Contact Information	School of Biosciences, University of Surrey Stag Hill Campus 15AX02, Surrey, GU2 7XH	b.lu@surrey.ac.uk	
Research Interests	Computational biology; machine learning; algorithms; pattern recognition; statistical inference; phylogenetics; population genetics; omics; platform and pipeline development; data analysis; method development and benchmarking; modelling; evolutionary dynamics; mutational pattern and process		
Employment	Surrey Future Fellow, University of Surrey	06/2023 -	
	Postdoctoral Fellow, University College London (UCL)	01/2019 - $05/2023$	
	<ul><li>Principle Investigator: Prof. Chris Barnes</li><li>Project: Dynamical modelling of somatic genomes</li><li>Developing methods to infer tumour sample phylogeny from copy number profiles</li><li>Modelling and analysis of chromosomal instability in experimental and real patient data</li></ul>		
	<b>Postdoctoral Fellow</b> , Genome Institute of Singapore (GIS)	09/2017 - $12/2018$	
	<ul><li>Principle Investigator: Prof. Weiwei Zhai</li><li>Project: Analyzing tumour heterogeneity and clonal evolution</li><li>Developing methods to simulate sequencing data of heterogeneous tumour samples</li><li>Analysis of intra-tumour heterogeneity in lung and liver cancer patient data</li></ul>		
Education	<b>Ph.D. Computer Science</b> , National University of Singapore (NUS)	08/2013 - 09/2017	
	Supervisor: Prof. Hon Wai Leong Project: Analyzing lateral gene transfer with computational methods - Predicting genomic islands in a microbial genome with machine learning - Developing algorithms related to phylogenetic networks		
	M.Eng. Computer Software and Theory, East China Normal University	09/2010 - 07/2013	
	<ul> <li>Supervisor: Prof. Zhenbing Zeng, Prof. Tieliu Shi</li> <li>Project: Developing high-throughput biological data analysis platform</li> <li>Platform for transcriptomic (RNA-Seq) data analysis</li> <li>Platform for proteomics (MS/MS) data analysis</li> </ul>		
	<b>B.Eng. Software Engineering</b> , East China Normal University (ECNU)	09/2005 - 07/2009	
Teaching	Teaching Assistant, Department of Computer Science, NUS	08/2015 - 11/2015	
	Module: Programming Methodology (undergraduate) Role: Laboratory guidance on programming exercises; After class support; Assignment marking		
	<b>Teaching Assistant</b> , Software Engineering Institute, ECNU Module: Discrete Mathematics (undergraduate) Role: Tutorial on explaining solutions to exercises; Assignment and exam ma	02/2011 - 06/2011 arking	
Student Mentoring	Christos Magkos (MSc Genetics of Human Disease), $01/2021 - 10/2021$ , UCL, Comparison of phylogenetic tree reconstruction methods using metastatic breast cancer data		
	Abbie Duan (Undergraduate), $10/2020 - 05/2021$ , UCL, Finding underlying patterns in copy number alteration data of gastric and colon cancers		
	Lisa Doetsch (Undergraduate summer project), $07/2020 - 10/2020$ , UCL, Finding underlying patterns in copy number alteration data of gastric and colon cancers		
	Simone De Angelis (MSc Genetics of Human Disease), $01/2020 - 09/2020$ , UCL, Evaluating tree building methods for tumour copy number profiles		
	Samuel Winnall (MSc Biological Physics), $10/2019 - 09/2020$ , UCL, Dynamical modelling of the evolution of structural variation in cancer genomes		
	Rachel Muir (MSc Genetics of Human Disease), $01/2019 - 09/2019$ , UCL, Evaluating phylogenetic inference of multi-region tumour data using copy number profiles		

Abner Herbert Lim (Undergraduate intern project), 01/2018 - 06/2018, GIS, Simulating next-generation sequencing data of tumour samples with PSiTE

Mengge Wang (Undergraduate intern project), 11/2016 - 05/2017, NUS, Analysing features related to the prediction of genomics islands

Naiqi Qiu (Undergraduate), 12/2012 - 05/2013, ECNU, The integration of PRIDE-tools and the MS-analysis-platform based on Galaxy

Publications Full list at google scholar

**Bingxin Lu**, Kit Curtius, Trevor A. Graham, Ziheng Yang, Chris P. Barnes (2023). CNETML: Maximum likelihood inference of phylogeny from copy number profiles of spatio-temporal samples. *Genome Biology*, 24, 144.

