manubot project. 2019 is my first PSB and I'm exciting to backpack around the Big Island following the conference.

My main area of research is integrating biomedical knowledge using hetnets [13,14]. However, I've also studied Sci-Hub, finding that it provides access to nearly all paywalled scholarly literature [15]. Perhaps my biggest discovery was observing an epidemiological association that higher elevation counties have lower rates of lung cancer, suggesting that oxygen is an inhaled carcinogen (Figure 1) [16,17].

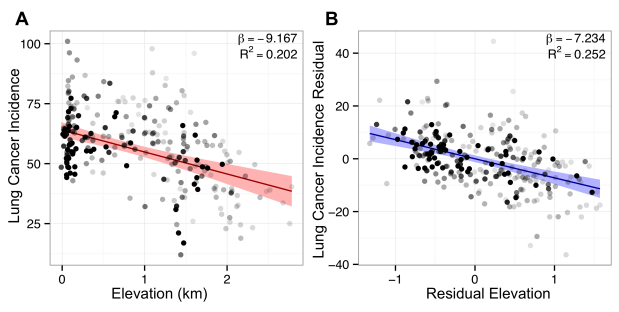


Figure 1: The association between elevation and lung cancer across Western U.S. counties. This figure is reused from here under its CC BY 4.0 License.

I haven't done much text mining, but I did enjoy extracting attendee names for PSB from the online PDF. Converting the PDF to text in Python was as easy as:

```
# https://stackoverflow.com/a/48673754
import tika.parser
parsed = tika.parser.from_file('attendees.pdf')
text = parsed["content"]
```

Lawrence Hunter

I'm a cofounder of the PSB conference, and a professor at the University of Colorado School of Medicine. You can find information about my lab at http://compbio.ucdenver.edu/Hunter. One of my early papers is 18.

Trang Le

Hello from the Moore lab at the University of Pennsylvania! I'm a mathematician who's currently excited about automated machine learning.

Here goes the self-citations:

- My own favorite study: Generalization of the Fermi Pseudopotential [19] a piece of mathematical physics work I got to do when procrastinating writing my dissertation.
- My first (first-author) scholarly publication: Differential privacy-based evaporative cooling feature selection and classification with relief-F and random forests [20]. Check out the Github repo for this study here.

• Code snippet I'm most proud of:

```
M = dec2bin(0:2^{n*n}-1,n*n)
```

I will be impressed if you could tell what the language is. This is my answer to a question on Math StackExchange.

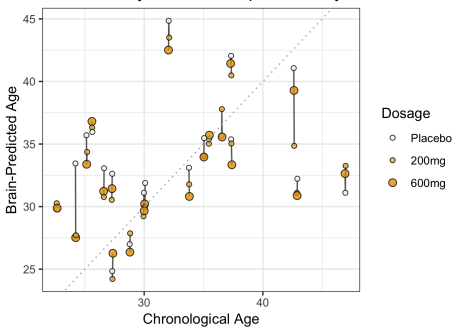
• I have too many favorite mathematical equations, but here's one:

$$a^p \equiv a \mod p$$

Anyone recognize this theorem?

• And a figure with a caption:

Plot of brain-predicted age vs. chronological age of subjects in the ibuprofen study



This is an improved version of my main figure in this interesting study [21].

Luca Pinello

Aloah from the Pinello Lab!

I am a computational biologist studying the role of chromatin structure/dynamics and non-coding regions including enhancers, promoters, insulators and their role in gene regulation. The mission of my lab is the integration of omics data to explore and better understand the functional mechanisms of the non-coding genome and to provide accessible tools for the community to accelerate discovery in this field. The long-term goal of my research is to develop innovative computational approaches and to use cutting-edge experimental assays, such as single cell and genome editing, to systematically analyze sources of genetic and epigenetic variation that affect gene regulation in different human traits and diseases. I believe this will further our understanding of disease etiology

involving these poorly characterized regions and will provide a foundation for the development of new drugs and more targeted treatments.

