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# A Bioinformatics Approach to Synthetic Lethal Interactions in Breast Cancer with Gene Expression Data

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a thesis submitted for the degree of Doctor of Philosophy at the University of Otago, Dunedin, New Zealand.

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#### Abstract

#### Background

Synthetic lethal genetic interactions are re-emerging in the post-genomics era due to their potential for use in precision medicine against cancers. Synthetic lethal drug design exploits the functional redundancy of genes disrupted in cancers (including tumour suppressors) to develop specific treatments against them. *CDH1*, which encodes E-cadherin, is a tumour supressor gene with loss of function in breast and stomach cancers. Experimental screens have identified candidate synthetic lethal interactions for drug target triage, which can be further supported with bioinformatics analysis. Furthermore, gene expression data enables investigation of synthetic lethal pathways and graph structure of synthetic lethal genes within them.

#### Approach

A computational methodology, the Synthetic Lethal Prediction Tool (SLIPT) has been developed to detect synthetic lethal interactions in gene expression data. This methodology was demonstrated on interactions with *CDH1* in breast and stomach cancer data from The Cancer Genome Atlas (TCGA) project. Synthetic lethal genes and pathways were further investigated with unsupervised clustering, gene set over-representation analysis, metagenes, and permutation resampling. In particular, analyses focused on comparing SLIPT gene candidates to an experimental siRNA screen Telford *et al.* (2015). Network analysis methods were applied to the most supported pathways to test for pathway structure among between synthetic lethal candidates. Simulation and modelling was used to assess the statistical performance of SLIPT, including simulated data with correlation structures derived from graph structures.

#### **Findings**

Many candidate synthetic lethal partners of CDH1 were detected in both TCGA breast cancer. These genes clustered into several distinct groups, with distinct biological functions and elevated expression in different clinical subtypes. While the number of genes detected by both approaches was not significant, these contained significantly enriched pathways. In particular,  $G_{\alpha i}$  signalling, cytoplasmic microfibres, and extracellular fibrin clotting were robustly supported by both approaches, which is consistent with the known cytoskeletal and cell signalling roles of E-cadherin and validation of GPCR pathways performed by Telford  $et\ al.\ (2015)$ . Many of these pathways were replicated in stomach cancer data. The pathways supported only by SLIPT included regulation of immune signalling and translational elongation which were not expected to be detected in an isogenic cell line model but are still candidates for further investigation.

Synthetic lethal candidates detected by SLIPT and siRNA were compared within graph structures of the candidate synthetic lethal pathways. These genes did not differ with respect to network metrics of importance or connectivity in the pathway. There was little support, across pathways, that SLIPT gene candidates were consistently upstream or downstream of siRNA gene candidates within pathways.

A model of synthetic lethality was used to simulate gene expression data with synthetic lethal partners of a gene. The SLIPT methodology had high statistical performance, detecting few synthetic lethal partners, which diminished with more synthetic lethal partners or lower sample size. The SLIPT methodology performed better than Pearson correlation or the  $\chi^2$ -test. In particular, it performed well with high specificity for datasets containing thousands of genes or genes positively correlated with the query gene (as expected to occur in expression data). SLIPT was robust across correlation structures, including those derived from complex pathway structures and often distinguished synthetic lethal genes from those positively or negatively correlated with them. Therefore SLIPT is appropriate to identify synthetic lethal genes within pathways and use candidate synthetic lethal genes (and their correlates) to identify synthetic lethal pathways.

#### Summary

Thus this thesis has developed, evaluated, and refined a bioinformatics approach to discovery of synthetic lethal genes solely from gene expression data. This approach has been demonstrated to detect biologically informative and clinically relevant candidate partners for *CDH1* in breast and stomach cancers. These investigations have also involved the development of network analysis and simulation procedures which may be more widely applicable.

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どうもありがとう由ちゃん。頑張った!もうすぐ行きます。!また来月!

