Gregory Gloor, PhD

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I Education and Training

- 1988-1990 Postdoctoral Fellow. University of Wisconsin Madison Laboratory of Genetics. Supervisor: Dr. William Engels.
- 1988 Ph.D, University of Western Ontario, Department of Biochemistry. Supervisor: Dr. George Chaconas. Dissertation: *Characterization of the Integrative Precursor Protein-DNA Complex of Bacteriophage Mu*.

1983 HBSc, University of Western Ontario, Genetics

2 Employment

2019-present, Chair-Department of Biochemistry

2002-present, Professor of Biochemistry University of Western Ontario, Faculty of Medicine now the Schulich School of Medicine and Dentistry

1997-2002, Associate Professor of Biochemistry University of Western Ontario, Faculty of Medicine

1993-1997, Assistant Professor of Biochemistry University of Western Ontario, Faculty of Medicine

1990-1992, Assistant Professor Medical Genetics Memorial University of Newfoundland, Faculty of Medicine

3 Faculty Development

- 2018, Four day applied course on Resolving Microbial Communities At Strain-Level using Metagenomic Assembly, University of Exeter, UK
- 2014, Five day theory and applied course on Compositional Data Analysis (UdG, Spain) Accredited by European Statistical Society
- 2007, Leadership Workshop

Offered by Continuing Education of Shulich School of Medicine and Dentistry for Medical School Accreditation Leaders

1995, Course on Teaching at the University Level Offered through the Faculty of Medicine Development Office.

1990, Faculty Orientation Day Memorial University of Newfoundland
This covered teaching tips and grant writing skills for new faculty at MUN

4 Awards, Honours, Fellowships

2014, Faculty Development Award: Attended week-long course on Compositional Data Analysis (UdG, Spain)

2011-2013, Faculty Scholar

2016-17, University Student's Council Teaching Honor Roll

2011-12, University Student's Council Teaching Honor Roll

2010-11, University Student's Council Teaching Honor Roll

2009-10, University Student's Council Teaching Honor Roll

2007, University Student's Council Teaching Honor Roll

2005, Schulich School of Medicine Teaching Award

2004, WL Magee Teaching Award, Biochemistry, UWO

1993 - 1998, Salary Award, Medical Research Council of Canada (MRC) Development Grant in Molecular Biology

1984 - 1988, K. M. Hunter Fellowship, National Cancer Institute of Canada.

1983, Graduate Entrance Scholarship, UWO.

5 Expertise and Research Interests

Composition and function of the human and other microbiomes. I use and develop tools to examine 16S rRNA gene composition, gene expression of mixed population samples, and metabolomic analysis of clinical samples. I teach a graduate course on the use of compositional data analysis techniques to examine transcriptomes, microbiomes and other types of complex data sets derived from high throughput sequencing.

Protein evolution. We use and develop tools to examine how protein structure and function is maintained in response to sequences changes. We have a special interest in identifying the role that variable positions play in protein evolution. I teach an undergraduate course in protein sequence alignment and proteins sequence-structure alignment.

Computational biology and that application of techniques for compositional data analysis to the above problems. Our primary contributions so far have been the ALDEx2 tool in Bioconductor for the analysis of high throughput experiments that generate counts per sequence tag: 16S rRNA gene sequencing, transcriptomics and selex-type experiments. I have further tools under development, and have contributed new visualization methods (effect-size plots) to the field.

6 Scholarly and Professional Activities

6.1 Scholarly Presentations and Invitations

2022, Invited speaker, CoDaWork 2022, Toulouse, France

2021, Presentation, CM Statistics (Zoom presentation)

2021, Invited speaker, National Eye Institute Workshop on the Microbiome, (Zoom presentation)

2020, Invited speaker, Genentech (Zoom presentation)

2020, Invited speaker and organiser, ISAPP Annual Meeting

2019, Invited workshop speaker, Computational and Methodological Statistics 2019, London, England

2019, Invited Participant/Speaker, Emerging Challenges in Microbiome Data Analysis, BIRS, Banff, Canada

2019, Invited Speaker, Barcelona Microbiome Debates, Barcelona, Spain

2019, Invited Speaker, Department of Bioinformatics, North Carolina State University, Charlotte

2019, Invited Speaker, Department of Biology, North Carolina State University,

- 2019, Invited Speaker, Program in Bioinformatics, Duke University,
- 2019, Invited Webinar Presenter, International Society for the Application of Probiotics and Probiotics (ISAPP)
- 2019, Invited Speaker, North American Microbiome Congress, Washington, USA
- 2018, Invited Webinar Presenter, International Life Sciences Institute (ILSI)
- 2018, Invited industrial speaker (KGK Biosciences), Supply Side West, Las Vegas, USA
- 2018, Invited speaker, Department of Physiology and Pharmacology, UWO
- 2018, Invited speaker, Department of Epidemiology, SUNY Buffalo, USA
- 2018, Invited speaker, EMBL-EBI, EBI Seminar Series, Hinxton, Cambridgeshire, UK
- 2018, Invited speaker, Gairdner Foundation Symposium on health through food and microbes, London, CA
- 2018, Invited speaker Gynecological Research Symposium, The Society of Gynecologic Oncology of Canada, Toronto, CA
- 2018, Workshop organizer and presenter, Compositional Data analysis methods, NGS'18, Barcelona, Catalonia
- 2017, Keynote, Microbial Ecology 2017, Toronto, Ontario
- 2017, Workshop, Compositional Data analysis methods, Microbial Ecology 2017, Toronto, CA
- 2017, Invited speaker, EMBL-EBI Industrial Program Workshop The human microbiome: challenges and opportunities for novel therapeutics, Hinxton, England
- 2017, Invited speaker Canadian Society of Microbiology, Waterloo, Ontario
- 2017, Canadian Statistical Sciences Institute Microbiome Planning Meeting speaker and discussion leader, Winnipeg, Manitoba
- 2017, Contributed Oral Presentation (2), Great Lakes Bioinformatics, Chicago, Illinois
- 2017, Invited speaker in the Microbiology & Immunology Department, Western University, London, CA
- 2017, Invited speaker in the Health Sciences Department, Carleton University, Ottawa, CA
- 2016, Invited speaker in the Biostatistics and Epidemiology Department, Western University, London, CA
- 2016, Invited speaker at Exploring Human Host-Microbiome Interactions in Health and Disease 2016, Cambridge, UK
- 2016, Invited workshop organizer at Exploring Human Host-Microbiome Interactions in Health and Disease 2016, Cambridge, UK
- 2016, Invited speaker at Symposium on Synthetic Biology, Western University, London, CA
- 2016, Invited workshop presenter, The Human Microbiome and Epidemiology, 2016 Epidemiology Congress of the Americas, Miami, USA
- 2016, Invited presentation/workshop, Infection, Inflammation and Immunity course, The Arctic University of Norway, Tromso, NO