I am excited to share during the workshop Reading between the genes: Interpreting noncoding DNA in high throughput a new computational methods we have recently developed to analyze CRISPR tiling screen called CRISPR-SURF. You can read more on the manuscript that was recently published in Nature Methods [22].

Jaclyn Taroni

I'm a data scientist at the Childhood Cancer Data Lab (CCDL), an initiative of Alex's Lemonade Stand Foundation. I'm interested in how diverse collections of publicly available transcriptomic data can help us learn about the biology of rare diseases. As a graduate student, I studied systemic sclerosis [23]. In the PSB 2019 Text Mining and Machine Learning for Precision Medicine Workshop, I'll present our MultiPLIER project [9]. With the CCDL, I've been working on refine.bio, a project for uniformly processing transcriptomic data from multiple species.

Ryan Whaley

Hi, I'm Ryan and I'm one of the technical leads for PharmGKB. I'm also helping to run the A/V desk during this presentation.

I'm trained in software development and started by career as a DBA. Over the past decade I've switched to Java and then web application development. I've contributed to PharmGKB [24], CPIC [25], and other PGx consortia.

References

1. Open collaborative writing with Manubot

Daniel S. Himmelstein, David R. Slochower, Venkat S. Malladi, Casey S. Greene, Anthony Gitter *Manubot Preprint* (2018-12-31) https://greenelab.github.io/meta-review/

2. A System for Accessible Artificial Intelligence

Randal S. Olson, Moshe Sipper, William La Cava, Sharon Tartarone, Steven Vitale, Weixuan Fu, Patryk Orzechowski, Ryan J. Urbanowicz, John H. Holmes, Jason H. Moore Genetic Programming Theory and Practice XV (2018) https://doi.org/gfsptm

DOI: 10.1007/978-3-319-90512-9_8

3. ADAGE-Based Integration of Publicly Available Pseudomonas aeruginosa Gene Expression Data with Denoising Autoencoders Illuminates Microbe-Host Interactions

Jie Tan, John H. Hammond, Deborah A. Hogan, Casey S. Greene *mSystems* (2016-01-19) https://doi.org/gcgmbq

DOI: 10.1128/msystems.00025-15 · PMID: 27822512 · PMCID: PMC5069748

4. Unsupervised Extraction of Stable Expression Signatures from Public Compendia with an Ensemble of Neural Networks

Jie Tan, Georgia Doing, Kimberley A. Lewis, Courtney E. Price, Kathleen M. Chen, Kyle C. Cady, Barret Perchuk, Michael T. Laub, Deborah A. Hogan, Casey S. Greene *Cell Systems* (2017-07) https://doi.org/gcw9f4

DOI: 10.1016/j.cels.2017.06.003 PMID: 28711280 PMCID: PMC5532071

5. Extracting a biologically relevant latent space from cancer transcriptomes with variational autoencoders

Gregory P. Way, Casey S. Greene

Biocomputing 2018 (2017-11-17) https://doi.org/gfspsd

DOI: 10.1142/9789813235533 0008

6. Machine Learning Detects Pan-cancer Ras Pathway Activation in The Cancer Genome Atlas

Gregory P. Way, Francisco Sanchez-Vega, Konnor La, Joshua Armenia, Walid K. Chatila, Augustin Luna, Chris Sander, Andrew D. Cherniack, Marco Mina, Giovanni Ciriello, ... Armaz Mariamidze *Cell Reports* (2018-04) https://doi.org/gfspsb

DOI: 10.1016/j.celrep.2018.03.046 PMID: 29617658 PMCID: PMC5918694

7. Genomic and Molecular Landscape of DNA Damage Repair Deficiency across The Cancer Genome Atlas

Theo A. Knijnenburg, Linghua Wang, Michael T. Zimmermann, Nyasha Chambwe, Galen F. Gao,