# Contents

$\mathbf{G}$	lossa	$\mathbf{r}\mathbf{y}$			xix
$\mathbf{A}$	crony	ms		3	xxvi
1	Intr	oducti	ion and Literature Review		1
	1.1	Cance	er Research in the Post-Genomic Era		1
		1.1.1	Cancer is a Global Health Issue		2
			1.1.1.1 The Genetics and Molecular Biology of Cancers		3
		1.1.2	The Genomics Revolution in Cancer Research		3
			1.1.2.1 High-Throughput Technologies		4
			1.1.2.2 Bioinformatics and Genomic Data		5
		1.1.3	Genomics Projects		5
			1.1.3.1 The Cancer Genome Project		6
			1.1.3.2 The Cancer Genome Atlas Project		6
		1.1.4	Genomic Cancer Medicine		8
			1.1.4.1 Cancer Genes and Driver Mutations		8
			1.1.4.2 Precision Cancer Medicine		9
			1.1.4.3 Molecular Diagnostics and Pan-Cancer Medicine		9
			1.1.4.4 Targeted Therapeutics and Pharmacogenomics		10
		1.1.5	Systems and Network Biology		11
	1.2		etic Lethal Cancer Medicine		12
		1.2.1	Synthetic Lethal Genetic Interactions		13
		1.2.2	Synthetic Lethal Concepts in Genetics		13
		1.2.3	Synthetic Lethality in Model Systems		15
			1.2.3.1 Synthetic Lethal Pathways and Networks		15
			1.2.3.2 Evolution of Synthetic Lethality		16
		1.2.4	Synthetic Lethality in Cancer		17
		1.2.5	Clinical Impact of Synthetic Lethality in Cancer		18
		1.2.6	High-throughput Screening for Synthetic Lethality		20
			1.2.6.1 Synthetic Lethal Screens		21
		1.2.7	Computational Prediction of Synthetic Lethality		22
			1.2.7.1 Bioinformatics Approaches to Genetic Interactions .		22
			1.2.7.2 Comparative Genomics		23
			1.2.7.3 Analysis and Modelling of Protein Data		26
			1.2.7.4 Differential Gene Expression		28
			1.2.7.5 Data Mining and Machine Learning		29

			1.2.7.6 Mutual Exclusivity and Bimodality	. 32
			1.2.7.7 Rationale for Further Development	. 33
	1.3	E-cad	herin as a Synthetic Lethal Target	. 33
		1.3.1	The CDH1 gene and its Biological Functions	. 34
			1.3.1.1 Cytoskeleton	. 34
			1.3.1.2 Extracellular and Tumour Micro-environment	. 34
			1.3.1.3 Cell-Cell Adhesion and Signalling	. 35
		1.3.2	CDH1 as a Tumour (and Invasion) Suppressor	. 35
			1.3.2.1 Breast Cancers and Invasion	. 35
		1.3.3	Hereditary Diffuse Gastric (and Lobular Breast) Cancer	
		1.3.4	Cell Line Models of <i>CDH1</i> Null Mutations	. 37
	1.4	Summ	nary and Research Direction of Thesis	. 38
		1.4.1	Thesis Aims	
2	Met	thods	and Resources	41
	2.1	Bioinf	Formatics Resources for Genomics Research	. 41
		2.1.1	Public Data and Software Packages	. 41
			2.1.1.1 Cancer Genome Atlas Data	
			2.1.1.2 Reactome and Annotation Data	. 43
	2.2	Data 1	Handling	. 43
		2.2.1	Normalisation	. 43
		2.2.2	Sample Triage	. 44
		2.2.3	Metagenes and the Singular Value Decomposition	
		2.2.4	Candidate Triage and Integration with Screen Data	
	2.3	Techn	iques	. 47
		2.3.1	Statistical Procedures and Tests	. 47
		2.3.2	Gene Set Over-representation Analysis	. 48
		2.3.3	Clustering	. 48
		2.3.4	Heatmap	
		2.3.5	Modelling and Simulations	. 49
			2.3.5.1 Receiver Operating Characteristic Curves	. 50
		2.3.6	Resampling Analysis	. 50
	2.4	Pathw	vay Structure Methods	
		2.4.1	Network and Graph Analysis	. 51
		2.4.2	Sourcing Graph Structure Data	. 52
		2.4.3	Constructing Pathway Subgraphs	. 52
		2.4.4	Network Analysis Metrics	
	2.5	Imple	mentation	. 54
		2.5.1	Computational Resources and Linux Utilities	. 54
		2.5.2	R Language and Packages	. 55
		2.5.3	High Performance and Parallel Computing	
3	Met	thods	Developed During Thesis	60
	3.1		thetic Lethal Detection Methodology	
	3.2	Synth	etic Lethal Simulation and Modelling	. 62
		$3\ 2\ 1$	A Model of Synthetic Lethality in Expression Data	63