- 2016, Oral Presentation, Great Lakes Bioinformatics / Canadian Computational Biology Conference, Toronto, CA
- 2015, Invited visiting scholar, Tiyani Health Sciences Research Institute, Zenjiang, China
- 2015, Invited speaker at Exploring Human Host-Microbiome Interactions in Health and Disease 2015, Cambridge, UK
- 2015, Invited paper at CoDaWork 2015, Girona, Spain
- 2015, Applying compositional data framework to microbiome datasets, Canadian Society of Microbiology workshop 2015, Saskatoon, Canada
- 2015, Invited speaker, University of Guelph Bioinformatics group
- 2014, Invited seminar, Dept. of Biochemistry, University of Calgary
- 2014, Invited participant at NIH sponsored Microbiome Quality Control Initiative: only Canadian group invited, Rockville, MD, USA
- 2013, Invited speaker at Fondation Merieux Conference on Better Foods for Better Health, Annecy, France
- 2013, Invited speaker at the Institute of Genome Sciences seminar series, University of Maryland, Baltimore, USA
- 2013, Invited expert participant at African International Conference and Workshop on the Microbiome and Probiotics, Nairobi, Kenya
- 2011, Invited speaker at the RePOOPulating the gut: therapeutic microbial preparations to eradicate recurrent C.difficile infections in Canada, Toronto
- 2011, Invited expert participant at International Society for the Application of Probiotics and Prebiotics, Berkley, CA
- 2010, Invited platform speaker at the Ontario Illumina Users Group, Toronto Ontario
- 2008, Invited speaker at University of North Carolina-Charlotte Department of Bioinformatics Seminar Series
- 2003, Invited speaker at First Canadian Workshop on Statistical Genomics, The Fields Institute, Toronto
- 2000, Department of Genetics seminar series, Harvard Medical School, Boston, Mass.
- 1999, London and Regional Cancer Center seminar series, London Ontario
- 1997 Department of Molecular Biology and Genetics seminar series, University of Guelph
- 1994 Mogenson Research Forum, UWO Faculty of Medicine
- 1992 Department of Genetics, University of Alberta seminar series Alberta Heritage Foundation for Medical Research Sponsored Speaker
- 1992 Lunchtime Seminar Series, Faculty of Medicine, Memorial University of Newfoundland
- 1992 Department of Biochemistry seminar series, Queens University
- 1992 Department of Biochemistry seminar series, UWO
- 1991 Janeway Hospital Residents Training seminar series, Faculty of Medicine, MUN
- 1991 Molecular Biology Research Discussion Group seminar series, Faculty of Medicine, MUN

1991 Department of Biochemistry seminar series, MUN

1990 Faculty of Medicine seminar series, MUN

6.2 Media Interviews and Public Presentations

I: radio or TV interview

W: written interview

P: press release

S: social media

R: public presentation or outreach

W Western News, Oct 2019: The promise of CRISPR for microbiome modulation

I Western News, Mar 2019: Study betters health by expanding gut knowledge

I Western News, Jan 2018: Molecular weapon targets bad bacteria

I BBC Radio, Oct 2017: The Science Hour http://www.bbc.co.uk/programmes/w3csv1fg

I BBC Radio, Oct 2017: Health Check http://www.bbc.co.uk/programmes/w3csty7v

I CTV Health, Oct 2017: http://www.ctvnews.ca/health/healthy-gut-linked-to-healthy-aging-in-new-research-1.3629433

W Medical News Today, Oct 2017 https://www.medicalnewstoday.com/articles/319756.php

W Medicalresearch.com, Oct 2017 https://medicalresearch.com/author-interviews/gut-microbiome-of-health-very-old-similar-to-younger-adults/37533/

P Science Daily, Oct 2017 https://www.sciencedaily.com/releases/2017/10/171011123728.htm

P Medscape, Oct 2017 https://www.medscape.com/viewarticle/887559

P Good Times, Oct 2017 https://goodtimes.ca/trust-gut-might-key-healthy-aging/

P Psychology Today, Oct 2017 https://www.psychologytoday.com/blog/the-athletes-way/201710/study-links-gut-microbiome-ridiculously-healthy-aging

I CTV London TV, Oct 2017 CTV video http://london.ctvnews.ca/video?clipId=1229909

I HealthLine, Oct 2017 https://www.healthline.com/health-news/does-healthy-gut-equal-long-healthy-life

I CBC Radio London, Oct 2017 - Afternoon Drive with host Chris Delatorre

I AM1290-London/AM800 Windsor, Oct 2017: TKO SHOW with Kara Ro

P Reader's Digest, Oct 2017: https://www.rd.com/health/wellness/healthy-gut-microbiome-longevity/

W Oxygen Magazine: Date of publication to be determined

S Science Reddit (to Nov 7, 2017) (196 comments, 3064 upvotes, 94%upvoted): https://www.reddit.com/r/science/comments/764rbg/ridiculously_healthy_elderly_have_the_same_gut/

- S Reddit AMA (to Nov 7, 2017)(99 comments, 184 upvotes, 89% upvoted): https://www.reddit.com/r/science/comments/79myv8/science_ama_series_were_the_researchers_from/
- S SchulichMedDent Twitter (to Nov 7, 2017) (10 retweets, 13 likes): https://twitter.com/SchulichMedDent/status/918117822865264641
- **S** Lawson Twitter (to Nov 7, 2017) (6 retweets, 12 likes): https://twitter.com/lawsonresearch/status/918222731849748480
- **S** Western Facebook post (to Nov 7, 2017) (109 shares, 371 reactions this also had a reach of 50.4K, and 2.1K clicks): https://www.facebook.com/WesternUniversity/
- I CTV London, Dec 2012:

http://london.ctvnews.ca/video?clipId=1015906

P Globe and Mail Jan 2013:

http://www.theglobeandmail.com/life/health-and-fitness/health/fake-poop-created-to-treat-c-diff/article7190494/

I CTV Kitchener, London, Kingston Jan 2016: http://kitchener.ctvnews.ca/video?playlistId=1.1111065