Andrew D. Cherniack, Huihui Fan, Hui Shen, Gregory P. Way, Casey S. Greene, ... Armaz Mariamidze

Cell Reports (2018-04) https://doi.org/gfspsc

DOI: 10.1016/j.celrep.2018.03.076 · PMID: 29617664 · PMCID: PMC5961503

8. Oncogenic Signaling Pathways in The Cancer Genome Atlas

Francisco Sanchez-Vega, Marco Mina, Joshua Armenia, Walid K. Chatila, Augustin Luna, Konnor C. La, Sofia Dimitriadoy, David L. Liu, Havish S. Kantheti, Sadegh Saghafinia, ... Armaz Mariamidze

Cell (2018-04) https://doi.org/gc7r9b

DOI: 10.1016/j.cell.2018.03.035 · PMID: 29625050 · PMCID: PMC6070353

9. MultiPLIER: a transfer learning framework reveals systemic features of rare autoimmune disease

Jaclyn N Taroni, Peter C Grayson, Qiwen Hu, Sean Eddy, Matthias Kretzler, Peter A Merkel, Casey S Greene

Cold Spring Harbor Laboratory (2018-08-20) https://doi.org/gfc9bb

DOI: 10.1101/395947

10. Parameter tuning is a key part of dimensionality reduction via deep variational autoencoders for single cell RNA transcriptomics

Qiwen Hu, Casey S Greene

Cold Spring Harbor Laboratory (2018-08-05) https://doi.org/gdxxjf

DOI: 10.1101/385534

11. Sci-Hub provides access to nearly all scholarly literature

Daniel S Himmelstein, Ariel Rodriguez Romero, Jacob G Levernier, Thomas Anthony Munro, Stephen Reid McLaughlin, Bastian Greshake Tzovaras, Casey S Greene *eLife* (2018-03-01) https://doi.org/ckcj

DOI: 10.7554/elife.32822 · PMID: 29424689 · PMCID: PMC5832410

12. Opportunities and obstacles for deep learning in biology and medicine

Travers Ching, Daniel S. Himmelstein, Brett K. Beaulieu-Jones, Alexandr A. Kalinin, Brian T. Do, Gregory P. Way, Enrico Ferrero, Paul-Michael Agapow, Michael Zietz, Michael M. Hoffman, ... Casey S. Greene

Journal of The Royal Society Interface (2018-04) https://doi.org/gddkhn

DOI: 10.1098/rsif.2017.0387 · PMID: 29618526 · PMCID: PMC5938574

13. Heterogeneous Network Edge Prediction: A Data Integration Approach to Prioritize Disease-Associated Genes.

Daniel S Himmelstein, Sergio E Baranzini

PLoS computational biology (2015-07-09) https://www.ncbi.nlm.nih.gov/pubmed/26158728

DOI: 10.1371/journal.pcbi.1004259 · PMID: 26158728 · PMCID: PMC4497619

14. Systematic integration of biomedical knowledge prioritizes drugs for repurposing

Daniel Scott Himmelstein, Antoine Lizee, Christine Hessler, Leo Brueggeman, Sabrina L Chen, Dexter Hadley, Ari Green, Pouya Khankhanian, Sergio E Baranzini

eLife (2017-09-22) https://doi.org/cdfk

DOI: 10.7554/elife.26726 · PMID: 28936969 · PMCID: PMC5640425

15. Sci-Hub provides access to nearly all scholarly literature

Daniel S Himmelstein, Ariel Rodriguez Romero, Jacob G Levernier, Thomas Anthony Munro, Stephen Reid McLaughlin, Bastian Greshake Tzovaras, Casey S Greene *eLife* (2018-03-01) https://www.ncbi.nlm.nih.gov/pmc/articles/PMC5832410/