		3.2.2	Simulation Procedure
	3.3	Detect	ing Simulated Synthetic Lethal Partners
		3.3.1	Binomial Simulation of Synthetic Lethality
		3.3.2	Multivariate Normal Simulation of Synthetic Lethality 72
			3.3.2.1 Multivariate Normal Simulation with Correlated Genes 74
			3.3.2.2 Specificity with Query-Correlated Pathways 80
	3.4	Graph	Structure Methods
		3.4.1	Upstream and Downstream Gene Detection
			3.4.1.1 Permutation Analysis for Statistical Significance 85
			3.4.1.2 Hierarchy Based on Biological Context 85
		3.4.2	Simulating Gene Expression from Graph Structures
	3.5		mised Functions and Packages Developed
		3.5.1	Synthetic Lethal Interaction Prediction Tool 91
		3.5.2	Data Visualisation
		3.5.3	Extensions to the iGraph Package
			3.5.3.1 Sampling Simulated Data from Graph Structures 94
			3.5.3.2 Plotting Directed Graph Structures
			3.5.3.3 Computing Information Centrality 95
			3.5.3.4 Testing Pathway Structure with Permutation Testing . 95
			3.5.3.5 Metapackage to Install iGraph Functions 96
4	Syn	thetic	Lethal Analysis of Gene Expression Data 96
	4.1	Synthe	etic Lethal Genes in Breast Cancer
		4.1.1	Synthetic Lethal Pathways in Breast Cancer
		4.1.2	Expression Profiles of Synthetic Lethal Partners 100
			4.1.2.1 Subgroup Pathway Analysis
	4.2	Compa	aring Synthetic Lethal Gene Candidates
		4.2.1	Primary siRNA Screen Candidates
		4.2.2	Comparison with Correlation
		4.2.3	Comparison with Primary Screen Viability
		4.2.4	Comparison with Secondary siRNA Screen Validation 110
		4.2.5	Comparison to Primary Screen at Pathway Level
			4.2.5.1 Resampling Genes for Pathway Enrichment 113
		4.2.6	Integrating Synthetic Lethal Pathways and Screens
	4.3	Synthe	etic Lethal Pathway Metagenes
	4.4	Replic	ation in Stomach Cancer
	4.5	Discus	sion
		4.5.1	Strengths of the SLIPT Methodology
		4.5.2	Synthetic Lethal Pathways for E-cadherin
		4.5.3	Replication and Validation
			4.5.3.1 Integration with siRNA Screening
			4.5.3.2 Replication across Tissues
	4.6	Summ	

5	Syn	thetic	Lethal Pathway Structure	128
	5.1	Synth	etic Lethal Genes in Reactome Pathways	128
		5.1.1	The PI3K/AKT Pathway	
		5.1.2	The Extracellular Matrix	131
		5.1.3	G Protein Coupled Receptors	134
		5.1.4	Gene Regulation and Translation	134
		5.1.5	Gene Connectivity and Vertex Degree	135
		5.1.6	Gene Importance and Centrality	137
			5.1.6.1 Information Centrality	137
			5.1.6.2 PageRank Centrality	139
	5.2	Relati	onships between Synthetic Lethal Genes	141
		5.2.1	Hierarchical Pathway Structure	141
			5.2.1.1 Contextual Hierarchy of PI3K	141
			5.2.1.2 Testing Contextual Hierarchy of Synthetic Lethal Gene	s 141
		5.2.2	Upstream or Downstream Synthetic Lethality	144
			5.2.2.1 Measuring Structure of Candidates within PI3K	145
			5.2.2.2 Resampling for Synthetic Lethal Pathway Structure	147
	5.3	Discus	ssion	148
	5.4	Summ	nary	150
6	<b>Sim</b> 6.1		n and Modelling of Synthetic Lethal Pathways etic Lethal Detection Methods	
		0.1.1	6.1.1.1 Correlated Query Genes affects Specificity	157
		6.1.2	Alternative Synthetic Lethal Detection Strategies	159
		0.1.2	6.1.2.1 Correlation for Synthetic Lethal Detection	160
			6.1.2.2 Testing for Bimodality with BiSEp	161
	6.2	Simula	ations with Graph Structures	162
	0.2	6.2.1	Performance over Graph Structures	163
		0.2.2	6.2.1.1 Simple Graph Structures	
			6.2.1.2 Constructed Graph Structures	
		6.2.2	Performance with Inhibitions	168
		6.2.3	Synthetic Lethality across Graph Structures	174
		6.2.4	Performance within a Simulated Human Genome	177
	6.3	Simula	ations in More Complex Graph Structures	182
		6.3.1	Simulations over Pathway-based Graphs	183
		6.3.2	Pathway Structures in a Simulated Human Genome	185
	6.4	Discus	ssion	188
		6.4.1	Simulation Procedure	188
		6.4.2	Comparing Methods with Simulated Data	189
		6.4.3	Design and Performance of SLIPT	190
		6.4.4	Simulations from Graph Structures	192
	6.5		-	193

7	Discussion	195
	7.1 Synthetic Lethality and <i>CDH1</i> Biology	
	7.1.1 Established Functions of <i>CDH1</i>	196
	7.1.2 The Molecular Role of <i>CDH1</i> in Cancer	
	7.2 Significance	
	7.2.1 Synthetic Lethality in the Genomic Era	
	7.2.2 Clinical Interventions based on Synthetic Lethality	
	7.3 Future Directions	
	7.4 Conclusions	202
	Bibliography	<b>20</b> 4
$\mathbf{A}$	Sample Quality	228
	A.1 Sample Correlation	228
	A.2 Replicate Samples in TCGA Breast Cancer Data	231
В	Software Used for Thesis	235
$\mathbf{C}$	Mutation Analysis in Breast Cancer	<b>24</b> 4
	C.1 Synthetic Lethal Genes and Pathways	244
	C.2 Synthetic Lethal Expression Profiles	245
	C.3 Comparison to Primary Screen	248
	C.3.1 Resampling Analysis	250
	C.4 Compare SLIPT genes	252
$\mathbf{D}$	Metagene Analysis	<b>25</b> 4
	D.1 Pathway Signature Expression	
	D.2 Somatic Mutation	
	D.3 Synthetic Lethal Reactome Metagenes	
	D.4 Expression of Somatic Mutations	266
$\mathbf{E}$	Intrinsic Subtyping	269
$\mathbf{F}$	Stomach Expression Analysis	271
	F.1 Synthetic Lethal Genes and Pathways	271
	F.2 Comparison to Primary Screen	
	F.2.1 Resampling Analysis	277
	F.3 Metagene Analysis	279
$\mathbf{G}$	Synthetic Lethal Genes in Pathways	280
н	Pathway Connectivity for Mutation SLIPT	288
Ι	Information Centrality for Gene Essentiality	292
J	Pathway Structure for Mutation SLIPT	295