- R 2000-2004: Tour Organizer for Regina Mundi College Gr12 students (∼30 per year)
- **R** Nov 2001: Presentation on Emerging Career Opportunities in Biotechnology to the post-graduate Corporate Communications and Public Relations course at Fanshawe College
- **R** March 2001: Presentation on Genetic Research opportunities to entire A.B. Lucas Secondary School Student body (1400 students)
- I CBC TV Nov 1996: *Undercurrents* Documentary program: interviewed and hosted by Wendy Mesley. Rebroadcast on CBC News world
- **R** Nov 1996: Presenteration on Gene Therapy at the joint meeting of National Science Teachers Association-Science Teachers of Ontario Meeting
- R June 1996: Presentation on Medical Research to Students at Tilsonburg High school
- I Feb 1996: Led press conference outlining partnership between the FFGCT and the MRC for the funds raised by Jesse's Journey
- R Nov 1995: Presentation on Molecular Genetic Research to Saunder's Secondary School students
- **R** Oct 1995: Presentation to South London Chapter of Phi-delta- epsilon on Duchenne Muscular Dystropy and the promise of gene therapy
- R Sept 1995: Presentation at Jesse Davidson Appreciation Night at the London City Press Club
- R Aug 1995: Presentation at the corporate fundraising launch for Jesse's Journey
- R Aug 1991: Gene Therapy, Approaches and Limitations Resident's Rounds, Janeway Children's Hospital, St. John's Newfoundland

6.3 Peer Reviewed Papers:

- [147] D. T. Ham, T. S. Browne, P. N. Banglorewala, T. L. Wilson, R. K. Michael, G. B. Gloor, and D. R. Edgell, "A generalizable cas9/sgrna prediction model using machine transfer learning with small high-quality datasets," *Nat Commun*, vol. 14, p. 5514, Sep 2023.
- [146] G. M. Pellegrino, T. S. Browne, K. Sharath, K. A. Bari, S. J. Vancuren, E. Allen-Vercoe, G. B. Gloor, and D. R. Edgell, "Metabolically-targeted dcas9 expression in bacteria," *Nucleic Acids Res*, vol. 51, pp. 982–996, Jan 2023.
- [145] G. B. Gloor, "amicompositional: Simple tests for compositional behaviour of high throughput data with common transformations," *Austrian Journal of Statistics*, vol. 52, no. 4, pp. 180–197, 2023.
- [144] D. J. Giguere, A. T. Bahcheli, S. S. Slattery, R. R. Patel, T. S. Browne, M. Flatley, B. J. Karas, D. R. Edgell, and G. B. Gloor, "Telomere-to-telomere genome assembly of phaeodactylum tricornutum," *PeerJ*, vol. 10, p. e13607, 2022.
- [143] S. S. Slattery, D. J. Giguere, E. E. Stuckless, A. Shrestha, L.-A. K. Briere, A. Galbraith, S. Reaume, X. Boyko, H. H. Say, T. S. Browne, M. I. Frederick, J. T. Lant, I. U. Heinemann, P. O'Donoghue, L. Dsouza, S. Martin, P. Howard, C. Jedeszko, K. Ali, G. Styba, M. Flatley, B. J. Karas, G. B. Gloor, and D. R. Edgell, "Phosphate-regulated expression of the sars-cov-2 receptor-binding domain in the diatom phaeodactylum tricornutum for pandemic diagnostics," *Sci Rep*, vol. 12, p. 7010, Apr 2022.
- [142] J. R. Wu, J. M. Macklaim, B. L. Genge, and G. B. Gloor, *Finding the Centre: Compositional Asymmetry in High-Throughput Sequencing Datasets*, pp. 329–346. Cham: Springer International Publishing, 2021.
- [141] L. A. Grieves, G. B. Gloor, M. A. Bernards, and E. A. MacDougall-Shackleton, "Preen gland microbiota covary with major histocompatibility complex genotype in a songbird," *R Soc Open Sci*, vol. 8, p. 210936, Oct 2021.
- [140] L. A. Grieves, G. B. Gloor, T. R. Kelly, M. A. Bernards, and E. A. MacDougall-Shackleton, "Preen gland microbiota of songbirds differ across populations but not sexes," *J Anim Ecol*, vol. 90, pp. 2202–2212, 09 2021.
- [139] D. Nouri-Nejad, B. L. O'Donnell, C. S. Patil, R. E. Sanchez-Pupo, D. Johnston, S. Sayedyahossein, K. Jurcic, R. Lau, L. Gyenis, D. W. Litchfield, M. F. Jackson, G. B. Gloor, and S. Penuela, "Pannexin 1 mutation found in melanoma tumour reduces phosphorylation, glycosylation, and trafficking of the channel-forming protein," *Mol Biol Cell*, p. mbcE19100585, 01 2021.
- [138] I. Erb, G. B. Gloor, and T. P. Quinn, "Editorial: Compositional data analysis and related methods applied to genomics—a first special issue from NAR Genomics and Bioinformatics," *NAR Genomics and Bioinformatics*, vol. 2, 12 2020. lqaa103.
- [137] R. R. Cochrane, S. L. Brumwell, A. Shrestha, D. J. Giguere, S. Hamadache, G. B. Gloor, D. R. Edgell, and B. J. Karas, "Cloning of thalassiosira pseudonana's mitochondrial genome in saccharomyces cerevisiae and escherichia coli," *Biology (Basel)*, vol. 9, p. 358, Oct 2020.
- [136] K. F. Al, J. D. Denstedt, B. A. Daisley, J. Bjazevic, B. K. Welk, S. E. Pautler, G. B. Gloor, G. Reid, H. Razvi, and J. P. Burton, "Ureteral stent microbiota is associated with patient comorbidities but not antibiotic exposure," *Cell Reports Medicine*, vol. 1, no. 6, p. 100094, 2020.

