Carlos Oliver, PhD

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SUMMARY

My expertise and interests span all aspects surrounding the question of understanding biomolecular structure & function. As a computer scientist, I approach this problem by combining machine learning and graph algorithms. Since my PhD, I have completed projects in a wide range of topics from *drug discovery*, *structure prediction*, and *pattern mining*.

WORK EXPERIENCE

Postdoctoral Researcher

Basel, Switzerland

October 2021 - Present

full-time

KPERIENCE ETH Zürich

- **Department**: Machine Learning and Computational Biology Group, Department of Biosystems Science and Engineering
- Advisor: Karsten Borgwardt
- Researching graph machine learning methods and their applications to biological problems

Technical Advisor

May 2022 - Present

Simmunome Inc.

remote, part-time

Montreal, Canada

 Advisory board member for biotech startup focusing on disease modeling and drug discovery.

Consultant & Co-Founder

January 2019 - Present

Ozeki Inc.

remote, part-time

Montreal, Canada

• Consultancy service for startups; specializing in AI and blockchain implementations and advising.

Lead Scientist in Residence

March 2020 - August 2021

NextAI

part-time

Montreal, QC, Canada

• Supporting incubator startups in multiple industries with technical planning, troubleshooting, hiring and education.

Expert AI Advisor

May 2021 - October 2021

Creative Destruction Labs

remote, part-time

Toronto, ON, Canada

- Conducted interviews with startup founders to select incubator cohort.
- Assessed startups on technical soundness and potential.

 $Expert\ AI\ Advisor$

August 2020 - July 2021

Québec Ministry of Immigration

remote, part-time

Montreal, QC, Canada

• Evaluating Québec residence applications with respect to artificial intelligence proficiency.

Course Lecturer

September 2018-2019

School of Computer Science, McGill University Montreal, Canada part-time

• Taught and prepared learning material for undergraduate and graduate-level computer science courses (> 600 students). Topics: 'Introduction to Computer Science', and 'Fundamentals of Programming for Biology'

EDUCATION

PhD - Computer Science

McGill U. & Montreal Institute for Learning Algorithms (MILA) Montreal, Canada

2016-2021

- Advisor(s): Jerome Waldispuhl, William L. Hamilton
- Thesis: Origins, structure, and patterns of complex RNA structures.
- Research Areas: Computational Biology, Machine Learning on Graphs, Representation Learning, Network Science, Drug Discovery

M.Sc - Molecular Biology McGill U. Montreal, Canada

2014-2016

- Advisor(s): Jackie Vogel
- Thesis: Molecular Dynamics of the γ -Tubulin C-Terminal Tail
- Research Areas: Molecular Dynamics, Intrinsically Disordered Proteins, Systems Biology

B.Sc - Computer Science & Biology McGill U.
Montreal, Canada

2010-2014

• Minor: Italian Studies

RESEARCH HIGHLIGHTS

Fuzzy Network Motif Mining

- Fully automated tool for mining quasi-isomorphic and frequent network motifs in biological data.
- Integrates concepts from graph kernel methods, unsupervised graph representation learning, and exact graph algorithms (See [4] in Publications section)
- Source Code: https://github.com/cgoliver/vernal
- Motif Gallery: http://vernal.cs.mcgill.ca

Automated Drug Discovery for RNA

- First learning platform for target-based drug discovery on RNA molecules.
- Graph representation learning and kernel methods revealed insights for understanding key interactions in biomolecules (See [8] in Publications).
- Source Code: https://github.com/cgoliver/vernal
- Webserver: http://rnamigos.cs.mcgill.ca