$\mathbf{K}$	Per	formance of SLIPT and $\chi^2$	298
	K.1	Correlated Query Genes affects Specificity	304
${f L}$	Sim	ulations on Graph Structures	310
		L.0.1 Simulations from Inhibiting Graph Structures	311
	L.1	Simulation across Graph Structures	314
	L.2	Simulations from Complex Graph Structures	318
		L.2.1 Simulations from Complex Inhibiting Graphs	321
	L.3	Simulations from Pathway Graph Structures	327

# List of Tables

1.1 1.2 1.3	Methods for predicting genetic interactions	23 24 25
2.1 2.2 2.3 2.4 2.5 2.6	Excluded samples by batch and clinical characteristics.  Computers used during thesis	44 54 55 56 56 58
4.1 4.2 4.3 4.4 4.5	Candidate synthetic lethal gene partners of <i>CDH1</i> from SLIPT Pathways for <i>CDH1</i> partners from SLIPT	98 99 104 107
4.6 4.7 4.8 4.9	genes against secondary short interfering RNA (siRNA) screen Pathways for <i>CDH1</i> partners from SLIPT and siRNA Pathways for <i>CDH1</i> partners from SLIPT	111 112 115 116 1120
5.1 5.2 5.3 5.4 5.5	ANOVA for synthetic lethality and vertex degree	136 138 140 144 147
B.1	Complete list of R packages used during this thesis	235
C.1 C.2 C.3 C.4 C.5 C.6	Candidate synthetic lethal gene partners of <i>CDH1</i> from mtSLIPT Pathways for <i>CDH1</i> partners from mtSLIPT Pathways for clusters of <i>CDH1</i> partners from mtSLIPT Pathways for <i>CDH1</i> partners from mtSLIPT and siRNA	244 245 247 249 250 251
D.1	Candidate synthetic lethal metagenes against CDH1 from mtSLIPT	265

E.1	Comparison of intrinsic subtypes	269
F.1	Synthetic lethal gene partners of <i>CDH1</i> from SLIPT in stomach cancer	271
F.2	Pathways for <i>CDH1</i> partners from SLIPT in stomach cancer	272
F.3	Pathways for clusters of <i>CDH1</i> partners in stomach SLIPT	274
F.4	Pathways for <i>CDH1</i> partners from SLIPT and siRNA	276
F.5	Pathways for <i>CDH1</i> partners from SLIPT in stomach cancer	277
F.6	Pathways for $CDH1$ partners from SLIPT in stomach and siRNA	278
F.7	Synthetic lethal metagenes against $\mathit{CDH1}$ in stomach cancer	279
H.1	ANOVA for synthetic lethality and vertex degree	291
	ANOVA for synthetic lethality and information centrality	
H.3	ANOVA for synthetic lethality and PageRank centrality	291
I.1	Information centrality for genes and molecules in the Reactome network	293
J.1	ANOVA for synthetic lethality and PI3K hierarchy	295
J.2	Resampling for pathway structure of synthetic lethal detection methods	297

# List of Figures

1.1 1.2	Synthetic genetic interactions	
1.2	Synthetic lethality in cancer	1
2.1	Read count density	5
2.2	Read count sample mean	5
3.1	Framework for synthetic lethal prediction	1
3.2	Synthetic lethal prediction adapted for mutation 69	2
3.3	A model of synthetic lethal gene expression	4
3.4	Modelling synthetic lethal gene expression	5
3.5	Synthetic lethality with multiple genes	6
3.6	Simulating gene function	8
3.7	Simulating synthetic lethal gene function	8
3.8	Simulating synthetic lethal gene expression	9
3.9	Performance of binomial simulations	1
3.10	Comparison of statistical performance	1
3.11	Performance of multivariate normal simulations	3
3.12	Simulating expression with correlated gene blocks	5
3.13	Simulating expression with correlated gene blocks	6
3.14	Synthetic lethal prediction across simulations	8
3.15	Performance with correlations	9
3.16	Comparison of statistical performance with correlation structure 80	0
	Performance with query correlations	1
3.18	Statistical evaluation of directional criteria	2
3.19	Performance of directional criteria	3
	Simulated graph structures	7
	Simulating expression from a graph structure	8
	Simulating expression from graph structure with inhibitions 89	9
3.23	Demonstration of violin plots with custom features	2
3.24	Demonstration of annotated heatmap	2
	Simulating graph structures	5
4.1	Synthetic lethal expression profiles of analysed samples	1
4.2	Comparison of SLIPT with siRNA	6
4.3	Comparison of SLIPT and siRNA genes with correlation 100	6
4.4	Comparison of SLIPT and siRNA genes with correlation	8
4.5	Comparison of SLIPT and siRNA genes with screen viability 109	9

