Curriculum Vitae Gabriel Etan Leventhal

Forch 2 Phone: +41(78)708 49 08
8574 Oberhofen Email: gabriel@leventhal.ch
Switzerland URL: http://www.leventhal.ch

Language Skills: English (native), Swiss German (native), German (fluent), French (fluent)

Current Position

Mar 2020 - present VP Data Science

Pharmabiome AG

Areas of Specialization

Microbial Ecosystems; Evolutionary Theory; Interaction Networks; Community Evolution; Evolutionary Ecology; Computational Biology; Phylogenetics; Statistical Inference; Ecological Networks; Epidemiology; HIV Dynamics

Academic Positions and Education

Nov 2015 – Feb 2020 Postdoctoral Fellow, Massachusetts Institute of Technology (MIT), U.S.A.

Cordero Lab, Department of Civil and Environmental Engineering

Sep 2014 – Oct 2015 Postdoctoral Researcher, ETH Zürich, Switzerland

Theoretical Biology Group, Institute of Integrative Biology

Sep 2009 – Aug 2014 PhD Thesis, ETH Zürich, Switzerland

Title: Modeling the ecology and evolution of infectious diseases

Supervisor: Prof. Sebastian Bonhoeffer

Mar 2008 – Jun 2008 Research Assistant, EPFL, Lausanne, Switzerland

Laboratory of Statistical Biophysics, Department of Physics

Oct 2006 - Feb 2008 MSc in Physics, EPFL, Lausanne, Switzerland

Aug 2007 – Feb 2008 Master's Thesis, Indiana University Bloomington, U.S.A.

Title: Spectral Coarse Graining in Ising Spin Systems

Supervisors: Prof. Alessandro Flammini (IUB) and Prof. Paolo De Los Rios (EPFL)

May 2006 – Aug 2006 Research Assistant, Hong Kong University of Science and Technology

Kwok Yip Szeto Group, Department of Physics

Aug 2002 - May 2006 BSc in Physics, EPFL, Lausanne, Switzerland

Sep 2005 – May 2006 Undergraduate Exchange, Hong Kong University of Science and Technology

One-year international exchange program as part of BSc in physics

Publications

I have published 22 original research articles in peer-reviewed journals (10 as first, co-first, or equal contribution, as well as 2 review articles. Selected important publications are followed by a brief summary in red font.

† equal contribution; * student advisee

Microbial Ecology & Evolution

2020 LS Bittleston, M Gralka, <u>GE Leventhal</u>, I Mizrahi, OX Cordero. Context-dependent dynamics lead to the assembly of functionally distinct pitcher-plant microbiomes. Nature Communications 10.1038/s41467-020-15169-0

In this paper, I contributed to the analysis and interpretation of the compositional time series of ten distinct natural microbial that were propagated over weeks in controlled laboratory conditions. We show that while some aspects of the dynamics are conserved across starting communities, the intial composition matters for the dynamics in the same laboratory conditions. This paper sets the ground work for a follow-up experiment that I lead, in which we continued to propagate the communities during a whole year to measure the evolutinary dynamics over extended periods of time.

2019 <u>GE Leventhal</u>, M Ackermann, K Schiessl. Why microbes secrete molecules to modify their environment: the case of iron-chelating siderophores. Journal of the Royal Society Interface 16(150) 10.1098/rsif.2018.0674

This paper addresses the question as to why some microbes secrete compounds to aid in foraging for resources. Because such compounds become public goods, they have been extensively discussed from the point of view of cheating and competition. Here I quantified what the benefits to a microbe of secretion versus keeping such compounds for example membrane-bound when resources are aggregated in large clumps. My results have important consequences for understanding the length scale of competition and cooperation in communities.

2018 <u>GE Leventhal</u>, C Boix*, U Kuechler, T Enke, E Sliwerska, C Holliger, OX Cordero. Strain-level diversity drives alternative community types in millimetre-scale granular biofilms. Nature Microbiology 3:1295–1303 10.1038/s41564-018-0242-3

This paper is one of the first to investigate the fine-scale population genomic strucutre across replicate microbial communities. By leveraging the milimeter-scale granular biofilms from an activate granular sludge reactor, I showed that the dominant organism maintained clear and diverse genetic structure across replicate communities, and that this structure was correlated with the overall community composition. This is of large important to understanding the role of strain diversity and specificity in microbial community assembly.