- [135] S. Yang, G. Reid, J. R. G. Challis, G. B. Gloor, E. Asztalos, D. Money, S. Seney, and A. D. Bocking, "Effect of oral probiotic lactobacillus rhamnosus gr-1 and lactobacillus reuteri rc-14 on the vaginal microbiota, cytokines and chemokines in pregnant women," *Nutrients*, vol. 12, Jan 2020.
- [134] D. J. Giguere, J. M. Macklaim, B. Y. Lieng, and G. B. Gloor, "omicplotr: visualizing omic datasets as compositions," *BMC Bioinformatics*, vol. 20, p. 580, Nov 2019.
- [133] L. Grieves, G. Gloor, M. Bernards, and E. MacDougall-Shackleton, "Songbirds show odour-based discrimination of similarity and diversity at the major histocompatibility complex," *Animal Behaviour*, vol. 158, pp. 131 138, 2019.
- [132] T. A. Hamilton, G. M. Pellegrino, J. A. Therrien, D. T. Ham, P. C. Bartlett, B. J. Karas, G. B. Gloor, and D. R. Edgell, "Efficient inter-species conjugative transfer of a crispr nuclease for targeted bacterial killing," *Nat Commun*, vol. 10, p. 4544, Oct 2019.
- [131] M. Laforet, T. A. McMurrough, M. Vu, C. M. Brown, K. Zhang, M. S. Junop, G. B. Gloor, and D. R. Edgell, "Modifying a covarying protein-dna interaction changes substrate preference of a site-specific endonuclease," *Nucleic Acids Res*, vol. 47, pp. 10830–10841, Nov 2019.
- [130] T. P. Quinn, I. Erb, G. Gloor, C. Notredame, M. F. Richardson, and T. M. Crowley, "A field guide for the compositional analysis of any-omics data," *Gigascience*, vol. 8, Sep 2019.
- [129] C. Elwood, A. Y. K. Albert, E. McClymont, E. Wagner, D. Mahal, K. Devakandan, B. L. Quiqley, Z. Pakzad, M. H. Yudin, J. E. Hill, D. Money, VOGUE Research Group, A. Bocking, T. Dumonceaux, G. Gloor, S. Hemmingsen, J. E. Hill, M. Links, D. Money, K. O'Doherty, G. Reid, J. van Schalkwyk, P. Tang, and M. H. Yudin, "Different and diverse anaerobic microbiota were seen in women living with hiv with unsuppressed hiv viral load and in women with recurrent bacterial vaginosis: a cohort study," *BJOG*, Sep 2019.
- [128] M. D. Berg, D. J. Giguere, J. S. Dron, J. T. Lant, J. Genereaux, C. Liao, J. Wang, J. F. Robinson, G. B. Gloor, R. A. Hegele, P. O'Donoghue, and C. J. Brandl, "Targeted sequencing reveals expanded genetic diversity of human transfer rnas," RNA Biol, vol. 16, pp. 1574–1585, Aug 2019.
- [127] Z. Awamleh, G. B. Gloor, and V. K. M. Han, "Placental micrornas in pregnancies with early onset intrauterine growth restriction and preeclampsia: potential impact on gene expression and pathophysiology," *BMC Med Genomics*, vol. 12, p. 91, 06 2019.
- [126] D. G. Wootton, M. J. Cox, G. B. Gloor, D. Litt, K. Hoschler, E. German, J. Court, O. Eneje, L. Keogan, L. Macfarlane, S. Wilks, P. J. Diggle, M. Woodhead, M. F. Moffatt, W. O. C. Cookson, and S. B. Gordon, "A haemophilus sp. dominates the microbiota of sputum from uk adults with non-severe community acquired pneumonia and chronic lung disease," *Sci Rep*, vol. 9, p. 2388, Feb 2019.
- [125] A. Almeida, A. L. Mitchell, M. Boland, S. C. Forster, G. B. Gloor, A. Tarkowska, T. D. Lawley, and R. D. Finn, "A new genomic blueprint of the human gut microbiota," *Nature*, vol. 568, p. 499, Feb 2019.
- [124] B. R. Joris and G. B. Gloor, "Unaccounted risk of cardiovascular disease: the role of the microbiome in lipid metabolism," *Curr Opin Lipidol*, Jan 2019.
- [123] M. D. Berg, J. Genereaux, Y. Zhu, S. Mian, G. B. Gloor, and C. J. Brandl, "Acceptor stem differences contribute to species-specific use of yeast and human tRNAser," *Genes (Basel)*, vol. 9, p. 612, Dec 2018.

- [122] T. A. McMurrough, C. M. Brown, K. Zhang, G. Hausner, M. S. Junop, G. B. Gloor, and D. R. Edgell, "Active site residue identity regulates cleavage preference of LAGLIDADG homing endonucleases," *Nucleic Acids Res*, vol. 46, pp. 11990–12007, Oct 2018.
- [121] M. I. Petrova, J. M. Macklaim, S. Wuyts, T. Verhoeven, J. Vanderleyden, G. B. Gloor, S. Lebeer, and G. Reid, "Comparative genomic and phenotypic analysis of the vaginal probiotic lactobacillus rhamnosus GR-1," *Front Microbiol*, vol. 9, p. 1278, 2018.
- [120] G. B. Gloor, R. G. Wong, E. Allen-Vercoe, V. Dinculescu, M. Pignanelli, C. Bogiatzi, G. Reid, and J. D. Spence, "Data on the gut and saliva microbiota from a cohort of atherosclerosis patients determined by 16s rrna gene sequencing," *Data Brief*, vol. 19, pp. 481–485, Aug 2018.
- [119] M. Pignanelli, C. Bogiatzi, G. Gloor, E. Allen-Vercoe, G. Reid, B. L. Urquhart, K. N. Ruetz, T. J. Velenosi, and J. D. Spence, "Moderate renal impairment and toxic metabolites produced by the intestinal microbiome: Dietary implications," *J Ren Nutr*, vol. 29, pp. 55–64, Aug 2018.
- [118] M. Pignanelli, C. Just, C. Bogiatzi, V. Dinculescu, G. B. Gloor, E. Allen-Vercoe, G. Reid, B. L. Urquhart, K. N. Ruetz, T. J. Velenosi, and J. D. Spence, "Mediterranean diet score: Associations with metabolic products of the intestinal microbiome, carotid plaque burden, and renal function," *Nutrients*, vol. 10, Jun 2018.
- [117] A. McMillan, S. Rulisa, G. B. Gloor, J. M. Macklaim, M. Sumarah, and G. Reid, "Pilot assessment of probiotics for pregnant women in rwanda," *PLoS One*, vol. 13, no. 6, p. e0195081, 2018.
- [116] J. M. Macklaim and G. B. Gloor, "From RNA-seq to biological inference: Using compositional data analysis in meta-transcriptomics," *Methods Mol Biol*, vol. 1849, pp. 193–213, 2018.
- [115] C. Bogiatzi, G. Gloor, E. Allen-Vercoe, G. Reid, R. G. Wong, B. L. Urquhart, V. Dinculescu, K. N. Ruetz, T. J. Velenosi, M. Pignanelli, and J. D. Spence, "Metabolic products of the intestinal microbiome and extremes of atherosclerosis," *Atherosclerosis*, vol. 273, pp. 91–97, Jun 2018.
- [114] K. F. Al, J. E. Bisanz, G. B. Gloor, G. Reid, and J. P. Burton, "Evaluation of sampling and storage procedures on preserving the community structure of stool microbiota: A simple at-home toilet-paper collection method," *J Microbiol Methods*, vol. 144, pp. 117–121, Jan 2018.
- [113] J. J. Egozcue, V. Pawlowsky-Glahn, and G. B. Gloor, "Linear association in compositional data analysis," *Austrian Journal of Statistics*, vol. 47, pp. 3–31, january 2018.
- [112] G. B. Gloor, J. M. Macklaim, V. Pawlowsky-Glahn, and J. J. Egozcue, "Microbiome datasets are compositional: And this is not optional," *Front Microbiol*, vol. 8, p. 2224, 2017.
- [111] R. Sinha, G. Abu-Ali, E. Vogtmann, A. A. Fodor, B. Ren, A. Amir, E. Schwager, J. Crabtree, S. Ma, Microbiome Quality Control Project Consortium, C. C. Abnet, R. Knight, O. White, and C. Huttenhower, "Assessment of variation in microbial community amplicon sequencing by the microbiome quality control (mbqc) project consortium," Nat Biotechnol, vol. 35, pp. 1077–1086, Nov 2017.
- [110] G. Bian, G. B. Gloor, A. Gong, C. Jia, W. Zhang, J. Hu, H. Zhang, Y. Zhang, Z. Zhou, J. Zhang, J. P. Burton, G. Reid, Y. Xiao, Q. Zeng, K. Yang, and J. Li, "The gut microbiota of healthy aged chinese is similar to that of the healthy young," *mSphere*, vol. 2, no. 5, pp. e00327–17, 2017.
- [109] S. L. Martz, M. Guzman-Rodriguez, S.-M. He, C. Noordhof, D. J. Hurlbut, G. B. Gloor, C. Carlucci, S. Weese, E. Allen-Vercoe, J. Sun, E. C. Claud, and E. O. Petrof, "A human gut ecosystem protects against c. difficile disease by targeting tcda," *Journal of Gastroenterology*, vol. 52, pp. 452–465, Apr 2017.