4.6 4.7	Comparison of SLIPT genes with siRNA screen viability Resampled intersection of SLIPT and siRNA candidate genes	109 114
5.1	synthetic lethality in the PI3K cascade	130
5.2	synthetic lethality in Elastic Fibre Formation	132
5.3	Synthetic lethality in Fibrin Clot Formation	133
5.4	Synthetic lethality and vertex degree	136
5.5	Synthetic lethality and centrality	139
5.6	Synthetic lethality and PageRank	140
5.7	Hierarchical structure of PI3K	142
5.8	Hierarchy score in PI3K against synthetic lethality in PI3K	143
5.9	Structure of synthetic lethality in PI3K	144
5.10	Structure of synthetic lethality resampling in PI3K	146
6.1	Performance of $\chi^2$ and SLIPT across quantiles	155
6.2	Performance of $\chi^2$ and SLIPT across quantiles with more genes	156
6.3 6.4	Performance of $\chi^2$ and SLIPT across quantiles with query correlation . Performance of $\chi^2$ and SLIPT across quantiles with query correlation	157
	and more genes	158
6.5	Performance of negative correlation and SLIPT	161
6.6	Simple graph structures	164
6.7	Performance of simulations on a simple graph	165
6.8	Performance of simulations is similar in simple graphs	166
6.9	Performance of simulations on a pathway	167
6.10	Performance of simulations on a simple graph with inhibition	169
	Performance is higher on a simple inhibiting graph	171
	Performance of simulations on a constructed graph with inhibition	172
	Performance is affected by inhibition in graphs	173
	Detection of synthetic lethality within a graph structure	175
	Performance of simulations including a simple graph	179
	Performance on a simple graph improves with more genes	180
	Performance on an inhibiting graph improves with more genes	181
	Performance of simulations on the PI3K cascade	184
	Performance of simulations including the PI3K cascade	186
6.20	Performance on pathways improves with more genes	187
A.1	Correlation profiles of removed samples	229
A.2	Correlation analysis and sample removal	230
A.3	Replicate excluded samples	231
A.4	Replicate samples with all remaining	232
A.5	Replicate samples with some excluded	233
C.1	Synthetic lethal expression profiles of analysed samples	246
C.2	Comparison of mtSLIPT to siRNA	248
C.3	Compare mtSLIPT and siRNA genes with correlation	252
C.4	Compare mtSLIPT and siRNA genes with correlation	252
C.5	Compare mtSLIPT and siRNA genes with siRNA viability	253

D.1	Pathway metagene expression profiles	256
D.2	Expression profiles for constituent genes of PI3K	258
D.3	Expression profiles for estrogen receptor related genes	259
D.4	Pathway metagene expression profiles	260
D.5	Expression profiles for p53 related genes	261
D.6	Expression profiles for BRCA related genes	262
D.7	Somatic mutation against the PI3K metagene	263
D.8	Somatic mutation against PIK3CA metagene	266
D.9	Somatic mutation against PI3K protein	267
D.10	Somatic mutation against AKT protein	268
F.1	Synthetic lethal expression profiles of stomach samples	273
F.2	Comparison of SLIPT in stomach to siRNA	$\frac{275}{275}$
1.2	Comparison of SER 1 in Stomach to SIRVA	210
G.1	Synthetic lethality in the PI3K/AKT pathway	280
G.2	Synthetic lethality in the PI3K/AKT pathway in cancer	281
G.3	Synthetic lethality in the Extracellular Matrix	282
G.4	Synthetic lethality in the GPCRs	283
G.5	Synthetic lethality in the GPCR Downstream	284
G.6	Synthetic lethality in the Translation Elongation	285
G.7	Synthetic lethality in the Nonsense-mediated Decay	286
G.8	Synthetic lethality in the 3' UTR	287
TT 4		200
H.1	Synthetic lethality and vertex degree	288
H.2	Synthetic lethality and centrality	289
H.3	Synthetic lethality and PageRank	290
I.1	Information centrality distribution	294
J.1	Synthetic lethality and heirarchy score in PI3K	295
J.2	Heirarchy score in PI3K against synthetic lethality in PI3K	296
J.3	Structure of synthetic lethality in PI3K	296
J.4	Structure of synthetic lethality resampling	
J.4	Structure of Synthetic lethanty resampling	231
K.1	Performance of $\chi^2$ and SLIPT across quantiles	298
K.2	Performance of $\chi^2$ and SLIPT across quantiles	300
K.3	Performance of $\chi^2$ and SLIPT across quantiles with more genes	302
K.4	Performance of $\chi^2$ and SLIPT across quantiles with query correlation .	304
K.5	Performance of $\chi^2$ and SLIPT across quantiles with query correlation .	306
K.6	Performance of $\chi^2$ and SLIPT across quantiles with query correlation	
	and more genes	308
L.1	Performance of simulations on a simple graph	310
L.2	Performance of simulations on an inhibiting graph	311
L.3	Performance of simulations on a constructed graph with inhibition	312
L.4	Performance of simulations on a constructed graph with inhibition	313
L.5	Detection of synthetic lethality within a graph structure	314
L.6	Detection of synthetic lethality within an inhibiting graph	316
4.0		$ \mathbf{v}_{\perp}$ $\mathbf{v}_{\parallel}$