- 2018 TN Enke, <u>GE Leventhal</u>, M Metzger, J Saavedra, OX Cordero. Microscale ecology regulates particulate organic matter turnover in model marine microbial communities. Nature Communications 10.1038/s41467-018-05159-8
- preprint <u>GE Leventhal</u>[†], L Wang[†], RD Kouyos. Real-world Interaction Networks Buffer Impact of Small Evolutionary Shifts On Biodiversity. bioRxiv 10.1101/013086

Microbiome & Health

- preprint E Katkeviciute, L Hering, A Montalban-Arques, P Busenhart, JC Aranda, K Atrott, S Lang, G Rogler, E Naschberger, VS Schellerer, M Stürzl, A Rickenbacher, M Turina, A Weber, S Leibl, <u>GE Leventhal</u>, M Levesque, O Boyman, M Scharl, MR Spalinger. Targeting protein tyrosine phosphatase non-receptor type 2 in immune cells converts immune-silent into highly immunogenic tumors. Submitted
 - 2019 MR Spalinger, M Schwarzfischer, L Hering, A Shawki, A Sayoc, A Santos, C Gottier, S Lang, K Bäbler, A Geirnaert, C Lacroix, **GE Leventhal**, X Dai, D Rawlings, AA Chan, G Rogler, DF McCole, M Scharl. Loss

of PTPN22 abrogates the beneficial effect of cohousing-mediated fecal microbiota transfer in murine colitis. **Mucosal Immunology** 10.1038/s41385-019-0201-1

Pathogen Evolution

- F Bertels, A Marzel, <u>GE Leventhal</u>, V Mitov, J Fellay, HF Günthard, J Böni, S Yerly, T Klimkait, V Aubert, M Battegay, A Rauch, M Cavassini, A Calmy, E Bernasconi, P Schmid, A Scherrer, V Müller, S Bonhoeffer, RD Kouyos, RR Regoes. *Dissecting HIV Virulence: Heritability Of Setpoint Viral Load, CD4+ T Cell Decline And Per-Parasite Pathogenicity.* Molecular Biology and Evolution 35(1):27-37 10.1093/molbev/msx246
- 2017 N Bachmann, T Turk, C Kadelka, A Marzel, M Shilaih, J Böni, V Aubert, T Klimkait, <u>GE Leventhal</u>, HF Günthard, RD Kouyos. Parent-offspring regression to estimate the heritability of an HIV-1 trait in a realistic setup. Retrovirology 14(33) 10.1186/s12977-017-0356-3
- 2016 <u>**GE Leventhal**</u>, S Bonhoeffer. *Potential pitfalls in estimating viral load heritability.* **Trends in Microbiology** 24(9):687-698 10.1016/j.tim.2016.04.008
- S Bonhoeffer, C Fraser, <u>GE Leventhal</u>. Heritability and the distribution of set point viral load in HIV carriers. PLoS Pathogens 11(2):e1004634 10.1371/journal.ppat.1004634
- 2014 C Fraser, K Lythgoe, <u>GE Leventhal</u>, G Shirreff, TD Hollingsworth, S Alizon, S Bonhoeffer. *Virulence and Pathogenesis of HIV-1 Infection: An Evolutionary Perspective*. Science 343(6177):1243727 10.1126/science.1243727
- 2014 <u>**GE Leventhal**</u>†, SR Dünner^{†,*}, S Barribeau. Delayed virulence and limited costs promote fecundity compensation upon infection. American Naturalist 103(4):480-493 10.1086/675242
- 2013 A Hool^{†,*}, **GE Leventhal**[†], S Bonhoeffer. Virus-induced target cell activation reconciles set-point viral load heritability and within-host evolution. **Epidemics** 7:35-35 10.1016/j.epidem.2013.09.002
- 2012 RD Kouyos, <u>GE Leventhal</u>, T Hinkley, M Haddad, J Whitcomb, C Petropoulos, S Bonhoeffer. *Exploring the Complexity of the HIV-1 Fitness Landscape*. **PLoS Genetics** 8(3):e100255110.1371/journal.pgen.1002551

Network Epidemiology

- 2017 JI Liechti, <u>GE Leventhal</u>, S Bonhoeffer. Host population structure impedes reversion to drug sensitivity after discontinuation of treatment. <u>PLoS Computational Biology</u> 13(8):e1005704 10.1371/journal.pcbi.1005704
- 2016 W Delva, **GE Leventhal**, S Helleringer. Connecting the dots: network data and models in HIV epidemiology. **AIDS** 30(13):2009-2020 10.1097/QAD.000000000001184
- GE Leventhal[†], AL Hill[†], M Nowak, S Bonhoeffer. Evolution and emergence of infectious diseases in theoretical and real-world networks. Nature Communications 6 10.1038/ncomms7101