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- [107] G. Ettinger, J. P. Burton, G. B. Gloor, and G. Reid, "Lactobacillus rhamnosus GR-1 attenuates induction of hypertrophy in cardiomyocytes but not through secreted protein msp-1 (p75)," *PLoS One*, vol. 12, no. 1, p. e0168622, 2017.
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6.4 Non Peer Reviewed Manuscripts

Giguere DJ, Bahcheli AT, Joris BR, Paulssen JM, Gieg LM, Flatley MW, Gloor GB (2020) Complete and validated genomes from a metagenome biorXIV: 10.1101/2020.04.08.032540

Toby Kenney, Glen Satten, Shyamal Peddada, Gregory Gloor, Hong Gu. Emerging Statistical Challenges and Methods for Analysis of Human Microbiome Data, 2019. Technical Report for the Banff International Research Centre (2019)

Russell J Dickson and Gregory B Gloor. Xorro: Rapid paired-end read overlapper. arXiv preprint arXiv:1304.4620, 2013.

Russell J Dickson and Gregory B Gloor. The MIp toolset: an efficient algorithm for calculating mutual information in protein alignments. arXiv preprint arXiv:1304.4573, 2013.

6.5 Software releases

ALDEx2. ALDEx tool to examine compositional high-throughput sequence data with Welch's t-test and Wilcoxon rank test. https://github.com/ggloor/ALDEx2, and http://www.bioconductor.org/packages/release/bioc/html/ALDEx2.html last update October 2023

Languages and utilities: R, bash, Perl, awk, LATEX, Markdown, HTML, git, svn

6.6 Patent filings

CIS CONJUGATIVE PLASMID SYSTEM, 2018, with David Edgell, Bogumil Karas SUPERIOR CONJUGATIVE PLASMIDS DELIVERED BY BACTERIA TO EUKARYOTIC CELLS, 2023 with Bogumil Kara, David Edgell

6.7 Academic trivia

- H-index: 58, past 5 years 48 (Google Scholar)
- Clarivate/Web of Science Highly Cited Researcher 2022, 2023 (top 1% worldwide in my field)
- Exaly profile
- mean iCite ratio: 3.1, past 10 years 3.9 (icite.od.nih.gov: >80th percentile NIH funded authors, >90th percentile of all authors)
- Erdos number 3 (Gloor Sheng Yu Andrew Granville Paul Erdos)
- Bacon number 4 (Gloor Wendy Mesley (Undercurrents) multiple paths multiple paths Kevin Bacon)
- Academic lineages: T.H. Morgan (Genetics through Engels), G. Khorana (Biochemistry through Chaconas)

6.8 Research Funding History

NSERC Alliance Grant, 2022-2024: Rapid scaling of viral spike protein production for SARS-CoV-2 testing using *Phaeodactylum tricornutum*

PI: Edgell DR, Co-PI Gloor GB, Karas B. Suncor

total: \$800000

Mitacs Accelerate, 2020: Rapid scaling of viral spike protein production for SARS-CoV-2 testing using *Phaeodactylum tricornutum*

PI: Gloor GB, Co-PI Edgell DR, Partner organization: Suncor

total: \$40000

NSERC Alliance Grant 2020-2024: Development and validation of rapid point of care food safety assays for Salmonella and Campylobacter - Rapid detection facilitated by CRISPR

PI; Rieder, M, Co-applicants Gloor, GB, Edgell, DR and Karas, B. Industrial Partner is International Point of Care Ray, D and Styba, G.

total: \$730999 + 1:1 matching from IPOC

Weston Family Microbiome Initiative 2020-2022: A Jekyll and Hyde role for gut bacterial proteases in the regulation of pain

Co-PI: Lomax, AEG, Reed, DE, Co-applicants Gloor, GB, Vanner, SJ, Allen-Vercoe E, Reed, DE, Ross, AC, Sheth, P, Bercik, P, De Palma, G

Applied research into the microbial causes of pain in IBD patients. Gloor role is metagenomic and metatranscriptomic profiling of patient microbiome samples.

total: \$3213000

CIHR Project Grant 2019-2022: Metabolic and Metagenomic effects of repopulation of the intestinal microbiome in patients with severe unexplained atherosclerosis (extended due to covid-19)

PI: Spence, DJ, Co-applicants Azarpazhooh, Mahmoud Reza; Burton, Jeremy P; Gloor, Gregory B; Silverman, Michael; Urquhart, Bradley L

Clinical trial to determine the effect of stool transplant on progression of atherosclerosis in patients predisposed coronary artery disease. Gloor role is metagenomic and metatranscriptomic profiling of patient microbiome samples.

total: \$200000

CIHR Project Grant, 2019-2024: Modulation of pain in IBD by microbial proteases

PI: Lomax, AEG, Co-PI Gloor, GB, Allen-Vercoe E, Reed, DE, Ross, AC, Vanner, SJ

Basic research into the microbial causes of pain in IBD patients. Gloor role is metagenomic and metatranscriptomic profiling of patient microbiome samples.