L.7	Detection of synthetic lethality within an inhibiting graph	317
L.8	Performance of simulations on a branching graph	318
L.9	Performance of simulations on a complex graph	319
L.10	Performance of simulations on a large graph	320
L.11	Performance of simulations on a branching graph with inhibition	321
L.12	Performance of simulations on a branching graph with inhibition	322
L.13	Performance of simulations on a complex graph with inhibition	323
L.14	Performance of simulations on a complex graph with inhibition	324
L.15	Performance of simulations on a large constructed graph with inhibition	325
L.16	Performance of simulations on a large constructed graph with inhibition	326
L.17	Performance of simulations on the $G_{\alpha i}$ signalling pathway	327
L.18	Performance of simulations including the $G_{\alpha i}$ signalling pathway	328

# Glossary

allele A gene variant with a specific sequence and

phenotype.

bioinformatics Statistical or computational approaches to

biological data or research tools.

bisulfite-Seq Epigenomic data from sequencing bisulfite

treated DNA.

CAGE-Seq Transcriptome data from cap analysis of gene

expression.

cancer A class of diseases, formally "malignant neo-

plasm", of abnormal cellular growth and

spread to other organs.

cancer gene A gene which is involved in the malignancy

of some cancers, encompassing oncogenes and tumour suppressors, which have molecular aberrations in cancer or variants which predis-

pose individuals to cancer.

centrality A network metric which identifies important

vertices.

chemoprevention The use of drugs to prevent early-stage can-

cers, generally applied to high-risk mutation

carriers.

chemotherapy The use of cytotoxic drugs to treat cancers, in

combinations, generally applied to advanced

stage cancers.

ChIP-Seq Epigenome data from chromatin immuno-

preciptation sequencing.

compound screen A high-throughput screen performed using a

library of chemical compounds.

computational biology Applying computational or mathematical

modelling to understanding biological systems

and relationships.

conditional essentiality A gene becoming essential to viability un-

der certain environmental conditions, including presence of compounds which inactivate

other genes.

copy number The number of copies of DNA, typically two

copies for diploid organisms but subject to

variation.

de novo A bioinformatics sequence assembly conduc-

ted entirely from raw genomics data without

a reference sequence.

diagnosis The identification of disease by clinical, cellu-

lar, and molecular characteristics.

driver mutation A mutation which promotes cancer growth.

E-cadherin Epithelial cadherin (calcium-dependent ad-

hesion), a cell-adhesion protein encoded by

CDH1.

edge or link A relationship connecting a pair of elements of

a graph structure or network, may be weighted

or directional.

epigenome An analysis of epigenetic modifications of all

genes in the genome.

epistasis (biological) The effects of a gene modifying or masking the

phenotype of another gene.

epistasis (statistical) A divergence of the observed double mutant

phenotype from that expected based on the respective phenotypes of single mutant (Fisher,

1919).

essential A gene which is required to be functional or

expressed for a cell or organism to be viable,

grow or develop.

exome A sequencing approach designed to generate

data enriched for coding genes within the gen-

ome.

familial A trait recurrently occurring in families, not

necessarily with a genetic cause.

functional redundancy Genes which perform a common function, also

known as genetic redundancy.

gene expression A measure of the relative expression of each

gene from the mRNA extracted from (pooled)

cells.

genetic robustness A system of biological pathways which (has

evolved to) continue to function as a whole under various conditions, including the inac-

tivation of various individual genes.

genome All of the DNA sequence in the genome.

genomic The use of data from all genes in the genome.

The use of genomic information to tailor medi-

cine treatment to the genetics of an individual.

germline mutation A mutation that occurred in germline cells and

is passed between generation.

graph or network A mathematical structure modelling or depict-

ing the relationships between elements.

hallmark of cancer An underlying characteristic of cancer as part

of a rational approach devised by (Hanahan

and Weinberg, 2000).

hereditary A trait or disease which has a genetic cause

and is inherited from family members.

high-throughput screen An experimental procedure to perform a large

scale series of chemical, genetic, or pharmaco-

logical tests.

hub A central or highly connected component of a

network.

in silico An investigation conducted using computa-

tions, typically simulations or analyses.

in vitro An investigation conducted using a controlled

experimental system to examine biomolecules.

in vivo An investigation conducted using in the con-

text of a biological cell or organism, including

pre-clincal models and clinical trials.

induced essentiality A gene becoming essential to viability under

certain conditions, including inactivation of a

synthetic lethal partner.

information centrality A network centrality metric which uses the im-

pact of removing a vertex or node on connec-

tions in the network.

intrinsic subtype Distinguishing cancer by molecular and ge-

netic features.