DOI: 10.7554/elife.32822 · PMID: 29424689 · PMCID: PMC5832410

16. Lung cancer incidence decreases with elevation: evidence for oxygen as an inhaled carcinogen

Kamen P. Simeonov, Daniel S. Himmelstein

PeerJ (2015-01-13) https://doi.org/98p

DOI: 10.7717/peerj.705 · PMID: 25648772 · PMCID: PMC4304851

17. Unraveling the Ties of Altitude, Oxygen and Lung Cancer

George Johnson

The New York Times (2016-01-25) https://www.nytimes.com/2016/01/26/science/unraveling-the-ties-of-altitude-oxygen-and-lung-cancer.html

18. EDGAR: Extraction of Drugs, Genes And Relations from the Biomedical Literature

Thomas C. Rindflesch, Lorraine Tanabe, John N. Weinstein, Lawrence Hunter *Biocomputing 2000* (1999-12) https://doi.org/gfsptq

DOI: 10.1142/9789814447331 0049

19. Generalization of the Fermi Pseudopotential

Trang T. Le, Zach Osman, D. K. Watson, Martin Dunn, B. A. McKinney *arXiv* (2018-06-14) https://arxiv.org/abs/1806.05726v1

20. Differential privacy-based evaporative cooling feature selection and classification with relief-F and random forests

Trang T Le, W Kyle Simmons, Masaya Misaki, Jerzy Bodurka, Bill C White, Jonathan Savitz, Brett A McKinney

Bioinformatics (2017-05-04) https://doi.org/f96b8d

DOI: 10.1093/bioinformatics/btx298 · PMID: 28472232 · PMCID: PMC5870708

21. Effect of Ibuprofen on BrainAGE: A Randomized, Placebo-Controlled, Dose-Response Exploratory Study

Trang T. Le, Rayus Kuplicki, Hung-Wen Yeh, Robin L. Aupperle, Sahib S. Khalsa, W. Kyle

Simmons, Martin P. Paulus

Biological Psychiatry: Cognitive Neuroscience and Neuroimaging (2018-10) https://doi.org/gfsprv DOI: 10.1016/j.bpsc.2018.05.002 · PMID: 29941380

22. CRISPR-SURF: discovering regulatory elements by deconvolution of CRISPR tiling screen data.

Jonathan Y Hsu, Charles P Fulco, Mitchel A Cole, Matthew C Canver, Danilo Pellin, Falak Sher, Rick Farouni, Kendell Clement, Jimmy A Guo, Luca Biasco, ... Luca Pinello Nature methods (2018-12) https://www.ncbi.nlm.nih.gov/pubmed/30504875

DOI: 10.1038/s41592-018-0225-6 · PMID: 30504875

23. A novel multi-network approach reveals tissue-specific cellular modulators of fibrosis in systemic sclerosis

Jaclyn N. Taroni, Casey S. Greene, Viktor Martyanov, Tammara A. Wood, Romy B. Christmann, Harrison W. Farber, Robert A. Lafyatis, Christopher P. Denton, Monique E. Hinchcliff, Patricia A. Pioli, ... Michael L. Whitfield

Genome Medicine (2017-03-23) https://doi.org/gfsptx

DOI: 10.1186/s13073-017-0417-1 · PMID: 28330499 · PMCID: PMC5363043

24. Pharmacogenomics knowledge for personalized medicine.

M Whirl-Carrillo, EM McDonagh, JM Hebert, L Gong, K Sangkuhl, CF Thorn, RB Altman, TE Klein *Clinical pharmacology and therapeutics* (2012-10) https://www.ncbi.nlm.nih.gov/pubmed/22992668 DOI: 10.1038/clpt.2012.96 · PMID: 22992668 · PMCID: PMC3660037

25. CPIC: Clinical Pharmacogenetics Implementation Consortium of the Pharmacogenomics Research Network.

MV Relling, TE Klein

Clinical pharmacology and therapeutics (2011-01-26) https://www.ncbi.nlm.nih.gov/pubmed/21270786

DOI: 10.1038/clpt.2010.279 · PMID: 21270786 · PMCID: PMC3098762