total: \$395990

CIHR Project Grant, 2018-2023: ConCRISPR: Conjugative delivery of a hybrid CRISPR/Tev nuclease system for specific microbiota modulation in the mammalian gut

PI: Edgell, DE, Co-PI Gloor, GB Heinrichs, D and Karas B

Basic research into the technology platform to target and eradicate specific taxa or genetic components of the human gut microbiome.

total: \$891000

Weston Family Foundation Microbiome Initiative, 2017-2018: Microbiota modulation with a hybrid CRISPR nuclease

PI: Gloor, GB, Co-PI Edgell, DE and Karas B

Developing technology to target and eradicate specific taxa or genetic components of the human gut microbiome.

total: \$149000

NSERC Discovery, 2015-2020: Molecular covariation in protein families

PI: Gloor, GB

Goal is to examine how covarying positions affect the structure and function of protein families that can be used as gene editing reagents.

total: \$155000 - pays for student and supplies

NIH R21 Dec 2015-2017, NIH R33 2017-2018: Microbes that matter

PI: Elaine Petrof (Queen's U), Allen-Vercoe (Guelph), Gloor (UWO)

Developing a synthetic stool substitute for the treatment of recurrent *C. difficile* infection

direct to Gloor: \$70000 - partially pays for one student and sequencing costs

CIHR 2013-2016: Role of intestinal microbiota in non-alcoholic fatty liver disease pre and post bariatric surgery

PI: ALLARD, Johane P (U. Toronto): , Comelli Elena M; GLOOR, Gregory B; JACKSON, Timothy D; LOU, Wen-Yi W; OKRAINEC, Allan

Characterization of the stool microbiota in a cohort of patients undergoing treatment for non-alcoholic fatty liver disease

total: 522169, direct to Gloor: \$25000/yr - partially paid for one student and sequencing costs

CIHR 2014-1016: Intestinal microbiome and extremes of atherosclerosis

PI: SPENCE, J. David, ALLEN-VERCOE, Emma; GLOOR, Gregory B; REID, Gregor

Characterization of the stool microbiota in a cohort of patients screened for risk of atherosclerosis

total: 211600, direct to Gloor: \$25000 - partially paid for one student and sequencing costs

CIHR 2013-1016: Non-Alcoholic Fatty Liver Disease: Role of Intestinal Microbiota and n-3 Polyunsaturated Fatty Acid Supplementationtle

PI: ALLARD, Johane P, COMELLI, Elena M; GLOOR, Gregory B; LOU, Wen-Yi W

Characterization of the stool microbiota in a cohort of patients treated for non-alcoholic fatty liver disease with fish oils

total 211600, direct to Gloor: \$17000 - partially paid for one student

CIHR Team grant, 2010-2015: The Vaginal Microbiome Project Team

PI: MONEY, Deborah M, BOCKING, Alan D; HEMMINGSEN, Sean M; HILL, Janet E; REID, Gregor (Co-PIs), co-investigators: DUMONCEAUX, Timothy J; GLOOR, Gregory B; LINKS, Matthew G; O'DOHERTY, Kieran C; TANG, Patrick K; VAN SCHALKWYK, Julianne E; YUDIN, Mark H

Collection and analysis of large vaginal microbiota cohorts to identify determinants of health and disease in the Canadian population

total:1745341, direct to Gloor: 15000/year - partially paid for one student. My role was tool development

Southeastern Ontario Academic Medical Organization 2014-2015: Elucidating the factors that determine success in fecal transplant therapy for *C. difficile* infection

PI: PETROF, E, ROPELSKI, Mark, ALLEN-VERCOE, Emma, GLOOR, Gregory

Identifying mechanisms of microbial ecosystem inhibition of C. difficile

total:92000, direct to Gloor: %14000 - paid for sequencing costs

CIHR Network grant 2012-2015 Maternal-Infant Microbiome and Immunity (MIMI) Network

PI: KOLLMANN, Tobias R, GLOOR, Gregory B; REID, Gregor

Team grant to further training and planning of microbiome effects on proper health and development in Africa.

total:600000, direct to Gloor: 200000 - paid for PDF, student and conference costs

NSERC ENGAGE, 2014-2015: Function of maize endophytic microbiome

PI: GLOOR, GREGORY with A&L Biologicals

Total: \$25000 - sequencing and sample processing costs

Determine the functional differences between high and normal yield sites by examining the endophytic corn sap meta-transcriptome.

Ontario Centre of Excellence, 2014-2015: Meta-transcriptome of high-yield corn endophytic microbiome

PI: GLOOR, GREGORY, with A&L Biologicals

Determine the functional differences between high and normal yield sites by examining the endophytic corn sap meta-transcriptome.

Total: \$25000 - sequencing and sample processing costs

CIHR Team grant in Bariatric Care, 2014-2019: Exploiting the therapeutic effects of the fecal microbiome in bariatric care

PI: ALLARD, Johane P, GAISANO, Herbert Y, BANKS, Kate; COMELLI, Elena M; GLOOR, Gregory B; HOTA, Susy S; JACKSON, Timothy D; LOU, Wen-Yi W; OKRAINEC, Allan; PHILPOTT, Dana J; POUTANEN, Susan M

Fecal microbiome transplant as a treatment for obesity

direct to Gloor: \$0 - I have chosen to withdraw from this research team

Agriculture and Agrifoods Canada, Agricultural Innovation Program, 2015-2017: Developing molecular methods as diagnostic tools to identify biological factors contributing to crop productivity and soil health

PI: Dr. George Lazarovitz (A&L Biologicals), Gloor, Gregory

Monitoring and understanding corn endophytic communities to maximize crop yield

Total: \$600000, direct to Gloor: \$120000 - pays for one PDF and sequencing costs

Academic Development Fund UWO, 2012: Major request for Illumina MiSeq Instrument

PI: Gloor, Hegele, Edgell, Singh

ADF money to buy an Illumina MiSeq for the Robarts sequencing core facility

Total: \$132500 - Paid for purchase and installation.