MCF10A cell line A non-tumorigenic epithelial cell line derived

from breast tissue.

metabolome All the metabolites and enzymes in the cell.

metagene A consistent signal of expression for a collec-

tion of genes such as a biological pathway, derived from singular value decomposition.

metagenome All of the genes and genomes in a community.

Mathematical Association and the genes and genomes in a community.

A secondary growth of a tumour or spread of

cancer to other organs.

methylation A measure of the epigenetic regulation of DNA

at CpG dinucleotide (CpG) sites.

microarray A high-throughput technique to measure pres-

ence or abundance of nucleic acid sequences

from binding to probes.

microRNA Short RNA molecules generally regarded to

regulate gene expression by binding to mRNA.

molecular profile A combination of genetic and biochemical

measures which identifies characteristic traits

of a tumour.

molecular subtype A classification of cancers based on an identi-

fication using molecular properties.

mutant A variant or dysfunctional phenotype arising

from a mutation in a gene.

mutation A change in DNA sequence that disrupts gene

function.

network biology The application mathematical and computa-

tional approaches to networks in understand-

ing biological relationships.

network medicine The use of network biology to understand, pre-

vent, or treat diseases.

non-oncogene addiction The dependence of a cancer cell on functioning

non-mutant genes.

'omics A combination of approaches to generating

biological data with high-throughput procedures such as genomics, proteomics or meta-

bolomics.

oncogene A gene that potentially causes cancer, typic-

ally by over-expression or mutant gene vari-

oncogene addiction The dependence of a cancer cell on a specific

oncogenic pathway.

PageRank centrality A network centrality metric which uses eigen-

vectors with a scaling factor (Brin and Page,

1998).

A focus on the molecular and genetic features pan cancer

across cancers in different tissues.

passenger mutation A mutation that occurs in cancers but does

not affect the growth of cancers.

pathway A series of biomolecules that produces a par-

ticular product or biological function.

lar targets or biological pathways.

pleiotropy When a gene has multiple biological functions. polypharmacology

The design of drugs to target multiple molecu-

precision medicine The application of prevention and treatment

measures to target diseases by molecular and

genetic features.

The estimation of disease progression and paprognosis

tient outcome.

proteome All the proteins expressed from the genome.

The non-mutant variant or precursor to a proto-oncogene

mutant oncogene.

recurrent mutation The repeated occurrence of mutations in a

particular gene across cancers.

RNAi screen A high-throughput screen performed using a

RNA interference (RNAi).

RNA-Seq The generation of transcriptome data from se-

quencing RNA.

Sanger sequencing A dideoxy chain termination method for DNA

sequencing (named after Fred Sanger).

scale-free A property of a network which has a power

law vertex degree distribution, that is several highly connected hub genes and many with

very few connections.

shortest path A path with the fewest possible edges which

connects two particular vertices.

small world A property of a network which is highly

connected and has a low characteristic path length, derived from the mean shortest path

length across all pairs of nodes.

somatic mutation A mutation that occurs in somatic cells, dur-

ing a patient's lifespan.

sporadic cancer Cancers which do occur in patients with a fam-

ily history or carry a high-risk genetic variant. When multiple drugs have more effect than

expected from the effect of each separately.

synthetic dosage lethal A synthetic genetic interaction (SGI) ana-

logous to synthetic lethality where where one gene is inactivated and the other over-

expressed.

synergy

synthetic lethal Genetic interactions where inactivation of

multiple genes is inviable (or deleterious) which are viable if inactivated separately.

synthetic lethal screen A high-throughput screen performed on iso-

genic cell lines to detect genes for which inhibition specifically deleterious to the null mutant

genotype.

synthetic rescue A synthetic genetic interaction when the com-

bined mutations restores the wild-type the

phenotype of one of the mutations.

synthetic sick Genetic interactions where inactivation of

multiple genes is deleterious which are viable

if inactivated separately.

synthetic suppression A synthetic genetic interaction when the com-

bined mutations (partially) suppresses the mutant phenotype of one of the mutations.

targeted therapy Cancer treatment that specifically acts against

a molecular target, in contrast to standard

chemotherapy.

transcriptome All of the genes expressed in the genome.

treatment Medical procedures for a disease to improve

patient outcomes.

tumour An abnormal lump of tissue or growth of cells,

may be cancerous.

tumour suppressor A gene potentially causes cancer, typically by

disruption of functions which protect the cell

from cancer.

vertex degree A network metric of connectivity of vertices

which uses the number of edges connected to

each vertex or node.

vertex or node An element of a graph structure or network.

wild-type A natural phenotype of a trait or the normally

functional allele which encodes it.