Kasper Karlsson, Moritz Przybilla, Eran Kotler, Aziz Khan, Hang Xu, Kremena Karagyozova, Alexandra Sockell, Wing H. Wong, Katherine Liu, Amanda Mah, Yuan-Hung Lo, **Bingxin Lu**, et al. (2023). Deterministic evolution and stringent selection during preneoplasia. *Nature*, 618, 383–393.

Weiwei Zhai<sup>\*</sup>, Hannah Lai<sup>\*</sup>, Neslihan Arife Kaya<sup>\*</sup>, Jianbin Chen<sup>\*</sup>, Hechuan Yang<sup>\*</sup>, **Bingxin Lu<sup>\*</sup>**, et al. (2022). Dynamic phenotypic heterogeneity and the evolution of multiple RNA subtypes in Hepatocellular Carcinoma: the PLANET study. *National Science Review*, 9(3), nwab192. (\* co-first authors)

Yannik Bollen, Ellen Stelloo, Petra van Leenen, Bas Ponsioen, Myrna van den Bos, **Bingxin Lu**, et al. (2021). Reconstructing single-cell karyotype alterations in colorectal cancer identifies punctuated and gradual diversification patterns. *Nature Genetics*, 53(8), 1187-1195.

Jianbin Chen, Hechuan Yang, Audrey Su Min Teo, Lidyana Bte Amer, Faranak Ghazi Sherbaf, Chu Quan Tan, Jacob Josiah Santiago Alvarez, **Bingxin Lu**, et al. (2020). Genomic landscape and ethnic specificity of lung adenocarcinoma in Asia. *Nature Genetics*, 52(2), 177-186.

Hechuan Yang, **Bingxin Lu**, Lan Huong Lai, Abner Herbert Lim, Jacob Josiah Santiago Alvarez, Weiwei Zhai (2019). PSiTE: A phylogeny guided simulator for tumor evolution. *Bioinformatics*, 35(17), 3148-3150.

Andreas D.M. Gunawan, **Bingxin Lu**, Louxin Zhang (2018). Fast methods for solving the Cluster Containment Problem for phylogenetic networks. *arXiv*:1801.04498.

**Bingxin Lu**, Hon Wai Leong (2018). GI-Cluster: detecting genomic islands via consensus clustering on multiple features. *Journal of Bioinformatics and Computational Biology*, 16(03), 1840010.

Jinwen Feng, Chen Ding, Naiqi Qiu, Xiaotian Ni, Dongdong Zhan, Wanlin Liu, Xia Xia, Peng Li, **Bingxin Lu**, et al. (2017). Firmiana: Towards a one-stop proteomic cloud platform for data processing and analysis. *Nature Biotechnology*, 35(5), 409-412.

**Bingxin Lu**, Louxin Zhang, Hon Wai Leong (2017). A program to compute the soft Robinson–Foulds distance between phylogenetic networks. *BMC Genomics*, 18(2), 111.

Andreas D.M. Gunawan<sup>\*</sup>, **Bingxin Lu**<sup>\*</sup>, Louxin Zhang (2016). A program for verification of phylogenetic network models. *Bioinformatics*, 32(17), i503-i510. (\* co-first authors)

**Bingxin Lu**, Hon Wai Leong (2016). Computational methods for predicting genomic islands in microbial genomes. *Computational and Structural Biotechnology Journal*, 14:200-206.

**Bingxin Lu**, Hon Wai Leong (2016). GI-SVM: A sensitive method for predicting genomic islands based on unannotated sequence of a single genome. *Journal of Bioinformatics and Computational Biology*, 14(01), 1640003.

Zhao Fang, **Bingxin Lu**, Mingyao Liu, Meixia Zhang, Zhenghui Yi, Chengping Wen, Tieliu Shi (2013). Evaluating the pharmacological mechanism of Chinese medicine Si-Wu-Tang through multi-level data integration. *PLoS ONE*, 8(11), e72334.

Geng Chen, Charles Wang, Leming Shi, Weida Tong, Xiongfei Qu, Jiwei Chen, Jianmin Yang, Caiping Shi, Long Chen, Peiying Zhou, **Bingxin Lu**, Tieliu Shi. Comprehensively identifying and characterizing the missing gene sequences in human reference genome with integrated analytic approaches. *Human Genetics*, 132(8):899-911, 2013.