Ontario Genomics Institute Summer Research, 2010: Metagenomic error rate analysis of rare biomes

PI: Gloor

Summer research project examining metagenomic sequencing error rates

Total: \$5000 - partially paid for one summer student

NSERC Discovery, 2010-2015: Molecular covariation in protein families

PI: Gloor

Total: \$255564 - Core Research Funding

Academic Development fund, UWO, 2010: Deep resequencing of single genes

PI: Gloor

Developing methodologies for resequencing on the Illumina MiSeq Sequencing Platform

direct to Gloor: \$7500 - partially pays for one student and sequencing costs

CIHR Operating, 2006-2010: MODULATORS OF DOUBLE-STRAND BREAK REPAIR IN DROSOPHILA

PI: Gloor

Core research funding for examining DSB repair in Drosophila somatic cells

Total: \$405312 - Core Lab Research funding

Academic Development Fund, UWO, 2008: IDENTIFYING AND VALIDATING COEVOLVING POSI-

TIONS

PI: Gloor, Wahl, Dunn

direct to Gloor: \$8500 - partially paid for one PDF

CANCER RESEARCH SOCIETY, 1997-1999: GENES INVOLVED IN SOMATIC-CELL DOUBLE-STRAND

REPAIR

PI: Gloor

Core research lab funding

Total: \$85145 - partially pays for one student and sequencing costs

CIHR Operating, 2005-2008: MODULATORS OF DOUBLE-STRAND BREAK REPAIR IN DROSOPHILA

PI: Gloor

Core research lab funding

Total: \$361580 - partially pays for one student and sequencing costs

CIHR Operating, 2000-2006: Double strand break repair in Drosophila somatic and germline cells

PI: Gloor

Core research lab funding

Total: \$148000 - Core Lab research funding

MRC of Canada, 1997-2000: TEMPLATE-DEPENDENT REPAIR OF DOUBLE STRAND CHROMOSOME BREAKS IN Drosophila

PI: Gloor

Total: \$105005 - Core Lab research funding

MRC of Canada, 1993-1998: Transposon induced gene targeting

PI: Gloor

direct to Gloor: \$156920 - Core Lab research funding

Cancer Research Society, 1992-1994: Gene targeting to arbitrary sites in the genome

PI: Gloor

Total: \$96000 - Core Research Lab Funding

Where, years: Title

PI:, co-apps

Total: \$70000 - Core Research Lab Funding

MRC of Canada, 1991-1994: P element regulation and double strand break repair in Drosophila

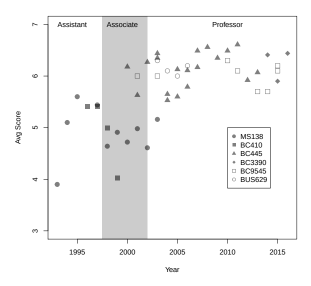
PI: Gloor

Total: \$197806 - Core Research Lab Funding

7 Teaching and Training

7.1 Course-based teaching

Figure 1: Teaching Ratings by Year and Course



I have been involved in teaching all or major portions of seven courses at Western, in addition to the occasional guest lecture. The six main courses are:

MS138: (1993-2003) First year medical school introduction to biochemistry. I taught 8-10 lectures that covered nucleotide metabolism, DNA replication and DNA repair as it related to cancer and HIV to first year medical students. I developed the material for delivery, following the broad theme given by the course co-ordinator at the time. The introductory didactic was removed from the undergraduate medical curriculum, along with most basic science, and replaced with small group teaching following a review of the undergraduate medical curriculum. I co-led a basic science review group for this process. I participated in small group teaching for several years afterward, but no evaluations were performed.

BC410: (1996-1999) I taught $\frac{1}{3}$ of this course, focussing on the basics of molecular biology pertaining to DNA structure, repair and recombination. The primary teaching tool was discussion of papers of current research interest with the objective of teaching the scientific discovery process. Included in this was one year of Biochemistry 453, the same course, different number.

BC445: (2000-2014) I developed and taught this course from scratch, often 2-3x per year. The course broadly covered 'bioinformatics' and had a focus on sequence and structure alignment, and at the time of development was rather novel being the first of its kind outside of a computer science department to cover the theory and practice of sequence alignment. The philosophy of the of this course was to teach how sequence alignment algorithms work in an intuitive way. The students would learn how the algorithm worked by doing it by hand, then we would move on to applying computers to make it faster. In addition, students learned the basics of Perl or Awk scripting and how to use a UN*X interface on a computer. Much of this material was rolled into BC3390.

BC3390: (2015 - now) A continuation, and simplification of the material in BC445 mandated when the undergraduate Biochemistry department curriculum was revamped.

BUS629: (2003-2006) I developed and taught this course in the Ivey MBA Biotechnology stream. I was responsible for all the material and for arranging guest experts. The thrust of the course was to teach non-science students in the MBA program how to quickly evaluate the business case for a biotechnology company based on the underlying science. This course was case-based and we discussed one case per week. I wish that Theranos was active at the time — it would have been a great example of foolish money chasing bad science. I received a citation from the Ivey Dean for teaching excellence.

BC9545: (2000 - now) This course initially was a more advance presentation of BC445 for graduate students. Since taking the course on compositional data analysis in 2014, I have redesigned it to be a course on the theory and practice of analyzing high throughput sequencing datasets — transcriptomics and metagenomics. I teach the theory and practice of probabilistic compositional data analysis for such datasets alongside the more traditional methods. My focus is on getting the students to understand and recognize when and why traditional methods go off the rails, and how they can use probabilistic compositional data analysis to double check their results.

The seventh course was a graduate computer science course.

CS881: (1996 - 2001) "Topics in computing and biology" Graduate Course. Six hours of lectures on the basics of DNA and on manipulations of DNA that could be used for primitive computational operations.

7.2 Summary of HQP

Graduate Students: 15; Undergraduate Student: 30; Postdoctoral Fellow: 2;

PhD Thesis Defence Examiner: 37 MSc Thesis Defence Examiner: 44 Thesis qualifying examiner: 48.

7.3 Graduate Students Supervised

Year	Student	Role	Description
2018-IPR	Joris	MSc (BC)	Defining correlated features in microbiomes
2017-IPR	Giguere	PhD (BC)	Integrative 'omic analysis of microbiomes
2017	Giguere	MSc (BC)	Visual exploration of multivariate multi-omic datasets
2017	Gajudhur	MSc (BC)	Visual exploration of multivariate multi-omic datasets
2013-2018	Macklaim	PDF	Microbial community composition of environmental datasets
2013-2016	Wong	MSc (BC)	Microbial community composition of patient cohorts
2012-2013	DiBella	MSc (MI, co)	Role of antisense transcription in bacterial gene regulation
2011-2017	McMurrough	PhD (BC, co)	Coevolution in LAGLIDAG endonucleases
2008-2013	Macklaim	PhD (BC)	Meta-omics of the vaginal microbiome
2008-2013	Dickson	PhD (BC)	Coevolution in protein families
2007-2012	Fernandes	PDF	Statistical tools for microbiome analysis
2002-2004	Gao	MSc (BCE, co)	Cell Surface Expression of Vitreoscilla Hemoglobin
2001-2008	Weedmark	PhD (BC)	DNA repair in Drosophila
1998-2005	Coveny	PhD (BC)	Polycomb genes and DSB
1997-1999	Ding	MSc (BCE, co)	Membrane display of an INP fusion protein
1994-1998	Dray	PhD (BC)	Targeting heterologous sequences in Drosophila melanogaster
1993-1998	Keeler	PhD (BC)	element induced double-strand break repair
1993-1995	Raynor	MSc (BC)	P element excision-induced double strand gap repair
1991-1995	Andrews	MSc (BC)	KP leucine zipper in the regulation of P transposition