PUBLICATIONS

- 1. Dexiong C., Fan B., Oliver, C. & Borgwardt, K. (2022). "Joint Multidimensional Scaling" (NeurIPS 2022 submission, preprint in preparation)
- 2. Oliver, C., Dexiong C., Mallet V., Philoppopoulos P., & Borgwardt, K. (2022). "Approximate Network Motif Mining Via Graph Learning" (NeurIPS 2022 submission, arXiv preprint arXiv:2206.01008.)
- 3. Mallet, V., **Oliver, C.**, Broadbent, J., Hamilton, W. L., & Waldispühl, J. (2022). "RNAglib: a python package for RNA 2.5 D graphs." *Bioinformatics*, 38(5), 1458-1459.
- 4. Oliver, C., Mallet, V., Philippopoulos, P., Hamilton, W. L., & Waldispühl, J. (2022). "Vernal: a tool for mining fuzzy network motifs in RNA." *Bioinformatics*, 38(4), 970-976.
- 5. Mallet, V., **Oliver, C. G.**, & Hamilton, W. L. (2021). "Edge-similarity-aware Graph Neural Networks." *arXiv preprint* arXiv:2109.09432.
- Boitreaud, J., Mallet, V., Oliver, C., & Waldispühl, J. (2020). OptiMol: optimization of binding affinities in chemical space for drug discovery. ACS Journal of Chemical Information and Modeling, 60(12), 5658-5666.
- 7. Philippopoulos, P., Ricottone, A., & Oliver, C. G. (2020). "Difficulty Scaling in Proof of Work for Decentralized Problem Solving." *Ledger*, 5.
- 8. Sarrazin-Gendron, R., Yao, H. T., Reinharz, V., Oliver, C. G., Ponty, Y., & Waldispühl, J. (2020, May). "Stochastic sampling of structural contexts improves the scalability and accuracy of RNA 3d module identification." In International Conference on Research in Computational Molecular Biology (pp. 186-201). Springer, Cham.
- Oliver, C., Mallet, V., Gendron, R. S., Reinharz, V., Hamilton, W. L., Moitessier, N., & Waldispühl, J. (2020). "Augmented base pairing networks encode RNAsmall molecule binding preferences." Nucleic Acids Research, 48(14), 7690-7699.
- 10. Oliver, C. G., Reinharz, V., & Waldispühl, J. (2019). "On the emergence of structural complexity in RNA replicators." RNA, 25(12), 1579-1591.
- 11. Vincent Mallet, Carlos G. Oliver, Nicolas Moitessier, Jérôme Waldispühl. "Leveraging binding-site structure for drug discovery with point-cloud methods". arXiv preprint arXiv:1905.12033 (2019)
- 12. Sarrazin-Gendron, R., Reinharz, V., **Oliver, C. G.**, Moitessier, N., & Waldispühl, J. (2019). "Automated, customizable and efficient identification of 3D base pair modules with BayesPairing." *Nucleic Acids Research*, 47(7), 3321-3332.
- 13. Harris, J., Shadrina, M., **Oliver, C.**, Vogel, J., & Mittermaier, A. (2018). "Concerted millisecond timescale dynamics in the intrinsically disordered carboxyl terminus of γ -tubulin induced by mutation of a conserved tyrosine residue." *Protein Science*, 27(2), 531-545.
- 14. **Oliver, C.G.** ., Ricottone A., Philippopoulos, P. "Proposal for a fully decentralized blockchain and proof-of-work algorithm for solving NP-complete problems." *ArXiV preprint* arXiv:1708.09419 (2017).
- 15. Bouchard, Johnathan, Carlos Oliver, and Paul M. Harrison. "The distribution and evolution of Arabidopsis thaliana cis natural antisense transcripts." *BMC Genomics* 16.1 (2015): 444.

AWARDS 2017 FRQNT Doctoral Fellowship 2017 John D. Thompson TechIdea Pitch Competition 3rd Place 2016 Graduate Excellence Award 2014 Cellular Dynamics Graduate Fellowship 2014 Dean's Multidisciplinary Undergraduate Researcher List

INVITED & REFEREED TALKS

- Deep Graph Learning Library Developer Group Virtual (Invited Talk)
- RECOMB 2021 Virtual (Oral Presentation for [4])
- RNA Society Meeting 2020 Online Vancouver, Canada (Selected Talk)
- Machine Learning in Drug Discovery 2018, University of Leuven, Belgium (Selected Talk)
- RiboClub Meeting 2018, Sherbrooke, Candada (Selected Talk)
- Computational Approaches to RNA Structure and Function, Benasque, Spain 2018 (Invited Speaker)

OPEN SOURCE CONTRIBU-TIONS

- GitHub: https://github.com/cgoliver/
- torch-pdb: co-creator and developer of largest deep learning-ready protein structure database; including newly published AlphaFold datasets.
- NetworkX (+ 4k users) graph comparison algorithms (WL-isomorphism, and Graph Edit Distance).
- Neural Subgraph Learning Library(+ 4k users) minor code enhancements.