 I extended the mathematical network framework commonly used in epidemiology to include an evolutionary process. Networks are often used to describe the precise way in which individuals in a host population encounter each other and transmit disease. I the field of network epidemiology, it is well known that the structure of these networks—for example the presence of super spreaders—strongly determines the likelihood and speed of an epidemic outbreak. How this contact structure influences the evolution of an infectious disease, however, had only been poorly understood. Here, I showed that the properties of host contact networks that increase the probability and speed of the spread of a single disease strain, also slow down the evolution of the disease. We have identified a first-come-first-serve effect, where host populations that are connected in such a way that facilitates the initial emergence of disease strain, also decrease the probability that the disease strain will be replaced by another one.
- 2012 <u>GE Leventhal</u>, RD Kouyos, T Stadler, VV Wyl, S Yerly, J Böni, C Cellerai, T Klimkait, HF Günthard, S Bonhoeffer. *Inferring Epidemic Contact Structure from Phylogenetic Trees.* PLoS Computational Biology 8(3):e1002413 10.1371/journal.pcbi.1002413
 - In this paper, I developed a mathematical approach to infer epidemic contact struture from phylogenetic trees. Contact structure is known to strongly influence the spread of an infectious diesease, but measuring contact structure is often difficult or impossible. Here, I develop

a statistical technique to infer population structure based on the shape of the phylogenetic tree reconstructed from genomic data.

Mathematical Modelling & Statistical Inference

- ²⁰¹⁹ TG Vaughan[†], <u>GE Leventhal</u>[†], DA Rasmussen, AJ Drummond, D Welch, T Stadler. *Estimating epidemic incidence and prevalence from genomic data*. <u>Molecular Biology and Evolution</u> 36(8):1804–1816 10.1093/molbev/msz106
- 2017 O Ratmann, EB Hodcroft, M Pickles, A Cori, M Hall, S Lycett, C Colijn, B Dearlove, X Didelot, S Frost, A Hossain, JB Joy, M Kendall, D Kühnert, <u>GE Leventhal</u>, R Liang, G Plazzotta, AF Poon, DA Rasmussen, T Stadler, E Volz, C Weis, AJ Leigh Brown, C Fraser. *Phylogenetic Tools for Generalized HIV-1 Epidemics: Findings from the PANGEA-HIV Methods Comparison*. <u>Molecular Biology and Evolution</u> 34(1):185–203 10.1093/molbev/msw217
- 2016 L du Plessis, **GE Leventhal**, S Bonhoeffer. How good are statistical models at approximating complex fitness landscapes?. **Molecular Biology and Evolution** 33(9):2454-2468 10.1093/molbev/msw097
- T Stadler, TG Vaughan, A Gavryushkin, S Guindon, D Kühnert, <u>GE Leventhal</u>, AJ Drummond. How well can the exponential-growth coalescent approximate constant-rate birth-death population dynamics?. Proceeding of the Royal Society B: Biological Sciences 282(1806):20150420 10.1098/rspb.2015.0420
- 2014 <u>GE Leventhal</u>, H Günthard, S Bonhoeffer, T Stadler. Using an epidemiological model for phylogenetic inference reveals density-dependence in HIV transmission. <u>Molecular Biology and Evolution</u> 31(1):6-17 10.1093/molbev/mst172

Software & Other Publications

- 2013-2016 **GE Leventhal**. R package expoTree to calculate the density dependent likelihood of a phylogenetic tree. Available on CRAN..
- 2012-2019 **GE Leventhal**, L Schulé, J Geering. iRiSS: a free online journal TOC aggregator that helps you stay informed about the latest work produced in your field.