**Bingxin Lu**, Zhenbing Zeng, Tieliu Shi (2013). Comparative study of de novo assembly and genomeguided assembly strategies for transcriptome reconstruction based on RNA-Seq. *Science China Life Sciences*, 56(2), 143-155.

Papers submitted	<b>Bingxin Lu</b> , Samuel Winnall, William Cross, Chris P. Barnes (2023). Cell-cycle dependent DNA repair and replication unifies patterns of chromosome instability. <i>bioRxiv</i> : 2024.01.03.574048.		
	William Cross <sup>*</sup> , Salpie Nowinski <sup>*</sup> , George Cresswell <sup>*</sup> , Maximilian Mossner <sup>*</sup> , Abhirup Banerjee <sup>*</sup> , <b>Bingxin Lu</b> <sup>*</sup> , et al. (2022). Evidence for stabilising selection causing grossly altered but broadly stable karyotypes in metastatic colorectal cancer. <i>bioRxiv</i> : 2020.03.26.007138. (* co-first authors)		
Posters	A computational model of double strand breaks and repair characterizes the generation of structural variants, <b>Bingxin Lu</b> and Chris P. Barnes, <i>ECCB</i> , Barcelona, Spain, 09/2022.		
	Tumour phylogeny reconstruction from copy number profiles of temporal multi-region samples, <b>Bingxin Lu</b> , Kit Curtius, Kane Smith, Ibrahim Al Bakir, Trevor A. Graham, Chris P. Barnes, <i>Evolution and Ecology of Cancer</i> , Hinxton, UK, 07/2019. (with lightning talk)		
	Fast Algorithms for the Cluster Containment Problem, Andreas D.M Gunawan, <b>Bingxin Lu</b> , and Louxin Zhang, <i>RECOMB</i> , Paris, France, 04/2018.		
	Computer program for verification of phylogenetic networks, Andreas D.M Gunawan, <b>Bingxin Lu</b> , Hon Wai Leong and Louxin Zhang, <i>RECOMB</i> , Hong Kong, China, 05/2017.		
	GI-Cluster: Detecting genomic islands in newly sequenced microbial genomes via consensus clustering on multiple features, <b>Bingxin Lu</b> and Hon Wai Leong, <i>RECOMB</i> , Hong Kong, China, 05/2017.		
Talks	A computational model of double strand breaks and repair characterizes the structural variants. <i>CSHA</i> , Suzhou, China	generation of somatic $12/2023$	
	GI-Cluster: detecting genomic islands via consensus clustering on multiple fea $GIW,$ Seoul, Korea	atures. $10/2017$	
	A program to compute the soft Robinson–Foulds distance between phylogenet $APBC$ , Shenzhen, China	tic networks. 01/2017	
	A program for verification of phylogenetic network models. $ECCB$ , The Hague, Netherlands	09/2016	
	GI-SVM: A sensitive method for predicting genomic islands based on unan single genome. <i>CSbio</i> , Bangkok, Thailand	notated sequence of a $11/2015$	
Awards &	Travel Fellowship, ECCB, The Hague, Netherlands	2016	
Funding	Student Travel Bursary, Winter School, The University of Queensland	2014	
	Research Scholarship, National University of Singapore	2013 - 2017	
	Excellent Graduate Student, East China Normal University	2009	
Service	Peer Review: APBC 2020; BMC Supplements (ICIBM 2021); CSBJ		
	Editing: PLOS One, guest editor	2023 -	
	Co-organizing a monthly joint journal club on cancer genomics with the researce Secrier and Dr. Simone Zaccaria, UCL	ch groups of Dr. Maria $11/2019 - 05/2022$	
	Volunteer for CGM (Chinese Genomics Meet-up) Europe	11/2022 - 07/2023	
Public	Volunteer for Soapbox Science, London	05/2023	
Engagement	Volunteer for Scratch training workshop and the Junior Category Hackathon @SG, Singapore $07/2015$		
Industry Experience	Software developer (C#), Jingchu Technologies Inc. (Beijing)	05/2010 - 08/2010	
	Software developer (C++), Luban Software Co. Ltd. (Shanghai)	04/2009 - 08/2009	
	Software developer (Java), Carestream Health, Inc. (Shanghai)	07/2008 - 10/2008	

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