8 Service to the University and Broader Community

8.1 University Committees

Year	Role	Description
2016	Member	Western NSERC Science Review Board
2012	Member	Law Graduate Program Review Committee
2011-2014	Member	Computer Science Promotion and Tenure Committee
2011-2013	Member	SUPR-G Committee
2011	Member	Faculty of Law and Ivey Business School Joint Promotion and Tenure Committee
2005-2006	Member	Ivey Business School Promotion and Tenure Committee
2005	Reviewer	External reviewer for Developmental Biology Graduate Program
2001-2004	Member	Promotion and Tenure - Biology Department
2001-2002	Member	UWO Bioinformatics Tier 2 CRC Search Committee
2000-2002	Member	Senate Subcommittee on Computing and Network Security
2000-2003	Chair	Senate Subcommittee on the WWW (SUWWW)
2000-2001	Member	Accessibility Subcommittee of SUWWW
1999-2000	Member	Biochemical Engineering Chair Selection Committee
1998-1999	Member	Senate Subcommittee on Computing and Network Services
1998-1999	Member	Senate Subcommittee on Computing and Information Technology
1997-1999	Member	Scientific Computing Research Advisory Subcommittee
1996-2000	Member	UWO Homepage Design and Implementation Subcommittee of SUWWW
1996-2000	Member	Senate Subcommittee on the WWW (SUWWW)
1995-2001	Chair	Molecular Biology Interfaculty Program Steering Committee
1995-2001	Chair	Molecular Biology Interfaculty Program Course Committee
1994	Co-Chair	Molecular Biology Interfaculty Program Open House
1993	Member	National Scholarship Selection Committee

8.2 Faculty Committees

Year	Role	Description
2010-2013	Member	Microbiology & Immunology Promotion and Tenure Committee
2007-2008	Member	Epidemiology and Biostatistics Chair Selection Committee
2007	Member	Ad hoc Faculty member disciplinary committee
2005-2007	Chair	Information Resources Institutional Self Study Task Force
2004-2006	Member	Promotion and Tenure - Pharmacology and Physiology Department
2001-2003	Member	Human Molecular Genetics Selection Committee
1999-2000	Chair	Information Resources Institutional Self Study Task Force
1997-2001	Member	UMEC Appeals Committee
1996-1998	Member	Medical Science Building Study Group
1996-1999	Member	Molecular Genetics Task Force
1996-1999	Co-Chair	Life Sciences group for the Medical School Curriculum renewal process
1996-2001	Member	ACMC Medical Informatics Group
1995-1996	Member	Computer and WWW Task Force
1994-2001	Member	Dean's advisory Committee on Genetics

8.3 Department Committees

Year	Role	Description
2019-present	Chair	Department of Biochemistry
2015-2017	Member (Seconded)	Biochemistry Promotion and Tenure Committee
2012-2015	Member	Research Committee
2010-2013	Member	Undergraduate Committee
2009-2012	Member	Biochemistry Promotion and Tenure Committee
2009-2012	Member	Appointments Committee
2004-2008	Chair	Biochemistry Graduate Studies
2002-2004	Member	Biochemistry Graduate Studies
2001-2004	Chair	Visiting Speakers Committee
1998-2004	CoChair	Department Outreach Committee
1997	Member	Biochemistry Promotion and Tenure Committee
1996	Chair	Nominating Committee
1995-2005	Member	Ad hoc Computing Advisory Committee
1995-1998	Member	Area, Safety, Equipment and Animal Care Committee
1993-1996	Member	Nominating Committee
1993-1996	Member	Graduate Studies Committee

8.4 Board memberships

2001-2002, Member, Board of Directors for Partners in Research

1995-2001 Member, Board of Directors, The Foundation for Gene and Cell Therapy (FFGCT).

I was the lead negotiator for the FFGCT in working out a partnership with the MRC to fund up to 9 post-doctoral fellows in the area of gene therapy. This partnership provided approximately \$900,000 in new money to the MRC.

1995, Member of the Home Team for Jesse's Journey (Internet and Science Advisor).

I posted and updated a World Wide Web map page so that people from around the world could follow John and Jesse's progress.

8.5 Grants and Awards Panels, Editorial membership, External Committees

2021-Present, Consulting Editor, Microbiome

2020, external reviewer, Fonds de la Recherche Scientifique - FNRS (Belgium)

2020, external reviewer, European Science Foundation

2019-2021, Member CodaWorks Student Grants Committee

2019-2021, Member, CodaWorks Meetings Committee

2019-now, Associate Editor, Nucleic Acids Bioinformatics and Genomics

2018-2021, Associate Editor, Microbiome

2018-2020, external reviewer Swiss National Research Agency

2016-present, Chair, CIHR Project Grant Scheme (GMX committee)

2016, Member, Agence Nationale de la Recherche, Preindustrial Biotechnology Demonstrator, Paris, France

2017-present, Member, CIHR College of Reviewers (First round invitee)

2017, Member, Canadian Crohns and Colitis Review panel

2016-2018, Section Editor, Microbiome

2016, Western Science and Engineering Review Board Member

2016-present CIHR Operating Grant Review Panel Chair

2015-2016, Associate Editor, Microbiome

2015, Ontario Genomics Institute: Large Scale Applied Research Program review panel

2014-present, CRC College of Reviewers

2012-2015, Editorial Board member Microbiome

2010-2014, Member CIHR Genetics panel

2008-present, IODE Doctoral Scholarship committee

2006-2010, NCIC Model Organisms Panel B2

1998, 1999, 2000, 2003, 2004, 2005, 2006, 2007, 2008 MRC/CIHR BMB/Genetics/Genomics invitee

1999, Chair OGS Biochemistry/Biophysics panel item 1997-1999, OGS Biochemistry/Biophysics panel

1997–2000, NCIC Virology and Molecular Biology Committee Panel F

1995-2001, Foundation for Gene and Cell Therapy (Jesse's Journey) Chair and Review Organizer 1995 - 2001