Teaching and Mentoring

- 2016, 2018 Co-Instructor, Computational Ecology. MIT, Cambridge, MA, USA
 - 2018 Co-Supervisor, Exchange graduate student (Jacob Russel). MIT, Cambridge, MA, USA
 - 2016 Co-Supervisor, Rotation student (Carles Boix). MIT, Cambridge, MA, USA
- 2013 2015 Co-Lecturer, Infectious Disease Dynamics . ETH Zürich, Switzerland
 - 2015 Co-Supervisor, Master's thesis (Adriano Pagano). ETH Zürich, Switzerland
 - 2015 Supervisor, Semester student (Adriano Pagano). ETH Zürich, Switzerland
 - 2015 Co-Supervisor, Master's thesis (Martin Müller). ETH Zürich, Switzerland
 - 2012 Co-Supervisor, Master's thesis (Anna Hool). ETH Zürich, Switzerland
 - 2012 Co-Supervisor, Semester student (Robert Dünner). ETH Zürich, Switzerland
- 2011 2012 Co-Lecturer, English for nurses: an introduction to academic reading . Zurich University of Applied Sciences, Switzerland
- 2007 2008 Tutor for undergraduate calculus. Indiana University Bloomington, IN, U.S.A.
- 2005 2007 Teaching assistant in physics. EPFL, Lausanne, Switzerland
- 2003 2004 Teaching assistant in scientific programming, EPFL, Lausanne, Switzerland

Selected presentations

- Jul 2019 ETH Zurich Food Biotechnology Seminar (talk) Zurich, Switzerland
- Feb 2019 Univeristy of Minnesota Seminar (talk) St. Paul, MN, USA
- Aug 2018 Penn State Microbiome Center Seminar (talk) State College, PA, USA
- Aug 2018 ISME Conference (talk) Leipzig, Germany
- Jul 2018 HFSP Fellows Meeting 2018 (poster) Toronto, Canada
- Jun 2018 International Sourdough Symposium (poster) Cork, Ireland
- Jun 2018 NYU Genomics Symposium (talk) New York, NY, USA
- Jan 2018 MIT Ecology Meeting (talk) Cambridge, MA, USA
- Nov 2017 Workshop, Symbiosis in the microbial world: from ecology to genome evolution)(talk) West Sussex, UK
- Jul 2017 Gordon Research Conference: Microbial Population Biology (poster) Andover, NH, USA
- Jul 2017 Gordon Research Seminar: Microbial Population Biology (talk) Andover, NH, USA
- Mar 2017 Winter q-Bio Conference (talk) Kauai, HI, USA
- Oct 2016 Weizman Genome Evolution Conference (talk) Rehovot, Israel
- Aug 2016 ISME Conference (poster) Montreal, Canada
- Aug 2015 ESEB Conference (talk) Lausanne, Switzerland
- Jul 2015 SMBE Conference (poster) Vienna, Austria
- May 2015 HIV Dynamics and Evolution Conference (talk) Budapest, Hungary
- Feb 2014 New Zealand Phylodynamics Meeting (talk) Waiheke, New Zealand
- Nov 2013 Epidemics Conference (poster) Amsterdam, Netherlands
- May 2013 MCBE Conference (poster) Montpellier, France
- May 2013 HIV Dynamics and Evolution Conference (poster) Utrecht, Netherlands
- Jul 2012 Gordon Research Conference: Drug Resistance Evolution (poster) Easton, MA, USA
- Jan 2012 EE2 Workshop: Facing the challenge of infectious diseases (poster) Aosta, Italy
- Jul 2011 Gordon Research Conference: Microbial Population Biology (poster) Andover, NH, USA
- Nov 2011 EAWAG Aquatic Ecology and Macroevolution Seminar (talk) Kastanienbaum, Switzerland

Other Acadmic Activities

Grant Referee

Swiss National Science Foundation

Journal Referee

eLIFE; PNAS; Ecology Letters; Environmental Microbiology; Communications Biology; Proceedings of the Royal Society B; The American Naturalist; PLoS Computational Biology; Molecular Biology and Evolution; Journal of the Royal Soceity Interface; mSystems; Bioinformatics; Epidemics; Journal of Theoretical Biology; Theoretical Population Biology; Journal of Acq. Immune Deficiency Syndromes; Scientific Reports; PLoS ONE; Applied Mathematics and Computation; International Health

Seminar Organizing Committee

Parson's Microbial Systems Seminar, Massachusetts Institute of Technology Ecology, Evolution, Environment, Behavior (E3B) Seminar, University of Zurich/ETH Zurich Zurich Interaction Seminar, University of Zurich/ETH Zurich

Other activities

Maintainer at Brewsci/bio. Bioinformatics formulae for the Linuxbrew and Homebrew package managers.. https://brewsci.github.io/homebrew-bio/

References

Prof. Otto Cordero, Massachusetts Institute of Technology, ottox@mit.edu

Prof. Sebastian Bonhoeffer, ETH Zurich, seb@env.ethz.ch

Prof. Christophe Fraser, University of Oxford, christophe.fraser@bdi.ox.ac.uk