# Acronyms

1KGP 1000 genomes project.

ADP Adenosine Diphosphate.

AMP Adenosine Monophosphate.

AMPK AMP-activated Protein Kinase.

ANOVA Analysis of Variance. ATP Adenosine Triphosphate.

AUROC Area Under the Receiver Operating Charac-

teristic (curve).

Bash Bourne Again Shell.

BioPAX Biological Pathway Exchange.BiSEp Bimodal Subsetting Expression.BMP Bone Morphogenic Protein.

cAMP Cylic AMP.

CCLE Cancer Cell Line Encyclopaedia. cDNA Complementary DNA (from mRNA).

CGP Cancer Genome Project.

ChIP Chromatin Immunopreciptation.

ChIP-Seq Chromatin Immunopreciptation Sequencing.

CNV Copy Number Variation.

COSMIC Catalogue Of Somatic Mutations In Cancer.

CpG 5'-C-phosphate-G-3'.

CPM Counts Per Million mapped reads.

CPU Central Processing Unit.

CRAN comprehensive R archive network.

CXCR Chemokine Receptor.

DAISY Data Mining Synthetic Lethal Identification

Pipeline.

DDBJ DNA Data Bank of Japan.
DNA Deoxyribonucleic Acid.

EMBL European Molecular Biology Laboratory.

EMT Epithelial-Mesenchymal Transition.
ENA The European Nucleotide Archive.
ENCODE Encyclopaedia of DNA Elements.

ER Estrogen Receptor.

exprSL Synthetic Lethality (expression).

FANTOM Functional Annotation Of Mammalian gen-

ome.

FDR False Discovery Rate.

GEO Gene Expression Omnibus.

GO Gene Ontology.

GPCR G Crotein Coupled Receptor.

HDAC Histone Deacetylase.

HDGC Hereditary Diffuse Gastric Cancer.

HLRCC Hereditary Leiomyomatosis and Renal Cell

Carcinoma.

HPC High Performance Computing.

ICGC International Cancer Genome Consortium.

IHC Immunohistochemistry.

InDel Insertion or Deletion (in DNA sequence).

JAK Janus Kinase.

lncRNA Long Non-Coding RNA.

METABRIC Molecular Taxonomy of Breast Cancer Inter-

national Consortium.

microRNA Micro RNA. mRNA Messenger RNA.

MSI Microsatellite Instability. mtSL synthetic Lethality (mutation).

mtSLIPT Synthetic Lethal Interaction Prediction Tool

(against mutation).

NCBI National Center for Biotechnology Informa-

tion (in the USA).

NCI National Cancer Institute (in the USA). NeSI New Zealand eScience Infrastructure. NGS Next-Generation Sequencing.

NHGRI National Human Genome Research Institute

(in the USA).

NIG National Institute of Genetics (in Japan).

NIH National Institutes of Health (in the USA).

NMD Nonsense-Mediated Decay.

PAM50 Prediction Analysis of Microarray 50.

PARP Poly-ADP-Ribose Polymerase.

PCR Polymerase Chain Reaction.

PDE Phosphodiesterase.

PI3K Phosphoinositide 3-kinase.

PIP<sub>2</sub> Phosphatidylinositol-(4,5)-bisphosphate. PIP<sub>3</sub> Phosphatidylinositol-(3,4,5)-trisphosphate.

PPI Protein-Protein Interaction.

PR Progesterone Receptor.

qPCR Quantitative (real-time) Polymerase Chain

Reaction.

RFLP Restriction Fragment Length Polymorphism.

RGS G-protein Signalling.

RHO Ras Homolog Family.

RMA Robust Multiarray Averaging (normalisation.

RNA Ribonucleic Acid.

RNAi RNA Interference.

ROC Reciever Operating Characteristic (curve).

RPKM Reads Per Kilobase per Million mapped reads.

RPPA Reverse Phase Protein Arrays.

RRBS Reduced Representation Bisulfite Sequencing.

rRNA Ribonucleic acid.

RSEM RNA-Seq by Expectation Maximization (nor-

malisation.

SGA Synthetic Gene Array (technique).

SGI Synthetic Genetic Interaction.

shRNA Short Hairpin RNA.

siRNA Short Interfering RNA.

SL Synthetic Lethal.

SLIPT Synthetic Lethal Interaction Prediction Tool.

Slurm Simple Linux Utility for Resource Manage-

ment.

SNP Single Nucleotide Polymorphism.

SOCKS Socket Secure.

SR Synthetic Rescue (or viability).

SS Synthetic Suppression.

SSL Synthetic Sick.

TCGA The Cancer Genome Atlas (genomics project).

TGF $\alpha$  Transforming Growth Factor  $\alpha$ .

TMM Trimmed Mean of M values (normalisation.

tRNA Transfer RNA.

UCSC University of California, Santa Cruz. UTR Untranslated Region (of mRNA).

WNT Wingless-Related Integration Site.

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